TatA and TatB generate a hydrophobic mismatch important for the function and assembly of the Tat translocon in Escherichia coli

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The twin-arginine translocation (Tat) system serves to translocate folded proteins across energy-transducing membranes in bacteria, archaea, plastids, and some mitochondria. In Escherichia coli, TatA, TatB, and TatC constitute functional translocons. TatA and TatB both possess an N-terminal transmembrane helix (TMH) followed by an amphipathic helix. The TMHs of TatA and TatB generate a hydrophobic mismatch with the membrane, as the helices comprise only 12 consecutive hydrophobic residues; however, the purpose of this mismatch is unclear. Here, we shortened or extended this stretch of hydrophobic residues in either TatA, TatB, or both and analyzed effects on translocon function and assembly. We found the WT length helices functioned best, but some variation was clearly tolerated. Defects in function were exacerbated by simultaneous mutations in TatA and TatB, indicating partial compensation of mutations in each by the other. Furthermore, length variation in TatB destabilized TatBC-containing complexes, revealing that the 12-residue-length is important but not essential for this interaction and translocon assembly. To also address potential effects of helix length on TatA interactions, we characterized these interactions by molecular dynamics simulations, after having characterized the TatA assemblies by metal-tagging transmission electron microscopy. In these simulations, we found that interacting short TMHs of larger TatA assemblies were thinning the membrane and—together with laterally-aligned tilted amphipathic helices—generated a deep V-shaped membrane groove. We propose that the 12 consecutive hydrophobic residues may thus serve to destabilize the membrane during Tat transport, and their conservation could represent a delicate compromise between functionality and minimization of proton leakage.

Tat systems serve the purpose to transport folded proteins across energy-transducing membranes in bacteria, archaea, plastids, and some mitochondria (1–3). They minimally consist of two components, TatA and TatC (4), but three-component TatABC systems are very common and found in the model Tat systems of Escherichia coli and plant plastids (5). TatA and TatB are structurally similar and evolutionary related (6). Both are membrane anchored by a 13 to 15 residues long N-terminal transmembrane helix (TMH) that contains only 12 consecutive hydrophobic residues. This helix is connected to an amphipathic helix (APH) via a short hinge at the cytoplasmic surface of the membrane (7–10) (Fig. 1A). The APH is followed by a negatively charged patch of residues in TatA (11), and more C-terminal regions are neither conserved nor functionally essential (12). TatC has six transmembrane domains with both termini on the cytoplasmic side (13, 14). TatB interacts tightly with TatC, whereas TatA gradually dissociates during purifications (15–17). From studies employing cross-linking and fluorescent protein tagging, it is known that the TatA interaction with TatBC changes during the translocation cycle, and larger TatA assemblies at TatABC complexes are believed to enable transport (18–23). A subpopulation of TatA protomers interacts with TatBC in a way that is not affected by mutations in TatC that abolish substrate binding (15), and resting-state-contacts of TatA to TatC have been identified (24). TatA assemblies undergo substrate-induced conformational changes that relate to Tat transport (25, 26), and the TatABC components rearrange during the translocation cycle (27, 28).

Tat-dependently transported proteins possess an N-terminal signal peptide that contains a highly conserved eponymous twin-arginine motif (1, 29). This motif is recognized by TatBC complexes that catalyze the membrane insertion of the signal peptide (30), whereas TatA is primarily required for the translocation step (18). TatA shows a high tendency to self-associate (31–34), but high-resolution structures are only available for detergent-solubilized TatA (7, 8, 10), and the mode of TatA self-interactions in membranes is still unclear. Spin-labeling studies indicated that the TMHs of TatA laterally interact (35), but only circular arrangements of aligned TatA have been suggested (8), and the orientation of the APH has not been included in the latest models (36). Interestingly, a dimeric structure of detergent-solubilized TatA indicated that also APHs of neighboring TatA protomers laterally interact (10).

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Roles of the hydrophobic mismatch generated by Tat systems

TMHs need to span the ca. 3 nm thick hydrocarbon core of membranes, and about 20 hydrophobic residues could in principal do this, but TMHs are in average 24.0 ± 5.6 residues long (37), which is generally due to flexing or tilting of helices in membranes (38). It is therefore very unusual that TatA and TatB have membrane anchors with TMHs of 13 to 15 residues that contain only 12 consecutive hydrophobic residues, and that the length of this 12-residues stretch is strictly conserved from archaea to bacteria (Fig. 1A). As this short length does not suffice to span a membrane lipid bilayer of normal thickness, the TMHs of TatA and TatB generate a hydrophobic mismatch. It has been found that the TMH of TatA per se can destabilize membranes (25), but potential effects of helix shortenings or extensions have never been systematically analyzed.

We therefore changed the hydrophobic mismatch of TatA and TatB by generating helix shortenings and extensions and investigated effects on Tat functionality and translocon assembly. Results indicate that both components tolerate smaller shortenings or extensions, but the natural 12 consecutive hydrophobic residues appear to function best. We demonstrate that, in case of TatB, this hydrophobic mismatch is important for the stable interaction of TatB with TatC, and therefore the short helices most likely trigger also the association of TatA with TatBC. To also analyze potential effects on the so far uncharacterized larger TatA assemblies that are present at Tat translocons during the translocation cycle, we visualized and characterized such assemblies in whole cells by metal-tagging transmission electron microscopy (METTEM), which to our

Figure 1. Constructs for the analysis of the 12-residues hydrophobic stretch in TatA and TatB. A, sequence logos of the functionally important region of TatA and TatB. Highlighted are the 12-residues hydrophobic stretch in the transmembrane helix of TatA and TatB (yellow), the preceding polar position (bluish purple), and the amphipathic helix (turquoise). The sequence logos have been obtained using enoLOGOS (88) based on sequence alignments by Clustal Omega (89) of sequences from a wide range of organisms, including α-, β-, γ-, δ-, and ε-proteobacteria, firmicutes, actinobacteria, cyanobacteria, chlorobi, chloroflexaceae, thermodesulfobacteria, aquificae, archaea, and plant plastids. B, TatA and TatB variants used in this study. The color code is the same as in (A). Inserted Leu residues are in red. C, carbonate wash analysis for the assessment of membrane integration of the TatA and TatB variants used in this study. TatA or TatB variants were detected by SDS-PAGE/Western blotting in the untreated membrane fraction (M), supernatant after carbonate wash (CS), and membranes after carbonate wash (CM). Molecular weight marker positions (kDa) are indicated. Note that all analyzed TatA and TatB variants are membrane inserted, but TatB variants are more stably membrane inserted than TatA variants, most likely due to their tighter interaction with TatC. The shorter the membrane anchor of TatA, the more was extracted by carbonate treatment.
knowledge is the first direct visualization of membrane protein interactions in whole cells, and then carried out molecular dynamics (MD) simulations. As a result of these simulations, we found that the short length of the TMH, together with the APH, has the potential to locally thin and destabilize the membrane. This is expected to cause membrane stress, which we could experimentally confirm by the analysis of the Psp membrane stress response induction.

Our study gives a biochemical and structural explanation for the membrane destabilization that is locally generated at Tat systems to permit the transport of folded proteins in all domains of life. The strictly conserved length of the TMH in TatA and TatB is not an obligate functional requirement and rather represents a compromise between a mechanistically important membrane destabilization and membrane stress minimization.

Results

**Tat systems tolerate some shortening and extension of the stretch of 12 consecutive hydrophobic residues in the TMHs of TatA and TatB**

To analyze the potential role of the short stretch of 12 consecutive hydrophobic residues in the TMHs of TatA and TatB, we shortened the hydrophobic helix by removing 1, 3, 5, or 7 residues in TatA, and 1 or 3 residues in TatB. We also extended the helix by 1, 2, 3, 4, 5, 6, or 7 leucine residues in TatA, and by 3 or 4 leucine residues in TatB (Fig. 1B). In the following, the nomenclature of the constructs is facilitated for the reader, with “A+1” standing for the construct with one Leu inserted into the helix of TatA and “B-3” standing for the construct with three hydrophobic residues deleted in the helix of TatB, as examples. To ensure the correct ratio of the Tat proteins, all genetic constructs were generated with the tatABC operon under control of its natural promoter in the Tat complementation vector pABS-tatABC (39). We used the tat-deletion strain DADE (40) in combination with these vectors, which permitted functional analyses as well as Blue-Native Polyacrylamide Gel Electrophoresis (BN-PAGE) detections of the Tat complexes. All TatA and TatB variants were stably produced, and carbonate washes revealed that all of them were membrane-integrated (Fig. 1C). The membrane integration of the very short TatA membrane anchors, such as A-5 or A-7, was unexpected and can only be explained by a major contribution of the APH. As expected, a significant portion of all TatA variants was released from the membranes, which is already known for TatA and may be due to the presence of these proteins in destabilized membrane regions (25). The shorter the membrane anchor, the more TatA was released by carbonate washes, indicating that membranes could not stably retain these TatA variants, possibly due to local membrane weakening (Fig. 1C, compare A-1, A-3, A-5, and A-7). As TatA can destabilize membranes, these results may indicate that shorter membrane anchors enhance the membrane-destabilizing effect of TatA. TatB and its variants were more stably membrane-integrated, which is expected as TatB tightly associates with TatC. We then analyzed Tat functionality with the described Tat systems. As Tat-deficient strains become SDS-sensitive, which relates to cell wall defects due to Tat-requirement for transport of the cell wall amidases AmiA and AmiC (41), we assessed Tat functionality by monitoring SDS resistance of the respective strains (Fig. 2A).

Only the A-5 and A-7 constructs were inactive among the TatA variants with shortenings, and only the A+6 and A+7 constructs were inactive among the extension variants. The A-5 and A-7 deletion variants showed higher sensitivity toward SDS than the noncomplemented tat deletion strain, indicating that these variants were not only nonfunctional but also had an additional negative physiological effect.

In case of TatB, the B-1 and B-3 deletion variants were both active in this assay, indicating that shorter hydrophobic helices in principle suffice to establish functional Tat translocons. This was unexpected as TatB tightly interacts with TatC for function, and we therefore expected less tolerance. The Tat system also tolerated the 3-residues extension in TatB but not anymore the 4-residues extension. As TatA and TatB possibly could have overlapping functions and therefore could complement defects of each other, we also tested combinations of single and triple deletions in TatA and TatB and triple extensions in TatA and TatB. Notably, the combined deletions were functional, indicating that indeed the deletions did not inactivate the proteins. The combined extension (A+3/B+3) showed a markedly reduced SDS resistance, indicating that some functional overlap had masked the effect of single +3 extensions. Nevertheless, the partial SDS resistance proofed residual activity of A+3 and B+3 variants.

A second way to monitor functionality is the chain formation phenotype, which is similarly based on the absence of AmiA and AmiC in the periplasm of Tat-deficient strains. If these amidases are not transported into the periplasm, the murein is not efficiently hydrolyzed between separating cells, resulting in chains of cells. In full agreement with the SDS-sensitivity measurements, the chain formation phenotype was only observed in those strains that were also SDS-sensitive (Fig. 2B). The strain with the A+3/B+3 combination, which had reduced but not abolished SDS resistance, showed no chain formation phenotype, indicating that sufficient amidases were transported for cell separation and confirming the residual activity of the Tat system with the +3 extensions in TatA and TatB. In line with that, the SDS-sensitive B+4 strain showed no obvious chain formation, indicating that transported amidases sufficed for cell separation but not for SDS resistance.

To recognize weaker effects on protein transport, we then carried out biochemical analyses of the transport of the Tat model substrate HiPIP, which is an iron-sulfur cluster-containing protein that strictly requires the Tat system for transport (15). Subcellular fractions were prepared from exponentially growing cells and analyzed by SDS-PAGE/Western blotting using antibodies specifically recognizing HiPIP (Fig. 3). We used a vector for low-level constitutive HiPIP production, which results in complete translocation of HiPIP into the periplasm in case of fully functional Tat systems (42). In this assay, the mature periplasmic HiPIP band.
indicates transport, and any precursor in the cytoplasm indicates reduced translocation relative to the fully functional Tat system. The WT Tat system, which served as positive control, showed a strong band of transported mature HiPIP in the periplasm and no accumulation of precursor in the cytoplasm. Fractionation controls showed that there was no leakage of cytoplasm into the periplasmic fraction, and the only faint mature HiPIP band in the cytoplasm indicated little proteolytic cleavage of the signal peptide or residual periplasm in the cytoplasmic fraction. The Tat-deficient empty vector control strain did not show any transport and instead the bands that are indicative for cytoplasmic accumulation of HiPIP, which are a strong cytoplasmic precursor band and a band that is due to partial degradation of the signal peptide. All TatA variants with shortenings showed detectable cytoplasmic precursor bands, with only very little precursor accumulating in A-1 variants and larger quantities accumulating in A-3 variants. The A-5 and A-7 variants were inactive, with no detectable mature protein in the periplasm. The TatA variants with extensions showed only slightly less transport in the A+1 variant and only very little cytoplasmic precursor in the A+2, A+3, and A+4 variants, but a sudden almost complete block of transport with the A+5 variant. The A+6 and A+7 variants were inactive. With the TatB variants, we observed little

Figure 2. Physiological Tat functionality assays with mutated Tat systems. Complementation of SDS sensitivity (A) and chain formation (B) phenotypes of the Tat-deficient strain DADE by Tat systems with indicated single or combined TatA and TatB variants. Scale bars in the micrographs correspond to 5 μm. Error bars of the SDS-sensitivity assays indicate SDs as deduced from three or more technical replicates. Controls: ‘no Tat’, empty vector control; ‘wt’, WT Tat system.

Roles of the hydrophobic mismatch generated by Tat systems

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accumulation with the B-1 and B-3 variants, indicating only a minor defect, whereas the B+3 variant strongly accumulated HiPIP in the cytoplasm and the B+4 variant hardly transported HiPIP anymore. Interestingly, the combinations A-1/B-1, A-3/B-3, and A+3/B+3 all showed a stronger effect than the single variants, indicating that the defects were additive or that there was indeed some partial compensation of defects of TatA by TatB or vice versa detectable in this semi-quantitative assay. Such effects were more difficult to detect by the apparently less-sensitive amidase-based assays, in which few transported enzymes may often suffice for physiological complementations (compare with the SDS sensitivity assay in Fig. 2).

Together, the activity measurements indicate that the 12-residues length of the hydrophobic helix in WT TatA and TatB results in highest transport efficiency. In case of TatA, shortenings by up to three residues and extensions by up to four residues are tolerated, and some residual transport is even detectable with a five-residues extension. TatB similarly tolerates shortenings and truncations of three residues, with only little residual transport detectable in case of the four-residues extended B+4 variant.

**Variation in the TatB TMH length, not in the TatA TMH length, interferes with TatBC complex stability**

It is well described that TatB interacts with the short helix 5 of TatC, whereas TatA can interact with the similarly short helix 6 of TatC (28, 36). It was therefore possible that functional effects of length variations in the hydrophobic TMH of TatA or TatB are simply due to disturbed interactions with TatC. To examine this aspect, we analyzed the stability of the known TatC-containing complexes (27).

Membrane fractions were solubilized by digitonin and analyzed by BN-PAGE/Western blotting to detect Tat complexes TC1 and TC2 (Fig. 4). While the complexes were unaffected by the length variations in TatA (Fig. 4A), the B+3 and B+4 variants caused disassembly of the complexes (Fig. 4B). In case of B-1, only the TC2 band was affected and changed to a very diffuse band, whereas the TC1 band remained unaffected. With B-3, the diffuse band of the size of TC2 was the only remaining complex, and TC1 had completely disappeared.

Some disassembly of Tat complexes is always observed and has been described previously (15). Some aggregated or not fully solubilized material is usually observed that remains at the top of the separating gel (17). Together, these data clearly indicate a severe effect of the TatB variants on TatBC interaction and consequently on TatBC complex stability. As also the functional TatB variants showed this destabilizing effect, sufficient TatBC complexes must have been present in living cells to support Tat transport with this expression system.

**METTEM-analysis of TatA clusters in the cytoplasmic membrane at WT expression level**

In the above shown BN-PAGE analyses, we could only detect effects of TatB variants on the Tat complexes, which is not surprising as most TatA dissociates from the detectable...
TatBC-containing complexes upon detergent solubilization, and only a regular ladder of multiple homooligomeric TatA associations is detected that does not permit analyses of the TatBC-bound TatA (31, 32). To reveal potential effects of the mutations on TatA, the lack of structural information in principle could be overcome by MD simulations. However, to create a reasonable starting point for MD simulations, we required some information about the organization of the large TatA assemblies at translocation sites. So far, it is only known from thylakoidal cross-linking analyses that at least up to 16 TatA protomers can be cross-linked at active translocation sites over a 4-min time span (22), which is evidence for an association of this number of protomers. Another analysis had indirectly calculated a median of 25 interacting TatA-YFP protomers in such complexes, based on mobility and fluorescence intensity measurements (23). The pixel size of 50 nm in these analyses certainly could not resolve single protomers or even whole TatA assemblies. We thus first aimed to examine the proposed larger TatA associations in the E. coli system by a higher-resolution method. To our knowledge, electron microscopy is the only currently available technique that can give sufficient positional resolution within cells. We chose MET-TEM for these analyses, using metallotheinonein-gold tags (43–48). Metallotheinonein-gold tags have the advantage of directly labeling the protein of interest with rather small electron microscopy–detectable electron-dense domains. In initial experiments with abundant recombinantly produced Tat systems, we found that three metallotheinoneins in row (= MT3) permitted clearly visible electron-dense granules of ± 1 nm in diameter in E. coli cells, and the detected particles could be identified as gold by Parallel Electron Energy-Loss Spectroscopy (PEELS) analysis (Fig. 5). Note that the recombinant protein levels that have been used for the PEELS analysis resulted in large clusters of interconnected associations that sometimes deformed the membrane, but this measurement only served to proof that the gold label of the MT3 tags had worked, and no other conclusion is drawn here.

Having the method established, we attempted to localize the Tat components at WT expression levels in strains that had the modified tat operons chromosomally integrated. With TatB-MT3 and TatC-MT3, no gold was detectable at WT level, but it was possible to detect TatA-MT3 in these experiments, which is likely due to the higher abundance of TatA relative to TatB and TatC in E. coli, which are present in an estimated 50:2:1 ratio, respectively (49). The WT level of the TatA-MT3-abundance was confirmed by SDS-PAGE/Western blot analysis (Fig. 6A). Substrates interact with the cytoplasmic face of TatA (25), and it is known that YFP tags (ca. 26 kDa) at the cytoplasmically located C-terminus of TatA inactivate TatA in the TatABC system of E. coli when produced at WT level (21). Similarly, we found that our MT3 tag (ca. 21 kDa) rendered TatA almost inactive, as a slight activity was likely due to some degradation to mature size (Fig. 6A). MT3 and TatC-MT3, no gold was detectable at WT level, but it was possible to detect TatA-MT3 in these experiments, which is likely due to the higher abundance of TatA relative to TatB and TatC in E. coli (25). As there so far do not exist any protein tags that can be fused to TatA without compromising activity at WT levels, this result was not surprising. Importantly, the tag did not prevent TatA–TatA interactions, and the number of TatA protomers in these associations was comparable with the number concluded from cross-linking of active nontagged TatA (see below, 22). Therefore, the effect on activity was most likely due to steric inhibitions of substrate interactions, as substrates need to pass through the TatA assemblies. In any case, these analyses can be taken as first direct visualization of WT level TatA assemblies in any organism.

TatA-MT3 formed gold assemblies near cell poles of irregular shape when seen in quasi top-view projection and roughly normal to the electron beam (Fig. 6, C and D). The mean length of the gold-assemblies was 20 nm ± 6.0 nm (N = 40, min = 10 nm, max = 37 nm). The width of the assemblies was about 10 nm. To determine the number of TatA-MT3 protomers in such assemblies, we counted the number of MT3-gold particles in 35 regions containing MT3-gold and calculated from this the average number of 17 ± 6 MT3-gold particles per 200 nm², which is the dimension of an average TatA-MT3 assembly. Interestingly, several of these assemblies visually appeared to be

![Figure 4. Analysis of Tat complexes in strains with mutated Tat systems. A, BN-PAGE/Western blot detection of Tat complex 1 (TC1) and 2 (TC2) in strains containing Tat systems with indicated TatA variants (A) or with TatB variants alone or in combination with TatA variants (B). Blots were developed with antibodies against TatB (upper blots) or TatC (lower blots). The position without the TC1 band in B-3 Tat systems is highlighted by open circles. B+3 and B+4 extensions have more general destabilizing effects. For direct comparison, the A+3 system is analyzed in parallel with B+3 and the A+3/B+3 combination. Positions of TC1, TC2, and molecular weight markers (440 kDa, 669 kDa) are indicated on the left.](image-url)
in contact, suggesting that they were somehow interdependent or connected (Fig. 6, E–H), which might be due to a clustering with TatBC complexes. Note that the MT3-gold intensity was variable, most likely due to individual kinetics of gold aggregate formation within MT3 tags, which may depend on the local environment (Fig. 7, A–C). Sometimes, TatA-MT3–gold-assemblies appeared in longer linear arrangements in the membrane. However, tilting of the sample revealed these arrangements to result from the superposition of individual assemblies when they are oriented in parallel to the electron beam within the ultrathin section (Fig. 7D).

In conclusion, TatA-MT3 formed longitudinal assemblies of irregular shape that were usually positioned near cell poles, the subcellular location of Tat translocons that has been observed in previous fluorescence-based studies (20, 50). The number of subunits in such assemblies agreed with previous calculations (22, 23). An often observed close clustering of several of these TatA assemblies suggested their indirect interaction, possibly mediated by TatBC, which explains the previous detection of a significant number of larger associations by fluorescence microscopy that was limited to 50 nm pixel resolution (23).

**MD simulation indicates membrane thinning and lipid disorder generation by the short TMHs and aligned APHs of TatA clusters**

Based on the METTEM analyses, which had directly shown the interacting TatA protomers and their predominantly linear
Roles of the hydrophobic mismatch generated by Tat systems

Figure 6. Detection of TatA-MT3-gold at WT expression level. A, detection of TatA by SDS-PAGE/Western blot analysis in subcellular fractions of strains with MT3-tags at TatA, TatB, or TatC as encoded by single copy chromosomally integrated tatABC-strep operons. WT *Escherichia coli* and the Tat-deficient strain DADE are analyzed for comparison. Positions of molecular weight markers are indicated on the left. B, SDS-sensitivity measurements of strains with TatA(MT3)BC, TatABC (wt), or no Tat system. C–H, clusters of TatA assemblies at WT expression level in strain DADE with a chromosomally integrated tatA(MT3)BC operon. (C and D) show overview images of cells with gold-labeled TatA-MT3-assemblies. Areas boxed in the respective overviews are shown in (E and F) at higher magnification. (E and G) are original images, and (F and H) were topographically 'relief'-filtered to discriminate MT3-gold granules in pseudo-3D within TatA-MT3 assemblies (lower edge = 100 nm). White 1 nm-dots (encircled in C–F) have been included as a direct scale for 1 nm MT3-gold-granules. C, cytoplasm; M, membrane; P, periplasm.
Roles of the hydrophobic mismatch generated by Tat systems

Figure 7. Detailed qualitative aspects of TatA-MT3-gold-labeled assemblies in Escherichia coli at WT level. A, survey view of an individual cell with a TatA-MT3 cluster, labeled with electron-dense gold (squared frame) in the subpolar region. B, detailed view of the area framed in (A), showing the TatA-MT3 assemblies of the cluster at higher magnification. Note the electron-dense granules of individual TatA-MT3-gold tags, which are about 1 to 1.5 nm in size. An isoscale scheme of the MD-simulated TatA assembly of 22 protomers (arrow, see Fig. 8A) is included for comparison. Note that the MT3 domains are 69 residues apart from the modeled APHs (green). C, a tilted, 3D-surface plot shows the range of gray level variance above the cytoplasmic background within the assemblies, which is indicative for MT3-gold. High peaks with black caps represent gold granules, as they can be readily recognized in (B). The range of different peak heights from high to low as seen in (C) indicates a variability of gold-loading of MT3 tags during the incubation of the cells with tetrachloroauric acid, possibly caused by differences in the local environment and unequal accessibility of gold. D, TEM-Tilting analysis of polar TatA assemblies.
Roles of the hydrophobic mismatch generated by Tat systems

assemblies, we were able to carry out 1500 ns MD simulations with 22 TatA protomers in the cytoplasmic membrane lipid bilayer, which is in the upper range of the observed TatA assemblies and therefore might reflect functional associations at Tat translocons. The MD simulations were challenged by the fact that, in living cells, larger TatA assemblies require TatBC interactions for their formation (20, 21). As no TatBC complex structure is known, and as we therefore could not include TatBC complexes in the simulations, we attempted to overcome this problem by placing linear arrangements of TatA protomers in close proximity with opposite APH-orientation. We had four reasons to expect a linear arrangement of TatA in the TatA assemblies: (1) our METTEM analyses had detected longitudinal assemblies (see Fig. 6), (2) an NMR structure of a TatA dimer showed laterally aligned APHs, which is only possible in linear arrangements (10), (3) circularly arranged TatA protomers had been reported to collapse within 40 ns in MD simulations, unless the circles were filled with a lipid bilayer that would need to be removed to permit translocation (8), and (4) a linear arrangement would permit regular interactions between TMHs of two oppositely oriented TatA protomers. This approach was successful, as the TMHs readily moved into staggered positions, and the resulting complex remained stable over time in the 1500 ns simulation (Fig. 8). Most APHs laterally contacted neighboring APHs, with some interactions interrupted (Fig. 8A), which agrees with the NMR evidence for lateral APH association (10), and these aligned APHs formed an angle of in average 109° with the TMH, which agrees with solid state NMR analyses on Bacillus subtilis TatA (51).

Also, the hinge was positioned deep in the membrane, which agrees with accessibility analysis data (25). The aligned APHs formed a deep V-shaped groove at the membrane surface (Fig. 8B and Video S1). As a result of this constellation, the membrane was thinned to about half its natural thickness at the hinge region. A view on the lipids in this simulation revealed that the lipid head groups were excluded from areas of aligned APHs, and the acyl groups were therefore disordered and partly oriented in parallel to the APHs. The lipid disorder in the membranes could be quantified employing the lipid tail order parameter Szz (Fig. 8C). The data confirmed that the lipid order was lowered in the environment of the APHs.

We then carried out the simulations with the TatA variants A-5, A-3, A-1, A+1, A+2, A+4, and A+6. As now expected, the shortenings resulted in further thinning of the membrane, whereas the extensions of the TMH resulted in a thicker membrane, which becomes evident when comparing the simulations A-5, wt, and A+6 (Fig. 8B). Accordingly, the lipid order further decreased with a shorter TMH and increased with a longer TMH (Fig. 8C). The TatA simulations therefore had added a potential new aspect to the role of the short length of the TMH, as they suggested that this exact length might not only be relevant for interactions (as shown for TatB; see Fig. 4) but also for membrane destabilization, which mechanically could be important (see Discussion).

As the MD simulations had indicated that large TatA assemblies can thin the membrane, we addressed whether the number of TatA protomers in the association could be relevant for membrane thinning. We repeated the simulations with 15, 10, and 5 TatA protomers and found that the membrane-thinning effect required the lateral association of multiple APHs, which was the reason why only Associations of >10 TatA protomers efficiently formed the V-shaped groove in the membrane (Fig. 9, A and B). In case of only five protomers, the lateral associations of APHs were too few to impose a V-shaped groove on the membrane lipids, which is why such small associations had less effect on the membrane bilayer, as quantified by the tail order parameter (Fig. 9C and Video S2).

Changing the length of the TatA TMH can affect cellular growth

The MD simulations suggested a second potential reason for the exact TMH length, which is the compromise between a mechanistically required membrane destabilization and an energetically required membrane stability. It has already been shown several years ago in in vitro experiments that TatA can destabilize the membrane and substrate-binding to TatA enhances this membrane destabilization, resulting in proton leakage (25). We thus wondered whether a too short TMH could result in extreme membrane destabilization in vivo, which could represent a selective pressure for maintaining the minimum of 12 consecutive hydrophobic residues in TatA (and consequently also in TatB and the short helices 5 and 6 in TatC, which need to assemble with TatA at the translocation site). As proton leakage might affect growth, we analyzed growth of all strains used in our study (Fig. 10).

Growth curves indicated a delay of growth by A-5, A-7, A+6, A+7, B-1, and A-3/B-3 systems, which may have been caused by cellular adaptations to stressed membranes (Fig. 10A). Some cases, especially A-5, A+6, and A+7, resulted in slow growth, indicating insufficient adaptations and persistent harm. The Psp membrane stress-response system is most likely involved in the adaptation to the membrane destabilization by the hydrophobic mismatch, as PspA is known to suppress proton leakage (52), and PspA of E. coli as well as its homolog LiaH in B. subtilis have been shown to interact with Tat systems (53, 54). We therefore tested this and found that shortened and elongated TatA variants as well as the B-1 variant induced the Psp membrane stress response, as monitored by an increase in PspA abundance (Fig. 10B). Interestingly, A-3 had a stronger effect than A-5 or A-7, and B-1 had a stronger effect than B-3, indicating that the orientation of the APH relative to the TMH influences stressor sensing by the Psp system. This upregulation represents most...
likely the adaptation that permits normal growth of strain A-3. In contrast, insufficient PspA induction in case of the extremely short A-5 and A-7 constructs most likely caused the clear growth phenotype. As the slow growth of the A+6 and A+7 constructs correlated also with the membrane stress response, this phenotype may be due to proton leakage as result of an influence on structures of other membrane proteins. The B+3 and B+4 constructs, which had a strongly reduced activity (Figs. 2 and 3) due to a diminished TatC interaction (Fig. 4), did not cause any significant growth phenotype, as the TatB elongations did not cause membrane destabilization (Fig. 10B). In general, constructs that did not have much effect on activity or whose effect on membrane stability could be compensated by the membrane stress response did not have strong effects on growth (Fig. 10), indicating that not compensated membrane stress likely is the reason for the observed growth phenotypes.

**Dominance of nonfunctional TatA variants**

To examine whether the inactive TatA variants influence the activity of functional Tat translocons, we examined Tat transport of the tat WT strain MC4100 in the presence or
absence of the recombinant inactive Tat systems A-5, A-7, A+6, and A+7 (Fig. 11). A microscopic examination already indicated that the recombinant A-5 Tat system suppresses Tat transport by the WT system of MC4100 (Fig. 11A). The more sensitive HiPIP transport assay then clearly showed that Tat transport is significantly inhibited in all four strains (Fig. 11B). In combination, these two assays indicate that the four tested nonfunctional TatA variants most likely mix into functional Tat systems, thereby rendering them almost inactive. The A-5 variant has the strongest inhibitory effect, resulting even in a chain-formation phenotype, whereas the residual activity suffices to enable normal cell separation in the other three cases. We also analyzed the induction of the Psp membrane stress response in these constructs (Fig. 11C). There was already a detectable upregulation by nonmutated recombinant Tat systems in strain MC4100, and this was markedly enhanced by A-5, A-7, A+6, or A+7 Tat systems, with A+7 resulting in the strongest membrane stress response. In summary, inactive TatA variants caused membrane stress also in MC4100 and exerted a dominant negative effect on the active WT Tat system in MC4100, most likely by mixing into the active translocons and thereby reducing their activity.

Figure 9. MD simulations with TatA complex of 15, 10, and 5 protomers. A, cross sections through TatA complexes. TMH, yellow; APH, turquoise; Q8, blue; hinge region, dark gray; N- and C-termini, gray; lipids, light and dark (head groups) brown. B, top views on the TatA complexes. C, lipid tail order parameter for lipids in the vicinity of the TMH (associated) and lipids in the bulk phase (nonassociated). Bars indicate 95% confidence intervals.
Discussion

Helices that pass the bacterial cytoplasmic membrane have to traverse a hydrophobic environment of about 3 nm (55). However, the TMH of TatA is predicted to consist of only 15 residues (10) and that of TatB consists of only 13 residues (9), and in these helices, there is only a stretch of 12 consecutive hydrophobic residues. A helix of 12 residues has a length of 1.8 nm, which does not suffice to span a lipid bilayer and therefore causes a hydrophobic mismatch. Such mismatches often have functional implications, such as the regulation of the mechanosensitive channel McsL or the aquaporin GlpF (55, 56), and hydrophobic mismatches can drive membrane protein interactions (57). This study now indicates that hydrophobic mismatches are also important for Tat functionality and Tat component interactions. The length of 12 residues is not a strict requirement for Tat transport but rather an optimal length, as little elongations and extensions of the TMH only reduce but not completely abolish function and interactions (Figs. 2 and 3). In case of TatB, our data show that all length variations affect TatBC complex stability (Fig. 4), and activity reductions are therefore likely due to weaker TatBC interactions. It is interesting that specific variations of the TMH in TatB can have selective effects on the different complexes, which might relate to conformational equilibria of the Tat complexes, such as previously described for mutations in TatC (27). Observations of residual activity with extended or shortened hydrophobic stretches in TatB are suggestive for significant flexibility of interactions within TatBC complexes, which likely are not rigid but rather dynamic oligomers. Also in case of TatA, a compromised interaction with TatB and/or TatC could contribute to the observed transport defects.

Interestingly, the combination of extensions or shortenings in TatA and TatB showed more severe defects than the mutations in single components, indicating that there exists some detectable functional overlap between TatA and TatB. The TatA TMH can occupy the same TatC-binding site as TatB (28), which may account for this functional overlap, and as the TatC interaction is likely important also for TatA, length variation in the TMH of TatA is expected to influence activity of the system, as does the length variation in the TMH of TatB.

As BN-PAGE is not helpful to address potential effects on TatA associations, which is due to the multiple associations that TatA forms in the presence of detergent, we sought to model such TatA associations with WT or mutated membrane anchors by MD simulations. Before doing any MD simulation, we required a more direct evidence for oligomeric TatA assemblies in E. coli, and the METTEM analysis served that purpose, showing that TatA-MT3 can indeed form large associations of an average 17 ± 6 protomers, which in turn can cluster to even larger associations, suggesting bridging interactions with other components, such as TatBC. In the end, our METTEM data fully agree with the cross-linking data of the thylakoid system, in which interacting components had been trapped over minutes (22), and they are also in remarkable agreement with the mean number of 25 protomers that have been calculated from fluorescence intensity and mobility

Figure 10. Growth analysis of strains containing the Tat systems analyzed in this study. A, A-5, A-7, A+6, A+7, B-1, A-3/B-3 inhibit cellular growth. For comparison, strains with the WT Tat system (wt) and without a Tat system (no Tat) are included in the growth analyses. All curves from constructs with growth in the range between the WT (blue line) and the Tat-deficient strain (no Tat, dashed blue line) are in gray and not specifically labeled for clarity reason. Stronger delayed or diminished growth phenotypes are labeled, and the corresponding curves are colored. Below the growth curves, the growth phenotypes of strains are compared, using the OD600 at the 3.36 h time point, as this time point is close to the end of the exponential growth of the fastest growing strains. Error bars indicate the SD of technical triplicates. Note that the OD600 was recorded by a 96-well multititer plate reader with 5.3 mm initial path length. B, SDS-PAGE/Western-Blot detection of PsPA (upper blot) and YidC (membrane protein control, lower blot) in membrane fractions of indicated strains, demonstrating the induction of the PsP response by membrane stress as generated by TMH length variations. Positions of molecular weight markers (kDa) are indicated on the right side of identical blot sections.
measurements of TatA-YFP foci (23). The latter fluorescence study also reported much larger TatA-YFP associations (at least up to 100 TatA-YFP), which may correspond to the somehow interconnected TatA assemblies in our METTEM analyses. Bridged-TatA assemblies likely diffuse together and therefore were not distinguishable from individual TatA assemblies within the 50 nm pixels in these fluorescence analyses (23). In contrast, METTEM resolved the 1 nm MT3-gold tags and therefore had a higher resolution than the current limit of C20 nm in super-resolution fluorescence techniques (58). To our knowledge, this is the first application for METTEM in localization of membrane protein associations in any organism.

Having directly seen and characterized WT level TatA associations in cells, we first simulated a larger cluster with 22 subunits, which was well in the size range of typical TatA assemblies in METTEM analysis (Fig. 7) and should be in the range of sizes of TatA associations at TatBC complexes. These simulations readily yielded stable TatA associations, with TMHs interacting with each other on a staggered position (Fig. 8). The assemblies were rather flexible, often showing small kinks due to some irregularities in the arrangements.

Most importantly, the MD simulations revealed that a potential second function of the short TMH could be membrane destabilization. The lipid tail order parameter clearly indicates a correlation of TMH length and membrane destabilization (Fig. 8C). Also important is the finding that, in these MD simulations, a larger association of TatAs is required for an efficient membrane destabilization (Fig. 9) and that the generation of the V-shaped groove comes along with generation of membrane curvature (Figs. 8 and 9). The simulations fully agree with published data: the APHs laterally interact, as previously evidenced by NMR (10), the TMHs laterally interact, as suggested by previous spin-labeling electron paramagnetic resonance studies (35), the hinge domain is positioned deep in the membrane, as supported by fluorescence quencher accessibility studies (25), and the APHs have a tilted orientation in the membrane, as seen in solid state NMR studies with B. subtilis TatA (51). It is noteworthy that the average angle of the APH relative to the TMH in the simulations is near 109°, which is in good agreement with the angle determined by solid state NMR (51).

Considering the obvious mechanistic tolerance of some length variation (Figs. 2 and 3), the strict conservation of the short length of 12 consecutive hydrophobic residues in TatA and TatB from bacteria, archaea, and plastids (Fig. 1) is unexpected, especially if only the functionality of protein–protein interactions is assumed to generate the selective pressure for the specific length. However, if membrane destabilization is a second function of these short TMHs, which may be of fundamental importance for the transport mechanism (59), then the hydrophobic mismatch as generated by the strict 12-residue length could be a delicate compromise between transport efficiency and proton leakage. Our MD simulations clearly show that the short TMHs and aligned tilted APHs of TatA have the potential to destabilize membranes, and the simulations of the TatA assemblies with extended or shortened TMHs show clear effects of such alterations on membrane thickness and lipid order (Fig. 8). Proton-tight membranes are of key importance for energy metabolism,
and bacteria have (often multiple) protective systems, which compensate for membrane destabilizations. First growth experiments already indicate that major adaptation processes are required in response to A-5 and B-3 systems, and adaptations to minor changes in TMH length on the membrane energetization may be sufficient to suppress growth phenotypes (Fig. 10, A and B). In agreement with this hypothesis, we found that the Psp membrane stress response was induced by A-3 and A-5 systems, and this induction was much stronger in A-3 than in A-5, which is in agreement with the growth data. The Psp system is known to stabilize membranes (52) and PspA already has been shown to interact with TatA in Tat systems of Gram negative and Gram positive bacteria (53, 54).

In light of the results of this study, our data strongly support the hypothesis that TatA, TatB, and TatC generate locally destabilized membranes that permit the membrane passage of folded proteins (25, 60). The “membrane weakening and pulling” hypothesis proposed that this process could depend on membrane destabilization by multiple aligned TatA TMHs (60), which was later supported by an experimental demonstration of membrane destabilization, MD simulations, and recently by electrochromic shift measurements (8, 25, 59). A conformational switch of the TatA APH in response to substrate binding has been first demonstrated in thylakoids and then in the E. coli system and has been suggested to contribute to this membrane destabilization (25, 26).

Hydrophobic mismatches are not only generated by the TMHs of TatA and TatB, but also by the short TMHs 5 and 6 of TatC. Notably, the TMHs of TatA and TatB are both shown to interact with these two TatC helices in a resting state (24, 28, 36), which is likely the reason for the shown importance of the specific short length of the TatB TMH for stable TatBC associations (Fig. 4). As substrate binding has been shown to trigger the association of TatA (20–22, 61), as TatA association has been reported to precede the translocation of TatA-bound Tat substrates (18, 19), as substrate has been shown to trigger membrane permeabilization by TatA in vitro (25), and as TatA assemblies have the capacity to destabilize the membrane (Fig. 8), we would like to suggest that larger TatA assemblies may form at TatABC complexes to enable Tat transport (see Fig. 12). In principle, such an interaction would explain why most TatA is lost during purification of TatBC complexes (15), and this model also explains how a simple conformational shift of TatC helices 5 and 6 in response to substrate binding may enhance the hydrophobic mismatch at the TatBC core to induce the binding of a large number of TatA protomers to the translocation site, which induces membrane destabilization where it is needed (Fig. 12).

The model also gives an idea about the region of membrane destabilization, which is essentially the region underneath the APHs. There are two plausible pathways through the destabilized membrane: (1) the two staggered rows of TatA, which are vis-à-vis positioned in the cluster, could open and permit the passage of the Tat substrate or (2) the TMHs remain in the center of the association and the substrate passes through the thinnest point of the destabilized membrane close to the center of the TatA cluster. As Tat substrates can be large in diameter, it is likely that both happen as follows: TMHs may lose their contact and APHs may dissociate to permit the passage of globular proteins. In such a situation of disorder, it could well be that TatA exchanges with TatB at its TatC-binding site, especially when the system is saturated with transported substrates (28). It is also possible that an opening of TatBC complexes can contribute to the TatA recruitment, such as proposed previously (2, 24, 36). Our simulations show that TatA can form stable clusters with two interacting rows of TMHs in the center. From the energetic point of view, MD simulations indicate that there will not be any stable ring-shaped TatA assembly with an aqueous hole (8). Instead, the hydrophobic mismatch can be expected to trigger associations of the TMHs, which preferentially should be linear to accommodate aligned APHs at the membrane surface that contribute to membrane destabilization (Figs. 8 and 9). During the passage, transient aqueous holes would be formed that would cause uncontrolled proton flux for a very short time span. Such a proton leakage would agree with the finding that about 80,000 protons pass through the membrane per
Roles of the hydrophobic mismatch generated by Tat systems

transported protein (62), albeit proton flux does not energize the transport (63, 64). The membrane-thinning assemblies of TatA are expected to alter the diffusion rate in the membrane, and this has been observed in a recent study that is currently published as preprint in BioRxiv (65). Although it is clear from many studies that the electric potential is required at certain steps of the pathway, one of which may be the proposed conformational transition of the TatBC core complex that induces the recruitment of TatA (which is PMF-dependent; (20, 21)), the driving force for the transport remains to be clarified. In principle, it would be possible that the transmembrane signal peptide insertion as catalyzed by TatBC complexes (30) constitutes this driving force, as the signal peptide remains bound to the TatBC complex during transport of the mature domain (66). Future experiments surely will shed further light on this highly fascinating transport system.

Experimental procedures

Bacterial strains and growth conditions

The E. coli ΔtatABCD ΔtatE strain DADE (40), and its parental strain MC4100 (67), were used for localization and complementation studies. E. coli XL1-Blue Mrf' Kan (Stratagene) or E. coli DH5α λ pir' were used for cloning. Unless otherwise stated, bacteria were grown aerobically at 37 °C on LB medium (1% (w/v) tryptone, 1% (w/v) NaCl, 0.5% (w/v) yeast extract) in the presence of the appropriate antibiotics (25 μg/ml chloramphenicol, 100 μg/ml ampicillin, 12.5 μg/ml tetracycline, 15 μg/ml kanamycin). For subcellular fractionations, all cultures were normalized to an OD600 of 1.0. For growth curves, the OD600 was measured with technical triplicates in 15-min intervals with cultures grown at 37 °C in shaking 96-well plates (culture volume: 200 μl), using the SpectraMax iD3 Microplate Reader (Molecular Devices).

Genetic methods and plasmids

For analyses of Tat functionality and for optimization of the MT3-gold-labeling method, the TatABC components were constitutively overproduced using the natural P_pstA promoter and tatABC operon organization in plasmid pABS-tatABC and derivatives thereof (39). The shortenings and extensions of the hydrophobic helices in TatA and TatB were carried out by QuikChange mutagenesis. Primers are listed in Table 1.

To analyze MT3-gold-labeled Tat components at WT level, the tatABC operon and its derivatives were cloned into pAH120-Ptat-tatABC-strep (see primer table), which is a tatABC-containing derivative of pAH120-Ptat-tatA-strep-ΔBamHI (68). Briefly, the MT3-tag (including the linker) was amplified from the pMAL-c2x-MT3 (48) using primers MT-I-F and MT-R, cut with BglII/BssHII, and ligated into the backbone of the plasmid pAH120-Ptat-tatABC-strep cut with BamHI/BssHII. This resulted in pAH120-Ptat-tatABC-MT3, which was then cut with Ndel/BamHI and ligated into the corresponding sites of pABS-tatABC-H6 (27), resulting in pABS-tatABC-MT3. To construct pABS-tatA-MT3, tatA was amplified from pABS-tatABC-MT3 using primers pABS-tatA-MT3-R and pABS-tatABC-Sall-F, and the fragment was cut with Sall/SaCl and ligated into the plasmid backbone obtained by restriction of pABS-tatABC-H6 with the same enzymes. The same procedure was followed to construct pABS-tatAB-MT3, except that the primers utilized were pABS-tatB-MT3-R and pABS-tatABC-Sall-F. To append the tatBC-genes to the tatA-MT3-construct, these were amplified from the pABS-Ptat-tatABC-MT3 plasmid using primers pABS-tatB-BamHI-F and pABS-tatC-ClaI-R, and the fragment was cut with BamHI/ClaI and ligated into the respective sites of the pABS-Ptat-tatA-MT3 backbone. The same steps were taken to construct the pABS-tatAB-MT3-tatC plasmid, only that primers pABS-tatC-F and pABS-tatC-ClaI-R were used and the fragment was ligated into the backbone of pABS-Ptat-tatAB-MT3. Finally, both the tatA-MT3-BC and tatAB-MT3-tatC constructs were transferred back into pAH120 by restriction of pAH120-Ptat-tatABC-MT3, pABS-tatA-MT3-tatBC, and pABS-tatAB-MT3-tatC with Ndel/Xbal and ligation of the tat-fragsments into the pAH120-backbone. Vectors were integrated into the chromosomal lambda attachment site according to the protocol of Haldiman and Wanner (69). pRK-hip was used for constitutive low-level expression of the hip gene from its own promoter (42).

Biochemical methods

Subcellular fractionations into periplasm, membranes, and cytoplasm were performed with 50 ml cultures as described previously (68). SDS-PAGE analysis was carried out by standard methods (70). BN-PAGE was performed as described previously (32), but without β-mercaptoethanol in the buffers. TatABC complexes were solubilized with 1% digitonin. For immunoblots, proteins were semi-dry blotted on nitrocellulose membranes, and blots were developed using antibodies directed against synthetic C-terminal peptides of TatA, TatB, TatC, or the chain formation phenotype was assessed by phase contrast microscopy. SDS sensitivity was determined by aerobic growth in LB medium containing 4% (w/v) SDS, using the quotient of the OD600 with/without SDS after 3 h of growth (41). For carbonate washes of membranes, samples were treated as described before (25).

Electron microscopy – sample preparation

Bacterial cells (20 ml culture, 5% inoculum) were grown at 37 °C at 140 rpm to OD600 ~0.5, then 0.4 ml of an aqueous 25 mM tetrachloroauric acid solution were added, and growth was continued for 90 min at 37 °C and 120 rpm. Cells were sedimented (10 min, 3820g, 4 °C), resuspended in 1.5 ml 100 mM Hepes pH 7.0, 90 mM sucrose (buffer A), and incubated for 5 min at room temperature (RT). Cells were sedimented again (2 min, RT, 6800g) and the incubation/
Roles of the hydrophobic mismatch generated by Tat systems

Table 1

| Mutagenesis primers<sup>a</sup> | Sequence (5<sup>′</sup>-3<sup>′</sup>) |
|-------------------------------|----------------------------------|
| Deletions                     |                                  |
| tatA-1-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatA-3-F                     | GTATTTGGGCTTAATAGTCGCTGATCTG    |
| tatA-5-F                     | GGTACATTTGGGCTTAATAGTCGCTGATCTG |
| tatA-7-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatB-1-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatB-3-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatA+1-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatA+2-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatA+3-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatA+4-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatA+5-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatA+6-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatA+7-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatB+3-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatB+4-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatA+1-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatA+2-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatA+3-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatA+4-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatA+5-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatA+6-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatA+7-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatB+3-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| tatB+4-F                     | CAGTTTGGGCTTAATAGTCGCTGATCTG    |
| Fusion Primers                |                                  |
| MT-F                         | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| MT-R                         | TATCGGCGCGCTCTGAATTACCGCGCAAGCGCCGAGG |
| pABS-tatB-BamHI-F            | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| pABS-tatC-ClaI-R             | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| pABS-tatABC-MT3-R            | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| pABS-tatABC-MT3              | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| Sall-P                       | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| pABS-tatC-F                  | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| Primers for cloaking of       |                                  |
| pAH120-Prat-tatABC-strep     | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| tatA-1-F                     | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| tatA-3-F                     | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| tatA-5-F                     | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| tatA-7-F                     | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| tatB-1-F                     | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| tatB-3-F                     | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| tatA+1-F                     | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| tatA+2-F                     | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| tatA+3-F                     | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| tatA+4-F                     | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| tatA+5-F                     | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| tatA+6-F                     | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| tatA+7-F                     | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| tatB+3-F                     | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |
| tatB+4-F                     | CCGAGGAGGATCCAAATCGCGTCGCAACAAACAC |

<sup>a</sup> Primer sequences are given in 5<sup>′</sup>-3<sup>′</sup> direction.
<sup>b</sup> Corresponding reverse primers covered the same sequence.
<sup>c</sup> Templates for larger deletions or insertions were the constructs with smaller deletions or insertions.

resuspension was repeated. Cells were then resuspended in 1 ml buffer A containing 2.5% glutaraldehyde, incubated for > 20 min at RT, washed with 1 ml buffer A, sedimented (2 min, 6177g, RT), and washed twice in 100 mM Hepes pH 6.9, 90 mM sucrose, 10 mM MgCl2, 10 mM CaCl2, sedimented again, and embedded either in SPURR epoxy resin (71) or in LR-White resin (72). Seventy nanometer ultrathin sections were cut using an ultramicrotome with a diamond knife (Ultracut; Leica Biosystems) and picked up with neoprene-collargol. Seventy nanometer ultrathin sections were cut using an ultramicrotome with a diamond knife (Ultracut; Leica Biosystems) and picked up with neoprene-coated Cu-grids (300 mesh, hexagonal). TEM image registration and processing: Unstained samples were analyzed in the elastic bright-field mode with an in-column energy-filter transmission electron microscope (Libra 120 plus, Zeiss). The energy-selecting slit aperture was set to 15 eV energy width, and images were recorded with a cooled bottom-mount 2 × 2 k CCD (SharpEye, Tröndle) at nominal magnifications from x12,500 to x50,000 near Gaussian or at 500 to 1500 nm underfocus. Data acquisition and spectrum/image-analysis were done using the iTEM software package (Olympus Soft Imaging Solutions Ltd). Images were further processed with the Corel Draw Graphics Suite X7 (Corel Ltd) and ImageJ Fiji software (https://fiji.sc/) (73). PEELS: PEELS was done with unstrained 35 nm ultrathin sections of epoxy resin-embedded E. coli strain DADE/pABS- tatAB(MT3)-C with MT3-gold as the only electron-dense intracellular material. PEELS registration was performed in the energy range from 30 to 139 eV, specific for the Au-O<sub>2</sub> edge, with a 100 μm spectrum entrance aperture and five acquisition cycles pro record. Spectrum magnification was set to 200-fold, and the illumination aperture to 0.16 mrad with 4 μA emission current; exposure time: 0.2 s, nominal magnification: ×31,500. Electron spectroscopic element mapping: Data acquisition was performed on well-defined Au-MT3 clusters and the energy-selective slit was set to 8 eV width. Jump-ratio images were registered at 66 eV (=Emax) and 38 eV (=first window). These energy settings were taken from PEEL spectrum data of the MT3-gold assemblies of interest. Exposure time: 2 s, illumination aperture: 0.63 mrad, nominal magnification: ×31,500.

MD simulation

MD simulations were performed using the GROMACS simulation package, version 2019.5 (74). The MARTINI v2.1 coarse-grained force field (75–77) was applied to simulate lipids, peptides, and solvent. In all simulations, the system was coupled to a constant temperature bath by v-rescale algorithm (78) with a relaxation time of 1.0 ps. Simulations were performed at 320 K. Periodic boundary conditions were applied to simulate bulk behavior. The time step used in the simulation was 20 fs. The dielectric constant in the simulations was 15. The pressure was coupled to 1 bar employing the Parrinello-Rahman barostat (79) with a relaxation time of 12.0 ps. Unless stated otherwise, the TatA membrane system consisted of 22 TatA proteins in a lipid bilayer with 75% phosphatidylethanolamine, 19% phosphatidylglycerol, and 6% cardiolipin, which is a usual ratio of these lipids in the cytoplasmic membrane of E. coli (80). For the MD simulations of TatA associations, a coarse-grained TatA model up to residue Glu47 was constructed based on the known NMR structure (PDB 2MN7, 10). To model the linear assembly of TatA proteins observed in the TEM experiments, 22 copies of the equilibrated TatA structure were placed vis-à-vis in two rows of 11 subunits and inserted in the membrane via deletion of all overlapping membrane lipids. The overall dimension of these 22-mers, about 20 nm in length, closely match to the ones observed in METTEM. This system was then equilibrated for 500 ns to enable adaption of the membrane. The TMHs readily associated in a staggered way with all TatA proteins included in the linear complex. For all simulations, the TatA sequences up to position Lys49 were used, as C-terminal truncation analyses have shown that a TatA of exactly that length is still functional (12). MODELLER 9.24 (81) was used to create the TatA variants. The resulting data were evaluated by MDA-analysis 0.20.1 (82). The lipid tail order parameter S<sub>zz</sub> = ½.
(3(b)n)^2−1) was determined to quantify the average orientation of tail bond vectors b with respect to the bilayer normal n (83). Order parameter time series were checked for equilibration and decorrelated using the pymbar package (84–86). Angles between the TMH and the APH were calculated between two vectors connecting the center of masses (COMs) of residues Gly21, Thr22, and Lys23 for the hinge either with the COM of residues Ala13, Val14, and Ile15 for the TMH or with the COM of residues Ile28, Gly29, and Ser30 for the APH (WT numbering). Images of the simulations were generated using NGLview v2.7.7 (87).

**Data availability**

All data are contained within the article.

**Supporting information**—This article contains supporting information.

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**Abbreviations**—The abbreviations used are: APH, amphipathic helix; BN-PAGE, Blue-Native Polyacrylamide Gel Electrophoresis; COM, center of mass; METTEM, metal-tagging transmission electron microscopy; PEELS, Parallel Electron Energy-Loss Spectroscopy; Tat, twin-arginine translocation; TMH, transmembrane helix.

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