

# Fission yeast myosin I facilitates PI(4,5)P<sub>2</sub>-mediated anchoring of cytoplasmic dynein to the cortex

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Several key processes in the cell, such as vesicle transport and spindle positioning, are mediated by the motor protein cytoplasmic dynein, which produces force on the microtubule. For the functions that require movement of the centrosome and the associated nuclear material, dynein needs to have a stable attachment at the cell cortex. In fission yeast, Mcp5 is the anchor protein of dynein and is required for the oscillations of the horsetail nucleus during meiotic prophase. Although the role of Mcp5 in anchoring dynein to the cortex has been identified, it is unknown how Mcp5 associates with the membrane as well as the importance of the underlying attachment to the nuclear oscillations. Here, we set out to quantify Mcp5 organization and identify the binding partner of Mcp5 at the membrane. We used confocal and total internal reflection fluorescence microscopy to count the number of Mcp5 foci and the number of Mcp5 molecules in an individual focus. Further, we quantified the localization pattern of Mcp5 in fission yeast zygotes and show by perturbation of phosphatidylinositol 4-phosphate 5-kinase that Mcp5 binds to phosphatidylinositol 4,5-bisphosphate [PI(4,5)P<sub>2</sub>]. Remarkably, we discovered that the myosin I protein in fission yeast, Myo1, which is required for organization of sterol-rich domains in the cell membrane, facilitates the localization of Mcp5 and that of cytoplasmic dynein on the membrane. Finally, we demonstrate that Myo1-facilitated association of Mcp5 and dynein to the membrane determines the dynamics of nuclear oscillations and, in essence, dynein activity.

Mcp5 | PIP<sub>2</sub> | cytoplasmic dynein | myosin I | cortical anchoring

In fission yeast, during meiotic prophase, the fused nucleus of the zygote undergoes several hours of oscillatory movements from one cell pole to the other (1–4). Abrogation of these oscillations leads to improper chromosome pairing and recombination, leading to sporulation defects (2). The key players required for the movement of the nucleus are (i) the minus-end-directed motor protein cytoplasmic dynein, which is the force generator (2, 5); (ii) the anchor protein of dynein at the cortex, meiotic coiled-coil (CC) protein 5 (Mcp5), which acts as a link between dynein and the cell membrane (6, 7); and (iii) the microtubule cytoskeleton, which transduces the force produced by dynein to the spindle pole body (SPB, centrosome in yeast) and thereby to the nucleus underlying it (1). In fission yeast, cells lacking either dynein or Mcp5 are viable in interphase but do not exhibit oscillations during meiosis and thus show defects in sporulation (2, 6–8).

The minus-end-directed cytoplasmic dynein protein complex, a ~1-MDa protein, is a homodimer of two heavy chains that performs the functions of transport of cargo and nuclear organization in several systems (9). In fission yeast, dynein expression is temporally restricted to prophase I of meiosis, when it is required for powering the oscillatory movement of the fused nucleus (2). The population of dynein that is responsible for producing force is that which is bound stably to the underlying cortex via the anchor protein Mcp5 (6, 7).

The anchor protein Mcp5 is the homolog of budding yeast Num1, with an N-terminal CC and a C-terminal pleckstrin homology (PH) domain (6, 7). As its name suggests, Mcp5 is also expressed only

during meiotic prophase I (10). Both the CC and PH domains are essential for anchoring dynein at the cortex and hence for nuclear oscillations (6, 7). The CC domain is required for the attachment between Mcp5 and dynein, whereas the PH domain is required for Mcp5's membrane localization (6, 7). Using single-molecule imaging of cytoplasmic dynein, it was discovered that the attachment of dynein to the cortical anchors occurs in a two-step process, with the first step being the attachment to the microtubule (11). In addition, although dynein diffuses in the cytoplasm and on the microtubule, directional movement of dynein is activated upon binding to cortical Mcp5 (11). Recently, similar activation was also observed in budding yeast, wherein binding to Num1 released the inhibition of the dynein motor (12). The PH domain of Num1 was concluded to bind to PI(4,5)P<sub>2</sub> based on in vitro pull-down experiments as well as studies involving a temperature-sensitive mutant of the budding yeast phosphatidylinositol 4-phosphate 5-kinase [PI(4)P5K], Mss4 (13, 14). Functional homologs of Mcp5 in higher animals are ternary complexes that have divided the functions of dynein attachment and membrane localization among the three different proteins NuMA/LGN/G $\alpha$ i (15).

In budding yeast, recent work has been instrumental in understanding Num1's interaction with the membrane and the effect of this attachment with the associated cytoplasmic dynein motors in metaphase spindle movements (14, 16). In fission yeast during nuclear oscillations, the lipid molecule underlying Mcp5's association with the membrane has been speculated to be PI(4,5)P<sub>2</sub> (6), similar to that in budding yeast, but has not been tested. In addition, the precise localization of Mcp5 along the membrane and that of PI(4,5)P<sub>2</sub> in relation to Mcp5 is unknown.

## Significance

Molecular motors perform a variety of functions such as muscle movement and nuclear positioning during cell division. One such motor protein, cytoplasmic dynein, is responsible for oscillations of the nucleus during meiosis of the unicellular eukaryote, fission yeast. In this study, we used genetics, live-cell fluorescence microscopy, and image processing to identify the key players that keep dynein at the cortex while it performs the function of back-and-forth nuclear movement by producing force on the microtubule. We discovered that the membrane phospholipid PI(4,5)P<sub>2</sub> and, surprisingly, the myosin I motor Myo1 are both indispensable for nuclear oscillations in fission yeast. This study thus demonstrates one of the few instances of interplay between different classes of motor proteins in the cell.

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The authors declare no conflict of interest.

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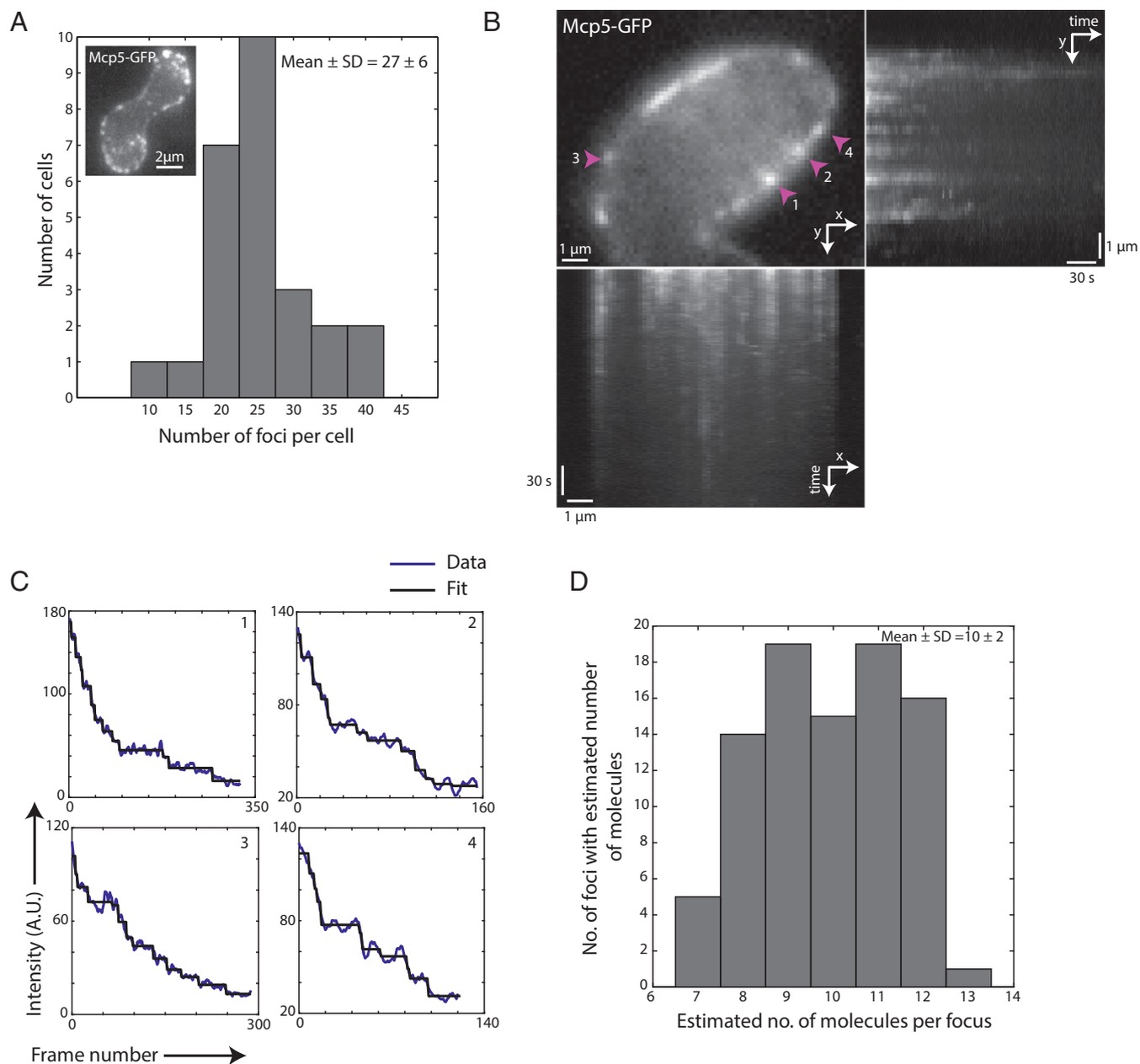
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The only known myosin I protein in fission yeast, Myo1, is required for actin regulation and maintaining cell shape (17, 18). Additionally, Myo1 has been demonstrated to be required for the organization of membrane lipids, with cells devoid of the non-filamentous, single-headed motor exhibiting aberrant sterol localization but normal actin cytoskeleton (19). Further, PI(4,5)P<sub>2</sub> in mammalian cells has been seen to localize to cholesterol-dependent rafts (20–22). Hence, Myo1 could serve as a good starting point for testing the role of membrane organization in dynein localization and activity at the cortex.

Here, we use fluorescence microscopy to quantify the number of Mcp5 foci at the membrane and the number of proteins per focus. By using a temperature-sensitive mutation of PI(4)P5K, we show that Mcp5 binds to PI(4,5)P<sub>2</sub> on the membrane. In addition, we demonstrate that perturbation of the membrane domains in a Myo1 deletion background affects Mcp5's association to the membrane, thereby affecting dynein localization and activity, leading to aberrant nuclear oscillations. We have thus demonstrated a remarkable interplay between different motor groups, with the activity of a myosin motor being required for the activity of cytoplasmic dynein.



**Fig. 1.** Each cluster of Mcp5 contains ~10 molecules. (A) *Inset* shows a representative maximum intensity projection of a fission yeast zygote (strain FY16854xFY16855; see Table S1) expressing Mcp5-GFP imaged using a confocal microscope; plot is the histogram of number of foci counted per cell versus number of cells with given number of foci. The mean ± SD of the number of foci per cell was estimated to be 27 ± 6 ( $n = 26$  cells). (B) Enlarged portion of a zygote expressing Mcp5-GFP (strain FY16854xFY16855; see Table S1) imaged using a TIRF microscope. The maximum intensity projection onto the  $y$  axis along time is shown on *Right* of the image and the maximum intensity projection onto the  $x$  axis along time is shown *Below*. (C) The points 1, 2, 3, and 4 marked with the magenta arrowheads in *B* are shown as representative data for bleaching step analysis. The raw five-frame sliding average data (blue) are plotted alongside the fits obtained from STEPFINDER software (black). (D) The histogram of the estimated number of molecules per Mcp5 focus is plotted to obtain a mean ± SD of 10 ± 2 molecules per focus ( $n = 89$  foci).

## Results

### Mcp5 Clusters into Spots of ~10 Molecules at the Membrane.

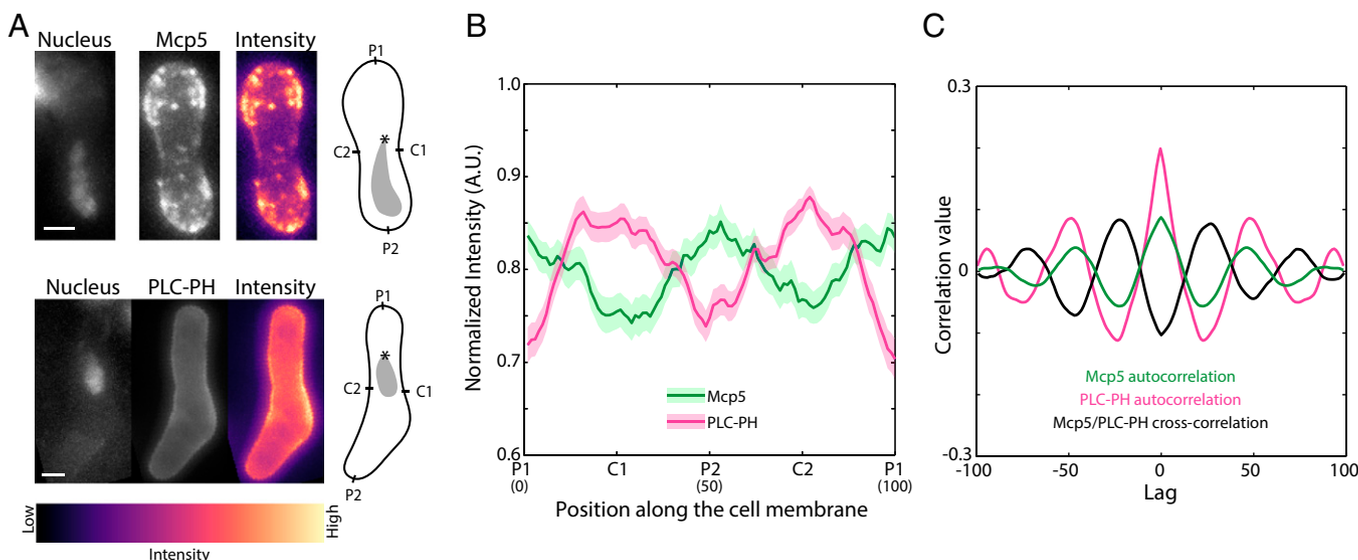
Mcp5 that is expressed during meiotic prophase clusters into distinct foci at the cell membrane (Fig. 1A, *Inset* and *Movie S1*). We first set out to count the number of such cortical Mcp5 foci per cell. We used a rolling ball average protocol with a diameter of 50 pixels to subtract background, and using intensity-based thresholding to include the top 2–4% of pixel intensities, we estimated  $27 \pm 6$  (mean  $\pm$  SD) Mcp5 foci per cell ( $n = 26$  cells). The Mcp5 spots were fairly stable over a period of minutes, which was ascertained by examination of the intensity profile of Mcp5 at the membrane over the course of the time lapse (Fig. S1A).

Because Mcp5 localizes to the membrane, we used Total Internal Reflection Fluorescence (TIRF) microscopy to count the number of Mcp5 molecules per focus (Fig. 1B). Time-lapse streams consisting of 5,000 frames with an exposure time of 20 ms were obtained of zygotes expressing fluorescent Mcp5. To reduce the background noise, a five-frame sliding average of the time lapse was taken. The intensity of the Mcp5 foci for each time point was then estimated using the Low Light tracking tool plugin for Fiji (23), which in addition also tracked the foci in x and y over time. The intensity of the foci was subjected to analysis using the STEPFINDER software (24) to count the number of bleaching steps of GFP (Fig. 1C and *SI Materials and Methods*). The number of steps obtained corresponded to the number of Mcp5 molecules, as each Mcp5 molecule contained a GFP tag (25). We thus estimated the number of Mcp5 molecules per cluster to be  $10 \pm 2$  (mean  $\pm$  SD,  $n = 89$  foci; Fig. 1D). Mean squared displacement (MSD) analysis of the tracking data yielded extremely slow diffusion of the Mcp5 clusters, thereby ruling out dynamic Mcp5 clusters at shorter time scales (Fig. S1B).

**PLC-PH and Mcp5 in a Zygote Show Preferential Localization to the Centers and the Poles of the Cell, Respectively.** The localization of the anionic phospholipid PI(4,5)P<sub>2</sub> is, to the best of our knowl-

edge, unreported in fission yeast zygotes during nuclear oscillations. In addition, the location of Mcp5 foci with respect to PI(4,5)P<sub>2</sub> is unknown. So we observed the localization of Mcp5 tagged with GFP and that of the PH domain of human phospholipase C (PLC $\delta$ 1) tagged with GFP (henceforth called PLC-PH) using confocal microscopy (Fig. 2A). The latter binds specifically to PI(4,5)P<sub>2</sub> (26) and thus acts as a proxy for PI(4,5)P<sub>2</sub> localization. The nucleus of the cell was stained using Hoechst to assign the leading and the lagging sides at any given time point. We observed that both Mcp5 and PLC-PH signals were enriched at the cell membrane (Fig. 2A and Fig. S2A). We assigned the leading pole as P1 and the lagging pole as P2 (Fig. 2A, schematic). C1 and C2 were the centers of the cell going clockwise from P1 and P2, respectively. We plotted the intensity profiles of Mcp5 and PLC-PH along these membrane positions. Intriguingly, we observed that locations that exhibited higher intensity of Mcp5 were accompanied by low intensity of PLC-PH, and conversely, wherever the intensity of Mcp5 was low along the membrane, the intensity of PLC-PH was found to be high (Fig. 2B and Fig. S2B and C). We define and henceforth refer to this recurrence of Mcp5 and PLC-PH intensities at poles and center of the cell, respectively, as “spatial periodicity.”

Specifically, on an average, the localization of Mcp5 was quantified to be higher at the poles than at the center of the cell, which is consistent with previous observations by Saito et al. and Yamashita et al. (6, 7). Conversely, PLC-PH was enriched at the membrane toward the cell center rather than at the poles (Fig. 2B). The spatial periodicity of the Mcp5 and PLC-PH signals was apparent in an autocorrelation analysis (Fig. 2C; see *SI Materials and Methods* for more details). The difference between Mcp5 and PLC-PH localization at the membrane was ascertained using cross-correlation analysis, with the two signals being inversely correlated (Fig. 2C and Fig. S2D). Additionally, the preferential localization of Mcp5 to the poles was found to be independent of the actin and microtubule cytoskeleton (Fig. S2E–H).



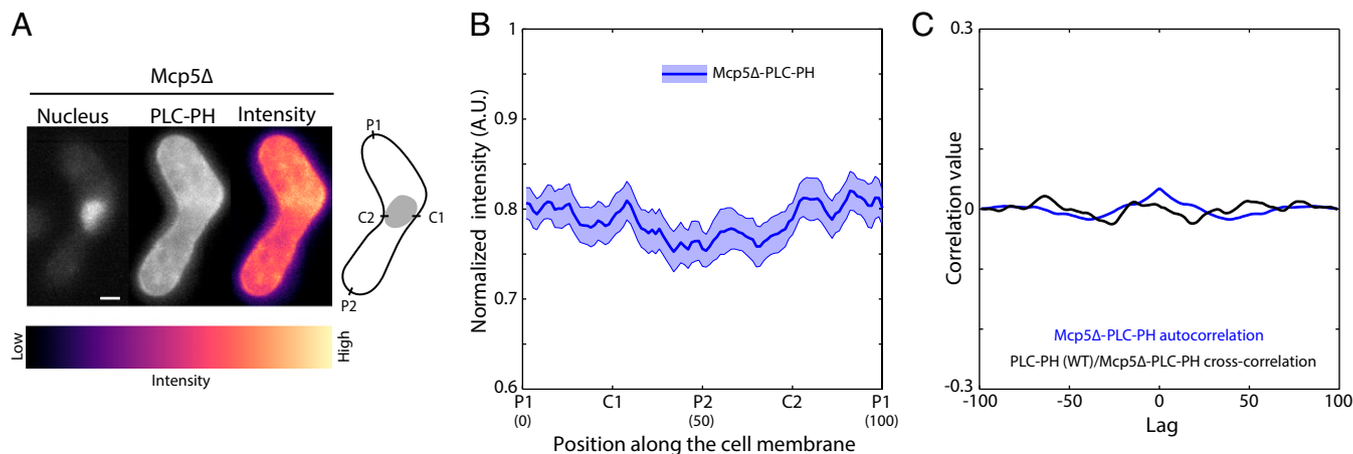
**Fig. 2.** Localization patterns of PLC-PH and Mcp5 on the membrane are complementary. (A) Confocal microscope images showing the maximum intensity projection of the Hoechst-stained nucleus (*Top and Bottom Left*), the summed intensity projection (*Top and Bottom Center*), and the intensity map (*Top and Bottom Right*) of a cell expressing fluorescent Mcp5 (strain FY16854xFY16855; see *Table S1*) and PLC-PH (strain AJC-D40xL972; see *Table S1*), respectively, and the schematics depicting the locations P1, C1, P2, and C2 along the membrane of the cell. The intensity range is shown below the images. The asterisk marks the position of the SPB with respect to the nucleus (gray). (Scale bar, 2  $\mu$ m.) (B) The mean normalized intensity profiles of Mcp5 (green,  $n = 41$  cells) and PLC-PH (magenta,  $n = 51$  cells) along the positions P1, C1, P2, and C2 are plotted. The lighter shaded regions depict the SEM of the normalized intensity profiles. The numbers in parentheses below the plot indicate the bin number (see *SI Materials and Methods* for details). (C) Autocorrelation analyses of Mcp5 signals (green) and PLC-PH signals (magenta) exhibit spatial periodicity, with Mcp5 signal being repeated at the poles and the PLC-PH signal at the centers. The intensity signals of Mcp5 and PLC-PH are inversely correlated (black), as can be seen from the negative correlation at zero lag.

**Mcp5 and PLC-PH Compete for the Same Binding Spots on the Membrane.** Although the inverse correlation of PLC-PH and Mcp5 localization at the membrane was surprising, it could have resulted from both molecules competing for PI(4,5)P<sub>2</sub> on the membrane. To test this hypothesis, we constructed a strain that expressed PLC-PH in the background of Mcp5 deletion (Mcp5Δ-PLC-PH, strain VA024; see Table S1; Fig. 3A). We observed that the PLC-PH intensity along the membrane in this strain did not exhibit preferential localization at the center (Fig. 3B). The spatial periodicity that was exhibited at the centers by wild-type PLC-PH localization had also disappeared in the autocorrelation plot (Fig. 3C). Cross-correlation analysis between wild-type PLC-PH intensity profile and Mcp5Δ-PLC-PH intensity profile confirmed this finding, with the signals showing no correlation (Fig. 3C).

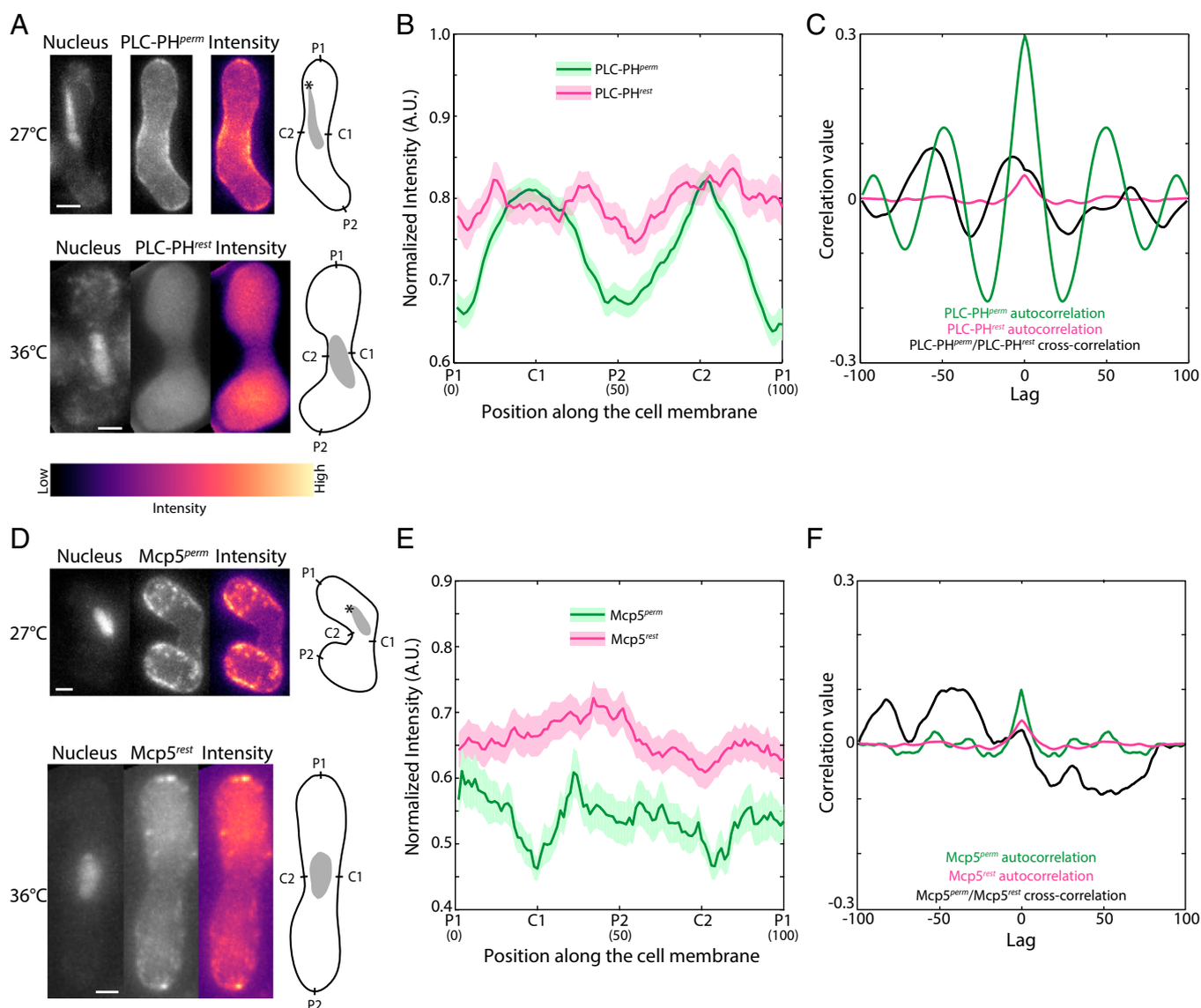
***its3-1* Zygotes Exhibit Perturbed Localization of PLC-PH and Mcp5 at the Membrane.** To confirm that Mcp5 indeed binds to PI(4,5)P<sub>2</sub>, we set out to deplete PI(4,5)P<sub>2</sub> in fission yeast zygotes by using the temperature-sensitive mutant of PI(4)P5K, *its3-1*. *Its3* in fission yeast was identified to be the sole kinase for the production of PI(4,5)P<sub>2</sub> (27). *its3-1* cells produce only 10% of PI(4,5)P<sub>2</sub> compared with wild-type cells even at permissive temperatures of 25–27 °C and almost no detectable PI(4,5)P<sub>2</sub> at a restrictive temperature of 33–36 °C (27). PI(4,5)P<sub>2</sub> in fission yeast is required for actin organization (28), and during meiosis, F-actin is essential for the progression of cell fusion (29). As a result, we did not observe zygote formation in crosses made and maintained at restrictive temperatures. Therefore, to induce meiosis, *its3-1* cells expressing PLC-PH were first crossed at 27 °C and then shifted to restrictive temperatures of 36 °C 3–4 h before imaging (Fig. 4A). In these cells, both the *its3-1* mutation and the expression of PLC-PH, which binds to free PI(4,5)P<sub>2</sub> and makes them unavailable for binding by other proteins, led to depletion of PI(4,5)P<sub>2</sub>. In such a background, we observed perturbation of the cortical localization of PLC-PH (Fig. 4A), with no specific preference for the cell centers (Fig. 4B and C), which is also apparent in the cross-correlation analysis between localization of PLC-PH at permissive temperature and PLC-PH of zygotes shifted to restrictive temperature (Fig. 4C). Additionally, the majority of PLC-PH signal was in the cytoplasm.

Similarly, the association of Mcp5 with the membrane was perturbed in zygotes with *its3-1* mutation at restrictive temperature, with preferential attachment to the poles being abrogated in these cells (Fig. 4D–F). At restrictive temperature, we observed no significant differences from wild type in Mcp5/PLC-PH distribution or Mcp5 foci number in other strains without the temperature-sensitive *its3-1* mutation (Fig. S3). In contrast, the number of Mcp5 clusters in PI(4,5)P<sub>2</sub>-depleted cells was significantly reduced from the wild-type estimate and that of *its3-1* cells grown at permissive temperature—that is, ~30 per cell, to  $7 \pm 5$  per cell (mean  $\pm$  SD,  $n = 27$  cells; Fig. S4A).

Then, to directly test whether Mcp5 localization was affected by PI(4,5)P<sub>2</sub> depletion, we transformed the *its3-1* strain with Mcp5-GFP and Mcp5-PH-GFP expression vectors (Fig. S4; see Table S1). Interphase cells in fission yeast do not normally express Mcp5 (6, 7), but cells transformed with the vectors showed clusters of Mcp5 and Mcp5-PH at the cortex, similar to those found in meiotic zygotes (Fig. S4B, D, F, and H). Cells were first grown at permissive temperature of 27 °C, fixed, and imaged using confocal microscopy (Fig. S4B, D, F, and H, Top). The cells were then shifted to restrictive temperature of 36 °C for 5.5 h, fixed, and then imaged in the same fashion (Fig. S4B, D, F, and H, Bottom). In *its3-1* strain expressing Mcp5, membrane localization of Mcp5 was observed after being grown in permissive temperature (Fig. S4E, green), whereas after 5.5 h in restrictive temperature, the cells did not exhibit Mcp5 along the membrane (Fig. S4E, magenta). The clustering of Mcp5 at the membrane was perturbed, with a reduction in total number of clusters as well as higher diffuse signal in the cytoplasm. This switch in localization pattern did not occur in wild-type cells at elevated temperatures (Fig. S4B and C). Additionally, the signal of Mcp5-PH in *its3-1* cells was apparent at the membrane at permissive temperature (Fig. S4I, green) and underwent a dramatic redistribution from the membrane into the cytoplasm after growth at restrictive temperatures (Fig. S4I, magenta), with a significant localization at the nucleus as well. The wild-type control did not exhibit this switch in distribution of Mcp5-PH at restrictive temperatures (Fig. S4F and G). This suggests that PH domain of Mcp5 depends on PI(4,5)P<sub>2</sub> for membrane localization.



**Fig. 3.** Preferential localization of PLC-PH to the cell centers is abolished in a cell devoid of Mcp5. (A) Confocal microscope images showing the maximum intensity projection of the Hoechst-stained nucleus (Left), the summed intensity projection (Center), and the intensity map (Right) of a Mcp5Δ cell expressing fluorescent PLC-PH (strain VA024; see Table S1) and the schematic depicting the nucleus (gray), and locations P1, C1, P2, and C2 along the membrane of the cell. (Scale bar, 2 μm.) (B) The mean normalized intensity profile of PLC-PH ("Mcp5Δ-PLC-PH," blue,  $n = 27$  cells) along the positions P1, C1, P2, and C2 is plotted. The lighter shaded region depicts the SEM of the normalized intensity profile. The numbers in parentheses below the plot indicate the bin number (see SI Materials and Methods for details). Unlike in Fig. 2B, PLC-PH's membrane localization does not show preference to the center. (C) Autocorrelation analysis of PLC-PH signals (blue) exhibits no spatial periodicity. The cross-correlation analysis of wild-type PLC-PH and PLC-PH in Mcp5Δ background (black) does not show any significant correlation.

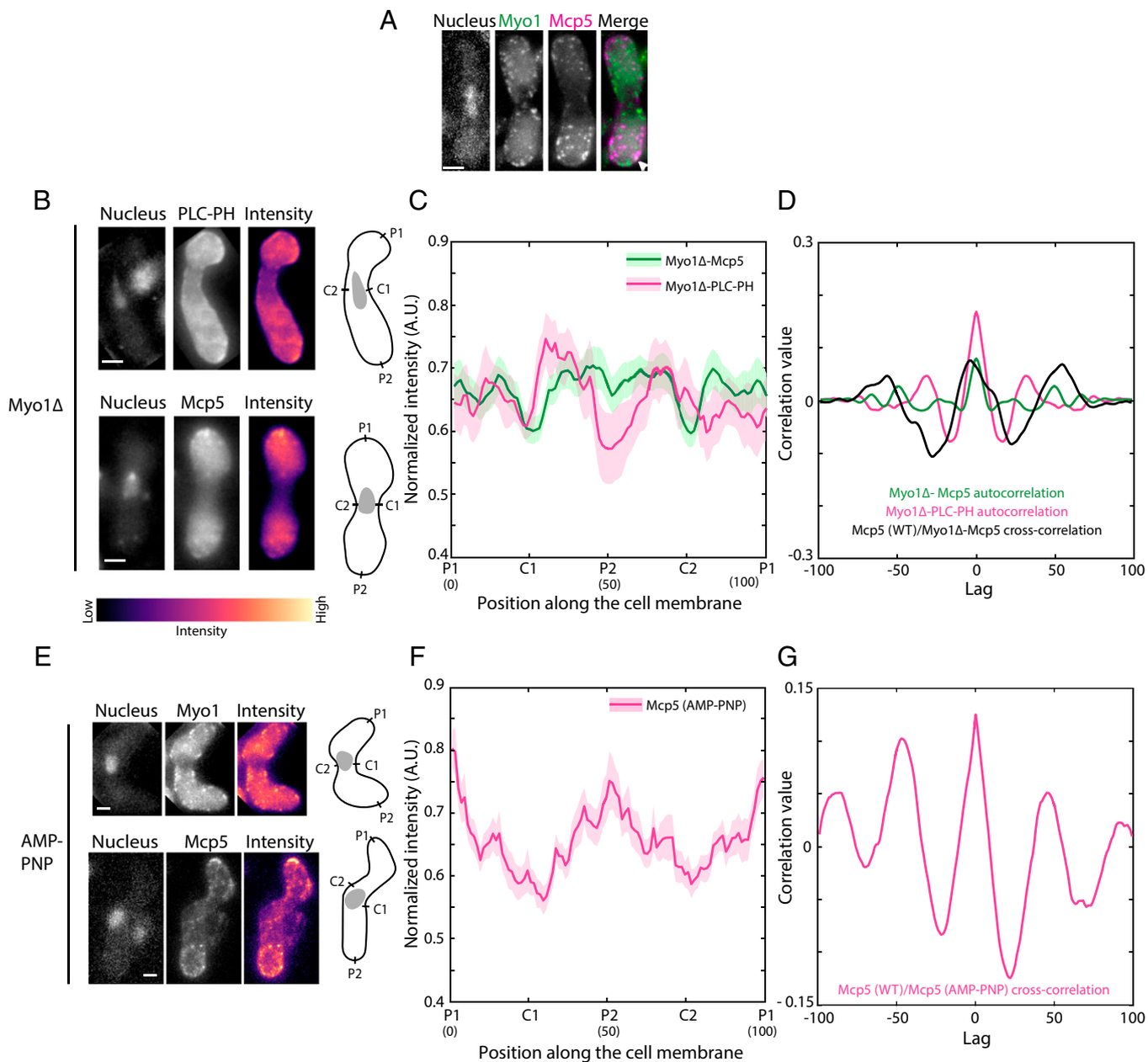


**Fig. 4.** PLC-PH and Mcp5 localization patterns are perturbed in *its3-1* zygotes. (A) Confocal microscope images showing the maximum intensity projection of the Hoechst-stained nucleus (Top and Bottom Left), the summed intensity projection (Top and Bottom Center), and the intensity map (Top and Bottom Right) of an *its3-1* zygote expressing fluorescent PLC-PH at permissive temperature of 27 °C ("PLC-PH<sup>perm</sup>", strain KP167xAJC-D40; see Table S1) and after shift to restrictive temperature of 36 °C ("PLC-PH<sup>rest</sup>", strain KP167xAJC-D40; see Table S1), respectively, and the schematics depicting the locations P1, C1, P2, and C2 along the membrane of the cell. The intensity range is shown below the images. (B) The mean normalized intensity profiles of PLC-PH<sup>perm</sup> (green,  $n = 76$  cells) and PLC-PH<sup>rest</sup> (magenta,  $n = 19$  cells) along the positions P1, C1, P2, and C2 are plotted. The lighter shaded regions depict the SEM of the intensity profiles. The numbers in parentheses below the plot indicate the bin number (see SI Materials and Methods for details). (C) Autocorrelation analysis of control PLC-PH<sup>perm</sup> signal (green) exhibits spatial periodicity at the cell centers, whereas PLC-PH<sup>rest</sup> signal (magenta) exhibits no such spatial periodicity. Cross-correlation analysis (black) reveals an aberrant correlation between the two signals, indicating a perturbation in PLC-PH<sup>rest</sup> localization upon shift to restrictive temperature. (D) Fluorescence microscope images showing the maximum intensity projection of the Hoechst-stained nucleus (Top and Bottom Left), the summed intensity projection (Top and Bottom Center), and the intensity map (Top and Bottom Right) of an *its3-1* zygote expressing fluorescent Mcp5 at permissive temperature of 27 °C ("Mcp5<sup>perm</sup>", strain KP167xVA026; see Table S1) and after shift to restrictive temperature of 36 °C ("Mcp5<sup>rest</sup>", strain KP167xVA026; see Table S1), respectively, and the schematics depicting the locations P1, C1, P2, and C2 along the membrane of the cell. (E) The mean normalized intensity profiles of Mcp5<sup>perm</sup> (green,  $n = 25$  cells) and Mcp5<sup>rest</sup> (magenta,  $n = 37$  cells) along the positions P1, C1, P2, and C2 are plotted. The lighter shaded regions depict the SEM of the intensity profiles. The numbers in parentheses below the plot indicate the bin number (see SI Materials and Methods for details). (F) Autocorrelation analysis of control Mcp5<sup>perm</sup> signal (green) shows a spatial periodicity that is absent in the Mcp5<sup>rest</sup> signal (magenta). Cross-correlation analysis (black) reveals improper correlation between the two signals upon shift to restrictive temperature. The asterisk marks the position of the SPB with respect to the nucleus (gray). Note that the SPB position is unknown in the PLC-PH<sup>rest</sup> and Mcp5<sup>rest</sup> cell, as nuclear oscillations were not apparent. (Scale bar, 2  $\mu$ m.)

#### Myo1 Is Essential for Localization of PLC-PH and Mcp5 at the Cortex.

Although we observed that Mcp5 occupies the poles more so than the centers of the zygotic membrane, it is not clear why this is the case. Myo1, which participates in the organization of sterol-rich domains in the fission yeast membrane (30), was then

tested for its role in ensuring restriction of Mcp5 localization primarily to the poles. Hence, we constructed a strain that expresses Myo1 tagged with GFP and Mcp5 tagged with 3mCherry to check for localization of Mcp5 with respect to Myo1 (strain VA046; see Table S1). We counted  $23 \pm 5$  Myo1 foci (mean  $\pm$  SD,



**Fig. 5.** Myo1 is required for proper localization of PLC-PH and Mcp5 to the membrane. (A) Fluorescence microscope images of the Hoechst-stained nucleus (Left), fluorescent Myo1 (green, Second From Left), fluorescent Mcp5 (magenta, Third From Left), and the merge of the three signals (Right) (strain VA046; see Table S1). A representative region of colocalization of Myo1 and Mcp5 signal is marked with the white arrowhead. (B) Fluorescence microscope images showing the maximum intensity projection of the Hoechst-stained nucleus (Top and Bottom Left), the summed intensity projection (Top and Bottom Center), and the intensity map (Top and Bottom Right) of Myo1 $\Delta$  zygotes expressing fluorescent PLC-PH and Mcp5 (Top and Bottom, respectively, strain VA050 and strain VA044xVA045; see Table S1), and the schematics depicting the locations P1, C1, P2, and C2 along the membrane of the cell. The intensity range is shown below the images. (C) The mean normalized intensity profiles of Myo1 $\Delta$ -Mcp5 (green,  $n = 27$  cells) and Myo1 $\Delta$ -PLC-PH (magenta,  $n = 11$  cells) along the positions P1, C1, P2, and C2 are plotted. The lighter shaded regions depict the SEM of the intensity profiles. The numbers in parentheses below the plot indicate the bin number (see SI Materials and Methods for details). (D) Autocorrelation analyses of Myo1 $\Delta$ -Mcp5 (green) and Myo1 $\Delta$ -PLC-PH signal (magenta) show spatial periodicity that is different from wild-type Mcp5 and PLC-PH signals (Fig. 2C). Cross-correlation analysis (black) between wild-type Mcp5 and Myo1 $\Delta$ -Mcp5 intensities yields an aberrant correlation. (E) Fluorescence microscope images showing the maximum intensity projection of the Hoechst-stained nucleus (Top and Bottom Left), the summed intensity projection (Top and Bottom Center), and the intensity map (Top and Bottom Right) of AMP-PNP-treated zygotes (see SI Materials and Methods for details) expressing fluorescent Myo1 and Mcp5 (Top and Bottom, respectively, strain FY13579xL975 and strain FY16854xFY16855; see Table S1) and the schematics depicting the locations P1, C1, P2, and C2 along the membrane of the cell. (F) The mean normalized intensity profiles of AMP-PNP-treated Mcp5 (SV91, magenta,  $n = 13$  cells) along the positions P1, C1, P2, and C2 are plotted. The lighter shaded regions depict the SEM of the intensity profiles. The numbers in parentheses below the plot indicate the bin number (see SI Materials and Methods for details). (G) Cross-correlation analysis (magenta) between wild-type Mcp5 and Mcp5 in AMP-PNP-treated cells. (Scale bar, 2  $\mu$ m.)

$n = 9$  cells) per cell, which is similar to the number of Mcp5 foci in a wild-type cell. However, we only observed weak colocalization of the two proteins (Fig. S4 and Fig. S5), likely because of the dynamic attachment of Myo1 to the membrane (19). We then constructed

strains that expressed fluorescently tagged PLC-PH and Mcp5 in the background of Myo1 deletion (strain VA050 and strain VA044xVA045; see Table S1). Similar to the case where PI(4,5)P<sub>2</sub> was depleted, both these constructs exhibited perturbed localization

of PLC-PH and Mcp5 compared with wild-type cells (Fig. 5B). The preferential associations of PLC-PH and Mcp5 to the centers and to the poles, respectively, were affected (Fig. 5C and D). Additionally, the number of Mcp5 clusters in the Myo1Δ zygotes was also significantly different from wild type with  $9 \pm 3$  (mean  $\pm$  SD,  $n = 23$  cells) clusters per cell (Fig. S6A), which is similar to the number of clusters in zygotes where PI(4,5)P<sub>2</sub> was depleted (Fig. S4A).

**Myo1's Motor Activity Is Not Essential for Myo1 and Mcp5's Membrane Localization.** Takeda et al. (19) have previously demonstrated that a Myo1 construct containing the motor domain and the tail homology 1 (TH1) domain of Myo1, without domains TH2 and TH3, is capable of organizing sterol-rich regions in the membrane of fission yeast, similar to that seen in cells expressing full-length Myo1. We visualized the localization of Mcp5 in this background where TH2 and TH3 domains of Myo1 were deleted (strain VA053; see Table S1). As expected, we observed localization of Mcp5 similar to that seen in wild type in these zygotes, with enrichment specifically at the poles ( $25 \pm 4$  clusters per cell,  $n = 10$  cells; Fig. S6B–D). However, it is not known whether Myo1's motor activity is required for its association to the membrane and consequently that of Mcp5. To test this, we first used a deletion strain of the calmodulin light chain 2 (Cam2), which is required for enhancing Myo1's motor motility in fission yeast (31). We did not observe a difference from wild type in Myo1 or Mcp5 localization in the Cam2Δ background ( $23 \pm 3$  clusters per cell,  $n = 5$  cells, and  $22 \pm 2$  clusters per cell,  $n = 9$  cells, respectively) and observed the characteristic cortical foci of dynein during the oscillations of the nucleus (Fig. S6E–G). To specifically block ATP-driven motor activity, we used AMP-PNP, a nonhydrolyzable analog of ATP. As seen previously (32), a concentration of 2 mM AMP-PNP was sufficient to prevent elongation of fission yeast mitotic spindle (Fig. S6H and I), indicating that AMP-PNP was successful in abrogating all motor activity. So too, when we treated fission yeast zygotes undergoing nuclear oscillations with AMP-PNP for 30 min, we first observed cessation of nuclear oscillations concomitant with a complete loss of dynein foci from the membrane ( $n = 13$  cells; Fig. S6J). However, treatment with AMP-PNP did not prevent wild type-like localization of Myo1 or Mcp5 (Fig. 5E–G) to the membrane, with  $20 \pm 5$  clusters ( $n = 7$  cells) and  $24 \pm 2$  clusters ( $n = 6$  cells), respectively, indicating that Myo1's motor activity was not essential for its association with the membrane or for Mcp5's Myo1-facilitated pole-specific localization at the cortex.

**Myo1 Deletion Affects Association of Dynein with the Cortex.** Because cells lacking Myo1 showed altered association patterns of Mcp5 with the membrane, we set out to test the role of Myo1 in ensuring cortical localization of dynein during nuclear oscillations. To this end, we constructed a strain that expressed fluorescently tagged dynein heavy chain (Dhc1) in the background of Myo1 deletion (strain VA049; see Table S1). In 90% of wild-type zygotes during nuclear oscillations, Dhc1 was observed on the SPB, as bright spots on the cortex and as fainter spots on the microtubule (Fig. 6, Top;  $n = 27/30$  cells). The cortical dynein spots in the wild-type cell are responsible for force production and, as a result, nuclear oscillations. In the wild-type cells exhibiting nuclear oscillations, we observed  $2 \pm 1$  (mean  $\pm$  SD) cortical dynein spots per cell. However, in the Myo1Δ background, Dhc1 was mostly absent from the cortex and the microtubule, being concentrated primarily on the SPB (Fig. 6, Bottom;  $n = 25/30$  cells). This confirms that by perturbing localization of Mcp5 to the membrane via deletion of Myo1, the association of dynein with the cortex is also affected.

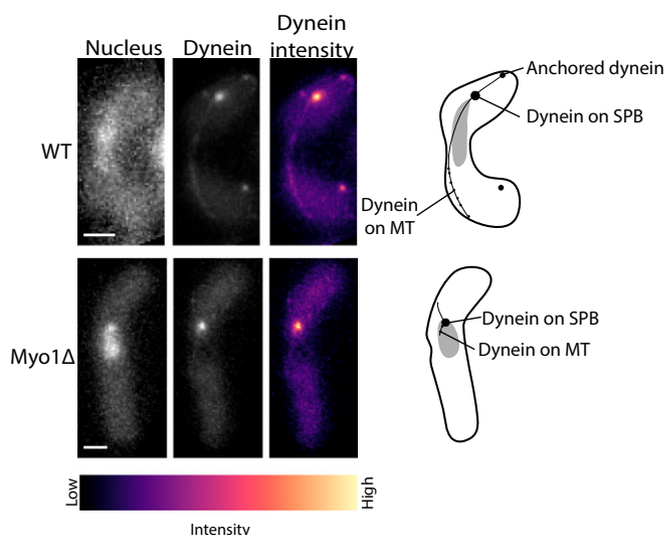
**PI(4,5)P<sub>2</sub> Depletion and Myo1 Deletion Abrogate Nuclear Oscillations.** Finally, we set out to test the effect of PI(4,5)P<sub>2</sub> depletion and Myo1 deletion on nuclear oscillations in fission yeast. Because

Mcp5 localization is affected in both cases, we expected irregularities in the nuclear oscillations in both these backgrounds. Wild-type cells exhibit robust oscillations from one cell pole to the other, with a period of oscillation of about 10 min and amplitude of around 5 μm (Fig. 7A and Fig. S7A). As before, we induced meiosis in cells expressing PLC-PH in the *its3-1* background at 27 °C and shifted them to the restrictive temperature of 36 °C 3–4 h before the horsetail nuclear movement stage. In this case, we observed that the oscillation of the horsetail nucleus was completely abrogated (Fig. 7B and Fig. S7B;  $n = 19$  cells), similar to what was observed in cells lacking Mcp5 altogether (Fig. 7D and Fig. S7B;  $n = 27$  cells). So too, deletion of Myo1, which leads to mislocalization of dynein, caused cessation of the nuclear oscillations (Fig. 7C and Fig. S7B;  $n = 24$  cells).

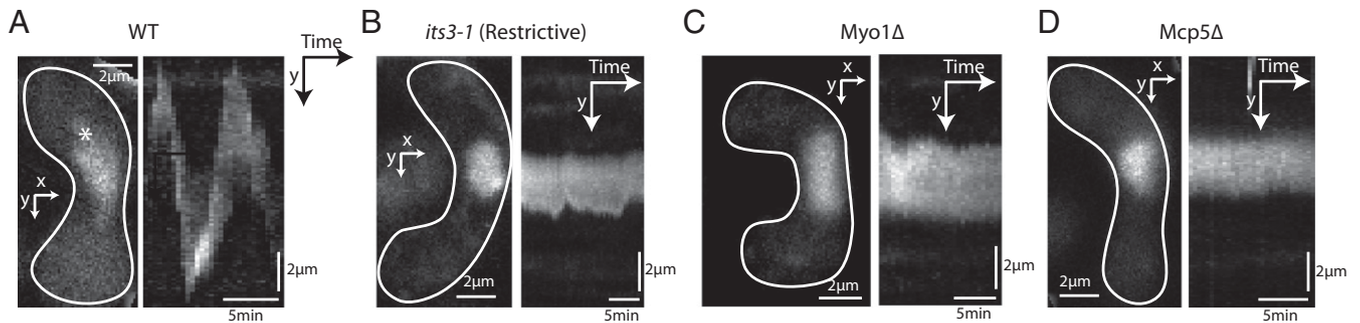
## Discussion

Our results indicate that the anchor proteins of dynein at the cortex in fission yeast are organized as distinct foci, about 30 in number per cell. Using TIRF microscopy, we identified that each of the Mcp5 foci in turn consists of around 10 molecules. The maximum number of dynein molecules that cluster into spots at the membrane during meiotic nuclear oscillations was roughly estimated to be around 30 per spot (11), which is the same order of magnitude as the number of Mcp5 molecules per cluster estimated here. This would imply a one-to-one interaction between dynein and Mcp5. In addition, we did not observe any significant dynamics or turnover of the Mcp5 clusters at time scales of ~1 min and ~10 min, similar to what was observed previously (7). However, we cannot rule out dynamics at longer time scales, as studies in budding yeast using Num1 tagged with photo-activatable GFP have demonstrated redistribution of Num1 among clusters in about 30 min (33).

We next identified and quantified the localization pattern of Mcp5 and PLC-PH on the membrane of fission yeast zygotes. We observed that the signals of both Mcp5 and PLC-PH showed spatial periodicity along the membrane, with the former appearing prominently at the poles and the latter appearing at the center of



**Fig. 6.** Dynein dissociates from the cortex in cells lacking Myo1. Fluorescence images of Hoechst-stained nucleus (Top and Bottom Left), fluorescent Dhc1 ("Dynein," Top and Bottom Center), and the intensity map of Dhc1 in wild-type (Top) and Myo1Δ (Bottom) fission yeast zygotes (strain SV111 and strain VA049, respectively; see Table S1). The intensity range is shown below the images. The schematics that appear alongside the images depict the localization of dynein at different locations in the cell. In a Myo1Δ background, dynein is absent from the cortex and the microtubules and appears almost exclusively on the SPB. (Scale bar, 2 μm.)



**Fig. 7.** Nuclear oscillations are affected in cells depleted of PI(4,5)P<sub>2</sub> and lacking Myo1. (A) Image of the nucleus stained with Hoechst (Left) and the maximum intensity projection onto the y axis along time (Right) of a representative control cell (strain FY16854xKP167; see Table S1), showing robust nuclear oscillations. The amplitude and period of oscillations are quantified in Fig. S7A. (B) Image of the nucleus stained with Hoechst (Left) and the maximum intensity projection onto the y axis along time (Right) of a representative *its3-1* cell shifted to restrictive temperature 3–4 h before the nuclear oscillations phase (strain KP167xAJC-D40; see Table S1), showing impaired nuclear oscillations. (C) Image of the nucleus stained with Hoechst (Left) and the maximum intensity projection onto the y axis along time (Right) of a representative Myo1Δ cell (strain FY13572; see Table S1), showing cessation of nuclear oscillations. (D) Image of the nucleus stained with Hoechst (Left) and the maximum intensity projection onto the y axis along time (Right) of a representative Mcp5Δ cell (strain VA024; see Table S1), showing abrogation of nuclear oscillations.

the cell. We discovered that the signal intensities of Mcp5 and PLC-PH surprisingly showed complementarity in their localizations, with a higher intensity of Mcp5 where PLC-PH intensity was low and vice versa. We demonstrated that this pattern was likely formed due to competition between Mcp5 and PLC-PH for PI(4,5)P<sub>2</sub> on the membrane by quantification of the localization pattern of PLC-PH in the background of zygotes lacking Mcp5. Previous research in budding yeast has identified the homologous protein Num1's PH domain to have the highest affinity for PI(4,5)P<sub>2</sub> among all of the proteins containing PH domains in *Saccharomyces cerevisiae* (13). Therefore, the complementarity in Mcp5 and PLC-PH localizations at the membrane is likely due to Mcp5's high affinity for PI(4,5)P<sub>2</sub>. In budding yeast, the formation of clusters was dependent on the N-terminal CC of Num1, which was identified to have a Bin-Amphiphysin-Rvs (BAR)-like domain, capable of homodimerization, similar to BAR domain-containing proteins (16). Accordingly, the formation of Num1 clusters is proposed to occur at regions of high curvature on the membrane via their BAR-like domains (16). Perhaps, Mcp5 could use a similar strategy for clustering at the membrane. To answer this, future experiments that identify whether such a domain exists in Mcp5 will have to be performed.

Further, we identified PI(4,5)P<sub>2</sub> to be the likely binding partner of Mcp5 at the cortex. First, we expressed PLC-PH in a strain with a temperature-sensitive mutation of *its3*, the PI(4)P5K in fission yeast. By shifting zygotes to restrictive temperatures just before meiotic prophase, we depleted PI(4,5)P<sub>2</sub> in the cells and consequently observed that the localization patterns of both PLC-PH and Mcp5 were perturbed. Second, we transformed *its3-1* cells with Mcp5 and compared its localization at permissive and restrictive temperatures. Although at permissive temperature Mcp5 exhibited cortical localization, after 5.5 h at restrictive temperature, the cortical localization was abrogated. Third, *its3-1* cells were transformed with the PH domain of Mcp5 tagged with GFP to visualize its specific interaction with the membrane. Indeed, growth at permissive temperature exhibited Mcp5-PH molecules on the membrane, whereas shift to restrictive temperature showed a dramatic relocation of Mcp5-PH to the cytoplasm. Taken together, these results indicated that Mcp5 depends on PI(4,5)P<sub>2</sub> for cortical localization in fission yeast. In HeLa cells during anaphase, dynein has been observed to interact with phosphatidylinositol 4-phosphate and PI(4,5)P<sub>2</sub> on the membrane via the ternary complex protein NuMA (34). As discussed previously, Num1's PH domain also depends on PI(4,5)P<sub>2</sub> for membrane localization (13, 14, 16). In all these cases, a stable interaction between the motor protein and the membrane is essential for nuclear movement and positioning. In other systems

that use an association between the membrane and dynein for nuclear organization, such as in *Drosophila* neuroblasts, *Caenorhabditis elegans* one-celled embryo, and metaphase of mammalian cells, myristoylation of the Gai protein in the ternary complex is essential for membrane localization of dynein (35–38). Recently, it was discovered that dynein clusters on cholesterol-rich microdomains on late phagosomes, which enables them to move in a persistent minus-end-directed fashion toward lysosomes (39). However, it is unknown whether the localization of dynein to these microdomains is mediated by PI(4,5)P<sub>2</sub>.

We identified the myosin I motor in fission yeast, Myo1, to be indispensable for the organization of Mcp5 and consequently that of cytoplasmic dynein at distinct locations on the cortex. At present, reports of interaction between actin-based motors and microtubule-based motors have been restricted to their roles in cargo transport, such as the melanophore transport in fish by coordination of myosin V, kinesin, and cytoplasmic dynein (40). Here, we have discovered an interplay between the two motor groups, wherein the force-generating function of cytoplasmic dynein at the cortex is facilitated by the activity of Myo1 that serves to organize the membrane. Although it is not yet clear how Myo1 brings about membrane organization, we discovered that the ATP-driven motor activity of Myo1 is not required for its membrane organization function and hence for localization of Mcp5 and dynein. The sterol organization by Myo1 could effect preferential enrichment of PI(4,5)P<sub>2</sub> at the poles, which in turn would increase Mcp5's association with the poles via its PH domain. Moreover, local enrichment of PI(4,5)P<sub>2</sub> in microdomains has been proposed to be capable of inducing positive curvature in the membrane (41). If Mcp5 does indeed contain a BAR-like domain at its N terminus, clustering of Mcp5 could be further enabled by its targeting to regions of high curvature on the membrane. In our experiments, we observed only a slight increase in PLC-PH intensity at the poles in the absence of Mcp5 (Fig. 3), likely due to restrictions imposed by using PLC-PH as a proxy for PI(4,5)P<sub>2</sub> localization (42). Future experiments with improved probes for PI(4,5)P<sub>2</sub> will provide better insight into the Myo1–PI(4,5)P<sub>2</sub>–Mcp5–dynein axis.

Finally, we demonstrated that either depletion of PI(4,5)P<sub>2</sub> or deletion of Myo1 essentially renders dynein inactive and leads to cessation of nuclear oscillations, much like the phenotype observed in cells lacking dynein or Mcp5 (2, 6, 7).

## Conclusions

In conclusion, using advanced light microscopy and image analysis, we quantified the dynamics and organization of both Mcp5 and

PLC-PH at the membrane. We discovered that the anchor protein Mcp5 is dependent on the phospholipid PI(4,5)P<sub>2</sub> and the myosin motor Myo1 for proper cortical localization. Cortical localization of cytoplasmic dynein, which is required for force production, is abolished in a Myo1 deletion background. Perturbation of PI(4,5)P<sub>2</sub> levels or deletion of Myo1 leads to aberrant nuclear oscillations and, in effect, perturbed dynein activity.

## Materials and Methods

**Strains and Media.** The fission yeast strains used in the study are listed in Table S1. Cells were grown on Yeast Extract (YE) medium or Edinburgh Minimal Medium (EMM) (43) with the appropriate supplements. Unless otherwise indicated, cells were grown at a temperature of 27 °C in an incubator (Ascension Innovations).

**Construction of Strains.** Strain VA024 was constructed by crossing strain AJC-D40 ( $h^+$  *leu1-padh1-PPH-GFP-leu1-32 ura4-D18*; see Table S1) with strain FY16831 ( $h^-$  *ade6-M26 ura4- D18 mcp5::ura4+*; see Table S1) following the Random Spore Analysis (RSA) protocol (43). Similarly, strain VA026 was constructed by crossing strain AJC-D40 (see Table S1) with strain IF960 ( $h^{90}$  *num1-3mCherry-hphR ade6-M216 leu1*; see Table S1), strains VA044 and VA045 were constructed by crossing strain FY13570 ( $h^-$  *myo1::kanr ade6-M216 leu1*; see Table S1) with strain AY277 ( $h^{90}$  *num1-3mCherry-hphR ade6-M216 leu1 ura4-D18*; see Table S1); strain VA046 was constructed by crossing strain FY13579 ( $h^-$  *myo1+-GFP<<kanr ade6-M210 leu1*; see Table S1) with strain AY277; strain VA049 was constructed by crossing strain SV111 ( $h^{90}$  *dhc1-mCherry-nat r leu1-32 lys1 ura4-D18*; see Table S1) with strain FY13570; strain VA050 was constructed by crossing strain AJC-D40 with strain FY13570 (see Table S1); strain VA051 was constructed by crossing strain SV91 ( $h^{90}$  *mcp5-mCherry-kan r dhc1-GFP-Leu2 leu1-32 lys1 ura4-D18*) with strain FY12794 ( $h^{90}$  *leu1-32 ura4-D18 ade6-M210 cam2::ura4+*); and strain VA053 was constructed by crossing strain AY277 with strain FC1261 [ $h^-$  *myo1 (H1TH1)-kanMX leu1-32 ura4-D18 ade6*] and selecting the appropriate strain using RSA.

**Induction of Meiosis and Preparation of Cells for Imaging.** Meiosis was induced in  $h^{90}$  strains by suspending a loopful of cells in 100  $\mu$ L of 0.85% NaCl and spotting on to malt extract agar (MEA) plates (43). For a cross between  $h^+$  and  $h^-$ , equal amounts of both strains were taken and resuspended in NaCl before spotting onto a MEA plate. The plate was incubated for 8 h and 15 h at room temperature for  $h^{90}$  and  $h^+h^-$  cross, respectively, to obtain zygotes. In experiments involving temperature-sensitive mutations, the cross was maintained at specific temperatures for specific times as mentioned in Results. For imaging, a toothpick full of cells was resuspended in 200  $\mu$ L of EMM-N and aspirated onto a 2 mg/mL lectin (Sigma-Aldrich, cat. no. L2380)-coated, 0.17-mm glass-bottom dish (SPL, cat. no. 100350). Cells were allowed to adhere to the glass bottom for 15–20 min. Unattached cells were washed out, and cells were imaged in EMM-N.

**Staining of Nuclei with Hoechst.** The nuclei of fission yeast zygotes were stained with Hoechst 33342 (Sigma-Aldrich, cat. no. B2261). Cells that were induced to undergo meiosis on MEA were first washed with water and then resuspended in

1–2  $\mu$ g/mL Hoechst for 20 min at room temperature. Cells were washed again after Hoechst staining and resuspended in EMM-N for imaging.

**Microscopy.** TIRF imaging was carried out on a custom-built TIRF microscope with stage body and controller from Applied Scientific Instrumentation, with an Olympus (Tokyo, Japan) UASPlanApo 100 $\times$ /1.49 N.A. TIRF objective with a correction collar. Images were captured on an Andor iXon Ultra DU-897 BV back-illuminated EMCCD. The pixel separation was 160 nm. GFP excitation was carried out with a 488-nm Sapphire laser (Coherent), and emission was filtered with a 536/40-nm bandpass filter. The system was controlled using Micro Manager 1.420 software. For counting the number of Mcp5 in each cluster on the membrane, 30 mW of the 488-nm laser excitation was used. The cells were focused in TIRF mode so as to visualize the Mcp5 foci, and a stream of 5,000 images was obtained with an exposure time of 20 ms for each frame. The intensities over each five frames of the movie were averaged to obtain a higher signal-to-noise ratio, and the resulting 1,000-frame movie was used for tracking.

Confocal microscopy was performed on an INCell Analyzer 6000 (GE Healthcare) with 60 $\times$ /0.7 N.A. objective, fitted with an sCMOS 5.5 Mp camera with an x–y pixel separation of 108 nm. In addition, fluorescence imaging was carried out using an inverted epifluorescence microscope (Olympus IX83) with stage body and controller from Applied Scientific Instrumentation and 100 $\times$ /1.4 N.A. objective. Excitation wavelength was selected through Spectra X light engine (Lumencor Inc.), and band pass filters in a high-speed filter wheel (ASI Inc.) were used for emission wavelength. The images were acquired using Evolve 512 EMCCD camera (Photometrics). The system was controlled using Slide Book6 (ver.6.0.3) software. For Hoechst, GFP, and mCherry imaging, the 405-, 488-, and 561-nm laser lines and bandpass emission filters 455/50, 525/20, and 605/52nm, respectively, were used. For obtaining the intensity profiles, fission yeast zygotes were selected, and a z-stack spanning the entire cell (9–11 slices, with a step size of 0.5  $\mu$ m) was obtained. The summed intensity projection of the stack was then used for analysis. For visualizing the nuclear oscillations in Fig. 7, a three-slice z-stack with 0.5- $\mu$ m step size was obtained for a total of 25 time frames, with a 25-s interval between each time frame.

**Image Analysis and Plotting.** Tracking of Mcp5 clusters in TIRF and intensity measurements were performed using the Low Light Tracking Tool plugin for Fiji/ImageJ (23). Bleaching step identification was performed using STEPFINDER software (24). Intensity profiles of Mcp5, PLC-PH, Myo1, and dynein were obtained using Fiji/ImageJ software. MSD analysis was performed using custom functions written in Matlab (Mathworks). All plots were created using Matlab.

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