Interspecies Transmission of CMY-2-Producing Escherichia coli Sequence Type 963 Isolates between Humans and Gulls in Australia

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ABSTRACT Escherichia coli sequence type 963 (ST963) is a neglected lineage closely related to ST38, a globally widespread extraintestinal pathogenic ST causing urinary tract infections (UTI) as well as sepsis in humans. Our current study aimed to improve the knowledge of this understudied ST by carrying out a comprehensive comparative analysis of whole-genome sequencing data consisting of 31 isolates from silver gulls in Australia along with another 80 genomes from public resources originating from geographically scattered regions. ST963 was notable for carriage of cephalosporinase gene \( \text{bla}^{CMY-2} \), which was identified in 99 isolates and was generally chromosomally encoded. ST963 isolates showed otherwise low carriage of antibiotic resistance genes, in contrast with the closely related \( E. \text{coli} \) ST38. We found considerable phylogenetic variability among international ST963 isolates (up to 11,273 single nucleotide polymorphisms [SNPs]), forming three separate clades. A major clade that often differed by 20 SNPs or less consisted of Australian isolates of both human and animal origin, providing evidence of zoonotic or zooanthropogenic transmission. There was a high prevalence of virulence F29:A-B10 pUTI89-like plasmids within \( E. \text{coli} \) ST963 (\( n = 88 \)), carried especially by less variable isolates exhibiting \( \leq 1,154 \) SNPs. We characterized a novel 115,443-bp pUTI89-like plasmid, pCE2050_A, that carried a \( \text{traS}^5 \text{S5} \) insertion absent from pUTI89. Since \( \text{S5} \) was also present in a transposition unit bearing \( \text{bla}^{CMY-2} \) on chromosomes of ST963 strains, \( \text{S5} \) insertion into pUTI89 may enable mobilization of the \( \text{bla}^{CMY-2} \) gene from the chromosome/transposition unit to pUTI89 via homologous recombination.

IMPORTANCE We have provided the first comprehensive genomic study of \( E. \text{coli} \) ST963 by analyzing various genomic and phenotypic data sets of isolates from Australian silver gulls and comparison with genomes from geographically dispersed regions of human and animal origin. Our study suggests the emergence of a specific \( \text{bla}^{CMY-2} \)-carrying \( E. \text{coli} \) ST963 clone in Australia that is widely spread across the continent by humans and birds. Genomic analysis has revealed that ST963 is a globally dispersed lineage with a remarkable set of virulence genes and virulence plasmids described in uropathogenic \( E. \text{coli} \). While ST963 separated into three clusters, a unique specific clade of Australian ST963 isolates harboring a chromosomal copy of AmpC \( \beta \)-lactamase encoding the gene \( \text{bla}\text{CMY-2} \) and originating from both humans and wild birds was identified.

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This phylogenetically close cluster comprised isolates of both animal and human origin, thus providing evidence of interspecies zoonotic transmission. The analysis of the genetic environment of the AmpC β-lactamase-encoding gene highlighted ongoing evolutionary events that shape the carriage of this gene in ST963.

**KEYWORDS** β-lactamases, Escherichia coli, ST963, WGS, Australia, transmission, comparative genomics, phylogenetic analysis, silver gulls, humans

Extended-spectrum β-lactamases (ESBL) and AmpC type β-lactamases are enzymes that are able to hydrolyze beta-lactams including extended-spectrum cephalosporins and are frequently found in Enterobacterales, such as Escherichia coli and Klebsiella pneumoniae, of human and animal origin (1). The most common plasmid-mediated AmpC β-lactamase in Enterobacterales is CMY-2 (2). The gene encoding CMY-2 (bla_{CMY-2}) likely originates from a chromosomal AmpC gene from Citrobacter freundii (3) and was translocated onto plasmids by IS_{Ecp1} (4). Consequently, the gene has spread to different hosts by conjugal plasmids belonging to various incompatibility (Inc) groups, including A, C2, II, F, N, K/B, and K2 (5–10).

*E. coli* sequence type 963 (ST963) is a relatively understudied lineage belonging to clonal complex (CC) 38 and phylogroup D. *E. coli* ST38 is an extraintestinal pathogenic (ExPEC) bacterium that is a documented cause of urinary tract infections and sepsis in humans (11). Moreover, *E. coli* ST38 is geographically widespread and is associated with multidrug resistance (MDR) and ESBL production, making it an important pandemic ST globally (11). To our knowledge, there is a lack of genomic studies comprising broader collections of ST963 isolates. *bla_{CMY-2}*-carrying isolates of ST963 sporadically appeared in larger studies related to detection of CMY-2-producers, and *bla_{CMY-2}* was located on plasmids in most instances (5, 7). Only a single study reported evidence for chromosomally encoded *bla_{CMY-2}* in a single human-derived ST963 isolate in Germany (8).

*E. coli* ST963 has been sporadically reported in wildlife, in companion animals, and very occasionally, in humans (5, 7, 8, 12). It came to our attention as part of an extensive study of *E. coli* strains with resistance to extended-spectrum beta-lactams and derived from gull chicks nesting on three sites in Australia undertaken in 2012. Whole-genome sequences (WGS) of 425 *E. coli* isolates (13) show that ST457 (14), ST216 (15), and ST963 were the dominant *E. coli* sequence types in that study. While our studies suggest that ST457 (14) and ST216 (15) are important emerging *E. coli* STs, little is known about ST963.

Here, we performed WGS-based comparative analysis of a collection of 111 *E. coli* ST963 isolates comprising 31 isolates from silver gulls in Australia and 80 GenBank sequences of various source and country origins. The isolates of human and animal origin predominantly originate from Australia and carry *bla_{CMY-2}* on the chromosome. A detailed phylogenetic analysis shed lights on their zoonotic and zooanthropogenic potential.

**RESULTS**

Phylogenetic analyses of *E. coli* ST963. A phylogenetic study based on core genome alignments of all 111 ST963 genomes showed a branching pattern consisting of 3 separate clades. A category of evolutionarily distant isolates (*n* = 13) lacking *bla_{CMY-2}* or encoding it on plasmids formed clade 1 (highlighted by pink color in Fig. 1). Only one isolate from clade 1 (GenBank SRA run accession no. SRR6376575) carried a chromosomal copy of *bla_{CMY-2}*, but with a genetic context different from remaining isolates that bear chromosomal *bla_{CMY-2}* (variant D, Fig. 2). Isolates from within clade 1 were obtained from all three sources and originated in the United States, Canada, Mexico, New Zealand, and Australia without clear clustering patterns. They all exhibited more than 3,000 single nucleotide polymorphisms (SNPs) against our CE2050 reference sequence. The remaining isolates (*n* = 98) formed a divergent clade which further subdivided into an Australian clade of CMY-2-producing isolates (clade 2, highlighted in green) and another clade of isolates from outside Australia which did not carry *bla_{CMY-2}* as well as isolates carrying...
blaCMY-2 on the chromosome or on an I1 plasmid (clade 3), shown in violet in Fig. 1. Clade 2 consisted of 84 isolates, of which 82 were from Australia, 1 was from New Zealand, and the remaining isolate was from the United States (SRR8689680).

To further discriminate between closely related isolates, another tree of 81 isolates with SNP counts below 100 against the CE2050 chromosome was constructed based on SNPs called from mapped sequencing read data (see Fig. S1 in the supplemental material). Of the 81, 43 were obtained from humans, 35 from gulls, and 3 from cows. Most of these closely related isolates originated in Australia ($n = 76$), with the remaining 5 coming from Europe ($n = 3$ from Netherlands, $n = 1$ each from the United Kingdom and Germany). Closely related isolates of animal and human origin, many with less than 20-SNP differences were detected, supporting the hypothesis of their zoonotic and/or zooanthropogenic transmission potential (see "SNP dist" sheet in Table S1). Australian isolates did not appear to cluster according to their territory of origin. Phylogenetic analysis clearly showed that there is a specific CMY-2-producing Australian clone circulating within the country that is spread by humans and animals (humans and gulls were the only available sources of WGS of E. coli ST963 from Australia). The fact that the same transposition unit (variant A) was found on chromosomes of isolates from Germany, Netherlands, United Kingdom, and the United States suggests these isolates shared a recent common ancestor.
Genomic features of a reference genome of *E. coli* ST963. Our closed reference genome CE2050 comprised a chromosome of 4,979,906 bp and two plasmids, pCE2050_A, a 115,443-bp F29:A-:B10 plasmid, and pCE2050_B, a 222,008-bp IncHI2 plasmid typed as ST3. While pCE2050_A was carried by most *E. coli* ST963 isolates described in this study (see below, Fig. 1), pCE2050_B was only present in isolate CE2050. One intact prophage region was identified on the chromosome (nucleotides [nt] 2050804 to 2082987 in GenBank accession no. CP073621), showing a high level of similarity to prophage PHAGE_Entero_WPhi_NC_005056. There was also cephalosporinase gene *bla*CMY-2 found on the chromosome along with 10 different virulence factors (see Table S1). Plasmid pCE2050_A encodes Col156, IncFIB, and IncFII plasmid replicons, and its replicon sequence type (RST) is F29:A-:B10. The same plasmid ST is a reliable predictor of plasmids that bear resemblance to virulence plasmid pUTI89 (16). These two plasmids are nearly identical in length (115,443 bp versus 114,230 bp, respectively) and gene content and exhibit few SNPs. Similar to other pUTI89-like plasmids, pCE2050_A did not carry any antibiotic resistance gene (ARGs); however, it carried an IS26 and a virulence factor, traT. Notably, pCE2050_A carried a traS5iSS sequence (nt 6770 to 7964 in pCE2050_A) absent from pUTI89, flanked by a CTAA 4-bp direct repeat (DR) on either end of the IS5 element. pCE2050_B is closely related to other IncHI2 plasmids, including pCE1681-A (94% coverage and 100% identity) typed as ST3, which was carried by *E. coli* ST216 recovered from a silver gull in Australia (GenBank accession no. MT180430) (15) in 2012. pCE2050_B harbored a 1,131-bp fragment that shows similarities to sequences previously described in IncN plasmids (17). This fragment included the *repA* gene and a part of the iteron region. pCE2050_B carries an MDR region with a *Tn1721*-specific tetracycline module with the tet(A) gene inserted in the same site as in other IncHI2 plasmids (18) and an In369 class 1 integron with *dfrA1b* and *aadA1b* gene cassettes.

**Antibiotic resistance, virulence genes, and plasmids of the ST963 collection.** All ST963 isolates from our Australian gull collection (*n* = 31) were resistant to ampicillin, cefazolin, ceftazidime, and amoxicillin, which is in accordance with their AmpC phenotype. They all also exhibited resistance or intermediate susceptibility to aztreonam and streptomycin. Analysis of this gull collection, and an additional 80 ST963 WGS data sets from EnteroBase, showed that *bla*CMY-2 was present in all isolates from our Australian silver gull collection (31/31) and in 85% of EnteroBase sequences (68/80). *bla*CMY-2 was identified in 52 out of 60 isolates from clinical samples of humans, 43/45 isolates obtained from wild animals, and 4 of 5 isolates from livestock/companion animals. Other resistance genes were present in much lower frequencies. The second most prevalent antibiotic resistance gene (ARG) in the collection was *bla*TEM-1B (encoding narrow-spectrum β-lactamase) found in 10 of 111 isolates, followed by
genes for resistance to sulfonamides, sul2 (9/111) and sul1 (8/111), and aminoglycosides, aph(3’)-lb (7/111) and aph(6)-ld (7/111). A complete list of predicted ARGs can be found in Table S1. On average, isolates carried 1.7 ARGs, indicating that while E. coli ST963 frequently harbors blaCMY-2, genotypic MDR is unusual. IS26 (99/111; 89%) was frequently identified in the collection. A total of 14 (13%) isolates carried the int1 gene for class 1 integrase.

Virulence gene analysis showed that all isolates harbored fimH, fyuA, irp2, and silA on the chromosome and several copies of ipaH and yeeT located on the chromosome as well. There was also high prevalence of chromosomally encoded sitA (n = 107) and plasmid-carried traT (n = 101) among the isolates under investigation, while other putative virulence factors were present in lower frequencies. A complete list of virulence-associated genes (VAGs) is given in Table S1.

Most (88/111; 79%) ST963 isolates, which included representation from humans (48/60; 80%) and wildlife (34/45; 76%) as well as livestock/companion animals (4/5), carried either the 115,443-bp F plasmid pCE2050_A or closely related variants with more than 90% of the pCE2050_A sequence. Analysis of all 88 pCE2050-like bearers showed that the ΔtraSI5 insertion was only present in three closely related isolates (CE2050, CE2010, and CE1902, all showing ≈3 SNP differences when compared to each other). pCE2050_A showed 99% coverage and 99.95% identity with a pUTI89-like plasmid named XXX from the GenBank database (accession no. LR730401) that was carried by E. coli ST73 from vasculated human blood in Germany. Apart from pCE2050_A replicons, PlasmidFinder analysis showed another 29 FII replicons, followed by 17 I1 and 15 FIB replicons that were present within the studied sample set. The FAB formulas for F plasmids along with a complete list of plasmid replicons that were identified in the study can be found in Table S1.

Genetic context of blaCMY-2. Of the 89% (99/111) of ST963 which carry blaCMY-2, the vast majority (94/99) had the gene located chromosomally (Fig. 1). In the remaining five isolates, it was carried by various IncI1-I(a) plasmids, which belonged to clonal complex 2 (CC2; two had ST2, two had ST23) or ST19. Two I1/ST23 plasmids, including the 92,387-bp pCE1747_I1 from our silver gull collection and the 89,653-bp pCH12 from a human (GenBank SRA run accession no. SRR6455977), originated in Australia and were closely related to the 94,698-bp plasmid pCE1628_I1 (14) from E. coli ST457 (GenBank accession no. MT468651), isolated from a silver gull in Australia, suggesting that plasmid sharing among E. coli STs occurs in gull colonies. pCE1628_I1 showed 99% coverage and 99.95% identity with a pUTI89-like plasmid from the GenBank database, while pCE1747_I1 exhibited 97% coverage and 99.97% identity with pCE1628_I1 using BLASTn. I1/ST2 plasmid pECOL-18-VL-SD-1A-0003 (88,569 bp) from a canine source in the United States (GenBank SRA run accession no. SRR7788245) shows 99% coverage and 99.97% sequence identity with an 88,229-bp pAR-0430-1 (GenBank accession no. CP044137) from an E. coli O157 isolate with ST11. Metadata regarding the source of isolation or country of origin is unavailable. An I1/ST19 plasmid with an approximate size of 94 kbp (GenBank SRA run accession no. SRR3290000) obtained from a human in the United States shows high similarity to the 94,170-bp plasmid p4540-1 (GenBank accession no. CP041533) recovered from E. coli ST963 from an unrelated human in the United States. Using BLASTn, the I1/ST19 plasmid exhibited 97% sequence coverage and 99.68% nucleotide identity of with p4540-1. The remaining I1/ST2 plasmid pECOL-19-VL-WA-KY-0014, recovered from a wolf in the United States (GenBank SRA run accession no. SRR10535897), showed 100% sequence coverage and 99.99% sequence identity with the 94,881-bp plasmid p95 (GenBank accession no. CP023356) carried by an E. coli typed as ST963 that was obtained from a canine source in the United Kingdom in 2002. Metadata relating to isolates carrying blaCMY-2-positive plasmids can be found in Table S1.

Analysis of regions flanking blaCMY-2. BlaCMY-2 was identified in four different genetic environments in the chromosome of ST963, here referred to as variants A to D (Fig. 2). Most (94/111; 85%) isolates, including those from humans, wildlife and livestock, and companion animals (51/60, 40/45, and 3/5, respectively), carried a similar genetic arrangement found
on the chromosome of _C. freundii_ (bla<sub>CMY-2</sub>-blc-sugE) (19). The transposition unit was likely mobilized by ISEcp1 (IS1380 family) truncated by insertion of an ISS element (Fig. 2A). The vast majority of isolates carried bla<sub>CMY-2</sub> as part of a transposition unit defined here as variant A on the chromosome (n = 93). Variant A was mostly located within the yehB gene, which is part of the yehABCD fimbral operon, and was flanked by a TATAAA DR. An identical transposition unit was also present on an I1/ST19 plasmid from a human isolate from our Enterobase _E. coli_ ST963 collection originating in the United States (GenBank SRA run accession no. SRR3290000). BLASTn analysis of the 3,931-bp-long region comprising ISEcp1-ISS-ΔISEcp1-bla<sub>CMY-2</sub>-ΔyehABCD (nt 1417672 to 1421692 in GenBank accession no. CP073621) showed only four high-scoring hits, all of them having a bla<sub>CMY-2</sub>-blc-sugE module. Two hits belonged to _E. coli_ ST963 chromosomal sequences (GenBank accession no. CP051733 and LR130562), one was to I1/ST19 plasmid p4540-1 (GenBank accession no. CP041533) from _E. coli_ ST963 obtained in 2012 from infected human blood in the United States, and the remaining match belonged to IncC-ST3 plasmid p25358-2 (GenBank accession no. CP051443) from _Salmonella enterica_ (20) isolated in 2002 from turkey in the United States.

8 variants were identified in three isolates from a human, a silver gull, and a canine, all three of which carried bla<sub>CMY-2</sub> on different IncI1-I(a) plasmids. In each of these isolates a copy of IS1294 followed by a small, truncated portion of ISEcp1 was identified downstream of bla<sub>CMY-2</sub>. Variant B showed similarity to a region found in plasmid pS10584 from _Salmonella enterica_ from China (GenBank accession no. KX058576). Various B variants differed from each other in the flanking region upstream of the IS1294-ΔISEcp1-bla<sub>CMY-2</sub>-ΔyehABCD module (Fig. 2). Variant C was identified in a single ST963 isolate from a wolf in the United States (GenBank SRA run accession no. SRR10535897). Here, bla<sub>CMY-2</sub> was detected on another IncI1-I(a) plasmid. In variant C, ISSbo1 (an IS91 family element) followed by a truncated portion of ISEcp1 was located upstream of bla<sub>CMY-2</sub>. Direct repeats indicating transposition of the bla<sub>CMY-2</sub>-carrying transposition unit into the specific genetic element were not detected. A single isolate obtained from a wild bird (GenBank SRA run accession no. SRR6376575) carried a chromosomally localized bla<sub>CMY-2</sub> gene also associated with ISEcp1 (variant D). bla<sub>CMY-2</sub> along with small portions of blc was inserted into the lysR gene and was flanked by TATAA DRs. Metadata associated with these genomic assemblies and their bla<sub>CMY-2</sub>-carrying transposition units is available in Table S1.

_Closely related ST963 strains are shared by humans and gulls._ ST963 isolates are phylogenetically diverse; nearly three quarters (81/111; 73%) of all ST963 genomes displayed fewer than 100 SNPs compared to the CE2050 chromosome, while 13% (14/111) showed more than 1,000 SNPs against our reference chromosome. Isolates with more than 3,000 SNPs compared to the CE2050 chromosome reside in clade 1 (shown in pink color in Fig. 1). The remaining ST963 isolates represented a dominant clade that was further divided into two subclades (clade 2, shown in green and clade 3, shown in violet). In clade 2 the _E. coli_ ST963 genomes, including those from gulls from three sites and from humans, contained most (at least 98.5%) of the CE2050 chromosome. Interestingly, most ST963 isolates with >1,000 SNP differences from the reference chromosome either did not carry bla<sub>CMY-2</sub> (8/14; 57%) or bla<sub>CMY-2</sub> was located on a plasmid (4/14; 29%). Plasmid-mediated bla<sub>CMY-2</sub> was detected in samples originating in all three source categories (human, wildlife, and livestock/companion animals), namely, human, gull, wolf, and dog. There were only two isolates with >1,000 SNPs that encoded bla<sub>CMY-2</sub> in the chromosome. An isolate with 1,154 SNPs, obtained from a human in Australia (SRR11094161), carried a variant A bla<sub>CMY-2</sub>-carrying genetic arrangement (Fig. 1, clade 2). Another isolate, with 3,190 SNPs, isolated from a bald eagle in the United States (SRR6376575), carried a variant D bla<sub>CMY-2</sub>-carrying genetic context (Fig. 1, clade 1).

The cohort was separated into two groups, one of close relatives (defined as ≤100 SNPs against the CE2050 chromosome and referred to as “group1”) and one of more distant relatives (defined as >100 SNPs, referred to as “group2”). Site-wise aggregation of SNP counts was undertaken and visualized (see Fig. S2). The distribution of SNPs within group1 appeared uniform except for a highly variable region...
which contained over 400 SNPs (Fig. S2A). Most of those SNPs were located in a sequence encoding GTPase Era, which was found as a family comprising four paralogs in the CE2050 chromosome. Due to possible mis-alignment of reads originally belonging to other copies of Era genes, SNPs found in this region were discarded from the alignment file used for the construction of a detailed phylogeny inference (Fig. S1). When scrutinizing group2 genomes, four regions each with an elevated number of SNPs were observed. Detailed genomic information about these regions can be found in the Table S3.

All ST963 isolates \((n = 81)\) exhibiting fewer than 100 SNPs to the reference CE2050 chromosome, as well as the reference strain itself, carried \(\text{bla}_{\text{CMY-2}}\) on the chromosome, except for one isolate where we identified a single SNP that led to a change in the \(\text{bla}_{\text{CMY-2}}\) allele to \(\text{bla}_{\text{CMY-143}}\). Chromosomally encoded \(\text{bla}_{\text{CMY-2}}\) was present in most isolates (11/15; 73%) with SNP counts in the range of 100 to 1,000 (isolates mostly from clade 3, shown in violet color in Fig. 1). All isolates with SNP counts below 1,000 that carried \(\text{bla}_{\text{CMY-2}}\) on the chromosome carried variant A (see Fig. 1). We also noticed a correlation between chromosomal SNP counts and carriage of F plasmid pCE2050_A (or its close variants) within ST963 genomes. While they were identified in 79% \((88/111)\) of ST963 isolates in total, these closely related plasmids were carried by 88% \((72/82)\) of isolates with fewer than 100 SNPs to the reference, by 80% \((12/15)\) isolates exhibiting SNP counts in the range of 100 to 1,000, and only by 29% \((4/14)\) of isolates showing more than 1,000 SNPs.

**DISCUSSION**

ST963 is a phylogroup D \(E.\ coli\) strain closely related to ST38 in CC 38. At the time of writing (May 2022), a search of the PubMed database matches did not return a hit to the term “\(E.\ coli\) ST963,” highlighting the lack of information on this sequence type. Nonetheless, ST963 has been reported in studies of \(E.\ coli\) recovered from wild and urban-adapted birds (12, 21). Here, we characterized 111 ST963 isolates originating in humans \((n = 60)\), wildlife \((n = 45\) [31 of which are from Australian silver gulls]), as well as livestock/companion animals \((n = 5)\). The isolates were recovered between 1984 and 2019 and predominantly sourced from Australia \((n = 89)\). While we contributed 31 \(E.\ coli\) ST963 strains from Australian silver gulls, the remaining 80 \(E.\ coli\) ST963 genomes were from geographically dispersed regions that were deposited in EnteroBase (http://enterobase.warwick.ac.uk/).

All \(E.\ coli\) ST963 genomes harbored multiple VAGs, including \(\text{fimH}\), \(\text{fyuA}\), \(\text{irp2}\), \(\text{silA}\), \(\text{ipaH}\), and \(\text{yeeT}\), and many also carried \(\text{sitA}\), \(\text{traT}\), and the insertion element \(\text{IS26}\). Cocarcriage within each isolate of iron-acquisition genes \(\text{fyuA}\) and \(\text{irp2}\) suggests the presence of the \(Yersinia\) high-pathogenicity island, a chromosomal genomic island associated with increased capacity for avian infection and uropathogenesis (22). \(\text{senB}\), a virulence gene used as a marker gene for the presence of virulence plasmid pUTI89, was indeed present in all isolates \((n = 88)\) that carried pUTI89-like plasmid pCE2050_A (or its close variants) within ST963 genomes. While they were identified in 79% \((88/111)\) of ST963 isolates in total, these closely related plasmids were carried by 88% \((72/82)\) of isolates with fewer than 100 SNPs to the reference, by 80% \((12/15)\) isolates exhibiting SNP counts in the range of 100 to 1,000, and only by 29% \((4/14)\) of isolates showing more than 1,000 SNPs.
into the coding sequence of ISEcP1 could explain the inactivation of the insertion element ISEcP1, which has been originally associated with the mobilization of the blacMY-2 resistance gene. The fact that the insertion site of the blacMY-2-carrying genetic determinant was identical in the majority of the studied isolates further supports the above-described hypothesis. Insertion elements are known to alter the expression of clinically important ARGs. In this regard, ISEcP1 is not only involved in gene mobilization, but it also enhances expression of blCTX-M lactamases (25). IS5 was previously associated with increased blacMY-2 expression through a tandem gene amplification mechanism described in clinical E. coli strains (26). Overall low carriage of ARGs along with frequent presence of blacMY-2 within E. coli ST963 genomes suggests that this E. coli lineage is possibly early in its evolutionary trajectory as a potential human antimicrobial resistance threat.

All ST963 genomes that were closely related (<100 SNPs) to our reference genome CE2050 harbored a chromosomal copy of blacMY-2. blacMY-2 was also present on the chromosome of most isolates (11/15; 73%) with SNP counts against CE2050 chromosome between 100 and 1,000, suggesting that its location on the chromosome is ancestral. In contrast, isolates divergent from CE2050 (>1,000 SNPs) either lack blacMY-2 or it was located on various I1 plasmids. Only two such divergent isolates encoded blacMY-2 chromosomally. One carried our predominant transposition unit (variant A), while another isolate carried variant D, and as such, it contained an intact copy of ISEcP1 and lacked ISS as well as large portion of blc and all of sugE, probably due to further mobilization events that occurred after the initial transposition unit was incorporated into the chromosome. These data highlight some of the ongoing evolutionary events that shape carriage of blacMY-2 in ST963.

We also noted a correlation between chromosomal SNP counts and carriage of the 115,443-bp F plasmid pCE2050_A from our closed reference genome. pCE2050_A and variants of it were borne by 87% of isolates with SNP counts below 1,000 compared to the CE2050 chromosome, while it was present only in 29% of isolates that showed more than 1,000 SNPs against our reference. pCE2050_A is an F29:A-:B10 with high nucleotide identity with pUTI89. However, it is notable that pCE2050_A carries a traS:ISS insertion absent from pUTI89. The truncation of the Tra region by ISS could result in the loss of the conjugation ability of pCE2050_A. This insertion event was not detected within the NCBI nucleotide collection; however, screening our collection identified it in three pCE2050_A-like bearers that were closely related to each other. This suggests a recent incorporation of the ISS element into pUTI89, potentially enabling further mobilization of blacMY-2 from the chromosome/transposition unit to pUTI89 via homologous recombination. This insertion sequence and its junction with traS therefore constitute a distinct, trackable genetic motif for pCE2050_A and related plasmids.

pUTI89 is a globally dispersed F virulence plasmid with an F29:A-:B10 replicon type found in several top 20 E. coli STs causing clinical disease (27). pUTI89 was first reported in strain UTI89, an ST95 E. coli from a patient with an acute bladder infection (28), and has been assessed for its ability to cause disease in a mouse urinary tract infection (UTI) infection model (29, 30). Specifically, pUTI89 and close variants of it have been found in some but not all sublineages of ST95 (27, 31), ST127 (32), and ST131 (33), E. coli STs that are dominant clinical uropathogens (11, 34). Despite a copy of IS26, a key element involved in the rearrangement and spread of multiple ARGs, being present in pUTI89, these plasmids rarely carry ARGs; however, they harbor important virulence factors, including the cjrABC-senB operon and traT gene.

We did not observe any correlation between carriage of pCE2050_A-like plasmids and source of isolation. Our closed reference genome CE2050 also contained a 222,008-bp IncHI2-N plasmid, pCE2050_B, that consisted of an IncHI2 plasmid backbone and a 1,131-bp fragment showing similarities to sequences previously described in IncN plasmids. pCE2050_B is similar to HI2 plasmid pCE1681-A (15) obtained from E. coli ST216 from our Australian gull chick nesting collection. Strain
CE2050 was recovered from Montague Island (also known as Barunguba Island), while ST216 strain CE1681, which bears a very similar HI2 plasmid, was sourced from Five Islands Nature Reserve. These two islands are separated by approximately 200 km along the eastern coastline of New South Wales, Australia.

Our analyses revealed significant variability in SNP counts (3 to 11,273 SNPs) when sequencing reads were mapped against the CE2050 chromosome. Isolates with <100 SNPs showed a generally uniform distribution of SNPs relative to the reference CE2050 genome, while SNP frequency distribution in isolates exhibiting more than 100 SNPs revealed four SNP hot spot regions within approximately 300 kbp (Fig. S2A and B). Those were found mainly in divergent isolates carrying more than 1,000 SNPs against our reference genome. This extensive SNP heterogeneity of E. coli ST963 strains suggests the need for the employment of typing methods with higher discriminatory power than standard multilocus sequence typing (MLST) schemes have, such as core genome MLST (cgMLST) or even the reference-independent phylogenetic approach to avoid reference bias.

In summary, phylogenetic analyses identified a unique specific clade of Australian ST963 isolates harboring a chromosomal copy of blaCMY-2. This Australian clade comprised phylogenetically close clusters (<20 SNP differences from the reference genome) of isolates of both animal and human origin, providing evidence of interspecies transmission. The fact that isolates originating outside Australia segregate to different clades, separated from the major Australian clade, supports the hypothesis of the emergence of a specific blaCMY-2-carrying E. coli ST963 clone in Australia that is widely spread across the continent by humans and birds. The presence of transposition unit ΔISEcp1-155-ΔISEcp1-blaCMY-2-bloc-sugE, inserted into the same location within the chromosome of isolates from Australia, Germany, Netherlands, United Kingdom, and the United States point to a shared recent common ancestor.

MATERIALS AND METHODS

Strain collection. The collection examined here comprised 35 E. coli isolates cultivated on MacConkey agar with cefotaxime (2 mg/L) from cloacal swabs from silver gulls (Chroicocephalus novaehollandiae) that were collected in three nesting colonies in New South Wales (NSW), Australia, in 2012 (35) and subjected to Illumina sequencing. Illumina WGS data of a further 81 E. coli ST963 strains were gathered from the GenBank SRA database based on information found in EnteroBase (http://enterobase.warwick.ac.uk; the search was conducted in February 2020). In total, there were 60 sequences originating from human clinical isolates, 50 from wild animals, 5 from livestock/companion animals, and a single isolate with missing information regarding the source. Isolates were collected between 1984 and 2019; however, most strains were collected in 2012 or later (n = 102); three isolates lacked the year of isolation. Countries of origin included Australia (n = 89), the United States (n = 16), Netherlands (n = 4), Canada (n = 2), New Zealand (n = 2), the United Kingdom (n = 1), Germany (n = 1), and China (n = 1). After an initial quality check, duplicate genomes (n = 5; 4 isolates from our gull collection and a single gull isolate from EnteroBase) were discarded from downstream analyses, leaving 111 samples in the final data set. Detailed metadata are available in Table S1.

Antibiotic susceptibility testing, DNA extraction, and sequencing. The disk diffusion methodology with a set of 21 antibiotics (Oxoid, Hants, UK) was performed according to European Committee on Antimicrobial Susceptibility Testing (EUCAST) recommendations (36). Inhibition zone diameters of the tested isolates were measured and interpreted according to EUCAST breakpoints (36) or using breakpoints defined by CLSI in 2017 for antibiotics (azithromycin, cefazolin, tetracycline, nalidixic acid, sulfonamide compounds, and streptomycin) with no defined breakpoints in EUCAST (36, 37). Susceptibility to colistin was accessed using the Colispot test (38).

The production of ESBL/AmpC enzymes was assessed using a D68C1 AmpC and ESBL detection set (Mast Diagnostics, UK). Results of susceptibility testing for all tested antibiotics and their interpretation can be found in Table S1. Genomic DNA was extracted and purified using NucleoSpin columns (NucleoSpin tissue; Macherey-Nagel, Germany). Fragment libraries were constructed using Nextera XT kits followed by paired-end sequencing (NovaSeq; Illumina) according to the manufacturer’s instructions. For the E. coli CE2050 strain, long-read sequencing was performed on the Sequel I platform (Pacific Biosciences, Menlo Park, CA, USA). A microbial multiplexing protocol was used for the library preparation according to the manufacturer’s instructions for sheared DNA. DNA shearing was performed using Covaris g-TUBES (Covaris, USA). No size selection was performed during the library preparation.

Assembly of sequencing reads. Raw Illumina sequencing reads were trimmed using Trimmomatic v0.36 (39), assembled with SPAdes v3.12.0 (40), and polished with Pilon v1.23 (41).
Illumina reads mapped against overlapping regions. Locations with uncertain connections were indicated by 100 N characters.

For isolate CE2050, for which long-read sequencing data were generated, the assembly and circularization of PacBio reads was performed using the Microbial Assembly pipeline offered by the SMRT Link v8.0 software using minimum seed coverage of 30×. The closed genome was polished with Pilon v1.23 (41).

**Comparative analysis, SNP calling, and phylogenetics.** The assembled data were annotated using Prokka v1.14.1 (45) and analyzed for the presence of antimicrobial resistance genes and plasmid replicons using ResFinder (46) and PlasmidFinder (47), respectively. **bla**<sub>CTX</sub>-carrying I1 plasmid sequences as well as F plasmid replicons were subjected to plasmid MLST (pMLST) and replicon sequence typing (RST) analysis, respectively (47). Genomes were also checked for the presence of virulence-associated genes using our custom database of ExPEC virulence gene sequences (see Table S2). Genomes were interrogated for the presence of prophage regions via PHASTER (48), and insertion sequences, via ISfinder (49). Initial comparative analysis was conducted using QUAST v5.0.2 (43) with manual inspection of genomic discrepancies via Icarus (50). SNP hot spot regions were annotated using the RAST annotation pipeline, and predicted genes were functionally categorized with SEED (51).

The assembled genomes were screened for the presence of pUTI89-like plasmids by mapping contigs against the closed reference plasmid pCE2050_A from strain CE2050. The presence of the truS5S sequence in pUTI89-like plasmids within our collection was then investigated manually by visual checks of mapped sequencing reads against the pCE2050_A reference via Tablet v1.20.12 (44). The Burrows-Wheeler Aligner (BWA) MEM algorithm (52) was used for contig as well as read mapping.

Quality-trimmed Illumina reads were mapped to the closed chromosome of CE2050 using Bowtie 2 v2.3.4.2 (53). SNPs were then detected with VarScan v2.4.3 (54) using the following parameters: minimum read depth of 8, minimum base quality of 20, variant allele frequency of ≤0.08. Variant sites occurring in phage and repetitive/homologous regions as well as sites in which at least one sample had a read depth below 8 were discarded from subsequent phylogenetic analysis. SNP distances among samples were calculated with the snp-dist tool v0.7.0 (https://github.com/tseemann/snp-dist). Based on called SNP sites, QUAST metrics and PlasmidFinder results, several samples (three pairs and a single trio) were considered identical. Duplicate genomes (n = 5; 4 isolates from our gull collection and a single gull isolate from Enterobase) were discarded.

Filtered SNP sites from isolates with fewer than 100 SNP (n = 81) differences from the CE2050 chromosome were concatenated and analyzed using jModelTest v2.1.10 (55) to estimate the best-fitting model of nucleotide substitution. Using Akaike as well as Bayesian criteria, the general time-reversible (GTR) substitution model was determined as the best fit for the maximum-likelihood (ML) analysis. ML tree topology was then inferred using RAxML v8.2.10 (56) with 500 rapid bootstrap replications. Using Akaike as well as Bayesian criteria, the general time-reversible substitution model was determined as the best model of nucleotide substitution. ML tree topology was then inferred using RAxML v8.2.10 (56) with 500 rapid bootstrap replications. Using Akaike as well as Bayesian criteria, the general time-reversible substitution model was determined as the best model of nucleotide substitution.

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