Genetic basis of the association of resistance genes \textit{mef}(I) (macrolides) and \textit{catQ} (chloramphenicol) in streptococci

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In streptococci \textit{mef}(I) and \textit{catQ}, two relatively uncommon macrolide and chloramphenicol resistance genes, respectively, are typically linked in a genetic module designated IQ module. Though variable, the module consistently encompasses, and is sometimes reduced to, a conserved \textasciitilde 5.8-kb \textit{mef}(I)-\textit{catQ} fragment. The prototype IQ module was described in \textit{Streptococcus pneumoniae}. IQ-like modules have subsequently been detected in \textit{Streptococcus pyogenes} and in different species of viridans group streptococci, where \textit{mef}(E) may be found instead of \textit{mef}(I). Three genetic elements, one carrying the prototype IQ module from \textit{S. pneumoniae} and two carrying different, defective IQ modules from \textit{S. pyogenes}, have recently been characterized. All are integrative and conjugative elements (ICEs) belonging to the \textit{Tn5253} family, and have been designated \textit{ICESpy}029IQ, \textit{ICESpy}029IQ and \textit{ICESpy}005IQ, respectively. \textit{ICESpy}029IQ and \textit{ICESpy}005IQ were the first \textit{Tn5253} family ICEs to be described in \textit{S. pyogenes}. A wealth of new information has been obtained by comparing their genetic organization, chromosomal integration, and transferability. The origin of the IQ module is unknown. The mechanism by which it spreads in streptococci is discussed.

Keywords: \textit{mef}(I), \textit{catQ}, macrolide resistance, chloramphenicol resistance, streptococci, IQ module, integrative and conjugative elements (ICEs)

The macrolide resistance gene \textit{mef}(I) and the chloramphenicol resistance gene \textit{catQ} are relatively uncommon genetic determinants accounting respectively for macrolide and chloramphenicol resistance. In streptococci, however, they are typically associated and genetically linked, since neither has yet been detected without the other. This review illustrates current data on the association and discusses its genetic basis.

\textbf{\textit{mef} GENES}

In streptococci, \textit{mef}-class genes encode efflux-mediated macrolide resistance. Compared to \textit{erm}-class genes encoding methylase-mediated target site modification, \textit{mef} genes usually produce lower-level resistance, only affecting 14- and 15-membered macrolides (M phenotype; Sutcliffe et al., 1996). In streptococci, \textit{mef} genes include a number of subclasses, of which \textit{mef}(A) and \textit{mef}(E) are the most significant. \textit{mef}(A), the first subclass to be discovered (Clancy et al., 1996), is widespread in \textit{Streptococcus pyogenes} but is also common in \textit{Streptococcus pneumoniae}. \textit{mef}(E), discovered in \textit{S. pneumoniae} (Tait-Kamradt et al., 1997), is frequently found in this and other \textit{Streptococcus} species, but is uncommon in \textit{S. pyogenes} (Klaassen and Mouton, 2005). \textit{mef}(I), discovered in \textit{S. pneumoniae} (Cochet et al., 2005; accession no. AJ971089) and subsequently detected in \textit{S. pyogenes} (Blackman Northwood et al., 2009) and viridans group streptococci (VGS; Brenciani et al., 2014), is the most widely investigated among the other \textit{mef} subclasses; these include \textit{mef}(O), \textit{mef}(B), and \textit{mef}(G), detected respectively in \textit{S. pyogenes} (Sangvik et al., 2005), \textit{Streptococcus agalactiae} (Cai et al., 2007), and group G streptococci (Amezaga and McKenzie, 2006; Cai et al., 2007). An \textit{msr}-class gene with homology to \textit{msr}(A), involved in macrolide efflux in \textit{Staphylococcus aureus} (Ross et al., 1990), is located immediately downstream of the \textit{mef} gene: it is usually designated \textit{msr}(D), although a number of variants are associated with different \textit{mef} genes. Cotranscription of \textit{mef}(E) and \textit{msr}(D) in \textit{S. pneumoniae} suggested that the products of the two genes may act as a dual efflux system (Gay and Stephens, 2001).

\textbf{\textit{cat} GENES}

The first and still predominant mechanism of bacterial resistance to chloramphenicol is enzymatic inactivation of the drug by different chloramphenicol acetyltransferases (CATs) encoded by \textit{cat} genes. Two types of CATs have been recognized, A and B, each including a number of structurally and phylogenetically diverse groups (Schwarz et al., 2004). The \textit{catQ} gene, encoding a type-A CAT assigned to a distinct group (Schwarz et al., 2004), was originally described (Rood et al., 1989) and then sequenced (Bannam and Rood, 1991; accession no. M55620) in \textit{Clostridium perfringens} CW531. However, no further \textit{catQ} gene from this species has been described or is available in GenBank, even though genes with 77–92% DNA identities to \textit{catQ} from \textit{C. perfringens} CW531 can be found in several other \textit{Clostridium} species by \textit{in silico} analysis. Different type-A CATs encoded by a number of \textit{cat} genes have been described in streptococci (Trieu-Cuot et al., 1993; Schwarz et al., 2004). With regard to \textit{catQ}, an early study reported a \textit{catQ}-like gene in a \textit{S. agalactiae} strain (Trieu-Cuot et al., 1993). More recently, Mingoia et al. (2007) described \textit{catQ} in two \textit{S. pneumoniae} isolates and found that the \textit{catQ} gene from \textit{S. pneumoniae} Spn529 (the reference isolate) displayed 96.2% DNA identity
(accession no. AJ971089) to the original catQ from *C. perfringens* CW531.

**THE IQ MODULE**

When catQ was first detected in *S. pneumoniae*, it was found to be linked to mef(I) in an ~15.1-kb DNA fragment, containing two identical *tnp*1 transposase genes at either end, that was designated ‘IQ element’ (Mingoia et al., 2007). Here, we prefer to call it ‘IQ module’, since the term ‘element’ might be confusing. Two defective IQ modules were subsequently described in *S. pyogenes*: one was characterized by a shorter region upstream of mef(I), formed only by *tnp*1 and a truncated recombinase gene rec2; the other consisted of the sole mef(I)-catQ fragment (Del Grosso et al., 2011). Very recently, a number of variously defective IQ-like modules have been detected in different VGS species (Brenciani et al., 2014). An interesting characteristic of VGS modules is that they most often bear mef(E) instead of mef(I). Moreover, all IQ-like modules detected in VGS lack the right *tnp*1, i.e., the transposase gene downstream of catQ that is found in the IQ module of *S. pneumoniae*. Despite variability upstream of the mef gene and downstream of catQ, all reported IQ modules appear to share a conserved mef-catQ fragment [six open reading frames (ORFs), ~5.8 kb; Figure 1].

The origin of the IQ module is unknown. The prototype IQ module of pneumococcal origin has been proved to be capable of undergoing excision in circular form, but was unable to be transferred to a recipient (Mingoia et al., 2014).

### FROM THE 5216IQ COMPLEX TO A Tn5253 FAMILY ICE

The first attempt to clarify the genetic basis of the mef(I)-catQ association dates back to 2007, when Mingoia et al. (2007) identified in *S. pneumoniae* a composite structure, that they designated 5216IQ complex (~30.5 kb); it consisted of two moieties, one represented by the IQ module and the other formed by fragments of transposons Tn5252 and Tn916. The Tn5252 fragment corresponds to the conjugal transfer-related (CTR) functional module of Tn5252 (Alarcon-Chaidez et al., 1997); the Tn916 fragment contains a silent tet(M), unexpressed because it lacks the promoter, the ribosome-binding site, and part of the leader peptide. As realized later (Mingoia et al., 2014) also thanks to the new notion of integrative and conjugative element (ICE; Seth-Smith and Croucher, 2009; Walder, 2010), the so-called 5216IQ complex is not in fact a genetic element (Varaldo et al., 2009), but rather part of a larger element, a Tn5253 family ICE.

### GENETIC ELEMENTS CARRYING IQ MODULES

The genetic location of the IQ module has recently been investigated in three strains: one *S. pneumoniae* (Spn529, serotype 11A [Mingoia 07]) and two *S. pyogenes* (Spy029 and Spy605, respectively emm types 25.0 and 12.42 [Del Grosso 11]). All three IQ modules were found to be inserted in Tn5253 family ICEs, which were respectively designated ICESpn529IQ, ICESpyp029IQ, and ICESpy605IQ (Mingoia et al., 2014). It is worth noting that this was the first time that Tn5253 family ICEs were described in natural isolates of *S. pyogenes*. The family prototype, Tn5253, is a long-established conjugative composite element resulting from the insertion of Tn5251 (virtually identical to Tn916) into Tn5252 (Ayoubi et al., 1991). Several studies have recently addressed the heterogeneity of Tn5253-like composite elements (Croucher et al., 2009, 2011; Ding et al., 2009; Henderson-Begg et al., 2009; Mingoia et al., 2011; Wyres et al., 2013), whose integrase genes may belong to one of two types: *int*5252 (the gene of the Tn5253 prototype) or *int*Spyp23 (first described in ICESpyp23ST81 [Croucher et al., 2009]). The complete sequence of Tn5253 is available (accession no. EU351020) and has been extensively analyzed (Iannelli et al., 2014).

ICESpn529IQ has been completely sequenced (accession no. HG965092): it is 59,466 bp long; its G+C content is 36%; and sequence analysis disclosed 66 ORFs. The IQ module spans from *orf*22 to *orf*36 (which correspond to the two identical *tnp*1 transposase genes located at either end of the module); catQ is *orf*23 and mef(I) is *orf*28. ICESpy029IQ and ICESpy605IQ have been
characterized based on PCR mapping, partial DNA sequencing, and restriction analysis. The three ICEs are shown and compared in Figure 2.

ICE_Spn916IQ and ICE_Spy029IQ appear to share a very similar organization. Both carry the integrase gene intSp23 and an identical Tn916 fragment. Aside from the different size of their IQ modules, which nevertheless share the same position and insertion site in the respective ICE, the only noticeable difference between the two ICEs is a transposase gene tnp1 adjacent to intSp23; as a result ICE_Spy029IQ carries a third tnp1 in addition to those found at the ends of the IQ module. In contrast, ICE_Spy005IQ has the integrase gene intSp252 and no Tn916 fragment, and its defective IQ module is found in a different position compared to the IQ modules of ICE_Spn916IQ and ICE_Spy029IQ.

All three ICEs lack the linearized plasmid pC194, which contains the chloramphenicol resistance determinant catC194 and is usually a distinctive cargo of the Tn5252-like moiety of Tn5253 family ICEs. This might be a fitness-related characteristic; as if the ICE no longer needed the plasmid since chloramphenicol resistance is already assured by catQ. It is well known since early studies that the DNA region containing the linearized pC194 is flanked by direct repeats whose recombination may lead to spontaneous curing of the region (Ayoubi et al., 1991; Kiliç et al., 1994). Recent analysis of the sequence of Tn5253 (Iannelli et al., 2014) disclosed a 7,627-bp \( \Omega \)cat(pC194) flanked by two 1,169-bp direct repeats, which contain a toxin-antitoxin gene pair (Palmieri et al., 2013) that has previously been detected in the pathogenicity island PPI-1 of \( S. \) pneumoniae strain TIGR4 (Khoo et al., 2007). In all three ICEs the lack of the integrated plasmid pC194 is associated with the presence in the corresponding region of a single copy of this toxin-antitoxin operon (TA pair), which likely represents the recombination site of \( \Omega \)cat(pC194) and might be the outcome of repair after plasmid excision. Of the three ICEs, only ICE_Spy005IQ has been detected as a free circular form, a critical condition for an ICE to undergo conjugal transfer (Wozniak and Waldor, 2010).

**TRANSFERABILITY OF mef(I) AND catQ**

The transferability of mef(I) and catQ varies with the transfer mechanism (conjugation or transformation) and the strain harboring the IQ module. Using \( S. \) pneumoniae donors, transfer of mef(I) and catQ has never been obtained to any recipient by conjugation, whereas mef(I)- and catQ-positive transformants have been obtained by transformation (Nielsen et al., 2010; Mingoia et al., 2014). It is worth noting that, in experiments using \( S. \) pneumoniae Spn529 as the donor, transformants were obtained at a very low frequency (around \( 10^{-9} \)) and only using a crude lysate as the transforming DNA: this suggests that DNA may undergo changes during the extraction process, which may account for the early negative results (Mingoia et al., 2007). Conversely, using \( S. \) pyogenes donors, conjugation assays yielded no transconjugants from Spy029, whereas transconjugants harboring the entire ICE_Spy005IQ were obtained at high frequencies from Spy005 (Del Grosso et al., 2011; Mingoia et al., 2014). Transformants were obtained with both \( S. \) pyogenes donors, but at a high frequency from strain Spy005 and at a very low frequency from strain Spy029.

**SPREAD OF mef(I) AND catQ IN STREPTOCOCCI**

It has been argued by analogy with other mef gene subclasses (especially in pneumococcal populations) that the spread of mef(I) — and of the IQ module, where it is consistently found — could largely depend on transformation mechanisms (Mingoia et al., 2014). In \( S. \) pneumoniae, the two major mef determinants, mef(E) and mef(A), are carried respectively by the mega element (Gay and Stephens, 2001) and Tn1207.1 (Santagati et al., 2000), whose relationship with the relevant mef gene is roughly comparable to that of the IQ module with mef(I). Similar to the IQ module, the mega element and Tn1207.1 are transferable by transformation rather than by conjugation. While the mega element has a variety of insertion sites in the pneumococcal chromosome and its dissemination is erratic, Tn1207.1 is integrated into the chromosome at a specific site and its dissemination is mainly clonal (Del Grosso et al., 2002). Non-clonal dissemination of the IQ module is suggested by the wide range of serotypes of the \( S. \) pneumoniae isolates that have so far been found to be mef(I)-positive (Cochetti et al., 2005; Reinert et al., 2008; Nielsen et al., 2010); moreover \( S. \) pyogenes strains Spy029 and Spy005 also belong to different \textit{emm} types (Del Grosso et al., 2011). Altogether, these data suggest that the behavior of mef(I) is likely to be more similar to that of mef(E) than of mef(A).
CONCLUDING REMARKS

In streptococci, the macrolide resistance determinant mef(I) and the chloramphenicol resistance determinant catQ, both relatively uncommon within the respective class of resistance genes, are typically linked in a genetic module, designated IQ module. Although the IQ module is uncommon in streptococci, it is likely to be unique to them: it has not been reported outside the genus Streptococcus, and its origin is unknown. Though variable, the IQ module consistently encompasses, and may be reduced to, a conserved ~5.8-kb DNA fragment spanning from mef(I) (or mef(E) in some VGS) to catQ. After early report in S. pneumoniae (Mingoia et al., 2007), IQ-like modules have been detected in S. pyogenes (Del Grosso et al., 2011) and VGS (Breciani et al., 2014); moreover, Mingoia et al. (2014) have recently reported on ongoing studies of isolates of other Streptococcus species (S. agalactiae) harboring linked mef(I) and catQ genes.

In recent years, neither mef(I) nor catQ have been described in the documented absence of the other in streptococci. However, the association is unlikely to have been the rule in earlier investigations. Indeed, the S. agalactiae strain where a catQ-like gene was first described in a streptococcus was erythromycin susceptible (Trieu-Cuot et al., 1993). The same is probably true of clostridia: C. perfringens CW531, where catQ was originally detected and sequenced even though its genetic environment was not investigated, was reported to be resistant, besides chloramphenicol, to tetracycline, but not to erythromycin (Rood et al., 1989); and there is no evidence of associated mef genes in the sequenced clostridial genomes or contigs containing catQ-related genes.

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