Transposon Insertion Site Sequencing of *Providencia stuartii*: Essential Genes, Fitness Factors for Catheter-Associated Urinary Tract Infection, and the Impact of Polymicrobial Infection on Fitness Requirements

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**ABSTRACT** *Providencia stuartii* is a common cause of polymicrobial catheter-associated urinary tract infection (CAUTI), and yet literature describing the molecular mechanisms of its pathogenesis is limited. To identify factors important for colonization during single-species infection and during polymicrobial infection with a common cocolonizer, *Proteus mirabilis*, we created a saturating library of ~50,000 transposon mutants and conducted transposon insertion site sequencing (Tn-Seq) in a murine model of CAUTI. *P. stuartii* strain BE2467 carries 4,398 genes, 521 of which were identified as essential for growth in laboratory medium and therefore could not be assessed for contribution to infection. Using an input/output fold change cutoff value of 20 and \( P \) values of <0.05, 340 genes were identified as important for establishing single-species infection only and 63 genes as uniquely important for polymicrobial infection with *P. mirabilis*, and 168 genes contributed to both single-species and coinfection. Seven mutants were constructed for experimental validation of the primary screen that corresponded to flagella (*fliC* mutant), twin arginine translocation (*tatC*), an ATP-dependent protease (*clpP*), \( \alpha \)-alanine-\( \beta \)-alanine ligase (*ddIA*), type 3 secretion (*yscl* and *sopB*), and type VI secretion (*impJ*). Infection-specific phenotypes validated 6/7 (86%) mutants during direct cochallenge with wild-type *P. stuartii* and 3/5 (60%) mutants during coinfection with *P. mirabilis*, for a combined validation rate of 9/12 (75%). Tn-Seq therefore successfully identified genes that contribute to fitness of *P. stuartii* within the urinary tract, determined the impact of coinfection on fitness requirements, and added to the identification of a collection of genes that may contribute to fitness of multiple urinary tract pathogens.

**IMPORTANCE** *Providencia stuartii* is a common cause of polymicrobial catheter-associated urinary tract infections (CAUTIs), particularly during long-term catheterization. However, little is known regarding the pathogenesis of this organism. Using transposon insertion site sequencing (Tn-Seq), we performed a global assessment of *P. stuartii* fitness factors for CAUTI while simultaneously determining how coinfection with another pathogen alters fitness requirements. This approach provides four important contributions to the field: (i) the first global estimation of *P. stuartii* genes essential for growth in laboratory medium, (ii) identification of novel fitness factors for *P. stuartii* colonization of the catheterized urinary tract, (iii) identification of core fitness factors for both single-species and polymicrobial CAUTI, and (iv) assessment of conservation of fitness factors between common uropathogens. Genomewide assessment of the fitness requirements for common uropathogens during single-species
and polymicrobial CAUTI thus elucidates complex interactions that contribute to disease severity and will uncover conserved targets for therapeutic intervention.

**KEYWORDS** Proteus mirabilis, Providencia stuartii, Tn-Seq, catheter, essential genes, pathogenesis, polymicrobial, transposon, urinary tract infection

*Providencia stuartii* is an opportunistic pathogen frequently isolated from the urine of patients with long-term indwelling urinary catheters (1–3). *P. stuartii* is well adapted for colonizing the catheterized urinary tract, where it persists for months (4, 5). As *P. stuartii* isolates are often resistant to multiple antibiotics (1, 6–8) and frequently cocolonize with other organisms (4, 5, 9–13), they contribute to the spread of antimicrobial resistance determinants to other organisms during cocolonization and to the risk of treatment failure. *P. stuartii* urine colonization is also a common source of bacteremia in long-term care facilities (8, 14–18). Up to 51% of *P. stuartii* bacteremias are polymicrobial (16, 19), suggesting that *P. stuartii* polymicrobial interactions promote development of bacteremia.

Despite the clinical significance of this organism, no previous studies explored *P. stuartii* genes essential for growth in vitro and little is known regarding *P. stuartii* virulence factors. Adherence, biofilm formation, host cell invasion, and motility are important aspects of urinary tract colonization for numerous uropathogenic bacteria, and in vitro and in silico evidence suggests that these properties may contribute to *P. stuartii* pathogenesis (20–26). However, their contribution to colonization and pathogenesis has not been assessed experimentally.

The polymicrobial nature of catheter-associated urinary tract infection (CAUTI) is an important consideration in investigating bacterial virulence factors. *Proteus mirabilis*, another common CAUTI pathogen, frequently cocolonizes the catheterized urinary tract with *P. stuartii* (4, 5, 9–12). In a mouse model of CAUTI, coinfection with *P. stuartii* and *P. mirabilis* increased inflammation and infection severity and altered the fitness factors required by *P. mirabilis* to colonize and persist within the urinary tract (23, 27). For instance, *P. mirabilis* did not require branched-chain amino acid biosynthesis during single-species infection, but coinfection with *P. stuartii* resulted in a requirement for these genes due to its high-affinity branched-chain amino acid importer (27). For common pathogens of polymicrobial infections, it is therefore imperative to define core genetic requirements for pathogenicity during both single-species infection and polymicrobial infection.

In this study, we utilized transposon insertion site sequencing (Tn-Seq) to identify *P. stuartii* genes essential for growth in rich medium, the full arsenal of fitness factors that contribute to single-species infection in a CAUTI model, and how fitness requirements change during polymicrobial infection. *P. stuartii* was found to carry a core set of 168 genes that contribute to colonization of the urinary tract during both single-species and polymicrobial infection, as well as accessory genes that contribute to only one infection type. Notably, several core genes were also identified as *P. mirabilis* fitness factors by Tn-Seq that were indicative of common fitness requirements for multiple urinary tract pathogens. These common fitness requirements may represent targets for treating or preventing CAUTI.

**RESULTS**

**Estimation of essential genes.** Based on the size of the *P. stuartii* BE2467 genome (4.635 Mbp, ~4,398 genes [23]), 39,099 transposon mutants are required for >99.9% probability of full-genome coverage (28). We therefore generated five pools containing $1 \times 10^4$ transposon mutants each for a total of 50,000 mutants. Insertion randomness was verified by Southern blotting (see Fig. S1 in the supplemental material). A total of 5 to 10 CBA/J mice were transurethrally inoculated with $1 \times 10^5$ CFU of one of five transposon library pools for single-species infection or with $1 \times 10^5$ CFU of a 1:1 mixture of the transposon library and wild-type (WT) *P. mirabilis* HI4320 for coinfection

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Thus, for each input pool, single-species infections and coinfections were conducted in parallel to utilize the same input inoculum. Genome saturation was achieved for the \textit{P. stuartii} BE2467 chromosome, plasmid PS1, and plasmid PS2, with most insertion sites represented in at least two input pools (Fig. 2). A Bayesian mixture model identified candidate genes essential for growth under laboratory conditions based on absence or underrepresentation of transposon insertions in these genes within the input pools (see Materials and Methods). The model identified 521 genes (12% of the genome) as potentially essential for growth in LB medium (see Table S1 in the supplemental material), representing a number comparable to the 436 essential genes (11.6% of the genome) of \textit{P. mirabilis} HI4320 (27). Twelve of the genes estimated to be essential were carried on pPS1, including 8 hypothetical genes, an antitoxin gene (\textit{higA}), a gene encoding a DNA-binding protein (\textit{HU}), an arsenate reductase gene, and an arsenical resistance operon repressor; 8 were carried on pPS2, all of which are hypothetical. The potential essentiality of these genes should be interpreted with caution, as it is possible that the mariner transposon was unable to efficiently target these locations or that their loss impacts plasmid stability.

Among the 521 genes identified as essential, 402 could be grouped based on classification of gene category within the Clusters of Orthologous Groups (COG) database (Fig. 3), including genes associated with rRNAs, ribosomal proteins, tRNA synthetases and reductases, ATP synthase, the Sap peptide transport system, the Lol lipoprotein transport system, the Lpt lipopolysaccharide transport system, the Sec protein translocase system, and factors involved in DNA replication, cell division, and cell wall and membrane biogenesis. Several genes encoding metabolic enzymes were also identified, including the majority of the pentose phosphate pathway and phos-
phoglycerate kinase, which is involved in glycolysis, gluconeogenesis, and glycerol degradation. Many of the same factors were identified as essential for growth of *P. mirabilis* in our prior Tn-Seq study (27). Several unexpected genes were also identified, such as those encoding copper resistance (*copC*), a urea channel (*ureI*), a biofilm regulator (*bssS*), and two fimbrial chaperone proteins. While these genes may contribute to growth of *P. stuartii* in LB broth, another possibility is that the transposon could not efficiently target these locations for other reasons, such as a low number of TA sites or restricted access by secondary structure.

**Primary screen of *P. stuartii* transposon mutant libraries.** Infection studies were conducted to screen for *P. stuartii* fitness factors during single-species and polymicrobial CAUTI as described in the Fig. 1 legend. In all cases, a 4-mm-long segment of sterile silicone catheter tubing was inserted into the bladder during inoculation and was

**FIG 2** Saturation of the *Providencia stuartii* BE2467 genome and plasmids with transposon insertions. Chromosomal and plasmid maps indicate the location of all transposon insertions contained in all five of the input pools (not to scale). Each line represents a single insertion site from the input samples, and the length of the line represents the log10 number of reads recovered from each insertion site. The color of the line indicates the number of pools in which each insertion site was identified as follows: light blue, all five pools; dark blue, four pools; brown, three pools; light green, two pools; red, one pool. Regions without insertions represent estimated essential genes and intergenic regions.

**FIG 3** Functional categories of *P. stuartii* genes estimated to be essential for growth in LB medium. A total of 402 of the 521 genes estimated to be essential for growth of *P. stuartii* strain BE2467 had a Cluster of Orthologous Groups (COG) category. The size of each wedge of the pie chart corresponds to the percentage of genes belonging to each COG.
Catheter segments were not removed prior to homogenization of bladder samples, so the CFU numbers recovered from bladder samples represent colonization of the catheter and bladder. Infection with *P. stuartii* transposon mutant pools resulted in comparable bladder and kidney bacterial burdens at 4 days postinoculation (dpi) for single-species infection and coinfection (Fig. 4A and B). The majority of coinfected mice were highly colonized by both bacterial species, as expected (Fig. 4C). Four mice per infection type per input pool were chosen for sequencing based on colonization levels of *P. stuartii* and high-quality DNA during sample preparation. A fitness index was calculated for each gene as previously described (27, 29).

**P. stuartii** fitness factors during single-species infection. A total of 508 genes (11% of the genome) were identified by Tn-Seq as candidate fitness factors for single-species infection, with fold changes in insertion coverage for input/outputs ranging from 20 to 1,841 (Table S1). Among the genes, 85 encoded fitness factors for colonization of both bladder and kidneys, 414 encoded kidney-specific fitness factors, and 9 were bladder specific. Thus, *P. stuartii* appears to undergo an initial selection process in the catheterized bladder before ascending to the kidneys.

Among the 508 fitness factors for single-species infection, 378 were present in COG (Fig. 5A). Notably, initial Tn-Seq results confirmed our prior investigations of *P. stuartii* urease. Six of the eight genes in the *P. stuartii* plasmid-carried urease operon were underrepresented in bladder and kidney outputs compared to inputs, but the fold change values ranged from 2 to 10 and did not meet our stringent cutoff of 20. The lack of a significant fitness defect for the urease operon may have been due to the ability of other urease-positive mutants in the pool to compensate for the loss of urease activity. However, our prior study of *P. stuartii* CAUTI indicated that loss of the entire urease operon resulted only in a nonsignificant trend toward reduced colonization of the bladder and kidneys (23). Thus, *P. stuartii* urease is not critical for CAUTI in this model.

**P. stuartii** fitness factors during polymicrobial infection. Tn-Seq analyses of fitness factors from coinfection of *P. stuartii* BE2467 with *P. mirabilis* HI4320 revealed 231 candidate fitness factors (5% of the genome), with fold changes in insertion...
A total of 74 were important for colonization of both the catheterized bladder and the kidneys, 86 were kidney-specific fitness factors, and 71 were bladder specific (Fig. 5A). Preliminary assessment revealed concordance with our prior investigation of branched-chain amino acid import in *P. stuartii*. Leucine-specific binding protein LivK was previously found to contribute to bladder colonization during coinfection with *P. mirabilis* (27) and was identified as a fitness factor for bladder colonization during coinfection in the current study (25-fold change, $P < 0.05$).

**Comparison of *P. stuartii* fitness factors for single-species versus polymicrobial CAUTI.** In total, 571 genes were identified by Tn-Seq as being fitness factors for *P. stuartii* CAUTI; 340 were specific for single-species infection (11 carried on pPS1 and 7 on pPS2), 63 were specific for polymicrobial infection (1 carried on pPS1 and 1 on pPS2), and 168 contributed to both infection types (1 carried on pPS2). A diagram of overlap of the fitness factors for each infection type and organ is presented in Fig. 5B.

Overall, there appear to be more similarities than differences in COG categories of fitness factors required by *P. stuartii* for single-species infection versus polymicrobial infection. One notable exception pertains to adhesins; 8 of the fitness factors specific to single-species infection encode putative fimbrial proteins and 3 of the fitness factors for both infection types were fimbrial proteins, but none were uniquely identified for coinfection. Thus, coinfection with *P. mirabilis* may alleviate the requirement for certain adhesins. This is in contrast to *P. mirabilis*, in which distinct fimbrial types were identified as fitness factors for coinfection compared to single-species infection (27). On the basis of these observations, *P. mirabilis* may require an array of adhesins to bind
various sites within the urinary tract and form extracellular bacterial clusters and biofilms, while *P. stuartii* may predominantly take advantage of structures established by *P. mirabilis* during coinfection.

The identification of 168 genes as candidate fitness factors for *P. stuartii* during both single-species infection and polymicrobial infection with *P. mirabilis* indicates that these genes likely encode critical functions for colonization of the catheterized urinary tract. For simplicity, we refer to this set of genes as “core” fitness factors to denote their importance during both single-species infection and coinfection in at least one organ (Table S1). To determine if core fitness factors are prevalent and conserved in other *P. stuartii* isolates, we conducted a genome comparison of *P. stuartii* strain BE2467 and 15 other *P. stuartii* complete genome sequences (Fig. S2). Even though *P. stuartii* strain BE2467 was isolated in the 1980s, there is a remarkable degree of genome conservation between this and other strains, including an isolate from 2016 (Fig. S2), and the majority of proteins encoded by *P. stuartii* BE2467 share ≥90% sequence identity with proteins encoded by other *P. stuartii* complete genome sequences (Fig. 6A). Furthermore, 164/168 core fitness factors are present with ≥95% protein sequence identity in all other *P. stuartii* complete genome sequences (Fig. 6B). Core fitness genes are therefore highly conserved in other *P. stuartii* isolates, including recent isolates from geographically distinct locations.

A total of 148 of the 168 core fitness factors were also present in the genome of *P. mirabilis* HI4320, and 20 of these were similarly identified as *P. mirabilis* core fitness factors (27). These factors include a ferredoxin, an iron-sulfur cluster assembly protein (SufB), an alkyl hydroperoxide reductase, the phosphate transport system, a repressor...

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**FIG 6** Proteome conservation between *P. stuartii* isolates. (A) The Pathosystems Resource Integration Center (PATRIC) Proteome Comparison tool was used to compare *P. stuartii* BE2467 (outer ring) with 9 other *P. stuartii* protein sequences, one from each node of the phylogenetic tree in Fig. S2. The three gaps in the blue location indicator separate the chromosome sequence from the two plasmid sequences. (B) A total of 166 candidate fitness factors for *P. stuartii* colonization of the bladder and/or kidneys during both single-species and polymicrobial CAUTI were compared across the same 9 *P. stuartii* isolates as follows (from outside to inside): *P. stuartii* BE2467, NCTC12257, 961029, ATCC 25827, FDAARGOS_87, MRSN 2154, AR_0026, INSRA21868, YD789-2, FDAARGOS_291. The color of each block indicates the protein sequence percent identity compared to *P. stuartii* BE2467.
of the glycerol-3-phosphate regulon, putrescine import, serine import, molybdenum transport, a transporter required for assembly of bd-type respiratory oxidases, and the twin arginine translocation system protein TatC. Identification of these genes as fitness factors in two uropathogens suggests that they may encode key factors required by multiple bacterial species to successfully colonize and persist within the catheterized urinary tract.

Of 168 core fitness factors identified in P. stuartii, 15 were important for colonization in both organs (bladder and kidneys) of all mice, regardless of infection type. This list includes pyruvate kinase (encoded by pykF), anaerobic glycerol-3-phosphate dehydrogenase (glpB), carbamoyl-phosphate synthase (carA), glycoprotein-polysaccharide metabolism (ybaY), the phosphotransferase system (ptsH), thioredoxin (trxA), δ-alanine-δ-alanine ligase (ddlA), ATP-dependent proteases (clpP), and molybdopterin synthase (moaD). Among the members of this list are also three core fitness factors for P. mirabilis: butanediol dehydrogenase (encoded by butA), δ-aminoacyl-tRNA deacylase (dtd), and twin arginine translocation (tatC) (27). These 15 factors therefore pertain to critical functions required for colonization of the catheterized urinary tract, regardless of changes in the bladder environment during polymicrobial infection.

Validation of P. stuartii candidate fitness factors for CAUTI. To validate the primary screen, genes of interest were selected from following categories: (i) fitness factors for single-species infections (flagella [encoded by fliC]), (ii) core fitness factors for both single-species infections and polymicrobial infections (twin arginine translocation [tatC], an ATP-dependent protease [clpP], δ-alanine-δ-alanine ligase [ddlA], and type 3 secretion [T3S] [yscC, yscY, and sopB]), and (iii) fitness factors for polymicrobial infection (type VI secretion [impJ]). Expression of each gene was first assessed after a 2-h incubation in LB broth or filter-sterilized human urine, and recA was utilized as a housekeeping gene for normalization (Fig. 7). All genes were expressed under both sets

![Graph](http://msphere.asm.org/)
of conditions (Fig. S3; see also Fig. 7A). Relative to recA, expression of tatC was significantly increased in urine compared to LB medium, while ddlA and yscI expression levels were significantly decreased (Fig. 7B). Mutants were generated in seven genes of interest for further validation experiments. Importantly, no mutants exhibited significant growth defects in LB medium (Fig. 7C) or minimal medium (Fig. 7D). The following subsections discuss validation of each mutant in further detail.

**Contribution of flagella to *P. stuartii* fitness during single-species CAUTI.** Flagella are well-known virulence factors of several urinary tract pathogens, including *Escherichia coli* and *P. mirabilis*, but their importance to infection has never been assessed for *P. stuartii* beyond a role in swarming motility *in vitro* (22). The *P. stuartii* BE2467 genome encodes production and secretion of flagella in a cluster of 21 genes (BGK56_00565 to BGK56_00660). The flagellar filament (*fliC*) was identified as a candidate fitness factor for kidney colonization during single-species infection, and other genes of the cluster had fitness defects in bladder colonization that ranged from 2-fold to 16-fold and kidney defects that ranged from 5-fold to 33-fold. We therefore generated a *fliC* mutant on MOT agar. Both strains were cultured overnight in LB medium and subjected to stab inoculation into 22 to 24 MOT agar plates for each of three independent experiments. Strain BE2467 was found to be predominantly nonmotile in MOT agar, although motility was observed in 13% of replicate plates after 7 days of incubation. In contrast, motility was never detected in the *fliC* mutant. Panel B shows the combined number of isolates tested and the percentage that exhibited motility. (C) A panel of *P. stuartii* clinical isolates from the urine of catheterized nursing home residents were assessed for motility using motility test medium with triphenyltetrazolium chloride (TTC) and were compared to *P. stuartii* BE2467, the *fliC* mutant, and a motile revertant of BE2467. Images are representative of three independent experiments. Motility scores are indicated as follows: −, no motility; +/−, intermediate motility; +, low motility; ++, moderate motility; +++, high motility. (D) CBA/J mice (*n* = 20) were transurethrally inoculated with 1 × 10^8 CFU of a 1:1 mixture of wild-type *P. stuartii* and the *fliC* mutant, and a 4-mm segment of catheter tubing was retained in the bladder for the duration of the study. After 96 h, mice were sacrificed and the catheterized bladder (B), kidneys (K), and spleen (S) were homogenized and plated onto LB agar with and without hygromycin to determine bacterial burden of wild-type *P. stuartii* and the mutant. A competitive index was then calculated on a per-mouse basis using the ratio of mutant to wild-type CFUs in each organ divided by the ratio of mutant to wild-type CFUs from the inoculum (see Materials and Methods). Statistical significance was assessed by Wilcoxon signed-rank test (*, *P* < 0.05; **, *P* < 0.01).
BE2467, exhibited intermediate motility; and 7 (41%) exhibited classic motility. Importantly, loss of fliC abrogated the intermediate motility of BE2467, while motile isolates recovered from the plates in Fig. 8A exhibited classic motility, suggesting that BE2467 is capable of modulating motility.

The fliA gene sequence of wild-type P. stuartii BE2467 has a two-nucleotide insertion (TG) 468 nucleotides into the coding sequence of the gene compared to fliA sequence from several other P. stuartii genomes, resulting in a frameshift that may explain the intermediate motility exhibited by this strain (Fig. S4). However, a P. stuartii strain that was isolated from human urine in 2014 exhibits 100% fliA nucleotide identity to strain BE2467, indicating that this two-nucleotide insertion in fliA is not unique to BE2467. It is also notable that both motile revertants of P. stuartii BE2467 had a two-nucleotide deletion within fliA, shifting it back into frame (22 nucleotides downstream of the TG in isolate 1 and 3 nucleotides downstream of the TG in isolate 2, as shown in Fig. S4). Thus, P. stuartii BE2467 can spontaneously revert to classic motility in vitro and this may also occur for other P. stuartii isolates.

Notably, wild-type P. stuartii BE2467 still expresses fliC during incubation in LB medium and urine (Fig. 7B). It has also been demonstrated in other bacterial species that flagella can have an additional role in pathogenesis that is independent of motility. For instance, in Pseudomonas aeruginosa, production of an intact flagellum is important for resistance to surfactant protein A (SP-A), which facilitates uptake of the bacterium by phagocytic cells (30). We therefore investigated the contribution of fliC to pathogenicity. Female CBA/J mice were transurethrally inoculated with 1 × 10^5 CFU of a 1:1 mixture of the fliC mutant and wild-type P. stuartii, the catheterized bladder and kidneys were homogenized for determination of bacterial burden after 96 h, and a competitive index (CI) value was calculated (Fig. 8C). As CAUTI can progress to secondary bacteremia, the spleen was also homogenized for determination of bacterial burden as an indicator of dissemination to the bloodstream. The fliC mutant exhibited a significant fitness defect in the catheterized bladder and kidneys during coculture with the wild-type strain, confirming the results of the primary Tn-Seq screen. Thus, while in vitro motility is variable in P. stuartii BE2467, flagella contribute to motility in P. stuartii BE2467 CAUTI. The mutant did not exhibit a fitness defect in the spleen, indicating that flagella do not contribute to development of bacteremia or survival once within the bloodstream. Flagella similarly do not contribute to Proteus mirabilis survival in the bloodstream (29).

### TABLE 1 Assessment of motility in P. stuartii clinical isolates

<table>
<thead>
<tr>
<th>Isolate ID</th>
<th>Motility score</th>
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<tbody>
<tr>
<td>BE2467</td>
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</tr>
<tr>
<td>BE2467 ΔfliC</td>
<td>-</td>
</tr>
<tr>
<td>BE125</td>
<td>-</td>
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<tr>
<td>HI122</td>
<td>-</td>
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<tr>
<td>206v0MAC4</td>
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</tr>
<tr>
<td>KU146</td>
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<tr>
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<tr>
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<td>-</td>
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<td>101v0MAC3</td>
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<tr>
<td>955</td>
<td>+++-</td>
</tr>
<tr>
<td>BE2467 motile</td>
<td>+++</td>
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*aMotility scores (--, no motility; +/-, intermediate motility; +, low motility; ++, moderate motility; ++++, high motility) are based on three independent assessments of each isolate. ID, identifier.*
Contribution of twin arginine translocation to *P. stuartii* fitness during single-species and polymicrobial CAUTI.

The twin arginine translocase (Tat) system is used by numerous Gram-negative bacterial species to export prefolded proteins carrying a twin arginine motif signal sequence. Tat substrates are generally periplasmic proteins and enzymes, many of which are involved in binding redox cofactors (31). The protein export system is comprised of three proteins (TatA, TatB, and TatC), of which *tatA* and *tatC* were identified as core fitness factors. Prior work in *P. stuartii* strain XD37 revealed that this species produces a TatA protein with an N-terminal extension that must be cleaved by rhomboid-type protease AarA to be active and that both *aarA* and *tatC* are required for production of a melanin-like pigment as well as for aerobic growth on MacConkey agar plates (32). Interestingly, *aarA* (BGK56_18970) was not identified as a fitness factor in *P. stuartii* BE2467, likely because this strain encodes two additional rhomboid-type proteases that may also be capable of processing TatA (BGK56_13380 and BKG56_12285). We therefore generated a *tatC* mutant for validation of the importance of twin arginine translocation to *P. stuartii* CAUTI.

The *tatC* mutant of *P. stuartii* BE2467 exhibited expected in vitro phenotypes of reduced pigment production (Fig. 9A) and a growth defect on MacConkey agar (Fig. 9B) as previously observed in *P. stuartii* strain XD37 (32). Upon direct competition with wild-type *P. stuartii* BE2467, the *tatC* mutant exhibited a significant defect in colonization of the catheterized bladder, kidneys, and spleen, indicating that Tat substrates are critical for *P. stuartii* fitness within the catheterized urinary tract (Fig. 9C). To determine if the Tat system also contributes to polymicrobial infection, CBA/J mice were transurethrally inoculated with 1 × 10^5 CFU of a 1:1 mixture of wild-type *P. stuartii*, the *tatC* mutant, and wild-type *P. mirabilis* HI4320. A competitive index was again calculated for the *tatC* mutant relative to wild-type *P. stuartii* BE2467 to determine fitness of the mutant during coinfection. Statistical significance was assessed by Wilcoxon signed-rank test (*, P < 0.05; **, P < 0.01).

Contribution of an ATP-dependent protease to *P. stuartii* fitness during single-species and polymicrobial CAUTI.

Most energy-dependent protein degradation in...
bacteria is mediated by five ATP-dependent proteases: FtsH (HflB), the Clp proteases (ClpAP, ClpXP, and ClpYQ/HslVU), and the Lon protease (33). P. stuartii BE2467 carries \textit{ftsH}, \textit{clpA}, \textit{clpP}, \textit{clpX}, \textit{hslU}, and \textit{hslV}, of which \textit{clpP} and \textit{hslV} were identified as core fitness factors. To validate the importance of ATP-dependent proteases to \textit{P. stuartii} fitness, we generated a mutant in \textit{clpP} to disrupt both ClpAP and ClpXP. The \textit{clpP} mutant was assessed to determine if loss of ClpAP and/or ClpXP would impact the ability to survive heat stress (Fig. 10A). Wild-type \textit{P. stuartii} and the \textit{clpP} mutant grew comparably at 37°C, but the \textit{clpP} mutant exhibited reduced viability during incubation at 45°C for 4 h, which became more pronounced after 24 h (Fig. 10A). Thus, loss of \textit{clpP} results in sensitivity to heat stress.

To validate the contribution of ClpP to \textit{P. stuartii} fitness during CAUTI, the \textit{clpP} mutant was directly competed against wild-type \textit{P. stuartii} during single-species infection (Fig. 10B) or during coinfection with \textit{P. mirabilis} (Fig. 10C). The \textit{clpP} mutant was significantly outcompeted in all organs during both types of infection, underscoring the central importance of ATP-dependent protein degradation to fitness during infection. This finding is notable as all five ATP-dependent protease types were found to represent either essential genes or fitness factors in our prior Tn-Seq study of \textit{P. mirabilis} (27).

**Contribution of \textit{d}-alanine-\textit{d}-alanine ligase to \textit{P. stuartii} fitness during single-species CAUTI.** The peptidoglycan layer of the bacterial cell envelope consists of linear chains of polysaccharides cross-linked by short peptides, each unit of which is first synthesized in the cell cytoplasm and then flipped across the membrane (reviewed in reference 34). During synthesis in the cytoplasm, short peptides are capped by two \textit{d}-alanine residues via the action of \textit{d}-alanine-\textit{d}-alanine ligase (\textit{ddl}), and these terminal \textit{d}-alanine residues are critical for peptidoglycan cross-linking by transpeptidases. The \textit{Ddl} enzymes therefore serve an essential role in peptidoglycan biosynthesis, some of which provide antibiotic resistance due to altered substrate specificity (such as vancomycin-resistant \textit{Ddl} enzymes VanA, VanB, and VanC) (35).

In our prior Tn-Seq investigation of \textit{P. mirabilis} HI4320, we determined that \textit{P. mirabilis} carries only one \textit{ddl} gene (\textit{ddIA}), which was identified as essential for growth in LB medium (27, 36). In contrast, \textit{P. stuartii} BE2467 carries two \textit{ddl} genes, one that was...
**FIG 11** d-Alanine–d-alanine ligase contributes to *P. stuartii* fitness during single-species CAUTI but not during polymicrobial infection. *P. stuartii* BE2467 and the isogenic *ddlA* mutant were assessed for sensitivity to d-cycloserine (A), fitness during single-species infection (B), and fitness during coinfection with *P. mirabilis* HI4320 (C). (A) Optical density of bacterial cultures following incubation with increasing concentrations of d-cycloserine. Box-and-whisker plots show all data points with minimum, maximum, and mean values for results from 4 independent experiments. (B) CBA/J mice (n = 9) were transurethrally inoculated with 1 × 10^6 CFU of a 1:1 mixture of wild-type *P. stuartii* and the *ddlA* mutant, and a competitive index was calculated as described above. (C) CBA/J mice (n = 6) were transurethrally inoculated with 1 × 10^6 CFU of a 1:1:2 mixture of wild-type *P. stuartii*, the *ddlA* mutant, and wild-type *P. mirabilis* HI4320, a 4-mm segment of catheter tubing was retained in the bladder for the duration of the study, and a competitive index was calculated as described above. Statistical significance was assessed by Wilcoxon signed-rank test (*, P < 0.05; **, P < 0.01).
BGK56_12525) with 48% amino acid sequence identity to that of Salmonella dublin FMB1, the species strain from which this protein was originally identified (42), and sopB was identified as a candidate fitness factor for kidney colonization during single-species infection. We therefore sought to determine the contribution of T3S to intracellular survival of P. stuartii in macrophages, in addition to pathogenesis in vivo (Fig. 12).

We first assessed the contribution of T3S to P. stuartii survival in the RAW 264.7 murine macrophage cell line. The wild-type strain and the yscI mutant were internalized by the RAW cells to similar degrees, and we observed no differences in intracellular survival of the mutant compared to the wild-type strain after 4h (Fig. 12A). Thus, the T3S system of P. stuartii did not appear to contribute to uptake by phagocytic cells or survival in vitro under our experimental conditions.

To assess the contribution of T3S to P. stuartii fitness within the catheterized urinary tract, the yscI mutant was directly competed against wild-type P. stuartii during single-species infection (Fig. 12B) or during coinfection with P. mirabilis (Fig. 12C). Interestingly, loss of yscI resulted in a significant fitness defect only during single-species infection and not during coinfection with P. mirabilis. Thus, T3S may be dispensable during coinfection, possibly due to the increased tissue damage that occurs with P. mirabilis (23). Considering that the yscI mutant exhibited a significant defect in the kidney during single-species infection, we next assessed the potential contribution of the putative T3S effector sopB to fitness during single-species infection (Fig. 12D). However, the sopB mutant did not exhibit a significant fitness defect in any organ. The lack of phenotype for sopB during single-species infection may have been due to compensation by other T3S effectors.

Contribution of type VI secretion to polymicrobial CAUTI. Type VI secretion (T6S) systems are common in Gram-negative bacteria and can mediate microbe-microbe interactions as well as host-microbe interactions through injection of effector proteins. A minimum of 13 genes are required for the assembly and function of the T6S apparatus (now universally named TssA-TssM), but T6S gene clusters can contain more than 30 genes due to the presence of additional accessory factors and effectors (43, 44). The T6S gene cluster of P. stuartii BE2467 contains 18 genes (BGK56_15180 to

![Type 3 secretion contributes to P. stuartii fitness during single-species CAUTI but not during polymicrobial infection. P. stuartii BE2467 and the isogenic yscI mutant were assessed for intracellular survival in RAW macrophage-like cells (A), fitness during single-species infection (B), and fitness during coinfection with P. mirabilis H4320 (C). (A) Log_{10} CFU/ml of viable bacteria recovered from the input inoculum, intracellular bacteria 15 min after removal of extracellular bacteria via amikacin treatment, and intracellular bacteria 4h after removal of extracellular bacteria. Gray bars indicate the means of results from 2 independent experiments performed with 4 replicates each. No significant differences were detected by a nonparametric Mann-Whitney test. (B) CBA/J mice (n = 20) were transurethrally inoculated with 1 × 10^5 CFU of a 1:1 mixture of wild-type P. stuartii and the yscI mutant, a 4-mm segment of catheter tubing was retained in the bladder for the duration of the study, and a competitive index was calculated as described above. (C) CBA/J mice (n = 8) were transurethrally inoculated with 1 × 10^5 CFU of a 1:1:2 mixture of wild-type P. stuartii, the yscI mutant, and wild-type P. mirabilis H4320, and a competitive index was calculated as described above. (D) CBA/J mice (n = 13) were transurethrally inoculated with 1 × 10^5 CFU of a 1:1 mixture of wild-type P. stuartii and the sopB mutant, and a competitive index was calculated as described above. Statistical significance was assessed by Wilcoxon signed-rank test (*, P < 0.05; **, P < 0.01).
While nosing genes in the cluster met our stringent fold change cutoff, 2 genes were estimated to be essential due to a lack of transposon insertions in the input pool and the remaining 16 genes all had statistically significant (P < 0.001) fitness defects ranging from 2.6-fold to 19.8-fold in the bladder and kidneys during both single-species and polymicrobial infection. The potential importance of T6S to \emph{P. stuartii} pathogenesis is notable, as our prior CAUTI Tn-Seq study revealed that all five T6S effector operons of \emph{P. mirabilis} HI4320 contained candidate fitness factors for coinfection with \emph{P. stuartii} \cite{27}. We therefore sought to determine the importance of T6S to fitness of both \emph{P. stuartii} and \emph{P. mirabilis} during single-species and polymicrobial CAUTI.

\textbf{ImpJ (TssK) is part of the baseplate of the T6S apparatus and essential for sheath polymerization, and disruption of \textit{impJ} prevents T6S function} \cite{43, 45, 46}. We generated an \textit{impJ} mutant in \emph{P. stuartii} BE2467 by disrupting BGKS5\_12575 and tested fitness during cochallenge with the parental isolate for single-species CAUTI and coinfection with \emph{P. mirabilis} HI4320 (Fig. 13A and B). The \emph{P. stuartii} \textit{impJ} mutant did not exhibit an obvious fitness defect during monomicrobial infection, but it was significantly outcompeted by wild-type \emph{P. stuartii} in the catheterized bladder during polymicrobial infection with wild-type \emph{P. mirabilis}, suggesting that T6S contributes to \emph{P. stuartii} fitness during coinfection.

To conduct the reciprocal experiment with \emph{P. mirabilis}, we utilized a previously constructed mutant in \emph{P. mirabilis} \textit{impJ} (PMI0742) that was confirmed to lack T6S functionality \cite{47}. Interestingly, the \emph{P. mirabilis} \textit{impJ} mutant was outcompeted by wild-type \emph{P. mirabilis} in the kidneys during single-species infection and was outcompeted in bladder and kidneys during coinfection with \emph{P. stuartii} (Fig. 13C and D). Thus, T6S appears to contribute to the fitness of both bacterial species during coinfection and may contribute to \emph{P. mirabilis} fitness during single-species CAUTI.

\textbf{Assessing \emph{P. stuartii} candidate fitness factors by independent challenge.} The experimental design of Tn-Seq is aimed at uncovering genes that contribute to the relative fitness of a bacterium competed against thousands of other mutants. However, fitness factors that are ideal targets for potential therapeutic intervention would also be expected to exhibit a defect during an independent challenge. We therefore utilized an independent challenge setup to assess colonization of each mutant that exhibited a significant fitness defect during single-species infection (\textit{fliC}, \textit{yscl}, \textit{clpP}, \textit{ddlA}, and \textit{tatC}) for comparison to wild-type BE2467 (Fig. 14).
While none of the mutants were completely cleared from the urinary tract, four of five showed significant colonization defects in at least one organ: the\(\text{yscI}\) mutant exhibited reduced kidney and spleen colonization, the\(\text{clpP}\) mutant exhibited reduced bladder colonization, the\(\text{ddlA}\) mutant exhibited reduced bladder and spleen colonization, and the\(\text{tatC}\) mutant exhibited a defect in all organs. Infection with the\(\text{tatC}\) mutant also resulted in colonization levels at or below the limit of detection in a significant proportion of mice, while infection with the other three mutants with colonization defects resulted only in a significant decrease in the proportion of mice with bacteremia. Based on these observations, disruption of\(\text{fliC}\) appears to have impacted relative fitness only during cochallenge against the parental isolate, while loss of the other fitness factors also resulted in some degree of colonization defect independently of competition against the wild-type isolate. Factors such as\(\text{ddlA}\) and\(\text{tatC}\) may therefore represent preferable targets for potential therapeutic intervention.

**DISCUSSION**

\(P.\ stuartii\) is a clinically relevant opportunistic pathogen with a high prevalence of antibiotic resistance, and yet little is known regarding the pathogenesis of this organism during infection. Here, we utilized Tn-Seq to identify \(P.\ stuartii\) genes that are essential for growth under standard laboratory conditions and to identify fitness factors for single-species versus polymicrobial CAUTI. Major strengths of this study included the following: (i) use of a genetically tractable CAUTI isolate with an available genome sequence (23), (ii) use of multiple transposon library pools and multiple mice per infection pool, (iii) stringent cutoffs for analysis of insertion site reads, and (iv) assessment of the impact of coinfection on fitness requirements. Limitations of the study included inability to assess the fitness contribution of genes that were poorly represented in the input pools, inability to accurately assess the fitness contribution of genes encoding secreted factors under conditions in which loss can be complemented by other mutants within the input pools, and potential polar effects of transposon insertions within operons, all of which are common limitations of Tn-Seq studies.

Despite these limitations, our approach successfully uncovered numerous \(P.\ stuartii\) fitness determinants for single-species and polymicrobial CAUTI, along with subsets of genes that contribute to fitness only during infection with each distinct type. By generating mutants in genes of interest and assessing fitness during direct cochallenge
with wild-type \( P. stuartii \), 6/7 (86%) fitness factors tentatively identified by Tn-Seq were validated as contributing to single-species CAUTI and 3/5 (60%) during coinfection with \( P. mirabilis \). Furthermore, 4/5 fitness factors tested also exhibited colonization defects compared to wild-type \( P. stuartii \) in an independent challenge model, indicating that they may represent valid targets for therapeutic intervention.

There are several potential explanations for the failure of certain mutants to achieve validation during direct cochallenge, including the overall level of coverage of TA insertion sites within these genes, the specific location of the transposon insertions in a gene versus replacement of the majority of the gene with a hygromycin resistance cassette for generation of the mutants, the fold change and \( P \) value cutoffs used in the analysis, and potential polar effects of transposon insertions. It is also possible that some of the mutants might exhibit significant defects only when dramatically under-represented in the input inoculum, such as in the case of the \(~1:10,000\) ratio that would have been present during the Tn-Seq screen but not during a 1:1 cochallenge. Regardless of the source of discrepancy, the validation rate resulting from combined assessments of mutants during single-species and polymicrobial infection was 9/12 (75%), which is consistent with the validation rates reported in other Tn-Seq studies (ranging from 75% to 86%) (27, 29, 48–51).

An additional consideration regarding the validation experiments is that the mutagenesis strategy utilized to construct targeted mutants of interest has the potential to result in polar effects due to replacement of the gene with a hygromycin cassette. Of the seven genes chosen for validation studies, only three, \( tatC \), \( yscI \), and \( impJ \), are present in operons; among the three, \( tatC \) is the final gene in the operon and unlikely to exhibit polar effects whereas \( yscI \) and \( impJ \) are in the middle of their respective operons. It is therefore possible that disruption of \( yscI \) and \( impJ \) would also result in loss of downstream gene products involved in their respective secretion systems, which might contribute to the fitness defects observed for the mutants. Future investigations of the role of each secretion system during \( P. stuartii \) CAUTI will therefore need to include an assessment of the relative contributions of these genes in production of a functional secretion apparatus.

In conclusion, \( P. stuartii \) carries a core set of genes and pathways that contribute to colonization during both single-species and polymicrobial infection, and those genes and pathways are both prevalent and highly conserved among isolates. Further research is needed to determine the \textit{in vivo} importance of these core genes and pathways during both single-species and polymicrobial CAUTI, including coinfections with other common cocolonizing species. The underlying mechanisms of these core fitness requirements likely represent the most promising candidates for development of potential therapeutic interventions aimed at reducing \( P. stuartii \) colonization or minimizing risk of progression to severe infection and urosepsis. Furthermore, continued genome-wide assessment of multiple uropathogens is expected to uncover fitness factors shared between bacterial species, thereby elucidating common pathways for colonization and persistence.

**MATERIALS AND METHODS**

**Ethics statement.** All animal protocols were approved by the Institutional Animal Care and Use Committee (IACUC) at the University of Michigan Medical School (PRO00005052) and the University of Buffalo (MIC31107Y), in accordance with the guidelines specified by the Office of Laboratory Animal Welfare (OLAW) and the United States Department of Agriculture (USDA), as well as guidelines specified by the Association for Assessment and Accreditation of Laboratory Animal Care, International (AAALAC, Intl.). Mice were anesthetized with a weight-appropriate dose (0.1 ml for a mouse weighing 20 g) of ketamine/xylazine (80 to 120 mg ketamine/kg of body weight and 5 to 10 mg xylazine/kg) by IP injection. Mice were euthanized by inhalant anesthetic overdose (Michigan) or \( CO_2 \) asphyxiation (Buffalo), followed by vital organ removal.

\( Providencia stuartii \) isolates were collected from the urine of individuals with indwelling urinary catheters in a study approved by the University at Buffalo Institutional Review Board (STUDY00002526). All participants (or approved decision makers) provided written informed consent prior to initiation of investigation, and all participants also assented to being in the study.

**Bacterial strains and culture conditions.** \( Providencia stuartii \) strains BE2467, BE125, HI122, KU146, DE266, RO106, GO42, MC118, BR143, TA101, NI114, and 955 and \( Proteus mirabilis \) strain HI4320 are part of the study.
of a strain collection of urine isolates that were previously obtained from the urine of catheterized nursing home residents (10, 23, 52). P. stuartii strains BE101v0MAC1, BE101v0MAC2, 201v0MAC1, 202v0MAC2, and 206v0MAC4 were isolated from the urine of catheterized nursing home residents under STUDY00002526, and the isolates were verified to represent Providencia stuartii by the use of API-20E test strips (BioMérieux, Marcy-l’Étoile, France).

Bacteria were routinely cultured at 37°C with aeration in 5 ml low-salt LB broth (10 g/liter tryptone, 5 g/liter yeast extract, 0.5 g/liter NaCl) or on low-salt LB medium solidified with 1.5% agar. Transposon mutants were cultured in LB medium containing 150 μg/ml hygromycin (Research Products International). Additional P. stuartii mutants were constructed for validation of conditionally essential genes by the use of a previously described allelic exchange vector to replace the gene of interest with a hygromycin resistance cassette (27). Resulting mutants were screened by hygromycin selection and PCR (all primers used in this study for generation and verification of mutants are listed in Table S2 in the supplemental material).

Construction of pAOJ12. To generate a transposon vector suitable for mutagenesis in P. stuartii, the hygromycin resistance cassette from pSIM18 (53) was subjected to PCR amplification with primers AOJ_123 and AOJ_124 and cloned into pCR2.1-TOPO (Thermo Fisher Scientific) to create pAOJ8. Plasmid pAOJ8 was digested with MfeI and SpeI restriction enzymes and cloned between the XbaI and MfeI sites on plasmid pSAM_Ec (54), resulting in the hygromycin resistance cassette being flanked by Mariner transposon inverted repeats. The resulting plasmid was named pSAM_Hygromycin.

A conditionally replicating plasmid was made by digesting pRK415 (55) with MfeI and religating, removing an ~2.8-kb fragment containing the trAI/A2 replication protein open reading frames. This plasmid, pRK415ΔMfeI, and all derivatives were transformed into E. coli EP1300, which expresses the Traf protein under the control of the arabinose promoter (56), and were cultured in media containing 0.5% arabinose. The temperature-sensitive lacB57 repressor from pSIM18, using primers AOJ_173/AOJ_174; (ii) the Himar 1C9 transposase from pSAM_Ec, using AOJ_175/AOJ_176; and (iii) the strong rho-independent spy transcriptional terminator from E. coli Top 10 (57), using AOJ_177/AOJ_178. These fragments were stitched together into a single unit using NEBuilder HiDNA assembly master mix. The entire assembled fragment was amplified using AOJ_173 and AOJ_178, which also contain Nhel and MfeI restriction sites. The AOJ_173/AOJ_178 amplified fragment was digested with these two enzymes and ligated into the EcoRI-XbaI site of pRK415ΔMfeI to generate pAOJ11. To insert the Ap* promoter into pAOJ11, an ~300-bp fragment was created by PCR amplification of pSIM18 with primers AOJ_195/AOJ_196, which add NdeI and KpnI sites, and were cultured in media containing 0.5% arabinose. The resulting plasmid was named pSAM_Hygromycin.

One verified by restriction digestion, pAOJ12 was transferred into E. coli S17-1::pir for biparental mating. Plasmid pAOJ12 contains an RK2-based suicide origin that can replicate only with Traf supplied in trans, an RK2 origin of transfer for mating, a hygromycin resistance transposon, and a hyperactive Himar1C9 transposase to drive transposition when expression is induced from the Ap* promoter by temperature elevation. As the Ap* promoter is active in many distally related Gram-negative bacteria, the RK2 conjugation system can transport DNA into a large variety of species, and hygromycin B has the useful features of being broadly toxic but not used clinically. Thus, we expect that this transposon vector could be useful for transposon mutagenesis in many Gram-negative species beyond P. stuartii.

Generation and validation of transposon mutants. P. stuartii BE2467 and E. coli S17-1::pir(pAOJ12) were cultured overnight in LB medium containing appropriate antibiotics at 30°C with aeration at 225 rpm. P. stuartii BE2467 (250 μl) and E. coli S17-1::pir(pAOJ12) (500 μl) were inoculated into separate conical tubes, each containing 25 ml fresh LB medium prewarmed to 30°C with antibiotics appropriate for S17-1::pir(pAOJ12). Cultures were incubated with shaking for 2 h at 30°C before centrifugation to pellet was performed, and the cell pellets were washed once with phosphate-buffered saline (PBS; 0.128 M NaCl, 0.0027 M KCl, pH 7.4). The cell pellets were resuspended in 0.5 ml PBS, combined into a single microcentrifuge tube, and centrifuged to pellet. The supernatant was removed, and the pellet consisting of the combined cells was incubated at ambient temperature for 5 min, after which the pellet was gently resuspended in 250 μl LB medium, spread onto a sterile 0.45-μm-pore-size filter resting on a LB agar plate, and incubated at 30°C for 90 min. Next, the mating mixture was removed from the filter and placed into 2.5 ml LB medium with 10 μg/ml chloramphenicol (stopping the growth of E. coli) and heated to 42°C for a 10-min heat shock. The culture was then transferred to 37°C with shaking to recover for 90 min, followed by plating on LB medium containing 150 μg/ml hygromycin B and 10 μg/ml chloramphenicol. Inoculated plates were incubated overnight at 37°C, and colonies were enumerated the following day. Roughly 50,000 colonies were scraped from the resulting plates into 5 pools of 10,000 colonies each for generation of glycerol freezer stocks, which were frozen at −80°C until further use.

The randomness of the insertions was verified by extracting genomic DNA from subcultures of individual colonies, digesting with EcoRV, and Southern blotting with a digoxigenin (Roche)-labeled probe targeting the hygromycin resistance cassette within the transposon, which was amplified by primers AOJ_199 and AOJ_200 (see Fig. S1 in the supplemental material).

Mouse model of CAUTI. Infection studies were carried out as previously described (21, 27) using a modification of the Hagberg protocol (58). Transposon mutant pools (1 ml volume) were thawed in 9 ml fresh LB medium with hygromycin and cultured at 37°C for no more than 10 h. Cultures were adjusted to an optical density at 600 nm (OD600) of ~0.2 (2 × 10⁸ CFU/ml) and were diluted 1:100. Female CBA/J mice (Envigo or Jackson Laboratory) (6 to 8 weeks of age) were inoculated transurethrally with 50 μl of 2 × 10⁸ CFU/ml (1 × 10⁸ CFU/mouse), and a 4-mm-long segment of sterile silicone tubing (Braintree...
Scientific, Inc.) (0.64-mm outer diameter [O.D.], 0.30-mm inner diameter [I.D.]) was carefully advanced into the bladder during inoculation and retained for the duration of the study as described previously (23, 29–61).

For coinfection experiments, mice were inoculated with 50 μl of a 1:1 mixture of the P. stuartii transposon mutant pools and WT P. mirabilis H4320. For each transposon pool input, 5 to 10 mice were inoculated for single-species infection and 5 to 10 mice were coinfected with P. mirabilis. Mice were euthanized at 4 days postinoculation (dpi), and bladders, kidneys, and spleens were harvested into PBS. Notably, catheter segments were not removed from the bladder samples for homogenization, so CFU data for bladder samples represent a catheterized bladder. Tissues were homogenized using either an Omni TH homogenizer (Omni International) or a Bullet Blender 5 Gold homogenizer (Next Advance), and a 150-μl aliquot was removed and plated using either an Autoplate 4000 spiral plater (Spiral Biotech) or an EddyJet 2 spiral plater (Neutec Group) for enumeration of colonies using either a QCount automated plate counter (Spiral Biotech) or a ProtoCOL3 automated colony counter (Synbiosis). The remaining bladder and kidney homogenates were spread plated in their entirety, and colonies were collected, pelleted, and frozen for sequencing. A competitive index (CI) value was calculated for all samples in which bacterial burden was described as being above the limit of detection as follows:

\[
CI = \frac{\text{Strain A output}}{\text{Strain A input}} \\
\frac{\text{Strain B output}}{\text{Strain B input}}
\]

A log10CI value of 0 indicates that the ratio of the strains in the output was similar to that in the input and that neither strain had an advantage. A log10CI value of >0 indicates that strain A had a competitive advantage over strain B. A log10CI value of <0 indicates that strain B had a competitive advantage over strain A.

For independent challenge experiments, groups of 5 to 6 mice were inoculated with 50 μl of 2 × 108 CFU/ml (1 × 108 CFU/mouse) of a strain of interest, and a catheter segment was inserted as described above. Mice were euthanized 4 days postinoculation (dpi) as described above, and organ homogenates were plated for enumeration of colonies.

**Illumina sequencing.** Genomic DNA was isolated from the five inocula replicates, and from the bladder and kidney homogenates of each individual mouse (outputs) by cetyltrimethylammonium bromide (CTAB) precipitation (62). Samples were enriched for transposon insertion junctions as outlined previously by Goodman et al. (63), and concentration and purity were confirmed by TapeStation analysis. Samples were multiplexed and subjected to V4 Single-End 50 HiSeq-2500 high-output sequencing as follows: (i) 5 input samples with 2 replicates each were multiplexed and sequenced on two lanes; (ii) P. stuartii single-species outputs from 20 mouse bladders were multiplexed and sequenced on two lanes; (iii) P. stuartii single-species outputs from 20 mouse kidneys were multiplexed and sequenced on two lanes; (iv) outputs from P. stuartii coinfection with P. mirabilis from 20 mouse bladders were multiplexed and sequenced on two lanes, and (v) outputs from P. stuartii coinfection with P. mirabilis from 20 mouse kidneys were multiplexed and sequenced on two lanes. Each lane was spiked with 15% bacteriophage ϕX174 DNA to overcome effects of the presence of low-diversity sequences. Sequencing was performed at the University of Michigan DNA core facility. All raw sequence reads are available in the Sequence Read Archive under BioProject PRJNA578390 as accession numbers SAMN13059443 to SAMN13059447 and SRA accession numbers SRR10312361 to SRR10312365. The barcodes associated with each unique sample are provided in Table S3.

**Mapping of transposon insertion sites.** The chromosome and plasmid sequences of P. stuartii BE2467 (NCBI accession numbers CP017054, CP017055, and CP017056) (23) were combined into a single genome sequence with 1,000 N’s added between each pair of contigs, and the coordinates of genes were adjusted accordingly. The Goodman In-Seq pipeline (63) was applied on the raw reads to perform read filtration, transposon nucleotide removal, debarcoding, alignment, and insertion calling, following a script that was previously used to map insertions onto the P. mirabilis genome (27, 29).

**Identification of essential genes.** Essential genes were identified as described previously (27). Briefly, input samples were corrected for potential read count bias, and a Bayesian mixture model was used to estimate the rate for insertion counts in each gene by assuming that the counts followed a Poisson distribution. JAGS (64) was used to obtain a posterior probability estimate for evaluating the likelihood of each gene being essential. Genes identified as having a 90% probability or greater of being essential were classified as “essential genes.” Protein sequences of estimated essential genes were annotated for orthologous groups (GO terms, KO terms, and COG categories) using the eggNOG-mapper function of EggNOG version 4.5.1 via HMmer search restricted to the “bacteria” HMM database (65).

**Identification of candidate fitness factors for single-species and polymicrobial CAUTI.** Fitness factors were identified using the approach previously published for P. mirabilis (27, 29). Briefly, the output samples from 20 mice (including 4 mice per input pool) were combined for estimation of the fitness contribution to a given organ for a given infection type (bladder versus kidneys and single-species versus coinfection). Individual genes were assessed for fitness contribution only when the mean of the sum of insertion site reads was >1,000 and the number of insertions in that gene was >5. The fitness contribution of each gene was then estimated as previously described (27, 29) using an R package called Tn-SeqDiff (66), which can be installed from the Comprehensive R Archive Network (CRAN). Significant genes for further analysis were selected based on an adjusted P value of <0.05 and a >20-fold ratio of input over output.

**Proteome analysis.** To identify homologs of candidate core fitness factors first, a file was generated containing the FASTA sequences of each fitness factor and then the proteomes of other P. stuartii strains available from the Pathosystems Resource Integration Center (PATRIC) were compared to that sequence...
using the PATRIC proteome comparison tool (67). The sequence identity was limited to ≥10% over a minimum of 30% sequence for this comparison.

**Growth curves.** Overnight cultures of *P. stuartii* mutants washed 1 × in PBS and diluted 1:100 in growth medium, and 100-μl aliquots were dispensed into at least 4 replicate wells of a 96-well plate. A BioTek Synergy H1 96-well plate reader was utilized to generate growth curves during incubation at 37°C with continuous double-orbital shaking for aeration. OD₅₆₀ readings were taken every 15 min for 18 h.

**RNA isolation.** *P. stuartii* BE2467 was cultured overnight at 37°C with shaking at 225 rpm in LB broth with 25 μg/ml kanamycin. A 500-μl volume of the culture was centrifuged to pellet, resuspended in 0.5 ml of sterile saline solution (0.9% NaCl [wt/vol]), and washed twice. The final suspension was diluted 1:100 into 5 ml of prewarmed filter-sterilized human urine or fresh LB medium and incubated for 2 h at 37°C with shaking at 225 rpm. Four 1-ml aliquots were then removed, centrifuged to pellet, and resuspended in 1 ml of RNA STAT-60 (Tel-Test, Inc.). Chloroform-isooamyl alcohol (Calbiochem) (24:1; 0.2 ml) was added to each aliquot, and microcentrifuge tubes were inverted several times for mixing and then centrifuged at 18,000 × g for 30 min at 4°C. The aqueous layer was removed and transferred to a new microcentrifuge tube, to which 0.5 ml isopropanol was added (Fisher Scientific), and the reaction mixture was then mixed by inversion and incubated overnight at −20°C to precipitate RNA. The next day, samples were centrifuged at 18,000 × g for 30 min at 4°C, supernatants were discarded, and the nucleic acid pellet was washed three times with 1 ml 75% ethanol and centrifuged at 18,000 × g for 10 min after each wash. After the final wash, the supernatant was carefully removed and the pellet was air-dried and then dissolved in 50 μl of molecular-biology-grade water (Corning). The nucleic acid concentration was measured by the use of a NanoDrop instrument and adjusted with water to 200 ng/μl in a 50-μl total volume. DNA was removed with two separate rounds of digestion via the use of an Invitrogen DNA-free kit, according to the manufacturer’s protocol. DNase-treated RNA was frozen at −20°C until use.

**CDNA synthesis and RT-qPCR.** A 1-μg volume of DNase-treated RNA in a total volume of 20 μl was used to generate CDNA via the use of an iScript CDNA synthesis kit (Bio-Rad) per manufacturer’s instructions. After CDNA synthesis, sample volumes were increased to 40 μl by adding 20 μl TE buffer (10 mM Tris [pH 8], 1 mM EDTA), and CDNA was frozen at −20°C until use. Three technical replicate reaction mixtures for each condition were prepared by combining 1 μl CDNA (approximately 25 ng of RNA template), 2 μl primers (10 pmol/μl), 7 μl water, and 10 μl qPCRBio SyGreen Blue Mix Lo-Rox (PCR Biosystems). Reactions were initiated on a Bio-Rad CFX-Connect real-time system with the default CFX reverse transcriptase quantitative PCR (RT-qPCR) protocol. For analysis, recA was chosen as the reference gene for normalization between samples as it exhibited a low level of variation between the urine and LB samples. Data were analyzed according to the relative quantification (RQ) method as described previously by Pfaffl (68), in which LB medium was considered the “Control” growth condition and urine was considered the “Sample” growth condition, in the following equation:

\[
\text{Relative Quantification ratio} = \left( \frac{E_{\text{target}}}{E_{\text{reference}}} \right)_{\text{Sample}} \left( \frac{E_{\text{target}}}{E_{\text{reference}}} \right)_{\text{Control}}^{-1}
\]

In this equation, *E* refers to the primer efficiency of the target gene (*E*<sub>target</sub>) or the reference gene (*E*<sub>reference</sub>), and CP refers to the cycle point, or the cycle number at which the signal exceeds the threshold.

**Motility assay.** Swimming motility agar (MOT) plates (10 g/liter tryptone, 5 g/liter yeast extract, 5 g/liter NaCl, 3 g/liter agar) were subjected to stab inoculation with an overnight culture of *P. stuartii* BE2467 or the isogenic *fliC* mutant. MOT plates were incubated without inversion at 37°C for 7 days for assessment of flagellin-mediated motility. In addition, overnight cultures of *P. stuartii* strains of interest were subjected to stab inoculation into motility test medium with triphenyltetrazolium chloride (TTC; Hardy Diagnostics) and incubated 37°C for 24 h. In this medium, motile bacteria extend outward from the stab line and reduce the TTC, generating a diffuse red pigment, while nonmotile organisms grow only along the stab line, generating a concentrated line of red pigment and leaving the surrounding medium clear.

**fliA sequence investigation.** Sequences of the *fliA* gene of *P. stuartii* BE2467 and six other *P. stuartii* isolates were obtained from the PATRIC database and aligned using the free sequence alignment tool from Benchling.com. Genomic DNA was extracted from *P. stuartii* BE2467 and two motile revertants, and *fliA* was subjected to PCR amplification using primers AOJ_325 and AOJ_326 (Table S2) and to Sanger sequencing for alignment with the other *fliA* sequences in Benchling. The resulting alignment is displayed in Fig. S4.

**Assessment of pigment production.** Overnight cultures of wild-type *P. stuartii* BE2457 and the isogenic *tatC* mutant were subjected to drip plating onto non-low-salt LB agar (10 g/liter tryptone, 5 g/liter yeast extract, 10 g/liter NaCl, 1.5% agar) supplemented with 0.4 g/liter tyrosine and 0.25 g/liter iron (II) sulfate heptahydrate. Pigment production was visually assessed after incubation for ~18 h at 37°C.

**Assessment of growth on MacConkey agar.** Overnight cultures of *P. stuartii* BE2467 and the isogenic *tatC* mutant were adjusted to an OD of 2.0 (2 × 10⁸ CFU/ml) and cultured in LB broth. Viable bacteria were enumerated after incubation for ~18 h at 37°C or 30°C.

**Assessment of heat sensitivity.** *P. stuartii* BE2467 and the isogenic *clpP* mutant were cultured overnight in LB broth at 37°C with aeration. Cultures were then adjusted to an OD of 2.0 (2 × 10⁸ CFU/ml) and subjected to serial dilution, and 10 μl of each dilution was spotted onto LB agar. Plates were incubated at either 37°C or 45°C for 20 h prior to growth assessment.

**MIC of D-cycloserine.** *P. stuartii* BE2467 and the isogenic *ddk* mutant were cultured overnight in LB broth at 37°C with aeration. Cultures were adjusted to an OD of 0.2 (2 × 10⁸ CFU/ml) and incubated with increasing concentrations of D-cycloserine for 18 h at 37°C with aeration, followed by measurement of MIC of D-cycloserine.
optical density at 600 nm. For cultures with an OD_{600} reading of >1.0, the culture was diluted 1:10 and remeasured, and the reading was then multiplied by the dilution factor.

**Macrophase-like cell survival assay.** Survival rates of wild-type *P. stuartii* BE2467 and the yscI mutant were measured using the RAW 264.7 murine macrophase-like cell line for an antibiotic protection assay, as described previously (S1). RAW cells were cultured in antibiotic-free RPMI 1640–10% heat-inactivated fetal bovine serum. Amikacin was used at a concentration of 100 µg/ml in place of gentamicin due to the high level of gentamicin resistance observed for *P. stuartii* BE2467. Following antibiotic treatment, the RAW cells were lysed with 1% saponin–PBS (pH 7.4) to quantify viable intracellular bacteria.

**Statistical analysis.** Significance was assessed using a combination of two-way analysis of variance (ANOVA), Student’s *t* test, the nonparametric Mann-Whitney test, and the Wilcoxon signed-rank test. These analyses were performed using GraphPad Prism, version 7 (GraphPad Software, San Diego, CA). All *P* values are two-tailed at a 95% confidence interval.

**Data availability.** Tn-Seq raw reads are freely available in the Sequence Read Archive under BioProject PRJNA578390 as accession numbers SAMN13059443 to SAMN13059447 and SRA accession numbers SRR10312361 to SRR10312365, and the complete data set is listed in Table S1.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**FIG S1**, TIF file, 1.5 MB.
**FIG S2**, TIF file, 1.9 MB.
**FIG S3**, TIF file, 0.4 MB.
**FIG S4**, TIF file, 0.3 MB.
**TABLE S1**, XLSX file, 1.1 MB.
**TABLE S2**, XLSX file, 0.01 MB.
**TABLE S3**, XLSX file, 0.01 MB.

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**REFERENCES**

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