Structural and molecular determinants for the interaction of ExbB from *Serratia marcescens*
and HasB, a TonB paralog.

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Running title: Interaction specificity between ExbB and TonB/HasB

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Abstract. ExbB and ExbD are cytoplasmic membrane proteins that associate with TonB to convey the energy of the proton-motive force to outer membrane receptors in Gram-negative bacteria for iron uptake. The opportunistic pathogen *Serratia marcescens (Sm)* possesses both TonB and a heme-specific TonB paralog, HasB. ExbB<sub>Sm</sub> has a long periplasmic extension absent in other bacteria such as *E. coli (Ec)*. Long ExbB’s are found in several genera of Alphaproteobacteria, most often in correlation with a hasB gene. We investigated specificity determinants of ExbB<sub>Sm</sub> and HasB. We determined the cryo-EM structures of ExbB<sub>Sm</sub> and of the ExbB-ExbD<sub>Sm</sub> complex from *S. marcescens*. ExbB<sub>Sm</sub> alone is a stable pentamer, and its complex includes two ExbD monomers. We showed that ExbB<sub>Sm</sub> extension interacts with HasB and is involved in heme acquisition and we identified key residues in the membrane domain of ExbB<sub>Sm</sub> and ExbB<sub>Ec</sub>, essential for function and likely involved in the interaction with TonB/HasB. Our results shed light on the new class of inner membrane energy machinery formed by ExbB, ExbD and HasB.
Transport of nutrients across the Gram-negative outer membrane is either a diffusion-facilitated or an active process. In the latter case, the process is powered by the energy of the proton-motive force (pmf) transmitted through the periplasm to specialized outer membrane (OM) transporters (the so-called TBDT’s, TonB dependent transporters). A complex of three cytoplasmic membrane proteins, TonB, ExbB and ExbD, that together form the TonB complex (see (1) for a review) conveys the energy of the pmf to the TBDT. In *Escherichia coli* K12, there is only one set of *tonB, exbB* and *exbD* genes, whereas there are nine TBDT’s (2), all energized by the same complex. ExbB and ExbD respectively belong to the MotA-TolQ-ExbB and MotB-TolR-ExbD protein families. Those proteins are involved in power generation and transmission in various processes (MotAB drives flagellar rotation, the ExbB-ExbD complex referred to as ExbBD energises active transport of molecules across OM receptors and TolQR is involved in cell division), at the expense of pmf dissipation across the cytoplasmic membrane (3-6). They form complexes that associate respectively with the flagellar rotor, TonB and TolA. It has also been shown that ExbD does not accumulate in the absence of ExbB, and that TonB does not accumulate in the absence of the ExbBD complex (7). The C-terminal domains of TonB and ExbD reside in the periplasm and interact with each other (8, 9). ExbD TM (Trans-Membrane segment) has one strictly conserved aspartate residue (10), which is thought to undergo cycles of protonation/deprotonation (coupled to pmf dissipation) and is essential to its function.

TBDT’s comprise a 22-stranded β-barrel anchored in the outer membrane. An N-terminal domain folded inside the barrel, and called the plug, contains the main region of interaction with TonB, the so-called TonB box, located in the N-terminal periplasm-exposed part of TBDT.
The N-terminus of TonB is localized in the cytoplasm, followed by a single transmembrane segment, and by a Pro-Lys rich region long enough to span the periplasmic space and ends with a structured C-terminal domain interacting with the TBDT TonB box. The substrate binds the extracellular side of the TBDT receptor, triggering conformational changes that are transmitted to the periplasmic side and allow the interaction between TonB and the TonB box of the TBDT, leading to the substrate entry into the periplasm by a yet unknown mechanism. A rearrangement of the plug domain has been proposed to occur creating a path for the substrate across the TBDT.

In *Serratia marcescens*, a close relative of *E. coli*, in which about twenty potential TBDT’s were identified, there are at least two TonB homologs: an ortholog of TonB$_{Ec}$ (48% aminoacid identity with TonB$_{Ec}$) and HasB, a TonB paralog that is strictly dedicated to its cognate outer membrane receptor HasR. HasB has the same topology as TonB, and its C-terminal domain interacts specifically with HasR with a circa (ca.) 40-fold higher affinity than the corresponding TonB domain. The Has (heme acquisition system) system includes the TBDT HasR, which recognizes both free heme and the high-affinity extracellular heme-binding HasA hemophore. HasB is encoded within the *has* locus and displays low sequence identity with either *E. coli* or *S. marcescens* TonB. The Has system has been functionally reconstituted in *E. coli*. Unlike TonB$_{Sm}$, HasB could not complement TonB$_{Ec}$ functions, nor was it able to drive heme entry via the HasR receptor in the presence of ExbBD$_{Ec}$. A gain of function mutation in the TM domain of HasB was however isolated in *E. coli*, restoring heme entry via HasR in the presence of ExbBD$_{Ec}$. This mutation corresponds to a 6 base-pair duplication leading to a longer TM segment for HasB by inserting AL into CLVLVLALHLLVAALLWP resulting in CLVLVLALALHLLVAALLWP.
Recently several structures of *E. coli* ExbB and ExbBD have been solved, either by X-ray crystallography or cryo-EM (14-16). All samples studied included ExbB and ExbD, but not all showed an ordered structure of ExbD. In these structures, ExbBD\textsubscript{Ec} (14) appeared as a pentamer. Each monomer of ExbB has three TM helices that extend into the cytoplasm, and with a highly polarized charge distribution on the cytoplasmic side. The central pore is apolar, lined by TM helices 2 and 3 of each monomer, creating a large hydrophobic cavity inside the structure.

Another study using X-ray crystallography, single-particle cryo-EM and electron diffraction on two-dimensional crystals concluded that ExbB could undergo a pH-dependent pentamer to hexamer transition, and that the hexameric ExbB could accommodate three ExbD TM segments in its pore (15). A more recent cryo-EM study of ExbBD reconstituted in nanodiscs however confirmed the ExbB\textsubscript{2}ExbD\textsubscript{2} stoichiometry with two TM helices of ExbD identified in the ExbB pore (16). This is consistent with previous DEER (Double Electron Electron Resonance) results (15). Similarly, *Pseudomonas savastanoi* ExbBD exhibits the same stoichiometry (17).

Structures of the flagellar MotAB motor complexes (related to ExbBD) from several bacteria (*Campylobacter jejuni, Bacillus subtilis, Clostridium sporogenes, Vibrio mimicus, Shewanella oneidensis*, PomAB from *Vibrio alginolyticus*) were recently published. They all display a MotA\textsubscript{5}MotB\textsubscript{2} stoichiometry and share some topology elements with ExbBD complex (17, 18).

The discovery of HasB and its functional specificities prompted us to identify the ExbBD complex that would function with it in *S. marcescens*. In this study, we identified the orthologous ExbBD\textsubscript{Sm} and found that this complex is active with both HasB and TonB. We characterised a new family of ExbB proteins with a long N-terminal extension whose presence is strongly correlated to the presence of a *hasB* gene ortholog in the genome. We purified ExbB\textsubscript{Sm} alone and ExbBD\textsubscript{Sm} and determined their structures by single-particle cryo-EM at 3.1 and 3.96Å resolution, respectively. We show that in both cases, ExbB\textsubscript{Sm} behaves as a stable
pentamer; in the ExbB<sub>Sm</sub> complex, we observe two TM helices of ExbB<sub>Sm</sub> in the pore of ExbB pentamer, as it was shown for the <i>E. coli</i> case. Using NMR measurements, we also show that the N-terminal periplasmic extension specific to ExbB<sub>Sm</sub> interacts with the C-terminal globular domain of HasB. Finally, via mutagenesis studies and bacterial growth assays, we show that the first transmembrane helix TM1 of ExbB contains specificity determinants for interaction with HasB/TonB and that the periplasmic extension of ExbB does play a role in heme acquisition via the Has system.

**Results**

1. Orthologs of <i>E. coli</i> <i>exbB</i> and <i>exbD</i> in <i>Serratia marcescens</i> define a new ExbB family with an N-terminal extension.

Sequence analysis of strain Db11, a fully sequenced <i>S. marcescens</i> isolate (GenBank: <i>Serratia marcescens</i> subsp. marcescens Db11, complete genome. ACCESSION HG326223, https://www.ncbi.nlm.nih.gov/nuccore/HG326223.1), identified one putative operon encoding orthologs of <i>E. coli</i> ExbB and ExbD (SMDB11_3479(ExbD) and SMDB11_3480(ExbB)). In <i>E. coli</i>, the <i>exbBD</i> operon is surrounded on the 5’ side, and in the opposite direction, by <i>metC</i> (encoding a cystathionine lyase) and on the 3’ side by <i>yhgA</i> (encoding an aldehyde reductase).

In <i>S. marcescens</i> Db11, the <i>exbB</i>-like gene is close to and in the opposite orientation to a <i>metC</i> homolog, as in <i>E. coli</i>. At the 3’ end of the operon, and in the opposite direction there are genes related to sucrose metabolism. Comparison of the coding sequences indicated that the <i>exbD</i> gene encoded a 140 residue-long protein (71% identity with the 141 residue-long <i>E. coli</i> sequence). In contrast, the putative <i>exbB</i> gene encodes a much longer protein than its <i>E. coli</i> counterpart (325 residues instead of 244 residues in <i>E. coli</i>, 73% identity in the common part).

The additional stretch of residues present in ExbB<sub>Sm</sub> is located at the N-terminus and...
corresponds to a putative signal peptide followed by a ~50 residue N-terminal extension of the mature protein (Figure 1A).

A BLAST search (19) on bacterial genomes indicated that such “extralong” ExbB’s are found in different Gammaproteobacteria, including *Serratia*, *Yersinia*, *Pseudomonas*, *Erwinia/Dickeya*, and many genera in the Alphaproteobacteria class. Table I lists representative species of Alphaproteobacteria in which such long ExbB’s are found. Interestingly, about 90% of the long ExbB-containing species listed in this table also have a *hasB* gene ortholog. An alignment of these ExbB aminoacid sequences (represented as a Logo on Figure 1B) also shows that the mature N-terminal extensions are of variable lengths between a few and 150 residues with an average of 50 residues. These extensions are quite rich in Ala (24.6%) and Pro (14.5%) residues and therefore likely to be unstructured but could be involved in protein-protein interactions as proline-rich regions often are in signalling processes (20-22).

The Conserved Domain Database (23) currently contains two ExbB subfamilies in the cl00568 MotA_ExbB superfamily, TIGR02797 (containing *E. coli* ExbB, as well as longer ExbB’s) and TIGR02805 (containing *Haemophilus influenzae* ExbB, with a very short cytoplasmic domain between TM1 and TM2). Based on sequence data, TIGR02797 can be divided into 2 groups: those with “extralong” ExbB’s in one subfamily, and *E. coli*-like sequences in the other. Along with the existence of HasB in the bacterial species, the identification of this N-terminal addition prompted us to further characterise the ExbB-ExbD complex from *S. marcescens*.

2. ExbBD<sub>Sm</sub> complements ExbBD<sub>Ec</sub> for iron acquisition and functionally associates with both TonB and HasB for heme acquisition through HasR.
As a first step in the characterization of the identified ExbBD<sub>Sm</sub>, we tested whether this complex complements ExbBD<sub>Ec</sub>. To this end, the plasmids pBADExbBD<sub>Sm</sub> and pBADExbBD<sub>Ec</sub> were constructed, introducing the two genes from <i>S. marcescens</i> or <i>E. coli</i>, into pBAD24 vector under the control of arabinose-inducible <i>P<sub>arab</sub></i> promotor. By using growth under iron starvation conditions as a test, we could show that <i>exbBD</i><sub>Sm</sub> complements <i>exbBD</i><sub>Ec</sub>. The <i>E. coli C600</i> ∆<i>exbBD</i> strain is indeed more sensitive than its wild-type counterpart to iron starvation (induced for example by the Fe<sup>2+</sup> chelator dipyridyl, DiP) as ExbBD are required to transduce pmf to TonB for siderophore uptake. Its growth in the presence of DiP is restored by the expression of either pBADExbBD<sub>Ec</sub> or pBADExbBD<sub>Sm</sub> (Figure 2).

We then tested the functionality of ExbBDSm in heme acquisition via the Has system reconstituted in a heme auxotroph <i>E. coli</i> strain. To avoid interference from the chromosomal <i>exbBD</i> operon and from <i>tonB</i>, we used the C600∆<i>hemA</i>∆<i>exbBD</i> and the C600∆<i>hemA</i>∆<i>exbBD</i>∆<i>tonB</i> strains, transformed with various recombinant plasmid pairs, one bringing the complete Has locus, with or without the <i>hasB</i> gene, the other one with the <i>exbBD</i> operon or its derivatives under the control of the <i>P<sub>arab</sub></i> promotor. In this kind of experiment, the bacterial growth reflects both the expression of the various Has components under the control of their own regulatory elements (24), and the efficiency of the heme uptake process itself. Two kinds of tests were carried out, one on solid substrate in Petri dishes, allowing to see haloes of bacterial growth around wells punched in the agar and containing the heme source, the other one in liquid medium in microplates with an absorbance plate reader, allowing to record growth at regular intervals over extended periods. The latter type of tests allows a more precise and quantitative description of the phenotypes as it is realized in a homogeneous medium, which is not the case with Petri dishes. Six strains were constructed: one set deleted from TonB C600∆<i>hemA</i>∆<i>exbBD</i>∆<i>tonB</i> (pAMHasISRADEB + pBAD24),
C600ΔhemAΔexbBDΔtonB (pAMHasISRADEB + pBADExbBD<sub>Ec</sub>),
C600ΔhemAΔexbBDΔtonB (pAMHasISRADEB + pBADExbBD<sub>Sm</sub>), and one set with TonB
C600ΔhemAΔexbBD (pAMHasISRADE + pBAD24), C600ΔhemAΔexbBD (pAMHasISRADE
+ pBADExbBD<sub>Ec</sub>), and C600ΔhemAΔexbBD (pAMHasISRADE + pBADExbBD<sub>Sm</sub>). The
ExbBD constructs for Ec and Sm are schematized on the first two lines of Figure 3A. The
results are reported in Figures 3B (Petri dishes, overnight observation), 3C (Petri dishes, 36h
observation) and 3D (microplates, recording over 66h). As expected, control strains (with
pBAD24) did not grow. ExbBD<sub>Sm</sub> was functional with both HasB and TonB (middle series of
holes in Figure 3C indicated with orange and blue dots, and orange and blue dots curves in
Figure 3D), although with quite dramatically different kinetics and yield, the onset of growth
occurring at 5-6hrs for the HasB-ExbBD<sub>Sm</sub> pair, and at 22-23hrs for the TonB-ExbBD<sub>Sm</sub> pair. Moreover, the final OD of TonB-ExbBD<sub>Sm</sub> is half that of HasB-ExbBD<sub>Sm</sub>. Under our
experimental setup, ExbBD<sub>Ec</sub> was also functional with both TonB and more surprisingly with
HasB (bottom series of holes in Figure 3C, and green and grey dots curves in Figure 3D), the
onset of growth occurred at around 15-16hrs for TonB, and 18hrs for HasB; however, the
maximal OD was lower for the HasB-ExbBD<sub>Ec</sub> pair (0.43 vs. 0.72). Finally, the ExbBD<sub>Sm</sub>-
HasB pair seems less sensitive to iron starvation than all the other ones, since similar results
are obtained at 100 and 200µM Dipyridyl (Figure 3B), which is not the case for the other pairs,
as already observed in the TonB-ExbBD<sub>Ec</sub> case (25). These results might seem at odds with the
previous results obtained on HasB (12) which showed that HasB was nonfunctional with
ExbBD<sub>Ec</sub>. However, the experimental setup was quite different, both in terms of plasmids used,
strains and conditions of observation. Therefore, ExbBD<sub>Sm</sub> is the <i>E. coli</i> ExbBD ortholog, able
to associate both with HasB and TonB<sub>Ec</sub>.
3. Characterization of ExbB<sub>Sm</sub> and ExbBD<sub>Sm</sub>: Specific function of the N-terminal extension of ExbB<sub>Sm</sub>

3.a Purification and mass-spectrometry analyses of protein and associated lipids

To gain further insight into the possible differences between ExbBD<sub>Sm</sub> and ExbBD<sub>Ec</sub>, we purified both ExbB<sub>Sm</sub> and ExbBD<sub>Sm</sub> (as C-terminal 6-His-tagged proteins on ExbB for ExbB purification, and C-terminal 6-His tag on ExbD for ExbBD purification, see Materials and Methods section) in LMNG (lauryl maltose neopentyl glycol) micelles (Figure 4, A and B). Both ExbB<sub>Sm</sub> and ExbBD<sub>Sm</sub> appear as homogeneous oligomers at the last size exclusion chromatography (SEC) step.

Mass spectrometry analysis showed that purified ExbD<sub>Sm</sub>His6 has a mass of 16 131 Da close to its theoretical mass of 16261.7 Da. Purified ExbB<sub>Sm</sub> has a measured mass of 29 557 Da and its predicted mature sequence has a theoretical mass of 29556.8 Da. Together with the determination of the first amino-acid residue for ExbB, this confirmed that the predicted signal sequence of ExbB<sub>Sm</sub> is absent from the mature sequence and has therefore been processed. It also showed that ExbD initial Met residue was cleaved off (see Supplementary Figure S1).

More interestingly, mass spectrometry analysis of chloroform/methanol extracts of the purified proteins also showed that both complexes contained native lipids (Supplementary Figure S2), mostly phosphatidylglycerol (PG) and phosphatidylethanolamine (PE), and that ExbBD also contained some cardiolipin (CL) (Supplementary Figure S3). Further analysis of aliphatic chain composition of lipids shows evidence of specific composition with a majority of PG with 34 carbons and 2 unsaturations (Supplementary Figure S4). Considering that the ExbBD<sub>Sm</sub> complex is functional in <i>E. coli</i>, and that the phospholipid composition of <i>S. marcescens</i> is quite similar to that of <i>E. coli</i> (26), this affinity for lipids may disclose a specific recognition that may be important for the function.
3b NMR shows an interaction between HasB<sub>CTD</sub> and ExbB<sub>Sm</sub> periplasmic extension.

The presence in ExbB<sub>Sm</sub> of an N-terminal extension residing in the periplasm led us to investigate its function. This extension contains more than 50% of Ala and Pro residues and is predicted to be unstructured by disorder prediction servers such as IUPRED2 (27). Indeed, NMR analysis of a 44 residue-long synthetic peptide corresponding to this extension (ExbB<sub>Sm</sub>1-44) did not show any indication of secondary structure (our unpublished results). Since this region is predicted to be in the periplasm where the C-terminal domain of HasB, HasB<sub>CTD</sub> is also located, we investigated the potential interaction of the ExbB<sub>Sm</sub>1-44 synthetic peptide with HasB<sub>CTD</sub> by NMR. The analysis of chemical shift perturbations (CSP) of amide resonances of HasB<sub>CTD</sub> upon addition of the peptide, showed that the chemical environment of their corresponding residues was modified because of their interaction (Figure 5 A and Supplementary Figure S5). Perturbed residues are mainly located on the helical face of HasB<sub>CTD</sub> forming a continuous surface of interaction (Figure 5B and 5C) (R175, R178-K180, K184, Q192, T200, L201, Q204, H206, A232, A240, G246). Interestingly, this face is on the opposite side of the third beta strand of HasB that was previously shown to interact with HasR (28). In addition, the residues of a small pocket at the C-terminus of HasB (D255, R259) show also high CSP and might be involved in the interaction with ExbB or subject to a conformation change induced by this interaction.

3c An ExbB<sub>Sm</sub> mutant devoid of the N-terminal extension is affected in its function.

To assess the potential function of the N-terminal periplasmic extension of ExbB<sub>Sm</sub>, we engineered a ExbB<sub>Smdelextss</sub> mutant lacking this extension (Figure 3A) (see Materials and Methods for details) and tested its effect on the growth of a C600ΔhemAΔexbBDΔtonB strain harbouring plasmids pAMHasISRADEB and pBADexbBD<sub>Sm</sub> or its derivatives in liquid
culture. We observed that the lag between the start of the experiment and the onset of growth is at least 2 hours longer for the ExbB_sm deletes mutant strain, as compared to the WT strain (Figure 6A, compare orange and yellow curves) and a control strain (medium blue). In addition, the maximum OD reached after growth is slightly but significantly decreased (Figure 6A). To ensure that this difference was not due to a difference of expression of the two constructs, we compared their amounts in membrane preparations by immunodetection. As our anti-ExbB_sm antibody was not sensitive enough for this measurement, we used the His-tagged version of ExbB_sm readily detectable with anti-His6 antibodies. Coomassie blue stained SDS-PAGE showed that the 3 conditions have similar total amounts of protein (Figure 6B). The Western blot shows that the amount of the ExbB_smHis6 variant deleted of its N-terminal extension is at least equal, if not slightly higher, to the amount of the wild-type protein (Figure 6C, lanes 2 and 3), ruling out a possible decrease in protein concentration. The N-terminal periplasmic extension of ExbB_sm is therefore likely involved in the functioning of the Has system, whether at the transcription activation of the Has locus, and/or at any later step.
**4 ExbB_{Sm} and ExbBD_{Sm} structural analysis by cryo-EM** show ExbB₂ and ExbB₃D₂ stoichiometries.

**4a ExbB_{Sm} structure**

As ExbB_{Sm} presented specific features compared to ExbB_{Eco}, we set out to determine its structure by single-particle cryo-EM, see Material and Methods for details. The 3D class average model clearly showed a pentameric structure. It was refined and polished to obtain a resolution of 3.1Å using the Fourier shell correlation (FSC) gold-standard criterion at 0.143. Maps were refined with and without C5 symmetry and showed a 98% correlation in density. Therefore, we chose to use the map based on C5 symmetry for model building.

The structure solved here and shown in Figure 7 A (side view) and B (cytoplasm view), has the same α-helix bundle topology and is very similar to that of *E. coli* ExbB 5SV0 structure (that was co-purified with ExbD but did not show any ordered density for ExbD (14)), with a 1.3Å root mean square deviation (rmsd) over all Cα atoms (Table II). This shows that ExbB_{Sm} is stable as a pentamer on its own. The periplasmic N-terminal extension did not yield any visible density, precluding its structure determination. At the C-terminus of each monomer, however, density was clearly defined, allowing structure determination for an additional 10 residues in a helical conformation up to the last one (helix α8 finishing with Gly 283, see Figure 7C) before the His-tag that is present but disordered. In the 5SV0 X-ray structure, a calcium ion, present in the crystallisation solution, is bound to the five Glu 106 (of helix α2 in the TM region) on the cytoplasmic side. In ExbB_{Sm}, this residue is replaced by an Asn, and we do not observe any density that could be assigned to a metal ion.

The structure of the whole pentamer with one monomer coloured as a function of sequence conservation in this sub-class of ExbB (Supplementary Figure S6) shows that the highest
residue conservation is inside the transmembrane channel, indicative of high functional constraints, while the TM residues located at the membrane surface are more variable. The same observation was made for ExbB sequences lacking N-terminal signal sequence and periplasmic extension (14).

The cryo-EM density of ExbB_{Sm} shows vents located at the interface of adjacent monomers and at the height of the cytoplasmic junction with the inner membrane leaflet (Figure 7A). These vents may allow solvent or ion passage. Additional density was also clearly observed on the external surface, and we attributed it to the phospholipids present in the preparation. Each ExbB monomer appears to be associated with the equivalent of two PG molecules, located at the inner leaflet of the cytoplasmic membrane (Figure 7A, D and E). However, we were able to model only one PG molecule per monomer with confidence. Three PE (phosphatidylethanolamine, the major *E. coli* phospholipid) molecules and one PG molecule were identified as associated to the ExbBD pentamer in the *E. coli* complex after reconstitution in nanodiscs (16). The cryoEM density map, when displayed at a level that shows the detergent belt, shows density inside the ExbB membrane pore (supplementary figure S7A). This density is too noisy to allow model building (supplementary figure S7B). However, it corresponds to a region with positive electrostatic charge on the top and bottom of the pore and neutral or hydrophobic in the middle (supplementary figure S7C). This density could be due to the presence of lipid or detergent. Interestingly, it is located at a different height as compared to the detergent belt and external lipids. In summary, our cryo-EM structure shows that ExbB_{Sm} is stable as a pentamer and associates strongly with specific phospholipids coming from the inner leaflet of the membrane.

**4b ExbBD_{Sm} structure**

We also solved the structure of the ExbBD_{Sm} complex by cryo-EM (Figure 8 and Table II). As for ExbB, the purified complex exhibited a symmetric peak on SEC (see Figure 4, B). Cryo-
EM data were collected and processed as described in the Material and Methods section. In contrast to ExbB<sub>Sm</sub>, 2D classes showed preferential orientations with 92% top views and only 8% side views. Due to this strong bias in particle orientation, the resolution achieved was 3.96 Å, precluding reliable positioning of the side chains. However, our ExbB<sub>Sm</sub> model fitted readily into the density. As observed for ExbB<sub>Sm</sub> alone, in the ExbBD<sub>Sm</sub> complex, ExbB<sub>Sm</sub> behaves as a pentamer, and no sign of other assemblies was found. Similarly, no density could be attributed to the periplasmic extension. Two clear densities inside the ExbB<sub>Sm</sub> pore were assigned to the pair of ExbD<sub>Sm</sub> TM segments (Figure 8 B and C). The charge distribution is highly polarised on the cytoplasmic side: the region close to the membrane is positively charged and the distal part is negatively charged as observed for ExbB<sub>Ec</sub> (Supplementary Figure S8A). The central pore of ExbB<sub>Sm</sub> is apolar, lined by TM helices 2 and 3 of each monomer, creating a large hydrophobic cavity inside the structure (Supplementary Figure S8B). The ExbD helices were initially oriented similarly to ExbD<sub>Ec</sub> in structure 6TYI and remained stable during refinement.

Interestingly, the ExbD TM helices are at the same height as the density observed in the pore of ExbB alone, and closer to the periplasm than the membrane bilayer. In the refined model, Asp 25 from ExbD<sub>Sm</sub> monomer chain G faces Thr 218 from ExbB<sub>Sm</sub> chain C, while Asp 25 from the ExbD<sub>Sm</sub> chain F faces the interface between two ExbB<sub>Sm</sub> monomers A and E (Supplementary Figure S8C and D). The estimation of pKa, by using the program Propka (29) server, shows that Asp 25 has a pKa of 7.3 for chain F and 7.4 for chain G, both very peculiar for solvent-accessible acidic residues but that can be found in buried active sites or membrane proteins (30). This pKa should allow protonation and deprotonation of ExbD Asp 25 at physiological pH.

As compared to the structure of ExbB<sub>Sm</sub>, there is a slight «opening» towards the periplasmic side (the distance of Ala 197 from one subunit to Leu 204 of the facing subunit varying from
25.5Å in ExbB<sub>Sm</sub> to 29.8Å in ExbBD<sub>Sm</sub>, much greater than any error expected due to the difference in resolution of the two structures). This opening of the structure is limited to the periplasmic part (Supplementary Figure S9). The rmsd between ExbB<sub>Sm</sub> alone and in complex with ExbD<sub>Sm</sub> is 2Å over all Cα of the pentamer, and the rmsd between ExbB<sub>Sm</sub> in complex with ExbD<sub>Sm</sub> and ExbB<sub>Ec</sub> in complex with ExbD<sub>Ec</sub> (structure 6TYI chains A-E) is 1.7Å.

As compared to the 6TYI structure of <i>E. coli</i> ExbBD complex, we observe significant differences that may be related to the specificity of function of ExbB<sub>Sm</sub>. Both ExbB<sub>Sm</sub> and ExbBD<sub>Sm</sub> inner pores are slightly wider than their <i>E. coli</i> counterparts at the periplasmic entrance. Consequently, two channels cross the membrane region, (extending from the periplasmic entrance to the Asp 25 residue from the ExbD TM deeply embedded in the ExbB pore) that are clearly seen in the ExbBD<sub>sm</sub> (as detected by the MoleOnline (31) Server, Figure 9, A and C; the average diameter is around 3Å) while the <i>E. coli</i> structure exhibits a single, much thinner channel as detected with the same parameters of MoleOnline (Figure 9, B and D; average diameter around 2Å). It is therefore possible that this structure represents another state of the ExbBD motor complex, whereby the periplasmic side allows solvent access to the Asp 25 residue of ExbD TM. The different physico-chemical conditions (nanodiscs vs. detergent micelles, 200 vs. 100mM NaCl, pH7.4 vs. pH8) might also influence such parameters.

In summary, we show that ExbBD from <i>S. marcescens</i> has a 5:2 stoichiometry with a larger channel allowing solvent or proton transport from the periplasmic side as compared with the <i>E. coli</i> structure.

5. Swapping residues in TM1 between ExbB<sub>Sm</sub> and ExbB<sub>Ec</sub> strongly suggests that ExbB TM1 interacts with HasB/TonB.
As already mentioned, TBDT function requires a productive association between TonB or its orthologs/paralogs with ExbBD complexes. It is known that the periplasmic domains of ExbD and TonB interact, and there are both genetic and structural pieces of evidence for the interaction between the TM helix of TonB and the first TM helix of ExbB. Our experimental system gave us the opportunity to investigate the specificity determinants between HasB/TonB and ExbB. Previous work (12) lends support to the absence of interaction between HasB and ExbBD Ec. However, our data rather favor a less efficient interaction between HasB and ExbBD Ec than with ExbBD Sm. E. coli can grow with the HasB-ExbBD Ec pair, although more slowly and with a lower yield: the final OD was 0.78 for HasB-ExbBD Sm and 0.43 for HasB-ExbBD Ec (Figure 3 D orange and grey curves). As it is known that ExbBD is required to stabilize TonB, we first tested whether HasB was stabilized by ExbBD. We could show that, although HasB does not accumulate in the absence of ExbBD, it is readily present in comparable amounts with either ExbBD Sm or ExbBD Ec (Figure 10). This observation therefore indicates that most likely HasB also interacts with ExbBD Ec so as to be stabilized (i. e., withstanding proteolytic degradation), but in a less functional manner than with ExbBD Sm.

A general framework put forward for the Mot complexes indicates that MotA (equivalent of ExbB) rotation around MotB (equivalent of ExbD) drives the rotation of the flagellum basal ring (17, 18). We tested the possibility that ExbB residues exposed at the surface of the protein might be involved in a functional interaction with HasB/TonB, as has been proposed in the case of the TonB-ExbBD complex from P. savastanoi (17). The superposition of residue conservation on the structure of ExbB showed that the membrane-facing residues of the TM region of helix α2 and the cytoplasmic residues were the least conserved (Figures 1 and S6). We noted one conspicuous stretch of residues located in the cytoplasmic leaflet of TM helix 1 (see Supplementary Figure S6):
residues 76–88 of ExbB_{Sm} TILFAKGSELLRA corresponding to residues 39–51 of ExbB_{Ec} AIFFSKSVEFFNQ

(identical residues are underlined) that is the least conserved one between the two proteins. In this stretch, the conserved I, K and E residues point toward the innermost part of the protein and the non-conserved residues point to the outside and are available for protein-protein interaction (Figure 11 E and F).

We speculated that this region of ExbB might establish interaction with HasB. We made a chimeric ExbB_{Ec} protein, with its 39-51 region replaced by residues 76-88 from ExbB_{Sm}, named ExbB_{Ec-Sm76-88}, and asked how this chimeric ExbB_{Ec} protein would allow growth in the presence of HasB (see Figure 3A for the constructions), as compared to ExbB_{Ec}. As shown in Figure 6A, this chimeric protein ExbB_{Ec-Sm76-88} (green dots) was more active with HasB than ExbB_{Ec} (grey), the onset of growth took 14hrs compared to 18hrs for ExbBD_{Ec} and the maximum OD was 0.74 compared to 0.43 for ExbBD_{Ec}. We also made the inverse change where the 76-88 region of ExbB_{Sm} was replaced by the 39-51 region from ExbB_{Ec} (ExbB_{Sm-Ec39-51}, magenta). This mutant has a degraded behaviour as compared to ExbB_{Sm}, since the onset of growth occurs at 19hrs compared to 6hrs, and the final OD is 0.47, compared to 0.78. To better locate the region responsible for specificity, we produced one additional mutant where only residues 76-84 (TILFAKGSE) from ExbB_{Sm} were exchanged for residues 39-47 (AIFFSKSVE) from ExbB_{Ec} yielding ExbB_{Ec-Sm76-84} (see Figure 3A). ExbB_{Ec-Sm76-84} was quite comparable to ExbB_{Ec-Sm76-88} (see deep-blue and green curves on Figure 6A). This 9-residue stretch is therefore sufficient to alter ExbB_{Ec} to be better adapted to HasB. This set of experiments shows that the intramembrane functional zones of ExbB are crucial for the growth of E. coli in our experimental set up and therefore likely govern the interaction between ExbB and HasB.
In summary, we showed using NMR and culture growth assays that ExbB\textsubscript{Sm} N-terminal extension (section 3b) and a few residues in the \(\alpha2\) helix TM region (section 5) are likely involved in specific interactions with HasB.

Discussion

1) ExbBD\textsubscript{Sm} is a 5:2 complex at pH 8.

Our structures were solved at pH 8, and examination of the 2D class averages only showed pentameric structures and not hexameric ones, contrarily to what has been observed in the Maki-Yonekura study, where ExbB was mostly seen as a hexamer in the high pH regime (15). In their study, the solubilizing detergent was DDM, that was subsequently exchanged for either C8E4 or C10E5. These shorter chain detergents might have led to some destabilization of the pentameric structure and provoked the conversion to the hexameric form. Our data agree with those published by Lloubes, Buchanan et al. (14, 16). In vivo, a stoichiometry of 7ExbB, 2ExbD and 1TonB has been determined in \textit{E. coli} (32). Although one cannot completely rule out some bias in this indirect measurement, a physiologically relevant reason for an excess of ExbB would be to provide a permanent scaffold to which TonB and/or ExbD could associate with, upon TonB complex dissociation, that might occur during the catalytic cycle, as has already been proposed (33).

2) ExbB and ExbBD are co-purified with endogenous lipids.

In their last study, Celia \textit{et al.} reconstituted ExbBD from \textit{E. coli} into nanodiscs with added phospholipids from \textit{E. coli}, and found mostly PE bound to ExbBD (16). It is unclear whether this phospholipid was a genuine tightly bound, co-purified lipid or whether it came from the lipids added during reconstitution in the nanodiscs. In our case, we did not add any lipid and our mass spectrometry identification of co-purified lipids found mostly PG and some PE in the
ExbB case (and CL for ExbBD). The cryo-EM structure shows that they are located in the inner
leaflet, as in the *E. coli* complex. PG and CL are negatively charged, whereas PE is zwitterionic.
This might be pertaining to the functioning of the complex, as the lipids are bound to a highly
positively charged interface, very close to the fenestrations seen in the structure and PE might
therefore be less tightly bound than PG or CL. Further analysis by mass spectrometry of the
aliphatic chains of the lipids present in our sample also revealed a specific composition in their
length and unsaturated nature.

One point also worth mentioning is the existence of “channels” inside the ExbBD tunnel,
potentially allowing the passage of proton/hydronium ion up to the Asp 25 residue of ExbD,
deeply embedded in the apolar medium of ExbB, and therefore with a pKa close to
physiological pH, allowing easy protonation/deprotonation steps of Asp 25 side chain. The
identified channels connect the cytoplasm to the periplasm via Asp 25, allowing us to propose
a possible trajectory for the proton transfer via those channels.

3) Motor model proposal

Regarding the coupling of the pmf dissipation with the mechanical work carried out by
MotAB/TolQR/ExbBD complexes, several models have been put forward that might be more
or less easy to reconcile with the structural data. A detailed mechanism was proposed for *C.
jejuni* MotAB complex, where cycles of protonation/deprotonation of the conserved Asp of the
MotB TM (equivalent to the conserved Asp 25 of the ExbD TM) are coupled to the rotation in
discrete steps of the MotA pentamer around MotB axis, provided that the periplasmic domain
of MotB is anchored to the peptidoglycan layer via its peptidoglycan binding site (18). Our
ExbBD structure indeed shows that ExbD monomers have two orientations relative to ExbB:
Asp 25 from one monomer faces Thr 208 of ExbB and Asp 25 from the second monomer faces
a hydrophobic region at the interface of two ExbB monomers. In the first state previously
protonated Asp25 could be deprotonated in contact with polar Thr208 while the second, deprotonated Asp 25, could pick a proton from the periplasm via the channel. Rotation of the ExbB pentamer around the two ExbD chains would then lead to a new protonation/deprotonation cycles accompanied by a change in the environment upon rotation. How is this rotative energy conveyed to the TBDT?

The binding of an iron-loaded substrate on the extracellular binding site of a TBDT triggers a reaction cascade ultimately leading to the substrate entry into the periplasm. The inner membrane TonB(HasB) complex conveys the energy of the pmf to the TBDT most likely by a rearrangement of the plug inside the barrel to allow substrate access to the periplasm and its capture by periplasmic binding proteins. Specific interactions between the TonB(HasB) box of the receptor and the C-terminal domain of TonB(HasB) are essential in this process (28). A wealth of data has also accumulated documenting specific interactions between TonB<sub>CTD</sub> and ExbD<sub>CTD</sub>, depending upon the energy state of the cell. It is also known that ExbD and TonB interact via their periplasmic parts (34) and in particular residue 150 of TonB (positioned just upstream the C-terminal globular domain) can make cross-links to the C-terminal domain of ExbD (35). There have also been indications and suggestions that both TonB<sub>CTD</sub> and ExbD<sub>CTD</sub> could interact with the peptidoglycan sacculus, providing anchor points to allow force transmission. Molecular modelling works hypothesized that the C-terminal domain of TonB (36) and the C-terminal domain of ExbD (37) have specific binding sites for the peptidoglycan network. Atomic force microscopy experiments also showed that by exerting a pulling force on the C-terminal domain of TonB bound to the BtuB TBDT, a partial unfolding of the TBDT plug occurs, potentially leading to the entry of the substrate in the periplasm (38). Our NMR results also show that ExbB interacts via its alanine and proline-rich periplasmic N-terminal region with the HasB periplasmic domain on the side opposite to that of the TonB box interaction, thus...
rendering a tripartite interaction possible. Our in vivo growth results with the ExbB<sub>Sm</sub>delextss mutant devoid of the periplasmic extension point to a possible role of the N-terminal periplasmic extension in the activation of the transcription of the Has locus (figure 6).

The MotA-MotB model also posits that MotA outer region interacts with the rotor of the flagellum and thus that MotA rotation drives the rotation of the flagellum. Similarly, the MotAB rotating model can be extrapolated to the ExbBD complex. In this model, ExbB would rotate around ExbD, driving the rotation of TonB/HasB thanks to the specific interaction between TonB/HasB TM and ExbB TM α2. This kind of interaction is supported by our mutagenesis data, as we could strongly increase the efficiency of ExbB<sub>Ec</sub> with HasB by exchanging a short stretch of residues between ExbB<sub>Sm</sub> and ExbB<sub>Ec</sub>. It is also in line with a previous TonB TM point mutant partially suppressed by an ExbB α2 point mutant (39). Structure comparison of the swapped regions shows that ExbB<sub>Sm</sub> has smaller residues that may be better accommodated and less specific than the E. coli sequence (Figure 11, compare structures on figures 11C and 11D and helical wheels on figures 11E and 11F). Similar interactions have been proposed by Deme et al. on the TonBExbBD complex from P. savastanoi (17), where an extra cryo-EM density is seen outside the ExbB pentamer and was tentatively assigned to the TM domain of TonB. The residues from ExbB<sub>Ec</sub>TM1 (S34 and A39) identified as co-evolving with TonB TM region (18) are only a little deeper in the bilayer than the residues we exchanged between ExbB<sub>Ec</sub>TM1 (39-51) and ExbB<sub>Sm</sub>TM1, that are closer to the cytoplasm. More specifically, whereas S34 is conserved between ExbB<sub>Ec</sub> and ExbB<sub>Sm</sub> (S71), A39 is replaced by T76. Moreover, the hydrophobic core of HasB TM is likely shorter than that of TonB as seen in the sequence alignment of HasB and TonB TM domains:
This difference in length might influence the orientation of HasBTM inside the bilayer and potentially explain the gain of function of HasB6 mutant (in which the hydrophobic core has two more residues (39)) to be better suited to ExbBEc than HasB. The presence of a proline residue in the TonB TM may also change its shape and the interaction with ExbB.

Several models have been proposed to account for the functioning of the TonB complex, in conjunction with the entry of a substrate bound to the extracellular side of a given receptor. In a first model, piston-like movements of TonB drive the unfolding of the plug inside the receptor barrel. *In vivo* proteolysis studies indicated that the C-terminal domain of TonB can change conformations during the catalytic cycle of protonation and deprotonation of the conserved Asp residue in ExbD TM (40). In addition, the periplasmic linker between the TM helix and the C-terminal domain of TonB is rich in Pro and Lys residues and might adopt an extended conformation of a sufficient length to span the periplasm (41). We therefore propose a model in which the force generated through the rotation of HasB/TonB, driven by ExbB rotation around ExbD, is not directly transmitted to the TonB box of the receptor, but could be mediated by the C-terminal domain of ExbD, that might act as an anchor point allowing force transmission and converting the rotation into a pulling force exerted on the TonB/HasB box of the receptor. Further studies are needed to test this model, in particular concerning protein-peptidoglycan interactions, the force needed to trigger TBDT plug opening, and how to distinguish between a rotation and a piston movement. Finally, given the wide range of lag periods we observe in our growth curves using different combinations of ExbB TM1 mutants...
and TonB/HasB, one may hypothesize that the membrane interaction between ExbB and TonB/HasB influences the rate of transcription activation of the Has locus. It is likely that the in vivo entry of a siderophore is a rather slow process: during a cell division time a bacterial cell must take up ca. 300000 iron atoms from its surroundings. Under full induction, there are roughly 10-15000 FepA siderophore receptors per cell and 1500 TonB complexes (32), meaning that each receptor has to undergoes 20 cycles during a generation estimated to 30 minutes, leading to a turnover time of the TonB complex of about 5-10 seconds. As compared to flagellar motor that can operate at extremely high speeds (several hundred of rotations per second), even though the basic mechanisms are likely to be conserved with the ExbBD/TonB-HasB complex, it is much slower, which likely points to different coupling mechanisms.

Material and Methods

Strains and plasmid construction

Strains, plasmids, and oligonucleotides used are shown Table III.

-Plasmid pBADExbBD_{Sm} was constructed after amplification on the genomic DNA from strain S. marcescens Db11 of a ca. 1.42kb fragment using primers ExbBDSm5’ and ExbBDSm3’. The PCR product was purified digested with EcoRI and SphI, and ligated with pBAD24 digested with EcoRI and SphI. Correct clones were selected after sequencing of the insert.

-Plasmid pBADExbBD_{Sm}His6 (encoding ExbB_{Sm} and a C-terminally His-tagged version of Exb{D}_{Sm}) was constructed by first amplifying on pBADExbBD_{Sm} a ca. 0.4kb fragment with the following oligonucleotides SphIHisCtexbDSm and BglIIexbDSm; after amplification, the fragment was purified, digested with BglII and SphI, and ligated with pBADExbBD_{Sm} digested
with the same enzymes. The correct clones were selected after sequencing of the insert. In
biological tests, this plasmid was indistinguishable from its parent plasmid pBADExbBDSm.
-Plasmid pBADExbBD_{Ec} was constructed by amplification on genomic MG1655 DNA of a ca.
1.7kb fragment with the following oligonucleotides ExbBD5c, and ExbBD3c; the PCR product
was purified, digested with EcoRI and SphI, and ligated with pBAD24 digested with the same
enzymes. Correct clones were selected after sequencing.
-Plasmid pBADExbB_{Sm}His6 (encoding a C-terminally His-tagged version of ExbB_{Sm}) was
constructed as follows: a PCR fragment was amplified on pBADExbBDSm with the oligos
PBADFOR and ExbBHis6, digested with EcoRI and SphI, and ligated with pBAD24 digested
with the same enzymes. Correct clones were selected by sequencing.
Plasmid pBADExbBD_{Ec-Sm76-88} (encoding a chimeric ExbB_{Ec} protein with its 39-51 residues
exchanged for the 76-88 residues from ExbB_{Sm} and ExbD_{Ec}) was constructed as follows:
Plasmid pBADExbBD_{Ec} was amplified with the two following 5’ phosphorylated
oligonucleotides: ExbBEcSm76-88.1 and ExbBEcSm76-88.2. After digestion with DpnI and
self-ligation, correct clones were selected by sequencing and the mutated fragments recloned
in pBAD24.
Plasmid pBADExbBD_{Ec-Sm76-84} (encoding a chimeric ExbB_{Ec} protein with its 39-47 residues
exchanged for the 76-84 residues from ExbB_{Sm} and ExbD_{Ec}) was constructed in the same
manner, with the following oligonucleotides pairs: ExbBEcSm 76-84.1 and ExbBEcSm 76-
84.2.
In the same manner, plasmid pBADExbBD_{Sm-Ec39-51} (encoding a chimeric ExbB_{Sm} protein with
its 76-88 residues exchanged for the residues 39-51 from ExbB_{Ec} and ExbD_{Sm}) was constructed
by PCR on the pBADExbBD_{Sm} plasmid with the following couple of 5’ phosphorylated
oligonucleotides: ExbBSmEc39-51.1 and ExbBSmEc39-51.2.
Plasmid pBADExbBD_{Smdelextss} (encoding ExbB_{Sm} deleted of its periplasmic extension and ExbD_{Sm}) was constructed similarly, by using as a template pBADExbBDSm, with the two phosphorylated oligonucleotides ExbBdelextss1 and ExbBdelextss2.

To construct pBADExbB_{Smdelextss}His6, a ca. 0.6kb *Eco*RI-*Kpn*I fragment from pBADExbBD_{Smdelextss} was exchanged for the corresponding fragment of pBADExbBHis6.

Plasmid pAMHasISRADEB (encoding HasI, HasS, HasR, HasA, HasD, HasE and HasB), was constructed by digesting pAMHasISRADE (encoding HasI, HasS, HasR, HasA, HasD and HasE) and pSYC7 (encoding HasD, HasE and HasB) (42) by *Kpn*I and *Hind*III, and purifying fragments of respectively ca. 9 and 7 kbases, ligating them together to obtain a plasmid with the whole *has* locus on a low copy number plasmid (pAM238) under the control of its endogenous regulation signals (a fur box upstream of *hasI* and *hasR*, the *hasI* and *hasS* genes respectively encoding the has specific sigma and anti-sigma factors, and the *hasS* box upstream of *hasS* and *hasR*).

All constructions were carried out in *E. coli* strain XL1-Blue.

Strains and other plasmids used in this study are shown in Table III and are from the laboratory collection.

**Protein expression and purification**

The BL21DE3(pExbBD_{Sm}His6/pBAD24) or BL21DE3(pExbB_{Sm}His6/pBAD24) were grown at 37°C either in TB or MDM medium, and induced with 0.2% arabinose at 1.5-2OD600nm (TB) or 5-6OD600nm (MDM) and the incubation continued for 3hrs. The cells were harvested by centrifugation (10000g 20minutes 4°C), washed once in 20mMTris-HCl pH8.0, flash-frozen in liquid N\_2 and kept at -80°C. Cells were broken in a Cell disruptor (Constant, UK) at 1kbar (10g of cells in 40ml final of 20M Tris pH 8.0 containing protease inhibitor (Roche, EDTA...
Benzonase was added and after ca. 15 minutes, the solution was centrifuged for 1 hr at 100000g at 4°C. The pellet (crude membrane preparation) was resuspended in 20mM Tris pH 8.0 plus protease inhibitor cocktail (Roche EDTA free), flash-frozen in liquid N₂ and kept at -80°C.

The crude membrane preparation was solubilized in 20mM Tris pH 8.0, 20mM Imidazole, 100mM NaCl, 10% Glycerol, 0.8% LMNG (10g of equivalent whole-cell pellet solubilized in 40ml), plus protease inhibitor cocktail (Roche EDTA-free) for 30 minutes at 15°C. After centrifugation (1 hr 100000xg), the supernatant was incubated with 2.5ml Ni-Agarose beads (Thermo-Fisher His-Pure Ni-NTA #88222) preequilibrated with the same buffer except for the detergent concentration (0.0015% LMNG). After three hours of incubation on a rotating wheel at 4°C, the beads were washed three times with 25ml of pre-equilibration buffer and then eluted with two times 25ml of the pre-equilibration buffer containing 200mM Imidazole. The eluate was concentrated and washed on 100kDa cut-off centrifugal device, in pre-equilibration buffer without NaCl and Imidazole. The resulting sample was loaded on a monoQ HR10-100 column equilibrated with 20mM Tris pH 8.0, 10% glycerol, 0.0015%LMNG and eluted with a gradient from 0 to 1M NaCl in the same buffer. Peak fractions were collected and concentrated as before. The concentrated sample was then loaded on a Superose 6 increase column equilibrated with 20mM Tris pH8.0, 100mM NaCl, 0.0015%LMNG. The peaks fractions were collected, concentrated and their concentration determined by using the theoretical absorption coefficient of either ExbB5D2 (113790M⁻¹cm⁻¹), or ExbB5 (104850M⁻¹cm⁻¹). They were then kept frozen at -80°C in aliquots until use. The SEC profiles as well as a gel of a representative sample after purification are shown in Figure 4. Previous attempts with DDM instead of LMNG yielded similar results, and LMNG was chosen, owing to its very low CMC and its lower background in cryo-EM.
Activity tests

Three types of tests were used, either in liquid medium or on agar plates.

1. Growth tests in liquid medium: *E. coli* C600 and its ΔexbBD derivatives were transformed with specified plasmids and their growth tested in liquid LB medium at 37°C, at various dipyridyl concentrations to induce iron starvation. OD$_{600nm}$ was measured after overnight growth.

2. Growth tests on plates: the relevant plasmids were also transformed into *E. coli* C600 ΔhemA, a heme auxotroph strain (and derivatives thereof), and growth of the strains was assayed as follows. Briefly, cells were first grown in LB medium (supplemented with delta-aminolevulinic acid (50µg/ml) to bypass the effect of the ΔhemA mutation) at 37°C up to an OD$_{600nm}$ of 1, and then mixed with melted top agar (0.6%agar in LB), and poured onto LB agar plates containing the appropriate antibiotics with arabinose at a concentration of 40µg/ml to induce expression of the genes under the $P_{araBAD}$ promoter(43). Wells, punched with Pasteur pipettes were filled with heme-albumin (at 5, 1 or 0.2µM), at two dipyridyl concentrations (100 or 200µM) to induce iron starvation. Plates were incubated overnight at 37°C and scored for growth around the wells. All experiments were performed in triplicate.

3. Growth curves in liquid medium: a few colonies of *E. coli* C600 ΔhemAΔtonBΔexbBD(pAMHasISRADEB + pBAD24 or derivatives thereof) were first inoculated in 4ml of LB medium at 37°C with the corresponding antibiotics, and 100µM dipyridyl, 4µg/ml arabinose but without delta-aminolevulinic acid. Once the culture reached an OD$_{600nm}$ of ca. 1.2-1.5, it was diluted and inoculated in 48 well Greiner plates, in the same medium to which was added 0.4µM He-BSA, as a heme source. The initial OD$_{600nm}$ of the cultures was 0.001. Each well contained 300µl of growth medium. Duplicates of each strain
were made, and the plate was incubated at 37°C with vigorous shaking (500rpm) in a Clariostar Plus Microplate reader. OD$_{600nm}$ was recorded every 30 minutes for 60-70 hours.

### N-terminal sequencing

The N-terminus of ExbB$_{Sm}$ was determined at the Plateforme Protéomique de l’institut de microbiologie de la Méditerranée (Marseille), after blotting on a PVDF membrane of a purified sample of ExbBD$_{His6}$ run on SDS-PAGE and determined to be APAAN.

### Lipid extraction

Chloroform (0.20 mL) and methanol (0.40 mL) were sequentially added to a sample of the ExbB or ExbBD complex (0.1 mL). The sample was vortexed for 10 min at room temperature and chloroform (0.2 mL) and water (0.2 mL) were further added. The organic phase was collected and the extraction procedure was repeated on the remaining aqueous phase. Combined organic layers were evaporated to dryness under argon and stored at -20°C.

### Mass Spectrometry

Dry lipid extracts obtained from ExbB, the ExbBD complex or from an *E. coli* lysate were solubilized in chloroform/methanol (50/50). Samples were then nano-electrosprayed using a TriVersa NanoMate (Advion Biosciences, Ithaca, USA) coupled to a Synapt G2-Si mass spectrometer (Waters Corporation, Manchester, UK). The instrument was calibrated in negative ion mode (for lipids) and positive ion mode (for proteins) from 50 m/z to 2000 m/z with NaI (50 mg/ml) with an accuracy of 0.8 ppm in resolution mode. The following settings were chosen: sampling cone 40 V, source offset 40 V, source temperature 80°C, trap gas flow 5 mL/min, helium cell gas flow 180 mL/min. MS/MS spectra were recorded using CID (Collision
Induced Dissociation) with a normalized collision energy set up to 30. To measure the mass of the intact proteins of the ExbBD and purified ExbB complexes, a desalting step was performed using micro Bio-Spin™ 6 (BIO-RAD) with 500mM ammonium acetate. Both samples were analyzed in denaturing conditions after a two-fold dilution with acetonitrile 4% formic acid. Mass spectra were acquired in positive ion mode. All molecular weights were measured after MaxEnt1 software deconvolution into neutral species.

**Cryo-EM grid preparation and data acquisition**

3 µL of either purified ExbB$_{Sm}$ or ExbBD$_{Sm}$ complex at ca. 1 mg/mL was applied to C-Flat 1.2/1.3 holey carbon grids (Protochips Inc., USA) previously glow-discharged in air for 30s. Grids were blotted for 2s at blot force 1 and vitrified in liquid ethane using a Vitrobot mark IV (FEI company) operated at 10 °C and 100% relative humidity.

All data collection was performed with a Titan Krios (ThermoFisher Scientific) operated at 300 kV equipped with a K2 Summit direct electron detector (ThermoFisher Scientific) at the European synchrotron research facility, ESRF (Grenoble, France). Movies were recorded in electron counting mode with EPU software (ThermoFisher Scientific), aligned with MotionCorr2 (44) and aligned images were processed with Gctf (45) using Scipion interface(46).

For ExbB data collection, 4567 movies were collected at a magnification of 165000x with a nominal pixel size of 0.827Å using a defocus range from -1.5 to -2.5µm. Movies of 56 frames were acquired using a dose rate of 8 electrons/Å$^2$/second over 7 seconds, yielding a cumulative exposure of 55.95 electrons/Å$^2$. 

For ExbBD data collection, 4043 movies were collected at a nominal magnification of 139000 x with a pixel size of 1.067Å. Movies of 48 frames were acquired using a dose rate of 4.6 electrons/Å²/second over 12 seconds yielding a cumulative exposure of 55.2 electrons/Å².

Cryo-EM image processing and analysis

a) ExbBD

For ExbB, aligned movies were processed with Gctf (45) and only images with a resolution higher than 4Å were kept; after visual inspection of the remaining images, processing was carried out with Relion-3 (47) (Supplementary Figure S10). Particles were extracted using a 2-fold binning, issued from a manual picking and a 2-D classification of particles picked out from 50 images. Automatic extraction was performed using the selected 2D class averages. After several rounds of 2D and 3D classification, 161k particles were selected for 3D refinement. They were corrected for local motion using Bayesian polishing option in Relion-3 and a post-refined map produced a 3.1Å overall resolution with a 5-fold symmetry.

A homology model was built with the Serratia marcescens ExbB sequence using Phyre2 server from 5SV0 monomer and docked into the refined map. Refinement was done with Phenix Real Space Refine option with secondary structure, “non-crystallographic symmetry” and Ramachandran restraints (48) and graphically adjusted with Coot (49). Lipid phosphatidyl glycerol starting structure and geometry were built using Phenix eLBOW (50) and was fit in the map using Coot then refined along with the protein model in Phenix.real_space_refine.

b) ExbBD

The data processing is summarised in Supplementary Figure S11. Movies were drift-corrected and dose-weighted using MotionCorr2 (44). Aligned dose weighted averages were imported
into Cryosparc2 (51) and contrast transfer function was estimated using CTFFIND4.1 (52).

Micrographs with poor CTF estimation statistics or high drift profiles were discarded. The remaining 3028 micrographs were used for automated particle picking. Particles were extracted, Fourier cropped to 2 Å/px and 2D classified. The best 2D classes were used as templates for automated particle picking resulting in 1.3 million particles. After several rounds of classification, the best 600k particles were submitted to 3D classification by means of multi-class ab-initio Reconstruction and Heterogeneous Refinement. 158 k particles belonging to the best resolved classes were corrected for local motion, re-extracted and used in Non-Uniform Refinement. The resulting refined map has a nominal resolution of 4.56 Å.

Based on the previous map a soft mask lining the micelle was designed in UCSF Chimera (53) and used to signal subtract the corresponding micelle density of particles in the refined map.

Localized refinement of the signal subtracted particles produced a map of the complex with an estimated resolution of 3.96 Å judging by FSC at 0.143 criterion. Data have been deposited both at the EMDB and PDB databases (EMDB 10789 and 11806, PDB 6YE4 and 7AJQ, for ExbBsm and ExbBDsm respectively, see Table II for refinement statistics).

Unless otherwise specified, the structure figures were made using UCSF Chimera software (53).

**NMR experiments**

The C-terminal periplasmic domain of HasB, HasB_{CTD} comprising residues 133-263, was produced and purified as previously reported (28, 54). The peptide corresponding to the periplasmic extension of ExbB_{sm} \((\text{A}_1\text{PAANPAVTESVAPTTAPAPAAAAPESITPVNPAPTIQPPETRG}_{44}^{-})\) numbering with reference to the mature protein was synthetized by Proteogenix.
NMR experiments were acquired at 293 K on a 600 MHz Bruker Avance III spectrometer equipped with a TCI cryoprobe. The spectra were processed with NMRpipe (55) and analyzed with CcpNmr Analysis 2.4 software (56). Proton chemical shifts were referenced to 2,2-dimethyl-2-silapentane-5 sulfonate as 0 ppm. $^{15}$N were referenced indirectly to DSS (57) (Wishart et al., 1995). $^1$H–$^{15}$N HSQC experiments were acquired on 0.15 mM HasB$_{CTD}$ in 50mM sodium phosphate pH 7, 50 mM NaCl with or without the peptide. Aliquots from a solution of peptide at 25mg/ml in the same buffer were added to the protein sample at 2:1 and 10:1 ratios. Chemical shift perturbations (CSPs) of backbone amide cross-peaks were quantified by using the equation CSP = $[\Delta \delta H^2 + (\Delta \delta N \times 0.15)^2]^{1/2}$, where $\Delta \delta H$ and $\Delta \delta N$ are the observed $^1$H and $^{15}$N chemical shift differences between the two experimental conditions. The $^1$H and $^{15}$N resonance assignments were from Lefevre et al 2007 (58).

Other biochemical methods

SDS-PAGE and immunodetection with anti-HasB or anti-His6 antibodies (Abcam [HIS.H8] (ab18184)) on whole cells or membrane preparations were carried out following standard protocols. Secondary antibodies were coupled to alkaline phosphatase.

BLAST search

ExbB$_{Sm}$ was used as a search for BLAST for orthologs in complete bacterial genomes, focusing on “long” ExbB’s. Top hits were in Serratia, Yersinia, Dickeya, Erwinia and Pseudomonas. Those genera were later excluded from successive BLAST searches to obtain orthologs in other genera with higher p-value.

Author contribution
PD and VB conceived the study and wrote the manuscript. All authors contributed to the manuscript and approved it. PD produced protein samples and performed microbiology experiments. MC performed electron microscopy sample screening and data collection. RJDA, PDC, VYNE and VB processed cryo-EM data. VB built structural models. VB and PD interpreted models. BL installed and tested programs. CM and JCR performed mass spec analyses. NIP and GCA performed NMR analyses. HS provided advice with data collection and processing.

**Data availability**

Protein structures were deposited to the protein data bank with ID 6YE4 and 7AJQ. Electron microscopy maps were deposited to the EMDB with ID 10789 and 11806.

**Acknowledgments** This work was supported by grants from the ANR (HEMESTOCKEXCHANGE ANR-12-BSV3-0022-01 and LABEX DYNAMO ANR-11-LABEX-0011-01). We thank Nathalie Dautin for helpful suggestions and critical reading of the manuscript, as well as Andrew Thompson. We acknowledge Emmanuel Frachon and Christophe Thomas from the Institut Pasteur platform “Production of recombinant proteins” for large scale cultures used in this work, European Synchrotron Radiation Facility for the provision of time on the Titan Krios at beamline CM01, the Institut Pasteur Biological NMR Technological Platform for the use of the NMR spectrometers, the staff at the ESRF CM01 facility in Grenoble for help in the data collection, the staff at the Marseille proteomics facility and Djalila Djouadi and Diana Basto who initially contributed to this project. We thank M. Nilges for interest in this work and B. Miroux for constant support.
References


### Table I: a list of selected species of Alphaproteobacteria where “extralong” ExbB’s are found.

All those species also contain HasB orthologs, except for the ones in red.
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Table II: Data collection, processing, and refinement statistics for ExbB<sub>Sm</sub> and ExbBD<sub>Sm</sub>.
Strains:

989  E. coli K12 C600
990  E. coli K12 C600ΔhemA::KmΔexbBD
991  E. coli K12 C600ΔhemA::KmΔexbBDΔtonB
992  E. coli K12 XL1-Blue
993  E. coli K12 JP313
994  E. coli K12 JP313ΔexbBDΔtonB
995  E. coli K12 C600ΔexbBD
996  E. coli K12 JP313ΔexbBDΔtonB
997  E. coli BL21DE3
998  S. marcescens Db11
999

Plasmids:

1000  pBAD24: lab collection
1001  pBAD33: lab collection
1002  pAM238: lab collection
1003  pAMHasISRADE: lab collection
1004  pAMHasISRADEB: this work
1005  pBADExbBDEc: this work
1006  pBADExbBDSm: this work
1007  pBADExbBDSm-ExbBD-EC: this work
1008  pBADExbBDSm-ExbBD-EC-84: this work
1009  pBADExbBDdelextss: this work
1010  pBADHasB: lab collection
1011  pBADExbBdelextssHis6: this work
1012  pBADExbBSmHis6: this work
1013  pBADExbB5cHis6: this work
1014  pBADExbBdelextssHis6: this work
1015  Oligonucleotides:

1016  ExbBD5c  5’-GGAGGAATTCACCATTGAAAAACGCTGGCAAGAAT-3’
1017  ExbBD6c  5’-AAGCTTGCATGCTTACTTCGCTTTGGCGGTTTCTT-3’
1018  SphIHisCtexbDSm  5’-TTGCATGCCTAATGGTGATGGTGATGGTGTTTGGCGCGCTTTCC-3’
1019  BglIIexbDSm  5’-GGCCTTAAATGAAGATCTGGACGACAGCGG-3’
1020  ExbBD5c  5’-AAGCTTGACTTTGCGGTTTCTT-3’
1021  ExbBD6c  5’-AAGCCTTGACTTTGCGGTTTCTT-3’
1022  PBADFOR  5’-CTGACGCTTTTTATCGCAAC-3’
1023  ExbBHis6  5’-AAGCTTGACTTTGCGGTTTCTT-3’
1024  ExbBDecEcSm76-84  5’-GGCAGTGAACTGCTGCGCGCCAAGCGTCGCCTTAAGCGCGAG-3’
1025  ExbBDecEcSm76-88  5’-TTTAGCGAACAAAATGGTCCAGGTGACTACGGAGGCCAAAAT-3’
1026  ExbBDecEcSm84-88.1  5’-CTAAAGGCAGTGAATTCTTCAATCAGAAGCGTCG-3’
1027  ExbBDecEcSm84-88.2  5’-CGAACAAAATGGTCAGGTGACTACGGAGGCCAAAAT-3’
1028  ExbBDecEcSm39-51  5’-GCGTAGAGTTCTTCAATCAGAAGCGCCGTCTGCGTCGCGA-3’
1029  ExbBDecEcSm39-51.2  5’-TCTTACTGAAAGGATTTGCCAGCTACGATAGACGCCAG-3’
1030  ExbBDecEcSm45-51  5’-GAAACCCGCGGCATGACCCCGCAGCTGTG-3’
1031  ExbBDecEcSm45-51.2  5’-GCCAGCTGTG-3’
1032  Table III: list of strains, plasmids and oligonucleotide sequences used in this work
**Figure Legends**

**Figure 1:** ExbB sequence analysis. A: alignment of ExbB<sub>Ec</sub> (top) with ExbB<sub>Sm</sub> (bottom), made with DNA Strider v1.4x-2b, CEA. In red is shown the signal sequence from ExbB<sub>Sm</sub>, and in blue the periplasmic extension, not present in ExbB<sub>Ec</sub>. The consensus sequence is shown on the middle line, where + indicates similar amino-acid residues. B: Weblogo (24) representation of 131 « long » ExbB aminoacid sequences aligned with Clustal omega (25); those with much shorter periplasmic extensions have been excluded from the alignment. The orange boxes show the position of the three TM segments from the *E. coli* 5SV0 structure.

**Figure 2:** ExbBD<sub>Sm</sub> complements ExbBD<sub>Ec</sub>. overnight growth in LB broth of *E. coli* C600ΔexbBD(pBAD24) (blue), C600ΔexbBD(pBAD24ExbBD<sub>Ec</sub>) (orange), and C600ΔexbBD(pBAD24ExbBD<sub>Sm</sub>) (grey), without (0) or with 100 and 200µM iron chelator di-pyridyl (DiP). One representative experiment is shown. The vertical axis represents mDO<sub>600nm</sub> absorbance units.

**Figure 3:** Role of HasB and TonB on bacterial growth. A: representation of the constructions used in Figure 3 B, 3C, 3D and elsewhere in this work. ss refers to the signal sequence, ext to the periplasmic extension of ExbB<sub>Sm</sub>, and tm1, tm2 and tm3 to the first, second and third ExB transmembrane segments, respectively. B and C: growth around wells of *E. coli* C600ΔhemAΔtonBΔexbBD harbouring pAMHasISRADE and specified recombinant plasmids (B and C left parts) or *E. coli* C600ΔhemAΔexbBD harbouring pAMHasISRADE and specified recombinant plasmids (B and C right parts); the pictures were taken after overnight growth at 37°C (B), or after 36hrs growth (C). Arabinose concentration was 40µg/ml. Dipyridyl (DiP) iron chelator concentration was either 100 or 200µM. Heme-BSA concentrations inside the wells were 10, 2 and 0.4µM. D: growth curves in microplates of the same strains as in B/C; DiP concentration was 100µM, arabinose concentration 4µg/ml, Heme-BSA 1µM. Colored dots in B and C refer to the equivalent growth conditions in D.

**Figure 4:** Homogeneity assessment of ExbB and ExbBD. Size exclusion chromatography profiles of representative purification of respectively ExbB<sub>Sm</sub>His6 (A) and ExbB<sub>Sm</sub>His6 (B) on a Superose 6 10/300 column. The 280nm absorbance is plotted as a function of elution volume. A Coomassie-stained gel of the pooled peak fractions is shown on the right of each profile together with a molecular weight ladder on the right (respectively 15, 25, 35, 40, 55, 70, 100, 130 and 170kDa). The faint band present in the Exb B<sub>Sm</sub>His6 sample above 100kDa was identified as AcrB using mass spectrometry.

**Figure 5:** Interaction of HasB<sub>CTD</sub> with the periplasmic fragment of ExbB<sub>Sm</sub>, as detected by NMR. A: Superposed 1H–15N HSQC spectra of 0.15 mM 15N-labelled HasB<sub>CTD</sub> in 50mM sodium phosphate, pH 7, 50 mM NaCl in the presence (red) or absence (black) of the periplasmic peptide of ExbB<sub>Sm</sub> residues 1-44. B and C: HasB<sub>CTD</sub> residues (PDB code 2M2K) exhibiting the highest CSP in the presence of the periplasmic peptide of ExbB<sub>Sm</sub> are coloured red (B: surface representation; C: cartoon representation). The residues showing the highest CSP are indicated.
Figure 6: ExbB TM1 interacts with HasB. A: bacterial growth curves of *E. coli*

*C600*Δ*hemA*Δ*exbBD*Δ*tonB* harbouring pAMHasISRADEB plasmid together with either

pBAD24 (medium-blue), pBADExbBD<sub>Sm</sub> (orange) or pBADExbBD<sub>Smdelextts</sub> (yellow),

pBADExbBD<sub>Ec</sub> (grey), pBADExbBD<sub>Ec</sub>Sm<sub>76-88</sub> (green), pBADExbBD<sub>Ec</sub>Sm<sub>76-84</sub> (dark-blue),

pBADExbBD<sub>Sm</sub>Ec<sub>39-51</sub> (magenta) in the presence of 1µM He-BSA as heme source, 4µg/ml arabinose to induce ExbBD expression and 100µM dipyrindyl to induce iron starvation (see Materials and Methods for further details). The optical path was ca. 3mm long, and each curve represents the mean of four replicates of the same culture, recorded every 30 minutes for 66 hours. Bottom: Coomassie-blue stained gel (B) and immunodetection (C) with anti-His<sub>6</sub> antibody of membrane preparations of *E. coli* C600 harbouring either pBAD24 plasmid (1), pBADExbBD<sub>Sm</sub> (2) or pBADExbBD<sub>Smdelextts</sub> (3). The equivalent of 0.6 OD<sub>600nm</sub> was loaded in B, and of 0.3 OD<sub>600nm</sub> in C.

Figure 7: Representation of the pentameric structure of ExbBD<sub>Sm</sub> solved by cryo-EM. A: the cryo-EM density map is shown, each monomer with distinct colours, view from the side. The yellow/grey regions represent non-protein density. The yellow regions were modeled as PG molecules. B: view from the cytoplasmic side. C: superposition of a monomer of ExbBD<sub>Ec</sub> (dark grey) and ExbBD<sub>Sm</sub> (red), represented as ribbon. The membrane thickness is represented by the dotted lines and the transmembrane segments TM1 (part of α2 helix), TM2 (part of α6 helix) and TM3 (part of α7 helix) are indicated. D: close-up representation of one PG density; E: fitting of the PG molecule inside the density; nearby ExbB residues are depicted as sticks using the Coot software (49).

Figure 8: Structure of ExbBD<sub>Sm</sub> solved by cryo-EM. The same color code as in Figure 7 for is used the ExbB monomers and the two ExbD monomers are colored yellow and gold. A: side view; B: view from the periplasmic space. C: clipped side view, with the helix of each ExbD monomer represented (yellow and gold). For B and C, the insets show the clipping planes.

Figure 9: Visualisation of “tunnels” inside ExbBD structures from *S. marcescens* (A and C) and *E. coli* 6TY1 structure (B and D), as calculated by the MoleOnline server with the same parameters. The Sm complex is colored as in Figure 8 and the Ec complex is colored in shades of grey for ExbB and orange and red for ExbD. The tunnels are the green volumes running through the structure, viewed from the periplasmic space (A and B), and from the side (C and D). The average tunnel diameter is 3Å for *S. marcescens* and 2Å for *E. coli*.

Figure 10: ExbBD co-expression is necessary for HasB stabilisation. Immunodetection of HasB in whole cells of *E. coli* JP313A*exbBD*Δ*tonB*(pHasB33) also harbouring pBAD24 (control), pBAD24ExbBD<sub>Ec</sub>, pBAD24ExbBD<sub>Sm</sub> or pBAD24ExbBD<sub>Ec</sub>Sm<sub>76-88</sub>, in the presence of either glucose (1mg/ml) or arabinose (40µg/ml), indicated by the + signs. The equivalent of 0.2OD<sub>600nm</sub> was loaded in each lane.

Figure 11: ExbB α2 helix: comparison between Sm and Ec. ExbB-ExbD structure with residues 76-88 coloured black and showing the aminoacid side chains. A, general view of the Sm pentamer; B, an enlarged view of A; C and D, two perpendicular close-up views of the exchanged region 76-88 in Sm (C) and 39-51 in Ec (D). E and F: Helical wheel representations of the swapped regions between ExbBD<sub>Sm</sub> and ExbBD<sub>Ec</sub> TM1. E: ExbBSm 76-88; F: ExbBEc 39-51. The red boxes show the helical face in interaction with the membrane.
Figure 2
Figure 3
Figure 4
Figure 5
Figure 6

A

OD_{600nm}

T (hrs)

B

C

MW, kDa

ExbB_{Sm}

ExbB_{Smdeletss}
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Figure 10
Figure 11