Recent progress in structure and dynamics of dual-membrane-spanning bacterial nanomachines
Vicki Gold\textsuperscript{1} and Mikhail Kudryashev\textsuperscript{2,3}

Advances in hard-ware and soft-ware for electron cryo-microscopy and tomography have provided unprecedented structural insights into large protein complexes in bacterial membranes. Tomographic volumes of native complexes \textit{in situ}, combined with other structural and functional data, reveal functionally important conformational changes. Here, we review recent progress in elucidating the structure and mechanism of dual-membrane-spanning nanomachines involved in bacterial motility, adhesion, pathogenesis and biofilm formation, including the type IV pilus assembly machinery and the type III and VI secretion systems. We highlight how these new structural data shed light on the assembly and action of such machines and discuss future directions for more detailed mechanistic understanding of these massive, fascinating complexes.

**Addresses**
\textsuperscript{1} Department of Structural Biology, Max Planck Institute of Biophysics, Max-von-Laue Str. 3, 60438 Frankfurt am Main, Germany
\textsuperscript{2} Max Planck Institute of Biophysics, Max-von-Laue Str. 3, 60438 Frankfurt am Main, Germany
\textsuperscript{3} Buchmann Institute for Molecular Life Sciences, Goethe University of Frankfurt, Max-von-Laue Str. 17, 60438 Frankfurt am Main, Germany

Corresponding authors: Gold, Vicki (vicki.gold@biophys.mpg.de) and Kudryashev, Mikhail (misha.kudryashev@biophys.mpg.de)

**Current Opinion in Structural Biology** 2016, 39:1–7
This review comes from a themed issue on Membranes
Edited by Jacqui Gulbis and Gabriel Waksman
For a complete overview see the Issue and the Editorial
Available online 17th March 2016
http://dx.doi.org/10.1016/j.sbi.2016.03.001
0959-440X/\textcopyright 2016 The Authors. Published by Elsevier Ltd. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

**Introduction**
A distinctive feature of all Gram-negative bacteria is that they have two membranes—a relatively porous outer membrane and an impermeable cytoplasmic membrane. Sandwiched between them is the densely crowded periplasm and shape-determining peptidoglycan layer. Bacteria have evolved highly specialized protein machineries that span both membranes, enabling complex functions, such as cell movement \cite{1} and molecular targeting, including protein secretion and DNA uptake \cite{2}. Pathogenic and symbiotic Gram-negative bacteria interact with other cells by secreting effector proteins across both bacterial membranes, into the extracellular matrix or directly into host cells \cite{3}. The mechanistic complexity of these nanomachines is achieved by assembling multiple copies of more than 20 different proteins, including both soluble and membrane-anchored subunits as structural and regulatory elements \cite{4}. Understanding their molecular mechanism requires a combination of structural biology with genetic tools and functional studies. Some of the most pertinent questions are: how do dual-membrane spanning nanomachines assemble in order to perform their biological function? What is their mode of action? How is communication between different subunits regulated and controlled?

Recent advances in single-particle electron cryo-microscopy (cryo-EM) have made it possible to determine structures of isolated proteins or complexes at near-atomic resolution \cite{5}. However, a major technical challenge associated with studies of large multi-component membrane protein complexes is the difficulty of purifying them in an intact state. Electron cryo-tomography (cryo-ET) enables large protein complexes inside cells to be observed in 3D at a resolution of several nanometers, by tilting a sample through a series of increments \cite{6,7}. The resolution may be further improved by sub-tomogram averaging (StA), whereby multiple copies of the same protein or protein complex are mutually aligned and added together (Figure 1). This can yield sub-nanometer resolution, sufficient to distinguish protein domains \cite{8} or even secondary structure \cite{9}. Integration of higher-resolution information obtained by single particle cryo-EM or X-ray crystallography into the StA density maps provides detailed mechanistic understanding at the level of the entire complex \cite{8,10,11}. In addition, trapping of specific physiological states or computational classification of different protein subsets can yield intermediates of assembly and function. Here, we review recent exciting insights into structure and dynamics of the type IV pili (T4P) assembly machinery, and of the type III and VI secretion systems (T3SS/T6SS) in bacteria \cite{3}, obtained by cryo-EM and cryo-ET.

**Requirements for \textit{in situ} structural analysis**
Several technical aspects need to be considered for structural determination of bacterial membrane protein complexes \textit{in situ}. Firstly, sample thickness places a strict limit on the quality of tomographic volumes. In practice this means a maximum thickness of 400–500 nm. It is possible to reduce cell thickness by optimizing culture conditions \cite{12}, genetically engineering thin cells or small minicells \cite{13}, or by physical sectioning of the frozen bacteria with a
diamond knife [14] or a focused ion beam [15]. Secondly, cryo-EM has benefitted greatly from direct electron detector cameras [16] and phase plates [17], generating high quality data in a high-throughput and automated manner. Thirdly, an inherent problem with cryo-ET is the ‘missing wedge’ of information, resulting in resolution anisotropy in the direction of the electron beam [18]. The missing wedge results from the current physical inability to rotate the sample in the microscope by a full 180°. Effects of uneven sampling are alleviated or eliminated by averaging particles in different orientations by StA [19]. Fourthly, large membrane protein complexes are often flexible, resulting in variable low-resolution maps. Conformational flexibility may be reduced biochemically, for example by cross-linking [20], or genetically, by introduction of stabilizing mutations or truncations, which carry the disadvantage of compromising the native state of the complex. Alternatively, computational methods of classification can be used to quantify and account for flexibility [21]. Finally, a prerequisite for both classification and high resolution is the collection of very large particles data sets. Parallel and GPU-enhanced software, and thorough image analysis helps to obtain unbiased structures and structural intermediates [21–24].

**Bacterial type IV pilus assembly machinery**

The type IV pilus is a surface-exposed filamentous protein polymer that is several μm long, anchored to cells of evolutionarily divergent Gram-negative and Gram-positive bacteria [25]. In Gram-negative bacteria, the assembly machinery forms a multimeric dual-membrane-spanning protein supercomplex [26]. T4P play an important role in cell motility, enabling cells to adhere to and move along surfaces, form colonies and biofilms [1], and can even act as nanowires carrying electrical current [27]. Pathogenic proteobacteria, such as *Pseudomonas aeruginosa* and *Neisseria meningitidis*, use T4P to mediate adhesion to host cells prior to infection [28,29]. The machinery that assembles T4P is also implicated in DNA uptake, which is critical for lateral gene transfer and adaptive evolution [2]. The T4P is
related to both the archaean equivalent (the archaellum) [30], and the bacterial type II secretion system (T2SS) [31], which in some bacteria secretes proteins and toxins into the extracellular environment.

The composition of pili varies between species, but they are predominantly formed of an oligomer of the major pilin protein (PilA in *Pseudomonas* and *Thermus*, PilE in *Neisseria*) plus a few copies of different minor pilins [32]. X-ray structures of several major pilins have been reported [32], demonstrating a conserved fold and a characteristic ‘lollipop’ shape. A pseudo-atomic model of a filamentous pilus from *Neisseria gonorrhoeae* was built by docking X-ray structures of pilin subunits into a cryo-EM map of isolated pili, which were ~6 nm wide with a narrow central channel of ~1 nm [33]. Studies of multiple species, including *N. meningitidis* [34] and *Thermus thermophilus* [2], have shown that pre-pilins mature by the action of a peptidase (PilD), then assemble at the inner membrane by a AAA-ATPase (PilF) with the help of additional proteins, including PilM, PilN and PilO. The pilus is directed across the outer membrane through the pore of a large multimeric ~1 MDa secretin protein (PilQ), which in *Thermus* is formed by a membrane-embedded cone-like structure with a series of 6 stacked rings in the periplasm [35]. Bacterial T4P are unique in that they can be rapidly retracted through the action of a cytoplasmic AAA-ATPase (PilT). Retraction results in forces of 100 pN per single motor [36], making the T4P machinery the most powerful molecular machine studied to date. It is this combined action of assembly, surface adherence and then retraction that enables cells to move in a jerky fashion, referred to as ‘twitching’ [37,38].

Whilst there have been numerous structural studies of the bacterial flagellar motor and associated motility [39,40], very little was known about the in situ structure and supramolecular assembly of a twitching machinery until recently. Using cryo-ET of whole bacterial cells, Gold and colleagues investigated the intact T4P machinery of *T. thermophilus in situ* [41**] (Figures 1 and 2a). In order to render the cells thin enough for reliable particle identification and StA, the sample was treated with EDTA, which revealed large transmembrane T4P complexes bridging both membranes at the cell poles [41**]. By analysis of different conformational states, the structure of the machinery was determined in the closed (pilus retracted) and open (pilus assembled) conformation. Evidently, the PilQ secretin is extremely dynamic and undergoes a large 30 Å shift of periplasmic N-terminal domains and widening of the central pore on channel opening to make way for the assembled pilus (Figure 2a). The diameter of the pilus was narrower than previously reported [33] suggesting that different forms may exist. Additional protein density was visible in the cytoplasm in the open state of the complex only. It follows that the ATPases (PilF/PilT) required for pilus assembly and retraction may contribute to this density.

A prerequisite for complete understanding of the assembly and dynamics of the various protein components in the T4P machinery is the assignment of specific densities within the electron density maps, and determination of complete high-resolution structural information. X-ray structures are available for a few proteins, their homologues or small fragments [42–49]. Identification of these proteins within the supercomplex will require a combination of genetic manipulation and/or protein labeling [50,51] with functional studies and further structural determination. Fitting high-resolution structures into their corresponding density in cryo-ET maps will provide important future insights into the composition and functionality of these highly dynamic and versatile molecular machines.

**Bacterial type III secretion systems**

The type III secretion system (T3SS or injectisome) is a machinery that is commonly used to deliver bacterial effectors from the cytoplasm directly into the cytosol of a target cell [52,53] in an ATP-dependent manner. The T3SS is composed of a basal body spanning the bacterial periplasm (StcC, SctD and SctJ according to the unified nomenclature [54]), a hollow needle extending into the extracellular space (SctF) and a cytoplasmic part containing the T3S export apparatus (SctV, SctR, SctS and SctT), a sorting platform (SctQ, SctL and SctK) [55] and an ATPase SctN (Figure 2b). The cytoplasmic part of the T3SS is evolutionarily related to its counterpart from the bacterial flagellar motor, responsible for assembly and secretion [56]. The ATPase SctN is structurally similar to both F-type and V-type ATPases [57] and is thought to be responsible for detachment of chaperones and unfolding the exported substrate [58]. Intact injectisomes were recently observed by cryo-ET inside native *Yersinia enterocolitica* [59], *Salmonella enterica* [60] and *Shigella flexneri* [59,61**]. In all cases the best resolution was obtained with bacterial micelles, which were shown to have active T3SS in *Salmonella* [62]. The periplasmic component of *Yersinia* injectisomes varied in length by ~20%, suggesting that its flexibility may be important for the secretion process. The 27 Å resolution structure of the *S. flexneri* injectisome by Hu and colleagues [61**] allowed unambiguous localization of the basal body. The proteins of the sorting platform that were poorly resolved in previous structures were clearly visible: hexameric SctN is axially aligned to the C-ring, composed of SctQ, by six radial spokes of SctL [61**]. Such positioning allows transport of partially unfolded proteins from SctN to the pore in the export apparatus protein SctV [61**] and further through the needle to the host cell.

Recent fluorescent studies in *Yersinia* demonstrating turnover of SctQ suggest that the T3SS is a highly dynamic.
Cross section models of dual-membrane-spanning bacterial protein machines in the inactive and active state. (a) The T4P machinery, based on the in situ structure from *T. thermophilus*. The PilQ secretin (blue) is shown embedded in the outer membrane and extends to the centre of the periplasm and peptidoglycan layer (grey line). Two gates enclose a periplasmic vestibule, which open on pilus assembly (green). Additional protein densities, of which the identities are currently unclear, are seen to associate with PilQ to build the entire machinery (C1, P1, P2) as previously assigned [41]. Protein densities P1 and P2 (yellow) form rings, which are likely attached to PilQ by linker domains (orange) that are not resolved in the sub-tomogram average. Protein C1 (grey) may be comprised of the soluble part of inner membrane anchored proteins M1 (dark blue), such as PilC, PilN or PilO, which are thought to form part of the inner membrane assembly platform for pilus biogenesis [2,44,45]. A large cytoplasmic protein density C2 (red) is visible in the open state only. Candidate proteins include PilM and the ATPases PilF and PilT1/2. (b) Model of the T3SS based on the in situ structure of the complex from *Chlamydia trachomatis*: the basal body consisting of SctC (blue), SctD and SctJ (green) contracts during secretion. The needle, composed of SctF (green) extends into the extracellular space, reaching the host cell membrane with the needle tip (top, orange). The extracellular plate (red) is *Chlamydia* – specific and not observed in other studies of T3SS; peptidoglycan layer (grey line) is observed in most of T3SS-containing bacteria, but not in C. trachomatis. Upon activation, the cytoplasmic density of the sorting platform, which aligns the SctN ATPase and the export apparatus, becomes more ordered. (c) Model for T6SS secretion. Left: T6SS in a resting state contains a periplasm-spanning part comprised of TssJ, TssL and TssM [76], an inner tube made of the Hcp protein containing effectors (yellow) and an extended sheath made of VipA/VipB (purple). Prompted by an as yet unknown trigger, the sheath contracts to release the Hcp tube into the host cell. The following steps are disassembly of the contracted sheath by a CtpV ATPase and re-building of a new T6SS.

Machine [63]. This turnover rate was faster for secreting injectisomes and required the ATPase SctN for stabilization in the non-secreting state [63]. Finally, upon activation of secretion, assembly of new injectisomes occurs close to the existing ones [64]. Nans and colleagues visualized large conformational changes associated with activation of injectisomes of *Chlamydia trachomatis* during host membrane contact with the cultured human cells [65**]. The periplasmic basal body contracted by 4–5 nm leading to an extension of the needle outside the bacteria reaching the host-cell membrane (*Figure 2b*). In the secreting state the sorting platform was significantly better defined on the cytoplasmic side, suggesting that ordering of the sorting platform is related to the ‘pump-like’ contraction of the basal body.

A combination of higher-resolution structures in both secreting and non-secreting states with computational analysis [59,66,67] and genetic manipulation will advance mechanistic understanding of injectisomes and
how secretion is regulated. However, there is as yet no obvious model system. The injectisome from *S. flexneri* has already been visualized in direct contact with a red blood cell [61**] and its proteins are easy to modify, providing an opportunity to genetically dissect the machinery and mechanism. Data collection for structural analysis will be challenging due to the limited number of injectisomes per *S. flexneri* minicell. *Chlamydia* on the other hand, whilst more difficult to modify genetically, are naturally small, contain numerous injectisomes and may be imaged inside thin human cells [65**,68]. In the future it may be possible to analyze structural changes associated to T3S *in situ* based on a combination of model systems.

**Bacterial type VI secretion systems**

Pathogenic bacteria, such as *Vibrio cholerae* and *P. aeruginosa*, use the T6SS to kill both eukaryotic and prokaryotic cells [69]. 25% of all Gram-negative bacteria have at least one T6SS gene cluster [70]. Based on remote sequence similarity to the tailed bacteriophage T4 [71], the T6SS is described as an inverted phage located inside the bacterial cytoplasm (Figure 2c). It contains a baseplate spanning the two bacterial membranes, followed by a long inner tube covered by an outer sheath. Analogous to phages, the energy for a single secretion `shot' has been suggested to be stored in the conformational energy of the sheath. An unknown trigger causes the energy to be converted into mechanical motion [71]. Basler and colleagues revealed dynamics of the *V. cholerae* T6SS by combining fluorescence microscopy and cryo-ET. The sheath polymerizes at the cytoplasmic membrane, forming ~1 μm filled tubes. Upon contraction, the length of the sheath decreases roughly twofold, the radius increases and the tube appears empty, suggesting that the contents of the inner tube are released to the outside of the cell [72].

Structures of entire T6SS *in situ* are not yet available, due to the large thickness of cells that typically contain this system. However, important structural insights have been gained by single-particle cryo-EM. Atomic structures of native contracted sheathes purified from *V. cholerae* [73*] and *Francisella tularensis* [74*] demonstrated a high degree of similarity between the inner domains of the T6SS and bacteriophage sheaths, while the outer domains were more divergent. The interaction between sheath subunits is organized in a 6-start helix, by a network of tightly interacting beta strands; in both cases this network was shown to be critical for assembly and contraction of the T6SS. Using cryo-EM, Ge *et al.* [75] revealed the contraction mechanism of R-type pyocins, which share high similarity to the inner domains of the both T6SS and phage sheaths; this data may be used to model the conformational changes occurring during sheath contraction. Remaining questions include a mechanistic understanding of effector packing into the T6SS, details of selective recycling of contracted sheaths and conformational changes occurring in the periplasmic baseplate related to effector secretion. Finally, the membrane-spanning core complex composed of TssJ, TssL, and TssM from pathogenic *E. coli* revealed by negative stain EM was shown to be five-fold symmetric [76], raising the question of how it can be attached to a sheath with C6 symmetry. *In situ* structural approaches may shed light on these questions, however, again this depends on a good model system. The classical model systems *V. cholerae* and *P. aeruginosa* are thicker than 500–700 nm, limiting high-resolution cryo-ET.

In a recent SfA structure of the *Myxococcus xanthus* type IVa pilus machinery [77], the locations of individual proteins were identified in the complex, leading to a working model for mechanism of action.

**Future directions**

*In situ* structural analysis is now one of the key tools to study the structure and function of large membrane complexes in a close-to-native state. Combining this approach with high-resolution structures that result from the wider use of advanced instrumentation and genetic and biochemical manipulations such as gene knock-outs [5] and electron-density labeling [50,51] will aid in the identification of individual components in density maps. In particular, structural analysis in defined functional states will help to understand the conformational changes associated with function. Supportive biochemical functional studies, fluorescence imaging and molecular simulations should then enable a complete mechanistic understanding of action. We describe how observation of large multimeric complexes *in situ* is extremely informative and may promote application of similar methodology to study other dynamic membrane protein systems.

**Note added in proof**

In a recent SfA structure of the *Myxococcus xanthus* type IVa pilus machinery [77], the locations of individual proteins were identified in the complex, leading to a working model for mechanism of action.

**Conflict of interest statement**

Nothing declared.

**Acknowledgements**

We thank Werner Kühbländer for support and helpful discussions, and Paolo Lastrico for help with graphics. We gratefully acknowledge Beate Averhoff, Marek Basler, Andrea Nans and Bertram Daum for their comments on this manuscript. This work was supported by the Max Planck Society and a Sofja Kovalevskaja Award from the Humboldt Foundation (to MK).

**References and recommended reading**

Papers of particular interest, published within the period of review, have been highlighted as:

- of special interest
- of outstanding interest


23. Pymol. The PyMOL Molecular Graphics System, Version 1.7.4 Schrödinger, LLC.


In this study the authors determine the first in situ structure of a type IV assembly machinery by cryo-ET and STA. Structures were determined for the closed (pilus retracted) and open (pilus assembled) state, which revealed a large conformational change associated with channel opening.

42. Trindade MB, Job V, Contreras-Martel C, Pelicic V, Dessen A: Structure of a widely conserved type IV pilus biogenesis factor


The authors used cryo-ET and STA to visualize the structure of the T3SS in fine detail, allowing unambiguous positioning of the basal body in the membrane and visualization of the T3 export apparatus inside the native assembled complex.