Genomic analysis of trimethoprim-resistant extraintestinal pathogenic Escherichia coli and recurrent urinary tract infections

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Abstract
Urinary tract infections (UTIs) are the most common bacterial infections requiring medical attention and a leading justification for antibiotic prescription. Trimethoprim is prescribed empirically for uncomplicated cases. UTIs are primarily caused by extraintestinal pathogenic Escherichia coli (ExPEC) and ExPEC strains play a central role in disseminating antimicrobial-resistance genes worldwide. Here, we describe the whole-genome sequences of trimethoprim-resistant ExPEC and/or ExPEC from recurrent UTIs (67 in total) from patients attending a regional Australian hospital from 2006 to 2008. Twenty-three sequence types (STs) were observed, with ST131 predominating (28%), then ST69 and ST73 (both 7%). Co-occurrence of trimethoprim-resistance genes with genes conferring resistance to extended-spectrum β-lactams, heavy metals and quaternary ammonium ions was a feature of the ExPEC described here. Seven trimethoprim-resistance genes were identified, most commonly dfrA17 (38%) and dfrA12 (18%). An uncommon dfrB4 variant was also observed. Two bla<sub>CTX-M</sub> variants were identified – bla<sub>CTX-M-15</sub> (16%) and bla<sub>CTX-M-14</sub> (10%). The former was always associated with dfrA12, the latter with dfrA17, and all bla<sub>CTX-M</sub> genes co-occurred with chromate-resistance gene chrA. Eighteen class 1 integron structures were characterized, and chrA featured in eight structures; dfrA genes featured in seventeen. ST131 H30Rx isolates possessed distinct antimicrobial gene profiles comprising aac(3)-Ia, aac(6)-Ib-cr, aph(3’)-Ia, aadA2, bla<sub>CTX-M-15</sub>, bla<sub>SUL1</sub>, and dfrA12. The most common virulence-associated genes (VAGs) were fimH, fyuA, irp2 and sitA (all 91%). Virulence profile clustering showed ST131 H30 isolates carried similar VAGs to ST73, ST405, ST550 and ST1193 isolates. The sole ST131 H27 isolate carried molecular predictors of enteroaggregative E. coli/ExPEC hybrid strains (aatA, aggR, fyuA). Seven isolates (10%) carried VAGs suggesting ColV plasmid carriage. Finally, SNP analysis of serial UTI patients experiencing worsening sequelae demonstrated a high proportion of point mutations in virulence factors.

DATA SUMMARY
The 67 draft genomes from the whole-genome sequencing described here have been submitted to the National Center for Biotechnology Information with the accession numbers SAMN14547713 to SAMN14547779, under BioProject PRJNA623470.

INTRODUCTION
Urinary tract infections (UTIs) are the most common bacterial infections to require medical attention and incur an estimated economic burden of ~$2 billion per annum (~£1.51 billion; $1=£0.76) in the USA alone [1]. In the UK, in 2013–2014, the National Health Service spent £434 million in treating 184000 patients with unplanned admissions relating to UTIs [2]. In Australia, UTIs caused 69823 hospitalizations in 2017–2018, resulting in 234455 inpatient days and treatment costs as high as $AU6400 (£3500) per patient [3, 4]. Additionally, UTIs are the leading rationale behind antibiotic prescription by general practitioners [5]. The most common aetiological agent of UTIs are extraintestinal pathogenic Escherichia coli (ExPEC), responsible for ~75–95% of cases [6]. Epidemiological studies utilizing multi-locus sequence typing (MLST) indicate that certain pandemic
ExPEC can enter the intestinal tract via contact with poultry, companion animals, the environment and through sexual contact [9]. Additionally, travellers are at higher risk of ingesting ExPEC strains [10]. The subsequent introduction of ExPEC into the urinary tract via the urethra can occur due to host behaviours, physiological abnormalities or medical interventions, such as catheterization [6]. After contaminating the urethra, ExPEC can ascend to infect the bladder (cystitis), the kidney (pyelonephritis) or enter the bloodstream (urosepsis) with potentially dire outcomes as ExPEC-associated sepsis has a mortality rate of up to 30% [11]. Indeed, each year at least 1.7 million adults in the USA develop sepsis, and 1 in 3 patients who die in hospital have sepsis [12, 13].

ExPEC possess an array of virulence-associated genes (VAGs) that enable ascension, colonization and persistence within a hostile and nutrient-deficient environment. These VAGs – commonly located on chromosomal pathogenicity islands (PAIs) or encoded on virulence plasmids – include uropathelial adhesins, such as P- fimbriae (pap genes), S- fimbriae (sfa genes), F1C-fimbriae (foc genes), and Dr adhesins (dra genes, afu genes), toxins including secreted autotransporter toxin (sat) and cytotoxic necrotizing factor 1 (cnf1), capsule (kps M II), and several iron-acquisition systems, such as aerobactin (iuc genes, iutA), salmochellin (iro genes) and yersiniabactin (ybt genes, irp1, irp2, fyuA) [14, 15]. ExPEC have also been associated with a greater abundance of bacteriocins, toxins that inhibit the growth of other E. coli [16]. These bacteriocins are frequently carried on virulence plasmids, such as CoIV plasmids [17]. While no singular VAG can be attributed to ExPEC pathogenesis, a molecular definition of ExPEC has been proposed as being E. coli that contain at least two of the following VAGs: papA and/or papC, sfa/foc, afu/draBC, kps M II and iutA [18]. Furthermore, the presence of certain VAGs have been associated with specific uro-clinical syndromes, such as P- and S-fimbriae, Dr adhesins, and the invasin iheA with pyelonephritis [6], and yersiniabactin irp2 and fyuA with sepsis [19]. FyuA is also directly linked to ExPEC biofilm formation in human urine [20], as is the salmochelin receptor iroN [21].

In Australia, severe pyelonephritis and sepsis are typically treated with intravenous gentamicin and amoxicillin, while cystitis is typically treated empirically with either nitrofurantoin or trimethoprim [22]. However, a 5 year study on antimicrobial resistance (AMR) in urinary E. coli from an Australian metropolitan hospital found significant increases in trimethoprim, nitrofurantoin, amoxicillin and gentamicin resistance, as well as significant extended-spectrum β-lactam resistance [23]. ExPEC, particularly ST131 strains, have been central to a worldwide increase in extended-spectrum β-lactamase (ESBL) -producing E. coli. The ST131 lineage consists of three major clades, each of which is strongly associated with specific fimH alleles. Strains from clade C typically carry fimH30 and are currently the most prominent subtype identified in human infections [24]. Clade C is further divided into two subclades – C1 and C2. C1 mostly comprises strains with gyrA and parC SNPs conferring fluoroquinolone resistance, designated H30R, while the latter consists of strains with both fluoroquinolone resistance SNPs and carriage of plasmid-associated bla CTX-M genes, designated H30Rx [25–27]. E. coli ST131 first emerged in the literature in the early 2000s and has been linked to a 300% increase in USA hospital admissions due to ESBL-producing E. coli in the following decade [28, 29]. The global rapid increase in ESBL-producing E. coli has had serious knock-on effects, driving carbapenem prescription and in turn promoting the spread of potentially untreatable carbapenemase-producing E. coli [30].

Like all E. coli, ExPEC have highly flexible genomes and a proclivity to capture and disseminate genes through horizontal gene transfer [31]. Horizontal gene transfer can occur via plasmid conjugation, phage transduction and via small mobile genetic elements (MGEs), such as transposons, insertion sequences (IS) and integrons, allowing for inter- and intra-species movement of genetic information [32]. As observed with the global dissemination of bla CTX-M – a single MGE capture event can have worldwide repercussions [30].

AMR surveillance of ExPEC populations within high selective pressure environments, such as hospitals, could provide meaningful insights and inform future policy making and practices. In addition to the aforementioned 5 year study on AMR within ExPEC isolated in an Australian hospital, a
recent Australian government report also stated that *E. coli* resistances to ESBLs and fluoroquinolones are climbing [4]. While these reports highlight national trends, they tend to focus on hospitals situated in metropolitan areas, while more remote and rural areas get neglected. Furthermore, genomic surveillance of antimicrobial genes (ARGs) and VAGs within Australian hospital *E. coli* populations is currently limited.

Here, we used whole-genome sequencing (WGS) to characterize 67 ExPEC strains from patients with UTIs collected in a large Western New South Wales (NSW) hospital catering to regional, remote and rural Australian communities. The genomes were typed using Clermont phylogrouping, e-serotyping and MLST, and screened for the presence of ARGs, VAGs, PAIs and plasmid replications, and several AMR associated class 1 and class 2 integrons found in this collection were characterized. Furthermore, isolates from recurrent infections, including those with a subsequently more severe clinical sequelae, were interrogated for SNPs leading to point mutations.

**METHODS**

**Sample collection and selection criteria for WGS**

Urine specimens were collected by clinical staff members of participating health-care centres using a standardized protocol. Semi-quantitative culture was performed on horse blood, MacConkey and chromogenic agars, followed by conventional identification. Isolates were stored in 50% (v/v) glycerol in trypticase soy broth at −70 °C.

**Phenotypic resistance testing**

The isolates were tested for susceptibility to 14 antibiotics as per the disc diffusion method specified by the Clinical and Laboratory Standards Institute (CLSI) [33], using Neo-Sensitabs discs (Rosco). The antibiotics tested were (disc content) amikacin (30 µg), amoxicillin–clavulanate (60 µg), ampicillin (25 µg), ceftazidime (30 µg), ceftriaxone (30 µg), cephalexin (30 µg), ciprofloxacin (10 µg), gentamicin (10 µg), imipenem (10 µg), nalidixic acid (30 µg), nitrofurantoin (300 µg), norfloxacin (10 µg), tetracycline (30 µg) and trimethoprim–sulfamethoxazole (5 µg). The double-disc diffusion test was used to detect the production of ESBLs [34].

**DNA isolation and WGS**

DNA extraction and WGS were performed as described previously [35]. Briefly, DNA was extracted using the ISOLATE II genomic DNA kit (Bioline) and stored at −20 °C. Short-read sequencing was performed using an Illumina HiSeq 2500 v4 sequencer in rapid PE150 mode.

**Genome assemblies**

Raw reads were used to assemble draft genome sequences via Shovill software using default settings (https://github.com/tseemann/shovill). Assemblies underwent quality control using assembly-stats software (https://github.com/sanger-pathogens/assembly-stats). Assembly statistics for this collection are available in Data S1 (available with the online version of this article).

**Phylogenetic analysis**

A maximum-likelihood phylogenetic tree of this collection was reconstructed using IQ-TREE 2 (http://www.iqtree.org/) [36] and a SNP-based phylogenetic tree was reconstructed using snplord (https://github.com/maxcummins/pipelord/snplord/tree/master/snplord), an automated pipeline that utilizes snippy (https://github.com/tseemann/snippy), Gubbins [37] (https://github.com/sanger-pathogens/gubbins) and SNP-sites (https://github.com/sanger-pathogens/snp-sites). The SNPs-based tree of the entire collection was built using 76.0% (3525942 bp/4639675 bp) of the reference genome K12 MG1655 (GCA_000005845.2) and consisted of 35038 SNPs (141270 SNPs before recombination filtering). Trimethoprim-sensitive ExPEC isolates ERR434278, ERR434751, ERR434270, ERR434273 and ERR434271 [38], and trimethoprim-sensitive enteroaggregative *E. coli* (EAEC) isolates SRR5470250, SRR5024242 and SRR3574247 [39], all representing the major STs in this collection, were added to both the phylogenetic analysis and gene screening as controls. An additional SNP-based ST131 only tree was built using 92.03% (4831235 bp/5249449 bp) of reference genome ST131 EC958 (GCA_000285655.3) to identify SNP sites. Recombination filtering reduced 7460 SNPs to 386 SNPs. The ST131 SNP-based tree was reconstructed using IQ-TREE 2 and a recombination filtered alignment of 478 bp produced by Gubbins. All trees were visualized using the Interactive Tree of Life (iTOL) v4 online-based software [40] (https://itol.embl.de/). The ExPEC pangenome was calculated using Roary v3.11.2 [41] and visualized using Phandango [42]. Clonal samples from serial patients were removed from this analysis (total of 57 isolates used) to prevent deflation of the core genome.

**Serial patient SNP analysis**

The snplord pipeline was run on isolates originating from a single patient using the isolate with the earliest isolation date as a reference in each instance. SNPs called by this pipeline were checked manually in .gbk files generated by Prokka [43] using SnapGene. Gubbins output was used to determine which SNPs were the result of homologous recombination.

**Gene screening**

Isolate STs, serogroups and phylogroups were determined *in silico* using MLST v2.0 [44] (https://cge.cbs.dtu.dk/services/MLST/), SerotypeFinder v2.0 [45] (https://cge.cbs.dtu.dk/services/SerotypeFinder/) and ClermontTyping [46] (http://enterobase.warwick.ac.uk/), respectively. ST131 isolates additionally underwent FimH typing using FimTyper [47] (https://cge.cbs.dtu.dk/services/FimTyper/). The Ariba read-mapping tool [48] was used the screen for ARGs, plasmid replications and VAGs using the following reference databases: ResFinder [49] (https://cge.cbs.dtu.dk/services/ResFinder/), PlasmidFinder [50] (https://cge.cbs.dtu.dk/services/PlasmidFinder/) and VirulenceFinder [51] (https://cge.cbs.dtu.dk/services/...
VirulenceFinder/). However, as VirulenceFinder screens for only one \( \text{papA} \) allele, an additional BLASTN search was also performed using 14 \( \text{papA} \) alleles [52]. An additional custom database with AMR-associated ISs, class 1 and 2 integrases, and additional ExPEC-associated VAGs was also utilized and can be accessed at https://github.com/CJREID/custom_DBs. ARIBA data were processed using a bespoke script accessible at https://github.com/maxlcummins/pipelord/tree/master/aribalord and visualized using the following R packages: ggplot2 v3.3.0 (https://github.com/tidyverse/ggplot2) and ggtree v2.2.1 [53] (https://github.com/YuLab-SMU/ggtree).

To infer ColV plasmid and PAI carriage, short reads from each isolate were mapped to the ColV reference plasmid pCERC4 (KU578032), PAI I \(_{\text{CFT073}}\) (AE014075.1, start 3462265 bp, end 3469205 bp), PAI II \(_{\text{AM}}\) (AJ494981), PAI III \(_{\text{AM}}\) (AF301153) and PAI IV \(_{\text{HPI}}\) (high pathogenicity island) using the Burrows–Wheeler Aligner (bwa) v0.7.17 [54] and converted to BAM file format using SAMtools v0.1.18 [55]. A bespoke script, available at https://github.com/maxlcummins/pipelord/tree/master/plasmidlord, was used to produce a histogram of read-depth as a function of reference coordinate and clustered based on their Euclidean distances, and used to generate a heatmap.

BLASTN [56] was used to determine whether integrons characterized in this collection are present in other genomes available in public databases, and associated metadata were pulled from GenBank [56] and Enterobase [57]. Integron B structure comparison was achieved using EasyFig [58]. All gene schematics were generated using SnapGene (www.snapgene.com/).

**Statistical analysis**

R software v4.0.2 was used for the reconstruction of correlation heat maps and classical (metric) multidimensional scaling (MDS) analysis, using the ggplot2 v3.3.0 (https://ggplot2.tidyverse.org/) package for visualization, and the ggcorrplot v0.1.3 (https://github.com/kassambara/ggcorrplot) package for correlation calculation. Two MDS analyses were performed, one for identified virulence genes and one for AMR genes, using the standard R package functions dist and cmdscale in conjunction with a presence/absence matrix (with one for presence and zero for absence).

**RESULTS AND DISCUSSION**

Genomic surveillance using WGS can provide high-resolution data on ARG presence, persistence, evolution and potential for horizontal transfer [59], as well as inform on population diversity and VAG carriage, and provide in-depth analyses of SNP-mediated mutations. Yet to date, genomic analyses on UTI-associated ExPEC within Australian hospitals has been limited. Here, we provide a retrospective WGS study on ExPEC resistant to first-line trimethoprim treatment and/or isolates collected from serial UTIs between 2006 to 2008 from a regional hospital in Western NSW, Australia. We aimed to provide a baseline for future studies tracking the evolution of ExPEC AMR and virulence potential in Australian UTI-associated ExPEC populations by reporting on ARG, VAG and MGE carriage, characterizing 18 class 1 integrons structures identified in this collection and identifying SNPs occurring in recurrent UTI isolates.

**Demographic and clinical characteristics**

From clinical samples collected between 2006 and 2008, 76 isolates were selected for genome sequencing due to phenotypic trimethoprim resistance and/or originating from a serial UTI patient. After quality-control measures, the final collection comprised 67 \( \text{E. coli} \) draft genomes (mean contig size of 23477 bp, with a mean number of contigs of 234) originating from 37 samples taken from patients with cystitis (55%), 23 samples from patients with pyelonephritis (34%) and 7 samples from patients with urosepsis (10%) (Fig. 1a). Recurrent UTIs are common, particularly in women, with 27% reporting a recurrence within 6 months [53]. In this

![Fig. 1. Sample population and clinical characteristics. (a) Diagnoses of patients presenting with UTIs over the study period. For the patients with multiple samples taken for the same diagnosis (cross-hatched stripes): for cystitis, the mean number of days between collections was 126 days; for pyelonephritis, 63 days; and for urosepsis, 68 days. For patients with a previous cystitis diagnosis and subsequent pyelonephritis or urosepsis diagnoses (sloped stripes): the mean length between collections was 174 days, and 181 days for urosepsis (n=1). (b) Age and biological sex of patients. Females are represented by grey, males by shading.](image-url)
collection, 12 out of 51 patients were sampled more than once due to multiple, or ongoing, UTIs during the study period. Women are more likely to contract UTIs than men at a ratio of 8:1, and one in three women will experience at least one UTI necessitating antibiotics by age 24 [60]. The risk of UTIs tends to increase with age in both sexes [61]; however, the prevalence of UTIs in women over the age of 65 is double that of the overall population [62]. This collection is reflective of UTI epidemiology with more isolates derived from female patients (44 vs 7; 86 vs 14%) and with patients’ ages ranging from 13 months to 91 years, the majority (64%) being above 66 years. The mean age for female patients was 66 years, and 74 years for male patients (Fig. 1b). No patient was identified as pregnant.

**Phylogenetic analysis**

While ExPEC are phylogenetically diverse, a recent systematic review and meta-analysis of 217 studies flagged a handful of globally dominant STs, the most prominent being ST131 (phylogroup B2), followed by ST69 (D), ST10 (A), ST405 (D), ST38 (D), ST95 (B2), ST648 (D) and ST73 (B2) [7]. Consequently, B2 and D constitute the phylogroups most frequently associated with ExPEC infections. To ascertain the evolutionary relationships between samples in this collection, a maximum-likelihood phylogenetic tree was reconstructed using the complete genome of E. coli strain K12-MG1655 as a reference (Fig. 2a). The tree demonstrates clear clustering based on phylogroup and ST (Fig. 2a). The collection comprised 23 STs, and despite selecting for trimethoprim resistance, the collection followed global trends with ST131 (O25:H4) being the most common (n=19; 28%), followed by ST69 (ONT:H18 and O11:H18) and ST73 (O22:H1, O25:H1, O6:H1, and ONT:H1) with five isolates each. Isolates from phylogroup B2 were the most prevalent (n=38; 56%), and the least prevalent were from B1 (n=3; 4%).

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**Fig. 2.** Maximum-likelihood phylogenetic trees showing genetic relatedness of ExPEC strains. Tree scale bars represent number of substitutions per site of alignment. (a) Mid-point rooted maximum-likelihood phylogenetic tree, inferred using iq-TREE 2 and K12-MG1655 as a reference, containing the 67 ExPEC isolates sequenced in this study. Coloured circles represent uro-clinical syndrome (black, cystitis; white, pyelonephritis; red, urosepsis). The inner-most ring represents the phylogroup (A, red; B1, green; B2, light blue; D, purple). The next ring represents ST. The two outermost rings apply to ST131 isolates only (marked in red shaded area), the inner ring highlights fimH alleles (orange, fimH30; green, H27) and outer ring shows the presence or absence of blaCTX-M-15 (blue, present; yellow, absent). Letters around the perimeter represent multiple isolates taken from a single patient, one letter per patient. (b) SNP-based phylogenetic tree, inferred using iq-TREE 2, resolving ST131 isolates into clades. H30R, blue shaded area; H30Rx, green shaded area. Trimethoprim-susceptible ExPEC isolates ERR434278, ERR434751, ERR434270, ERR434273 and ERR434271, and trimethoprim-susceptible EAEC isolates SRR5470250, SRR5024242 and SRR3574247 were added as controls.
The fact that ST131 isolates, which are associated with ESBL resistance, outnumbered ST69 isolates, which are associated with trimethoprim resistance [63], is reflective of the high number of UTIs caused by ST131, particularly in association with pyelonephritis, within this regional NSW community [64–66]. To resolve the ST131 isolates into clades, we screened for \textit{fimH} alleles in conjunction with a separate SNP-based phylogenetic tree (Fig. 2b). Only one ST131 isolate, HOS40, carried H27; thus, it belonged to clade B, a clade associated with animal to human transmission [67]. All other ST131 isolates carried H30, thereby belonging to the globally dominant clade C [24]. Of the ST131 H30 isolates, 7 (37%) were H30R (clade C1) and 11 (61%) were ESBL \textit{bla}_{CTX-M-15}^- associated H30Rx (clade C2) [25]. SNP analyses showed that the H27 isolate differed from H30 by a mean of 277 SNPs, and that H30R isolates differed from H30Rx by a mean of 93 SNPs. The H30Rx isolates originated from different patients, situated in seven different postcodes (~300 km between the two most distant postcodes), caused either cystitis or pyelonephritis, and were isolated over a period of 569 days. Yet despite these differences, the H30Rx isolates differed by only seven SNPs on average (across 99.6% of the reference H30Rx HOS54 genome), indicating a persistent clone in this community. Conversely, H30R isolates differed by a mean of 58 SNPs, due to the presence of two distinct branches (Fig. 2b; i and ii), and the SNP difference between H30R and H30Rx branches was 140. The pangenome for this collection consisted of 14091 genes making up a core genome of 2886 genes and an accessory genome of 11205 genes, leading to a total pangenome length equal to 11.32 Mb, and can be viewed in Data S2.

\textbf{Phenotypic resistance}

ExPEC have been a driving force behind a worldwide increase in ESBL-producing \textit{Enterobacteriaceae} [68]. Furthermore, ExPEC isolates have been reported as carrying transmittable resistances to carbapenem and colistin [69, 70], both being last-line antibiotics for multidrug-resistant (MDR) Gram-negative bacteria. The World Health Organization (WHO) has flagged increasing cases of MDR bacteria worldwide as one of the most serious public-health threats and deemed surveillance as a strategic priority of the Global Action Plan on AMR [71]. Despite trimethoprim remaining a first-line empirical treatment option for UTIs, resistance in uropathogens is increasing worldwide. In Australia, resistance to trimethoprim in \textit{E. coli} has increased from 16.6% in 2004 [72] to 31.2% in 2017 [73]. These figures and trends are similar to those in other developed countries, though lower than in developing countries where trimethoprim resistance in UTI-associated ExPEC ranges from 54 to 82% [74]. Isolates were tested against 11 antibiotics (Fig. 3). The mean number of antibiotics an isolate was resistant to was five, with two isolates (HOS56 and HOS70, both ST131 H30Rx) being resistant to 10/11 tested (all bar the carbapenem imipenem). The most common resistance phenotype, aside from trimethoprim

![Figure 3](image-url)
and trimethoprim/sulphonamide, was ampicillin \((n=50; 75\%)\). High levels of ampicillin resistance in UTIs have been described worldwide, leading to governing bodies recommending its disuse [4]. The least common phenotypic resistances were to imipenem \((n=2; 3\%)\) and the anaerobic DNA inhibitor nitrofurantoin \((n=4; 6\%)\). This hopeful observation indicated that both a first and a last-line treatment option for UTIs remained largely efficacious, at least in the past, though low levels of nitrofurantoin and imipenem resistance continue to be reported in UTI isolates currently [75, 76]. Antibiotic resistance varied by ST (Fig. 3a) and uro-clinical syndrome. Isolates originating from patients with cystitis and pyelonephritis shared similar phenotypic resistance profiles, however, isolates from urosepsis patients tended to be more sensitive and no isolate was resistant to the aminoglycoside gentamicin, the cephalosporin ceftoxime, nitrofurantoin or imipenem (Fig. 3b). A high co-occurrence (>30%) of trimethoprim and ESBL resistance was recently reported in the USA [77]. Furthermore, Mulder et al. [78] demonstrated that previous use of extended-spectrum \(\beta\)-lactams in patients with UTIs was significantly associated with trimethoprim resistance. These data are reflective of treatment practices and suggests a step-wise acquisition of resistances to antimicrobials over time, having become resistant to first-line treatment and then increasingly to ESBLs.

**Genetic screening**

This collection was screened for ARGs, ISs associated with AMR, class 1 and 2 integrases \((\text{intI}1, \text{intI}2)\), heavy-metal-resistance genes, VAGs and plasmid replicons using ariba (Fig. 4). A full list of identified genes and associated metadata for each isolate is available in Data S3.

**ARGs**

AMR in uropathogens complicates the treatment of UTIs. An advantage of WGS in AMR surveillance is the level of detail and precision gleaned from identifying specific ARG alleles, and co-resistances not tested on standard antibiotic panels, including disinfectant and heavy-metal-resistance genes [59]. Here, we identified a total of 40 ARGs (Fig. 4), with the majority of isolates \((n=56; 84\%)\) carrying at least three ARGs.

The most common mechanism for acquiring trimethoprim resistance is through the acquisition of \(dfr\) genes [79]. There are currently more than 40 identified \(dfr\) variants [80]. These are often associated with MGEs, such as plasmids and transposons, and are almost exclusively observed as resistance gene cassettes within the variable regions of class 1 and 2 integrons in human [81], animal [35, 82, 83] and environmental [84] \textit{E. coli} isolates, resulting in their rapid spread. Seven trimethoprim-resistance genes were identified in this collection, the most common being \(dfrA17\) \((n=26; 38\%)\), followed by \(dfrA12\) \((n=12; 18\%)\). Several ESBL genes were also detected, the most common being of the \(\text{bla}_{\text{CTX-M}}\) type \((n=18, 26\%)\). \(\text{bla}_{\text{CTX-M}}\) progenitor genes are thought to have originated from the chromosomes of soil bacteria [85] and their subsequent capture by predominately IncF family plasmids has led to their global dissemination in both clinical and non-clinical settings [30]. IncF family replicons were the most prevalent replicons detected in this collection, the foremost

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**Fig. 4.** Genotypic profile of 73 ExPEC isolates. Tip label colour corresponds to phylogroups (red, A; green, B1; teal, B2; purple, D). The adjacent heatmap represents phenotypic resistances and intact genes present in each isolate (orange, phenotypic resistance; light blue, ARG; dark blue, IS and class 1 and 2 integrase \((\text{intI}1, \text{intI}2)\); red, VAG; green, plasmid replicon). Along the top of the heatmap, the numbers in brackets after gene names represent the total number of isolates with trait present). Trimethoprim-susceptible ExPEC isolates ERR434278, ERR434751, ERR434270, ERR434273 and ERR434271 and trimethoprim-susceptible EAEC isolates SRR5470250, SRR5024242 and SRR3574247 (marked by asterisks) from other studies were added as controls. The tree on the left side is the maximum-likelihood tree seen in Fig. 2 (reference K12 MG1655 omitted) presented in cladogram form.
being IncFIB (n=53; 79%), followed by IncFII (n=51; 76%). Additionally, IncFIA replicons were detected in 30 isolates (44%). While bla\textsubscript{CTX-M} ESBLs comprise a heterogeneous family of enzymes, the bla\textsubscript{CTX-M-15} and bla\textsubscript{CTX-M-14} variants are currently most prevalent worldwide and are strongly associated with ST131 [30], though other STs have also contributed to their spread [86]. In this historic collection, bla\textsubscript{CTX-M-15} (n=11; 16%) and bla\textsubscript{CTX-M-14} (n=7; 10%) were the only two bla\textsubscript{CTX-M} variants identified. All bla\textsubscript{CTX-M-15} genes were carried solely by ST131 H30Rx isolates (Fig. 2a). Conversely, the bla\textsubscript{CTX-M-14} genes were present in four ST38 isolates, the sole examples of ST648 and ST773 and an H30R ST131 isolate. UTI isolates carrying bla\textsubscript{CTX-M} genes have previously been noted exhibiting high co-resistance to trimethoprim [77]. In this collection, all isolates carrying bla\textsubscript{CTX-M-15} also carried dfrA12, and all isolates carrying bla\textsubscript{CTX-M-14} co-occurred with dfrA17. Three additional ESBL genes were identified, bla\textsubscript{OXA-10}, bla\textsubscript{TEM-214} [87] and bla\textsubscript{TEM-57} [88], occurring in one ST95 isolate, one ST131 H30R isolate and one ST448 isolate, respectively.

In addition to trimethoprim- and ESBL-resistance genes, other common ARGs identified were β-lactamase gene bla\textsubscript{TEM-1B} (n=49; 73%), sulphonamide-resistance gene sul1 (n=46; 69%), aminoglycoside-resistance gene aadA5 (n=25; 37%), tetracycline-resistance gene tetA (n=22; 30%) and macrolide-resistance gene mphA (n=31; 46%). Fluoroquinolone-resistance genes aac(6)-Ib-cr (n=11; 16%) and qepA1 (n=1; ST448) were also detected. Isolates belonging to phylogroup A carried the highest number of ARGs overall (n=14). An MDS analysis demonstrated that ST131 H30Rx isolates possessed the most distinct ARG profiles (Fig. 5, red area), which could be attributed to the strong correlation of aac(3)-IIa, aac(6)-Ib-cr, aph(3′)-la, aadA2, bla\textsubscript{CTX-M-15}, bla\textsubscript{OXA-1} and dfrA12 in this ST and clade (Data S4). Notably, all the trimethoprim-sensitive ExPEC and EAEC isolates used as controls in this study clustered together with four trimethoprim-sensitive isolates from this collection (Fig. 5, blue area), with the exception of one control ExPEC ST131 isolate (Fig. 5, red arrow).

Heavy-metal- and detergent/disinfectant-resistance genes are often carried by the same MGEs harbouring ARGs, sparking concern and growing evidence that selective pressures induced by their extensive use in industry, agriculture and health-care facilities, as well as via contamination, can co-select for AMR [92–94]. The hospital wherein our isolates originated caters for a catchment area significantly associated with agriculture,
including food animals (bovine, sheep, pig and poultry), and various crops, such as fruits and nuts [95]. Thus, the isolates were screened, and two detergent-resistance genes – \textit{qacE\textsubscript{A1}} (\(n=47; 70\%\)) and \textit{qacH} (\(n=3; 4\%\)) – conferring resistance to quaternary ammonium compounds, and six heavy-metal-resistance genes were identified. All isolates carried \textit{zntA}, which confers resistance to zinc, lead and cadmium. We also identified \textit{chrA} (\(n=32; 48\%\)), corresponding to chromium resistance, and \textit{merA} (\(n=16; 24\%\)), which confers mercury resistance. Silver- and copper-resistance genes \textit{silA} and \textit{pcoA} were present in only three isolates (4%; all ST216) and tellurium-resistance gene \textit{tera} was observed in a single ST200. Heavy-metal-resistance genes, including \textit{zntA}, \textit{merA}, \textit{pcoA} and \textit{chrA}, are known to be significantly associated with detergent resistance genes, and \textit{sul}, \textit{tet}, \textit{bla\textsubscript{TEM}} \textit{bla\textsubscript{SHV}} and \textit{bla\textsubscript{CTX}} Variants [93]. Furthermore, chromium resistance has been positively correlated with relative \textit{bla\textsubscript{CTX-M}} and \textit{bla\textsubscript{OXA}} gene abundance [96]. Here, 100\% of isolates carrying \textit{bla\textsubscript{CTX-M}} genes also carried \textit{chrA}, and 91\% of isolates with \textit{bla\textsubscript{OXA}} also harboured \textit{chrA}.

### Class 1 and 2 integrons

The presence of class 1 or 2 integrons is considered a reliable proxy for an MDR genotype [97]. Class 1 integrons are more prevalent than class 2 integrons [97] and typically contain two conserved regions – 5′-CS and 3′-CS – flanking a variable gene cassette. The \textit{intI1} integrase gene, responsible for capturing, removing and rearranging genes within the variable cassette, is located in the 5′-CS region. The 3′-CS region typically contains a truncated but functional \textit{qacE\textsubscript{A1}} gene fused to a \textit{sul1} gene [98]. Class 1 and 2 integrons are mobilized and spread by MGEs, such as transposons and conjugative plasmids. Additionally, the insertion element IS26 is renowned for its ability to capture and assemble ARGs in complex resistance regions [32] and alter the structure of class 1 integrons [35, 98]. Here, IS26 was identified in 63 (94\%) isolates, including all 45 (67\%) that carried \textit{intI1}.

Class 1 and 2 integron loci are often rich in repetitive sequences, posing challenges to assembling complete structures using short-read sequencing data. Despite this limitation, we resolved 18 class 1 integron structures, including ARGs and MGEs found downstream of the typical 3′-CS region (Fig. 6a) and 1 class 2 integron (Fig. 6b). BLASTN and GenBank were used to determine whether these integrons had been previously deposited into public databases (Table 1).

In general, the integrons from this collection are also found in ExPEC-associated STs, most commonly ST131 and ST617 [of ST10 clonal complex (CC)]. They are most frequently seen on IncF plasmids of varying plasmid multilocus sequence types (pMLSTs), in a range of hosts, including humans, dogs and gulls, and from a range of geographical locations, including Europe, Asia, North America and Australia (Table 1). The most common genes located in the variable gene cassette were \textit{dfra17} and \textit{aadA5} (Fig. 6a).

Integron \textit{k} was the most common structure in this collection, present in 12 isolates (from ST131 and ST405), and carried \textit{dfra12} (trimethoprim), \textit{aadA2} (aminoglycoside), \textit{qacE\textsubscript{A1}} (quaternary ammonium compounds), \textit{sul1} (sulphonamide) and \textit{mphA} (macrolide) resistance genes. The same integron was described in an ST131 H30Rx isolate taken from a French patient a year after contracting pyelonephritis while travelling in Nepal [99]. The integron \textit{k} structure also carried the aforementioned chromium resistance gene \textit{chrA}, which in turn was also present in seven other integron structures (\(d, e, f, g, l, m, \text{and } r\)). Class 1 integrons are often translocated via mercury-resistance transposons carrying genes of the \textit{mer} operon [81], and such transposons were present in integrons \(i, j, \text{and } n\) structures. Integrons \(n\) and \(r\) also carried a recently characterized \textit{dfra4} gene conferring high-level resistance to trimethoprim. Both isolates that carry these integrons (ST1193 and ST448) predate the first report of this gene in a clinical sample (UTI) in 2017 [100]. Additionally, integron \textit{r} carried an uncommon plasmid-mediated fluoroquinolone-resistance gene, \textit{qepA}. Integron \(b\) was the only structure to carry an ESBL gene (\textit{bla\textsubscript{OXA-10}}), chloramphenicol (\textit{cmlA1}), aminoglycosides (\textit{aadA1}) and sulphonamides (\textit{sul1}). An identical structure to integron \(b\) could not be found in public databases; however, a similar structure was observed chromosomally in \textit{Pseudomonas aeruginosa} strain SP4527 (ST357) isolated from a patient with bacteremia in India, in 2016 (Fig. 6b). Integron \(c\) consists of a short \textit{dfra5-IS26} configuration that has been detected in, and is identical to, \textit{E. coli} strains sourced from cattle located on several NSW properties [98, 101], and in commensal \textit{E. coli} from Australian pigs [35]. Furthermore, the signature has been used to track CoV plasmids carrying a complex resistance locus in patients with UTI and urosepsis in Australia in \textit{E. coli} ST58 [102, 103]. These studies highlight how plasmids that play an important role in carrying intestinal pathogenic \textit{E. coli} virulence genes [101] and extraintestinal pathogenic virulence gene cargo [102, 103] can be tracked by the presence of this unique signatures. Moreover, it shows that transposon belonging to the \textit{Tn3} family of mercury-resistance transposons can mobilize complex resistance loci on diverse plasmid backbones. Integron \(l\) has been described within a larger salmonella genomic island 1 (SGI1) structure in \textit{Proteus mirabilis} [104]. The integron a structure carried the sole example of a \textit{sul3} gene, which has been associated with Australian \textit{E. coli} isolates from pig farms [35, 83]. This integron, found in our ST10 isolates, also uniquely carried \textit{dfra16}, \textit{bla\textsubscript{TEM}} and \textit{qacH}. Integron \(a\) has been located on IncFIA and IncI plasmids in CC10 (ST617) \textit{E. coli} isolates originating from both human clinical samples from Germany and the USA, a broiler chicken sample from Belgium (Table 1), as well as farmed red deer from Spain [105]. Incidentally, the patients from whom these isolates originated resided in proximity to two red deer farms. Lastly, the class 2 integrase \textit{intI2} gene was identified in one ST372 and one ST73 isolate (Fig. 6b). The \textit{intI2} from this class of integron is inactive, impeding its ability to modify gene cassettes. Thus, class 2 integron cassettes are highly conserved [97]. The \textit{dfra15-sat2-aadA1} cassette found in this collection represents the most common
Fig. 6. Class 1 and class 2 integron structures. (a) Class 1 (a–r) and class 2 integrons structures identified in this collection. ST131 H30R i and ii refer to the two H30R subclades resolved by the SNP-based phylogeny presented in Fig. 2(b). (b) EasyFig comparison of integron b and the chromosome-associated integron structure from Pseudomonas aeruginosa strain SP4527.
### Table 1. Summary of BLAST hits to integrons found in this collection

Rows in bold indicate integrons with no 100% matches. The closest integron match is provided.

| Integron Accession no. | Coverage; Identity | Integron Location | Inc group (pMLST) | Isolation date | Species (ST)/ [STs from this collection] | Host | Host disease | Host location |
|------------------------|--------------------|-------------------|-------------------|----------------|------------------------------------------|------|--------------|--------------|
| MK070495.1             | 100%; 100%         | Plasmid           | I1 (CC-3)         | 2013           | *E. coli* (na)/[ST10]                     | Chicken | NA           | Belgium      |
| CP018105.1             | 100%; 100%         | Plasmid           | F (F-;A18:B-)     | 2016           | *E. coli* (ST617)                        | Human | NA           | Germany      |
| CP034409.1             | 93%; 100%          | Chromosome        | -                 | 2016           | *P. aeruginosa* (ST357)/[ST95]           | Human | Sepsis       | India        |
| CP026940.1             | 100%; 100%         | Plasmid           | F (F2:A-;B1)      | 2010           | *E. coli* (ST58)/[ST58, ST117, ST141]    | Cow   | Mastitis     | France       |
| CP041338.1             | 84%; 100%          | Plasmid           | N (A2;N5;J-)      | NA             | *E. coli* (ST131)/[ST131, ST127]         | NA    | NA           | NA           |
| MK295829.1             | 100%; 100%         | Plasmid           | F (F1:A6;B20)     | 2015           | *E. coli* (ST131)                        | Dog   | NA           | Israel       |
| MK295827.1             | 100%; 100%         | Plasmid           | F (F1:A6;B20)     | 2015           | *E. coli* (ST131)                        | Human | NA           | Israel       |
| MK295830.1             | 100%; 100%         | Plasmid           | F (F1:A6;B20)     | 2007           | *E. coli* (ST131)/[ST38]                 | Human | NA           | Israel       |
| CP021734.1             | 84%; 100%          | Plasmid           | N (A2;N5;J-)      | NA             | *E. coli* (ST617)/[ST131]                | NA    | NA           | NA           |
| CP041557.1             | 100%; 100%         | Plasmid           | F (F2:A1;B-)      | NA             | *E. coli* (ST131)/[ST131]                | Human | NA           | USA          |
| CP041272.1             | 100%; 100%         | Plasmid           | F (F2:A1;B1)      | 2015           | *E. coli* (ST131)                        | Yak   | NA           | China        |
| CP024718.1             | 100%; 100%         | Plasmid           | F (F2:A1;B1)      | 2015           | *E. coli* (ST131)                        | Yak   | NA           | China        |
| CP021289.1             | 100%; 100%         | Plasmid           | F (F51:A-;B10)    | 2010           | *E. coli* (ST95)/[ST73, ST95, ST550]     | Human | Sepsis       | Australia    |
| CP023845.1             | 100%; 100%         | Plasmid           | F (F22:A1;B20)    | 2009           | *E. coli* (ST131)/[ST131, ST405]         | Human | Post UTI     | Sweden       |
| LC056386.1             | 100%; 100%         | Contig            | -                 | 2013           | *E. coli* (na)                            | Environment | NA           | India        |
| CP028322.1             | 100%; 100%         | Plasmid           | F (F73:A-;B-)     | 2000           | *E. coli* (ST73)                          | Human | NA           | USA          |
| CP022632.1             | 100%; 100%         | Plasmid           | F (F2:A-;B-)      | 2008           | *E. coli* (ST95)                          | Human | Sepsis       | USA          |
| CP041523.1             | 100%; 100%         | Plasmid           | F (F29:A-;B10)    | 2014           | *E. coli* (ST69)/[ST69]                  | Human | Sepsis       | USA          |
| CP032145.1             | 100%; 100%         | Chromosome        | -                 | 2015           | *E. coli* (ST38)/[ST38]                  | Gull   | NA           | Turkey       |
| CP040263.1             | 100%; 100%         | Chromosome        | -                 | 2014           | *E. coli* (ST38)                          | Mollusc | NA          | Norway       |
| CP040390.1             | 100%; 100%         | Chromosome        | -                 | 2016           | *E. coli* (ST38)                          | Gull   | NA           | USA          |
| CP016546.1             | 97%; 99.97%        | Chromosome        | -                 | 2015           | *E. coli* (ST359)/[ST1193]               | Human | Healthy      | Netherlands  |
| CP019014.1             | 100%; 100%         | Plasmid           | F (F2:A1;B-)      | 2013           | *E. coli* (ST131)/[ST773]                | Human | NA           | USA          |
| CP041393.1             | 100%; 100%         | Plasmid           | F (F36:A4;B-)     | 2018           | *E. coli* (ST167)/[ST167]                | Dog   | NA           | USA          |
| MG764548.1             | 100%; 100%         | Plasmid           | F (F33;A4;B-)     | NA             | *E. coli* (na)                            | NA    | NA           | China        |
| CP019076.1             | 100%; 100%         | Plasmid           | F (F36:A4;B-)     | 2013           | *E. coli* (ST617)                        | Human | NA           | China        |
| CP021737.1             | 100%; 100%         | Plasmid           | F (F36:A4;B-)     | NA             | *E. coli* (ST617)                        | NA    | NA           | NA           |
| KU043115.1             | 100%; 100%         | Plasmid           | F (F36:A4;B-)     | 2013           | *E. coli* (ST617)                        | Human | UTI          | China        |
| CP023854.1             | 100%; 99%          | Plasmid           | F (F48:A1:-B49)   | 2009           | *E. coli* (na)/[ST405]                   | Human | UTI          | Sweden       |

Continued
array, and is particularly prevalent in UTI-associated Proteus species [106].

**VAGs**

ExPEC employ a range of VAGs that enable ascension, colonization and persistence within the urinary tract. Like ARGs, many VAGs are also carried on MGEs, including plasmids as well as chromosomally situated PAIs [15]. Thus, the virulence potential of ExPEC strains is constantly evolving and tracking VAGs improves understanding of ExPEC pathogenesis. Using the VirulenceFinder database, a total of 84 VAGs were identified in this collection, with the number of VAGs per isolate spanning from 2 to 36, with a mean of 20 and a median of 19.

To combat the iron-scarcity encountered in the urinary tract, ExPEC have acquired a number of siderophore and ferrous iron uptake systems to scavenge Fe^{3+} and Fe^{2+}, which are vital ExPEC have acquired a number of siderophore and ferrous iron-utilization systems to scavenge Fe^{3+} and Fe^{2+}, which are vital to their growth, persistence and establishment of full virulence [15]. ExPEC-associated iron-acquisition systems include yersinia-bacterin (ybt, irm, fyua), salmochelin (iro), aerobactin (iuc, iut), ferric citrate transport system (fec) and the Sit ferrous iron utilization system (sit). In this study, the most common iron-acquisition genes were fyua, irm2 and sita (all n=61, 91%), followed by iucD and fecA (both n=45; 67%). The high carriage of fyua is consistent with previous reports tracking VAGs in ExPEC [14, 107, 108]. While there are contradictory reports regarding fyua and mortality [107, 109, 110], immunization with FyuA has been shown to protect against pyelonephritis in mice [111].

Several adhesins and fimbriae demonstrate specificity for binding uroepithelium. Of these, the type 1 fimbriae and P-fimbriae are most prevalent in UTI-associated ExPEC [15]. Type 1 fimbriae bind uroepithelial associated α-D-mannosylated proteins via fimbrial adhesin H (fimH). Thus, the fimH gene plays a pivotal role in ExPEC urothelial adhesion [15]. P-fimbriae (pap genes) bind α-D-galactopyranosyl(1→4)-β-D-galactopyranoside-containing receptors found in the upper urinary tract and, therefore, have been associated with pyelonephritis [112]. Here, the most common adhesins identified were fimH (n=61; 91%), papB (n=47; 70%) and papI (n=45; 67%). However, other P-fimbriae operon genes were less common, such as papC, papD, papI and papK (each n=20; 30%), papF (n=21; 31%), papH and papG (both n=19; 28%), and papA (n=17; 25%).

In addition to factors facilitating adhesion and growth, ExPEC express toxins contributing to host tissue damage, as well as immune evasion molecules [113]. Regarding toxins, several serine protease autotransporters of Enterobacteriaceae (SPATE) genes were identified including sat (secreted autotransporter toxin) (n=39; 58%), which has been shown to compromise gap junctions in uroepithelium [112], vat (n=18; 27%) and pic (n=8; 12%). Enterotoxins senB (n=21; 31%) and astA (n=4; 6%) were also observed. Regarding immune evasion, VAGs encoding increased serum survival protein (iss), VirB5-like protein TraT (traT) and outer-membrane protein T (ompT) were identified in 52 (78%), 50 (75%) and 5 (7%) isolates, respectively.

Despite studies reporting various VAGs linked to pyelonephritis and sepsis [15, 114, 115], we found no single gene nor plasmid replicon strongly associated with a particular pathology (correlation heat map in Data S5), nor was there a significant difference between the number of VAGs present and uro-clinical syndrome (Fig. 7a). The discrepancy with the literature may have arisen due to sampling based on trimethoprim resistance and/or the relatively small sample size, limiting statistical power. Nevertheless, an MDS analysis showed some clustering of STs based on virulence profiles (Fig. 7b). Notably, isolates formed two distinct diagonal groups, which could be attributed to the distribution of pap genes with papC, papD, papI and papK present in all isolates within the distal group and absent in all isolates in the top group. Other pap genes were also more prevalent in the bottom group including papGIII (0% top, 78% bottom), papGIII (0, 17%), papH (0, 96%), papE (4%, 0%) and papF (4, 96%). Also of note was that the ST131 H30 isolates clustered with two emerging pathogen STs, ST405 [116] and ST1193 [117], as well an ST550 isolate [same CC as ST1193 – CC41] and one ST73 isolate that deviated from other ST73 isolates (Fig. 7b, red area). The ST131 H30 isolates fell under virotype C [afa operon (−), iroN (−), ibeA (−), sat (+)], which is the most represented ST131 virotype internationally and associated with a higher frequency of infection [118]. The other isolates within this cluster could also be categorized as virotype C. The only ST131 H27 isolate did not cluster with its ST and carried the molecular predictors of EAEC/ExPEC hybrid strains (aatA, aggR, fyua), as did one ST200 isolate. The EAEC isolates sourced from outside this collection (Fig. 7b, blue arrows) all deviated from ExPEC isolates of the same ST; however, they did not form a separate cluster and each carried ExPEC-associated genes including fyua (Fig. 4). These EAEC isolates were not previously described as hybrid strains and originated from cases of diarrhoeal disease in England [40]. However, hybrid EAEC/ExPEC strains are known to cause UTI outbreaks [119, 120]. EAEC strains carry most of their virulence cargo on plasmids and are renowned for prolific biofilm formation [121]. Indeed, EAEC/ExPEC hybrid strains

| Table 1. Continued |
|-------------------|
| **Integron** | **Accession no.** | **Coverage: Identity** | **Integron location** | **Integron (pMLST)** | **Isolation date** | **Species (ST)/ [STs from this collection]** | **Host** | **Host disease** | **Host location** |
| r | CP014320.1 | 100%; 99% | Plasmid | F (F36.A–B32) | 2007 | *E. coli* (ST131)/[ST448] | Human | UTI | USA |

NA: Not Applicable

*Only 100% matches from *E. coli* are shown.
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have shown significantly higher levels of biofilm formation, as well as enhanced adhesion to uroepithelium cells, compared to non-hybrids [122].

**PAIs**

The PAI marker genes malX and usp were seen in 44 (66%) and 38 (57%) isolates, respectively. The acquisition of PAIs via horizontal transfer can vastly contribute to the evolution of *E. coli* pathogens as they contain potent combinations of VAGs, including the aforementioned P-fimbriae, salmochelin, aerobactin and yersiniabactin operons, as well as shigella-fimbriae (S-fimbriae, *sfa*) and haemolysin toxin (*hly*) operons [123, 124]. As such, to indicate the presence of ExPEC-associated PAIs, we used short-reads derived from each isolate mapped to reference sequences PAI I CFT073, PAI II 536 and PAI III 536. PAI IV 536 [also known as high pathogenicity island (HPI)] was also screened for; however, possibly due to allelic differences, the read-depth was relatively lower for this PAI (Data S6) so *blast* was used in conjunction with assembled draft genomes to indicate its presence. While all isolates had at least partial hits to genes and gene clusters within PAI I CFT073, PAI II 536 and PAI III 536, most demonstrated extensive deletions of regions, and few had near intact examples (Fig. 8).

The most represented VAG operon was aerobactin in PAI I CFT073 and the least were S-fimbriae in PAI III 536. However, PAI IV 536 – which consists of the yersiniabactin operon (*ybt* genes, *irp1, irp2, fyuA*; total 11 genes), a P4-like integrase and an uncharacterized protein Yeel nestled between tRNA-Ser and tRNA-Asn – was identified over >95% length at >98% identity in 58 isolates (Data S7). In addition to being a potent siderophore system, yersiniabactin can protect against copper toxicity, redox-based phagocytosis and is a prerequisite for sepsis in some *E. coli* strains [19].

**ColV plasmid markers**

ColV plasmids are considered a defining feature of APEC (avian pathogenic *E. coli*) [125]; however, they have also been associated with human ExPEC strains [102, 126]. While the expression of colicin V may benefit ExPEC strains by reducing competition for nutrients [127], ColV plasmids are intriguing in that they share high sequence homology to PAIs, and it is theorized that they and other plasmids and phages are the progenitors of ExPEC-associated PAIs [123]. As ColV plasmids are heterogeneous in their VAGs and pMLST, Liu *et al.* [67] defined ColV plasmids as having at least one gene from four or more of the following: (i) *cvaABC* and *cvi* (ColV operon); (ii) *sitABCD* (ferrous iron utilization system); (iii) *hlyF* (regulator of outer membrane vesicle biogenesis) and *ompT*; (iv) *iucABCD* and *iutA* (aerobactin operon); (v) *iroB-CDEN* (salmochelin operon); and (vi) *etsABC* (novel ABC transport system). In this study, aribA gene screening identified seven isolates (10%) meeting this criterion – HOS66 (ST73, cystitis), HOS45 (ST73, cystitis), HOS74 (ST95, cystitis), HOS17 (ST117, pyelonephritis), HOS36 (ST117,
pyelonephritis), HOS48 (ST58, cystitis) and HOS53 (ST448, cystitis). Three of these isolates (HOS17, HOS36 and HOS48) also carried the aforementioned integron \(\text{dfrA5-IS26}\) that is used to track ColV plasmids containing complex AMR regions. Future long-read sequencing studies are needed to provide deeper insight into these initial observations (plasmid short-read mapping is provided in Data S8).

**SNP analysis of serial patient isolates**

Phenotypic diversification via adaptive evolution is driven by horizontal gene transfer and by mutation [128]. A single SNP-mediated point mutation can lead to an amino acid replacement with significant adaptive effects, as witnessed in single amino acid replacements in \(\text{parC/E}\) and \(\text{gyrA}\) genes causing fluoroquinolone resistance [24, 25], and in \(\text{fimH}\), which increases binding to uroepithelium and biofilm formation [129]. Such adaptive mutations can increase bacterial fitness and provide rapid adaption to niche-specific conditions [128]. ExPEC are exposed to a variety of niches in addition to their primary reservoir, the intestinal tract, and as a result may be under greater selective pressure for adaptive mutations. Indeed, UTI-associated ExPEC are known to have significantly higher mutation rates compared to other \(\text{E. coli}\) pathotypes [130]. While studies have been conducted comparing SNP-mediated point mutations in UTI and faecal isolates [131], relatively little is known about the SNPs that occur in recurrent UTIs, particularly when subsequently more serious sequelae occur. In this study, 13 patients had more than one presentation of symptomatic UTI during the study period (indicated by letters in Fig. 2). Six patients had subsequent UTIs caused by the same ST, and seven patients had different STs. Here, we focus on the patients who had multiple infections with the same ST (Fig. 9), and on SNP-mediated mutations with a higher probability of inducing functional changes as predicted by the BLOSUM62 substitution matrix [132]. A catalogue of all SNPs can be viewed in Data S9.

Patient A was a 37-year-old female who in the first instance presented with cystitis with an ST10 isolate (HOS1) as the causative agent and then pyelonephritis (HOS2; ST10) 63 days later. The ST10 isolates had a difference of 29 SNPs, with most occurring in genes involved in DNA processing, transport and virulence (Fig. 9). However, mutations with negative BLOSUM62 scores (indicating a higher probability that the original function is not conserved) occurred most frequently in VAGs and DNA processing. Potentially increasing the significance of the observed mutations, another patient, a 69-year-old male living in the same postcode, presented with pyelonephritis 16 days after patient A’s second presentation. The causative agent was also an ST10 (HOS34), which shared 71% \((n=24)\) of the same SNPs found in HOS2. Both HOS2 and HOS34 had the same SNPs causing non-synonymous amino acid substitutions with negative BLOSUM scores in the following genes: \(\text{irp1}\) (D2717A), \(\text{ybtS}\) (P184A), \(\text{asta}\) (G103V), \(\text{aroC}\) (G152E), \(\text{rnhB}\) (R98L), \(\text{hrpB}\) (G644E), \(\text{traC}\) (G644W), \(\text{lsrA}\) (E296G) and \(\text{recC}\) (S794C). Interestingly, \(\text{recC}\) was previously shown to be under positive selective pressure for adaptive mutations [130]. Though not present in HOS34, HOS2 possessed a SNP causing A432D (negative BLOSUM62 score) in DNA-repair protein Mfd (mutation-frequency-decline). Mfd promotes mutagenesis and accelerates AMR development to multiple antibiotic classes [133]. Fascinatingly, point mutations produce Mfd variants that express greater activity than wild-type [134].

Patient B was a 21-year-old female who first presented with cystitis (HOS3) then pyelonephritis 284 days later (HOS4, 13 SNP difference), then pyelonephritis again 128 days later (HOS5, 19 SNP difference from HOS3). The causative agent in

![Fig. 8](#) Mapping of short-reads indicating the presence of ExPEC-associated PAIs. Clustering of rows is based on the similarity between isolate PAI coverage profiles. (a) Schematic and heatmap for PAI \(\text{ICFT072}\) coverage. (b) Schematic and heatmap for PAI \(\text{II}_{536}\) coverage. (c) Schematic and heatmap for PAI \(\text{III}_{16}\) coverage. Trees alongside the heatmaps were reconstructed by hierarchical clustering using Euclidean agglomeration method.
each episode was ST393; however, unlike the high percentage of conserved SNPs in ST10 pyelonephritis isolates, HOS4 and HOS5 ST393 only shared 23% of all SNPs against the HOS3 reference, possibly due to the longer period between isolations, and only shared two negative BLOSUM62 score amino acid substitutions in chromosome partition protein MukF (F401V and T394H). Most SNPs occurred in intergenic regions, which may be significant as these regions often contain riboswitches, promoters, terminators and regulator binding sites, and SNPs can cause significant phenotypic changes [135].

Patient C was a 64-year-old female with cystitis and two samples were taken on the same day. Both isolates (HOS8 and HOS9) were ST144 and differed by five SNPs. While no SNPs occurred in VAGs (Fig. 9), two mutations with potential for functional change were identified in transposase TnpB (G11R) and environmental stress protein Ves (G7R). Conversely, in patient H, a 67-year-old male who presented with cystitis (HOS33; ST131 H30Rx) and then cystitis again a week later (HOS34; ST131 H30Rx), the two causative agents were indistinguishable.

Patient F was a 72-year-old female with cystitis followed by urosepsis 181 days later (HOS26; ST131 H30R; 53 SNPs or 20 SNPs without recombination), followed by urosepsis 101 days later (HOS27; ST131 H30R; 61 or 27 SNPs) and then a subsequent sample was taken 3 days later (HOS28; ST131 H30R; 62 or 26 SNPs). The vast majority of SNPs were a result of recombination, which is 50–100 times more likely to occur than mutations [136], and occurred in aerobactin biosynthesis protein iucC and iron transport system genes sitABC. However, three point mutations with potential change to function were identified in all urosepsis isolates: R18L in capsule protein KpsM, R268L in molybdoenzyme biosynthesis protein MoeA and M58R in molybdoenzyme chaperon protein YcdY. Intriguingly, HOS50 (also ST131 H30R), originating from a 50-year-old patient presenting with pyelonephritis 21 days prior to patient F’s initial presentation, also possessed these three precise amino acid substitutions produced by the same SNPs. These identical substitutions may be indicative of a shared source, particularly as both patients resided in the same postcode, or may be examples of pathoadaptive mutations that increase fitness in the upper urinary tract. However, no recombination events were detected in HOS50.

Lastly, patient K was a 2-year-old female who presented with urosepsis (HOS41; ST69) and subsequent urosepsis 100 days later (HOS42; ST69; 13 SNPs). Similar to isolates derived from patient B, most SNPs occurred within intergenic regions (Fig. 9); however, R133L in yersiniabactin protein ybtT and R337S in K⁺/H⁺ antiporter CvrA may have incurred functional changes. Future characterization of all these SNP-mediated mutations is required to ascertain any functional changes.

**Fig. 9.** Serial patient isolates – SNP distribution and consequence to amino acid sequence. Each panel represents an UTI patient from whom multiple isolates of the same ST were derived. Sunburst graphs represent the distribution of SNPs within functional groups. The number in the centre represents the total number of unique SNPs (accumulative if more than one recurrent infection occurred). Stacked column graphs represent the consequence of SNPs to protein sequence. Color legend for the stacked columns located bottom right corner. Unstacked column graphs represent the distribution of specific SNPs in recurrent infections from the same patient, and in the case of patient A and F, two different patients. 
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Author contributions
H.L. contributed roles: data curation, formal analysis, investigation, methodology, software, validation, visualization. C. J. R. contributor roles: formal analysis, methodology, software. T. K. contributor roles: data curation, resources. V. M. J. contributor roles: investigation, methodology, project administration, supervision, validation, writing – original draft. S. P. D. contributor roles: conceptualization, funding acquisition, project administration, supervision. All authors had a role in writing – review and editing.

Conflicts of interest
The authors declare that there are no conflicts of interest

Ethical statement
The project was approved by Charles Sturt University and Sydney West Area Health Service research ethics committees. Since clinical information for patients with UTIs was provided anonymously by clinicians, patient consent was not required.

References
1. Simmering JE, Tang F, Cavanaugh JE, Polgreen LA, Polgreen PM. The increase in hospitalizations for urinary tract infections and the associated costs in the United States, 1998–2011. Open Forum Infect Dis 2017;4:ofw281.
2. Medical Technology Group. Admissions of Failure: the Truth About Unplanned NHS Admissions in England (https://mtg.org.uk/wp-content/uploads/2016/07/Admissions-of-Failure-2015.pdf). London: Medical Technology Group; 2015.
3. Australian Commission on Safety and Quality in Health Care. Atlas 2017 – 1. Chronic Disease and Infection: Potentially Preventable Hospitalisations (www.safetyandquality.gov.au/our-work/healthcare-variation/atlas-2017/atlas-2017-1-chronic-disease-and-infection-potentially-preventable-hospitalisations). Sydney: Australian Commission on Safety and Quality in Health Care; 2017.
4. Australian Commission on Safety and Quality in Health Care. AURA 2019: Third Australian Report on Antimicrobial Use and Resistance in Human Health (www.safetyandquality.gov.au/publications-and-resources/resource-library/aura-2019-third-australian-report-antimicrobial-use-and-resistance-human-health). Sydney: Australian Commission on Safety and Quality in Health Care; 2019.
5. Abbou LM, Hooton TM. Antimicrobial stewardship and urinary tract infections. Antibiotics 2014;3:174–192.
6. Foxman B, Brown P. Epidemiology of urinary tract infections: transmission and risk factors, incidence, and costs. Infect Dis Clin North Am 2003;17:227–241.
7. Manges AR, Geum HM, Guo A, Edens TJ, Fibke CD et al. Global extraintestinal pathogenic Escherichia coli (ExPEC) lineages. Clin Microbiol Rev 2019;32:e00135-18.
8. Yamraj R, Rubin J, Thys E, Friedman CR, Riley LW. Persistent pandemic lineages of uropathogenic Escherichia coli in a college community from 1999 to 2017. J Clin Microbiol 2018;56:e01834-17.
9. Manges AR, Johnson JR. Reservoirs of extraintestinal pathogenic Escherichia coli. Microbiol Spectr 2018;5:UTI-0006-2012.
10. Kantele A, Lääveri T, Mero S, Häkkinen IMK, Kirveskari J et al. Despite predominance of uropathogenic/extraintestinal pathotypes among travel-acquired extended-spectrum β-lactamase-producing Escherichia coli, the most commonly associated clinical manifestation is travelers’ diarrhea. Clin Infect Dis 2020;70:210–218.
11. Russo TA, Johnson JR. Medical and economic impact of extraintestinal infections due to Escherichia coli: focus on an increasingly important endemic problem. Microbes Infect 2003;5:449–456.
12. Epstein L, Dantes R, Magill S, Fiore A. Varying estimates of sepsis mortality using death certificates and administrative codes — United States, 1999–2014. MMWR Morb Mortal Wkly Rep 2016;65:342–345.
13. Rhee C, Dantes R, Epstein L, Murphy DJ, Seymour CW et al. Incidence and trends of sepsis in US hospitals using clinical vs claims data, 2009–2014. JAMA 2017;318:1241–1249.
14. Daga AP, Koga VL, Soncini JGM, de Matos CM, Perugini MRE et al. Escherichia coli bloodstream infections in patients at a university hospital: virulence factors and clinical characteristics. Front Cell Infect Microbiol 2019;5:9:191.
15. Dale AP, Woodford N. Extra-intestinal pathogenic Escherichia coli (ExPEC): disease, carriage and clones. J Infect 2015;71:615–626.
16. Micenková L, Bosák J, Vrba M, Ševčíková A, Šmajs D. Human extraintestinal pathogenic Escherichia coli strains differ in prevalence of virulence factors, phylogroups, and bacteriocin determinants. BMC Microbiol 2016:16:218.
17. Johnson TJ, Nolan LK. Pathogenicity of the virulence plasmids of Escherichia coli. MMBR 2009;73:750–774.
18. Peirano G, Mulvey GL, Armstrong GD, Pitout JDD. Virulence potential and adherence properties of Escherichia coli that produce CTX-M and NDM β-lactamases. J Med Microbiol 2013;62:525–530.
19. Johnson JR, Magistro G, Clabots C, Porter S, Manges A et al. Contribution of yersiniabactin to the virulence of an Escherichia coli sequence type 69 (‐clonal group A) P. cystisolate in murine models of urinary tract infection and sepsis. Microb Pathog 2018;128:128–131.
20. Hancock V, Ferrières L, Klemm P. The ferric yersiniabactin uptake receptor FyuA is required for efficient biofilm formation by urinary tract infectious Escherichia coli in human urine. Microbiology 2008;154:167–175.
21. Magistro G, Hoffmann C, Schubert S. The salmochelin receptor IroN itself, but not salmochelin-mediated iron uptake promotes biofilm formation in extraintestinal pathogenetic Escherichia coli (ExPEC). Int J Med Microbiol 2015;305:435–445.
22. Alam M, Bastakoti B. Therapeutic guidelines: antibiotic, version 15. Aust Prescr 2015;38:137.
23. Fasugba O, Mitchell BG, Mnatzaganian G, Das A, Collignon P et al. Five-year antimicrobial resistance patterns of urinary Escherichia coli at an Australian tertiary hospital: time series analyses of prevalence data. PLoS One 2016;11:e0164306.
24. Nicolas-Chanoine M-H, Bertrand X, Madec J-Y. Escherichia coli ST131, an intriguing clonal group. Clin Microbiol Rev 2014;27:543–574.
25. Price LB, Johnson JR, Aziz M, Clabots C, Johnston B et al. The epidemic of extended-spectrum-β-lactamase-producing Escherichia coli ST131 is driven by a single highly pathogenic subclone, H30-Rx. mBio 2013;4:e00377-13.
26. Paltansingh S, Kraakman MEM, Ras JMC, Wessels E, Bernards AT. Characterization of fluoroquinolone and cephalosporin resistance mechanisms in Enterobacteriaceae isolated in a Dutch teaching hospital reveals the presence of an Escherichia coli ST131 clone with a specific mutation in parE. J Antimicrob Chemother 2013;68:40–45.
27. Johnson JR, Tchesnokova V, Johnston B, Clabots C, Roberts PL et al. Abrupt emergence of a single dominant multidrug-resistant strain of Escherichia coli. J Infect Dis 2013;207:919–928.
28. Poolman JT, Wacker M. Extraintestinal pathogenic Escherichia coli, a common human pathogen: challenges for vaccine development and progress in the field. J Infect Dis 2016;213:6–13.
29. Zilberberg MD, Shorr AF. Secular trends in gram-negative resistance among urinary tract infection hospitalizations in the United States, 2000–2009. Infect Control Hosp Epidemiol 2013;34:940–946.

30. Bevan ER, Jones AM, Hawkey PM. Global epidemiology of CTX-M β-lactamases: temporal and geographical shifts in genotype. J Antimicrob Chemother 2017;72:2145–2155.

31. Bajaj P, Singh NS, Virdi JS. Escherichia coli β-lactamases: what really matters. Front Microbiol 2016;7:417.

32. Partridge SR, Kwong SM, Firth N, Jensen SO. Mobile genetic elements associated with antimicrobial resistance. Clin Microbiol Rev 2018;31:e00088–17.

33. Clinical and Laboratory Standards Institute. Performance Standards for Antimicrobial Susceptibility Testing. Wayne, PA: Clinical and Laboratory Standards Institute; 2017.

34. Clinical and Laboratory Standards Institute. Performance Standards for Antimicrobial Disk Susceptibility Tests for Bacteria That Grow Aerobically: Approved Standard, M02-A10. 10th edn. Wayne, PA: Clinical and Laboratory Standards Institute; 2009.

35. Reid CJ, Wyrsch ER, Chowdhury PR, Zingali T, Liu M et al. Porcine commensal Escherichia coli: a reservoir for class 1 integrons associated with IS26. Microb Genom 2017:3:mgen.00014.13.

36. Minh BQ, Schmidt HA, Chernomor O, Schrempf D, Woodhams MD et al. IQ-TREE 2: new models and efficient methods for phylogenetic inference in the genomic era. Mol Biol Evol 2020;37:1530–1534.

37. Croucher NJ, Page AJ, Connor TR, Delaney AJ, Keane JA et al. Rapid phylogenetic analysis of large samples of recombinant bacterial whole genome sequences using Gubbins. Nucleic Acids Res 2015;43:e15.

38. Kallonen T, Brodick HJ, Harris SR, Corander J, Brown NM et al. Systematic longitudinal survey of invasive Escherichia coli in England demonstrates a stable population structure only transiently disturbed by the emergence of ST131. Genome Res 2017;27:1437–1449.

39. Nascimento V, Day MR, Doumith M, Hopkins KL, Woodford N et al. Comparison of phenotypic and WGS-derived antimicrobial resistance profiles of enteroaggregative Escherichia coli isolated from cases of diarrhoeal disease in England, 2015–16. J Antimicrob Chemother 2017;72:3288–3297.

40. Letunic I, Bork P. Interactive Tree Of Life (iTOl) v4: recent updates and new developments. Nucleic Acids Res 2019;47:W256–W259.

41. Page AJ, Cummins CA, Hunt M, Wong VK, Reuter S et al. Roary: rapid large-scale prokaryote pan genome analysis. Bioinformatics 2015;31:3691–3693.

42. Hadfield J, Croucher NJ, Goater RJ, Abudahab K, Aanensen DM et al. Phandango: an interactive viewer for bacterial population genomics. Bioinformatics 2018;34:292–293.

43. Seemann T. Prokka: rapid prokaryotic genome annotation. Bioinformatics 2014;30:2068–2069.

44. Larsen MV, Cosentino S, Rasmussen S, Friis C, Hasman H et al. Multilocus sequence typing of total-genome-sequenced bacteria. J Clin Microbiol 2012;50:1395–1361.

45. Joensen KG, Tetzchner AMM, Iguchi A, Aarestrup FM, Scheutz F. Rapid and easy in silico serotyping of Escherichia coli isolates by use of whole-genome sequencing data. J Clin Microbiol 2015;53:2410–2420.

46. Clermont O, Christensen JK, Denamur E, Gordon DM. The Clermont Escherichia coli phylo-typing method revisited: improvement of specificity and detection of new phylo-groups. Environ Microbiol Rep 2013;5:58–65.

47. Camacho C, Coulouris G, Avagyan V, Ma N, Papadopoulos J et al. BLAST+: architecture and applications. BMC Bioinformatics 2009;10:421.

48. Hunt M, Mather AE, Sánchez-Busó L, Page AJ, Parkhill J et al. ARIBA: rapid antimicrobial resistance genotyping directly from sequencing reads. Microb Genom 2017;3:mgen.000131.

49. Zankari E, Hasman H, Cosentino S, Vestergaard M, Rasmussen S et al. Identification of acquired antimicrobial resistance genes. J Antimicrob Chemother 2012;67:2640–2644.

50. Carattoli A, Zankari E, García-Fernández A, Voldby Larsen M, Lund O et al. In silico detection and typing of plasmins using plasmidfinder and plasmid multilocus sequence typing. Antimicrob Agents Chemother 2014;58:3985–3993.

51. Joensen KG, Scheutz F, Lund O, Hasman H, Kaas RS et al. Real-time whole-genome sequencing for routine typing, surveillance, and outbreak detection of verotoxigenic Escherichia coli. J Clin Microbiol 2014;52:1501–1510.

52. Johnson JR, Stell AL, Scheutz F, O’Bryant TT, Russo TA et al. Analysis of the F antigen-specific papAlleles of extraintestinal pathogenic Escherichia coli using a novel multiplex PCR-based assay. Infect Immun 2000;68:1587–1599.

53. Yu G, Smith DK, Zhu H, Guan Y, Lam Tommy Tszan-Yuk. ggtree: an r package for visualization and annotation of phylogenetic trees with their covariates and other associated data. Methods Ecol Evol 2017;8:28–36.

54. Li H, D turb R. Fast and accurate short read alignment with Burrows-Wheeler transform. Bioinformatics 2009;25:1754–1760.

55. Li H, Handsaker B, Wysoker A, Fennell T, Ruan J et al. The Sequence Alignment/Map format and SAMtools. Bioinformatics 2009;25:2078–2079.

56. Leray M, Knowlton N, Ho S-L, Nguyen BN, Machida RJ. GenBank is a reliable resource for 21st century biodiversity research. Proc Natl Acad Sci USA 2019;116:22651–22666.

57. Zhou Z, A likhan N-F, Mohamed K, Fan Y, Agama Study Group, et al. The Enterobase user’s guide, with case studies on Salmo nella transmissions, Yersinia pestis phylogeny and Escherichia core genomic diversity. Genome Res 2019;19:251678.

58. Sullivan MJ, Petty NK, Beaton SA. Easyfig: a genome comparison visualizer. Bioinformatics 2011;27:1099–1101.

59. Hendriksen RS, Bortolaia V, Tate H, Tyson GH, Aarestrup FM et al. Using genomics to track global antimicrobial resistance. Front Public Health 2019;7:242.

60. Al-Badr A, Al-Shaikh G. Recurrent urinary tract infections management in women. Sultan Qaboos Univ Med J 2013;13:359–367.

61. Schmiemann G, Kniehl E, Gebhardt K, Matejczyk MM, Hummers-Pradier E. The diagnosis of urinary tract infection: a systematic review. Dtsch Arzteblatt Int 2010;107:361–367.

62. Chu CM, Lowder JL. Diagnosis and treatment of urinary tract infections across age groups. Am J Obstet Gynecol 2018;219:40–51.

63. Johnson JR, Menard M, Johnston B, Kuskowski MA, Nichol K et al. Epidemic clonal groups of Escherichia coli as a cause of antimicrobial-resistant urinary tract infections in Canada, 2002 to 2004. Antimicrob Agents Chemother 2009;53:2733–2739.

64. Kudinha T, Johnson JR, Andrew SD, Kong F, Anderson P et al. Escherichia coli sequence type 131 as a prominent cause of antibiotic resistance among urinary Escherichia coli isolates from reproductive-age women. J Clin Microbiol 2013;51:3270–3276.

65. Kudinha T, Johnson JR, Andrew SD, Kong F, Anderson P et al. Genotypic and phenotypic characterization of Escherichia coli isolates from children with urinary tract infection and from healthy carriers. Pediatr Infect Dis J 2013;32:543–548.

66. Kudinha T, Johnson JR, Andrew SD, Kong F, Anderson P et al. Distribution of phylogenetic groups, sequence type ST131, and virulence-associated traits among Escherichia coli isolates from men with pyelonephritis or cystitis and healthy controls. Clin Microbiol Infect 2013;19:E173–E180.

67. Liu CM, Stegger M, Aziz M, Johnson TJ, Waits K et al. Escherichia coli ST131-H122 as a foodborne thoge. mBio 2018;9:mBio.00470–18.

68. Mathers AJ, Peirano G, Pitout JDD. The role of epidemic resistance plasmins and international high-risk clones in the spread of multidrug-resistant Enterobacteriaceae. Clin Microbiol Rev 2015;28:565–591.
69. Ewers C, Göttig S, Bülte M, Fiedler S, Tietgen M et al. Genome sequence of avian _Escherichia coli_ strain HITH25637, an extraintestinal pathogenic _E. coli_ strain of ST131 encoding colistin resistance determinant MCR-1. Genome Announc 2016;4:e00863-16.

70. Johnson TJ, Hargreaves M, Shaw K, Snippes P, Lynfield R et al. Complete genome sequence of a carbapenem-resistant extraintestinal pathogenic _Escherichia coli_ strain belonging to the sequence type 131 H30R subclade. Genome Announc 2015;3:e00272-15.

71. World Health Organization. Global Action Plan on Antimicrobial Resistance (www.who.int/antimicrobial-resistance/publications/global-action-plan/en/). Geneva: World Health Organization; 2015.

72. Pearson J, Turnidge J, Frankini C, Bell J. Australian Group on Antimicrobial Resistance. Prevalence of antimicrobial resistances in common pathogenic Enterobacteriaceae in Australia, 2004; report from the Australian Group on Antimicrobial Resistance. Commum Dis Intell Q Rep 2007;31:106–112.

73. Bell JM, Gottlieb T, Daley DA, Coombs GW. Australian Group on Antimicrobial Resistance (AGAR) Australian gram-negative sepsis outcome programme (GNSOP) annual report 2017. Commun Dis Intell Q Rep 2019;43:cdi.9156-155.

74. Kot B. Antibiotic resistance among uropathogenic _Escherichia coli_. Pol J Microbiol 2019;68:403–415.

75. Thänter R, Reske KA, Hink T, Wallace MA, Wang B et al. Comparative genomics of antibiotic-resistant uropathogens implicates three routes for recurrence of urinary tract infections. mBio 2019;10:e01977-19.

76. Mortazavi-Tabatabaei SAR, Ghaderkhanji J, Nazari A, Sayehmiri K, Sayehmiri F et al. Pattern of antibiotic resistance in urinary tract infections: a systematic review and meta-analysis. Int J Prev Med 2019;10:169.

77. Crichtley IA, Cotroneo N, Pucci MJ, Mendes R. The burden of antimicrobial resistance among urinary tract isolates of _Escherichia coli_ in the United States in 2017. PLoS One 2019;14:e0220265.

78. Mulder M, Verbon A, Lous J, Goessens W, Stricker BH. Use of other antimicrobial drugs is associated with trimethoprim resistance in patients with urinary tract infections caused by _E. coli_. Eur J Clin Microbiol Infect Dis 2019;38:2283–2290.

79. Sepulveda V, Piovonis J, Ruzauskas M, Pavilonis A, Suziediene E. Prevalence of trimethoprim resistance genes in _Escherichia coli_ isolates of human and animal origin in Lithuania. J Med Microbiol 2010;59:315–322.

80. Alcock BP, Raphenya AR, Lau TTY, Tsang KK, Bouchard M et al. CARD 2020: antibiotic resistance surveillance with the comprehensive antibiotic resistance database. Nucleic Acids Res 2020;48:D517–D525.

81. Djordjevic SP, Stokes HW, Chowdhury PR. Mobile elements, zoonotic pathogens and commensal bacteria: conduits for the delivery of resistance genes into humans, production animals and soil microbiota. Front Microbiol 2013;4:86.

82. Cummins ML, Reid CJ, Chowdhury PR, Bushell RN, Esbert N et al. Whole genome sequence analysis of Australian avian pathogenic _Escherichia coli_ that carry the class 1 integrase gene. Microb Genomics 2019;5:e000250.

83. Jarocki VM, Reid CJ, Chapman TA, Djordjevic SP. _Escherichia coli_ ST302: genomic analysis of virulence potential and antimicrobial resistance mediated by mobile genetic elements. Front Microbiol 2020;10:3098.

84. Reid CJ, Blau K, Jechalke S, Smalla K, Djordjevic SP. Whole genome sequencing of _Escherichia coli_ from store-bought produce. Front Microbiol 2020;10:03050.

85. Poirel L, Kämpfer P, Nordmann P. Chromosome-encoded ambler class A _β_-lactamase of _Klebsiella pneumoniae_, a probable progenitor of a subgroup of CTX-M extended-spectrum _β_-lactamas. Anti- microb Agents Chemother 2002;46:4038–4040.

86. Cormier A, Zhang PLC, Chalmers G, Weese JS, Deckert A et al. Diversity of CTX-M-positive _Escherichia coli_ recovered from animals in Canada. Vet Microbiol 2019;231:71–75.

87. El-Badawy MF, Tawakol WM, Maghrabi IA, Mansy MS, Shohayeb MM et al. Iodometric and molecular detection of ESBL production among clinical isolates of _E. coli_ fingerprinted by ERIC-PCR: the first Egyptian report declares the emergence of _E. coli_ O25b-ST131 clone harboring bla _Ges_. Microb Drug Resist 2017;23:703–711.

88. Yuan L, Liu J-H, Hu G-Z, Pan Y-S, Liu Z-M et al. Molecular character- ization of extended-spectrum _β_-lactamase-producing _Escherichia coli_ isolates from chickens in Henan Province, China. J Med Microbiol 2009;58:1449–1453.

89. Farajzadeh Sheikh A, Goodarzi H, Yadadjí MJ, Aslani S, Amin M et al. Virulence–associated genes and drug susceptibility patterns of uropathogenic _Escherichia coli_ isolated from patients with urinary tract infection. Infect Drug Resist 2019;12:2039–2047.

90. Ramirez-Castillo FY, Moreno-Flores AC, Avelar-González FJ, Márquez-Díaz F, Harel J et al. An evaluation of multidrug-resistant _Escherichia coli_ isolates in urinary tract infections from Aguascalientes, Mexico: cross-sectional study. Ann Clin Microbiol Anti- microbial 2018;17:34.

91. Ochoa SA, Cruz-Córdova A, Luna-Pineda VM, Reyes-Grajeda JP, Cózares-Dominguez V et al. Multidrug- and extensively drug- resistant uropathogenic _Escherichia coli_ clinical strains: phylo- genetic groups widely associated with integrons maintain high genetic diversity. Front Microbiol 2016;7:20042.

92. Li L-G, Xia Y, Zhang T. Co-occurrence of antibiotic and metal resistance genes revealed in complete genome collection. ISME J 2017;11:651–662.

93. Yang S, Deng W, Liu S, Yu X, Mustafa GR et al. Presence of heavy metal resistance genes in _Escherichia coli_ and _Salmonella_ isolates and analysis of resistance gene structure in _E. coli_ E308. J Glob Antimicrob Resist 2020;21:420–426.

94. Nguyen CC, Hugie CN, Kile ML, Navab-Daneshmand T. Association between heavy metals and antibiotic-resistant human pathogens in environmental reservoirs: a review. Front Environ Sci Eng 2019;13:46.

95. Department of Primary Industries. Central West Region Pilot Area Agricultural Profile (www.dpi.nsw.gov.au/__data/assets/pdf_file/0010/457588/Agricultural-profile-central-west-region.pdf). Sydney: NSW Government; 2012.

96. Knapp CW, McCluskey SM, Singh BK, Campbell HD, Hudson G et al. Antibiotic resistance gene abundances correlate with metal and geochemical conditions in archived Scottish soils. PLoS One 2011;6:e27300.

97. Gillingers MR. Integrons: past, present, and future. Microbiol Mol Biol Rev 2014;78:257–277.

98. Dawes FE, Kuzevski A, Bettelheim KA, Hornitzky MA, Djordjevic SP et al. Distribution of class 1 integrons with IS26-mediated deletions in their 3-conserved segments in _Escherichia coli_ of human and animal origin. PLoS One 2010;5:e12754.

99. Brolund A, Rajer F, Giske CG, Melefors Ö, Titelman E et al. Dynamics of resistance plasmids in extended-spectrum- _β_- lactamase-producing _Enterobacteriaceae_ during postinfection colonization. Antimicrob Agents Chemother 2019;63:e02201-18.

100. Toulouse JL, Edens TJ, Alejadre L, Manges AR, Pelle- tier JN. Integron-associated DfrB4, a previously uncharacterized member of the trimethoprim-resistant dihydrofolate reductase B family, is a clinically identified emergent source of antibiotic resistance. Antimicrob Agents Chemother 2017;61:e02665-16.

101. Venturini C, Hassan KA, Chowdhury PR, Paulsen IT, Walker MJ et al. Sequences of two related multiple antibiotic resistance virulence plasmids sharing a unique IS26-related molecular signature isolated from different _Escherichia coli_ pathotypes from different hosts. PLoS One 2013;8:e78862.

102. McKinnon J, Chowdhury PR, Djordjevic SP. Genomic analysis of multidrug-resistant _Escherichia coli_ ST58 causing urosepsis. Int J Antimicrob Agents 2018;52:430–435.

103. McKinnon J, Chowdhury PR, Djordjevic SP. Molecular analysis of an IncF ColV-Like plasmid lineage that carries a complex
resistance locus with a trackable genetic signature. Microb Drug Resist 2020;26:787–793.

104. Bie L, Fang M, Li Z, Wang M, Xu H. Identification and characterization of new resistance-conferring SGⅠs (Salmonella genomic island 1) in Proteus mirabilis. Front Microbiol 2018;9:03172.

105. Alonso CA, González-Barrio D, Tenorio C, Ruiz-Fons F, Torres C. Antimicrobial resistance in faecal Escherichia coli isolates from farmed red deer and wild small mammals. Detection of a multiresistant E. coli producing extended-spectrum beta-lactamase. Comp Immunol Microbiol Infect Dis 2016;45:34–39.

106. Mendes Moreira A, Couvé-Deacon E, Bouquet P, Chainer D, Jové T et al. Proteae: a reservoir of class 2 integrons? J Antimicrob Chemother 2019;74:1156–1158.

107. Mora-Rillo M, Fernández-Romero N, Navarro-San Francisco C, Díez-Sebastián J, Romero-Gómez MP et al. Impact of virulence genes on sepsis severity and survival in Escherichia coli bacteremia. Virulence 2019;6:93–100.

108. Bonacorsi S, Houdouin V, Mariani-Kurkdjian P, Mahjoub-Messai F, Bingen E. Comparative prevalence of virulence factors in Escherichia coli causing urinary tract infection in male infants with and without bacteremia. J Clin Microbiol 2006;44:1116–1158.

109. Lefort A, Panhard X, Clermont O, Woerther P-L, Branger C et al. Host factors and portal of entry outweigh bacterial determinants to predict the severity of Escherichia coli bacteremia. J Clin Microbiol 2011;49:777–783.

110. Jauréguy F, Carbonnelle E, Bonacorsi S, Clec’h C, Casassus Pet al. Host and bacterial determinants of initial severity and outcome of Escherichia coli sepsis. Clin Microbiol Infect 2007;13:854–862.

111. Brumbaugh AR, Smith SN, Mobley HL. Immunization with the yersiniabactin receptor, FyuA, protects against pyelonephritis in a murine model of urinary tract infection. Infect Immun 2013;81:3309–3316.

112. Wright KJ, Hultgren SJ. Sticky fibers and uropathogenesis: bacterial adhesins in the urinary tract. Future Microbiol 2006;1:75–87.

113. Majilovic H, Mac Aogán M, Collins CJ, Rogers TR, Smith SGJ. Characterization of Escherichia coli bloodstream isolates associated with mortality. J Med Microbiol 2016;65:71–79.

114. Dale AP, Pandey AK, Hesp RJ, Belogiannis K, Laver JR et al. Genomes of Escherichia coli bacteremia isolates originating from urinary tract foci contain more virulence-associated genes than those from non-urinary foci and neutropenic hosts. J Infect 2018;77:534–543.

115. Croxen MA, Finlay BB. Molecular mechanisms of Escherichia coli pathogenicity. Nat Rev Microbiol 2010;8:26–38.

116. Chowdhury PR, McKinnon J, Liu M, Djordjevic SP. Multidrug resistant uropathogenic Escherichia coli ST405 with a novel, composite IS26 transposon in a unique chromosomal location. Front Microbiol 2018;9:3212.

117. Johnson JR, Johnston BD, Porter SB, Clabots C, Bender TL et al. Rapid emergence, subsidence, and molecular detection of Escherichia coli sequence type 1193-fimH64, a new disseminated multidrug-resistant commensal and extraintestinal pathogen. J Clin Microbiol 2019;57:e01664–18.

118. Blanco J, Mora A, Maman R, López C, Blanco M et al. Four main virotypes among extended-spectrum-β-lactamase-producing isolates of Escherichia coli O25b:H4-B2-ST131: biological, epidemiological, and clinical characteristics. J Clin Microbiol 2013;51:3358–3367.

119. Olesen B, Scheutz F, Andersen RL, Menard M, Boisen Net al. Enteroaggregative Escherichia coli 078:H10, the cause of an outbreak of urinary tract infection. J Clin Microbiol 2012;50:3703–3711.

120. Boll EJ, Stegger M, Hasman H, Roer L, Overballe-Petersen S et al. Emergence of enteroaggregative Escherichia coli within the ST131 lineage as a cause of extraintestinal infections. Front Microbiol 2017;8:146.

121. Lara FBM, Nery DR, de Oliveira PM, Araujo ML, Carvalho FRQ et al. Virulence markers and phylogenetic analysis of Escherichia coli strains with hybrid EAEC/UPEC genotypes recovered from sporadic cases of extraintestinal infections. Front Microbiol 2018;9:3594.

122. Boll EJ, Struve C, Boisen N, Olesen B, Stahlhut SG et al. Role of enteroaggregative Escherichia coli virulence factors in uropathogenesis. Infect Immun 2013;81:1164–1171.

123. Dobrindt U, Chowdary MG, Krumbholz G, Hacker J. Genome dynamics and its impact on evolution of Escherichia coli. Med Microbiol Immunol 2010;199:145–154.

124. Calhau V, Ribeiro G, Mendonça N, Da Silva GJ. Prevalent combination of virulence and plasmidic-encoded resistance in ST 131 Escherichia coli strains. Virulence 2013;4:726–729.

125. Rodriguez-Siek KE, Giddings CW, Doekkert C, Johnson TJ, Nolan LK. Characterizing the APEC pathotype. Vet Res 2005;36:241–256.

126. Johnson TJ, Wannemuehler Y, Johnson SJ, Stell AL, Doekkert C et al. Comparison of extraintestinal pathogenic Escherichia coli strains from human and avian sources reveals a mixed subset representing potential zoonotic pathogens. Appl Environ Microbiol 2008;74:7043–7050.

127. Gérard F, Pradel N, Wu LF. Bacterial activity of colicin V is mediated by an inner membrane protein, SdaC, of Escherichia coli. J Bacteriol 2005;187:1945–1950.

128. Chattopadhyay S, Weissman SJ, Minin VN, Russo TA, Dykhui-zen DE et al. High frequency of hotspot mutations in core genes of Escherichia coli due to short-term positive selection. Proc Natl Acad Sci USA 2009;106:12412–12417.

129. Denamur E, Bonacorsi S, Giraud A, Duriez P, Hilali F et al. High frequency of mutator strains among human uropathogenic Escherichia coli isolates. J Bacteriol 2002;184:605–609.

130. Chen SL, Hung C-S, Xu J, Reigstad CS, Magrini V et al. Identification of genes subject to positive selection in uropathogenic strains of Escherichia coli: a comparative genomics approach. Proc Natl Acad Sci USA 2006;103:5977–5982.

131. Nielsen KL, Stegger M, Godfrey PA, Feldgarden M, Andersen PS et al. Adaptation of Escherichia coli traversing from the faecal environment to the urinary tract. Int J Med Microbiol 2013;306:595–603.

132. Eddy SR. Where did the BLOSUM62 alignment score matrix come from? Nat Biotechnol 2004;22:1035–1036.

133. Ragheb MN, Thomason MK, Hsu C, Nugent P, Gage J et al. Inhibiting the evolution of antibiotic resistance. Mol Cell 2019;73:157–165.

134. Selby CP, Mfd protein and transcription-repair coupling in Escherichia coli. Photochem Photobiol 2017;93:280–295.

135. Thorpe HA, Bayliss SC, Hurst LD, Feil EJ. Comparative analyses of selection operating on nontranslated intergenic regions of diverse bacterial species. Genetics 2017;206:363–376.

136. Guttmann DS, Dykhuizen DE. Clonal divergence in Escheri-chia coli as a result of recombination, not mutation. Science 1994;266:1380–1383.