An *Acinetobacter* trimeric autotransporter adhesin reaped from cells exhibits its nonspecific stickiness via a highly stable 3D structure

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Trimeric autotransporter adhesins (TAAs), cell surface proteins of Gram-negative bacteria, mediate bacterial adhesion to host cells and extracellular matrix proteins. However, AtaA, a TAA in the nonpathogenic *Acinetobacter* sp. strain Tol 5, shows nonspecific, high adhesiveness to abiotic material surfaces as well as to biotic surfaces. AtaA is a homotrimer of polypeptides comprising 3,630 amino acids and forms long nanofibers; therefore, it is too large and structurally complex to be produced as a recombinant protein. In this study, we isolated AtaA's passenger domain (AtaA PSD), which is translocated to the cell surface through the C-terminal transmembrane domain and exhibits biological functions, using a new method. We introduced a protease recognition site and reaped AtaA nanofibers 225 nm in length from the cell surface through proteolytic cleavage with a specific protease. Biochemical and biophysical analyses of the purified native AtaA PSD revealed that it has a stable structure under alkaline and acidic conditions. Temperatures above 80 °C, which disrupted AtaA's higher-order structure but maintained the full-length AtaA polypeptide, inactivated AtaA's nonspecific adhesiveness, suggesting that the stickiness of AtaA requires its 3D structure. This finding refutes the widespread but vague speculation that large unfolded polypeptides readily stick to various surfaces.

In autotransporters, extracellular proteins in diverse Gram-negative bacteria, the transmembrane domain (TM) hosts the autotransport function, also called type V secretion\(^1\). During this process, the passenger domain (PSD) is exported to the bacterial cell surface through a pore formed by the TM, with the assistance of periplasmic chaperones and the β-barrel assembly machinery\(^2\). Although classical autotransporters comprise one polypeptide with a signal peptide, a PSD at the N-terminus of the mature protein, and a C-terminal TM, trimeric autotransporter adhesins (TAAs) form homotrimeric structures with a common N-terminus–head–stalk–TM–C-terminus architecture after eliminating a signal peptide\(^3\)–\(^5\). The TM, which is formed by a 12-stranded β-barrel (four strands from each polypeptide × three polypeptides) at the outer membrane (OM), anchors the head–stalk domain, the PSD of TAAs, to the OM, after secreting it through the β-barrel pore. The PSDs are assembled from a set of analogous, structurally conserved domains, and the evolutionary rearrangement of these domains has resulted in considerable diversity in the configuration and length of PSDs\(^4\). Most TAAs have been reported to be involved in the initial attachment of bacterial cells to host cells and/or extracellular matrix proteins, as well as in autoagglutination, colonization, biofilm formation, and serum resistance\(^6\)–\(^19\). The TAAs in *Burkholderia* species are currently attracting attention for their ability to drive actin-based motility in host cells by mimicking host actin polymerases\(^20\). The PSDs are responsible for these biological functions.

AtaA, a TAA from the nonpathogenic toluene-degrading bacterium *Acinetobacter* sp. Tol 5\(^21\), has unique adhesive properties that are not shared by other reported TAAs: AtaA mediates strikingly tenacious, nonspecific adhesion of the bacterial cells to abiotic surfaces and has dramatically high autoagglutination ability\(^22\). Through AtaA's adhesive properties, large clumps of the bacterial cells are rapidly immobilized onto various material surfaces.

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surfaces, ranging from hydrophobic plastics to hydrophilic glass and stainless steel, in a manner independent of cell growth\(^2\). A novel and highly efficient method for microbial cell immobilization using AtaA has been developed, and its effectiveness has been demonstrated\(^{24,25}\). AtaA is composed of 3,630 amino acids and is one of the largest TAA family members. To unravel the mechanism of the striking adhesive properties of AtaA, direct characterization of the AtaA molecule using purified protein is indispensable. The effects of other factors, including the surface charge and hydrophobicity of cells and other nanofibers on the cell surface\(^2\), can be eliminated using purified protein. However, it is virtually impossible to produce the entire multi-domain AtaA membrane protein as a recombinant protein in a correctly folded state. Alternatively, extensive experiments would be required to find a suitable method for disruption of the membrane envelope and to select the optimum surfactant for the purification of full-length, native AtaA from Tol 5 cells.

In contrast with TAAs, which are anchored to the OM, many classical autotransporters release their PSDs into the extracellular space by proteolytic cleavage, using either autoproteolytic activity or a membrane-bound protease\(^{29-35}\). Furthermore, heterologous passenger proteins fused to the TMs of classical autotransporters have been released to the external environment by OmpT protease\(^{29,36}\). An OmpT protease recognition site was also inserted at the fusion site of a heterologous epitope with the TM to release the heterologous passenger polypeptide by proteolytic cleavage\(^37\).

Here, we show the first example of applying proteolytic cleavage to a TAA for the isolation of its PSD. A specific protease site was artificially introduced to obtain the full-length extracellular portion of AtaA, which forms nanofibers. Long AtaA PSD nanofibers were successfully reaped directly from the cell surface and used to characterize the AtaA molecule.

Results

Molecular engineering to obtain the AtaA passenger domain from the cell surface. We designed an isolation method that enables reaping of the PSD nanofiber directly from the cell surface by cleaving the AtaA protein at the base of the PSD using a protease with high specificity (Fig. 1). According to the annotation provided by the daTAA program, AtaA has five FGG motifs, which are numbered from the N-terminus of AtaA (FGG_1 to FGG_5); FGG_5 is situated at the N-terminal side adjacent to the TM. A previously determined X-ray crystal structure that included FGG_5 (PDB ID = 3WPA) showed that it has a tri-glycine loop exposed on the molecular surface of the fiber (Fig. 1B)\(^37\). We selected a position between the leucine residue and the tri-glycine residues in this FGG motif to insert the protease cleavage site so as to form an expanded flexible loop. By digesting at this position, 98.5% of the full-length fiber of the trimeric AtaA PSD, which contains 3428 amino acids in each polypeptide, would be reaped from the cell surface. It is 1016 kDa as the trimer based on the calculated molecular weight and has a theoretical isoelectric point of 4.76. For the site-specific cleavage of the PSD, we selected a human rhinovirus 3C (HRV 3C) protease, which recognizes a unique amino acid sequence with high specificity. A nucleotide sequence encoding the protease recognition site and a tri-glycine linker was inserted into the full-length ataA gene at the target site (Fig. 1C). The resultant recombinant DNA encoding the modified AtaA (3CAtaA) was used to transform the ΔataA mutant strain Tol 5 4140\(^{38}\) for its exclusive expression. Production of 3CAtaA protein and its cell-surface display were analyzed by immunoblotting and flow cytometry, respectively. Immunoblotting analysis of the whole-cell lysates demonstrated the production of 3CAtaA protein in Tol 5 4140 (p3CataA) cells with induction of 3CataA gene expression by arabinose (3C\(^+\)) but not in the cells without the induction (3C\(^−\)) (Fig. 2A). Flow cytometric analysis using cells stained with anti-AtaA antiserum revealed that Tol 5 4140 (p3CataA) cells with induction of 3CataA gene expression exhibited a shift of the histogram toward the stronger fluorescent side, implying much 3CAtaA protein displayed on the cell surface, compared with even surface display of AtaA protein on Tol 5 wild-type cells. The functionality of 3CAtaA protein on the cell was examined by adherence assays using polystyrene (PS) and glass plates. As shown in Fig. 2C, cells expressing 3CAtaA (3C\(^+\)) exhibited higher adhesion to both material surfaces than not only cells without expressing 3CAtaA (3C\(^−\)) but also Tol 5 wild-type cells expressing the original AtaA, due to higher expression of 3CAtaA from the plasmid than that of AtaA from the genome. These results verified that 3CAtaA was properly displayed on the cell surface and kept the adheriveness of the original AtaA in spite of the insertion of the HRV 3C recognition site.

Reaping AtaA PSD nanofibers from the cell surface by proteolytic cleavage. Tol 5 4140 transformant cells, which express 3CAtaA, were treated with HRV 3C protease to specifically cleave AtaA present on the cell surface. After the proteolytic digestion, the cells were harvested by centrifugation to examine the cleavage efficiency, and the supernatant was collected for the purification of AtaA PSD released in the reaction buffer. The cells treated with or without the protease were stained with anti-AtaA antiserum and observed under a confocal laser scanning microscope (CLSM). As shown in insets of Fig. 3A,B, treatment of protease drastically reduced the fluorescence of the cells, implying effective cleavage of AtaA displayed on the cells. For more quantitative evaluation of the proteolytic cleavage efficiency, we performed flow cytometric analysis using the same samples as used for CLSM. The protease treatment reduced the percentage of the fluorescent labeled cells displaying 3CAtaA from 73.5% (Fig. 3B) to 17.1% (Fig. 3A). Therefore, the cleavage efficiency was calculated to be 77%.

As shown in Fig. 3C, AtaA PSD could be recovered from the supernatant of the sample treated with the protease but could not be recovered from the control sample that did not undergo protease treatment. However, other proteins were present as contaminants in the supernatant, even though it contained mainly AtaA PSD. For further purification, the proteins in the supernatant were precipitated in the 30% solution of the saturated concentration of ammonium sulfate. The majority of the AtaA PSD was recovered in the precipitate with high purity.
(Fig. 3D). To confirm that the AtaA PSD recovered from the precipitate was dissociated from the OM, a sample was subjected to sucrose density gradient ultracentrifugation, which has been used to purify the OM\textsuperscript{39–41}. Upon fractionation on a 5–50% sucrose density gradient, the purified AtaA PSD was found mainly in the fraction with a density of 1.060 g/mL. Little AtaA was detected in the high-density fractions or in the bottom fraction, where OM should have been distributed (Fig. 3E). Thus, AtaA PSD could be isolated en bloc from the surface of cells displaying 3CAtaA by proteolytic digestion with HRV 3C protease and could be purified simply by ammonium sulfate fractionation. The AtaA PSD protein in the sample after ammonium sulfate fractionation was concentrated by ultrafiltration with a 100-kDa cut-off filter. We ultimately obtained 5 μg of purified AtaA PSD from 20 mL of culture solution.

Morphological characteristics of the purified AtaA passenger domain. Dynamic light scattering (DLS) analysis was performed to confirm that the purified AtaA PSD maintained its trimeric structure. DLS analysis is often used to determine the size and molecular weight of protein. The molecular weight of the AtaA PSD, modeled as a linear polymer, was estimated to be 1071 ± 138 kDa (Fig. 4A, Peak1). This value almost coincides with the value (1016 kDa) calculated using its constituent amino acids. A peak detected at very high molecular mass might reflect the aggregation of AtaA PSD molecules in the sample (Fig. 4A, Peak2).

We next analyzed the morphological characteristics of AtaA PSD using transmission electron microscopy (TEM). Many nanofibrous structures were observed in the fraction of purified AtaA PSD. AtaA PSD nanofibers were uniform in size, at 225 ± 1.7 nm in length and 4 ± 0.4 nm in thickness (mean ± SEM of 10 fibers, chosen at random), were curved in places, and had two globular structures: one at the fiber tip and the other near the...
opposite end (see Supplementary Figs S1 and S2). The former globular structure was predicted to correspond to the globular tip of the AtaA nanofiber, that is, N-terminal Ylhead (Nhead). To confirm the topological direction of the nanofibers observed, we constructed another modified AtaA, 3CFGG2AtaA, in which a HRV 3C protease recognition site was inserted at FGG_2 (see Supplementary Fig. S1). After proteolytic cleavage, a shorter version of the AtaA PSD fiber (AtaA PSD short), which did not include the C-terminal Ylhead (Chead), was obtained. TEM revealed that AtaA PSD short fibers were 180 ± 1.8 nm in length (mean ± SEM of six fibers, chosen at random), had only one globular structure at the fiber tip, and lost the other globular structure (see Supplementary Fig. S1). These data suggest that the globular structures at the fiber tip and near the opposite end corresponded to Nhead and Chead, respectively. The TEM image of a single-molecule fiber of AtaA PSD was then aligned to the predicted domain configuration (Fig. 4B).

Biophysical and biochemical characterization of the AtaA passenger domain. The effects of temperature and pH on the structure of AtaA PSD were examined by measuring the circular dichroism (CD) spectra. CD is commonly used to estimate the secondary structure of protein. This measurement is widely used to determine the stability of the structure of a protein in various physicochemical conditions. After incubation of AtaA PSD dissolved in deionized water (dH₂O) at temperatures 25–98 °C for 5 min, CD spectra were measured at a room temperature (Fig. 5A). The CD spectrum of the native AtaA PSD exhibited two negative peaks at 209 and 222 nm that were attributed to α-helices, and one negative peak at 218 nm that was attributed to β-strands; their contents in this intact AtaA PSD were 31% and 10%, respectively. The AtaA PSD treated at 70 and 80 °C showed CD spectra with only small differences from native AtaA PSD at 25 °C. However, the CD spectrum of AtaA PSD changed drastically after incubation at temperatures over 90 °C; The CD signals became weaker, implying the disruption of secondary structures. The ellipticity of the CD peak at 222 nm gradually decreased with increasing temperature below 70 °C, but it drastically decreased above 80 °C, reflecting the disruption of secondary structures within AtaA PSD (Fig. 5B). Next, to examine the effect of pH, the CD spectra of AtaA PSD dissolved in buffers at different pH or in 0.1 M HCl (pH 1.0) were directly measured. The CD spectra of all samples at pHs from 2 to 12, and even in 0.1 M HCl, almost overlapped, implying that the secondary structure of AtaA PSD is mostly unaffected by pH change in this range (Fig. 5C).
We also examined the morphological changes that occurred in AtaA PSD nanofibers using the same samples used for CD spectroscopy after thermal or pH treatment. TEM revealed no morphological changes resulting from the treatment at 80 °C, relative to the morphology in the control treated at 25 °C. However, the fibrous structure of AtaA PSD shrank, shortened, and was difficult to observe upon thermal treatment at 90 and 98 °C (Fig. 6A), which correspond to temperatures that caused drastic changes in the CD spectrum (Fig. 5A,B). Apparent disruption of the fiber structure was not observed for any pH conditions (Fig. 6B), as expected from the unchanged morphological structure.
CD spectra under acidic and alkaline conditions (Fig. 5C). Even in 0.1 M HCl, AtaA PSD maintained a distinctly visible fiber morphology, as well as its secondary structure, although bent fibers were observed.

**Functional analysis of the AtaA passenger domain.** Obtaining purified AtaA PSD allowed us to directly measure the interaction of the protein molecule with a surface. Quartz crystal microbalance (QCM) is a highly sensitive and label-free mass measurement sensor, which enables quantifying molecules interacting with a quartz crystal sensor chip as a frequency change. We examined the adhesiveness of intact and denatured AtaA PSD using QCM. As shown in Fig. 7A, the resonance frequency rapidly declined with time when AtaA PSD pre-treated at 25 or 70 °C was applied to the QCM sensor. This implies the rapid adhesion of protein molecules to the sensor chip surface. In contrast, the decline in the resonance frequency was faint when bovine serum albumin was applied, emphasizing the stickiness of AtaA PSD. The decline in resonance frequency decreased with increasing treatment temperature for denaturation of the protein molecules and became faint at 98 °C. The bar chart in Fig. 7A shows the frequency change (dF) caused by the adhesion of protein molecules to the QCM sensor when the frequency-shift curve became almost flat (10 min). The absolute value of the dF was almost the same after incubation at 70 °C as it was at 25 °C; it suddenly diminished at 80 °C, gradually diminished with increasing treatment temperature, and became very small at 98 °C, implying that adhesiveness was almost lost in AtaA PSD that was denatured at the high temperature. However, treatment with acidic or alkaline solution did not affect its adhesiveness to the QCM sensor chip at all, and thus the dF remained the same (Fig. 7B). These QCM data suggest that AtaAs high adhesiveness to an abiotic surface requires its structural integrity. To determine whether the heat denaturation that causes the loss of adhesiveness degrades the AtaA polypeptide itself, AtaA PSD samples after thermal treatment were subjected to sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE). This revealed that the molecular sizes of the monomers in these AtaA PSD samples were unchanged, implying no degradation of the polypeptide by thermal treatment (Fig. S3). Therefore, loss of the adhesive function of AtaA PSD must have been caused by disruption of the 3D structure of the protein.

**Discussion**

TAAs have most often been studied using fragmented recombinant proteins expressed in *Escherichia coli* or using bacterial cells expressing intrinsic or foreign TAAs on their cell surfaces. These studies have uncovered the structure and function of TAAs. However, their molecular properties remain unclear because the properties of a fragmented recombinant protein do not always reflect those of the native, full-length molecule. By contrast, proteins isolated using our enzymatic reaping method are guaranteed to be properly folded and assembled because they are actual parts of the native protein secreted by the original carrier cells. As for classical autotransporters that release their PSDs into the extracellular space, PSDs purified from culture supernatants have been used for biochemical characterization or determination of their crystal structures. To the best of our knowledge, however, the artificial insertion of a protease recognition site into an autotransporter has never been employed to obtain PSDs for their characterization. In this study, we first inserted a protease recognition site into a TAA to isolate its PSD by proteolytic cleavage, and then purified the native PSD proteins from the extracellular space. The expected molecular weight from DLS analysis (Fig. 4A) suggested that the AtaA PSD purified by this method appropriately formed a trimer. TEM revealed its nanofibrous structure, which corresponded to the predicted domain configuration having two globular Ylhead domains at the N-terminus of the distal end of the fiber (Nhead) and the C-terminal region near the fiber base (Chead), and a long stalk between the two Ylheads (Fig. 4B). The Chead was not reasonably found on the AtaA PSD short fiber (see Supplementary Fig. S1). These results demonstrate that AtaA PSD was cleaved by HRV 3C protease at the inserted protease recognition site, as designed. The fibrous morphology of the AtaA PSD molecules was disrupted and became difficult to recognize when the AtaA PSD molecules were denatured (Fig. 6A). TEM images of the AtaA PSD obviously changed after incubation of the protein at temperatures over 90 °C, but were unchanged after incubation at temperatures lower than 80 °C. When the temperature exceeded 80 °C, the CD spectrum changed drastically and the ellipticity at 222 nm indicated that the α-helix content decreased sharply, implying drastic denaturation of the protein that resulted in the disruption of its fibrous structure.
In denaturation studies, AtaA PSD showed high structural stability under conditions of extreme pH. It was reported that TAAs maintain their trimeric structure under the standard sample preparation conditions used for SDS-PAGE, and that dissociation into monomers requires boiling in 6–10 M urea solution, 95% formic acid, or extraction with hot phenol\(^9\),\(^{52-54}\). Compared with the other TAAs reported in these articles, AtaA does not seem so resistant to high temperatures. However, it should be noted that the AtaA PSD does not contain a TM, which is important for trimer stability\(^{53}\). Nevertheless, AtaA PSD showed thermal stability that is considerably higher than that of typical proteins. Recently, we determined the X-ray crystal structure of the C-terminal PSD of AtaA, which exhibits the toughness of AtaA nanofibers\(^{27}\). The crystal structure revealed that interchain interactions of the three polypeptides forming the trimer, especially topological chain exchange, contribute to the toughness, which also should give the AtaA PSD high stability. The high stability is also significant for utilization of AtaA in our newly developed method for microbial immobilization\(^{24,25}\). The robustness due to the stability is expected to become an advantage in bioprocesses for industrial use.

QCM measurements revealed that AtaA PSD denatured by thermal treatment at high temperatures considerably decreased its adhesiveness (Fig. 7A). By contrast, alkaline and acid treatments did not affect the adhesiveness of AtaA PSD at all (Fig. 7B). However, the heat denaturation that caused the loss of adhesiveness did not degrade the AtaA polypeptide itself (see Supplementary Fig. S3); only the higher-order structure was disrupted. These

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**Figure 5.** Effect of temperature and pH on secondary structure of AtaA PSD measured by CD spectroscopy. (A) CD spectra of AtaA PSD after thermal treatment at various temperatures for 5 min. The measurement was carried out at a room temperature. (B) Ellipticity at 222 nm as a function of temperature. (C) CD spectra of AtaA PSD in acid or alkaline. The protein samples dissolved in buffer solutions at different pH or in 0.1 M HCl were directly measured by CD spectroscopy.

**Figure 6.** Effect of temperature and pH on the fibrous morphology of AtaA PSD observed by TEM. (A) TEM images of AtaA PSD after thermal treatment at various temperatures for 10 min. After the treatment, the samples were subjected to negative staining for TEM. Scale bars: 100 nm. (B) TEM images of AtaA PSD treated in different pH solutions. After the treatment, the sample was subjected to negative staining for TEM. Scale bars: 100 nm.
results lead us to conclude that the 3D structure is essential for the AtaA molecule to exhibit its unique adhesive character. The uniqueness of the adhesive character of AtaA is its nonspecific interactions, which have no specific ligand. In specific interactions, the stereostructure of the binding site for the ligand of a protein is important. In fact, TAAs require stable trimer formation to exhibit specific adhesion to host cells and tissues52,53. In contrast, nonspecific interactions do not necessarily require such precise stereostructures. Generally speaking, unfolded polypeptides are believed to exhibit nonspecific adhesiveness or a broad binding specificity due to their disordered structures, which allow enlargement of the capture radius and surface exposure of amino acid residues, depending on surface properties. Especially, large polypeptides are thought to be sticky due to multipoint adsorption on an extensive interface55–57. It has been speculated that the huge size of AtaA is responsible for its high adhesiveness. This idea was ruled out by the results showing that AtaA PSD almost completely lost its adhesive character after treatment at 98 °C, despite maintaining the full length of the polypeptide. It should be stressed that maintaining a proper 3D fiber structure is crucially important for the high, nonspecific adhesiveness of the AtaA molecule, and that the adhesiveness cannot be achieved from the large size of the protein alone.

In conclusion, we established a methodology that uses specific protease digestion to prepare the very large PSD of AtaA while retaining its native properties. We observed, for the first time, the molecular shape of purified AtaA PSD and found that its fibrous structure has high stability in strict pH conditions. The 3D structure forming this fiber is necessary for AtaA PSD to exhibit its nonspecific adhesiveness.

Methods
Bacterial strains and culture conditions. The bacterial strains used in this study are listed in Supplementary Table S1. Acinetobacter sp. Tol 5 and its derivative strains were maintained as previously described22. Luria-Bertani (LB) medium was supplemented with ampicillin (500 μg/mL) and gentamicin (10 μg/mL), when required. Arabinose (0.5%, wt/vol) was added to induce expression of 3CataA and 3CFGG2ataA. Production of the expressed proteins was confirmed by SDS-PAGE, followed by immunoblotting using anti-AtaA699–1014 antiserum, as previously described22.
Construction of p3CAtaA and p3CFGG2AtaA. The procedures for the construction of p3CAtaA and p3CFGG2AtaA, which were used to generate AtaA with an inserted HRV 3C protease recognition site, are schematically shown in Supplementary Figs S4 and S5, respectively. Insertion of the sequence encoding the protease recognition site into ataA was performed using overlap-extension PCR. The primers used in this study are listed in Supplementary Table S2.

To construct p3CAtaA, an initial PCR was performed with template pTA2::ataA vector and two primer sets: Bgl II ataA-F/HRV3C ataA-R and HRV3C ataA-F/Xba I ataA-R. A second round of PCR was performed using the DNA fragments amplified by the first PCR as templates using the primer set Bgl II ataA-F/Xba I ataA-R. The amplicon was inserted into the vector pTA2, generating pTA2::3CataA fragment. After DNA sequencing for confirmation, the 3CataA fragment was excised again with Bgl II and Xba I and sub-cloned for the Bgl II-Xba I fragment of pTA2::ataA, generating p3CataA. Thus, the 3CataA gene was constructed, and then sub-cloned into the pAR3 vector22, generating p3CAtaA. Transformation of the ΔataA mutant Tol 5 414038 was carried out by conjugal transfer from E. coli S17-1 ΔataA34, as previously described22.

To construct of p3CFGG2ataA, plasmid pDONR::ataA was digested with Kpn I and the linearized plasmid was re-circularized by self-ligation. The resulting circular plasmid was digested with Xcm I and the linearized plasmid was again re-circularized by self-ligation, generating pDONR::ataA (ΔXcm I, Kpn I). This plasmid was used as the template for overlap PCR with two primer sets: In-Fusion FGG2-F/3CataA-FGG2-R and 3CataA-FGG2-F/ In-Fusion FGG2-R. A second round of PCR was performed using DNA fragments amplified by the first PCR as templates and the primer set In-Fusion FGG2-F/In-Fusion FGG2-R. The amplicon was fused to pDONR::ataA (ΔXcm I, Kpn I), which was linearized by double digestion with Xcm I and Kpn I using the In-Fusion HD cloning Kit (Takara Bio, Shiga, Japan), generating pDONR::3CFGG2ataA (ΔXcm I, Kpn I). Finally, two ataA gene fragments, a Kpn I fragment and an Xcm I fragment, were inserted into the Kpn I and Xcm I restriction sites of pDONR::3CFGG2ataA (ΔXcm I, Kpn I), generating pDONR::3CFGG2ataA. Thus, 3CFGG2ataA was constructed and then sub-cloned into the vector pAR322, generating p3CFGG2ataA.

Obtaining AtaA passenger domain by protease digestion. Bacterial cells were harvested by centrifugation, suspended in 10 mL of 50 mM HEPES buffer (pH 7.0) supplemented with 80 units of HRV 3C protease (Merck Millipore, MA, USA), incubated at 4°C for 2 days in a Proteosave SS 15-mL centrifuge tube (Sumitomo Bakelite, Tokyo, Japan), and then centrifuged at 10,000 × g at 4°C for 10 min. The 10 mL of supernatant was dispensed into new Proteosave SS 1.5-mL microtubes and fractionated by precipitation in the 30% solution of 2-propanol (DMSO) and 90% solution of dH2O. Precipitates were washed several times with dH2O and finally dispersed in particles of different sizes was estimated as a linear polymer by using this relationship embedded in the Zetasizer Nano software (Malvern Instruments).}

Separation of AtaA passenger domain by sucrose density gradient centrifugation. Purified AtaA PSD (100 μL) was layered on a linear sucrose gradient (3 to 50% sucrose in dH2O) prepared using a gradient maker (Hitachi Kouki, Tokyo, Japan) in a 13-mL ultracentrifuge tube (PA13; Hitachi Kouki). This tube was centrifuged at 200,000 × g for 18 h. Gradient fractions of 1 mL were collected from the top of the tube (11 fractions in total) and 30 μL of sample from each aliquot was subjected to SDS-PAGE followed by Coomassie Brilliant Blue (CBB) staining.

Cell analysis. Flow cytometric analysis of cell-surface 3CAtaA was performed as previously described22, with slight modifications. Before fixation with 4% paraformaldehyde solution, pre-fixation was conducted by mixing pre-fixation was conducted by mixing 4% paraformaldehyde and suspension culture in a 1:1 ratio. The cells were washed with dH2O after fixation. FlowJo software (Tomy Digital Biology, Tokyo, Japan) was used to create histograms and to compare the fluorescence intensities among the samples. For CLSM, the same cell samples immunoreacted with the anti-AtaA699–1014 antiserum as the first antibody and Alexa Fluor 488-conjugated anti-rabbit antibody as the secondary antibody were placed onto a slide glass for 15 min. After being washed with NET buffer (150 mM NaCl, 5 mM EDTA, 50 mM Tris-HCl, 0.05% Triton X-100; pH 7.6) twice and dH2O once, cells were observed under a CLSM (FV1000, OLYMPUS, Tokyo, Japan).

Adherence assays were performed as previously described23, with slight modifications. Aliquots (200 μL) of a bacterial cell suspension in BS-N medium27 at an optical density at 660 nm of about 0.5 were placed into a 96-well PS plate (353072; Becton, Dickinson and Company, NJ, USA) or a 96-well glass plate (FB-96; Nippon Sheet Glass Co., Ltd., Tokyo, Japan). After incubation for 2 h at 28°C, cells adhering to the well surface were stained with 1% crystal violet solution for 15 min. The stain was eluted from the cells with 70% ethanol solution, and the A590 of the eluate was measured using a micro plate reader (ARVO X3; PerkinElmer, MA, USA).

Biophysical and biochemical analyses of the AtaA passenger domain. The DLS of purified AtaA PSD in dH2O (0.09 mg/mL) was measured using a Zetasizer Nano ZSP (Malvern Instruments, UK), yielding the particle size distribution per volume data and their average hydrodynamic diameter. The relationship between hydrodynamic diameters and molecular weights is expressed by the following equation:

\[ R_H = (a \times M_w)^b \]  

where \( R_H \) is a hydrodynamic diameter measured by DLS, \( M_w \) is a molecular weight and a, b are coefficients fixed by a molecular type: globular protein, linear polymer, or dendrimer. The molecular weight of AtaA PSD distributed in particles of different sizes was estimated as a linear polymer by using this relationship embedded in Zetasizer Nano software (Malvern Instruments).
To conduct electron microscopy on purified AtaA PSD nanofibers, 10 μL of the sample solution (3–30 μg/mL) and 2 μL of 50 mM KCl solution were placed onto a copper grid with an ultra-high resolution carbon substrate (UHR-C10; Okensoji, Tokyo, Japan). The sample was hydrophilized by plasma treatment using a plasma ion bombarder (PIB-10; Vacuum Device, Ibaraki, Japan) immediately before use. When observing AtaA PSD treated at different pHs or in 0.1 M HCl were used directly for the QCM measurement. After equilibration, 1 ping 20 times into 10 mM KCl solution and three times into dH2O. Negative staining of the proteins on the grid grids were then incubated for 10 min at room temperature to allow adsorption of the proteins and washed by dipping 20 times into 10 mM KCl solution and three times into dH2O. Negative staining of the proteins on the grid was performed with the organo-tungstate compound Nano-W (Nanoprobes, NY, USA). These sample preparation procedures were conducted in a moisture box to avoid desiccation. The specimen was observed using a TEM system (JEM-1400EX; JEOL, Tokyo, Japan) operated at 100 kV.

For denaturation analysis of AtaA PSD at different pHs, buffers of pH 2, 7, and 12 were prepared by mixing 50 mM citric acid solution, 200 mM boric acid solution, and 100 mM sodium phosphate. AtaA PSD in dH2O and the above-mentioned buffers were mixed in a 19:1 ratio (final concentration of AtaA was 0.15 mg/mL) on micro-sampling discs for CD spectroscopy. For the extreme low pH condition, 0.1 M HCl solution was used instead of the buffers. For denaturation by thermal treatment, 0.15 mg/mL of AtaA PSD in dH2O was incubated in a Proteosave SS 1.5-mL microtube at different temperatures for 5 min. The heat-treated samples were then centrifuged for 1 min at 15,000 g and 4 °C to remove debris, and the temperature of the samples was brought back to a room temperature before conducting further studies.

The CD spectrum of purified AtaA PSD was recorded using a J-725 spectropolarimeter (JASCO, Tokyo, Japan). The parameters (wavelength range, scan rate, scan interval, response, and band width) were set at 190–250 nm, 100 nm/min, 0.1 nm, 0.25 s, and 1.0 nm, respectively. For heat-treated samples, 10 μL of the sample (0.15 mg/mL) was applied to the micro-sampling disc after cooling to room temperature. The spectra from the buffer or dH2O were recorded as blanks. The α-helix and β-strand contents of AtaA were calculated using the DichroWeb59,60 program with the CONTINLL algorithm and Protein Reference Data Sets 7.

The adhesiveness of AtaA PSD was measured using a QCM system (AFFINIX Q8; ULVAC, Kanagawa, Japan). A gold-coated electrode (QCM01S-01; ULVAC) was cleaned with piranha solution (H2SO4:30% H2O2 equilibrated with 99 (0.15 mg/mL) was applied to the micro-sampling disc after cooling to room temperature. The spectra from the sample solution (3–30 μg/mL) was applied to the micro-sampling disc after cooling to room temperature. The spectra from the buffer or dH2O were recorded as blanks. The α-helix and β-strand contents of AtaA were calculated using the DichroWeb59,60 program with the CONTINLL algorithm and Protein Reference Data Sets 7.

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22. Ishikawa, M., Nakatani, H. & Hori, K. AtaA, a new member of the trimeric autotransporter adhesins from Acinetobacter sp. Tol 5 mediating high adhesiveness to various abiotic surfaces. PLoS One 7, e48830 (2012).

23. Ishikawa, M., Shigemori, K., Suzuki, A. & Hori, K. Evaluation of adhesiveness of Acinetobacter sp. Tol 5 to abiotic surfaces. J. Biosci. Bioeng. 113, 719–725 (2012).

24. Ishikawa, M., Shigemori, K. & Hori, K. Application of the adhesive bacteriananofiber AtaA to a novel microbial immobilization method for the production of indigo as a model chemical. Biotechnol. Bioeng. 111, 16–24 (2014).

25. Hori, K., Ohara, Y., Ishikawa, M. & Nakatani, H. Effectiveness of direct immobilization of bacterial cells onto material surfaces using the bacteriananofiber protein AtaA. Appl. Microbiol. Biotechnol. 99, 5025–5032 (2015).

26. Szczesny, P. & Lupas, A. Domain annotation of trimeric autotransporter adhesins—dTaTA. Bioinformatics 24, 1251–1256 (2008).

27. Koivai, K., Hartmann, M. D., Linde, D., Lupas, A. N. & Hori, K. Structural basis for toughness and flexibility in the C-terminal domain of a bacterial outer membrane trimeric autotransporter adhesin. J. Biol. Chem. 291, 3705–3724 (2016).

28. Hori, K. et al. Production of peritracheal bacteriananofibers and their proteinaceous components by Acinetobacter sp. Tol 5 cells affected by growth substrates. J. Biosci. Bioeng. 111, 31–36 (2011).

29. Klauser, T., Pohlner, J. & Meyer, T. F. Selective extracellular release of cholera toxin B subunit by Escherichia coli: dissection of Nettetria lga beta-mediated outer membrane transport. EMBO J. 11, 2327–2335 (1992).

30. Desvaux, M., Parham, N. J. & Henderson, I. R. The autotransporter secretion system. Res. Microbiol. 155, 53–60 (2004).

31. Barnard, T. J., Dautin, N., Lukacik, P., Bernstein, H. D. & Chakraborty, T. K. Autotransporter structure reveals intra-barrel cleavage followed by conformational changes. Nature Structural and Molecular Biology 14, 1214–1220 (2007).

32. Henderson, I. R., Navarro-Garcia, F., Desvaux, M., Fernandez, R. C. & Ala`Aldeen, D. Type V protein secretion pathway: the autotransporter story. Microbiol. Mol. Biol. Rev. 68, 692–744 (2004).

33. Scaglione, P. et al. Structural characteristics of the plasmid-encoded toxin from enterogaeggerative Escherichia coli. Biochemistry (Moscow). 47, 9582–9591 (2008).

34. Fink, D. L.,COPE, L. D., Hanssen, E. J. & Geme, J. W. 3rd. The Hemophilus influenzae Hip autotransporter is a chymotrypsin-like serine protease and undergoes autoproteolysis via an intermolecular mechanism. J. Biol. Chem. 276, 39492–39500 (2001).

35. Henderson, I. R., Navarro-Garcia, F. & Nataro, J. P. The great escape: structure and function of the autotransporter proteins. Trends Microbiol. 6, 370–378 (1998).

36. Maurer, J., Jose, I. &Meyer, T. F. Autodisplay: one-component system for efficient surface display and release of soluble recombinant proteins from Escherichia coli. J. Bacteriol. 179, 794–804 (1997).

37. Ruiz-Olvera, P. et al. Display and release of the Plasmodium falciparum circumsporozoite protein using the autotransporter MisL of Salmonella enterica. Plasmid 50, 12–27 (2003).

38. Ishikawa, M. & Hori, K. A new simple method for introducing an unmarked mutation into a large gene of non-competent Gram-negative bacteria by FLI/FRT recombination. BMC Microbiol. 13, 86 (2013).

39. Blanco, D. R. et al. Isolation of the outer membranes from Treponema pallidum and Treponema vincentii. J. Bacteriol. 176, 6088–6099 (1994).

40. Gotoh, N., White, N. J., Chaoawagul, W. & Woods, D. E. Isolation and characterization of the outer-membrane proteins of Burkholderia (Pseudomonas) pseudomallei. Microbiology 140, 797–805 (1994).

41. Tefsen, B., Gerstung, J., Beckers, E., Tommassen, J. & C&cock, H. Lipopolysaccharide transport to the bacterial outer membrane in spheroplasts. J. Biol. Chem. 280, 4504–4509 (2005).

42. Szczesny, P. et al. Structure of the head of the Bartonella adhesin BadA. PLoS Pathog 4, e1000119 (2008).

43. Leo, J. C., Elovaa, H., Brodsly, B., Skurnik, M. & Goldman, A. The Yersinia adhesin YadA binds to a collagenous triple-helical conformation but without sequence specificity. Protein Eng. Des. Sel. 21, 475–484 (2008).

44. Leo, J. C. & Grant, E. T. Characterization of the E. coli lgD-binding protein D suggests a general model for bending and binding in trimeric autotransporter adhesins. Structure 19, 1021–1030 (2011).

45. Leo, J. C. & Goldman, A. The immunoglobulin-binding Eib proteins from Escherichia coli are receptors for IgGFc. Mol. Immunol. 46, 1860–1866 (2009).

46. Riess, T. et al. Bartonella adhesin mediates a proangiogenic host cell response. J. Exp. Med. 200, 1267–1278 (2004).

47. Yu, C., Minta, K. P. & Ruiz, T. Investigation of the three-dimensional architecture of the collagen adhesion EmaA of Aggregatibacter actinomycetemcomitans by electron tomography. J. Bacteriol. 191, 6253–6261 (2009).

48. Mil-Homens, D., Leca, M. I., Fernandes, F., Pinto, S. N. & Fialho, A. M. Characterization of BCAM0224, a multifunctional trimeric autotransporter from the human pathogen Burkholderia cenocepacia. J. Bacteriol. 196, 1968–1979 (2014).

49. Renn, J. P. & Clark, P. L. A conserved stable core structure in the passenger domain beta-helix of autotransporter virulence proteins. Biopolymers 89, 420–427 (2008).

50. Khan, S., Mian, H. S., Sandercock, L. E., Chirgadze, N. Y. & Pai, E. F. Crystal structure of the passenger domain of the Escherichia coli autotransporter EspP. J. Mol. Biol. 413, 985–1000 (2011).

51. Domingo Meza-Aguilar, J. et al. X-ray crystal structure of the passenger domain of plasmid encoded toxin(Pet), an autotransporter enterotoxin from enterogaeggerative Escherichia coli (EAEC). Biochem. Biophys. Res. Commun. 445, 439–444 (2014).

52. Cotter, S. E., Surana, N. K., Grass, S. & St Geme, J. W. 3rd. Trimeric autotransporters require trimerization of the passenger domain for stability and adhesive activity. J. Bacteriol. 188, 5400–5407 (2006).

53. Schutz, M. et al. Trimer stability of YadA is critical for virulence of Yersinia enterocolitica. Infect. Immun. 78, 2677–2690 (2010).

54. Sandt, C. H. & Hill, C. W. Four different genes responsible for nonimmune immunoglobulin-binding activities within a single strain of Escherichia coli. Infect. Immun. 68, 2205–2214 (2000).

55. Gunasekaran, K., Tsai, C. J., Kumar, S., Zanuy, D. & Nussinov, R. Extended disordered proteins: targeting function with less scaffold. Trends Biochem. Sci. 28, 81–85 (2003).

56. Wright, P. E. & Dyson, H. J. Linking folding and binding. Curr. Opin. Struct. Biol. 19, 31–38 (2009).

57. Shoemaker, B. A., Portman, J. I. & Wolynes, P. G. Speeding molecular recognition by using the folding funnel: the fly-casting mechanism. Proc. Natl. Acad. Sci. USA. 97, 8866–8873 (2000).

58. Simon, R., Priester, U. & Puhler, A. A broad host range mobilization system for in vivo genetic engineering: transposition mutagenesis in Gram negative bacteria. Biotechnology. (N.Y.) 1, 784–791 (1983).

59. Whitmore, L. & Wallace, B. A. DICHOMEWEB, an online server for protein secondary structure analyses from circular dichroism spectroscopic data. Nucleic Acids Res 32, W668–673 (2004).

60. Whitmore, L. & Wallace, B. A. Protein secondary structure analyses from circular dichroism spectroscopy: methods and reference databases. Biopolymers 89, 392–400 (2008).

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Author Contributions
K.H. designed the study and S.Y., H.N. and K.H. wrote the paper. H.N. constructed the p3CAtaA, analyzed cells by flow cytometry and CLSM, and conducted TEM. H.N. and S.Y. purified AtaA PSD. S.Y. conducted biochemical and biophysical analyses of AtaA PSD. S.Y. assayed adherence of cells. K.I. constructed the p3CFG2ataA plasmid and purified AtaA PSD short. All authors reviewed the manuscript.

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