

# The OPAQUE1/DISCORDIA2 myosin XI is required for phragmoplast guidance during asymmetric cell division in maize

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## Abstract

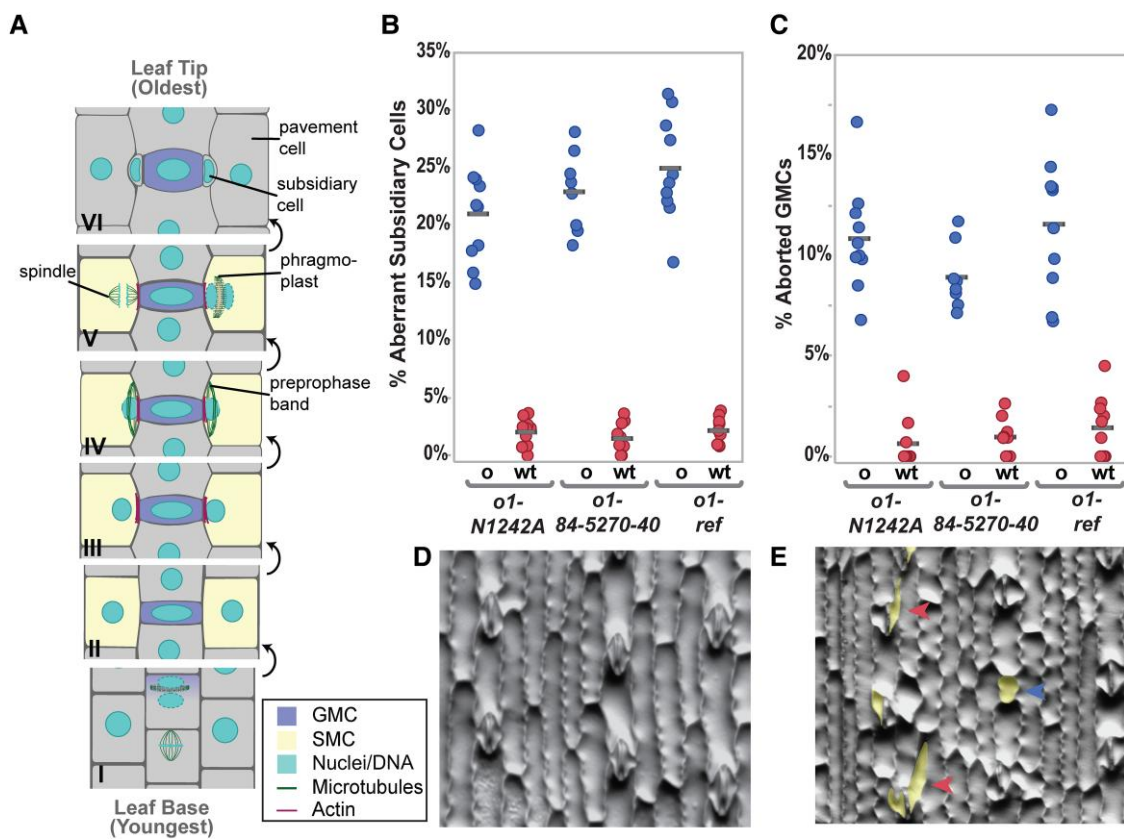
Formative asymmetric divisions produce cells with different fates and are critical for development. We show the maize (*Zea mays*) myosin XI protein, OPAQUE1 (O1), is necessary for asymmetric divisions during maize stomatal development. We analyzed stomatal precursor cells before and during asymmetric division to determine why *o1* mutants have abnormal division planes. Cell polarization and nuclear positioning occur normally in the *o1* mutant, and the future site of division is correctly specified. The defect in *o1* becomes apparent during late cytokinesis, when the phragmoplast forms the nascent cell plate. Initial phragmoplast guidance in *o1* is normal; however, as phragmoplast expansion continues *o1* phragmoplasts become misguided. To understand how O1 contributes to phragmoplast guidance, we identified O1-interacting proteins. Maize kinesins related to the *Arabidopsis thaliana* division site markers PHAGMOPLAST ORIENTING KINESINS (POKs), which are also required for correct phragmoplast guidance, physically interact with O1. We propose that different myosins are important at multiple steps of phragmoplast expansion, and the O1 actin motor and POK-like microtubule motors work together to ensure correct late-stage phragmoplast guidance.

## Introduction

Asymmetric divisions in plants and animals occur when a cell divides to give 2 daughters that differ from each other. Asymmetric divisions take place concomitantly with fate specification and are important in multicellular organisms to generate cell types of various fates, and to ensure the proper relative orientations of cells required for tissue patterning. Stomatal divisions from many plant species have served as a model for asymmetric cell division and have led to the

discovery of a suite of proteins required for asymmetric division and fate regulation.

Stomata are always composed of 2 guard cells and may also consist of a variable number of subsidiary cells (Gray et al. 2020). Stomata from maize (*Zea mays*) and other grasses are made of 4 cells: 2 inner guard cells laterally flanked by a pair of outer subsidiary cells (Facette and Smith 2012; Hepworth et al. 2018; McKown and Bergmann 2020; Nunes et al. 2020). Three types of divisions are required to create



**Figure 1.** OPAQUE1 is required for normal subsidiary cell formation in maize. **A**) Division sequence of stomatal development in maize and other grasses. **B**) Percentage of abnormal subsidiary cells and **C**) percentage of aborted GMCs in *o1* mutants (o) and their wild-type siblings (wt). In **(B and C)** seeds were scored based on the appearance of their endosperm (o, seeds that are phenotypically opaque; wt, translucent seeds). Each data point represents the percentage of abnormal cells from 1 plant; between 100 and 200 cells were counted per plant. Gray horizontal bars indicate means. ANOVA comparing each mutant to wild-type sibling returned  $P < 0.0001$  for all 3 alleles. **D**) Methacrylate impression of the leaf epidermis of a wild-type sibling showing normal stomatal complexes. **E**) *o1-ref* mutant showing abnormal subsidiary cells (red arrowheads) and aborted GMC (blue arrowhead). Abnormal cells in **(E)** are highlighted in yellow.

a stomatal complex in grasses: (i) asymmetric division of a protodermal cell to form a guard mother cell (GMC) and an interstomatal cell (Fig. 1A, panel I); (ii) asymmetric divisions of 2 subsidiary mother cells (SMCs) flanking the GMC (Fig. 1A, panels III–V); and (iii) the symmetric but oriented division of the GMC (not shown in Fig. 1A). Subsidiary cells are believed to contribute to the rapid stomatal movements observed in grass species (Franks and Farquhar 2007; Gray et al. 2020; Nunes et al. 2020).

Stomatal development requires many coordinated cellular processes before and during the formative divisions. Mutants in grass species that fail to correctly form stomata have been identified in maize, rice (*Oryza sativa*), and purple false brome (*Brachypodium distachyon*). Grass mutants with abnormal stomatal development generally fall into 2 classes: mutants defective in genes important for cell fate specification (Liu et al. 2009; Raissig et al. 2016, 2017; Wang et al. 2019; Wu et al. 2019), or genes encoding proteins important for physically executing the asymmetric division. Mutants that fail to correctly execute stomatal divisions are classified based on which universal and temporally distinct phase of

asymmetric division is defective: cell polarization, division plane establishment, division plane maintenance, or cytokinesis. The asymmetric division of the SMC, in particular, is useful as a model for dissecting the processes that occur during asymmetric division, as they have conspicuous polarization markers and the daughter cells are morphologically distinct.

Mutations in maize genes leading to defects in cell polarization include *brick1* (*brk1*), *brk2*, and *brk3* (Frank et al. 2003; Facette et al. 2015); *pangloss2* (*pan2*) and *pan1* (Cartwright et al. 2009; Zhang et al. 2012); and *rho gtpase of plants2* (*rop2*), *rop4*, and *rop9* (Humphries et al. 2011). These mutations lead to a nuclear polarization defect and aberrant division planes. In *B. distachyon*, BdPolar has the opposite localization of PAN1 (it is excluded from the GMC–SMC interface) and is required for correct SMC polarization (Zhang et al. 2022). Mutants in which SMCs polarize correctly, but fail in subsequent steps include the *discordia* (*dcd*) and *tangled1* (*tan1*) mutants (Smith et al. 1996; Gallagher and Smith 1999; Martinez et al. 2017).

The microtubular preprophase band (PPB) is an early division site marker that disappears prior to metaphase.

Identification of the microtubule-binding protein TANGLED1 (TAN1) answered a long-standing question of how the cortical division site was maintained throughout mitosis after the disappearance of the PPB (Smith et al. 1996; Walker et al. 2007). TAN1 and other division site markers are important for division plane maintenance and continuously mark the division site from prophase until telophase (Cleary and Smith 1998; Walker et al. 2007; Xu et al. 2008; Stöckle et al. 2016; Martinez et al. 2017). The TAN1-interacting partners PHRAGMOPLAST ORIENTING KINESIN1 (POK1) and POK2 are kinesin proteins that have been characterized in *Arabidopsis* (*Arabidopsis thaliana*) and mark the division site (Müller et al. 2006; Lipka et al. 2014; Herrmann et al. 2018). POK2 and TAN1 localize to the phragmoplast as well as the cortical division site (Herrmann et al. 2018; Buschmann and Müller 2019; Bellinger et al. 2021; Mills et al. 2022).

Cortical division site markers explain how the division site is maintained throughout mitosis. However, for division plane fidelity, the phragmoplast must be correctly guided to the correct division site. The phragmoplast is composed of membranes, microtubules, actin, and other accessory proteins, which starts as a disc in the center of the cell (Smertenko et al. 2018; Buschmann and Müller 2019). Vesicles targeted to the phragmoplast fuse to form the cell plate. The phragmoplast expands in circumference until it eventually meets the existing cell wall at the cortical division site marked by TAN1, POK, and other division site markers (Smertenko et al. 2018). Phragmoplast expansion (reviewed in Smertenko 2018; Smertenko et al. 2018; Buschmann and Müller 2019; Lee and Liu 2019)) requires microtubule turnover and many different mutants affect phragmoplast stability, morphology, or guidance.

Phragmoplast guidance occurs in stages. The initial rapid expansion of the disc-phragmoplast is likely guided by actin networks (Molchan et al. 2002; van Oostende-Triplet et al. 2017). Phragmoplast expansion slows when 1 edge meets the cell cortex and eventually, through expansion, the entire circumference of the phragmoplast reaches the cell cortex (van Oostende-Triplet et al. 2017). In this later stage, microtubules at the cell cortex are incorporated into the phragmoplast, where TAN1 (and probably other division site-localized proteins) are important for the incorporation of cortical telophase microtubules into the phragmoplast (Bellinger et al. 2021). After phragmoplast expansion is complete, the newly formed cell plate fuses with the existing cell wall (Smertenko et al. 2017; van Oostende-Triplet et al. 2017; Smertenko 2018; Lee and Liu 2019). Importantly, early phragmoplast guidance, late phragmoplast guidance, and phragmoplast integrity are distinct from one another. Despite recent progress using mutants and time-lapse imaging, the precise mechanism and protein–protein interactions that promote correct phragmoplast guidance remain unclear.

Using maize stomatal precursors as a model, we wanted to understand how plant cells correctly execute asymmetric divisions. Actin–myosin networks are prominent during plant cell division generally (Sadot and Blancaflor 2019), and

specifically during the asymmetric division of the maize SMC (Facette et al. 2015). Actin accumulates in the PPB and the spindle of plant cells, and accumulates on either side of the division zone (Palevitz 1987; Sano et al. 2005; Yasuda et al. 2005; Van Damme et al. 2007; Panteris 2008; Kojo et al. 2013). F-actin, myosin VIII, and myosin XI localize to the spindle and phragmoplast (Wu and Bezanilla 2014; Abu-Abied et al. 2018; Sun et al. 2018). Actin–myosin networks play additional roles specific to asymmetric SMC divisions. Indeed, in grasses and monocots with similar stomatal development, nuclear migration is driven by actin networks (Cho and Wick 1991; Kennard and Cleary 1997; Apostolakos et al. 2018). Actin is polarized at the GMC–SMC interface, and the SCAR/WAVE complex—which promotes actin nucleation—is required for the polarization of PAN proteins (Facette et al. 2015). These observations indicate that the actin motors (i.e. myosins) are likely important for SMC divisions, and could potentially play roles during polarization, division plane establishment, and/or cytokinesis. Therefore, we investigated the role of the previously identified OPAQUE ENDOSPERM1 (O1) protein in asymmetric divisions.

The *opaque* class of mutants was originally identified based on their seed phenotype of opaque endosperm (Neuffer et al. 1968; Gibbon and Larkins 2005). O1 encodes a myosin XI protein required for normal endoplasmic reticulum (ER) and protein body morphology in developing seeds, although the gene is expressed throughout the plant (Wang et al. 2012). Notably, O1 is very similar to *Arabidopsis* MYOXI-I, which is required for nuclear movement and shape (Tamura et al. 2013; Zhou et al. 2015; Muroyama et al. 2020). In *Arabidopsis* stomatal precursors, pre-mitotic nuclear migration is driven by microtubule (rather than actin) networks, but postmitotic migration of the nucleus is driven by actin networks and MYOXI-I (Muroyama et al. 2020).

We hypothesized that O1 might play a role in the asymmetric division of maize SMCs, perhaps during pre-mitotic polarization of the nucleus. Indeed, we determined that asymmetric divisions of both SMCs and GMCs are abnormal in *o1* mutants. However, division defects in *o1* are not a result of cell polarization defects, but rather late-stage phragmoplast guidance defects. To gain insight into how O1 promotes correct phragmoplast guidance, we identified proteins that physically interact with O1, leading to the identification of the maize orthologs of POK1 and POK2, in addition to actin-binding proteins and other myosins. Given their physical interaction, and the similarity of *pok* mutant phenotypes in *Arabidopsis* to the phenotypes we observed in maize *o1* mutants, we hypothesize these 2 cytoskeletal motors work together to promote phragmoplast guidance.

## Results

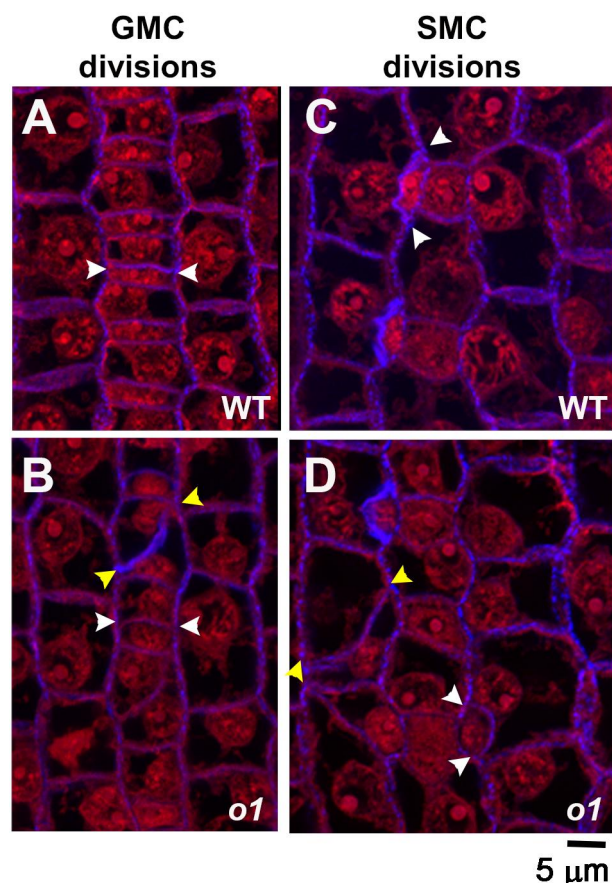
### *Opaque1* is required for stomatal divisions

To determine if the myosin OPAQUE1 was involved in stomatal divisions, we examined the morphology of subsidiary

cells in fully expanded juvenile leaves (leaf 4). We phenotyped segregating F2 plants for opaque seeds and calculated the corresponding frequency of abnormal subsidiary cells in wild-type (both *o1*+/+ and +/++) and *o1* sibling plants. We examined 3 different *o1* mutant alleles. Between 20% to 30% of subsidiary cells were abnormal in *o1*, while siblings with translucent seed (i.e. wild type) had less than 5% abnormal subsidiary cells (Fig. 1B). This phenotypic penetrance was consistent with that seen in other mutants such as *brk*, *pan*, and *dcd*, which also show ~25% abnormal subsidiary cells (Gallagher and Smith 1999; Cartwright et al. 2009; Zhang et al. 2012; Facette et al. 2015). All 3 alleles also displayed an increased frequency of aborted GMCs (Fig. 1C). We classified GMCs as aborted when a single cell with the same morphology as a GMC was present instead of a normal 4-celled stomatal complex (blue arrow, Fig. 1E).

Aberrant subsidiary cells and aborted GMCs in *o1* suggest a role for this myosin in both types of stomatal formative asymmetric divisions. To confirm the defect, we compared *o1* and wild-type siblings at the early stages of leaf development, when stomatal divisions are taking place. We stained cell walls and nuclei with propidium iodide (PI), and plasmodesmata and new cell plates with aniline blue. In *o1* mutant plants, recently formed GMCs showed abnormal division planes (Fig. 2, A and B). We propose that aborted GMCs are present in expanded leaves when the GMC progenitor divides abnormally, resulting in failed GMC fate specification. We also observed abnormal division planes in recently formed SMCs (Fig. 2, C and D). Importantly, we noticed no multinucleate cells or cell wall stubs, as we only observed abnormal division planes. This result suggests that this specific myosin is not required for cell plate integrity, but rather it is the division plane that is affected. These data indicate a role for O1 during the asymmetric division of stomatal precursors.

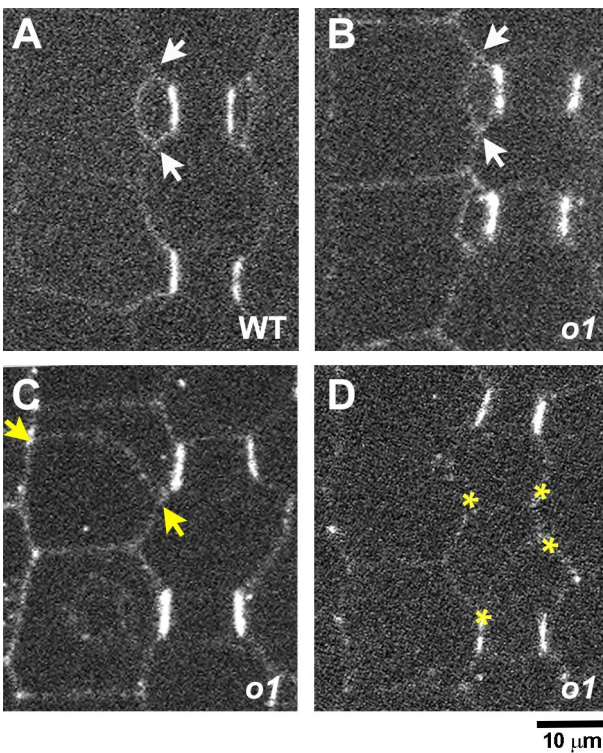
At what point does O1 play a role in determining the division plane: cell polarization, division plane establishment, division plane maintenance, or cytokinesis? We examined *o1* SMCs to determine if failed polarization caused the division plane defect. Polarization in SMCs occurs in a series of ordered steps with BRK1, PAN2, PAN1, and ROP proteins each becoming polarized sequentially, and each protein is required for the next to become polarized (Cartwright et al. 2009; Humphries et al. 2011; Zhang et al. 2012; Facette et al. 2015). Actin patch formation and nuclear migration are the last steps of cellular polarization (Facette et al. 2015). We assayed PAN1-YFP polarization, actin patch formation, and nuclear migration in *o1* and wild-type siblings to determine if cell polarization was normal. PAN1-YFP becomes polarized in SMCs prior to nuclear migration and remains polarized throughout division until after the subsidiary cell is formed (Cartwright et al. 2009). We analyzed recently divided SMCs for PAN1-YFP polarization and classified the daughter as having correct or incorrect division planes. PAN1-YFP polarized normally in all recently divided SMCs in *o1* (209/209), regardless of whether the SMC divided



**Figure 2.** Stomatal lineage cells have abnormal division planes in *o1*. The region of developing leaf 4 undergoing stomatal divisions was dissected, fixed, and stained with PI (red) and aniline blue (blue). Segregating wild-type (WT; A, B) and homozygous sibling *o1*-N1242A mutants (o1; C, D) are shown. White arrowheads mark correct divisions; yellow arrowheads mark incorrect divisions. A, B) Recently formed GMCs. C, D) Recently formed subsidiary cells, formed from SMCs. Z-projection of 3 confocal images. Scale bar, 5  $\mu$ m.

normally (136 cells) or abnormally (73 cells) (Fig. 3, *Supplemental Table S1*). When a GMC progenitor cell divided abnormally and failed to form a morphologically normal GMC, adjacent cells did not polarize PAN1-YFP—presumably because GMC fate was not correctly specified (Fig. 3D).

We also assayed whether the polar accumulation of actin and nuclear migration to the division site occurred normally in *o1* (Fig. 4, *Supplemental Fig. S1*). We hypothesized that nuclear migration, in particular, may be affected in *o1* mutants, as different myosin XI isoforms have been previously shown to be important for nuclear positioning (Tamura et al. 2013; Ali et al. 2020; Muroyama et al. 2020). We assayed actin and nuclear polarization at different developmental stages. As stomatal development proceeds, GMC width increases. Therefore, GMC width can be used as a proxy for the developmental state (Fig. 4, A and C). We quantified the number of SMCs with polarized actin patches (Fig. 4, A and B) and



**Figure 3.** PAN1-YFP polarizes correctly in *o1*. Recently formed subsidiary cells from *o1*-ref plants and wild-type siblings expressing PAN1-YFP were assayed for PAN1-YFP polarization. Arrows in (A–C) indicate correctly (white) or incorrectly (yellow) oriented cell walls generated from an SMC division. **A**) Recently divided SMC with polarized PAN1-YFP at the GMC-subsidiary cell interface. **B**) Correctly formed subsidiary cell from *o1*. PAN1-YFP is correctly polarized. **C**) Incorrectly oriented cell wall generated from an aberrant SMC division. PAN1-YFP is correctly polarized. **D**) Incorrectly oriented cell wall generated from an aberrant division of the GMC progenitor cell. Yellow asterisks in (D) indicate 4 corners of a cell formed from an aberrant GMC-generating division. Scale bar, 10  $\mu$ m.

polarized nuclei (Fig. 4D, Supplemental Fig. S1) in *o1* and wild-type siblings and observed normal polarization in *o1*. Actin patch formation and nuclear polarization are the last known steps of polarization and depend on the polarization of earlier factors, implying earlier factors that were not examined (BRK1, PAN2, and ROP) also polarized normally. Together, the data indicate that polarization is normal in *o1*, and the defect that leads to abnormal asymmetric divisions in *o1* SMCs occurs post-polarization.

### OPAQUE1 localizes to the phragmoplast

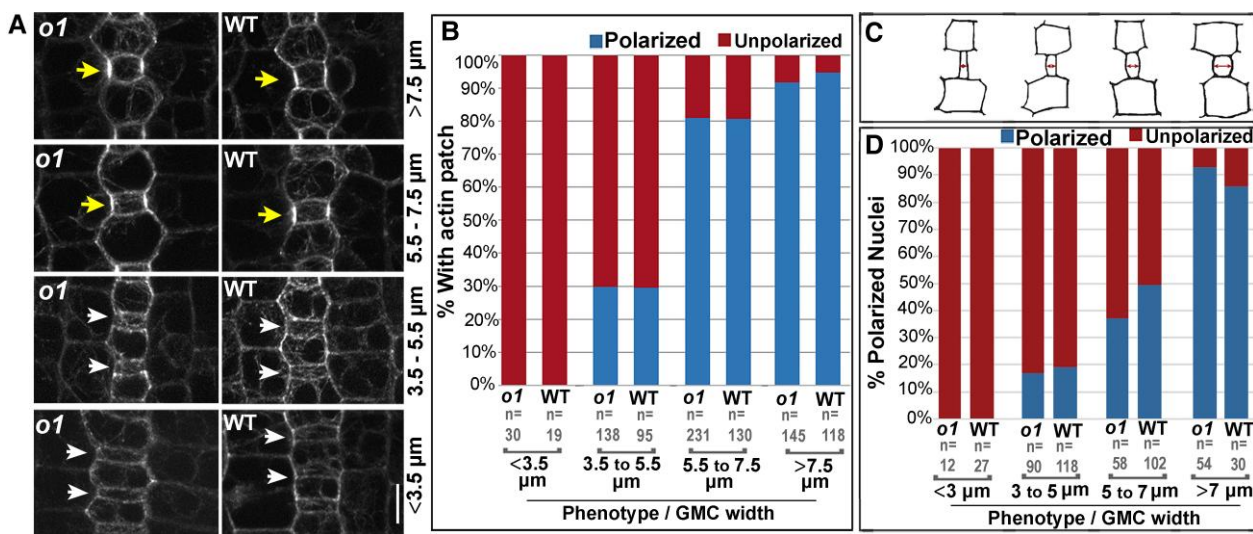
If O1 is not required for cell polarization, it must be required at a later stage of asymmetric cell division—specifically during division plane establishment, division plane maintenance, and/or cytokinesis. We used immunofluorescence to determine O1 localization during cell division, to ascertain at what stage of the cell cycle it might be important. We generated an O1-specific peptide antibody and performed

co-immunostaining for O1 and microtubules (Fig. 5, A to C; Supplemental Fig. S2). We observed specific staining, present only in wild-type but not mutant siblings, at the phragmoplast midline. O1 localized to phragmoplasts in all dividing cells we examined, including symmetrically dividing pavement cells (Fig. 5A), asymmetrically dividing GMC progenitor cells (Fig. 5B), and asymmetrically dividing SMCs (Fig. 5C). Previously, myosin XI isoforms were shown to localize to the phragmoplast, spindle, and cell cortex (Abu-Abied et al. 2018; Sun et al. 2018); phragmoplast localization was previously shown for a myosin VIII protein (Wu and Bezanilla 2014). We did not observe reproducible or specific staining in the PPB, spindle, or at the cortical division zone in dividing cells (Supplemental Fig. S2). However, the background signal was high and, therefore, any faint staining (such as in the spindle midzone or cell cortex) would be difficult to detect. These data suggest that O1 plays a role during cell division, especially in the phragmoplast during cytokinesis.

### Phragmoplasts are misguided in *o1* mutants

Since O1 localizes to the phragmoplast and the mutant has a post-polarization defect, we wanted to know if any division structures—especially the phragmoplast—were abnormal in *o1* mutants. We examined microtubule division structures using immunofluorescence on SMCs from developing leaf 4 (Fig. 6, D to K). We observed no abnormal PPBs in immunostained *o1* cells (0/55) (Fig. 2, D and F). Spindles persist only briefly and, therefore, were rare, but were always normal in *o1* (0/8) (Fig. 2, E and G). We observed abnormal late-stage phragmoplasts in some *o1* SMCs (18/52 = 35%) (Fig. 2, H to K). Abnormal phragmoplasts were misguided and did not localize at the expected site of division. However, the phragmoplast midline and microtubule alignment within the phragmoplast appeared normal and we saw no evidence for destabilized or fragmented phragmoplasts. This result is consistent with the absence of cell wall stubs in *o1* and suggests normal phragmoplast assembly. The misguided phragmoplasts showed either small deviations or large deviations in the division plane. Notably, all misguided phragmoplasts appeared to be correctly oriented at 1 edge, i.e. 1 edge of the phragmoplast was always anchored at the expected SMC division site. Because O1 is an actin motor, we also examined actin in *o1* mutants. The maize ACTIN-BINDING DOMAIN 2 (ABD2)-YFP marker used to assess actin patch formation does not localize to phragmoplasts (Sutimantanapi et al. 2014); therefore, we used fluorescently labeled phalloidin staining on fixed cells. Phragmoplast structure and orientation in phalloidin-stained cells was similar to microtubule-stained cells—while some were normal, a subset of phragmoplasts were misguided (Supplemental Fig. S3).

The *discordia* class of mutants displays post-polarization defects during SMC divisions (Gallagher and Smith 1999). Both *dcd1* and *dcd2* mutants are similar to *o1* in that they have aberrant GMC divisions, and normal nuclear polarization in SMCs (Gallagher and Smith 1999, 2000). The *Dcd2* gene has not yet been identified, but was mapped to



**Figure 4.** Actin patch formation and nuclear migration is normal in *o1*. **A)** The stomatal division zone was examined in leaf 4 of *o1*-N1242A mutant plants and wild-type siblings expressing the ABD2-YFP marker. GMCs in early developmental stages, found at the leaf base (lower panels), are narrow and width increases towards the leaf tip as development proceeds (upper panels). Early SMCs flanking narrow GMCs do not form an actin patch (white arrows). SMCs at later developmental stages, flanking wider GMCs, have an actin patch (yellow arrows). **B)** Percentage of SMCs with a polarized actin patch at progressive developmental stages in *o1*-N1242A mutant plants and their corresponding wild-type siblings. **C)** Cartoons depicting representative cell outlines at increasing GMC widths. Red arrows indicate where GMC width was measured. **D)** Percentage polarized nuclei in SMCs at progressive developmental stages in *o1*-N1242A mutant plants and their corresponding wild-type siblings. Fisher's exact tests comparing *o1* mutants to their respective wild-type siblings indicate no differences between mutants and wild type at each developmental stage ( $P > 0.05$  in all cases).

chromosome 4, and O1 lies within the mapping interval (Supplemental Fig. S4). Examination of *dcd2* seeds revealed that their endosperm is opaque. Complementation crosses between *dcd2* and 2 *o1* mutant alleles indicated that *dcd2*, which was identified based on subsidiary cell defects (Gallagher and Smith 1999), is allelic to *o1* (Supplemental Fig. S4).

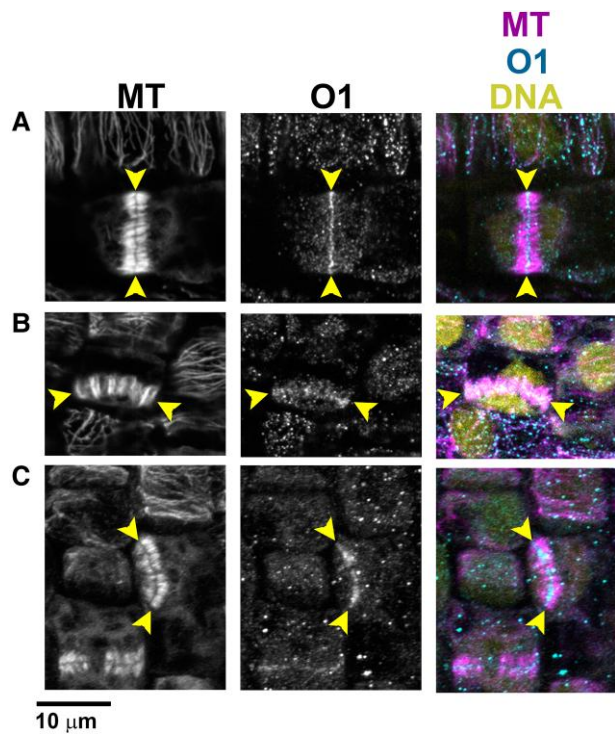
#### Live cell imaging of *opaque1* mutants indicates that phragmoplasts are misguided

To confirm that the abnormally divided cells seen in *o1* were a result of abnormal phragmoplast guidance, and to determine when phragmoplasts become misguided, we performed time-lapse imaging of dividing SMCs. CFP-TUBULIN (CFP-TUB) or YFP-TUB were crossed into *o1* mutants. We directly compared the location of the PPB, phragmoplast, and newly formed cell wall in dividing SMCs. These live markers also allowed us to observe PPBs and spindles to confirm our immunostaining results. We imaged cells in developing leaf 5 or 6 from 2 *o1* alleles and their corresponding wild-type siblings from prophase until the end of cytokinesis (Fig. 7 and Supplemental Fig. S5). In wild type, all dividing SMCs formed new cell walls that aligned with the former location of the PPB ( $n = 76$ ) (Fig. 7A and Supplemental Movie S1). Division proceeded normally (including normal PPBs and spindles) in all *o1* mutant cells until telophase. In *o1*-N1242A mutants, ~35% ( $n = 28/81$ ) of divisions displayed misguided phragmoplasts (Fig. 7B). In all cases, the initial site of contact between

the phragmoplast edge was always aligned with the site predicted by the PPB. This finding was consistent with our immunofluorescence data (Fig. 6), where 1 edge of the phragmoplast was always correctly oriented. However, as the phragmoplast continued to expand, the phragmoplast would “fall off track” in some cells and become misguided. The most severe division plane defects occurred when the phragmoplast became misguided shortly after initial contact with the existing cell wall (Supplemental Movie S2); small defects occurred when the phragmoplast became misguided near the completion of expansion (Supplemental Movie S3). We obtained similar results with the *o1*-5270-84 allele (Supplemental Fig. S5). These data indicate that (i) initial phragmoplast guidance, prior to first contact with the cortex, is separable from late-stage guidance and (ii) in *o1*, only late-stage phragmoplast guidance is defective.

#### The myosin O1 interacts with the maize orthologs of the kinesins POK1 and POK2

To gain insight into how O1 might be influencing phragmoplast guidance, we identified proteins that interact with O1 using co-immunoprecipitation followed by mass spectrometry (co-IP/MS). We extracted membrane and membrane-associated proteins from the stomatal division zone of *o1* plants and their wild-type siblings. We performed 3 biological replicates using the same antibody used in immunostaining. We also performed a second set of 3 independent biological replicates using a second anti-O1

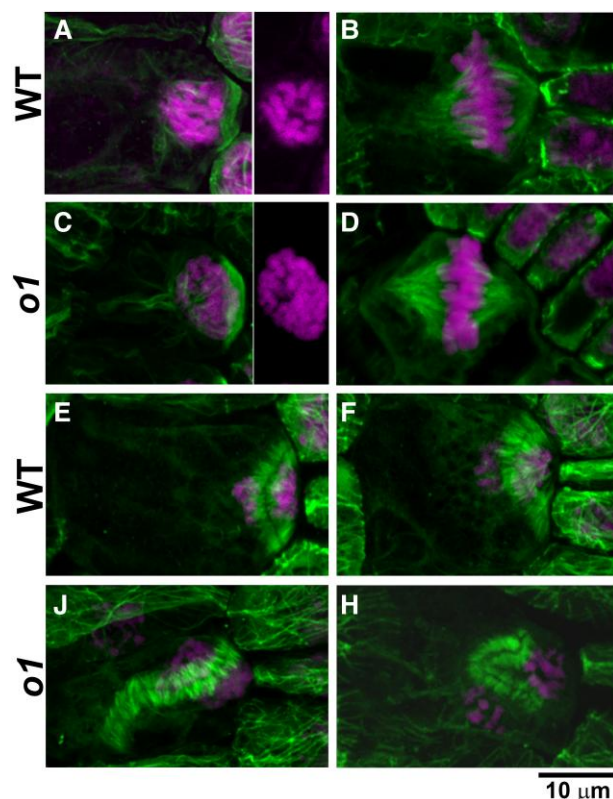


**Figure 5.** O1 localizes to phragmoplasts. Immunofluorescence detection of microtubules and O1 in wild-type cells (**A** to **C**). All samples are from the division zone of developing leaf 4. O1 is detected in phragmoplasts of symmetrically dividing cells (**A**), asymmetrically dividing stomatal lineage cells that will form GMCs (**B**), and asymmetrically dividing SMCs (**C**). Yellow arrowheads indicate phragmoplast ends. Scale bar, 10  $\mu$ m.

antibody generated from an independent rabbit. Both antibodies identified 2 protein bands present in wild type but not in their *o1* mutant siblings (Supplemental Fig. S6). Our MS analyses isolated 2 isoforms of O1 that have predicted sizes corresponding to these 2 bands. We considered proteins among the immunoprecipitates showing a more than 2-fold increase in abundance in the wild-type samples relative to those from their *o1* siblings, using both antibodies, as likely O1-interactors (Supplemental Data Set 1).

High-confidence interactors included many actin-associated proteins such as other myosins (including both myosin VIII and myosin XI family members), Actin Related Protein 2/3 complex (ARP2/3) proteins, and villin (Supplemental Data Set 1). We did not identify other actin-associated proteins such as fimbrin, NETWORKED, formins, or cofilin/actin-depolymerizing factors, suggesting that the observed interactions are likely specific. Notably, we identified the known myosin interactor MadA1 (myosin adaptors of family A) (Kurth et al. 2017).

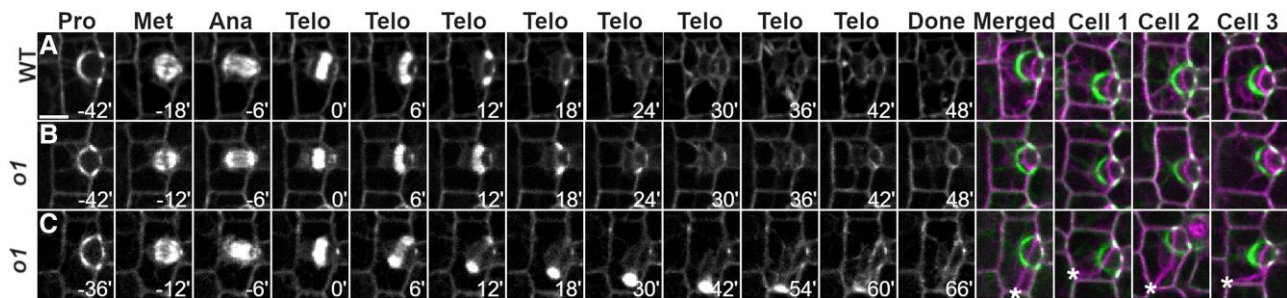
While we identified many actin-associated proteins, the only microtubule-associated proteins identified were 2 paralogous kinesins, KIN12C and KIN12D. We detected a third closely related kinesin, KIN12E, in the immunoprecipitates from 1 antibody, but was just below the threshold cutoff



**Figure 6.** O1 is required for normal phragmoplast guidance. Immunofluorescence detection of microtubules (green) and DAPI-stained nuclei (magenta) in wild-type (**A**, **B**, **E**, **F**) and *o1* cells (**C**, **D**, **J**, **K**). PPBs in SMCs (**A**, **C**) appear similar in wild-type siblings and *o1*-N1242A. Side panels in (**A** and **C**) show DAPI channel only to show condensed chromosomes. Spindles in SMCs (**B**, **D**) were similar in wild-type and *o1* SMCs. In *o1*, SMC phragmoplasts appeared as in wild type (**E**, **F**) or misguided (**J**, **H**). Z-projections of 40 to 60 images. Scale bar, 10  $\mu$ m applies to all images.

for the second antibody, making it another but lower-confidence potential interactor (Table 1). These 3 proteins are related to *Arabidopsis* POK1, POK2, and KIN12E (Müller et al. 2006; Lipka et al. 2014; Herrmann et al. 2018; Herrmann et al. 2021). Notably, these 3 maize proteins were all previously identified as direct interactors of maize TAN1 (Müller et al. 2006). TAN1 (in both *Arabidopsis* and maize), AtPOK1, and AtPOK2 all positively mark the cortical division site during cell division. These 3 proteins all localize to the phragmoplast and have phragmoplast guidance defects (Müller et al. 2006; Walker et al. 2007; Martinez et al. 2017; Herrmann et al. 2018; Mills et al. 2022). It is possible that the interaction between O1 and maize POK orthologs is important for phragmoplast guidance.

Although we replicated the interaction of O1 and POK-like KIN12 proteins with 2 different antibodies, we also determined if these proteins can interact using a yeast 2-hybrid (Y2H) assay (Supplemental Fig. S7). Myosin and kinesin proteins are large; in our hands, the full-length cDNA of O1 was lethal to *E. coli*. The KIN12C and KIN12D coding sequences are



**Figure 7.** Time-lapse imaging confirms a phragmoplast guidance defect in *o1*. CFP-TUB was used to observe progression of cell division SMCs from leaf 5 or 6 in *o1*-N1242A and corresponding wild-type siblings. **A)** Wild-type cell division. **B)** Correctly oriented *o1* cell division. **C)** Misoriented *o1* cell division. Pro, prophase; Met, metaphase; Ana, anaphase; Telo, telophase; Done, completed division; Merged, overlay of prophase (green) and completed division. Cells 1 to 3 show 3 additional representative cells. Time (min) is listed at the bottom of each image. Misoriented cell walls are indicated by asterisks. Z-projections of 6 images. All cells displayed at the same magnification; scale bar in (A), 10  $\mu$ m.

**Table 1.** Co-IP/MS results for O1 as bait protein and 3 related kinesin-like proteins

Accession	Name	Antibody 1		Antibody 2	
		WT peptides	<i>o1</i> peptides	WT peptides	<i>o1</i> peptides
Zm00001d052110_P042	OPAQUE1	546/435/546	0/0/0	474/493/458	0/0/0
Zm00001d052110_P007	OPAQUE1	258/229/268	0/0/0	230/232/213	0/0/0
Zm00001d022276_P012	KINESIN-LIKE PROTEIN 12C	29/3/18	0/0/0	36/56/8	1/0/0
Zm00001d041353_P002	KINESIN-LIKE PROTEIN 12D	12/6/9	0/0/0	16/24/5	0/0/0
Zm00001d034030_P017	KINESIN-LIKE PROTEIN 12E	3/3/3	0/0/0	7/9/6	4/2/0

Co-IPs were performed using two independently generated antibodies against O1. For each antibody, 3 biological replicates were conducted using wild-type plants and *o1*-N1242A mutants as a negative control. The number of peptides identified for each replicate are separated by a slash. At least 2 isoforms of O1 are present in developing leaves; no peptide corresponding to O1 was identified in homozygous *o1* mutants. The POK-like kinesins KIN12C and KIN12D were found only in wild-type samples and are considered plausible interactors. Fewer peptides were found for the related KIN12E and did not meet the threshold for a probable high-confidence interactor.

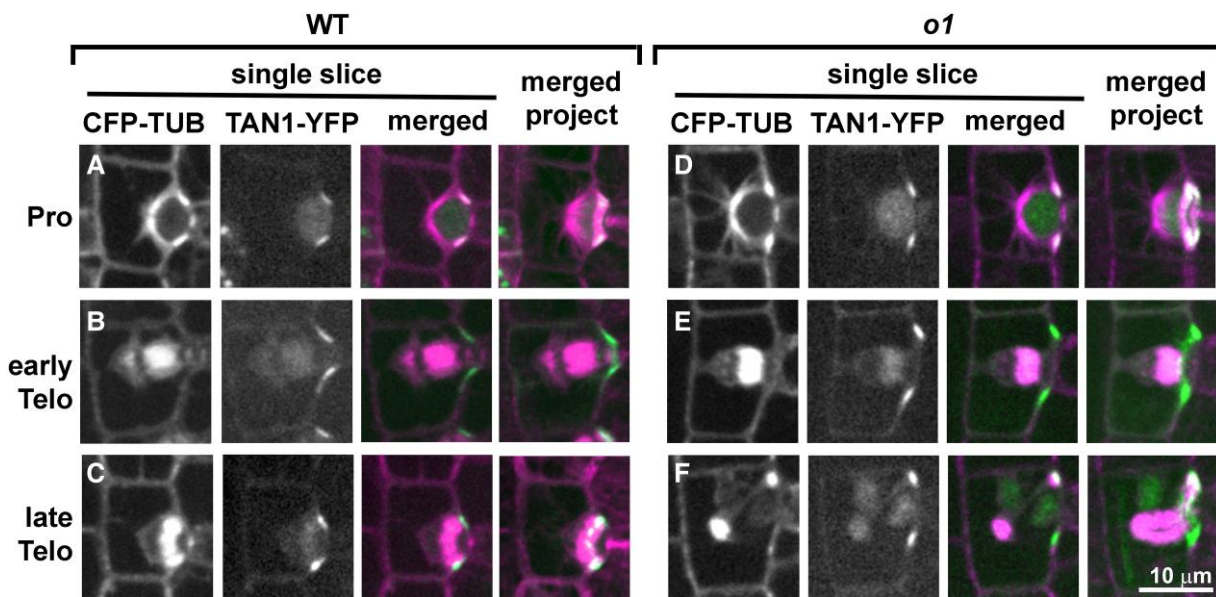
also large with 5,766 and 8,163 bp, respectively. Therefore, we used the C-terminal tails of O1, KIN12C, and KIN12D, which excludes the N-terminal motor domains. However, the C-terminal portion of KIN12C autoactivated in our Y2H assay, and the C-terminal portion of KIN12D did not show an interaction with O1. We used the full-length version of KIN12E, which is much smaller than KIN12C and KIN12D. O1 and KIN12E showed a positive interaction in our Y2H conditions (Supplemental Fig. S7).

Since POK proteins are division site markers, we wanted to know if the division site was being correctly maintained in *o1*. A failure to maintain the division site might explain the late-stage phragmoplast guidance defect. We used a previously characterized *ZmTAN1*-YFP marker line (Martinez et al. 2017) co-expressed with CFP-TUBULIN to determine if division plane maintenance was normal in *o1* mutants in different phases of mitosis. Importantly, in *Arabidopsis*, TAN1 localization to the division site is dependent on POK (Lipka et al. 2014). Therefore, if TAN1 localization is normal, we speculate that POK localization and division site specification in general is also normal. We separated telophase cells into early telophase, before the phragmoplast initially meets the cortex; and late telophase, when the defect in *o1* occurs. In wild-type SMCs, TAN1-YFP always correctly marked the predicted division plane throughout mitosis from prophase (Fig. 8A) through early telophase (Fig. 8B) and late telophase

(Fig. 8C; Supplemental Table S2;  $n = 154$  total cells). In *o1* mutants, TAN1-YFP always marked the correct division site from prophase (Fig. 8D) through early telophase (Fig. 8E) and late telophase (Fig. 8F), even in 44 late telophase cells where the phragmoplast became misguided (Supplemental Table S2;  $n = 231$  total cells). In rare cases, we observed TAN1-YFP at an additional site during early telophase (2/231 cells; Supplemental Fig. S8). Since this additional TAN1-YFP localization occurred only rarely, and we always saw correct TAN1-YFP localization in late telophase cells (when we see the phragmoplast guidance defect), ectopic TAN1 localization cannot be the primary cause of the *o1* phragmoplast guidance defect. Since the asymmetric division of GMC progenitor cells is also abnormal, we also examined TAN1 and phragmoplast localization during these divisions. Similar to SMC divisions, the division plane was correctly marked in divisions of GMC progenitor cells (Supplemental Fig. S9). These data indicate that O1 is not required for correct division site maintenance and specification. Rather, O1 is required for the phragmoplast to be guided to the specified division site during cytokinesis.

## Discussion

We characterized the myosin XI protein OPAQUE1/DCD2 as essential for late-stage phragmoplast guidance during maize

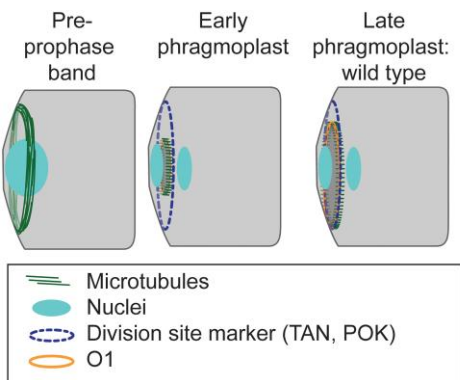


**Figure 8.** TAN1-YFP correctly marks the division plane during normal and abnormal *o1* SMC divisions. Dividing SMCs from leaf 5 or 6 in wild-type siblings (**A** to **C**) or *o1*-N1242A (**D** to **F**) cells co-expressing CFP-TUB (magenta) and TAN1-YFP (green). Single planes are shown in the first 3 panels and a full projection is shown in the last panel. In wild-type cells, TAN1-YFP correctly marked the predicted division plane throughout mitosis including prophase (**A**;  $n = 33/33$ ), metaphase ( $n = 30/30$ ), anaphase ( $n = 12/12$ ), early telophase (**B**;  $n = 21/21$ ), and late telophase (**C**;  $n = 58/58$ ). In *o1*-N1242A mutant cells, TAN1-YFP always correctly marked the division plane in prophase (**D**;  $n = 85/85$ ), metaphase ( $n = 20/20$ ), anaphase ( $n = 13/13$ ), and early telophase (**E**;  $n = 32/32$ ). In 2/32 cases, TAN1-YFP was also seen at an additional site during early telophase (see *Supplemental Fig. S10*). During late telophase, TAN1-YFP was at the cortical division site in *o1* SMCs with correctly oriented phragmoplasts ( $n = 38/38$ ) and incorrectly oriented phragmoplasts (**F**;  $n = 44/44$ ). Scale bar, 10  $\mu$ m applies to all images.

stomatal asymmetric divisions. Myosin XI family proteins are implicated in organelle trafficking and motility, cytoplasmic streaming, tip growth, auxin response, gravitropism, growth and division plane orientation (Sparkes et al. 2008; Avisar et al. 2009; Ueda et al. 2010; Vidali et al. 2010; Madison et al. 2015; Talts et al. 2016; Abu-Abied et al. 2018; Nebenführ and Dixit 2018; Olatunji and Kelley 2020). O1 was originally isolated as a mutant with opaque seed endosperm (Wang et al. 2012), which has abnormal protein bodies that are potentially caused by defects in ER motility. Recently, it was discovered that O1 also has defects in plant height and tassel branching (Zebosi 2022). Together with the data presented here, this suggests O1 influences multiple cellular processes. Although myosin XIs participate in nuclear positioning in other cell types (Tamura et al. 2013; Muroyama et al. 2020), O1 was not required for pre-mitotic nuclear migration in maize SMCs—either because O1 does not play a role, or its role is masked by genetic redundancy. The asymmetric division plane defect we observed in O1 was attributable to a late-stage phragmoplast guidance defect, consistent with localization of O1 to the phragmoplast midline. This phragmoplast guidance defect is phenotypically similar to *Arabidopsis* *pok1 pok2* double mutants (Lipka et al. 2014).

In addition to O1, many other proteins localize to the phragmoplast. Cell plate or phragmoplast localization has been observed for both myosin VIII and XI proteins—some of these myosins also localize to the PPB, cortical division

site, spindles, nuclei, and other organelles (Miller et al. 1995; Reisen and Hanson 2007; Sattarzadeh et al. 2008; Wu and Bezanilla 2014; Zhou et al. 2015; Abu-Abied et al. 2018; Duan and Tominaga 2018). It was previously observed that *Arabidopsis* MYOSIN XI-K localizes to the cortical division site, spindle, and phragmoplast; in the same article, a knockout in 3 myosin XI genes led to aberrant division planes in roots (Abu-Abied et al. 2018). However, no clear mechanism was advanced to explain how XI-K might lead to aberrant division planes. Indeed, given the complex localization pattern of this protein family, there are several potential roles for myosins during division. We showed here that in the case of O1, the division plane defect was specifically due to late-stage phragmoplast guidance. Although the role for O1 appears to be specific, it is plausible that myosins have other roles in cell division and that multiple myosins might fulfil the same role. Like XI-K, a myosin VIII from moss (*Physcomitrium patens*) localized to multiple mitotic structures, and the corresponding mutants also had a phragmoplast guidance defect, including slower vesicle delivery (Wu and Bezanilla 2014). Treatment with a general myosin inhibitor alters division planes (Molchan et al. 2002), cellular polarity (Holweg et al. 2003), and even resulted in incomplete divisions (Molchan et al. 2002). We did not observe polarity defects or failed divisions; together, these data suggest multiple roles for myosins during division and cytokinesis. Interestingly, we observed physical interactions between O1 and multiple other myosins, including both class VIII



**Figure 9.** Participation of O1 in late-stage phragmoplast guidance. The cortical division site is initially marked by the PPB and later by division site markers such as TAN1 and POK proteins (which are also present in the phragmoplast). In wild-type cells, the phragmoplast is guided to the division site by actin filaments and myosin VIII (Wu and Bezanilla 2014). After meeting the cortex, the phragmoplast continues to expand and interactions are stabilized by microtubules (Bellinger et al. 2021). In wild type, the phragmoplast fuses with the existing cell wall along the established division site, which is mediated by POK, TAN, and O1. In *o1* mutants, after initial contact the phragmoplast becomes misguided, resulting in abnormal division planes.

and XI myosins. Careful phenotypic analyses of different myosin mutants will help unravel the roles that each actin–myosin network plays during cell division.

Our data confirm that phragmoplast guidance is dynamic and that different proteins likely play different roles at different stages. Dividing cells treated with caffeine are unaffected in early-stage phragmoplast guidance, but the phragmoplasts disintegrate at later stages (Valster and Hepler 1997), often resulting in multinucleate cells. This observation led to the conclusion that phragmoplast guidance occurs in (at least) 2 steps (Valster and Hepler 1997) and is coupled with the observation that actin filaments connect the phragmoplast leading edge and the cell cortex (Valster and Hepler 1997; Wu and Bezanilla 2014). Indeed, studies of phragmoplast guidance in tobacco (*Nicotiana tabacum*) BY-2 cells indicate that rates of phragmoplast expansion vary, and slow considerably at the last stage of expansion when the leading edge first strikes the cortex (van Oostende-Triplet et al. 2017). At this final stage, the phragmoplast is more sensitive to actin depolymerization via latrunculin B (van Oostende-Triplet et al. 2017). It is also during this late stage that cortical telophase microtubules are incorporated into the phragmoplast (Mills et al. 2022). Since the phragmoplast always correctly meets the cortical division site at the initial site of contact in *o1* mutants, it is this last stage of expansion where O1 plays its role.

The identification of O1-interacting partners provides clues as to O1 functions. Since co-IP data will report both direct and indirect interactions, some observed interactors may be indirect. O1 interacted with many proteins, including actin-binding proteins, confirmed myosin binding partners, and other myosins. Our data are consistent with prior

observations of interactions between myosin and MadA1 (Kurth et al. 2017) myosin and (super)villin (Smith et al. 2013) and myosin and calmodulin/calmodulin-like (Shen et al. 2016). We also observed interactions between O1 and 2 closely related kinesin 12 proteins that are similar to *Arabidopsis* POK1. Even though many kinesins localize to the phragmoplast midline (Smertenko et al. 2018), we only identified KIN12D and KIN12E as O1-interactors. Moreover, the only other 2 microtubule-associated binding proteins we observed were IQ-domain proteins, recently shown in *Arabidopsis* to interact with POKs (Kumari et al. 2021). Mutations in *pok1* and *pok2* lead to misguided phragmoplasts similar to those observed in *o1* (Lipka et al. 2014; Herrmann et al. 2018). Mutations in other kinesins, such as *Arabidopsis* PAKRP1 (PHRAGMOPLAST-ASSOCIATED KINESIN-RELATED PROTEIN 1, also named KIN12a) and PAKRP2 (orphan kinesin) and *P. patens* KINID1a (KINESIN FOR INTERDIGITATED MICROTUBULES 1a) and KINID1b (orphan kinesins) lead to severe phragmoplast structural defects not seen in *pok* or *o1*, despite similar localization of all these proteins to the phragmoplast (Pan et al. 2004; Lee et al. 2007; Hiwatashi et al. 2008). Indeed, mutations in several genes encoding phragmoplast-localized proteins result in defects that alter phragmoplast structure, often leading to cell wall stubs and multinucleate cells (Müller et al. 2004; Bannigan et al. 2007; Ho et al. 2011; Zhang et al. 2018, 3; Schmidt and Smertenko 2019). The similar phenotypes of *pok* and *o1* coupled with their physical interaction suggest that they work together to ensure correct phragmoplast guidance.

How might O1 and the POK-like kinesins work together to ensure proper phragmoplast guidance and disassembly? During late and slow phragmoplast expansion (van Oostende-Triplet et al. 2017), the phragmoplast falls “off-track” and becomes misguided in *o1* (Fig. 9). Since POK proteins mark the cortical division site, and O1 localizes to the phragmoplast midline, a plausible model is that physical interactions between POKs at the division site and O1 at the phragmoplast midline help the phragmoplast “zip-up” around the cell cortex at the time of cell plate fusion. O1 and POK proteins may mediate interactions between the actin and microtubule cytoskeletons, to promote fusion of the phragmoplast at the cell wall. An alternative (non-mutually exclusive) model is that POK-like kinesins interact with O1 within the phragmoplast to coordinate microtubule and actin functions that promote slow phragmoplast expansion or phragmoplast disassembly once it reaches the cell cortex.

## Materials and methods

### Plant material and growth conditions

The 3 maize (*Z. mays*) *o1* mutant alleles (*o1-ref*, *o1-N1242A*, and *o1-84-5270-40*) used for this study were obtained from the Maize Genetics Cooperation stock center. Mutant alleles were backcrossed to the maize inbred line B73 1 to 4 times

and then selfed. In all experiments, segregating *o1* mutants were analyzed and compared with their corresponding wild-type siblings (grown side-by-side) as controls. The progeny from segregating families were classified based on their seed phenotype.

Mutant *o1* plants were crossed with various fluorescent protein-tagged maize lines generated by the Maize Cell Genomics Project (described at <http://maize.jcvi.org/cellgenomics/index.php>). PAN1-YFP (Humphries et al. 2011), the actin marker line YFP-ABD2-YFP (Mohanty et al. 2009), TAN1-YFP (Martinez et al. 2017), or the tubulin marker lines CFP- $\beta$ -tubulin and YFP- $\alpha$ -tubulin (Mohanty et al. 2009) were crossed to *o1* homozygous plants, and the F1 progeny were backcrossed to *o1* homozygous plants to obtain progeny expressing the fluorescent marker and segregating homozygous mutant (*o1/o1*) and phenotypically wild-type heterozygous (*o1/+*) individuals for experiments.

Plants used for phenotypic analysis and imaging were grown for 10 to 14 d in a greenhouse maintained between 22°F and 32°F under natural light illumination in greenhouses at the University of New Mexico, the University of Massachusetts at Amherst, or the University of California, Riverside. Plants were grown in Pro-mix Professional soil supplemented with Peters Excel 15-5-15 Cal Mag (weekly) and chelated iron (Southern Ag) as needed.

### Stomatal defects in expanded leaves

To quantify subsidiary cell and GMC defects, *o1* homozygous plants and their wild-type siblings from self-pollinated *o1-ref/+*, *o1-N1243/+*, and *o1-N1478A/+* were classified via their seed phenotypes. Impressions of fully expanded leaf 4 of *o1* homozygous and corresponding wild-type siblings were prepared using cyanoacrylate glue (Allsman et al. 2019) and imaged on a Nikon stereo microscope.

### Confocal microscopy

Instruments are described here; details on specific experimental protocols are given below.

For O1 immunostaining and nuclear polarization:

Images were collected using a Zeiss LSM710 with a 63 $\times$  (1.4 NA) oil immersion objective. Aniline blue was excited at 405 nm with a violet blue laser, and PI was excited using the 568-nm laser line and emission filter 620/60.

For PAN-YFP and ABD2-YFP localization:

ABD2-YFP-ABD2 and PAN1-YFP images were acquired with a custom spinning-disk confocal microscope (3i) equipped with a Yokogawa W1 spinning disk with 50- $\mu$ m pinholes, iXon Life 888 EM-CCD camera (Andor) using 150 EM-CCD intensification, ASI piezo stage, and solid-state lasers. YFP Images were acquired using a 60 $\times$  (1.2NA) silicone immersion objective, YFP fluorescence was excited by a 514-nm 100 mW solid-state laser at 6% with a dichroic excitation filter (Chroma), and a 542/27 emission filter (Semrock).

For microtubule immunostaining and phalloidin staining:

Immunolocalization and actin localization experiment images were collected with a Nikon A1R with a 60 $\times$  (1.40

NA) oil immersion objective. Alexa Fluor 488 and Alexa Fluor 568 were excited at the appropriate wavelengths of 488 and 568 nm, respectively, emission filters were 525/50 nm for Alexa Fluor 488 and 595/50 for Alexa Fluor 568.

For live imaging of CFP-TUBULIN, YFP-TUBULIN, and TAN1-YFP:

Time-lapse imaging was performed using a custom-built spinning-disk confocal microscope (Solamere Technology) with a Yokogawa W1 spinning disk (Yokogawa), EM-CCD camera (Hamamatsu 910 °c), and an Eclipse Ti-U (Nikon) inverted microscope. A 60 $\times$  water immersion lens (1.2 NA) was used with perfluorocarbon immersion liquid (RIAAA-678, Cargille). The stage was controlled by Micromanager software ([www.micromanager.org](http://www.micromanager.org)) with ASI Piezo (300  $\mu$ m range) and 3 axis DC servo motor controller. Solid-state lasers (Obis from 40 to 100 mW) and standard emission filters (Chroma Technology) were used. For CFP-TUBULIN, a 445-nm laser with an emission filter 480/40 was used. For YFP-TUBULIN and TAN1-YFP, a 514-nm laser with emission filter 540/30 was used.

All image analyses and figure preparations including cell measurements and processing were performed using ImageJ/FIJI (Schindelin et al. 2012) Adobe Photoshop CS6 or GIMP using only linear adjustments.

### Polarization measurements

To analyze PAN1 polarization, the basal 0.5 to 2.5 cm of leaf 4 of *o1-ref* homozygous and heterozygous wild-type sibling plants expressing PAN1-YFP were examined. Z-stacks of the stomatal division zone were collected using a spinning-disk confocal microscope (3i) described above using a 60 $\times$  silicone-oil immersion lens, YFP settings, and a 500-ms exposure. Recently formed subsidiary cells adjacent to a GMC that had not yet divided were assayed for polarity. PAN1-YFP polarization was scored by eye by comparing fluorescence intensity at the GMC-SMC interface and the adjacent SMC cell membrane. Cells were scored as having divided "normally" if both ends of the newly formed cell wall met the angled wall of the subsidiary cell.

To analyze F-actin polarization, the basal 0.5 to 2.5 cm of leaf 4 of *o1-N1242A/o1-N1242A* and wild-type *o1/+* expressing YFP-ABD2-YFP (Mohanty et al. 2009) were examined using a spinning-disk confocal microscope (3i) described below using YFP settings and a 100-ms exposure. Actin polarization was scored by eye by comparing fluorescence intensity at the GMC-SMC interface and the adjacent SMC cell membrane. GMC widths were measured using FIJI (Schindelin et al. 2012). Cell counts were then binned by GMC width and the % of cells with an actin was calculated.

To analyze nuclear polarization, double staining using aniline blue and PI of fixed tissue was performed. The stomatal division zone (basal 0.5 to 2.5 cm of unexpanded leaves) from leaf 4 of *o1* plants and wild-type siblings was isolated and fixed with FAA (3.7% formaldehyde, 5% acetic acid, 50% ethanol, w/v/v) for 1 h. Tissues were stained with 0.1% (w/v) aniline blue in phosphate-buffered saline (PBS) at pH

11 for 30 min. After rinsing with PBS, the tissues were stained with PI (10 µg mL<sup>-1</sup> in water) and mounted onto slides.

### Anti-O1 antibody generation

Custom rabbit antibodies were obtained from Pacific Immunology (Ramona, CA). Two peptides (Cys-NSEPKH<sup>I</sup>YESPTPTK and NSEPKH<sup>I</sup>YESPTPTK-Cys) were co-injected into 2 separate rabbits. The resulting sera were affinity-purified against both peptides, according to Cartwright et al. (2009). The 2 antibodies were named O1-11759 (Antibody 1) and O1-11760 (Antibody 2).

### Immunolocalization and phalloidin staining

Dual labeling of O1 and microtubules was performed as previously described, with minor modifications (Cartwright et al. 2009; Nan et al. 2019). The basal 0.5 to 2.5 cm of leaf 4 from *o1*-N1242A and wild-type siblings or *o1*-N1242A and wild-type siblings was used. Immunolocalization and phalloidin staining were performed separately. For immunolocalization, the dilutions used for rabbit anti-O1 (antibody O1-11759) and mouse anti-tubulin (Sigma Aldrich) antibodies were 1:1,000. Alexa Fluor 488-conjugated anti-mouse and Alexa Fluor 568-conjugated anti-rabbit (Invitrogen) were used at a dilution of 1:500. Samples were mounted in ProLong Gold Antifade with DAPI (ThermoFisher). For phalloidin staining, the basal 0.5 to 2.5 cm of leaf 4 from *o1* and wild-type siblings was fixed and stained with Alexa fluor 488-phalloidin (ThermoFisher) as described previously (Cartwright et al. 2009; Nan et al. 2019). Nuclei and cell walls were stained using 10 µg mL<sup>-1</sup> PI (ThermoFisher).

### CFP-TUBULIN, YFP-TUBULIN, and TAN1-YFP imaging

Time-lapse imaging was performed by taking a Z-stack every 6 min and assessing the morphology of the mitotic structure. The start of metaphase was counted from the first time the spindle was observed until the anaphase spindle was observed. This time point became the first time point for anaphase. Telophase timing was measured from the first time point when a phragmoplast was observed until the phragmoplast was completely disassembled. Leaf 5 or 6 from 12- to 14-d-old maize seedlings were used. Samples were prepared as described before (19).

### Dcd2-O mapping

*dcd2*-O was mapped to chromosome 4 using a near-isogenic line analysis after 4 backcrosses to B73. *dcd2* mutants in the B73 background were crossed to Mo17 and W64 inbred lines creating mapping populations for positional cloning. Markers on chromosome 4 were evaluated in over 1,300 *dcd2* mutants from the 2 mapping populations. Markers used for fine-mapping on chromosome 4 included single sequence repeat (SSR) markers from Sigma's Maize SSR Polymorphic Primer Set (umc2038, umc1620, bnlg1189, umc1871, and bnlg2162) and discovered single nucleotide polymorphism (SNP) markers

(N19-SNP: amplify with 5'cggagagaaagggttggtt and 5' ctcatcggttccgtttggtt and cut with MboII; F07-SNP: amplify with 5'tgaaataaaccagcttgc and 5'gccaaaggatgccttctc and cut with StuI; and AC185621: amplify with 5'aagtcaacctgtcgttcc and 5'cgccctctgattcaccatct and cut with PvuII).

### Co-IP/MS

Co-IP/MS experiments were performed as previously described with some modifications (Facette et al. 2015). Families segregating *o1*-N1242A and wild-type siblings were used, with 3 biological replicates per genotype. The experiment was entirely conducted twice, once with the O1-11759 antibody and once with the o1-117560 antibody. The cell division zone (0.5 to 2.5 cm from the leaf base) was isolated from unexpanded leaves 4 to 6 of 10- to 14-d-old seedlings; 4 to 10 seedlings were pooled per replicate to obtain 1.5 g of tissue. The leaf tissues were ground in liquid nitrogen. Extraction buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 5 mM EGTA, 5 mM EDTA, 0.3% [v/v] β-mercaptoethanol, 1% [v/v] Plant Protease Inhibitor Cocktail) was added as 1 mL per 0.25 g of tissue and the mixture was homogenized for 3 × 15 s, with 30-s breaks in between. Extracts were centrifuged at 25,000 × g at 4 °C in a microcentrifuge 2 times, then the supernatant was transferred and centrifuged at 110,000 × g at 4 °C for 45 min in an ultracentrifuge. After spinning, the supernatant was removed and the pellet was resuspended in 500 µL (per 0.25 g of starting tissue) of solubilization buffer (50 mM Tris-HCl pH 7.5, 150 mM NaCl, 1% [v/v] NP-40, 10% [v/v] glycerol, 0.05% [w/v] sodium deoxycholate). Samples were sonicated 2 × 15 s on ice and left rotating at 4 °C for 1 to 2 h. The extracts were centrifuged again at 110,000 × g at 4 °C for 45 min, and the supernatant was transferred to a new tube. Dynabeads coupled with anti-O1 antibody 11759 or 117560 were prepared according to the Dynabeads kit (Thermo Fisher) and added to the supernatant. The sample was incubated by rotating at room temperature for 30 min. The Dynabeads-Co-IP complex was washed according to the instructions of the Dynabeads kit.

The Dynabeads-Co-IP complexes were digested overnight at 37 °C with 400 ng of trypsin (Promega) per sample in 50 mM NH<sub>4</sub>CO<sub>3</sub> buffer. After digestion, peptides were reduced with 1 mM dithiothreitol at room temperature for 30 min and then alkylated with 5 mM iodoacetamide at room temperature in the dark for 30 min. Formic acid was added to a final concentration of 0.1% (v/v) and peptides were extracted from the beads and desalted using the C18-Stage-Tip method and then vacuum-dried. The dried peptides were reconstituted in 20 µL of 5% (v/v) formic acid and 5% (v/v) acetonitrile and 3 µL of each sample was injected into an LC column with a run time of 60 min of the gradient method for each run for MS analysis. Samples were run in technical triplicates on a Q-Exactive mass spectrometer with instrument and chromatography settings as described previously (Markmiller et al. 2018). The RAW files were analyzed using Andromeda/MaxQuant (version 1.6.0.16) (Cox and Mann 2008) with default settings except that match between the runs and label-free quantification settings were

enabled. Data were searched against a concatenated target-decoy database comprised of forward and reverse FASTA peptide sequences from the B73 RefGen\_v4, AGPv4 (Jiao et al. 2017). Data generated from samples from Experiment 1 (using antibody O1-11759) and Experiment 2 (using O1-11760) were searched against the database independently. A similar number of peptides was mapped for each sample from the different experiments. Experiment 1 resulted in 112,927 mapped peptides, with 21,900, 18,310, and 22,319 total in the 3 wild-type samples and 20,624, 14,777, and 14,977 in the 3 *o1* samples. Experiment 2 resulted in 158,803 mapped peptides, with 29,038, 29,219, and 28,476 total in the 3 wild-type samples and 24,262, 24,423, and 23,385 in the 3 *o1* samples.

### Y2H analysis

Total RNA was isolated from the stomatal division zone and cDNA was prepared using a Protoscript II First Strand DNA synthesis kit (NEB). A cDNA fragment encoding the C-terminal region of O1 was PCR-amplified with primers O1TAIL\_NO\_INTRO\_GWF and O1\_NOSTOP\_GWR. The full-length *KIN12E* cDNA was cloned using primers KIN12E\_GWF and KIN12E\_GWR. Fragments of *KIN12C* and *KIN12D* were cloned using primers KIN12C\_Cterm\_GWF and KIN12C\_GW\_R or KIN12D\_Cterm\_GW\_F and KIN12D\_GW\_R, respectively. Primer sequences are listed in **Supplemental Table S3**. The cDNA fragments were subcloned into pDONR221 (ThermoFisher) using BP Clonase II, and then cloned into either pAS-GW or pACT-GW (Nakayama et al. 2002) using LR Clonase II (ThermoFisher). Sanger sequence-verified plasmids were co-transformed into yeast strain AH109 according to (Gietz and Woods 92002). Co-transformants were selected on a synthetic defined (SD) medium without leucine and tryptophan (SD-LT) (Clontech). Between 12 and 80 colonies from each independent transformation were randomly picked and replated on SD medium without leucine, tryptophan, histidine, and adenine (SD-LTHA). Plates were incubated at 28 °C. Growth was assayed after 3 d.

### Accession numbers

The accession number for O1 is Zm00001d052110 (B73 RefGen\_v4, AGPv4) or Zm00001eb193160 (B73 RefGen\_v5). The accession number for *Kin12C* is Zm00001d022276 (B73 RefGen\_v4, AGPv4) or Zm00001eb328750 (B73 RefGen\_v5). The accession number for *Kin12D* is Zm00001d041353 (B73 RefGen\_v4, AGPv4) or Zm00001eb135060 (B73 RefGen\_v5). The accession number for *Kin12E* is Zm00001d034030 (B73 RefGen\_v4, AGPv4) or Zm00001eb057370 (B73 RefGen\_v5). The accession number for *ZmTan1* is Zm00001d038060 (B73 RefGen\_v4, AGPv4) or Zm00001eb286860 or (B73 RefGen\_v5).

### Acknowledgments

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### Author contributions

Q.N., H.L., J.M., A.J.W., E.J.B., C.G.R., and M.R.F. designed the research. Q.N., H.L., J.M., L.L., A.F., and M.R.F. performed the research; Q.N., H.L., J.M., L.L., A.F., E.J.B., C.G.R., and M.R.F. analyzed the data; A.W. contributed the *dcd2* allele; M.R.F., C.G.R., H.L., and Q.N. wrote the paper with input from other authors; M.R.F., C.G.R., and E.J.B. provided funding.

### Supplemental data

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

**Supplemental Figure S1.** Nuclear migration is normal in *o1*.

**Supplemental Figure S2.** O1 localizes to phragmoplasts.

**Supplemental Figure S3.** Actin in phragmoplasts of wild-type and *o1* mutant cells.

**Supplemental Figure S4.** *Dcd2* is allelic to O1.

**Supplemental Figure S5.** Time-lapse imaging of wild-type and *o1-84-5270-40* mutant subsidiary mother cell division.

**Supplemental Figure S6.** Validation of anti-O1 antibodies and co-IP/immunoblot results.

**Supplemental Figure S7.** The C-terminal region of O1 interacts with KIN12E.

**Supplemental Figure S8.** Unexpected TAN1-YFP localization in *o1* telophase SMCs.

**Supplemental Figure S9.** TAN1-YFP localization during division of GMC progenitor cells.

**Supplemental Table S1.** PAN1-YFP is polarized in *o1* subsidiary mother cells that have divided normally and abnormally.

**Supplemental Table S2.** Total number of SMCs examined for TAN1-YFP localization at different steps of the cell cycle.

**Supplemental Table S3.** Primers used in this study for yeast 2-hybrid analysis.

**Supplemental Data Set 1.** Peptides identified in co-IP/MS experiments in the *o1* mutant and wild-type siblings using 2 different antibodies.

**Supplemental Data Set 2.** Variables, degrees of freedom, and P-values for statistical tests.

**Supplemental Movie S1.** Normal asymmetric division of a wild-type SMC.

**Supplemental Movie S2.** Division of an *o1-N1242A* maize leaf subsidiary mother cell with a misguided phragmoplast.

**Supplemental Movie S3.** Division of an *o1-N1242A* maize leaf subsidiary mother cell with a misguided phragmoplast.

**Supplemental Movie S4.** Localization of TAN1-YFP in *o1-N1242A* SMC with phragmoplast

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