

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

João Ramalheira Ferreira<sup>1†</sup>, Ruilan Xu<sup>1†</sup>, Zach Hensel<sup>1\*</sup>

<sup>1</sup>ITQB NOVA, Universidade NOVA de Lisboa, Av. da República, Oeiras, 2780-157, Portugal.

\*Corresponding author(s). E-mail(s): [zach.hensel@itqb.unl.pt](mailto:zach.hensel@itqb.unl.pt);

†These authors contributed equally to this work.

## Abstract

Latent infection by *Mycobacterium tuberculosis* impedes effective tuberculosis therapy and eradication. The protein PerM is essential for chronic *M. tuberculosis* infections in mice and acts via the divisome protein FtsB to modulate cell division. Using transgenic co-expression in *Escherichia coli*, we studied the *M. tuberculosis* PerM-FtsB interaction in isolation from other *M. tuberculosis* proteins, engineering PerM to enhance expression in the *E. coli* membrane. We confirmed the reported instability of *M. tuberculosis* FtsB and linked FtsB instability to a segment of FtsB predicted to bind cell-division proteins FtsL and FtsQ. Though narrowly conserved, the FtsB-PerM interaction emerges as a potential target as part of therapy targeting persistent infections by disregulating cell division. Using fluorescence microscopy, we found that the 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:** keyword1, Keyword2, Keyword3, Keyword4

## Introduction

*Mycobacterium tuberculosis* is a major contributor to preventable deaths by infectious disease and has infected approximately a quarter of the global population (Houben and Dodd, 2016). Most infections progress to latent tuberculosis infection (LTBI), characterized by an immunological response to *M. tuberculosis* antigens without clinical signs of active TB (WHO, 2018). 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 (Dale et al, 2021). Protocols that reduce the duration and adverse effects of treatment can reduce LTBI burden by improving upon completion rates for preventative treatment (Assefa et al, 2023). Furthermore, imperfect treatment over long periods can contribute to acquisition of drug resistance (Li et al, 2021). Consequently, it is imperative to better understand how *M. tuberculosis* infections persist to establish LTBI 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 *M. tuberculosis* persists through host-induced stress (Dartois and Rubin, 2022). The actinomycete protein PerM was one of 21 genes identified in a screen of transposon *M. tuberculosis* 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 (Vandal et al, 2008). 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 (Goodsmith et al, 2015). 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 (Wang et al, 2019). 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 (Meade et al, 2023).

Recently, a covalent inhibitor of divisome formation based upon the structure of FtsB-FtsQ was developed and found to be active in vivo against drug-resistant *E. coli* (Paulussen et al, 2022). The *M. tuberculosis* proteome includes homologs of the five core *E. coli* divisome proteins FtsQ, FtsL, FtsB, FtsW, and FtsI (Wu et al, 2018), suggesting that a similar approach can target protein-protein interactions in the *M. tuberculosis* divisome. However, it is unknown whether PerM directly interacts with FtsB (Wang et al, 2019) and, if it does, whether there is a regulatory mechanism for PerM beyond impacting FtsB stability. PerM lacks known orthologs outside of Actinomycetota (Goodsmith et al, 2015), so PerM interactions could potentially be targets of specific therapy. 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 *Mycolicibacterium smegmatis* (Wang et al, 2019). Furthermore, proteins such as

PerM with a high number of transmembrane helices pose challenges for recombinant expression (Gräve et al, 2022; Korepanova et al, 2005).

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 (Baek et al, 2021; Evans et al, 2022) facilitated rapid structure prediction of the core *E. coli* divisome (Attaibi and den Blaauwen, 2022; Craven et al, 2022). 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 (Käshammer et al, 2023). All-atom MD simulations identified this conformational change within 1  $\mu$ s, and further predicted interactions between the core divisome and *E. coli* FtsN (Britton et al, 2023). 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 (Park et al, 2023).

In this work, we employed a combination of structure prediction, MD, and fluorescence microscopy to probe the predicted interaction between *M. tuberculosis* PerM and FtsB proteins. In our approach, *M. tuberculosis* PerM and FtsB were expressed in *E. coli* in order to directly attribute observations to changes in the *M. tuberculosis* 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 *M. tuberculosis* divisome complexes suggestive of PerM regulatory complexity beyond FtsB stabilization.

## Results

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Paragraph introducing approach and perhaps pointing to methods for sequences used and plasmids generated and so on.

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

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Qualitative description and definition of TM (DeepTMHMM prediction), LQ (region with predicted interactions with FtsQ and FtsL and homologous to experimentally observed interactions), H (hydrophobic, helical region predicted to interact with PerM). Check out Fig. 1A for description of regions of FtsB.

Note similar predictions are observed for diverse homologous sequences (supplementary fig with pLDDT shown for different complexes)

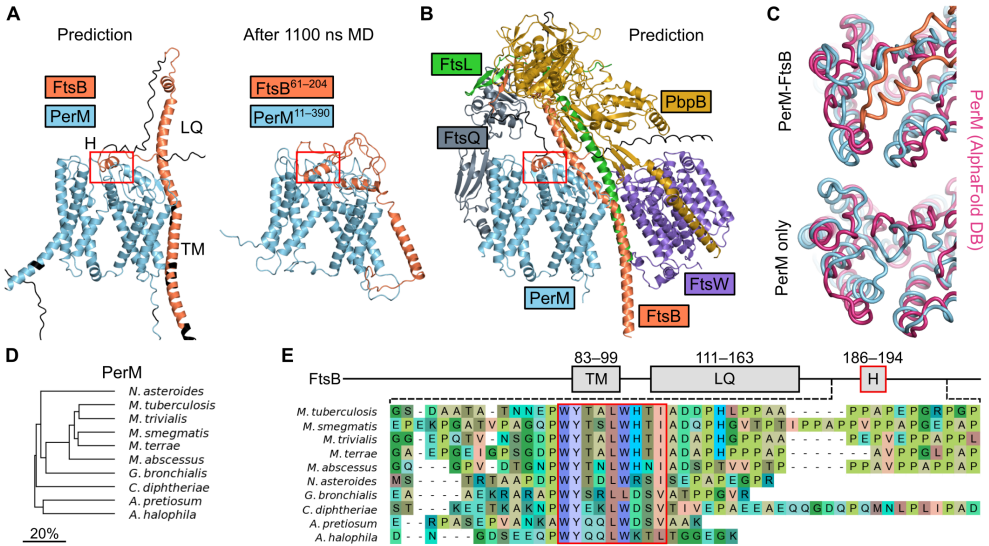
Summary aMD protocol and describe FtsBLQ collapse and loss of predicted conformation with FtsB<sup>TM</sup> while maintaining and appearing to strengthen interaction with FtsB<sup>H</sup>. All simulations using *M. tuberculosis* sequences (ref supp table) Fig. 1A shows aMD results for PerM<sup>61–204</sup> and FtsB<sup>11–390</sup>

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PerM predicted in context of full divisome using full-length sequences. Shown in Fig. 1B.

Fig. 1C compares final conformers after 1.1 ns aMD for simulations of PerM alone and PerM-FtsB to a PerM structure prediction from the AlphaFold DB (REF AF DB?). Describe qualitatively differences. Quantify RMSD. Raise a question here prompted by observation that PerM-FtsB looks more like AFDB than PerM alone MD... what is FtsB stabilizing?

Reproducible MD results Check out Fig. 2! Describe binding pocket maturation. Describe degree of conservation of G228. Describe hydrogen bonds to residues identified in 1E as strongly conserved. Yet these residues are not hydrogen bonded in



**Fig. 1 A conserved, predicted *M. tuberculosis* 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 ns 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 *M. tuberculosis* 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 FtsB 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. **(E)** Top: Diagram of *M. tuberculosis* 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.

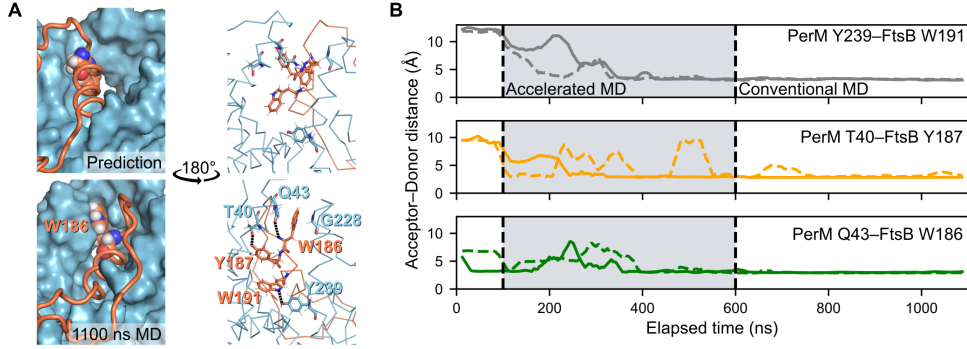
the predicted dimer structure. Instead both the W186 binding pocket and hydrogen bonding with conserved FtsB residues emerge during MD simulation. Both were reproducible in replication of the aMD protocol **2B**.

## FtsB and PerM expression

Introduce short names for proteins and describe strains and experimental approach. Reference supplemental table with strain details.

Check out Fig. 3!

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**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 ns MD (bottom). Left: a large predicted periplasmic binding pocket in PerM evolves to tightly bind FtsB W186. Right: Following MD, FtsB W186 interacts with conserved PerM residue G228 and hydrogen bonds are formed between PerM and conserved FtsB residues. (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.

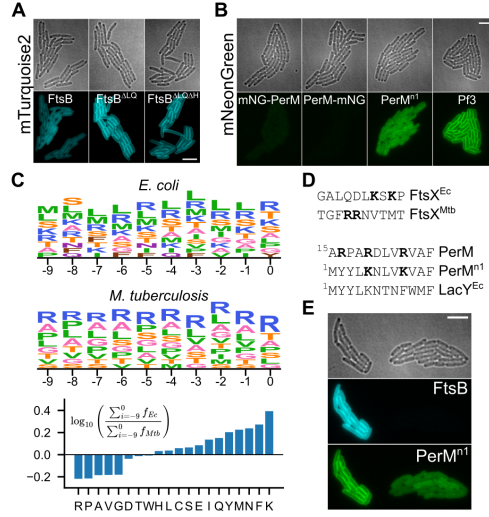
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### PerM N-terminal modification

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### 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* micro-colonies (Fig. 4). In these experiments, expression of PerM<sup>N1</sup>-mNG was induced with

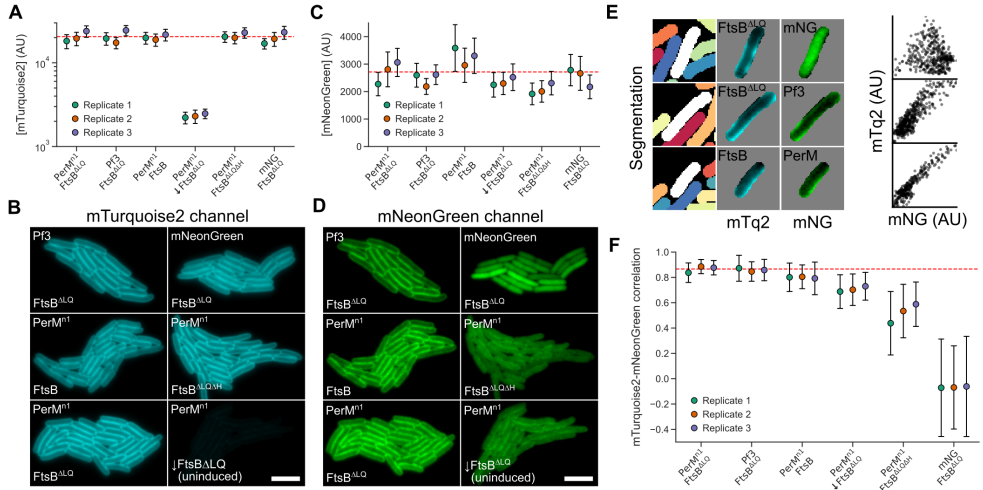


**Fig. 3 Expression of *M. tuberculosis* FtsB and PerM in *E. coli*.** (A) Expression and membrane localization of FtsB and variants with deletions of FtsB<sup>LQ</sup> or with deletion of both FtsB<sup>LQ</sup> and FtsB<sup>H</sup>. Identical minimum and maximum intensities for mTurquoise2 images. (B) Expression and membrane localization for mNG-PerM, PerM-mNG, PerM<sup>n1</sup>-mNG, and Pf3-mNG. Identical minimum and maximum intensities for mNeonGreen images. (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 *M. tuberculosis*. (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  $\mu$ m.

100  $\mu$ M IPTG. Membrane-localization controls mNeonGreen (cytoplasmic) and Pf3-mNG (membrane) were induced with 60  $\mu$ M and 240  $\mu$ M 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) suggests that PerM<sup>n1</sup> stabilizes FtsB when expressed in *E. coli*, as was observed in *M. tuberculosis* (Wang et al., 2019). 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>.



**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 uM 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.

There was no significant change when FtsB<sup>ΔLQ</sup> was replaced with wild-type FtsB ( $p = 0.29$ ). This suggested that FtsB increases PerM stability in *E. coli*, and that this depends on the presence of FtsB<sup>H</sup>. However, inspection of Fig. 4C suggests that our sample size is overpowered ( $N = 10\,930$  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 mNG-PerM<sup>n1</sup> levels despite apparently large impacts of different FtsB constructs on localization suggested that fluorescent mNeonGreen often remained in the cytoplasm following mNG-PerM<sup>n1</sup> 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 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 drops 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

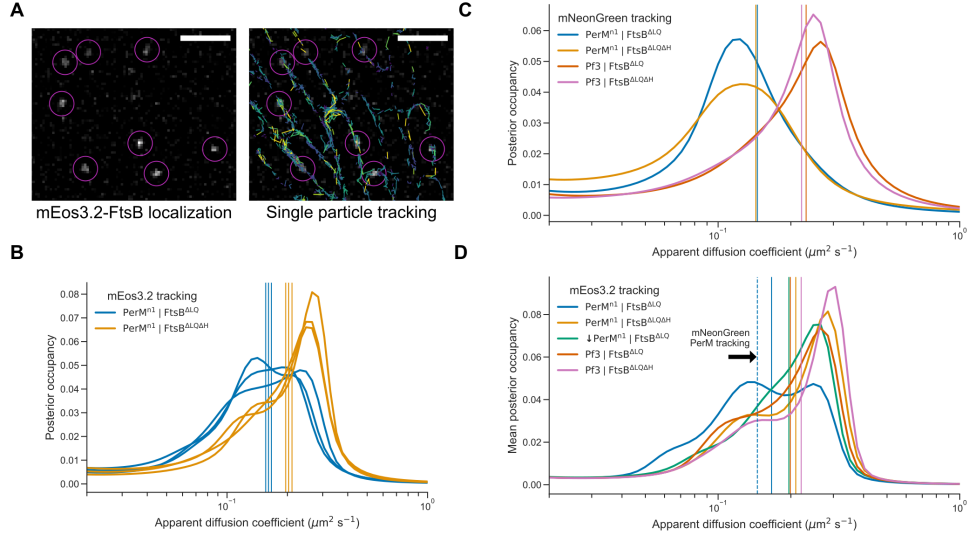
Our correlation analysis strongly suggested that FtsB interaction with PerM via FtsB<sup>H</sup> is key for expression of PerM<sup>n1</sup> in the membrane. Since we observed this for transgenic expression in *E. coli*, it is unlikely that PerM-FtsB interaction is mediated by host proteins. However, 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 *M. tuberculosis* divisome as suggested by depletion phenotypes and midcell localization (Goodsmith et al, 2015; Wang et al, 2019), we hypothesized that this implied long-lived FtsB-PerM interactions that could be detected by single-molecule tracking of FtsB diffusion. FtsB has a single transmembrane helix and PerM is predicted to have 8 (Fig. 1A), suggesting an expected reduction in the FtsB diffusion coefficient from approximately 0.7 to 0.3  $\mu\text{m}^2 \text{s}^{-1}$  upon PerM binding (Lucena et al, 2018).

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Check out Fig. 5!

### Complex MD (fig 5)

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**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 (dashed 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.

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Note region initially predicted and used for MD.

## Discussion

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## Methods

### Structure prediction

colabfold. mmseqs2. AF2-multimer. Pymol for visualization.

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## Molecular dynamics

PerM-FtsB simulations. System details. Simulation methods. Analysis RMSD. Analysis H bond distance.

Divisome simulation. System details. Simulation methods. Active site residue positions. Angle and PerM distance calculations.

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## Sequence analysis

Identification of diverse species with PerM-FtsB interactions by searching for PerM homologs. Multiple sequence alignment.

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## Strains

Cloning details. Strain list.

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## **Microscopy**

Cell growth is same for both types of images. Sample preparation. Microscope details.

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## **Microcolony fluorescence imaging**

Sample size.

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## **Single-molecule tracking**

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## **Image analysis**

ImageJ/Fiji reference for image preparation.

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## **Inference of diffusion coefficient distributions**

SASPT details. Brief discussion of apparent 2D diffusion and challenge of identifying perfect method. Details needed to define reported values.

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## **Microcolony fluorescence correlation**

Background estimation and subtraction. Does not effect correlation by definition. Segmentation of brightfield images with Omnipose. Image registration. Important.

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## Statistical analysis

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 ( $\text{PerM}^{\text{n}1}$  and  $\text{FtsB}^{\Delta\text{LQ}}$ , 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.

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**Acknowledgements.** FUNDING. HELP IN LAB (SMC). HELP IN WRITING AND REVIEW. ANY PLASMIDS/REAGENTS?

**Resource availability.** Plasmids described in this manuscript will be made available through AddGene shortly. Analysis code on GitHub. Data available on request and will be archived to Zenodo following peer review.

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

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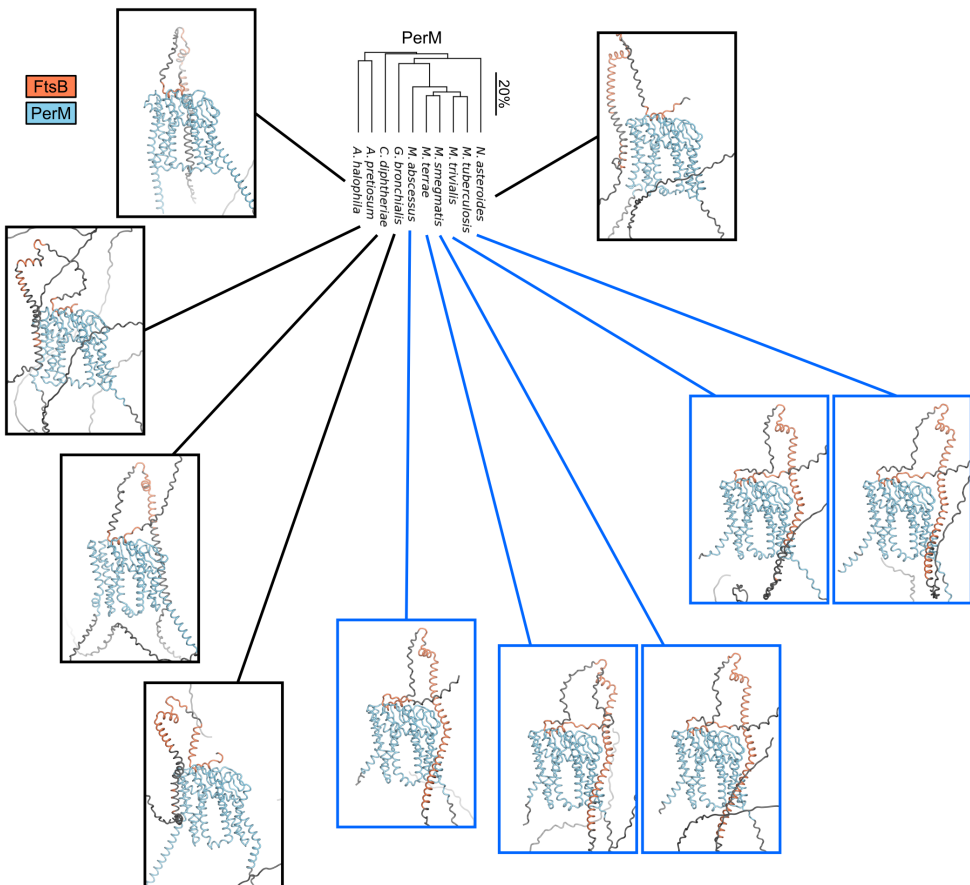
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## Supplementary Figures and Tables



**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 *M. tuberculosis* 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  $pLDL < 50$  are colored gray.