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## The HtrA/DegP family protease MamE is a bifunctional protein with roles in magnetosome protein localization and magnetite biomineralization

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### Summary

Magnetotactic bacteria contain nanometer-sized, membrane-bound organelles, called magnetosomes, which are tasked with the biomineralization of small crystals of the iron oxide magnetite allowing the organism to use geomagnetic field lines for navigation. A key player in this process is the HtrA/DegP family protease MamE. In its absence, *Magnetospirillum magneticum* str AMB-1 is able to form magnetosome membranes but not magnetite crystals, a defect previously linked to the mislocalization of magnetosome proteins. In this work we use a directed genetic approach to find that MamE, and another predicted magnetosome-associated protease, MamO, likely function as proteases *in vivo*. However, as opposed to the complete loss of *mamE* where no biomineralization is observed, the protease-deficient variant of this protein still supports the initiation and formation of small, 20 nm-sized crystals of magnetite, too small to hold a permanent magnetic dipole moment. This analysis also reveals that MamE is a bifunctional protein with a protease-independent role in magnetosome protein localization and a protease-dependent role in maturation of small magnetite crystals. Together these results imply the existence of a previously unrecognized “checkpoint” in biomineralization where MamE moderates the completion of magnetite formation and thus committal to magneto-aerotaxis as the organism’s dominant mode of navigating the environment.

### Keywords

bacterial organelle; HtrA/DegP protease; magnetosome; magnetite; biomineralization; heme-binding

### Introduction

Organelles have long been hailed as a defining feature of eukaryotic cells, a definition that is changing as we gain more insight into the membrane- and protein-bound compartments found in bacteria and archaea (Shively, 2006, Fuerst, 2005, Murat *et al.*, 2010a). The

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existence of organelles in bacteria presents interesting evolutionary questions about the conservation of mechanisms of organelle formation, maintenance and organization. Bacterial organelles have also received attention because they often compartmentalize otherwise inefficient or toxic processes, a feature which may be exploited to carry out useful, but potentially cytotoxic, reactions.

One such organelle is the magnetosome of magnetotactic bacteria, which facilitates biomineralization processes that *in vitro* require harsh cytotoxic conditions. The magnetosome is a membrane-bound compartment that directs biomineralization of nanometer-sized, fixed single domain crystals of iron oxide (magnetite, Fe<sub>3</sub>O<sub>4</sub>) or iron sulfide (greigite, Fe<sub>3</sub>S<sub>4</sub>). In a species-dependent manner, cells contain anywhere from ten to hundreds of magnetosomes, which are organized into one or several chains, allowing the cells to align with the earth's magnetic field lines. This passive alignment is thought to facilitate the bacterium's search for favored low oxygen environments, a process referred to as magnetoaerotaxis (Frankel *et al.*, 1997, Smith *et al.*, 2006, Komeili, 2007). The magnetosome membrane is enriched for a specific set of proteins termed magnetosome proteins (Grunberg *et al.*, 2004, Tanaka *et al.*, 2006). Many of these proteins, as well as other factors implicated in magnetosome formation, are encoded by a genomic island, the magnetosome island (MAI), which is a region essential for magnetosome formation conserved in all magnetotactic bacteria studied to date (Fukuda *et al.*, 2006, Richter *et al.*, 2007, Grunberg *et al.*, 2001). The proteins found at the magnetosome are thought to facilitate magnetosome membrane formation, crystal formation, and chain formation and it is hypothesized that they impart the species-specific size and shape of the magnetic mineral.

In *Magnetospirillum magneticum* AMB-1 (AMB-1), magnetosomes are invaginations of the inner membrane formed in a step-wise fashion, where membrane invagination is followed by sorting of magnetosome proteins and magnetite biomineralization (Komeili, 2007, Murat *et al.*, 2010b). Of particular interest is the question of how magnetotactic bacteria generate such uniformly sized and shaped crystals of magnetite, since traditional recapitulation of magnetite synthesis *in vitro* yields a heterogeneous mixture of crystals. How bacteria can exert such tight control over the process of magnetite formation is of interest for many proposed applications in medicine and biotechnology (Schuler & Frankel, 1999) and has driven much of the research on magnetotactic bacteria. In addition, understanding the steps and proteins involved in magnetite biomineralization may also inform how other processes of biomineralization, such as those of teeth or bone, are accomplished.

Although little is known about the molecular mechanisms underlying magnetite biomineralization, recent biochemical and genetic studies have implicated several factors in this process. For instance, addition of the magnetosome protein Mms6 to *in vitro* magnetite synthesis reactions confers more shape and size homogeneity, thus illustrating both the ability of magnetosome proteins to impart shape to magnetite crystals and the potential that specific proteins present for future applications (Arakaki *et al.*, 2003, Arakaki *et al.*, 2010). In *Magnetospirillum gryphiswaldense* MSR-1 (MSR-1), the absence of the *mamCDFG* operon results in cells that form crystals approximately 75% of the size of wildtype crystals, and it was suggested that the activity of these four genes in concert may exercise regulatory or accessory functions in crystal formation (Scheffel *et al.*, 2008). Several factors important for magnetite biomineralization were also recently identified in a genetic dissection of the MAI in AMB-1. Confirming the importance of the *mamCDFG* and *mms6* gene clusters, the loss of a large region of the MAI, R3, which contains both of these gene clusters, resulted in a severe defect in crystal size. In addition, other genes, including *mamP*, *mamR*, *mamS* and *mamT*, were implicated in control of crystal size, number and/or shape. Lastly, four genes, *mamM*, *mamN*, *mamE* and *mamO*, were shown to be essential for the early steps of

biomineralization, as the independent deletion of each gene led to a complete absence of minerals within magnetosomes (Murat *et al.*, 2010b).

Two of these genes, *mamE* and *mamO* encode putative HtrA/DegP family proteases. The members of this family of serine proteases are found in bacteria, archaea and eukaryotes and are known for their involvement in essential housekeeping functions such as the degradative removal of unfolded proteins during the periplasmic stress response, the initiation of the  $\sigma^E$  stress response and the removal of peroxisomal targeting signals (Kim & Kim, 2005, Clausen *et al.*, 2002, Schuhmann *et al.*, 2008). In AMB-1, several HtrA/DegP family proteases are encoded outside of the MAI. MamE and MamO, however, are encoded by the MAI suggesting a specific connection to magnetosome formation rather than a general role in cellular homeostasis. Accordingly, in the absence of *mamE* or *mamO*, cells are viable and form magnetosome membranes, but fail to produce minerals within these membranes (Murat *et al.*, 2010b). This phenotype could be due to a specific defect in mineral formation or result from a more general defect in the localization of biomineralization proteins to the magnetosome. The latter was suggested for MamE since two cytoplasmic magnetosome-associated proteins are mislocalized in the *mamE* deletion strain (Murat *et al.*, 2010b). Also suggestive of less canonical functions for MamE and MamO is the presence of additional functional domains (Clausen *et al.*, 2002). MamE contains two putative *c*-type cytochrome CXXCH heme-binding motifs, and MamO contains a domain of unknown function (DUF81) predicted to code for seven transmembrane domains. Whether the putative protease activities of MamE and MamO are required for magnetosome formation and whether the unusual additional domains are of importance to their function was unknown. We thus undertook a mechanistic dissection of MamE and MamO to further define their roles in magnetosome biogenesis and to begin to develop an understanding of magnetosome formation at the molecular level.

Using a site directed mutagenesis, we show that MamE and MamO are likely to act as proteases *in vivo* and that the additional domains, uncommon for HtrA/DegP family proteases, are important for their functions. By dissecting MamE's functional domains, we find that MamE is a bifunctional protein with a previously unidentified role in biomineralization that can be decoupled from its role in the proper localization of magnetosome proteins. Based on these results, we propose that MamE's protease activity is required for a previously unidentified crystal size transition from 20nm crystals too small to hold a fixed magnetic dipole moment, to large crystals that can contribute to the cell's magnetic response.

## Results

### MamE and MamO act as proteases *in vivo*

MamE and MamO are putative HtrA/DegP family proteases encoded by genes found within the MAI. In their absence, cells are non-magnetic yet still possess the ability to form magnetosome membranes (Murat *et al.*, 2010b, Yang *et al.*, 2010). Members of the HtrA/DegP family of proteins are trypsin-like serine proteases that share a high degree of sequence homology within their protease domains, including a highly conserved active site triad (Kim & Kim, 2005). Primary sequence alignment of MamE and MamO with *E. coli* DegP allowed identification of their putative active site residues. MamE shares the conserved histidine-aspartate-serine active site (Fig. 1A), whereas MamO has a threonine in place of the serine as the predicted active site nucleophile (Fig. 1B). To determine whether MamE and MamO's putative protease functions are required for magnetite crystal formation, we generated the mutant constructs, *mamE<sup>P</sup>* and *mamO<sup>P</sup>*, by site-directed mutagenesis of the predicted active site triad residues to alanines. As a preliminary assay for the activity of these variants *in vivo*, we relied on the  $C_{mag}$  measurement, a differential

spectrophotometric assay that quantifies the ability of the bacteria to orient in an external magnetic field (Schuler *et al.*, 1995). In this assay, *mamO<sup>P</sup>* fully complemented the *mamO* deletion, and *mamE<sup>P</sup>* partially complemented the *mamE* deletion (Fig. 1D), suggesting that the putative protease functions are not essential for crystal formation.

The MAI, however, encodes paralogues of both MamE and MamO, Amb1002 and Amb1004, respectively (Figs. 1A and B) which we have renamed Like-MamE (LimE) and Like-MamO (LimO). Since HtrA/DegP proteases are known to function as oligomeric assemblies (Clausen *et al.*, 2002, Kim & Kim, 2005) we hypothesized that these paralogues could form hetero-oligomeric complexes with MamE and MamO and thus provide active protease domains *in trans*. To test this hypothesis, we deleted *mamE* and *mamO* in the R9 deletion background. R9 is a region of the MAI containing both *limE* and *limO* that can be deleted in the wildtype background without a detectable phenotype (Murat *et al.*, 2010b). The resulting double deletion strains,  $\Delta R9\Delta mamE$  and  $\Delta R9\Delta mamO$ , could be complemented to the same degree as the single deletion strains by wildtype *mamE* and *mamO*, respectively. However, the putative protease-inactive MamE mutant could not restore a measurable magnetic response when expressed in the  $\Delta R9\Delta mamE$  strain (Fig. 1D). Constructs of MamO in which all putative active site triad residues of the protease domain are mutated to alanines also did not restore a magnetic phenotype to the  $\Delta R9\Delta mamO$  strain. As noted above, however, MamO has a threonine in the position of the conserved active site serine residue. This threonine can be changed to an alanine without reduction of MamO's ability to complement  $\Delta R9\Delta mamO$  (Fig. S1). A single point mutation of the putative active site histidine, however, decreased MamO activity, a phenotype that would be expected if this residue were part of the serine protease active site triad (Fig. S1).

This mutational analysis suggests that the two proteins may function as proteases *in vivo* and that they may act within hetero-assemblies with LimE and LimO, respectively. To build on these initial results we chose to focus on MamE, with the aim of defining the specific function of its putative protease activity in magnetosome protein targeting and magnetite formation. Additionally, to avoid background effects from the unusual cross-complementation observed above, all subsequent experiments were performed in the  $\Delta R9\Delta mamE$  background.

### MamE protease mutant produces small magnetite crystals

To further characterize the phenotype of the  $\Delta R9\Delta mamE/mamE^P$  strain, cells were imaged by transmission electron microscopy (TEM). As described above, this mutant could not restore a magnetic response in the  $\Delta R9\Delta mamE$  strain. However, TEM analysis revealed that it still allowed for synthesis of chains of small electron dense structures (Fig 2). Under the same growth conditions, the  $\Delta R9\Delta mamE$  strain complemented with wildtype *mamE* showed a bimodal crystal size distribution with peaks centered in the 36–40 nm and 51–55 nm size ranges (Fig. 2). In contrast, when *mamE<sup>P</sup>* was expressed in the  $\Delta R9\Delta mamE$  strain, particle size distribution was centered at 16–20 nm, and only 3% were larger than 35 nm. Magnetite crystals smaller than 30 nm are too small to hold a permanent magnetic dipole moment (Butler & Banerjee, 1975), perhaps explaining the inability of the *mamE<sup>P</sup>* mutant to align in a magnetic field. High resolution TEM (HRTEM) confirmed that these 16–20 nm particles, and even those that were irregular in shape, were indeed crystalline (Fig. 3B and C, Fig S2). HRTEM allows for visualization of the lattice fringes of crystalline material and the spacing between these fringes can be characteristic of the crystallographic planes of various minerals. Lattice fringe images of most of the crystalline particles, formed when the *mamE<sup>P</sup>* mutant was expressed in the  $\Delta R9\Delta mamE$  strain, showed a characteristic *d*-spacing of 0.48 nm corresponding to {111} plane of magnetite (Figs. 3B and S2). The XRD data of magnetite in the mineral database <<http://rruff.info>> was used as reference for these measurements. Other crystals showed measurements that were also consistent with other

planes of magnetite. In addition, spectra from energy dispersive x-ray spectroscopy (EDS) of these small crystals are consistent with the presence of magnetite (Fig. 3D). While the possibility that other Fe-containing phases are produced in this mutant cannot be completely ruled out, the most parsimonious explanation is that the crystals produced in the absence of the putative protease activity of MamE are indeed magnetite. Furthermore, small crystals were observed within mature sized magnetosome membranes, suggesting that their growth is not restrained by smaller magnetosome membranes (Fig. 3A).

Recently, a second MamE paralog, MamE-like, was annotated outside of the MAI as part of the newly identified magnetosome islet (Rioux *et al.*, 2010). To rule out the possibility that MamE-like can cross-complement MamE protease activity and is responsible for the 20 nm crystals observed in the  $\Delta R9\Delta mamE/mamE^P$  strain, the triple *mamE* deletion strain  $\Delta R9\Delta mamE\Delta mamE\text{-like}$  was generated. The triple deletion strain could be complemented with wildtype *mamE* to the same extent as the  $\Delta R9\Delta mamE$  and  $\Delta mamE$  strains. *mamE^P* still supported the growth of small 20 nm crystals in this background (Fig. S3), suggesting that unlike LimE, MamE-like is not capable of cross-complementing protease activity. Together, these data suggest that MamE's putative protease function is not essential for initiation of biomineralization.

### MamE's protease activity is required for a novel crystal size transition step

The above results suggest that the putative protease activity of MamE is required for continued growth of magnetite crystals beyond the 16–20 nm size range. The results could also be consistent with a delay in the initiation of biomineralization rather than a change in the kinetics of crystal growth. To distinguish between these two models, we examined the development of the magnetic phenotype and biomineralization of magnetite over time. Briefly, we passaged strains carrying different versions of *mamE* in the absence of iron until no electron-dense structures were detectable by TEM. Cultures were then moved to iron-containing medium, and their optical densities and C<sub>mag</sub> values were monitored over time. Wildtype complemented  $\Delta R9\Delta mamE$  became magnetic four hours post addition of iron and reached its maximum C<sub>mag</sub> of ~2 after 1335 minutes. As expected, cells expressing only MamE<sup>P</sup> do not become magnetic (Fig. 4A).

To observe the morphological characteristics associated with the biomineralization of magnetite at the early stages, samples obtained at the following points in time were prepared for TEM: (i) before any strain becomes magnetic; (ii) the point at which the wildtype strain becomes magnetic; and (iii) late time points after both strains have been growing in iron for some time. After one hour of growth in iron-containing media, the earliest point in our time-course, both strains had produced electron-dense structures of similar size and shape (Fig. 4B), suggesting that at the time resolution of our experiments, MamE<sup>P</sup> was capable of initiating biomineralization properly. After three hours, TEM showed large crystals in the  $\Delta R9\Delta mamE$  cells expressing *mamE*, whereas the same strain carrying *mamE^P* showed only small electron-dense structures (Fig. 4B). In fact in the presence of MamE<sup>P</sup>, cells were unable to build large crystals, confirming the arrest at the 16–20 nm growth stage seen in the steady-state cultures (Fig 4B and C). Furthermore, the crystal size distributions the  $\Delta R9\Delta mamE$  strain complemented with wildtype *mamE* or *mamE^P* are remarkably similar at the one hour timepoint (Fig 4C), suggesting that the putative protease function of MamE is not required for initiation of biomineralization but for a potentially unrecognized crystal size transition step. Although the overall trend is similar, for unknown reasons, size distributions measured for these time-courses are slightly shifted towards larger crystals as compared to distributions seen in end point experiments shown in Figure 2. It should also be noted that a small number of crystals (~3%) seem to escape this arrest and attain sizes of greater than 35nm and thus fixed single domain sizes (Fig. 2 and Fig. 4C).

### MamE's putative heme-binding motifs are required for wildtype biomineralization

Unlike most other Deg family proteases, MamE is predicted to have additional functional domains. Specifically, MamE contains two putative CXXCH heme-binding motifs (Fig. 1C), which in *c*-type cytochromes are known to bind heme covalently via thioether bonds formed through the two conserved cysteines (Bowman & Bren, 2008). To determine whether MamE's CXXCH motifs are required for magnetosome formation, we used site-directed mutagenesis to change the conserved cysteine residues of these motifs to alanines. This construct, *mamE<sup>C2</sup>*, can only partially complement magnetite formation in the  $\Delta R9\Delta mamE$  strain (Fig. 1D), restoring a Cmag of  $1.14\pm 0.03$  as opposed to  $1.84\pm 0.05$  when complemented with wildtype *mamE*. This suggests a role for MamE's putative heme-binding motifs in crystal formation.

TEM analysis revealed that this mutant is capable of forming large, wildtype-sized crystals. However, complementation with *mamE<sup>C2</sup>* yielded only ~30% of crystals larger than 35 nm, compared to 82% when the  $\Delta R9\Delta mamE$  strain was complemented with wildtype *mamE*. The crystal size distribution of *mamE<sup>C2</sup>* was shifted to smaller crystal sizes with a bimodal crystal size distribution centered at 16–20 nm and 36–40 nm (Fig. 2). The 16–20 nm peak observed in this strain clearly overlapped with the crystal size distribution of MamE<sup>P</sup> strain. Additionally, HRTEM and EDX spectroscopy confirmed that the 16–20 nm crystals in the *mamE<sup>C2</sup>*-complemented strain are of similar composition to those found in the *mamE<sup>P</sup>*-complemented strain (data not shown). Interestingly, in time-course experiments, the strain expressing *mamE<sup>C2</sup>* was always delayed in the onset of its magnetic response as compared to one bearing wildtype *mamE* (Fig. 4A), even though growth rates of these strains were not significantly different (data not shown). TEM analysis showed that similar to the strain carrying *mamE<sup>P</sup>*, the one with *mamE<sup>C2</sup>* does not show obvious defects in the early steps of biomineralization leading to 20 nm crystals. Instead, MamE<sup>C2</sup>'s decreased number of large magnetite crystals is due to a defect in late steps of biomineralization (Fig. 4B and C).

This suggests that these 20 nm crystals may be an early intermediate in magnetite biomineralization and further supports the hypothesis that MamE is required for the transition from 20 nm crystals to larger crystals that can contribute to a cell's magnetic response.

### MamE is a bifunctional protein with roles in protein localization and biomineralization

The absence of crystals in the  $\Delta mamE$  strain had previously been attributed to a defect in magnetosome protein localization as two soluble magnetosome proteins, MamA and MamJ, were mislocalized in this mutant strain (Murat *et al.*, 2010b). To determine whether this defect extends to integral magnetosome membrane proteins, we generated C-terminal GFP fusions to MamC and MamF using a ten-glycine linker, as previously described (Lang & Schuler, 2008). We also assayed localization of GFP-MamI (Murat *et al.*, 2010b). All three proteins form continuous linear structures in wildtype AMB-1 that are reminiscent of the localization of the magnetosome chain. This pattern is disrupted in the  $\Delta R9\Delta mamE$  (Fig. 5 and Fig. S4A) and  $\Delta mamE$  strains (Fig. S4B). Several localization patterns were observed for the GFP constructs in these deletion backgrounds, ranging from one or multiple foci to evenly distributed membrane localization. This suggests that MamE is required for the proper localization of not only soluble but also membrane-bound magnetosome proteins. It should be noted that in a small percentage of cells, less than 1%, foci were arranged in a linear pattern that could be suggestive of some degree of magnetosome localization (Fig. S4C).

It thus seemed likely that MamE's protease function is required for the proper localization of magnetosome proteins. To test this hypothesis, localization of MamC-GFP and GFP-MamI

were assessed in the  $\Delta R9\Delta mamE$  strain complemented by wildtype *mamE*, *mamE<sup>P</sup>*, or *mamE<sup>C2</sup>*. Wildtype *mamE* complemented both the magnetic response (C<sub>mag</sub>) and protein localization defects of the  $\Delta R9\Delta mamE$  mutant (Table 1 and Fig 5). However, to our surprise, *mamE<sup>P</sup>*, which cannot restore a magnetic response in the  $\Delta R9\Delta mamE$  strain, could complement the protein localization defect. Similar levels of restoration of protein localization were also observed for the strain complemented with *mamE<sup>C2</sup>* (Table 1 and Fig. 5). This suggests that MamE is a bifunctional protein with independent roles in protein localization to the magnetosome and in biomineralization.

## Discussion

To date, most molecular studies of magnetosome formation have focused on identifying magnetosome-associated proteins and determining the phenotypes of deletion or disruption mutants (Grunberg *et al.*, 2004, Tanaka *et al.*, 2006, Fukuda *et al.*, 2006, Komeili *et al.*, 2004, Okuda *et al.*, 1996). These studies have been powerful in providing a list of candidate proteins involved in this intricate process and have laid the foundations for more detailed mechanistic studies. In this work, we used a mutational dissection of the HtrA/DegP family protease MamE to show that this protein acts at two functionally distinct steps of magnetosome formation. First, MamE is required for proper localization of magnetosome proteins. This role does not require MamE protease activity, and proper targeting of magnetosome proteins is not sufficient to produce large magnetic minerals. Since localization of magnetosome proteins does not require MamE protease activity, it could be accomplished through physical interaction of MamE with one or more magnetosome proteins at the magnetosome. Alternatively, it may be mediated by a chaperone-like activity of MamE, which has been attributed to some HtrA/DegP proteins and is separable from their protease activity (Spiess *et al.*, 1999, Rizzitello *et al.*, 2001). Once magnetosome proteins are correctly localized, MamE's protease activity is required for its second role in magnetosome formation, the maturation of small, 20 nm crystals into larger single-domain crystals with fixed dipole moments. We also show that MamO, a second HtrA/DegP family protease encoded by the MAI, likely has protease activity *in vivo*. However, further biochemical characterization is required to determine conclusively whether MamE and MamO possess protease activity and if so, what their substrates might be.

Canonical HtrA/DegP proteases consist of a protease domain coupled to one or multiple PDZ domains (Clausen *et al.*, 2002, Kim & Kim, 2005). Strikingly, MamE and MamO have additional functional domains and we show these domains to be important for their function. MamO contains a seven transmembrane spanning domain of unknown function (DUF81). MamO's paralogue, LimO, is identical to MamO except that it lacks the DUF81 domain (Fig. 1C). Previous proteomic work has suggested that LimO is expressed in wild-type cells (Tanaka *et al.*, 2006). We show that LimO can cross-complement the protease function of MamO when only a protease-inactive version of MamO is expressed in AMB-1. Thus, if LimO is expressed in the  $\Delta mamO$  strain, as it is the  $\Delta R9\Delta mamO/mamO^P$  strain, the non-magnetic phenotype of  $\Delta mamO$  can be attributed to the lack of a DUF81 domain in LimO. This is consistent with the inability of *mamO* lacking DUF81 to complement a *mamO* disruption strain in *Magnetospirillum gryphiswaldense* MSR-1, a species that does not have a homolog of *limO* and thus only encodes one copy of *mamO* (Yang *et al.*, 2010). The function of the DUF81 domain remains mysterious. While it has been suggested to function as an anion transporter in some systems (Weinitschke *et al.*, 2007, Mampel *et al.*, 2004), direct experimental evidence for such an activity is lacking. Alternatively, it is possible that this portion of MamO acts as a localization determinant to bring the protein to the magnetosome.

Similar to MamO, MamE is an unusual HtrA/DegP family protease, in that it also has additional functional domains, two putative CXXCH heme-binding motifs. These motifs are known to bind heme covalently via thioether bonds between the cysteines of the CXXCH motif and the heme vinyl group, and are most commonly associated with *c*-type cytochromes (Bowman & Bren, 2008). *c*-type cytochromes have been shown to function in the oxidation and reduction of metals (Paquete & Louro, 2010) and to act as gas sensors (Takayama *et al.*, 2006). MamE could thus act in the reduction and/or oxidation of iron required for magnetite formation. However, MamE cannot be the sole player in this role, since the heme-binding mutant is still capable of biomineralization of mature magnetite crystals. Alternatively, it is possible that MamE's CXXCH domains are regulatory in nature. Interestingly, two other magnetosome proteins implicated in biomineralization, MamP and MamT, also have CXXCH motifs, raising the possibility that a network of redox activity may be at the center of magnetite biomineralization. Another possible function for the CXXCH motifs is that they could modulate MamE's protease activity. Time-course experiments showed that although *mamE<sup>C2</sup>* is capable of synthesizing large fixed single-domain magnetite crystals, it is delayed or slowed in this process and forms a significant number of small 20 nm crystals similar to those found in *mamE<sup>P</sup>*. This similarity in phenotype to *mamE<sup>P</sup>* suggests that MamE's CXXCH motifs may serve to activate or enhance proteolysis required for crystal size transition.

While it is clear that protease activity of MamE is linked to a specific step of biomineralization, the mechanisms by which this is achieved remain mysterious. We could envision that once a crystal has reached the 20 nm transition point, MamE (i) degrades one or several inhibitors of biomineralization (Fig. 6A), or (ii) proteolytically activates proteins essential for further crystal growth (Fig. 6B). If proteolysis by MamE activates biomineralization factors, then these should be proteins that act in the post-nucleation steps of biomineralization and their deletion phenotypes should be similar to that of the MamE protease mutant. Thus, possible substrates of MamE, based on our current knowledge of biomineralization, could be MamS, MamR, MamT and the proteins encoded by the R2 and R3 genomic regions, which include the *mms6* and *mamCDFG* gene clusters (Arakaki *et al.*, 2003, Scheffel *et al.*, 2008, Murat *et al.*, 2010b). Inhibitors of crystal size transition would have eluded identification by methods currently used to screen biomineralization mutants, since their deletion or disruption would presumably lead to larger magnetite crystals, a phenotype that likely would not have caused a noticeable difference in the cells' magnetic response. Interestingly, MamE and MamO are not the only examples of serine proteases essential for biomineralization. Enamel proteases have long been known for their role in tooth formation, where they act to remove an organic matrix from nucleated enamel crystallites allowing for growth of the crystallites into mature-sized enamel (Bartlett & Simmer, 1999). Additionally, some serine proteases are capable of precipitating metal oxides *in vitro*, raising the intriguing possibility that MamE or MamO could play a direct role in the formation of iron oxide crystals (Smith *et al.*, 2009). Thus, identification of MamE and MamO's substrates may shed light on the potential similarities between the biomineralization pathways of magnetotactic bacteria and those found in other organisms.

The work presented here helps define the functional relevance of MamE and MamO in more detail. We speculate that our results may also have uncovered a previously unrecognized "checkpoint" step in magnetite biomineralization where the protease activity of MamE is required for the development of mature magnetite crystals. The transition from small superparamagnetic crystals to larger single domain crystals is an important decision point for the bacterium, one that will trap the cell in a forced biased swim guided by the geomagnetic field, leaving it to use magnetoaerotaxis as its sole mode of exploring the environment. Thus, if MamE's protease activity could be modulated, cells could arrest biomineralization at the 20 nm stage and be primed for the formation of larger crystals



without committing themselves to magnetoaerotaxis. When desirable conditions are reached, biomineralization could be resumed by simply activating MamE protease activity. Such a system would allow cells to align with the earth's magnetic field lines when favorable without being committed to a forced directional swim under all environmental conditions. In addition, the two-fold increase in the diameter of the crystal represents an approximately eight-fold increase in its volume, meaning that significant resources must be dedicated to build a magnetosome chain after this transition point. MamE's protease activity could integrate the availability of iron with the need to maximize the production of large single domain crystals. It remains to be determined whether MamE's protease activity is modulated *in vivo*, potentially via its putative heme-binding motifs. Regardless of the physiological role of this transitional step in biomineralization, our work has uncovered intriguing possibilities for genetic engineering of MTB where artificial control over MamE protease activity might allow for external control of crystal size and magnetoaerotaxis.

## Experimental Procedures

### Growth conditions

Cells were grown in defined minimal media (MG medium) supplemented with both Wolfe's Vitamin Solution and iron (3 mM iron chloride 9 mM malate) at 1/100 (Murat *et al.*, 2010b). Additional iron sources were omitted when preparing Wolfe's Mineral Solution. For fluorescence microscopy cells, were grown in 10 mL of MG with 20 mL of head space in a microaerobic chamber maintained at 30°C and less than 10% oxygen. To assay complementation and to measure crystal size distributions, cells were grown in a 30°C incubator in sealed Balch tubes containing 10 mL of MG and 20 mL headspace. Balch tubes were flushed briefly with N<sub>2</sub> after autoclaving and for 10 minutes after addition of iron and vitamins before cells were added. No additional air was added to the tubes. For time course experiments, cells were grown in Balch tubes, as described above, except that the culture volume was increased to 15ml. For these experiments, all glassware was soaked in oxalic acid for 12 hours before use, and the cells were passaged twice in MG medium containing no iron until the cultures were non-magnetic and no crystals or inclusions were observable by TEM. Solid media plates contained 7g agar per liter medium. Antibiotics were used at the following concentrations: kanamycin at 15 µg/mL in solid media, and at 10 µg/mL in liquid media; 7 µg/mL in liquid when plasmid was integrated on the chromosome; and carbenicillin 2 0µg/mL on solid and liquid media.

### Plasmids, Primers and Strains

All primers, plasmids and strains used in this study are listed in Supporting Information tables S1 and S2 and S3.

### Time-course experiments

For time-course experiments cells were passaged twice in 10 mL MG medium containing no iron in 20 mL tubes and incubated in a microaerophilic chamber. Cells in exponential phase were then passaged into 15 mL MG medium plus iron malate in Balch tubes and the increase in C<sub>mag</sub> was monitored (Komeili *et al.*, 2004). To follow crystal growth, cells were collected by filtration, washed in PBS and fixed in 2.5% glutaraldehyde in 0.1 M sodium cacodylate buffer for TEM analysis. Crystal size was measured by hand using GIMP software and the long axis is reported as crystal size.

### Generation of $\Delta R9\Delta mamE$ , $\Delta R9\Delta mamO$ and $\Delta R9\Delta mamE\Delta mamE$ -like

All deletions were generated using the two-step recombination method previously described (Murat *et al.*, 2010; Komeili *et al.*, 2004). *mamE* and *mamO* were deleted in the  $\Delta R9$

background using the plasmids pAK241 and pAK243 described in Murat *et al.*, 2010. To delete *mamE-like* in the  $\Delta R9\Delta mamE$  background, regions flanking the gene were amplified and combined into the deletion plasmid pAK455 by fusion PCR.

### Generation of MamC-GFP and MamF-GFP

All PCR except for QuikChange® site directed mutagenesis were carried out using the Promega GoTaq® Green Master Mix. MamC was C-terminally GFP-tagged by fusion PCR. MamC was amplified using a forward primer adding an EcoRI site and a reverse primer deleting the stop codon and adding a BamHI site followed by 10 C-terminal Glycines as a linker. GFP was amplified using a forward primer adding 10 N-terminal Glycines and a reverse primer adding a SpeI site. The MamC-BamHI-10Glycine-GFP construct was then generated by fusion PCR using the MamC forward and GFP reverse primers. This fragment was EcoRI/SpeI cloned into pAK22 to generate pAK452. To confer carbenicillin resistance, the ampicillin resistance gene (*bla*) placed downstream of the *tac* promoter was isolated from pAK237 by SpeI digestion and then SpeI cloned into pAK452 to generate pAK453. MamF was amplified adding an N-terminal EcoRI and a C-terminal BamHI site, deleting the stop codon. This fragment was cloned into pAK452 to replace MamC generating pAK454.

### Fluorescence microscopy

Cells were imaged in early stationary phase on 1% agarose pads using a Nikon Eclipse 80i microscope. Images were acquired at 1000X magnification using a QImaging® RETIGA 2000R Fast 1394 camera.

### Generation of *mamO* and *mamE* complementation plasmids

The previously described plasmids pAK263 and pAK250 (Murat *et al.*, 2010b) were used for wildtype complementation of the *mamO* and *mamE* deletion strains respectively. To identify *mamO* and *mamE*'s putative active site residues, their amino acid sequence was aligned to *E. coli* DegP, DegQ and DegS using ClustalW2 (Chenna *et al.*, 2003). Putative active site residues were changed to alanines using Stratagene QuikChange® mutagenesis as directed by the manufacturer. MamE's putative heme-binding motifs were inactivated using the same strategy to change the two cysteines of the CXXCH domains to alanines. The *mamO* deletion strains were complemented with *mamO* on a plasmid expressed off of the *tac* promoter (pAK263) whereas *mamE* deletion strains were complemented with *mamE* expressed from its endogenous promoter by integrating *mamE* into the chromosome (pAK250) (Murat *et al.*, 2010b).

### Complementation assays

Cmag measurements were performed with cultures grown in 10ml of MG in sealed 20 mL Balch tubes after two days of growth at 30°C. For *mamO* complementation, Cmags were measured in the presence of kanamycin. For *mamE* complementation, Cmags were measured in the absence of any antibiotics. Complementation of protein localization was assayed by growing complemented cells in 10 mL of MG with carbenicillin in 20 mL tubes in a microaerobic chamber.

### TEM

For imaging of whole mounts in TEM, a 5  $\mu$ L drop of the suspended cells was adsorbed onto 200-mesh Cu grid coated with Formvar film. Samples were imaged with an FEI Tecnai 12 TEM equipped with a Gatan Bioscan (1k  $\times$  1k) CCD Camera Model 792 at an accelerating voltage of 120 kV.

## Cryo-ultramicrotomy and EDS spectroscopy

Cryo-ultramicrotomy was performed as previously described. The sections were imaged with an FEI Tecnai G<sup>2</sup> F20 cryo-S/TEM equipped with a Gatan Ultrascan 1000 (4k × 4k) CCD camera and a EDAX Genesis microanalytical system at an accelerating voltage of 200 kV.

## Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

## Acknowledgments

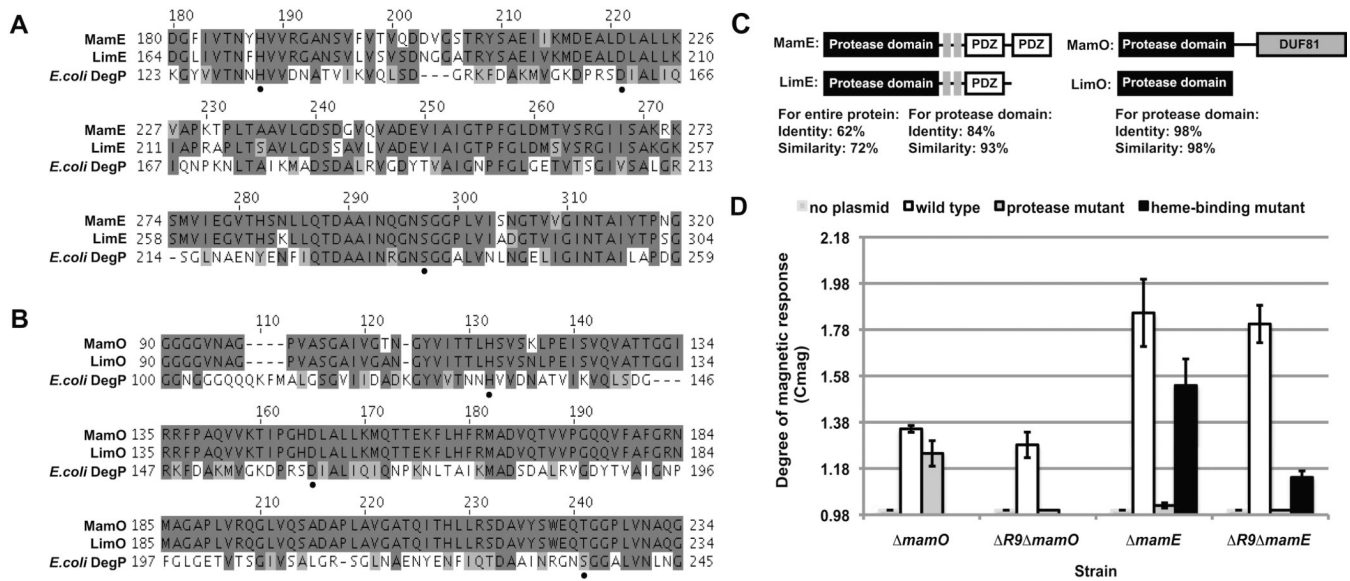
We thank Komeili laboratory members, K. Ryan (also at UC Berkeley) and R.J. Quinlan for their valuable input and discussion both into this work and into preparation of this manuscript. A.Q. was supported by the Winkler Family Foundation. A.K. was additionally funded by the David and Lucille Packard Foundation and the National Institute of Health (5R01GM084122). H.V. acknowledges the Natural Sciences and Engineering Research Council of Canada for financial support.

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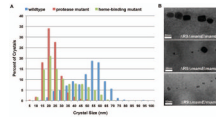
**Fig. 1. AMB-1 magnetic response (Cmag) depends on MamE and MamO protease activity**

**A.** Alignments of MamE and LimE protease domains with *E. coli* DegP.

**B.** Alignments of MamO and LimO protease domains with *E. coli* DegP. Active site triad residues of MamE and MamO, indicated by black dots, were identified based on homology with DegP. Identical residues are highlighted in dark grey and conserved substitutions in light grey.

**C.** Domain architecture of MamE, LimE, MamO and LimO. MamE heme-binding motifs are indicated by grey boxes. For MamE and LimE, identity for both the entire protein and for the protease domains only are shown. For MamO and LimO identity of the protease domain is shown.

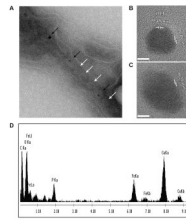
**D.** Complementation of magnetic response of the single and double *mamO* and *mame* deletion strains with wildtype and protease-inactive (*mame<sup>P</sup>* and *mamO<sup>P</sup>*) constructs. For MamE, complementation by the heme-binding-deficient construct (*mame<sup>C2</sup>*) is also shown. Error bars represent one standard deviation from  $\geq 10$  independent cultures.



**Fig. 2. MamE protease domain is essential for crystal maturation**

**A.** Crystal size distribution of  $\Delta R9\Delta mamE$  complemented with wildtype, protease inactive ( $mamE^P$ ), and heme-binding deficient  $mamE$  ( $mamE^{C2}$ ). Crystal size is plotted as the percent of the total number of crystals that fall into each 5nm bin.  $n > 500$  crystals were measured for each strain from two independent  $\Delta R9\Delta mamE/mamE$ , three independent  $\Delta R9\Delta mamE/mamE^P$ , and four independent  $\Delta R9\Delta mamE/mamE^{C2}$  cultures.

**B.** Representative TEM images of crystals for each of the three complemented strains.  $>100$  fields were visualized by TEM from  $>5$  independent cultures.



**Fig. 3. Electron-dense structures within MamE protease mutant are crystalline**

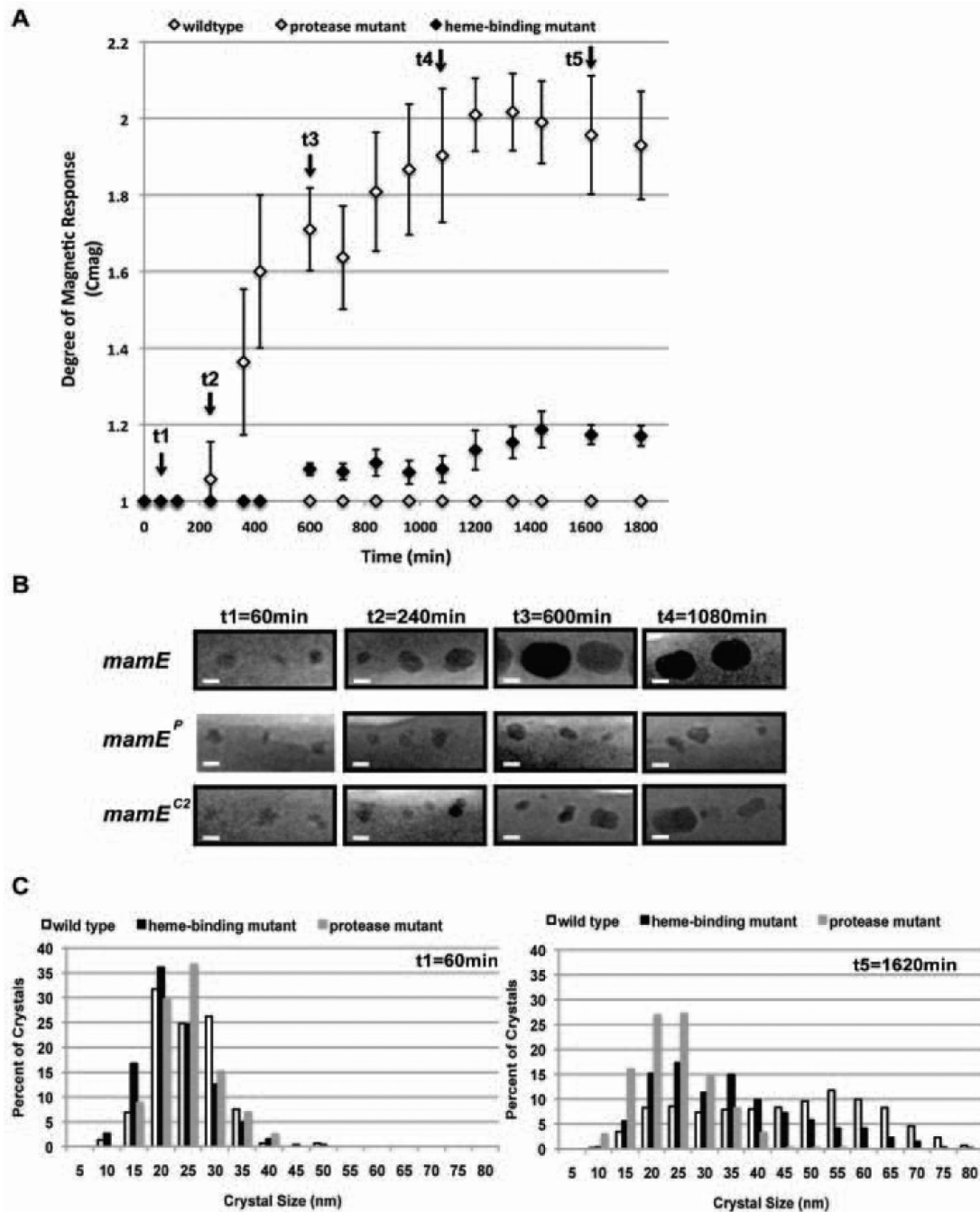
**A.** Representative TEM image of a cryo-sectioned  $\Delta R9\Delta mamE$  cell complemented with the MamE protease mutant ( $mamE^P$ ) showing that the small electron-dense 20nm inclusions grow within mature-sized magnetosome membranes. A chain of both empty (white arrows) and inclusion-containing (black arrows) magnetosomes can be observed.

**B.** HRTEM shows that 20nm electron-dense structures formed when  $mamE^P$  is expressed in the  $\Delta R9\Delta mamE$  strain, are crystalline, as indicated by the characteristic appearance of the lattice fringes as parallel lines. In the image shown, a measurement of 0.48 nm is consistent with the {111} plane of magnetite. Dotted white lines indicate linear pattern. Scale bar represents 5 nm.

**C.** HRTEM shows even amorphous electron-dense structures found in this strain are crystalline. Similar to part B, the lattice fringe measurement of 0.25 nm is consistent with the {311} plane of magnetite. Scale bar represents 5 nm.

**D.** EDX spectra of 20nm inclusions show iron and oxygen peaks consistent with the presence of magnetite. Copper signal is due to copper grids, and phosphorous and carbon peaks from media/cells.



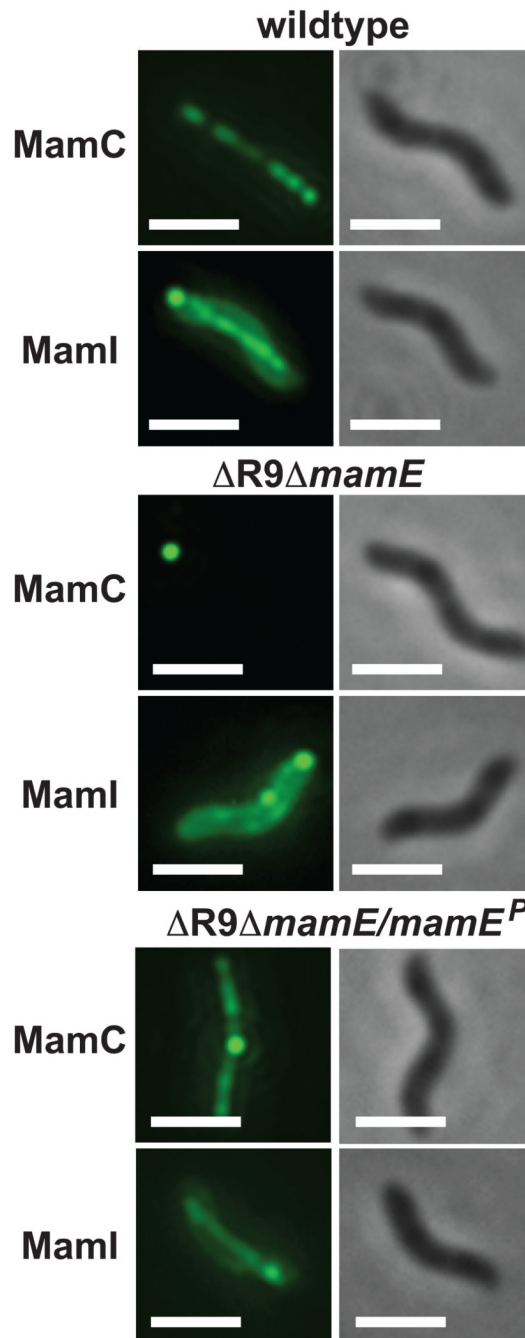


**Fig. 4. MamE protease and heme-binding mutants are not defective in early steps of biomineralization**

**A.** Time-course of increase in magnetic response of complemented strains. Cells were passaged in the absence of iron and then transferred into iron-containing media to initiate magnetite crystal formation. In contrast to the *mamE* protease mutant the *mamE* heme-binding mutant becomes magnetic but is delayed in the onset of its magnetic response as compared to wildtype complemented  $\Delta R9\Delta mamE$ . Error bars represent one standard deviation of three independent cultures. Arrows indicate points at which cells were fixed for TEM imaging (see B) and crystal size determination (see C).

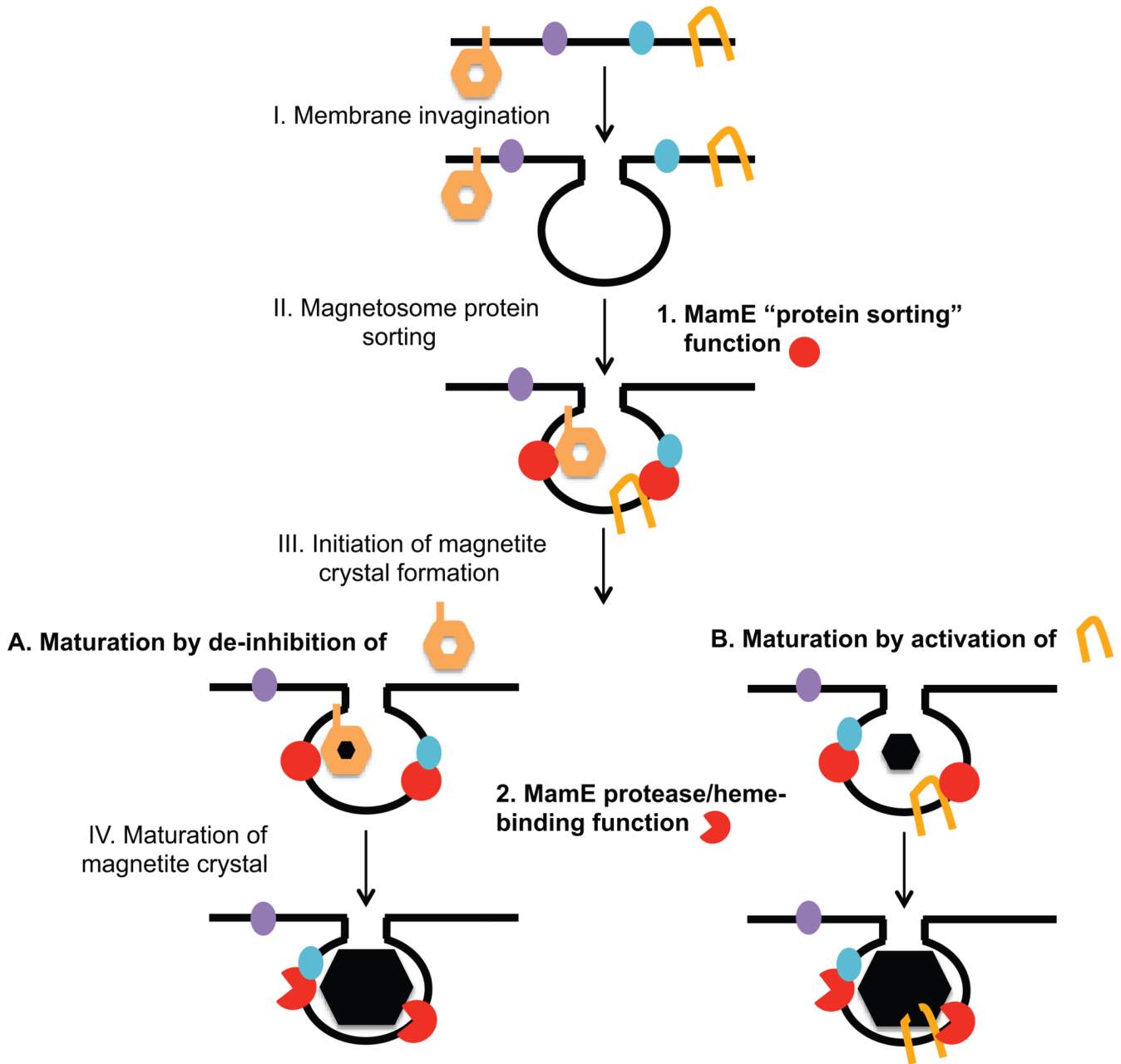
**B.** Representative TEM images of electron dense inclusions from  $\Delta R9\Delta mamE$  strain complemented with either wildtype *mamE*, or protease- (*mamE<sup>P</sup>*) or heme-binding-deficient (*mamE<sup>C2</sup>*) derivatives. At 60 minutes, all three strains produce similar-sized electron-dense inclusions. >150 crystals were visualized and measured at each timepoint.

**C.** Size distribution of electron dense structures formed by the three strains at early (60 minutes post addition of iron) and late (1620 minutes post addition of iron) time points. Cells were fixed and imaged by TEM at each time point. Crystals were measured and the long axis of the crystal is reported as crystal size. n>150 for 60 minute. n>350 for 1620 minute time point.



**Fig. 5. MamE is a bifunctional protein with a protease-independent role in magnetosome protein localization**

Representative images of localization pattern of MamC-GFP and GFP-MamI and in wildtype AMB-1, in  $\Delta R9\Delta mamE$  and in  $mamE^P$  complemented  $\Delta R9\Delta mamE$ . These magnetosome proteins form linear structures reminiscent of the localization of the magnetosome chain in wildtype cells that are disrupted in  $\Delta R9\Delta mamE$ . Wildtype localization patterns can be restored with  $mamE^P$ . >200 cells were imaged. Scale bar represents 1  $\mu$ m.



**Fig. 6. Model for MamE's role in magnetosome formation**

After magnetosome membranes are formed MamE is required in a protease independent fashion for the proper localization of at least a subset of magnetosome proteins. This activity is sufficient for the formation of 20nm crystals of magnetite. MamE's protease function is then required to mature these crystals into large fixed single domain crystals of magnetite. This could be achieved either by (A) proteolytically removing one or several inhibitors of magnetosome formation or an inhibitory matrix (indicated by the orange hexagon), or by (B) proteolytically activating one or several biomineralization promoting proteins. Blue oval: magnetosome membrane protein; purple oval: inner membrane protein; orange symbol: inhibitor or activator of biomineralization; red circle: MamE protease independent function;

red pacman: MamE protease-dependent activity. I.-IV.: proposed steps in magnetosome formation.

**Table 1**

Cmag and MamC-GFP and GFP-MamI localization in complemented  $\Delta R9\Delta mamE$  strains. Magnetic response and percent of cells with continuous linear, wildtype-like localization patterns are reported. n is the number of cells scored for GFP localization. Error is reported as one standard deviation representative of variation between different cultures.

<b>MamC-GFP localization</b>	<b>Cmag</b>	<b>% of cells with linear localization</b>	<b>n</b>
$\Delta R9\Delta mamE$	1.0±0	1±1	259
$\Delta R9\Delta mamE/mamE$	1.52±0.07	85±12	400
$\Delta R9\Delta mamE/mamE^P$	1.0±0	75±7	464
$\Delta R9\Delta mamE/mamE^{C2}$	1.10±0.03	63±6	631
<b>GFP-MamI localization</b>	<b>Cmag</b>	<b>% of cells with linear localization</b>	<b>n</b>
$\Delta R9\Delta mamE$	1.0±0	0±0	523
$\Delta R9\Delta mamE/mamE$	1.5±0.15	82±8	1278
$\Delta R9\Delta mamE/mamE^P$	1.0±0	73±10	1061
$\Delta R9\Delta mamE/mamE^{C2}$	1.07±0.03	63±7	1018