Aerococcus urinae isolated from women with lower urinary tract symptoms: In vitro aggregation and genome analysis

Evann E Hilt  
*Loyola University Chicago*

Catherine Putonti  
*Loyola University Chicago*

Krystal Thomas-White  
*Loyola University Chicago*

Amanda L Lewis  
*Washington University School of Medicine in St. Louis*

Karen L Visick  
*Loyola University Chicago*

*See next page for additional authors*

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Authors
Evann E Hilt, Catherine Putonti, Krystal Thomas-White, Amanda L Lewis, Karen L Visick, Nicole M Gilbert, and Alan J Wolfe
Aerococcus urinae Isolated from Women with Lower Urinary Tract Symptoms: In Vitro Aggregation and Genome Analysis

Evann E. Hilt,** Catherine Putonti,a,b,c,d Krystal Thomas-White,a Amanda L. Lewis,a Karen L. Visick,a Nicole M. Gilbert,a, f Alan J. Wolfe*a

aDepartment of Microbiology and Immunology, Stritch School of Medicine, Loyola University Chicago, Maywood, Illinois, USA
bDepartment of Biology, Loyola University Chicago, Chicago, Illinois, USA
cDepartment of Computer Science, Loyola University Chicago, Chicago, Illinois, USA
dBioinformatics Program, Loyola University Chicago, Chicago, Illinois, USA
eCenter for Women’s Infectious Disease Research, Department of Molecular Microbiology, Washington University School of Medicine, St. Louis, Missouri, USA
fCenter for Reproductive Health Sciences, Department of Obstetrics and Gynecology, Washington University School of Medicine, St. Louis, Missouri, USA

ABSTRACT Aerococcus urinae is increasingly recognized as a potentially significant urinary tract bacterium. A. urinae has been isolated from urine collected from both males and females with a wide range of clinical conditions, including urinary tract infection (UTI), urgency urinary incontinence (UUI), and overactive bladder (OAB). A. urinae is of particular clinical concern because it is highly resistant to many antibiotics and, when undiagnosed, can cause invasive and life-threatening bacteremia, sepsis, or soft tissue infections. Previous genomic characterization studies have examined A. urinae strains isolated from patients experiencing UTI episodes. Here, we analyzed the genomes of A. urinae strains isolated as part of the urinary microbiome from patients with UUI or OAB. Furthermore, we report that certain A. urinae strains exhibit aggregative in vitro phenotypes, including floccing, which can be modified by various growth medium conditions. Finally, we performed in-depth genomic comparisons to identify pathways that distinguish floccing and nonfloccing strains.

IMPORTANCE Aerococcus urinae is a urinary bacterium of emerging clinical interest. Here, we explored the ability of 24 strains of A. urinae isolated from women with lower urinary tract symptoms to display aggregation phenotypes in vitro. We sequenced and analyzed the genomes of these A. urinae strains. We performed functional genomic analyses to determine whether the in vitro hyperfloccing aggregation phenotype displayed by certain A. urinae strains was related to the presence or absence of certain pathways. Our findings demonstrate that A. urinae strains have different propensities to display aggregative properties in vitro and suggest a potential association between phylogeny and floccing.

KEYWORDS Gram positive, aggregation, bladder, lower urinary tract symptoms, urinary microbiome

The urinary tract is one of the most common sites of bacterial infection. In addition to detecting recognized uropathogens such as Escherichia coli, hospital microbiology labs routinely find “clinically insignificant” numbers of bacteria in urine. Recent studies have revealed evidence of diverse microbes (the microbiota) in the urinary bladders of many adults (1–4). Furthermore, these studies have found that the microbiota of adult female bladders differ between women with and without lower urinary tract symptoms, including those of urgency urinary incontinence (UUI) and urinary tract infection (UTI) (5, 6). Several bacterial genera, often the Gram-positive Actinobaculum (formerly Actinobaculum), Streptococcus, and Aerococcus, are being referred to as “emerging uropathogens,” since they have been isolated in large numbers in urine.


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Address correspondence to Nicole M. Gilbert, gilbert@wustl.edu, or Alan J. Wolfe, awolfe@luc.edu.

* Present address: Evann E. Hilt, Department of Pathology and Laboratory Medicine, University of California Los Angeles, Los Angeles, California, USA; Krystal Thomas-White, Department of Microbiology & Immunology, Stanford University School of Medicine, Stanford, California, USA.

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samples from patients exhibiting lower urinary tract symptoms and have been increasingly implicated as causes of UTI (7). However, relatively little is known about the molecular features that enable these species to cause bacteriuria, especially in comparison to our understanding of *E. coli* UTI.

*Aerococcus urinae* is a Gram-positive alpha-hemolytic bacterium first isolated and identified from urine from patients with UTI (8). Since its discovery, *A. urinae* has been isolated from a wide variety of patients, both male and female, ranging from children to the elderly (9). In addition to being a suspected cause of UTI, *A. urinae* has been implicated in cases of bacteremia, musculoskeletal infections, infective endocarditis, and necrotizing soft tissue infections (10–13). Even though *A. urinae* is increasingly recognized and identified by clinical microbiology laboratories, it can still be mischaracterized as either streptococci or staphylococci, due to similarities in morphology and Gram stain results (14). Misidentification is a significant clinical concern, because *A. urinae* is highly resistant to many sulfonamides and other antibiotics used to treat UTIs. When undiagnosed or misdiagnosed, *A. urinae* can cause invasive and life-threatening sepsis (9, 15, 16).

Previous studies have focused on *A. urinae* isolated from cases of UTI. In our previous studies, *A. urinae* was frequently detected in women with conditions associated with lower urinary tract symptoms, including urgency urinary incontinence (UUI) and overactive bladder (OAB) (5, 6, 17). Here, we characterized 24 *A. urinae* strains, isolated primarily from women experiencing lower urinary tract symptoms but not experiencing a UTI (as classically defined), for two aggregation in vitro phenotypes. We performed a comprehensive bioinformatic analysis on the whole genomes of these strains and examined the relationship between the genotypes and in vitro aggregation phenotypes.

**RESULTS**

*Aerococcus urinae* displays in vitro aggregation phenotypes. Aggregation and biofilm formation are key features for many pathogenic bacteria, including those of the urinary tract, because they aid in protection from the immune response and antimicrobial agents and thus allow for persistence (18). A previous study found that *A. urinae* induces platelet aggregation and forms biofilms on plastic in standing culture, demonstrating the aggregative capacity of this bacterium (19). We found that *A. urinae* exhibits two additional in vitro aggregation phenotypes (Fig. 1 and Table 1). The first is a “flocking” phenotype, whereby the bacteria stick to each other and aggregate during static growth in liquid culture. Of the 24 *A. urinae* strains we tested, 21% (5/24) flocked in the two different media tested, brain heart infusion broth (BHIB) and tryptic soy broth (TSB) (Fig. 1A, middle), 25% (6/24) flocked in one of the media, while 54% (13/24) did not flock regardless of the medium composition we tested (Table 1; Fig. 1A, left). Three strains exhibited a “hyperflocking” phenotype in both the TSB and BHIB liquid media (Fig. 1A, right); we define hyperflocking as larger aggregates that remain intact after vortexing for up to 1 min and when cells are washed in sterile phosphate-buffered saline (PBS) (see Fig. S1 in the supplemental material).

A second aggregation characteristic of *A. urinae* was revealed when we performed the so-called “hockey puck test,” in which a bacterial colony is pushed across the surface of an agar plate supplemented with glucose. This test is used as a diagnostic tool for the respiratory pathogen *Moraxella catarrhalis* (20). In this test, some *A. urinae* colonies maintained their shape when pushed across the surface of an agar plate (Fig. 1B, top), while *A. urinae* colonies negative for the phenotype were unable to maintain shape (Fig. 1B, bottom). The majority of the 24 tested *A. urinae* strains (54% [13/24]) formed the “hockey puck” phenotype regardless of the medium composition (Table 1), 29% (7/24) formed hockey pucks on one medium, and only 17% (4/24) did not form a hockey puck on either medium. When comparing the two in vitro phenotypes, some strains (5/24) were positive for both phenotypes regardless of medium composition, including the 3 hyperflocking strains; in contrast, a few strains (3/24) were negative for both phenotypes regardless of medium composition. Together, these data demon-
FIG 1. *A. urinae* urinary strains display in vitro aggregation phenotypes. (A) The flocking phenotype. *A. urinae* forms aggregates in a liquid culture when grown statically in TSB or BHI for 48 h under 5% CO₂ atmospheric conditions. *A. urinae* cells gather at the bottom of the tube, but once the tube is disturbed by flicking, the ability of *A. urinae* cells to form aggregates becomes apparent. (B) Hockey puck phenotype. When *A. urinae* is grown on BHI + Glu (or TSA + Glu [not shown]) agar plates for 48 h under 5% CO₂ atmospheric conditions (strain UMB0080), the resultant colonies maintain their shape when pushed across the agar surface (indicated by the arrow and inset of top image). When *A. urinae* does not have the hockey puck phenotype (UM8722), the colonies are unable to maintain their shape (indicated by the arrow and inset of bottom image). Experiments to observe the in vitro phenotypes were performed in triplicates more than 20 times.

strate that *A. urinae* exhibits self-adhesive properties and that phenotypic differences in these properties vary between individual strains.

**Genomic diversity among *A. urinae* strains.** To gain further insight into the potential genomic basis for the phenotypic differences we observed in aggregative

<table>
<thead>
<tr>
<th>Strain collection ID*</th>
<th>Disease*</th>
<th>Species</th>
<th>Flocking phenotype on: TSB</th>
<th>BHI</th>
<th>Hockey puck phenotype on: TSA+Glu</th>
<th>BHI+Glu</th>
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*ID, identifier.

*UUI, urgency urinary incontinence; OAB, overactive bladder; UTI, urinary tract infection; SUI, stress urinary incontinence.
properties between *A. urinæ* strains, we performed whole-genome sequencing. We sequenced the genomes of the 24 *A. urinæ* strains and those of 3 other *Aerococcus* species. For the 24 *A. urinæ* strains, the number of *de novo* assembled scaffolds ranged from 16 to 661; the genome sizes averaged 2.16 ± 0.26 Mb, with on average 1,927 ± 272 protein-coding regions (see Table S1). At the time we completed this study, only five *A. urinæ* genomes were publicly available through GenBank. These five publicly available *A. urinæ* strains were isolated from a variety of sources from human patients with diverse diagnoses (see Table S2). Genome annotation of the five previously sequenced genomes and our 24 newly sequenced genomes revealed a core *A. urinæ* genome composed of 1,355 genes (see Table S3). The vast majority of the functionality encoded in the core genome remains unknown, as only 34% (456/1355) of these genes have a predicted function. Nevertheless, among the proteins with annotated functions include several putative virulence factors, including a member of the hemolysin II family, and homologs of fibronectin-binding protein A, conserved virulence factor B, and choline-binding protein.

To compare the genomes of the 24 *A. urinæ* strains isolated from the female urinary bladder to those of the five publicly available strains, we performed a phylogenetic analysis as described in Materials and Methods. All 29 *A. urinæ* strains were highly related (Fig. 2; see also Fig. S2). Strains within the phylogenetic trees were overlaid with color coding based on the flocking and hockey puck *in vitro* phenotypes. Associations were not immediately apparent with the hockey puck phenotype, as the negative strains were distributed across the tree (Fig. 2, right). In contrast, the strains positive for flocking clustered together, with the hyperflocking strains being most closely related. This suggests a potential association between phylogeny and the flocking phenotype (Fig. 2, left).

*A. urinæ* is cytotoxic to human urothelial cells. In an effort to understand the consequence of *A. urinæ* interaction with the bladder urothelium, we asked if *A. urinæ*
was cytotoxic to human urothelial cells (HUCs). To test the cytotoxicity effects of *A. urinae* to HUCs, we infected monolayers of HUCs with *A. urinae* strains UMB0080, UMB0722, and UMB3669 at a multiplicity of infection (MOI) of 300 (10⁴ cells). We compared *A. urinae* cytotoxicity to that of uropathogenic *E. coli* (UPEC), group B *Streptococcus* (GBS), and *Lactobacillus crispatus* (LC40) at the same MOI. At 3 h post-inoculation (hpi), HUCs were stained with trypan blue solution to determine the percentage of cells that remained viable (Fig. 3). These results demonstrate that all three *A. urinae* strains, regardless of their ability to flock, caused a statistically significant decrease in HUC viability compared to that of untreated HUCs, with viability ranging between 45% and 60%. The level of HUC cytotoxic caused by *A. urinae* was similar to those of UPEC strains UTI89 and NU14 and GBS strains UMB1026 and UMB1047 (Fig. 3). In contrast, *L. crispatus* (LC40) was not cytotoxic to HUCs (Fig. 3). This finding is consistent with a previous study that detected *L. crispatus* almost exclusively in asymptomatic women, supporting the hypothesis that *L. crispatus* is a commensal bacterium in the bladder (5). The observation that *A. urinae* is cytotoxic aligns with the observed presence of putative hemolysin genes and their known alpha-hemolytic behavior on blood agar and further suggests that the putative hemolysin may also be cytotoxic to the bladder epithelium.

**Functional annotation comparison of hyperflocking strains versus nonflocking strains.** Since the three hyperflocking strains clustered together in our phylogenetic analysis (Fig. 2), we performed a more in-depth genomic analysis to determine whether these strains harbored a certain gene(s) or pathways that were absent from nonflocking strains. We assessed the gene annotations for each genome at each of the following functional classification levels: category, subcategory, subsystem, and role. Compared to the 13 nonflocking strains, the 3 hyperflocking strains were significantly enriched for coding DNA sequences (CDS) predicted to encode proteins involved in β-glucoside metabolism and l-ascorbate utilization (Fig. 4). Specifically, with respect to β-glucoside metabolism, there are 11 genes present in the hyperflocking strains that encode proteins in the pathway, and all 11 genes are absent in the nonflocking strains (see Table S4). The majority of the 11 encoded proteins are members of two phosphotrans-
FIG 4 Functional annotation comparison of hyperflocking strains versus nonflocking strains. (A) Coding sequence difference. Average coding sequence difference between the hyperflocking and nonflocking strains. The graph is pared down to the seven specific subsystems that depict either CDS being higher in the hyperflocking strains than in nonflocking strains or completely absent from the hyperflocking strains. The colored dot under each pair of bars corresponds to the name of the subsystem in panel B. (B) Phylogenetic tree with specific subsystem comparison. This phylogenetic tree is a modified version of the trees shown in Fig. 2. This tree focuses on only the hyperflocking and nonflocking strains. Next to each strain is an array of colored dots that correspond to the specific subsystems that are present in the genome of that strain.

ferase systems (PTSs) used for importing and metabolizing β-glucoside sugars. Conversely, the hyperflocking strains were depleted for CDS involved in tryptophan synthesis, auxin biosynthesis, d-sorbitol (d-glucitol) and l-sorbose utilization, d-tagatose and galactitol utilization, and mannose metabolism (Fig. 4). In conclusion, genome comparisons revealed significant differences between hyperflocking and nonflocking strains and highlighted multiple pathways that could potentially contribute to in vitro aggregation phenotypes which can serve as a basis for future mechanistic investigations.

DISCUSSION

The recent implementation of mass spectrometry-based identification of bacteria in the clinical microbiology lab has highlighted novel agents of disease known as emerging uropathogens (7). These are primarily Gram-positive bacteria that have been repeatedly isolated in high numbers from urine samples of patients reporting lower urinary tract symptoms. One emerging uropathogen of particular interest is A. urinae. In the present study, we explored the in vitro ability of 24 clinical strains to display in vitro aggregative phenotypes, sequenced and analyzed their genomes, and determined whether the in vitro phenotypes correlated with their cytotoxicity, phylogeny, or genomic functional pathways.

By analyzing 24 clinical strains from cases of UUI or OAB, we observed two in vitro biofilm-associated phenotypes that support and extend a previous report that A. urinae can form biofilms (19). Depending on the growth conditions, the strains exhibited an array of phenotypic profiles. The flocking phenotype was observed in liquid media without the addition of any sugars. In contrast, the hockey puck phenotype was observed only when agar plates were supplemented with glucose. These results suggest the existence of a signal associated with sugar metabolism that induces the hockey puck phenotype. Future studies to explore the nature of those signals are needed.

The first complete genome sequence of A. urinae was made available in 2016 for an A. urinae strain (CCUG 36881T) isolated from an individual experiencing a UTI episode. An additional study reported whole-genome sequences and phylogenetic characterization of A. urinae strains isolated from individuals experiencing episodes of UTI and bacteremia and included more male than female patients (21, 22). To our knowledge,
the present study is the first comprehensive bioinformatic analysis of *A. urinae* strains isolated from the urine of women with other conditions associated with lower urinary tract symptoms (UUI and OAB). Furthermore, much of the existing literature reports *A. urinae* in males, especially in cases where it has been known to cause more invasive disease (9). This apparent male dominance likely results from a paucity of studies, as our team and others have repeatedly detected *A. urinae* in perimenopausal women with lower urinary tract symptoms (5, 6, 23). In all of our studies combined (844 women), we have cultured *A. urinae* from a nonsymptomatic woman only twice. The vast majority have been cultured from women experiencing lower urinary tract symptoms, most often those associated with UUI and OAB. At present, it is unclear whether *A. urinae* is a cause of or contributor to these symptoms. While the present study focused on *A. urinae* strains isolated from patients with UUI or OAB, future studies are warranted to compare these to *A. urinae* strains isolated from patients with UTI and bacteremia, both at the genomic level and with respect to *in vitro* aggregation phenotypes.

Although the 24 sequenced *A. urinae* strains were all highly related (Fig. 2), we identified some potentially important differences between strains based on their *in vitro* aggregative characteristics. Because the 3 hyperflocking strains clustered phylogenetically, we compared their genomes to those of the 13 strains that did not flock and found significant differences in their predicted metabolic capacities. The first of these differences was the complete absence from the nonflocking strains of an operon that is predicted to encode two systems for the import and initial metabolism of β-glucosides. The epithelial lining of the bladder, known as the urothelium, is lined with a polysaccharide glycosaminoglycan (GAG) layer. This GAG layer is known to contain β-linked disaccharide repeating units (24). It is possible that the ability of hyperflocking strains to metabolize these β-disaccharide units could enhance their survival *in vivo*. Another difference is that nonflocking strains have a complete tryptophan and auxin biosynthesis pathway, while the hyperflocking strains lack the tryptophan synthase α chain. The tryptophan synthase α chain catalyzes the last step in *de novo* biosynthesis of tryptophan and is thought to contribute to auxin biosynthesis (25); the absence of this gene in the hyperflocking strains suggests that they cannot synthesize tryptophan or auxin. Auxin, also known as indole-3-acetic-acid (IAA), is a phytohormone important for plant cell growth and development and is produced by both symbiotic and pathogenic plant bacteria (26). Indole has been shown to be an intercellular signaling molecule between bacteria in the gut microbiome (27), and it is possible that similar signaling events occur between *A. urinae* and the bladder microbiota. Additional studies are needed to determine the role of β-glucoside metabolism and auxin in *A. urinae* uropathogenesis.

Many well-characterized uropathogens have aggregative properties that contribute to virulence (7). Future studies using *in vivo* small animal models, such as mice, should examine whether the aggregative behaviors displayed by *A. urinae* *in vitro* are related to the ability to infect the urinary tract. It is possible that aggregative *A. urinae* strains could be better able to attach to the urothelium or avoid phagocytosis by neutrophils. It is also possible that growth under flocking-inducing conditions differentially regulates certain metabolic pathways that render *A. urinae* more capable of colonizing and surviving in the urinary tract. With the information gained from genomic sequencing and pathway analysis, future studies can also examine whether differing metabolic capabilities between *A. urinae* strains may also contribute to their ability to acquire nutrients and persist in the bladder. All together, these studies highlight the potential importance of *A. urinae* aggregative properties and provide important tools to further study this important urinary bacterium.

**MATERIALS AND METHODS**

**Aerococcus urinae** strain isolation, identification, and characterization. Twenty-four clinical isolates of *Aerococcus urinae* were collected from transurethral catheter urine samples of 21 female patients between 2013 and 2016. Following Loyola University Medical Center (LUMC) Institutional Review Board approval, patients gave verbal and written consent for chart abstraction and urine collection with analysis for research purposes. Patients were recruited as part of separate studies (5, 23, 28, 29).
Matrix-assisted laser desorption ionization–time of flight mass spectrometry (MALDI-TOF MS) with the MALDI Biotyper 3.0 software program (Bruker Daltonics, Billerica, MA) was used to identify the bacterial strains, as described elsewhere (2).

The 24 clinical A. uinae strains were isolated from women with and without lower urinary tract symptoms (Table 1). Nine strains were collected from seven patients diagnosed with overactive bladder (OAB), nine strains were from eight patients diagnosed with UUI, and three strains were from three patients diagnosed with stress urinary incontinence (SUI). One A. uinae strain was cultured from a case of UTI, as defined by patient symptoms and culture of ≥100,000 CFU/ml of A. uinae as the sole uropathogen, verified by the clinical microbiology lab using MALDI-TOF MS as described above. Finally, two strains were cultured from two asymptomatic control patients. Of note, the three hyperflocking strains were isolated from different patients, collected months apart.

In addition to the 24 clinical strains of A. uinae, strains from 3 other Aerococcus species were selected for the genome comparison analysis described below. All three were isolated via the same method and during the same time period as indicated for the 24 A. uinae strains. Aerococcus christensenii (UMBO884) was cultured from an asymptomatic control patient, while Aerococcus sanguinicola (UMBO139) and Aerococcus viridans (UMBO240) were cultured from two different patients with OAB.

**In vitro aggregation phenotypes.** To observe in vitro phenotypes, all 24 A. uinae strains were first grown on a 5% sheep blood agar plate (BAP) (BD BBL prepared plated medium) and incubated under 5% CO2 atmospheric conditions at 35°C for 48 h. To observe the floccing phenotype, A. uinae was subcultured in either tryptic soy broth (TSB) or brain heart infusion broth (BHI) and then grown statically under 5% CO2 atmospheric conditions at 35°C for 48 h. To observe the hockey puck phenotype, A. uinae was subcultured onto either tryptic soy agar (TSA) plates supplemented with 0.5% glucose (TSA + Glu) or brain heart infusion (BHI) agar plates supplemented with 0.5% glucose (BHI + Glu) and grown under 5% CO2 atmospheric conditions at 35°C for 48 h. Experiments to observe the in vitro phenotypes were performed in triplicates more than 20 times.

**DNA extraction.** Each of the 24 A. uinae strains and the three Aerococcus sp. strains were grown on 5% sheep blood agar plates (BAPs) (BD BBL prepared plated medium) under 5% CO2 conditions for 48 h and then pelleted. To extract genomic DNA, cells were resuspended in 0.5 ml DNA extraction buffer (20 mM Tris-CI, 2 mM EDTA, 1% Triton X-100, pH 8) followed by the addition of 50 µl lysozyme (20 mg/ml) and 30 µl mutanolysin. After a 1-h incubation at 37°C, 80 µl 10% SDS and 20 µl proteinase K were added, followed by 2 h of incubation at 55°C. Then, 210 µl of 6 M NaCl and 700 µl phenolchloroform were added. After a 60-min incubation with rotation, the solution was centrifuged at 13,500 rpm for 10 min, and the aqueous phase was discarded. An equivalent volume of isopropanol was added, after a 10-min incubation, the solution was centrifuged at 13,500 rpm for 10 min. The supernatant was decanted, and the final pellet was precipitated using 600 µl 70% ethanol. Following ethanol evaporation, the DNA pellet was resuspended in nuclease-free H2O and stored at −20°C.

**Genome sequencing, assembly, and annotation.** Genomic DNA was diluted in water to a concentration of 0.2 ng/µl. Library preparation was performed using the Nextera XT DNA library preparation kit (Illumina) according to the manufacturer’s instructions with 1 ng of input DNA. Each strain was sequenced using the Illumina MiSeq platform and the MiSeq reagent kit v2 (300 cycles) to produce 150-bp paired-end reads. Sequencing reads were parsed into individual folders according to the respective barcodes. The resulting sequence reads were trimmed with Sickle (30), and the assembly was performed using the SPAdes assembler (31).

**Phylogenetic analysis and average nucleotide identity analysis.** The phylogenetic analysis was conducted by extracting amino acid sequences for 40 universal single copy marker genes (32-33) from the Aerococcus genomes using SpecI (34). A. christensenii (UMBO884), A. sanguinicola (UMBO139), and A. viridans (UMBO240) were used to root the tree of all 24 A. uinae strains and the 5 publicly available strains of A. uinae. The protein sequences were concatenated and aligned with Mafft v. 7.20 (35), and maximum likelihood trees were constructed using FastTree (36) with default settings. All phylogenetic trees were visualized in iTOL (37).

**Cytotoxicity assay.** To test the cytotoxicity of A. uinae cells, HUCs were stained with trypan blue staining protocol. A. uinae cells were prepared by inoculation in liquid BHI cultures for 48 h and then standardizing the 48-h culture to an optical density at 600 nm (OD600) of 1.0 before resuspending in Dulbecco’s modified eagle’s medium (DMEM) supplemented with 5% fetal bovine serum (FBS), UPEC and GBS strains were prepared by inoculation in liquid BHI cultures for 24 h, standardization of the 24-h culture to an optical OD600 of 1.0, and then resuspension in DMEM plus 5% FBS. L. crispatus cells were prepared by inoculation in liquid cultures for 48 h, standardization of the 48-h culture to an optical OD600 of 1.0, followed by resuspension in DMEM plus 5% FBS. The HUC cell line used in these experiments (Huro18H7) was obtained from the bladders of consenting female patients described in a previous study (38). To prepare the HUCs, the day before inoculation, HUCs were counted with a hemocytometer and then diluted to 1×10⁷ cells/ml in thymic epithelial (TE) medium (38). Next, 1 ml aliquots of uterine cells were placed in a 24-well tissue culture-treated plate and incubated overnight at 37°C in 5% CO2 to allow the cells to adhere to the well. HUCs were treated with the resuspended A. uinae, UPEC, GBS, or L. crispatus. As a control, bacteria resuspended in 1 ml of DMEM plus 5% FBS were also inoculated in a well without uterine cells. For negative controls, uterine cells were treated with various toll-like receptor (TLR) agonists: lipopolysaccharide (LPS) (20 ng; Enzo Life Sciences), Pam3 (20 ng; InvivoGen), and Pam2 (30 ng; InvivoGen). After 3 h of treatment, the percent viability of the uterine cells was measured with trypan blue (Lonza) staining for dead cells. In addition, bacterial cells were plated for CFU prior to and after treatment of the uterine cells.
Functional genomic analysis. Each genome was annotated using RAST (39). A custom python script was written to conduct reciprocal blastp queries using the BLAST+ package (40) in order to identify homologous genes among the A. urinae genomes. Referring to RAST-predicted annotations, function assignments were made for each identified gene family. Custom python scripts were developed to identify gene families associated within phenotypic and phylogenetic groups.

Functional analysis also was performed using RAST. RAST gene annotations utilize the manually curated hierarchical subsystem library (41), which includes functional assignment covering all modules of cellular machinery. A custom python script was written to identify the number of unique coding regions for each genome and each level of functional classification: category, subcategory, subsystem, and role. Next, pairwise genome comparisons of the number of unique coding regions for each level of functional classification were conducted. Presence/absence statistics were calculated for all genomes for each functional classification and manually inspected further.

Data availability. Raw sequence data and assembled genomes were deposited in NCBI’s SRA and WGS databases, respectively. SRA record numbers are SRR6973109 through SRR6973128, and WGS record numbers are QMG100000000 through QMHM00000000.

SUPPLEMENTAL MATERIAL
Supplemental material is available online only.

SUPPLEMENTAL FILE 1, PDF file, 1.1 MB.

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