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Comprehensive analysis of type 1 fimbriae regulation in fimB -null strains from the multidrug resistant Escherichia coli ST131 clone

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- 1 Comprehensive analysis of type 1 fimbriae regulation in *fimB*-null strains
- 2 from the multidrug resistant *Escherichia coli* ST131 clone
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Summary

Uropathogenic *Escherichia coli* (UPEC) of sequence type 131 (*E. coli* ST131) are a pandemic multidrug resistant clone associated with urinary tract and bloodstream infections. Type 1 fimbriae, a major UPEC virulence factor, are essential for ST131 bladder colonization. The globally dominant sub-lineage of ST131 strains, clade C/H30-R, possess an ISEc55 insertion in the *fimB* gene that controls phase-variable type 1 fimbriae expression via the invertible *fimS* promoter. We report that the inactivation of *fimB* in these strains causes altered regulation of type 1 fimbriae expression. Using a novel molecular approach, we demonstrate that 'off' to 'on' *fimS* inversion is reduced in these strains and mediated by the *fimE* and *fimX* genes. Unlike typical UPEC strains, the nucleoid-associated H-NS protein does not strongly repress *fimE* transcription in these strains. Using a genetic screen to identify novel regulators of *fimE* and *fimX* in the clade C/H30-R ST131 strain EC958, we defined a new role for the *guaB* gene as a regulator of type 1 fimbriae and a mouse bladder colonization factor. Our results provide a comprehensive analysis of type 1 fimbriae regulation in the globally predominant group of ST131 strains, and highlight important differences in its control compared to non-ST131 UPEC.

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Uropathogenic Escherichia coli (UPEC) are the leading cause of urinary tract infection (UTI), resulting in over 150 million cases worldwide every year (1) and amounting to billions of dollars spent in direct and associated healthcare costs (2). The increased incidence of UTIs caused by multidrug resistant (MDR) strains, including strains that belong to high-risk globally pandemic clones such as E. coli sequence type 131 (E. coli ST131), presents significant new challenges for the management and treatment of UTI. E. coli ST131 is a globally disseminated MDR clone originally identified due to its close association with the spread of the $bla_{CTX-M-15}$ extended spectrum β -lactamase (ESBL) gene (3-5). Genomic analysis of E. coli ST131 has identified a globally dominant fluoroquinolone resistantfimH30 subgroup previously defined as H30-R (6) or clade C (7), as well as two additional less prevalent clades (referred to as A and B) (7). Type 1 fimbriae represent the best-characterized E. coli ST131 virulence factor and are required for ST131 colonization of the mouse bladder (8). Its pharmacological inhibition has been shown to prevent the establishment of acute MDR UTI and treat chronic bladder infection in mice (9).

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The *E. coli* type 1 fimbriae (*fim*) gene cluster contains nine genes, which encode the major (FimA) and minor (FimFGH) structural components, the chaperone-usher transport and assembly apparatus (FimCD) and two regulatory proteins (FimB and FimE) (10, 11). Type 1 fimbrial expression is phase variable due to inversion of the *fim* switch (*fimS*), a 314 bp invertible DNA element which contains a promoter that drives transcription of the *fimACDFGH* genes (12). 'On' or 'off' orientation of *fimS* results in a fimbriated or bald phenotype, respectively. Two tyrosine-like

recombinases, FimB and FimE, catalyze the inversion of *fimS* (13). The FimB recombinase possesses bidirectional switching activity ('off'-to-'on' and 'on'-to-'off'), while FimE primarily catalyzes 'on'-to-'off' *fimS* inversion (13, 14). FimE can also mediate 'off'-to-'on' inversion of *fimS* under some growth conditions, but at lower efficiency (15, 16). The transcription of the *fimB* and *fimE* genes is driven by individual promoters, both of which are repressed by the histone-like nucleoid associated protein H-NS in *E. coli* K-12 strains (17). The inversion of *fimS* is subject to a complex regulatory network that involves several other global regulators, including integration host factor (IHF) (18) and the leucine-responsive regulatory protein (Lrp) (19), as well as other proteins that directly or indirectly impact on *fimS* orientation (reviewed in (20)). The 'off'-to-'on' switching of *fimS* is enhanced by growth in static liquid culture, which results in the enrichment of type 1 fimbriated cells and the formation of a pellicle at the air-liquid interface (21-23).

In addition to *fimB* and *fimE*, three other genes encoding tyrosine-like recombinases have been identified in UPEC (24). The products of two of these genes, FimX (24-26) and IpuA (24) can mediate *fimS* inversion. It has been shown that FimX possesses specificity for *fimS* 'off' to 'on' inversion, while IpuA can mediate bidirectional inversion (24, 26). In the reference UPEC strain UT189, FimX mediates slow 'off' to 'on' *fimS* inversion *in vitro*, but rapid *fimS* 'on' switching during infection of the mouse bladder (25). The *fimX* gene is located adjacent to the *hyxR* gene at a site distal to the *fim* genes on the chromosome (25, 27). An invertible DNA switch analogous to *fimS* lies between the *fimX-hyxR* genes (27). FimX-mediated inversion of this *in cis* cognate switch influences the transcription of *hyxR*, which encodes a LuxR-like response regulator that controls tolerance to reactive nitrogen intermediates (27).

E. coli ST131 strains from the globally dominant clade C/H30-R group (hereafter referred to as clade C) possess a 1,895bp insertion element within the fimB recombinase gene (fimB::ISEc55; Fig. 1) (8, 28). Given the role of fimB in fimS 'on' switching, it is therefore likely that the presence of this insertion affects type 1 fimbriae expression in ST131. Indeed, the fimB::ISEc55 insertion in ST131 has been associated with a slower 'off'-to-'on' switching phenotype in vitro (8).

In this study, we investigated the impact of the *fimB*::ISEc55 insertion on type 1 fimbriae expression by comparing ST131 strains that possessed either an intact or disrupted *fimB* gene. We confirmed that strains containing the *fimB*::ISEc55 insertion display slow 'off' to 'on' type 1 fimbriae switching phenotype and demonstrated its association with increased transcription of the *fimE* gene. Notably, H-NS repression of *fimE* in these strains was also altered compared to non-ST131 strains, highlighting an important difference in type 1 fimbriae regulation between ST131 strains and other well-characterized UPEC reference strains. We also examined the role of the FimE and FimX recombinases in *fimS* inversion in the ST131 clade C reference strain EC958, and described the use of a novel molecular approach based on Illumina deep sequencing to quantitate *fimS* orientation. Finally, we show that in EC958, guanosine 5' triphosphate (GTP) homeostasis and the bacterial stringent response triggered by the (p)ppGpp alarmone contribute

to type 1 fimbriae expression via *fimE* and *fimX* regulation.

Results

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Most E. coli ST131 strains can express type 1 fimbriae regardless of their fimB status We previously sequenced a large collection of E. coli ST131 isolates from different geographical locations (7), including the high-quality reference genome of EC958, a representative clade C strain (8, 29). Despite most strains encoding intact and seemingly functional type 1 fimbriae operons, type 1 fimbriae expression was not a conserved trait among E. coli ST131 strains under standard laboratory growth (7, 8). This variability in type 1 fimbriae expression was significantly associated with the presence of an insertion element within the *fimB* recombinase gene that was identified in most clade C ST131 strains studied by us and others (7, 8, 28). Here, we extended these findings in a larger collection of 91 ST131 strains (71 previously published, detailed in Materials and Methods) for which we provide a comprehensive type 1 fimbriae expression profile. In this collection, 57 strains contained the fimB::ISEc55 insertion (and were from clade C) and 34 contained an intact fimB gene (and were from clades A and B). Using yeast cell agglutination, a standard method for monitoring the production of mannose-sensitive fimbriae (30), we determined type 1 fimbriae expression following growth in different conditions (Table 1). Following overnight shaking growth in LB broth at 37°C, only 14% of the fimB::ISEc55 containing strains expressed type 1 fimbriae compared to 70.6% of strains with an intact fimB gene, X^2 (1, N = 91) = 29.87, P < 0.0001. This significant difference was also observed following overnight static growth, X^2 (1, N = 91) = 9.063, P < 0.01, but no significant association was seen between fimB status and type 1 fimbriae expression when strains were further statically subcultured (Table 1; days 2 and 3). By three successive rounds of static subculture, the majority of ST131 strains were positive for type 1 fimbriae expression irrespective of fimB gene status. However, a step-wise increase in the proportion of type 1 positive strains was only seen for those

containing the *fimB::ISEc55* insertion, suggesting a slower rate of 'on' switching for most of these strains.

In our analysis, ten strains (five with an intact *fimB* and five with a *fimB::ISEc55* insertion) remained negative for type 1 fimbriae expression and were further subjected to six additional rounds of 24-hour static subculture. This resulted in the expression of type 1 fimbriae by four strains, three out of which had an intact *fimB* gene. Of the six remaining strains (S1, S18, S27, S36, S61, S77), S36 (Supporting Information Fig. S1) and S77 (7) had large deletions within coding regions of the type 1 fimbrial operon, accounting for their negative yeast agglutination phenotype. To test the capacity of *fimS* to invert to an 'on' orientation in S1, S18, S27 and S61, each strain was transformed with a plasmid containing the *fimB* gene from CFT073 (pFimB) and subjected to static culture followed by a screening PCR to determine *fimS* orientation (24). Overexpression of FimB in this manner resulted in *fimS* inversion to an 'on' orientation in three of the strains (S1, S18 and S61), however functional type 1 fimbriae expression as assessed by yeast cell agglutination could only be detected for S18 (Table 2). Sequence analysis of *fimS* from S27 revealed a single cytosine deletion (Supporting Information Fig. S2) 8bp downstream of the left inverted repeat in *fimS* (31), which may account for the 'locked off' status of *fimS* in this strain.

The transcription of fimE is enhanced in ST131 strains with a fimB::ISEc55 insertion

We hypothesized that the *fimB*::IS*Ec55* insertion may alter the transcription of the downstream *fimE* recombinase gene. In order to test this, we examined the level of *fimE* transcription using qRT-PCR in EC958 as well as four additional clade C ST131 strains with the same *fimB*::IS*Ec55* insertion (S4, S54, S60, S88), and compared this to the level of *fimE* transcription in four ST131

strains with an intact fimB gene (S17, S31, S55, S90). Strains were grown in M9 minimal
glucose medium with aeration to mid-log phase for analysis. In all cases, the level of fimE
transcription was significantly higher in strains with the fimB::ISEc55 insertion compared to
strains with an intact <i>fimB</i> gene (overall 6.26-fold increase in <i>fimE</i> ; Fig. 2A; <i>P</i> <0.05).

FimX is another accessory tyrosine recombinase that contributes to turning 'on' *fimS* in UPEC (32). Given that all nine ST131 strains used for this analysis also screened positive for the *fimX* gene by PCR, *fimX* transcript levels were also examined. There was no significant difference in *fimX* transcription between the two strain sets (Fig. 2B).

The fimB::ISEc55 insertion does not contain an additional fimE promoter

The transcription start site (TSS) of *fimE* has previously been mapped to a T residue 166 bases upstream of the GTG start codon in the E. coli K-12 strain VL751 (17). As we observed significantly higher *fimE* transcript levels in ST131 strains with the *fimB*::ISEc55 insertion, we hypothesized that there may be an additional promoter located within the insertion element that could drive fimE transcription. The fimE TSS was mapped using 5' RACE to a G residue 164 bases upstream of the start codon in all five of the fimB::ISEc55 ST131 strains tested (EC958, S4, S54, S60, S88; Fig. 3). As a control, we also mapped the TSS of *fimE* in one ST131 strain with an intact fimB gene (S90), and showed that it was identical. A consensus -35 (TTGTTA) and -10 (AAAATA) promoter sequence, separated by 18 bp, was conserved in all ST131 strains examined.

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Since the TSS was conserved in all ST131 strains tested, consensus sequences of fimE promoter and coding regions (1074 bp) from clades A, B and C strains were constructed from previously published genome sequences (7) to ascertain if there were any differences that could explain the higher *fimE* transcription observed in clade C strains. These were compared to corresponding fimE sequences from two non-ST131 UPEC strains (CFT073 and UTI89) and E. coli K-12 strain MG1655 by multiple sequence alignment (Supporting Information Fig. S3). Sequences from ST131 clade A and C strains were 100% identical and closely related to *fimE* from MG1655. In contrast, the sequence of the *fimE* promoter and coding region from ST131 clade B strains contained several nucleotide changes, and was most similar to the corresponding sequence from CFT073 and UTI89. The most parsimonious explanation for our observation in light of the known phylogeny of ST131 (7) is an independent recombination in the *fimE* region of clade B strains after divergence from clade C. This is distinct from the previously reported fimD-uxuR recombination event in clade C that encompassed the *fimH*30 allele (28). Nucleotide changes in the coding region were synonymous with only a single amino acid substitution (V198A) found in FimE from clade B ST131 strains (also present in UTI89, Supporting Information Fig. S3). These changes, however, are unlikely to account for the increased *fimE* transcript levels observed in strains with the *fimB*::ISEc55 insertion, as this is exclusively found in clade C ST131 strains.

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FimE and FimX are responsible for inverting fimS to the 'on' orientation in EC958

Despite the lack of an intact *fimB* gene, the majority of clade C ST131 strains still expressed functional type 1 fimbriae after multiple rounds of static growth, indicating that they possess an alternative recombinase capable of switching 'on' *fimS*. Based on genomic analysis, EC958 contains two tyrosine-like recombinase genes that have been shown to mediate inversion of *fimS*

in other <i>E. coli</i> strains; namely <i>fimE</i> (15) and <i>fimX</i> (24-26). We therefore constructed a series of
isogenic mutants in EC958 lacking fimE (EC958fimE), fimX (EC958fimX) or both genes
(EC958fimE fimX) and monitored their capacity to express type 1 fimbriae by yeast agglutination
and western blot analysis (using an antibody against the major fimbrial subunit FimA) following
static growth with repeated subculture over a period of 5 days (Fig. 4).

Type 1 fimbriae expression was detected in EC958 following 3 successive rounds of 24-hour static growth. In contrast, type 1 fimbriae expression by EC958fimE was delayed and only observed after 4 successive rounds of static growth, demonstrating a direct role for FimE in 'off' to 'on' fimS inversion (Fig. 4A). This also implicated a role for FimX in fimS inversion, and although the single EC958fimX mutant demonstrated a type 1 fimbriae expression profile identical to wild-type, the EC958fimE fimX double mutant failed to express type 1 fimbriae after prolonged static subculture for up to 5 days (Fig. 4A). This result was further validated by complementation of the EC958fimE fimX double mutant with a plasmid containing either fimE or fimX (pFimE or pFimX), both of which resulted in rapid type 1 fimbriae production within two days of static subculture (Fig. 4B). Thus, our data demonstrate that FimE is the major recombinase responsible for 'off' to 'on' fimS inversion in EC958, and that FimX can also mediate this switching mechanism, albeit less efficiently than FimE during in vitro static growth.

Direct quantitation of fim switching in EC958 using Illumina sequencing

In order to quantitate the efficiency of FimE- and FimX-mediated *fimS* inversion in EC958, we utilized a novel read-mapping method based on Illumina sequencing data that we refer to as DNA Invertible Switch Counter (or DISCus). EC958, EC958*fimE* and EC958*fimX* were grown

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under static conditions as described in the previous section. Aliquots from the air-liquid interface of each subculture were harvested on days 1-5, and genomic DNA was extracted and sequenced using Illumina Technology. Using DISCus, we determined the percentage of reads that corresponded to either the 'on' or 'off' orientation of fimS at each time point by read mapping to fimS pseudo-reference sequences representing both orientations. The starting inoculum for these experiments was an overnight shaking culture, which possessed only 1% fimS 'on' reads. In the static subculture time course, EC958 produced a steadily increasing fimS 'on' switching profile, with a maximum of 71% 'on' reads at day 5 (Fig. 5A). EC958fimX produced a similar fimS switching profile, with the exception that 'on' switching was slower on day 2 compared to wildtype (Fig. 5A). In contrast, fimS 'on' switching in EC958fimE occurred at a significantly slower rate. Only 11.6% of 'on' reads were detected at day 4 of static subculture, which then increased to a level comparative to EC958 and EC958fimX at day 5. DISCus analysis of the EC958fimE fimX double mutant did not identify any significant fimS 'on' switching, supporting our observation that FimE and FimX are the only recombinases capable of inverting fimS in EC958. Overall, the quantitative fimS switching data closely paralleled the type 1 fimbriae expression analysis for each strain based on yeast agglutination and FimA protein production (Fig. 4).

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To examine the specificity of FimE and FimX for *fimS* inversion in greater detail, we complemented the EC958*fimE fimX* double mutant by introduction of a plasmid containing *fimE* (pFimE) or *fimX* (pFimX). The EC958*fimE fimX* (pSU2718, empty vector control), EC958*fimE fimX*(pFimE) and EC958*fimE fimX*(pFimX) strains were subcultured by static growth over 3 days as described above, and DISCus was used to analyze the *fimS* orientation from Illumina sequences generated at each time point. Interestingly, data from these experiments demonstrated

246	that when the recombinases were over-expressed in this manner, FimX exhibited greater
247	propensity for <i>fimS</i> 'off' to 'on' inversion than FimE (Fig. 5B).
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249	FimE does not invert the fimX switch in EC958
- - - - - - - - - - - -	Time does not invert the junix switch in EC750
250	The fimX switch, located between the fimX-hyxR genes, comprises two 16 bp inverted repeats
251	flanking a 278 bp central region (27). In addition to mediating in trans inversion of fimS (27),
252	FimX also mediates in cis bidirectional inversion of this switch. To assess this in greater detail,
253	we analyzed the fimX switch inversion rate from the sequence data generated above. In the
254	experiments that involved repeated static subculture of EC958, EC958fimE and EC958fimX, the
255	fimX switch remained primarily in the 'off' orientation (92-97% 'off') throughout the 5-day
256	static subculture period (Fig. 6A). However, in the experiments involving subculture of the
257	complemented strains EC958fimE fimX(pFimE) and EC958fimE fimX(pFimX), and the control
258	strain EC958fimE fimX(pSU2718), we observed very rapid FimX-mediated fimX switch
259	inversion within a day (Fig. 6B). Taken together, these results demonstrate a rapid and strong
260	interaction of FimX with its cognate switch. Furthermore, these results also demonstrate for the
261	first time that FimE is unable to invert the <i>fimX</i> switch in EC958.
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263	H-NS is not a major repressor of fimE gene expression in EC958
264	The histone-like nucleoid structuring protein (H-NS) is a global transcriptional regulator (33). H-
265	NS has been shown to directly regulate type 1 fimbriae expression in E. coli K-12 by binding to
266	sequences adjacent to and within fimS (34) and also to the promoters of fimB and fimE (17, 35).
267	At 37°C, H-NS down-regulates fimE transcription and hns deletion leads to increased fimE
268	promoter activity (35). We constructed a hns deletion in EC958 and three non-ST131 UPEC

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strains (UTI89, 536 and IHE3034) and measured *fimE* transcript levels of all wild-type strains and corresponding *hns* mutants by qRT-PCR. At 37°C, deletion of *hns* led to significantly increased *fimE* transcription compared to the wild-type in all non-ST131 UPEC strains (Fig. 7B-D). However, in EC958, there was only a relatively small increase in *fimE* transcription in the *hns* mutant compared to the wild-type (Fig. 7A). Thus, H-NS does not strongly repress *fimE* transcription in EC958.

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Identification of genes that affect fimE and fimX promoter activity in EC958

In order to investigate the regulation of the fimE and fimX genes in EC958, two promoterreporter strains were generated. The *lacIZ* genes of EC958 were initially inactivated to generate strain EC958lac, and this strain was subsequently modified by inserting the lacZ gene as a chromosomally located transcriptional fusion to the fimE (EC958fimE::lacZ) or fimX (EC958fimX::lacZ) promoter. When grown on LB agar containing X-gal, EC958fimE::lacZ colonies were pale blue (indicating weak transcription) and EC958fimX::lacZ colonies were white (indicating no transcription). The two reporter strains were then subjected to Tn5 mutagenesis; in each case 50,000-60,000 transposon mutants were generated and visually screened based on their blue colour intensity following growth on LB X-gal agar. Individual colonies that were dark blue (compared to the control strain) were selected and further analyzed for β -galactosidase activity. Genes identified to contain transposon insertions that altered the β galactosidase activity of EC958fimE::lacZ or EC958fimX::lacZ are listed in Tables 3 and 4, respectively. Strikingly, multiple independent mutants containing insertions in the guaB gene were identified for both EC958fimE::lacZ and EC958fimX::lacZ; thus the remainder of this study focused on understanding the regulatory role of *guaB* on *fimE* and *fimX* transcription.

Deletion of guaB in EC958 leads to up-regulation of the fimE and fimX recombinase genes but

slower fimS 'on' switching In order to study how guaB regulates fimE and fimX, we constructed defined deletion mutants in EC958fimE::lacZ and EC958fimX::lacZ to generate the following strains: EC958fimE::lacZ guaB and EC958fimX::lacZ guaB. Activity from the fimE and fimX promoters was measured by determining the β-galactosidase activity from control and guaB mutant strains (Fig. 8A, B). Deletion of guaB resulted in significantly increased promoter activity from both EC958fimE::lacZ guaB (2-fold, P < 0.05, Fig. 8A) and EC958fimX::lacZ guaB (1.5-fold, P < 0.05, Fig. 8B). Furthermore, complementation of the mutant strains by over-expressing guaB in trans (via plasmid pGuaB) resulted in promoter activity comparable to the parental controls. These

results strongly suggest that the guaB gene product contributes to the regulation of fimE and fimX

in EC958.

To further assess the effect of *guaB* on *fimS* inversion, we generated an EC958*guaB* mutant and determined its switching profile using DISCus. Static subculture of EC958 and EC958*guaB* was performed over a 5-day period as described in previous sections. DISCus analysis of *fimS* was performed on samples at days 1, 3 and 5. Analysis of samples taken at days 3 and 5 revealed a significantly slower 'on' switching profile for EC958*guaB* compared to EC958 (Fig. 8C). Furthermore, this slower *fimS* switching rate was restored to wild-type level following complementation of EC958*guaB* with plasmid pGuaB (Fig. 8C). Taken together, this data strongly supports a role for the *guaB* gene in *fimS* inversion.

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The effect of guab deletion on fimE and fimX transcription is partially dependent on GTP homeostasis and mimics (p)ppGpp stress response conditions Guanosine tetra- and pentaphosphate, also known as (p)ppGpp, is a stress response alarmone that has been shown to increase type 1 fimbriae expression by up-regulating fimB and to a lesser extent, fimE transcription in E. coli (36). GuaB catalyzes the rate-limiting step in GMP biosynthesis and its deletion depletes the intracellular GTP pool (37, 38). Production of large 321 amounts of (p)ppGpp can also result in a sharp decline in the intracellular GTP level both by 322 direct GTP consumption and by (p)ppGpp-mediated GuaB inhibition (38, 39). Thus, (p)ppGpp is 323 involved in regulating intracellular GTP homeostasis (38). We hypothesized that the effect of guab deletion on fimE and fimX transcription in EC958 may be mediated through (p)ppGpp and tested this using pooled human urine as a growth medium. We predicted that guaB deletion would induce the stringent response as a result of reduced cellular GTP and trigger (p)ppGpp production by the action of purine salvage pathways (40) utilizing guanine and guanosine species present in human urine (41, 42). Elevated levels of (p)ppGpp, in turn, would cause increased transcription of fimE and fimX. To test this, we first used a sub-inhibitory concentration of serine hydroxamate (SHX, 0.2 mM) to induce (p)ppGpp generation within the cell via depletion of serine-charged t-RNA, leading to activation of the

333 amino acid starvation pathway (43). Stationary phase cultures of EC958fimE::lacZ and 334 EC958fimX::lacZ grown in the presence or absence of SHX were analyzed for β-galactosidase 335 activity (Fig. 9). The addition of SHX led to a significant de-repression of the *fimE* promoter

(Fig. 9A; P<0.05), albeit not to the level seen in EC958fimE::lacZ guaB. No significant change

was observed in *fimX* promoter activity (Fig. 9B).

Next, we attempted to replenish the depleted intracellular GTP pool in *guaB* deletion reporter strains by direct supplementation with GTP. The rationale behind this was that restoring intracellular GTP levels would rescue the cell from stress, leading to a decline in (p)ppGpp production and a subsequent decrease in *fimE* and *fimX* transcription. Thus, EC958*fimE::lacZ guaB* and EC958*fimX::lacZ guaB* were grown in pooled human urine supplemented with 0.5 mM GTP. For both strains, the addition of GTP led to a significant decrease in *fimE* (P<0.01) and *fimX* (P<0.05) promoter activities to levels similar to those observed in the parent EC958*fimE::lacZ* (Fig. 9C) and EC958*fimX::lacZ* strains (Fig. 9D). Taken together, these results suggest that GTP depletion in a *guaB* mutant background mimics conditions observed during (p)ppGpp stress and is at least partially responsible for increased transcription of *fimE* and *fimX*.

- The guaB gene is required for EC958 persistence in urine and bladder colonization in a mouse
- 351 UTI model
- In order to assess the contribution of guaB to the virulence of EC958, we tested the ability of the EC958guaB mutant to survive in the mouse urinary tract using a competitive infection assay. We employed an EC958lac strain as the wild-type to enable differentiation of both strains on MacConkey lactose medium; EC958lac colonized the mouse bladder in equivalent numbers to wild-type EC958 in a mixed competitive infection (Supporting Information Fig. S4A, B). EC958lac and EC958guaB inocula were prepared by four successive rounds of static subculture in LB broth (3x48 hours followed by 1x24 hours); under these conditions both strains expressed similar levels of type 1 fimbriae (Supporting Information Fig. S4C). Female C57BL/6 mice were

co-inoculated with 1:1 ratio of EC958lac and EC958guaB, and the colonization of each strain

was ass	sessed at 2	24 hours	post in	fection.	In t	these	experiments	, EC9	58guaB	was	signific	antly
outcomp	peted by E	C958lac	in the b	ladder (P<0.	.01) aı	nd urine (P<	0.05)	of infect	ed mi	ice (Fig.	10).



Discussion

The emergence and rapid spread of the UPEC ST131 clone has been well documented since 2008 with epidemiological reports from all over the world. Like other UPEC, *E. coli* ST131 strains utilize type 1 fimbriae for colonization of the urinary tract. However, most ST131 isolates from the globally predominant clade C sub-lineage possess an IS*Ec55* insertion element within the tyrosine-like recombinase *fimB* gene that is associated with reduced type 1 fimbriae expression (8, 28). Here, we define the molecular basis of *fimS* inversion in this clinically-relevant fluoroquinolone-resistant group of *E. coli* ST131 strains.

The high prevalence of the *fimB*::ISEc55 insertion in ST131 strains from clade C led us to assess the impact of this insertion on type 1 fimbriae expression. We have previously reported a delayed 'on' switching phenotype of type 1 fimbriae expression in a subset of the strains used in the current study (8). We therefore compared the type 1 fimbriae expression profile of a large collection of ST131 strains that possessed an intact or disrupted *fimB* gene over multiple rounds of static subculture. Overall, in accordance with previous observations (8), most ST131 strains containing the *fimB*::ISEc55 insertion were able to express type 1 fimbriae, but exhibited a slower 'off' to 'on' switching profile during static growth. Interestingly, in the present analysis, six strains were unable to express type 1 fimbriae after prolonged subculture, suggesting they lack the capacity to make functional type 1 fimbriae. Two of these strains had large deletions in the *fim* operon and transformation of the remaining four strains with a plasmid containing *fimB* induced type 1 fimbriae expression in only one strain. All three remaining strains harboured the *fimX* recombinase gene previously reported to switch *fimS* 'on' (24), indicating that they may contain other mutations that disrupt this phenotype.

The presence of the ISEc55 insertion element in *fimB* was associated with increased transcription of the downstream *fimE* gene in the clade C representative EC958 strain and four other clade C ST131 strains. Mapping of the *fimE* promoter in these strains revealed that ISEc55 did not contain an independent promoter driving *fimE* transcription. However, we did observe that deletion of H-NS, a global transcriptional regulator and a known repressor of *fimE* (35), did not have a major effect on *fimE* transcription in EC958. H-NS is known to bind preferentially to intrinsically-curved DNA and AT-rich sequences, usually found in promoter regions, to exert regulatory effects on its many target genes (44, 45). Our observations indicate an *in trans* effect of ISEc55 that could alter the local DNA structure and inhibit H-NS binding. Indeed, analogous alterations in the transcription of genes adjacent to other insertion elements have been documented in *E. coli* (46-48).

Analysis of *E. coli* ST131 genomes has revealed that the *fim* region is associated with recombination (7, 28). Here, we provide a comprehensive analysis of type 1 fimbriae regulation in fluoroquinolone-resistant clade C ST131 strains that contain the *fimB*::ISEc55 insertion. The expression of type 1 fimbriae by the majority of ST131 strains containing the *fimB*::ISEc55 insertion suggested that they possess an alternative recombinase/s capable of *fimS* inversion. Using a combined mutagenesis and complementation strategy, we showed that both FimE and FimX could mediate *fimS* inversion in EC958, with FimE demonstrating the greatest *fimS* inversion activity under the static growth conditions employed in this study. We did not observe any significant difference in *fimX* transcript levels in ST131 strains with an intact *fimB* and those with the *fimB*::ISEc55 insertion. We also developed DISCus, a novel read-mapping approach

based on Illumina sequencing, to quantitatively determine the percentage of *fimS* 'on' or 'off' populations. Our data suggest that *fimE* plays a more important role than *fimX* in regulating the orientation of *fimS* during *in vitro* culture. This is in agreement with a previous report that detected relatively low levels of FimX activity *in vitro*, but observed rapid expression of type 1 fimbriae upon experimental infection in strains with FimX as the only active recombinase (25). The role of FimX in the inversion of *fimS* in EC958 during *in vivo* infection remains to be elucidated. In this respect, DISCus could be used to monitor *fimS* orientation in such populations. We also note that DISCus could be applied more broadly to analyze DNA invertible elements in other bacteria.

To identify potential regulators of the *fimE* and *fimX* genes in EC958, we employed random transposon mutagenesis of the *fimE* and *fimX* reporter strains and identified several genes that when mutated caused increased activity of the *fimE* and *fimX* promoters. The *guaB* gene, which encodes an enzyme that catalyzes the first rate limiting step in *de novo* GTP/GDP biosynthesis and is responsible for maintaining GTP homeostasis (49), was identified in screens of both reporter strains and was the focus of subsequent experimental analysis (see below). Mutation of several other genes was also linked with increased *fimE* promoter activity (*yubO*, *pdxH*, *lrhA*, *dprA*) or increased *fimX* promoter activity (*betA*, *yjjA*). The LysR family transcriptional regulator *lrhA* has previously been shown to repress both *fimB* and *fimE* (50), and our identification of *lrhA* in this study as a repressor of *fimE* is consistent with these results. The *yubO* gene is located on a 135.6 kb IncF plasmid in EC958 and encodes a protein of unknown function (51). Three independent Tn5 mutants were identified in *yubO*, however specific mutagenesis of *yubO* failed to reproduce a significant increase in *fimE* transcription (data not shown). Thus, a definite role

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for *yubO* in type 1 fimbriae regulation remains to be demonstrated. The *dprA* gene encodes a putative DNA processing protein implicated in natural bacterial transformation (52) and has been previously reported to be expressed *in vivo* during UTI (53). However, the molecular mechanism by which *dprA* might affect *fimE* transcription, as well as the role of *pdxH* (encodes an enzyme required for vitamin B6 biosynthesis (54)), *betA* (encodes a choline dehydrogenase (55)) and *yjjA* (encodes a putative metal chaperone) in type 1 fimbriae expression, remains to be determined.

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We focused our molecular investigation on the role of the guaB gene in fimE and fimX regulation. While there is no direct link between guaB and expression of type 1 fimbriae in the literature, guaB has been shown to be upregulated in women with UTI (53) and is important for UPEC fitness in an experimental model of systemic infection (56). In other studies, the (p)ppGpp alarmone, synthesized from GTP, has been shown to increase type 1 fimbriae expression in E. coli (36). Strains lacking any (p)ppGpp exhibit reduced transcription from the fimB and fimE promoters, although the effect on the latter gene is approximately half of that reported for fimB (36). Under conditions of nutritional stress, (p)ppGpp controls global metabolic changes within the cell (57-59) via direct interaction with RNA polymerase (60, 61) and activation of stressassociated sigma factors (57). Synthesis of (p)ppGpp results in a reduction of the intracellular GTP pool (62-64). Recently, it has been reported that (p)ppGpp directly inhibits various GTP biosynthesis enzymes, thus further lowering the level of cellular GTP (38). We hypothesized that deletion of guaB would produce a GTP deficient state mimicking that observed during the activation of the stringent response pathway, and that this in turn would increase fimE and fimX promoter activity. Chemical induction of this pathway in EC958 increased fimE promoter activity and the addition of GTP to the culture media of guaB mutant strains significantly reduced both *fimE* and *fimX* promoter activity. Taken together, these data support the hypothesis that the cellular GTP concentration influences type 1 fimbriae expression by controlling the transcription of *fimE* and *fimX*. We suggest that this effect is partly mediated by the action of the (p)ppGpp alarmone on *fimE* promoter activity in strains that lack a functional copy of the *fimB* gene. Indeed, DISCus analysis of EC958guaB revealed an overall slower *fimS* 'on' switching phenotype compared to the wild-type EC958, possibly due to the strong natural bias for FimE to mediate 'on' to 'off' switching (14, 65). These findings are also consistent with our mouse colonization data, which showed that an EC958guaB mutant was significantly outcompeted by wild-type EC958 with respect to bladder colonization and persistence in urine.

In conclusion, this work has identified and characterized several novel molecular mechanisms associated with the regulation of type 1 fimbriae in *E. coli* ST131 strains that harbour an IS*Ec55* insertion in the *fimB* recombinase gene. Our data demonstrate that type 1 fimbriae regulation in these strains is different to that described for non-ST131 UPEC, highlighting the importance of understanding the control of this essential virulence factor in the context of a clinically dominant MDR resistant clone. Overall, the impact of this mutation on the fitness and global dissemination of ST131 remains to be determined. However, given that the only non-ST131 strain shown to harbour the *fimB*::IS*Ec55* insertion to date is the probiotic Nissle 1917 strain (8), it is tempting to speculate that such modulation of type 1 fimbriae expression may be associated with enhanced colonization of the gut.

Experimental Procedures

Ethics statement

Approval for mouse infection studies was obtained from the University of Queensland Animal Ethics Committee (SCMB/471/09/NHMRC (NF)). Experiments were carried out in strict accordance with the recommendations in the Animal Care and Protection Act (Queensland, 2002) and the Australian Code of Practice for the Care and Use of Animals for Scientific Purposes (8th edition, 2013). Approval for the collection of human urine was obtained from the University of Queensland Institutional Human Research Ethics Committee (2015000347) and the Griffith University Human Ethics Research Committee (MSC/06/08/HREC). Participation was voluntary and all individuals provided informed consent prior to participation in the study.

Bacterial strains and growth conditions

The *E. coli* strains and plasmids used in this study are listed in Table S1 in the Supporting Information. *E. coli* EC958 is a completely sequenced fluoroquinolone-resistant Clade C ST131 strain originally isolated in the United Kingdom in 2005 (7, 8, 29). Other *E. coli* ST131 strains were isolated from urine samples collected at hospitals in Manchester and Preston, Northwest England (n=54) and Brisbane, Australia (n=37). Most of the isolates (n=71) used in the present study have been described previously (7, 8, 29, 66-68). The additional 21 strains were collected from Brisbane, Australia as part of routine methods for UTI diagnosis. Strains were routinely cultured at 37°C on solid or in liquid lysogeny broth (LB) unless otherwise specified. Culture media was supplemented with appropriate antibiotics as required: gentamicin (Gent, 20 μg ml⁻¹), chloramphenicol (Cm, 30 μg ml⁻¹). Human urine, when used as culture media, was pooled from at least three healthy donors with no recent history of antibiotic use.

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DNA manipulations and genetic techniques

Chromosomal DNA purification, PCR and DNA sequencing was performed as previously described (69). PCR screening for the presence of *fimH* (70) and the insertion element within *fimB* was performed as previously reported (8). The sequence type of UTI isolates used in this study was determined by multilocus sequence typing as previously described (4, 66, 67). DNA extraction for Illumina sequencing was performed using the Ultraclean® Microbial DNA Isolation Kit (MO BIO Laboratories). Restriction enzymes were purchased from New England Biolabs (Genesearch, Australia). All mutants, tagged strains and plasmid constructs were confirmed by PCR followed by Sanger sequencing performed at the Australian Equine Genetics Research Centre (Queensland, Australia). Illumina sequencing was performed at the Australian Genome Research Facility (Melbourne, Australia).

Construction of plasmids

The plasmids used for *in trans* complementation were created by PCR amplification of the respective gene from EC958 (*fimB* was amplified from CFT073) using primers listed in Supporting Information Table S1. Relevant PCR products were digested with SacI-HindIII and ligated into SacI-HindIII-digested pSU2718 (71). The cloned genes were under the control of a *lac* promoter; cloned genes were induced by the addition of 1 mM IPTG (Sigma Aldrich) to the culture medium.

Construction of deletion mutants and complemented strains

EC958 mutants were constructed using λ -Red mediated homologous recombination as previously described (8) using primers listed in Table S1. A three-step PCR procedure was employed for the generation of products with 500 bp homology regions used in recombination (72). Plasmid pCP20 (73) was modified by the introduction of a gentamicin resistance cassette at the NcoI site to yield pCP20-Gent. For complementation of mutants, removal of the FRT-site-flanked chloramphenical cassette was mediated by the Flp recombinase expressed by pCP20-Gent. Mutant strains were then complemented by introduction of the respective genes cloned into the low-copy number pSU2718 plasmid. All *lacZ* reporter-tagged transcriptional fusions were generated in an *mKate2*-tagged EC958*lac* background using PCR products with a *lacZ-Cm* cassette flanked by 500 bp homology regions specific to *fimE* or *fimX*.

Type 1 fimbriae expression profile

A type 1 fimbriae expression profile was established for all 91 *E. coli* ST131 strains examined in this study. Strains were grown overnight on LB agar, and single colonies were picked and used to inoculate a 5 ml LB for overnight aerobic (shaking) growth. A volume of 1 μl of each overnight culture was inoculated into a new tube containing 5 ml LB and cultured for 24 hours under static conditions. Strains were sub-cultured from the air-liquid interface in a similar manner for further growth. Type 1 fimbriae expression was determined at each stage of subculture using a yeast cell agglutination assay as previously described (8). Type 1 fimbriae expression of EC958 wild-type and mutant derivatives was monitored similarly. The orientation of *fimS* was evaluated from single colonies by PCR as previously described with primers 2848, 2859 and 2850 listed in Table S1 (24). Cells from colonies with *fimS* in the 'off' orientation were used to inoculate 5 ml LB broth and cultured for 24 hours under static conditions; a volume of 10

μl was taken from the air-liquid interface and used to generate a new subculture every 24 hours over a 5-day period. For each subculture, type 1 fimbriae expression was monitored by yeast cell agglutination and western blot analysis of whole cell lysates using a FimA-specific antibody. A volume of 1 ml was also collected from the air-liquid interface at each 24-hour time point; cells were pelleted by centrifugation, genomic DNA was extracted and sequenced using Illumina technology for DISCus analysis.

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RNA isolation, cDNA synthesis and quantitative RT-PCR (qRT-PCR) analysis

RNA was extracted from bacteria grown aerobically at 37°C in M9 minimal medium supplemented with 0.2% glucose (w/v) (74). Mid-log phase cultures were standardized to OD₆₀₀ = 0.6 prior to RNA extraction using the RNeasy Protect Bacteria Mini Kit (Qiagen) following the manufacturer's guidelines. RNA was treated with RNase-free DNase I to remove contaminating DNA and re-purified using Qiagen RNeasy columns. RNA samples were quantified spectrophotometrically at 260nm. cDNA synthesis was performed with 1 µg total RNA and random hexamer primers using the SuperScript®III First Strand Synthesis System (Life Technologies) according to the manufacturer's instructions. All cDNA samples were diluted 10fold prior to semi-quantitative real time PCR (qRT-PCR) analysis. qRT-PCR reactions were performed using SYBR® Green Master Mix (Applied Biosystems) and 500 nM gene specific primers for fimE, fimX and gapA (Supporting Information Table S1) on a ViiATM 7 instrument (Life technologies) with the following cycling parameters: initial denaturation at 95°C for 10 min followed by 40 cycles of denaturation at 95°C for 15 s, annealing at 60°C for 15 s and elongation at 72°C for 30 s. Each reaction was performed in quadruplicate. Amplicon specificity was confirmed via gel and melting curve analysis. The $ViiA^{TM}$ 7 software determined the threshold

cycle (C_t) for each reaction. The expression of fimE and fimX was normalized to gapA expression
for each sample and the $2^{-\Delta\Delta Ct}$ method (75) was employed to calculate the difference in
expression of the fim recombinase genes between strains with an intact fimB and those with a
fimB::ISEc55 insertion.

Rapid amplification of cDNA ends (5' RACE)

The transcription start site (TSS) of *fimE* was mapped using the 5' RACE system (Life Technologies) according to manufacturer's guidelines. Briefly, total RNA was isolated as described above and first strand cDNA synthesis was performed using SuperScript™ II and a gene-specific primer (Supporting Information Table S1, primer 4342). The resulting cDNA was purified and a homopolymeric tail was added to its 3'-end using TdT and dCTP. The cDNA was then amplified by PCR using an anchor primer (supplied) and a gene-specific primer (Supporting Information Table S1, primer 4343) followed by another round of nested PCR (primer 4344). The purified PCR product was then sequenced to determine the TSS.

- Sequence analysis of fimE
- Nucleotide sequences of ST131 *fimE* promoter and coding regions (1074 bp) were obtained from previously published genomes of strains used in this study (n=26) (7). The corresponding sequences from two reference non-ST131 UPEC strains, UTI89 (GI: 91209055) and CFT073 (GI: 26111730) and the *E. coli* K-12 strain MG1655 (GI: 556503834) were included for comparison. Multiple sequence alignment was performed using Clustal X v2.0 (76) to construct a neighbour-joining phylogenetic tree which was then visualised with FigTree v1.4.2

[http://tree.bio.ed.ac.uk/software/figtree/]. Amino acid sequences of the FimE protein (198 AA) were also aligned using Clustal X2.

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Quantification of the fimS promoter switching frequency using DISCus

To accurately measure invertible DNA switching, a read mapping approach was developed to determine the frequency of Illumina sequence reads that mapped uniquely to the 'on' or 'off' orientation of the *fimS* invertible switch. To determine the ratio of 'off' to 'on' mapped reads, a pseudo-reference containing both fimS orientations and 1 kbp of flanking sequence was constructed using the relevant sequence from EC958 (GI:641682562). An analogous approach was taken to construct a pseudo-reference for the hyxR promoter region. Illumina 100 bp pairedend reads from ST131 strains were then mapped to the pseudo-reference sequences using Burrows-Wheeler Aligner (77), and inter-converted to SAM (Sequence Alignment/Map) and BAM (Binary Alignment/Map) files using SAMtools where necessary (78). To ascertain the number of reads overlapping the unique bordering regions of the 'off' and 'on' orientations, and thereby their relative switching frequencies, Bedmaps was used to count reads overlapping 10 bp pseudo-exons, which were created at both edges of the 'off' and 'on' orientations (79). The number of paired-reads, which traversed these bordering regions, was also counted. This was accomplished by assigning six regions in the pseudo-reference, corresponding to 'left flank', 'right flank' or 'switch region' for both the 'off' and 'on' orientation. Paired end read locations were determined from the BAM file and counted if they were found to traverse from a 'switch region' to either 'flank' for the 'off' or 'on' orientation respectively. Reads were not counted if they were unpaired or traversed from an 'on' region to an 'off' region. Final read counts were normalised as a percentage of total input reads to allow comparison between independent samples. Python and Bash scripts for generating pseudo-reference sequences and automating DISCus, respectively, are available in a public repository at https://github.com/LeahRoberts/DISCus.

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Transposon mutagenesis analysis

A mini Tn5-Cm transposon containing a chloramphenicol (Cm) cassette flanked by Tn5 mosaic ends was PCR amplified from NotI-digested pKD3 plasmid with primers 2279 and 2280 (Supporting Information Table S1). The amplicon was phosphorylated using T4 polynucleotide kinase (New England Biolabs) and mixed with EZ-Tn5TM transposase (Epicenter Biotechnologies) to create transposomes according to the manufacturer's guidelines. Electrocompetent cells of EC958 fimE and fimX promoter-lacZ transcriptional fusion strains were prepared as previously described (80). Cells were electroporated with transposomes and after a 2 h recovery period in SOC, were spread on LB plates supplemented with Cm and 5bromo-4-chloro-3-indolyl-D-galactoside (X-Gal). Approximately 50,000 to 60,000 Cm resistant mutants were obtained for each strain. Mutant colonies were visually screened for increased blue color intensity on X-Gal plates using the parental strains as controls. Selected mutants were further screened for lacZ expression by measurement of β -galactosidase activity. The transposon insertion site of mutants with altered β-galactosidase activity was determined using an arbitrary PCR (using primers 2209 and 1799; Supporting Information Table S1) followed by a nested PCR (using primers 2240 and 1801) and sequencing of the final amplicon. Mutation of the guaB gene in EC958fimE::lacZ-cm and EC958fimE::lacZ-cm was performed using λ-Red homologous recombination as described above.

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636	β-Galactosidase	assays
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β-Galactosidase assays were performed essentially as previously described (81). Briefly, strains carrying *fimE* and *fimX* promoter-*lacZ* transcriptional fusions were grown overnight in pooled human urine alone or supplemented with 0.2 mM serine hydroxamate (SHX) or 0.5 mM GTP as indicated. Cultures were diluted in Z-buffer (60 mM Na₂HPO₄, 40 mM NaH₂PO₄, 50 mM β-mercaptoethanol, 10 mM KCl, 1 mM MgSO₄ at pH 7.0) followed by the addition of 0.004% SDS (w/v) and chloroform to permeabilize the cells. The samples were incubated at 28°C, and reactions were initiated by the addition of *ortho*-Nitrophenyl-β-D-galactopyranoside (ONPG). The reactions were stopped with the addition of Na₂CO₃ (300 mM). The enzymatic activity (expressed in Miller units) was assayed in quadruplicate for each strain by measuring the absorbance at 420 nm.

Mouse model of UTI

The C57BL/6 mouse model of ascending UTI was employed as previously described (69). All strains used in this experiment were enriched for type 1 fimbriae expression by three successive rounds of static growth in LB for 48 hours followed by one round of static growth for 24 hours for inoculum preparation. All strains were positive for type 1 fimbriae expression as determined by yeast cell agglutination and western blot analysis. Infections were performed as competitive assays; the inoculum contained 1:1 strain mixture of EC958 WT (*lac*⁻ strain) and EC958*guaB*. Bacteria (5x10⁸ CFU in 30 µl PBS) were injected transurethrally into female C57BL/6 mice (8-10 weeks). Urine was collected from mice 24 h post challenge, after which they were euthanized by cervical dislocation. Bacterial loads in urine and bladder were enumerated as colony forming

658	units (CFU) counts. The two strains were differentiated by colony colour on MacConkey plates:
659	white (lac WT) or pink (EC958guaB).

Statistical Analysis

All statistical analyses were performed using the GraphPad Prism 6 software package. Significant associations between fimB status and type 1 fimbriae expression were investigated with the $\chi 2$ test. Statistical significance of fold-changes in fimE transcript levels between ST131 strains was determined using a two-tailed Mann Whitney test. The Wilcoxon matched pairs signed-rank test and repeated-measures one-way ANOVA with Dunnett's multiple comparisons test were used to analyse results from β -galactosidase assays. A Pearson's $\chi 2$ test with Yates' continuity correction was used to compare raw read counts generated by DISCus using RStudio (v0.98.501). The significance level was adjusted to 0.0167 (alpha = 0.05, Bonferroni adjusted for three pairwise comparisons). Data from mouse UTI experiments was analyzed using a two-tailed Wilcoxon matched pairs signed-rank test. In all instances, statistical significance was considered at P<0.05.

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References

- Russo TA, Johnson JR. 2003. Medical and economic impact of extraintestinal infections due to Escherichia coli: focus on an increasingly important endemic problem. Microbes Infect 5:449-456.
- Flores-Mireles AL, Walker JN, Caparon M, Hultgren SJ. 2015. Urinary tract infections: epidemiology, mechanisms of infection and treatment options. Nat Rev Microbiol 13:269-284.
- Coque TM, Novais A, Carattoli A, Poirel L, Pitout J, Peixe L, Baquero F, Canton R,
 Nordmann P. 2008. Dissemination of clonally related Escherichia coli strains expressing extended-spectrum beta-lactamase CTX-M-15. Emerg Infect Dis 14:195-200.
- 4. Lau SH, Kaufmann ME, Livermore DM, Woodford N, Willshaw GA, Cheasty T, Stamper K, Reddy S, Cheesbrough J, Bolton FJ, Fox AJ, Upton M. 2008. UK epidemic Escherichia coli strains A-E, with CTX-M-15 beta-lactamase, all belong to the international O25:H4-ST131 clone. Journal of Antimicrobial Chemotherapy 62:1241-1244.
- Nicolas-Chanoine MH, Blanco J, Leflon-Guibout V, Demarty R, Alonso MP,
 Canica MM, Park YJ, Lavigne JP, Pitout J, Johnson JR. 2008. Intercontinental
 emergence of Escherichia coli clone 025: H4-ST131 producing CTX-M-15. Journal of
 Antimicrobial Chemotherapy 61:273-281.
- 704 6. Price LB, Johnson JR, Aziz M, Clabots C, Johnston B, Tchesnokova V, Nordstrom L, Billig M, Chattopadhyay S, Stegger M, Andersen PS, Pearson T, Riddell K, Rogers P, Scholes D, Kahl B, Keim P, Sokurenko EV. 2013. The epidemic of extended-spectrum-beta-lactamase-producing Escherichia coli ST131 is driven by a single highly pathogenic subclone, H30-Rx. MBio 4:e00377-00313.
- 709 7. Petty NK, Ben Zakour NL, Stanton-Cook M, Skippington E, Totsika M, Forde BM,
 710 Phan MD, Gomes Moriel D, Peters KM, Davies M, Rogers BA, Dougan G,
 711 Rodriguez-Bano J, Pascual A, Pitout JD, Upton M, Paterson DL, Walsh TR,
 712 Schembri MA, Beatson SA. 2014. Global dissemination of a multidrug resistant
 713 Escherichia coli clone. Proc Natl Acad Sci U S A 111:5694-5699.
- 714 8. **Totsika M, Beatson SA, Sarkar S, Phan MD, Petty NK, Bachmann N, Szubert M,**715 **Sidjabat HE, Paterson DL, Upton M, Schembri MA.** 2011. Insights into a multidrug
 716 resistant Escherichia coli pathogen of the globally disseminated ST131 lineage:
 717 genome analysis and virulence mechanisms. PLoS One **6:**e26578.
- 718 9. **Totsika M, Kostakioti M, Hannan TJ, Upton M, Beatson SA, Janetka JW, Hultgren**719 **SJ, Schembri MA.** 2013. A FimH inhibitor prevents acute bladder infection and
 720 treats chronic cystitis caused by multidrug-resistant uropathogenic Escherichia coli
 721 ST131. J Infect Dis **208**:921-928.
- 722 10. Blomfield IC. 2001. The regulation of pap and type 1 fimbriation in Escherichia coli.
 723 Adv Microb Physiol 45:1-49.
- 724 11. **Klemm P, Jorgensen BJ, van Die I, de Ree H, Bergmans H.** 1985. The fim genes responsible for synthesis of type 1 fimbriae in Escherichia coli, cloning and genetic organization. Mol Gen Genet **199:**410-414.

- 727 12. **Abraham JM, Freitag CS, Clements JR, Eisenstein BI.** 1985. An invertible element of DNA controls phase variation of type 1 fimbriae of Escherichia coli. Proc Natl Acad Sci U S A **82:**5724-5727.
- 730 13. **Klemm P.** 1986. Two regulatory fim genes, fimB and fimE, control the phase variation of type 1 fimbriae in Escherichia coli. EMBO J **5:**1389-1393.
- 732 14. **Gally DL, Leathart J, Blomfield IC.** 1996. Interaction of FimB and FimE with the fim switch that controls the phase variation of type 1 fimbriae in Escherichia coli K-12. Mol Microbiol **21:**725-738.
- 735 15. **Stentebjerg-Olesen B, Chakraborty T, Klemm P.** 2000. FimE-catalyzed off-to-on inversion of the type 1 fimbrial phase switch and insertion sequence recruitment in an Escherichia coli K-12 fimB strain. FEMS Microbiol Lett **182:**319-325.
- 738 16. **Smith SG, Dorman CJ.** 1999. Functional analysis of the FimE integrase of Escherichia coli K-12: isolation of mutant derivatives with altered DNA inversion preferences. Mol Microbiol **34:**965-979.
- 741 17. **Olsen PB, Klemm P.** 1994. Localization of promoters in the fim gene cluster and the effect of H-NS on the transcription of fimB and fimE. FEMS Microbiol Lett **116:**95-100.
- 744 18. **Blomfield IC, Kulasekara DH, Eisenstein BI.** 1997. Integration host factor stimulates both FimB- and FimE-mediated site-specific DNA inversion that controls phase variation of type 1 fimbriae expression in Escherichia coli. Mol Microbiol **23:**705-717.
- 748 19. **Blomfield IC, Calie PJ, Eberhardt KJ, McClain MS, Eisenstein BI.** 1993. Lrp stimulates phase variation of type 1 fimbriation in Escherichia coli K-12. J Bacteriol **175:**27-36.
- 751 20. **Schwan WR.** 2011. Regulation of *fim* genes in uropathogenic *Escherichia coli*. World J Clin Infect Dis **1:**17-25.
- 753 21. **Old DC, Duguid JP.** 1970. Selective outgrowth of fimbriate bacteria in static liquid medium. J Bacteriol **103:**447-456.
- 755 22. **Hasman H, Schembri MA, Klemm P.** 2000. Antigen 43 and type 1 fimbriae determine colony morphology of Escherichia coli K-12. J Bacteriol **182:**1089-1095.
- Hung C, Zhou Y, Pinkner JS, Dodson KW, Crowley JR, Heuser J, Chapman MR, Hadjifrangiskou M, Henderson JP, Hultgren SJ. 2013. Escherichia coli biofilms have an organized and complex extracellular matrix structure. MBio 4:e00645-00613.
- 761 24. **Bryan A, Roesch P, Davis L, Moritz R, Pellett S, Welch RA.** 2006. Regulation of type 1 fimbriae by unlinked FimB- and FimE-like recombinases in uropathogenic Escherichia coli strain CFT073. Infect Immun **74:**1072-1083.
- 764 25. **Hannan TJ, Mysorekar IU, Chen SL, Walker JN, Jones JM, Pinkner JS, Hultgren**765 **SJ, Seed PC.** 2008. LeuX tRNA-dependent and -independent mechanisms of Escherichia coli pathogenesis in acute cystitis. Mol Microbiol **67:**116-128.
- 767 26. **Xie Y, Yao Y, Kolisnychenko V, Teng CH, Kim KS.** 2006. HbiF regulates type 1 fimbriation independently of FimB and FimE. Infect Immun **74**:4039-4047.
- 769 27. **Bateman SL, Seed PC.** 2012. Epigenetic regulation of the nitrosative stress response and intracellular macrophage survival by extraintestinal pathogenic Escherichia coli. Mol Microbiol **83:**908-925.

- 772 28. **Paul S, Linardopoulou EV, Billig M, Tchesnokova V, Price LB, Johnson JR,**773 **Chattopadhyay S, Sokurenko EV.** 2013. Role of homologous recombination in adaptive diversification of extraintestinal Escherichia coli. J Bacteriol **195:**231-242.
- Forde BM, Ben Zakour NL, Stanton-Cook M, Phan MD, Totsika M, Peters KM, Chan KG, Schembri MA, Upton M, Beatson SA. 2014. The complete genome sequence of Escherichia coli EC958: a high quality reference sequence for the globally disseminated multidrug resistant E. coli O25b:H4-ST131 clone. PLoS One 9:e104400.
- 780 30. **Schembri MA, Sokurenko EV, Klemm P.** 2000. Functional flexibility of the FimH adhesin: insights from a random mutant library. Infect Immun **68:**2638-2646.
- 782 31. **Gally DL, Rucker TJ, Blomfield IC.** 1994. The leucine-responsive regulatory protein binds to the fim switch to control phase variation of type 1 fimbrial expression in Escherichia coli K-12. J Bacteriol **176:**5665-5672.
- 785 32. **Bateman SL, Stapleton AE, Stamm WE, Hooton TM, Seed PC.** 2013. The Type 1 Pili Regulator Gene fimX and Pathogenicity Island PAI-X as Molecular Markers of Uropathogenic Escherichia coli. Microbiology doi:10.1099/mic.0.066472-0.
- 788 33. **Dorman CJ.** 2004. H-NS: a universal regulator for a dynamic genome. Nat Rev Microbiol **2:**391-400.
- 790 34. Schembri MA, Olsen PB, Klemm P. 1998. Orientation-dependent enhancement by
 791 H-NS of the activity of the type 1 fimbrial phase switch promoter in Escherichia coli.
 792 Mol Gen Genet 259:336-344.
- 793 35. **Olsen PB, Schembri MA, Gally DL, Klemm P.** 1998. Differential temperature modulation by H-NS of the fimB and fimE recombinase genes which control the orientation of the type 1 fimbrial phase switch. FEMS Microbiol Lett **162**:17-23.
- 796 36. **Aberg A, Shingler V, Balsalobre C.** 2006. (p)ppGpp regulates type 1 fimbriation of Escherichia coli by modulating the expression of the site-specific recombinase FimB. Mol Microbiol **60**:1520-1533.
- 799 37. **Ochi K.** 1987. Metabolic initiation of differentiation and secondary metabolism by Streptomyces griseus: significance of the stringent response (ppGpp) and GTP content in relation to A factor. J Bacteriol **169**:3608-3616.
- 802 38. **Kriel A, Bittner AN, Kim SH, Liu K, Tehranchi AK, Zou WY, Rendon S, Chen R, Tu**803 **BP, Wang JD.** 2012. Direct regulation of GTP homeostasis by (p)ppGpp: a critical
 804 component of viability and stress resistance. Mol Cell **48**:231-241.
- 39. **Gallant J, Irr J, Cashel M.** 1971. The mechanism of amino acid control of guanylate and adenylate biosynthesis. J Biol Chem **246**:5812-5816.
- 807 40. **Vogels GD, Van der Drift C.** 1976. Degradation of purines and pyrimidines by microorganisms. Bacteriol Rev **40**:403-468.
- Sander G, Topp H, Wieland J, Heller-Schoch G, Schoch G. 1986. Possible use of urinary modified RNA metabolites in the measurement of RNA turnover in the human body. Hum Nutr Clin Nutr 40:103-118.
- Andreoli R, Manini P, De Palma G, Alinovi R, Goldoni M, Niessen WM, Mutti A. 2010. Quantitative determination of urinary 8-oxo-7,8-dihydro-2'-deoxyguanosine, 8-oxo-7,8-dihydroguanine, 8-oxo-7,8-dihydroguanosine, and their non-oxidized forms: daily concentration profile in healthy volunteers. Biomarkers **15:**221-231.
- 816 43. **Shand RF, Blum PH, Mueller RD, Riggs DL, Artz SW.** 1989. Correlation between histidine operon expression and guanosine 5'-diphosphate-3'-diphosphate levels

- during amino acid downshift in stringent and relaxed strains of Salmonella typhimurium. J Bacteriol **171:**737-743.
- Yamada H, Yoshida T, Tanaka K, Sasakawa C, Mizuno T. 1991. Molecular analysis
 of the Escherichia coli hns gene encoding a DNA-binding protein, which
 preferentially recognizes curved DNA sequences. Mol Gen Genet 230:332-336.
- 45. Gordon BR, Li Y, Cote A, Weirauch MT, Ding P, Hughes TR, Navarre WW, Xia B,
 Liu J. 2011. Structural basis for recognition of AT-rich DNA by unrelated xenogeneic silencing proteins. Proc Natl Acad Sci U S A 108:10690-10695.
- Petersen C, Moller LB, Valentin-Hansen P. 2002. The cryptic adenine deaminase gene of Escherichia coli. Silencing by the nucleoid-associated DNA-binding protein, H-NS, and activation by insertion elements. J Biol Chem 277:31373-31380.
- 829 47. **Barker CS, Pruss BM, Matsumura P.** 2004. Increased motility of Escherichia coli by insertion sequence element integration into the regulatory region of the flhD operon. J Bacteriol **186:**7529-7537.
- 832 48. **Reynolds AE, Felton J, Wright A.** 1981. Insertion of DNA activates the cryptic bgl operon in E. coli K12. Nature **293:**625-629.
- 49. **Hedstrom L.** 2009. IMP dehydrogenase: structure, mechanism, and inhibition. Chem Rev **109:**2903-2928.
- 836 50. **Blumer C, Kleefeld A, Lehnen D, Heintz M, Dobrindt U, Nagy G, Michaelis K,**837 **Emody L, Polen T, Rachel R, Wendisch VF, Unden G.** 2005. Regulation of type 1
 838 fimbriae synthesis and biofilm formation by the transcriptional regulator LrhA of
 839 Escherichia coli. Microbiology **151**:3287-3298.
- Phan MD, Forde BM, Peters KM, Sarkar S, Hancock S, Stanton-Cook M, Ben Zakour NL, Upton M, Beatson SA, Schembri MA. 2015. Molecular Characterization of a Multidrug Resistance IncF Plasmid from the Globally Disseminated Escherichia coli ST131 Clone. PLoS One 10:e0122369.
- 52. **Johnston C, Martin B, Fichant G, Polard P, Claverys JP.** 2014. Bacterial transformation: distribution, shared mechanisms and divergent control. Nat Rev Microbiol **12**:181-196.
- Hagan EC, Lloyd AL, Rasko DA, Faerber GJ, Mobley HL. 2010. Escherichia coli global gene expression in urine from women with urinary tract infection. PLoS Pathog 6:e1001187.
- Zhao G, Winkler ME. 1995. Kinetic limitation and cellular amount of pyridoxine (pyridoxamine) 5'-phosphate oxidase of Escherichia coli K-12. J Bacteriol 177:883-852
 891.
- Styrvold OB, Falkenberg P, Landfald B, Eshoo MW, Bjornsen T, Strom AR. 1986.
 Selection, mapping, and characterization of osmoregulatory mutants of Escherichia coli blocked in the choline-glycine betaine pathway. I Bacteriol **165:**856-863.
- 856 56. **Subashchandrabose S, Smith SN, Spurbeck RR, Kole MM, Mobley HL.** 2013. Genome-wide detection of fitness genes in uropathogenic Escherichia coli during systemic infection. PLoS Pathog **9:**e1003788.
- Paul BJ, Barker MM, Ross W, Schneider DA, Webb C, Foster JW, Gourse RL. 2004. DksA: a critical component of the transcription initiation machinery that potentiates the regulation of rRNA promoters by ppGpp and the initiating NTP. Cell 118:311-322.

- Barker MM, Gaal T, Josaitis CA, Gourse RL. 2001. Mechanism of regulation of transcription initiation by ppGpp. I. Effects of ppGpp on transcription initiation in vivo and in vitro. J Mol Biol **305**:673-688.
- 866 59. **Murphy H, Cashel M.** 2003. Isolation of RNA polymerase suppressors of a (p)ppGpp deficiency. Methods Enzymol **371:**596-601.
- 60. **Chatterji D, Fujita N, Ishihama A.** 1998. The mediator for stringent control, ppGpp, binds to the beta-subunit of Escherichia coli RNA polymerase. Genes Cells **3:**279-287.
- 871 61. **Artsimovitch I, Patlan V, Sekine S, Vassylyeva MN, Hosaka T, Ochi K, Yokoyama**872 **S, Vassylyev DG.** 2004. Structural basis for transcription regulation by alarmone ppGpp. Cell **117:**299-310.
- 874 62. **Torok I, Kari C.** 1980. Accumulation of ppGpp in a relA mutant of Escherichia coli during amino acid starvation. J Biol Chem **255**:3838-3840.
- Johnson GS, Adler CR, Collins JJ, Court D. 1979. Role of the spoT gene product and manganese ion in the metabolism of guanosine 5'-diphosphate 3'-diphosphate in Escherichia coli. J Biol Chem **254:**5483-5487.
- 879 64. **Lopez JM, Dromerick A, Freese E.** 1981. Response of guanosine 5'-triphosphate concentration to nutritional changes and its significance for Bacillus subtilis sporulation. J Bacteriol **146**:605-613.
- 882 65. **McClain MS, Blomfield IC, Eisenstein BI.** 1991. Roles of fimB and fimE in sitespecific DNA inversion associated with phase variation of type 1 fimbriae in 884 Escherichia coli. J Bacteriol **173:**5308-5314.
- 885 66. **Lau SH, Reddy S, Cheesbrough J, Bolton FJ, Willshaw G, Cheasty T, Fox AJ, Upton M.** 2008. Major uropathogenic Escherichia coli strain isolated in the northwest of
 887 England identified by multilocus sequence typing. Journal of Clinical Microbiology
 888 **46:**1076-1080.
- 889 67. **Sidjabat HE, Derrington P, Nimmo GR, Paterson DL.** 2010. Escherichia coli ST131 producing CTX-M-15 in Australia. Journal of Antimicrobial Chemotherapy **65:**1301-1303.
- 68. **Gibreel TM, Dodgson AR, Cheesbrough J, Fox AJ, Bolton FJ, Upton M.** 2012. Population structure, virulence potential and antibiotic susceptibility of uropathogenic Escherichia coli from Northwest England. J Antimicrob Chemother **67:**346-356.
- 896 69. **Allsopp LP, Totsika M, Tree JJ, Ulett GC, Mabbett AN, Wells TJ, Kobe B, Beatson**897 **SA, Schembri MA.** 2010. UpaH is a newly identified autotransporter protein that contributes to biofilm formation and bladder colonization by uropathogenic Escherichia coli CFT073. Infect Immun **78:**1659-1669.
- Johnson JR, Stell AL. 2000. Extended virulence genotypes of Escherichia coli strains
 from patients with urosepsis in relation to phylogeny and host compromise. J Infect
 Dis 181:261-272.
- 903 71. **Martinez E, Bartolome B, de la Cruz F.** 1988. pACYC184-derived cloning vectors containing the multiple cloning site and lacZ alpha reporter gene of pUC8/9 and pUC18/19 plasmids. Gene **68:**159-162.
- 906 72. **Derbise A, Lesic B, Dacheux D, Ghigo JM, Carniel E.** 2003. A rapid and simple method for inactivating chromosomal genes in Yersinia. FEMS Immunol Med Microbiol **38**:113-116.

- 909 73. **Cherepanov PP, Wackernagel W.** 1995. Gene disruption in Escherichia coli: TcR and KmR cassettes with the option of Flp-catalyzed excision of the antibiotic-resistance determinant. Gene **158**:9-14.
- 912 74. **Sambrook J, Fritsch EF, Maniatis T.** 1989. Molecular cloning: a laboratory manual, 2nd edition., 2 ed. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY.
- 914 75. **Livak KJ, Schmittgen TD.** 2001. Analysis of relative gene expression data using real-time quantitative PCR and the 2(-Delta Delta C(T)) Method. Methods **25:**402-408.
- 76. Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, McWilliam H,
 Valentin F, Wallace IM, Wilm A, Lopez R, Thompson JD, Gibson TJ, Higgins DG.
 2007. Clustal W and Clustal X version 2.0. Bioinformatics 23:2947-2948.
- 920 77. **Li H, Durbin R.** 2009. Fast and accurate short read alignment with Burrows-921 Wheeler transform. Bioinformatics **25:**1754-1760.
- 922 78. **Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, Marth G, Abecasis G,**923 **Durbin R.** 2009. The Sequence Alignment/Map format and SAMtools.
 924 Bioinformatics **25:**2078-2079.
- Neph S, Kuehn MS, Reynolds AP, Haugen E, Thurman RE, Johnson AK, Rynes E,
 Maurano MT, Vierstra J, Thomas S, Sandstrom R, Humbert R,
 Stamatoyannopoulos JA. 2012. BEDOPS: high-performance genomic feature operations. Bioinformatics 28:1919-1920.
- 929 80. Langridge GC, Phan MD, Turner DJ, Perkins TT, Parts L, Haase J, Charles I, 930 Maskell DJ, Peters SE, Dougan G, Wain J, Parkhill J, Turner AK. 2009. Simultaneous assay of every Salmonella Typhi gene using one million transposon mutants. Genome Res 19:2308-2316.
- 933 81. **Miller JH.** 1992. A short course in bacterial genetics: a laboratory manual and handbook for *Escherichia coli* and related bacteria, vol 1. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY.
- 936 82. **Mobley HL, Green DM, Trifillis AL, Johnson DE, Chippendale GR, Lockatell CV,**937 **Jones BD, Warren JW.** 1990. Pyelonephritogenic Escherichia coli and killing of
 cultured human renal proximal tubular epithelial cells: role of hemolysin in some
 strains. Infect Immun **58:**1281-1289.
- 940 83. **Berger H, Hacker J, Juarez A, Hughes C, Goebel W.** 1982. Cloning of the chromosomal determinants encoding hemolysin production and mannose-resistant hemagglutination in Escherichia coli. J Bacteriol **152:**1241-1247.
- 943 84. **Mulvey MA, Schilling JD, Hultgren SJ.** 2001. Establishment of a persistent Escherichia coli reservoir during the acute phase of a bladder infection. Infect Immun **69:**4572-4579.
- 946 85. **Korhonen TK, Valtonen MV, Parkkinen J, Vaisanen-Rhen V, Finne J, Orskov F,**947 **Orskov I, Svenson SB, Makela PH.** 1985. Serotypes, hemolysin production, and
 948 receptor recognition of Escherichia coli strains associated with neonatal sepsis and
 949 meningitis. Infect Immun **48**:486-491.
- 950 86. **Allsopp LP, Beloin C, Ulett GC, Valle J, Totsika M, Sherlock O, Ghigo JM,** 951 **Schembri MA**. 2012. Molecular characterization of UpaB and UpaC, two new autotransporter proteins of uropathogenic Escherichia coli CFT073. Infect Immun **80:**321-332.

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954	87.	Phan MD, Peters KM, Sarkar S, Lukowski SW, Allsopp LP, Moriel DG, Achard
955		ME, Totsika M, Marshall VM, Upton M, Beatson SA, Schembri MA. 2013. The
956		Serum Resistome of a Globally Disseminated Multidrug Resistant Uropathogenic
957		Clone, PLoS Genet 9: e1003834.

- 958 88. **Allsopp LP, Beloin C, Moriel DG, Totsika M, Ghigo JM, Schembri MA.** 2012. Functional heterogeneity of the UpaH autotransporter protein from uropathogenic Escherichia coli. J Bacteriol **194:**5769-5782.
 - 89. **Chaveroche MK, Ghigo JM, d'Enfert C.** 2000. A rapid method for efficient gene replacement in the filamentous fungus Aspergillus nidulans. Nucleic Acids Res **28:**E97.

966 Tables

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Table 1. Type 1 fimbriae expression profile of *E. coli* ST131 strains

Number of yeast agglutination positive ST131 strains			
following overnight growth in LB broth at 37°C (%)			
Shaking			
culture	Static subcu	ılture	
Day 1	Day 1	Day 2	Day 3
8 (14)	27 (47.4)	34 (59.6)	52 (91.2)
24 (70.6)***	27 (79.4)*	27 (79.4)	29 (85.3)
	Shaking culture Day 1 8 (14)	following overnight growth in Shaking culture Static subculture Day 1 Day 1 8 (14) 27 (47.4)	following overnight growth in LB broth at Shaking culture Static subculture Day 1 Day 1 Day 2 8 (14) 27 (47.4) 34 (59.6)

^{*}P < 0.01, ***P < 0.0001; $\chi 2$ test.

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Table 2. Effect of FimB over-expression in type 1 fimbriae negative strains

_			fimS	Yeast
Strain	fimB status	Genotype	orientation	Agglutination*
S1	fimB::ISEc55	Wild-type	OFF	-
		+ pFimB	ON	-
S18	fimB::ISEc55	Wild-type	OFF	-
		+ pFimB	ON	+
S27	Intact	Wild-type	OFF	-
		+ pFimB	OFF	-
S61	fimB::ISEc55	Wild-type	OFF	-
		+ pFimB	ON	-

972 *Yeast agglutination negative (-), positive (+)

Table 3. Mutated genes associated with increased *fimE* promoter activity

EC958 locus tag	Gene	Product	No. independent Tn5 insertions	β-gal activity as Miller units ^a (mean±SD)	Fold-change relative to control ^b (mean±SD)
EC958_2815	guaB	inosine 5' monophosphate dehydrogenase	3	2102±56, 1321±28, 792±36	4.95±0.94 3.11±0.59 1.86±0.36
EC958_A0030	yubO	Unknown function, plasmid located	3	1175±139, 830±70, 702±40	2.76±0.61 1.95±0.40 1.65±0.32
EC958_1861	pdxH	pyridoxamine 5' phosphate oxidase	2	819±30, 1936±131	1.93±0.36 4.56±0.91
EC958_2624	lrhA	LysR homologue A	1	2348±365	5.52±1.35
EC958_3829	dprA	putative DNA processing protein	1	1358±59	3.20±0.62

976 SD, Standard deviation.

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977 ^aValues represent data from each independently isolated Tn5 mutant.

^bβ-gal activity of control strain EC958*fimE::lacZ* = 425 ± 80 .

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Table 4. Mutated genes associated with increased *fimX* promoter activity

EC958 locus tag	Gene	Product	No. independent Tn5 insertions	β-gal activity as Miller units (mean±SD)	Fold-change relative to control ^a (mean±SD)
EC958_2815	guaB	inosine 5' monophosphate dehydrogenase	1	94±7	1.52±0.15
EC958_0463	betA	choline dehydrogenase	1	108±9	1.74±0.18
EC958_0090	yjjA	putative metal chaperone	1	94±7	1.52±0.15

982 SD, Standard deviation.

983 ^aβ-gal activity of control strain EC958*fimX::lacZ* = 62 ± 4 .

Table 5. Strains and plasmids used in this study

Strain ^a	Description	Reference	
CFT073	Wild-type UPEC isolate	(82)	
536	Wild-type UPEC isolate	(83)	
UTI89	Wild-type UPEC isolate	(84)	
IHE3034	Wild-type UPEC isolate	(85)	
EC958	Wild-type UPEC isolate; E. coli ST131 ESBL	(8)	
S1 pFimB	E. coli ST131 strain S1 pFimB, ESBL, Cm ^r	This study	
S18 pFimB	E. coli ST131 strain S18 pFimB, ESBL, Cm ^r	This study	
S27 pFimB	E. coli ST131 strain S27 pFimB, ESBL, Cm ^r	This study	
S61 pFimB	E. coli ST131 strain S61 pFimB, ESBL, Cm ^r	This study	
CFT973hns	CFT073hns::kan, Kan ^r	(86)	
536hns	536hns::kan, Kan ^r	This study	
UTI89hns	UTI89hns::kan, Kan ^r	This study	
IHE3034hns	IHE3034hns::kan, Kan ^r	This study	
EC958hns	EC958hns::cm, ESBL, Cm ^r	(87)	
MS3198	CFT073lacIZ upaH:lacZ-zeo	(88)	
EC958 pKOBEG- Gent	EC958 pKOBEG-Gent, ESBL Gent ^r	(8)	
EC958fimE	EC958fimE::cm, ESBL Cm ^r	This study	
EC958fimX	EC958fimX::cm, ESBL Cm ^r	This study	
EC958fimE fimX	EC958 <i>fimE fimX</i> :: <i>cm</i> , ESBL Cm ^r	This study	
EC958fimE fimX pSU2718	EC958fimE fimX, ESBL Cm ^r	This study	
EC958fimE fimX pFimE	EC958fimE fimX pFimE, ESBL Cm ^r	This study	
EC958fimE fimX pFimX	EC958fimE fimX pFimX, ESBL Cm ^r	This study	
EC958lac::mKate2	EC958lac::mKate2::cm, ESBL Cm ^r	(87)	
EC958fimE::lacZ-cm	EC958lac::mKate2 fimE::lacZ-cm, ESBL Cm ^r	This study	
EC958fimX::lacZ-cm	EC958lac::mKate2 fimX::lacZ-cm, ESBL Cm ^r	This study	
EC958fimE::lacZ		This study	

EC958fimX::lacZ guaB	EC958lac::mKate2 fimX::lacZ guaB::cm, ESBL Cm ^r	This study	
EC958fimE::lacZ guaB (pGuaB)	EC958 <i>lac::mKate2 fimE::lacZ guaB::cm</i> pGuaB, ESBL Cm ^r	This study	
EC958fimX::lacZ guaB (pGuaB)	EC958lac::mKate2 fimX::lacZ guaB::Cm pGuaB, ESBL Cm ^r	This study	
Plasmid			
pKOBEG-Gent	λ-Red plasmid, replicates at 37°C, <i>araC</i> arabinose inducible promoter, Gent ^r	(8, 89)	
pCP20-Gent	Replicates at 30°C, encodes Flp recombinase, Gent ^r	This study, (73)	
pSU2718	Cloning vector, Plac promoter, replicates at 37°C, Cm ^r	(71)	
pFimB	pSU2718 fimB; fimB coding sequence from CFT073, Cm ^r	This study	
pFimE	pSU2718 fimE; fimE coding sequence from EC958, Cm ^r	This study	
pFimX	pSU2718 fimX; fimX coding sequence from EC958, Cm ^r	This study	
pGuaB	pSU2718 <i>guaB</i> ; <i>guaB</i> coding sequence from EC958, Cm ^r	This study	
^a In addition to the strains listed above, our <i>E. coli</i> ST131 strain collection comprised of 54 strains			
from the United Kingdom (S1-S54, including EC958; (7, 8, 29, 66, 68) and 37 strains from			
Australia (S55-S91; (7, 8, 67); Cm ^r -chloramphenicol, Gent ^r -gentamicin resistant, Kan ^r -			
kanamycin resistant.			

992	Figure	Legends
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- 993 Fig. 1. Schematic representation of the type 1 fimbrial operon. The 1895bp fimB::ISEc55
- insertion sequence element is depicted in its genomic context. CFT073, reference UPEC strain;
- 995 EC958, representative E. coli ST131 clade C strain; adapted from Totsika et al.(8).

- 997 **Fig. 2.** Quantification of *fimE* and *fimX* transcription in ST131 strains by qRT-PCR.
- 998 The level of fimE (A) and fimX (B) transcription relative to housekeeping gene gapA was
- 999 determined by qRT-PCR in mid-log phase ST131 strains with the *fimB*::ISEc55 insertion (n=5,
- including EC958) and compared to those with an intact *fimB* (n=4) to determine fold-difference
- in gene expression between the two groups using the $2^{-\Delta\Delta Ct}$ method. Bars represent group means
- ± standard deviation (SD). Asterisk indicates a statistically significant difference between the two
- groups (P<0.05) as determined by the two-tailed Mann Whitney test.

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- 1005 **Fig. 3.** Analysis of *fimE* promoter sequence.
- The transcription start site (TSS) of *fimE* was mapped by 5'RACE in EC958 and five other
- 1007 E. coli ST131 strains (S4, S54, S69, S88 and S90). Shown here is the consensus nucleotide
- sequence. The -35 and -10 promoter elements are underlined and the TSS mapped to a G residue
- 1009 164 bases upstream of the GTG start codon (indicated by the curved arrow).

- 1011 Fig. 4. Contribution of FimE and FimX to fimS inversion and type 1 fimbriae expression in
- 1012 EC958.

1013 Western blots demonstrating expression of the FimA major subunit protein of type 1 fimbriae 1014 using α-FimA antibody and yeast cell agglutination assays to monitor type 1 fimbriae expression 1015 (- negative, + positive fimbriae expression) in static cultures. 1016 A. Type 1 fimbriae expression was measured in EC958 (wild-type), EC958 fimE, EC958 fimX, 1017 and EC958fimE fimX over 5 days of static subculture. 1018 B. Type 1 fimbriae expression in EC958fimE fimX strains over 3 days of static subculture. 1019 EC958fimE fimX (pSU2718) was used as an empty vector control. Over-expression of FimE 1020 (pFimE) and FimX (pFimX) was induced by the addition of 1mM IPTG to the culture media. 1021 1022 Fig. 5. Quantification of fimS switch orientation bias in EC958 wild-type, mutants and 1023 complemented strains using DISCus. A. fimS orientation in EC958, EC958fimE, EC958fimX and EC958fimE fimX over 5 days of 1024 1025 static subculture quantified using DISCus. Samples from the air-liquid interface of each culture 1026 were taken on each day and used for genomic DNA extraction and Illumina sequencing. Bars 1027 represent *fimS* percentage 'on' population within each culture. 1028 B. DISCus analysis of *fimS* orientation following *in trans* complementation with FimE (pFimE) 1029 or FimX (pFimE) in the EC958fimE fimX double mutant over 3 days of static subculture. 1030 1031 Fig. 6. Quantification of fimX switch orientation bias in EC958 wild-type, mutants and 1032 complemented strains using DISCus. 1033 A. DISCus analysis of fimX switch orientation in EC958, EC958fimE and EC958fimX strains

culture population.

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over 5 days of static subculture. Bars represent percentage fimX switch 'on' reads within each

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1036 B. DISCus analysis of fimX switch orientation in EC958fimE fimX and in EC958fimE fimX 1037 following in trans complementation with FimE (pFimE) or FimX (pFimE) over 3 days of static 1038 subculture. 1039 1040 **Fig. 7.** Effect of *hns* deletion on *fimE* transcript level in different UPEC strains. 1041 A. Fold-change in *fimE* transcription in EC958hns compared to wild-type EC958 as determined 1042 by gRT-PCR. 1043 B-D. Fold-change in *fimE* transcription in non-ST131 wild-type UPEC strains and their 1044 corresponding hns deletion mutants, namely UTI89 (B), 536 (C) and IHE3034 (D) as determined 1045 by qRT-PCR. Vertical bars represent means±SD of three independent replicates. 1046 1047 **Fig. 8.** Effect of guaB deletion on fimE/fimX promoter activity and fimS switching in EC958. 1048 A-B. β-galactosidase activity of EC958 fimE (A) and fimX (B) promoter-lacZ reporter strains 1049 following overnight aerated growth in pooled human urine. All guaB mutant strains and 1050 complemented derivatives were constructed in their respective promoter-lacZ reporter 1051 backgrounds. Bars represent means \pm SD of at least three independent biological experiments. 1052 Asterisk indicates a statistically significant difference (P < 0.05) as determined by repeated-1053 measures one-way ANOVA with Dunnett's multiple comparisons test. 1054 C. DISCus analysis of the effect of guab deletion on fimS switching. EC958 1055 (wild-type), EC958guaB and EC958guaB(pGuaB) were grown statically in LB (37°C) over a 1056 period of 5 days. Over-expression of GuaB in EC958guaB(pGuaB) was induced by the addition

of 1mM IPTG. Genomic DNA was extracted after static subculture on days 1, 3 and 5, and

Illumina sequencing data was analyzed by DISCus. Bars represent percentage 'on' reads of *fimS*

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1059 in each culture population. Asterisks indicate a statistically significant difference (* P = 0.01-0.001; *** P < 0.0001) as determined by a Pearson's $\chi 2$ test with Yates' continuity correction. 1060

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1062 **Fig. 9.** Regulation of *fimE* and *fimX* promoter activity by (p)ppGpp.

1063 A-B. Effect of serine hydroxamate (SHX, 0.2 mM) induced amino acid starvation on the 1064 promoter activity of fimE (A) and fimX (B) in EC958fimE:lacZ and EC958fimX:lacZ strains, 1065 respectively. All guaB mutant strains and complemented derivatives were constructed in their 1066 respective promoter-lacZ reporter backgrounds. Strains were incubated overnight under static 1067 conditions in pooled human urine and promoter activity was determined by the β -galactosidase 1068 assay. Bars represent means ± SD of three independent biological experiments with four 1069 technical replicates. Asterisk indicates a statistically significant difference (P<0.05) as

1070 determined by the Wilcoxon matched pairs signed-rank test.

C-D. Effect of GTP (0.5 mM) on the promoter activity of fimE (C) and fimX (D) in EC958fimE::lacZ guaB and EC958fimX::lacZ guaB strains, respectively. Asterisks indicate statistically significant differences (**P<0.01, *P<0.05) in guaB reporter strains upon addition of GTP, as determined by a repeated-measures one-way ANOVA with Dunnett's multiple comparisons test.

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Fig. 10. Mouse urinary tract colonization by EC958 WT and EC958guaB. Female C57B/L6 mice were transurethrally inoculated with a 1:1 mixture of type 1 fimbriae enriched EC958 WT and EC958guaB strains in a competitive infection assay. Each marker represents total colony forming units (CFU) recovered from each mouse per 1 ml of urine or per 0.1 g of bladder tissue

(as labelled) on selective medium. Lines connect data points for the same mouse and horizontal bars represent median values. Asterisk indicates statistically significant difference between the strains for persistence in urine (P<0.05) as well as bladder colonization (P<0.01) as determined by the Wilcoxon matched pairs signed-rank test.



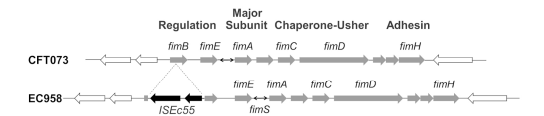


Fig. 1. Schematic representation of the type 1 fimbrial operon. The 1895bp fimB::ISEc55 insertion sequence element is depicted in its genomic context. CFT073, reference UPEC strain; EC958, representative E. coli ST131 clade C strain; adapted from Totsika et al.(8).

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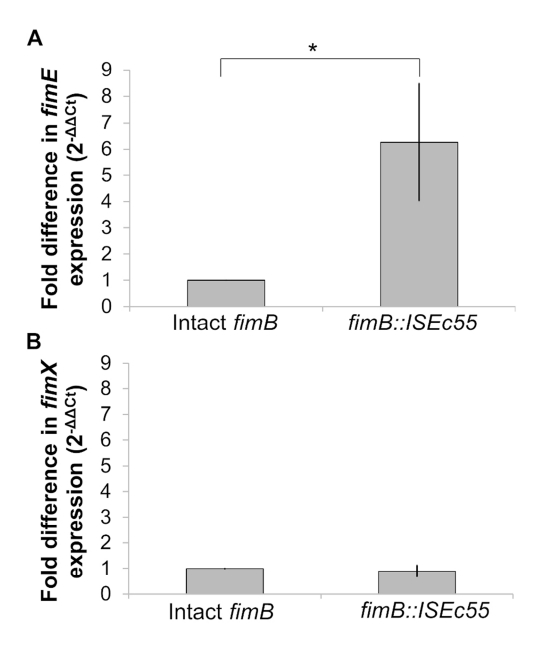


Fig. 2. Quantification of fimE and fimX transcription in ST131 strains by qRT-PCR. The level of fimE (A) and fimX (B) transcription relative to housekeeping gene gapA was determined by qRT-PCR in mid-log phase ST131 strains with the fimB::ISEc55 insertion (n=5, including EC958) and compared to those with an intact fimB (n=4) to determine fold-difference in gene expression between the two groups using the $2-\Delta\Delta$ Ct method. Bars represent group means \pm standard deviation (SD). Asterisk indicates a statistically significant difference between the two groups (P<0.05) as determined by the two-tailed Mann Whitney test. 78x95mm (300 x 300 DPI)

Fig. 3. Analysis of fimE promoter sequence. The transcription start site (TSS) of fimE was mapped by 5'RACE in EC958 and five other E. coli ST131 strains (S4, S54, S69, S88 and S90). Shown here is the consensus nucleotide sequence. The -35 and -10 promoter elements are underlined and the TSS mapped to a G residue 164 bases upstream of the GTG start codon (indicated by the curved arrow).

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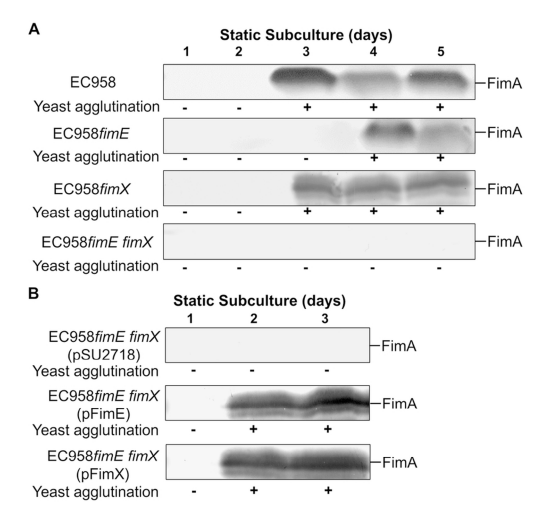


Fig. 4. Contribution of FimE and FimX to fimS inversion and type 1 fimbriae expression in EC958. Western blots demonstrating expression of the FimA major subunit protein of type 1 fimbriae using a-FimA antibody and yeast cell agglutination assays to monitor type 1 fimbriae expression (- negative, + positive fimbriae expression) in static cultures.

A. Type 1 fimbriae expression was measured in EC958 (wild-type), EC958fimE, EC958fimX, and EC958fimE fimX over 5 days of static subculture.

B. Type 1 fimbriae expression in EC958fimE fimX strains over 3 days of static subculture. EC958fimE fimX (pSU2718) was used as an empty vector control. Over-expression of FimE (pFimE) and FimX (pFimX) was induced by the addition of 1mM IPTG to the culture media.

76x74mm (300 x 300 DPI)

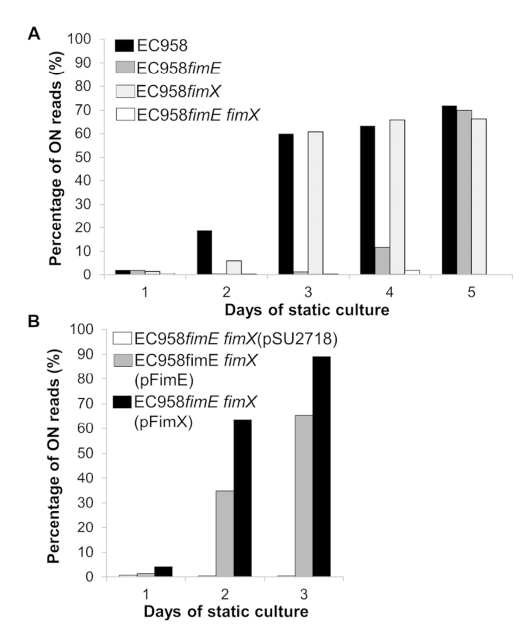


Fig. 5. Quantification of fimS switch orientation bias in EC958 wild-type, mutants and complemented strains using DISCus.

A. fimS orientation in EC958, EC958fimE, EC958fimX and EC958fimE fimX over 5 days of static subculture quantified using DISCus. Samples from the air-liquid interface of each culture were taken on each day and used for genomic DNA extraction and Illumina sequencing. Bars represent fimS percentage 'on' population within each culture.

B. DISCus analysis of fimS orientation following in trans complementation with FimE (pFimE) or FimX (pFimE) in the EC958fimE fimX double mutant over 3 days of static subculture.

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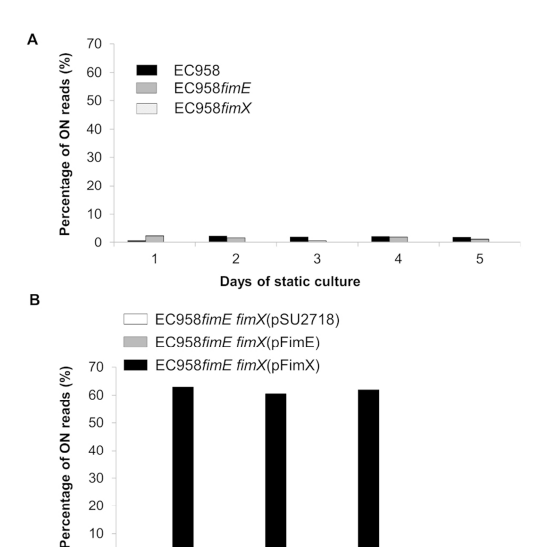


Fig. 6. Quantification of fimX switch orientation bias in EC958 wild-type, mutants and complemented strains using DISCus.

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Days of static culture

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- A. DISCus analysis of fimX switch orientation in EC958, EC958fimE and EC958fimX strains over 5 days of static subculture. Bars represent percentage fimX switch 'on' reads within each culture population.
- B. DISCus analysis of fimX switch orientation in EC958fimE fimX and in EC958fimE fimX following in trans complementation with FimE (pFimE) or FimX (pFimE) over 3 days of static subculture. 78x91mm (300 x 300 DPI)

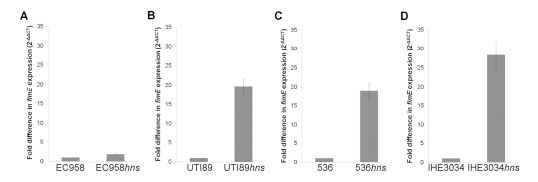


Fig. 7. Effect of hns deletion on fimE transcript level in different UPEC strains.

A. Fold-change in fimE transcription in EC958hns compared to wild-type EC958 as determined by qRT-PCR.

B-D. Fold-change in fimE transcription in non-ST131 wild-type UPEC strains and their corresponding hns deletion mutants, namely UTI89 (B), 536 (C) and IHE3034 (D) as determined by qRT-PCR. Vertical bars represent means±SD of three independent replicates.

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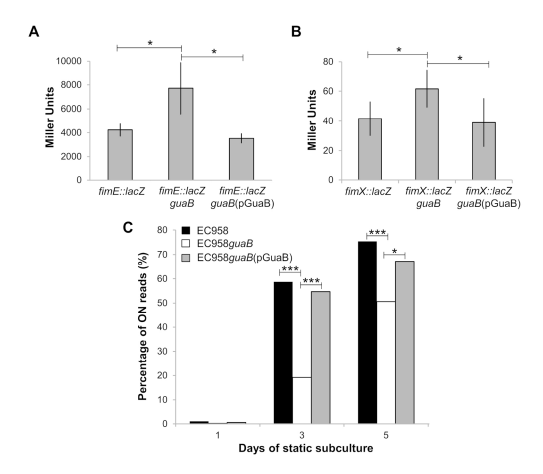


Fig. 8. Effect of guaB deletion on fimE/fimX promoter activity and fimS switching in EC958.

A-B. β-galactosidase activity of EC958 fimE (A) and fimX (B) promoter-lacZ reporter strains following overnight aerated growth in pooled human urine. All guaB mutant strains and complemented derivatives were constructed in their respective promoter-lacZ reporter backgrounds. Bars represent means ± SD of at least three independent biological experiments. Asterisk indicates a statistically significant difference (P<0.05) as determined by repeated-measures one-way ANOVA with Dunnett's multiple comparisons test.

C. DISCus analysis of the effect of guaB deletion on fimS switching. EC958 (wild-type), EC958guaB and EC958guaB(pGuaB) were grown statically in LB (37oC) over a period of 5 days. Over-expression of GuaB in EC958guaB(pGuaB) was induced by the addition of 1mM IPTG. Genomic DNA was extracted after static subculture on days 1, 3 and 5, and Illumina sequencing data was analyzed by DISCus. Bars represent percentage 'on' reads of fimS in each culture population. Asterisks indicate a statistically significant difference (* P = 0.01-0.001; *** P < 0.0001) as determined by a Pearson's χ2 test with Yates' continuity correction.

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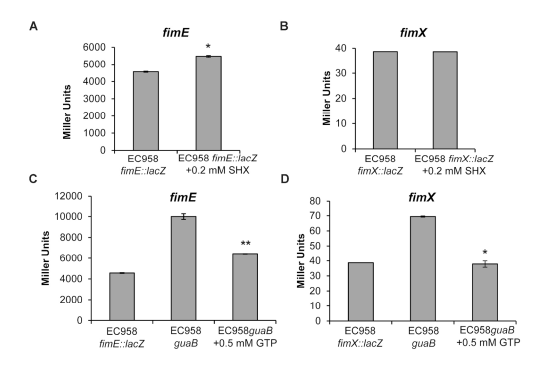


Fig. 9. Regulation of fimE and fimX promoter activity by (p)ppGpp. A-B. Effect of serine hydroxamate (SHX, 0.2 mM) induced amino acid starvation on the promoter activity of fimE (A) and fimX (B) in EC958fimE:lacZ and EC958fimX:lacZ strains, respectively. All guaB mutant strains and complemented derivatives were constructed in their respective promoter-lacZ reporter backgrounds. Strains were incubated overnight under static conditions in pooled human urine and promoter activity was determined by the β -galactosidase assay. Bars represent means \pm SD of three independent biological experiments with four technical replicates. Asterisk indicates a statistically significant difference (P<0.05) as

C-D. Effect of GTP (0.5 mM) on the promoter activity of fimE (C) and fimX (D) in EC958fimE::lacZ guaB and EC958fimX::lacZ guaB strains, respectively. Asterisks indicate statistically significant differences (**P<0.01, *P<0.05) in guaB reporter strains upon addition of GTP, as determined by a repeated-measures one-way ANOVA with Dunnett's multiple comparisons test.

determined by the Wilcoxon matched pairs signed-rank test.

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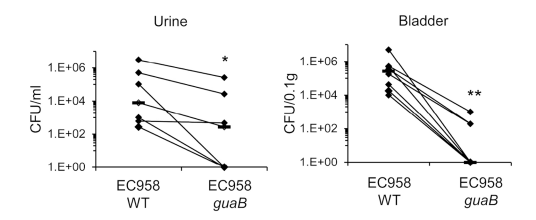


Fig. 10. Mouse urinary tract colonization by EC958 WT and EC958guaB. Female C57B/L6 mice were transurethrally inoculated with a 1:1 mixture of type 1 fimbriae enriched EC958 WT and EC958guaB strains in a competitive infection assay. Each marker represents total colony forming units (CFU) recovered from each mouse per 1 ml of urine or per 0.1 g of bladder tissue (as labelled) on selective medium. Lines connect data points for the same mouse and horizontal bars represent median values. Asterisk indicates statistically significant difference between the strains for persistence in urine (P<0.05) as well as bladder colonization (P<0.01) as determined by the Wilcoxon matched pairs signed-rank test. $150x61mm (300 \times 300 \text{ DPI})$