

The protein network of bacterial motility

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Motility is achieved in most bacterial species by the flagellar apparatus. It consists of dozens of different proteins with thousands of individual subunits. The published literature about bacterial chemotaxis and flagella documented 51 protein–protein interactions (PPIs) so far. We have screened whole genome two-hybrid arrays of *Treponema pallidum* and *Campylobacter jejuni* for PPIs involving known flagellar proteins and recovered 176 and 140 high-confidence interactions involving 110 and 133 proteins, respectively. To explore the biological relevance of these interactions, we tested an *Escherichia coli* gene deletion array for motility defects (using swarming assays) and found 159 gene deletion strains to have reduced or no motility. Comparing our interaction data with motility phenotypes from *E. coli*, *Bacillus subtilis*, and *Helicobacter pylori*, we found 23 hitherto uncharacterized proteins involved in motility. Integration of phylogenetic information with our interaction and phenotyping data reveals a conserved core of motility proteins, which appear to have recruited many additional species-specific components over time. Our interaction data also predict 18 110 interactions for 64 flagellated bacteria.

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Introduction

Motility in most bacterial species depends on a sophisticated molecular machine called the flagellum. The flagellar apparatus is made of dozens of different proteins and thousands of individual subunits. The bacterial flagellum is actually a mechanical nanomachine with a rotation frequency of 300 Hz, an energy conversion rate of nearly 100%, and the ability to self assemble (Macnab, 1999, 2003; Berg, 2003; Kojima and Blair, 2004).

Various efforts have been made to identify all components required for bacterial motility, resulting in a list of more than 60 proteins in *Escherichia coli* (Supplementary Table S1, Kanehisa *et al.*, 2004). Functionally, these proteins can be subdivided into several subsets: the chemotaxis system connects environmental stimuli to the direction of flagellar rotation and thus direction of movements. The chemotaxis system is connected to the basal body complex, which anchors the flagellum in the inner membrane and also incorporates a type-III-secretion system necessary for the self-assembly

process of the flagellum. Two motor proteins, MotA and MotB, convert an ion gradient (for most bacteria a proton gradient) into rotational energy of basal body components; these components are connected to the rod structure and then via a flexible hook to the filament of the flagellum, which operates like a propeller. However, whereas the overall structure has been known for decades, many of the mechanistic details responsible for the assembly and operation of the motor have yet to be worked out. In fact, it remains unclear whether all the protein components of the flagellar apparatus have been identified. Similarly, whereas at least 51 protein–protein interactions (PPIs) have been described in the literature (Supplementary Table S2), many more interactions are likely to be required for assembly and proper operation.

Despite the vast body of literature about bacterial motility, there have been few systematic attempts to identify the components of the flagellar apparatus and their function besides genome sequencing. Systematic analysis of hundreds of completely sequenced genomes, for example, has predicted many additional motility genes based on their location in

flagellar operons or gene clusters, yet their actual roles in motility often remain unknown.

In this study, we systematically identified genes essential for bacterial motility by testing the swarming capability of 3985 gene deletion strains of *E. coli* (Baba *et al*, 2006). In addition, we integrated data from similar screens carried out for *Bacillus subtilis* (Schumann *et al*, 2001) and mutant screens of *Campylobacter jejuni* (Golden *et al*, 2000; Hendrixson *et al*, 2001) and *Helicobacter pylori* (Salama *et al*, 2004).

Second, we screened all motility proteins recovered from the literature for PPIs. We reasoned that unknown motility proteins can be discovered by interactions with known flagellar and chemotaxis components. Protein interactions were identified by screening the proteomes of two small distantly related bacteria, *Treponema pallidum* and *C. jejuni*, using comprehensive array-based yeast-two-hybrid screens (Uetz *et al*, 2000; Parrish *et al*, in press). In addition, we compared our data to protein interaction data of *H. pylori*

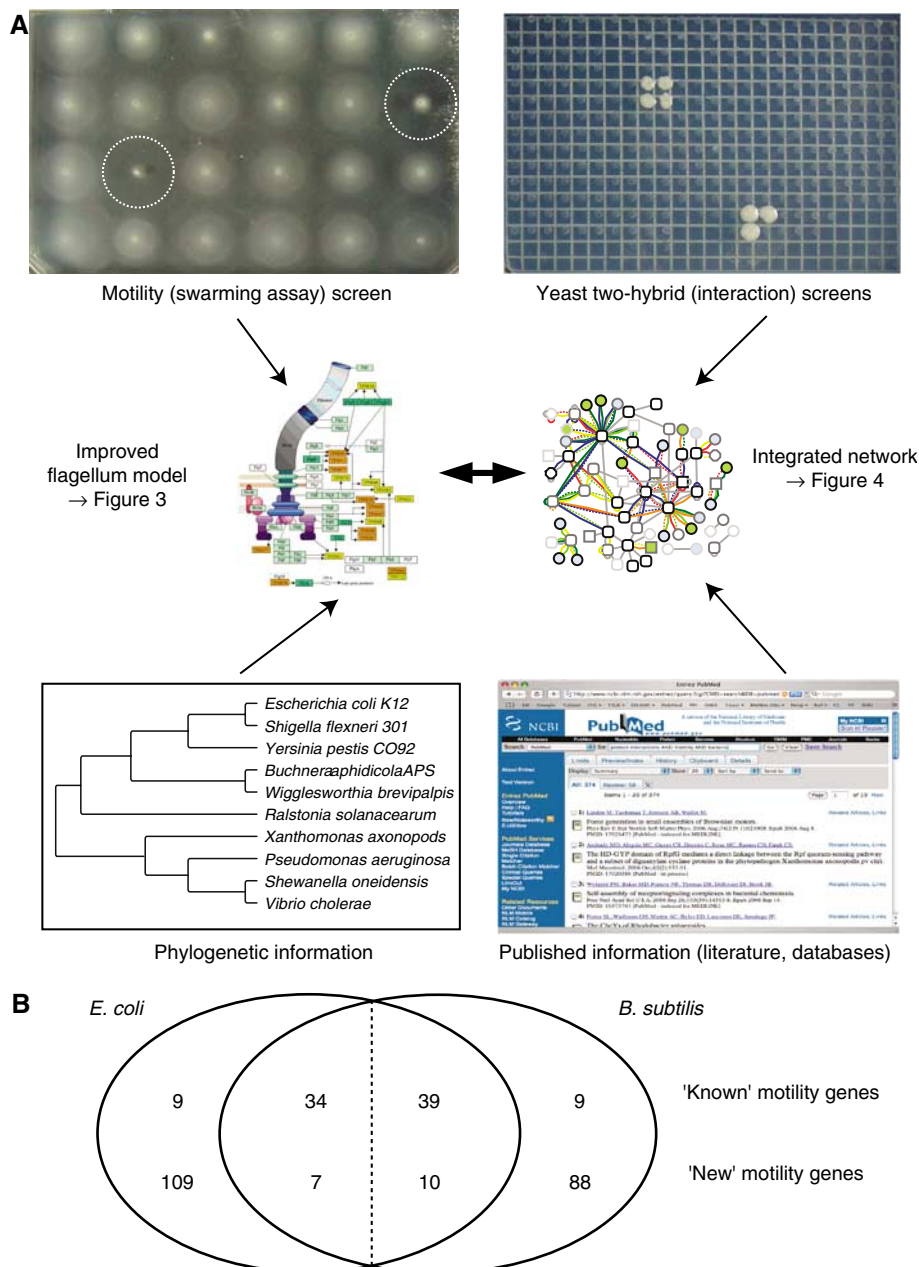


Figure 1 Strategy used in this study. **(A)** Motility genes were studied by screening 3985 *E. coli* mutants for motility defects (24 are shown in the top left panel) and by screening motility proteins for protein interactions (here: bait FlaB1 from *T. pallidum* returning preys FlaB1 and FliS). These data were integrated with literature interactions and large-scale interaction data sets to refine the current picture of the bacterial flagellum. **(B)** Comparison of motility genes identified in *E. coli* (this study) and *B. subtilis* (Schumann *et al*, 2001). There are 43 'known' motility genes reported in the literature for *E. coli*, 34 of which have orthologs in *B. subtilis*. Our mutant screens identified 116 additional 'new' motility genes of which only 7 have orthologs in *B. subtilis*. Equivalent numbers are given for *B. subtilis* (individual genes are listed in Supplementary Tables S1 and S3).

Table I Novel motility proteins

COG	Gene locus (ORF)	Mutant gene (ortholog)	Interacting proteins	Motility	Notes [Refs]
COG1463 COG0082	CJ1648 HP1464 CJ1634C	— <i>aroC</i> (ECO)	MotB FlhA, FlgE2, FliE, FlgK, FliD, FlgB	—/ + ^a	Rain <i>et al</i> (2001) See Figure 4
COG0055 COG2194 COG0642 COG2604 None	B3732 CJ0256 CJ1222c CJ1318 HP0488	<i>atpD</i> (ECO) CJ0256, <i>yjgX</i> (ECO) CJ1222c (CJE) CJ1318 (CJE) CJ1340c (CJE)	CheZ, CheW, Tsr (<i>E. coli</i>) FlgG-2 FliN FliN, FliQ FlgE, FlgB	—/ + ^a — ^a — — —	Arifuzzaman <i>et al</i> (2006) Golden <i>et al</i> (2000) Coloniz. ^b ↓ MacKichan <i>et al</i> (2004) Golden <i>et al</i> (2000) FlhA + FlhF ^c Rain <i>et al</i> (2001); Niehus <i>et al</i> (2004)
COG1923 COG0455	B4172 TP0712 (CJ0063c)	<i>hfq</i> (ECO) HP1034 (HPY)	FliS, FliL TP0464 (flhF [CJE])	—/ + ^a —	Arifuzzaman <i>et al</i> (2006) van Amsterdam and van der Ende (2004)
COG0457 COG0673 COG0791 None	HP1479 CJ0504c Cj1653c CJ0055c	HP1479 (HPY) <i>idh</i> (BSU) <i>lytF</i> (BSU) <i>rapG</i> (BSU), HP1479 (HPY)	FlgB FlaG FliL FliM	— — — —	Rain <i>et al</i> (2001); Salama <i>et al</i> (2004) Schumann <i>et al</i> (2001) Schumann <i>et al</i> (2001) Schumann <i>et al</i> (2001); Salama <i>et al</i> (2004)
COG0267 COG1774	TP0209 TP0046	<i>rpmJ</i> (ECO) <i>yaaT</i> (BSU)	FliG-2, CheR, FliY, FlgE, FliS CheR, FlgD, MetK, FlaB3, TP0959, CheW-2, Mcp2-3	—/ + ^a —/ +	Schumann <i>et al</i> (2001)
COG0084 COG0346 COG1512	TP0979 CJ1301 TP0561	<i>ycfH</i> (<i>tatD</i>) (ECO) <i>ydfO</i> (BSU) <i>ydjH</i> (BSU)	FliE FliM, FliY SigG, FliF, FlhB, FliR, FliQ, FliL	+ + ^a — —/ + ^a	See Figure 5 Schumann <i>et al</i> (2001)
COG1664 COG2001 COG3334	TP0048 TP0383 CJ1496c	<i>yhbE yhbF</i> (BSU) <i>yllB</i> (BSU) <i>ylxF</i> (BSU)	FliY, FliS FliS, FlgK FliL	—/ + ^a —/ + ^a —	See Figure 4B Kobayashi <i>et al</i> (2003) Operon, domain ^d Schumann <i>et al</i> (2001)
COG3391 COG1699	TP0421 TP0658	<i>yncE</i> (ECO) <i>yviF</i> (BSU)	TP0567 FlaB2, FlaB1, FlaB3	—/ + ^a — ^a	Titz <i>et al</i> (2006)

Conserved hypothetical proteins in the columns 'Gene locus (ORF)' and 'Mutant gene (ortholog)' are predicted to be novel flagellar components, based on their physical interactions with known motility proteins (in *T. pallidum*, *C. jejuni*, *E. coli*, or *H. pylori*) and their motility phenotype (in *E. coli* (ECO), *H. pylori* (HPY), *C. jejuni* (CJE), or *B. subtilis* (BSU)). The Motility phenotype refers to the Mutant gene (ortholog): reduced motility '—/ +', nonmotile '—', and increased motility '+ +'. Entries are sorted by 'Mutant gene'.

^aThis study (Motility phenotype).

^bDiminished capacity to colonize.

^cRegulated by FlhA and FlhF.

^dFlagellum operon, extracellular sensory domain.

(Rain *et al*, 2001) and *E. coli* (Arifuzzaman *et al*, 2006). Finally, we integrated these experimental data sets with predictions of functional associations from the STRING database (von Mering *et al*, 2003; Stein *et al*, 2005). The result is a list of known and new flagellar components, including 23 novel motility proteins (Figure 1 and Table I).

Many features of the bacterial flagellum have changed over the course of evolution. This is reflected in the surprisingly different composition and protein interaction patterns in the flagella of different species, which may reflect adaptations to species-specific motility needs (compare Figure 2 and Supplementary Figure S2). While the overall conservation allows us to predict ~18000 interactions for 64 proteomes of flagellated bacteria, it remains to be seen how many of them are functional.

Results

Genes important for bacterial motility

Several systematic mutant screens have been performed to find genes involved in bacterial motility (Hendrixson *et al*, 2001; Schumann *et al*, 2001; Golden and Acheson, 2002; Inoue *et al*, 2007). To generate a comprehensive motility mutant data set for the gram-negative model bacterium *E. coli*, we have used the gene deletion library constructed by Baba *et al* (Baba

et al, 2006). These mutants were plated out in arrays of 24 colonies on swarming agar and tested for swarming (Figure 1). Of 3985 mutants tested 159 deletions showed a swarming defect (Supplementary Table S3a).

Interestingly, a similar screen in *B. subtilis* yielded a similar number of 146 motility mutants (Schumann *et al*, 2001) (Supplementary Tables S3b and 3f). Thus about 4% of the nonessential genes in both species show an effect on motility under the conditions tested. Among them are 43 (30%) and 48 (27%) genes previously annotated as *bona fide* motility genes in *E. coli* and *B. subtilis*, respectively (Figure 1B). The other mutants with motility phenotypes are significantly enriched for proteins involved in 'motor activity' and 'macromolecule metabolism' (Supplementary Figure S1). Many of these genes may be required to provide energy to the flagellar motor or may be indirectly involved in the assembly of the flagellar apparatus, for example in restructuring the peptidoglycan to allow penetration of the cell wall.

Unexpectedly, only 7 (10) of the *E. coli* (*B. subtilis*) mutant genes that were previously not known to have a motility function, had a homolog with a phenotype in *B. subtilis* (*E. coli*) (Figure 1B and Supplementary Table S3g). Thus, there appear to be many proteins with a species-specific role in motility. Examples of such proteins are discussed further below.

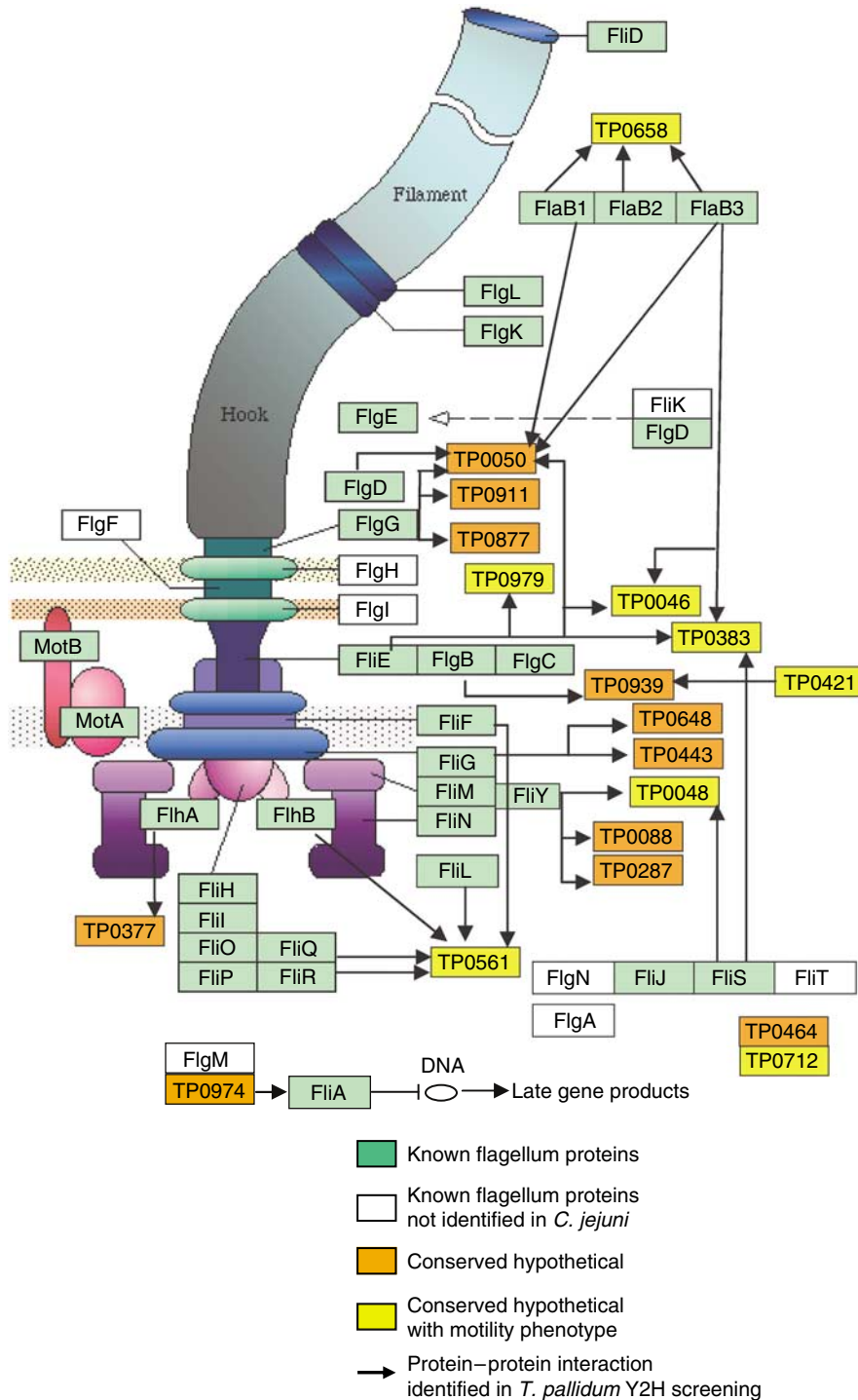


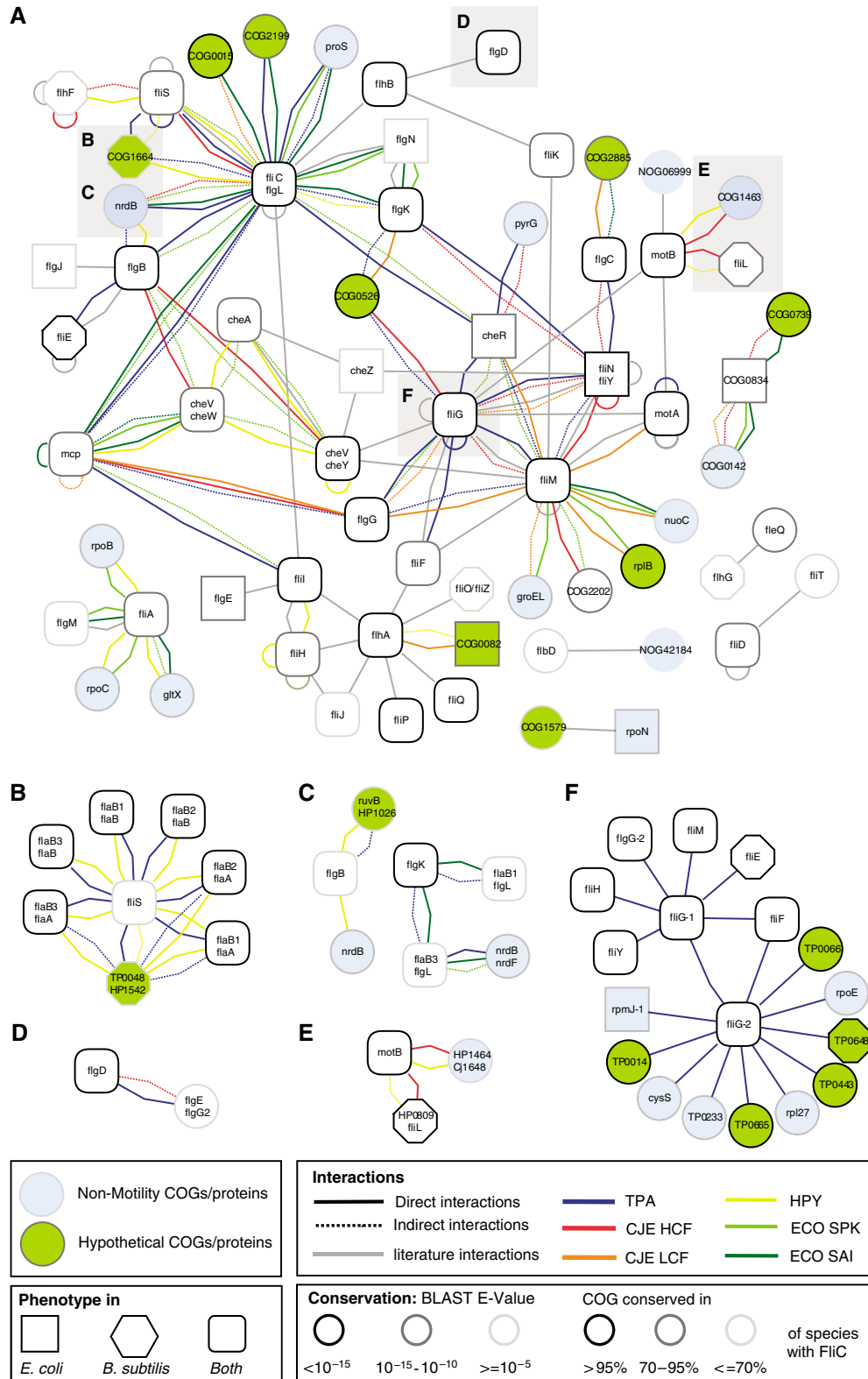
Figure 3 New motility proteins in the context of the *T. pallidum* flagellar apparatus. Model of the bacterial flagellar apparatus (based on KEGG; Kanehisa *et al*, 2006) with putative new motility proteins discovered in this study (yellow and orange). New motility proteins of *C. jejuni* are shown in Supplementary Figure S3. Compare to Figures 2 and 4.

(ECO SPK) versus the *Helicobacter* data to 25.0% for *Helicobacter* versus CJE HCF (Supplementary Table S5). Overall, ECO SPK has the weakest pairwise similarities. Thus the overlap between the different data sets appears to reflect both phylogenetic relationships as well as methodological differences between yeast two-hybrid and complex purification data sets.

As might be expected, interactions between motility proteins are common in the motility interaction maps (Figure 2 and Supplementary Figures S1 and S2). An overview of the number of proteins (nodes) and their interactions (edges) and additional properties of these networks can be found in Supplementary Table S6.

Finally, for a comparative analysis of motility interactions, we carried out a comprehensive review of the literature for published flagellar PPIs using PubMed and found 51 unique

interactions (Supplementary Table S2a). Of these 51, 39 had interologs in *T. pallidum* and 38 in *CJE ALL*, but only 9 and 5 were reproduced in *T. pallidum* and *CJE ALL*, respectively



(Supplementary Tables S2 and S5a). Only one interaction is common to both *T. pallidum* and CJE ALL screens, and thus a total of 13 interactions were recovered in either of our screens. That is, sampling of the two species recovered 33% of all published flagellar interactions. One reason for this relatively low coverage may be that most previous studies used different methods that may be better applicable to flagellar proteins. All literature comparisons can be found in Supplementary Table S5.

An integrated view of the flagellum

The diversity of information on different genomes, proteins, phenotypes and so on makes it difficult to keep track of all details. Therefore, we generated an integrated motility network, which combines a diverse set of interaction networks as well as phylogenetic and phenotyping data (Figure 4). This network combines protein–protein interactions of *T. pallidum*, *C. jejuni*, *H. pylori*, and *E. coli* as well as interactions curated from the literature with motility phenotyping data from *E. coli* and *B. subtilis*. It also displays clusters of orthologous groups (COGs) rather than individual proteins, which reduces complexity and improves the quality of links. These links represent direct interactions, indirect interactions (if proteins do not interact directly, but via a bridging protein), and literature interactions. Out of all interactions, 73% connect known motility COGs. In addition, 45% were predicted by STRING (highest confidence: $S > 0.9$) to be strongly associated. These numbers indicate that this integrated network is more reliable and biologically relevant than individual networks. In addition, links among orthologous groups can usually be transferred to proteins of other species. However, because of the stringent filtering not all interactions are included in this network.

Several insights into the internal organization of the bacterial flagellum can be obtained. For example, the aligned network shows that the flagellum filament protein, FliC, and its homolog FlgL, a hook-associated protein, are members of the same COG. FlgL is connected to the second hook-associated protein FlgK and both are stabilized by their export chaperone, FlgN. The interaction of FliC with its chaperone FliS is conserved in all species. The basal body complex with FliN/FliY, FliG, FliM, and FliF, forms another cluster, which is connected to the motor proteins, MotA and MotB, and to rod proteins such as FlgC and FlgG.

The chemotaxis protein cluster in the network is only connected to the flagellum switch complex. The interaction of CheY with FliM depends on CheY's phosphorylation, which is not detected in our yeast two-hybrid assays, because we do not coexpress the pertinent kinase CheA. Nevertheless, the integrated network reflects the fact that external signals are detected by homodimerizing methyl-accepting chemoreceptors (Mcps), which are linked by an adapter protein, CheW, to

the kinase CheA, which transfers the phosphate group to CheY. By its interaction with the basal body complex, phosphorylated CheW controls the rotation state of the flagellum. In addition to these previously known interactions, we find conserved links between chemotaxis proteins and rod proteins such as FlgB and FlgG, which are difficult to explain by the standard model of the flagellum, but allow for interesting speculations about the organization of chemotaxis signalling in the cytoplasm.

Another striking connection is the conserved MotB–FliL interaction in *C. jejuni* and *Helicobacter*. For *Proteus mirabilis*, FliL is thought to be involved in sensing of the actual flagellum status (Belas and Suvanasuthi, 2005). Here, we found evidence that this sensing is mediated by a direct interaction with the motor apparatus (Figure 4E).

Discussion

New bona fide motility proteins

A major goal of this study was to find novel flagellar components among the many proteins of still unknown function. In addition, we suspected that there must be previously characterized proteins whose role in motility remained unknown. Indeed, 28% and 33% of the interactions found in *T. pallidum* and *C. jejuni*, respectively, connected a known motility protein to a conserved hypothetical protein (Supplementary Table S4), suggesting that there are still unidentified proteins with a motility function. To identify potential novel bona fide motility proteins, we used our integrated data and identified 23 hitherto uncharacterized proteins (Table I).

For example, members of the orthologous group COG1664, such as TP0048 and HP1542 (Figure 4B), show interactions to the FliC–FliS cluster. Additional evidence for a role in motility comes from the double mutant of the *B. subtilis* orthologs *yhbE* and *yhbF*, which also shows reduced motility.

TP0658 (*yviF* in *B. subtilis*), a previously uncharacterized protein, was found to interact with all three flagellin proteins (FlaB1–B3) of *T. pallidum*. The deletion mutants of TP0658 orthologs in both *B. subtilis* (*yviF*) and *C. jejuni* (CJ1075) show a highly reduced motility phenotype (Golden *et al*, 2000; Titz *et al*, 2006). We have recently shown that TP0658 and *yviF* appear to stabilize flagellin in the cytoplasm, thus exhibiting properties of a chaperone (Titz *et al*, 2006). TP0658 and its orthologs thus appear to be flagellar assembly factors or factors involved in export of the filament protein FliC.

TP0561 is another hitherto uncharacterized protein that appears to be involved in flagellar protein export based on its interaction pattern; it interacts with multiple components of the export machinery such as FliR, FliL, FliQ, and FlhB (Figure 3, bottom left). In addition, a mutation in TP0561 results in significantly reduced motility.

Figure 4 Integrated motility network. Multispecies summary of interactions among motility proteins. (A) Homologous protein nodes of pairwise aligned networks were merged into orthologous groups (COGs). Nodes were labelled according to KEGG descriptions (Kanehisa *et al*, 2006); their shape corresponds to motility phenotypes in *E. coli* (squares), *B. subtilis* (octagons), or both (rounded squares). Edges represent either direct, indirect, or literature interactions among orthologous groups and are color-coded (see legend in Figure). The border shading of a COG nodes indicates its conservation ratio among 68 flagellated bacteria. (B–E) Aligned protein interactions showing COG interactions from (A) in more detail: each node represents two homologous proteins from either *T. pallidum* (TPA), *C. jejuni* (CJE), *H. pylori* (HPY), or *E. coli* (ECO). Edges represent either direct or indirect interactions and are color-coded (see legend in Figure). The border color of a protein node indicates the BLAST *E*-value of its homologous protein pair. (F) Interactions of paralogous Flig1 and Flig2 proteins taken from *T. pallidum* network.

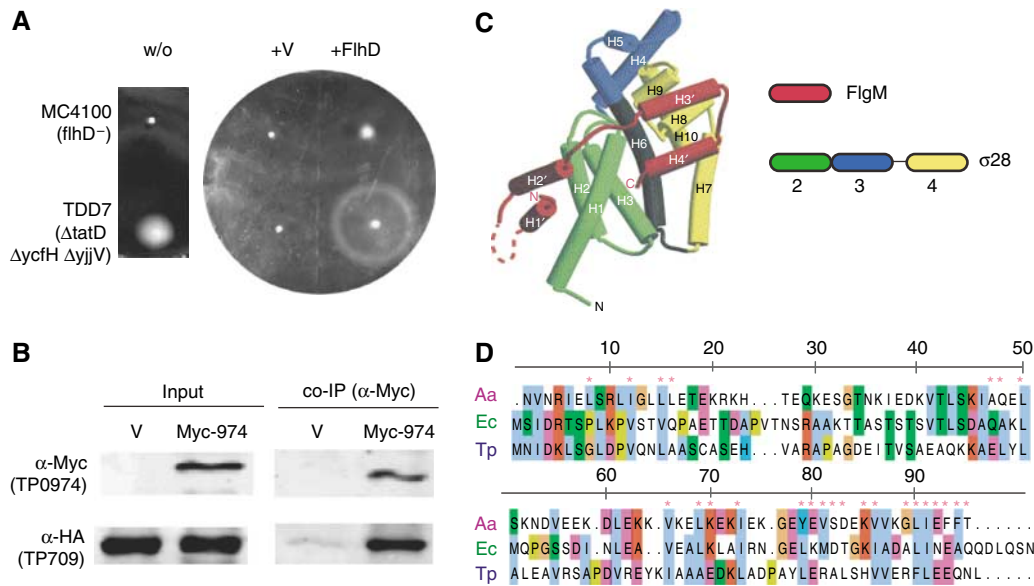


Figure 5 Lessons for individual proteins. **(A)** *E. coli* *tatD* mutants were tested in a swarming assay. The *E. coli* strain MC4100 (*flhD*⁻) and a *tatD* triple mutant (TDD7, all *TatD* paralogs deleted, i.e., *tatD*, *ycfH* and *yjiV* in a MC4100 background) were left untransformed (w/o) or transformed with an empty expression vector (+V) or the same vector expressing FlhD (+*flhD*). The *tatD* triple-mutant was much more motile than the parental strain. **(B)** TP0974 is the *T. pallidum* FlgM protein. Myc-tagged TP0974 (Myc-974) or a vector control (V) was co-immunoprecipitated with HA-TP0709 (sigma factor). Note that the results from (A) and (B) are mentioned in the discussion, not results section. **(C)** Crystal structure of the *Aquifex aeolicus* σ 28/FlgM Complex (Sorenson *et al*, 2004). The domain architecture and color-coding are illustrated schematically. FlgM is colored red. **(D)** The homolog of FlgM in *T. pallidum* (TP0974) aligned to FlgM from *A. aeolicus* and *E. coli* using ClustalW (Chenna *et al*, 2003) (Ae=*A. aeolicus*, Ec=*E. coli*, Tp=*T. pallidum*). The colors of amino acids denote chemical similarity. Numbering is according to the *A. aeolicus* sequence. A pink asterisk indicates residues mediating the interaction of FlgM with σ 28 (showing the conservation of the FlgM interaction epitope). The putative *C. jejuni* FlgM homolog CJ1464 was left out as its function is still unclear.

Due to its motility phenotype in *E. coli* *TatD* is an exceptional case. Its *Treponema* homolog TP0979 interacted with FliE. In *E. coli*, *tatD* has two paralogs, *ycfH* and *yjiV*, which are functionally unassigned, genomically unlinked, and show 29% and 24% amino-acid sequence identity to *TatD*, respectively. As *tatD* is localized in an operon with genes of the twin-arginine transport (Tat) system, a transport function of *TatD* was anticipated. But even a strain with all three *TatD* paralogs deleted did not show a *Tat*-related transport deficiency, leaving the question for *TatD*'s function unanswered (Wexler *et al*, 2000). As we found a very small, but significant, increase in motility for the *ycfH* single mutant, we tested the previously described triple mutant (all *TatD* paralogs deleted) for motility (Figure 5A). The triple mutant was constructed in a MC4100 strain background, a strain known to be nonmotile, presumably due to a point mutation in *FlhD*, a known master regulator for motility (Wexler *et al*, 2000). Unexpectedly, the triple mutant showed a slight rescue of motility. We investigated the relation between the *FlhD* point mutation and the *TatD* paralogs by expressing a functional *FlhD* construct both in the MC4100 strain and the triple mutant. A strong synergistic effect of *FlhD* expression and the triple mutation on motility was observed pointing to a regulatory antagonism of *FlhD* and *TatD* paralogs (Figure 5A). These findings indicate that *TatD* and/or its orthologs (COG0084) have a negative role in motility, perhaps mediated by its DNAse activity (Wexler *et al*, 2000).

A few proteins were previously annotated as having functions unrelated to motility, but appear to be involved

in flagellar function based on their motility phenotype and their interactions with other motility proteins. Among these proteins, ribosomal protein RpmJ/L36 (TP0209) is an unexpected case, because it not only interacted with multiple flagellar proteins, but also showed a reduction of motility to about 14% of the wild type when mutated. It is possible that RpmJ links protein synthesis to the flagellar protein export machinery. Another one, Rpe (TP0945) is a ribulose-phosphate 3-epimerase. Surprisingly, this protein also showed a strong reduction in motility when deleted. Rpe could have a role in providing energy for the flagellar motor.

Novel interactions and functions of known motility proteins

Overall, most parts of the flagellum are well conserved in motile bacteria. Nevertheless, evolutionary adaptation of several components can clearly be identified and range from the duplication of proteins, for example of flagellins, to the complete loss or gain of components, for example of export chaperones (Pallen and Matzke, 2006). Here, we will give three examples for such evolutionary processes on the interaction level.

Generation of rotational asymmetry by FliG paralogs

The flagellum of spirochetes possesses several unique features not found in other bacterial species. One unique feature, for example, is the periplasmic localization of two polar

flagellum bundles, which perform an asymmetrical rotation (Charon *et al*, 1992; Li *et al*, 2000). Interestingly, the molecular basis of this asymmetry is unknown. One hypothesis states that chemotaxis plays an essential role in this asymmetry (Berg and Anderson, 1973; Berg, 1975; Armitage, 1999). Another hypothesis assumes that the motor complexes at both cell poles are somehow differently organized. One candidate that might lead to this asymmetry is FliG, as it is the only duplicated basal body complex protein in spirochetes. The paralogs of FliG in *T. pallidum* are named FliG-1 (TP0026) and FliG-2 (TP0400). Strikingly, despite significant protein sequence identity (~30%), both proteins show a very different interaction pattern in our Y2H study (Figure 4F): both proteins interact with each other and with FliF, the flagellum rotor protein. In addition, FliG-1 interacts with FliY, FliM, FliH, and FliE, whereas FliG-2 interacts with TP0014, TP0066, TP0665, TP0443, and TP0648 (all proteins of unknown function), and an anti-sigma factor (TP0233). Although the differential interaction patterns do not clearly explain the asymmetric behavior, they suggest a possible basis for this asymmetry, for example, recruitment of different proteins to the two motor complexes at both ends of the cell. However, it remains to be shown that the two FliG variants are differentially localized and indeed have different effects on motor activity.

Sigma factor and anti-sigma factor interactions

During flagellar assembly in *E. coli* and *Salmonella*, the expression of the late flagellar genes is timed by the anti-sigma factor FlgM. At the beginning of assembly, FlgM binds to its sigma factor FliA/ σ^{28} and blocks its function (Figure 5C). Later, the assembled export machinery of the basal body exports FlgM, releases its inhibition of the sigma factor, and the sigma factor becomes free to activate transcription of the late flagellar genes. Initially, no copy of FlgM was identified in the genome of *T. pallidum* and *C. jejuni*, which might have implied that the timing of flagellum assembly is differently regulated in these species. Recently, Pallen *et al* (2005) computationally identified remote FlgM homologs in both species: TP0974 for *T. pallidum* and CJ1464 in *C. jejuni*. TP0974 has 23% and 24% sequence identity to its homologs in *E. coli* and *B. subtilis*, respectively, and both alignments require multiple gaps, despite its short sequence (around 90 amino acids) (Figure 5D). In our study, we experimentally confirmed the interaction of both putative FlgM homologs with their respective sigma factors: TP0974 interacts with the sigma-factor TP0709, which was confirmed by co-immunoprecipitation (Figure 5B), and CJ1464 interacts with the sigma factor CJ0061c (FliA, σ^{28}). Thus, we provide experimental evidence that FlgM (and its function) is conserved in *T. pallidum* and *C. jejuni*. However, it should be noted that flagellin transcription in *C. jejuni* requires both σ^{54} and σ^{28} and that CJ1464 does not appear to have a strong influence on σ^{28} -dependent transcription of flagellin (Hendrixson and DiRita, 2003). Thus, whereas TP0974 may function as an anti-sigma factor, CJ1464 may not. This example shows that large-scale interaction data sets can provide useful initial experimental evidence for functional predictions whose mechanistic details, however, have to be worked out by additional experiments.

Chaperones

Interestingly, two other proteins in the same operon as CJ1464, FlgK (CJ1466) and the remote homolog of FlgN (CJ1465), a chaperone of FlgK, interact with each other, suggesting that CJ1465 likely functions as an FlgK-specific chaperone in *C. jejuni*.

As judged by the differential phylogenetic distributions of different export chaperones (Pallen *et al*, 2005), different functions and substrate specificities are likely to be found in different bacterial species. FliS is thought to be a flagellin-specific chaperone (Ozin *et al*, 2003), whereas FlgN and FliT are substrate-specific flagellar chaperones that prevent oligomerization of the hook-associated proteins, or HAPs, in *S. typhimurium*. Interestingly, FlgN and FliT orthologs are not present in spirochetes (Pallen *et al*, 2005). The interactions between FliS and FlgK and between FliS and FlgB are the first experimental evidence that FliS may also function as a chaperone for FlgB and FlgK in spirochetes, partly substituting for FlgN and FliT.

Connections between motility and other functional classes

Whereas the flagellar apparatus is a well-defined nanomachine, it does not act in isolation. Besides the obvious link to the chemotaxis pathway, we noticed several interactions with proteins of other function (Supplementary Figure 1). For example, a link of motility proteins with 'nucleobase, nucleoside, nucleotide, and nucleic acid metabolism' (GO:0006139) is found in both *E. coli* and *T. pallidum* interaction sets, as well as the *E. coli* mutant phenotyping data. NrdB (ribonucleoside-diphosphate reductase), the key enzyme for the conversion of ribonucleosides into desoxy-ribonucleosides, interacts with two flagellar proteins, FliC and FlgB (Figure 4C). A functional link of NrdB to motility is provided by a study by Nishimura and Hirota (1989), where the authors found a reduction in flagellar protein expression upon *nrdB* deletion. Although the authors assumed a link on the transcriptional level, an additional post-translational link, as indicated by a direct protein interaction, becomes likely.

The functional link between electron transport (electron transport chain) and motility via a proton gradient (or sodium gradient) is well known and also reflected by an association with 'transport' (GO:0006810) in a motility gene expression data set (FlhD) (Pruss *et al*, 2003). We found a direct interaction between NuoC, (NADH dehydrogenase I) and FliM both in the *E. coli* and *C. jejuni* data sets (Figure 4A). This enzyme forms complex I of the electron transport chain and converts the oxidation of NADH into an electrochemical proton gradient. At least in these two species (*T. pallidum* does not have an electron transport chain), motility might be optimized by increasing the local proton concentration.

Motility is known to be regulated by environmental stimuli such as nutrients and this is reflected by the over-representation of 'response to stimulus' (GO:0050896) proteins among flagellar interactors. For example, motility is controlled by the second messenger cyclic-di-GMP, which is produced by the enzymatic activity of so-called GGDEF domains (Ryjenkov *et al*, 2005). Here, we find a conserved interaction of the GGDEF COG, COG2199, with FliC in *T. pallidum* and *E. coli* (Figure 4A), pointing to an important regulatory role of this interaction.

FliA, the sigma factor for several flagellar operons, interacts with two subunits of the RNA-polymerase (rpoB and rpoC) in the *E. coli* and the *H. pylori* interaction sets (Figure 4A). In addition, in the same species, an interaction of FliA with the glutamyl-tRNA synthetase, GltX, was found, suggesting a regulatory role of this interaction.

Thus, a close inspection reveals a number of interesting functional links, which are also supported by the integrated motility network. However, most of these interactions have to be analyzed in more detail to shed more light on their precise biological role and the mechanistic details.

The evolution of the flagellum

Given the amazing complexity of the bacterial motility system, we wondered whether our interaction data and phenotypes can contribute to the understanding of its evolution. As a first step into that direction, we first constructed a phylogenetic supertree of 30 species based on 35 flagellar protein families (Supplementary Figure 4). Our flagellum supertree strongly supports the monophyly of spirochetes, as well as γ and β , ϵ , and α proteobacteria. These relationships are similar to the previously reported phylogenies, for example, an rRNA tree (Olsen *et al*, 1994) and a tree which was based on 31 highly conserved protein families (Ciccarelli *et al*, 2006). This shows that the flagellar system evolved together with other cellular systems and not independently.

Evolution of the flagellum is also consistent with the fact that neither any flagellar proteins nor any of their interactions is conserved. In fact, our *Treponema* data set predicted 173 interactions for *C. jejuni*, of which we found only 49 (Supplementary Table S4d). This indicates that protein interactions may be evolutionarily less conserved than generally believed.

An evolutionary model also predicts that core proteins, which have been associated with the flagellum, should be tightly integrated, and thus have more interactions than peripheral proteins, which have been only recently recruited to the flagellar machinery. Indeed, we did find a weak, but statistically significant linear relationship between the number of interactions of an orthologous group and its conservation ratio among flagellated bacteria ($r=0.43$, $P<0.005$; Supplementary Figure S7). Therefore, our analysis supports the evolution of the flagellum from core components by adding additional ones over time (Pallen and Matzke, 2006).

Conclusions

In this study, we pursue an integrative systems biology approach to assemble a comprehensive picture of bacterial motility. Motility interaction data sets for *T. pallidum* and *Campylobacter pylori* and a genome-wide motility data set for *E. coli* are presented. Our data are combined with functional and interaction data from multiple species to reconstruct an integrated network of bacterial motility. Insights into the internal structure of the flagellum, its connections to other functional classes, and on potentially novel components of the flagellum have been obtained.

Due to the size of our data set, we were able to analyze only a few selected interactions in more detail. We confirmed

the presence of anti-sigma factors (FlgM) in *T. pallidum* and possibly *C. jejuni* based on their interactions with a flagellum-specific sigma factor. Recently, we have assigned a new function to TP0658 (now called FliW), a conserved protein of previously unknown function. We could show that this protein acts as a molecular chaperone and/or assembly factor of the bacterial flagellum (Titz *et al*, 2006).

The bacterial flagellum represents an interesting entity to study the evolution of complex biological machines. For an evolutionary view of the flagellum on the protein level, we constructed a phylogenetic supertree solely based on flagellar protein sequences. As anticipated, this tree closely recapitulates phylogenetic relationships identified, employing traditional phylogenetic marker molecules such as rRNAs.

Whereas it is generally believed that the motility machinery evolved from an ancient type III secretion system, the detailed steps leading to current structures have yet to be defined. A prediction from this theory would be that the conserved core proteins should exhibit more interactions than peripheral proteins. Indeed, proteins which are well conserved and part of all flagellar complexes have more conserved interactions (e.g., FliC, FliG, FliY, FliM, FliA, Mcp, CheW, and CheY) than proteins which are found only in a subset of motility complexes (e.g., FlhF or FlgJ; see Figure 4 and Supplementary Figure S7).

Similar to protein sequences and structures, interactions among proteins are often conserved in the course of evolution. In fact, the phylogenetic relationships of different species are partially reflected by the phylogenetic interaction profile of the integrated network (Supplementary Figure S4).

Finally, we could thus use our interaction data sets to predict interactions in other bacterial species. To obtain only high-confidence predictions, we used our integrated motility network, that is, all interactions found in more than one species or supported by other evidence from the literature, and predicted ~18 000 interactions for 64 flagellated bacteria (Supplementary Table S7). It remains to be seen which of these interactions do indeed occur and what specific role they play in each of these organisms.

Materials and methods

Collection of known motility proteins

We collected motility genes from three major classification systems: KEGG (including motility, chemotaxis, and flagellar assembly; Kanehisa *et al*, 2006), TIGR (including chemotaxis and motility; Peterson *et al*, 2001), and GO (including GO:0019861 flagellum; Camon *et al*, 2004). Data were compiled in March 2007. In total, we identified 293 proteins (in *T. pallidum*, *C. jejuni*, *H. pylori*, *E. coli*, and *B. subtilis*) with at least one classification evidence. Among those, 89% were classified by KEGG, 75% by TIGR, and 65% by GOA. As KEGG provides the most comprehensive classification, we have used the KEGG motility collection throughout this study and refer to its proteins as 'known motility proteins' (Supplementary Table S1).

Orthologous relationships

Throughout this study, we used COGs to infer orthologous relationships between proteins of different species.

COGs were taken from the NCBI COG database (downloaded March 2006) and complemented by COGs from the STRING version 3 database (including nonsupervised COGs) (von Mering *et al*, 2003).

Swarming (motility) assays and phenotyping data sources

A systematic single-gene knockout collection of *E. coli* of 3985 individual mutant strains (Baba *et al*, 2006) was tested for altered motility by a swarming assay. Each gene mutation was tested in two independent strains as provided by the Keio collection. Strains were grown to saturation in LB medium at 37°C (mutants with growth defects were not considered for the motility assay) and transferred to Omnitrays (Nunc) with swarming agar (LB medium with 0.25% Agar) in a 24 colonies per plate format by pin replication with a Biomek 2000 laboratory robot (Beckman-Coulter). The swarming diameters of the mutant strains were compared after ~8 h incubation at 37°C and mutants with reproducible reduced motility were retested in individual swarming assays. The swarming behavior of each mutant was classified as wild type, reduced (reduction by at least 50%), or nonmotile (reduction by at least 90%), as measured by the diameter of the bacterial colony (Figure 1A).

Additionally, we constructed a few individual gene deletions of *E. coli* and *B. subtilis* and tested them for motility: gene disruptions of *yjeK*, *yncE*, and *ycfH* were performed in *E. coli*, as described by Datsenko and Wanner (2000). The *B. subtilis* mutants of *yviF*, *ydjH*, and *yhbE/yhbF* (double mutant) were obtained by phleomycin-cassette integration, as described by Fabret *et al* (2002). The PCR primers used are listed in Supplementary Table S8. Information on gene deletions affecting motility in *B. subtilis*, *H. pylori*, and *C. jejuni* was taken from the literature (Golden *et al*, 2000; Schumann *et al*, 2001; Kobayashi *et al*, 2003; Salama *et al*, 2004).

The motility Y2H protein–protein interaction network of *T. pallidum*

Forty-nine *T. pallidum* proteins, which are part of our KEGG motility collection, were selected (Supplementary Table S1). Bait fusions (Gal4-DNA-binding domain) of these proteins were constructed by Cre-loxP mediated recombination of pUni clones (Liu *et al*, 2000; McKeivitt *et al*, 2003) with two bait vectors: pAS1 and pLP-GBKT7Amp (created by replacing Kan[®] by Amp[®] in pLP-GBKT7 (Clontech)). A systematic whole-genome prey library for *T. pallidum* was created by transferring all ORFs from their original pUni-vector vector (McKeivitt *et al*, 2003) to our prey vector, pLP-GADT7 (Clontech) by Cre-LoxP-mediated recombination. All prey and bait clones were then individually transformed into Y187 (MAT α) and AH109 (MAT α) (Harper *et al*, 1993; James *et al*, 1996) yeast strains, respectively, by a standard LiAc protocol. Prey strains were arrayed onto 384-well formatted Omnitray-agar plates (Nunc) and each bait strain was individually tested against the whole *T. pallidum* prey array using a previously described array-based Y2H procedure (Cagney *et al*, 2000).

The motility Y2H protein–protein interaction network of *C. jejuni*

Interactions involving the 46 *C. jejuni* proteins assigned to the motility category in the KEGG database (Kanehisa *et al*, 2002) (Supplementary Table S1) were identified in proteome-wide two-hybrid screens, using a pooled matrix approach as described previously (Zhong *et al*, 2003) (Parrish *et al*, in press).

The motility Y2H protein–protein interaction network of *H. pylori*

The motility PPI set '*Helicobacter*' (HPY) was generated by selecting all interactions of known motility proteins from Rain *et al* (2001) (also see motility filtering below). Note that Rain *et al* (2001) tested only 261 bait fusion proteins (out of 1590 ORFs) against a random prey library.

Literature interactions

To mine all the published PPIs of the known flagellum components, we carried out a comprehensive literature review for flagellum PPIs, using the PubMed query '(flagellum OR flagella) AND (interaction OR interact OR interacts OR bind OR binds)' on 13 January 2004. This analysis yielded ~700 abstract/articles from which 51 unique PPIs between flagellar components were manually curated (Supplementary Table S2).

Motility-related interactions derived from *E. coli* complex purification data

Motility-related protein interactions of *E. coli* were derived from Arifuzzaman *et al* (2006), who conducted a comprehensive complex purification study using a His-tagged ORF clone library. A total of 2667 out of 4339 proteins were successfully analyzed and their interacting partners were identified by MALDI-TOF in this study. Complex purification studies do not provide binary interaction data, but only lists of proteins that co-purified with the used bait protein involving both direct and indirect interactions. Arifuzzaman *et al* (2006) provided their results according to the spoke model. We used two models to predict direct interactions for flagellum/chemotaxis proteins (one among the pair of interacting proteins is a known flagellum/chemotaxis protein) from the complex data. The 'ECO SPK' interaction set assumes binary interactions between bait proteins and their co-purifying proteins (SPOKE model). The 'ECO SAI' interaction set is based on a model that has been proposed by Gavin *et al* (2006) to infer complexes from multiple overlapping purifications. Similar to the matrix model, it predicts PPIs among all proteins. However, the difference is that PPIs are weighted according to the pair's propensity to associate with each other relative to what would be expected from their frequency. Based on the cumulative percentage distribution of socio-affinities, we defined the top 25% of PPIs to be highly associated (socio-affinity score > 5). Both ECO sets have been filtered for interactions of known motility proteins (see motility filtering below) (Supplementary Table S2). Data from Butland *et al* (Nature 433: 531, 2005) have not been considered in this analysis as only one flagellar protein, FliY, appears to have worked as a bait in this study.

Motility filtering

T. pallidum, *C. jejuni*, *H. pylori*, and *E. coli* PPIs were filtered for motility interactions by retaining only PPIs, which contain at least one protein which is part of our KEGG motility collection

Integrated protein interaction network of different species

Construction of aligned protein networks

Pairwise alignments of the PPI sets were performed using the Network Comparison Toolkit (NCT, <http://chianti.ucsd.edu/nct/>), a Java implementation of the PathBLAST algorithm, as described previously (Kelley *et al*, 2004). Briefly, the algorithm integrates PPIs from two species with protein sequence homology to generate an 'aligned network'. Homologous proteins (one from each organism) are merged (aligned) into single nodes. We have defined proteins to be homologs if the geometric mean of their *E*-values is $\leq 10^{-5}$ normalized for each genome size based on manual inspections of *E*-values among orthologous interactions (Supplementary Table S5c 'Orthology overlap with *E*-values'). The rule for creating an edge is that one of the pairs of proteins (one species) must represent (have) an interaction (distance 1 edge), whereas the other pair (the other species) can be in one of three states: (i) the other pair is the same protein (distance 0 edge); (ii) the other pair represents an interaction (distance 1 edge); (iii) the two proteins in the other pair do not interact themselves, but interact with a common neighbor (distance 2 edge) also referred to as gap.

Construction of the integrated network

Homologous protein nodes of the pairwise aligned networks were merged into orthologous groups (COGs) if all proteins were members of the same COG. Nodes were labelled according to KEGG (Kanehisa

et al, 2006) (Supplementary Table S1 'KEGG motility COGs') and by manual inspection of the common names of the merged proteins. Edges were directly transferred from the pairwise aligned networks. Finally, we have incorporated interactions among orthologous groups found in our literature set (Supplementary Table S2). COG conservation is based on a COG's conservation ratio among flagellated species, reported in the STRING database (in total 68 species).

STRING confidence score

The confidence score of the StringDB (*S* score) is the approximate probability that a predicted link exists between two enzymes in the same metabolic map in the KEGG database. Confidence limits are as follows: low confidence (*S* score > 0.15) 20% (or better); medium confidence (*S* score > 0.4) 50%; high confidence (*S* score > 0.7) 75%; highest confidence (*S* score > 0.9) 95% (from <http://string.embl.de>).

Co-immunoprecipitation of Myc-TP0974 with HA-TP0709

Myc-tagged TP0974 and HA-tagged TP0709 were cloned into vectors of the pBAD series (Guzman *et al*, 1995) and were co-transferred into BL21 (DE3) *E. coli*. Protein expression was induced with 0.2% (w/v) L-Ara for 3 h at 37°C. The co-immunoprecipitation was performed with anti-Myc antibodies (Santa Cruz).

Availability of data

All interaction data from this study can be retrieved from the IntAct database (<http://www.ebi.ac.uk/intact/>) under the following accession numbers: EBI-1190357 (*C. jejuni* data set) and EBI-1190361 (*T. pallidum* data set).

Supplementary information

Supplementary information is available at *Molecular Systems Biology* website (www.nature.com/msb).

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