ANTIMICROBIAL RESISTANCE

Characterisation of drug resistance gene cassettes associated with class 1 integrons in clinical isolates of Escherichia coli from Taiwan, ROC

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The presence of class 1 integrons in clinical isolates of Escherichia coli was detected by PCR. Of 164 E. coli isolates from Kaohsiung, 54 (32%) carried class 1 integrons, with inserted DNA regions of 1–3 kb. These integrons were located on plasmids, as demonstrated by Southern hybridisation. DNA sequencing was used to identify the genetic content of the integron-variable regions. Different class 1 integrons contained various numbers, kinds and combinations of gene cassettes within their variable regions. These gene cassettes included those encoding resistance to trimethoprim (dfr5A, dfrV, dfr12 and dfr17), aminoglycosides (aadA1u, aadA2, aadA4 and aadB), chloramphenicol (cmA), erythromycin (ermA2) and β-lactams (blaP1). An integron carrying three inserted cassettes – dfr12-orfF-aadA2 – was present in 33 (61%) of the 54 isolates with class 1 integrons. Gene cassettes encoding resistance were expressed phenotypically. The results indicate that class 1 integrons are widespread in clinical E. coli isolates in Taiwan. The types, combinations and frequency of the gene cassettes in integrons may reflect the specific selective pressures to which the isolates were exposed and could provide useful surveillance data for relation to antibiotic usage information.

Introduction

The rapid dissemination of drug-resistant bacteria is an alarming and increasing problem, complicating the treatment of infections. Much of the problem is the result of antibiotic resistance genes transferring among bacterial species, carried by plasmids and transposons [1]. In recent years, a third mechanism for the dissemination of antibiotic resistance genes has been discovered; this involves integrons, novel DNA elements that mediate the integration of antibiotic resistance genes through site-specific recombination [1–3]. This recombination system depends on two determinants: an int gene encoding a site-specific DNA integrase (Int) [1, 2], and an adjacent site, attI, which is recognised by the integrase and acts as the receptor for the inserted genes [3]. The inserted genes, which mostly encode antibiotic resistances, are contained in mobile elements called gene cassettes [4]. Each gene cassette consists of a coding region and an integrase-specific recombination site known as a 59-base element [4, 5]. Most of the gene cassettes lack their own promoters, and integrons act as natural expression vectors by supplying a common promoter, Pint, located in the conserved sequences upstream of the inserted genes [2–4].

Four types of integrons, with different int genes, have been identified. Most integrons from clinical isolates belong to class 1 [6] and consist of two conserved segments. A 5′-conserved segment contains the int gene (int1), the attI site and the common promoters Pint [3, 4], whereas the 3′-conserved segment includes an antiseptic resistance gene (qacEΔ1), a sulphonamide resistance gene (sul1) and an open reading frame (orf5) of unknown function [2]. The central variable region includes different combinations of inserted gene cassettes. More than 40 different antibiotic resistance cassettes have been characterised from integrons, which consequently can allow their bacterial hosts to become resistant to broad spectra of antimicrobial agents [4].

Several studies have investigated the prevalence of integrons in clinical isolates and have found them to be widespread [7–9]. However, the kinds of gene cassettes

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inserted into the integrons were not identified, except in the work of Sallen et al. [7]. In Taiwan, resistance to various antibiotics is very common in clinical isolates, often more so than in Western countries [10]. However, the frequency, characteristics, and roles of integrons and gene cassettes have not yet been investigated. In this study, *Escherichia coli* isolates from Kaohsiung, Taiwan, were investigated to demonstrate the prevalence and characteristics of class 1 integrons and the gene cassettes in these organisms, relating the results to the phenotypic expression of resistance.

**Materials and methods**

**Bacterial strains**

A total of 104 *E. coli* isolates was collected consecutively in 1993 and 1994 from the Kaohsiung Medical University Hospital, a medical centre in southern Taiwan. They were isolated from various clinical specimens from diverse units of the hospital. Once identified, the isolates were preserved at −70°C in Tryptic Soy Broth (Difco Laboratories, Detroit, MI, USA) containing glycerol 15% v/v.

**PCR analysis**

Integrons were detected by PCR with primers specific for the 5′-conserved segment (5′-CS; 5′-GGCATCC AAGCAGAAG-3′) and the 3′-conserved segment (3′-CS; 5′-AAGCAGACTTGACCTGA-3′) [11]. The DNA templates for PCR were prepared as described by Bass et al. [12]. The PCR itself was performed in 20-μl volumes containing 200 μM dNTP, 1 U Taq DNA polymerase, 10× PCR buffer (TaKaRa, Shiga, Japan), 0.3 μM of each primer and template DNA in a Gene Amp 9700 PCR system (Perkin-Elmer, Norwalk, CT, USA). The PCR conditions consisted of denaturation at 95°C for 1 min, then 30 cycles of denaturation for 30 s at 95°C, annealing for 30 s at 55°C and extension for 1.5–2 min at 72°C. The PCR products were visualized after electrophoresis through an agarose 1.5–2% gel containing ethidium bromide. Plasmid pUB2401, which harbours transposon Tn21 containing the In2 class 1 integron [2–4], was used as a positive control. Plasmid CoEL1::Tn7, which harbours transposon Tn7 with a class 2 integron [2–4], was included as a negative control.

**Southern hybridisation**

Plasmid DNA was extracted by the method of Kado and Liu [13], subjected to electrophoresis through an agarose 0.8% gel (Boehringer-Mannheim GmbH, Mannheim, Germany), stained with ethidium bromide and photographed under UV transillumination. The gels were then partially depurinated in 0.25 M HCl and denatured in 0.5 M NaOH, after which the DNA was transferred to a nylon membrane. This was prehybridised, then hybridised with an oligonucleotide probe (5′-GGCTCAGACTCCTGCTGGTGC-3′) specific for the *intI1* gene [14]. This probe was labelled with DIG-dUTP. A positive hybridisation result was detected with a DIG Luminescent Detection Kit (Boehringer-Mannheim), used according to the manufacturer’s recommendations. Luminescence was documented by exposure to Kodak BioMax Light film (Kodak, Rochester, NY, USA).

**DNA sequencing and computer analysis of sequence data**

The genetic material inserted within the integron-variable regions was identified by direct nucleotide sequencing. The template amplicons were purified with a QIAquick PCR Purification Kit (QIAGEN GmbH, Hilden, Germany). When two amplicons of different lengths were obtained from a single isolate, each fragment was excised from the agarose gel and purified with a QIAquick Gel Extraction Kit (QIAGEN). Nucleotide sequencing was performed with the ABI PRISM™ Dye Terminator Cycle Sequencing Ready Reaction Kit (Perkin-Elmer), and the reactions were analysed on an ABI PRISM 377 DNA sequencer. Similarity searches with nucleotide sequences in the GenBank database were performed with the BLASTN program (National Center for Biotechnology Information, USA).

**Amplon cloning**

To further explore the expression of the integron-borne gene cassettes, an amplon representative of each of the integrons found with different gene cassettes was ligated into the pGEM-T Easy Vector (Promega, Madison, WI, USA) according to the manufacturer’s instructions and was then transformed into *E. coli* JM101 [supE thi A (lac-proAB) F' traD36 proAB+ lacIq lacZAM15] [15]. Transformants were selected on Luria-Bertani (LB) agar containing ampicillin 100 mg/L, 0.5 mM isopropylthiogalactoside and X-Gal 80 μg/ml.

**Antimicrobial susceptibility testing**

Antimicrobial susceptibility tests were performed by the disk diffusion method recommended by the National Committee for Clinical Laboratory Standards [16].

**Results**

**Presence and location of class 1 integrons in *E. coli* isolates**

Class 1 integrons were sought in 104 *E. coli* isolates from clinical specimens by PCR with the 5′-CS and 3′-CS primers, which correspond to the sequences flanking the integron-variable regions; 54 (52%) of the isolates carried detectable class 1 integron structures. The amplicon lengths, corresponding to the approximate sizes of the inserted cassette DNA, varied from 1 to 3 kb (Table 1 and Fig. 1). Among the 54 isolates, 46 yielded one amplon and eight yielded two amplicons of different sizes (Table 1 and Fig. 1). None of the *E. coli* isolates had only the integron conserved segments without inserted DNA; if such ‘empty’ integron structures were present, the resulting amplons would be c. 150 bp and would have been resolved
Table 1. Sizes and numbers of amplicons amplified from E. coli isolates with primers 5’-CS and 3’-CS

<table>
<thead>
<tr>
<th>Approximate length (kb)</th>
<th>Number of isolates (group)</th>
<th>Approximate lengths (kb)</th>
<th>Number of isolates (group)</th>
</tr>
</thead>
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<td>1.0</td>
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<td>1.0, 1.9</td>
<td>3 (6)</td>
</tr>
<tr>
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<tr>
<td>1.9</td>
<td>29 (3)</td>
<td>1.6, 1.9</td>
<td>1 (8)</td>
</tr>
<tr>
<td>2.0</td>
<td>1 (4)</td>
<td>1.6, 2.0</td>
<td>3 (9)</td>
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<tr>
<td>3.0</td>
<td>1 (5)</td>
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Fig. 1. PCR amplification of the integron-variable regions with the 5’-CS and 3’-CS primers. The amplicons were separated by electrophoresis through an agarose 1.5% gel. Lane M, size markers; lane 1, Tn251 (positive control); 2, ColE1:Tn7 (negative control); 3–11, amplicons from clinical E. coli isolates; 3, 1.0-kb amplicon; 4, 1.6-kb amplicon; 5, 1.9-kb amplicon; 6, 2.0-kb amplicon; 7, 3.0-kb amplicon; 8, 1.0-kb and 1.9-kb amplicons; 9, 1.0-kb and 2.1-kb amplicons; 10, 1.6-kb and 1.9-kb amplicons; 11, 1.6-kb and 2.0-kb amplicons.

by the electrophoresis conditions used. The PCR analysis showed that different insert sizes were present, indicating heterogeneity in the gene cassettes; nevertheless, 33 of the 54 isolates with class 1 integrons had 1.9-kb inserts.

Southern hybridisation with an oligonucleotide probe specific for the intI1 gene consistently revealed that the integrons were located on plasmids (Fig. 2).

Identification of integron-borne gene cassettes

Purified PCR products were subjected to DNA sequencing and the gene cassettes thereby identified are listed in Table 2. The integrons were classified into nine groups according to the length and the numbers of amplicons yielded by a single isolate (Table 1). Groups 1 and 2 were further divided into two subgroups, a and b, based on the kinds of gene cassettes identified (Table 2). The gene cassettes encoded antibiotic resistance determinants with the possible exception of one open reading frame encoding an unknown product, orfF. These antibiotic resistance gene cassettes included four trimethoprim resistance determinants (dfrA1, dfrV, dfr12 and dfr17) and three spectinomycin/streptomycin resistance determinants (aadA1, aadA2 and aadA4).

Other gene cassettes found included aadB, encoding gentamicin resistance; cmdA, encoding non-enzymic chloramphenicol resistance [17]; ereA2, encoding erythromycin resistance and bladP1, encoding PSE-1, a class A β-lactamase [18].

The integrons were found to contain one-to-three gene cassettes and the combinations of these are shown in Table 2. The 1.9-kb amplicon present in 33 isolates carried dfr12, orfF and aadA2 (group 3, 6 and 8). An integron containing two newly described gene cassettes, dfr17 and aadA4 [19], was found in the five isolates yielding a 1.6-kb amplicon (groups 2b and 8). The entire sequences of the dfr17 and aadA4 cassettes were deposited in GenBank, under accession no. AF170088. Eight isolates yielded two amplicons of different sizes with primers 5’-CS and 3’-CS; sequencing analysis confirmed the co-existence of two distinct integrons carrying different gene cassettes in these isolates (group 6, 7, 8 and 9).

Antimicrobial resistance in relation to integron carriage

When isolates contained a gene cassette, the corresponding antibiotic-resistant phenotype was detected.
consistently (Table 2). Further cloning and expression of the amplicons containing the cassette regions demonstrated that the transformants containing an amplicon expressed phenotypic antibiotic resistance encoded by the gene cassettes present in this amplicon. For instance, the transformants containing the 1.9-kb amplicon, comprising *dfr*12, *orfF* and *aadA2* cassettes, were resistant to trimethoprim, spectinomycin and streptomycin. Nevertheless, the integron-carried genes did not account for all the phenotypic resistances of the *E. coli* isolates. Thus, although 39 isolates were resistant to chloramphenicol and 22 were resistant to gentamicin, only one contained an integron carrying the *cmlA* and *aadB* cassettes (group 5 in Table 2). A few isolates (c. 2–7%) were resistant to cefmetazole, ceftazidime, moxalactam, aztreonam and netilmicin, drugs to which no integron-associated resistance determinants were found.

Discussion

Of 104 *E. coli* isolates, 54 (52%) had class 1 integron structures, as detected by PCR with primers specific for the CS regions flanking the inserted gene cassettes. Other reports have shown the prevalence of class 1 integrons in gram-negative clinical isolates to be c. 43% in Western and Central Europe [8], >50% in the Netherlands [20] and 59% in France [7]. Among aminoglycoside-resistant isolates from the USA, nearly 75% had integrons [11]. These results, together with those obtained in the present study, indicate that class 1 integrons are widespread in clinical isolates. International travel may contribute to their dissemination and that of their host strains. Most (61%) of the class 1 integrons found in this study carried *dfr*12, *orfF* and *aadA2* cassettes. This pattern was also reported in a urinary *E. coli* isolate from Turku, Finland [21], and in *Shigella* strains isolated in Finland, but originating from Asia [21]. The present study also revealed an integron carrying two newly characterised gene cassettes, *dfr*17 and *aadA4*. These cassettes were also found in a recent urinary *E. coli* isolate from Australia [22] and the same *aadA4* cassette (although designated as *aadA5*) was found in a class 1 integron in a porcine *E. coli* isolate from Denmark [23]. These data reveal that the same cassettes can disseminate in different bacterial species, in different human or animal hosts and in different areas of the world. Such observations underscore the concern that human, animal and
environmental microbial ecosystems are inextricably intertwined, with antibiotic resistance readily crossing ecological boundaries and spreading widely [24]. This makes control much more difficult.

Although the integrons contained different combinations of gene cassettes, some combinations in gene cassettes were frequently conserved such as dfrA-\textit{aadA1\textsubscript{1}} and dfrA2-\textit{orfF-aadA2}. This observation may indicate that these gene cassettes have become stably integrated over a long period of time. Although integrase-mediated insertion and excision of gene cassettes has been demonstrated [2, 4], Martinez-Freixo \textit{et al.} also proposed that many inserted gene cassettes may be preserved and stable [6]. Moreover, the transfer of the entire integrin, via a plasmid or transposon, is more frequent than single gene mobilisation or integration within the integrin [6]. In this study, integrons were located on plasmids, and conjugation experiments revealed that strain-to-strain transfer could occur.

All the 54 isolates with class 1 integrons contained at least one kind of \textit{aadA} cassette (\textit{aadA1\textsubscript{1}}, \textit{aadA2} or \textit{aadA4}), alone or in combination with other cassettes. \textit{aadA} genes confer resistance to spectinomycin and streptomycin and have been found frequently in integrons [6, 7]. Moreover, despite minimal use as therapeutic agents, resistances to streptomycin and spectinomycin remain prevalent among \textit{E. coli} isolates [12, 25]. It seems likely that, due to the lack of direct selection pressure, different \textit{aadA} variants have not been selected to the extent that has occurred with genes conferring resistance to trimethoprim and \(\beta\)-lactams. However, the finding of a novel spectinomycin/spectinomycin resistance gene indicated that variations in the \textit{aadA} gene do arise. Moreover, streptomycin is still used extensively in animal husbandry [26], perhaps selecting new \textit{aadA} gene variants that may spread to pathogenic bacteria isolated from man.

Some researchers propose that the \textit{aadA} gene may be the first cassette acquired by integrons [27, 28]. In the present study, most \textit{aadA} cassettes were combined with other kinds of gene cassettes, adjacent to the S\textsuperscript{5\prime}-conserved segment and thus upstream of the \textit{aadA} cassettes. To this extent the results could support the suggestion that \textit{aadA} is the first cassette acquired by integrons, because a gene cassette preferentially recombines with the \textit{attL} site, rather than into the \textit{attC} sites at the 3\textsuperscript{\prime}-ends of cassettes already present in the integron [3]. On the other hand, \textit{aadA} genes were rarely the sole cassettes in integrons. This finding is different from that of Rosser and Young [28], who found that \textit{aadA} was often the sole cassette in the variable region of integrons from coliform bacteria, \textit{Pseudomonas}-like and \textit{Vibrio}-like isolates from a natural estuarine environment. The disagreement might reflect the fact that clinical isolates are exposed to more antibiotic selection than environmental bacteria, promoting accumulation of multiple gene cassettes.
Besides aadA, other frequently found cassettes were those encoding resistance to trimethoprim (dfr), which were present in 43 (80%) of the isolates. Four dfr gene types cassettes were identified. All the 54 E. coli isolates with class 1 integrons showed resistance to sulphonamides, perhaps reflecting the fact that co-trimoxazole is frequently used to treat E. coli infection. Such specific selection pressure may favour the acquisition and maintenance of a trimethoprim resistance cassette by class 1 integrons also containing sul1 in the 3' region [23]. This selection pressure also may explain the increasing number of different types of cassette-encoded dihydrofolate reductase. For example, dfr17 is a novel trimethoprim resistance gene cassette. Furthermore, dfr cassettes were located directly behind the 3'-conserved segment. This site is the closest to the promoter, giving high-level expression and contingent resistance.

Antimicrobial susceptibility tests confirmed that the gene cassettes in the integrons contributed to resistance, as also found by others [7, 28]. Many isolates also had other antibiotic resistances not encoded by integron-associated resistance gene cassettes. These resistances may have resulted from chromosomal mutation, plasmid acquisition or the presence of other integrons besides class 1 types.

The present study revealed that class 1 integrons are widespread in clinical isolates. Their location on plasmids may contribute to horizontal dissemination of the antibiotic resistance gene cassettes. All the integrons contained antibiotic resistance gene cassettes. The study of integrons and their associated gene cassettes could provide information about which antibiotics should be used more carefully to prevent further accumulation of resistance.

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References