Global Phylogeny and F Virulence Plasmid Carriage in Pandemic Escherichia coli ST1193

Ethan R. Wyrsch,a Rhys N. Bushell,b Marc S. Marenda,b Glenn F. Browning,b Steven P. Djordjevic,a

aAustralian Institute for Microbiology & Infection, University of Technology Sydney, Ultimo, New South Wales, Australia
bAsia-Pacific Centre for Animal Health, Department of Veterinary Biosciences, Melbourne Veterinary School, Faculty of Veterinary and Agricultural Sciences, University of Melbourne, Parkville, Victoria, Australia

ABSTRACT Lower urinary tract, renal, and bloodstream infections caused by phylogroup B2 extraintestinal pathogenic Escherichia coli (ExPEC) are a leading cause of morbidity and mortality. ST1193 is a phylogroup B2, multidrug-resistant sequence type that has risen to prominence globally, but a comprehensive analysis of the F virulence plasmids it carries is lacking. We performed a phylogenomic analysis of ST1193 (n = 707) whole-genome sequences from EnteroBase using entries with comprehensive isolation metadata. The data set comprised isolates from humans (n = 634 [90%]), including 339 (48%) from extraintestinal infection sites, and isolates from companion animals, wastewater, and wildlife. Phylogenetic analyses combined with gene detection and genotyping resolved an ST1193 clade structure segregated by serotype and F plasmid carriage. Most F plasmids fell into one of three related plasmid subtypes: F2:A1:B10 (n = 444 [65.97%]), F2:A1:B1 (n = 84 [12.48%]), and F2:A1:B20 (n = 80 [11.89%]), all of which carry the virulence genes cjrABC colocalized with senB (cjrABC-senB), a trademark signature of F29:A2:B10 subtype plasmids (pUTI89). To examine the phylogenetic relationship of these plasmids with pUTI89, complete sequences of F2:A1 and F2:B20 plasmids were resolved. Unlike pUTI89, the most dominant and widely disseminated F plasmid that carries cjrABC-senB, F plasmids in ST1193 often carry a complex resistance region with an integron truncation (intI1\_245) signature embedded within a structure assembled by IS26. Plasmid analysis shows that ST1193 has F plasmids that carry cjrABC-senB and ARG-encoding genes but lack tra regions and are likely derivatives of pUTI89. Further epidemiological investigation of ST1193 should seek to confirm its presence in human-associated environments and identify any potential agricultural links, which are currently lacking.

IMPORTANCE We have generated an updated ST1193 phylogeny using publicly available sequences, reinforcing previous assertions that Escherichia coli ST1193 is a human-associated lineage, with many examples sourced from human extraintestinal infections. ST1193 from urban-adapted birds, wastewater, and companion animals are frequent, but isolates from animal agriculture are notably absent. Phylogenomic analysis identified several clades segregated by serogroup, all noted to carry highly similar F plasmids and antimicrobial resistance (AMR) signatures. Investigation of these plasmids revealed virulence regions with similarity to pUTI89, a key F virulence plasmid among dominant pandemic extraintestinal pathogenic E. coli lineages, and encoding a complex antibiotic resistance structure mobilized by IS26. This work has uncovered a series of F virulence plasmids in ST1193 and shows that the lineage mimics the host range and virulence attributes of other E. coli strains that carry pUTI89. These observations have significant ramifications for epidemiological source tracking of emerging and established pandemic ExPEC lineages.

KEYWORDS antimicrobial resistance, Escherichia coli, integron, pandemic, plasmids, virulence
Sequence type 1193 (ST1193) is a fluoroquinolone-resistant lineage of *Escherichia coli* phylogroup B2, within the ST14 clonal cluster (1, 2). Its closest relative is *E. coli* ST6460 (3). ST1193 is considered a hybrid lineage and is estimated to have arisen in 2005 by acquisition, via conjugation, of a single 976-kb chromosomal fragment from a distantly related ST10 *fimH*54 strain of *E. coli*. The conjugation event is remarkable because subsequent recombination events resulted in acquisition of multiple mutations in *gyrA* and *parC* and consequent resistance to fluoroquinolones (3–5). ST1193 isolates’ shared features include three chromosomal mutations conferring fluoroquinolone resistance, a *fimH*64 allele, and a non-lactose-fermenting phenotype (2).

ST1193 is known to carry diverse plasmid cargo, including large multidrug resistance plasmids (6), and the first carbapenem nonsusceptible ST1193 isolates carrying *bla*<sub>NDM</sub> (7) and *bla*<sub>KPC</sub> (8) were reported recently. The sequence type is remarkable because it has risen to prominence in a short period of time and has been recovered from multiple hosts, including humans (9, 10), dogs (11, 12), synanthropic birds (13, 14), and water sources (6). While it is likely that humans are an important gastrointestinal reservoir for ST1193, dogs are considered significant in the human-companion animal infection chain (1).

Reports of infections with ST1193 surfaced in 2012 from health care settings in Germany (15), England (16), France (17), Canada (18), China (10, 19, 20), the United States (21), South Korea (9), and Australia (11, 12, 22), raising awareness of its importance as a rapidly emerging human pathogen, but few studies have examined the lineage from a One Health perspective. In humans, ST1193 strains are a cause of urinary tract (3), bloodstream (9, 18, 21), and other extraintestinal and invasive infections, and many are multidrug resistant. In China, ST1193 is a cause of invasive infections in neonates, with most of the invasive isolates being resistant to ciprofloxacin, amoxicillin, sulfonamides, and tetracycline (10). Notably, a third of the invasive ST1193 isolates studied had acquired *bla*<sub>CTX-M-15</sub>, which encodes an extended spectrum of resistance to β-lactams (10). ST1193 can also persistently colonize the gastrointestinal tract of healthy women (3) but was not identified in the feces of healthy women in France prior to 2010 (23).

Australia was one of the first places to report ST1193 as a cause of extraintestinal disease in humans and dogs (11). Apart from being fluoroquinolone resistant, the Australian ST1193 isolates belonged to serogroup O75 and clonal cluster 14, displaying resistance to multiple antimicrobials, including piperacillin, gentamicin, tetracycline, and trimethoprim-sulfamethoxazole (11). The presence of these antimicrobial resistance genes (ARGs) indicates that ST1193 likely hosts class 1 integron structures located within complex resistance regions that are assembled by insertion elements, particularly IS<sub>26</sub> (24), which play important roles in the mobilization and capture of ARGs within *E. coli*.

One of the key influencers of extraintestinal pathogenicity in *E. coli* is F plasmid carriage, an assertion supported by the repeated observations of key virulence factors, particularly siderophores, which assist in iron capture during urinary tract infection (UTI) (25, 26), being mobilized by F family plasmids (27–30). An F plasmid replicon sequence typing (RST) scheme (31) allows subtyping of F plasmids from whole-genome sequence data, primarily with typing of RNAI-FI (designated “F”), *repFIA* (“A”), and *repFIB* (“B”) alleles, reported as F:A:B. Two particular groups of F virulence plasmid are closely associated with pandemic extraintestinal pathogenic *E. coli* (ExPEC) lineages: F29:A:B10 plasmids, typified by pUTI89 and carrying the predicted iron uptake system genes *cjrABC-senB* (26), and the comparatively broader family of ColV plasmids, characterized by the presence of numerous virulence operons (colicin *[cva]*, salmochelin *[iro]*, aerobactin *[iuc]*, ets, *sit*, *ompT*, and *hlyF*) (32), again primarily iron uptake systems, and typed frequently as F2, F18, and F24 plasmids (33). F29:A:B10 is one of the most frequently observed F RSTs and is widespread in ExPEC isolates from humans, wastewater, and companion and wild animals but is conspicuously missing in ExPEC from animal agriculture (28). ColV plasmids are also widespread in ExPEC isolates from diverse sources and have a notable link with animal agriculture, particularly poultry.
and swine. Several chromosomal islands are also significant contributors to pathogenicity in UTI models (34), particularly the high-pathogenicity island (HPI) carrying the yersiniabactin siderophore genes (irp and fyuA) (35) and the colibactin operon (36).

The perception of public health is being impacted by One Health approaches to infectious disease (37). Here, we undertook a comprehensive phylogenetic analysis of 707 E. coli ST1193 isolates from different countries and multiple sources, contributing two completed sequences from Australian companion animals and providing insight into the role of F plasmids in the success of E. coli ST1193 as a pandemic pathogen. Collectively, our analysis has consequences for how emerging Escherichia coli pathogens are identified and traced.

RESULTS

The genome sequences of ST1193 isolates (n = 707) depicted here were retrieved from EnteroBase and were selected based on having available isolation source, collection year, and country of isolation metadata. A further 553 genomes were excluded because metadata were incomplete. Antimicrobial resistance (AMR) and virulence genotypes, F plasmid RST subtypes, and carriage of the class 1 integrase gene intI1 and insertion sequence IS26 were also included. The intI1 allele used for this analysis can be found under accession number MZ396394.1. Using a different intI1 sequence can alter the reported hit sizes for this analysis by several base pairs, depending on the insertion site of IS26. The complete data set is available in Table S1 in the supplemental material.

Geographically, the isolates were from 32 countries, with Australia (n = 259 [36.5%]) and the United States (n = 135 [19.0%]) being the most prominent contributors of sequence data. Isolates were primarily from humans (n = 634 [89.7%]), frequently with a clinical presentation. The remainder were from companion animals (canine and feline), wild avian sources, wastewater, wastewater treatment plants (WWTP), and other water sources. As of 31 August 2022, EnteroBase included approximately 44,000 entries reported from livestock, animal feed, and food products, and despite this wealth of data, ST1193 genomes from agricultural sources were notably absent outside a single entry from bovine livestock. This isolate was excluded from analysis due to a lack of metadata (Table S2). The earliest available ST1193 sequence was from an Australian isolate obtained in 2007. Table 1 presents key metadata and genetic features of ST1193 isolates from countries that contributed at least 10 isolates.

A phylogenetic tree based on single nucleotide variations (SNVs) was generated and aligned with isolate metadata and genotyping (Fig. 1). The clade structure segregated according to serotype (where serotype could be identified), with clusters comprising types O75:H5 (n = 676), O45:H5 (n = 5), and O25:H5 (n = 13), and clade O18:H5 (n = 10) identified. ST1193 isolates in serogroup O75:H5 were the dominant lineage. Sublineages within this serogroup fell into three groups associated with F plasmid RST. The majority of O75:H5 clade isolates, as well as O6:H5 and O45:H5 clade isolates, were predicted to carry F-:A1:B10 plasmids. F-:A1:B1 plasmids were identified in O18:H5 and O25:H5 isolates, as well as a subclade of O75:H5, and sporadically in other clades. F-:A1:B20 plasmid carriage was generally limited to a discrete subclade of the O75:H5 isolates. In all cases, these differently typed subclades comprised isolates from multiple nations and sources, indicating that at least seven (eight if the distant O75:H5 group is included) discernible subgroups of ST1193 are in circulation globally.

While 673 of 707 isolates (95%) carried an F plasmid sequence represented by 34 plasmid subtypes, 29 of these subtypes were detected in five or fewer isolates. F-:A1:B10 plasmids were identified in 444/673 (65.97%) isolates, followed by F-:A1:B1 (n = 84; 12.48%) and F-:A1:B20 (n = 80; 11.89%) plasmids, and these three plasmid replicon types represented 90.34% of all isolates carrying an F plasmid. Many of the remaining plasmid subtypes appear to be related to this set of three plasmid RSTs, often having a novel FII RST subtype or lacking the FIA subtype (e.g., F95:A1:B20 or F-:A-:B10). Notably, 591 isolates carried the virulence factor gene senB, with BLASTn data
### TABLE 1 Key features of *Escherichia coli* ST1193

| Country (n) | Source(s) (n) | e-serotypes (n) | Isolation date(s) | intI1 hits (bp) | F plasmid RSTs (n)
|-------------|---------------|-----------------|-------------------|-----------------|------------------|
| Australia (259) | Human (237), wild avian (12), canine (10) | O75:H5 (118), O18:H5 (7), O6:H5 (5), O-H5 (129) | 2007–2020 | 1,014 (5), D746 (108), D850 (12), D798 (2), D682 (2), D548 (2), other (3) | F :A1:B10 (170), F :A1:B20 (46), F :A1:B3 (24), F :A1:B10 (3), F :A1:B2 (2), other (8) |
| Canada (41) | Human (38), canine (1), water (2) | O75:H5 (34), O-H5 (7) | 2010–2019 | 1,014 (12), D746 (13), other (1) | F :A1:B10 (27), F :A1:B1 (6), other (4) |
| China (46) | Human (46) | O75:H5 (40), O18:H5 (1), O25:H5 (3), O-H5 (2) | 2011–2018 | Δ746 (21), other (3) | F :A1:B10 (20), F :A1:B1 (12), F :A1:B20 (2), F12:A1:B10 (2), other (4) |
| Denmark (13) | Human (13) | O75:H5 (9), O-H5 (4) | 2018 | 1,014 (5), Δ746 (3) | F :A1:B10 (7), F :A1:B20 (2), other (4) |
| France (12) | Human (10), canine (1), feline (1) | O75:H5 (11), O-H5 (1) | 2014–2020 | Δ746 (9), other (1) | F :A1:B10 (9), F :A1:B20 (2), other (1) |
| Japan (11) | Human (4), water (7) | O75:H5 (9), O-H5 (2) | 2015–2018 | 1,014 (1), Δ746 (4) | F :A1:B10 (5), F :A1:B20 (2), F :A1:B1 (2), other (1) |
| New Zealand (23) | Human (23) | O75:H5 (16), O-H5 (7) | 2015–2016 | 1,014 (1), Δ746 (8), other (1) | F :A1:B10 (20), other (3) |
| Qatar (11) | Human (11) | O75:H5 (10), O-H5 (1) | 2016–2018 | 1,014 (1), Δ850 (4), other (1) | F :A1:B10 (8), F :A1:B2 (2), other (4) |
| Singapore (36) | Human (34), ND (2) | O75:H5 (24), O6:H5 (2), O-H5 (10) | 2012–2018 | 1,014 (7), Δ746 (11), Δ850 (2), Δ682 (2), other (1) | F :A1:B10 (28), F :A1:B1 (3), other (4) |
| Switzerland (14) | Human (11), water (2), ND (1) | O75:H5 (9), O6:H5 (1), O-H5 (4) | 2014–2020 | 1,014 (1), Δ746 (5), other (3) | F :A1:B20 (7), F :A1:B10 (5), other (2) |
| Thailand (16) | Human (16) | O75:H5 (11), O45:H5 (4), O18:H5 (1) | 2014–2018 | 1,014 (2), Δ746 (10), other (1) | F :A1:B10 (12), other (2) |
| United Kingdom (15) | Human (15) | O75:H5 (13), O18:H5 (1), O-H5 (1) | 2010–2021 | 1,014 (4), Δ746 (2), other (2) | F :A1:B10 (8), F :A1:B20 (2), F35: A1:B10 (2), other (2) |
| United States (135) | Human (125), canine (7), ND (3) | O75:H5 (66), O-H5 (69) | 2009–2021 | 1,014 (34), Δ746 (20), Δ667 (3), Δ666 (15), other (7) | F :A1:B10 (73), F :A1:B1 (28), F :A1:B10 (10), F :A1:B20 (7), other (8) |
| Vietnam (34) | Human (12), ND (22) | O75:H5 (19), O-H5 (15) | 2014–2018 | 1,014 (1), Δ746 (28), other (1) | F :A1:B10 (26), F :A1:B20 (3), F :A1:B1 (2), other (3) |

*Only counts of 10 or greater are included.
**ND, not described.
***Those with counts of 1 are grouped as ‘other,’ with 1,014-bp intI1 being an exception. Note that the 746-bp hit reported has a corrected annotation size of 745 bp.*
confirming colocalization with \textit{cjrABC} (\textit{cjrABC-senB}), as was originally described and characterized in \textit{pUTI89} (F29:2-B10). Within the whole collection, only one isolate encoding \textit{senB} lacked an F plasmid, consistent with observations of \textit{senB} being primarily distributed on F plasmids together with \textit{cjrABC}. Interestingly, just two isolates were predicted to carry an F29:2-B10 plasmid.

Siderophore genes detected included \textit{fyuA} (n = 703) and \textit{irp2} (n = 690), which are associated with the \textit{Yersinia} HPI, and the \textit{sit} operon (n = 705), which is associated with
The iut-iuc (aerobactin) operon ($n = 694$). The iro operon was detected in only four isolates. Thirty-three isolates from multiple serogroups were found to carry the colibactin/pks operon ($clb$), and 23 isolates carried complete $pap$ operons, with most isolates carrying at least some $pap$ genes (Table S1). All isolates tested negative for carriage of $ColV$, according to the criteria of Liu et al. (38), indicating that ST1193 isolates are not carriers of these important ExPEC-associated F virulence plasmids. The absence of $ColV$ F plasmids is consistent with the observation that ST1193 does not circulate in food production animals (28).

Antimicrobial resistance genes and class 1 integron structures were widely distributed in ST1193. Most isolates (685 [96.9%]) carry IS26, an element strongly associated with the capture and mobilization of antibiotic resistance genes. In total, 467 isolates (66%) carried a class 1 integron. In Fig. 1, BLASTn hit data for $intI1$ are presented to highlight integrons that are either truncated or broken due to whole-genome sequencing (WGS) assembly. A complete copy of $intI1$ (1,014 bp) was detected in 133 (28.5%) isolates. Notably, 256 (54.8%) isolates carried a 745-bp truncated copy of $intI1$ ($intI1^{D}_{745}$), which appeared as a BLASTn hit size of 746 bases.

To explore the structures of the most prevalent F plasmids within ST1193 and simultaneously resolve complex resistance structures hosting the $intI1^{D}_{745}$ element, we performed comparative analyses of five closed F plasmid sequences. Three hosted the integron under investigation: pAVS0096-a ($F^{-}$:A1:B10; CP076345.1), as well as pMVC530_A ($F^{-}$: A1: B1; JAMQKE000000000) and pMVC465_A ($F^{-}$: A1: B2; JAMQKF000000000) from strains MVC530 and MVC465, respectively (39). Two additional relevant plasmids were included in this analysis, pNMEC-O75A (ST1193; $F^{-}$:A1:B10; CP030112.1) and pUTI89 (ST95; F29:A2:B10; CP000244.1). Figure 2 depicts annotations and BLASTn comparisons of these plasmids.

Regarding the complex integron structure, the three resolved examples were variants of the same integron typed by the presence and location of sul1 and mphA, but a Tn3-family transposon, which typically mobilizes class 1 integrons (40), was not observed in the annotations. The structure, encompassing the integron and linked resistance gene cargo, appeared to be assembled and mobilized by IS26, with a terminal copy of IS26 truncating $intI1$. The integron hosted dfrA17 and adaA5 cassettes. A second tandem IS26-associated resistance region
carried sul2, strAB and tetAR, and a third carried blaTEM and aacC2 in the cases of pMVC370_A and pMVC465_A or blCTX,M-27 in pAVS0096-a. Another closed plasmid from ST1193, pNMEC-O75A, lacked the integron and sul2 modules, carrying only blaTEM and aacC2 (Fig. 2). Differences within observed gene counts in the overall ST1193 data set, such as dfrA17 (n = 417) and aadA5 (n = 273), or sul1 (n = 287) and mphA (n = 394), suggests that the structures we resolved here represent only some of the variation within the complex resistance structures to be found in ST1193.

Outside the aforementioned ARGs, differing integron gene cassette profiles and blCTX,M alleles were observed both sporadically within the data set and associated with small clusters of isolates. The occasional isolate was also positive for blCMY (n = 29), blCTX,A (n = 62), and/or tetB (n = 170). Low counts and multiple alleles of blaIMP (n = 2), blASP (n = 11), blAGMOM (n = 10), and blAGRh (n = 1) were identified (Table S1). Isolates lacking an F plasmid (n = 34) also often lacked most ARGs, and similarly, all isolates lacking IS26 (n = 22) also lacked an F plasmid, indicative of links between F plasmid carriage and mobilized multiple-drug-resistance carriage in ST1193. All isolates encoded the gyrA fluoroquinolone resistance mutations S83L and D87N. Investigation into blCTX,M (n = 325) showed three primary alleles: CTX-M-27 (n = 152), CTX-M-15 (n = 111), and CTX-M-55 (n = 38). Notably, the latter two alleles were colocalized with complete copies of ISEcp1, while CTX-M-27 was contextualized alongside ISEcp1 elements truncated by IS26, with IS26 bordering both ends of investigated contigs. Completed sequences will be required to fully resolve the context of blCTX,M in ST1193 and to determine if CTX-M-27 is now being mobilized by IS26 activity, as seemingly observed in pAVS0096-a.

The four senB-bearing plasmids depicted in Fig. 2 are representatives of the elements that were widely distributed throughout the ST1193 phylogeny (n = 606 [86%]) and in all serogroups. A linear comparison with plasmid pUTI89 is presented in Fig. 2, highlighting that not only is senB conserved between the plasmids, but also the entirety of the virulence gene region of the pUTI89 plasmid, as well as a short region of uncharacterized ORFs, is conserved as well. This virulence region is typically split into three sections (i) encoding copper and iron transporters, (ii) encoding iron-associated uncharacterized ORFs, and (iii) encoding a putative colicin J sensitivity (cjrABC) and carrying senB. These regions, including the diverse array of IS elements found within them, were present in all four plasmids, suggesting that F-:A1:B1/B10/B20 plasmids are closely related, but with distinct plasmid backbones. In pAVS0096-a and pMVC465_A, it was noted that the region carrying cjrABC-senB was reversed between IS3 family elements. The comparison also highlighted further similarity between F-:A1:B1 and F+: A1:B10 plasmids, with the novel F-:A1:B20 backbone region encoding a type 1 restriction modification system in place of a set of IS elements and VapBC toxin/antitoxin systems encoded by the B1/B10 plasmids.

To observe whether these plasmid assemblies were relevant to the WGS data, pAVS0096-a, pMVC465_A, pMVC370_A, and pUTI89 were aligned against representative F-:A1:B1, B10, and B20 plasmid subtypes found in ST1193 whole-genome sequences. A single representative from each of the 14 most prevalent countries of origin was included for each subtype, where available. The BRIG alignments with pUTI89 (Fig. 3A) confirmed the absence of the plasmid transfer region and typically showed full coverage of the virulence regions and their flanking sequence. The alignments demonstrated high conservation of their respective references in B1 and B10 subtype plasmids (Fig. 3B and C), and in B20 isolates overall, but variability across the B20 backbone region (Fig. 3D) was noted. Coverage of the antimicrobial resistance structure was variable, as expected given the promiscuous nature of IS26.

To provide additional resolution of the dominant O75:H5 sublineage of ST1193, a phylogenetic tree of all Australian representatives (n = 243) was generated (Fig. 4), excluding the distant group highlighted in Fig. 1 to increase average WGS inclusion up to 85% for the analysis (compared to 50% using the global collection against an ST14 reference). The major clustering again split most isolates based on F plasmid subtypes,
with isolates hosting B20 alleles forming the most distant clade of the data set at approximately 420 SNVs from the reference ESC_FB3738AA_AS. A single isolate typed as B1 was linked to this group. Isolates hosting B10 then formed the primary cluster of isolates at an average of 60 SNVs difference, with three subclades of B1 and one of B20 forming alongside them, ranging from 60 to 90 SNVs. Noticeable clustering, highlighted in Fig. 4, supported One Health associations, with as few as 3 SNVs between the closest pairs of isolates. These connections were observed between humans and companion animals (C1 and C2) or humans and wild avian species (C5, C6, C8, and C10). A pair of isolates obtained 4 years apart (C7) were noted to have truncated intI1 BLASTn signatures of 682 bp, and a larger clade (C10) had truncated intI1 hits of 850 bp. Similarly, cluster C9 had some isolates with a truncated intI1 hit size of 548 bp, another example of which can be seen in C10. Taken together, these data indicate that the lineage has undergone clonal expansion while hosting the intI1_D745 truncation on presumably an F2:A1:B10 plasmid, followed by diversification of the lineage, rather than arising through multiple capture events.

A smaller international clade was resolved in a separate phylogeny based around the HierCC H10 cluster 11740 (Fig. 5). This was the second largest H10 cluster in the collection, selected because it had numerous internal H5 groupings and included some of the earliest isolates sequenced. This FIB10 sublineage cluster comprised only human isolates, except one from a wild Australian bird, and the isolates were obtained at significantly different times and locations. Sporadic variations in integrons, AMR profiles, and senB carriage were occasionally associated with loss of the dominant F plasmids, indicating again that ST1193 appears to have gone through clonal expansion.
mobilizing the intI1,4745 as part of the F plasmid structure, followed by diversification and plasmid loss.

In summary, ST1193 is a pandemic lineage composed of several serogroups, and while there was observable phylogenetic distance between the serogroups, all clades included isolates with virulence plasmids of subtypes F-:A1:B1, B10, or B20, with further subgroupings of F plasmid types within the major O75:H5 cluster. These plasmids shared virulence content with pUTI89, a plasmid associated with globally disseminated E. coli uropathogens, but with a clear association with specific antimicrobial resistance structures mobilized by IS26.
Here, we describe a detailed phylogenetic analysis of 707 ST1193 genome sequences from 32 countries. Isolates from Australia (259 genomes) and the United States (135 genomes) contributed more than half (394 genomes [56%]) of the collection. The primary source was humans; however, data from companion animals (canine and feline), wastewater, water, and wild birds are indicative of anthropogenic transmission. Isolates from poultry, pigs, cattle, and other food animals are currently absent from public databases. While this may be due to sampling bias, this is a most unlikely scenario. As stated earlier, there are over 44,000 *E. coli* genomes representing 3,737 sequence types sourced from agricultural animals and related sources in EnteroBase, and ST1193 appears only once with a livestock source attribution. We did not find ST1193 in over 250 Australian *E. coli* isolates of porcine origin (41, 42), nor was it detected in *E. coli* from Australian poultry with symptoms of colibacillosis (29). In addition, we have recently reported that *E. coli* isolates that carry pUTI89 (F29:A2:B10), the *F* plasmid replicon sequence type that is frequently found in pandemic ExPEC lineages, are rarely detected among collections of agricultural isolates (28). The weight of evidence supports the contention that ST1193, and *E. coli* isolates hosting *F* plasmids harboring cjrABC-sen*B* in general, are unable to successfully colonize or persist in agricultural animals. The mechanism(s) that underpins host bias is poorly understood, however.

The overwhelming presence of ST1193, often associated with urinary tract infections, in human samples underscored the lineage’s pathogenic nature. Understanding the genetic factors that may have helped its success is crucial for identifying future potential ExPEC pathogens. Chromosomal virulence genes were consistently found across the collection, with a high prevalence of the yersiniabactin ferric uptake receptor gene (*fyuA*) and biosynthetic protein gene *irp2*, markers for the *Yersinia* HPI (43), and the *sit* operon. These genes are considered reliable markers of pathogenic potential in *E. coli*.
(35). Some other virulence attributes, such as the colibactin and pap operons, were also sporadically detected, an indication that the pathogenic potential of the lineage may vary and that despite the overwhelming carriage of key virulence plasmids and the HPI, the presence of other virulence elements may define the importance of specific ST1193 sublineages in the future.

The distribution of phylogenetic clusters within ST1193 matched the different serotypes identified, expanding on earlier observations conducted on a smaller cohort of isolates (2). A similar observation was reported for pandemic ExPEC lineage ST73 (44), highlighting the need for epidemiological investigations beyond the level of resolution that sequence typing can provide. Serotype O75:H5 dominated the ST1193 collection, but O45:H5, O25:H5, O18:H5, and O6:H5 isolates were just as clearly associated with the described virulence and AMR attributes. This suggests that all serogroups descended from a single common ancestor. All serotypes had a H5 flagellar antigen, but a substantial proportion of the collection (n = 264 [37%]) were O-nontypeable:H5, again reinforcing the need for further investigation (like HeirCC or phylogenetic analysis) to place these isolates appropriately.

Our observations here on ST1193 expand our understanding of F virulence plasmid variation, host range, and AMR carriage. ST1193 is a significant host of plasmids with similarities to pUTI89, though with heavily altered plasmid backbone sequences (including lost transfer and stability portions in comparison to pUTI89), giving rise to different F RSTs which are with association to cjABC-senB (2). The senB-encoding plasmids pAVS0096-a (F⁻:A1:B10), pMVC465 (F⁻:A1:B20), and pMVC530 (F⁻:A1:B1) are representatives of plasmids that are widely distributed throughout the ST1193 phylogeny and are hosted in all serogroups within the lineage, indicating that it was carried by the progenitor strain of the lineage. Comparisons with pUTI89 (F29:A⁻:B10) showed the virulence gene region and surrounding genetic environment are highly conserved in all plasmids, including mobile elements. However, unlike pUTI89 (F29:A⁻:B10), which often carries a single copy of IS26, these plasmids carry a complex resistance gene region (CRR) and often have a specific resistance gene structure containing a class 1 integron with a truncated intI1, mobilized solely by IS26. These observations led us to hypothesize that pAVS0096-a (F⁻:A1:B10), pMVC465 (F⁻:A1:B20), and pMVC530 (F1:A1:B1) are potential relatives or derivatives of pUTI89, but with distinct plasmid backbones, that have emerged within ST1193.

Similar to the pandemic lineages ST127 (45), ST95 (28, 46), and ST131 (32), which carry pUTI89, ST1193 isolates carry senB⁺ plasmids and are not found in poultry, swine, or cattle but rather are widely found in humans, dogs, cats, wastewater, and wildlife, particularly urban birds. Furthermore, all plasmids under analysis carried a copy of IS26 in the same position, abutting copper sensitivity suppression open reading frames (ORFs), although this position appears to be variable due to repeated IS26 activity. In addition, alignment of the four plasmid sequences showed that pAVS0096-a (F⁻:A1:B10), pMVC465 (F⁻:A1:B20), and pMVC530 (F1:A1:B1) share considerable sequence identity with pUTI89 but that the entire transfer (tra) module in pUTI89 is missing from the F plasmids found in ST1193. It is conceivable that the loss of the tra region in these ST1193 F virulence plasmids enabled the acquisition and stable integration of the CRR into the plasmid backbone. Stephens et al. (46) hypothesized that pUTI89, possibly in concert with chromosomal genes, may inhibit lineages of ST95 from acquiring other plasmid cargo, limiting their ability to acquire antimicrobial resistance. Consistent with this hypothesis, extensive phylogenomic analyses of ST95 (n = 668) showed that lineages that carry pUTI89 (F29:A⁻:B10) are pan-sensitive and lack evidence of class 1 integrons (28). It is not known if the F plasmids in ST1193 do the same, although carriage of additional plasmids in ST1193 isolates was noted, with the second most prominent plasmid subtype, I1, detected in 17% of isolates. These observations suggest that the genetic modifications found in the F plasmids hosted by ST1193 may enable the lineage to acquire other plasmid families. Further studies should decipher these evolutionary developments and seek evidence of these plasmids in other E. coli lineages, as...
they encode combinations of important virulence and antibiotic resistance gene cargo and pose an elevated threat to human health.

Distribution of ARGs within the data set was quite variable, but some key genotypes alongside truncations of intI1, caused by IS26, served as prominent markers that highlight the types of complex resistance structures found within ST1193. The key feature of the most prevalent integron, identified by the conserved 745-bp truncation of intI1, was that it was not associated with Tn21, the classical mobilizer of class 1 integrons associated with sul1 and mphA (40), and were instead mobilized by IS26 (47, 48). The variable presence of ARGs suggests that plasmids carried by ST1193 often undergo alterations to their ARG cargo, likely mediated by IS26. The IS26-mediated deletion event that featured prominently in ST1193 is likely to have rendered the integrons unable to capture, lose, or rearrange gene cassettes, while retaining their gene cassette expression functionality. This results in the “locking in” of gene cassettes in these truncated integron structures and their retention in place until they are activated by a functional intI1 present elsewhere in the genome or lost via homologous recombination. The intI1Δ745 truncation has been reported previously in wild Australian birds, albeit rarely, notably appearing in association with F29A^-B10 and F2: A^-B10 (both pUTI89-like) plasmids within E. coli ST131 and ST10 isolates in 2012 (33). This same study also reported intI1Δ682 BLASTn signatures from ST624, ST457, and occasionally other STs. This truncation size was also observed here in ST1193, similarly associated with carriage of dfrA17 and aadA5. These associations suggest that this integron structure is in circulation within ExPEC outside the F plasmids found within ST1193.

The global dissemination and clinical association of these IS26 mobilized integrons is of major concern and highlights the rapid adaptability of these urinary tract pathogens during antimicrobial treatment or, importantly, in synanthropic species, where exposure to selection pressures is more indirect. These developments demonstrate the next stage in the evolution of class 1 integrons and the resistance gene cargo they carry (33). Some other ARGs, blaCTX-M alleles being a key example, had a distribution disconnected from that of the class 1 integron, indicating that they were mobilized separately in ST1193. It is notable that chromosomal integration of blaCTX-M genes is on the rise in E. coli (49), including clinically relevant E. coli lineages that carry pUTI89, such as ST963 (50) and ST131 (32).

Multiple-drug resistance in E. coli that cause extraintestinal disease can lead to increased morbidity (extended hospital stays) and mortality and has a dominant influence on the epidemiology and management of infectious disease. E. coli ST1193 has spread globally at a high rate, given that the earliest entry of ST1193 in EnteroBase is 2007 and it was absent, for example, in 403 WGS of human commensal E. coli isolates collected between 1980 and 2010 (23). Understanding virulence, however, is complex because of the redundancy in the myriad of mobile elements that carry virulence gene cargo, especially in ExPEC. The number of ST1193 isolates recovered from human clinical cases serves as a strong indicator of its importance as a pandemic ExPEC lineage. Our study highlights the urgency of better understanding ExPEC lineages and genotype distributions, as well as identifying markers to differentiate and track future variation.

**Conclusion.** Members of the globally disseminated pandemic lineage ST1193 carry F plasmids encoding the pUTI89 virulence region comprising the copper and iron transporters, cjrABC and senB, and a putative hemin receptor but appear to differ markedly from pUTI89 in their capacity to be infiltrated by complex resistance regions assembled by the activity of IS26. These F plasmids, which appear to have developed within ST1193, are potential derivatives of pUTI89 but with smaller, distinct plasmid backbones and consistent carriage of ARGs. Our data contribute to the hypothesis that senB-carrying F plasmids are found in ExPEC lineages that colonize humans or have close human association, like companion animals and urban wildlife.

**MATERIALS AND METHODS**

**Long-read whole-genome sequencing.** The isolation and short-read sequencing of Escherichia coli isolates MVC465 and MVC530 from Australian canine companion animals was described previously (39). Both were isolated at the University of Melbourne Veterinary Hospital; MVC465 was taken from a canine...
urine sample in 2014, and MVC530 was obtained from urine through cystocentesis in 2015. Whole-genome sequencing of both isolates using a long-read platform was performed as described. Genomic DNA sequences were extracted from overnight cultures using the Promega Wizard HMW kit. Sequencing libraries were prepared from approximately 400 ng genomic DNA using the Oxford Nanopore rapid barcoding kit SQK-RBK004. The libraries were loaded onto a FLO-MIN106 R9.4 flow cell fitted on a MiniION Mk1B device. Live base calling of the MVC465 and the MVC530 raw reads was performed by MinKNOW versions 4.2.5 and 4.5.4, respectively, running Guppy versions 4.3.4 and 5.1.15, respectively, with the ‘fast’ R9.4.1 model. Demultiplexing and barcode trimming were performed on passed reads using Guppy barcoder version 5.1.15. The resulting fastq files were filtered for quality scores of >8 and lengths of >1,000 using NanoFilt (51). Completed assemblies were generated with Unicycler v0.5.0 using both short and long sequence reads.

**Data acquisition.** The whole-genome sequences, isolation metadata, e-serotypes and HincII (multilevel cluster assignment) (52) values for *E. coli* ST1193 (n = 707) were downloaded from EnteroBase (53) ([https://enterobase.warwick.ac.uk/](https://enterobase.warwick.ac.uk/)) on 20 August 2021. EnteroBase entries that did not include isolate source, year, and/or country of origin were excluded. Plasmid sequences were sourced from the GenBank nucleotide database ([https://www.ncbi.nlm.nih.gov/genbank/](https://www.ncbi.nlm.nih.gov/genbank/)).

**Genotyping and phylogenomic analyses.** Virulence genes, plasmid replicons, and mobilized antimicrobial resistance genes were detected using ABRicate ([https://github.com/tseemann/abricate](https://github.com/tseemann/abricate)) and the databases VirulenceFinder (54), PlasmidFinder (31) and ResFinder (55), respectively. ABRicate was also used for F plasmid subtyping using the F RST scheme (available at [https://pubmlst.org/organisms/plasmid-mlst/](https://pubmlst.org/organisms/plasmid-mlst/)). ABRicate was run using default settings outside custom database selection. The class 1 integrase gene intI1, IS26, ISEsp1, gyrA mutations, and the colibactin operon cib were identified separately using BLASTn (56) with no cutoff values set and then manually curated to 90% identity and differing coverage values depending on observed truncations. Phylogenetic trees and SNV counts were generated using Parsnp (with -x and -c flags) and gingr v1.2 from the Harvest Suite (57) ([https://github.com/marbl/parsnp](https://github.com/marbl/parsnp)) and visualized in iTOL (58) ([https://itol.embl.de/](https://itol.embl.de/)). Genomic alignments for comparative analysis of sequence data were performed using progressiveMauve (59). Circular BLASTn alignments were generated using BRIG (60), and linear alignments with EasyFig (61). Annotations were generated with RASTtk (62). SnapGene was used to handle and output sequence annotations. Typing data were cross-referenced with outputs from the Centre for Genomic Epidemiology ([http://www.genomicepidemiology.org/services/](http://www.genomicepidemiology.org/services/)).

**Data availability.** Data generated for this project can be found under GenBank BioProject accession PRJNA846578.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**SUPPLEMENTAL FILE 1**, XLSX file, 0.2 MB.

**SUPPLEMENTAL FILE 2**, XLSX file, 5.9 MB.

**SUPPLEMENTAL FILE 3**, PDF file, 0.1 MB.

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E.R.W. designed and performed data analysis and drafted and edited the manuscript. R.N.B. performed bacterial isolation and identification. M.S.M. generated genome sequences and edited the manuscript. G.F.B. assisted with analysis design and edited the manuscript. S.P.D. designed analysis, drafted and edited the manuscript, acquired funding, and provided supervision.

We declare that we have no competing interests.

**REFERENCES**

1. Johnson JR, Johnston BD, Porter SB, Clabots C, Bender TL, Thuras P, Trot DJ, Cobbold R, Mollinger J, Ferriero P, Drawz S, Banerjee R. 2019. Rapid emergence, subsidence, and molecular ejection of *Escherichia coli* sequence type 1193-fimH64, a new disseminated multidrug-resistant commensal and extraintestinal pathogen. J Clin Microbiol 57:e01664-18. [https://doi.org/10.1128/JCM.01664-18](https://doi.org/10.1128/JCM.01664-18).

2. Johnson TJ, Elnakeev E, Miller EA, Munoz-Aguayo J, Flores Figueroa C, Johnston S, Nielson DW, Logue CM, Johnson JR. 2019. Phylogenomic analysis of extraintestinal pathogenic *Escherichia coli* sequence type 1193, an emerging multidrug-resistant clonal group. Antimicrob Agents Chemother 63:e01913-18. [https://doi.org/10.1128/AAC.01913-18](https://doi.org/10.1128/AAC.01913-18).

3. Tchesnokova V, Radey M, Chattopadhyay S, Larson L, Weaver J.L, Kisiela D, Sokurenko EV. 2019. Pandemic fluoroquinolone resistant *Escherichia coli* clone ST1193 emerged via simultaneous homologous recombinations in 11 gene loci. Proc Natl Acad Sci U S A 116:14740–14748. [https://doi.org/10.1073/pnas.190302116](https://doi.org/10.1073/pnas.190302116).

4. Marcussen LL, Frimodt-Møller N, Hughes D. 2009. Interplay in the selection of fluoroquinolone resistance and bacterial fitness. PLoS Pathog 5: e1000541. [https://doi.org/10.1371/journal.ppat.1000541](https://doi.org/10.1371/journal.ppat.1000541).

5. Huseby DL, Pietsch F, Brandis G, Garoff L, Tegehall A, Hughes D. 2017. Mutation supply and relative fitness shape the genotypes of ciprofloxacin-resistant *Escherichia coli*. Mol Biol Evol 34:1029–1039. [https://doi.org/10.1093/molbev/msx052](https://doi.org/10.1093/molbev/msx052).
6. Biggel M, Moehrl S, Schmidt K, Frei A, Jung C, Steiger H. 2021. Complete genome sequence of Escherichia coli serotype ST1193. Microbiol Resour Announc 10:e0067-20. https://doi.org/10.1128/mrxr.0067-20.

10. Ding Y, Zhang J, Yao K, Gao W, Wang Y. 2021. Molecular characteristics of ST1193 microorganisms. Genome Res 31:1230-1236. https://doi.org/10.1101/gr.280625.120.

14. Findlay J, Gould VC, North P, Bowker KE, Williams MO, MacGowan AP. 2011. Molecular epidemiology of uropathogenic Escherichia coli from newborns in the United States. PLoS One 14:e0219352. https://doi.org/10.1371/journal.pone.0219352.

18. J Antimicrob Chemother 52:430–435. https://doi.org/10.1093/academic/jac/242.

22. Li D, Reid CJ, Kudinha T, Jarocki VM, Djordjevic SP. 2020. Molecular analysis of multidrug-resistant Escherichia coli ST58. Nat Commun 13:683. https://doi.org/10.1038/s41467-022-28342-4.

31. Carattoli A, Zankari E, Garcia-Fernandez A, Voldby Larsen M, Lund O, Börjesson S, Brouwer MSM, Hasman H. 2014. Prominence of an O75 clonal group (clonal complex 14) among non-ST131 fluoroquinolone-resistant Escherichia coli causing extraintestinal infections in humans and dogs in Australia. Antimicrob Agents Chemother 58:3898–3904. https://doi.org/10.1128/AAC.01210-11.

34. Desvaux M, Dalmasso G, Beyrouthy R, Barnich N, Delmas J, Bonnet R. 2007. ndle. 2017-18. J Antimicrob Chemother 72:1063-1069. https://doi.org/10.1093/jac/dkz397.

36. Dougherty MW, Jobin C. 2021. Shining a light on colibactin biology. Toxins 13:512-527. https://doi.org/10.3390/toxins13050512.

39. Elankumaran P, Cummins ML, Browning GF, Marenda MS, Reid CJ, Djordjevic SP. 2022. Second report of the new emerging global clone ST1193 Escherichia coli: salmochelin and aerobactin contribute more to virulence than heme in a chicken infection model. BMC Microbiol 12:143. https://doi.org/10.1186/s12866-012-143-3.

40. Findlay J, Gould VC, North P, Bowker KE, Williams MO, MacGowan AP. 2011. Molecular epidemiology of uropathogenic Escherichia coli from newborns in the United States. PLoS One 14:e0219352. https://doi.org/10.1371/journal.pone.0219352.

41. Platell JL, Trott DJ, Johnson JR, Heisig P, Heisig A, Clabots CR, Johnston B, Nelson TM, Uhlmann ML, Moller Aarestrup F, Hasman H. 2014. And trends in population structure of ESBL-producing Enterobacteriaceae among clinical isolates of extended-spectrum β-lactamase producing Escherichia coli causing urinary tract infections in children in France from 2014 to 2017. J Antimicrob Chemother 72:1063-1069. https://doi.org/10.1093/jac/dkz397.

42. Dougerty MW, Jobin C. 2021. Shining a light on colibactin biology. Toxins (Basel) 13:346. https://doi.org/10.3390/toxins13050346.

43. Elankumaran P, Cummins ML, Browning GF, Marenda MS, Reid CJ, Djordjevic SP. 2022. Second report of the new emerging global clone ST1193 Escherichia coli: salmochelin and aerobactin contribute more to virulence than heme in a chicken infection model. BMC Microbiol 12:143. https://doi.org/10.1186/s12866-012-143-3.

44. Findlay J, Gould VC, North P, Bowker KE, Williams MO, MacGowan AP. 2011. Molecular epidemiology of uropathogenic Escherichia coli from newborns in the United States. PLoS One 14:e0219352. https://doi.org/10.1371/journal.pone.0219352.

45. Findlay J, Gould VC, North P, Bowker KE, Williams MO, MacGowan AP. 2011. Molecular epidemiology of uropathogenic Escherichia coli from newborns in the United States. PLoS One 14:e0219352. https://doi.org/10.1371/journal.pone.0219352.

46. Findlay J, Gould VC, North P, Bowker KE, Williams MO, MacGowan AP. 2011. Molecular epidemiology of uropathogenic Escherichia coli from newborns in the United States. PLoS One 14:e0219352. https://doi.org/10.1371/journal.pone.0219352.

47. Findlay J, Gould VC, North P, Bowker KE, Williams MO, MacGowan AP. 2011. Molecular epidemiology of uropathogenic Escherichia coli from newborns in the United States. PLoS One 14:e0219352. https://doi.org/10.1371/journal.pone.0219352.

48. Findlay J, Gould VC, North P, Bowker KE, Williams MO, MacGowan AP. 2011. Molecular epidemiology of uropathogenic Escherichia coli from newborns in the United States. PLoS One 14:e0219352. https://doi.org/10.1371/journal.pone.0219352.

49. Findlay J, Gould VC, North P, Bowker KE, Williams MO, MacGowan AP. 2011. Molecular epidemiology of uropathogenic Escherichia coli from newborns in the United States. PLoS One 14:e0219352. https://doi.org/10.1371/journal.pone.0219352.

50. Findlay J, Gould VC, North P, Bowker KE, Williams MO, MacGowan AP. 2011. Molecular epidemiology of uropathogenic Escherichia coli from newborns in the United States. PLoS One 14:e0219352. https://doi.org/10.1371/journal.pone.0219352.

51. Findlay J, Gould VC, North P, Bowker KE, Williams MO, MacGowan AP. 2011. Molecular epidemiology of uropathogenic Escherichia coli from newborns in the United States. PLoS One 14:e0219352. https://doi.org/10.1371/journal.pone.0219352.
human ExPEC. Microbiol Spectr 10:e01291-22. https://doi.org/10.1128/spectrum.01291-22.
40. Liebert CA, Hall RM, Summers AO. 1999. Transposon Tn21, flagship of the floating genome. Microbiol Mol Biol Rev 63:507–522. https://doi.org/10.1128/MMBR.63.3.507-522.1999.
41. Reid CJ, Wyrsh ER, Roy Chowdhury P, Zingali T, Liu M, Darling AE, Chapman TA, Djordjevic SP. 2017. Porcine commensal Escherichia coli: a reservoir for class 1 integrons associated with IS26. Microb Genom. 3:e000143. https://doi.org/10.1099/mgen.0.000143.
42. Zingali T, Reid CJ, Chapman TA, Gaio D, Liu M, Darling AE, Djordjevic SP. 2020. Whole genome sequencing analysis of porcine faecal commensal Escherichia coli carrying class 1 integrons from sows and their offspring. Microorganisms 8:843. https://doi.org/10.3390/microorganisms8060843.
43. Schubert S, Rakin A, Karch H, Carniel E, Heesemann J. 1998. Prevalence of the “high-pathogenicity island” of Yersinia species among Escherichia coli strains that are pathogenic to humans. Infect Immun 66:480–485. https://doi.org/10.1128/IAI.66.2.480-485.1998.
44. Bogema DR, McKinnon J, Liu M, Hitchick N, Miller N, Venturini C, Iredell J, Darling AE, Roy Chowdury P, Djordjevic SP. 2020. Whole-genome analysis of extraintestinal Escherichia coli sequence type 73 from a single hospital over a 2 year period identified different circulating clonal groups. Microb Genom 6:e000255. https://doi.org/10.1099/mgen.0.000255.
45. Elankumaran P, Browning GF, Marenda MS, Reid CJ, Djordjevic SP. 2022. Close genetic linkage between human and companion animal extraintestinal pathogenic Escherichia coli ST127. Curr Res Microb Sci 3:e000143. https://doi.org/10.1128/MMBR.63.3.507-522.1999.
46. Stephens CM, Adams-Sapper S, Sekhon M, Johnson JR, Riley LW. 2017. Genomic analysis of factors associated with low prevalence of antibiotic resistance in extraintestinal pathogenic Escherichia coli sequence type 95 strains. mSphere 2:e00390-16. https://doi.org/10.1128/mSphere.00390-16.
47. Harmer CJ, Moran RA, Hall RM. 2014. Movement of IS26-associated antibiotic resistance genes occurs via a translocatable unit that includes a single IS26 and preferentially inserts adjacent to another IS26. mBio 5:e01801-14. https://doi.org/10.1128/mBio.01801-14.
48. Harmer CJ, Hall RM. 2015. IS26-mediated precise excision of the IS26-aphA1a translocatable unit. mBio 6:e01866-15. https://doi.org/10.1128/mBio.01866-15.
49. Gomi R, Yamamoto M, Tanaka M, Matsumura Y. 2022. Chromosomal integration of blaCTX-M genes in diverse Escherichia coli isolates recovered from river water in Japan. Curr Microbiol Res Sci 3:100144. https://doi.org/10.1016/j.jcmcrs.2022.100144.
50. Medvecky M, Papagiannitiss CC, Wyrsh ER, Bitar I, Cummings ML, Djordjevic SP, Dolejška M. 2022. Interspecies transmission of CMY-2-producing Escherichia coli sequence type 963 isolates between humans and gulls in Australia. mSphere 7:e00238-22. https://doi.org/10.1128/mSphere.00238-22.
51. De Coster W, D’Hert S, Schultz DT, Cruts M, Van Broeckhoven C. 2018. NanoPack: visualizing and processing long-read sequencing data. Bioinformatics 34:2666–2669. https://doi.org/10.1093/bioinformatics/bty149.
52. Zhou Z, Charlesworth J, Achtman M. 2021. HierCC: a multi-level clustering scheme for population assignments based on core genome MLST. Bioinformatics 37:3645–3646. https://doi.org/10.1093/bioinformatics/btab234.
53. Zhou Z, Alikhan NF, Mohamed K, Fan Y, Achtman M, Agama Study Group. 2020. The Enterobase user’s guide, with case studies on Salmonella transmissions, Yersinia pestis phylogeny, and Escherichia core genomic diversity. Genome Res 30:138–152. https://doi.org/10.1101/gr.251678.119.
54. Kleinheinz KA, Joensen KG, Larsen MV. 2014. Applying the ResFinder and VirulenceFinder web-services for easy identification of acquired antibiotic resistance and E. coli virulence genes in bacteriophage and prophage nucleotide sequences. Bacteriophage 4:e27943. https://doi.org/10.4161/bact.27943.
55. Zankari E, Hasman H, Cosentino S, Vestergaard M, Rasmussen S, Lund O, Aarestrup FM, Larsen MV. 2012. Identification of acquired antimicrobial resistance genes. J Antimicrob Chemother 67:2640–2644. https://doi.org/10.1093/jac/dks261.
56. Camacho C, Coulouris G, Avagyan V, Ma N, Papadopoulos J, Bealer K, Madden TL. 2009. BLAST+: architecture and applications. BMC Bioinformatics 10:421. https://doi.org/10.1186/1471-2105-10-421.
57. Treangen TJ, Ondov BD, Koren S, Phillippy AM. 2014. The Harvest suite for rapid core-genome alignment and visualization of thousands of intraspecific microbial genomes. Genome Biol 15:524. https://doi.org/10.1186/s13059-014-0524-x.
58. Letunic I, Bork P. 2019. Interactive Tree Of Life (iTOl) v4: recent updates and new developments. Nucleic Acids Res 47:W256–W259. https://doi.org/10.1093/nar/gkz239.
59. Darling AE, Mau B, Perna NT. 2010. progressiveMauve: multiple genome alignment with gene gain, loss and rearrangement. PLoS One 5:e11147. https://doi.org/10.1371/journal.pone.0011147.
60. Aarestrup FM,椋田 Mohji, Achtman M. 2021. Hierarchical clustering algorithm for population assignments based on core genome MLST. Bioinformatics 37:3645–3646. https://doi.org/10.1093/bioinformatics/btab234.
61. Sullivan MJ, Petty NK, Achtman M. 2011. Easyfig: a genome comparison visualization tool. Bioinformatics 27:1009–1010. https://doi.org/10.1093/bioinformatics/btr039.
62. Bretten T, Davis JJ, Dicz T, Edwards RA, Gerdes S, Olsen GJ, Olson R, Overbeek R, Parrello B, Pusch GD, Shukla M, Thomason JA, 3rd, Stevens R, Vonstein V, Wattam AR, Xia F. 2015. RASTtk: a modular and extensible rapid core-genome alignment and visualization of thousands of intraspecific microbial genomes. Genome Biol 15:524. https://doi.org/10.1186/s13059-014-0524-x.