Structure and Electrophysiological Properties of the YscC Secretin from the Type III Secretion System of Yersinia enterocolitica

Peter Burghout,1 Ria van Boxtel,1 Patrick Van Gelder,2 Philippe Ringler,3 Shirley A. Müller,3 Jan Tommassen,1,* and Margot Koster1

Department of Molecular Microbiology and Institute of Biomembranes, Utrecht University, 3584 CH Utrecht, The Netherlands1; Department of Ultrastructure, Flemish Interuniversity Institute of Biotechnology, Free University Brussels, 1050 Brussels, Belgium2; and Maurice E. Müller Institute, Biozentrum, University of Basel, CH-4056 Basel, Switzerland3

Received 23 December 2003/Accepted 5 April 2004

YscC is the integral outer membrane component of the type III protein secretion machinery of Yersinia enterocolitica and belongs to the family of secretins. This group of proteins forms stable ring-like oligomers in the outer membrane, which are thought to function as transport channels for macromolecules. The YscC oligomer was purified after solubilization from the membrane with a nonionic detergent. Sodium dodecyl sulfate did not dissociate the oligomer, but it caused a change in electrophoretic mobility and an increase in protease susceptibility, indicating partial denaturation of the subunits within the oligomer. The mass of the homo-oligomer, as determined by scanning transmission electron microscopy, was approximately 1 MDa. Analysis of the angular power spectrum from averaged top views of negatively stained YscC oligomers revealed a 13-fold angular order, suggesting that the oligomer consists of 13 subunits. Reconstituted in planar lipid bilayers, the YscC oligomer displayed a constant voltage-independent conductance of approximately 3 nS, thus forming a stable pore. However, in vivo, the expression of YscC did not lead to an increased permeability of the outer membrane. Electron microscopy revealed that the YscC oligomer is composed of three domains, two stacked rings attached to a conical domain. This structure is consistent with the notion that the secretin forms the upper part of the basal body of the needle structure of the type III secreton.

The outer membrane of gram-negative bacteria not only protects the cell against harmful compounds in the extracellular environment, such as antibiotics, detergents, and digestive enzymes, but also forms a barrier to the secretion of proteins and uptake of nutrients. The passage of small hydrophilic molecules across the outer membrane is facilitated by the presence of a large number of proteinaceous channels, formed by general and substrate-specific porins and TonB-dependent receptors (31, 41). Gram-negative bacteria have evolved several specialized pathways for the secretion of proteins. These pathways all require one or more integral outer membrane proteins, which in the cases of the type II and III protein secretion systems are related and belong to the family of secretins (19). Other members of this family are involved in the assembly of type IV pili or in the secretion of filamentous phages.

The best-studied secretins with respect to structure and channel formation are XcpQ of Pseudomonas aeruginosa and PulD of Klebsiella oxytoca, both components of type II secretion pathways, the filamentous phage protein pIV, and the PilQ protein of Neisseria meningitidis that is involved in the assembly of type IV pili. Secretins exist in the outer membrane as stable oligomeric complexes that have been purified and visualized by electron microscopy (EM). Top views of the oligomers show them to be large ring-like structures with an internal cavity that might function as an export channel (2, 4, 7, 35, 42). The fact that the PilQ protein of Neisseria gonorrhoeae is essential for a periplasmically assembled pilus to traverse the outer membrane (55) is consistent with this notion. Further, on some micrographs, the internal cavity of the purified secretin oligomers is occluded (4, 8, 43, 44). Side views of the oligomers reveal cylindrical structures composed of at least two stacked ring-like regions (35, 43).

Purified pIV (37), XcpQ (4), and PulD (42) oligomers have been shown to display pore-forming activity when inserted in phospholipid bilayers. The conductance of the channels was voltage dependent, suggesting that the secretins change their conformation in response to the transmembrane potential. The level of conductance was very irregular, especially for XcpQ and PulD, which also indicates that the channels are gated. In contrast, in vivo expression of secretins seems to have no effect on the permeability of the outer membrane (4, 37), which makes it likely that in the cell the secretin is closed. How the channel is opened to facilitate export is unknown, but probably interactions with substrates or other components of its cognate system induce a conformational change.

The homology between the members of the secretin family is particularly marked in their C-terminal halves. Deletion mutagenesis (20) and limited proteolysis (4, 43) have shown that this domain participates in oligomerization and pore formation, while the N terminus is probably involved in system-specific interactions (12, 49). A small nonconserved domain C terminal to the conserved part of the protein functions as the binding site for a small lipoprotein in a number of secretins (11, 13, 47, 49). These so-called pilot proteins protect the secretin against proteolysis and mediate its stable insertion into the outer membrane (10, 13, 16, 23, 32, 49). In the Pul systems derived from Yersinia, the YscC secretin fits this role. The C-terminal half of YscC shows homology with the C-terminal half of PulD and the C-terminal half of XcpQ (2, 15). However, a C-terminal domain of secretins is not always protease resistant (43, 44).
system of *K. oxytoca*, the PulS pilot protein remains associated with the secretin after its insertion into the membrane (42).

Secretins have been shown to be major components of type III secretion systems (33, 48, 51). These systems, which include approximately 20 proteins (9, 28), play a central role in the interaction between gram-negative bacteria and their eukaryotic hosts by allowing the bacteria to inject proteins directly into the cytosol of eukaryotic target cells (28). Secretins of two type III systems have been purified and visualized by EM and are ring-shaped oligomeric molecules (10, 32). However, these purified oligomers have not been analyzed further. In this study, we describe the pore formation and EM analysis of the YscC secretin of the type III secretion system of *Yersinia enterocolitica*.

### MATERIALS AND METHODS

**Bacterial strains and growth conditions.** *Escherichia coli* strains DH5α (21), S17-1β (50), and BL21(DE3) (Novagen) were used for routine gene cloning, conjugal transfer of plasmids to *Y. enterocolitica*, and expression of genes under T7 promoter control, respectively. *Y. enterocolitica* KG22703 is a *blc* mutant derivative of strain W22703, in which the gene encoding β-lactamase has been replaced by the *luxAB* genes (30). KG22703(pAA2003) carries the *aph*4-3 cassette in the *ysc* gene creating a nonpolar mutation (32). The pY-cured *E. enterocolitica* strain CE1525 was obtained by growing strain W22703 on calcium-depleted medium at 37°C. Fast-growing colonies were screened, and a strain from which no plasmid could be isolated was selected.

*E. coli* strains were routinely grown at 37°C in a modified Luria-Bertani broth (LB) (52), and *Y. enterocolitica* strains were grown at room temperature in LB supplemented with 0.4% glucose. To induce expression of *yscC* from which no plasmid could be isolated was selected.°

**Transformation of *E. coli*** and **induction of expression**. Plasmids were introduced into *E. coli* by transformation using the CaCl2 procedure (45) and into *Y. enterocolitica* essentially as described previously (45). Plasmids were introduced into *E. coli* BL21 (DE3) carrying pV3D3 that had been grown overnight was diluted 1:20 in LB and grown for 1 h at 37°C. After the addition of IPTG to a final concentration of 1 mM, the bacteria were grown for another 2 h at 37°C. Cells were collected by centrifugation, and the cell pellet was resuspended in a solution consisting of 100 mM NaCl, 1 mM EDTA, and 50 mM Tris-HCl (pH 8.0) and sonicated. The lysate was centrifuged for 25 min at 30,000 × *g* and the pellet into which inclusion bodies were solubilized in 8 M urea–50 mM Tris-HCl (pH 8.0). After centrigugation for 1 h at 100,000 × *g* at 15°C, solubilized *Hsc*-YscW from the supernatant was purified with Ni-nitrilotriacetic acid beads (Qiagen), following the manufacturer’s instructions. A rabbit was injected subcutaneously with 0.15 mg of purified protein in Freund’s complete adjuvant. Seven booster doses of the same protein preparation in Freund’s incomplete adjuvant were given, one every 4 weeks.

**Antibody preparation against YscW.** For overproduction and purification of *Hsc*-YscW, a culture of *E. coli* BL21 (DE3) carrying pV3D3 that had been grown overnight was diluted 1:20 in LB and grown for 1 h at 37°C. After the addition of IPTG to a final concentration of 1 mM, the bacteria were grown for another 2 h at 37°C. Cells were collected by centrifugation, and the cell pellet was resuspended in a solution consisting of 100 mM NaCl, 1 mM EDTA, and 50 mM Tris-HCl (pH 8.0) and sonicated. The lysate was centrifuged for 25 min at 30,000 × *g* and the pellet into which inclusion bodies were solubilized in 8 M urea–50 mM Tris-HCl (pH 8.0). After centrigugation for 1 h at 100,000 × *g* at 15°C, solubilized *Hsc*-YscW from the supernatant was purified with Ni-nitrilotriacetic acid beads (Qiagen), following the manufacturer’s instructions. A rabbit was injected subcutaneously with 0.15 mg of purified protein in Freund’s complete adjuvant. Seven booster doses of the same protein preparation in Freund’s incomplete adjuvant were given, one every 4 weeks.

**SDS-PAGE and immunoblotting.** Proteins were routinely separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) (34) with 0.2% SDS in the running gel. For seminative SDS-PAGE, the loading buffer contained no β-mercaptoethanol and only 0.1% SDS instead of 2% SDS, the gels contained no SDS, and electrophoresis was performed at 4°C. The detergent-solubilized YscC secretin was dialyzed with proteinase K (22). Alternatively, the YscG secretor was dissociated by treatment with trichloroacetic acid (TCA) and trifluoroacetic acid (TFA). One volume of cell culture was mixed with 1 volume of 10% TCA, and the mixture was incubated for 20 min on ice. Insoluble material was collected by centrifugation at 20,000 × *g* for 20 min at 4°C, washed twice with 2 volumes of ice-cold acetone, and dried. To increase the dissociation efficiency, 30 μl of TFA was added to the pellet, which was then immediately evaporated by a stream of air. Protein bands were stained either with Coomassie Brilliant blue or with silver (38). Alternatively, proteins were transferred onto nitrocellulose membranes by semidyeblotting. After incubation with primary antibodies and with horseradish peroxidase-coupled anti-rabbit immunoglobulin G (diluted 1:3,000) (BioSource), the immunoblots were developed by enhanced chemiluminescence (ECL). The primary antibodies used were 1:1,000 dilutions of rabbit antisera directed against a synthetic YscC peptide (32) or YscW.

**Stability assay for the purified YscC oligomer.** The stability of the purified YscC oligomer in the presence of detergents was evaluated by incubating the oligomer for 1 h at 40°C in 2% Enulion, octyl-polyoxyethylene (OPOE) (Alexis), sulfobetaine 12 (SB12) (Sigma), SDS (J. T. Baker), Triton X-100 (Serva), or Zwittergent 3-14 (Calbiochem) in chromatography buffer. Similarly, the stability of the oligomer in the presence of 20 mM dithiothreitol (DTT) was evaluated. After the incubations, the samples were placed on ice. The loading buffer for seminative gel electrophoresis was added just before the gel was loaded.

**Protease accessibility of the YscC oligomer.** Purified YscC oligomer was diluted 1:10 in chromatography buffer containing 2% SDS, and the solution was either boiled for 10 min or left untreated. The samples were incubated with different concentrations of protease K for 10 min at room temperature. Subsequent SDS-PAGE loading buffer was added, and the samples were heated for 10 min, and proteolytic degradation was immediately evaluated by SDS-PAGE.

**Mass determination by STEM.** The protein stock was diluted 1:2 with chromatography buffer without Enulion, and 6-μl aliquots were adsorbed for 60 s to glow-discharged scanning transmission electron microscopy (STEM) films, which are thin carbon films that span a thick fenestrated carbon layer covering 200-mesh/in., gold-plated copper grids. The grids were then blotted, washed on 9 drops of water, and freezedried. Ten μl of gradient buffer 0.3% Eluent, 250 mM NaCl, protein over night in the microscope. Tobacco mosaic virus (TMV) particles served as mass standard. These particles were similarly adsorbed (45 s) to separate
microscopy grids, washed on 4 droplets of 10 mM ammonium acetate, and air dried.

A STEM HB-5 vacuum generator interfaced to a modular computer system (Tietz Video and Image Processing Systems GmbH, Gauting, Germany) was employed (39). Series of dark-field images (512 by 512 pixels) were recorded from the unstained samples at an accelerating voltage of 80 kV and a nominal magnification of \( \times 200,000 \). The recording dose was \( 345 \pm 45 \) electrons/nm\(^2\). The digital images were evaluated using the program package IMPSYS (39). Accordingly, the projections were selected in circular boxes, and the total scattering of each region was calculated. The background scattering of the carbon support film was then subtracted, and the mass was calculated. The results were scaled according to the mass measured for TMV and corrected for beam-induced mass loss on the basis of the behavior of proteins in a similar mass range (40; unpublished data). The mass values were then displayed in histograms and described by Gauss curves.

**Negative stain EM.** The grids were prepared as described above for mass measurement except that after the washing step, they were negatively stained on 2 droplets of 2% uranyl acetate (pH 4.5), blotted, and air dried. The grids were examined in the STEM, and digital, dark-field images (512 by 512 pixels) were recorded at an accelerating voltage of 100 kV and a nominal magnification of \( \times 500,000 \). Bright-field images of larger regions of the same grids were subsequently recorded on photographic film using an Hitachi 7000 TEM at an accelerating voltage of 100 kV and a magnification of \( \times 50,000 \). The negatives were digitalized using a Primescan D7100 scanner at a sampling step size of 0.4 nm/pixel.

Projections were manually selected from the STEM images and examined; averaging procedures were not employed. Top- and side-view projections of the complexes were also manually selected from the TEM images and placed in galleries using the EMAN (36) and SEMPER image-processing packages (46). The galleries were analyzed individually. The side-view projections were angularly and translationally aligned to arbitrarily selected reference particles using the SEMPER programs. Projections with the specified correlation factor (\( \geq 0.3 \) and \( \geq 0.5 \) for side views of the single and double complexes, respectively) to the reference were subsequently superimposed and averaged. The SPIDER image-processing package (17) was employed to align the top-view projections without the use of a reference. Repeated alignment cycles yielded several class averages. Their rotational power spectra were determined using the SEMPER programs.

**Planar lipid bilayer experiments.** Planar lipid bilayer experiments were performed essentially as described previously (54) with 5 mM CaCl\(_2\) in the electrode buffer. Briefly, after formation of the lipid bilayer, 50 ng of YscC was added to the aqueous subphase. After a potential of 100 mV was applied over the membrane, insertions of the protein into the bilayer were monitored as stepwise increments of the current. To study the effect of calcium on channel activity, the electrolyte buffer was prepared without CaCl\(_2\). Voltage-ramp experiments were performed by applying an increasing potential from 0 to \( +200 \) or \( -200 \) mV over a time span of 100 s. Zero-current membrane potentials were determined after establishing a 10-fold KC gradient (1 M in the cis cell and 100 mM in the trans cell) across the bilayer.

**Uptake of chromogenic \( \beta \)-lactam antibiotics.** The pYV-cured \( Y. enterocolitica \) strain CE1525 carrying plasmid pSM3 (containing the yscW gene) with or without plasmid pRS6 (containing the yscW gene) were grown in BHI-OX or BHI-Ca\(^{2+}\), and expression of yscC was induced by the addition of IPTG as described above. Cells were washed and resuspended to an optical density at 600 nm of 0.5 in phosphate-buffered saline (45). After the addition of CENTA \( \beta \)-lactamase substrate (Merck) or nitrocefin (Calbiochem) to a final concentration of 50 \( \mu \)g/mL, the conversion of these substrates by periplasmic \( \beta \)-lactamase was monitored for 1 min by determining the optical density at 400 or 486 nm. Transport across the outer membrane is the rate-limiting step in the degradation of these antibiotics in this assay (55). As a control for possible leakage of periplasmic \( \beta \)-lactamase, the conversion of the \( \beta \)-lactam antibiotics in the supernatant of the cell suspension was measured.

**RESULTS**

**Purification of the YscC oligomer under nondenaturing conditions.** In the protocol for the purification of the YscC oligomer of Koster et al. (32), SDS was used for its extraction from cell envelopes of the \( Y. enterocolitica \) wild-type strain KNG22703. Although the oligomer did not dissociate, it is possible that the protein was partially denatured during the extraction. Therefore, a new purification method was developed in which the nonionic detergent Elugent was used to solubilize the cell membranes instead of SDS. A pYV-cured \( Y. enterocolitica \) strain (the virulence plasmid pYV carries the genes for all the components of the secretion machinery) was used to produce the YscC oligomer. Two plasmids, pSM3km encoding YscC and pRS6 encoding YscW, the pilot protein required for the production of high levels of YscC and proper localization of the YscC oligomer (5, 32), were introduced into the strain. Membrane proteins from this strain were solubilized in 3% Elugent, and YscC was purified using sucrose gradient centrifugation followed by ion-exchange chromatography (see Materials and Methods). A NaCl concentration of at least 100 mM had to be present in the purification buffers to prevent aggregation of the solubilized secretin. The final preparation contained almost exclusively the oligomeric form of YscC (Fig. 1A). Similar to what was reported previously (32), a small amount of the secretin was purified in its 64-kDa monomeric form (Fig. 1A), as confirmed by immunoblot analysis (data not shown). It is possible that unstable assembly intermediates of the YscC oligomer copurified during the procedure. Upon treatment with hot phenol, the oligomer dissociated into the constituent YscC monomers (Fig. 1B). No other proteins were detected, showing purified YscC to be homo-oligomeric. Im-

FIG. 1. Purification of the YscC oligomer with the nonionic detergent Elugent. (A) Protein fractions of the different steps of the purification procedure were analyzed on a 3 to 15% polyacrylamide gradient gel and stained with silver. Samples were adjusted to contain equal amounts of YscC oligomer. Lane 1, protein fraction from whole cells (lane 1); lane 2, protein fraction from cell envelopes; lane 3, soluble protein fraction obtained after membrane extraction with Elugent; lane 4, protein fraction obtained after sucrose gradient centrifugation; lane 5, purified YscC oligomer after ion-exchange chromatography. (B) The purified YscC oligomer was dissociated with hot phenol, loaded on an 11% polyacrylamide gel, and visualized with Coomassie brilliant blue staining. The positions of the molecular mass markers (in kilodaltons) and of the YscC oligomer and monomer are indicated to the left and right of the gels, respectively.
Interestingly, analysis of the purified YscC oligomer revealed two high-molecular-weight oligomers (Fig. 2, lane 1), which were both recognized by antibodies directed against YscC (data not shown) and thus represent two forms of the secretin oligomer. When boiled in the presence of 2% SDS, the more-abundant, faster-migrating form was completely converted into the slower-migrating form (Fig. 2, lane 2). This may be the result of SDS partially unfolding the polypeptides within the oligomer without dissociating it but thereby affecting its electrophoretic mobility. A small proportion of the purified YscC oligomer was always in the slower-migrating form after seminative SDS-PAGE (Fig. 2, lane 1) possibly due to the presence of 0.1% SDS in the sample buffer. The fast-migrating YscC oligomer was also the most prominent form in cell envelopes that were directly analyzed by seminative SDS-PAGE (Fig. 2, lane 3), indicating that this form represents the most native configuration of YscC. These results indicate that the YscC oligomer normally has a densely packed structure, which partially unfolds in the presence of SDS. Thus, even though SDS does not dissociate the YscC oligomer, it appears to strongly affect its structure.

**Modification of the YscC oligomer by temperature and SDS.** Interestingly, analysis of the purified secretin by seminative SDS-PAGE revealed two high-molecular-weight oligomers (Fig. 2, lane 1), which were both recognized by antibodies directed against YscC (data not shown) and thus represent two forms of the secretin oligomer. When boiled in the presence of 2% SDS, the more-abundant, faster-migrating form was completely converted into the slower-migrating form (Fig. 2, lane 2). This may be the result of SDS partially unfolding the polypeptides within the oligomer without dissociating it but thereby affecting its electrophoretic mobility. A small proportion of the purified YscC oligomer was always in the slower-migrating form after seminative SDS-PAGE (Fig. 2, lane 1) possibly due to the presence of 0.1% SDS in the sample buffer. The fast-migrating YscC oligomer was also the most prominent form in cell envelopes that were directly analyzed by seminative SDS-PAGE (Fig. 2, lane 3), indicating that this form represents the most native configuration of YscC. These results indicate that the YscC oligomer normally has a densely packed structure, which partially unfolds in the presence of SDS. Thus, even though SDS does not dissociate the YscC oligomer, it appears to strongly affect its structure.

**Stability of the native YscC oligomer.** The stability of the native form of the YscC oligomer was studied in more detail. In a buffer containing 0.1% Elugent, 100 mM NaCl, 0.1 mM EDTA, and 10 mM Tris-HCl (pH 7.8), the native form of the YscC oligomer remained stable during incubation for 10 min at temperatures up to 60°C (Fig. 3A). After incubation at 80 or 100°C, neither form of the oligomer was detected (Fig. 3A), and there was no concomitant increase in the amount of monomeric YscC observed (data not shown). Therefore, heat modification of the oligomer in this buffer most likely caused aggregation. When the YscC oligomer was incubated in the presence of 2% SDS, conversion of the native YscC oligomer into the slow-migrating form was already observed at temperatures above 20°C, and this conversion was detected at all higher temperatures tested (Fig. 3A). Hence, the presence of SDS promotes partial denaturation at lower temperatures and prevents aggregation of the oligomer at higher temperatures.

The stability of the native YscC oligomer was also determined in other detergents that are often used for the extraction of cell envelope proteins. The native form of the oligomer was still present after incubation for 1 h at 40°C in 2% Elugent, OPOE, SB12, Triton X-100, or Zwittergent 3-14 (Fig. 3B).

The susceptibility of the two forms to proteolytic degradation was studied by incubating the purified YscC oligomer with 2% SDS either at room temperature or at 100°C to obtain the native or nonnative form, respectively, followed by treatment with proteinase K. The nonnative form showed increased sensitivity to the protease compared to the native form (Fig. 4), which substantiates the notion that SDS induces partial unfolding of the YscC oligomer at high temperatures.

The nonconserved C-terminal domain of YscC contains four cysteine residues that form at least one intramolecular disulfide bond (29). To investigate whether this bond helps to stabilize the native oligomeric form of YscC, the purified oligomer was incubated in the presence of 20 mM DTT for 10 min at 40°C. The minor shift in electrophoretic mobility of the monomers obtained after dissociation of these oligomers with TCA and TFA indicates that at least one disulfide bond was broken by the DTT treatment (Fig. 5A). In its reduced state, the oligomeric secretin was still in its native form after incubation at 40°C (Fig. 5B), which implies that the disulfide bond is not essential to retain the YscC oligomer in this conformation.

**Mass analysis of the purified YscC oligomers.** The purified YscC oligomers were adsorbed to thin carbon films and imaged by STEM for mass measurements. Evaluation of the images yielded a histogram displaying a major mass peak and two minor mass peaks (Fig. 6A) and a few higher mass values (data not shown). More than 75% of all the particles had a mass of 985 ± 175 kDa (n = 546; standard error [SE] of ±7 kDa), the overall uncertainty being ±50 kDa considering the SE of the measurement and the 5% calibration uncertainty of the instrument. As shown in the left gallery of Fig. 6B, these particles generally had circular projections ~14 nm in diameter. The following smaller peak on the histogram indicates a

**FIG. 2.** Effects of heat and SDS on the purified YscC oligomer. The seminative 3 to 9% polyacrylamide gradient gel was loaded with purified YscC oligomer (lane 1), purified YscC oligomer that had been incubated for 10 min at 100°C in the presence of 2% SDS (lane 2), and cell envelopes from the pYV-cured Y. enterocolitica strain CE1525 carrying plasmids pSM3km and pRS6. Proteins were visualized by staining with silver. The positions of the native and partially denatured (nonnative) forms of the YscC oligomer are indicated to the right of the gel.

**FIG. 3.** Stability of the native YscC oligomer. (A) Elugent-solubilized YscC oligomer was incubated in the absence or presence of 2.0% SDS for 10 min at the indicated temperatures. (B) Elugent-solubilized YscC oligomer was incubated in the presence of 2% Elugent (e), OPOE (o), SB12 (sb), SDS (s), Triton X-100 (t), or Zwittergent 3-14 (z) for 1 h at 40°C. All samples were loaded on seminative 3 to 9% polyacrylamide gradient gels, subjected to electrophoresis, and stained with silver. The positions of the native and nonnative forms of the YscC oligomer are indicated to the right of the gels.
mass of $1,928 \pm 254$ kDa ($n = 97$; SE of $\pm 26$ kDa) and arises from the association of two oligomers. The corresponding projections were typically both elongated and of uniform mass density, implying that the oligomers were viewed side on (Fig. 6B, center gallery). The third peak indicates a mass of $3,079 \pm 374$ kDa ($n = 53$; SE of $\pm 51$ kDa) and arises from the association of three oligomers to form aggregates with distinctly triangular projections (Fig. 6B, right gallery).

Purely on the basis of the mass of $64.25$ kDa predicted from the sequence of the mature $YscC$ monomer, particles in the main population yielding the peak at $985$ kDa are composed of $15 \pm 1$ $YscC$ monomers. However, as $YscC$ is a membrane protein, detergent will also be present, making this number the upper limit.

**Negative stain EM.** Negatively stained samples of the purified $YscC$ oligomers were first examined by STEM. Most projections were ring-shaped, top views of the particles, which from the mass measurement results and image galleries are single oligomers (Fig. 6). The projections have outer and inner diameters of $\sim 14$ and $\sim 4.5$ nm, respectively (Fig. 7A). There was a considerable tendency to aggregate; large rosette-shaped aggregates were frequently detected (Fig. 7B). Various side views of the oligomer were also observed, and again there was a high tendency for aggregation. The overall shape of these projections was rectangular (Fig. 7C), star-like (Fig. 7D), or square (Fig. 7E). The rectangular structures have a total length of $\sim 23.5$ nm, whereas the star-like and square projections are composed of three or four $\sim 12$-nm-long units, respectively. In accordance with the mass measurements (Fig. 6), these dimensions suggest that the rectangular projections arise from dimers formed by head-to-head or tail-to-tail interactions and that the star-like structures are trimers. Further, they imply that the single $YscC$ oligomer is approximately $12$ nm long and $14$ nm wide. The square projections may arise from both tetramers and pentamers, as a central density with dimensions
consistent with a top view of the oligomer was sometimes observed (Fig. 7E).

A few side views of the single oligomers \(n = 16\) and further rectangular side views from dimers \(n = 16\) were detected when large areas of the grids were scanned by TEM. The average dimensions of the projections (Fig. 7F, top micrographs) are 10 to 11 and 20 to 23.5 nm long, respectively, and ~14 nm wide. These dimensions agree well with the STEM results and confirm the proposed assignments. Both the single oligomer and dimer side views reveal internal stain-filled regions up to 6 nm in cross-section. The reference-free alignment and classification of 81 top-view projections selected from the TEM images yielded the average projection shown in Fig. 7F (bottom right micrograph). The average projection was calculated from 58 projections, i.e., 72% of those selected, and has outer and inner diameters of ~14 and ~4 nm, respectively. Further, the calculated angular power spectrum displays a strong 13-fold angular harmonic (Fig. 7F, graph).

Both the individual STEM and the average TEM side-view projections reveal that the oligomeric secretin is composed of three domains as indicated in the cartoon (Fig. 7G): (i) a conical domain that shows some variation in form as illustrated in Fig. 7C, (ii) a lower ring, and (iii) a less prominent upper ring that is located at the interface of the dimer.

**Channel formation of YscC in lipid bilayers.** The pore-forming activity of the purified secretin was investigated in planar lipid bilayer experiments. Channel activity with conductance steps of three different size classes were detected at a potential of 100 mV (Fig. 8A and B). Interestingly, only openings and no closings were recorded. Two types of openings, having conductance increments of approximately 2.6 and 3.0 nS, were only slightly different from each other. The third type of opening had a conductance increment of approximately 0.5 nS and was often recorded after a 2.6-nS transition. This result suggests that the YscC oligomer has two conductance states and can undergo a conformational change, leading to the transition from the lower to higher conductance level. To determine whether this conformational change is similar to the one induced by SDS treatment, the purified YscC oligomer was incubated for 10 min in the presence of 2% SDS at 95°C prior to its addition to the aqueous subphase of the planar lipid bilayer system. The SDS-treated oligomer did not show any channel activity, suggesting that either the channel is blocked...
FIG. 8. Pore activity of the YscC oligomer. (A) Channel recording of multiple insertions of the purified YscC oligomer in a planar lipid bilayer at a membrane potential of 100 mV. The sizes of the transitions are indicated. (B) Amplitude histogram of channel openings ($n = 204$) at 100 mV. (C) Voltage-ramp analysis of multiple YscC channels from 0 to 200 mV and 0 to $-200$ mV over a total time span of 200 s.

In vivo pore activity. The uptake of chromogenic $\beta$-lactam antibiotics by intact cells was used to study the pore-forming activity of the YscC oligomer in vivo. In this assay, the diffusion of the antibiotics across the outer membrane is the rate-limiting step in their degradation by periplasmic $\beta$-lactamase. Simultaneous expression of yscC and yscW from pSM3 and pRS6, respectively, in the pYV-cured Y. enterocolitica strain CE1525 did not increase the rate of degradation of the antibiotics (data not shown). Also, when these cells were grown in calcium-depleted medium, the rate of degradation of the antibiotics was not affected (data not shown). Apparently, the permeability of the outer membrane did not change upon yscC expression, suggesting that the YscC oligomer was closed. Hence, in vivo, the secretion channel formed by YscC remains closed in the absence of the other components of the type III secretion system.

DISCUSSION

Here, we report structural and functional data for the YscC secretin of the type III protein secretion system of Y. enterocolitica. The use of a nonionic detergent for solubilization allowed the YscC secretin to be purified in its oligomeric form. Analysis of the oligomer by seminative SDS-PAGE always revealed two bands (Fig. 2), both recognized by YscC antibodies. The faster-migrating form was completely converted into the slower-migrating one upon incubation in SDS at temperatures above 20°C. In the absence of SDS, the faster-migrating form was stable at temperatures up to 60°C. The drastic shift in electrophoretic mobility may be explained by the denaturation of a normally tightly folded YscC domain. This finding that unfolding affects the electrophoretic behavior has been observed for other outer membrane proteins (25) and is often used to monitor the refolding of outer membrane proteins in vitro (15). Moreover, the shift in mobility was accompanied by an increase in protease sensitivity, which is also indicative of a change in the folding state of the YscC oligomer. Thus, although the oligomer does not dissociate in SDS, its structure appears to be drastically changed. Importantly, the SDS sensitivity of the YscC oligomer allowed us to monitor the conservation of the folded state during the purification procedure. Since the highly conserved C-terminal halves of secretins are involved in oligomerization, it is probably the N-terminal part of YscC that unfolds in response to heat and SDS. We did not detect a similar SDS-induced change in the electrophoretic mobility of the P. aeruginosa XcpQ and the N. meningitidis PilQ secretins that are involved in type II secretion and type IV pili assembly, respectively (unpublished observations), which could indicate that the N termini of these secretins already denature under the mild conditions of seminative SDS-PAGE.

As documented by Fig. 1B, the purified secretin is homooligomeric, being formed from YscC alone. Interestingly, in contrast to the pilot protein PulS, which copurifies with the secretin PulD (42), the pilot protein YscW did not copurify with YscC and is therefore not a stable constituent of the oligomeric structure.

A mass of 600 kDa was reported earlier for the YscC complex (32). However, in that case the size of the oligomer was estimated by SDS-PAGE, where it migrated well above the highest molecular mass marker protein, preventing the accumu-
rate determination of its size. The mass of the Elugent-puri-
fi
fi
fi
fi
d, YscC oligomers determined by STEM was signifi-
cantly higher, 985 kDa (Fig. 6). Since YscC is a membrane protein,
this value will include a contribution from the detergent, mak-
ing the indicated number of monomers (15 ± 1) an overesti-
mate. The amount of detergent retained by a membrane pro-
tein depends on both the size of the latter’s hydrophobic domain and the detergent used. STEM mass measurements
made on approximately 250-kDa photosystem II complexes solubilized in the nonionic detergent n-dodecyl-β-
alcohol indicated the association of one 50-kDa detergent micelle (24).
Since the detergent mixture used, Elugent, is of the same type,
several micelles are probably required to solubilize the much
larger YscC oligomer. Accordingly, the latter is most likely
composed of 13 or 14, rather than 15, YscC monomers.

As confirmed by mass measurements of unstained samples,
the single YscC oligomers adsorbed preferentially with their
long axis perpendicular to the air-glow-discharged carbon mi-
croscopy film, resulting in circular (unstained samples; Fig. 6)
or ring-shaped (negatively stained samples; Fig. 7C and F,
bottom right micrograph) projections. Subsequent analysis of
the latter recorded from negatively stained samples by TEM
yielded an average with strong 13-fold angular order (Fig. 7F,
graph). In agreement with the mass measured, this implies the
presence of 13 YscC monomers in the oligomer. While the
presence of 14 subunits has been reported for the pIV secretin
oligomer (37, 44), a dodecameric structure has been proposed for the PilQ (7) and PulD (42) secretins. Thus, despite their
homology, the number of subunits in the oligomer seems to
differ between members of the family of secretins.

The top views of negatively stained, Elugent-puriﬁed YscC
oligomers recorded by STEM (Fig. 7A) and the TEM projec-
tion average (Fig. 7F, bottom right micrograph) show the
secretin to have a stain-accessible central cavity and an outer
diameter of ~14 nm. The latter is considerably smaller than the
20 nm reported earlier for the SDS-puriﬁed YscC oligomer
(32). This suggests that the presence of SDS resulted in con-
formational rearrangements at the outer rim of the oligomer,
which is in agreement with the observed effect of SDS on the
electrophoretic mobility. From the average side-view projec-
tion of the single oligomer (Fig. 7F) and the side-view projec-
tions of two (Fig. 7C and F) or three (Fig. 7D) associated YscC
oligomers, the secretin has a total length of 11 to 12 nm and is
composed of two stacked rings and a conical domain (Fig. 7G)
that can be in a more or less open state as illustrated in Fig. 7C.
Similar conical domains have been found at the ends of other
secretin dimers (7, 35, 43).

Secretins of type III systems, such as YscC, are part of a
much larger structure, the needle complex, which comprises an
external needle and a basal body that traverses both cell mem-
branes and the peptidoglycan layer (14, 33, 48, 51). The latter
domain is formed by two upper and two lower stacked rings
linked by a rod-like structure, which seems to protrude through
the upper rings to form the needle-like extension. The two
stacked rings identified in the side views of the puriﬁed YscC
secretin (Fig. 7C and D; annotated U and L in Fig. 7G) would
appear to correspond to the upper rings, which, depending on
the bacterial host, have an outer diameter of 15 to 20 nm and
a length of approximately 10 nm. This assignment is strength-
ened by the observation that without their needle-like exten-
sions, the basal bodies of isolated secretion complexes aggre-
gate through their upper rings (33, 51), just as the puriﬁed
YscC oligomer tends to aggregate. On the basis of two-dimen-
sional averages of Shigella needle complexes (3), the conical
domain sometimes visible in the side views of the puriﬁed
oligomer (Fig. 7C) probably corresponds to the region between
the two upper and two lower rings of the basal body.

The puriﬁed YscC oligomer inserted into planar lipid bilay-
ers to form stable high-conductance channels. These channels
had two conductance levels, 2.6 and 3.0 nS. Conductance in-
crements of about 0.5 nS were often recorded after a 2.6-nS
transition, indicating that the 2.6-nS channels can convert into
the 3.0-nS form. It remains unknown whether this conversion is
related to the partial unfolding of the oligomer, since the
inability of SDS-treated oligomers to give channel activity could be the result of a failure to insert into the bilayer. The
channel activity of YscC has several unique characteristics
compared to those of three other secretins, XcpQ, PilV, and
PulD. First, the conductance was very regular, and no closings,
only openings, were observed. Other secretins, especially
XcpQ (4), display large variations in their conductance level.
Second, the current passing through the YscC channel in-
creased linearly as the transmembrane potential was increased.
In contrast, the channel activity of the other secretins is non-
linear at higher potentials, which suggests that these pores
change in a voltage-dependent manner (37, 42). Third, the
direction of the potential field had no influence on the con-
ductance of the YscC channel, while the other secretins show
an asymmetrical response (4, 37). The different electrophysio-
logical characteristics of the YscC channel compared to those
of the other secretins may be a consequence of YscC being part of
the type III protein secretion machinery, which is a stable
structure. Once incorporated into the needle complex, the
secretin may remain continuously open, gating occurring at a
different level within the machinery. In other systems, such as
type II secretion, the secretin might be a more dynamic struc-
ture, necessitating the secretin to close when not actively in-
volved in secretion of substrates.

In vivo, the expression of YscC alone did not lead to in-
creased permeability of the outer membrane, suggesting that
the channel was closed. This result is not consistent with the
electrophysiological data discussed above. One explanation
might be that the conformation of the YscC oligomer is slightly
altered during purification. Alternatively, interactions with
components of the cell envelope, such as lipopolysaccharide or
peptidoglycan, might affect the structure of the secretin in vivo.
However, the channel of the puriﬁed YscC oligomer does not
seem to have been in a completely open conformation in the
lipid bilayer either. The theoretical pore diameter can be de-
ferred from the channel activity in planar lipid bilayers, assum-
ing that the channel is an ideal cylinder (26). Since the overall
length of the YscC oligomer is ~12 nm (see above) and the
bulk conductance of the electrolyte is 19.6 S/m, the YscC
channel with a conductance of 3.0 nS would have a pore size of
1.6 nm. This is much smaller than the ~4-nm channel diameter
indicated by electron microscopy (Fig. 7F, bottom right micro-
graph). However, it is clear from both the individual and av-
gage side views (Fig. 7C and F, top micrographs) that the
secretin does not form an ideal cylinder and that the conical
domain restricts the channel diameter. Furthermore, protein
structures in the cavity of the oligomer might be covered by the negative stain and therefore remain undetected. Studies performed on the purified needle complexes of *Salmonella* and *Shigella* suggest that the needle traverses the secretin (3, 33, 51). In order to accommodate the needle, the diameter of the YscC channel should be 6 to 7 nm (27), which is larger than both the electrophysiological and STEM data suggest. Therefore, possibly, the YscC oligomer adopts its final conformation only upon contact with another component of the system, which could be the needle itself. Our attempts to open or widen the pore in vivo and in vitro by calcium depletion were unsuccessful.

Several secretins have been studied so far, and they share many characteristics. However, there are also important differences in the number of subunits forming the oligomer, protein folding, oligomer stability, and electrophysiological behavior of the channel formed. These differences probably reflect the role of the secretin in the various unrelated systems. This makes the study of secretins from different systems essential to obtain insight in the structure and function of this group of important and seemingly versatile transporters.

**ACKNOWLEDGMENTS**

We thank Vesna Olivieri for STEM, Marco Gregorini for help with the TEM image processing, and Andreas Engel for many helpful discussions and critical assessment of the manuscript. We thank R. Diaz-Avalos of the Institute of Molecular Biophysics at Florida State University for kindly providing the TMV particles.

This research was supported in part by grant 700-97012 from the Council for Chemical Sciences of The Netherlands Organization of Scientific Research (CW-NWO), grant HPRN-CT-2000-00075 from the European Community, grant NF 31-59 415.99 from the Swiss National Foundation, and the Maurice E. Muller Foundation of Switzerland.

**REFERENCES**


