The Polar Organizing Protein PopZ Is Fundamental for Proper Cell Division and Segregation of Cellular Content in *Magnetospirillum gryphiswaldense*

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**ABSTRACT** Magnetotactic bacteria (MTB) are of special scientific interest due to the formation of magnetosomes, intracellular membrane-enveloped magnetite crystals arranged into a linear chain by a dedicated cytoskeleton. Magnetotaxis relies on the formation and proper inheritance of these unique magnetic organelles, both of which need to be coordinated with the segregation of other cellular content such as chromosomes or motility and chemotaxis related structures. Thus, elaborated mechanisms are required in MTB to coordinate and maintain a high level of spatial and temporal subcellular organization during cytokinesis. However, thus far, underlying mechanisms and polarity determinants such as landmark proteins remained obscure in MTB. Here, we analyzed an ortholog of the polar organizing protein Z in the alphaproteobacterium *Magnetospirillum gryphiswaldense* termed PopZ Mgr. We show that deletion of the *popZmgr* gene causes abnormal cell elongation, minicell formation, DNA missegregation, and impairs motility. Overproduction of PopZ Mgr results in PopZ-rich regions near the poles, which are devoid of larger macromolecules, such as ribosomes, chromosomal DNA, and polyhydroxybutyrate (PHB) granules. Using superresolution microscopy, we show that PopZ Mgr exhibits a bipolar localization pattern throughout the cell cycle, indicating that the definition of new poles in *M. gryphiswaldense* occurs immediately upon completion of cytokinesis. Moreover, substitution of PopZ orthologs between *M. gryphiswaldense* and the related alphaproteobacterium *Caulobacter crescentus* indicated that PopZ localization depends on host-specific cues and that both orthologs have diverged to an extent that allows only partial reciprocal functional complementation. Altogether, our results indicate that in *M. gryphiswaldense*, PopZ plays a critical role during cell division and segregation of cellular content.

**IMPORTANCE** Magnetotactic bacteria (MTB) share the unique capability of magnetic navigation, one of the most complex behavioral responses found in prokaryotes, by means of magnetosomes, which act as an internal compass. Due to formation of these unique nanoparticles, MTB have emerged as a model to study prokaryotic organelle formation and cytoskeletal organization in conjunction with complex motility systems. Despite the high degree of subcellular organization required in MTB, less is known about cell-cycle-related factors or proteins responsible for spatiotemporal polarity control. Here, we investigate the function of the polar organizer PopZ in the magnetotactic alphaproteobacterium *Magnetospirillum gryphiswaldense*. Although PopZ is widely distributed among the alphaproteobacteria, its function in MTB belonging to this class has remained unexplored. Our results suggest that in *M. gryphiswaldense*, PopZ has a key role during cell division and subcellular organization. Furthermore, we show that PopZ localization and function differ from other nonmagnetotactic alphaproteobacterial model organisms.

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During cytokinesis, bacteria have to coordinate division with the equipartitioning or de novo synthesis of cellular content, such as chromosomes, intracellular storage granules, or (polar) structures, such as chemosensory clusters, flagella, or pili (1–5). In magnetotactic bacteria (MTB), this in addition has to be coordinated with the proper duplication and segregation of their key organelles, the magnetosomes (6–8). In the widely studied magnetotactic alphaproteobacterium *Magnetospirillum gryphiswaldense*, the magnetosomes consist of membrane-enclosed crystals of magnetite, which during their biosynthesis become arranged into linear chains to build a magnetic sensor sufficiently strong to align the cells in the weak Earth’s magnetic field (9, 10). During cell division, this magnetosome chain is then split and magnetosomes are equipartitioned to daughter cells (6, 7). To overcome magnetic forces during separation of magnetosome chains, cells divide by asymmetric septation (i.e., unidirectional constriction of the inner and outer membranes) (6). Both magnetosome chain formation and division are orchestrated by a multipartite and complex cytoskeleton (the “magnetoskeleton” [M. Toro-Nahuelpan, G. Giacomelli, O. Raschdorf, S. Borg, J. M. Plitzko, M. Bramkamp, D. Schüler, and F. D. Müller, submitted for publication]), involving the actin-like MamK protein, which forms dynamic filaments which position and relocate the magnetosome chain within the cell (11, 12). For proper magnetic navigation, the inherent magnetic polarity of the resulting cellular compass has to be coordinated with the organization of motility and chemotaxis structures, such as polar flagella and chemosensory clusters, resulting in a biased directionality of swimming motility (8, 13). However, despite the high degree of intra- and extracellular polar organization required in MTB, underlying cell-cycle-related factors and mechanisms have been only poorly characterized.

In other alphaproteobacteria, polar organizing protein Z (PopZ) has been found to play a major role in definition of cell polarity and spatiotemporal control of multiple proteins. In *Caulobacter crescentus*, PopZ (PopZ<sub>Cc</sub>) assembles into a putative filamentous network in chromosome-free regions at the cell poles, serving as a hub for many other cell-cycle-related proteins (14–18). During the asymmetric cell cycle of *C. crescentus*, PopZ first localizes to the old stalked pole, where it binds to the adaptor protein ParB, tethering the chromosome to the pole. In the second half of the cell cycle, the PopZ network duplicates at the opposite pole, to ensure capture of the sister chromosome upon replication, which avoids chromosome missegregation (17, 18). In addition, PopZ plays a critical role in cell division by polar retention of MipZ, which is a regulator ensuring proper midcell assembly of the FtsZ ring (19, 20). Therefore, deletion of *popZ* in *C. crescentus* causes severe cell division phenotypes, including abnormal cell elongation and formation of minicells (17, 18). Additionally, since PopZ is required for robust recruitment of proteins necessary for stalk synthesis, loss of *popZ* also affects formation of this organelle (15). In *Agrobacterium tumefaciens*, which grows predominantly by addition of peptidoglycan at one pole (the new “growth pole”) (21, 22), PopZ is located specifically at the growing pole. After septation, PopZ switches to the newly generated “growth poles” of both daughter cells (23, 24). Absence of *popZ* in *A. tumefaciens* resulted in asymmetric sites of cell constriction and cell branching (25, 26). In the pathogen *Brucella abortus*, PopZ is also unipolar and marks the new pole throughout the cell cycle (27).

Although PopZ is widespread among the alphaproteobacteria (28), its function has been studied in only a very few representatives, and its role in MTB from this class has remained entirely unexplored. Here, we characterized an ortholog of the PopZ polarity factor in *M. gryphiswaldense* (referred to as PopZ<sub>Mgr</sub>). Loss of *popZ<sub>Mgr</sub>* caused severe defects of growth, cell division, and motility. In contrast, the dynamic MamK pole-to-midcell treadmilling was independent of PopZ<sub>Mgr</sub>. Moreover, we observed remarkable differences in the cell-cycle-dependent localization pattern of PopZ between *M. gryphiswaldense*, *C. crescentus*, and *A. tumefaciens*. Substitution of PopZ orthologs between *M. gryphiswaldense* and *C. crescentus* indicated that the proteins from both strains can partially replace their respective functionalities, but differ to an extent that prevents full...
implementation within the cell-cycle-dependent interaction network of the heterologous host. In summary, our results reveal a key role of PopZ\textsubscript{Mgr} in subcellular organization and provide the first fundamental insights into its function in cell cycle control and polarity determination in MTB. Thus, this work also demonstrates the usefulness of \textit{M. gryphiswaldense} as a potential and emerging model to scrutinize the bacterial cell cycle and its coordination to spatiotemporal organelle organization.

**RESULTS**

\textbf{PopZ\textsubscript{Mgr} localizes to both cell poles.} In \textit{M. gryphiswaldense}, PopZ\textsubscript{Mgr} is encoded in a conserved genomic region, similar to \textit{C. crescentus}, next to putative genes coding for a valyl-tRNA synthetase and an outer membrane efflux protein (see Fig. S1A in the supplemental material). To study its localization pattern throughout the cell cycle, PopZ\textsubscript{Mgr} was translationally fused to green fluorescent protein (GFP) by integration of an \textit{M. gryphiswaldense} codon-optimized \textit{gfp} gene (mage\textit{gfp} [29]) within this genomic region. Expression of popZ\textsubscript{Mgr}\textit{gfp} from its native promoter was verified via immunoblotting using an antibody against GFP (not shown). \textit{In vivo} time-lapse fluorescence microscopy revealed that PopZ\textsubscript{Mgr}-GFP localized to both cell poles and exhibited a bipolar localization pattern throughout the cell cycle (Fig. 1A; see Movie S1 in the supplemental material). PopZ foci at the future new poles appeared at the end of the cell cycle (Fig. 1B). Since conventional wide-field microscopy did not allow us to judge with high confidence if cells with PopZ foci present at the division plane had already completed cytokinesis, we imaged dividing cells with superresolution three-dimensional structured illumination microscopy (3D-SIM). Using FM4-64 membrane staining, 3D-SIM revealed two adjacent PopZ foci (~250 nm apart) at the cell division site (Fig. 1Ci; see Fig. S2 in the supplemental material). In general, all cells with two PopZ foci present at the division site had already completed separation of their membranes (Fig. 1Ci and Fig. S2). In contrast, no PopZ foci were observed in cells with membranes and cytoplasm still connected, but which already had undergone partial membrane constriction (Fig. 1Cii and Fig. S2). These results indicated that formation of PopZ-rich zones at the new poles occurs very late during or shortly after completion of cytokinesis.

\textbf{Deletion of popZ\textsubscript{Mgr} causes severe cell division defects.} To study the effects of popZ absence in \textit{M. gryphiswaldense}, a markerless in-frame deletion mutant was constructed. The ΔpopZ\textsubscript{Mgr} strain was viable, but showed severely impaired growth and increased cell length (Fig. 2). Some cells were elongated up to 60 μm (Fig. 2Ai), equivalent to ~20-fold the length of a newborn wild-type cell (~3 μm). Elongated cells contained between 1 and 3 abnormally long magnetosome chains running in parallel, which were sometimes interspaced by segments of unknown origin and composition—i.e., parts of the cell body that appeared brighter in the electron microscope (Fig. 2Aii, black arrowheads). Elongated magnetosome chains were up to ~20 μm in length (allowing gaps not larger than 150 nm). Elongated cells with more than 700 particles were observed; however, cell length and number of magnetosomes were well correlated, resulting in ~12 particles/μm (Fig. 2B). For comparison, we determined that the wild type typically exhibited a median chain length of ~1 μm and median magnetosome numbers of 35 particles per cell. In contrast to the ΔpopZ\textsubscript{Mgr} strain, particle number and cell length were only poorly correlated in the wild type (Fig. 2B, inset), similar to previous correlative estimations of magnetosome particle numbers versus cell area (30). Moreover, even in the most highly elongated cells, magnetosome chains were mostly absent from the regions near the cell poles (Fig. 2Aiii). As is also commonly observed in the wild type (31), cells of the ΔpopZ\textsubscript{Mgr} strain contained large amounts of polyphosphate and polyhydroxybutyrate (PHB) granules (Fig. 2Ai to Aiii).

During time-lapse microscopy (Fig. 2C), the wild type divided at a median cell length of 3.3 μm, whereas in the ΔpopZ\textsubscript{Mgr} strain, the cell elongation and unequal division produced a much broader length distribution with a median cell length of 3.9 μm during division (Fig. 2D). Furthermore, the time required to complete a division cycle was less regular for the ΔpopZ\textsubscript{Mgr} strain (Fig. 2C and Movie S1). While wild-type cells divide approximately every 4 to 5 h, cell division in the ΔpopZ\textsubscript{Mgr} strain occurred with
variable timing and at ectopic positions, suggesting that the generation time as well as division septum positioning were affected. To measure potential asymmetry during cell division, the difference in length of both newborn daughter cells was calculated from time-lapse series ([Fig. 2E](#fig2e)). Thus, the median cell length differences for wild-type daughter cells were 0.29 and 0.94 μm for the ΔpopZMgr deletion strain, confirming that the ΔpopZMgr population contained cells with high variation in length caused by unequal division. Also, a slight asymmetry (~11% off-center) was detected for newborn daughter cells of the wild-type strain, in agreement with previous observations (on average, 15% off-center) by Katzmann et al. ([6](#ref6)).

**CET ultrastructural analysis of the ΔpopZMgr mutant reveals missegregation of cellular content and chemotactic receptor arrays, septum mislocalization, and minicell formation.** Prompted by the observed cell elongation and impairment of division, we performed cryo-electron tomography (CET) to further investigate the cell

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**FIG 1** Localization of PopZMgr-GFP in *M. gryphiswaldense*. (A) Time-lapse microscopy of cells expressing PopZMgr-GFP (popZ::popZ-gfp strain). First row, bright-field; second row, fluorescence channel; third row, overlay of bright-field and GFP channel. PopZMgr localizes to both cell poles. In dividing cells, an additional spot appears at the cell division site (fourth and ninth frames, white arrowheads). Generation time during time-lapse series was approximately 4 to 5 h. Numbers indicate hours and minutes. (B) Demograph of cells expressing PopZMgr-GFP (*n* = 642 cells). The appearance of the signal at midcell is marked with an arrowhead. (C) Structured illumination microscopy (3D-SIM) allows us to resolve two PopZ foci in close proximity at the division plane with a distance near the resolution limit of conventional epifluorescence microscopy (~250 nm). Micrographs are maximum-intensity projections of z-stack images from representative FM4-64-stained dividing cells. First row, bright-field (left image) and GFP channel (right image); second row, FM4-64 channel (left image) and overlay of FM4-64 and GFP channel (right image). Insets are magnified xy, xz, and yz projections of PopZ foci (GFP channel) and division plane (FM4-64 channel). Note cells with PopZ foci present at the site of division (Ci) had already completed compartmentalization and separation of their membranes, whereas no foci were observed in cells that were still connected (Cii). A representative number of dividing cells was imaged (*n* = 29 [additional cells are shown in Fig. S2]). All scale bars not indicated in the figure correspond to 3 μm.
Deletion of popZ in *M. gryphiswaldense* causes severe cell division defects. (A) Upon deletion of popZ::gfp, highly elongated cells (up to 60 μm) were observed, which contained long, sometimes interrupted (black arrowheads) magnetosome chains (M) traversing throughout the cell, except polar regions (Ai to Aiii). Bipolar positioning of flagella (F) was not affected in the ΔpopZ::gfp strain (Aii and Aiv), but occasionally additional flagella were observed at ectopic positions along the cell’s body (Ai and Aii). Scale bars correspond to 1 μm (Aii and Aiv) and 3 μm (Ai and Aiii). (B) A linear correlation between cell length and number of magnetosomes was revealed for the ΔpopZ::gfp strain, including highly elongated cells, with single chains consisting of more than 700 particles. Dotted lines indicate 95% confidence intervals (big dots) and prediction intervals (small dots). A total number of 47 cells for the ΔpopZ::gfp strain and 54 cells for the wild type (WT [inset]) were analyzed from at least 3 independent cultures grown under standard microoxic conditions. (C) Time-lapse microscopy of the *M. gryphiswaldense* wild-type and ΔpopZ::gfp strains. Division of the ΔpopZ::gfp cell occurs at ectopic positions. Cell division events are marked with white arrowheads. Time values are given in hours and minutes. Scale bar = 3 μm. (D) Cell length distributions and (E) difference in length of both newborn daughter cells (Continued on next page)
division defect of the ΔpopZ<sub>Mgr</sub> strain. A cryo-electron micrograph of an elongated ΔpopZ<sub>Mgr</sub> cell (Fig. 3A) indicates the areas analyzed by CET (cell pole and cell body) and confirms the formation of distinct minicells at the poles that were also observed in time-lapse imaging experiments (Fig. 2C and Movie S1). CET and further segmentation of the cell pole showed the typical arrangement of magnetosomes by the actin-like MamK filament (Fig. 3Bi and Bii). In addition, a chemoreceptor array was observed at the pole close to the lateral cytoplasmic membrane (inset from Fig. 3Bi and purple objects in Bii), as observed previously in wild-type cells (11). Additionally, a tomographic slice of the cell pole revealed the presence of chemoreceptor arrays also within the minicell (Fig. 3Bi and vi), highlighted in the tomogram segmentation (purple objects in Fig. 3Bii and Biii). Moreover, additional structures inside the minicell, such as magnetosome membrane vesicles, MamK filaments, and ribosomes, were visible (Fig. 3Biv to vi), indicating putative defects in segregation of cellular content. The presence of chemoreceptor arrays at both the pole and within the adjacent minicell (Fig. 3B), and the observation of double chemoreceptor arrays at the cell poles (see Fig. S3Ai in the supplemental material) argue for an improper polar chemoreceptor array localization (CET micrographs of chemoreceptor arrays in the wild-type strain can be found in reference 11 for comparison).

Localization of magnetosome chains did not seem to be affected in the ΔpopZ<sub>Mgr</sub> strain, as tomographic slices of the cell body showed a properly localized magnetosome chain across the long cell axis (Fig. 3Ci). Remarkably, a deep unidirectional constriction of the membrane located distant from midcell indicated a putative septum mispositioning (black-white arrowheads in Fig. 3A and Cii and white arrowheads in Fig. 3Cii and Civ). Furthermore, both the misplaced septum invaginations far-off midcell (Fig. S3Ai and Av and Bi, Bii, and Cii) and minicell formation (Fig. S3Bii to Biii) were commonly observed by CET. Therefore, minicell formation is likely caused by ectopic septum localization, confirming the cell division impairment in the ΔpopZ<sub>Mgr</sub> mutant. Thus, ΔpopZ<sub>Mgr</sub> cells are likely unable to properly control the FtsZ ring localization.

Since one of the major functions of PopZ in both C. crescentus and A. tumefaciens is also in regulation of chromosome segregation (17, 18, 24), we quantified DNA content of ΔpopZ<sub>Mgr</sub> minicells by staining with DAPI (4',6-diamidino-2-phenylindole), a dye specific for DNA (Fig. 4A). Remarkably, minicells had an approximately 2.8-fold or 1.5-fold reduced mean cell fluorescence (Fig. 4B) compared to either other cells or their polar regions (to account for possible volume differences of minicells, since cells get thinner toward the poles), further suggesting that PopZ in M. gryphiswaldense also contributes to proper chromosome segregation.

**Deletion of popZ<sub>Mgr</sub> impairs motility and magneto-aerotaxis.** In M. gryphiswaldense, magnetotaxis is tightly coupled to aerotaxis in order to govern directed swimming toward optimal low-oxygen levels (13, 32). Remarkably, it was also found that cells that perform polar magneto-aerotaxis and display a distinct swimming polarity bias within the magnetic field (preferentially north or south seeking) can be enriched within only few generations (13). Thus, during cytokinesis the proper segregation of magnetosome chains with an inherent physically imprinted magnetic polarity must be coordinated with the determination of the magnetotactic swimming direction. In order to analyze if the ΔpopZ<sub>Mgr</sub> strain is affected with respect to motility and magneto-aerotaxis, we performed tracking microscopy (Fig. 5A) and various soft-agar-based assays (Fig. 5B to D). Most cells of the ΔpopZ<sub>Mgr</sub> strain contained bipolar flagella as in
FIG 3  Cryo-electron tomography of the ΔpopZ<sub>Gm</sub> strain. (A) Cryo-electron micrograph of an elongated cell of the ΔpopZ<sub>Gm</sub> strain. Tomograms were acquired at the cell pole and cell body areas. Combined black and white arrowheads indicate the position of membrane invagination observed in panel Cii. Scale bar = 1 μm. (B) CET of the cell pole area. (Bi) A 15.7-nm-thick tomographic slice (average of 5 slices) through the central part of the cell. The black dashed rectangle indicates the area seen in the inset. (Inset) Base plate layer of a chemoreceptor array indicated by a black double arrowhead. (Bii and Biii) Three-dimensional rendering of the cell pole shown in panels A and Bi. Magnetite crystals are red, magnetosome membrane vesicles are yellow, the actin-like MamK filament is green, and the chemoreceptor arrays are purple. The cellular envelope inner and outer membranes are depicted in blue. (Biv) A 15.7-nm-thick tomographic slice of the minicell displaying magnetosome vesicles (black arrows), (Bv) the MamK filament (3.1-nm-thick slice [black arrowheads]), and (Bvi) the chemoreceptor array (red arrowhead, periplasmic chemoreceptor domains; black double arrowhead, chemoreceptor base plate layer). (C) CET of the cell body area. (Ci) A single 3.1-nm-thick tomographic slice through the central part of the cell displaying the magnetosome chain (electron-dense magnetite crystals arranged into a chain), (Cii) A 3.1-nm-thick slice near the cell edge showing a deep membrane unidirectional constriction at the cell body area. (Inset) A 3.1-nm slice in a different z position through the tomogram showing the continuity of the magnetosome chain from panel Ci. (Ciii and Civ) Three-dimensional rendering of the cell body area shown in panels A and Ci. White arrowheads show membrane invagination seen in panel Cii. Scale bars: panels Bi, Ci, and Cii = 200 nm; panels Bv to Bvi and insets = 100 nm. A total of 6 cells were analyzed.
the wild type (Fig. 2Aii and 2iv), but occasionally additional flagella were observed along the cell body (Fig. 2Aii). Elongated cells of the popZMgr deletion strain were motile and despite their highly increased length still aligned to an external magnetic field when observed by dark-field microscopy (Fig. 5Ai; see Movie S3 in the supplemental material) or optical measurement of their magnetic response ($C_{mag}$). In general, the $\Delta$popZMgr population was heterogeneous, consisting of smaller fast-swimming cells and more elongated cells that moved at a lower speed than the wild type (Movie S3 [movies of the wild-type strain can be found in reference 13]). Shorter cells, which had higher swimming speeds and traveled longer distances within the time frame of observation, were observed to swim in circular motions and were not well aligned within the magnetic field (Fig. 5Ai and Movie S3), which might be due to their altered cell length or the existence of no or only short magnetosome chains. In contrast, longer cells, which displayed low swimming speeds and traveled only short distances, were very well aligned within the magnetic field, presumably due to their overly elongated magnetosome chains. In summary, due to the higher number of highly motile short cells, the mean overall alignment of the $\Delta$popZMgr population was reduced compared to that of the wild type (Fig. 5Aiii), whereas the swimming speed distribution did not differ significantly from the wild type (Fig. 5Aiv). Accordingly, $\Delta$popZMgr cultures also reproducibly displayed a slightly lower $C_{mag}$ than the wild type (as determined from at least triplicate cultures [values are given in the legend to Fig. 5Aiii and 5Aiv]). In contrast to the wild type, the formation of aerotactic swim halos in semisolid medium was almost completely abolished (Fig. 5B). Moreover, in comparison to the wild type, which forms sharp aerotactic bands in soft agar tubes, the $\Delta$popZMgr mutant grew only in a diffuse zone close to the surface (Fig. 5C). When soft agar assays were performed in the presence of a magnetic field (Fig. 5D; see Fig. S4C in the supplemental material), spreading of the popZMgr deletion mutant parallel to the magnetic field was only observed after prolonged incubation (>4 days), confirming that cells are still able to align and distribute along the magnetic field lines, but in a slow and possibly only growth-dependent manner. We also failed to restore a swimming polarity bias in elongated $\Delta$popZMgr cells by magnetic selection. Whereas the wild-type and popZMgr::popZMgr-gfp strains displayed a south-seeking polarity bias upon repeated passaging in $O_2$ gradients within a superimposed vertical Southern Hemisphere-like magnetic field, $\Delta$popZMgr cells were rather equally distributed toward both magnetic poles (Fig. 5D and Fig. S4C). These findings were confirmed by the hanging drop assay (not shown). In general, only few cells of the $\Delta$popZMgr strain accumulated in equal proportions at the northern and southern magnetic pole (facing the air-adjacent borders of the drop), speaking for a general impairment of aerotaxis. In summary, deletion of popZMgr severely impaired motility and aerotaxis. Since we observed an increased tendency of highly elongated $\Delta$popZMgr cells to intertwine and aggregate (Fig. 5E), reduced motility in soft agar is also partially explained by the formation of cell clumps.
MamK filament dynamics is independent of PopZ<sub>Mgr</sub>. In *M. gryphiswaldense*, magnetosome chains are recruited to midcell to ensure equal partitioning of magnetosomes to both daughter cells. It has been recently found that the MamK filament has a particular dynamic behavior, growing from both cell poles, elongating toward midcell, and undergoing treadmilling (7).
Therefore, based on the PopZ
Mgr localization pattern, we asked whether PopZ
Mgr is involved in or influences the MamK dynamics. To examine this hypothesis, we performed photokinetic analysis of the MamK filament in Δ
popZ
Mgr cells. Fluorescence recovery after photobleaching (FRAP) of MamK filaments using an mCherry-MamK fusion showed a half-time fluorescence recovery ($t_{1/2}$) of 87.6 ± 17.9 s (Fig. 6A). Recently, it was reported that the mCherry-MamK translational fusion expressed in M. gryphiswaldense wild-type cells from a plasmid and chromosomally showed $t_{1/2}$ values of 71.8 ± 6.6 and 68.3 ± 4.8 s, respectively (7). A one-way analysis of variance (ANOVA) followed by a Tukey’s multiple-comparison test determined that the mCherry-MamK filament $t_{1/2}$ in the absence of popZ
Mgr is statistically not significant compared to the
previously reported values for the wild-type strain \( (P < 0.05) \). Furthermore, the MamK pole-to-midcell growth and its treadmilling behavior are not affected upon absence of \( \text{popZ}_{\text{Mgr}} \) (Fig. 6B). Thus, it can be concluded that the MamK filament dynamics, especially the directed pole-to-midcell growth, is independent of \( \text{PopZ}_{\text{Mgr}} \).

**PopZ\(_{\text{Mgr}}\) forms a polar exclusion zone devoid of macromolecules and chromosomal DNA.** To study the effect of \( \text{popZ}_{\text{Mgr}} \) overexpression, \( \text{PopZ}_{\text{Mgr}} \) and \( \text{PopZ}_{\text{Mgr}}\text{-GFP} \) were overproduced in \( \text{trans} \) under the control of \( \text{P}_{\text{rec}} \) (anhydrotetracycline-inducible promoter) in the \( M. \text{gryphiswaldense} \) wild type and \( \Delta \text{popZ}_{\text{Mgr}} \) strain. Upon reintroduction of \( \text{popZ}_{\text{Mgr}} \) (or \( \text{popZ}_{\text{Mgr}}\text{-gfp} \)), cell morphology of the wild type, formation of swim halos, and growth were restored in the \( \Delta \text{popZ}_{\text{Mgr}} \) strain (see Fig. S5 in the supplemental material). Furthermore, prolonged overexpression of \( \text{popZ}_{\text{Mgr}} \) or \( \text{popZ}_{\text{Mgr}}\text{-gfp} \) in the wild-type strain caused severe cell filamentation and delayed growth, whereas expression in the \( \Delta \text{popZ}_{\text{Mgr}} \) strain had a lesser effect on growth and cell length, likely due to the absence of endogenous \( \text{PopZ}_{\text{Mgr}} \) (Fig. 6D and Fig. 5S). Moreover, \( \text{PopZ}_{\text{Mgr}}\text{-GFP} \) overproduction in the \( M. \text{gryphiswaldense} \) wild type caused either (i) cells with two large polar foci and multiple smaller foci distributed across the cell and in between the PHB granules (stained with the lipophilic dye Nile red, specific for membranes and polyhydroyxbutyrate [PHB]) (Fig. 6C, inset) or (ii) cells with a large \( \text{PopZ} \) accumulation cluster expanding from one pole (Fig. 6C). In the latter, some cells had an additional smaller \( \text{PopZ}_{\text{Mgr}}\text{-GFP} \) cluster at the opposite pole. The \( \text{PopZ} \) expansion zone encompassed several micrometers in length and presented a reduced cell diameter, resulting in a tail-like appearance. Additional staining with DAPI and Nile red revealed that chromosomal DNA and PHB granules were excluded from the expanded \( \text{PopZ} \) area. Transmission electron microscopy (TEM) analysis confirmed that this zone was depleted of larger cytoplasmic structures such as PHB or polyphosphate granules (Fig. 6D). Furthermore, the brighter appearance indicated that the putative polar \( \text{PopZ} \)-rich region is mostly devoid of electron-dense cytoplasmic structures and macromolecules (e.g., ribosomes). Even upon \( \text{PopZ}_{\text{Mgr}} \) overexpression, magnetosome chains were still located at midcell, resembling the wild-type phenotype, but in a few cases were also embedded into the outermost part of the \( \text{PopZ} \) expansion zone. Of note, the formation of flagella at the \( \text{PopZ} \)-rich poles was not impaired (Fig. 6D). In summary, \( \text{PopZ}_{\text{Mgr}} \) forms a polar expansion zone that is depleted in larger macromolecules and organelles, similar to previously reported observations regarding \( \text{PopZ} \) in \( C. \text{crescentus} \) (15, 17).

**Bipolar \( \text{PopZ}_{\text{Mgr}} \) localization requires host-specific factors.** In \( C. \text{crescentus} \), \( \text{PopZ} \) (abbreviated \( \text{PopZ}_{\text{Cc}} \)) first localizes to the old pole and undergoes a transition from monopolar to bipolar after completion of cell division (17). The \( C. \text{crescentus} \) life cycle is highly asymmetric, generating a smaller and motile swarmer cell and a stalked cell that possesses a tubular extension at the old pole, required for surface attachment. The distinct bipolar localization pattern in \( M. \text{gryphiswaldense} \) (Fig. 1) prompted us to investigate the localization pattern of \( \text{PopZ}_{\text{Mgr}} \) in \( C. \text{crescentus} \) (Fig. 7; see Fig. S6A in the supplemental material). When \( \text{PopZ}_{\text{Mgr}}\text{-GFP} \) was heterologously produced in \( C. \text{crescentus} \) NA1000 (in \( \text{trans} \) expressed from \( \text{P}_{\text{tet}} \), in the presence of endogenous \( \text{PopZ}_{\text{Cc}} \)), a unipolar-to-bipolar transition pattern was revealed (Fig. 7A, 1-h time-point and Fig. S6A), similar to the localization pattern of \( \text{PopZ}_{\text{Cc}} \). \( \text{PopZ}_{\text{Mgr}} \) and \( \text{PopZ}_{\text{Cc}} \) are conserved in their N- and C-terminal regions (37.2% identity and 51.3% similarity, in a global alignment, including some of the most related orthologs [Fig. S1B to D]), which are known to be important in \( C. \text{crescentus} \) for interaction with the ParA/ParB chromosome segregation machinery and \( \text{PopZ} \) cluster formation, respectively. Thus, the observed localization pattern of \( \text{PopZ}_{\text{Mgr}} \) in \( C. \text{crescentus} \) might be explained by a direct interaction between \( \text{PopZ}_{\text{Mgr}} \) and \( \text{PopZ}_{\text{Cc}} \) and/or with other known \( \text{PopZ} \) interactors present in \( C. \text{crescentus} \), such as ParA/ParB. Upon prolonged expression of \( \text{popZ}_{\text{Mgr}}\text{-gfp} \) in \( C. \text{crescentus} \), cells became heavily elongated and aberrantly shaped (Fig. 7A and B), indicating that overexpression of \( \text{PopZ}_{\text{Mgr}} \) also interferes with cell division in \( C. \text{crescentus} \). In addition, heterologous \( \text{popZ}_{\text{Mgr}}\text{-gfp} \) overproduction in \( C. \text{crescentus} \) resulted in the appearance of multiple \( \text{PopZ} \) foci and large polar \( \text{PopZ} \) exclusion zones (Fig. 7A),
FIG 7 PopZMgr and PopZCc are able to partially replace their reciprocal functions. (A) Heterologous expression of popZMgr-gfp in trans from _P_. _lim._ in the presence (C. _crescentus_ NA1000 wild type) and absence (ΔpopZCc _Cc_ strain) of endogenous popZCc. Numbers indicate hours and minutes postinduction with 50 ng/ml anhydrotetracycline during growth in PYE (peptone-yeast extract) medium. Overproduction of PopZMgr-GFP in NA1000 (5- to 24-h time points) leads to formation of polar PopZ-rich regions and cell filamentation, whereas expression in the ΔpopZCc strain causes shortening of cells. The first row shows the fluorescence channel, and the second row shows an overlay image of DIC and the fluorescence channel. (B) Cell length distributions at selected time points for the two strains shown in panel A. (C) C. _crescentus_ NA1000 cells expressing popZMgr-gfp (23 h postinduction with 50 ng/ml anhydrotetracycline). Since only a few PHB granules were formed in _C. _crescentus_ during cultivation in rich PYE medium, the cells shown were cultivated in HIGG (Hutner base–imidazole-buffered–glucose–glutamate)minimal medium with limiting amounts of phosphate (0.01 mM), which did induce formation of large amounts of PHB storage granules in addition to stalk elongation. Note PopZMgr forms large polar exclusion zones devoid of DNA and PHB granules. Furthermore, overexpression of PopZMgr-GFP did cause cell elongation and affected stalk formation (cells with bipolar stalks [yellow arrowheads]). Cells were stained with DAPI and Nile red. (Di) Localization of PopZCc-GFP in _C. _crescentus_ when expressed as the sole source of PopZ from the endogenous popZCc promoter (popZCc::popZCc-gfp strain). Cells were grown in PYE medium. (Dii) Localization of mCherry-PopZCc in _M. _gryphiswaldense_ when expressed as sole source of PopZ from the endogenous popZCc promoter (popZCc::mCherry-popZCc _Cc_ strain). Cells were grown in flask standard medium (FSM). (E) Time-lapse microscopy of strains shown in panel D. (Continued on next page)
similar to the previously described observations regarding the overproduction of native PopZ<sub>cc</sub> in <i>C. crescentus</i> (15, 17). As observed for <i>M. gryphiswaldense</i> (Fig. 6C and D), PopZ<sub>Mgr</sub> exclusion zones in <i>C. crescentus</i> were devoid of DNA and PHB storage granules (Fig. 7C). Heterologous expression of popZ<sub>Mgr</sub>-gfp in the ΔpopZ<sub>cc</sub> background partially restored the cellular morphology of the <i>C. crescentus</i> wild-type strain (Fig. 7A), resulting in a reduced cell length close to wild-type-like levels (Fig. 7B). However, in some cells, we observed PopZ<sub>Mgr</sub>-GFP foci located at ectopic positions opposite to the stalked pole (Fig. 7A, 12-h time point, yellow arrowheads), indicating that the absence of PopZ<sub>cc</sub> was not fully transcomplemented by PopZ<sub>Mgr</sub>-GFP.

In order to avoid artifacts caused by altered expression levels (due to expression from a random ectopic locus under the control of P<sub>ter</sub> or the presence of endogenous PopZ<sub>cc</sub>), we constructed a <i>C. crescentus</i> strain harboring a site-specific chromosomal replacement of popZ<sub>cc</sub> against popZ<sub>Mgr</sub>-gfp. We also performed the reciprocal experiment and constructed an <i>M. gryphiswaldense</i> strain that carries a site-specific chromosomal replacement of popZ<sub>Mgr</sub> for an mCherry-popZ<sub>cc</sub> fusion. Notably, PopZ<sub>Mgr</sub>-GFP localized in a monopolar-to-bipolar fashion in <i>C. crescentus</i> (Fig. 7Di and Ei), when expressed from the endogenous popZ<sub>cc</sub>-promoter as the sole source of PopZ present. However, transition of PopZ<sub>Mgr</sub>-GFP to the new poles occurred with ectopic timings, and not all PopZ<sub>Mgr</sub> was retained in polar regions, as indicated by the diffuse cytoplasmic fluorescence signal (Fig. 7Di and Ei). In <i>M. gryphiswaldense</i>, mCherry-PopZ<sub>cc</sub> in general did localize to both cell poles (Fig. 7Di and Ei). However, one cell pole displayed much stronger fluorescence, and filamentous localization patterns were observed (Fig. 7Dii, elongated cell at the bottom left). Furthermore, the appearance of PopZ foci at the new cell poles was delayed, and disappearance of polar foci was observed in some cases (Fig. 7Eii). Together, these results indicated that bipolar PopZ localization is regulated by host-specific proteins in <i>M. gryphiswaldense</i> and that both PopZ orthologs differ to an extent that does not allow full functionality within the cell cycle of the heterologous host. The findings that bipolar PopZ subcellular localization is not inherent to the protein itself but rather host specific were further corroborated by expression of PopZ<sub>Mgr</sub>-GFP in various other bacterial hosts (as shown in Fig. S6B to E), including only distantly related <i>Escherichia coli</i> and the two other alphaproteobacteria Rhodobacter sphaeroides and Rhodospirillum rubrum, which contain no PopZ (R. <i>sphaeroides</i>) or an endogenous PopZ ortholog (R. <i>rubrum</i> [49% identical to PopZ<sub>Mgr</sub>]) (28). Expression of PopZ<sub>Mgr</sub>-GFP in <i>E. coli</i> WM3064 resulted in the formation of large fluorescent clusters in polar nucleoid-free regions, exhibiting unipolar localization at either the new or old pole (Fig. S6B), similar to the observations made upon expression of PopZ<sub>cc</sub> in <i>E. coli</i> (17, 18). When we studied the localization of PopZ<sub>Mgr</sub>-GFP in spheroid 2,6-diaminopimelic acid (DAP)-auxotrophic WM3064 cells formed after depletion of DAP, random foci were formed close to the cell periphery (Fig. S6C), indicating that localization of PopZ<sub>Mgr</sub> does not depend on geometrical constraints. In ovoid rod-shaped <i>R. sphaeroides</i> cells, PopZ<sub>Mgr</sub> generally localized only at the old pole (Fig. S6D). A similar bipolar pattern to that in <i>M. gryphiswaldense</i> was observed in

**FIG 7 Legend (Continued)**

Numbers indicate hours and minutes. Note both PopZ orthologs display delayed or erratic polar localization when expressed in the opposing parent strain. (F) Analysis of cell length of <i>C. crescentus</i> (Fi) and <i>M. gryphiswaldense</i> (Fii) strains harboring a chromosomal site-specific replacement of their PopZ ortholog compared to the respective wild-type and ΔpopZ strains. <i>C. crescentus</i> strains were grown in PYE to an OD<sub>600</sub> of ~0.17. <i>M. gryphiswaldense</i> strains were grown under microaerobic conditions to an OD<sub>600</sub> of ~0.2 in FSM medium. Note both PopZ orthologs are capable to partially replace their reciprocal functions with respect to the cell-length phenotype compared to strains harboring popZ deletions. (G) The <i>C. crescentus</i> wild-type NA1000, ΔpopZ<sub>cc</sub>, and popZ<sub>cc</sub>:popZ<sub>Mgr</sub>-gfp strains were grown in HIGG minimal medium containing 0.1 mM phosphate to an OD<sub>600</sub> of ~1.5 for the analysis of stalks. (Gi) Epifluorescence micrographs of all three strains stained with FM4-64 and DAPI. Note since FM4-64 did not reliably stain stalks of all cells, DIC microscopy and DAPI staining (which unspecifically stains stalks) were included to identify and measure stalks. (Gii and Giii) Analysis of (Gii) stalk length and (Giii) stalk number (given as percentage of cells with no stalks or mono- or bipolar stalks). Note PopZ<sub>Mgr</sub> is able to partially complement the function of PopZ<sub>cc</sub> with respect to proper stalk formation. Measurements shown in panel B were taken from one representative induction experiment. Experiments shown in panels C to E and G are maximum-intensity projections of deconvolved z-stacks. Fluorescence channels are indicated in the graph. Representative stalks and cell division events are exemplarily marked with yellow and white arrowheads, respectively. All scale bars = 3 μm.
spirillum-shaped *R. rubrum* cells, with two new PopZ foci emerging at the site of cell division (Fig. S6E). In summary, these results suggest that monopolar accumulation in DNA-free polar regions occurs by a mechanism that is inherent to PopZ$_{Cc}$, whereas bipolar localization apparently depends on distinct alphaproteobacterium-specific host factors.

To further investigate whether PopZ orthologs can replace their functionalities, we compared median cell lengths of *M. gryphiswaldense* and *C. crescentus* strains harboring reciprocal PopZ orthologs with the respective wild-type and ΔpopZ strains. The median cell length of the *C. crescentus* popZ$_{Cc}$::popZ$_{Mgr}$::gfp strain was 1.5-fold reduced compared to the ΔpopZ$_{Cc}$ strain (and 1.2-fold higher than that of the NA1000 wild-type strain [Fig. 7Fii]), whereas the median cell length of the *M. gryphiswaldense* popZ$_{Mgr}$::mCherry-popZ$_{Cc}$ strain was 1.4-fold lower than that of the ΔpopZ$_{Mgr}$ strain (but only 1.05-fold larger than that of the *M. gryphiswaldense* wild-type strain [Fig. 7Fii]). These results indicated that in both strains, the loss of the respective PopZ ortholog can be partially rescued by expression of the reciprocal ortholog.

To analyze whether PopZ$_{Mgr}$ is capable to accomplish functions with respect to stalk formation, which is impaired in the *C. crescentus* popZ deletion strain (17, 18), we also analyzed stalk length and frequency in cells grown under phosphate-limiting conditions, known to cause severe stalk elongation (33), to facilitate detection and analysis of stalks (Fig. 7G). In contrast to previous reports that the ΔpopZ$_{Cc}$ strain does not form stalks (3, 21), we found that the ΔpopZ$_{Cc}$ strain grown under phosphate starvation is still able to form stalks, but of 2.5-fold-reduced median length (Fig. 7Gi) and at a lower frequency (Fig. 7Giii) than the NA1000 wild-type strain. In comparison, the fraction of cells with monopolar stalks was 1.6-fold lower in the ΔpopZ$_{Cc}$ mutant than in the NA1000 wild-type strain. In contrast, the fraction of cells without stalks was 2.3-fold reduced in the popZ$_{Cc}$::popZ$_{Mgr}$::gfp strain compared to the ΔpopZ$_{Cc}$ strain, whereas the number of cells with monopolar stalks was 1.9-fold higher, bringing both values closer to wild-type levels. A minor fraction of cells with bipolar stalks was detected within the ΔpopZ$_{Cc}$ and popZ$_{Cc}$::popZ$_{Mgr}$::gfp populations, whereas no cells containing bipolar stalks were found in the NA1000 wild-type strain. Median stalk lengths of the popZ$_{Cc}$::popZ$_{Mgr}$::gfp strain were still 1.4-fold lower relative to the wild type but 1.8-fold larger in comparison to the ΔpopZ$_{Cc}$ strain. Notably, stalk formation was also restored upon expression of popZ$_{Mgr}$::gfp in the ΔpopZ$_{Cc}$ strain from P$_{tet}$ (Fig. 7A, 8 and 12 h, yellow arrowheads), and overexpression of PopZ$_{Mgr}$::GFP in NA1000 caused aberrant bipolar stalk formation (Fig. 7C, yellow arrowheads). In summary, these results argue that PopZ$_{Mgr}$ is able to partially accomplish functions inherent to PopZ$_{Cc}$ with respect to stalk formation in *C. crescentus*.

**DISCUSSION**

In *C. crescentus*, PopZ has been described as an important landmark protein, generating a polar hub domain for multiple proteins involved in cell cycle control and polar morphogenesis (14–18, 34, 35). In addition to *C. crescentus*, PopZ has been studied in *A. tumefaciens* (23–26), which exhibits unipolar growth by addition of peptidoglycan at the new "growth pole" (22). Here, we report that PopZ in the magnetotactic model organism *M. gryphiswaldense* plays a similar, but somewhat distinct role. In contrast to *C. crescentus* and *A. tumefaciens*, where cell division results in morphologically distinct cells and/or daughter cells that differ in cell cycle progression, division in *M. gryphiswaldense* gives rise to morphologically nearly equal daughter cells. Deletion and overexpression of popZ in *M. gryphiswaldense* resulted in severe cell division defects (Fig. 2, Fig. 3, Fig. S3, Movie S1, Movie S2, and Fig. 6C and D and Fig. S5A and B, respectively) and DNA missegregation (Fig. 4), consistent with previous observations in *C. crescentus* (17, 18) and *A. tumefaciens* (24, 25). However, we did not observe formation of ectopic poles and cell branching as in *A. tumefaciens* (25, 26). In accordance with reported results in *C. crescentus* (15, 17), we have observed formation of large exclusion zones upon overproduction of PopZ$_{Mgr}$ in *M. gryphiswaldense* (Fig. 6C and D). These results imply that PopZ$_{Mgr}$ may have an important role as a putative...
landmark protein and in the control of cell-cycle-related factors in \textit{M. gryphiswaldense}. As for now, it can only be speculated that the severe cell elongation and minicell formation of the \textit{$\Delta$popZ\textsubscript{Mgr}} strain are due to an indirect impairment in FtsZ ring positioning. In \textit{C. crescentus}, MipZ inhibits FtsZ polymerization by generating a gradient with the highest concentration in polar regions via ParB-PopZ-dependent retention of MipZ\textsuperscript{25}, thus creating a region with the lowest MipZ concentration at midcell with suitable conditions for FtsZ ring positioning and formation. Since orthologs of the ParA/ParB chromosome segregation system and MipZ spatial regulator are present in \textit{M. gryphiswaldense}, it is likely that PopZ\textsubscript{Mgr} contributes to stabilization of the MipZ gradient and, thereby, proper placement of the division site. However, the specific functions of ParA, ParB, and MipZ in \textit{M. gryphiswaldense} remain to be elucidated.

Deletion of \textit{popZ} in \textit{M. gryphiswaldense} severely affected motility and apparently polar magneto-aerotactic behavior (Fig. 5, Fig. S4, and Movie S3). Inference of a specific magnetotactic pole-seeking polarity was hypothesized to rely on a yet elusive superimposed mechanism of cellular polarity control, by defining a cellular polarity axis in addition to the magnetosome chain’s magnetic dipole (8, 13). However, the affected motility and loss of swimming polarity are supposedly not directly caused by the absence of \textit{popZ\textsubscript{Mgr}}. In contrast, the aforementioned phenotypes are likely explained by a general impairment of aerotaxis in the \textit{$\Delta$popZ\textsubscript{Mgr}} strain (i.e., due to improper localization of motility-related structures, as discussed below) and as an indirect effect due to formation of short cells that are highly motile, but only weakly aligned within the magnetic field, as well as severe cell elongation, which affects hydrodynamic properties of cells’ propulsion during swimming (as also previously observed for artificially elongated cells caused by cephalaxin treatment [6]). Furthermore, we observed an increased tendency of elongated \textit{$\Delta$popZ\textsubscript{Mgr}} cells to form aggregates (Fig. 5E), which might contribute to the strong motility phenotype observed in soft-agar-based assays (Fig. 5B, C, and D). An increased tendency of \textit{$\Delta$popZ} cells to aggregate has also been observed in \textit{A. tumefaciens} and might be caused by an altered formation of extracellular polysaccharides (25), but in the case of \textit{M. gryphiswaldense} also due to the helical nature of intertwined elongated cells. It can be further hypothesized that the disturbed aerotactic behavior in the \textit{$\Delta$popZ\textsubscript{Mgr}} strain may be due to a delayed or impaired signal transduction from the chemotactic machinery to the flagellar motors, since some cells contained improperly placed (Fig. 3B) or additional chemosensory clusters (as confirmed by fluorescence microscopy of various methyl-accepting chemotaxis proteins [MCPs] fused to GFP in the \textit{$\Delta$popZ\textsubscript{Mgr}} parent strain [results not shown]) as well as occasional flagella located in nonpolar regions (Fig. 2Ai). Altered localization of MCPs, chemoreceptor-associated histidine kinase CheA, and flagellar basal body proteins FliG and FliM upon \textit{popZ} deletion has been also reported for \textit{C. crescentus} (17) and \textit{A. tumefaciens} (25) or in artificially elongated cephalaxin-treated \textit{E. coli} cells (36, 37). However, only a mild effect on motility in swim plate assays has been observed upon \textit{popZ} deletion in \textit{A. tumefaciens} (25), and artificially elongated \textit{E. coli} cells were only affected in their swimming speed, but were still able to perform chemotaxis (36). Hence, due to their different flagellation patterns, cell shapes, and chemotactic behaviors, the experimental results among different strains are not directly comparable.

In addition, severe cell elongation upon \textit{popZ} deletion in \textit{M. gryphiswaldense} resulted in drastically elongated magnetosome chains and a highly increased number of particles per cell (Fig. 2A and B). Our results imply that magnetosome number and chain length are likely directly related to cell length, resembling previously published observations on artificially elongated cephalaxin-treated cells (6). Besides, it has recently been shown that increased gene dosage by genomic multiplication of the magnetosome island results in increased particle numbers as well (38), but with several chains running in parallel or cells closely packed with magnetosomes that lack an ordered chain-like arrangement. Presumably, elongated \textit{$\Delta$popZ\textsubscript{Mgr}} cells also possess an increased number of gene copies due to the presence of multiple chromosomes (albeit we were not able to identify distinct individual chromosomes by DAPI staining, without any specific treatment to condense DNA [Fig. 4A]). However, in contrast to the
overproducer strain (38), the amount of magnetosomes and gene copies per cell volume in elongated ΔpopZ_mgr cells can be assumed to be roughly in the same range as for the wild type.

Magnetosome synthesis, midcell positioning and proper segregation of magnetosome chains are controlled by the treadmilling behavior of the actin-like MamK, which forms dynamic filaments (6, 7). MamK-dependent repositioning of magnetosome chains was not affected in the ΔpopZ_mgr strain (Fig. 2A and Fig. 6A and B), suggesting that PopZ_mgr does not play a role in magnetosome organelle segregation or positioning by exerting direct control of the MamK dynamics. Since magnetosome chain segregation is tightly coupled to cell division, it can be hypothesized that PopZ_mgr may influence magnetosome segregation indirectly—likely by regulating the FtsZ ring localization. Thus, lack of PopZ_mgr causes unequal cell division and misdistribution of chains during cell division as a side effect.

Most strikingly, a consistent bipolar localization pattern of PopZ in M. gryphiswaldense was observed (Fig. 1, Fig. S2, and Movie S1), contrasting with the reported monopolar-to-bipolar transition in C. crescentus (17, 18) and unipolar localization in A. tumefaciens (23–25). Ortholog substitution experiments between C. crescentus and M. gryphiswaldense (Fig. 7 and Fig. S6) indicated that bipolar PopZ localization is not inherent to the protein itself but rather is host specific. PopZ_mgr and PopZ_Cc have conserved N and C termini (Fig. S1) and were capable of partially substituting their reciprocal functionalities (Fig. 7). Thus, the observed localization pattern of both orthologs may be explained by a direct interaction with the respective PopZ interactors present in each host, albeit our results also indicate that both orthologs have diverged to an extent that does not allow full conservation of all PopZ-dependent interactions. For C. crescentus, several factors for control of PopZ localization have been discussed (14, 15, 34, 39). Polar localization of PopZ relies on its self-assembly into higher-order structures in DNA-free polar regions, and the unipolar-to-bipolar transition is coupled to the asymmetric distribution of ParA during the cell cycle (14). The chromosome segregation system adaptor protein ParB and the ParA ATPase, which act together to spatially separate replicated chromosomes in C. crescentus (1), might be suitable candidates for control of bipolar PopZ localization in M. gryphiswaldense. Recently, the zinc finger protein ZitP (28, 40) and muramidase homolog SpmX (41) have been described as additional important factors to nucleate new PopZ microdomains in C. crescentus. An ortholog of ZitP (locus tag MGR_3358) is also encoded in the M. gryphiswaldense genome (23% identity and 39% similarity compared to ZitP_Cc, respectively), whereas no protein orthologous to SpmX is present. Further investigation is needed to identify PopZ interactors in M. gryphiswaldense and elucidate how they differ in function from those of other alphaproteobacteria.

In conclusion, protein functions depend on the genetic context, and can be implemented in different ways, even in closely related species. Thus, M. gryphiswaldense also serves as an appropriate and interesting model organism to study the function of cell cycle factors and its coordination with organelle synthesis and segregation. In the near future, these cell-cycle-related studies will also help to understand how polar magnetotaxis is functionally controlled and inherited in MTB.

MATERIALS AND METHODS

Bacterial strains, plasmids, and culture conditions. Bacterial strains and plasmids used in this study are listed in Text S1 in the supplemental material. Cells were grown using previously described standard procedures described in detail in Text S1.

Molecular and genetic techniques. Oligonucleotides (sequences are listed in Text S1) were purchased from Sigma-Aldrich (Steinheim, Germany). Plasmids were constructed by standard recombinant techniques (as described in Text S1), employing a homologous recombination-based counterselectable system for the construction of in-site deletion and insertion mutants (31) and a Tns-based anhydrotetracycline-inducible expression vector (29, 42) for the construction of transcomplementation and overexpression constructs. All constructs were sequenced by Macrogen Europe (Amsterdam, Netherlands).

Motility assay. Motility soft agar assays were performed as described by Popp et al. (13) and in Text S1. Single-cell tracking was performed at 24°C on a Nikon FN1 Eclipse microscope (Fig. S4A) equipped with an S Plan Fluor 20× differential inference contrast (DIC) N1 objective (NA0.5), a dark-field condenser
addition to DIC microscopy. always stain all cells) and DAPI staining (which we found to unspecifically stain stalks) were used, in

suspension was mixed with either 10

were tracked due to Brownian motion were deleted manually from the analysis. Values given in Fig. 5A are defined as follows: ‘line speed’ is the length of a straight line from the track origin to the current point (≡”line length”) divided by the time elapsed. “Swimming speed” is the track segment length divided by the amount of time elapsed between two positions. “Heading” is the angle between the direction of the velocity vector and the x axis. “Alignment” within the magnetic field (along the x axis) was calculated in Excel (Microsoft) with the following formula: abs(cos(”heading”)).

Fluorescence microscopy. To image fluorescent protein fusions, 3 μl of cell suspension were immobilized on MSR agarose pads (7) and covered with a coverslip. For fluorescent staining, 20 μl of cell suspension was mixed with either 10 μl of a Nile red solution (0.3 μg/ml in dimethyl sulfoxide (DMSO)) or 6 μl of MM64 (an FM4-64 derivative (16 μM in DMSO)) and/or 10 μl of a DAPI solution (50 μg/ml). Conventional epifluorescence microscopy was performed on an Olympus BX81 microscope equipped with a 100× UPLSAPO100×O objective (NA1.4) and an Orca-ER camera (Hamamatsu). Time-lapse imaging and fluorescence recovery after photobleaching (FRAP) were performed on a Deltavision Elite system (GE Healthcare) equipped with a U-Plan S-Apo 100× oil PSF objective (NA1.4) and a CoolSnap HQ2 charge-coupled device (CCD) camera as described previously (7). Additional time-lapse series were acquired on a Nikon Eclipse Ti2-E microscope equipped with a CFI SR Apo TIRF AC 100×H oil objective (NA1.49) and Retiga R1 CCD camera (QImaging). Further methodological details with respect to image deconvolution and the different systems used for epifluorescence microscopy, FRAP, and time-lapse imaging are given in Text S1.

Structured illumination microscopy. 3D-SIM (striped illumination at 3 angles and 5 phases) was performed on a Nikon Eclipse Ti2-E N-SIM E fluorescence microscope equipped with a CFI SR Apo TIRF AC 100×H NA1.49 oil lens objective. Samples (3 μl of FM4-64-stained cell suspension) were immobilized on MSR agarose pads (7). High-resolution coverslips of 0.17 mm thickness (no. 15H; Marienfeld) and immersion oil (type F 30cc; Nikon) with a refractive index of 1.518 (at 23°C) were used to minimize sample-induced spherical aberration. Samples were sealed with wax to provide long-term stability and avoid sample drift during imaging. Calibration of the SIM grating focus and motorized temperature change objective correction collar was performed at 22°C with a bead sample (T-7279 TetraSpeck microspheres) for high-quality image reconstructions at around 115-nm lateral (xy) spatial resolution with 32.5-nm reconstructed image pixel spacing. System and samples were eprequillibrated at 22°C before imaging to avoid temperature-induced changes in the refractive index. Fast piezo stage z-series images were taken at a total thickness of 1.5 to 2.0 μm with 200-nm z-step spacing with raw frame exposure times in the range 100 to 150 ms, avoiding detector saturation of the 16-bit 1.5-electron read noise Orca Flash4.0 LT Plus scCMOS camera (Hamamatsu). EM700/75 and EM525/50 emission filters and fluorescence excitation with 561- and 488-nm laser lines at 80% laser power were used for imaging of FM4-64 and PopZ

Statistical analysis. Statistical analysis was performed in Prism 7.04 (GraphPad) as described in the respective legend to each figure. Data sets were tested for normality using the D’Agostino and Pearson, Shapiro-Wilk, and Kolmogorov-Smirnov tests.
SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.02716-18.

MOVIE S1, AVI file, 10 MB.
MOVIE S2, MPG file, 18.1 MB.
MOVIE S3, MPG file, 1.5 MB.
TEXT S1, DOCX file, 0.1 MB.
FIG S1, PDF file, 2.6 MB.
FIG S2, PDF file, 2.4 MB.
FIG S3, JPG file, 2.7 MB.
FIG S4, JPG file, 2.7 MB.
FIG S5, JPG file, 2.8 MB.
FIG S6, JPG file, 2.8 MB.

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D.S., D.P., and M.T.-N. designed and outlined the study. D.P. constructed plasmids and strains. D.P. performed and analyzed most epifluorescence microscopy experiments. D.P. conducted TEM, 3D-SIM, motility, and transcomplementation experiments and analyzed the data. M.T.-N. performed time-lapse imaging of the wild-type, ΔpopZ, and popZ_Mgr::popZ_Mgr_gfp strains, and D.P. analyzed the data. M.T.-N. performed cryo-electron tomography experiments, and M.T.-N. and J.M.P. analyzed the data. M.T.-N. performed FRAP experiments, and M.T.-N. and M.B. analyzed the data. D.P., M.T.-N., and D.S. wrote the manuscript.

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