

Wag31, a membrane tether, is crucial for lipid homeostasis in mycobacteria

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eLife Assessment

Understanding bacterial growth mechanisms can potentially help uncover novel drug targets that are crucial for maintaining cellular viability, particularly for bacterial pathogens. In this **important** study, the authors investigate the role of mycobacterial Wag31 in lipid and peptidoglycan biosynthesis. A detailed analysis of Wag31 domain architecture revealed a role in membrane tethering, more specifically, the N-terminal and C-terminal domains appear to display distinct functional roles therein. Whilst the data presented are of use, the experimental evidence is currently **incomplete** and does not yet fully support the conclusions made.

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Abstract

The mycobacterial cytoskeletal protein Wag31 is necessary for maintaining cell shape and directing cellular growth and elongation. Wag31 has a characteristic N-terminal DivIVA-domain and a C-terminal coiled-coil domain. While the role of Wag31 in polar elongation is known, there is limited mechanistic insight on how it orchestrates growth and elongation. In this report, we delineate roles of the N- and C-terminal domains of Wag31 using genetics, state-of-the-art multi-omics, biochemical, and imaging approaches. We show that Wag31 predominantly interacts with several membrane-associated proteins involved in lipid metabolism, cell wall synthesis and division. Native levels of Wag31 are critical for the maintenance and distribution of membrane lipids. Both depletion and overexpression of Wag31 perturbs lipid homeostasis, leading to the formation of intracellular lipid inclusions (ILIs). Protein-lipid crosslinking and imaging studies reveal that purified Wag31 can bind and effectively tether Cardiolipin (CL)-containing liposomes. Despite retaining its ability to interact with partner proteins, the DivIVA domain-deleted Wag31 mutant shows defects in liposome tethering *in vitro* and non-polar localization of CL *in vivo*, which eventually causes lethality. Our study suggests that membrane tethering ‘licenses’ Wag31 to form scaffolds that help orchestrate protein-lipid and protein-protein interactions necessary for mycobacterial growth and survival.

Introduction

Cell division is a critical event in the life of a bacterium as it is the prime mode of bacterial growth and reproduction. Hence, the process of their division is tightly knitted. The key process in bacterial cell division, i.e., assembly of the FtsZ ring at the mid-cell (1), is tightly regulated by the Min and Nucleoid occlusion systems (2–6). *Mycobacterium*, which comprises clinically relevant pathogens-*Mycobacterium tuberculosis* (*Mtb*), *Mycobacterium leprae*, and *Mycobacterium abscessus* lacks both the regulators of FtsZ ring placement (7). Instead, they encode for a DivIVA protein, usually encoded by Firmicutes and other Actinobacteria (8–11). It is necessary that the bacterial shape, crucial for normal cellular physiology, is maintained during cell division. Rod-shaped bacteria such as *Escherichia coli* and *Bacillus subtilis* encode MreB and multiple MreB-like proteins that sustain the rod morphology of cells (12). *Mycobacterial spp.* lack MreB homologs (13, 14). The DivIVA protein of *Mtb* called Wag31 is known to regulate both cellular growth and morphology (10, 15), which makes it an exciting candidate to investigate.

DivIVA proteins were discovered three decades ago during the study of cell division mutants of *B. subtilis* (16). DivIVA proteins are rich in coiled-coil domains (17–21), which enable them to form higher oligomeric structures and, hence, scaffolds, making them suitable adaptor proteins (22, 23). They are known to play diverse roles in various species (24), e.g., they play dual roles of septum positioning and chromosome partitioning in *B. subtilis* (9, 25); cell-separation/resolution post-cell-division in *Listeria monocytogenes* (26, 27); pole formation and maturation in *Streptococcus spp.* (Fadda, 2007) In *Mycobacterium spp.*, Wag31 is an essential protein that senses negative membrane curvature found at poles and septum and localizes there (28). It plays roles in cellular elongation and septation in concert with other divisome and elongasome components (10, 15, 29). It is specifically found in higher amounts at the old pole, where it directs polar elongation by recruiting the acyl CoA carboxylase (ACCase) complex involved in fatty acid, mycolic acid, and methyl-branched chain lipid precursor synthesis (28, 30–32). It localizes to the septum after cytokinesis is over, perhaps for septal synthesis via its interactions with FtsI (32–34), though the process remains largely unknown. Consequently, loss of Wag31 leads to cell division defects-loss of polar elongation, which renders the cells 'round' in shape that occurs due to affected peptidoglycan (PG) synthesis (10, 15, 28). Though Wag31 participates in coordinating PG metabolism, the mechanism or the protein network by which it does so remains unknown.

Wag31 has also been associated with the maintenance of the intracellular membrane domain (IMD), a specialized compartment for lipid synthesis, localized predominantly at the old pole in consonance with pole elongation and spottily all around the plasma membrane where it may function to repair membrane defects (35–37). The loss of Wag31 leads to the delocalization of IMD and its protein markers, such as MurG and GlfT2, but it is not understood whether Wag31 plays a direct role in membrane-partitioning or it is just a consequence of localized membrane synthesis (34, 37). The role of Wag31 in PG and lipid synthesis that drives cell division remains unclear. Despite numerous studies, the details of how Wag31 orchestrates mega-complexes at the pole or partitions the membrane into two to govern cellular elongation remain elusive.

Here, we endeavoured to address the lacunae in the functionality of Wag31 by asking the following questions-i) what are the ultrastructural changes that occur in mycobacteria due to the absence of Wag31, ii) do Wag31 deficient cells undergo transcriptional rewiring to adjust to cell division defects, iii) is the change in cellular morphology from 'rod-to-round' accompanied by altered lipid levels, iv) what is the interactome of Wag31, v) what are the functions of its N and C-terminal domains and finally vi) what is its relationship with the mycobacterial membrane. The present study shows that Wag31 levels are critical for maintaining lipid homeostasis in

mycobacteria. Our results demonstrate that Wag31 binds and tethers the membrane together to orchestrate membrane-localized processes essential for maintaining normal cellular morphology and physiology.

Results

Loss of Wag31 leads to the formation of intracellular lipid inclusions (ILIs)

In the pursuit of bridging the knowledge gap existing in the functionality of Wag31 ([Fig. 1a](#)), we generated a *Wag31* conditional mutant in *Msm*, by integrating a copy of *wag31* under tetracycline (ATc-tet off) regulation at the L5 locus and replacing the native copy with a hygromycin cassette ([Fig. S1a](#)). The mutant, $\Delta wag31$, thus generated, was confirmed by PCRs ([Fig. S1b](#)) and Western blotting ([Fig. 1b](#)). The lack of Wag31 led to a drop in the turbidity of culture ([Fig. 1c](#)). Analysis of bacillary survival *in vitro* by enumerating CFUs 12h post-ATc addition, revealed a 2 log fold difference in the survival of $\Delta wag31$ ([Fig. 1d](#)). Given the essential role of Wag31 in causing polar growth of the cell, we investigated the effect of depletion of Wag31 on bacterial cellular morphology using electron microscopy (EM). Scanning EM (SEM) suggested that the absence of Wag31 caused a transition in the morphology of the cells from rod to round, as previously documented ([10](#), [15](#), [38](#)). Apart from round morphology (~36%), we also observed a rod-to-round transformation stage in which cells were bulged at one pole with the other pole intact ([Fig. 1e-g](#)). To obtain detailed insights, we performed transmission EM (TEM), which divulges information on the internal structure of cells. We discovered spherical, electron-lucent bodies (indicated by an arrowhead in [Fig. 1h](#)) in the cytosol of all the samples. While *Msm* and $\Delta wag31$ -ATc harbored a few (~2-3) electron-lucent bodies, their numbers were drastically higher in the case of $\Delta wag31$ +ATc ([Fig. 1h](#)). The cytosol of $\Delta wag31$ +ATc was filled with multiple variable-sized bodies which resembled Lipid bodies that *Mycobacterium tuberculosis* (*Mtb*) accumulates inside the host ([39](#), [40](#)). As the term 'lipid bodies/lipid droplets' is used specifically for the lipids acquired by *Mtb* from the host cell, we referred to them as intracellular lipid inclusions (ILIs). We categorized the cells based on the number of ILIs they contained. Their distribution remained reasonably similar within *Msm* and $\Delta wag31$ -ATc cells for all the classes ([Fig. 1i](#)). Whereas $\Delta wag31$ treated with ATc had as high as 85% cells harboring more than 5 ILIs and very few cells with less than 5 ILIs ([Fig. 1i](#)). Our data, therefore, reports a novel observation of lipid accumulation when Wag31 is depleted from mycobacteria.

Wag31 levels influence lipid homeostasis

Next, we set out to analyze if the ILIs are indeed lipidic in nature by staining cells with BODIPY (493/503), a dye that preferentially partitions into neutral lipid droplets ([41](#), [42](#)). We stained either untreated or ATc-treated *Msm* and $\Delta wag31$ with BODIPY and visualized them on a microscope. ATc-treated $\Delta wag31$ cells displayed the highest level of BODIPY fluorescence, indicating the maximum abundance of lipid bodies among the three strains ([Fig. 2a, c](#)). Further, we examined whether the appearance of ILIs is associated just with the loss of Wag31 or is a consequence of deviations from the native expression level of Wag31. Towards this, we overexpressed Wag31 episomally using an isovaleronitrile (IVN) inducible system (*Msm::wag31*) ([Fig. S2a](#)) and stained the cells with BODIPY. Unlike Wag31 depletion, which resulted in spherical cells ([Fig. 1e](#)), Wag31 overexpression resulted in the bulging of cells from one pole ([Fig. 2b](#)). Additionally, the overexpression of Wag31 also led to lipid aggregates in the cell with a few larger 'donut-like' aggregates in the bulged pole ([Fig 2b,d](#)).

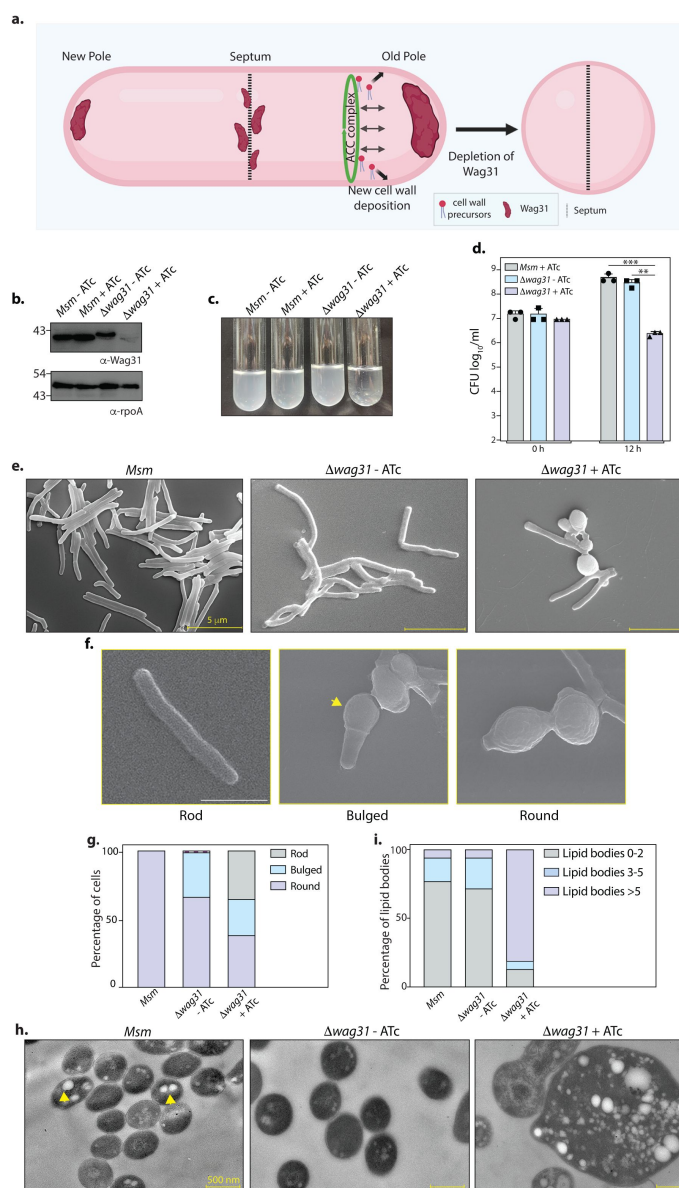


Fig. 1.

Loss of Wag31 leads to the formation of Intracellular Lipid Inclusions (ILIs).

(a.) Illustration depicting the known role of Wag31 in directing polar elongation and maintaining rod-shaped morphology of cells. (b.) Whole Cell Lysates (WCL) were prepared from *Msm* and $\Delta wag31$ (Table S1) either untreated or treated with 100 ng/mL ATc for 12 h, and 40 μ g lysate was resolved on a 12% gel, transferred on Nitrocellulose membrane, and probed with either α -Wag31 or α -rpoA antibody (loading control), respectively. (c.) Cultures showing growth of *Msm* and $\Delta wag31$ either untreated or treated with 100 ng/mL ATc for 12 h. (d.) *Msm*+ATc, $\Delta wag31$ -ATc, and $\Delta wag31$ +ATc were tracked for bacillary survival at 0 and 12 h by enumerating CFUs. Appropriate serial dilutions were plated on 7H11 plates (without antibiotic or ATc), and bar graphs demonstrating bacillary survival (CFU $\log_{10}$ /mL $\pm$ standard deviations (SD, represented by error bars) were plotted at indicated time points. Statistical significance was calculated using Two-way ANOVA (0.0021 (**), 0.0002 (***)). The experiment was performed twice independently, each performed in triplicates. (e.) SEM micrographs showing *Msm*, $\Delta wag31$ -ATc, and $\Delta wag31$ +ATc at 12 h post-treatment. (f.) Cells depicting rod, bulged or round morphology. (g.) 200 cells from each group were checked for either rod, bulged or round morphology, and the percentage of each morphology category was plotted in a bar graph. (h.) TEM micrographs showing *Msm*, $\Delta wag31$ -ATc, and $\Delta wag31$ +ATc at 12 h post-treatment. (i.) 250 cells, each from *Msm*, $\Delta wag31$ -ATc and 185 cells from $\Delta wag31$ +ATc, were divided into three classes (0-2, 2-5, and >5 ILIs) based on the number of ILIs and the percentage of each class in every strain was plotted in a bar graph. This figure was created using BioRender.com.

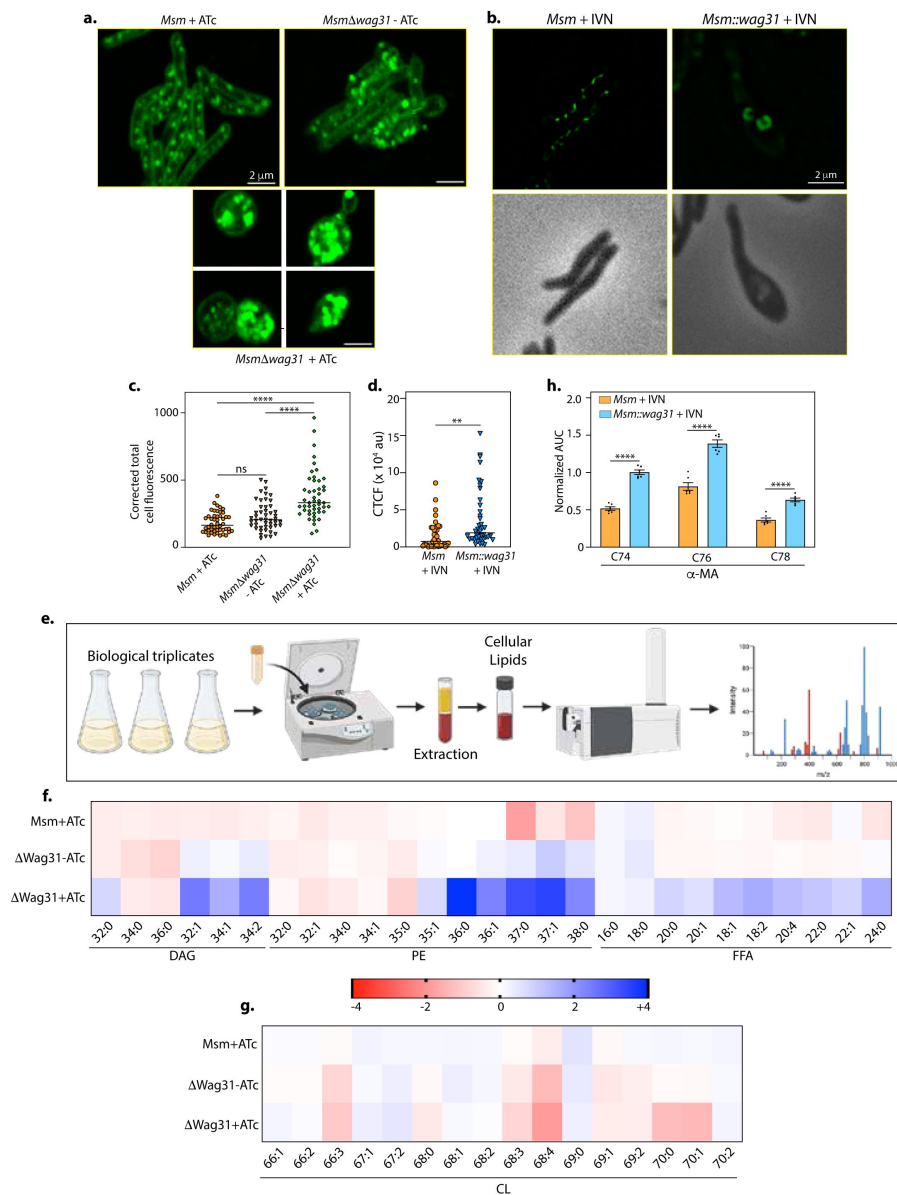


Fig. 2.

Wag31 levels perturb lipid homeostasis

Representative micrographs showing BODIPY staining in **(a.)** *Msm*+ATc, Δ*wag31*-ATc and, Δ*wag31*+ATc samples imaged 12 h post-ATc treatment **(b.)** *Msm*+ IVN, *Msm::wag31* + IVN samples imaged 12 h post 5 mM IVN treatment. **(c.)** Corrected total cell fluorescence was calculated in individual bacterial cells from *Msm*+ATc, Δ*wag31*-ATc, and Δ*wag31*+ATc strains (N=48) and plotted in a graph. Horizontal bar represents median CTFC. (Statistical analysis was performed using one-way ANOVA followed by Brown-Forsythe test (Tukey's multiple comparison test; 0.0021 (**), <0.0001 (****)). **(d.)** Corrected total cell fluorescence was calculated for 42 individual cells from *Msm*+IVN, and *Msm::wag31*+IVN strains and plotted in a graph (Statistical analysis was performed using Welch's t-test 0.0021 (**), **(e.)** Schematic depicting the workflow of Lipidomics (LC-MS). This panel was created using *BioRender.com* <https://www.biorender.com/>. **(f-g.)** Heatmap depicting log₂ fold change (FC) in indicated lipid classes of *Msm* + ATc, Δ*wag31*-ATc, and Δ*wag31*+ATc as compared to *Msm*-ATc. The experiment was performed with six biological replicates, and the abundances of identified lipid classes were normalized to *Msm*-ATc to yield relative abundances of lipids in other strains. Blue and red colours indicate upregulated and downregulated lipid classes; color intensity variation changes with log₂ FC. **(h.)** Bar graph (Mean ± SEM) depicting the abundance of α-MA in *Msm* and *Msm::wag31* treated with 5 mM IVN for 12 h. Values on the Y-axis represent area under the curve (AUC) normalised to an internal standard (FFA 15:0). X-axis represent various species (chain length variations) of α-MA. (Statistical analysis was performed using two-way ANOVA followed by a Tukey's multiple comparison's test; <0.0001 (****)).

Lipid accumulation in both cases led us to inspect whether lipid metabolism is closely tied to Wag31 expression levels in mycobacteria. To investigate that possibility, total cellular lipids were extracted from *Msm*, Wag31 depletion, and overexpression conditions and subjected to semi-quantitative LC-MS analysis (illustrated in [Fig. 2e](#)). Lipidomic analysis of Δ wag31+ATc revealed an imbalance in the levels of neutral lipids, phospholipids, and free fatty acids in comparison to *Msm*-ATc ([Fig. 2f-g](#)). Δ wag31+ATc cells had a significant accumulation of unsaturated diacylglycerols (~3-5 fold) and a very long chain containing (>C36) phosphatidylethanolamines (~5-32 fold). This was accompanied by increased levels of different chain lengths (C18-C24) of free fatty acids (~1.7-4 fold) and reduced levels of cardiolipins C70:0 & C70:1 (~3 fold). Wag31 overexpression, on the other hand, led to a 2-fold increase in the level of different species of alpha-mycolic acids (C74-C78) as compared to *Msm*+IVN ([Fig. 2h](#)), while other tested lipids remained unchanged ([Fig. S2](#)). Altogether, the results presented above demonstrate that Wag31 levels are crucial for maintaining lipid homeostasis in mycobacteria, the depletion and overexpression of which impact lipid metabolism differently.

Molecular interactions of Wag31 with membrane-associated proteins govern cellular homeostasis

We reasoned that Wag31, an adaptor protein, might maintain lipid homeostasis by interacting with proteins involved in these pathways. To examine the protein interaction network of Wag31, we performed an interactome analysis by utilizing Δ wag31, which expresses FLAG-Wag31 integrated at the L5 locus, and used FLAG-GFP as the control to differentiate from non-specific interactions. FLAG-Wag31 and FLAG-GFP from biological triplicates were immunoprecipitated using whole cell lysates from the log phase cells and subjected to mass spectrometry analysis ([Fig. S3a](#) & [Table S2](#)). This study identified both direct and indirect protein-partners of Wag31. The proteins identified were categorized based on their functions. It was found that Wag31 interacts with proteins involved in cell-wall elongation and division, lipid metabolism, intermediary metabolism and respiration and information pathways (translation, replication) ([Fig. 3a](#), [Fig. S4](#)). Interactions of Wag31 with AccA3 and AccD5, subunits of the ACCase complex, and Rne or RNaseE (Msm4626), a membrane-associated protein, have been reported previously ([28](#), [43](#)). Subunits of ACCase complex and Rne were among the top 5 hits in our MS/MS data, validating our approach ([Fig. 3a](#), marked with asterisk). The classification of the top 20 hits is represented in [Fig. 3a](#). The highest number of protein interactors, expectedly, fell in the cell wall and cell processes category ([Fig. 3a](#)). For the present study, we focussed on the interactors found in the cell wall elongation and cell division and lipid metabolism functional categories to probe the cause behind the lipid dysbiosis phenotype.

MurG, involved in PG metabolism ([36](#), [44-46](#)), and SepIVA, involved in regulating septation and the spatial regulator of MurG ([47](#)) also surfaced in the interactome of Wag31. Msm1813, or acyl CoA carboxylase (ACAD), crucial for nascent fatty acid precursor synthesis, was also enriched in the interactome. Other lipid metabolism interactors of Wag31 included acyl-CoA dehydrogenase (Msm2080), involved in lipid catabolism; GpsA, involved in phospholipid biosynthesis; and Msm0372, involved in fatty acid biosynthesis. To validate the high-throughput data, we chose MurG, SepIVA and Msm2092 and performed pull-down assays with His-tagged Wag31. Briefly, His-tagged Wag31 ([Fig. 3b](#)) was incubated with *Msm* lysates expressing 3X-FLAG tagged versions of the tested proteins ([Fig. 3c](#)) and pulled-down with the help of cobalt beads followed by a western blot analysis. Results obtained confirmed the interaction of Wag31 with MurG, SepIVA and Msm2092, thus validating our data ([Fig. 3d](#)).

Since Wag31 is a spatiotemporal regulator of other proteins, we were intrigued to discover the localisation of its interacting partners. We used the data curated by Zhu et al. ([48](#)) and determined that Wag31 majorly interacts with membrane-associated proteins with a few exceptions ([Fig. 3e](#), [Fig. S3b](#)). The mycobacterial plasma membrane has been recently shown to have a cell wall-associated compartment called PM-CW and a cell wall-free compartment called

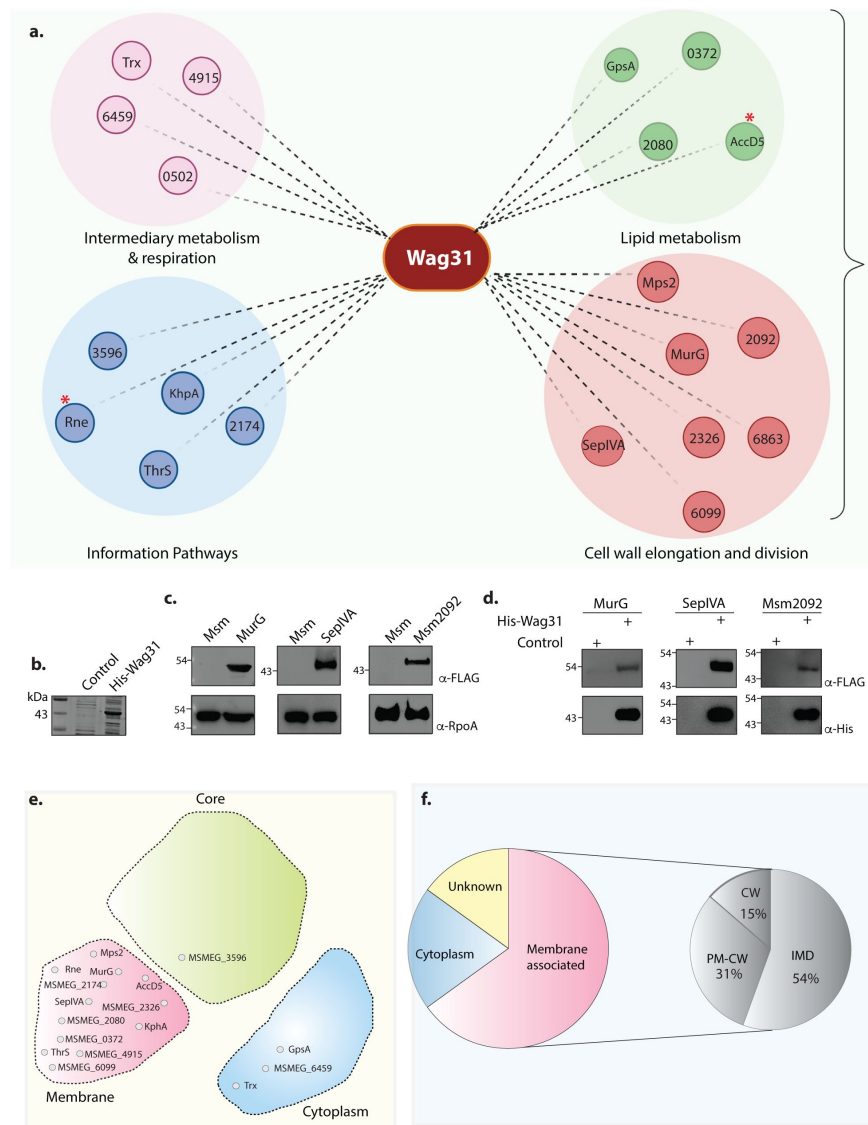


Fig. 3.

Molecular interactions of Wag31 with membrane-associated proteins govern cellular homeostasis.

(a.) Illustration showing the distribution of the top 20 interactors of Wag31 identified from MS/MS analysis from three independent immunoprecipitations. Interactors were classified on the basis of their functional category based on data available in Mycobrowser (66). MSMEG numbers or the identity of interactors are marked in smaller circles. (b.) Coomassie Brilliant blue (CBB) staining of *E. coli* lysates expressing either His-Wag31. *E. coli* lysate prepared from BL21-DE3 strain was used as the control. (c.) Representative western blots demonstrating expression of 3XFLAG tagged-MurG, -SepIVA, -Msm2092 and, -AccA3. WCLs were prepared from 0.5 mM IVN induced *Msm::murG*, *Msm::sepIVA*, *Msm::msm2092* and 40 mg of each was resolved on a 10 % SDS-PAGE, transferred onto a nitrocellulose membrane and probed with α-FLAG and α-RpoA (loading control). (d.) Representative western blots showing interaction of MurG, SepIVA, and Msm2092 with Wag31. *Msm* and *E. coli* lysates represented in (b & c) incubated together and pulled down with Cobalt beads were resolved on a 10-12 % SDS-PAGE, transferred onto nitrocellulose membrane and probed with α-His to detect the pulldown and α-FLAG to detect the interaction. The experiment was performed independently twice. (e.) Graphic representation (adapted from (48)) shows the distribution of the top 20 interactors in different compartments of a mycobacterial cell: membrane, cytoplasm, or core (involved in DNA replication and recombination). Interactors from each compartment are marked inside the respective domain. (f.) Pie-chart shows the distribution of interactors belonging to the IMD, PM-CW, or membrane proteins whose specific localization is not determined. Percentage of proteins (top 25 hits) belonging to each membrane-compartment is mentioned. This figure was created using BioRender.com.

PM_f or IMD, which is the site for lipid synthesis. Based on the available data (36, 48, 49), we found that Wag31 interacts more with IMD-proteins than with PM-CW proteins (Fig. 3f). In accordance with assigned roles (36), PM-CW residing proteins in the interactome are involved in cell wall and cell processes and intermediary metabolism and respiration. Similarly, IMD-homing proteins clustered in the lipid metabolism and cell wall synthesis category. These results suggest that Wag31 maintains lipid homeostasis and cellular morphology primarily through its interactions with the membrane proteins participating in such pathways.

Wag31 binds and tethers cardiolipin-containing membranes

Wag31 is a cytoplasmic protein that recognizes negative curvature regions in the cell, i.e., poles and septum (28). It also forms higher-order structural assemblies at the poles (21, 50). However, the precise mechanism by which Wag31 assemblies localize to the poles remains elusive. The predominant interactions of Wag31 with membrane-associated proteins (Fig. 3c) were intriguing and led us to investigate if it can directly bind to membranes. We reasoned that such membrane binding activity might render it capable of acting as a scaffold that anchors and organizes other proteins, thereby modulating their functions. To test this, we performed a lipid crosslinking-based PLiMAP assay (Fig. 4a). Purified Wag31 was incubated with ~100 nm extruded liposomes containing each of the abundant classes of bacterial lipids phosphatidylethanolamine (PE), phosphatidylglycerol (PG), or cardiolipin (CL). Liposomes contained trace amounts of a reactive fluorescent lipid (RFL), which crosslinks with membrane-bound proteins upon photoactivation. These assays revealed that Wag31 preferentially binds CL- but not PG- or PE-containing liposomes (Fig. 4b). Both CL and PG are anionic lipids, and the preferential binding of Wag31 to CL indicates that the binding cannot be attributed to a charge-based interaction. Remarkably, the sensitive fluorescence-based detection revealed the presence of SDS-resistant higher-order oligomers of membrane-bound Wag31 (Fig. 4b).

Wag31 is a homolog of the DivIVA protein from *B. subtilis*. The latter has been shown to bind to the membrane via the N-terminal DivIVA domain (N-DivIVA_{BS}) and localize other cell division proteins through its interactions with the C-terminal region (51, 52). However, the functional significance of the N- & C-terminal domains of mycobacterial Wag31 has not been dissected. Considering that the Wag31 N-terminal region shares 42% sequence identity with N-DivIVA_{BS} (50), we sought to inspect the effect of disruption of a particular residue (K20) in the N-terminus and the deletion of the entire N-terminus, encompassing the DivIVA domain (1-60 amino acids) on membrane binding (Fig. 55a). The K20 residue lies within a positively charged patch, which likely constitutes a membrane-associating surface in the intertwined loop region of Wag31 (50). Moreover, mutating the K20 residue has been shown to decrease the cell length and elongation rate by 18% and 22%, respectively (34). The K20A mutation showed a marginal defect in binding to CL-liposomes (Fig. 4c-d), indicating that K20 alone is insufficient for membrane binding. However, Wag31_{K20A} showed a marked defect in forming SDS-resistant higher-order oligomers, as seen in the binding assays. Interestingly, deletion of the entire N-terminal domain caused a significant defect in both membrane binding and forming SDS-resistant higher-order oligomers (Fig. 4c-d). Surprisingly, binding was not entirely abolished, suggesting that the C-terminus of Wag31 may also contribute to membrane binding. Taken together, our results indicate that Wag31 preferentially binds CL-containing membranes and that regions other than the N-terminal domain contribute to membrane binding. The latter feature distinguishes Wag31 from DivIVA_{BS}, and Wag31 likely contains membrane binding sites located at both the N- and C-terminal regions, which might lead to differences in the manner in which it interacts with the membrane and partner proteins compared to DivIVA_{BS}.

To further confirm membrane binding, we imaged 100 nm-extruded CL-containing liposomes with a membrane dye DiI in the presence of Wag31. We used a GFP-tagged Wag31 mixed with an untagged Wag31 at a 1:10 molar ratio for these experiments. In the absence of Wag31, liposomes appeared as faint but distinct fluorescent puncta (Fig. 4e). Surprisingly, in reactions containing Wag31, we observed liposomes clustered into bright fluorescent puncta that were positive for GFP-

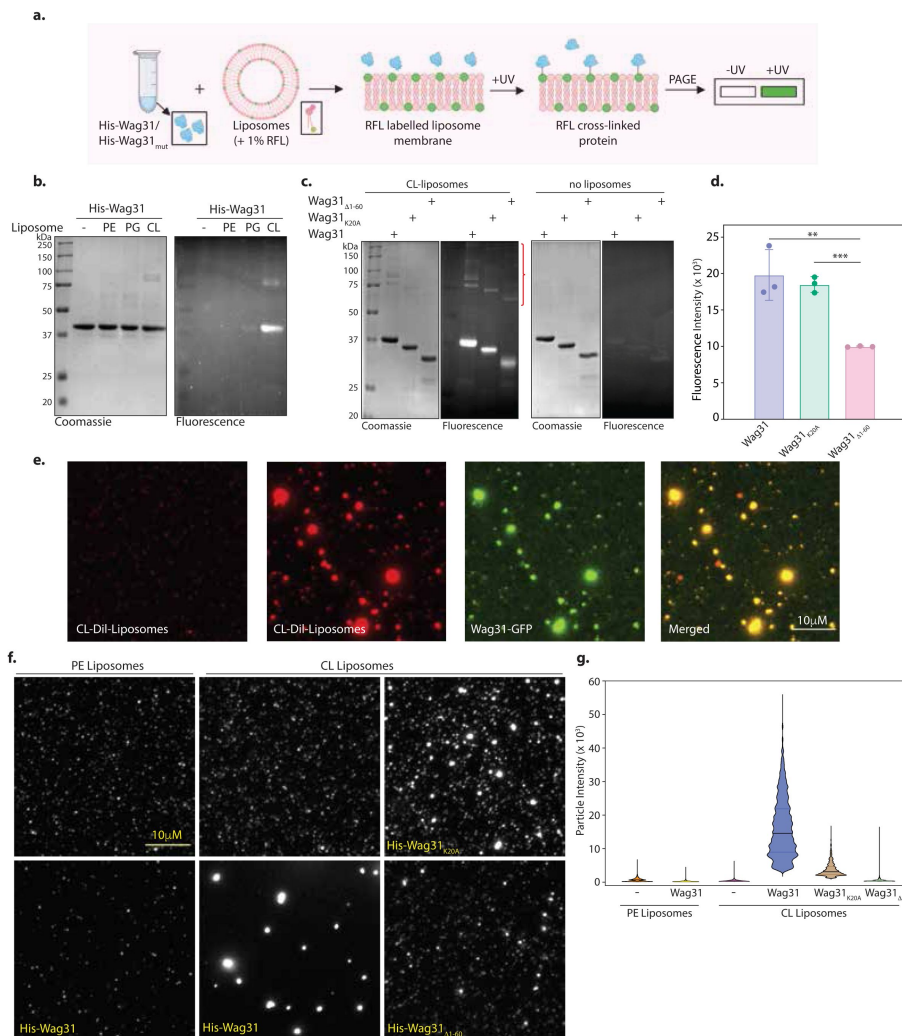


Fig. 4.

Wag31 binds and tethers mycobacterial membrane.

(a.) Schematic showing the workflow of the PLiMAP assay. This panel was created using *BioRender.com* <https://www.biorender.com/>. **(b.)** Coomassie Brilliant blue (CBB) staining of His-Wag31 and in-gel fluorescence (FI) from a representative PLiMAP experiment performed using μ M Wag31 and hundred-fold excess of 30 mol% of either 1,2-dioleoyl-sn-glycero-3-phosphoethanolamine (DOPE-), and 1',3'-bis[1,2-dioleoyl-sn-glycero-3-phospho]-glycerol (DOPG-), or CL-containing liposomes. **(c.)** CBB and FL gels from a representative PLiMAP experiment performed with 2 μ M Wag31, Wag31_{K20A}, or Wag31 $_{\Delta 1-60}$ either in the presence or absence of 30 mol% CL liposomes. Note the presence of SDS-resistant higher-order polymers in gel indicated by the red curly bracket. **(d.)** Quantitation of PLiMAP data from (c.) showing a difference in binding propensities of His-Wag31, His-Wag31_{K20A} and His-Wag31 $_{\Delta 1-60}$. Data represent the mean $\pm$ SD of three independent experiments. Statistical analysis was performed using a 2-tailed unpaired student's t test in Graph Pad Prism 9. Wag31 vs Wag31_{K20A} showed no significant difference (ns, p value = 0.5667) whereas Wag31 vs Wag31 $_{\Delta 1-60}$ (p value = 0.0081, **) and Wag31_{K20A} vs Wag31 $_{\Delta 1-60}$ (p value = 0.0002, ***) showed significant difference in binding **(e.)** Representative micrographs from binding and tethering assay showing CL-Dil-liposomes (~100 nm) before and after incubation with 1 μ M His-Wag31-GFP. Membrane and protein fluorescence are rendered in red and green, respectively. The experiment was performed independently twice. **(f.)** Representative micrographs from tethering experiments performed with His-Wag31, His-Wag31_{K20A}, or His-Wag31 $_{\Delta 1-60}$ proteins. The left panels indicate PE liposomes only (upper panel) and PE liposomes incubated with 1 μ M Wag31 (lower panel), the middle panels indicate CL liposomes (upper) or CL liposomes incubated with 1 μ M Wag31 (lower panel), the right panel indicates CL liposomes either incubated with His-Wag31_{K20A} (upper) and His-Wag31 $_{\Delta 1-60}$ (lower). For visualization, PE and CL-containing liposomes were doped with trace amounts of 3,3'-dilinoleoyloxycarbocyanine perchlorate (FAST DiO™) and 4-chlorobenzenesulfonate (FAST DiI™), respectively. **(g.)** Data from particle analysis showing the maximum intensity of liposomes for all the reactions from two independent experiments.

Wag31 (**Fig. 4e**). The Wag31 tetramer has been modelled as a long rod-shaped protein filament (21, 50) and such clustering of liposomes falls in congruence with the model. It indicates that these long filaments of Wag31 act to tether liposomes. We then systematically evaluated this effect by testing liposomes of different compositions and Wag31 mutants. Again, in the absence of Wag31, both PE- and CL-containing liposomes appeared as faint but distinct fluorescent puncta (**Fig. 4f**). The addition of Wag31 caused clustering of CL-containing liposomes into bright foci (**Fig. 4f**), which is quantified by a particle intensity analysis (**Fig. 4g**). This clustering was caused by membrane binding of Wag31 because PE-containing liposomes that do not bind Wag31 (**Fig. 4b**) showed no clustering (**Fig. 4f,g**). As expected, Wag31 $_{\Delta 1-60}$ was defective in membrane binding (**Fig. 4c,d**) and showed a severe tethering capacity defect (**Fig. 4f,g**). Surprisingly, Wag31 $_{K20A}$, which showed a marginal defect in membrane binding (**Fig. 4c,d**), showed a marked defect in tethering capacity (**Fig. 4f,g**). Together, these results suggest that membrane-binding and membrane-tethering are non-overlapping attributes in the structure of Wag31, and the N-terminal DivIVA-domain is indispensable for membrane-tethering activity.

Tethering is crucial for the survival of mycobacteria

Results presented in Fig. 4c indicate that Wag31 binds strongly to CL-containing membranes. CL is a negatively charged inverted cone-shaped lipid that has been shown to be concentrated at the sites of negative curvature in the bacterial cell, i.e., pole and septum (53, 54). Wag31 has been demonstrated previously to localize at sites of negative curvature (28). Their similar localization pattern, coupled with Wag31's affinity to bind to CL, led us to examine the effect of this binding *in vivo*. We hypothesized that modulations in Wag31 levels might alter CL distribution in the cell. To examine this, we utilized a fluorophore 10-N-Nonyl-acridine orange (NAO) that binds to anionic phospholipids (54, 55). We stained *Msm* and Wag31 depleted and overexpressing cells with NAO and imaged them using a fluorescence microscope. While the localization pattern of NAO was polar in *Msm*+ATc and Δ wag31-ATc, there was a redistribution of red fluorescence to the cytosol and along the perimeter of the cell in Δ wag31+ATc (**Fig. 5a-b**). We quantified the distribution of fluorescence using a line profile which yielded peaks of intensity at the poles and a basal level of fluorescence along the lateral cell body for both *Msm*+ATc and Δ wag31-ATc (**Fig. 5b**). Wag31 overexpression also led to a change in the distribution of NAO from the poles and septum to 'donut-like' fluorescent foci in the cytosol (**Fig. 5c-d**). Since our *in vitro* results indicate that Wag31 prefers to bind CL over PG (**Fig. 4b**), the change in the distribution of NAO fluorescence implies that the loss or overexpression of Wag31 causes the mislocalization of CL in mycobacterium.

Subsequently, we investigated the impact of tethering-deficient Wag31 mutants on cell survival. Wag31 has an N-terminal domain joined to the C-terminal via a linker region. The N-terminal domain contains a positively charged patch in the intertwined region (the region between two helices in the N-terminus) and membrane binding has been attributed to this region (50). This patch is lined with conserved arginine and lysine residues, K15, K20, and R21 (50) (**Fig. 5e**). As our *in vitro* results suggest the importance of K20 and the DivIVA-domain in tethering, we performed complementation assays with the tethering-defective mutants to examine cell survival and morphology. Δ wag31 was complemented with an integrative copy of either Wag31 or Wag31 $_{K20A}$ or Wag31 $_{\Delta 1-60}$. Δ wag31+ATc showed a two log-fold difference in survival and a 40 % decrease in the percentage of rod-shaped cells compared with Δ wag31-ATc (**Fig. 5f-g**). Complementation with Wag31 and Wag31 $_{K20A}$ rescued the loss of cell viability and restored the rod shape of the cells (**Fig. 5f-g**), whereas Wag31 $_{\Delta 1-60}$ behaved similar to Δ wag31+ATc. It neither restored the rod shape of the cells nor reversed the survival defect (**Fig. 5f-g**). These results highlight the importance of tethering for sustaining mycobacterial morphology and survival.

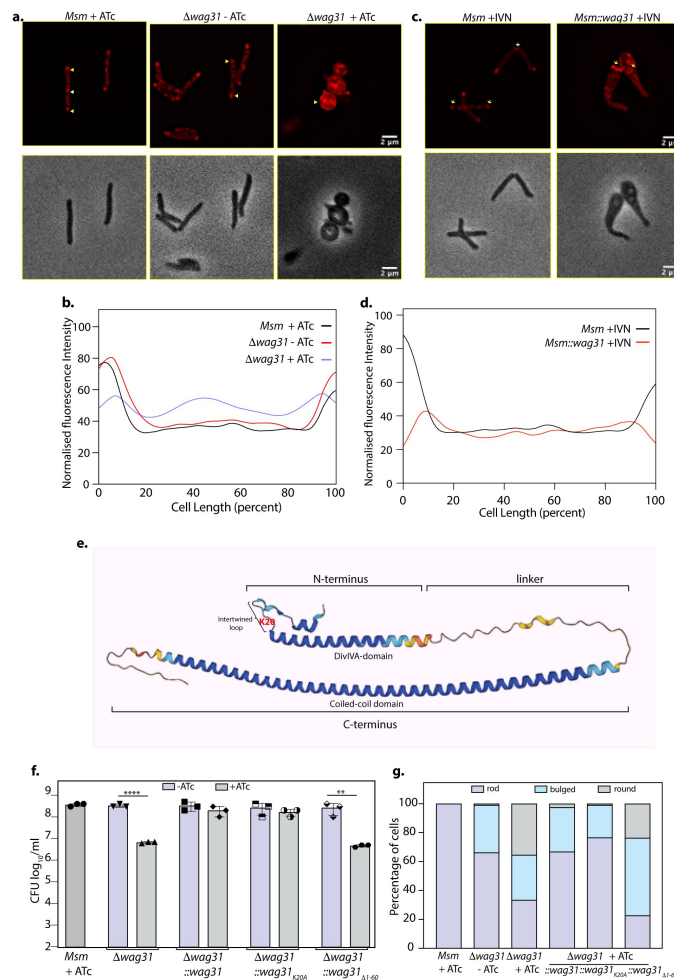


Fig. 5.

Tethering is crucial for the survival of mycobacteria

(a.) Top panel: Representative Fluorescent micrographs showing NAO intensity across *Msm*+ATc, Δ *wag31*-ATc and, Δ *wag31*+ATc samples imaged at 12 h post-ATc treatment. Bright red foci of polar CL are indicated with yellow arrowheads and septal CL foci with white arrowheads. Bottom panel: shows respective phase images. **(b.)** Line plot depicting CL distribution along the length of the cells across *Msm*+ATc, Δ *wag31*-ATc and, Δ *wag31*+ATc. N=100 cells across two independent biological replicates were analysed using Fiji and their normalised fluorescence intensities were plotted against cell length; fluorescence emitted from 0-20 % & 80-100 % of cell length was considered polar. **(c.)** Representative fluorescent micrographs showing NAO staining in *Msm*+IVN, *Msm::wag31*+IVN samples imaged at 12 h post 5 mM IVN treatment. Bright red foci of polar CL are indicated with yellow arrowheads, and septal CL foci with white arrowheads. The panels below show corresponding phase images. **(d.)** Line plot depicting CL distribution along the length of the cells in *Msm*+IVN, *Msm::wag31*+IVN. N=100 cells across two independent biological replicates were analysed using Fiji. The y-axis represents the normalized fluorescence intensities, and the x-axis represents cell length expressed as a percentage of total cell length. Fluorescence peaks in 0-20 % & 80-100 % of cell length were considered polar. **(e.)** Structure of Wag31 (MSMEG_4217) from AlphaFold protein structure database (67). The structure represents the N-terminal, linker, and C-terminal domains in Wag31. The N-terminal contains a short helix, followed by an intertwined loop harbouring positively charged amino acid K20 (highlighted in red) and the DivIVA domain. The N-terminus is linked to the C-terminus via a linker. **(f.)** *Msm*+ATc, Δ *wag31*, Δ *wag31::wag31*, Δ *wag31::wag31*_{K20A}, Δ *wag31::wag31*_{D1-60} either untreated or treated with ATc were scored for bacillary survival post 12 h ATc addition by enumerating CFUs. Appropriate serial dilutions were plated on 7H11 plates (without antibiotic or ATc), and bar graphs demonstrating bacillary survival (CFU log₁₀/mL $\pm$ standard deviations (SD), represented by error bars) were plotted and statistical significance was calculated using two-way RM ANOVA followed by Tukey's multiple comparison test, $\alpha=0.01$, GP: 0.1234 (ns), 0.0332 (*), 0.0021 (**), 0.0002 (***), <0.0001 (****). The experiment was performed twice independently, each with triplicates. **(g.)** Strains described in **(f.)** were scored for their morphology-rod, bulged or round at 12 h post ATc treatment. 308 cells of each strain across two independent experiments were analysed.

Tethering is correlated to lipid levels and CL localization in mycobacteria

As described above, Wag31-mediated membrane tethering is an absolute for the survival of mycobacteria. Perturbations in lipid homeostasis and mislocalization of CL due to altered Wag31 levels hint at a correlation between its ability to function as a homotypic tether and maintain lipid composition and localization. To derive the mechanism behind it, we performed fluorescence imaging experiments with native and tethering defective mutants of Wag31. To investigate a role of tethering in lipid homeostasis in mycobacteria, we stained wildtype and complementation strains with BODIPY and quantified the fluorescence intensities. We observed comparable staining intensity in *Msm* +ATc and, Δ wag31 -ATc (**Fig. 6a-b**). Δ wag31 +ATc expectedly had the highest BODIPY staining intensity, indicating higher than normal lipid levels (**Fig. 6a-b**).

Complementation with Wag31 fully restored the lipid levels to normal, whereas complementation with Wag31_{K20A} couldn't restore the lipid levels completely (**Fig. 6a-b**). Wag31 Δ 1-60 exhibited higher lipid levels than Δ wag31-ATc (**Fig. 6a-b**). These results highlight that the maintenance of lipid homeostasis by Wag31 is a consequence of its tethering activity.

Next, we examined the dependence of CL localisation on tethering by staining *Msm* +ATc, Δ wag31 -ATc, Δ wag31 +ATc, Δ wag31::wag31 +ATc, Δ wag31::wag31_{K20A} +ATc, and Δ wag31::wag31 Δ 1-60 +ATc with NAO. NAO staining showed a polar and septal (in dividing cells) localization pattern in the case of *Msm* and Δ wag31-ATc, whereas it stained the cytosol and perimeter of the Wag31 depleted cells (**Fig. 6c-d**). Complementation with Wag31 restored NAO localization completely, whereas Wag31_{K20A} restored it partially, which could be attributed to its limited tethering activity (**Fig. 6c-d**). Importantly, we observed a mis-localized pattern of NAO with Wag31 Δ 1-60 similar to Wag31 depletion (**Fig. 6c-d**). These results inform of the dependence of CL localization on membrane tethering activity of Wag31. Taken together, the data suggests that Wag31 regulates lipid homeostasis and localisation by the virtue of its tethering activity.

The N and C-terminal domains of Wag31 have distinct functions

From the results presented above, we established a role for the N-terminal region of Wag31 that comprises DivIVA-domain in membrane-tethering (**Fig. 4**). We also identified the protein-interaction network of Wag31 with the help of MS/MS analysis (**Fig. 3** & **Fig. S4**). This implored us to investigate the functionality of the C-terminal region of Wag31. We hypothesized that it might be responsible for mediating interactions of Wag31 with other proteins. We performed a pull-down assay utilizing *E. coli* lysates expressing either Wag31 or Wag31 Δ 1-60 (**Fig. 7a**). We chose novel interactors of Wag31 viz. MurG, SepIVA, and Msm2092 from our interactome database (**Fig. 3a**) and a known interactor of Wag31, AccA3. The *Msm* lysates expressing 3x-FLAG tagged versions of the above-mentioned proteins (**Fig. 7b**) were incubated with *E. coli* lysate or *E. coli* lysates expressing either Wag31 or Wag31 Δ 1-60, followed by pulldown with Cobalt beads. The western blot analysis revealed that all the tested interactors bound to Wag31 (**Fig. 7c**). Interestingly, interactions with all the proteins were retained in the case of Wag31 Δ 1-60 as well, thus indicating that Wag31 interacts with other proteins, largely via its C-terminal. Taken together, the results suggest that the N-terminal region of Wag31 is involved in membrane-tethering through interactions with CL, and the C-terminal region is involved in modulating interactions with other membrane-associated proteins. Thus N-terminal and C-terminal of Wag31 have distinct molecular functions that together are crucial for maintaining cell shape, lipid homeostasis and survival of mycobacteria.

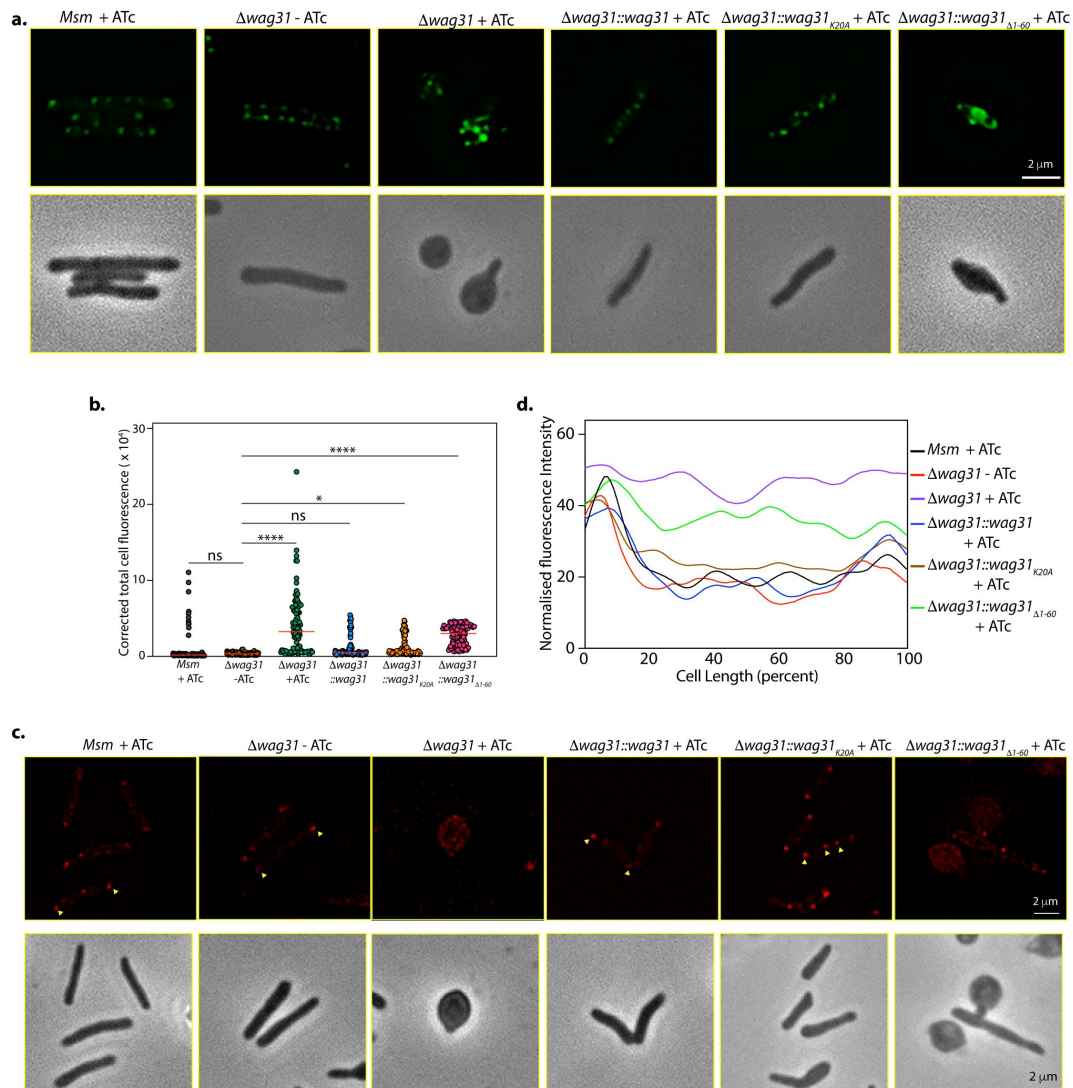


Fig. 6.

Tethering is correlated with lipid levels and CL localization in mycobacteria

(a.) Top panel: Representative fluorescent micrographs showing BODIPY staining across $\Delta w\text{ag}31\text{-ATc}$ and *Msm*, $\Delta w\text{ag}31$, $\Delta w\text{ag}31::w\text{ag}31$, $\Delta w\text{ag}31::w\text{ag}31_{K20A}$, $\Delta w\text{ag}31::w\text{ag}31_{\Delta 1-60}$ imaged at 12 h post ATc addition. Bottom panel: corresponding phase images. **(b.)** Corrected total cell fluorescence (CTCF) of BODIPY-stained cells was calculated from N=100 cells across two independent experiments and analysed using Fiji and plotted in a graph. The horizontal bar represents the median CTCF. (Statistical analysis was performed using one-way ANOVA followed by Brown-Forsythe test (Tukey's multiple comparison test; 0.1234 (ns), 0.0332 (*), <0.0001 (****)). **(c.)** Fluorescent micrographs showing NAO intensity of strains as mentioned above. Bottom panel: corresponding phase images. **(d.)** Line plot analysis of strains described in **(c.)** depicting CL localisation. N=100 cells across two independent experiments were analysed using Fiji and their normalised fluorescence intensities were plotted against cell length.

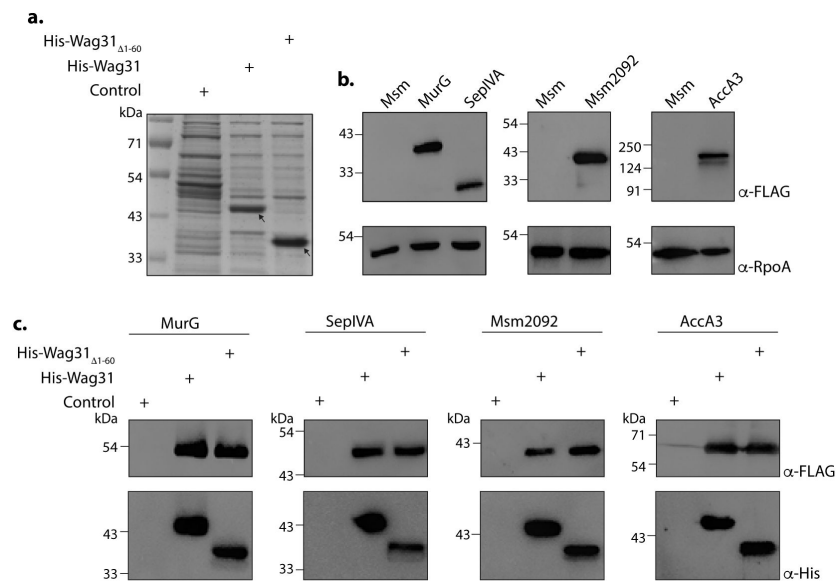


Fig. 7.

The N and C-terminal domains of Wag31 have distinct functions.

(a.) Coomassie Brilliant blue (CBB) staining of *E. coli* lysates expressing either His-Wag31. *E. coli* lysate prepared from BL21-DE3 strain was used as the control. **(b.)** Representative western blots demonstrating expression of 3XFLAG tagged-MurG, -SepIVA, -Msm2092 and, -AccA3. WCLs were prepared from 0.5 mM IVN induced *Msm::murG*, *Msm::sepIVA*, *Msm::msm2092*, *Msm::accA3* and 40 mg of each was resolved on a 10 % SDS-PAGE, transferred onto a nitrocellulose membrane and probed with α-FLAG and α-RpoA (loading control). **(i.)** Representative western blots showing interaction of MurG, SepIVA, Msm2092 and, Acca3 with Wag31 and Wag31_{Δ1-60}. *Msm* and *E. coli* lysates represented in (g & h) incubated together and pulled down with Cobalt beads were resolved on a 10-12 % SDS-PAGE, transferred onto nitrocellulose membrane and probed with α-His to detect the pulldown and α-FLAG to detect the interaction. The experiment was performed independently twice.

Discussion

Wag31 modulates mycobacterial cell division by acting as an adaptor protein that recruits the elongation machinery to the old pole for unipolar growth of cells (10, 15, 28). In the absence of Wag31, rod-shaped cells advance into a spheroplast-like stage, which is thought to arise from dysregulated PG metabolism (10, 11, 28). Recently, Wag31 has been implicated in maintaining the lipid-rich IMD compartment (37). But the molecular mechanisms by which Wag31 manages these functions has remained unclear. In this study, we discover an additional function of Wag31 as a membrane tether, which likely reconciles many of the previously ascribed functions of this protein into a unified model. To gain insights into defects associated with a change in cell shape, we imaged Wag31-deficient cells and found a dramatic accumulation of lipids in the form of ILIs. Interestingly, both *Msm* and Δ Wag31-ATc showed the presence of ILIs, but in much less abundance, indicating that ILIs are a normal occurrence that may function to regulate cellular functions (Fig. 1g-h).

The loss of Wag31 leads to the accumulation of neutral lipids while its overexpression increase the levels of alpha-mycolic acid (a-MA), implying that the maintenance of native Wag31 levels is critical for the proper balance of lipid composition (Fig. 2f-h). A protein's interactome is akin to its functional fingerprint. Here, we identify 116 novel interacting partners of Wag31 through mass spectrometry and validate three of these interactions, namely with MurG, SepIVA and Msm2092, through pull-down studies (Fig. 3 & 5i). Strikingly, the bulk of Wag31 interactors are membrane proteins, which might be attributed to the formation of Wag31 oligomers on the membrane. Previous studies have reported that the depletion of Wag31 delocalize MurG, but if this was caused by loss of direct interaction or was a consequence of change in cell shape was not known (34, 37), which our results clarify. Furthermore, our results identify SepIVA, which regulates MurG (47), the D-aminopeptidase Msm2092 and AmiB, both of which are PG remodellers, as Wag31 interactors. Together, these results contribute towards our understanding of how Wag31 affects PG synthesis (Fig 3a,d). AccA3 is an interactor of Wag31 (28) (34) and catalyses the first committed step in the synthesis of long-chain fatty acids by forming malonyl-CoA. In addition, Long-chain specific acyl dehydrogenase (LCAD), which we report here to be another interactor of Wag31, catalyses the first step in the β -oxidation of fatty acids, which might explain the accumulation of long chain PEs and long-chain fatty acids.

As mentioned above, the overexpression of Wag31 leads to an upregulation in a-MA levels (Fig. 2h). The localization of the a-MA transporter MmpL3 depends on polar PG levels (56, 57). MurG is an essential enzyme for PG synthesis (58) and we report here that it strongly interacts with Wag31 (Fig. 3a,d). It is tempting to hypothesize that homeostatic control on lipid composition arises from the ability of Wag31 to modulate the activity of enzymes involved in lipid metabolism and PG synthesis. That both depletion and overexpression of Wag31 affects lipid composition is likely also a manifestation of its ability to function as a homotypic tether, where the tethering ability is inherently sensitive to the tether concentration.

Our results are the first to establish a membrane tethering function of Wag31. DivIVA proteins possess an N-terminal membrane-binding domain and a C-terminal coiled-coil domain (50, 51). Our data show that disruption of the N-terminal domain (Wag31 Δ 1-60) markedly reduces membrane binding and completely abolishes membrane tethering (Fig. 4f, g). The N-terminal domain binds CL while the C-terminal coiled-coil domain facilitates multimerization, which together could explain the molecular basis for Wag31 to function in homotypic tethering of CL-containing membranes. Results from testing mutants defective in membrane tethering indicate that this function is important for the proper localisation of CL and eventually the survival of mycobacteria (Fig. 5f & 6c-d). Thus, alterations in the native levels of Wag31 results in a displacement of CL from the poles (Fig. 5a-d). However, given NAO's ability to bind to anionic

phospholipids, the observed staining could also typify a redistribution of anionic phospholipids in the cytosol and membrane. In the absence of Wag31 binding to the poles (due to gene silencing or deletion of the DivIVA-domain), the multiprotein scaffold required for cell envelope biosynthesis and cell elongation is not assembled properly, leading to changes in lipid and PG metabolism and consequently to cells failing to form poles where CL would normally concentrate. The failure of a Wag31-centered multiprotein scaffold formation likely cause of this phenotype is further supported by the *wag31* overexpression study, which shows similar changes in NAO staining. By affecting the stoichiometry of Wag31 relative to other interacting protein partners, Wag31 overexpression negatively impacts the formation and activity of the multiprotein scaffold.

While, complementation with Wag31 reverses the phenotypic changes observed upon Wag31 depletion, complementation with the membrane tethering mutant (Wag31 Δ_{1-60}) fails to do so (Fig. 5f-g & 6a-d). These results not only highlight the dependence of CL localization on membrane-tethering but also underscore the importance of tethering for maintaining cellular morphology, physiology and survival. CL micro-domains at the poles (53, 54) have been shown to be an important hub for protein localisation and activity. Several studies have implicated CL in modulating the activities of membrane-resident proteins involved in electron-transport, protein translocation, bacterial two-component system etc. (59–62).

Our work delineates the roles of the N-and C-terminal domains of Wag31. The N-terminal DivIVA-domain facilitates lipid binding and membrane tethering, and the C-terminal coiled-coil domain engages in protein-protein interactions and facilitates the multimerization of Wag31. This is evident from findings that deletion of the N-terminal DivIVA-domain affects lipid binding and tethering but not its ability to bind interacting partners (Fig. 4c-d, f-g & 7c). Based on our results, we propose a model in which the N-terminal of Wag31 binds and tethers CL ensuring its proper localization while the C-terminal is free to interact and recruit proteins to the membrane to facilitate cell growth and division (Fig. 8). We reason that the loss of cell-shape and viability in the absence of tethering occurs due to mis-localization of CL which in turn has a negative impact on membrane-localized processes at a global scale.

Materials and Methods

All the illustrations were made using BioRender. List of constructs and strains used in the study are given in Table S1. Mass spectrometry samples were processed and performed by Valerian Chem.

Bacterial strains and culturing

Middlebrook 7H9 medium (BD Biosciences) or 7H11 agar (BD Biosciences) was used to cultivate mycobacterial strains. 7H9 medium was supplemented with 10% ADC (albumin, dextrose, NaCl, and catalase) and 0.2% glycerol (Sigma) and, 0.05% Tween-80 (Sigma) or 7H11 agar (BD Biosciences) with 10% OADC (oleic acid added to ADC) and 0.2% glycerol were added to the media.

Generation of *wag31* conditional mutant and *in vitro* Growth kinetics

Recombineering method, as described in (63), was used to generate a Wag31 conditional gene replacement mutant. Detailed methodology is provided in the supplementary text. To perform *in vitro* growth kinetics, secondary cultures of *Msm* and Δ *wag31* were seeded at a density of $A_{600} \sim 0.05$ and either left untreated or treated with 100 ng/mL ATc for 12 h. At 0 and 12 h, appropriate serial dilutions were plated on 7H11 agar plate and incubated at 37°C for 2-3 days, and CFUs were enumerated. CFUs were plotted using GraphPad Prism 9 and analysed by performing Two-way ANOVA (0.0021 (**), 0.0002 (***)).

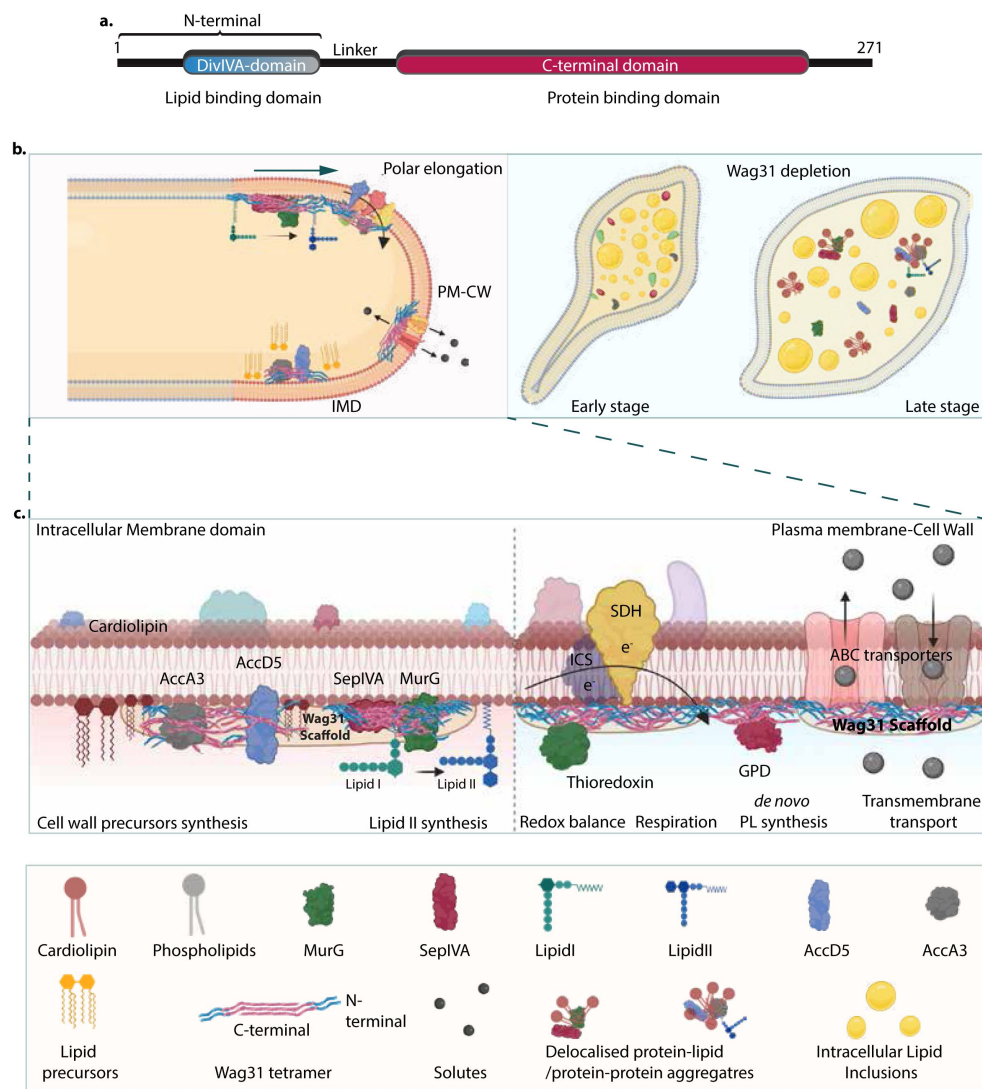


Fig. 8.

Model for Wag31 mediated tethering of the membrane

(a.) Model of Wag31 showing the N- and C-terminals. N-terminal houses the DivIVA-domain that is involved in membrane-binding and the C-terminal binds to various interactors of Wag31. **(b.)** Left: Illustration showing Wag31 scaffolds bound to cardiolipin in the membrane via N-terminal, tethering it and creating CL-microdomains at the poles. The C-terminal of Wag31 binds to protein-partners, localizing them to the membrane, facilitating several membrane-centric processes such as PG and lipid synthesis, that ensure mycobacterial survival. This panel was created using *BioRender.com*. The right panel shows the time-dependent consequences of Wag31 depletion i.e. early and late phase. The cells are bulged at one pole and start accumulating ILIs in the early phase and as time progresses, the effect of Wag31 depletion becomes severe rendering the cells round and full of ILIs. In the absence of Wag31, CL-microdomains become delocalised due to the loss of tethering, pulling away several membrane-associated proteins causing non-polar growth and loss of curvature. This panel was created using *BioRender.com*. **(c.)** A zoomed-in micrograph of the left panel of (b.) highlighting Wag31-mediated cellular activities that occur in IMD and PM-CW. The proteins shown in the front in dark solid colours are interactors of Wag31; light-coloured proteins in the background are other membrane proteins. From left: AccA3 & AccD5 (subunits of Acyl CoA complex) showing cell wall lipid-precursor synthesis; MurG is involved in Lipid II (precursor of PG) synthesis, SepIVA is a spatio-temporal regulator of MurG. Thioredoxin is an antioxidant critical for maintaining cellular redox balance, ICS (Iron-sulfur cluster) and SDH (Succinate Dehydrogenase), or respiratory complex II, are involved in bacterial respiration. GPD (NAD-dependent Glycerol 3-phosphate Dehydrogenase) engaged in *de novo* phospholipid biosynthesis. ABC transporters are involved in transmembrane transport (grey spheres represent solutes). This panel was created using *BioRender.com*.

Western blot analysis

Msm, *Msm::wag31* and *Δwag31* transformants grown in the presence or absence of IVN/ ATc for 12 h were resuspended in Phosphate Buffer Saline containing 5% glycerol and 1mM protease inhibitor-phenylmethylsulfonylfluoride, PMSF (Sigma) and whole cell lysates (WCL) were prepared with the help of bead-beating. Concentration of protein in each sample was estimated by Pierce BCA protein Assay kit (Thermo Fischer Scientific) and 40 mg WCLs were resolved on 10-12% SDS-PAGE gels, transferred onto nitrocellulose membrane, probed with desired antibodies and developed using ECL reagent (Millipore).

Sample preparation for SEM and TEM

Msm and *Δwag31* were incubated in the absence or presence of ATc for 12 h, and the samples were prepared for both SEM and TEM as described previously in (64 [64](#)). For SEM, the samples were imaged at 20,000X using FEI Nova NanoSEM 450. TEM sections were examined on a 120 kV transmission electron microscope (L120C, Talos, Thermo Scientific) at 17,500 magnification.

BODIPY labelling, imaging and analysis

1 mL cells of *Msm* and *Δwag31* either untreated or treated with 100 ng/mL ATc for 12 h were stained with 20 μg/mL BODIPY (Invitrogen). For examining lipids in Wag31 overexpression strain, *Msm* and *Msm::wag31* were treated with 5 mM IVN for 12 h and 1 mL cells of both strains were stained with 2.5 mg BODIPY. Staining was performed for 30 min protected from light in an incubator shaker followed by washing with sterilized Phosphate buffer saline (PBS).

Sample preparation for Lipidomics and analysis

Secondary cultures of *Msm* and *Δwag31* were seeded at a density of $A_{600} \sim 0.05$ and either left untreated or treated with 100 ng/mL ATc for 12 h until $A_{600} \sim 0.8-1.0$. Post 12 h incubation with ATc, cells corresponding to $A_{600} \sim 20$ were processed for lipid extraction as described (64 [64](#)). Detailed methodology is provided in supplementary text.

Immunoprecipitation (IP) and MS/MS analysis

250 mL secondary cultures of *Msm::gfp* and *Δwag31* were inoculated at an $A_{600} \sim 0.05$ and incubated in a shaker incubator until $A_{600} \sim 0.8-1.0$. Cells were collected and lysed using bead-beating to make Whole cell lysate (WCL) and passed through a 0.2 μM filter. 3 mg lysate from each group was immunoprecipitated overnight with anti-FLAG[®] M2 magnetic beads (Millipore), washed thrice, and heated in 4X Laemmli Buffer. One-fifth of the IP supernatant was resolved on a 10% gel, transferred onto a Nitrocellulose membrane, and probed with anti-FLAG Antibody (Sigma). Lysates from confirmed IPs (biological quadruplets) were taken further for mass spectrometry sample preparation and analysis as described in supplementary text.

Proximity-Based Labelling of Membrane-Associated Proteins (PLiMAP)

For carrying out the PLiMAP assay, Wag31 and its mutants were purified and desired liposomes were prepared as described in supplementary. PLiMAP was carried out and analysed as described (65 [65](#)). Briefly, 2 μM Wag31 or its mutants were incubated with liposomes containing 30 mol% CL, DOPG, or DOPE (200 μM total lipid) in a final volume of 30μL. The reaction was incubated for 30 min in the dark at room temperature followed by exposure to a 1 min long pulse of 365 nm UV light (UVP crosslinker CL-1000L) at an intensity of 200mJ cm⁻². The reaction was mixed with Laemmli sample buffer, boiled, and equal volumes for all reactions were loaded onto a 10%

polyacrylamide gel and resolved using SDS-PAGE. iBright1500 (Thermo Fischer Scientific) or Amersham Typhoon (GE) were used for imaging gels. Gels were first imaged for BODIPY fluorescence and later fixed and stained with Coomassie Brilliant Blue.

Tethering assays, fluorescence imaging, and image analysis

μM of Wag31 and its mutants were mixed with 10 μM of 100 nm extruded liposomes in Eppendorf. The mixture was then transferred to a LabTek chamber and imaged using 100x 1.4 NA oil-immersion objective on an Olympus IX73 inverted microscope connected to an LED light source (CoolLED) and an Evolve 512 EMCCD camera (Photometrics). Image acquisition was controlled by $\mu\text{Manager}$, and images were analysed using Fiji. For single liposome intensity analysis, all images from the same condition from two independent experiments were combined into one montage. The montage was duplicated and thresholded using Fiji's Otsu algorithm to generate a binary mask. The mask was overlaid onto the original montage to obtain the maximum intensity of liposomes.

NAO labelling, imaging and analysis

Msm, *Msm::wag31* were treated with 5 mM IVN for 12 h and *Msm* and Δwag31 were either left untreated or treated with 100 ng/mL ATc for 12 h. For complementation experiment, 1 mL of each strain was stained with 2 mM NAO for 1 h protected from light in an incubator shaker and washed thrice with sterilised 1X PBS. 2 ml of cells were spotted on a 1% agar pad (prepared in 7H9) and visualized using a 100X/1.32 NA oil immersion objective on a Leica THUNDER Imaging systems at 475nm:535nm (Excitation: Emission). About 20 Z-stack images, each spaced 0.1 μm apart, were acquired using a scientific CMOS K8 camera (Leica microsystems). For CL specific labelling, collection was also performed at 642nm.

Complementation experiment

Secondary cultures of *Msm*, Δwag31 , $\Delta\text{wag31}::\text{wag31}$, $\Delta\text{wag31}::\text{wag31}_{K20A}$, and $\Delta\text{wag31}::\text{wag31}_{D1-60}$ were set up at $A_{600} \sim 0.05$. For each strain, two types of cultures were set up in triplicates-untreated and treated with 100ng/mL ATc and grown for 12 h. At 0 and 12 h, appropriate serial dilutions were plated on 7H11 agar plates and incubated at 37°C for 2-3 days. CFUs were enumerated and plotted using GraphPad Prism 9. Statistical analysis was performed using two-way RM ANOVA followed by Tukey's multiple comparison test, $\alpha=0.01$, GP: 0.1234 (ns), 0.0332 (*), 0.0021 (**), 0.0002 (***), <0.0001 (****). For BODIPY-and NAO-labelling, the strains were grown as described above. At 12 h, 1 mL cultures of *Msm*+ATc, Δwag31 , Δwag31 +ATc, $\Delta\text{wag31}::\text{wag31}$ +ATc, $\Delta\text{wag31}::\text{wag31}_{K20A}$ +ATc, and $\Delta\text{wag31}::\text{wag31}_{D1-60}$ +ATc were taken and labelled with either BODIPY or NAO. The labelling, imaging and analysis for both fluorophores was performed as described in the respective sections above.

His pulldown assay

E.coli and *Msm* lysate preparation for the assay is described in supplementary. Briefly, 500 mg of *E.coli* lysates expressing Wag31 or mutant were mixed with 300 mg *Msm* lysates expressing Wag31 interactors in a total reaction volume of 700 μL and the samples were twirled at 4°C for 2 h. 50 mL cobalt beads/ sample (GoldBio) were washed twice with lysis buffer and incubated with 5% BSA solution for 2 h at 4°C. Subsequently, beads were washed with lysis buffer and incubated with samples for 2 h (twirled at 4°C). The pull downs were washed thrice in lysis buffer containing 1% Triton X-100 and finally resuspended in 2X-SDS sample buffer. Pulldown samples were resolved on 10-12% SDS-PAGE and probed with α -His and α -FLAG antibodies.

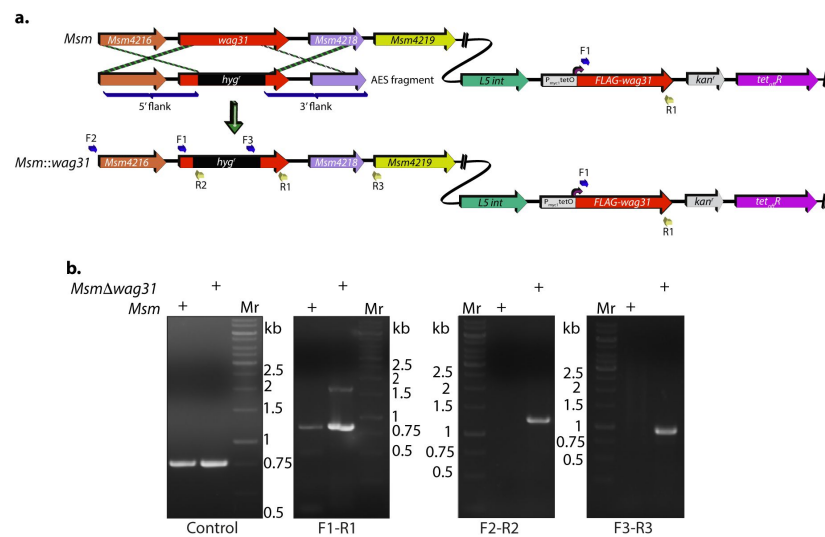


Figure S1.

Generation of Wag31 knockdown

(a.) Schematic showing the *wag31* locus in *Msm* and Δ *wag31* and illustrating the strategy utilized to replace *wag31* locus with hygromycin resistance cassette. Primers used for knockout confirmation are indicated as F1, R1, F2, R2, F3 and R3. (b.) 1% agarose gels showing confirmatory PCRs performed with different primer sets. The first PCR, or the control PCR, was done using SepIVA (738 bp) primers showing amplicon at ~750bp. It indicates an equal amount of gDNA isolated from *Msm* and Δ *wag31* and subsequently used for all PCRs. The next panel shows amplicons from the PCR performed with the F1-R1 pair, which bind to *wag31* locus (~819 bp). They yield an 819 bp amplicon in the case of *Msm* and 819 bp and an additional ~1.5 kb amplicon in the case of Δ *wag31*. The third and fourth panels show amplicons amplified using the F2-R2 set and F3-R3 set, which are expected only in the case of Δ *wag31* confirming legitimate recombination at the native locus.

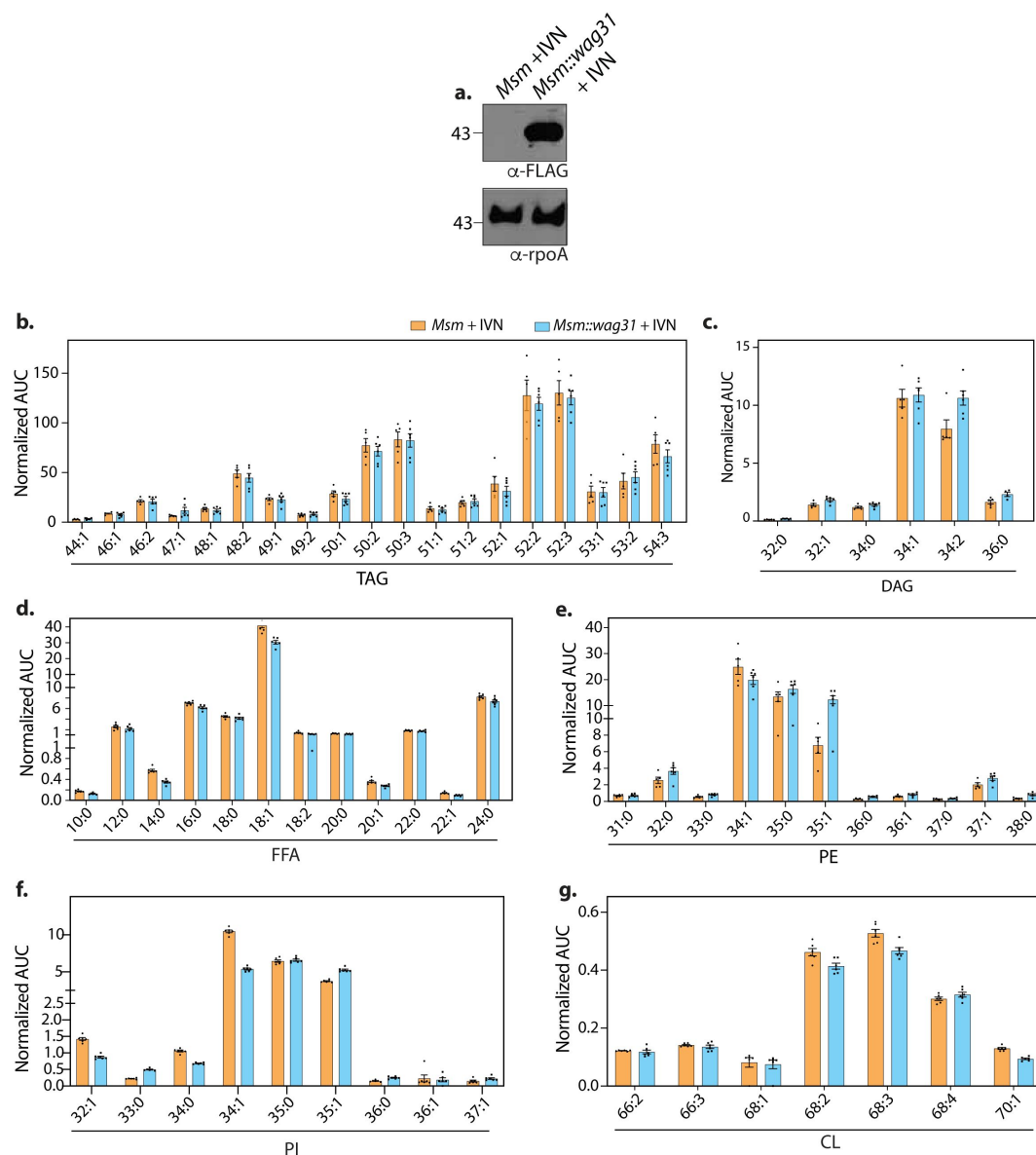


Fig. S2.

Wag31 levels perturb lipid homeostasis

(a.) Whole Cell Lysates (WCL) were prepared from *Msm* and *Msm::wag31* (Table S1) treated with 5 mM IVN for 12 h, and 40 µg lysate was resolved on a 12% gel, transferred on Nitrocellulose membrane, and probed with either α-FLAG or α-rpoA antibody (loading control), respectively. **(b-g)** Bar graphs (Mean ± SEM) depicting the abundance of various lipid classes **(b.)** TAG, **(c.)** DAG, **(d.)** FFA, **(e.)** PE, **(f.)** PI, **(g.)** CL in *Msm* and *Msm::wag31* treated with 5 mM IVN for 12 h. Values on the Y-axis represent area under the curve (AUC) normalised to an internal standard. X-axis represent various species (chain length variations) of the above mentioned lipid classes. The experiment was performed with six biological replicates, and the abundances of identified lipid classes were normalized to *Msm*-IVN to yield relative abundances of lipids in *Msm::wag31* + IVN.

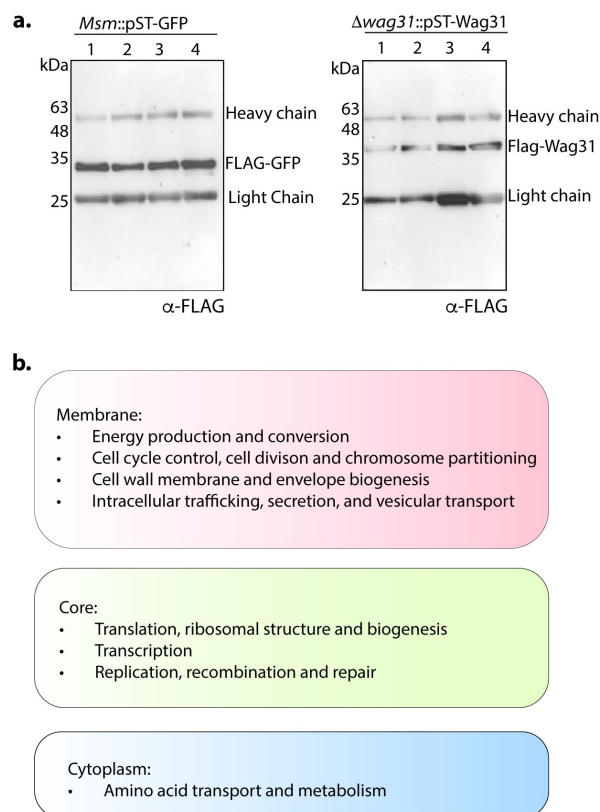


Figure S3.

Molecular interactions of wag31 with membrane proteins govern cellular homeostasis.

(a.) Western blots showing FLAG-GFP (left panel) and FLAG-Wag31 (right panel) immunoprecipitations. 3 mg lysate from each group was processed for overnight FLAG immunoprecipitations, resolved on a 10% gel, and probed with α -Flag antibody. HC and LC are short for Heavy chain and light chain, respectively. **(b.)** Classification of cellular processes that occur in different cellular compartments of *Msm*, as shown in (48 [48](#)).

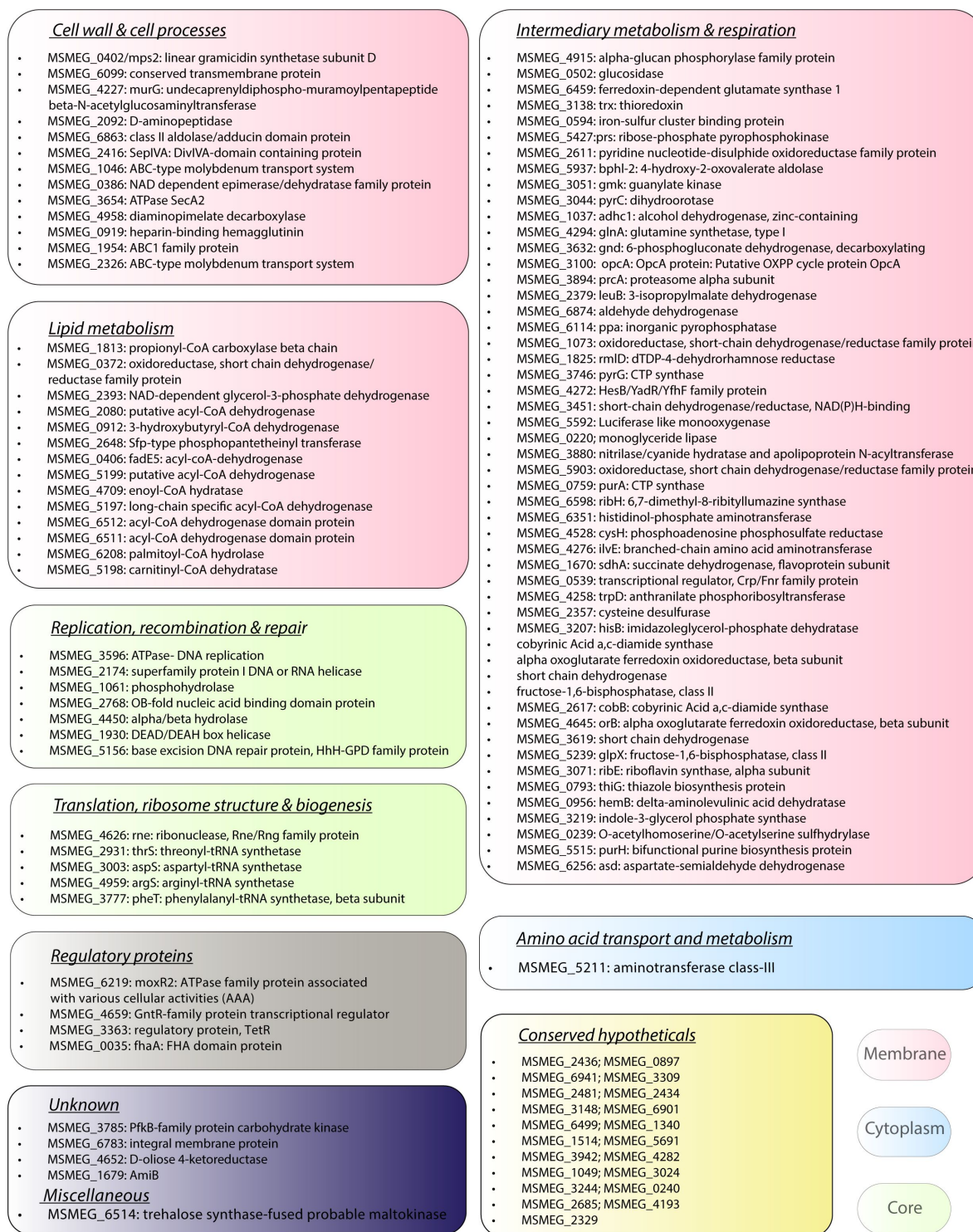


Figure S4.

Molecular interactions wag31 with membrane proteins govern cellular homeostasis.

The list of Wag31 interactors is classified according to functional categories based on the data available in Mycobrowser. The colour of the textbox indicates localization of the proteins: Pink-membrane, blue-cytoplasm, and core-green.

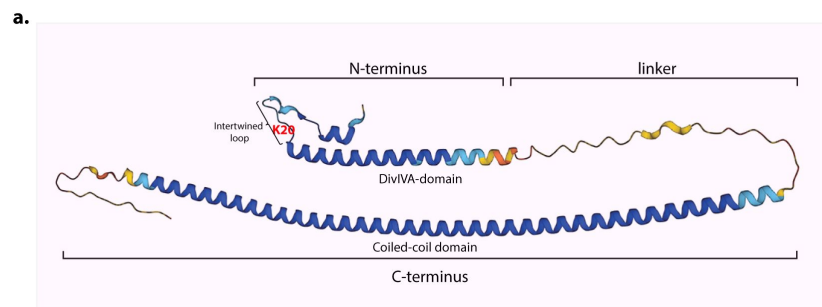


Figure S5.

Supplementary material

Materials and methods

Bacterial culturing

Antibiotic concentrations used: Hygromycin (Hyg) was used at 100 µg/mL, Kanamycin (Kan) at 25 µg/mL, and Apramycin (Apra) at 30 µg/mL for *Msm*. *E. coli* was cultured in Miller Luria Bertani Broth (Himedia) or Miller Luria Bertani Agar. Antibiotic concentrations used: Hyg was used at 150 µg/mL, Kan at 50 µg/mL, Apra at 30 µg/mL, Ampicillin (Amp) at 100 µg/mL for *E. coli*.

Generation of *wag31* conditional mutant

First, *wag31* was cloned under a tet regulatable promoter in pST-KiT_{off} (kan^{res}; [68](#)), which turns off transcription upon the addition of anhydrotetracycline. pST-*wag31* construct thus generated was electroporated into a recombineering proficient strain, *Msm::pJV53*. An allelic exchange substrate was generated by cloning 675 bp from upstream and 700 bp from downstream of *wag31* (including ~150 bp at both ends) with *hyg^{res}* cassette and *oriE*+*cosλ* and digested with SnaBI to release the donor LHS-*hyg^{res}*-RHS. The linear donor was electroporated into *Msm::wag31* expressing gp60 and gp61 recombinases. The colonies were selected on Hyg and ten colonies were screened with multiple primer pairs to confirm recombination at the native locus. Confirmed recombinants were passaged multiple times to cure them of pJV53 to generate conditional mutant *Δwag31*.

BODIPY imaging and analysis

For *Msm* and *Δwag31*, ~10 ml of cells were spotted on a 1% agar pad (prepared in 7H9) and visualized at 4X zoom through Confocal microscopy (LSM880) at 488 nm/510 nm (Excitation: Emission). Background correction in the collected images followed by Corrected Total Cell Fluorescence analysis was performed using Fiji ([69](#)) and plotted using Graph Pad Prism 9. For examining lipids in BODIPY-stained *Msm* and *Msm::wag31*, 2 ml of cells were spotted on a 1% agar pad (prepared in 7H9) and visualized using a 100X/1.32 NA oil immersion objective on a Leica THUNDER Imaging systems at 475nm:519nm (Excitation: Emission). About 20 Z-stack images, each spaced 0.1 µm apart, were acquired using a scientific CMOS K8 camera (Leica microsystems). Background correction and Thunder image processing was performed on the images using the Las-X software module (Leica microsystems). Processed images were then analysed for CTCF using Fiji ([69](#)) and plotted using Graph Pad prism 9.

Lipid extraction and analysis

The dried lipid extracts were re-solubilized in 200 µL of 2:1 CHCl₃: MeOH, and 10 µL was injected into an Agilent 6545 QTOF (quadrupole-time-of-flight) instrument for semi-quantitative analysis using high-resolution auto MS/MS methods and chromatography techniques. A Gemini 5U C-18 column (Phenomenex) coupled with a Gemini guard column (Phenomenex, 4x3 mm, Phenomenex security cartridge) was used for LC separation. The solvents used for negative ion mode were buffer A: 95:5 H₂O:MeOH + 0.1% ammonium hydroxide and buffer B: 60:35:5 Isopropanol:MeOH:H₂O + 0.1% ammonium hydroxide. The 0.1% ammonium hydroxide in each buffer was replaced by 0.1% Formic acid + 10 mM Ammonium Formate for positive ion mode runs. All LC/MS runs were for 60 min, starting with 0.3 mL/min 100% buffer A for 5 minutes, 0.5 mL/min linear gradient to 100% buffer B over 40 minutes, 0.5 mL/min 100% buffer B for 10 minutes, and equilibration with 0.5 mL/min 100% buffer A for 5 minutes. The following settings were used for the ESI-MS analysis: drying gas and sheath gas temperature: 320°C, drying gas and sheath gas flow

rate: 10 L/min, fragment or voltage: 150 V, capillary voltage: 4000 V, nebulizer (ion source gas) pressure: 45 psi and nozzle voltage: 1000 V. For analysis, a lipid library was employed in the form of a Personal Compound Database Library (PCDL), and the peaks were validated based on relative retention times and fragments obtained. This library was selectively curated from the MycoMass database and the LIPID MAPS Structure Database (LMSD). All robustly detected lipid species were quantified by normalizing areas under the curve (AUC) to the AUC of the relevant internal standard added and by normalizing to the total cell number. Log₂ fold changes were then plotted for each lipid compared to the Msm-ATc samples.

Immunoprecipitation (IP) sample preparation and MS/MS analysis

Sample preparation and, mass spectrometry analysis were outsourced to VProteomics, New Delhi. Briefly, the IP samples were reduced with 5 mM TCEP, followed by alkylation with 50 mM iodoacetamide. The samples were digested with Trypsin (1:50, Trypsin/lysate ratio) for 16 h at 37 °C. Digests were cleaned using a C18 silica cartridge and resuspended in a buffer containing 2% acetonitrile and 0.1% formic acid. 1 mg of peptide was used for analysis in an Easy-nlc-1000 system coupled to an Orbitrap Exploris mass spectrometer. The gradients were run for 110 min. MS1 spectra were acquired in the Orbitrap (Max IT = 60 ms, AGQ target = 300%; RF Lens = 70%; R=60K, mass range = 375–1500; Profile data). Dynamic exclusion was employed for 30 seconds, excluding all charge states for a given precursor. MS2 spectra were collected for the top 20 peptides. MS2 (Max IT=60ms, R= 15K, AGC target 100%). Proteome Discoverer (v2.5) was used to analyse the RAW data against the Uniprot *Msm* database. The precursor and fragment mass limitations for the dual Sequest and Amanda searches were established at 10 ppm and 0.02 Da, respectively. The false discovery rate for proteins and peptide spectrum matches was adjusted to 0.01 FDR. Proteins found in at least three replicates with minimum unique peptides #2 were considered interactors. To identify top hits, PSM cut-off was set to 18 and unique peptides to 5.

Protein expression and purification

BL21(DE3) cells transformed with pET-wag31, or pET-wag31K20A or pET-wag31_{D1-60} were cultured at 37°C until the A₆₀₀ ~0.6. 0.1 mM IPTG was added to the culture to induce protein expression, followed by incubating the culture at 18°C for 12 h. Cells were pelleted and stored at -40 °C. The bacterial pellet was resuspended in a lysis buffer (20 mM HEPES pH 7.4, 150 mM NaCl, and 20 mM imidazole with 1 mM phenylmethylsulphonyl fluoride (PMSF)) and lysed by sonication in an ice-water bath. Lysate was spun at 30,000 g for 20 min, and the supernatant obtained was incubated with TALON[®] Metal Affinity Resin (Takara Bio) 4 °C for 30 min. The resin was extensively washed with lysis buffer, and the bound protein was eluted with 20 mM HEPES pH 7.4, 150 mM NaCl, and 250 mM imidazole. The elution was dialyzed against 20 mM HEPES pH 7.4 150 mM NaCl and stored at 4 °C for the duration of the experiments. Aggregates were removed by centrifugation at 100,000 g. Concentration of the proteins was quantified by measuring A₂₈₀ using the molar extinction coefficient predicted by the ExPASy ProtParam tool.

Liposome preparation

1,2-dioleoyl-sn-glycero-3-phosphocholine (DOPC), 1,2-dioleoyl-sn-glycero-3-phosphoethanolamine (DOPE) and 1',3'-bis[1,2-dioleoyl-sn-glycero-3-phospho]-glycerol (sodium salt), CL were obtained from Avanti Polar Lipids. 1,1'-dilinoleyl-3,3',3'-tetramethylindocarbocyanine, 4-chlorobenzenesulfonate (FAST DiI[™]), and 3,3'-dilinoleyloxycarbocyanine perchlorate (FAST DiO[™]) were obtained from Invitrogen. The UV-activable, diazirine-containing reactive fluorescent lipid probe BODIPY-diazirine phosphatidylethanolamine (BDPE) was prepared as described earlier (68 [68](#), 69 [69](#)). For liposome preparation, lipids were aliquoted from chloroform stocks at desired ratios in a clean glass tube and dried under a high vacuum for an hour to a thin film. The dried lipids were used at a final concentration of 1 mM (deionised water was added to them). After hydrating at 50 °C for 30 minutes, lipids were vigorously vortexed and then extruded through polycarbonate filters with pore sizes of 100 nm (Whatman).. For PLiMAP assays, liposomes were

CL, DOPG, or DOPE with 69 mol% DOPC and 1 mol% BDPE. Liposomes used in tethering assays were made similarly with 30 mol% CL or DOPE and 1 mol% of the fluorescent lipid DiI or DiO) in the DOPC background.

NAO imaging analysis

Background correction and Thunder image processing was performed on the images using the Las-X software module (Leica microsystems). Processed images were then analysed using Fiji (69) and plotted using Graph Pad prism 9. Briefly, N=100 cells across two independent experiments were examined for their profile of CL distribution by normalizing the cell length and assigning pole with brighter NAO fluorescence intensity as 0 and the other pole as 100. The fluorescence intensity was normalised to the highest fluorescent value for each cell. Corresponding fluorescence intensity value for all cells for every point between 0 to 100 were calculated from the LOOKUP function of excel. The values representing CL fluorescence across 100 cells were averaged out and plotted against normalised cell length in GraphPad prism 9. 0th order smoothening with 4 neighbours on each size was performed on the curves obtained.

His pulldown assay-*E.coli* and *Msm* lysate preparation

BL21 (DE3) cells transformed with no plasmid, or pET-wag31 or pET-wag31_{D1-60} were cultured as described above and cell lysates were prepared. The expression of Wag31 or Wag31_{D1-60} was assessed by resolving the samples on 10% SDS-PAGE followed by Coomassie staining. The interacting partners, namely *murG*, *sepIVA*, *msm2092* and *accA3* were amplified from *Msm* genomic DNA using gene specific primers harbouring NdeI-HindIII sites and amplicons were digested and cloned into the corresponding sites in pNit-3X-FLAG shuttle vector. The constructs thus generated were electroporated in *Msm* to generate *Msm::murG*, *Msm::SepIVA*, *Msm::msm2092* and *Msm::accA3*. The cultures grown till A₆₀₀ ~0.8 were used for fresh inoculation at A₆₀₀ of 0.05 in 7H9 media containing 0.5 mM IVN. Cultures were grown for 12 h and lysates were prepared by bead-beating and 40 mg were resolved on 10% gels and probed with a-FLAG antibody.

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Plasmid	Description	Antibiotic resistance	Source
pJV53-Apra	pJV53 plasmid from Dr. Hatfull's lab wherein the <i>kan^r</i> antibiotic cassette was swapped with <i>apra^r</i> cassette.	apra	(3)
pST-KiT_{off}	L5 integrative pST-KiT plasmid harboring reverse tetR gene.	kan	(4)
pST-gfp	pST-KiT _{off} construct harboring gfp between NdeI-HindIII sites.	Kan	This study
pST-wag31	pST-KiT _{off} L5 integrative plasmid harboring FLAG- <i>wag31</i> -3X HA under the control of tetracycline induction	kan	This study
pET-wag31	pET28b harboring His- <i>wag31</i> -His	kan	This study
pET-wag31_{K20A}	pET28b harboring His- <i>wag31_{K20A}</i> -His	kan	This study
pET-wag31_{Δ1-60}	pET28b harboring His- <i>wag31_{Δ1-60}</i> -His	kan	This study
pET-wag31-gfp	pET28b harboring His- <i>wag31-gfp</i> -His	kan	This study
pSCG-wag31	Giles integrative plasmid harboring His- <i>wag31</i> -His under the control of <i>wag31</i> native promoter	apra	This study
pSCG- <i>wag31_{K20A}</i>	Giles integrative plasmid harboring His- <i>wag31_{K20A}</i> -His under the control of <i>wag31</i> native promoter	apra	This study
pSCG- <i>wag31_{Δ1-60}</i>	Giles integrative plasmid harboring His- <i>wag31_{Δ1-60}</i> -His under the control of <i>wag31</i> native promoter	apra	This study
pNiT-wag31	IVN inducible plasmid harboring 3X-FLAG- <i>wag31</i>	kan	This study
pNiT-<i>murG</i>	IVN inducible plasmid harboring 3X-FLAG- <i>murG</i>	kan	This study
pNiT-<i>sepIVA</i>	IVN inducible plasmid harboring 3X-FLAG- <i>sepIVA</i>	kan	This study
pNiT-<i>msm2092</i>	IVN inducible plasmid harboring 3X-FLAG- <i>msm2092</i>	kan	This study
pNiT-<i>accA3</i>	IVN inducible plasmid harboring 3X-FLAG- <i>accA3</i>	kan	This study

Table S1.

Plasmids and strains used in the study.

<i>E. coli</i> DH5α	<i>E. coli</i> strain used for cloning	none	Invitrogen
<i>E. coli</i> BL21DE3	<i>E. coli</i> strain used for protein expression	kan	Novagen
<i>Msm</i>	<i>M. smegmatis</i> mc ² 155 strain	none	ATCC
<i>Msm</i>::pJV53	<i>Msm</i> strain electroporated with pJV53 plasmid that expresses recJ and	apra	This study
<i>Msm</i>::gfp	<i>Msm</i> strain electroporated with pST-KiT-gfp		
<i>Msm</i>::pS-wag31	<i>gp60</i> and <i>gp61</i> recombinase expressing <i>Msm</i> ::pJV53 strain electroporated harboring integrative pST-KiT _{off}	apra, kan	This study
Δwag31	wag31 mutant in <i>Msm</i> in which the native wag31 gene was replaced with hyg ^r marker. The strain harbours integrative tet inducible pST-wag31 at L5 site. Cured to remove pJV53 episomal copy.	hyg, kan	This study
Δwag31::wag31	Δ wag31 electroporated with pSCG-wag31	hyg, kan, apra	This study
Δwag31::wag31_{K20A}	Δ wag31 electroporated with pSCG-wag31 _{K20A}	hyg, kan, apra	This study
Δwag31::wag31_{Δ1-60}	Δ wag31 electroporated with pSCG-wag31 _{Δ1-60}	hyg, kan, apra	This study
<i>Msm</i>::wag31	<i>Msm</i> electroporated with pNiT-wag31	kan	This study
<i>Msm</i>::murG	<i>Msm</i> electroporated with pNiT-murG	kan	This study
<i>Msm</i>::sepIVA	<i>Msm</i> electroporated with pNiT-sepIVA	kan	This study
<i>Msm</i>::msm2092	<i>Msm</i> electroporated with pNiT-msm2092	kan	This study
<i>Msm</i>::accA3	<i>Msm</i> electroporated with pNiT-accA3	kan	This study

Table S1. (continued)

Accession	Gene ID	# PSMs	# Unique Peptides	Localisation (Based on PMID: 34965429, 27114527, 20412585)	Functional category
A0QYB1	MSMEG_3596	185	37	Unknown	replication, recombination & repair
A0QTE7	MSMEG_1813	104	16	IMD	lipid metabolism
Q3L891	MSMEG_0402	58	16	PM-CW	cell wall & cell processes
A0R1Y2	MSMEG_4915	61	16	PM-CW	Intermediary metabolism and respiration
A0R152	:MSMEG_4626	37	13	IMD	Translation, ribosomal structure & biogenesis
A0QWG2	MSMEG_2931	44	14	PM-CW	Translation, ribosomal structure & biogenesis
A0QPS6	MSMEG_0502	34	10	unknown	Intermediary metabolism and respiration
A0R684	MSMEG_6459	29	11	Cytosolic	Intermediary metabolism and respiration
A0R582	MSMEG_6099	41	10	PM-CW	Cell wall and cell processes
A0R016	MSMEG_4227	34	9	IMD	Cell wall and cell processes
A0QQW2	MSMEG_0897	20	7	unknown	conserved hypothetical
A0QU64	MSMEG_2092	41	7	unknown	Cell wall and cell processes
A0R7C9	MSMEG_6863	25	5	unknown	cell wall & cell processes
A0QX15	MSMEG_3138	28	4	Cytosolic	Intermediary metabolism and respiration
A0QPE7	MSMEG_0372	24	6	IMD	Lipid metabolism
A0QV38	MSMEG_2436	21	4	IMD	unknown
A0QUZ5	MSMEG_2393	24	6	Cytosolic	Lipid metabolism
A0QV18	MSMEG_2416	13	3	Cell wall	cell wall & cell processes
A0QRA5	EG_1046, MSMEC	12	4	Cell wall	Cell wall and cell processes
A0QUE0	MSMEG_2174	25	7	Cell wall	replication, recombination & repair
A0R2R8	MSMEG_5211	23	6	unknown	Amino acid transport and metabolism
A0R7J6	MSMEG_6941	18	5	IMD	conserved hypotheticals
A0QU53	MSMEG_2080	18	6	IMD	lipid metabolism
A0QQ17	MSMEG_0594	16	6	IMD	Intermediary metabolism and respiration
A0R3C8	MSMEG_5427	20	7	unknown	Intermediary metabolism and respiration
A0QVL0	MSMEG_2611	18	4	unknown	Intermediary metabolism and respiration
A0QX10	MSMEG_3309	10	3	unknown	conserved hypothetical
A0QPG1	MSMEG_0386	7	3	unknown	Cell wall and cell processes
A0R4S6	MSMEG_5937	18	4	unknown	Intermediary metabolism and respiration
A0QV82	MSMEG_2481	11	4	unknown	conserved hypothetical
A0QWS9	MSMEG_3051	16	3	unknown	Intermediary metabolism and respiration
A0QWS2	MSMEG_3044	14	4	unknown	Intermediary metabolism and respiration
P0CH36	MSMEG_1037	16	4	unknown	Intermediary metabolism and respiration
A0R083	MSMEG_4294	10	3	unknown	Intermediary metabolism and respiration
A0QQX7	MSMEG_0912	14	4	unknown	lipid metabolism
A0QYE7	MSMEG_3632	10	3	PM-CW	Intermediary metabolism and respiration
A0QYG9	MSMEG_3654	13	4	PM-CW	cell wall & cell processes
A0QV36	MSMEG_2434	10	3	unknown	conserved hypotheticals
A0QVP7	MSMEG_2648	11	4	cell wall	lipid metabolism
A0QWX7	MSMEG_3100	14	4	unknown	Intermediary metabolism and respiration
Q3L887	MSMEG_0406	16	3	unknown	lipid metabolism
A0QX25	MSMEG_3148	15	4	Cell wall	conserved hypothetical
A0QZ46	MSMEG_3894	10	3	IMD	Intermediary metabolism and respiration
A0QUY3	MSMEG_2379	10	3	unknown	Intermediary metabolism and respiration
A0QRB9	MSMEG_1061	10	3	cell wall	replication, recombination & repair
A0R7D8	MSMEG_6874	13	3	unknown	Intermediary metabolism and respiration
Q9X5M1	MSMEG_4958	16	4	unknown	cell wall & cell processes
A0R7G3	MSMEG_6901	14	3	unknown	conserved hypothetical
A0R5K0	MSMEG_6219	12	1	PM-CW	regulatory protein
A0QW11	MSMEG_2768	19	3	unknown	transcription, replication, recombination & repair
A0R597	MSMEG_6114	10	3	unknown	Intermediary metabolism and respiration
A0QWN3	MSMEG_3003	10	2	unknown	Translation, ribosomal structure & biogenesis
A0QRD2	MSMEG_1073	14	4	IMD	Intermediary metabolism and respiration
A0R0N6	MSMEG_4450	10	3	IMD	replication, recombination & repair
A0QTF8	MSMEG_1825	10	4	unknown	Intermediary metabolism and respiration
A0R6C4	MSMEG_6499	10	2	unknown	conserved hypotheticals
A0QYQ7	MSMEG_3746	12	3	Cell wall	Intermediary metabolism and respiration
A0R184	MSMEG_4659	12	3	unknown	regulatory protein
A0R061	MSMEG_4272	10	2	cell wall	Intermediary metabolism and respiration

Table S2.

It depicts unique interacting partners found for Wag31.

A0QXW7	MSMEG_3451	10	2	unknown	Intermediary metabolism and respiration
A0QYU0	MSMEG_3785	9	3	unknown	Unknown
A0R2Q7	MSMEG_5199	9	2	unknown	lipid metabolism
A0R1D1	MSMEG_4709	8	3	unknown	lipid metabolism
A0QTQ8	MSMEG_1930	8	3	IMD	replication, recombination & repair
A0R2Q5	MSMEG_5197	8	3	unknown	lipid metabolism
A0R3T9	MSMEG_5592	8	2	unknown	Intermediary metabolism and respiration
A0R6D9	MSMEG_6514	8	2	unknown	Virulence, detoxification, adaptation
A0QNZ7	AMSMEG_0220	8	2	IMD	Intermediary metabolism and respiration
A0QZ33	MSMEG_3880	8	2	unknown	Intermediary metabolism and respiration
A0R4P1	MSMEG_5903	8	2	unknown	Intermediary metabolism and respiration
A0QS40	MSMEG_1340	8	2	unknown	conserved hypotheticals
A0QQH7	MSMEG_0759	8	2	unknown	Intermediary metabolism and respiration
A0QSK8	MSMEG_1514	8	2	unknown	conserved hypotheticals
A0R6M2	MSMEG_6598	8	2	unknown	Intermediary metabolism and respiration
A0R436	MSMEG_5691	7	3	unknown	conserved hypothetical protein
A0R5X8	MSMEG_6351	7	3	unknown	Intermediary metabolism and respiration
A0QT17	MSMEG_1679	7	2	Cell wall	unknown
A0QXN2	MSMEG_3363	7	2	unknown	regulatory protein
A0R0W2	MSMEG_4528	7	2	unknown	Intermediary metabolism and respiration
A0R066	MSMEG_4276	7	2	unknown	Intermediary metabolism and respiration
A0QT08	MSMEG_1670	6	3	PM-CW	Intermediary metabolism and respiration
A0QPW2	MSMEG_0539	6	3	PM-CW	Intermediary metabolism and respiration
Q3I5Q7	MSMEG_0919	6	3	unknown	Cell wall and cell processes
A0QZ93	MSMEG_3942	6	3	unknown	conserved hypothetical protein
A0R6D7	MSMEG_6512	6	2	unknown	lipid metabolism
A0R071	MSMEG_4282	6	2	IMD	conserved hypothetical protein
A0R048	MSMEG_4258	6	2	PM-CW	Intermediary metabolism and respiration
A0R753	MSMEG_6783	6	2	unknown	Unknown
A0QRA8	EG_1049;MSMEG	6	2	unknown	Conserved hypotheticals
Q9X5M0	MSMEG_4959	6	2	PM-CW	Translation, ribosomal structure & biogenesis
A0QUW2	MSMEG_2357	6	2	unknown	Intermediary metabolism and respiration
A0QNG7	MSMEG_0035	6	2	PM-CW	regulatory protein
A0QX83	MSMEG_3207	6	2	unknown	Intermediary metabolism and respiration
A0QVL6	MSMEG_2617	6	2	unknown	Intermediary metabolism and respiration
A0R170	MSMEG_4645	6	2	unknown	Intermediary metabolism and respiration
A0QYD4	MSMEG_3619	6	2	PM-CW	Intermediary metabolism and respiration
A0R2U7	MSMEG_5239	6	2	unknown	Intermediary metabolism and respiration
A0R6D6	MSMEG_6511	6	2	unknown	lipid metabolism
A0QWQ3	MSMEG_3024	5	2	unknown	Conserved hypotheticals
A0R177	MSMEG_4652	5	2	unknown	Unknown
A0QXB8	MSMEG_3244	5	2	unknown	conserved hypothetical protein
A0QTT2	MSMEG_1954	4	2	PM-CW	Cell wall and cell processes
A0QP17	MSMEG_0240	4	2	unknown	conserved hypothetical protein
A0QYT3	MSMEG_3777	4	2	unknown	Translation, ribosomal structure & biogenesis
A0R2L4	MSMEG_5156	4	2	unknown	replication, recombination & repair
A0R5I9	MSMEG_6208	4	2	unknown	lipid metabolism
A0QWU9	MSMEG_3071	4	2	PM-CW	Intermediary metabolism and respiration
A0QQL0	MSMEG_0793	4	2	unknown	Intermediary metabolism and respiration
A0QR20	MSMEG_0956	4	2	unknown	Intermediary metabolism and respiration
A0QVT2	MSMEG_2685	4	2	unknown	Conserved hypotheticals
A0R2Q6	:MSMEG_5198	4	2	unknown	lipid metabolism
A0QX95	MSMEG_3219	4	2	Cell wall	Intermediary metabolism and respiration
A0QP16	MSMEG_0239	4	2	unknown	Intermediary metabolism and respiration
A0QZY4	MSMEG_4193	4	2	Cytosolic	Conserved hypotheticals
A0R3L4	MSMEG_5515	4	2	Cytosolic	Intermediary metabolism and respiration
A0R5N7	MSMEG_6256	3	2	PM-CW	Intermediary metabolism and respiration

Table S2. (continued)

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Reviewer #1 (Public review):

This a comprehensive study that sheds light on how Wag31 functions and localises in mycobacterial cells. A clear link to interactions with CL is shown using a combination of microscopy in combination with fusion fluorescent constructs, and lipid specific dyes. Furthermore, studies using mutant versions of Wag31 shed light on the functionalities of each domain in the protein. My concerns/suggestions for the manuscript are minor:

- (1) Ln 130. A better clarification/discussion is required here. It is clear that both depletion and overexpression have an effect on levels of various lipids, but subsequent descriptions show that they affect different classes of lipids.
- (2) The pulldown assays results are interesting, but links are tentative.
- (3) The authors may perhaps like to rephrase claims of effects lipid homeostasis, as my understanding is that lipid localisation rather than catabolism/breakdown is affected.

<https://doi.org/10.7554/eLife.104268.1.sa3>

Reviewer #2 (Public review):

Summary:

Kapoor et. al. investigated the role of the mycobacterial protein Wag31 in lipid and peptidoglycan synthesis and sought to delineate the role of the N- and C- terminal domains of Wag31. They demonstrated that modulating Wag31 levels influences lipid homeostasis in *M. smegmatis* and cardiolipin (CL) localisation in cells. Wag31 was found to preferentially bind CL-containing liposomes, and deleting the N-terminus of the protein significantly decreased this interaction. Novel interactions between Wag31 and proteins involved in lipid metabolism

and cell wall synthesis were identified, suggesting that Wag31 recruits proteins to the intracellular membrane domain by direct interaction.

Strengths:

- (1) The importance of Wag31 in maintaining lipid homeostasis is supported by several lines of evidence.
- (2) The interaction between Wag31 and cardiolipin, and the role of the N-terminus in this interaction was convincingly demonstrated.

Weaknesses:

- (1) MS experiments provide some evidence for novel protein-protein interactions, however, the pull-down experiments are lacking a valid negative control.
- (2) The role of the N-terminus in the protein-protein interaction has not been ruled out.

<https://doi.org/10.7554/eLife.104268.1.sa2>

Reviewer #3 (Public review):

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This manuscript describes the characterization of mycobacterial cytoskeleton protein Wag31, examining its role in orchestrating protein-lipid and protein-protein interactions essential for mycobacterial survival. The most significant finding is that Wag31, which directs polar elongation and maintains the intracellular membrane domain, was revealed to have membrane tethering capabilities.

Strengths:

The authors provided a detailed analysis of Wag31 domain architecture, revealing distinct functional roles: the N-terminal domain facilitates lipid binding and membrane tethering, while the C-terminal domain mediates protein-protein interactions. Overall, this study offers a robust and new understanding of Wag31 function.

Weaknesses:

The following major concerns should be addressed.

- Authors use 10-N-Nonyl-acridine orange (NAO) as a marker for cardiolipin localization. However, given that NAO is known to bind to various anionic phospholipids, how do the authors know that what they are seeing is specifically visualizing cardiolipin and not a different anionic phospholipid? For example, phosphatidylinositol is another abundant anionic phospholipid in mycobacterial plasma membrane.
- Authors' data show that the N-terminal region of Wag31 is important for membrane tethering. The authors' data also show that the N-terminal region is important for sustaining mycobacterial morphology. However, the authors' statement in Line 256 "These results highlight the importance of tethering for sustaining mycobacterial morphology and survival" requires additional proof. It remains possible that the N-terminal region has another unknown activity, and this yet-unknown activity rather than the membrane tethering activity drives the morphological maintenance. Similarly, the N-terminal region is important for lipid homeostasis, but the statement in Line 270, "the maintenance of lipid homeostasis by Wag31 is a consequence of its tethering activity" requires additional proof. The authors should tone down these overstatements, or provide additional data to support their claims.

• Authors suggest that Wag31 acts as a scaffold for the IMD (Fig. 8). However, Meniche et. al. has shown that MurG as well as GlfT2, two well-characterized IMD proteins, do not colocalize with Wag31 (DivIVA) (<https://doi.org/10.1073/pnas.1402158111>). IMD proteins are always slightly subpolar while Wag31 is located to the tip of the cell. Therefore, the authors' biochemical data cannot be easily reconciled with microscopic observations in the literature. This raises a question regarding the validity of protein-protein interaction shown in Figure 7. Since this pull-down assay was conducted by mixing E. coli lysate expressing Wag31 and Msm lysate expression Wag31 interactors like MurG, it is possible that the interactions are not direct. Authors should interpret their data more cautiously. If authors cannot provide additional data and sufficient justifications, they should avoid proposing a confusing model like Figure 8 that contradicts published observations.

<https://doi.org/10.7554/eLife.104268.1.sa1>

Author response:

Public Reviews:

Reviewer #1 (Public review):

This a comprehensive study that sheds light on how Wag31 functions and localises in mycobacterial cells. A clear link to interactions with CL is shown using a combination of microscopy in combination with fusion fluorescent constructs, and lipid specific dyes. Furthermore, studies using mutant versions of Wag31 shed light on the functionalities of each domain in the protein. My concerns/suggestions for the manuscript are minor:

(1) Ln 130. A better clarification/discussion is required here. It is clear that both depletion and overexpression have an effect on levels of various lipids, but subsequent descriptions show that they affect different classes of lipids.

We thank the reviewer for the comments. We will improve Ln130 in the manuscript. The lipid classes that get impacted by the depletion of Wag31 vs overexpression are different. Wag31 is an adaptor protein that interacts with proteins of the ACCase complex (Meniche et al., 2014; Xu et al., 2014) that synthesize fatty acid precursors and regulate their activity (Habibi Arejan et al., 2022).

The varied response to lipid homeostasis could be attributed to a change in the stoichiometry of these interactions with Wag31. While Wag31 depletion would prevent such interactions from occurring and might affect lipid synthesis that directly depends on Wag31-protein partner interactions, its overexpression would lead to promiscuous interactions and a change in the stoichiometry of native interactions, ultimately modulating lipid synthesis pathways.

(2) The pulldown assays results are interesting, but links are tentative.

The interactome of Wag31 was identified through the immunoprecipitation of Flag-tagged Wag31 complemented at an integrative locus in Wag31 mutant background to avoid overexpression artifacts. We used Msm::gfp expressing an integrative copy (at L5 locus) of FLAG-GFP as a control to subtract non-specific interactions. The experiment was performed in biological triplicates, and interactors that appeared in all replicates were selected for further analysis. Although we identified more than 100 interactors of Wag31, we analyzed only the top 25 hits, with a PSM cut-off ≥ 18 and unique peptides ≥ 5 . Additionally, two of Wag31's established interactors, AccD5 and Rne, were among the top five hits, thus validating our data.

Though we agree that the interactions can either be direct or through a third partner, the fact that we obtained known interactors of Wag31 makes us believe these interactions are genuine. Moreover, we performed pulldown experiments for validation by mixing *E. coli* lysates expressing His-Wag31 full-length or truncated protein with *M. smegmatis* lysates expressing FLAG-tagged interacting proteins. The wash conditions used were quite stringent for these pull-down assays—the wash buffer contained 1% Triton X100, eliminating all non-specific and indirect interactions. However, we agree that we cannot conclusively state that the interactions are direct without purifying the proteins and performing the experiment. We will describe this caveat in the revised manuscript.

(3) The authors may perhaps like to rephrase claims of effects lipid homeostasis, as my understanding is that lipid localisation rather than catabolism/breakdown is affected.

In this manuscript, we are trying to convey that Wag31 is a spatiotemporal regulator of lipid metabolism. It is a peripheral protein that is hooked to the membrane via Cardiolipin and forms a scaffold at the poles, which helps localize several enzymes involved in lipid metabolism.

Homeostasis is the process by which an organism maintains a steady-state of balance and stability in response to changes. Depletion of Wag31 not only results in delocalisation of lipids in intracellular lipid inclusions but also leads to changes in the levels of various lipid classes. Advancement in the field of spatial biology underscores the importance of native localization of various biological molecules crucial for maintaining a steady-cell of the cell. Hence, we have used the word “homeostasis” to describe both the changes observed in lipid metabolism.

Reviewer #2 (Public review):

Summary

*Kapoor et. al. investigated the role of the mycobacterial protein Wag31 in lipid and peptidoglycan synthesis and sought to delineate the role of the N- and C- terminal domains of Wag31. They demonstrated that modulating Wag31 levels influences lipid homeostasis in *M. smegmatis* and cardiolipin (CL) localisation in cells. Wag31 was found to preferentially bind CL-containing liposomes, and deleting the N-terminus of the protein significantly decreased this interaction. Novel interactions between Wag31 and proteins involved in lipid metabolism and cell wall synthesis were identified, suggesting that Wag31 recruits proteins to the intracellular membrane domain by direct interaction.*

Strengths:

(1) The importance of Wag31 in maintaining lipid homeostasis is supported by several lines of evidence.

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Weaknesses:

(1) MS experiments provide some evidence for novel protein-protein interactions. However, the pull-down experiments lack a valid negative control.

We thank the reviewer for the comments. We will include a valid negative control in the experiment. We would choose ~2 mycobacterial proteins that are not a part of our interactome study and perform a similar pull-down experiment with them and a positive control (known interactor of Wag31).

(2) *The role of the N-terminus in the protein-protein interaction has not been ruled out.*

Previously, we attempted to express the N-terminal (1-60 aa) and the C-terminal (60-212 aa) proteins in various mycobacterial shuttle vectors to perform MS/MS experiments. Despite numerous efforts, neither was expressed with the N/C-terminal FLAG tag nor without any tag in episomal or integrative vectors due to the instability of the protein. Eventually, we successfully expressed the C-terminal Wag31 with an N and C-terminal hexa-His tag. However, this expression was not sufficient or stable enough for us to perform Ni affinity pull-down experiments for mass spectrometry. The N-terminal of Wag31 could not be expressed in *M. smegmatis* even with N and C-terminal Hexa-His tags.

To rule out the role of the N-terminal in mediating protein-protein interactions, we plan to attempt to express N-terminal of Wag31 with N and C-terminal hexa-His tag in *E. coli*. If this clone successfully expresses in *E. coli*, we will perform pull-down experiments as described in Figure 7.

Reviewer #3 (Public review):

Summary:

This manuscript describes the characterization of mycobacterial cytoskeleton protein Wag31, examining its role in orchestrating protein-lipid and protein-protein interactions essential for mycobacterial survival. The most significant finding is that Wag31, which directs polar elongation and maintains the intracellular membrane domain, was revealed to have membrane tethering capabilities.

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The following major concerns should be addressed.

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We thank the reviewer for the comments. Despite its promiscuous binding to other anionic phospholipids, 10-N-Nonyl-acridine orange is widely used to stain Cardiolipin and determine its localisation in bacterial cells and mitochondria of eukaryotes (Garcia Fernandez et al., 2004; Mileykovskaya & Dowhan, 2000; Renner & Weibel, 2011). This is because it has a stronger affinity for Cardiolipin than other anionic phospholipids with the affinity constant being $2 \times 10^6 \text{ M}^{-1}$ for Cardiolipin association and $7 \times 10^4 \text{ M}^{-1}$ for that of phosphatidylserine and phosphatidylinositol association (Petit et al., 1992). Additionally, there is not yet another stain available for detecting Cardiolipin. Our protein-lipid binding assays suggest that Wag31 preferentially binds to Cardiolipin over other anionic phospholipids (Fig. 4b), hence it is likely that the majority of redistribution of NAO fluorescence that we observe might be contributed by Cardiolipin mislocalization due to altered Wag31 levels, with smaller degree of NAO redistribution intensity coming indirectly from other anionic phospholipids displaced from the membrane due to the loss of membrane integrity and cell shape changes due to Wag31.

• Authors' data show that the N-terminal region of Wag31 is important for membrane tethering. The authors' data also show that the N-terminal region is important for sustaining mycobacterial morphology. However, the authors' statement in Line 256 "These results highlight the importance of tethering for sustaining mycobacterial morphology and survival" requires additional proof. It remains possible that the N-terminal region has another unknown activity, and this yet-unknown activity rather than the membrane tethering activity drives the morphological maintenance. Similarly, the N-terminal region is important for lipid homeostasis, but the statement in Line 270, "the maintenance of lipid homeostasis by Wag31 is a consequence of its tethering activity" requires additional proof. The authors should tone down these overstatements or provide additional data to support their claims.

We agree with the reviewer that there exists a possibility for another function of the N-terminal that may contribute to sustaining mycobacterial physiology and survival. We would revise our statements in the paper to accurately reflect the data. Results shown suggest that the tethering activity of the N-terminal region may contribute to mycobacterial morphology and survival. However, additional functions of this region can't be ruled out. Similarly, the maintenance of lipid homeostasis by Wag31 may be associated with its tethering activity, although other mechanisms could also contribute to this process.

• Authors suggest that Wag31 acts as a scaffold for the IMD (Fig. 8). However, Meniche et al. has shown that MurG as well as GlfT2, two well-characterized IMD proteins, do not colocalize with Wag31 (DivIVA) (<https://doi.org/10.1073/pnas.1402158111>). IMD proteins are always slightly subpolar while Wag31 is located to the tip of the cell. Therefore, the authors' biochemical data cannot be easily reconciled with microscopic observations in the literature. This raises a question regarding the validity of protein-protein interaction shown in Figure 7. Since this pull-down assay was conducted by mixing *E. coli* lysate expressing Wag31 and *Msm* lysate expression Wag31 interactors like MurG, it is possible that the interactions are not direct. Authors should interpret their data more cautiously. If authors cannot provide additional data and sufficient justifications, they should avoid proposing a confusing model like Figure 8 that contradicts published observations.

In the literature, MurG and GlfT2 have been shown to have polar localization (Freeman et al., 2023; Hayashi et al., 2016; Kado et al., 2023), and two groups have shown slightly sub-polar localization of MurG (García-Heredia et al., 2021; Meniche et al., 2014). Additionally, (Freeman et al., 2023) they showed SepIVA to be a spatio-temporal regulator of MurG. MS/MS analysis of Wag31 immunoprecipitation data yielded both MurG and SepIVA to be interactors of Wag31 (Fig. 3). Given Wag31 also displays polar localisation, it likely associates with the polar MurG. However, since a sub-polar localization of MurG has also been reported, it is possible that they do not interact directly, and another protein mediates their interaction. We will modify the model proposed in Fig. 8 based on the above.

We agree that for validation of interaction, we performed pulldown experiments by mixing *E. coli* lysates expressing His-Wag31 full-length or truncated protein with *M. smegmatis* lysates expressing FLAG-tagged interacting proteins. The wash conditions used were quite stringent for these pull-down assays—the wash buffer containing 1% Triton X100, which eliminates all non-specific and indirect interactions. However, we agree that we cannot conclusively state that the interactions are direct without purifying the proteins and performing the experiment. We will describe this caveat in the revised manuscript and propose a model reflecting our results.

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