A conserved cell-pole determinant organizes proper polar flagellum formation

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Abstract

- 35 The coordination of cell cycle progression and flagellar synthesis is a complex process in motile bacteria. In γ-proteobacteria, the localization of the flagellum to the cell pole is mediated by the SRP-type GTPase FlhF. However, the mechanism of action of FlhF, and its relationship with the cell pole landmark protein HubP remain unclear. In this study, we discovered a novel protein called FipA that is required for normal FlhF activity and function in polar flagellar
- 40 synthesis. We demonstrated that membrane-localized FipA interacts with FlhF and is required for normal flagellar synthesis in *Vibrio parahaemolyticus, Pseudomonas putida*, and *Shewanella putrefaciens*, and it does so independently of the polar localization mediated by HubP. FipA exhibits a dynamic localization pattern and is present at the designated pole before flagellar synthesis begins, suggesting its role in licensing flagellar formation. This discovery
- 45 provides insight into a new pathway for regulating flagellum synthesis and coordinating cellular organization in bacteria that rely on polar flagellation and FlhF-dependent localization.

Introduction

Many cellular processes depend on a specific spatiotemporal organization of its components.

- 50 The DNA replication machinery, cell division proteins and motility structures are some examples of elements that need to be positioned at a particular site during the cell cycle in a coordinated manner to ensure adequate cell division or proper function of the structures. To carry out this task, many cellular components of bacteria have a polar organization, *i. e.* they are asymmetrically positioned inside the cell, which is particularly apparent in monopolarly
- 55 flagellated bacteria. In the marine bacteria *Vibrio*, which constitutively express a single polar flagellum (McCarter 1995; Kim and McCarter 2000), it is crucial to coordinate the localization and timing of flagellar components, in order to guarantee that newly born cells only produce a flagellum around cell division. This process is coordinated with the chromosome replication and the chemotaxis clusters through a supramolecular hub that is tethered at the old cell pole
- 60 (Takekawa et al. 2016; Yamaichi et al. 2012). This organization depends on ATPases of the ParA/MinD family, which regulate the migration of the hub components to the new cell pole as the cell cycle progresses. In this way, the new cell has a copy of the chromosome and a set of chemotaxis clusters positioned next to what will be the site for the new flagellum. Central to this process is the protein HubP, which recruits to the pole the three ATPases responsible
- 65 for the localization of these organelles: ParA1 for the chromosome (Fogel and Waldor 2006), ParC for the chemotaxis clusters (Ringgaard et al. 2011) and FlhG for the flagellum (Correa, Peng, and Klose 2005; Arroyo-Pérez and Ringgaard 2021). Orthologs of HubP occur in several species, e.g., *Pseudomonas* (where it is called FimV), *Shewanella* or *Legionella*, where they similarly act as organizers (hubs) of the cell pole (Wehbi et al. 2011; Rossmann et al. 2015; Coil and Ann (2016).
- 70 and Anné 2010).

The bacterial flagellum is a highly intricate and complex subcellular structure. It is composed of around 25 different proteins, which have to be assembled in different stoichiometries in a spatiotemporally coordinated manner (Macnab 2003; Chevance and Hughes, 2008). Although

75 there may be significant differences among species, flagella in general are composed of a motor attached to the cytoplasmic membrane, a rod connecting it to the extracellular space, a hook and a filament. The basal body is composed of a cytoplasmic C-ring or switch complex (Francis et al. 1994), which receives the signal from the chemotaxis proteins and transfers it to the MS ring, which is embedded in the cytoplasmic membrane (Homma et al. 1987; Ueno,

- 80 Oosawa, and Aizawa 1992). Attached to it, on the cytoplasmic side, is the export apparatus, which allows secretion of rod, hook and filament components (Minamino 2014). Flagella can also be sheathed, if an extension of the outer membrane covers the filament, as is the case in many *Vibrio* species (Chen et al. 2017).
- The positioning and counting of the flagella are mediated in many bacteria by the interplay between two flagellar regulators, FlhF and FlhG. Studies in a wide variety of proteobacteria indicate that FlhG is a negative regulator that restricts the number of flagella that are synthesized. In organisms such as strains of *Vibrio, Shewanella putrefaciens* and *Pseudomonas aeruginosa*, the absence of MinD-like ATPase FlhG results in hyper-flagellated cells (Kusumoto
- 90 et al. 2006; Arroyo-Pérez and Ringgaard 2021; Schuhmacher et al. 2015; Blagotinsek et al., 2020; Campos-García et al. 2000; Murray and Kazmierczak 2006). In contrast, in polarly flagellated species such as *Campylobacter jejuni, Vibrio cholerae, Vibrio parahaemolyticus, P. aeruginosa, S. putrefaciens* and *Shewanella oneidensis*, the SRP-type GTPase FlhF is a positive regulator of flagellum synthesis and necessary for proper localization. A deletion of *flhF* results
- in reduced number and mis-localization of flagella (Hendrixson and Dirita 2003; Correa, Peng, and Klose 2005; Arroyo-Pérez and Ringgaard 2021; Pandza et al. 2000; Rossmann et al. 2015; Gao et al. 2015, Navarrete et al, 2019).

The current model predicts that GTP-bound dimeric FlhF (Bange et al. 2007; Kondo et al. 2018) localizes to the cell pole to where it recruits the initial flagellar building blocks, e.g., the MS-

- 100 ring protein FliF (Green et al., 2009). A long-standing question is how FlhF recognizes and localizes to the designated cell pole. In *P. aeruginosa* and *V. alginolyticus* it was demonstrated that FlhF localized polarly upon ectopic production in the absence of the flagellar master regulator FlrA (Green et al. 2009; Kondo, Homma, and Kojima 2017). It was therefore speculated that FlhF assumes its polar localization in the absence of other flagellar proteins or
- 105 even without any further additional protein factors. The underlying mechanism, however, remains enigmatic, given that neither in its monomeric nor its dimeric form, FlhF possesses any regions that would indicate a membrane association.

Here, we provide evidence demonstrating that FlhF does not localize independently. Instead, we show that FipA, a small integral membrane protein with a domain of unknown function, is

110 (co-)responsible for recruiting FlhF to the membrane at the cell pole and, at least in some species, it acts in concert with the polar landmark protein HubP/FimV. Using *V*.

parahaemolyticus and two additional polarly flagellated γ -proteobacteria, the monopolarly flagellated *S. putrefaciens* and the lophotrichous *P. putida*, we show that FipA universally mediates recruitment of FlhF to the designated cell pole. The spatiotemporal localization

115 behavior of FipA as well as its relationship to FlhF, support its role as a licensing protein that enables flagellum synthesis.

Results

Identification of an FIhF protein interaction partner: FipA

- 120 In order to identify potential factors required for FlhF function and its recruitment to the cell pole, we performed affinity purification of a superfolder green fluorescent protein (sfGFP)tagged FlhF (FlhF-sfGFP) ectopically expressed in wild-type V. parahaemolyticus cells followed by shotgun proteomics using liquid chromatography-tandem mass spectrometry analysis (LC-MS/MS). Among the proteins that were significantly enriched in FlhF-sfGFP purifications, eight
- 125 were structural components of the flagellum (Fig. 1A; Supplementary Table S1), but also a non-flagellar protein, VP2224, was significantly co-purified with FlhF (Fig. 1A; Supplementary Table S1). The homologue of VP2224 in *V. cholerae*, there named FlrD, was previously shown to exert the same activating role on flagellar gene regulation as FlhF (Moisi et al. 2009), suggesting a function related to FlhF. Notably, FlhF was also significantly co-purified in the
- 130 reciprocal co-IP-MS/MS experiment using VP2224-sfGFP as bait (Fig. 1B), suggesting a direct or indirect interaction of VP2224 and FlhF. To further investigate the co-purification data, bacterial two-hybrid (BACTH) assays in the heterologous host *E. coli* were carried out and suggested a direct interaction between FlhF and VP2224. FlhF and VP2224 were also found to self-interact (Fig. 1E). Additionally, these data indicated that FlhF and VP2224 form an
- interaction complex in both the native and a heterologous host organism. Thus, we identified VP2224 as a novel interaction partner of FlhF and named it FipA for Flh<u>F</u> Interaction <u>P</u>artner <u>A</u>.

FipA constitutes a new family of FlhF interaction partners

140 The gene encoding FipA is located immediately downstream of the flagellar operon that encodes FlhA, FlhF, FlhG, FliA and the chemotaxis proteins (Fig. 1C). In *V. parahaemolyticus*, FipA consists of 163 amino acids with a molecular mass of 18.4 kDa. *In silico* analysis and membrane topology mapping predicted that FipA consists of a short periplasmic N-terminal

part, consisting of amino acids 1-5, followed by a transmembrane region (6-28), and a cytoplasmic part harboring a coiled region (amino acids 31-58) and a domain of unknown function, DUF2802, positioned in the C-terminal half of the protein (from amino acids 68-135) (Fig. 1D; Supplementary Fig. S1).

InterPro database analyzes found that FipA homologues (*i. e.* membrane proteins consisting of a single cytoplasmic DUF2802 repeat) are widespread among γ-proteobacteria. Exceptions

- 150 are the *Enterobacteriaceae*, which do not possess any copies of either *fipA* nor of *flhF* and *flhG*. Actually, FipA is only present in genomes that also encode FlhF and FlhG (**Fig. 1G**), which prompted the question of whether FipA is involved in regulating the flagellation pattern in concert with the FlhF-FlhG system. By including in our analysis the flagellation pattern reported in the literature, we found that the species that encode FipA are all polar flagellates,
- either monotrichous or lophotrichous (Fig. 1G). FipA homologues are absent from bacteria that use the FlhF-FlhG system to produce different flagellation patterns, like the peritrichous *Bacillus*, the amphitrichous ε-Proteobacteria or Spirochetes (Supplementary Table S2). In the α-Proteobacteria, where FlhF homologues are only present in a few species, FipA is absent as well (Fig. 1G).
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Based on these analyzes, we hypothesized that FipA represents a new family of FlhF interaction partners important for the γ-proteobacteria. To test this hypothesis, we decided to analyze FipA function in our monotrichously flagellated model organism *V. parahaemolyticus*, the distantly related and lophotrichously flagellated *Pseudomonas putida* and the monotrichous *Shewanella putrefaciens*. BACTH analysis showed that the FipA orthologue from both *P. putida* (*Pp*FipA, PP_4331) and *S. putrefaciens* (*Sp*FipA, SputCN32_2550) interact with FlhF of their respective species and that they self-interact (**Fig. 1F; Supplementary Fig. S2**) – a result similar to that of FipA from *V. parahaemolyticus* (*Vp*FipA). This supported our hypothesis and indicated that FipA has a general function as an FlhF interaction partner, thus constituting a new class of FlhF interaction partners.

FipA is required for proper swimming motility and flagellum formation

We continued our studies of FipA in the three organisms *V. parahaemolyticus*, *P. putida* and *S. putrefaciens* in parallel. To avoid interference with the secondary non-polar flagellar system
of *S. putrefaciens*, we used a strain that is unable to form the secondary flagella (Δ*flaAB*₂), to

which we will refer to as *S. putrefaciens* wild type in this study. For the swimming assays in soft agar, a strain of *V. parahaemolyticus* deleted in the lateral flagellin gene ($\Delta lafA$) was used as the background.

We first generated strains with individual deletions of the fipA and flhF genes. Strikingly,

- absence of FipA in V. parahaemolyticus completely abolished swimming motility in soft-agar medium, to the same degree as cells lacking FlhF (Fig. 2A). Furthermore, single-cell tracking of planktonic V. parahaemolyticus cells confirmed that cells lacking FipA were completely nonmotile and behaved identical to cells lacking FlhF (Fig. 2B). Importantly, ectopic expression of FipA in the ΔfipA strain restored the strain's swimming ability to wild-type levels (Fig. 2B),
- 185 further supporting that it is the deletion of *fipA* that results in the phenotype and not polar effects resulting from the *fipA* deletion. The C-terminal FipA-sfGFP fusion used throughout this paper also restored the phenotype (**Fig. 2B**), indicating that the fusion protein is fully functional.

A similar phenotype was observed for P. putida cells lacking FipA, however, the phenotype

- 190 was not as pronounced as that observed in *V. parahaemolyticus* (Fig. 2A, 2C). While cells lacking FlhF were severely affected in their ability to spread through soft agar (around 40% of wild-type spreading, Fig. 2C), *fipA* deletion strains of *P. putida* were still able to spread, with only a ~25% reduction.
- A different phenotype was observed for *S. putrefaciens* mutants. Firstly, cells lacking *flhF* were
 still able to spread in soft agar about half as far as wild-type cells. Secondly, the deletion of
 fipA only had a minor effect on spreading, as the mutants almost reached 90% of wild-type
 spreading (**Supplementary Fig. S3**). Altogether, these results indicate that FipA is necessary
 for normal flagella-mediated swimming motility in all three model species, albeit at a different
 extent reaching from almost complete loss of swimming in *V. parahaemolyticus* to only a small
 effect in *S. putrefaciens*.

FipA is required for proper assembly of the polar flagellum

The swimming phenotypes in the absence of FipA could result from defects in either flagellum assembly or in the flagellar motor performance. To differentiate between these possibilities,
we examined *V. parahaemolyticus* by transmission electron microscopy (TEM) (Fig. 2D, E) and *P. putida* and *S. putrefaciens* by fluorescent labeling of the flagellar filament(s) (Fig. 2F, G; Supplementary Fig. S3). Planktonic *V. parahaemolyticus* cells lacking either FlhF or FipA,

showed a complete absence of flagella on the bacterial surface, while a single polar flagellum was observed in ~50 % of wild-type cells (**Fig. 2D, E**). Flagellum quantification in *P. putida*

- 210 showed that the deletion of *Pp*FlhF drastically decreased flagellar formation, but in contrast to *V. parahaemolyticus*, did not abolish flagellation completely. Delocalization of flagella was also observed in ~5 % of *P. putida* Δ*flhF* cells (**Fig. 2F-G**). The deletion of *Pp*FipA had a more modest effect than in *V. parahaemolyticus*, reducing the number of flagellated cells to about half of that of the wild type (down to ~36% from ~77%, **Fig. 2G**). In *S. putrefaciens*, deletion of
- 215 *fipA* also results in a decrease in flagellation, albeit again at a lower extent ($\Delta fipA$, 42%; wild type 57%). Similarly, as previously reported (Rossmann et al, 2015), loss of *Sp*FlhF only reduces the proportion of flagellated cells (to 35%). However, in most of the flagellated $\Delta flhF$ mutants, the flagella were displaced away from the pole (28%), which was not observed in $\Delta fipA$ mutants.
- 220 Based on this set of experiments, we concluded that FipA and FlhF are essential for normal formation of polar flagella in all three bacterial model species, but their precise function might be different.

FipA and HubP ensure proper localization of FlhF to the cell pole

- 225 As our previous results showed an interaction between FipA and FlhF, we analyzed if the intracellular localization of FlhF was influenced by FipA. In this regard, we used functional translational fusions of FlhF to fluorescent proteins expressed from its native site on the chromosome in *V. parahaemolyticus*, *P. putida* and *S. putrefaciens* (Supplementary Figs. S4 and S5; Arroyo-Pérez and Ringgaard 2021; Rossmann et al. 2015).
- In *V. parahaemolyticus*, FlhF localized either diffusely in the cytoplasm or to the cell pole as previously reported (Arroyo-Pérez and Ringgaard 2021). However, a significant delocalization of FlhF from the cell pole occurred in the absence of FipA (Fig. 3A-C). Particularly, FlhF was diffusely localized in ~37% of cells or localized to the cell pole in a uni- and bi-polar manner in ~45% and ~19%, respectively, compared to wild-type cells (Fig. 3B). Absence of FipA significantly reduced localization of FlhF to the cell pole with a concomitant increase in diffusely localized FlhF (70%; Fig. 3A-C). Furthermore, the foci of FlhF at the cell pole in the absence of FipA were significantly dimmer compared to wild-type FlhF foci (Fig. 3D), indicating
 - that the amount of FlhF localized to the cell pole is lower in the absence of FipA. Importantly, even though FlhF was still able to localize to the cell pole in a certain number of cells lacking

240 FipA, no flagellum was formed in this background (**Fig. 2D-E**), thus indicating that FipA not only is required for proper polar recruitment of FlhF, but also for the stimulation of flagellum formation.

Given the role of HubP in cell pole organization and its reported interaction with FlhF (Yamaichi et al, 2012), we also analyzed the localization of FlhF in the absence of HubP. As for

- FipA mutants, recruitment of FlhF to the cell pole was reduced in the absence of HubP (Fig. 3A-C). Strikingly, in the double deletion strain Δ*fipA* Δ*hubP*, FlhF did not localize as foci at the cell pole at all but was instead localized diffusely in the cytoplasm in 100% of cells (Fig. 3A-C). Immunoblot analysis showed that the difference in localization of FlhF-sfGFP was not due to differences in expression levels (Supplementary Fig. S5A).
- 250 The requirement for FipA and FimV (the HubP ortholog in *P. putida*) for FlhF-mCherry localization in *P. putida* was even stronger, with ~95 and ~90% cells with diffuse FlhF-mCherry signal in their absence, respectively (**Fig. 3E, F**). FlhF was also completely delocalized in the double mutant $\Delta fipA \Delta fimV$. Unfortunately, the signal from the mCherry fusion was too weak to reliably measure the foci intensities in these strains, which may partly be due to instability
- 255 of the fusion.

Furthermore, analysis of FlhF-mVenus localization in *S. putrefaciens* confirmed that FlhF localization was not affected in a $\Delta hubP$ background (**Supplementary Fig. S6**; Rossmann et al, 2015). In a *S. putrefaciens* mutant lacking *fipA*, monopolar localization of FlhF-mVenus decreased to 63% of the cells (wild type 86%; **Supplementary Fig. S6**) and in a number of cells

- 260 (35% compared to 3% in the wild type), localization was lost completely and the signal became diffuse in the cytoplasm. Moreover, in a mutant lacking both *hubP* and *fipA*, dislocalization of FlhF-mVenus became more pronounced, as only weak polar localization occurred in 45% of the cells, while in 35% of the cells polar fluorescence was completely lost and FlhF-mVenus was distributed in the cytoplasm.
- 265 Altogether, these data show that both FipA and HubP work together to promote normal polar localization of FlhF.

Membrane anchoring of FipA is required for its function in regulating flagellum formation and proper FlhF localization

270 In order to identify the regions of FipA mediating its role in flagellum regulation, we next analyzed the N-terminal transmembrane domain for FipA membrane anchoring and its function in mediating proper flagellum production. When expressed in *E. coli*, a C-terminally GFP-tagged version of FipA (*Vp*FipA-sfGFP) showed a clear membrane localization, while a FipA variant deleted for the predicted transmembrane (TM) domain between residues 7-27

- 275 (FipA Δ TM), was diffusely localized in the cytoplasm (**Fig. 4A**). These observations show that FipA is indeed a membrane protein anchored by the predicted transmembrane domain. To then study the function of FipA membrane localization, the native *fipA* locus was replaced by a gene encoding a FipA variant lacking the N-terminal TM domain, and the resulting strain (*fipA* Δ TM) was analyzed for flagellum production. The *V. parahaemolyticus* strain carrying this
- mutation did not produce any flagella at all (Fig. 4B), as did the Δ*fipA* strain (Fig. 2D). Similarly, the *fipA* Δ*TM* mutation in *P. putida* had the same swimming defect as the deletion of *fipA* (Fig. 4C), indicating that the attachment to the membrane is essential for FipA function also in this species.

In *S. putrefaciens*, loss of the FipA TM domain had rather little effect on flagella-mediated spreading through soft agar (**Supplementary Figs. S3A**. However, this was expected as in *S. putrefaciens* the full deletion of FipA alone exerts only little effect on swimming and FlhF localization (**Supplementary Figs. S3A**, **B**).

A conserved cytoplasmic domain in FipA mediates interaction with FlhF

- 290 After validating an essential role for membrane anchoring of FipA, we analyzed in more detail the cytoplasmic DUF2802 domain. By aligning various FipA homologs, we identified a motif of conserved amino acids, and three of them (G110, E126 and L129 of *Vp*FipA) were 100% conserved among FipA homologues (**Fig. 5A**). Therefore, we chose these amino acids for mutagenesis.
- Alanine substitutions of residues G110 and L129 abolished the interaction between VpFipA and VpFlhF in BACTH analysis, while the E126A substitution did not affect the interaction (Fig. 5B). All these variants were, however, still able to self-interact with wild-type VpFipA (Fig. 5B). The corresponding substitutions in PpFipA (G104A and L123A) also abolished the interaction with PpFlhF; the mutation in the glycine had a less pronounced effect as in V.
- 300 parahaemolyticus (Fig. 5C). None of these mutations affected the self-interaction of FipA (Fig. 5C). In S. putrefaciens, the corresponding residue substitutions in SpFipA (G106A and L118A) also suppressed interaction with FlhF, but not with itself (Supplementary Fig. S7A). Altogether, these results suggest that conserved residues in the DUF2802 domain support the

interaction between FipA and FlhF, and that the self-interaction is mediated by a different region.

Interaction between FipA and FlhF is required for FipA function on regulating flagellum formation and FlhF localization

We proceeded to evaluate the role of these residues in vivo. After introducing the mutations

- 310 in the *fipA* gene in its native loci, the effects on motility were almost indistinguishable from a Δ *fipA* mutation. In *V. parahaemolyticus,* this resulted in completely non-motile cells in planktonic cultures (**Fig. 5D**). In *P. putida,* the effect was again less pronounced, but there was a reduction in the swimming halo of the same magnitude as in the Δ *fipA* background. Only the G104A mutation had a less pronounced phenotype, which was still significantly different from
- the wild-type (**Fig. 5E**).

The localization of FlhF was also affected by the point mutations in *fipA*. Cells of *V*. *parahaemolyticus* expressing *Vp*FlhF-sfGFP natively had reduced polar localization when *Vp*FipA was substituted with either G110A or L129A (**Fig. 5F**), with almost 50% of cells presenting only diffuse *Vp*FlhF-sfGFP signal. The effect was more pronounced in *P. putida*,

- 320 with almost 90% percent of cells presenting diffuse PpFlhF-mCherry signal in the presence of PpFipA G104A or L123A, very close to the ~95% of cells in the $\Delta fipA$ background (**Fig. 5G**). The general effect of the point mutations in *S. putrefaciens* was also similar to the deletion of *fipA*, where it has a weaker effect than in the other two organisms (**Supplementary Fig. S7**). Thus, it seems that the DUF2802 domain of FipA is responsible for the effect of FipA on motility, and
- 325 that its effects occur primarily through the interaction with FlhF. Furthermore, the interruption of FipA-FlhF impedes the recruitment of FlhF to the cell pole.

Cell cycle-dependent polar localization of FipA

Given FipA's function on regulating correct recruitment of FlhF to the cell pole, we next analyzed the intracellular localization of FipA in our three model organisms, *V. parahaemolyticus*, *P. putida* and *S. putrefaciens*. To this end, a hybrid gene bearing a fusion of *fipA* to *sfgfp* was used to replace native *fipA* in all three species, resulting in stable and functional production of FipA C-terminally tagged with sfGFP (**Supplementary Fig. S8**). In wildtype strains of all three organisms, FipA formed distinct foci at the cell pole, either uni- or bi-

polarly (Figs. 6A, B and Supplementary Fig. S9). No delocalized foci were ever observed. A

remarkable difference between *P. putida* and *V. parahaemolyticus* is that in *P. putida*, FipA occurred as foci in virtually all cells (Fig. 6B), whereas in *V. parahaemolyticus*, it remained diffuse in half of the population (Fig. 6A). The ratio of bi-polar to uni-polar foci was also greater in *P. putida* (~1:2) than in *V. parahaemolyticus* (~1:5). This behavior was explored further by
following the proteins through the cell cycle in the two organisms. In *V. parahaemolyticus*, FipA foci disappear frequently (Fig. 6C; 14'). A new focus appears at the opposite pole, sometimes before cell division (Fig. 6C; 56'), sometimes after (Fig. 6C; 42'). On rare cases, the first focus persists until after the second focus appears, resulting in bipolar foci (Fig. 6C; 63'). On the other hand, in *P. putida*, the FipA foci were more stable, and foci at both poles often

occurred right after cell division (**Fig. 6D; 50'**), explaining the greater bi-polar to uni-polar ratio of FipA foci in this organism. These results are consistent with a protein that is recruited to the cell pole right before the start of the flagellum assembly after cell division, albeit the timing may vary between different species.

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Polar localization of FipA depends on the FipA-FlhF interaction and on HubP

Finally, we determined whether FlhF plays a role in FipA localization. To this end, FlhF was deleted in the strains expressing FipA-sfGFP from its native promoter. Almost no FipA foci were detected in this background for both *V. parahaemolyticus* (Fig. 7A-C) and *P. putida* (Fig.

7D-F), even though the protein levels of FipA were comparable to that in the wild type (Supplementary Fig. S8).

Furthermore, the FipA variants that do not interact with FlhF (**Fig. 5B, C**), were also labeled in the native *fipA* locus with sfGFP. Both *Vp*FipA G110A and *Vp*FipA L129A were incapable of forming polar foci (**Fig. 7B, C**). The corresponding variant in *P. putida*, *Pp*FipA L123A was also

360 mostly distributed in the cytoplasm, whereas the variant *Pp*FipA G104A did exhibit some polar localization, although reduced (Fig. 7E, F). This is the variant that did show some interaction with *Pp*FlhF (Fig. 5C, G).

Altogether, these results suggest that, in *V. parahaemolyticus* and *P. putida*, a direct interaction with FlhF, mediated by the residues in the DUF2802, is responsible for recruiting

365 FipA to the pole. Surprisingly, in *S. putrefaciens* mutants deleted in *flhF*, FipA displayed only a slight difference in polar localization (Supplementary Fig. S9). However, a *Sp*FipA G106A reduced the protein's localization (to ~50% cells with foci). It is likely that in *S. putrefaciens*,

HubP is (co-)mediating the localization of FipA, as in the absence of HubP, FipA-sfGFP at the cell pole drastically decreases (**Supplementary Fig. S9**).

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Discussion

In bacteria, numerous processes are localized to specific cellular compartments. This is particularly evident in polarly flagellated bacteria, where the intricate flagellar multicomplex needs to be synthesized in a spatiotemporally regulated fashion. Flagellar synthesis is initiated

- 375 by the assembly of the first flagellar building blocks within the cytoplasmic membrane, which in polar flagellates is targeted to the designated cell pole. In a wide range of bacterial species, the SRP-type GTPase FlhF and its antagonistic counterpart, the MinD-like ATPase FlhG, regulate flagellar positioning and number (Kazmierczak and Hendrixson 2013, Schuhmacher, Thormann, and Bange 2015). In particular, FlhF has been shown to function as a positive
- 380 regulator and a major localization factor for the initial flagellar building blocks of polar flagella. Upon production, GTP-bound dimeric FlhF localizes to the cell pole, but the mechanism by which the protein assumes its designated position at the cell pole, remained elusive. One candidate protein, the polar landmark HubP (FimV in *Pseudomonas* sp.) had been identified earlier and was shown to interact with FlhF in *V. parahaemolyticus* (Yamaichi et al, 2012).
- 385 Accordingly, polar localization of FlhF is decreased in mutants lacking *hubP* (this study; Rossmann, Brenzinger et al, 2015), however, in a number of cells FlhF localized normally and the cells showed normal flagellation. This was indicative that HubP does play a role in functionally recruiting FlhF, but that another factor was still missing. In this study we identified the protein FipA as a second polarity factor for FlhF.

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FipA consists of an N-terminal transmembrane domain and a cytoplasmic region with a conserved domain of unknown function (DUF2802). The corresponding gene is located immediately downstream of the motility operon that includes *flhF* and *flhG*. Notably, FipA is highly conserved among bacteria that use the FlhF/FlhG system to position the flagella, strongly suggesting that FipA has evolved together with FlhF and FlhG to regulate the flagellation pattern. In this study, we therefore investigated three distantly related *γ*-proteobacteria including one lophotrichous species, *V. parahaemolyticus*, *P. putida* and *S. putrefaciens*, in more detail. We found that FipA is conserved as a regulatory factor of FlhF and that its functions rely on similar features in all three species.

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FipA is essential for normal flagellum synthesis, however, the severity of a *fipA* deletion differed among the three species studied. In *V. parahaemolyticus*, a *fipA* deletion phenocopies the loss of *flhF*, and cells no longer synthesize flagella. Notably, in the absence of FipA in *V. parahaemolyticus*, FlhF is still localized at the pole in about a third of the cells. Despite this,

- these cells were still unable to form flagella, strongly suggesting that occurrence of FlhF at the pole alone is not sufficient to trigger flagellum synthesis in the absence of FipA.
 This strict requirement for FlhF has been reported in the closely related *V. alginolyticus* (Kusumoto et al, 2009). On the other hand, in *V. cholerae*, the requirement for FlhF on flagellation is less strict (Green et al, 2009), and in *P. putida* and *S. putrefaciens* even less so:
- 410 mutants deleted in *flhF* or *fipA* of both species can generally still produce flagella and swim (this work; Rossmann et al, 2015). In *P. putida*, a Δ *fipA* mutant has a phenotype reminiscent to a Δ *flhF* mutant as both drastically decrease the number of flagella, which may also be delocalized in *flhF* mutants (Pandza et al, 2003). In *S. putrefaciens*, deletion *fipA* decreases the number of flagella to a similar extent as a *flhF* mutation, however, while in the latter case the
- 415 flagella are frequently delocalized, the flagella remain at a polar postion, when *fipA* is missing. Taken together, these results strongly suggest that FipA does not act as a general polarity factor for the full flagellar machinery, but rather it stimulates the activity of FlhF to initiate polar flagellation.

420 FipA and HubP affect polarity of FlhF through different mechanisms

In all three species, FipA directly interacts with FlhF as demonstrated by reciprocal co-immuno precipitations and by bacterial two-hybrid assays. These experiments suggest that FipA is able to dimerize (or oligomerize) and that the FlhF-FipA interaction is dependent on conserved residues in the DUF2802 domain. Furthermore, these residues are not required for the ability

425 of FipA to interact with itself, suggesting that the two processes occur independently and at different protein interfaces. Microscopic and physiological assays showed that direct interaction between FlhF and FipA is required for FlhF targeting and function. In addition, a FipA mutant lacking its N-terminal transmembrane domain was non-functional, indicating that interaction of FlhF and FipA has to take place at the membrane.

In the absence of FipA, polar localization of FlhF was significantly decreased in all three species and almost completely diminished when *hubP* was deleted together with *fipA*. Removing HubP/FimV alone had a similar effect as removing FipA on the localization of FlhF, although to a lesser extent, as more cells still had polar FlhF in the $\Delta hubP/fimV$ mutant than in the $\Delta fipA$

- 435 mutant. Furthermore, the unipolar to bipolar ratio of FlhF foci increased in the $\Delta hubP$ mutant compared to the wild-type or the $\Delta fipA$ background. While the deletion of *fipA* increases the number of cells with diffuse localization, it does not affect the time frame in the cell cycle at which FlhF shifts from unipolar to bipolar in *V. parahaemolyticus*. In contrast, deleting *hubP* delays the time frame in which FlhF assumes a bipolar position. This indicates that HubP and
- 440 FipA represent two different pathways that stimulate polar localization of FlhF, and that both are required to bring sufficient (active) FlhF molecules to trigger MS-ring formation and start flagellum assembly.

Localization of FipA

- A notable feature of FipA is its dynamic localization pattern within the cell over the cell cycle in all three species studied, the nature of which is unclear so far. Foci of FipA may appear mono- or bipolarly and frequently, but not necessarily, disappear once the flagellar apparatus, or the flagellar bundle in the case of *P. putida*, is established. It remains to be shown whether the FipA protein is actively moving from one pole to another or if newly formed FipA is
- 450 recruited to the opposite pole while being degraded at the old position. Notably, in *V. parahaemolyticus* and *P. putida* FipA does not localize polarly in the absence of interaction with FlhF, suggesting that both proteins are recruited as a complex.
 In contrast, in *S. putrefaciens,* FipA still localizes to the cell pole also in the absence of FlhF as

long as the polar landmark HubP is present. This is indicating a more important role for HubP

- 455 in this species. However, also in *S. putrefaciens* active FlhF localizes to the cell pole and flagella are formed in the absence of HubP. The nature of the polar marker that recruits or guides FipA or a FipA/FlhF to the designated pole is still elusive. The discovery of FipA provides a new point where spatiotemporal organization is coordinated. However, it remains to be explored if the complex is anchored through yet another protein, such as the TonB_m-PocA-PocB complex in
- 460 *P. putida* (Ainsaar et al, 2019) or through an intrinsic feature of the cell envelope or cytoplasm during cell division.

How does FipA upgrade the current model of polar flagellar synthesis?

- Based on the findings in different species, our current model of FlhF/FlhG-mediated polar flagellar synthesis predicts, that upon production, GTP-bound dimeric FlhF is localizing or being recruited to the cell pole to where it recruits the first flagellar components to initiate the assembly. The monomeric form of the ATPase FlhG, the FlhF antagonist, is binding to the flagellar C-ring building block FliM and is transferred to the nascent flagellar structure, where it is released and dimerizes upon ATP binding (Schuhmacher et al, 2015; Blagotinsek et al,
- 470 2020). The ATP-bound FlhG dimer can associate with the membrane and interact with the GTP-bound FlhF dimer, thereby stimulating its GTPase activity. This leads to monomerization of FlhF and loss of polar localization (reviewed in Kojima, Terashima, and Homma 2020; Schuhmacher et al., 2015). Dimeric FlhG does also interact with the master regulator of flagella synthesis, FlrA (or FleQ in *Pseudomonas*) and prevents the synthesis of further early flagellar
- 475 building blocks (Dasgupta & Ramphal, 2001; Blagotinsek et al, 2020; Chanchal et al, 2021). By this, FlhG links flagella synthesis with transcription regulation and effectively restricts to number of polar flagella that are formed. Based on this model, FipA may recruit FlhF to the membrane and stimulate or stabilize GTP binding and dimerization of FlhF to promote polar localization and initiation of flagella synthesis. Thus, the role of FipA is to shift the equilibrium
- 480 to the active state of FlhF in order to start assembly of the MS ring. Accordingly, current studies address the structural basis of the FipA-FlhF interaction, the potential effect of FipA on GTP-binding and dimerization of FlhF, and the role of the polar marker HubP in this process. In addition, *fipA* expression is independent of the rest of the flagellar genes in other species of *Vibrio* (Moisi et al. 2009; Petersen et al. 2021) and *S. putrefaciens* (Schwan et al, 2022).
- 485 Thus, *fipA* transcription could provide another regulatory point to lead to the synthesis of a flagellum. Thus, the discovery of FipA opens several novel open questions in the field of flagellum regulation.

490 Materials and methods

Growth conditions and media

In all experiments *V. parahaemolyticus*, and *E. coli* were grown in LB medium or on LB agar plates at 37°C containing antibiotics in the following concentrations: 50 μ g/mL kanamycin, 100 μ g/mL ampicillin and 20 μ g/mL chloramphenicol for *E. coli* and 5 μ g/mL for *V*.

495 parahaemolyticus. S. putrefaciens CN-32 and P. putida KT 2440 were grown in LB medium or LB agar plates at 30 °C. When required, the media were supplemented with 50 µg/mL kanamycin, 300 µM 2,6-diaminopimelic acid and/or 10 % (w/v) sucrose.

Strains and plasmids

- 500 The strains and plasmids used in this study are listed in Tables S3 and S4, respectively. Primers used are listed in Table S5. *E. coli* strain SM10λ*pir* was used to transfer DNA into *V. parahaemolyticus* by conjugation (Miller & Mekalanos, 1988). For DNA transfer into *S. putrefaciens* and *P. putida*, *E. coli* WM3064 was used. *E. coli* strains DH5αλ*pir* and SM10λ*pir* were used for cloning. Construction of *V. parahaemolyticus* deletion mutants was performed
- 505 with standard allele exchange techniques using derivatives of plasmid pDM4 (Donnenberg & Kaper 1991). Chromosomal deletions and integrations in *S. putrefaciens* and *P. putida* were carried out by sequential crossover as previously described (Rossman et al. 2015) using derivatives of plasmid pNPTS138-R6K (Lassak et al. 2010).

510 **Construction of plasmids**

Plasmid pSW022: The regions flanking *vp2224* (*fipA*) were cloned with primers VP2224-del-a/ b and VP2224-del-c/ d, using *V. parahaemolyticus* RIMD 2210633 chromosomal DNA as template. The resulting products were fused in a third PCR using primers VP2224-del-a/ VP2224-del-d. The end product was digested with Xbal and ligated in the equivalent site of

- 515 vector pDM4, resulting in plasmid pSW022. The mutation in *V. parahaemolyticus* was confirmed with a PCR using primers VP2224-del-a/VP2224-check. *Plasmid pPM123:* The regions flanking aa 7-27 of *vp2224* (*fipA*) were cloned with primers VP2224-del-a/ del AA7-27 vp2224-b and del AA7-27 vp2224-c/ VP2224-del-d, using *V. parahaemolyticus* RIMD 2210633 chromosomal DNA as template. The resulting products were fused in a third PCR using primers
- 520 VP2224-del-a/ VP2224-del-d. The end product was digested with XbaI and ligated in the equivalent site of vector pDM4, resulting in plasmid pPM123. The mutation in *V. parahaemolyticus* was confirmed with a PCR using primers VP2224-del-a/VP2224-check. *Plasmid pPM178:* The gene *vp2224* (*fipA*) was amplified from *V. parahaemolyticus* RIMD 2210633 chromosomal DNA with primers C-term sfGFP-vp2224-a/-b; the downstream region
- 525 with primers C-term sfGFP-vp2224-e/f; the gene encoding sfGFP with C-term sfGFP-vp2224c/d from plasmid pJH036. The three products were fused together in another PCR using

primers C-term sfGFP-vp2224-a/f. The obtained product, encoding FipA fused in frame to sfGFP via a 5-residue linker, was digested with SpeI and SphI and ligated in vector pDM4 digested with XbaI and SphI, resulting in plasmid pPM178. The mutation in *V*.

- 530 *parahaemolyticus* was confirmed with a PCR using primers C-term sfGFP-vp2224-f/VP2224check. *Plasmid pPM179 & pPM180:* The region upstream of *vp2224 (fipA)* were amplified as for pSW022. The downstream region was amplified with downstream vp2224-cw/VP2224-deld from *V. parahaemolyticus* RIMD 2210633. The gene *vp2224* itself was amplified with primers vp2224 cw restore deletion/vp2224 ccw restore deletion, from plasmids pEP005 & pEP006,
- 535 carrying the mutations G110A and L129A. The products were fused in a third PCR using primers VP2224-del-a/ VP2224-del-d. The end product was digested with Xbal and ligated in the equivalent site of vector pDM4, resulting in plasmids pPM179 & pPM180. The the reinsertion in *V. parahaemolyticus* SW01 was confirmed with a PCR using primers VP2224-dela/VP2224-check. *Plasmid pPM187 & pPM191:* The insertion of sfGFP at the C-terminus of FipA
- 540 was cloned with the same strategy for pPM178, but using pPM179 & pPM180 as templates for the *vp2224* sequence, resulting in plasmids pPM191 & pPM187, respectively. *Plasmid pPM146:* The gene *vp2224* (*fipA*) was amplified from *V. parahaemolyticus* RIMD 2210633 chromosomal DNA with primers vp2224-cw-pBAD/vp2224-ccw-pBAD. The product was digested with enzymes XbaI and SphI, and inserted in the corresponding site in plasmid
- 545 pBAD33. *Plasmid pPM159:* The gene *vp2224* (*fipA*) was amplified from *V. parahaemolyticus* RIMD 2210633 chromosomal DNA with primers C-term sfGFP-vp2224-a/-b, and the gene encoding sfGFP with C-term sfGFP-vp2224-c/sfGFP-1-ccw from plasmid pJH036. The resulting products were fused in another reaction with primers C-term sfGFP-vp2224-a/sfGFP-1-ccw, and this final product was digested with XbaI and cloned in pBAD33. *Plasmid pPM194:* Same
- 550 as pPM159, but vp2224 ΔTM was amplified from pPM123 using primers vp2224 AA1-6/28end w/o Stop/sfGFP-1-ccw. Plasmids pSW74 & pSW119: The region of the gene vp2224 (fipA) encoding the cytoplasmic part (residues 28-163) was amplified with primers tr2224 put18C cw/ pUT18C/pKT25-vp2224-ccw from V. parahaemolyticus chromosomal DNA. The product was digested with KpnI and XbaI and ligated in the corresponding site in pKT25, to generate
- 555 pSW74, and in pUT18C, to generate pSW119. *Plasmid pPM118 & pPM119:* The cytoplasmic part of the gene *vp2224* (*fipA*) was amplified with primers pUT18/pKNT25-tr-vp2224-cw & pUT18/pKNT25-vp2224ccw from *V. parahaemolyticus* chromosomal DNA. The product was digested with KpnI and XbaI and ligated in the corresponding site in pUT18, to generate

pPM118, and in pKNT25, to generate pPM119. Plasmid pPM124 & pPM128: The gene vp2234

- 560 (*flhF*) was amplified with primers pUT18/pKNT25- vp2234-cw & pUT18/pKNT25-vp2234 -ccw from *V. parahaemolyticus* chromosomal DNA. The product was digested with KpnI and XbaI and ligated in the corresponding site in pUT18, to generate pPM124, and in pKNT25, to generate pPM128. *Plasmid pPM132 & pPM136:* The gene *vp2234* (*flhF*) was amplified with primers pUT18C/pKT25- vp2234-cw & pUT18C/pKT25-vp2234 -ccw from *V. parahaemolyticus*
- 565 chromosomal DNA. The product was digested with KpnI and XbaI and ligated in the corresponding site in pUT18C, to generate pPM132, and in pKT25, to generate pPM136. *Plasmids pPM160, pPM161 & pPM162:* Site directed mutagenesis was performed on plasmid pSW74 by the QuickChange method (Zheng, Baumann, and Reymond 2004), using primers vp2224-Gly-110Ala-cw/ccw, vp2224-Glu 126Ala-cw/ccw or vp2224-Leu 129Ala-cw/ccw. After
- 570 digesting the template with DpnI, the products were transformed into *E. coli* and the mutations were confirmed by sequencing. *Plasmid pPM106:* The gene vp2224 (fipA) was amplified with primers pUT18C/pKT25-vp2224-cw & pUT18C/pKT25-vp2224ccw from V. parahaemolyticus chromosomal DNA. The product was digested with KpnI and XbaI and ligated in the corresponding site in pKTop, to generate pPM106. *Plasmid pPM109 & pPM112:*
- 575 The phoA-lacZα fragment of pKTop was amplified with primers vp2224 C-term PhoA-LacZ cw & end -LacZ w/o STOP ccw. The full length vp2224(fipA) gene was amplified from plasmid pPM146 using primers LacZ to vp2224 w/o ATG & end vp2224 ccw. The vp2224 ΔTM allele was amplified from plasmid pPM123 using primers vp2224 AA1-6/28-end w/o Stop & end vp2224 ccw. The PCR products were fused in a second PCR using primers vp2224 C-term PhoA-
- 580 LacZ cw & end vp2224 ccw. The fusion product was digested with Xbal and HindIII and cloned in the corresponding site of plasmid pBAD33. *Plasmids for genetic manipulation of S. putrefaciens and P. putida:* The desired DNA fragments

were generated by PCR using appropriate primer pairs (see Supplementary Table S5) that in addition create overhangs suitable for subsequent Gibson assembly into EcoV-digested vector

585 pNTPS138-R6K (Gibson et al., 2009; Lassak et al, 2010). If necessary, primer overhangs were also used to generate base substitutions in the gene fragment to be cloned. *Plasmids for Bacterial Two Hybrid (BACTH) analysis* were similarly generated by amplification of the desired DNA-fragments, which were then introduced into the suitable vectors by Gibson assembly.

590 Soft-agar swimming assays

V. parahaemolyticus soft-agar swimming assays were essentially performed as described in (Ringgaard, et al, 2013) with the following modifications. Late exponential cultures of the required strains were used to prick LB plates with 0.3% agar, and incubated for 30 hours at 30° C. For *S. putrefaciens* and *P. putida* soft-agar swimming assays, 2 µl of an exponentially growing culture of the appropriate strains were spotted onto 0.25 % LB agar plates and

incubated at 30 °C (*S. putrefaciens*) or room temperature (*P. putida*) for about 18 hours. Strains to be directly compared were always spotted onto the same plate.

Bacterial-two-hybrid experiments, BACTH

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- BTH101 cells were made competent with calcium chloride. 15 μL aliquots were spotted in a 96 well plate, to which 2.5 μL of the corresponding pUT18(C) and pK(N)T25 derivative plasmids were added. After 30 minutes on ice, a heat shock at 42° C was applied for 30 s. The transformed cells were allowed to recover for 1 hour, after which the were grown in selective LB broth for another 3 hours. The resulting cultures were spotted on plates containing kanamycin, ampicillin, IPTG (0.25 mM) and X-Gal (80 μg/mL). The plates were photographed
- 605 kanamycin, ampicillin, IPTG (0.25 mM) and X-Gal (80 μg/mL). The plates were photographed after 48-72 hours at 30° C.

Video tracking of swimming cells

- Video tracking of swimming cells was performed essentially as described previously (Ringgaard, et al, 2013). Swimming cells were recorded using the streaming acquisition function in the Metamorph software and the swimming paths of individual cells were tracked using the MTrackJ plug-in for ImageJ. The swimming speed, displacement, and number of reversals of individual cells were then measured and the average plotted with error bars indicating the standard deviation. Video tracking was performed using a Zeiss Axio Imager M1
- 615 fluorescence microscope. Images were collected with a Cascade:1K CCD camera (Photometrics), using a Zeiss α Plan-Fluar 40x/1.45 Oil phase contrast objective.

Transmission-electron-microscopy (TEM) analysis

Cell cultures grown to an OD600 = 0.5–0.6 were spotted on a plasma-discharged carbon-coated copper
 grid (Plano, Cat#S162-3) and rinsed with 0.002% uranyl acetate. Afterwards they were rinsed with
 water and blotted dry with Whatman filter paper. TEM images were obtained with a JEOL JEM-1400
 Plus 120 KV transmission electron microscope at 80 kV.

Flagellum labeling

To fluorescently label flagellar filaments, maleimide-ligates dyes (Alexa Fluor 488 C5 maleimide fluorescent dye; Thermo Fisher Scientific) were coupled to surface-exposed cysteine residues, which were introduced into the flagellins (FlaA and FlaB) of the polar flagellar system in *S. putrefaciens* and *P. putida* as described (Kühn et al. 2017, Hintsche et al. 2017). Images were recorded as described in the microscopy section.

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Fluorescence microscopy

Florescence microscopy on *V. parahaemolyticus* was carried out essentially as previously described (Muraleedharan et al, 2018), using a Nikon eclipse Ti inverted Andor spinning-disc confocal microscope equipped with a 100x lens, an Andor Zyla sCMOS cooled camera, and an

- Andor FRAPPA system. Fluorescence microscopy on *S. putrefaciens* and *P. putida* was carried out as previously described (Kühn et al. 2017) using a custom microscope set-up (Visitron Systems, Puchheim, Germany) based on a Leica DMI 6000 B inverse microscope (Leica) equipped with a pco.edge sCMOS camera (PCO), a SPECTRA light engine (lumencor), and an HCPL APO 63×/1.4–0.6 objective (Leica) using a custom filter set (T495lpxr, ET525/50m;
- 640 Chroma Technology) using the VisiView software (Visitron Systems, Puchheim, Germany). Microscopy images were analyzed using ImageJ imaging software (http://rsbweb.nih.gov/ij) and Metamorph Offline (version 7.7.5.0, Molecular Devices). Demographs were generated as described by Cameron et al, 2014, and modified in Heering & Ringgaard 2016 and Heering, Alvarado and Ringgaard, 2017.

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Mapping interaction partners using co-immunoaffinity purification and mass spectrometry (co-IP-MS)

For sample preparation of the IP-MS experiments we were using a modified version of the protocol presented by Turriziani et al, 2014. Cells were centrifuged and cell pellets were

- washed with cold PBS. Cell pellets were resuspended in lysis buffer (50 mM HEPES (pH 7.5), 150 mM NaCl, 0.5 % NP50, 5 mM EDTA, compete mini protease inhibitors (Complete Mini (Roche)). Cell lysis was performed by repetitive sonication. After removing cell debris by centrifugation 10ul GFP-trap Sepharose (Chromotek) slurry was added to the lysate and incubation was carried out for 1.5 hours on a rotating shaker at 4 °C. Then the beads were
- pelleted, the supernatant removed and the beads washed 4x with 100 mM NH₄CO₃ to remove

remaining protease inhibitors and detergents. 200 μ l elution buffer (1 M urea, 100 mM NH_4CO_3 , 1 µg Trypsin (Promega)) was added to the beads and incubated at 1000 rpm on a thermomixer at 27°C for 45min. Beads were centrifuged and supernatant was collected. In order to increase peptide recovery 2 washing steps with 100µl elution buffer 2 (1 M urea, 100 mM NH₄CO₃, 5 mM Tris(2-caboxyethyl)phosphine)) was performed and the individual bead supernatants were collected into one tube. Tryptic digestion was carried out overnight at 30 °C. After digest alkylation was performed with 10 mM iodoacetamide at 25 °C (in the dark). Then the samples were acidified (1 % trifluoroacetic acid (TFA)) and C18 Microspin column (Harvard Apparatus) was carried out according to the manufacturer's instruction. The samples

were dried and recovered in 0.1 % TFA and applied to liquid chromatography-mass 665 spectrometry (LC-MS) analysis.

LC-MS analysis of digested lysates was performed on a Thermo QExactive Plus mass spectrometer (Thermo Scientific), which was connected to an electrospray ionsource (Thermo Scientific). Peptide separation was carried out using an Ultimate 3000 RSLCnano with Proflow

- upgrade (Thermo Scientific) equipped with a RP-HPLC column (75 μm x 42 cm) packed in-670 house with C18 resin (2.4 µm; Dr. Maisch) on an in-house designed column heater. The following separating gradient was used: 98 % solvent A (0.15 % formic acid) and 2 % solvent B (99.85 % acetonitrile, 0.15 % formic acid) to 35 % solvent B over 90 at a flow rate of 300 nl/min. The data acquisition mode was set to obtain one high resolution MS scan at a resolution of
- 675 70,000 full width at half maximum (at m/z 200) followed by MS/MS scans of the 10 most intense ions. To increase the efficiency of MS/MS attempts, the charged state screening modus was enabled to exclude unassigned and singly charged ions. The dynamic exclusion duration was set to 30 sec. The ion accumulation time was set to 50 ms (MS) and 50 ms at 17,500 resolution (MS/MS). The automatic gain control (AGC) was set to 3x10⁶ for MS survey

680 scan and $1x10^5$ for MS/MS scans.

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MS analyzed with MaxQuant 1.6.3.4) raw data was then (Version (https://www.nature.com/articles/nbt.1511) using a V.parahaemolyticus RIMD 2210633 uniprot database (www.uniprot.org). MaxQuant was executed in standard settings with activated "match between runs" option. The search criteria were set as follows: full tryptic specificity was required (cleavage after lysine or arginine residues); two missed cleavages were allowed; carbamidomethylation (C) was set as fixed modification; oxidation (M) and deamidation (N,Q) as variable modification. For further data analysis the MaxQuant LFQ

values were loaded into Perseus (<u>https://www.nature.com/articles/nmeth.3901</u>) and a Student's T-test was performed on LFQ values with false discovery rate 0.01 and S0: 0.1 as significance cut-off.

The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE (PubMed ID: 34723319) partner repository with the dataset identifier PXD045379 (<u>http://www.ebi.ac.uk/pride</u>).

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Membrane topology mapping of FipA

We experimentally determined the membrane orientation of FipA in the membrane using the dual pho-lac reporter system (Karimova et al, 2009), which consists of a translational fusion of the *E. coli* alkaline phosphatase fragment PhoA22-472 and the α -peptide of *E. coli* β -

galactosidase, LacZ4-60. A periplasmic localization of the reporter leads to high alkaline phosphatase activity and low β-galactosidase activity, whereas a cytosolic location of the reporter results in high β-galactosidase activity and low alkaline phosphatase activity. Pho-Lac-FipA and FipA-Pho-Lac fusion proteins were ectopically expressed in *E. coli* DH5α grown on a dual-indicator LB medium containing a blue indicator for phosphatase activity (X-Phos) and red indicator for β-galactosidase activity (Red-Gal) (see Supplementary Figure S1).

Bioinformatic analysis

Homologues of FipA were searched using BLAST against the KEGG database, with the sequence of the *V. parahaemolyticus* protein. All homologues containing a DUF2802 were

710 included. The search was later expanded to species known to encode a FlhF homologue (defined as the highest scoring result from a BLAST search using FlhF of *V. parahaemolyticus* as a query, that was also encoded upstream of an FlhG homologue). The flagellation phenotype was later corroborated in the description registered at the List of Prokaryotic names with Standing in Nomenclature (Parte et al. 2020).

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725 Conflict of interests

The authors declare no conflict of interests.

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Figure 1. FipA constitutes a new family of FlhF interaction partners. Volcanoplots representing Log-ratios versus significance values of proteins enriched in **(A)** FlhF-sfGFP or **(B)**

FipA-sfGFP purifications using shotgun proteomics and liquid chromatography-mass

- 810 spectrometry; sfGFP was used as control. The full list of pulled-down proteins can be found in the Supplementary Tables S1 and S2. C) Organization of the flagellar/chemotaxis gene region encoding FlhF and FipA in *V. parahaemolyticus*. D) Domain organization of FipA (TM, transmembrane region; DUF, domain of unknown function). (E, F) Bacterial two-hybrid confirming the interaction between FipA and FlhF from *V. parahaemolyticus* (E) and *P. putida*
- 815 (F). The indicated proteins (FipA, FlhF) were fused N- or C-terminally to the T18- or T25fragment of the Bordetella pertussis adenylate cyclase. In vivo interaction of the fusion proteins in Escherichia coli is indicated by blue color. The corresponding assay of S. putrefaciens is displayed in Supplementary Figure S2. G) Dendrogram of γ-proteobacteria, indicating the presence of FlhF or FipA homologues and the corresponding flagellation
- pattern. An extended version and sources are available in Supplementary Table S2.



Figure 2. FipA is required for correct flagellum formation. (A, C) Representative soft-agar swimming assay of *V. parahaemolyticus* (A) or *P. putida* (C) strains (left panels) and the corresponding quantification (right panels). For the latter, the halo diameter measurements were normalized to the halo of the wild type on each plate. Data presented are the from six (A) or three (C) independent replicates, asterisks represent a p-value <0.05 (according to ANOVA + Tukey tests). (B) Single-cell tracking of *V. parahaemolyticus*. Shown are representative swimming trajectories and quantification of swimming speed, total displacement and reversal rate. N indicates number of cells tracked among 3 biological replicates (ANOVA + Tukey test). (D) Representative electron micrographs of the indicated *V. parahaemolyticus* strains stained with uranyl acetate. (E) Quantification of flagellation pattern

in the populations of the indicated *V. parahaemolyticus* strains. (F) Flagellum stain of indicated *P. putida* strains with Alexa Fluor 488-C5-maleimide and (G) quantification of the
corresponding flagellation in the population. N indicates the number of cells counted among
3 biological replicates. For *S. putrefaciens*, see Supplementary Figure S3.



Figure 3. Localization of FlhF depends on FipA and HubP. (A) Representative micrographs of
the indicated strains of *V. parahaemoloyticus* expressing FlhF-sfGFP from its native promoter.
Upper panel shows the DIC image, the lower panel the corresponding fluorescence image.
Fluorescent foci are highlighted by white arrows. Scale bar = 2 μm. (B) Demographs displaying
FlhF-sfGFP fluorescence intensity along the cell length within the experiments shown in (A).
(C) Quantification of localization patterns and (D) foci fluorescence intensity of the
fluorescence microscopy experiment presented in (A). The data was combined from the given number (N) of cells combined from three biological replicates. (E) Representative micrographs of the indicated *P. putida* strains expressing FlhF-sfGFP from its native promoter. The upper panels show the DIC and the lower panels the corresponding fluorescence images. Fluorescent foci are marked by small white arrows. The scale bar equals 5 μm. The low intensity of the foci did not allow a quantitative analysis of foci intensities or the generation of demographs. (F)

Quantification of FlhF-sfGFP localization patterns in the corresponding strains of *P. putida* from the experiments shown in **(E)**. Corresponding data on the localization of FlhF in *S. putrefaciens* is displayed in **Supplementary Figure S6**.



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Figure 4. Activity of FipA depends on FlhF and on its transmembrane domain. (A) Micrographs of *E. coli* cells expressing FipA-sfGFP from *V. parahaemolyticus*, and a truncated version lacking the transmembrane domain (ΔTM). The left panels display the DIC and the right panels the corresponding fluorescence images. The scale bar equals 5 µm. (B) Electron micrographs of *V. parahaemolyticus* wild-type and mutanzs cells lacking the transmembrane domain of *fipA*, respectively. The corresponding quantification of the flagellation pattern is shown to the left of the micrographs. Note that the data for the wild-type cells is the same as in Fig. 2D, E. (C) Spreading behavior of the indicated *P. putida* strains (left) with the corresponding quantification (right). Loss of the FipA TM region phenocopies a complete *fipA* 865 deletion.



Figure 5. Conserved residues in the domain of unknown function of FipA are essential for interaction with FlhF. (A) Weight-based consensus sequence of the conserved region of DUF2802 as obtained from 481 species. The residues targeted in the FipA orthologs of *V. parahaemolyticus, P. putida* and *S. putrefaciens* are indicated along with their appropriate residue position. (B, C) Bacterial two-hybrid assay of FipA variants of *V. parahaemolyticus* (B) or *P. putida* (C) with an alanine substitution in the conserved residues indicated in (A). The constructs were tested for self-interaction and interaction with FlhF. *In vivo* interaction of the 875 fusions in *E. coli* is indicated by blue coloration of the colonies. (D) Left panel: Single-cell tracking of *V. parahaemolyticus* strains expressing FipA bearing the indicated substitution in the DUF2802 domain (see Figure 2B for wild-type behavior). Right panel: Corresponding

quantification of the swimming speed. Asterisks indicate a p-value <0.05 (ANOVA + Tukey test)

(F) Localization of VpFlhF in the absence of FipA or in cells with substitutions in the DUF2802
domain. Left: Micrographs showing the localization of FipA-sfGFP in the indicated strains; the upper panels display the DIC and the lower panels the corresponding fluorescence images. The scale bar equals 5 μm. Right: the corresponding quantification of the FlhF-sf-GFP patterns in the indicated strains. (E) Soft-agar spreading assays of *P. putida* wild-type and indicated mutant strains, asterisks display a p-value of 0.05 (*) or 0.01 (**) (ANOVA). (G) Localization of *P. putida* FlhF in strains bearing substitutions in the DUF2802 interaction site. Left: micrographs displaying the localization of FlhF-mCherry in the indicated strains. Upper panels show the DIC and lower panels the corresponding fluorescence images. The scale bar equals 5 μm. Right: Corresponding quantification of the FlhF localization pattern in the indicated

strains. Data for *S. putrefaciens* is displayed in **Supplementary Figure S7**.



Figure 6. The localization pattern of FipA. (A, B) Localization pattern of fluorescently labeled
FipA in *V. parahaemolyticus* and *P. putida*. (A) Representative micrographs of *V. parahaemolyticus* expressing FipA-sfGFP from its native promoter. Scale bar = 2 μm. The upper panel shows the DIC and the lower panel the corresponding fluorescence channel. To the right the localization was quantified accordingly. (B) The same analysis for *P. putida*. (C, D) Time lapse analysis of FipA-sfGFP localization over a cell cycle in *V. parahaemolyticus* (C) and *P. putida* (D). The numbers in the upper DIC micrographs show the minutes after start of the experiment. The scale bars equal 1 μm (C) and 5 μm (D).



Figure 7: Normal localization of FipA depends on interaction with FlhF. (A, B) Localization pattern of V. parahaemolitycus FipA-sfGFP in the indicated wild-type and mutant strains. The upper panels display the DIC micrographs, the middle panel the corresponding fluorescence 905 imaging (scale bar equals 5 μ m), and the lower panel the corresponding demograph showing the fluorescence of FipA-sfGFP along the cell length. (C) Quantification of the cell localization pattern from the experiment shown in (A, B) as combined from three biological replicates. (D, E, F) The same analysis for the corresponding P. putida strains as indicated. The scale bar equals 5 μm. The data for *S. putrefaciens* is displayed in **Supplementary Figure S9**.