

# *Mycobacterium tuberculosis* FtsB and PerM interact via a C-terminal helix in FtsB to modulate cell division

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## Abstract

Latent infection by *Mycobacterium tuberculosis* (Mtb) impedes effective tuberculosis therapy and eradication. The protein PerM is essential for chronic Mtb infections in mice and acts via the divisome protein FtsB to modulate cell division. Using transgenic co-expression in *Escherichia coli*, we studied the Mtb PerM-FtsB interaction in isolation from other Mtb proteins, engineering PerM to enhance expression in the *E. coli* membrane. We confirmed the reported instability of Mtb FtsB, and we linked FtsB instability to a segment of FtsB predicted to bind cell-division proteins FtsL and FtsQ. Though narrowly conserved, the PerM-FtsB interaction emerges as a potential target for therapy targeting persistent infections by disrupting regulation of cell division. Using fluorescence microscopy, we found that stability of both FtsB and PerM hinges on their interaction via a C-terminal helix in FtsB. Molecular dynamics results supported the observation that FtsB stabilized PerM, and suggested that interactions at the PerM-FtsB interface differ from our initial structure prediction in a way that is consistent with PerM sequence conservation. Integrating protein structure prediction, molecular dynamics and single-molecule microscopy, our approach is primed to screen potential inhibitors of the PerM-FtsB interaction and can be straightforwardly adapted to explore other putative interactions.

**Keywords:** *Mycobacterium tuberculosis*, cell division, single-molecule microscopy, molecular dynamics

## Introduction

*Mycobacterium tuberculosis* (Mtb) is a major contributor to preventable deaths by infectious disease and has infected approximately a quarter of the global population [1]. Most infections progress to latent tuberculosis infection (LTBI), characterized by an immunological response to Mtb antigens without clinical signs of active TB [2]. Preventing establishment and reactivation of LTBI is a growing priority for TB elimination strategies, although uncertainties continue to limit measurements of the relative burdens of new TB infections and LTBI reactivation [3]. Protocols that reduce the duration and adverse effects of treatment can reduce LTBI burden by improving completion rates for preventative treatment [4]. Furthermore, imperfect treatment over long periods can contribute to acquisition of drug resistance [5]. Consequently, it is imperative to better understand how Mtb infections persist to establish LTBI in order

to inform strategies for novel LTBI treatments with higher efficacy, shorter durations, and reduced risk of drug resistance.

A potential strategy to prevent establishment and reactivation of LTBI is to characterize and target mechanisms through which Mtb persists through host-induced stress [6]. The actinomycete protein PerM was one of 21 genes identified in a screen of transposon Mtb mutants with attenuated growth at pH 4.5 in media containing Tween 80, with most of the identified genes being associated with cell wall synthesis [7]. A subsequent study focused on PerM, finding that Mtb PerM knockout reduced growth during chronic mouse infection and increased  $\beta$ -lactam antibiotic susceptibility. Localization of fluorescently tagged PerM to dividing septa suggested a connection to cell division [8]. Further work demonstrated that PerM associates with the Mtb divisome, a protein complex orchestrating cell wall remodeling during

division, and that PerM depletion can be complemented by overexpression of the divisome component FtsB [9]. PerM was also recently identified by transposon sequencing to play an even more significant role in infecting mice in a background with weakened adaptive immune response [10].

Recently, a covalent inhibitor of divisome formation based upon the structure of FtsB-FtsQ was developed and found to be active against drug-resistant *E. coli* in an animal infection model [11]. The Mtb proteome includes homologs of the five core *E. coli* divisome proteins FtsQ, FtsL, FtsB, FtsW, and FtsI [12], suggesting that a similar approach can target protein-protein interactions in the Mtb divisome. However, it is unknown whether PerM directly interacts with FtsB [9] and, if it does, whether PerM has a regulatory role beyond impacting FtsB stability. PerM lacks known orthologs outside of actinomycetes [8], so PerM interactions could potentially be targeted by specific therapies. However, no experimental structural data have been published on the PerM-FtsB interaction or on PerM alone to guide experimental design. Expression and purification of Mtb PerM and FtsB for structural study is likely to pose difficulties given that FtsB stability depends upon PerM co-expression in Mtb and *Mycobacterium smegmatis* [9]. Furthermore, proteins such as PerM with a high number of transmembrane helices pose challenges for recombinant expression [13, 14].

Recent insights into the molecular structure and regulatory mechanisms of the gram-negative divisome have been enabled by the confluence of protein structure prediction, cryogenic electron microscopy (Cryo-EM), and molecular dynamics (MD) simulations. The extension from prediction of monomers to protein complexes [15, 16] facilitated rapid structure prediction of the core *E. coli* divisome [17, 18]. Analysis of a Cryo-EM structure of the *Pseudomonas aeruginosa* divisome was largely consistent with predicted protein-protein interfaces, but also revealed a global conformational change absent in structure predictions [19]. All-atom MD simulations identified a similar conformational change within 1  $\mu$ s, and further predicted interactions between the core divisome and *E. coli* FtsN [20]. However, limitations of this MD approach, such as uncertainty in structure predictions and limited MD timescales, call for experimental validation. Importantly, *in silico* predictions for FtsN were corroborated by experimental results in living cells [21].

In this work, we employed a combination of structure prediction, MD, and fluorescence microscopy to probe the predicted interaction between Mtb PerM and FtsB proteins. In our approach, Mtb PerM and FtsB were expressed *E. coli* in order to directly attribute observations to changes in the Mtb proteins.

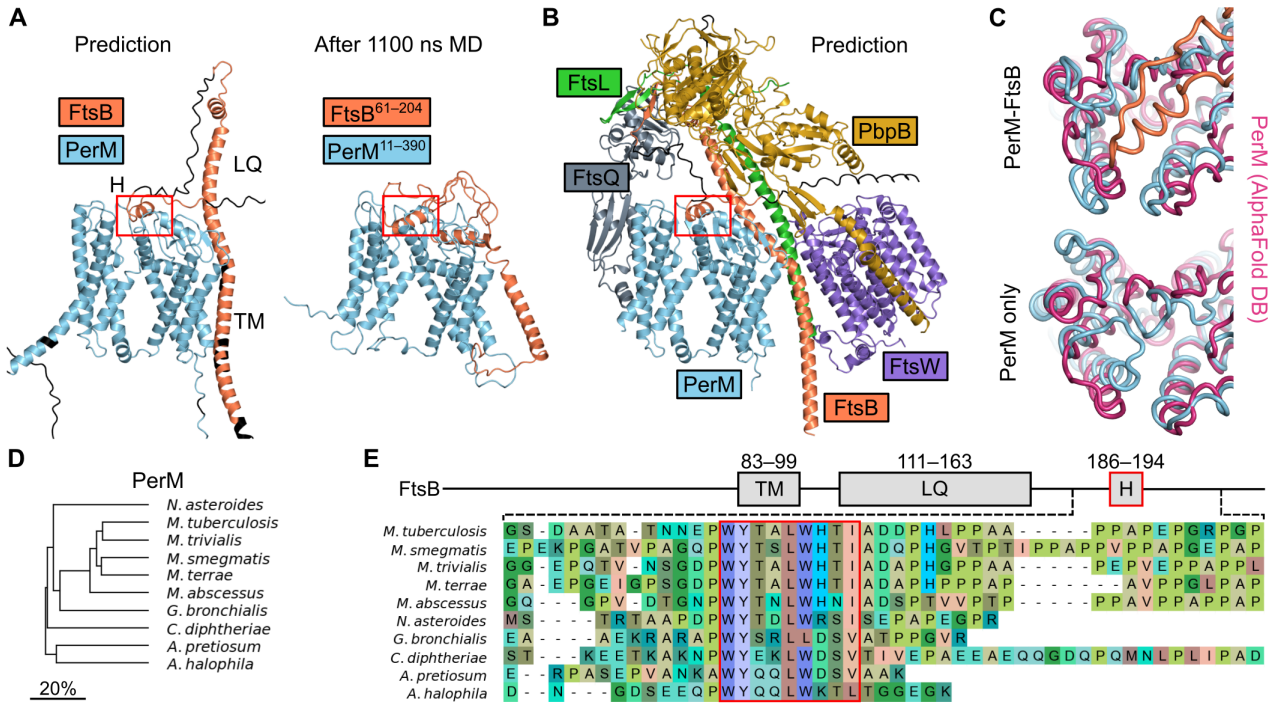
First, we investigated conservation and dynamics at the predicted PerM-FtsB interface, identifying roles for conserved residues that are absent in structure predictions. Second, we showed that FtsB instability (when expressed in *E. coli*) depends on a region predicted to bind FtsL and FtsQ, and that PerM expression in *E. coli* can be enhanced by strategically modifying its N-terminal signal sequence. This enabled the quantification of the PerM-FtsB interaction via fluorescence correlation and single-molecule tracking. We found PerM expression in the *E. coli* membrane to depend on the FtsB sequence predicted to mediate PerM-FtsB interaction. Lastly, we investigated MD simulations of Mtb divisome complexes suggested PerM regulatory complexity beyond FtsB stabilization.

## Results

### Structure prediction and molecular dynamics of PerM-FtsB interaction

In preliminary structure predictions, we identified a high-confidence predicted interaction between Mtb PerM and FtsB ( $pDockQ \approx 0.6$  [22]). The predicted structure of PerM was largely consistent with previous predicted topology [8] except that two regions with hydrophobic residues in predicted transmembrane helices did not span the membrane in structure predictions (Fig. 1A). The topology remains N-in, C-in, with the predicted PerM structure consisting of two halves, each with the same topology (three transmembrane helices with a buried extracellular loop between the first two). Preliminary molecular dynamics (MD) simulations showed that the predicted PerM-FtsB interaction persisted on the microsecond timescale. To test stability of the structure of the predicted complex further, we carried out a three-stage MD protocol with 100 ns of equilibrium MD followed by 500 ns of accelerated molecular dynamics (aMD), and a final 500 ns of equilibrium MD. Terminal residues lacking high local prediction confidence ( $pLDDT < 50$ ) were omitted in constructing the simulation system of FtsB<sup>61–204</sup> and PerM<sup>11–390</sup>. Fig. 1A shows how the predicted extended structure of FtsB collapsed in the absence of interactions with other divisome components [17–20]. Interaction between transmembrane helices of PerM and FtsB was not predicted with high confidence and also did not exhibit persistent, specific interactions in MD. Conversely, the predicted interaction between FtsB and a specific pocket in the periplasmic face of PerM was maintained throughout the aMD protocol and in a simulation replicate.

We also found that a prediction of the core Mtb divisome with the addition of PerM was consistent



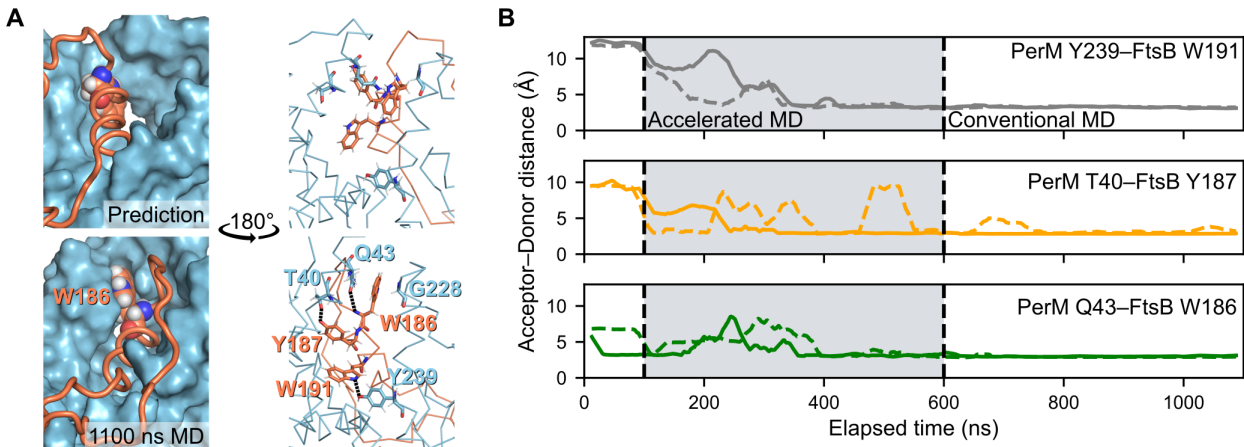
**Fig. 1 A conserved, predicted Mtb PerM-FtsB interaction stabilizes PerM in MD.** (A) Left: predicted PerM-FtsB complex in which an  $\alpha$  helix, FtsB<sup>H</sup>, interacts with the periplasmic face of PerM. Transmembrane helix FtsB<sup>TM</sup> and the region interacting with FtsL and FtsQ, FtsB<sup>LQ</sup>, are indicated. Residues with  $pLDLDT < 50$  colored black. Right: final conformer following 1.1  $\mu$ s MD; PerM-FtsB<sup>H</sup> interact persists, FtsB<sup>LQ</sup> collapses, and FtsB<sup>TM</sup> moves away from PerM. (B) Prediction for the Mtb divisome with inclusion of PerM. Terminal regions with  $pLDLDT < 50$  are omitted except for the FtsB C-terminus. All complexes in A and B are aligned by PerM C $\alpha$  atoms; the red box is placed at the same position relative to PerM for comparison of FtsB<sup>H</sup> movement. (C) Final MD conformers for PerM-FtsB and for PerM alone are aligned by PerM C $\alpha$  atoms and compared to a PerM prediction (AlphaFold DB P9WKN3-F1-model\_v4). (D) Phylogenetic tree for actinomycete species with predicted PerM-FtsB interactions. Branch length scaled by divergence in amino acid identity in pairwise sequence alignments of PerM. (E) Top: Diagram of Mtb FtsB defining the FtsB<sup>TM</sup>, FtsB<sup>LQ</sup>, and FtsB<sup>H</sup> regions. Bottom: Multiple sequence alignment of a region near the C-terminus of FtsB for actinomycete species illustrates conservation of residues in FtsB<sup>H</sup> and diversity in FtsB C-termini.

with PerM interacting with the core divisome without obviously disrupting interactions between core divisome components (Fig. 1B). Within this complex, PerM was only predicted to interact with FtsB. Based on these predictions, we identified three regions of interest in Mtb FtsB: the predicted transmembrane helix, FtsB<sup>TM</sup> (FtsB<sup>83–99</sup>), the region predicted to interact with FtsL and FtsQ, FtsB<sup>LQ</sup> (FtsB<sup>111–163</sup>), and a span of hydrophobic residues forming an  $\alpha$  helix predicted to interact with PerM, FtsB<sup>H</sup> (FtsB<sup>186–194</sup>). Although C-terminal residues following FtsB<sup>H</sup> were predicted to thread between the PpbB anchor and head domains, there were no high-confidence predictions for this region.

In order to see the impact of FtsB on PerM, we replicated the three-stage MD protocol using PerM alone. Final conformers for PerM-FtsB and PerM simulations are shown in Fig. 1C and compared to a recent PerM structure prediction from the AlphaFold Protein Structure Database [23]. Surprisingly, we found that the final conformer in the monomeric PerM simulation differed from the predicted PerM structure

( $RMSD = 3.47 \text{ \AA}$ ) to a greater extent than final conformers in PerM-FtsB simulation replicates ( $2.50 \text{ \AA}$  and  $2.21 \text{ \AA}$ ). The largest structural changes were seen in the predicted FtsB<sup>H</sup> binding pocket.

Since our structure predictions were informed by information encoded in multiple sequence alignments, we searched for and identified PerM orthologs in actinomycete species (Fig. 1D). For each, we also identified the corresponding FtsB sequence and predicted the structures of PerM-FtsB complexes. In every species tested there was a predicted interaction between PerM and FtsB<sup>H</sup> with the same orientation of FtsB<sup>H</sup> on the periplasmic face of PerM (Supplementary Fig. 1). Further, there is little difference in the lengths of regions linking FtsB<sup>LQ</sup> and FtsB<sup>H</sup>. PerM-FtsB complexes differed, however, in the predicted position of FtsB<sup>TM</sup> relative to PerM. We constructed a multiple sequence alignment of FtsB to investigate the basis for conserved, predicted PerM-FtsB interaction and found that FtsB<sup>H</sup> was highly conserved (Fig. 1E) with near perfect conservation of hydrophobic residues predicted to interact with PerM (W186, Y187, L190, and W191).



**Fig. 2 Reproducible dynamics at the PerM-FtsB binding interface.** (A) The shape of the FtsB binding pocket for PerM is compared between the predicted interface structure (top) and the final conformer following 1.1  $\mu$ s MD (bottom). Left: a large predicted periplasmic binding pocket in PerM (top left) evolves to tightly bind FtsB W186 (bottom left). Right: Following MD (bottom right), FtsB W186 interacts with conserved PerM residue G228 and hydrogen bonds are formed between PerM and conserved FtsB residues. The same region is shown rotated by 180°. (B) Maturation of PerM-FtsB interactions were reproducible in MD. Solid and dashed lines show acceptor-donor distance (25 ns moving average; two MD replicates) for hydrogen bonds at the PerM-FtsB interface that are absent in the predicted structure.

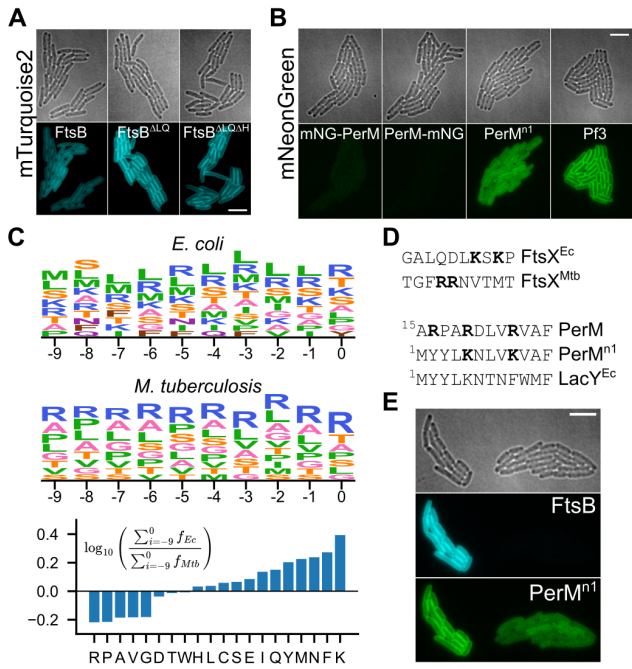
Next, we investigated the structural basis for specific PerM-FtsB interaction. Fig. 2 illustrates the evolution of the FtsB binding pocket in PerM during MD. In both simulation replicates, we observed that a large, hydrophobic binding pocket tightened around FtsB<sup>H</sup>, and around FtsB W186 in particular. Residues in the W186 binding pocket include PerM G228, which was perfectly conserved in PerM sequences in species that we analyzed. Remarkably, specific hydrogen bonds between PerM and conserved FtsW residues emerged, none of which were present in structure predictions. Fig. 1B shows that evolution of the PerM-FtsB binding interface was reproducible.

### Transgenic expression of fluorescent FtsB and PerM constructs in *E. coli*

Despite the apparent specificity and stability of the predicted PerM-FtsB interface in MD, we wanted to confirm that this interaction does not depend on other Mtb components to be stable on timescales inaccessible by MD and, if so, to investigate whether FtsB<sup>H</sup> mediated the interaction. We constructed variants of FtsB with an N-terminal fusion of mTurquoise2 (mTq2-FtsB, mTq2-FtsB<sup>ΔLQ</sup>, and mTq2-FtsB<sup>ΔLQΔH</sup>; Fig. 3A). Deletion of FtsB<sup>LQ</sup> clearly increased the FtsB expression level, suggesting instability of Mtb FtsB in the absence of FtsL and FtsQ. However, FtsB instability in Mtb may result from other mechanisms [9], so we did not investigate this in depth. We did not observe any further effect on mTurquoise2 fluorescence when deleting FtsB<sup>H</sup> in addition to FtsB<sup>LQ</sup>.

Conversely, initial attempts to construct mNeonGreen-labeled PerM failed for both N- and C-terminal fusion constructs (mNG-PerM and PerM-mNG; Fig. 3B). We manually inspected the predicted PerM structure as well as sequences of integral transmembrane protein orthologs in *E. coli* and Mtb, hypothesizing that an engineered construct, PerM<sup>n1</sup>, would exhibit higher expression. Specifically, we noticed an apparent “LK” motif in *E. coli* that rarely appeared in Mtb and was reminiscent of part of the twin-arginine translocation motif [24]. Our strategy in designing the PerM<sup>n1</sup> construct was to modify PerM with aspects of the short LacY N-terminal sequence in ways that would not perturb the predicted PerM structure. While expression was indeed much higher for PerM<sup>n1</sup>-mNG, it failed to exhibit the same degree of membrane localization as Pf3-mNG, which we used as a control given its efficient translocation and uniform transmembrane orientation [25].

We were surprised at how much the modifications in PerM<sup>n1</sup> increased expression levels and systematically investigated sequences in residues preceding the first predicted transmembrane helix in all proteins including at least one transmembrane helix in the *E. coli* and Mtb proteomes (Fig. 3C). We found large differences in the frequencies of some residues, most strikingly for differences in the relative frequencies of lysine and arginine. Fig. 3D shows how this is reflected in orthologous sequences of FtsX as well as in PerM<sup>n1</sup> compared to Mtb PerM. Finally, we proceeded to co-express PerM<sup>n1</sup>-mNG and mTq2-FtsB and look for evidence of PerM-FtsB interaction. Serendipitously, one of two adjacent *E. coli* micro-colonies in a co-expression experiment lost the plasmid



**Fig. 3 Expression of Mtb FtsB and PerM in *E. coli*.** (A) Expression and membrane localization of FtsB and variants with deletions of FtsB<sup>LQ</sup> or with deletion of both FtsB<sup>LQ</sup> and FtsB<sup>H</sup>. Identical minimum and maximum intensities used to produce figures. (B) Expression and membrane localization for mNG-PerM, PerM-mNG, PerM<sup>n1</sup>-mNG, and Pf3-mNG. Identical minimum and maximum intensities. (C) Top: residue frequency for positions preceding the first predicted transmembrane helix, inclusive of residues found at frequencies above 5%. Letter height is proportional to residue frequency with more frequent residues on top. Bottom: relative frequencies in *E. coli* and Mtb. (D) Top: residues preceding the first transmembrane helix in paralogs of FtsX, highlighting lysine and arginine residues. Bottom: comparison of PerM to PerM<sup>n1</sup> and *E. coli* LacY. (E) Two adjacent microcolonies from a strain with co-expression of mTq2-FtsB and PerM<sup>n1</sup>-mNG. The microcolony on the right has coincidentally lost the plasmid encoding mTq2-FtsB, and exhibits reduced membrane localization of PerM<sup>n1</sup>-mNG. Scale bars 5 μm.

for mTq2-FtsB expression (this occasionally happened without including selective antibiotics in agarose gel pads; we note that we used a YFP filter set for mNeon-Green imaging to avoid crosstalk from mTurquoise2). The large difference in PerM<sup>n1</sup>-mNG membrane localization between these two microcolonies suggested an unexpected role for FtsB in stabilizing PerM in the *E. coli* membrane.

### PerM stabilization in *E. coli* membrane by FtsB depends on FtsB<sup>H</sup>

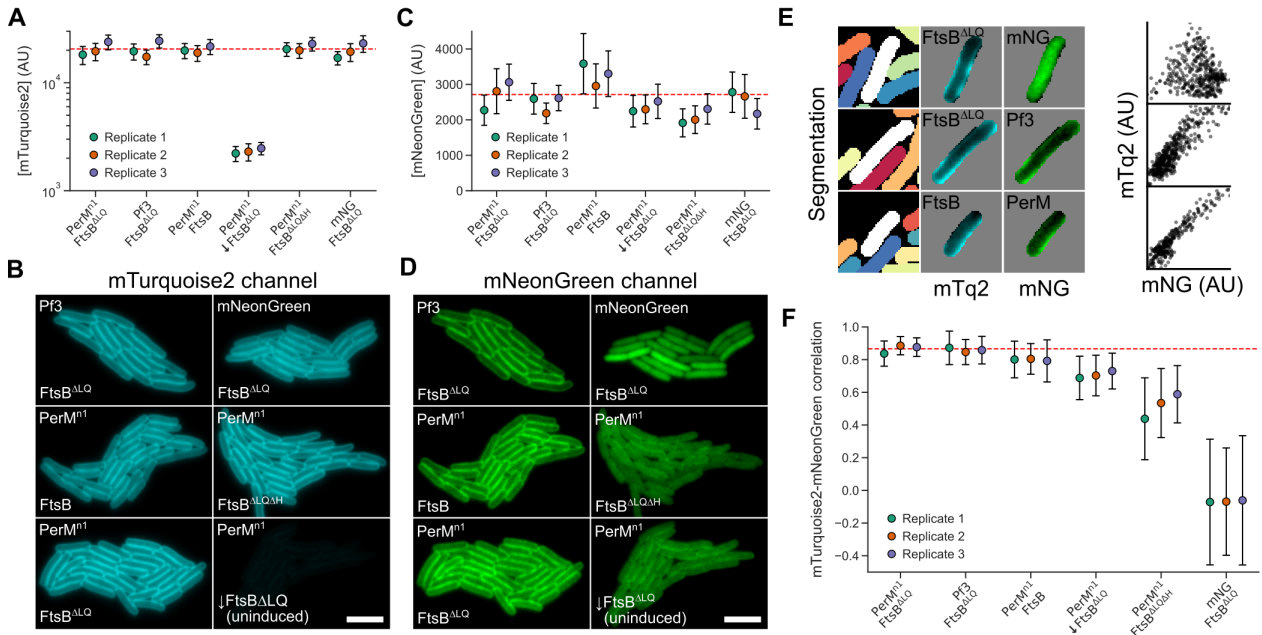
In order to confirm that FtsB increases membrane-localized PerM expression and, if so, to investigate the role of FtsB<sup>H</sup> in this phenomenon, we co-expressed mNeonGreen- and mTurquoise2-labeled constructs and analyzed fluorescence in *E. coli* microcolonies

(Fig. 4). In these experiments, expression of PerM<sup>n1</sup>-mNG was induced with 100 μM IPTG. Localization controls mNeonGreen (strong cytoplasmic localization) and Pf3-mNG (strong membrane localization) were induced with 60 μM and 240 μM IPTG, respectively, to match typical PerM<sup>n1</sup>-mNG expression levels. Expression of mTurquoise2-labeled FtsB constructs was induced with 10 nM ATc in all conditions except for one condition with no induction (indicated with down arrows in Fig. 4). Brightfield images of *E. coli* cells were segmented, and protein concentration was estimated to be proportional to total integrated fluorescence after subtracting background and normalizing by cell size.

Relative to the reference condition (PerM<sup>n1</sup> and FtsB<sup>ΔLQ</sup>, 10 nM ATc), there was an 89% ( $p = 3.3 \times 10^{-12}$ ) decrease in mTq2-FtsB<sup>ΔLQ</sup> expression in the absence of induction, with no significant differences observed for other conditions. The absence of any significant impact on wild-type FtsB expression ( $p = 0.97$ ) compared to when FtsB was expressed in the absence of PerM (compare Fig. 4B to Fig. 3A) suggested that PerM<sup>n1</sup> stabilized FtsB when expressed in *E. coli*, as was observed in Mtb [9]. There was also no obvious difference in localization of any FtsB variant in any condition tested.

In contrast, there was a significant reduction in PerM<sup>n1</sup>-mNG expression of 13.3% ( $p = 3.8 \times 10^{-2}$ ) in the absence of induction of mTq2-FtsB<sup>ΔLQ</sup> expression, and a reduction of 23.5% ( $p = 5.5 \times 10^{-4}$ ) when FtsB<sup>ΔLQ</sup> was replaced with FtsB<sup>ΔLQΔH</sup>. There was no significant change when FtsB<sup>ΔLQ</sup> was replaced with wild-type FtsB ( $p = 0.29$ ). This suggested that FtsB increased PerM stability in *E. coli*, and that this depended on the presence of FtsB<sup>H</sup>. However, inspection of Fig. 4C suggests that our sample size is overpowered ( $N = 10930$  cells in total from 3 replicates;  $607 \pm 44$  cells per condition per replicate) given variation between replicates in mNeonGreen expression levels.

In preliminary experiments, we observed a large decrease in spatial correlation between mTurquoise2 and mNeonGreen fluorescence when FtsB<sup>ΔLQ</sup> was depleted or replaced by FtsB<sup>ΔLQΔH</sup> (Fig. 4D). The observation of small reductions in PerM<sup>n1</sup>-mNG levels despite apparently large impacts of different FtsB constructs on localization suggested that fluorescent mNeonGreen often remained in the cytoplasm following PerM<sup>n1</sup>-mNG degradation. We hypothesized that quantifying spatial correlation would be a more sensitive measurement less susceptible to variation in protein expression levels. Fig. 4E shows scatter plots of mNeonGreen and mTurquoise2 intensities for pixels in typical cells with cytoplasmic mNeonGreen, membrane-localized Pf3-mNG, and PerM<sup>n1</sup>-mNG. We calculated Spearman correlation coefficients from such



**Fig. 4** FtsB stabilizes PerM in the *E. coli* membrane. **(A)** Distribution of single-cell mTurquoise2 concentration (mean  $\pm$  standard deviation) for FtsB constructs in different conditions, each with replicates. Dashed red line indicates the mean value for the reference condition (PerM<sup>n1</sup>-mNG, mTq2-FtsB<sup>ΔLQ</sup>, 100  $\mu$ M IPTG, 10 nM ATC). Uninduced FtsB<sup>ΔLQ</sup> is indicated with a down arrow. A logarithmic scale is used to allow comparison of high- and low-expression conditions. **(B)** Example data from microcolonies for each condition. The uninduced FtsB<sup>ΔLQ</sup> condition has very low fluorescence in example data because identical minimum and maximum intensities were used. Scale bar 5  $\mu$ m. **(C, D)** Distribution of single-cell mNeonGreen concentrations and example data for Pf3-mNG, mNeonGreen alone, or PerM<sup>n1</sup>-mNG in different conditions; prepared identically to **A** and **B**. **(E)** Left: example data analysis showing cell segmentation and isolated, single-cell intensities. Right: Scatter plots of single-pixel intensities for each cell show clear correlation for Pf3-mNG and PerM-mNG, but not for mNeonGreen alone. **(F)** Distribution of single-cell Spearman correlation coefficients (mean  $\pm$  standard deviation) for three replicates. Conditions ordered by decreasing mean correlation.

distributions for segmented cells and Fig. 4F shows the mean and standard deviation of correlation coefficients for each condition and replicate.

In the reference condition (PerM<sup>n1</sup>-mNG, mTq2-FtsB<sup>ΔLQ</sup>, 10 nM ATC), correlation was high ( $\rho = 0.87$ , 0.84–0.89 95% confidence interval) and indistinguishable from that when Pf3-mNG was expressed instead ( $p = 0.72$ ). All other conditions had significant reductions in correlation. Wild-type FtsB was marginally less well correlated ( $\rho = 0.80$ , 0.77–0.83). The apparent reductions in PerM<sup>n1</sup>-mNG expression levels discussed above were more strongly supported by analysis of spatial correlation, with a drop in correlation in the absence of mTq2-FtsB<sup>ΔLQ</sup> induction ( $\rho = 0.71$ , 0.69–0.73) and a larger drop when mTq2-FtsB<sup>ΔLQ</sup> was replaced with mTq2-FtsB<sup>ΔLQΔH</sup> ( $\rho = 0.52$ , 0.46–0.58). While FtsB<sup>LQ</sup> was not absolutely essential for membrane localization of PerM<sup>n1</sup>, this was also observed in the absence of any FtsB expression (Fig. 3B).

## Single-molecule FtsB tracking reveals that FtsB<sup>H</sup>-dependent PerM binding

Our correlation analysis strongly suggested that FtsB interaction with PerM via FtsB<sup>H</sup> is key for expression of PerM<sup>n1</sup> in the membrane. Since we observed

this for transgenic expression in *E. coli*, it is unlikely that PerM-FtsB interaction is mediated by host proteins. However, transient PerM-FtsB interaction could be sufficient for PerM stability without a large fraction of molecules being bound at any time. If PerM is a significant component in the core Mtb divisome as suggested by structure prediction, depletion phenotypes, and midcell localization [8, 9], we hypothesized that this implied long-lived PerM-FtsB interactions that could be detected by single-molecule tracking of FtsB diffusion. FtsB has a single transmembrane helix, while PerM is predicted to have 8 (Fig. 1A); based on the dependence of diffusion of integral membrane proteins on the number of transmembrane helices, we expected a reduction in the FtsB diffusion coefficient from approximately 0.7 to 0.3  $\mu\text{m}^2 \text{s}^{-1}$  upon PerM binding [26].

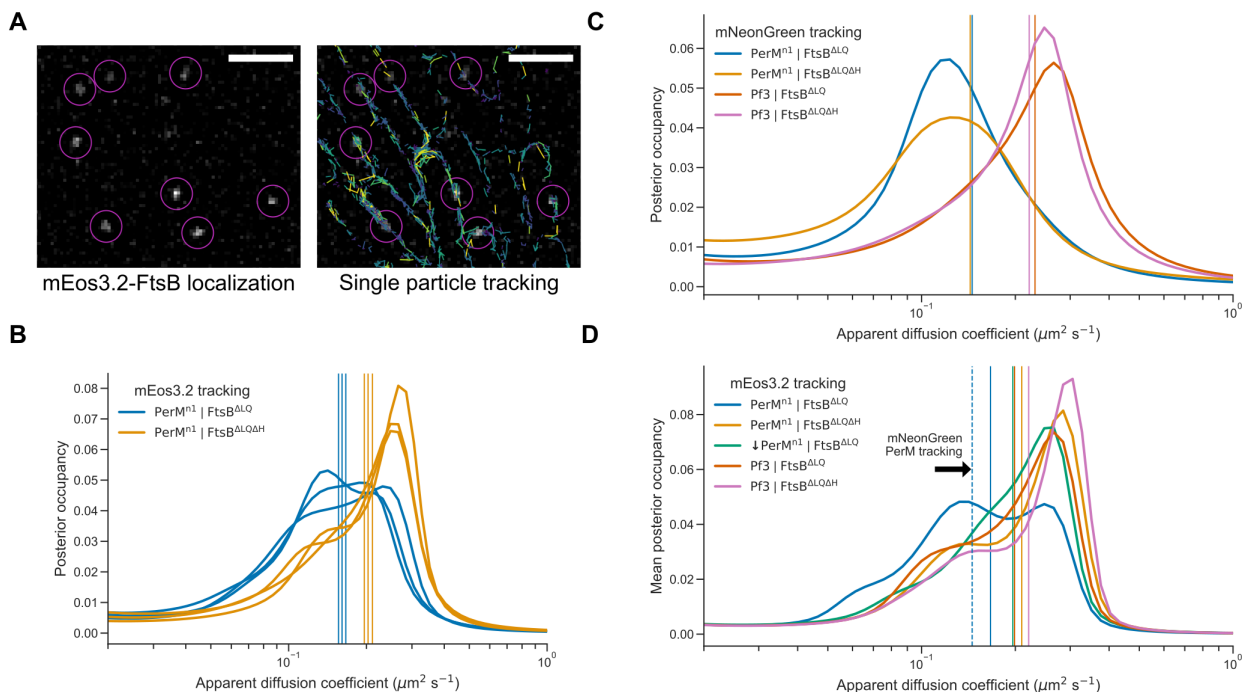
We constructed mEos3.2-tagged variants of FtsB<sup>ΔLQ</sup> and FtsB<sup>ΔLQΔH</sup>, and co-expressed them with either PerM<sup>n1</sup>-mNG or Pf3-mNG. Fig. 5A shows a typical frame from single-molecule imaging data, localized mEos3.2 molecules, and single-molecule tracks gathered in one movie showing that tracked molecules were detected near the middle plane of *E. coli* cells (approximately 0.5  $\mu$ m from the microscope coverslip). In analysis of preliminary data, we

found that analysis of either mean squared displacement or fitting jump-length distributions was sensitive to variable trajectory length. Thus, we utilized SASPT [27] to infer distributions of *apparent* diffusion coefficients using a method robust against short trajectories, variable localization error, and defocalization [28]. Note that we refer to an *apparent* diffusion coefficient, as diffusion within our images is largely confined to the one-dimensional perimeter at the middle plane and, further, confined at the cell poles. So, while this is sufficient to compare relative diffusion rates in different conditions, precise estimates of diffusion coefficients will require more sophisticated methods and would benefit from 3D tracking over the entire membrane.

In tracking experiments, mEos3.2-FtsB constructs were expressed without induction so that PerM would be in excess and maximize potential binding. We inferred distributions of diffusion coefficients after marginalizing out localization error for three independent replicates (Fig. 5B). We observed reproducible differences in mean diffusion coefficients between FtsB<sup>ΔLQ</sup> ( $0.161 \pm 0.004 \mu\text{m}^2 \text{s}^{-1}$ ; mean  $\pm$  standard deviation) and FtsB<sup>ΔLQΔH</sup> ( $0.203 \pm 0.006 \mu\text{m}^2 \text{s}^{-1}$ ;

mean  $\pm$  standard deviation). Distributions for both FtsB<sup>ΔLQ</sup> and FtsB<sup>ΔLQΔH</sup> suggested the presence of a slow-diffusion population at approximately  $0.14 \mu\text{m}^2 \text{s}^{-1}$  that is more prominent for FtsB<sup>ΔLQ</sup>. We also collected single-molecule tracking data for PerM<sup>n1</sup>-mNG and Pf3-mNG in single experiments. Despite limitations of this data (fewer localizations and variable spot density as mNeonGreen photo-bleaches over time), we inferred average diffusion coefficients of  $0.144 \mu\text{m}^2 \text{s}^{-1}$  and  $0.226 \mu\text{m}^2 \text{s}^{-1}$  for mNG-PerM and mNG-Pf3, respectively (Fig. 5C).

To estimate the most likely apparent diffusion coefficients, we inferred distributions from all data from each condition (Fig. 5D). This confirmed that mEos3.2-FtsB<sup>ΔLQ</sup>, when expressed with PerM, clearly exhibits slower diffusion than when PerM is replaced with Pf3 or in the absence of FtsB<sup>H</sup>. Peaks at approximately  $0.14$  and  $0.25 \mu\text{m}^2 \text{s}^{-1}$  are consistent with a substantial decrease in the rate of diffusion upon PerM binding. Out of all other conditions, mEos3.2-FtsB<sup>ΔLQ</sup> in the absence of induction of PerM<sup>n1</sup>-mNG exhibited the slowest diffusion (green line in Fig. 5D). However, the effect size was clearly



**Fig. 5 PerM-FtsB binding detected by FtsB single-molecule tracking.** (A) Left: single, 33-ms frame from a fluorescence microscopy movie of mEos3.2-FtsB<sup>ΔLQ</sup> diffusion in *E. coli*. Right: Single-molecule localization and tracking results for the entire movie show membrane-localized diffusion. Scale bar 2  $\mu\text{m}$ . (B) Posterior occupancies in a model of regular Brownian motion and localization error were marginalized over localization error to estimate the distribution of apparent 2D diffusion coefficients. Distributions and their means (vertical lines) are shown for three replicates of experiments combining PerM<sup>n1</sup>-mNG with either mEos3.2-FtsB<sup>ΔLQ</sup> or mEos3.2-FtsB<sup>ΔLQΔH</sup>. (C) Estimated distributions of diffusion coefficients inferred from single-molecule tracking of either PerM<sup>n1</sup>-mNG or Pf3-mNG with either FtsB<sup>ΔLQ</sup> or FtsB<sup>ΔLQΔH</sup> (single experiment). Vertical lines indicate the mean estimated diffusion coefficient for each condition. (D) Estimated distribution of diffusion coefficients for either mEos3.2-FtsB<sup>ΔLQ</sup> or mEos3.2-FtsB<sup>ΔLQΔH</sup> co-expressed with either Pf3-mNG or PerM<sup>n1</sup>-mNG (with or without addition of 100  $\mu\text{M}$  IPTG), inferred from combining data from three replicates. Vertical lines indicate mean estimated diffusion coefficients and the dashed vertical line is replicated from C.

low and further work is required to determine the concentration dependence of this interaction.

## PerM-FtsB interaction could restrict conformational flexibility of the Mtb divisome

Our observation that FtsB<sup>H</sup>-dependent PerM-FtsB interaction was sufficiently long-lived to produce large changes in FtsB diffusion prompted us to explore how this interaction could impact the Mtb divisome. We conducted MD simulations to compare divisome constructs in which FtsB was truncated to FtsB<sup>205</sup> (removing the uncertain C-terminal residues) or truncated to FtsB<sup>185</sup> (additionally remove FtsB<sup>H</sup>). First, we performed MD for the Mtb divisome with PerM and full-length FtsB for 500 ns. FtsB N-terminal residues and other terminal residues of subunits without confident structure predictions were omitted. A period of initial MD was required since our structure prediction did not have all subunits at ideal relative orientations; this was especially significant for FtsQ since its transmembrane helix is distant from others. The final conformer after 500 ns was used to build systems with truncation to FtsB<sup>205</sup> (FtsB<sup>ΔC</sup>) and to FtsB<sup>185</sup> (FtsB<sup>ΔCΔH</sup>), and each system was simulated for 1 μs. The final conformer for FtsB<sup>ΔC</sup> is shown in Fig. 6A.

Comparing the final conformers of the FtsB<sup>ΔC</sup> and FtsB<sup>ΔCΔH</sup> simulations, we examined the angle formed as the vector between the putative active site residues FtsW<sup>D311</sup> and PbpB<sup>S386</sup> moves during MD simulation (Fig. 6B). This angle is a proxy for the tilt of the PbpB transpeptidase domain relative to FtsW that has been used to analyze Cryo-EM data as well as MD simulation. We observed a 19.2° tilt for FtsB<sup>ΔC</sup>, similar to that observed for *P. aeruginosa* FtsI comparing a structure prediction to its Cryo-EM structure [19] and for *E. coli* comparing a structure prediction to conformers following MD [20]. Conversely, with the loss of PerM-FtsB interaction mediated by FtsB<sup>H</sup>, the tilt angle reverted to a value within only 4.8° of the predicted structure. The evolution of these angles over the course of MD is shown in Fig. 6C.

Relative to studies on the divisome in model organisms, there is a paucity of experimental data to relate our simulations to phenotypes associated with mutations to Mtb divisome components. Thus, we did not analyze interactions between Mtb divisome sub-units in depth or conformations in and around FtsW and PbpB active sites. However, we note substantial differences the relative conformations of divisome components in final conformers for the FtsB<sup>ΔC</sup> and FtsB<sup>ΔCΔH</sup> simulations; tilt of the PbpB transpeptidase domain was associated with twisting of the PbpB head domain, FtsL, and FtsQ relative to FtsW. Lastly,

in Fig. 6D we return to the final conformer of a PerM-FtsB simulation following aMD to highlight residues of interest in PerM that are suggestive of potential regulatory mechanisms. We also show how a phosphatidylglycerol molecule occupies a channel in PerM and interacts with FtsB<sup>H</sup>.

## Discussion

With our integrative approach combining fluorescence microscopy with molecular dynamics starting from predicted structures of protein complexes, we showed that FtsB<sup>H</sup> directly mediates PerM-FtsB binding. Our results are suggestive of significance of PerM-FtsB interaction, going beyond FtsB stabilization [9] to potentially impacting the structure of the core Mtb divisome. Given the role of PerM in persistent Mtb infection and that it is essential in *M. smegmatis*, it will be interesting to learn whether FtsB also stabilizes PerM in Mtb, as it does in *E. coli*, and whether the PerM-FtsB interaction that we have identified can be targeted to disrupt regulation of cell division.

Our approach is straightforward to apply to other predicted protein-protein interactions. For example, Mtb PerM is located in the same operon as Rv0954 which similarly localizes to the Mtb divisome yet has no reported deletion phenotype [29]. Intriguingly, while PerM is associated with the divisome component FtsB, Rv0954 exhibited physical and genetic interactions with elongasome components such as RodA and PpbA. However, division and elongation machinery in mycobacteria are not perfectly analogous to those in other bacteria [30]. The cell-stress phenotypes of PerM depletion in Mtb suggest that conditional phenotypes may be identified for Rv0954 as well.

In Fig. 6D, we highlight PerM residues that could play roles in integrating environmental stress in proximity to FtsB<sup>H</sup>. These include buried histidines that are intriguing with respect to the Mg<sup>2+</sup>-dependent phenotype of PerM, suggesting sensitivity to nutrient depletion [8], methionine residues that make hydrophobic interactions with FtsB<sup>H</sup> and could be sensitive to oxidation, and histidine residues near the membrane surface that could sensitize PerM-FtsB interaction to pH. Furthermore, in MD we routinely observed lipids to penetrate the first half of PerM and come into contact with FtsB, suggesting the possibility that bilayer composition can directly impact PerM-FtsB interaction.

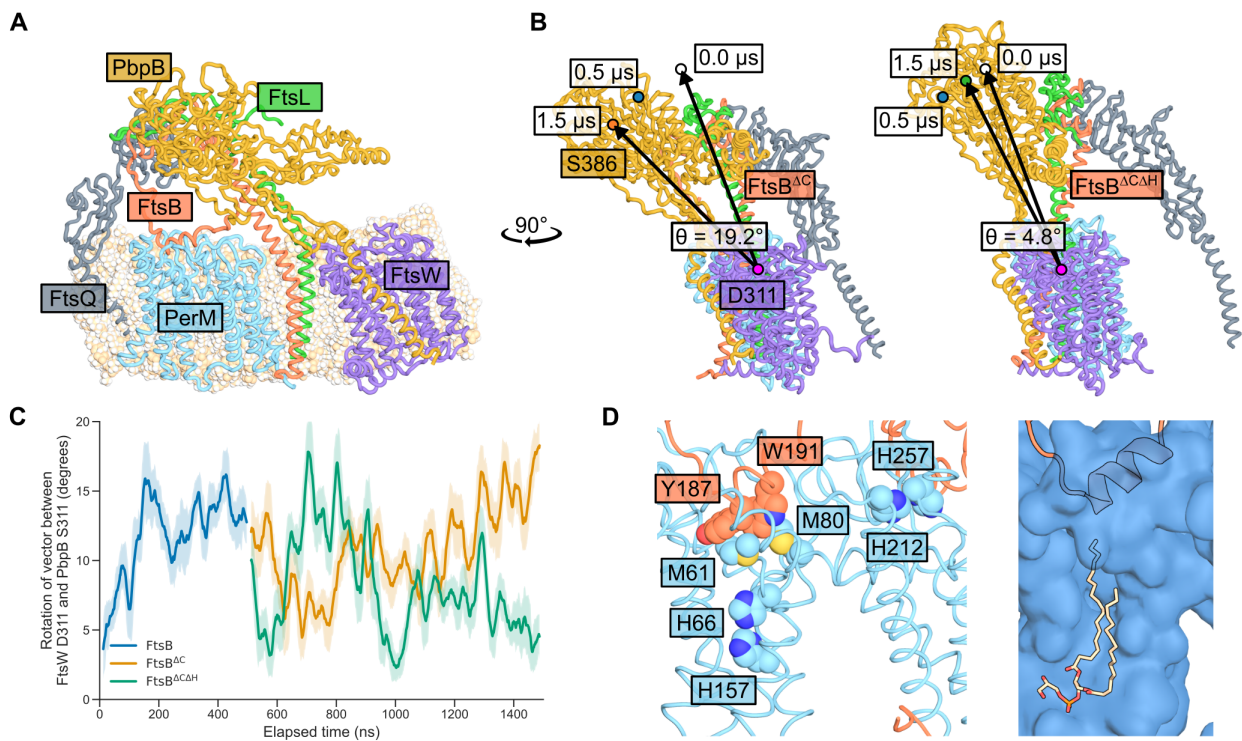
In our experimental system, transgenic expression of Mtb PerM and FtsB in *E. coli* makes it likely that we observed effects of direct PerM-FtsB interaction. This is an advantage of our system as it suggests that PerM-FtsB binding does not depend on phosphorylation or other post-translational modification specific to mycobacteria. While our single-molecule

tracking experiments are not easily extensible to high-throughput investigation, PerM-FtsB and other Mtb protein-protein interactions can be explored by adopting our approach to develop FRET interaction reporters based on structures of predicted complexes. It will also be interesting to see whether our anecdotal success in engineering improved membrane protein expression in *E. coli* is applicable to other Mtb membrane proteins.

Our approach also carries the inevitable limitations of exploring an interaction in a surrogate system. It is critical to not extrapolate too much from our experiments before confirming results in Mtb. Our models of the Mtb divisome make countless predictions that are testable given the variety of tools for manipulating mycobacterial gene expression and recombineering. For example, our MD simulations omitted terminal residues in divisome components that are likely functional in some cases. This is only one example of interactions that we did not predict, but also cannot rule out. As another example, a specific FtsB-PerM interface was not predicted and did

not emerge in MD, but chimeras replacing FtsB<sup>TM</sup> with an alternative transmembrane helix could determine whether it contributes to PerM binding. One prediction arising from our work is that overexpression of FtsB<sup>H</sup> in the Mtb or *M. smegmatis* periplasm could produce PerM-depletion phenotypes [9] via competitive inhibition of PerM-FtsB interaction.

Lastly, our observation that PerM-FtsB interaction constrains PpbB in a tilted conformation can be considered in light of results and discussion emerging from the Cryo-EM structure of the *P. aeruginosa* divisome [19], where an elongated conformation was considered to potentially reflect the catalytically active state. Recent work combining *in vitro* single-molecule and peptidoglycan polymerization assays with characterization of *in vivo* phenotypes supported this model for the gram-negative elongasome [31]. Within the context of this model and in light of our results, the role of PerM would be to reduce divisome activity by constraining PpbB in a conformation with relatively low transpeptidase activity. This is an apparent paradox given that PerM is essential in some conditions in



**Fig. 6 PerM-FtsB interaction constrains the Mtb divisome in MD simulations.** (A) PerM simulated in context of Mtb divisome. Final conformer after 0.5  $\mu$ s of MD with full-length FtsB followed by 1  $\mu$ s of MD with FtsB<sup>ΔC</sup>. Note C-terminal truncation of FtsB relative to the predicted structure in Fig. 1A that is uncertain in this region. (B) Difference in tilt of PpbB transpeptidase domain relative to its conformation in structure prediction. Final conformers for FtsB<sup>ΔC</sup> and FtsB<sup>ΔCΔH</sup> simulations are shown aligned to FtsW and points show locations of putative active-site residues FtsW<sup>D311</sup> and PpbB<sup>S386</sup>. Points at 0  $\mu$ s and 0.5  $\mu$ s are the initial and final positions of PpbB<sup>S386</sup> in initial simulation using full-length FtsB and are plotted at the same position relative to FtsW. (C) Dynamics of tilt angles defined in B show that the FtsB<sup>ΔCΔH</sup> simulation returned towards the elongated conformation observed in structure predictions. A 25 ns moving average of the tilt angle is shown and shaded areas indicate one standard deviation above and below the average angle. (D) Final conformer of PerM-FtsB after 1.1  $\mu$ s MD. Left: residues in PerM near FtsB-binding interfaces suggest that PerM could potentially be sensitive to environmental cues. Right: a lipid often fills a channel in PerM in MD; here, in the final conformer of the FtsB<sup>ΔC</sup> simulation, a phosphatidylglycerol molecule interacts with FtsB<sup>H</sup>.

**Table 1** GenPept accession numbers for PerM and FtsB sequences used for different actinomycete species.

Species	PerM accession	FtsB accession
<i>Mycobacterium tuberculosis</i>	NP_215470	NP_215540
<i>Mycolicibacterium smegmatis</i>	WP_011730590	WP_029104417
<i>Mycolicibacillus trivialis</i>	WP_085109558	WP_085109605
<i>Mycolicibacter terrae</i>	WP_085262653	WP_085262505
<i>Mycobacteroides abscessus</i>	WP_012296377	WP_005066469
<i>Nocardia asteroides</i>	WP_022566234	WP_019049142
<i>Gordonia bronchialis</i>	WP_012833219	WP_041919790
<i>Corynebacterium diphtheriae</i>	WP_014318836	WP_021334958
<i>Actinosynnema pretiosum</i>	WP_157767928	WP_253846810
<i>Actinopolyspora halophil</i>	WP_026152723	WP_017976585

Mtb and in *M. smegmatis*. However, the paradox can be resolved if PerM plays both a role in promoting FtsB stability (which can be bypassed with FtsB over-expression [9]) and also a conditionally essential role in regulating divisome activity that could contribute to persistent Mtb infection.

## Methods

### Structure prediction

PerM and FtsB sequences used for structure prediction are listed in Table 1 and additional Mtb sequences used for complex structure prediction are listed in Table 2. FtsB and PerM were identified in various actinomycete species and, after multiple sequence alignment, have between 19 % and 61 % (PerM) and 27 % and 62 % (FtsB) amino acid identity with Mtb orthologs. Protein complex structures were predicted using ColabFold [32]. Multiple sequences alignments were provided by MMseqs2 [33] using both paired and unpaired sequences in reference and environmental sequence databases. AlphaFold-Multimer [16] (Version 3, Model 4, 12 recycles) was used for structures compared in Fig. 1 and Supplementary Fig. 1 and for structure predictions utilized to build MD systems. PyMOL [34] was used for all structure visualizations and its align function was used to align FtsW structures and to calculate RMSD (using cycles=0).

### Sequence analysis

To compare PerM and FtsB for different species, multiple sequence alignments were generated using

Clustal Omega [35] and visualized using Biotite [36]. To compare amino acid frequencies at transmembrane helices, we used DeepTMHMM [37] to predict the positions of the first transmembrane helix for all proteins in reference proteomes for *E. coli* and Mtb (UniProt ID UP000000625 and UP000001584). Amino acid frequencies and corresponding sequence logos for the 10 residues preceding the first residue in the first predicted transmembrane helix were calculated and plotted using Logomaker [38].

### Molecular dynamics

Molecular dynamics systems were constructed from structures of predicted complexes (or from the final conformer of the initial 500 ns divisome complex simulation) with the CHARMM-GUI Membrane Builder [39]. Shared simulation parameters chosen in CHARMM-GUI were: 150 mM KCl, CHARMM36m forcefield [40], 310 K temperature, NPT ensemble, and capping all non-native N- and C-terminal residues with ACE and CT3, respectively. Scripts for OpenMM [41] output by CHARMM-GUI were modified and OpenMM was used for MD simulation. Simulation system sizes are detailed in Table 3.

Simulation systems differed in a few ways (other than small differences in residues included, shown in Table 4). First, bilayers simulations for the PerM-FtsB complex and PerM alone included a 3:1 POPE:POPG ratio commonly used for simulations of *E. coli* membranes since MD results were compared to *E. coli* microscopy data. Simulations for divisome complexes

**Table 3** Number of total atoms and numbers and types of lipid residues in MD systems.

System	Atoms	Residues	
		POPE	POPG
PerM-FtsB	156168	210	70
PerM only	105155	213	71
Complex (FtsB)	724208	1246	—
Complex (FtsB <sup>ΔC</sup> )	709754	1239	—
Complex (FtsB <sup>ΔCΔH</sup> )	709645	1239	—

**Table 4** Residues included in MD simulation systems.

Component	PerM-FtsB and PerM only	FtsB	Divisome systems	
			FtsB <sup>ΔC</sup>	FtsB <sup>ΔCΔH</sup>
PerM	11–390	10–385	10–385	10–385
FtsB	61–204	74–228	74–205	74–185
FtsQ	—	92–314	92–314	92–314
FtsL	—	120–232	120–232	120–232
FtsW	—	41–456	41–456	41–456
PbpB	—	79–679	79–679	79–679

used pure POPE bilayers, and future work may investigate differences in more complex model membranes [42]. Second, simulations of the PerM-FtsB complex and PerM alone followed a three-stage aMD protocol described in the main text with 2 fs timesteps. For the aMD stage, OpenMM scripts from CHARMM-GUI [43] were modified to utilize empirically determined dual boost potentials as previously described for MD of transmembrane proteins [44] and to output a log of applied boost potentials. We monitored boost potentials applied to dihedral angles and total energy and found the magnitudes of the two boost potentials were similar. Lastly, divisome complex simulations utilized hydrogen mass repartitioning [45] and 4 fs timesteps.

Atoms of interest for analysis shown in Fig. 2B and Fig. 6C were identified by investigating sequence alignments to orthologs of FtsB, PbpB, and FtsW to determine conserved FtsB<sup>H</sup> residues and putative active-site residues. The MDAnalysis package [46, 47] was used to extract trajectories wrapped around the center of mass of protein atoms, to align divisome complex trajectories to FtsW<sup>60–407</sup> (omitting more mobile cytoplasmic residues), and to extract atomic coordinates. All-atom coordinates of initial and final conformers as well as protein-only trajectories are available as supplementary data.

## E. coli strain construction

In addition to Mtb PerM and FtsB (and variations described in the main text), capsid protein G8P from bacteriophage Pf3 (UniProt identifier P03623; referred to as “Pf3” elsewhere in this manuscript) was utilized as a control based upon its small size, efficient translocation, and uniform transmembrane orientation [25]. Proteins were expressed as translational fusions with fluorescent proteins mTurquoise2 [48], mNeonGreen [49], or mEos3.2 [50]. No linkers were added between Mtb protein sequences and fluorescent proteins as we expected that native termini were disordered based on structure predictions. Sequences for all proteins were codon-optimized for *E. coli* expression. Plasmids for FtsB-expressing constructs were constructed by isothermal assembly from low-noise, tetracycline-inducible, ampicillin-resistant vectors [51]. Plasmids

for expression of mNeonGreen as well as mNeonGreen fusions to PerM variants and Pf3 were constructed from IPTG-inducible, spectinomycin-resistant vectors [52]. All plasmids and induction conditions used in this study are summarized in Table 5.

Plasmids were transformed into *E. coli* strain MG1655 and selected for growth on 50 mg L<sup>-1</sup> carbenicillin and/or spectinomycin in LB or SOB media. For microscopy experiments, cells were grown overnight from LB plates in rich defined medium [53] with 0.2% glucose (ForMedium Neidhardt Basal Salt Mixture, Glucose 20 g L<sup>-1</sup>, and Neidhardt Supplement Mixture, Complete), adjusted to pH 7.0 with KOH. All cell growth and imaging took place at 37 °C. Consistent with our earlier work, our plasmids allowed for independent induction of co-expressed genes [52]. In preliminary experiments, we confirmed that mTurquoise2- and mNeonGreen-labeled proteins could be independently induced, that inducing expression did not impact growth at expression levels utilized in this work, and that there was no significant crosstalk between mTurquoise2 (CFP filter set) and mNeonGreen (YFP filter set).

## Fluorescence microscopy

All imaging was done on a Leica DMI6000 inverted microscope using illumination from either a Leica EL6000 source or with laser excitation at 514 nm (mNeonGreen tracking) or combined 405 nm and 561 nm laser excitation (mEos3.2 tracking), adjusting 405 nm laser intensity to achieve acceptable densities of photoactivated mEos3.2 molecules. Data was acquired using a 100×/1.46 a-plan apochromat oil immersion objective with additional 1.6× magnification, Leica type F immersion oil, and an Evolve 512 EM-CCD camera (Photometrics), giving a 100 nm pixel size and a narrow depth of field. Fluorescence images were acquired with three different filter sets: Chroma 49001 (mTurquoise2), Semrock LF514-B (mNeonGreen), and a Chroma custom dual-laser filter set (mEos3.2; ZET405/561x, ET610/75m, ZT405/561/657rpc-UF2). Images of example data were composed using Fiji [54], with linear scaling

**Table 5** *E. coli* plasmids utilized in this study and their respective induction conditions. Mutations are described relative to wild-type Mtb protein sequences in Table 1 and Table 2

Plasmid	Selection	Induction	Expressed protein	Mutation
pJRF002	Carbenicillin	10 nM ATc	mTq2-FtsB	—
pJRF007	Carbenicillin	10 nM ATc	mTq2-FtsB <sup>ΔLQ</sup>	FtsB <sup>Δ111–163</sup>
pJRF007_helix	Carbenicillin	10 nM ATc	mTq2-FtsB <sup>ΔLQΔH</sup>	FtsB <sup>Δ111–163Δ186–194</sup>
pZH904	Carbenicillin	10 nM ATc	mEos3.2-FtsB <sup>ΔLQ</sup>	FtsB <sup>Δ111–163</sup>
pLG906	Carbenicillin	10 nM ATc	mEos3.2-FtsB <sup>ΔLQΔH</sup>	FtsB <sup>Δ111–163Δ186–194</sup>
pJRF001	Spectinomycin	100 μM IPTG	mNG-PerM	—
pJRF004	Spectinomycin	100 μM IPTG	PerM-mNG	—
pJRF004_n1_cWT	Spectinomycin	100 μM IPTG	PerM <sup>n1</sup> -mNG	PerM <sup>Δ1–23</sup> :MYYLKKNLVK
pJRF010	Spectinomycin	240 μM IPTG	Pf3-mNG	—
pZH813	Spectinomycin	60 μM IPTG	mNeonGreen	—

and maintenance of minimum and maximum intensity values for all comparable images.

Microscope samples were prepared by sandwiching agarose gel pads (rich, defined media with 2% Invitrogen UltraPure agarose) of approximately 100 μL between microscope slides and acid-cleaned #1.5H coverslips (Marienfeld). For all experiments, samples were reinoculated from overnight cultures at a 1:1000 dilution in media including IPTG and/or ATc at specified concentrations, grown for 3 h, added to a gel pad prepared from the same media, and grown for an additional 1.5 h before acquiring data. Microscope sample temperature was held at 37 °C using a combination of a heated microscope sample chamber and an objective heater. This protocol was based on empirical observations that cells reached steady state fluorescence and growth in less than 1 h after sample preparation, and that excessive cell density led to reduced growth and fluorescence after approximately 3 h. In preliminary experiments, we also noted that this combination of growth media and agarose exhibited high background with substantial variation between experiments, found that this could be attributed to agitation while melting agarose. We subsequently minimized agitation while preparing gel pads.

To obtain comparable data for constructs expected to have different translation initiation rates, we conducted preliminary experiments and determined that 60 μM and 240 μM IPTG induction for plasmids expressing mNeonGreen and Pf3-mNG, respectively, were approximately equivalent to 100 μM IPTG for plasmids expressing PerM<sup>n1</sup>-mNG. Protein expression was always induced with IPTG and ATc concentrations listed in Table 5 except in the uninduced conditions indicated by down arrows in figures. To increase reproducibility in fluorescence correlation and single-molecule tracking analysis, we aimed to image the middle plane of *E. coli* cells approximately 500 μm from the microscope coverslip. For fluorescence correlation analysis, we obtained images of 20 fields of cells for each condition in each of 3 independent replicates.

For single-molecule tracking analysis, we obtained streaming movies (33 ms exposure time) of 5 fields of cells for each condition in each of 3 independent replicates (except for mNeonGreen tracking for which we did not replicate the experiment). Replicates were obtained on separate days, and sample preparation steps were staggered in time to minimize differences in how samples were handled for each condition.

## Fluorescence correlation analysis

Brightfield images were segmented using Omnipose [55]. The low and variable contrast in our bright-field data produced variation in widths of segmented objects, so objects were morphologically thinned to single-pixel width and then thickened to a uniform width that was more consistent with observed cell morphology. Sub-pixel channel alignment was achieved by maximizing pairwise cross-correlation of low-pass filtered fluorescence images with inverted brightfield images (*i.e.* mTurquoise2 and mNeon-Green images were aligned to brightfield images—not to each other). Background intensity was estimated as the mode of a histogram of low-intensity pixels outside of *E. coli* cells; in all conditions including uninduced mTq2-FtsB<sup>ΔLQ</sup>, cellular autofluorescence was very low relative to signal from fluorescent proteins. Fluorescence intensities in pixels corresponding to segmented cells after alignment and background subtracted were compared as described in the main text.

For mNeonGreen and mTurquoise2 concentration and correlation comparisons, a mixed linear model was fit to data points from single cells with random slopes and intercepts to allow for day-to-day variation. Two-tailed p-values were calculated for comparison to the reference condition (PerM<sup>n1</sup> and FtsB<sup>ΔLQ</sup>, 100 nM ATc) and adjusted for multiple comparisons using the Holm-Šidák method. Effect sizes are reported for significant results only (adjusted  $p < 0.05$ ). Unless stated otherwise, errors in the

manuscript are 1 standard error of the mean. We followed our designed analysis protocol and manually screened data to identify and exclude image fields that could contribute to outliers (large shifts in sample position, poor focus, or unusual levels of expression for one or both proteins in a microcolony; 32 of 360 in total) and also excluded cells falling outside the middle 95th percentile in either fluorescence intensity or cell area. However, we found that removing all data filtering steps did not significantly impact differences we observed between experimental conditions.

## Inference of diffusion coefficient distributions

We extracted single-molecule trajectories from image stacks using TrackMate [56]. Spots were detected using DogDetector (250 nm spot radius) and tracked using SimpleSpareseLAPTracker (500 nm maximum linking distance). Thresholds were determined empirically for mEos3.2 (160) and mNeonGreen (200) data, and the first 250 frames of mNeonGreen movies were omitted to exclude data before sufficient photobleaching had occurred to acquire single-molecule tracks of mNeonGreen molecules. Trajectories were  $3.56 \pm 0.03$  and  $3.65 \pm 0.11$  frames long for mEos3.2 and mNeonGreen, respectively (mean and standard error of the means for 15 mEos3.2 and 4 mNeonGreen experiments; 5 streaming movies acquired in each experiment). We then used SASPT [27] to infer posterior distributions of track diffusion coefficients and localization errors, marginalizing out localization distributions for analysis related to Fig. 5. Posterior distributions were inferred for log-spaced diffusion coefficients between  $0.01$  and  $10 \mu\text{m}^2 \text{s}^{-1}$ . Distributions were normalized to occupancies between 0.02 and  $1 \mu\text{m}^2 \text{s}^{-1}$  to exclude states with highly variable occupancy at the extremes of the diffusion coefficient range.

Our primary analysis used default SASPT parameters (200 iterations of refining state occupancy, trajectories longer than 10 frames split into trajectories 10 frames long or less) and we increased the `sample_size` parameter to avoid subsampling data. Based on our observation that single-molecule tracks clearly show the perimeter of *E. coli* cells at the middle plane, we used 500 nm for the `focal_depth` parameter. We also note that we manually inspected maximum projections of all image stacks to verify that cells were well focused. To check for sensitivity to SASPT analysis parameters, we repeated posterior occupancy inference with various conditions: (1) excluding trajectories that were relatively long ( $> 10$  frames which can arise from fluorescent objects that do not photobleach) or short (2-frame trajectories that can arise from false positive localizations), (2) fixing

localization error to 25 nm, and (3) increasing state array refinement to 1000 iterations. We found that SASPT results were robust against parameter choices and that the largest difference was somewhat narrower distribution peaks with increased iterations.

**Acknowledgements.** ZH is supported by Fundação para a Ciência e a Tecnologia (FCT) through MOSTMICRO-ITQB (DOI 10.54499/UIDB/04612/2020; DOI 10.54499/UIDP/04612/2020) and LS4FUTURE Associated Laboratory (DOI 10.54499/LA/P/0087/2020). ZH and RX are supported by a MOSTMICRO-ITQB Exploratory Projects grant. ZH and JRF thank Sara F. Costa for assistance and insight on this project. ZH thanks Luísa Guerreiro and Margarida Pereira who worked on this project during internships in our lab.

**Data availability.** Plasmids described in this manuscript will be made available through AddGene. Data and analysis notebooks will be made available shortly after preprint publication at <https://github.com/smmlab/mtb-perm-ftsb>. Raw microscopy data and MD trajectories (protein atoms only) will be deposited at the Zenodo repository shortly and can be found by visiting the URL above.

**Author contributions.** JF: molecular cloning, data acquisition, preliminary data analysis and interpretation. RX: data acquisition, data analysis. ZH: structure predictions and simulation, data acquisition, data analysis. All authors wrote and edited the manuscript.

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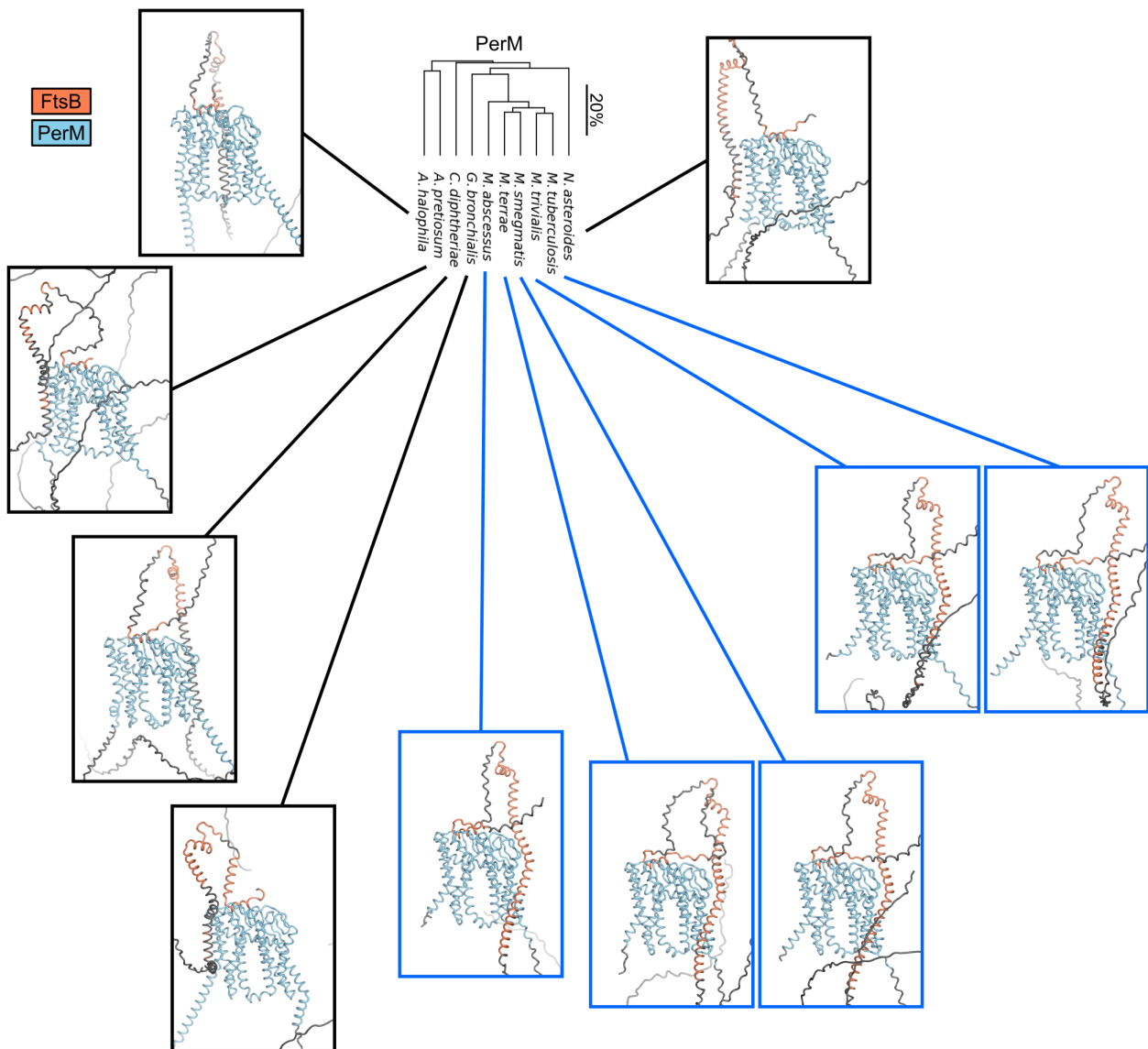
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## Supplementary Figures



**Fig. 1** The PerM phylogenetic tree from Fig. 1 is reproduced and compared to PerM-FtsB complexes predicted from full-length sequences for the same species. The predicted orientation of FtsB<sup>TM</sup> relative to PerM observed for Mtb is only observed for more closely related species (blue lines). However, FtsB<sup>H</sup> interaction with PerM is predicted for all species. Residues with  $pLDDT < 50$  are colored gray.