Molecular characterization of a 21.4 kilobase antibiotic resistance plasmid from an hemolytic Escherichia coli O108:H-human clinical isolate

Fay E. Dawes
University of Wollongong, fdawes@uow.edu.au

Dieter M. Bulach
Monash University

Alexander Kuzevski
University Of Melbourne

Karl A. Bettelheim
University Of Melbourne

Carola Venturini
University of Queensland, cv98@uow.edu.au

See next page for additional authors

Publication Details
Molecular characterization of a 21.4 kilobase antibiotic resistance plasmid from an hemolytic Escherichia coli O108:H-human clinical isolate

Abstract
This study characterizes the 21.4 kilobase plasmid pECTm80 isolated from Escherichia coli strain 80, an α hemolytic human clinical diarrhoeal isolate (serotype O108:H-). DNA sequence analysis of pECTm80 revealed it belonged to incompatibility group X1, and contained plasmid partition and toxin-antitoxin systems, an R6K-like triple origin (ori) replication system, genes required for replication regulation, insertion sequences IS1R, ISEc37 and a truncated transposase gene (Tn3-like ΔtnpA) of the Tn3 family, and carried a class 2 integron. The class 2 integron of pECTm80 contains an intact cassette array \textit{dfrA1-sat2}, encoding resistance to trimethoprim and streptothricin, and an \textit{aadA1} gene cassette truncated by the insertion of IS1R. The complex plasmid replication system includes α, β and y origins of replication. Pairwise BLASTn comparison of pECTm80 with plasmid pE001 reveals a conserved plasmid backbone suggestive of a common ancestral lineage. Plasmid pECTm80 is of potential clinical importance, as it carries multiple genes to ensure its stable maintenance through successive bacterial cell divisions and multiple antibiotic resistance genes.

Keywords
plasmid, resistance, antibiotic, kilobase, 4, 21, characterization, h, hemolytic, escherichia, coli, human, clinical, isolate, molecular, o108, CMMB

Disciplines
Life Sciences | Physical Sciences and Mathematics | Social and Behavioral Sciences

Publication Details

Authors
Fay E. Dawes, Dieter M. Bulach, Alexander Kuzevski, Karl A. Bettelheim, Carola Venturini, Steven P. Djordjevic, and Mark J. Walker

This journal article is available at Research Online: http://ro.uow.edu.au/scipapers/4433

Fay E. Dawes¹, Dieter M. Bulach², Alexander Kuzevski³, Karl A. Bettelheim³⁴, Carola Venturini⁴, Steven P. Djordjevic⁵⁶, Mark J. Walker¹⁴*

¹ School of Biological Sciences, University of Wollongong, New South Wales, Australia, ²Department of Microbiology and Victorian Bioinformatics Consortium, Monash University, Clayton, Victoria, Australia, ³Microbiological Diagnostic Unit, Department of Microbiology and Immunology, University of Melbourne, Parkville, Victoria, Australia, ⁴School of Chemistry and Molecular Biosciences and the Australian Infectious Diseases Research Centre, University of Queensland, Queensland, Brisbane, Australia, ⁵Microbiology and Immunology Section, Industry and Investment NSW, Elizabeth Macarthur Agricultural Institute, Menangle, New South Wales, Australia, ⁶The Ithree Institute, University of Technology, Sydney, New South Wales, Australia

Abstract

This study characterizes the 21.4 kilobase plasmid pECTm80 isolated from Escherichia coli strain 80, an α-hemolytic human clinical diarrheal isolate (serotype O108:H-). DNA sequence analysis of pECTm80 revealed it belonged to incompatibility group X1, and contained plasmid partition and toxin-antitoxin systems, an R6K-like triple origin (ori) replication system, genes required for replication regulation, insertion sequences IS1R, IS3Ec37 and a truncated transposase gene (Tn3-like \( \lambda \text{tnpA} \)) of the Tn3 family, and carried a class 2 integron. The class 2 integron of pECTm80 contains an intact cassette array \( \text{dfrA1-sat2} \), encoding resistance to trimethoprim and streptothricin, and an \( \text{aadA1} \) gene cassette truncated by the insertion of \( \text{IS15} R \). The complex plasmid replication system includes \( \alpha \), \( \beta \) and \( \gamma \) origins of replication. Pairwise BLASTn comparison of pECTm80 with plasmid pE001 reveals a conserved plasmid backbone suggestive of a common ancestral lineage. Plasmid pECTm80 is of potential clinical importance, as it carries multiple genes to ensure its stable maintenance through successive bacterial cell divisions and multiple antibiotic resistance genes.


Introduction

The dissemination of multi-drug resistant bacteria is a serious and growing global health threat. Infections caused by multi-drug resistant pathogens that fail to respond to treatment, often result in prolonged illness and an increased risk of death. Understanding the molecular mechanisms that facilitate the clustering and horizontal transfer of antibiotic resistance genes is essential to the development of strategies that address this growing problem in the treatment of infectious diseases.

Horizontal gene transfer of antibiotic resistance genes may occur via various genetic elements including transformation or conjugation of plasmids, mobilizable plasmids, conjugative transposons and phages [1,2]. Plasmids belonging to the incompatibility (Inc) group X have been implicated in the acquisition and spread of antibiotic resistance-transposons such as Tn7, Tn3 and Tn21 in pathogenic enterobacteria [3,4]. IS elements may facilitate the dissemination of resistance genes and participate in chromosomal and plasmid rearrangement, integration and excision [5]. Integrons play an important role in the emergence of multi-drug resistant pathogens by functioning as mobile gene cassette capture and expression systems.

Class 2 integron screening studies frequently identify the Tn7-type cassette array of \( \text{dfrA1-sat2-aadA1} \), which confers resistance to trimethoprim, streptothricin, streptomycin and spectinomycin. Class 2 integrons are typically located at a unique site near the end of the non-replicative transposon Tn7 or related transposons including Tn1825, Tn1826 and Tn4132, which provide a means for their mobilization [6,7]. This study characterizes an Escherichia coli isolate that harbors an antibiotic resistance plasmid containing a class 2 integron.

Methods

Strain isolation and characterization

The Escherichia coli isolate characterized in this study, designated strain 80, was recovered from a patient with clinical diarrheal and submitted to the Microbiological Diagnostic Unit (MDU, Public Health Laboratory, Department of Microbiology and Immunology, University of Melbourne, Victoria, Australia). The strain was identified as E. coli by culture in specialized media, whilst O- and H-serotyping was performed as previously described [8]. Carriage of virulence determinants by E. coli strain 80 was examined by
Class 2 integron detection

Class 2 integron carriage was detected by PCR screening for the intI2 gene using primers [12] and cycling conditions [13] described previously. Confirmation of DNA integrity and strain identification as E. coli was achieved by PCR of the E. coli-specific universal stress protein A (uspA) gene. PCR amplification of the uspA gene was carried out simultaneously with amplification of the intI2 gene using previously described primers EC2 [14] and FD-uspAF [13]. E. coli strain DH5α harboring the plasmid pMAQ612 (ampicillin\(^{\text{r}}\); intI2 from Tn7 cloned into pUC18) was used as the positive control for PCR [15]. Southern hybridization of plasmid DNA versus total genomic DNA was performed to establish the genomic location of the intI2 gene. Genomic DNA was extracted using the DNeasy tissue kit (Qiagen) and digoxigenin (DIG)-labelled intI2 PCR product amplified using the primers Int2.F and Int2.R [12] was used as a probe.

Nucleotide sequence and annotation of pECTm80

Plasmid DNA isolated from E. coli strain 80 was transformed by electroporation into E. coli JM109 using standard methods [16] and the complete DNA sequence of the plasmid, designated pECTm80 was determined. DNA sequencing was performed according to the manufacturer’s instructions using the BigDye Terminator v3.1 cycle sequencing kit (Perkin-Elmer) and the 3130 Terminator v3.1 cycle sequencing kit (Perkin-Elmer) and the 3130 Genetic Analyzer capillary sequencer (Applied Biosystems). Both strands of the plasmid were sequenced by employing a primer walking strategy. To facilitate sequencing, HindIII digested plasmid fragments of 2.6 kb, 6.8 kb and 12.6 kb were cloned into pUC18 (Fermentas). Sequencing of pECTm80 proceeded preferentially from the original plasmid and from the recombinant plasmids containing pECTm80 inserts as required. Plasmid DNA was extracted using the plasmid maxi kit (Qiagen) or the Wizard Miniprep DNA Purification System (Promega). Oligonucleotide primers were designed with the Primer3 program [17] and synthesized by Sigma, Sydney, Australia.

Contig Express by Vector NTI Advance 10 (Invitrogen) was utilized to align contiguous sequences. The CDS (coding sequences) were identified using the gene finder program GeneMarkS [18]. Annotation of CDS was achieved by performing BLAST searches available from NCBI and IS BLAST server (http://www.is.biotoul.fr). Annotated sequences were visualized using Artemis (version 13.2) [19]. Comparative DNA analysis was performed by visualizing pairwise BLASTn comparisons with the Easyfig program available at http://easyfig.sourceforge.net/ [20].

Plasmid conjugation assay

The ability to transfer resistance by conjugation was performed as previously described [21]. Matings were carried out with the transformant E.coli strain JM109 (pECTm80) and recipient E. coli strain 294 Rif\(^{\text{r}}\) Nal\(^{\text{r}}\) (rifampicin and nalidixic acid resistant; \(\beta\)-galactosidase positive) [22] with and without the addition of E.coli HB101 containing the conjugal helper plasmid pRK600 Cm\(^{\text{r}}\) (chloramphenicol resistant) [23].

Results

O- and H-serotyping revealed that E. coli strain 80 possessed the serotype O108:H-. Strain 80 was found to produce \(\alpha\)-haemolysin but did not produce either of the Shiga toxins. E. coli strain 80 displayed resistance to multiple antibiotics including streptomycin (25 µg ml\(^{-1}\)), tetracycline (20 µg ml\(^{-1}\)), sulfathiazole (550 µg ml\(^{-1}\)), trimethoprim (50 µg ml\(^{-1}\)), kanamycin (10 µg ml\(^{-1}\)), nalidixic acid (50 µg ml\(^{-1}\)), spectinomycin (50 µg ml\(^{-1}\)), gentamicin (2.5 µg ml\(^{-1}\)) and ciprofloxacin (2 µg ml\(^{-1}\)).
Table 1. Identification of CDS in the nucleotide sequence of pECTm80.

<table>
<thead>
<tr>
<th>Gene/ORF</th>
<th>Strand</th>
<th>pECTm80 coordinates</th>
<th>Function/name of protein</th>
<th>Accession no.</th>
<th>Plasmid</th>
<th>% Identity</th>
</tr>
</thead>
<tbody>
<tr>
<td>ddp3 (taxO)</td>
<td>_</td>
<td>3–425</td>
<td>DNA distortion protein 3- DNA transfer element</td>
<td>JF776874.1</td>
<td>1</td>
<td>100</td>
</tr>
<tr>
<td>bis</td>
<td>_</td>
<td>496–939</td>
<td>Replication initiation protein</td>
<td>JF776874.1</td>
<td>1</td>
<td>100</td>
</tr>
<tr>
<td>repB (repX)</td>
<td>_</td>
<td>967–1818</td>
<td>Iteron binding initiator of plasmid replication</td>
<td>JF776874.1</td>
<td>1, 2, 3</td>
<td>100</td>
</tr>
<tr>
<td>stbD</td>
<td>+</td>
<td>2800–3128</td>
<td>Stability protein StbD</td>
<td>JF776874.1</td>
<td>1, 2, 3, 4, 5, 6</td>
<td>100</td>
</tr>
<tr>
<td>stbE</td>
<td>+</td>
<td>3121–3399</td>
<td>Stability protein StbE</td>
<td>JF776874.1</td>
<td>1, 4</td>
<td>100</td>
</tr>
<tr>
<td>orf01</td>
<td>+</td>
<td>3548–3871</td>
<td>Conserved hypothetical protein</td>
<td>JF776874.1</td>
<td>1, 4, 6</td>
<td>100</td>
</tr>
<tr>
<td>orf02</td>
<td>+</td>
<td>3904–4157</td>
<td>Conserved hypothetical protein</td>
<td>JF776874.1</td>
<td>1, 4</td>
<td>100</td>
</tr>
<tr>
<td>orf03</td>
<td>+</td>
<td>4145–4366</td>
<td>Conserved hypothetical protein</td>
<td>JF776874.1</td>
<td>1, 4</td>
<td>100</td>
</tr>
<tr>
<td>orf04</td>
<td>+</td>
<td>4360–4788</td>
<td>Conserved hypothetical protein</td>
<td>JF776874.1</td>
<td>1, 4</td>
<td>99</td>
</tr>
<tr>
<td>orf05</td>
<td>+</td>
<td>4831–5010</td>
<td>Conserved hypothetical protein</td>
<td>JF776874.1</td>
<td>1, 4</td>
<td>100</td>
</tr>
<tr>
<td>orf06</td>
<td>+</td>
<td>5032–5229</td>
<td>Conserved hypothetical protein</td>
<td>JF776874.1</td>
<td>1, 4, 5, 6</td>
<td>100</td>
</tr>
<tr>
<td>orf07</td>
<td>+</td>
<td>5242–5511</td>
<td>Conserved hypothetical protein</td>
<td>JF776874.1</td>
<td>1, 2, 4</td>
<td>100</td>
</tr>
<tr>
<td>orf08</td>
<td>_</td>
<td>5485–5789</td>
<td>Conserved hypothetical protein</td>
<td>JF776874.1</td>
<td>1</td>
<td>100</td>
</tr>
<tr>
<td>ddp2 (taxA)</td>
<td>+</td>
<td>5840–6382</td>
<td>DNA distortion protein 1-DNA transfer auxiliary protein</td>
<td>JF776874.1</td>
<td>1</td>
<td>100</td>
</tr>
<tr>
<td>ddp2 (taxC)</td>
<td>+</td>
<td>6382–7359</td>
<td>DNA distortion protein 2- DNA transfer relaxase</td>
<td>JF776874.1</td>
<td>1</td>
<td>100</td>
</tr>
<tr>
<td>ISeC37 trpA</td>
<td>_</td>
<td>7545–8729</td>
<td>Putative NLS-Histone family</td>
<td>AE005674.2</td>
<td>NA</td>
<td>97</td>
</tr>
<tr>
<td>mdBA</td>
<td>+</td>
<td>9113–9385</td>
<td>Putative DNA-binding protein</td>
<td>DQ115387.2</td>
<td>5, 6</td>
<td>100</td>
</tr>
<tr>
<td>upfA</td>
<td>+</td>
<td>9439–9699</td>
<td>UpfA conserved hypothetical protein</td>
<td>JF776874.1</td>
<td>1, 3</td>
<td>100</td>
</tr>
<tr>
<td>orf09</td>
<td>_</td>
<td>10269–10528</td>
<td>Conserved hypothetical protein</td>
<td>JF776874.1</td>
<td>1</td>
<td>100</td>
</tr>
<tr>
<td>orf10</td>
<td>+</td>
<td>10458–10601</td>
<td>Conserved hypothetical protein</td>
<td>SCH_O83</td>
<td>JF776874.1</td>
<td>1</td>
</tr>
<tr>
<td>orf11</td>
<td>+</td>
<td>10671–10847</td>
<td>Conserved hypothetical protein</td>
<td>JF776874.1</td>
<td>1, 2, 3, 5, 6</td>
<td>100</td>
</tr>
<tr>
<td>orf12</td>
<td>_</td>
<td>10933–11751</td>
<td>Conserved hypothetical protein</td>
<td>JF776874.1</td>
<td>1</td>
<td>100</td>
</tr>
<tr>
<td>orf13</td>
<td>_</td>
<td>12001–12525</td>
<td>Conserved hypothetical protein</td>
<td>JF776874.1</td>
<td>1</td>
<td>100</td>
</tr>
<tr>
<td>Tn3-like D tnpA E</td>
<td>_</td>
<td>12697–13368</td>
<td>Transposase truncated N-terminus by Tn7</td>
<td>EU330199</td>
<td>NA</td>
<td>100</td>
</tr>
<tr>
<td>int2</td>
<td>_</td>
<td>14309–15283</td>
<td>Class 2 integrase</td>
<td>AB188272</td>
<td>NA</td>
<td>100</td>
</tr>
<tr>
<td>dfrA1</td>
<td>+</td>
<td>15615–16088</td>
<td>Dihydrofolate reductase</td>
<td>AB188272</td>
<td>NA</td>
<td>100</td>
</tr>
<tr>
<td>sat2</td>
<td>+</td>
<td>16183–16704</td>
<td>Streptothricin acetyltransferase</td>
<td>AB188272</td>
<td>NA</td>
<td>100</td>
</tr>
<tr>
<td>JaaadA1</td>
<td>+</td>
<td>16765–17244</td>
<td>Aminoglycoside adenylytransferase</td>
<td>AB188272</td>
<td>NA</td>
<td>100</td>
</tr>
<tr>
<td>IS1R insA</td>
<td>+</td>
<td>17295–17569</td>
<td>InsA transcription repressor</td>
<td>CF002890.1</td>
<td>NA</td>
<td>98</td>
</tr>
<tr>
<td>IS1R insB</td>
<td>_</td>
<td>17549–19792</td>
<td>None</td>
<td>AE005674.2</td>
<td>NA</td>
<td>98</td>
</tr>
<tr>
<td>IS1R insAB'</td>
<td>+</td>
<td>17295–17548</td>
<td>IS1R transposase InsAB' (frame shift product)</td>
<td>CP002890.1</td>
<td>NA</td>
<td>98</td>
</tr>
<tr>
<td>orf14</td>
<td>_</td>
<td>18365–19018</td>
<td>Partitioning (par)-Resolvase</td>
<td>JF776874.1</td>
<td>1</td>
<td>100</td>
</tr>
<tr>
<td>parA (yafB)</td>
<td>_</td>
<td>19381–20040</td>
<td>Plasmid partition protein A</td>
<td>JF776874.1</td>
<td>1, 5</td>
<td>100</td>
</tr>
<tr>
<td>yafA</td>
<td>+</td>
<td>20116–20421</td>
<td>Conserved hypothetical protein</td>
<td>YafA</td>
<td>JF776874.1</td>
<td>1</td>
</tr>
<tr>
<td>orf15</td>
<td>+</td>
<td>20449–21423</td>
<td>Conserved hypothetical protein</td>
<td>JF776874.1</td>
<td>1</td>
<td>100</td>
</tr>
</tbody>
</table>

*Accession numbers provided represent the results of BLAST searches (NCBI and IS BLAST server) showing the highest identity to the query sequence. The accession number for pE001 is given when there is greater than one BLAST hit at 100% ID to pECTM80. BPlasmids showing 100% ID to pECTM80 are represented as follows: 1, pE001 (JF776874.1); 2, R485 (HE577112.1); 3, pMAS52027 (FJ666132.1); 4, pOLA52 (EU370913.1); 5, pOU1114 (DQ115387.2); 6, pSE34 (EU219533.1); and 7, pMCC7-H23 (EFS387837.1). Boundaries of mobile elements in the nucleotide sequence of pECTm80 are as follows: CISEc37, 7355–9182; DTn3-like, 12671–18314; FIS1R, 17240–18007. ETn3-like D tnpA gene showed 64% to 76% identity to transposase genes of the Tn3 family of transposons (subgroups Tn501 and Tn3) and the IS elements IS5ba14, and IS5so9 of the Tn3 family.

doi:10.1371/journal.pone.0034718.t001
antitoxin systems, a complex replication system, genes required for conjugative transfer and replication regulation, a truncated Tn7 class 2 integron, insertion sequences and a truncated Tn3 family IS element or transposon remnant. An IS1 isoform that showed 98% nucleotide identity to IS1R (GenBank J01730) and a new IS element ISEc37 belonging to the IS91 family were identified in pECTm80. ISEc37 is 95% identical to IS91 (GenBank X17114) and the encoded transposases show 95% amino acid sequence identity.

An IS1R-mediated deletion is observed in the Tn7-like class 2 integron. This class 2 integron contains intact gene cassettes dfrA1 and att2 encoding resistance to trimethoprim and streptomycin, and an adaA1 cassette gene (AdaA1) truncated by the insertion of IS1R. The open reading frames (ORFs) ybaA (orfY), ybfA, ybfB, and Tn7 transposition genes tsaABCDE, which are usually found at the 3' end of class 2 integrons, were also deleted by IS1R insertion. Consistent with other class 2 integrons, the integrase gene intI2 contains a premature in-frame stop codon, which typically encodes a defective IntI2 protein [24].

A Tn3 family transposase gene (Tn3-like Δp64), located at the 5' end of the class 2 integron, contains a Tn7-mediated truncation. An 8 bp IR at the left end of Tn7 (IR57) is adjacent to the Tn3-like Δp64 deletion site. The complete ATn3-like Δp64-A Tn7 class 2 integron-IS1R gene configuration is flanked by Tn9 38 bp IRs (IR9p-Tn9 and IR9p-Tn9c). The 5 bp direct repeats (TATAT) characteristic of Tn3 insertion are located adjacent to the Tn9 38 bp IRs. BLAST results (NCBI and IS BLAST server), indicate Tn3-like Δp64 is a remnant of a transposon or IS element of the Tn3 family. Tn3-like Δp64 showed 64–76% identity to transposase genes of transposons belonging to the Tn3 family (subgroups Tn501 and Tn3; GenBank: X90708,2 and Y00502,1) and IS elements ISShal1,4, and ISShal9 (IS Finder; GenBank NC_009502,1 and NC_004349) also of the Tn3 family. The highest identity was displayed to ISShal1,4 and Tn2501 (76% identity and 87% query coverage).

The replication system identified in pECTm80 consists of an origin (ori) of plasmid replication region that spans 7.6 kb. This region contains α, β and γ origins of replication, three DNA distortion protein genes (ddp1, ddp2 and ddp3) encoding proteins involved in conjugative transfer and the regulation of replication [25,26], and the initiation transfer genes bis and repB [27]. oriTα and oriTβ were identified within the α and β long inverted repeat nucleotide sequences of pECTm80.

Nucleotide sequence of pECTm80 showed 99–100% identity (65% pECTm80 coverage) with a 38.6 kb IncX1 plasmid, pE001, isolated from E. coli strain 2161 in broiler meat in Denmark (GenBank JF776874.1) [28]. A diagram of the pairwise BLASTn comparison between plasmids pECTm80 and pE001 is given in Figure 2. pECTm80 also showed 90–98% identity to the 34.5 kb plasmid pOU1114 (GenBank NC_010421) isolated from Salmonella enterica serovar Dublin (49% pECTm80 coverage and 10 CDS) [29] and 89–99% identity to the 51.6 kb plasmid pOLAS2 (GenBank EU370913) isolated from Escherichia coli (39% pECTm80 coverage and 14 CDS) [30]. The plasmids pECTm80 and pE001 shared a common backbone showing 99.9% nucleotide sequence identity. Plasmid pE001 did not contain Tn3-like Δp64, a class 2 integron, ISEc37 nor IS1R found in pECTm80 while pECTm80 did not contain the puaX operon, found on pE001 and other conjugative IncX1 plasmids [29].

**Discussion**

This study characterizes pECTm80, a 21.4 kilobase IncX1 plasmid containing a truncated class 2 integron from an *Escherichia coli* O108:H- human clinical isolate. *E. coli* strain 80 serogroup O108 was non-motile (H-) and produced α-hemolysin, a potent enterotoxin that is known to enhance virulence in a number of clinical infections [31], although strain 80 did not produce either of the Shiga toxins. DNA sequence analysis of pECTm80 revealed a Tn7 class 2 integron with complete dfaA1 and att2 gene cassettes and an IS1R-mediated deletion resulting in truncation of the adaA1 gene cassette and deletion of the ORFs ybaA (orfY), ybfA and ybfB, and Tn7 transposition genes tsaABCDE.

Several key features of the integron-containing plasmid pECTm80 include the presence of plasmid partitioning and segregational stability genes to ensure the stable maintenance of the plasmid through successive bacterial cell divisions and a highly-regulated DNA replication system consisting of three distinct origins of replication α, β and γ, that are controlled and activated by plasmid- and host-encoded genes [32]. Plasmid pECTm80 contains plasmid partitioning genes and toxin-antitoxin system genes, which ensure its vertical transfer. The plasmid partitioning protein ParA is encoded by the parA (gyrB) gene and a partitioning (par)-resolvase by orf14 [33]. The toxin-antitoxin system, encoded by stbD and stbE genes, promotes segregational stability by compromising the survival of plasmid-free daughter cells.

**Figure 2. Comparison of pECTm80 with the *E. coli* plasmid pE001.** Pairwise BLASTn comparisons between the plasmids pECTm80 and pE001 were visualized using the Easyfig program [23]. Regions of nucleotide identity are connected by yellow blocks. The yellow color gradient indicates the extent of similarity as shown in the color scale on the right. Functions of CDS in pE001 have been taken from BLAST matches and existing annotation. Functional categories of predicted CDSs include: insertion sequence/transposon transposases (red); conjugal transfer (green); plasmid maintenance and stability (blue); gene expression modulation (light blue); class 2 integron intI2 gene and gene cassettes (yellow); type IV secretion system (mauve); blaTEM-52 (yellow) and hypothetical proteins of unknown function (grey). Scale bar represents 3000 base pairs.

doi:10.1371/journal.pone.0034718.g002
cells that may arise during cell division [34,35,36]. The complex replication system of pECTm80 is a strategy for increasing plasmid mobility to a range of hosts [37]. A study characterizing the host-range of an IncX plasmid of R6K lineage found the plasmid was established in 9 of 16 species tested [30]. Plasmid pECTm80 belongs to the IncX1 group, showing 96% identity to the IncX1 plasmid R485 replication origin region (GenBank M11688.1). Division of the IncX group into IncX1 and IncX2 arose following discovery of in vivo incompatibility between the IncX plasmids R485 and R6K, and lack of extensive similarity over long nucleotide stretches in these replicons [35]. R6K is the sole well-characterized member of the IncX2 group [39], although a second IncX2 plasmid was recently identified [40].

Nucleotide sequence analysis of pECTm80 also revealed this plasmid carries genes and sites necessary for plasmid mobilization including ddx1, ddx2, ddx3, bsi and repβ genes, and α and β oriTs. However pECTm80 was neither conjugative nor mobilized by the helper plasmid pRK600. pECTm80 did not contain genes of the pilX operon that encode a Type IV secretion system essential for mate-pair formation and conjugal transfer, which are found on conjugative IncX1 plasmids including pE001, pOLA52 and pOU1114 [29,30,41]. The absence of pilX genes may explain the non-conjugative properties of pECTm80. The deletion of the pilX operon may have resulted due to insertion of the IS37 element into pECTm80.

Close association of the Tn7 class 2 integron with other mobile DNA elements including insertion sequences and transposons is reported in this study. The complex and dynamic interaction of mobilizable elements described here, act to increase their potential for gene shuffling [42]. The IS7/α-mediated truncation of the Tn7 class 2 integron and the IS7/α-mediated truncation of the Tn3-like ÑtpA are observed. The mosaic Tn3-like ÑtpA-ΔTn7 class 2 integron-IS7/α gene configuration is flanked by Tn7 38 bp IRs (IRb-Tn3 and IRa-Tn7) and 5 bp target-site direct repeats (TATA). The presence of IS7/α and a Tn3 family transposon IS remnant with the class 2 integron implicates these DNA elements in the evolution of this unique integron. Sequential addition of Tn3-like (IS/transposon), Tn7 and IS7/α and the resulting insertion-mediated deletions are hypothesized to have occurred in pECTm80. The sequential addition of Tn3 and Tn7 to IncX1 plasmids leading to the creation of complex R-plasmids has been previously described [39].

Comparative DNA sequence analysis of pECTm80 with the conjugal IncX1 plasmid pE001, also isolated from an E. coli strain, provides insight into how this plasmid and its resident class 2 integron evolved. pECTm80 displayed a conserved backbone with archetypical IncX1 regions for replication and plasmid stability found in pE001 [29]. The shared essential regions for replication and plasmid stability and the high-level of identity described support the hypothesis that these plasmids have arisen from a common ancestral plasmid. The absence of IS37, IS7/α, the Tn3 family transposon/IS remnant, and the class 2 integron in pE001 may suggest the integration of these elements into a common ancestral plasmid occurred to create pECTm80. Insertion sequences IS37/α and IS7/α are intact in pECTm80 and may represent recent insertions into the plasmid. Although the exact mechanism of evolution of the IncX1 plasmid pECTm80 and its resident class 2 integron remains unknown, multiple insertion, deletion and rearrangement events are likely to have occurred, as suggested by the proximity of several IS elements.

Acknowledgments

The authors acknowledge Prof. H. Stokes (University of Technology Sydney, NSW, Australia), Dr. S. Partridge, Assoc. Prof. J. Iredell and Prof. R. Hall (University of Sydney, NSW, Australia) for the provision of a class 2 integron positive control E. coli strain. The authors thank Assoc. Prof. G. Hogg, director of MDU Public Health Laboratory University of Melbourne, Melbourne, VIC, Australia.

Author Contributions

Conceived and designed the experiments: FED SPD MJW. Performed the experiments: FED AK KAB CV. Analyzed the data: FED DMB AK KAB SPD MJW. Contributed reagents/materials/analysis tools: DMB AK KA SPD MJW. Wrote the paper: FED.

References

13. Davies FE, Kuzevski A, Bettelheim KA, Hornitzky MA, Djordjevic SP, et al. (2010) Distribution of class 1 integrons with IS26-mediated deletions in their-


