AtaA, a New Member of the Trimeric Autotransporter Adhesins from Acinetobacter sp. Tol 5 Mediating High Adhesiveness to Various Abiotic Surfaces

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Abstract

Acinetobacter sp. Tol 5 exhibits an autoagglutinating nature and noteworthy adhesiveness to various abiotic surfaces from hydrophobic plastics to hydrophilic glass and stainless steel. Although previous studies have suggested that bacteriannonfibers on Tol 5 cells are involved in the adhesive phenotype of Tol 5, the fiber that directly mediates Tol 5 adhesion has remained unknown. Here, we present a new member of trimeric autotransporter adhesins designated AtaA, which we discovered by analyzing a less adhesive mutant of Tol 5, T1, obtained by transposon mutagenesis. AtaA forms thinner and shorter nanofibers than fimbriae on Tol 5 cells. We performed target disruption of ataA by allelic marker exchange, and the resulting ΔataA strain was complemented with ataA on the Escherichia coli-Acinetobacter shuttle vector, which was newly constructed. These results proved that AtaA is essential for Tol 5's autoagglutinating nature and high adhesiveness to surfaces of various materials. In addition, the adhesiveness to solid surfaces mediated by AtaA is notably higher than that mediated by YadA of Yersinia enterocolitica WA-314. Moreover, and importantly, these characteristics can be conferred to the non-adhesive, non-agglutinating bacterium Acinetobacter sp. ADP1 in trans by transformation with ataA, with expected applications to microbial immobilization.

Introduction

On bacterial cell surfaces, adhesive nanofibers, which are filamentous appendages of several nanometers in diameter, not only bind to host biomolecules and mediate infectious disease but also tether cells to abiotic surfaces [1]. Many specific interactions between pathogenic bacteria and host cells and tissues are mediated by proteinaceous "bacteriannonfibers", which can be divided into two major classes, fimbriae assembled from hundreds of subunits [2,3], and non-fimbrial fibers having a simple monomeric or oligomeric structure [4].

Trimeric autotransporter adhesins (TAAs) belong to a subclass of non-fimbrial autotransporter adhesins and have recently attracted our attention as virulence factors of Gram-negative pathogens. TAAs form homotrimeric structures with a N-terminal-head-neck-stalk-membrane anchor-C-terminal architecture [5,6]. The C-terminal anchor domain forms a 12-stranded β-barrel at the outer membrane (OM) and the head-neck-stalk domain, called a passenger domain, is secreted to the cell surface through the β-barrel pore. Although head and stalk domains alternate in the sequence of some TAAs, the anchor domain is always localized at the C-terminus, is homologous in all TAAs, and defines this family [7]. TAAs have been reported to mediate bacterial adhesion to host cells and/or extracellular matrix (ECM) proteins, such as collagen, fibronectin, and laminin, invasion of host cells, serum resistance, autoagglutination, and biofilm formation [4,6,8,9,10,11,12,13,14,15,16].

Generally speaking, nonspecific interactions between bacteriannonfibers and abiotic surfaces are initially weak and reversible and enable many bacteria to crawl, glide, or twitch over solid surfaces [17]. Subsequently, bacterial cells attain strong and irreversible adhesion through interactions with surfaces at multiple sites of cell surface biomolecules, including nanofibers. Thereafter, during growth, bacteria on the surfaces form microcolonies, secrete exopolymeric substances (EPS), and develop biofilms, resulting in tenacious adhesion. Thus, bacterial cells require sufficient time to achieve tenacious adhesion to abiotic surfaces through formation of a biofilm. It has however been reported that in certain bacteria, such as Caulobacter crescentus with an adhesive extension of the cell envelope [18] and Aggregatibacter actinomycetemcomitans with Flp fimbriae [19], nanofibers directly mediate rapid and tenacious adhesion to abiotic surfaces.

The toluene-degrading bacterium Acinetobacter sp. Tol 5 previously isolated from a biofiltration process demonstrated an autoagglutinating nature and noteworthy adhesiveness through the nanofibers on the cell surface [20,21,22]. Tol 5 cells are intrinsically adhesive to surfaces of various abiotic materials including hydrophobic plastics, hydrophilic glass, and stainless steel [23]. A large number of Tol 5 resting cells rapidly adhere to solid surfaces independently of cell growth. Tol 5’s initial
attachment ability is quite high and distinguishable from its biofilm formation ability. At least three types of peritrichate nanofibers have been found on Tol 5 cells [24]. Interestingly, production of the peritrichate nanofibers was affected by the available growth substrate. Thick, long, straight nanofibers on cells grown on toluene, lactate, and ethanol were not observed on cells grown on triacylglycerol (TAG). In contrast, cells grown on TAG were covered with long, curved nanofibers, which only existed sparsely on cells grown on toluene, lactate, and ethanol. Thin, short, straight nanofibers were found densely covering the margin of cells grown on all four growth substrates. The analyses of cell adhesiveness and the expression levels of cell surface proteins relevant to the supplied carbon source suggested that the first and second types of nanofibers are type 1 and Fil fimbriae, respectively, and that these fibers are not responsible for the high adhesiveness of Tol 5.

To determine which nanofiber and which of its component molecules directly mediate adhesion of Tol 5 cells, we performed random gene disruption by transposon insertion [22]. Five strains that showed significantly decreased adhesiveness were obtained from culture suspension after cultivation of a mass of random insertion mutants in the presence of a polyurethane (PU) foam support. Of the five less-adhesive mutants obtained, T1 was the least adhesive. When Tol 5 cells were grown in the presence of the PU foam support, most of the wild type (WT) cells adhered to the support while T1 cells were individually dispersed in the culture broth. In the present study, we identified the gene disrupted in T1 as a new member of the TAA genes and demonstrated that this gene product is responsible for the high adhesiveness and autoagglutinating nature of Tol 5 cells.

Results

Discovery of a new TAA member in Acinetobacter sp. Tol 5

Transposon insertion sites on the chromosome of the less-adhesive mutant T1 were analyzed by Southern blot analysis using the Tn5 tetA gene [22] as a probe. On the chromosomal DNA of T1 digested with HindIII, a 5.1-kb fragment hybridized with the tetA probe. The fragment was sequenced, and the DNA sequence from the Tol 5 chromosome within the fragment was determined. The region of the Tol 5 WT chromosome adjacent to the transposon insertion site was amplified by inverse PCR and sequenced. Owing to the presence of long repeat sequences, it was difficult to determine the DNA sequence of the whole gene disrupted by Tn5 insertion. The sequencing strategy is described in Fig. S1. Finally, it was revealed that the structural gene encoding a protein belonging to the TAA family was disrupted in T1. This gene was designated ataA (Acinetobacter trimeric auto-transporter adhesin), and its sequence data have been deposited with DDBJ/EMBL/GenBank under accession no. AB542908.

AtaA is predicted to consist of 3,630 amino acid residues, which makes it one of the largest TAAs known to date (Fig. 1). Although AtaA follows the general N-terminus-head-stalk-anchor-C-terminal organization, an additional head domain localizes in the C-terminal region. The stalk domain of AtaA is notably longer than that of other TAAs and contains peptide repeats that are mosaically arranged. Although the boundaries of these repeat sequences are fuzzy, they can be divided into six groups when non-conserved amino acids are minimized in the respective groups; in each repeat sequence group, amino acid sequences are entirely conserved, or only a few amino acid residues are non-conserved (Fig. S2). Further domain annotation using the dnaTAA program [7] revealed that the two heads of AtaA were annotated as the YIhead domain, the head of Yersinia enterocolitica YadA, which is the most well-studied TAA (Fig. 1). AtaA also has twelve Trp-rings, in which the tryptophan residue of the highly conserved Gly-Trp motif in many TAA Trp-ring domains is replaced by leucine or phenylalanine. As the Trp-ring consists mainly of β-structures and motifs, the AtaA stalk domain is also anticipated to contain more β-structures than coiled coils, although coiled coils are abundant in most TAAs. In fact, in the ultrastructure of AtaA, coiled coils were predicted to be infrequent at the N-terminal stalk domain according to analysis performed using the COILS program (Fig. S3) [25].

To identify which nanofiber is composed of AtaA, Western-blot analysis and immunoelectron microscopy were carried out using an anti-AtaA antibody against a part of the stalk domain (AtaA_301, 1014). AtaA expression and depletion were confirmed at the OM of the Tol 5 WT and the mutant strain T1, respectively (Fig. 2A). Immunoelectron microscopy revealed that AtaA composes the thinnest and shortest nanofiber (200 nm long, 3.5±0.4 nm diameter) of the three types of peritrichate nanofibers previously reported [24] (Fig. 2B and 2C). The antibody labels did not line each fiber but were localized solely near the distal end of the fiber, showing that AtaA architecture is different from fimbriae constructed by the accumulation of a single protein in the hundreds. In our previous study, the larger nanofibers, which often obscured the smallest AtaA fibers, were deduced to be dominant type 1 fimbriae and sparse Fil fimbriae [24]. Furthermore, the lack of AtaA fibers on T1 was confirmed by immunoelectron microscopy (Fig. S4).

Involvement of AtaA in the high adhesiveness of Tol 5 cells

To clarify that disruption of ataA causes a lack of high adhesiveness in Tol 5 cells, allelic marker exchange was carried out for target disruption of ataA. The resulting ΔataA mutant was confirmed to neither produce AtaA protein at the OM by Western-blot analysis (ataA) nor display AtaA fibers on the cell surface by immunoelectron microscopy (Fig. S4) and flow cytometry (Fig. 3B). Expression levels of components of type 1 fimbriae and Fil fimbriae, as well as respective fiber production in the ΔataA mutant, were also confirmed to be similar to those in the WT (Fig. S5).

Figure 4 shows the results of the adherence assay using 48-well polystyrene (PS) plates and type I collagen-coated PS plates under a resting cell condition. Although the photographs resemble those of a biofilm formation assay, the adherence assay in the present study differed from a biofilm formation assay and directly reflected the initial attachment ability of bacteria to solid surfaces, as reported previously [23]. To compare adhesiveness mediated by AtaA with that mediated by a typical TAA, two Y. enterocolitica strains, WA-314 and WA-C, were subjected to the adherence assay with Tol 5 and its derivative strains. The strain WA-314 expresses YadA and exhibits YadA-dependent adhesion to type I collagen [26], whereas the strain WA-C is a YadA-deficient derivative of WA-314 formed by curing of the virulence plasmid pYV08 [27]. Tol 5 WT cells showed high adhesiveness to both surfaces, regardless of collagen-coating. Although the strain WA-C scarcely adhered to both surfaces, the strain WA-314 expressing YadA preferred the collagen-coated surface (P<0.01) to the abiotic surface; but its adhesiveness was trivial compared with Tol 5 WT cells. Cells of the ΔataA mutant adhered only slightly to both surfaces, implying that the deficiency of AtaA nanofibers results in the poor adhesiveness of Tol 5 cells.

For further confirmation of AtaA involvement in Tol 5 adhesion, we tried to complement the ΔataA mutant with ataA...
on a plasmid vector. However, Tol 5 cells were difficult to transform due to their low competency, we constructed the novel *Acinetobacter-Escherichia coli* shuttle vector pARP3. This expression plasmid vector can be introduced into recipients by conjugation, is capable of replicating in *Acinetobacter* strains, and possesses an arabinose-inducible promoter (Fig. S6). The expression level of a gene may be important when the physiology of a bacterium (e.g., pathogenicity) is the primary focus, because the responsibility of the target gene for physiology under natural conditions is important. However, when functions or properties of gene products themselves are the focus, researchers demonstrate the functions and properties under elicitable conditions. In fact, for the characterization of individual TAAs, other researchers on TAAs also have used inducible promoters [13,28,29,30]. Thus, the ataA locus containing the Shine-Dalgarno (SD) sequence was cloned under the control of the arabinose-inducible promoter of pARP3, generating the ataA-expression vector pAtaA. The ΔataA mutant was transformed with pARP3 (ΔataA, pARP3) or pAtaA (ΔataA, pAtaA), and the OMs of these transformants were subjected to Western-blot analysis. Although no AtaA was detected at the OM of ΔataA, pARP3 cells (negative control), ΔataA, pAtaA cells restored AtaA production to a similar level as that in the WT (Fig. 3A). These transformants were also subjected to flow cytometry to analyze the surface display of AtaA. The resultant histograms of flow cytometry showed that AtaA displayed on the cell surface of the complemented mutant (ΔataA, pAtaA) was restored to a level similar to that in the WT (Tol 5 WT), regardless of the use of the inducible promoter in the multiple copied plasmid (Fig. 3B). The adherence assay of the complemented mutant, whose ataA expression level was almost equal to that of the WT, demonstrated that complementation of ΔataA with pAtaA entirely restored the high adhesiveness to both PS and collagen surfaces (Fig. 4). These results prove that AtaA is essential for the high adhesiveness of Tol 5, that the AtaA-mediating adhesiveness is notably higher than the YadA-mediating adhesiveness, and that AtaA interacts with abiotic surfaces to at least the same degree as it does with a biotic collagen surface.

**Adhesive features mediated by AtaA**

The Tol 5 strain, despite its hydrophobic cell surface, can adhere to hydrophilic glass and metal surfaces as well as to hydrophobic plastic surfaces [23]. To confirm that this nonspecific adhesiveness is mediated by AtaA, we examined the adhesiveness of the ΔataA mutant, its transformant with the empty vector pARP3 (ΔataA, pARP3), and the complemented mutant (ΔataA, pAtaA).
pAtaA) to four abiotic surfaces: polychloroprene (PP), polystyrene (PS), and hydrophilic metal (SUS). While ΔataA and ΔataA, pARP3 strains showed low adhesiveness to these surfaces, the ΔataA, pAtaA strain restored the nonspecific and high adhesiveness of Tol 5 WT, indicating that AtaA is responsible for the nonspecific adhesion of Tol 5 cells (Fig. 5).

T1 cells obtained by Tn5 random insertion not only show low adhesiveness but also lack the autoagglutinating nature of Tol 5 [31], and some TAA s are involved in autoagglutination, which is self-adhesion of cells [6]. Therefore, this feature was also examined for the ΔataA mutant and its derivatives. The cell suspension was prepared by the same procedure as the adherence assay to prevent cell growth and subjected to the tube-settling assay to quantify the sedimentation of cell clumps. While the WT and ΔataA, pAtaA strains showed high autoagglutination ratios, these ratios of ΔataA and ΔataA, pARP3 remained very low during the course of the assay (Fig. 6). Although strain WA-314 showed a higher autoagglutination ratio than strain WA-C, the value was much lower than that of Tol 5. Therefore, AtaA mediates the autoagglutination as well as the high adhesiveness of Acinetobacter sp. Tol 5.

Expression of ataA in another non-adhesive, non-autoagglutinating bacterium

To conclude that the adhesive features described above are intrinsic in nature of AtaA, we finally examined the capability of AtaA to confer the adhesive features to another strain of Gram-negative bacteria. Another non-adhesive, non-autoagglutinating Acinetobacter species, strain ADP1, which is useful for biotechnological applications due to its capability to degrade and produce many important chemicals [32], was transformed with pAtaA. Since this strain is highly competent for natural transformation, resulting in ease of genetic modification of this strain [33], it is an ideal model organism for genome engineering and is superior to Escherichia coli in this respect [34]. The resultant transformant (ADP1, pAtaA) produced AtaA protein at the OM (Fig. 7A) and displayed AtaA nanofibers on ADP1 cells (Fig. 7B, C). Thus, we succeeded in the production of AtaA nanofibers on cells of a different bacterial strain from the original Tol 5.

The ADP1, pAtaA strain, the original ADP1, and the transformant harboring the empty vector pARP3 (ADP1, pARP3) were evaluated for their adhesiveness to various solid surfaces composed of biotic collagen, hydrophobic plastics, hydrophilic glass, and hydrophilic metal, as described above for Tol 5 and its derivatives. Although we previously reported the adhesion preferences of ADP1 for PP, PVC, and SUS surfaces [23], strain ADP1, pAtaA showed nonspecific adhesion to all examined surfaces (Fig. 8A). Exogenous AtaA conferred the high adhesiveness of ADP1 cells to type I collagen, PS, and glass surfaces in a level similar to that of Tol 5 cells, although the original ADP1 cells could adhere only slightly or not at all to these surfaces. The adhesiveness of ADP1 cells to PP, PVC, and SUS surfaces was also elevated by AtaA to a level comparable with that of Tol 5 in relation to the adhesiveness of the original cells, which was less than half or one-third that of Tol 5.

ADP1 and its derivatives were also subjected to the tube-settling assay for quantification of autoagglutination. ADP1 cells intrinsically showed a non-agglutinating nature, but the ADP1, pAtaA strain acquired the autoagglutinating nature through the expression of exogenous ataA (Fig. 8B). Thus, it was proven that AtaA has a nature of exhibiting high adhesiveness to various solid surfaces and autoagglutinating bacterial cells and that this nature could be conferred in trans to another bacterial strain.

Discussion

In the present study, we discovered AtaA, a new member of the TAA family, in Acinetobacter sp. Tol 5, which intrinsically exhibits a high adhesive phenotype to various abiotic surfaces. Most reports on TAAs have focused on their ability to adhere to biotic surfaces, although several TAAs have been reported to be responsible for biofilm formation on various abiotic surfaces [8,13,29,35,36]. However, biofilm formation ability is different from the initial attachment ability exhibited by AtaA [23]. Biofilm formation includes multiple biological factors as well as the direct interaction between adhesins and surfaces. During formation of biofilms, EPS, which contains a variety of biopolymers, such as polysaccharides, proteins, and DNA, are secreted from bacterial cells and construct the matrix of biofilms. Furthermore, involvement of TAAs in biofilm formation on abiotic surfaces might be attributed to the
autoagglutination function rather than to initial attachment, as reported in UpaG of E. coli [13]. Recently, Muller et al analyzed the initial attachment of bacterial cells to solid surfaces via three TAAs, BadA of Bartonella henselae, Vomps of Bartonella quintana, and YadA of Y. enterocolitica, under static and flow conditions [37]. In their study, the TAAs, particularly BadA, mediated relatively nonspecific adherence to “biotic” surfaces as well as adherence to a plastic surface under the static condition, although YadA did not mediate adherence to fibronectin surfaces. Our data demonstrate that adhesiveness mediated by AtaA to “abiotic” surfaces is nonspecific and notably higher than that mediated by YadA to both PS and type I collagen surfaces. Furthermore, and importantly, this phenotype can be conferred to other bacterial species by introducing the single gene ataA. This potential leads us to expect that AtaA can be utilized for direct immobilization of bacterial cells to abiotic support surfaces.

The most frequently used technique for microbial immobilization involves entrapping cells in polymer gel; however, such an approach adversely affects the mass transfer rate in the inner part of the gel [38,39]. Therefore, direct immobilization of bacterial cells onto the surface of a support by utilizing AtaA would represent a great advance for bioengineering employing whole microbial cells. In theory, the introduction of AtaA into industrially relevant Gram-negative bacteria, including other Acinetobacter strains, would provide a novel and effective means for the immobilization of bacteria in the production of chemicals, biomass energy, and other applications.

Our data demonstrate that AtaA also mediates autoagglutination (Figs. 6 and 8B). Autoagglutination has been reported to be mediated by various types of nanofibers and to be important for biofilm formation on both biotic and abiotic surfaces [3,6,15,19,40]. However, it is regarded as a form of specific binding since it is mediated by homologous recognition of an adhesin or interaction with other target molecules on the cell surface [6,15]. The functional sites mediating the high adhesiveness of AtaA to solid surfaces and those for autoagglutination may be different.

With regard to features of the primary structure of AtaA, multiple head domains are seen not only in AtaA but also in some TAAs [13,41,42,43,44]. Although mosaically arranged repeat sequences are also a common feature of TAAs [7], the AtaA stalk that consists of extremely conserved, long (50-243 residues) repeats is remarkable. Recently, Cha adhesin in Haemophilus cryptic genospecies was also found to contain a series of entirely conserved tandem repeats, but they consist of only 28 residues in contrast with the long repeat sequences in AtaA [12]. Although the tandem repeats in Cha affect the adherence activity of the Haemophilus strain [44], the function of the long repeat sequences in AtaA remains to be studied.

On the basis of our results alone, the possibility that other proteins cooperate with AtaA in Tol 5 adhesion cannot be denied. However, other bacterial species that are intrinsically non-adhesive and non-autoagglutinating acquired the adhesive properties by introduction of the single gene ataA. This strongly suggests that AtaA is the primary mediator of the high adhesiveness and autoagglutinating nature of bacteria.

Materials and Methods

Culture conditions

The bacterial strains used in this study are listed in Table 1. Acinetobacter sp. Tol 5 and its derivative mutants were grown in basal salt (BS) medium at 28°C as described previously [22] or in Luria-Bertani (LB) medium. Acinetobacter sp. ADP1 and its derivative mutants were grown in LB medium at 30°C. Antibiotics were used at the following concentrations when required: ampicillin (1,000 µg/ml), tetracycline (10 µg/ml), gentamicin (10 µg/ml) for Tol 5 derivative mutants; ampicillin (100 µg/ml) and gentamicin (10 µg/ml) for ADP1 derivative mutants. Arabinose was added to a final concentration of 0.5% (wt/vol) for the induction of ataA under the control of the PataA promoter. For expression of YadA, Y. enterocolitica was grown overnight in LB medium at 27°C, the overnight culture was diluted to 1:20 in LB medium, and then incubated at 37°C for 3 h, as described previously [26].
DNA manipulation

General DNA manipulations, such as PCR, restriction enzyme digestion, and ligation, were performed using standard protocols. The plasmids and the primers used in this study, with the exception of those used for primer walking during DNA sequencing, are detailed in Table 1 and Table S1, respectively. Transformation of Tol 5 T1 was carried out by bacterial conjugation with donor strain *E. coli* S17-1 harboring the target plasmid [45]. Transformation of *Acinetobacter* sp. ADP1 was performed by natural transformation as described previously [34].

Construction of pARP3 and pAtaA

A mobile plasmid, pJQ200sk [46], was used as the plasmid backbone. A fragment containing an *Acinetobacter* origin of replication was amplified by PCR from pWH1266 [47] using the primers *Acinetobacter*-oriF/*Acinetobacter*-oriR, digested with *Sph*I, and then ligated into the *Sph*I site of pJQ200sk to generate pSAE1. To remove the *sacB* gene and introduce a new *Pvu*II site in pSAE1, inverse PCR was performed using the primers dele-sacBF/dele-sacBR, and the amplified PCR product was digested with *Pvu*II and then self-ligated to generate pSAE2. Although Tol 5 and T1 have potential ampicillin resistance, both are sensitive to high...
was slightly transparent compared with the partially formed cell clumps. Cell suspension of WA-314 strain of cells, whereas cell suspensions of strains lacking functional

The P3-Apr cassette was amplified from pP3 by PCR derivatives, and performed in two steps. First, the fragment containing the deduced promoter P3 and confers resistance to high concentrations of ampicillin, could survive in LB medium containing 1,000 μg/ml gentamicin and 10 μg/ml tetracycline, and supplemented with toluene. Following, positive selection of allelic exchange mutants was performed on a BS agar plate containing 100 μg/ml gentamicin and 10 μg/ml tetracycline, and supplemented with toluene. Inactivation of ataa and the antibiotic resistance of the allelic exchange mutants were confirmed by PCR and the plating test (GmS, TcR, SucS (sucrose sensitive) colonies were selected on a BS agar plate containing 100 μg/ml gentamicin and 10 μg/ml tetracycline, and supplemented with toluene. Preparations of outer membrane proteins (OMPs), bacterial cells were grown to the stationary phase, aerobically in 100 ml of LB medium for 6 h at 28°C (Tol 5 and its derivatives) or for 12 h at 30°C (ADP1 and its transformants), in the presence of 0.5% arabinose, with shaking at 115 rpm. The cultures (OD660 = 2.1) were harvested at centrifugation at 6,000 g for 10 min, washed with 50 mM phosphate buffer (pH 7.4), and then resuspended in 10 ml of the same buffer. The cells were lysed by sonication and centrifuged at 6,000 g for 10 min to remove unlysed cells and debris. The supernatant was centrifuged at 100,000 g for 1 h to pellet total membrane proteins which were then resuspended in a 0.5% N-lauroylsarcosine solution and incubated at room temperature for 30 min to lyse inner membrane proteins. The sample was centrifuged again at 100,000 g for 1 h and OMPs were collected as a pellet which was resuspended in 50 mM phosphate buffer (pH 7.4) or directly lysed in SDS-PAGE sample buffer.

Production of GST-AtaA fusion protein and generation of an anti-AtaA antibody

Recombinant AtaA99-1014 was expressed in E. coli as a glutathione S-transferase (GST) fusion protein to generate an anti-AtaA antibody. The DNA fragment encoding AtaA99-1014 was amplified by PCR with the primer set AtaA-stalkF/AtaA-stalkR. The PCR product was digested with BamHI and NotI, ligated into the BamHI-NotI site of the pGEX-4T-2 plasmid (GE Healthcare), and E. coli BL21 was transformed with the constructed plasmid. The transformation was cultured at 37°C until an OD600 of 1.5 was reached, and the expression of the GST fusion gene was then induced by adding 0.1 mM IPTG, followed

concentrations of ampicillin (the minimum inhibitory concentration for growth is 120 μg/ml). The Tol 5 transformant harboring plasmid pP3, which expresses the bla gene under the control of promoter P3 and confers resistance to high concentrations of ampicillin, could survive in LB medium containing 1,000 μg/ml ampicillin. The P3-Ap' cassette was amplified from pP3 by PCR using the primers p3-ApF/p3-ApR, digested with PshII, and then cloned into the PshII site of pSAE2 to generate pSAE3 which facilitated selection of Tol 5 derivative transformants. The araC-PRAD system was excised from the pJN105 plasmid [48] as a SalI-KpnI fragment and cloned into the SalI-KpnI site of the pSAE3, generating pARP3 (Fig. S6), which served as an arabinose-inducible expression vector capable of replicating in Acinetobacter strains.

Cloning ataa under the control of promoter PRAD of pARP3 was cloned in two steps. First, the fragment containing the deduced Shine-Dalgarno (SD) sequence and the entire ataa gene was amplified by PCR using two primer sets, attB1-ataA1F/attB2-ataA1R and attB1-adapter/attB2-adapter. The PCR product was inserted into pDONR221 using BP Clonase II enzyme mix (Invitrogen), and both the SD sequence and the entire ataa gene were then inserted into the EcoRI-XhoI site of pARP3 as an EcoRI-XhoI fragment to generate plasmid pAtaA.

Target mutagenesis of ataa by allelic marker exchange

To construct a ataa mutant, we employed a double selection strategy using sacB, which is lethal to various Gram-negative bacteria in the presence of sucrose, as a negative selection marker. For construction of a suicide vector to knockout ataa, we amplified a 1,712 bp-internal fragment containing a part of ataa (corresponding to bases 9,060–10,771 of ataa) by PCR using primers ataa9060F/ataa10771R and cloned it into the pTA2 vector by using TArget clone-plus- (TOYOBO) according to the protocol provided by the manufacturer. A selectable marker was inserted in vitro using the EZ:TN <TET-1> insertion kit (Epicentaur Biotechnologies), and the <TET-1> insertion site was confirmed by sequencing. The ataa fragment integrated by <TET-1> was amplified by PCR using primers infusion-ataa9060F/infusion-ataa10771R and cloned into the BamHI site of pJQ200ks by using the In-Fusion 2.0 Dry-down PCR Cloning kit (Clontech). The resultant suicide vector was transferred into Tol 5 cells from E. coli S17-1 via bacterial conjugation, and GmR, TcR, SucS (sucrose sensitive) colonies were selected on a BS agar plate containing 100 μg/ml gentamicin and 10 μg/ml tetracycline, and supplemented with toluene. Following, positive selection of allelic exchange mutants was performed on a BS agar plate containing 5% sucrose supplemented with toluene. Inactivation of ataa and the antibiotic resistance of the allelic exchange mutants were confirmed by PCR and the plating test (GmS, TcS, SucR (sucrose resistant), respectively.

Protein manipulation

Preparation of nanofiber proteins was performed as described previously [24]. For the preparation of outer membrane proteins (OMPs), bacterial cells were grown to the stationary phase, aerobically in 100 ml of LB medium for 6 h at 28°C (Tol 5 and its derivatives) or for 12 h at 30°C (ADP1 and its transformants), in the presence of 0.5% arabinose, with shaking at 115 rpm. The cultures (OD660 = 2.1) were harvested at centrifugation at 6,000 g for 10 min, washed with 50 mM phosphate buffer (pH 7.4), and then resuspended in 10 ml of the same buffer. The cells were lysed by sonication and centrifuged at 6,000 g for 10 min to remove unlysed cells and debris. The supernatant was centrifuged at 100,000 g for 1 h to pellet total membrane proteins which were then resuspended in a 0.5% N-lauroylsarcosine solution and incubated at room temperature for 30 min to lyse inner membrane proteins. The sample was centrifuged again at 100,000 g for 1 h and OMPs were collected as a pellet which was resuspended in 50 mM phosphate buffer (pH 7.4) or directly lysed in SDS-PAGE sample buffer.

Figure 6. Autoagglutination assay of Acinetobacter sp. Tol 5, its derivatives, and Y. enterocolitica strains. (A) Strains expressing ataa (WT; ataaA, pAAaA) autoagglutinate and form cell clumps that settle at the bottom of test tubes. After standing for 3 h without agitation, suspensions of these strains were transparent due to the sedimentation of cells, whereas cell suspensions of strains lacking functional ataa (∆ataA; ∆ataA, pAAaA) remained cloudy. Yersinia strain expressing YadA (WA-314) partially formed cell clumps. Cell suspension of WA-314 strain was slightly transparent compared with the yadaA-mutant strain (WA-C). (B) Autoagglutination observed in (A) was quantified by a decrease in optical density at 660 nm (OD660). Graph bars show the autoagglutination ratio (%). Data are expressed as mean and SEM (n = 3) values. Statistical significance, P < 0.001. doi:10.1371/journal.pone.0048830.g006
by incubation at 25°C for 3 h. The cells were harvested by centrifugation, resuspended in PBS containing 1 mM dithiothreitol, lysed by sonication, and centrifuged at 6,000 g for 10 min to remove the unlysed cells and debris. The GST fusion protein was purified from the supernatant using GSTrap FF (GE Healthcare). The purified protein was inoculated into rabbits for generating a polyclonal anti-AtaA antiserum and antibody.

Western-blot and immunodetection

OMPs were separated on 7.5% acrylamide gels and transferred to PVDF membrane with a constant voltage of 50 V overnight at 10°C using a Criterion Trans-Blot Cell (Bio-Rad) and blotting buffer (25 mM Tris; 192 mM Glycine; 0.1% SDS; 5% methanol). The blotted membrane was blocked using a 5% skim milk solution for 1 h at room temperature and then treated with anti-AtaA699–1014 antiserum at a 1:50,000 dilution in 5% skim milk solution.

The proteins bound by the antiserum were detected with a horseradish peroxidase-conjugated anti-rabbit antibody (GE Healthcare) at a 1:10,000 dilution in 5% skim milk solution and visualized using ECL Western Blotting Detection Reagents (GE Healthcare).

Flow cytometry

Acinetobacter strains were inoculated into 2 ml of LB medium and grown overnight. The overnight cell cultures were diluted to 1:100 with 20 ml of fresh LB medium and grown aerobically to the stationary phase (around 2.1 of OD660) for 6 h at 28°C (Tol 5 and its derivatives), or for 12 h at 30°C (ADP1 and its transformants) in the presence of 0.5% arabinose, with shaking at 115 rpm. Cells were washed with PBS, fixed with 4% paraformaldehyde, washed twice with PBS, and resuspended in PBS. Samples were treated with anti-AtaA699–1014 antiserum at a 1:10,000 dilution in PBS containing 0.05% Tween 20 (Calbiochem) for 30 min at room temperature. After incubation, cells were washed twice with NET buffer (150 mM NaCl, 5 mM EDTA, 50 mM Tris-HCl, 5% Triton X-100 (Calbiochem); pH 7.6), treated with Alexa Fluor 488 conjugated to anti-rabbit antibody (Cell Signaling Technology) at 1:10,000 dilution in PBS containing 0.05% Tween 20 (Calbiochem). The proteins bound by the antiserum were detected with a horseradish peroxidase-conjugated anti-rabbit antibody (GE Healthcare) at a 1:10,000 dilution in 5% skim milk solution and visualized using ECL Western Blotting Detection Reagents (GE Healthcare).

Figure 7. Expression of ataA in another Acinetobacter strain, ADP1. (A) Immunodetection of AtaA using anti-AtaA antiserum against OM proteins prepared from Acinetobacter sp. ADP1, ADP1 transformed with the empty vector pARP3 (ADP1, pARP3), and ADP1 transformed with ataA in pARP3 (ADP1, pAtaA). (B) Flow cytometry analysis of AtaA displayed on the cell surface of ADP1 and its transformants. ADP1, pAtaA (red line) shows a large number of AtaA proteins on its cell surface. ADP1 (dashed line, gray fill) and ADP1, pARP3 (black line) strains are shown as negative controls. (C) Confirmation of AtaA fibers on strains expressing ataA (ADP1, pAtaA) by immunoelectron microscopy.
a 1:500 dilution in NET buffer for 30 min at room temperature, and washed three times with NET buffer. Finally, the cells were resuspended in pure water, and surface-displayed AtaA was measured by flow cytometry on a FACSCanto II (Becton Dickinson). Histograms were created using WinMDI 2.9 (J. Trotter) software.

Figure 8. Adherence and autoagglutination assays of Acinetobacter sp. ADP1 and its transformants. (A) Adherence assay of ADP1 and its transformants (ADP1; ADP1, pARP3; ADP1, pAtaA) to 48-well PS and type I collagen-coated plates, to 96-well plates made of PP, PVC, and glass, and to a sample cup made from SUS. Data are expressed as mean ± SEM (n = 3) values. Statistical significance, *P < 0.01. (B) Autoagglutination assay of ADP1 and its transformants (ADP1; ADP1, pARP3; ADP1, pAtaA). Graph bars show the autoagglutination ratio (%). Data are expressed as mean ± SEM (n = 3) values. Statistical significance, *P < 0.001.
Adherence and autoagglutination assays

Cells of *Acinetobacter* strains were grown by the same procedure as flow cytometry. *Y. enterocolitica* strains were cultured and treated as described above for the induction of *yadA*. The grown cells were suspended in BS-N medium [49] containing no carbon nor nitrogen sources and were subjected to the adherence and autoagglutination assays under the resting cell condition, as described previously [23]. The cell suspension was incubated at 28°C for 2 h in the adherence assay. In the autoagglutination assay, test tubes containing the cell suspension were left to stand at room temperature for 3 h (tube-settling assay) [15,50].

Electron microscopy

Nanofibers on cell surfaces were observed using transmission electron microscopy (TEM) after staining cells with 1% methylamine tungstate (MAT) as described previously [24]. For immunoelectron microscopy, anti-AtaA<sub>699-1014</sub> antibody and colloidal gold conjugated with a goat anti-rabbit IgG were used for the immunolabeling of the AtaA fibers. Carbon-coated copper grids were hydrophilized using an ion bombarder (PIB-10, Vacuum Device). Specimen preparation was performed in a moisture box to avoid desiccation. For each sample, 1 ml of cell culture with an OD<sub>660</sub> of 0.6 was placed into a 1.5 ml tube and centrifuged at 6,000 g for 1 min. The pellet was resuspended in phosphate buffered saline (PBS) after rinsing the cells twice with PBS. One drop of cell suspension was placed on the hydrophilized carbon-coated copper grid and allowed to stand for 5 min. The excess liquid was removed using filter paper and the grid was rinsed twice with PBS. For blocking, one drop of PBS containing 1% BSA was placed on the grid and left for 1 h. For detection of the nanofiber formed by AtaA, the grid was incubated with an anti-AtaA antibody at a 1:100 dilution in PBS containing 1% BSA for 2 h. After washing, the grid was stained with 1% MAT and observed under a TEM (Hitachi H-7100) operated at 100 kV.

Supporting Information

Figure S1 Sequencing strategy for *ataA* containing several highly conserved, long repeat sequences. To determine the DNA sequence of the gene disrupted in the transposon Tn5-inserted mutant T1, nine DNA fragments were obtained by inverse PCR, southern hybridization, and normal

| Table 1. Bacterial strains and plasmids used in this study. |
|---------------------------------|---------------------------------|------------------|
| **Strain or plasmid**            | **Description**                 | **Reference**    |
|---------------------------------|---------------------------------|------------------|
| *Acinetobacter* sp.             |                                 |                  |
| Tol 5                           | Wild type strain                | [20]             |
| Tol 5 T1                        | Tol 5 mutant lacking adhesive and autoagglutinating properties | [22]             |
| *ataA*                          | Tol 5 mutant with defect in *ataA* by allelic marker exchange | This study       |
| ADP1                            | A representative strain of *Acinetobacter* | [33]             |
| E. coli                         |                                 |                  |
| DH5s                            | Host for routine cloning        | Takara           |
| DH10B                           | Host for routine cloning        | Invitrogen       |
| S17-1                           | Donor strain for conjugation    | [45]             |
| BL21                            | Host for protein expression     | GE Healthcare    |
| *Y. enterocolitica*              |                                 |                  |
| WA-314                          | *Y. enterocolitica* serotype O:8 expressing YadA | [27]             |
| WA-C                            | Plasmidless derivative of WA-314, YadA<sup>+</sup> | [27]             |
| **Plasmid**                     |                                 |                  |
| pUC118                          | Cloning vector, Ap<sup>+</sup>  | Takara           |
| pCR-XL-TOPO                     | Cloning vector, Km<sup>+</sup>  | Invitrogen       |
| pDONR221                        | Donor vector for Gateway technology, Km<sup>+</sup> | Invitrogen       |
| pGEX4T-2                        | Expression vector for GST fusion protein, Ap<sup>+</sup> | GE healthcare    |
| pHWH1266                        | *E. coli-Acinetobacter* shuttle plasmid, Tc<sup>+</sup>, Ap<sup>+</sup> | [47]             |
| pJ2Q200sk                       | Mobile plasmid, SacB, Gm<sup>+</sup> | [46]             |
| pP3                             | Source of P3-Ap<sup>+</sup> cassette; high resistance to ampicillin, Km<sup>+</sup>, Ap<sup>+</sup> | T. Kurihara     |
| pSxE1                           | pJ2Q200sk::*Acinetobacter* origin, SacB, Gm<sup>+</sup> | This study       |
| pSXE2                           | pSXE::PvuII site, ΔSacB, Gm<sup>+</sup> | This study       |
| pSXE3                           | pSXE::P3-Ap<sup>+</sup>, ΔSacB, Gm<sup>+</sup>, Ap<sup>+</sup> | This study       |
| pJN105                          | Source of araC<sub>P</sub>-araBAD<sub>Gm</sub> system, Gm<sup>+</sup> | [48]             |
| pARP3                           | *E. coli-Acinetobacter* shuttle expression vector, pSXEΔSacB::P3 Ap<sup>+</sup>araC<sub>P</sub>-araBAD<sub>Gm</sub>, Ap<sup>+</sup> | This study       |
| pAtaA                           | *ataA*-expression vector, pARP3::ataA | This study       |

<sup>a</sup>Ap, ampicillin; Km, kanamycin; Tc, tetracycline; Gm, gentamycin.

doi:10.1371/journal.pone.0048830.t001
PCR. Initially, the Tn5 insertion site was identified from the genomic DNA of T1 by southern hybridization with the Tn5 tetA gene as a DNA probe, and the 5-kb flanking DNA region of the Tn5 insertion site was then amplified by inverse PCR. As a result of sequencing this fragment with the primer walking method, an incomplete structural gene (ataA), which showed partial homology with TAs, was identified. The overlapping regions containing the ataA gene were cloned as seven DNA fragments (three Pst I and four Hind III fragments) by repeating southern hybridization. As these DNA fragments contained several long repeat sequences which disturbed sequencing with the primer walking method, deletion fragments were prepared using exonuclease III to avoid this problem, and their sequences were then determined. Finally, a DNA fragment containing full-length ataA was amplified by PCR to reconfirm the sequence with the primer walking method. The sequencing primers used in the repeat-rich regions were individually designed so that their 3’ terminal nucleotide was inconsistent with that of other repeat sequences. The 20 black flags in the PCR fragment show the positions of the primers that were used to confirm the sequence and the direction of the sequencing from the respective primers.

(TIF)

Figure S2 Full-length AtaA amino acid sequence. Long repeat sequences can be classified into six groups colored light blue (243 aa), orange (50 aa), pink (115 aa), blue (86 aa), light green (79 aa), and green (130 aa) which minimize non-conserved amino acid residues. The non-conserved residues in each group are indicated by characters in bold font. Underlined characters indicate the amino acid residues that were expressed as a recombinant protein for generating anti-AtaA antibody.

(TIF)

Figure S3 Prediction of AtaA deficiency in coiled coils. The full-length amino acid sequence of AtaA was analyzed using COILS [http://www.ch.embnet.org/software/COILS_form.html] to determine the coiled coil probability score using a MTDIck matrix and a 28 residue window. AtaA was compared with several typical TAs: YadA of T. enterocolitica, BadA of Bartonella henselae, UspA1 of Moraxella catarrhalis, and Hia of Haemophilus influenzae. Although the coiled coil structure is abundant in most TAs, this structure is infrequent in the N-terminal stalk region of AtaA.

(TIF)

Figure S4 Confirmation of the specificity of an anti-AtaA antibody on immunoelectron microscopy. Acinetobacter sp. Tol 5 T1 (T1) and ΔataA mutant (ΔataA) were observed by immunoelectron microscopy using anti-AtaA<sub>99-1014</sub> antibody. No nanofibers including AtaA were observed on T1 cells grown on tolulene. The antibodies did not bind to nanofibers, which were different from AtaA, on ΔataA cells.

(TIF)

Figure S5 Confirmation of the production of type 1 and Fil fimbral component proteins by SDS-PAGE and their deduced fibers by TEM in the ΔataA strain. (A) Samples were prepared from WT and ΔataA cells grown on tolulene. Monomeric FimA protein was detected by SDS-PAGE, and the deduced type 1 fimbriae were observed on the cell surface of ΔataA. (B) Samples were prepared from WT and ΔataA cells grown on triacylglycerol. Monomeric FilA protein was detected by SDS-PAGE, and the deduced Fil fimbriae were observed on the cell surface of ΔataA. The bands of FimA and FilA were previously identified by Edman degradation.

(TIF)

Table S1 Primers used in this study (excluding those for DNA sequencing).

Acknowledgments

We thank Y. Hotta and K. Yamamoto of Aichi University for technical support during the TEM observation of bacterial cells. We thank A. Higuchi and H. Ebi for TEM analysis, S. Takada for DNA sequencing, and T. Kurihara of Kyoto University for providing the pP3 vector. We are grateful for receiving vectors from the cloning vector collection distributed by "National BioResource (NIG, Japan): E. coli".

Author Contributions

Conceived and designed the experiments: KH MI. Performed the experiments: KH MI. Analyzed the data: KH MI. Contributed reagents/materials/analysis tools: KH MI. Wrote the paper: KH MI.

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