Elevated urine IL-10 concentrations associate with *Escherichia coli* persistence in older patients susceptible to recurrent urinary tract infections

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**Abstract**

**Background:** Age is a significant risk factor for recurrent urinary tract (rUTI) infections, but the clinical picture is often confused in older patients who also present with asymptomatic bacteriuria (ASB). Yet, how bacteriuria establishes in such patients and the factors underpinning and/or driving symptomatic UTI episodes are still not understood. To explore this further a pilot study was completed in which 30 male and female community based older patients (mean age 75y) presenting clinically with ASB / rUTIs and 15 control volunteers (72y) were recruited and monitored for up to 6 months. During this period symptomatic UTI episodes were recorded and urines collected for urinary cytokine and uropathogenic *Escherichia coli* (UPEC) analyses.

**Results:** Eighty-six per cent of patients carried *E. coli* (10^2 ≥ 10^5 CFU/ml urine) at some point throughout the study and molecular typing identified 26 different *E. coli* strains in total. Analyses of urine samples for ten different cytokines identified substantial patient variability. However, when examined longitudinally the pro-inflammatory markers, IL-1 and IL-8, and the anti-inflammatory markers, IL-5 and IL-10, were significantly different in the patient urines compared to those of the controls (P < 0.0001). Furthermore, analysing the cytokine data of the rUTI susceptible cohort in relation to *E. coli* carriage, showed the mean IL-10 concentration to be significantly elevated (P = 0.04), in patients displaying *E. coli* numbers ≥10^5 CFU/ml.

**Conclusions:** These pilot study data suggest that bacteriuria, characteristic of older rUTI patients, is associated with an immune homeostasis in the urinary tract involving the synthesis and activities of the pro and anti-inflammatory cytokines IL-1, IL-5, IL-8 and IL-10. Data also suggests a role for IL-10 in regulating bacterial persistence.

**Keywords:** Urinary tract infection, *Escherichia coli*, Cytokines, Ageing, Antibiotics

**Background**

Recurrent urinary tract infections (rUTI) are a major clinical problem, particularly in older age groups, impacting significantly on patient well-being and global healthcare systems [1, 2]. Such infections are classified as complicated or uncomplicated, depending on the presence or absence of structural or functional abnormalities of the urinary tract, but are linked to the persistent breaching of the host innate urinary tract defences by uropathogenic bacteria [3]. In treating rUTIs, patients are often prescribed repeated short-term antibiotic treatments or receive long-term, low-dose prophylaxis therapies. Continuous use of these antimicrobial agents has been shown to impact UTI frequency, but at the cost of bacterial resistance, which is a major public health concern [4–6].

Age is a significant risk factor for rUTI [2, 7]. However, the clinical picture is frequently confused by older
patients who present with significant bacteriuria, but without the symptoms or other adverse effects associated with an UTI [2]. This harmless condition is termed asymptomatic bacteriuria (ASB) and it has been suggested that the microbial strains colonising the urinary tract, and associated with bacteriuria evolve from their virulent predecessors [8, 9]. Still, how bacteriuria establishes, the level of host-pathogen communication that occurs and the factors underpinning and/or driving symptomatic UTI episodes are not well understood.

Asymptomatic bacteriuria would not normally be treated with antibiotics [7, 10]. Yet many older patients, particularly those with cognitive impairments where history-taking is clinically challenging, are often subjected to frequent, but needless antibiotic treatment regimens that do not cure or eradicate the bacteriuria, but actually drive microbial resistance [11]. However, the clinical dilemma is considerable because leaving a suspected UTI untreated in such patients may allow the infection to progress resulting in pyelonephritis, septicaemia and in some cases death [2, 12], but if treated unnecessarily can predispose individuals to opportunistic infections such as Clostridium difficile antibiotic induced diarrhoea [11]. This conundrum illustrates the need for further investigations in older patient groups, specifically focussing on host-microbial interactions during periods of bacteriuria or asymptomatic carriage and infection.

Asymptomatic bacteriuria is associated with a range of bacterial species including the Enterobacteriaceae Escherichia coli, Klebsiella pneumoniae and Proteus mirabilis and Gram-positive bacteria including Enterococcus [3, 7]. Of the species isolated from urine E. coli is the most common identified agent with uropathogenic E.coli or UPEC linked to >75% of all reported UTIs [2, 3]. Protection of the lower urinary tract from microbial assault is mediated through innate defence mechanisms that include physical factors such as urine pH, ionic composition and flow. These, in conjunction with innate immune responses characterised by the constitutive or inducible synthesis of urothelial host defence molecules including antimicrobial agents, chemokines and cytokines, function to prevent infection [13]. In healthy individuals these factors work rapidly and collectively to contain, and eliminate uropathogens [13, 14] from the urinary tract, but in older groups, especially those with incomplete bladder emptying, and comparable groups including those afflicted by neural or spinal pathologies, a bacterial presence described as ASB is common.

In establishing diagnostic tools to differentiate ASB from an UTI the focus has been on urinary cytokines and chemokines, which are easily measured and presumed to reflect the immune response of the urinary tract. To date only IL-6, a pro-inflammatory cytokine, has been nominated as a potential biomarker of infection in older patients, but the diagnostic thresholds remain confusing [15–17]. To address this combinations of IL-6 and, for example, leukocyte esterase have been suggested to help improve diagnosis [17]. The key aim of this pilot study was to further understand host-bacteriuria interactions that may help facilitate the development of robust diagnostics that direct appropriate antibiotic treatment regimens. The study was designed to allow the host cytokine response of a mixed population of 30 male and female community-based patients aged 65+ years of age, presenting with a clinical history of uncomplicated rUTIs, to be examined over a 6-month study period. Urine was collected at 14-day intervals, in an unbiased manner and irrespective of UTI status or clinical treatments, with the objectives of exploring host-bacterial, specifically E. coli, interactions and identifying potential bacterial persistence and urine infection biomarkers.

Results
Study population characteristics and baseline data
The target population for this 6-month study was mixed, males and females, aged 65 years plus that were community dwelling. Recurrent UTI patients were selected specifically because of their clinical history of uncomplicated rUTIs, all attended an UTI out-patients clinic at Freeman Hospital, Newcastle upon Tyne, and lack of co-morbidities. The cohort included 23 females and 7 males with an average age of 74.0 and 76.7 years, respectively (Table 1 and Fig. 1a, ANOVA $P = 0.26$). Patients with structural abnormalities were included if clinical records indicated no evidence of a functional defect in the urinary tract impeding bladder voidance. The study group included 11 patients with either vaginal prolapse (Female $n = 5$), enlarged prostate (Male $n = 3$), urethral stenosis ($n = 2$) or phimosis ($n = 1$). Diabetic patients or females using either topical or systemic oestrogen were also not excluded (Table 1).

The control healthy volunteers were selected because they had no clinical history of UTIs over the previous 3 years or longer (Fig. 1a). Five female and 10 male volunteers were recruited with an average age of 68.0 and 73.9 years, respectively. The average age of the group was not significantly different from the rUTI participants (ANOVA $P = 0.16$), although the average age of the female control cohort was younger, statistically, when compared to their rUTI counterparts (ANOVA $P = 0.02$).

Correlation of diagnostic criteria
A combination of criteria are used to establish an UTI diagnosis [18, 19]. These include assessment of clinical symptoms, the use of a dipstick assay, where a urine
coli and counts of E. study focussed on the most common uropathogen, number of different uropathogens can cause UTIs this 
ulations [18, 19]. While it is acknowledged that diagnosis of an UTI as opposed to 
italic{ASB} is challenging and particularly within older pop-

As expected from the inclusion criteria no control vol-

tions reported any symptoms, although one returned 
itive strains were identified as being negative for E. coli were in receipt of antibiotics at some point during the study period (Fig. 2a). In fact, antibiotic treatment amongst all 
ility significant thresholds, but did not elim-

Strain shifts and antibiotic therapy
Microbiological analyses of each urine sample revealed that 86% of the rUTI patients (26/30) had a positive urine E. coli culture, CFU ranging between $10^5$ to $10^8$ CFU/ml, at least once during the study period. Two of the three patients whose urines were identified as being negative for E. coli were in receipt of antibiotics at some point during the study period (Fig. 2a). In fact, antibiotic treatment amongst all 
P. coli loads was common and shown to be benefi-

E. coli is a versatile bacterial species that exhibits a 

Cytokine profiles
To identify potential biomarkers associated with E. coli carriage and infection, the urine concentrations of an
array of pro and anti-inflammatory cytokines, linked to UTIs and including IL-1β, IL-4, IL-5, IL-6, IL-8, IL-10, IL-12, IL-17A, TNFα and IFNγ were measured [17, 24–30]. Quantification of these ten innate effectors in rUTI patient and control urines highlighted substantial variability, but statistical analyses of the longitudinal data supported the mean concentrations of four cytokines, the pro-inflammatory markers, IL-1β and IL-8, and the anti-inflammatory markers, IL-5 and IL-10, to be significantly elevated in the rUTI patient cohort compared to the controls (Fig. 3, *P* < 0.001).

Of the 360 rUTI patient urines analysed, 184 were positive for *E. coli* of which 80 displayed >10^2 < 10^5 CFU/ml, and 104 samples registered ≥10^5 CFU/ml, which, clinically, is defined as ‘significant bacteriuria’ or indicative of an UTI [31]. Comparing urine samples that were negative for *E. coli* to those that harboured *E. coli* at counts of >10^5 CFU/ml revealed no significant differences in the urinary IL-1β, IL-5, IL-6, IL-8 and IL-10 cytokine concentrations (Table 3) although statistically these data did suggest a link between IL-10 and the presence of *E. coli* (*P* = 0.08). However, when the cytokine concentrations of rUTI urine samples negative for *E. coli* were compared to those containing ≥10^5 CFU/ml bacteria a significant increase in IL-10 (6.45 ± 12.26 pg/ml versus 10.57 ± 20.85 pg/ml [*P* = 0.04]) (Fig. 4 and
Table 4), was observed. Presented longitudinally these urine data support significantly elevated IL-10 concentrations in rUTI patients carrying ≥10^5 CFU/ml E. coli loads (Fig. 4a and b). The impact of antibiotic treatment with respect to IL-10 showed no statistical significance (Table 4).

E. coli persistence and infection have been reported to be more common in patients presenting with diabetes and/or a clinical history of bladder/kidney surgery [2]. When rUTI patients with diabetes, taking oestrogen supplements, or a history of vaginal prolapse, or prostate enlargement (n = 19), were excluded from the analyses the IL-10 data relating to E. coli loads ≥10^5 CFU/ml remained significant (P = 0.00002) (Fig. 4c and d, Table 4).

IL-6 has been nominated as a biomarker in older patients that discriminates ASB from symptomatic infection [15–17]. Threshold IL-6 concentrations indicative of an UTI have been proposed as > 25 pg/ml [15] or > 30 pg/ml [17]. Applying a cut-off of > 25 pg/ml the rUTI patient IL-6 urine data identified only 13/360 measurements (Fig. 5) indicative of an UTI although only seven of these values were associated with E. coli counts of ≥10^5 CFU/ml urine. Analyses of the IL-6 urine concentrations in conjunction with either IL-8 concentrations, positive dipstick (leukocyte esterase) or symptomatic data (Fig. 5a – c) revealed trends, but there was no consistency between these three sets of sample data that clarified the ASB / UTI diagnosis. However, it was noted that the two urine samples characterised by elevated IL-6 and IL-8 concentrations (Fig. 5a) also correlated with a positive dipstick outcome and the patient in question informing the study of suspected UTI symptoms.

**Fig. 2** Schematic representation of study data for seven rUTI patients. E. coli loads, antibiotic treatments and symptom reports are defined in the attached key. The numbers in the E. coli boxes represent the sequence types derived from the MLST analysis. a Case examples of patients who received antibiotics and did not present with urinary E. coli loads during the study. b Case examples of patients who did not receive antibiotic treatments but displayed urinary E. coli loads throughout the study. c Case examples of patients on short-term (3–7 day courses) of antibiotics. d Two case examples of patients on prophylactic antibiotics and where consistent E. coli loads were observed.
Discussion

Asymptomatic bacteriuria (ASB) and UTIs are common in older people, yet non-specific symptoms, often compounded by cognitive problems and the lack of good diagnostic tools to discriminate between the two conditions, can compromise the clinical management of such patients [18]. This frequently results in cautious, but unnecessary treatment regimens that achieve little clinically and challenge good antibiotic stewardship. Focussing on an older, yet cognitively sound, mixed-sex community-based population of 65+ years, and existing non-invasive methods of diagnosing an UTI, we similarly found poor concordance between self-reported patient symptoms, dip-stick, urinary cytokine and microbiological measurements. Complicating the diagnosis was the constant presence and variable numbers of bacteria in the urine of these patients. While bacteriuria is well-known to affect many older and indeed younger patient groups there is still a lack of understanding of its pathology, which continues to compromise the ASB / UTI diagnosis [7]. Therefore, to progress diagnosis and treatments for rUTI in older patients the physiological, microbiological and immunological mechanisms underpinning their bacteriuria need to be explored and unravelled.

Taking a more general approach we analysed our six-month patient and control cytokine data longitudinally, which identified the mean concentrations of four urine markers IL-1β, IL-5, IL-8 and IL-10 to be elevated within the ASB / rUTI susceptible patient cohort.

Table 2 E. coli sequence types (ST) isolated and sorted by isolation statistics

| Phylogroup | Sequence Type | Frequency Isolated | Patients |
|------------|---------------|--------------------|----------|
| B1         | 677           | 9                  | 2        |
|            | 3640          | 5                  | 1        |
|            | 442           | 2                  | 1        |
|            | 602           | 1                  | 1        |
|            | 1571          | 1                  | 1        |
| B2         | 73            | 54                 | 9        |
|            | 12            | 15                 | 4        |
|            | 127           | 11                 | 2        |
|            | 420           | 11                 | 1        |
|            | 131           | 10                 | 3        |
|            | 404           | 8                  | 1        |
|            | 144           | 7                  | 1        |
|            | 95            | 6                  | 2        |
|            | 355           | 6                  | 1        |
|            | 91            | 5                  | 1        |
|            | 625           | 2                  | 1        |
|            | 681           | 2                  | 1        |
|            | 80            | 1                  | 1        |
|            | 421           | 1                  | 1        |
|            | 583           | 1                  | 1        |
| D          | 69            | 12                 | 6        |
|            | 362           | 2                  | 1        |
|            | 38            | 1                  | 1        |
| E          | 335           | 1                  | 1        |
| F          | 354           | 9                  | 1        |
|            | 59            | 1                  | 1        |

*Historical phylogroups or clades of the E. coli species utilized across the microbiology community [21]

Isolation frequency based on how many times each sequence type (ST) was identified in the strain collection

Number of patients harbouring specific STs

Table 3 rUTI patient cohort mean ± SD cytokine concentrations for IL-1, IL-5, IL-6, IL-8 and IL-10 with respect to E. coli carriage

| Cytokine | Average Concentration | ANOVA P-value |
|----------|-----------------------|---------------|
| IL-1     | 6.5 ± 14.9            |               |
| IL-5     | 12.4 ± 30.0           | 0.23          |
| IL-6     | 6.4 ± 27.9            | 0.30          |
| IL-8     | 42.9 ± 104.8          | 0.98          |
| IL-10    | 64 ± 12.2             | 0.26          |

*ANOVA P < 0.05
Interestingly, these four molecules encompass two pro-inflammatory cytokines, IL-1β and IL-8, and two anti-inflammatory cytokines, IL-5 and IL-10 (Fig. 3), which suggested the host innate response of this older ASB / rUTI cohort had specifically adapted to tolerate urinary E. coli colonisation. This was further supported by the longitudinal analyses revealing cytokine responses to be maintained during periods of intermittent colonisation which were linked often, but not always, to antibiotic treatment (Table 2). However, a study limitation was that we focussed specifically on E. coli, therefore it was possible that other bacterial species were present and similarly impacting the host innate response.

These findings appear to suggest that in older patients with a history of ASB / rUTIs the urothelial innate defences respond and adapt to the constant microbial challenge by establishing and maintaining an urinary microbiome, defined clinically as ASB [32]. These data also suggest that this urobiome is selected and tolerated through the local production and interactions of specific host pro- and anti-inflammatory cytokines. Essentially the host creates an immune homeostasis in the lower urinary tract that supports bacterial persistence. While it could be argued that this host urobiome is a constant source of potential infection [32, 33], the counter argument is that it protects and plays a role in preventing UTIs, with misguided or precautionary short-term antibiotic use causing a dysbiosis that increases susceptibility to infection. Although our microbiological data was limited specifically to E. coli it was noted that some patients undergoing short-course antimicrobial treatments were characterised by E. coli strain shifts, albeit linked to antibiotic resistance patterns, that clinically could expose them to an increased risk of infection.

This study was designed as a pilot study with a patient cohort of 30. While the study power was justified for a pilot study [34] the small size of the patient and control cohorts was a study limitation, which arguably was further complicated by a significant difference in the average age of the female rUTI and control groups (74 compared to 68 yrs). UTI disease prevalence and pathology differ between males and females therefore comparable but larger single-sexed patient studies are needed to explore whether these IL-1β, IL-5, IL-8 and IL-10 cytokine / ASB observations are impacted by gender.

It is also acknowledged that host genetics plays a role in patient susceptibility to rUTIs [29, 35]. In children and young women (18–49 years) it has been reported that a TLR2_G2258A SNP associates with an increased risk of ASB [29, 36]. Our patient cohort size did not contain the power to examine the effects of host gene polymorphisms on microbial colonisation. However, it was noted that only 2/30 patients carried the TLR2_G2258A SNP (data not shown), suggesting that physiological or molecular events associated with ageing may either impact or supersede those due to genetics. Interestingly, age dependent
differences in uropathogen susceptibility and colonisation have recently been reported in different mice strains used to model UTIs with colonisation characterised by specific cytokine / chemokine profiles [37].

While acknowledging that differences in patient genetics, lifestyle and disease pathology influence susceptibility to UTIs [29, 38, 39] it was interesting that 26 unique *E. coli* STs were identified in the rUTI patient urines with some patients, during the six-month study, carrying more than one ST. The majority of strains isolated associated with one phylogenetic clade, namely B2, but a number of STs from other clades were also identified (Fig. 2 and Table 2). Similar clade distributions of UTI isolates have been reported [40, 41] with different clade members shown to be robust (B1 and B2), deficient (A) or variable colonisers of mice bladders [41]. This study showed that antibiotic use also impacted ST prevalence but did not, however, impact cytokine production (Table 4). However, these data strongly suggest interplay between host and microbial factors, genetic and phenotypic, underpin the establishment and persistence of bacteriuria and the presentation of UTI symptoms [33].

The population targeted during this pilot study was purposely chosen to be mixed sex, aged 65+ years, susceptible to rUTIs, and community dwelling and therefore represented a relatively healthy ageing population. This could be argued as a study limitation with future studies needing also to consider other at-risk groups such as the frail, less mobile or cognitively impaired ageing individuals. A previous study in older patients, also a mix of community based males and females showing susceptibility to rUTIs, identified the urinary markers IL-6 and IL-8, both pro-inflammatory cytokines, to be elevated in bacteriuria, and significantly (4-fold) increased during acute cystitis [17]. Longitudinally the mean IL-6 urine concentration of our study participants was comparable to that of the controls (ANOVA $P = 0.123$), suggesting that this cytokine did not reflect either bacteriuria or rUTIs. However, it is acknowledged that statistical analysis of the longitudinal data may have masked elevated IL-6 values, seen as a cluster of values in Fig. 3, which reflected genuine UTIs. Further analyses (Fig. 5), did support cytokine fluctuations, but no consistent cytokine patterns were identified that specifically diagnosed an UTI. Microbiologically these urine measurements were not always associated with increased *E. coli* numbers although again it is recognised that the study focussed specifically on the association of *E. coli* with ASB although ASB can encompass a mix of organisms [7].

It was of particular interest that microbial colonisation and significant urinary *E. coli* loads in the ASB /

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**Table 4** Mean ± SD IL-10 concentrations with respect to antibiotic treatment and *E. coli* carriage

| Condition | Average IL-10 Concentration (pg/ml) | ANOVA P-values |
|-----------|------------------------------------|----------------|
| All rUTI Patients ($n = 30$) | | |
| No Antibiotics | 7.7 ± 15.4 | 0.85 |
| Antibiotics | 8.0 ± 13.6 | 0.56 |
| *E. coli* | 6.4 ± 12.2 | 0.21* |
| < $10^5$ CFU/ml | 7.3 ± 10.1 | |
| ≥ $10^5$ CFU/ml | 10.5 ± 20.8 | 0.04* |
| Patients without complicated urinary tract history* ($n = 11$) | | |
| *E. coli* | 3.6 ± 8.2 | 0.3 |
| < $10^5$ CFU/ml | 6.0 ± 10.3 | 0.05* |
| ≥ $10^5$ CFU/ml | 19.3 ± 27.5 | 0.00002* |

* P-value stated is for No *E. coli* versus ≥ $10^5$ CFU/ml data sets

*aUTI Patients recruited to study who were not diabetic, taking oestrogen supplements, or a previous clinical history of vaginal prolapse / prostate enlargement

*p-value stated is for < $10^5$ CFU/ml versus ≥ $10^5$ CFU/ml data sets

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![Fig. 5](image-url) Urine concentrations of IL-6 in relation to a IL-8, b dipstick outcome and c self-declared symptoms. The highlighted region in (a) represents the required threshold for an acute UTI diagnosis based on IL-6 (> 25 pg/ml) and IL-8 (> 2000 pg/ml) as defined by Sunden et al. (2017). Both points shown in (a) are associated with patient UTI337 shown in Fig. 2c.
rUTI susceptible cohort (>10⁵ CFU/ml), were marked by the synthesis of the immunomodulatory cytokine IL-10 [42, 43]. Observations in humans and in IL-10 deficient mice, support a key role for IL-10 in defending the urinary tract from UPEC infection [43, 44]. Moreover recent studies exploring UT microbial colonisation in older female mice have specifically identified IL-10 as a significant factor in their susceptibility to colonisation by E. coli [37]. The cytokine IL-10 is known to function in the host immune response to infectious disease [42, 43] and shown to over-ride the host inflammatory responses to infection, meaning it is closely associated with microbial persistence [45]. In the bladder markedly high IL-10 concentrations have been proposed to promote an immunosuppressive environment, which function to dampen any auto-immune responses [46] and allow the prompt regeneration of damaged epithelia [46]. Interestingly, IL-10 induction has been associated with the early phases of an infection [43] whereas our data links elevated IL-10 levels to long-term colonisation / persistence. Further evidence, albeit from young mouse UTI models, has suggested that IL-10 is synthesised primarily by migrating mast cells (MC) [27, 46] with bacterial persistence associated with elevated bladder MC numbers [46]. At present how IL-10 functions in the susceptibility and / or protection of these older patients in relation to urinary tract microbial colonisation is not known, but data does suggest a key role for the innate host response involving IL-10 in promoting bacterial persistence and potentially tolerance.

**Conclusion**

Our longitudinal cytokine data from a pilot study involving 65+ year old male and female patients susceptible to ASB / rUTIs suggest that the urinary tract innate response of this ageing population has adapted to create an immune homeostasis with the IL-1β, IL-5, IL-8 and IL-10 cytokine profiles, playing a key role in the pathology of bacterial colonisation and persistence. Further patient studies are needed to confirm these observations, understand the local immune changes, particularly those involving IL-10 that underpin ABU and accompany the progression to UTI, and the impacts of therapies including antibiotics.

**Methods**

**Study design**

Patients were recruited with written informed consent through UTI clinics led by the Urology Department at Freeman Hospital, Newcastle upon Tyne, UK. Inclusion criteria for patients aged 65 or over with rUTI were: 1) two or more symptomatic UTIs within the last 6 months or three or more within 12 months; 2) UTIs supported clinically by urine samples positive for bacteria; 3) each diagnosed UTI associated with antibiotic treatment. Males and females were eligible, but exclusion criteria included urinary tract reconstruction, history of bladder cancer, presence of indwelling catheter or need for intermittent self-catheterisation, all of which related to a complicated UTI. Each patient in the rUTI cohort was studied for 6 months and agreed to supply a mid-stream catch urine sample every 14 days (12 samples/patient). With each urine sample the patients provided answers to a series of previously validated questions relating to their UTI status [20]. No clinical decisions were made by the research team. All rUTI patients were encouraged to seek assessment from their regular health care provider hence no urines were provided at the time points when participants approached their health-care provider in relation to an UTI. Instead clinical records were used to record the incidence of an UTI +/- 3 days of a study sample being provided and any UTI specific antibiotics prescribed.

An ethical amendment was approved in March 2016 to recruit control volunteers aged 65 years or over, with no previous history of rUTI. Male and females were enrolled utilising the volunteer network VoiceNorth (http://www.voice-global.org/) and supplied three mid-stream catch urine samples at 14-day intervals. Inclusion criteria stipulated no UTI history for the preceding 3 years which participants confirmed verbally. Screening of all urine samples, from rUTI and control cohorts, was performed in an unbiased manner. Study data was only analysed when all the urine samples had been collected and analysed.

**Analyses of urine**

Urine samples from patients and controls were analysed within 4 h of collection for nitrite and leukocyte esterase using Multistix® 10SG (Siemens) according to the manufacturer’s instructions. Urines with positive nitrite and leukocyte results, were regarded as having urinalysis suggestive of an UTI. Semi-quantitative urine culture was performed according to current UK National Standard Methods of Investigation, using both 1µl and 10µl aliquots. Urine samples were plated onto CPS ID 3 (CPS3) or CPS Elite chromogenic agar plates (bioMérieux), incubated at 37°C, in room air for 18–24 h. The presence of presumptive E. coli was noted, and the bacterial counts enumerated. The remaining urine was filtered using a 0.45 µm filter (GE Healthcare Life Sciences), and aliquots stored at ~80°C for cytokine analyses.

Single colony isolates of E. coli were typed using a multi-locus sequence typing scheme [23]. Genomic DNA was isolated using the standard procedure from the Bacterial Genome Kit (Sigma). PCR products
generated using published primers, at the recommended temperatures, were purified using commercial PCR Purification kits (Sigma) prior to sequencing (Source-Bioscience). Sequence results were processed using the pubmlst.org website by choosing the Achtman database to identify the allele number of each gene and the corresponding sequence type.

ELISA based cytokine detection
Ready-Set-Go® ELISA kits (Affymetrix eBioscience) were used for the cytokine assays, which avoided frequent freeze-thawing of urine samples. All ELISAs were completed in Nunc MaxiSorp® flat-bottom 96 well plates (Affymetrix eBioscience) and the manufacturer’s protocol was followed for each of the kits. Plates were quantified at 450 and 571 nm using an Infinite® F50 / Robotic Absorbance Microplate Reader (Tecan Trading AG, Switzerland).

Statistical analysis and data presentation
Figure generation and analysis for statistical significance was performed using a combination of R, Microsoft Excel, StatPlus and Adobe Illustrator. Statistical significance throughout this study was determined using ANOVA (Analysis of Variance). \( P \)-values quoted represent the actual \( P \)-value of the returned \( F \)-value generated by ANOVA [47].

Abbreviations
ANOVA: Analysis of Variance; ASB: Asymptomatic bacteriuria; CFU: Colony forming units; ELISA: Enzyme linked immunosorbent Assay; MC: Mast cells; PCR: Polymerase chain reaction; rUTI: recurrent UTI; UPEC: Uropathogenic Escherichia coli; UT: Urinary tract; rUTI: Urinary tract infections

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Authors’ contributions
RP, JH, AA and PA conceived the study, LD, WR, CM, JP, CH were involved in patient recruitment and sample management, LD, JH and PA collated and analysed the data. LD, CM, JH and PA prepared the presentation of data in the figures. LD, JH and PA drafted the manuscript, which was critically reviewed and edited by WR, CM, JP, AA, CH, KW and RP to develop the final version. All authors read and approved the final manuscript.

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Availability of data and materials
The datasets generated and/or analysed during the current study are not publicly available due to privacy reasons but are available in anonymized form from the authors on reasonable request.

Ethics approval and consent to participate
The study protocol was approved in March 2014 by the Newcastle Research Ethics Committee (ref: REC-14-NE-0026) and sample collections took place between April 2014 and November 2015. Patients were recruited with written informed consent through UTI clinics led by the Urology Department at Freeman Hospital, Newcastle upon Tyne, UK.

Consent for publication
Not applicable.

Competing interests
LD, WR, CM, JH, PA declare no conflicts of interest. CH declares advisory/consultancy roles for Pierre Fabre, AMS and Astellas and speaker roles for Astellas, Pfizer, Ferring, Allergan and Medtronic.

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