ISCR Elements: Novel Gene-Capturing Systems of the 21st Century?

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INTRODUCTION

Bacterial antibiotic resistance is, once again, high on the clinical and scientific agenda. The ability of bacteria to evolve antibiotic-resistant forms is impressive, as attested to by a continuing concern that untreatable “superbugs” will continue to emerge in clinical medicine. Bacteria are noteworthy for their remarkable ability to adapt to changes in their environments, not least for their adaptability in reassortting and redistributing genetic information. Bacteria possess an impressive set of tools with which to adjust the blueprint of the cell, the bacterial genome, enabling the cell to alter, add, or lose genetic information. Nowhere is this potential for change better illustrated than the bacterial riposte to antibiotics, antiseptics, disinfectants, and heavy metals used in clinical medicine. Although mutation has an important role to play in the evolution of antibiotic resistance, the predominant factor for the escalation of antibiotic resistance in more than half a century is the acquisition of antibiotic resistance genes.

The acquisition and spread of antibiotic resistance genes among bacteria that are intimately associated with humans and their domesticated animals are well documented. Data from studies examining this horizontal gene transfer indicate that it is driven by multiple systems that involve both cell-to-cell transfer and gene transfer from one DNA molecule to another. The latter can take place regardless of similarity between the donating and recipient molecules (38, 73) and includes systems such as transposons and site-specific recombination systems termed integrons (8, 9, 19). Recently it has become clear that while these systems can account for much of the movement of resistance genes between DNA molecules, they fail to explain the spread of a substantial and growing subset of resistance genes. These resistance genes are linked to sequences termed “common regions” (CRs), which are often found beyond but close to the 3’ conserved sequences of class 1 integrons. A comparative analysis of these CRs has revealed that they are related to each other and resemble an atypical class of insertion sequences (ISs), designated IS91-like. IS91 and like ele-

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FIG. 1. Genetic context of IS\textsubscript{CR1} elements found in association with antibiotic resistance genes as part of complex class 1 integrons. IS\textsubscript{CRs} are represented by yellow boxes. The qac\textsubscript{sul} 3' ends of the class 1 complex integrons are denoted by gray and black boxes, respectively. The blue box denoting the int gene (encoding the integrase) represents the 5' end of the class 1 integron. Gene cassettes forming parts of class 1 integrons are represented by white boxes with aqua vertical lines. Genes that are not part of cassettes have been given their own individual color as indicated by the key in Fig. 4. Open reading frames are indicated with open boxes and the direction of their transcription indicated with arrows. See the text for a detailed description of class 1 integrons. (A) Complex class 1 integrons In6 and In7, in which IS\textsubscript{CR1} elements were first observed. (B to F) Genetic context of IS\textsubscript{CR1} associated with trimethoprim resistance dfr\textsubscript{G} genes, ciprofloxacin resistance qnr\textsubscript{A} genes, aminoglycoside resistance genes, class C \beta-lactamase genes, and class A \beta-lactamase genes, respectively. Accession numbers of the nucleotide sequences of the various gene arrays are included on the right.
ments differ from the insertion sequence paradigm (59) in that they lack terminal inverted repeats (IRs) and are thought to transpose by a mechanism termed rolling-circle (RC) transposition (because the mechanism involves rolling-circle replication). As a consequence, they can transpose adjacent DNA sequences, mediated by a single copy of the element (96). This is unlike most IS elements, where two copies (one of which must be intact) need to flank the mobilized gene (6), as in the mobilization of the \textit{bla}_{CTX-M} genes by IS\textit{Ecp1} (75, 76). We will argue that CRs are members of an extended family of IS\textit{91} -like elements that transpose by RC transposition and are responsible for the mobilization of virtually every class of antibiotic resistance genes, including those encoding extended-spectrum β-lactamases (ESBLs), carbapenemases, and enzymes conferring broad-spectrum aminoglycoside resistance, florfenicol/chloramphenicol resistance, and resistance to trimethoprim and quinolones.

**CRs AND THEIR DISCOVERY**

Common regions were first discovered and reported in the early 1990s as a DNA sequence of 2,154 bp that was found in two complex class 1 integrons, In6 and In7 (93). These arrays possess partial duplications of the 3’ conserved sequence of the class 1 integrons (Fig. 1), between which are located the CR and antibiotic resistance genes. The sequence was initially described as a common region to distinguish it from the 5’ and 3’ conserved sequences (CS) of class 1 integrons and was found immediately downstream of a shortened 3’ CS (Fig. 1A).

Class 1 integrons are genetic elements that possess a specific recombination site, \textit{attI}, into which resistance genes, in the form of gene cassettes, can be inserted by site-specific recombination. Gene cassettes are small circular DNA molecules, approximately 1 kb in size, comprising a single gene together with a recombination site termed a 59-base element (59be) (6, 19). Each 59be has imperfect terminal IRs of approximately 20 bp, and the different sizes of 59bes reflect the fact that the central sequences differ markedly from one element to another. Each terminal IR comprises oppositely oriented putative integrase binding sites of 7 or 8 bp, designated 1L and 2L at the 5’ end and 1R and 2R at the 3’ end, separated by 5 bp and 5 or 6 bp, respectively (94). Cassette integration usually occurs at the \textit{attI} site of the integron, and the original cassette 59be is reformed whenever the gene cassette is excised from
the integron. Thus, resistance genes found on gene cassettes are potentially highly mobile (7–9, 19).

However, the gene arrays of In6 and In7 are therefore somewhat different from the more common form of class 1 integrons, which contain only one copy of the 3' CS and no CR, and are often called complex class 1 integrons in the literature (9). These common regions were also interesting in that they are intimately associated with different resistance genes carried on each integron. In the case of In6 the CR is found immediately upstream of a chloramphenicol resistance gene (catAII) (Fig. 1A), while the CR in In7 is associated with a novel trimethoprim resistance gene (dfrA10) (Fig. 1A). The particularly striking aspect of these two CR-associated resistance genes is the fact that neither is linked to 5'bes, as is typical for the overwhelming majority of resistance genes embedded in class 1 integrons. Integrons generally evolve by integration and excision of a variety of gene cassettes. Searches of available databases in 1993 detected only one other sequence with high identity to the CR, i.e., a 349-bp section of the small nonconjugative plasmid, RSF1010 (93). This plasmid is a broad-host-range (incQ) plasmid that confers resistance to sulfonamide and streptomycin and has been the basis of numerous broad-host-range vectors for recombinant DNA manipulation. This fragment of the CR is the result of truncation at both ends relative to the sequence found in In6 and In7. The sequence is related but not identical to the CR and is now known to be a degenerate copy of a closely related variant of the original CR sequence (now designated CR2 and CR1, respectively). The truncated CR in RSF1010 is also associated with a gene (sul2) encoding resistance to sulfonamide (Fig. 2) (accession no. D37825). Integrons In6 and In7 were discovered on plasmids pSa and pDG0100, respectively, with pDG0100 being a 120-kb incC group plasmid and pSa a 39-kb incW group plasmid (93). Recently, a retrospective study found an integron that is virtually identical to In6 on the incU group plasmid pAR-32, which confers resistance to sulfonamide, chloramphenicol, and streptomycin. This plasmid, which is globally distributed and identical to the
incU reference plasmid RA3, was found in a resistant *Aeromonas salmonicida* isolate that came from Japan in 1970 (92) (Fig. 1A).

The CRs in integrons In6 and In7 and the homologous CR segment of RSF1010 comprised, therefore, the sum of the information known about these elements in 1993, and for several years accumulation of information concerning them was slow (90, 93). Recently, however, there has been a major proliferation of information regarding these important genetic elements and the various antibiotic resistance genes that are intimately associated with them. Most of this information has been reported in the last 2 to 3 years, and it probably represents just a small portion of these elements associated with resistance genes. CR elements have now been found in numerous gram-negative organisms of clinical importance, as well as in a few gram-positive bacteria (2) (Fig. 1 to 4). Various studies have shown CRs to be present on both plasmids and chromosomes, and they have been found in bacteria worldwide. CRs are now known to be closely associated with many antibiotic resistance genes, including those encoding extended-spectrum class A β-lactamases, extended-spectrum class C β-lactamases, the metallo-β-lactamase (MBL) *bla*<sub>SPM-1</sub>, the extended-spectrum class D β-lactamase *bla*<sub>OXA-45</sub>, and plasmid-mediated quinolone resistance (*qnr*), as well as various trimethoprim resistance genes, the chloramphenicol resistance gene *catAII*, broad-spectrum aminoglycoside resistance genes, tetracycline resistance genes, the florfenicol resistance gene *floR*, the macrolide resistance gene *ereB*, and others (Fig. 1 to 4).

There is substantial evidence linking the origin of a subset of these resistance genes to the chromosomes of environmental bacteria (1, 86, 103), and there is considerable circumstantial evidence that CR elements are responsible for the mobility and dissemination of many genes (23, 107). CRs have also been implicated in the construction of the multidrug resistance (MDR) regions of the integrative conjugative element SXT and the *Salmonella* genomic island 1 (SGI1) and their variants (4, 15). If this is true, as we will argue, then the potential of CRs to mobilize adjacent antibiotic resistance genes may pose a considerable threat to already beleaguered anti-infective regimens, particularly for gram-negative bacteria. We contend that CRs represent a family of related mobile genetic elements that are similar to the rather unusual IS element IS91 (34). This review will itemize the available information concerning these elements and the resis-
tance genes associated with them and will attempt to give some insight into their mode of movement and their origins.

**CHARACTERIZATION OF CRs**

CRs comprise a set of related sequences. Each CR identified so far accommodates a single open reading frame (ORF). The predicted products of these putative genes, all approximately 500 amino acids long, are closely related to each other, differing by 4 to 88% in amino acid identity (Fig. 5 and 6). However, CR identity extends beyond the ORFs to include nucleotide sequences in both directions. To the left-hand end of the CR ORFs, the sequences show some homology for 240 to 250 bp before ending in a short section of conserved sequence (Fig. 7). This end of a CR sequence will be referred to as the 3' end of the CR because it is 3' to the ORF. The other end, therefore, is the 5' end.

Initial screening of the EMBL prokaryote nucleotide sequence databases with the CR sequence of In6 identified many sequences with different levels of identity to the search sequence. These were found to fall into a number of subgroups. Many sequences have been found to display identity throughout the single gene accommodated on each CR, while others

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**FIG. 5.** Phylogenetic tree (A) and sequence identity comparisons (B) of all extant ISCR elements, based on a CLUSTAL alignment with the PAM 250 matrix prepared using Lasergene DNAsStar software. Alignments were undertaken with partial amino acid sequences (250 amino acids) encompassing a central section of the putative ISCR transposases of ISCR1 to -12 (complete protein sequences are available only for ISCR1 to -5). Comparative identities of ISCR elements are given below the phylogenetic tree, and sequences of individual ISCR elements displaying most and least identity are highlighted.
show a significant degree of identity only to the sequence downstream, in some cases encompassing just the 29 bp at the 3' end of the probe sequence.

**CRs Are IS91-Like Transposable Elements**

The CR embedded in the complex class 1 integrons In6 and In7 is 2,154 bp long and accommodates an ORF, Orf513, encoding a putative product of 513 amino acids. This was originally misidentified as an ORF of 341 codons and accordingly designated Orf341. The mistake was corrected when it was discovered (71) that the CR sequence is used to interrogate the nucleotide databases, the only sequences that show significant degrees of homology are CRs. The search provides no convincing clues as to CR identity. This is also the case when the amino acid sequence of Orf513 is used to interrogate the protein databases. No convincing identity is revealed, although Cloeckaert et al. noted that Orf513 displays some identity with the transposition protein of IS\(_{801}\) but that the identity is insufficient to be confident of the match in the absence of other data (17). The putative identity of CRs was also suggested by other studies after the publication of that by Cloeckaert et al. (17, 29, 60, 70, 71). Significant identity is seen only between Orf513 and the putative products of other CR ORFs. Given that CRs are frequently associated with antibiotic resistance genes that clearly are mobile, and given their sizes, it is possible that CRs are atypical transposable elements. However, there is a trio of somewhat unusual IS elements, typified by IS\(_{91}\) and including IS\(_{801}\) and IS\(_{1294}\), that employ transposition proteins and lack the DDE motifs of standard transposases (34). Instead, the activities of these proteins are tyrosine based, and they are related to the Rep proteins of *Staphylococcus aureus* plasmids and to the replication protein of coliphage \(\phi\)H9278X174 (64). When the amino acid sequences of the putative products of the CR ORFs are aligned with those of the transposition proteins of IS\(_{91}\) and its relatives, IS\(_{801}\) and IS\(_{1294}\), although overall amino acid identity between the CR-encoded proteins and the transposition proteins of IS\(_{91}\), IS\(_{801}\), and IS\(_{1294}\) is very low, key amino acid motifs of the transposition proteins can be seen to be present in the CR gene products (Fig. 8), indicating they also may indeed be transposition proteins, consistent with the idea that CRs are transposable elements (34, 96).

**FIG. 6. Alignment of the transposase genes of ISCR1 to -5, based on a CLUSTAL alignment with the PAM 250 matrix prepared using Lasergene DNAstar software. Residues found in the majority of the sequences are highlighted, and the consensus sequence is given at the top of the alignment.**
in that they lack the terminal IRs typical of most IS elements (10, 33). Instead, their termini are distinctive and have different functions (34). This is because their transposition mechanism differs from that of most transposable elements in that it involves RC transposition; accordingly, it has been termed RC transposition (34, 65). The single protein encoded by each element is the cognate transposase, which is responsible for initiating replication of the element and is also believed to be involved in terminating replication prior to the final recombination step of the transposition event (34, 96).

With these elements, transposition is a processive system in which one end of the element, oriIS, serves as an origin of replication while the other, terIS, is a replication terminator (34, 96). The oriIS sequence is 5'H11032 GxxTTTxAATTCCTATx3/C AT3/H11032 and is located approximately 300 bp downstream from the transposase gene in IS91 and IS1294 but only 179 bp downstream from the transposase gene in IS801. Sequences in a similar position relative to CR ORFs, i.e., 240 to 250 bp downstream and at the 3'H11032 end of the CR sequence, have the consensus sequence 5'H11032 GCG TTngxT CTTT/CAC GCGC/CACG xxs/C AT3/H11032 (Fig. 7). This sequence is strikingly similar to the oriIS consensus sequence (identity between the nucleotide sequence of the consensus of the 3' end of CRs and that of the oriIS of IS91-like elements is indicated in boldface), to the extent that, given the likely identity of CR-encoded proteins, it is not unreasonable to conclude that this end of a CR sequence is also an origin for RC replication (34, 65, 96). Hence, CR sequences have two key features of IS91-like elements that are similarly located. We conclude, therefore, that these sequences are IS91-like transposable elements. Accordingly, we propose that they be designated ISCR elements, and will so refer to them hereafter, so as to highlight both their identities as insertion sequences and the history of their discovery. Further, the numbers presently used to distinguish CRs are retained to distinguish different ISCRs. Therefore, CR1 becomes ISCR1, CR2 becomes ISCR2, etc.

The third feature of IS91 that is strongly conserved both in IS801 and in IS1294 is terIS (96), located approximately 300 bp upstream from the start of the transposase gene in IS91 and IS1294 but only 101 bp from the start of the transposase gene.

![FIG. 7. Alignment of the 3’-terminal DNA sequences of the various ISCR elements. The stop codons of the various ISCR elements are indicated with asterisks. Sequences found in the majority of the ISCR elements are highlighted and the consensus sequence given above. The 29-bp sequence found at the 3’ ends of the various ISCR elements is underlined.](image)

![FIG. 8. Alignment of motif regions of IS91 group protein sequences with the transposase sequences of ISCR1 to -4 elements. All ISCR elements have the five motifs found within IS91 group elements, and all residues conserved throughout the various IS91 elements are found within ISCR1 to -4 elements. Resides conserved in all IS91 group elements are highlighted.](image)
The IS\textsubscript{91} family element IS\textsubscript{294} has been studied in detail (96), and it has been shown that it can move adjacent sequences when the rolling-circle replication mechanism mis-identifies terIS and proceeds to replicate the DNA adjacent to terIS. This has been calculated to occur in between 1% and 10% of normal transposition events for IS\textsubscript{294} (96). In particular, IS\textsubscript{294} was shown to mobilize the kanamycin resistance gene of pUB2380 to another plasmid, R388, at a frequency of approximately $10^{-3}$ to $10^{-6}$. IS\textsubscript{294} is unusual in that a single intact copy can promote transposition of sequences adjacent to one end of the element, specifically to those linked to the 3\textsuperscript{'} terminus of the element, which has also been shown for IS\textsubscript{91} (65). This is very different from the case for most other insertion sequences, which can only mobilize additional DNA found between two copies of the insertion element (6–8, 38). IS\textsubscript{294} therefore gives us a precedent for a single copy of an insertion element moving adjacent DNA sequences. Thus, we can hypothesize that ISCR elements mobilize chromosomal genes by first transposing into a position adjacent to them, which is followed by a second but aberrant transposition event that mobilizes the adjacent sequence onto a conjugative plasmid and then, via conjugation, into other species/genera of bacteria. We hypothesize that, like IS\textsubscript{294}, ISCR elements present a very powerful mobilization system which can, in theory, mobilize any section of DNA. In this case, they are likely to be more important than the integron system, which can mobilize DNA only in the form of gene cassettes.

FIG. 9. Alignment of DNA sequences found upstream of IS\textsubscript{CR2} elements, depicting the 5\textsuperscript{'} end of the IS\textsubscript{CR2} element. Nucleotide bases found in the majority of sequences are highlighted and consensus sequence given above the alignment. The start codon of the IS\textsubscript{CR2} transposase gene is indicated with asterisks, and the palindromic ter\textsubscript{IS} sequence is underlined. Accession numbers are given at the ends of the sequences.

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FIG. 10. Alignment of the oriIS and terIS of IS\textsubscript{91} family elements and ISCR elements. (Top panel) Alignment of the complement of the last 21 bp of the various ISCR elements with the first 21 bp of IS\textsubscript{91}, IS\textsubscript{294}, and IS\textsubscript{801}, showing oriIS. Nucleic acids found in the majority of sequences are highlighted. (Bottom panel) Alignment of terIS of IS\textsubscript{91} family elements with the sequences found at the equivalent termini of IS\textsubscript{CR2} and IS\textsubscript{CR4}. Inverted repeats are underlined. The direction of rolling-circle replication is indicated with arrows. Accession numbers are given on the right.
**ISCR ELEMENTS AND ASSOCIATED GENES**

ISCR elements are notable for their close association with a wide variety of antibiotic resistance genes.

**ISCR1**

**ISCR1** association with trimethoprim resistance genes. In addition to In7, ISCR1 is associated with four trimethoprim resistance genes in five other complex class 1 integrons (Fig. 1A). All of these were identified in the mid-1990s; three were detected in *Salmonella enterica* serovar Typhimurium strains isolated in Italy, Albania, and Belgium, and the fourth was identified in a strain of *Citrobacter freundii* from Argentina and in a *Klebsiella pneumoniae* strain from Sydney, Australia. The novel trimethoprim resistance gene *dfra23* was found on a nonconjugative plasmid, t-st4, of 110 kb in *S. enterica* serovar Typhimurium from Italy (104); *dfra18* was found on the *incC* group plasmid pDG0100 (93) and on a conjugative *incF1* plasmid of 140 kb in *S. enterica* serovar Typhimurium from Albania (102); and *dfra10* was found in In34 (71) and as a component of two variants (SG1A and SG1D) of the *Salmonella* genomic island SG1 on the chromosome of *Salmonella enterica* serovar Agona (see below) (15). The *dfra3b* gene was found linked to ISCR1 in the *C. freundii* isolate from Argentina (2).

**ISCR1** association with plasmid-mediated quinolone resistance. Plasmid-mediated decreased susceptibility to quinolones, which was subsequently associated with the gene *qnrA*, was first detected in 1994 in a *K. pneumoniae* isolate collected in Birmingham, Alabama (43, 61). However, it has recently emerged in China, Korea, and Europe, suggesting that it is likely to become much more common in the future (47, 50, 60, 68, 107). In all cases tested, the *qnrA* gene was found immediately downstream of ISCR1 as part of a complex class 1 integron that included the 3’ CS duplication (Fig. 1C). The *qnrA* genes isolated in the different geographical locations are virtually identical: i.e., the *qnrA* gene isolated in Europe is the same as the one identified in the United States, and these have only one silent nucleotide difference from those of the Chinese isolates (60). This suggests a common source for this gene irrespective of where it is isolated, which has recently been identified as *Shewanella algeae* (70). The genetic contexts of the *qnrA* gene in several different isolates suggest that it has been mobilized on more than one occasion. The association of ISCR1 with the *qnrA* gene in unrelated isolates is also striking. The gene assemblies are found on plasmids of different sizes and have been seen to move from one plasmid to another during conjugation experiments (107). Several β-lactamase genes have also been found on *qnrA*-carrying plasmids, such as *bla* *PER-3* and *bla* *VEB-3* from Japan (112). Aminoglycosides are among the most commonly used broad-spectrum antibiotics for the treatment of infectious diseases caused by gram-negative bacteria. They cause bacterial cell death by binding irreversibly to the 30S ribosomal subunit (54). The 16S rRNA methylases are particularly powerful resistance mechanisms, because by altering the ribosome they can confer high-level resistance to almost all clinically important aminoglycosides (31). This is in contrast to the more commonly found aminoglycoside resistance genes that encode aminoglycoside-modifying enzymes that individually confer resistance to a much more limited range of drugs (78).

The ISCR1 element was found upstream of *armA* in isolates of *C. freundii* from Poland, *Escherichia coli* and *Serratia marcescens* from Japan, and *K. pneumoniae* from France. The isolates from France (31) and Poland (accession no. AF550415) were shown to harbor the gene array on large plasmids. The *C. freundii* plasmid was sequenced in its entirety, and this revealed the ISCR1 element as part of a complex class 1 integron but without the second copy of the 3’ CS (accession no. AF550415) (35). The *armA* gene was the second gene beyond ISCR1 and distal to the integron and was separated from ISCR1 by a gene with high identity (76%) to a transposase gene from *Providencia rettgeri* (Fig. 1D). Recently, a number of studies have found the *armA* resistance determinant on plasmids in several clinical isolates, including isolates of *C. freundii*, *E. coli*, *K. pneumoniae*, *Enterobacter cloacae*, *S. enterica*, *Shigella flexneri*, and *S. marcescens* collected in France, Bulgaria, and Poland (32) and isolates of *E. coli* and *K. pneumoniae* collected in Taiwan (111), suggesting mobility. It has also been found on an *incN* R-46-like plasmid in *E. coli* isolated from pigs in Spain (35, 36).

**ISCR1 association with class A β-lactamases.** ISCR1 has also been found to be associated with several class A β-lactamase genes, including those belonging to two of the four groups of extended-spectrum *bla* *CTX-M* genes (2, 87), the extended-spectrum class A β-lactamase gene *bla* *VEB-3* from China (48), and the extended-spectrum *bla* *PER-3* gene isolated from *Aeromonas punctata* in France (accession no. AY740681) (Fig. 1E).

**ISCR1** association with *bla* _CTX-M_ genes. CTX-M β-lactamases initially emerged as resistance determinants in Europe and South America at the end of the 1980s and have become the dominant ESBLs (39, 73, 97). For example, in Argentina, the *bla* _CTX-M-2_ enzyme alone now accounts for almost 69% of all ESBLs in *Enterobacteriaceae* that are analyzed in Buenos Aires (30). In both instances, the resistance genes are closely associated with ISCR1 (13, 89).

Genetic analysis of the *bla* _CTX-M_ genes in pathogens in Argentina began with the *bla* _CTX-M-2_ gene in a *Proteus mirabilis* strain isolated in 1993 (1). The *bla* _CTX-M-2_ gene was found on a plasmid (pMAR-12) in an integron, In35, containing the 3’ CS duplication typical of complex class 1 integrons but not as part of a gene cassette. In this array the *bla* _CTX-M-2_ gene is immediately downstream of a copy of ISCR1 (Fig. 1E). Interestingly, this copy of *bla* _CTX-M-2_ is preceded by 266 bp of DNA displaying 96% identity to *bla* _KLU-1_, encoding the class A β-lactamase of *Kluyvera ascorbata*. In addition, 1,043 bp of sequence downstream of *bla* _CTX-M-2_ also displays high identity to *Kluyvera* DNA, a finding that provides evidence that the *bla* _CTX-M-2_ gene almost certainly originates from *K. ascorbata* (42). It therefore seems likely that the associated ISCR1 element was involved in sequestering this section of DNA from the chromosome of *K. ascorbata* or a near relative into a
plasmid carried by the host cell (42). A similar situation was found with an isolate of S. enterica serovar Infantis that was collected in 1997. Here the bla\textsuperscript{CTX-M-2} gene was again found on a plasmid (73 kb) together with 266 bp of \textit{Kluyvera} DNA upstream of the \(\beta\)-lactamase gene and adjacent to a copy of \textit{ISCR1} in a complex class 1 integron, In521, but associated with gene cassettes different from those in In55 (22). In addition, a screen of 130 clinical isolates, which included gram-positive and gram-negative isolates collected between 1993 and 2000 from various hospitals in Buenos Aires, identified \textit{ISCR1} next to \textit{bla\textsuperscript{CTX-M-2}} in all \textit{bla\textsuperscript{CTX-M-2}}-containing isolates, strongly implicating \textit{ISCR1} in the emergence and dissemination of this particular resistance gene in South America (2). The \textit{bla\textsuperscript{CTX-M-2}} gene has been found on plasmids of different sizes and associated with different class 1 integrons. The association of \textit{bla\textsuperscript{CTX-M-2}} and \textit{ISCR1} has been identified in 10 different gram-negative pathogens (including \textit{Acinetobacter} spp., \textit{Enterobacter cloacae}, \textit{E. coli}, \textit{P. mirabilis}, \textit{Pseudomonas aeruginosa}, \textit{Salmonella} spp., and \textit{Serratia marcescens}) (2), in \textit{K. pneumoniae} (62), and in \textit{Morganella morganii} (78). In addition, \textit{bla\textsuperscript{CTX-M-2}} has also been found in isolates of the gram-positive species \textit{Enterococcus faecium} and serogroup \textit{G Streptococcus agalaciae} (2), again in close association with \textit{ISCR1}.

In bacteria isolated in Europe, the \textit{bla\textsuperscript{CTX-M-2}} variant most commonly associated with \textit{ISCR1} elements is \textit{bla\textsuperscript{CTX-M-9}}. The different structural subgroups probably reflect the different origin of each subgroup from source organisms of different \textit{Kluyvera} spp. Thus, \textit{ISCR1} would appear to have been involved in sequestration and mobilizing \(\beta\)-lactamase genes intrinsic to bacterial chromosomes from different species of \textit{Kluyvera}.

The only incidence of \textit{bla\textsuperscript{CTX-M-9}} that has been fully analyzed with respect to its genetic context is from an \textit{E. coli} isolate collected in 1996 in Spain (85). In this instance, the \textit{bla\textsuperscript{CTX-M-9}} allele is linked to sequences displaying significant identity to \textit{Kluyvera} spp. The \textit{bla\textsuperscript{CTX-M-9}} gene and its downstream sequence share 81% and 78% identity, respectively, with the \textit{bla\textsuperscript{KLU-A1}} \(\beta\)-lactamase gene and an adjacent sequence, \textit{orf3}, from the chromosome of \textit{K. ascorbata} (86). Further, the \textit{bla\textsuperscript{CTX-M-9}} allele is next to a copy of \textit{ISCR1}, which together comprise part of a complex class 1 integron, In60 (Fig. 1E). This same arrangement has been found in 30 other Spanish \textit{E. coli} isolates and in 4 \textit{S. enterica} isolates collected between 1996 and 1999. The \textit{bla\textsuperscript{CTX-M-9}} allele has also been found associated with \textit{ISCR1} in bacterial isolates from hospitals in Paris and southwestern France. However, in the former location \textit{bla\textsuperscript{CTX-M-9}} was found to be mostly associated with the insertion element \textit{ISEcp1} rather than \textit{ISCR1}, although evidence of \textit{ISCR1} activity in the form of short \textit{ISCR1} terminal sequences was identified upstream of \textit{bla\textsuperscript{CTX-M-9}} in an \textit{E. coli} isolate, and similar sequences were found in two \textit{P. mirabilis} isolates harboring either \textit{bla\textsuperscript{CTX-M-9}} or \textit{bla\textsuperscript{CTX-M-20}} (88). Further analysis should elucidate whether these short sequences reflect intact copies of \textit{ISCR1} or simply fragments. The \textit{bla\textsuperscript{CTX-M-9}}-harboring isolates from southwestern France are of particular interest because they were isolated from both human and poultry sources, suggesting an animal reservoir of \textit{ISCR1} and \textit{bla\textsuperscript{CTX-M-9}} alleles. The \textit{bla\textsuperscript{CTX-M-9}} alleles were again found adjacent to copies of \textit{ISCR1} in nine isolates of \textit{S. enterica} originating from a single hatchery supplying chicks to six farms (108). The integron structure in these isolates was 99% identical to that of In60, found in Spain, and in all isolates was on a large plasmid (>126 kb). Since this integron was first reported in Spain, it has also been found in France, Turkey, Brazil, and China (108).

\textbf{ISCR1 association with other class A \(\beta\)-lactamase genes.} Other class A \(\beta\)-lactamase genes that are associated with a copy of \textit{ISCR1} are \textit{bla\textsuperscript{VEB-3}} (48) and \textit{bla\textsuperscript{PER-3}} (accession no. AY740681). The \textit{bla\textsuperscript{VEB-3}} gene has been found in numerous \textit{E. cloacae} isolates in Shanghai, China, and has features that suggest that it is part of a gene cassette. However, it could not be linked to the class 1 integrons that were found in these strains; rather, it was shown to be linked to the 3’ end of a fragment of \textit{ISCR1} found upstream of the \textit{bla\textsuperscript{VEB-3}} gene. The sequence following this \textit{ISCR1} fragment displays high identity to \textit{IS6100}, and therefore it seems likely that \textit{bla\textsuperscript{VEB-3}} was initially mobilized by \textit{ISCR1}, which was then fragmented by an \textit{IS6100} insertion. The \textit{bla\textsuperscript{PER-3}} gene was identified in an isolate of \textit{Aeromonas punctata} in France adjacent to a copy of \textit{ISCR1}, as part of a complex class 1 integron. Again, the gene \textit{bla\textsuperscript{PER-3}} does not appear to have the associated 59-base element that is characteristic of a gene cassette.

\textbf{ISCR1 association with class C \(\beta\)-lactamase genes.} Class C \(\beta\)-lactamases, unlike the extended-spectrum class A \(\beta\)-lactamases, mediate resistance to cephamycins as well as broad-spectrum cephalosporins (39, 97). However, unlike those of class A ESBLs, the activities of class C \(\beta\)-lactamases are not generally inhibited by clavulanic acid or tazobactam. Genes encoding class C \(\beta\)-lactamases are found on the chromosomes of many enteric bacteria and also are intrinsic to some nonfermenting gram-negative bacteria such as \textit{P. aeruginosa} and \textit{Acinetobacter} spp., where they may be silent and under tight transcriptional control (58, 91). However, recently resistance to cephamycins and cephalosporins has been acquired by \textit{Klebsiella} spp. and \textit{E. coli}, mediated by plasmids carrying genes encoding enzymes closely related to chromosomal AmpC \(\beta\)-lactamases. It appears likely that \textit{ISCR1} has also played a major role in the mobilization of these genes into human bacterial pathogens via plasmids and other mobile genetic elements (T. R. Walsh, unpublished data).

\textit{ISCR1} elements have been found adjacent to a number of cephalosporinase genes, including \textit{bla\textsuperscript{DHA-1}} (originating from the chromosome of \textit{M. morganii}) (103), \textit{bla\textsuperscript{CMY-3}} and \textit{bla\textsuperscript{CMY-8}} to \textit{bla\textsuperscript{CMY-11}} (3, 23, 55, 110) (thought to originate from an \textit{Aeromonas} sp.), and a variant of \textit{bla\textsuperscript{CMY-type}} genes, \textit{bla\textsuperscript{MOX-1}} (41) (Fig. 1F). Of these class C \(\beta\)-lactamase genes, the genetic loci of only the \textit{bla\textsuperscript{CMY-9}} and \textit{bla\textsuperscript{DHA-1}} genes have been sequenced and analyzed in any detail. In the case investigated, \textit{bla\textsuperscript{CMY-9}} was plasmid encoded in an \textit{E. coli} isolate, HKYM68, collected in Japan in 1995 and displaying high-level resistance to ceftazidime, cefpirome, and moxalactam (23). Cloning and sequencing of 8 kb around the \textit{bla\textsuperscript{CMY-9}} gene (Fig. 1F) revealed that the gene is adjacent to a copy of \textit{ISCR1} and downstream of a class 1 integron but that there is no second copy of the 3’ CS, as is usual when \textit{ISCR1} is linked to a class 1 integron. The \textit{bla\textsuperscript{CMY-9}} allele shows 78% identity to \textit{cepH} of \textit{Aeromonas hydrophila}. Comparison of the nucleotide sequence upstream of \textit{bla\textsuperscript{CMY-9}} and those of the other related cephalosporinase genes \textit{bla\textsuperscript{CMY-10}} and \textit{bla\textsuperscript{CMY-11}} with the corresponding region of the \textit{A. hydrophila cepH} locus reveals 64%
identity, again suggesting that the origins of these particular β-lactamase genes are probably A. hydrophila or closely related strains (23).

It is interesting that all ISCR1-associated bla<sub>CMY</sub>-like structure and were probably acquired recently (4). Antibiotics actually promotes mobilization (5). The resistance (4) (Fig. 2). It seems likely, therefore, that ISCR1 (CR1) found beside the trimethoprim resistance gene dhfR18 (Fig. 2) of the element, leading to deletions, have given rise to SX1 variants that differ with respect to their resistance gene complements (4; compare Fig. 2, accession no. AY055428 and AB114188).

**ISCR2**

ISCR2 (encompassing the gene known as orfA) is the primary representation of the second subgroup of ISCR elements. It is part of the MDR-encoding genetic element SXT, as well as being found on numerous plasmids carrying chloramphenicol/flofenicol and sulfonamide resistance genes (Fig. 2). The ISCR2 transposase shares 65% amino acid identity with its ISCR1 equivalent but differs dramatically from ISCR1 in its genetic contexts. While ISCR1 has hitherto been found as a component of complex class 1 integrons, ISCR2 has never been found associated with class 1 integrons but is often associated with the sul2 gene (compare Fig. 1A to Fig. 2).

**Involvement of ISCR2 with SXT.** Before 1992 the major cause of cholera on the Indian subcontinent was Vibrio cholerae El Tor. However, in 1992, V. cholerae El Tor was replaced by V. cholerae serogroup 0139 as the predominant cause of cholera (40). This strain was also different from V. cholerae El Tor in being resistant to several antibiotics, i.e., sulfamethoxazole, trimethoprim, chloramphenicol, flofenicol, and streptomycin. The resistance genes were located on a novel genetic element of the integrative conjugative element group named SXT. This element is now present in virtually all clinical V. cholerae serovars (40). SXT is a conjugative, self-transmissible element that integrates into the prfC gene on the chromosome of V. cholerae and has also been found on the chromosome of Providencia alcalifaciens (4). Integration is necessary for replication of the element, and SXT can be transferred into several other gram-negative organisms, including E. coli, by conjugation. Of interest and concern is the fact that challenge with other gr-

**ISCR2 association with other resistance mechanisms.** The ISCR2 element has also been found next to resistance mechanisms in other gram-negative organisms. ISCR2 is present on several plasmids, often in truncated form, including plasmids which have been studied for several decades. As previously stated, it is closely associated with the sulfonamide resistance gene sul2 in the conjugative plasmid RSF1010 (81). ISCR2 has also been found associated with a flofenicol resistance gene found on plasmids in E. coli isolated from cattle, poultry, and pigs (12). This has spread to the major target bacterium for cattle, Pasteurella multocida (52). Flofenicol was licensed in Europe for the control of bacterial respiratory tract infections in cattle and pigs in 1995 and 2000, respectively, and is active against chloramphenicol-resistant isolates with chloramphenicol acetyltransferases and efflux systems (12). The first flofenicol resistance gene was detected in 1996 on a plasmid in the fish pathogen Pasteurella damselae subsp. piscida (formerly known as Pasteurella piscida) (46) and has recently been found in the Salmonella genomic island SGI1 in S. enterica serovar Typhimurium DT104 (14) and in the SXT element of V. cholerae (18) (Fig. 2). The close association between ISCR2 and sul2 prompted us to study Stenotrophomonas maltophilia isolates that were resistant to trimethoprim-sulfamethoxazole (TMP-SMX). Out of 25 TMP-SMX-resistant isolates collected from various different geographic regions, the ISCR2 element was found in 6 isolates (M. A. Toleman, unpublished data). In each case, the ISCR2 element was found adjacent to a truncated phosphoglucomutase gene, ΔglmM, and the sul2 gene. This arrangement is identical to that found in the previously mentioned E. coli strains isolated from cattle in France and Germany, together with on plasmid pRVS1 isolated from V. salmonicida in Norway, on a plasmid from an S. enterica strain isolated in Japan, and on the chromosome of an S. flexneri strain isolated in the United States. The arrangement of ΔglmM and sul2 has also been found in plasmids isolated from marine psychrophilic bacteria from Norway (accession numbers AJ306553 and AJ306554), but in these cases the ISCR2 element was absent. Two of the six S. maltophilia isolates also carried a copy of a floR gene immediately upstream of ISCR2 and showed the arrangement incorporating ISCR2 seen in the E. coli ISCR2-carrying strains from cattle in Germany and France. The two S. maltophilia isolates came from Turkey and the United States, indicating widespread dissemination of the floR resistance gene on plasmids in association with ISCR2 (Toleman, unpublished data).

It has been suggested recently that a section of DNA encompassing the floR gene and an intact copy of ISCR2 is part of a novel transposon, TnfloR (28), that transposes via a circular intermediate by a mechanism similar to that proposed for the staphylococcal transposon Tn554 (28). The authors of that paper correctly recognize that the presence of the floR gene on the chromosome or on different plasmids in a number of unrelated Salmonella enterica and E. coli isolates is characteristic of movement by transposition. The evidence provided to substantiate the claim that floR is part of a transposon is production of a PCR product that contains both ends of the proposed element, using primers that direct replication out of the linear form of the sequence, indicating a circular form of the sequence in the cell. This is interpreted to indicate formation of a circular intermediate in the transposition process. However, no account has been taken of the fact that the floR gene in this particular strain is flanked by a complete copy of ISCR2 on one side and a truncated version on the other. Accordingly, homol-
ogous recombination between the repeated sequence would generate a circular structure of the form reported. Given that the strain used in the experiment was a wild-type *E. coli* strain and not a *recA* mutant deficient for homologous recombination, generation of free “ISCR2-*floR*” circles is to be expected. Strikingly, the circular “intermediate” form of the proposed transposon was detected in only one of six florfenicol-resistant *E. coli* strains and, specifically, in the only one in which the *floR* gene is flanked by a sequence duplication. Furthermore, transposition of the novel “transposon” was not actually demonstrated. Accordingly, definitive proof of the existence of *Tn*21 was not presented. We would argue that the *floR* gene has indeed been transposed on several occasions, mediated by ISCR2 via RC transposition, but that the combination ISCR2-*floR* does not constitute a transposable element.

**ISCR3**

The third subgroup of ISCR elements is typified by ISCR3. This subgroup was again originally identified by the transposable gene, namely, *orf2*. This version encodes a protein that shares approximately 55% and 57% identity with the transposases of ISCR1 and ISCR2, respectively. To date it has been found associated with the SG1I element (and variants thereof) (Fig. 3) and is also linked to an *erm* gene encoding erythromycin resistance and the aminoglycoside resistance gene *rmtB* (Fig. 3). We have also recently identified the ISCR3 in TMP-SMX-resistant *S. maltophilia* strains isolated in Spain, but the genetic locus has not yet been determined (Toleman et al., unpublished data).

**Involvement of ISCR3 and ISCR1 with the Salmonella genomic island 1 genetic element.** SG1I is a genetic element of approximately 43 kb (15). It has been associated mainly with MDR isolates of *S. enterica* serovar Typhimurium phage type DT104 that are resistant to ampicillin, chloramphenicol, streptomycin, sulfonamides, and tetracycline. This pathogen emerged in the last decade as a global animal and human health problem (15). Outbreaks of MDR *S. enterica* serovar Typhimurium DT104 have occurred in poultry, beef, and pigs and their food products, as well as in dairy products and salad ingredients. MDR *Salmonella* are very common in the United Kingdom and increasingly prevalent in many other countries (15). Since its initial discovery in *S. enterica* serovar Typhimurium DT104, SG1I has also been found in other *S. enterica* serovar Typhimurium phage types, i.e., DT120, DT12, DT1, and U302, and in other serovars such as Agona, Paratyphi B, Albany, Meleagridis, Newport, Emek, Cerro, Derby, Dusseldorf, Infantis, and Kiambo (15, 26, 56). These observations are indicative of SG1I transfer, and transduction of SG1I between *S. enterica* serovar Typhimurium DT104 isolates has been demonstrated (11). However, attempted transfer to other *S. enterica* serovars in vitro was not successful, and SG1I excision has not been detected experimentally (14). It has been suggested, therefore, that the presence of this element in different serovars of *S. enterica* and in different phage types of *S. enterica* serovar Typhimurium is due to separate emergence events rather than horizontal movement between *S. enterica* strains, but the source of the genomic island is as yet unknown. SG1I is found at the same place on the chromosomes of all of these organisms, inserted between *thdF* and *yidY* (26). DT104 isolates also have the addition of another element, described as a cryptic retrophage inserted between *thdF* and the left end of SG1I (14). Whether this element is involved in SG1I movement is not known.

SG1I contains a resistance module of approximately 14 kb at one end. This consists of *floR* and *tetR* bracketed by two class 1 integrons, one carrying the gene cassette *aadA2* and the other carrying the *bla*<sub>pse-1</sub> cassette (Fig. 3). Recently, seven variants of SG1I in isolates originating from Belgium and Scotland (SG1IA to -G) (15) to Australia (SGIA I and J) (56) have been described. These variants differ mostly in the resistance genes that they carry and in the presence or absence of CR elements (15, 56).

**ISCR3** is the ISCR variant most often found in SG1I. It is present in SG1I variants SG1IA, SG1IE, SG1I-I, and SG1I-J, as well as in SG1I (Fig. 3). However, recently ISCR1 has also been detected in SG1I variants in isolates originating from poultry in Belgium (27), specifically, in variants SG1IA, SG1ID, and SG1IG. Hence, SG1IA harbors both ISCR1 and ISCR3, whereas variants SG1IB and SG1IC lack an ISCR element (Fig. 3). In all variants containing ISCR1, the element is found immediately upstream of the trimethoprim resistance gene *dfrA10*, in a complex class 1 integron and downstream of a class 1 integrase/*groEL* gene fusion (Fig. 3); i.e., it is found with *dfrA10* between repeat copies of the 3’ CS of class 1 integrons. The ISCR1 element has also been found in deletion versions of SG1I where it is not associated with any antibiotic resistance gene (27). In these cases, the entire SG1I sequence normally downstream of ISCR1 has been deleted, bringing the chromosomal gene *yieE* or *yieF* into juxtaposition with ISCR1 (Fig. 3, accession numbers AY434091 to -3). Such a deletion has occurred on at least three occasions, involving deletion of sequences of 7.4 kb, 7.7 kb, and 8.5 kb. Such deletions are characteristic of recombination events between direct copies of insertion elements and are likely to have been ISCR1 mediated (27).

In SG1I ISCR3 is located between the two class 1 integrons found on the genomic island in association with *floR* and *tetA/R* (Fig. 3). An exception to this is found in variant SG1IE, which has a copy of ISCR3 located at one end of the genomic island, downstream from the second class 1 integron (Fig. 3). This may reflect an inversion event involving the insertion element *IS6100*, copies of which are found at the ends of the inverted section of DNA. It is therefore very likely that both ISCR1 and ISCR3 elements have been involved individually and/or corporately in the construction of SG1I and its variant types, as well as being instrumental in incorporating further resistance genes.

ISCR3 has also been found adjacent to the 16S rRNA methylase gene *rmtB* in a clinical strain of *S. marcescens* (23) (Fig. 3). This methylase gene confers high-level resistance to various aminoglycosides, including kanamycin, tobramycin, amikacin, arbekacin, gentamicin, sisomicin, and isepamicin, but not neomycin, streptomycin, and hygromycin B. The *rmtB* gene is likely to have been mobilized to *S. marcescens* from an environmental microorganism by ISCR3. The deduced amino acid sequence of RmtB is 29% identical to that of the ArmA 16S rRNA methylase found associated with ISCR1 (Fig. 1D). Therefore, similar broad-spectrum aminoglycoside genes have been found adjacent to ISCR1 and ISCR3.

In all cases investigated so far, the left end of the ISCR3
element is flanked by the groEL gene, displaying 78% identity with Xanthomonas campestris. This same association is seen in both the SG11 of S. enterica and the mntB locus of S. marcescens. Interestingly, the same groEL gene is found upstream of the blaSPM-1 gene found in close association with ISCR4.

ISCR4

The fourth subgroup of ISCR elements has recently been identified, associated with the MBL gene blaSPM-1, which confers resistance to all β-lactam antibiotics except from aztreonam (67, 77, 98) (Fig. 4). P. aeruginosa strains harboring this resistance mechanism were found to be resistant to every available antibiotic, with the sole exception of polymyxin B. The blaSPM-1 MBL gene has been found in nonfermenting gram-negative bacteria collected from several institutions throughout Brazil, and its product is a major contributor to the high level of resistance to carbapenems seen in South America, which are currently among the highest in the world (51). The blaSPM-1 gene was found on a large plasmid in various strains of P. aeruginosa with distinct ribogroups, implying horizontal dissemination (51). blaSPM-1 is not part of a gene cassette, nor is it found in the vicinity of a class 1 integron (98) as found for other MBL genes, but it is located beside the ISCR variant ISCR4. Again, this element was first detected by its transposase gene, previously named orf495 (77). The transposase encoded by ISCR4 shows 79%, 57%, and 53% amino acid identity with those of ISCR3, ISCR2, and ISCR1, respectively. A partial repeat of ISCR4 has been found downstream of blaSPM-1, similar to the cases of several of the ISCR elements (Fig. 4). Upstream of ISCR4 and downstream of blaSPM-1, sequences displaying 94% nucleotide identity to each other were found. These were identified as groEL. The sequence upstream of ISCR4 encodes a predicted amino acid sequence with 89% and 87% identity with GroEL proteins from Desulfitobacterium hafniense and S. maltophilia, respectively (77). The sequence downstream of blaSPM-1 encodes a protein with 73% identity to the GroEL protein of Xanthomonas campestris (98) (Fig. 4). The second copy of ISCR4 was truncated by the cloning process.

ISCR5

To date ISCR5 is uniquely associated with the class D oxacillinase gene blaOXA-45 (99). This resistance gene was cloned from P. aeruginosa strain 07-406, which was collected via the SENTRY Antimicrobial Surveillance Program from the Anderson Medical Center in Texas (99). The sequence immediately downstream of blaOXA-45 shows 88% nucleotide sequence identity with ISCR3 (Fig. 4). It has been found both on a small plasmid (99) and on the chromosome of strain 07-406 (Toleman, unpublished data), indicating that ISCR5 is mobile. The ISCR5 ORF displays 91%, 78%, 57%, and 52% identity (entire sequence) with the transposase genes of ISCR3, ISCR4, ISCR2, and ISCR1, respectively (Fig. 5).

ISCR6

ISCR6 was found adjacent to an aminoglycoside-modifying gene, ant(4′)-Hb, that confers resistance to amikacin but not netilmicin (87). This arrangement was detected in six epidemiologically unrelated clinical strains of P. aeruginosa that were isolated between 1992 and 1998 in Bulgaria. Unfortunately, only the 3′ end of the ISCR6 transposase gene was cloned on the fragment with the aminoglycoside resistance gene (Fig. 4). The recovered sequence encodes the last 247 amino acids of the ISCR6 transposase and the orfIS of the element. The ISCR6 transposase sequence is 89% and 88% identical to analogous sections of the transposases of ISCR3 and ISCR5, respectively. The mobility of the aminoglycoside resistance gene was determined by Southern blotting. Analysis of the six clinical P. aeruginosa strains with an ant(4′)-Hb probe detected the gene on the chromosomes of four of the strains and on a large (>300-kb) plasmid in the other two (87).

ISCR7 and ISCR8

The sequences that identify ISCR7 and ISCR8 appear to be incomplete. Both were found in Pseudomonas spp., close to genes encoding degradative enzymes involved with catabolism of haloalkanes and monocyclic nitroaromatic compounds, respectively (20, 74). The truncated elements were identified by their predicted products, which display approximately 30% amino acid identity with other ISCR proteins. That of ISCR8 contains all the key peptide motifs identified in the IS91 transposase, while the predicted product from ISCR7 lacks motif 1. Both putative transposase genes appear to have suffered a deletion and sustained double insertions, in comparison to other ISCR genes, suggesting that both putative transposases are inactive. In keeping with the conclusion that these sequences indicate degenerate copies of ISCR elements, neither ORF has an accompanying version of orfIS or a recognizable terIS.

Other ISCR Elements

In addition to the reasonably well characterized elements described above, several other ISCRs are currently being characterized. ISCR9 and ISCR10 were discovered in strains of S. maltophilia by using a PCR strategy employing degenerate primers designed against the aligned DNA sequences of ISCR1 to -5 (Toleman, unpublished data). The amplified sections encoded amino acid sequences 95% identical to each other and possessing identities of 30%, 48%, and 74% to the same sections of the transposases of ISCR2, ISCR3, and ISCR5, respectively.

Using the same generic method, we have also detected ISCR elements in several strains of P. aeruginosa and Acinetobacter baumannii that harbor MBL genes. ISCR2 was discovered in a P. aeruginosa isolate isolated in Brazil that also harbored the MBL gene blaIMP-1, while ISCR3 was discovered in two strains of P. aeruginosa isolated in Italy that harbored the MBL blaVIM-1: ISCR11 was discovered in two A. baumannii isolates from Germany that have the MBL blaVIM-2 and in a P. aeruginosa isolate from Greece that produces the MBL VIM-1. This new ISCR family member encodes a putative transposase that shows highest identity (79%) to the ISCR3 transposase. Finally, a strain of P. aeruginosa that produces the MBL SPM-1 carries two ISCR elements, ISCR4 and a new element, ISCR12, that encodes a putative transposase with 89% identity to the
ISCR3 transposase and 75% identity to the ISCR4 transposase (100). We have also recently detected >50 ISCR5-like copies in the A. hydrophila genome (TIGR). The sequences in the A. hydrophila genome display approximately 54% nucleotide identity to ISCR5 and probably represent at least one further family member. ISCR13, an ISCR-like sequence in this array, is found adjacent to a gene with high identity to an aminoglycoside resistance gene (unpublished results).

**ISCRs MOBILIZE AND CONSTRUCT COMPLEX CLASS 1 INTEGRONS**

ISCR1 appears to be unusual in the sense that it is almost always found in a complex class 1 integron structure. The sequence upstream of ISCR1 is always the same: i.e., the junction between ISCR1 and the sequence adjacent to the 3' CS of the class 1 integrons is invariant. This suggests that a single event has been responsible for this feature of all complex class 1 integrons. Given that ISCR1 is significantly shorter than other ISCR and IS91-like elements, it is also likely that the terIS end of the element has been deleted following transposition (Fig. 11). This would fuse the remainder of the ISCR1 element (the transposase gene, orf513, and oriIS) to the 3' CS end of the class 1 integron and provide the integron with the basis for genetic mobilization, since the oriIS and transposase gene remain intact. This ISCR1-class 1 integron fusion would also require that in the event of transposition being initiated, an alternative termination site would have to be utilized due to the fact that the original termination site, terIS-1, has now been deleted (Fig. 11). One consequence of the generation of ISCR1 by deletion of its terIS is that it would be expected to cotranspose the adjacent sequence, i.e., class 1 integron sequences, more often. This, then, allows the construction of a model in which ISCR1 has mediated the assembly of a variety of complex class 1 integrons (Fig. 12).

At some time in the past, possibly in the first half of the 20th century, the progenitor of ISCR1 transposed into a site close to the 3' CS of a class 1 integron (Fig. 12). This transposition was followed by a deletion that removed the terIS terminus of the
element and some of the 3’ CS normally associated with class 1 integrons, i.e., orf5 and orf6 (Fig. 12). This variant, ISCR1, then mediated a series of secondary transposition events that translocated ISCR1 and various sections of the adjacent class 1 integron into sites next to other, noncassette resistance genes, such as catA, dfrA, qnr, and various CMY genes (Fig. 12). Once in these locations, the ability of IS91-like elements to generate free circular forms was manifest. Circular entities carrying ISCR1 and sequences adjacent to it and distal to orfIS, including at least the truncated 3’ CS and the linked resistance gene, were generated. These were then, in turn, rescued by homologous recombination into either the 3’ CS of conventional class 1 integrons or that of an integron-ISCR1 variant. The difference in the arrangements that would arise from these recombination events is that in the former case the array would carry only a single copy of ISCR1 adjacent to the class 1 integron between direct repeats of the 3’ CS of the class 1 integron, while in the latter case there would be a direct duplication of the ISCR1 element following the duplication of the 3’ CS, a particular arrangement that has been reported by Mammeri et al. (60). In that study, two unusual sequences were reported, where two copies of “Orf513” flanking a qnrA/qacDE/sul region were present on different plasmids (pMG252 and pQR1) from the same E. coli strain (pMG252 and pQR1) (60). It is therefore possible that two copies of ISCR1 have been transposed independently onto these plasmids via an RC mechanism. Furthermore, given that the genetic contexts of ISCR1 on pMG252 and pQR1 are virtually identical, it is possible that ISCR1 has mobilized itself and adjacent DNA from pMG252 to pQR1 or vice versa. According to our model, the terminal full-length copy of the 3’ CS on a complex class 1 integron that lacks an ISCR1 duplication represents that originally linked to the int gene cassette array depicted on the left of the complex integron in Fig. 12. The internal, shortened version of the 3’ CS, together with the adjacent copy of ISCR1 and any other gene(s), has been acquired by rescue, via ho-
mologous recombination, of an ISCR1-generated circularized DNA fragment. The reverse event, i.e., deletion of sequences flanked by extensive direct repeats, would also be expected on occasion, and indeed it has been demonstrated experimentally that this can happen (71).

It is possible that ISCR3 and ISCR4 mobilize adjacent DNA sequences which are subsequently rescued by homologous recombination via flanking groEL sequences rather than the 3′ CS of class 1 integrons as seen for ISCR1. ISCR4 also appears to have lost its original terIS sequence, possibly by a deletion event linking groEL to the 5′ end of ISCR4. This has caused the groEL termination sequence to be in a position similar to where the original terIS should be, and this may function as an alternative terIS (Fig. 10). A similar event also appears to have happened with ISCR3, and groEL is now found immediately upstream of ISCR3 in SG1 and variants as well as upstream of ISCR3 associated with the aminoglycoside resistance mmtB gene (Fig. 3).

SIMILARITIES AND DIFFERENCES BETWEEN ISCR AND IS91-LIKE TRANSPOSASES

An alignment of the predicted amino acid sequences of the transposase proteins of ISCR1 to -5 (Fig. 6) reveals that they differ in amino acid identity from 53% to 95% (Fig. 5). However, within the principal motifs, the alignment reveals key conserved residues that have been implicated as being essential for transposition of IS91 and like elements. The most notable exception to the match is Y249, which in ISCR1, ISCR2, and ISCR4 is replaced by arginine and in ISCR3 is replaced by lysine. In the case of IS91, it has been proposed that RC transposition involves two conserved tyrosine residues (Y249 and Y253), one of which serves as a nucleophile to attack the oriIS sequence of the IS element while the other attacks the target site, both effecting strand cleavage and the formation of a covalent bond between the amino acid residue and the 5′ phosphate group released by DNA strand cleavage (34, 34).

The nucelophilic centers provided by the hydroxyl groups of the tyrosine residues are clearly critical to the proposed reactions. When Y249 or/and Y253 in the IS91 transposase were mutated to phenylalanine or serine, transposition activity was no longer detected, consistent with the view that both residues are essential to the transposase activity (33). However, it is yet to be proved that both tyrosine residues do participate directly in DNA strand cleavage, as proposed by Garcillan-Barcia et al. (33, 34), in that altering the Y249 amino acid residue could simply adversely affect the overall active-site architecture. The ISCR substitution of Y249 brings into question the proposed role of Y249 in the IS91 transposase; however, it is possible that ISCRs have found alternative residues to facilitate RC transposition. Interestingly, the Rep proteins encoded by plasmid pT181 possess the equivalent of Y253 but not Y249 and act as dimers, with the second molecule nicking the displaced old-to-new leading strand junction to initiate termination of replication (32, 49). Other Rep proteins, such as those mediated by plasmid pC194, require a glutamate residue as well as tyrosine for enzyme activity; the former is thought to catalyze the release of the protein from the intermediate at the termination step (69). Recently, the crystal structure of a TnpA, an RC replication protein, has been solved (84). TnpA also appears to possess the single tyrosine and, as demonstrated by the cocrystal structure with DNA stem-loops, acts as a dimer.

Clearly, more research into the active sites of IS91-like molecules is needed. The other notable exception in terms of conserved amino acid residues is evident from the alignment of ISCR transposases with those of IS91 and IS1294 (CR1) is H149Q; where glutamine has replaced histidine in all the ISCR-encoded proteins (Fig. 6). This may be a change enforced by the Y249K/L substitution needed to retain transposase activity.

SPECIFICITY OF INSERTION OF IS91 AND ISCR ELEMENTS

IS91 and the related family members IS1294 and IS801 target and insert at a specific tetranucleotide sequence (GTTC or minor variants thereof), and similar sequences are also recognized at terIS. These elements insert into the target DNA 5′ to this tetranucleotide so that the target site is then found immediately adjacent to the oriIS sequence of the element and is necessary for further transposition events (63). This sequence is also precisely the cleavage site for the PC194 plasmid family and for ϕX174-like phages (34).

Interestingly, in all examples of ISCR1 and ISCR2 available in the sequence databases, there appears to be no obvious common tetranucleotide found adjacent to the oriIS. It is therefore possible that the ISCR transposase no longer relies on the junction between target site and oriIS for RC transposition origin or that the majority of transposition events are dead ends in the sense that retransposition is prevented. In support of the former suggestion, the tetranucleotide sequences that are thought to be involved in target site recognition in IS91 are the same amino acids that are changed in ISCR elements (63). Perhaps, therefore, ISCR elements are no longer target site dependent. However, nonfunctional insertions may be rescued if a second copy of the element with the correct oriIS/tetranucleotide junction is present in the cell, as has been shown for IS91 (63).

ISCR ELEMENTS AND PROMOTER SEQUENCES

Insertion elements are known, on occasion, to provide promoters for expression of otherwise silent genes by transposing into a site immediately upstream of the promoterless gene: e.g., the cfiA MBL gene of Bacteroides fragilis is normally silent, but ceftazidime-resistant isolates are occasionally found (106). When these are genetically analyzed, they are found to have an insertion element upstream of the cfiA gene that provides a promoter sequence for its expression and, as a result, acquisition of clinical resistance. Promoter sequences have been found at the 5′ end of the ISCR1 element (60, 107). A hybrid promoter was also suggested to have been formed between a ~10 sequence that is found naturally upstream of bla<sub>SPM-1</sub> and a ~35 sequence found within the right-hand boundary of ISCR4. The spacing between these sequences is 17 bp. This combination of mobility and expression via ISCR elements is loosely comparable to the integron structure, where promoterless genes are expressed by virtue of proximity to promoters in the 5′ end of the integrase gene. However, the ISCR elements do not have to provide a promoter for the genes they associate.
with because many of these resistance genes have their own promoters.

**ORIGINS OF CR ELEMENTS**

A number of observations suggest that the ISCR1 element may have originated from a waterborne bacterium. Both SXT and the SG1I elements are found in *V. cholerae* and *S. enterica* serovar Typhimurium, which often inhabit aquatic environments. ISCR1 is found on plasmids closely associated with the waterborne organism *A. salmonicida* (82). Moreover, the ciprofloxacin resistance gene *qnrA* originates from a waterborne organism, *S. algae*, and ISCR1 has been found adjacent to several members of the *bla*CMY-1 group genes, which are suggested to have come from an *Aeromonas* sp. However, they have not been found associated with the *bla*CMY-2 group members, which are thought to have originated from enteric organisms such as *C. freundii*, *blaPER, 3* is closely associated with ISCR1 and was isolated from *A. punctata* in France (accession no. AY740681). GenBank screens of incomplete microbial genome sequence databases with ISCR elements have also revealed many positive results (>50%) in the *A. hydrophila* genome sequence that are ~54% identical to ISCR5. This organism may be the source of another ISCR family member, provisionally named ISCR13. It is also noteworthy that the first florfenicol resistance gene, *flor*, was detected on a plasmid from the fish pathogen *P. damselae* subsp. *piscicida* (formerly known as *P. piscicida*), which may also suggest a marine environmental source of ISCR2.

The various ISCR elements vary in their G+C contents from 54% for ISCR1 to 69% G+C for ISCR5 (Table 1), which suggests that the different ISCR family members have originated from different source organisms. ISCR1 and ISCR2 have the lowest percent G+C of all ISCR family members isolated so far, with 54% and 59.5% G+C, respectively. The rest of the ISCR family members have G+C contents of above 60%. ISCR3 and ISCR4 are both found adjacent to partial *groEL* genes in their hosts *S. enterica* serovar Typhimurium/S. marc-eceur and *P. aeruginosa*, respectively. These partial *groEL* genes both display highest identity with the *groEL* genes of *Xanthomonas campestris* and *S. maltophilia*. Both of these organisms are found in the soil and have high G+C content, which is consistent with the hypothesis that the *groEL* genes and ISCR3/ISCR4 were acquired at the same time, possibly from a soil/plant rhizosphere-associated organism. The significant differences in G+C content shown for the various ISCR elements may indicate that their sequences have been diverging for some considerable time.

**IS91 AND VIRULENCE FACTORS**

IS91, the prototype element of this expanding family of IS91-like insertion elements, was first discovered in plasmids encoding the α-hemolytic function in *E. coli* in 1982 (113) and was subsequently identified in all plasmids and chromosomal pathogenicity islands harboring the α-hemolysin operon (Hly) (34, 53), which codes for synthesis, activation, and export of α-hemolysin. The Hly plasmids belonged to different incompatibility groups, which suggested that IS91 was involved in the dissemination of these pathogenicity determinants (21, 37, 53, 114). IS91 or very closely related isoforms have also been found adjacent to various other virulence genes in enteropathogenic, enterohemolytic, and enterotoxigenic strains of *E. coli* (16, 42, 109), including the *eltAB* toxin-encoding genes of the heat-labile enterotoxin (89). Furthermore, direct movement of *eltAB* genes by IS91 has been demonstrated (89). Similarly, IS1294 has been associated with virulence genes in *E. coli* (96), and IS801 has been discovered in close association with virulence genes in the plant pathogen *Pseudomonas syringae* (66, 83, 95).

**CONCLUSIONS**

Not less than 10 years ago there were numerous examples of IS91 found adjacent to virulence genes but not found adjacent to antibiotic resistance genes. This phenomenon has prompted Garcia-Baeria et al. (34) to speculate that IS91-like elements are best suited to movement of virulence genes and that transposons and integrons are more suited to the dissemination of antibiotic resistance genes. However, there is no reason why these interesting elements cannot mobilize any piece of adjacent DNA. Therefore, it is not surprising that this powerful gene mobilization mechanism has connected with such easily selectable markers as antibiotic resistance genes. In fact, the presence of ISCR1 as part of the complex class 1 integrons In6 and In7 on the plasmids pS6 and pDG1000 indicates that they were already associated with chloramphenicol and trimethoprim resistance genes in the late 1950s and early 1960s, shortly after these antibiotics were first used. Clinically, the most worrying aspect of ISCR elements is that they are increasingly being linked with more potent examples of resistance, i.e., MBL in *P. aeruginosa* and co-trimoxazole resistance in *S. maltophilia*. Furthermore, if ISCR elements do move via “unchecked RC transposition,” as has been speculated for ISCR1, then this mechanism provides antibiotic resistance genes with a highly mobile genetic vehicle that could eclipse the effects of previously reported mobile genetic mechanisms.

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