

Conjugative Transfer and *cis*-Mobilization of a Genomic Island by an Integrative and Conjugative Element of *Streptococcus agalactiae*

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Putative integrative and conjugative elements (ICEs), i.e., genomic islands which could excise, self-transfer by conjugation, and integrate into the chromosome of the bacterial host strain, were previously identified by *in silico* analysis in the sequenced genomes of *Streptococcus agalactiae* (M. Brochet et al., J. Bacteriol. 190:6913–6917, 2008). We investigated here the mobility of the elements integrated into the 3' end of a tRNA^{Lys} gene. Three of the four putative ICEs tested were found to excise but only one (ICE_515_tRNA^{Lys}) was found to transfer by conjugation not only to *S. agalactiae* strains but also to a *Streptococcus pyogenes* strain. Transfer was observed even if recipient cell already carries a related resident ICE or a genomic island flanked by *attL* and *attR* recombination sites but devoid of conjugation or recombination genes (CIs-Mobilizable Element [CIME]). The incoming ICE preferentially integrates into the 3' end of the tRNA^{Lys} gene (i.e., the *attR* site of the resident element), leading to a CIME-ICE structure. Transfer of the whole composite element CIME-ICE was obtained, showing that the CIME is mobilizable in *cis* by the ICE. Therefore, genomic islands carrying putative virulence genes but lacking the mobility gene can be mobilized by a related ICE after site-specific accretion.

Mobile genetic elements (MGEs) and genomic islands play a key role in bacterial genome evolution by disseminating new genes and phenotypes to the recipient cells (1). The multiplication of bacterial genome sequencing projects in the last few years provides a remarkable opportunity to explore the pool of bacterial genetic mobile elements or "mobilome" (2, 3).

Bacterial genome analyses pointed out a novel class of widespread MGEs, called integrative and conjugative elements (ICEs), and related genomic islands (4-7). These chromosomal elements can excise by site-specific recombination, transfer by conjugation to another bacterial cell and integrate into the chromosome of the recipient cell (4, 7). ICEs are characterized by a combination of modules that can either be involved in their dissemination and maintenance (recombination, conjugation, and regulation modules) or confer adaptive functions to their host (catabolic properties, virulence, and antibiotic resistance) (8, 9). They evolve by acquisition, deletion, and exchange of these modules between different ICEs or with other MGEs (5, 8). The excision of most ICEs relies on an integrase of the tyrosine recombinase family, which catalyzes the site-specific recombination between identical sequences carried by attL (left attachment site) and attR (right attachment site) recombination sites flanking the element. This leads to the excision of a circular form of the ICE harboring an attI site (attachment site of ICE) and to a chromosomal attB (bacterial attachment site) empty site. After transfer, the circular ICE generally integrates into the chromosome of the recipient cell by sitespecific recombination between identical sequences carried by the attI site and the attB site, including the 3' end of a tRNA gene, of a gene encoding ribosomal protein or another gene encoding a conserved protein (4).

In Gram-negative bacteria, the transfer of conjugative plasmids is initiated by a relaxase that nicks plasmid at the origin of transfer (*oriT*) and interacts with a coupling protein to convey DNA to a transport channel crossing the bacterial cell envelope (10). Little is known about the components of the conjugation machinery of conjugative plasmids and ICEs from *Firmicutes* (7). The conjugation machinery of *Firmicutes* has been studied extensively for two conjugative plasmids: pIP501 of *Streptococcus agalactiae* and pCW3 of *Clostridium perfringens* (10). Major proteins of these conjugative systems include a relaxase ensuring DNA processing, an ATPase that likely energizes the DNA transport process, a coupling protein that links the relaxosome with the transport apparatus, and a peptidoglycan hydrolase to facilitate the assembly of the transport channel in the membrane (10). For ICE*Bs1* of *Bacillus subtilis*, the ATPase ConE colocalized with excised ICE*Bs1* DNA at or near the cell poles, suggesting that the conjugation machinery assembles at the donor cell poles (11).

Genomic islands, which carry a recombination module but encode only some the proteins required for conjugation (relaxase, other proteins of the relaxosome, and sometimes the coupling protein), could excise and use the transport apparatus of unrelated ICEs and conjugative plasmids to transfer into a recipient cell and were thus called integrative mobilizable elements (IMEs) (5). Other elements that lack recombination and conjugation modules but are flanked by recombination sites and derived from ICEs have been reported (8, 12). We recently demonstrated in *Streptococcus thermophilus* that a related ICE can integrate in these recombination sites and mobilize in *cis* the element (thus called CIs-Mobilizable Element [CIME]) (13).

We have previously detected 35 different ICEs, IMEs, and related genomic islands in eight sequenced genomes of *S. agalactiae* (group B streptococci) (14), an opportunistic pathogen responsi-

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ble for invasive bacterial diseases in human neonates (15) and causing infections in various animals (16, 17). In the present study, we examined the functionality of putative ICEs of *S. agalactiae* that are integrated at the 3' end of a gene encoding a tRNA lysine (CTT anticodon) and carry a conjugation module distantly related to those of the ICEs RD2 from *S. pyogenes* and ICE*St3* from *S. thermophilus*. We first test their excision as a circular form and then their conjugative transfer inside the *S. agalactiae* species and to other *Firmicutes* species. We also examine the ability of the ICEs to *cis*-mobilize a genomic island integrated into the same locus.

MATERIALS AND METHODS

Bacterial strains and culture conditions. The original strains, spontaneous resistant strains, and plasmids used to obtain recipient or donor strains are listed in Table 1. Strains which were modified or mutated were named according to the modification. For example, the *S. agalactiae* Nem316 Rif^r Str^r strain refers to a spontaneous mutant selected from *S. agalactiae* Nem316 strain which is resistant to rifampin and spectinomycin.

S. agalactiae, Streptococcus uberis, and Streptococcus dysgalactiae subsp. dysgalactiae strains were grown in brain heart infusion (BHI; Difco) broth at 37°C with shaking at 150 rpm. Streptococcus pyogenes and Enterococcus faecalis strains were grown in the same conditions but without shaking. Solid cultures of these species were made on tryptic soy plates supplemented with defibrinated horse blood (5%). S. thermophilus strains were grown in reconstituted skim milk (10% [wt/vol]), M17 broth supplemented with 0.5% lactose (LM17; Oxoid) at 42°C in anaerobic conditions (GENbox Anaer atmosphere generators and incubation jars from bio-Mérieux). Streptococcus salivarius strains were grown in M17 broth supplemented with 0.5% glucose (GM17; Oxoid) at 37°C in anaerobic conditions. Streptococcus mutans strains were grown in Todd-Hewitt broth supplemented with 0.1% of yeast extract (Oxoid) at 37°C in anaerobic conditions. Cultures were supplemented with the following antibiotic when required: chloramphenicol, $16 \,\mu g \,ml^{-1}$; erythromycin, $50 \,\mu g \,ml^{-1}$; rifampin, 75 μ g ml⁻¹; spectinomycin, 500 μ g ml⁻¹; or streptomycin, 250 $\mu g m l^{-1}$.

Escherichia coli strains were grown in Luria-Bertani (LB) broth at 37°C with shaking at 200 rpm. Recombinant plasmids derived from the pG+host9 vector were transformed into *E. coli* EC101 strain, a strain which contains a chromosomal copy of the pWV01 *repA* gene (18), and selected at 37°C on LB medium containing 150 μ g of erythromycin ml⁻¹.

DNA manipulations, PCR, and pulsed-field gel electrophoresis. Preparation of chromosomal and plasmid DNAs was performed according to standard protocols (28).

PCRs and high-fidelity PCRs were carried out according to the manufacturer's instructions using the ThermoPol PCR kit (New England Bio-Labs) and the Phusion high-fidelity DNA polymerase (Thermo Scientific), respectively. PCRs were performed with 4 μ g of DNA template ml⁻¹ and included 30 cycles of amplification. The annealing step was executed at 5°C below the melting temperature (T_m) of the primers for standard PCR (ThermoPol PCR kit) and at a 3°C above the T_m of the primers for high-fidelity PCR (Phusion high-fidelity DNA polymerase). The primers used in the present study were purchased from Eurogentec and are listed in Table S1 in the supplemental material.

Transconjugants were confirmed by PCR using primers located in the SAG2026 (corresponding to the SAL_2079 gene in strain 515, 100% identical to the SAG2026 gene) gene. Transconjugants of *S. pyogenes* were also confirmed by PCR using a primer located in the ICE integrase gene (*int*ICESa03 Fwd) and a primer located in the tRNA^{Lys} gene (tRNA^{Lys}) (see Table S1 in the supplemental material).

The characterization of the CIME-ICE composite elements in the transconjugants of strain Nem316 was performed using three sets of primers: SAG2026-1 and *int*ICESa03 Fwd (for integration in the *attL* site), *int*ICESa03 Fwd and SAG1986 Rev (for integration in the *attI* site), and SAG1986 Fwd and SAG2026-1 (for integration in the *attR* site). Two pairs

of primers (SAG2026-1/tRNAMet Rev and SAG2026-1/tRNA16S Rev) were also used to examine integration in the two other tRNA^{Lys} genes.

Nested PCR were performed to determine the presence of an *attI* site and an *attB* site using two steps. The first step corresponds to a standard PCR except that 25 cycles of amplification were made instead of 30. In the second step, template corresponds to the PCR product obtained in the first step and primers used are internal to this fragment. Primers location and orientation are indicated in Fig. 2 and 5. Conditions used are those of the standard PCR described above. The *attI* and *attB* fragments obtained by PCR were sequenced to confirm their specificity.

Pulsed-field gel electrophoresis experiments were carried out as described previously using SmaI as the restriction enzyme (25).

ICE tagging. ICE was tagged at its left hand side by a resistance gene using the pG+host9 spc vector. Vector pG+host9 spc is a derivative of the pG+host9 plasmid carrying a spectinomycin resistance gene (X. Bellanger, unpublished data). The resistance cassette was obtained from the pSET4S plasmid by digestion with the SpeI restriction enzyme and ligated to XbaI-digested pG+host9 plasmid (Bellanger, unpublished).

The locus tagged is the SAG2026 (SAL_2079) gene encoding the ATPase subunit of an ABC transporter. This gene belongs to an operon homologous to the *yydFGHIJ* operon of from *Bacillus subtilis* (32), which likely encodes a system of synthesis, maturation, and export of a signaling and/or antibacterial peptide. This operon is complete only in strain COH1; two genes are missing in the three other strains.

To construct the mutant, the 5' and 3' ends of SAG2026 gene were independently amplified by PCR (using the primers SAG2026-1-HindIII, SAG2026-2-AvrII, SAG2026-3-AvrII, and SAG2026-4-EcoRI) and subcloned in the pSL1180 vector. An erythromycin resistance gene amplified from pG+host9 was inserted in the internal AvrII site, and the whole insert (containing the two SAG2026 fragments separated by the *ery* cassette) was then cloned into pG+host9spc to give pG+host9spc-SAG2026ery, which was used to transform *S. agalactiae* by electroporation (33). Two crossovers, upstream and downstream from the tagged region, were selected as described previously to obtain replacement of the gene by the *ery* resistance gene (26).

Attempts to obtain strains of *S. agalactiae* devoid of elements integrated in tRNA^{Lys}. In order to find a strain devoid of element integrated into the 3' end the tRNA^{Lys CTT} gene to be used as recipient strain during mating experiments, a large collection of strains of *S. agalactiae* (n = 70) was screened by PCR. The primers used are specific for SAG1986 (SAG1986 Fwd and SAG1986 Rev) and SAG1993 (*intICESa03* Fwd and *intICESa03* Rev) integrase genes of ICE_2603_tRNA^{Lys} and specific for the SAG2026 gene (SAG2026-1 and SAG2026-4).

Two strategies were also tested for curing cells from their resident element. First, plasmids expressing the integrase and excisionase genes and carrying *attR* recombination site of ICE_515_tRNA^{Lys} were constructed and electroporated in the Nem316 and 515 strains. The purpose was to increase the excision of the elements and to select cells which would have lost them after 100 bacterial generations, as previously obtained for ICE*St3* (26). The second strategy relied on exposition of the cells to mitomycin C in order to increase the excision of the ICE, as described for the distantly related ICEs ICESt3 and RD2. This strategy was tested on strain 515. Since the loss of the element could be hampered due to a putative toxin-antitoxin system encoded by ICE_515_tRNA^{Lys} (SAL_2044-SAL_2045 ORF), we also constructed a mutant deleted in the toxin gene to try to obtain cells cured of their ICEs.

Selection of spontaneous rifampin- and streptomycin-resistant mutants to be used as recipient cells in filter-mating experiments. In a first step, rifampin-resistant mutants were selected by plating the parental strain on appropriate media containing rifampin (75 μ g ml⁻¹). In a second step, rifampin- and streptomycin-resistant mutants were selected by plating the rifampin-resistant mutants obtained in the first step on appropriate media containing streptomycin (250 μ g ml⁻¹). All of the selected mutants were confirmed by sequencing of the *sodA* gene (34) (see Table S1 in the supplemental material).

TABLE 1 Strains and plasmids used in this study

Strain or plasmid	Relevant phenotype or genotype ^{<i>a</i>}	Source or reference
Strains		
S. agalactiae		
515	Wild-type strain, carrying a putative ICE (ICE_515_tRNA ^{Lys})	19
2603V/R	Wild-type strain, carrying a composite ICE-IME element (ICE_2603_tRNA ^{Lys}), encoding a	20
	truncated conjugal transfer protein OrfD (ATPase VirB4 homolog)	
COH1	Wild-type strain, carrying a putative ICE (ICE_COH1_tRNA ^{Lys}), encoding a truncated	19
	conjugal transfer protein OrfD (ATPase VirB4 homolog)	
18RS21	Wild-type strain, carrying a putative ICE (ICE_18RS21_tRNA ^{Lys}), encoding three	19
	truncated conjugal proteins (OrfC, OrfD, and OrfL)	
Nem316	Wild-type strain, carrying a putative CIME (CIME_Nem_tRNA ^{Lys})	21
A909	Wild-type strain, carrying a putative IME (IME_A909_tRNA ^{Lys})	19
515 (ICE_515_tRNA ^{Lys} ery)	Strain carrying ICE_515_tRNA ^{Lys} tagged by an Ery ^r cassette, Ery ^r	This study
515 (ICE_515_tRNA ^{Lys} cat)	Strain carrying ICE_515_tRNA ^{Lys} tagged by a Cm ^r cassette, Cm ^r	This study
2603V/R (ICE_2603V/R_tRNA ^{Lys} ery)	Strain carrying ICE_2603V/R_tRNA ^{Lys} tagged by an Ery ^r cassette, Ery ^r	This study
COH1 (ICE COH1 tRNA ^{Lys} ery)	Strain carrying ICE COH1 tRNA ^{Lys} tagged by an Ery ^r cassette, Ery ^r	This study
18RS21 (ICE 18RS21 tRNA ^{Lys} erv)	Strain carrying ICE 18RS21 tRNA ^{Lys} tagged by an Ery ^r cassette, Ery ^r	This study
Nem316 (CIME Nem $tRNA^{Lys}cat$)	Strain carrying carrying CIME Nem $tRNA^{Lys}$ tagged by a Cm ^r cassette, Cm ^r	This study
COH1 Rif ^r Str ^r	Spontaneous rifampin- and streptomycin-resistant mutant	This study
Nem316 Rif ^r Str ^r	Spontaneous rifampin- and streptomycin-resistant mutant	This study
A909 Bif ^r Str ^r	Spontaneous rifampin- and streptomycin-resistant mutant	This study
S mutans	spontaneous maniphi and sreptomychi resistant mutant	This study
UA159	Wild-type strain no element integrated in the tRNA ^{Lys CTT} gene	22
UA159	Spontangous rifemating and strentomy in resistant mutant	ZZ This study
CA159 KII Sti	Spontaneous manipin- and streptomycm-resistant mutant	This study
CID102502	Mild type attain	In stitut Destaur
CIP102505	wild-type strain	Institut Pasteur
11) (0222		collection
JIM8///	Wild-type strain, with a putative IME integrated in the tRNA ²⁷⁰ err gene	23
CIP102503 Rif' Str	Spontaneous ritampin- and streptomycin-resistant mutant	This study
JIM8777 Rif Str	Spontaneous ritampin- and streptomycin-resistant mutant	This study
S. thermophilus		
LMG18311	Wild-type strain, with a putative CIME-IME composite element integrated in the tRNA ^{Lys CTT} gene	24
LMG18311 Rif ^r Str ^r	Spontaneous rifampin- and streptomycin-resistant mutant	This study
S. uberis		
20388	Wild-type strain	25
21458	Wild-type strain	25
20388 Rif ^r Str ^r	Spontaneous rifampin- and streptomycin-resistant mutant	This study
21458 Rif ^r Str ^r	Spontaneous rifampin- and streptomycin-resistant	This study
S. dysgalactiae		,
14998	Wild-type strain	25
16192	Wild-type strain	25
14998 Rif ^r Str ^r	Spontaneous rifampin- and streptomycin-resistant	This study
16192 Rif ^r Str ^r	Spontaneous rifampin- and streptomycin-resistant mutant	This study
S progenes	opontaneo ao mampin' ana orreptom/em resistant matant	1 mo otday
ATCC 12202	Wild-type strain	ATCC
ATCC 12202 Biff	Spontaneous rifampin-resistant mutant	26
F faecalic	Spontaneous maniphi-resistant mutant	20
	Diff Eucl	27
JII2-2	Kii , Fus	Z/ This study
JHZ-2 KII SU	Spontaneous streptomycm-resistant mutant	This study
E. coli	auto E had 5 the (las and AD) E (the DC and AD la 10 1- 73 (15) and A definition of the ' TO 1	10
ECIUI	supe nsa-5 thi (lac-proAB) F (traD6 proAB lacl ⁴ lacZ M15) repA, derivative of strain TG	18
DH5α	supE44 IacU169 (\$80 IacZ M15) hsdR17 endA1 gyrA96 thi-1 relA1	28
Plasmids	2.9 like mMM701 tomo the announcities and instance with (T) () MM70000 F	20
	2.0 kb, pwvv01-type thermosensitive replication origin (1s) from pvE6002, Ery	29
p5E14s	3.8 KD, $pwv01$ -type 1s from $pvE6002$, $lacZ'$, Spc'	30
pSET5s	3.8 kb, pW v01-type Ts from pVE6002, <i>lacZ'</i> , Cm ⁴	30
pSL1180	3.4 kb, pMB1 origin, superpolylinker for vector construction, Amp ^r	31

^{*a*} Abbreviations: Amp^r, ampicillin resistance; Cm^r, chloramphenicol resistance; Ery^r, erythromycin resistance; Fus^r, fusidic acid resistance; Rif^r, rifampin resistance; Spc^r, spectinomycin resistance; Str^r, streptomycin resistance.



FIG 1 Open reading frame (ORF) organization and comparison of the elements integrated into the 3' end of the tRNA^{Lys CTT} gene in *S. agalactiae* 18RS21, 515, COH1, 2603V/R, A909, and Nem316. ORFs appear as arrows (truncated genes are indicated by a delta letter and underlined). The name of the gene where the element is integrated appears in red. Genes of the conjugation module are indicated with blue arrows, genes of the regulation module are indicated with red arrows. When a putative domain has been found for the gene product or a putative function can be assigned to its product, the gene is named accordingly (*rel* for relaxase, CP for coupling protein, ATPase, *xis* for excisionase, and *int* for integrase). Other genes were named according to their similarity to ICE*St1*/*St3* of *S. thermophilus*. The different genes encoding integrase, excisionase, and *relaxase* have been given different numbers. The putative toxin-antitoxin system (SAL2044-2045 and homologs in the other ICEs) appear as a yellow ORFs. The putative *oriT* is indicated by a star. Recombination sites are drawn as vertical rectangles. Black rectangles indicate identical sequences found in *attL*, *attR*, and *attI* sites. Yellow rectangles indicate the arm of *attR* sites and the related arm of *attI* sites, and rel creatingles indicate the arm of *attL* sites and the related arms of *attI* sites. Protein identity higher than 80% is indicated in gray. Gaps in the genome due to missing contigs are indicated by a double slash.

Filter-mating experiments. The filter-mating protocol used was described previously (26). Briefly, both donor and recipient strains were grown overnight. A 15-ml culture in the relevant broth was inoculated with 150 µl of overnight culture of the recipient or the donor strain. The cultures were grown at the relevant temperature until mid-exponential phase (optical density at 600 nm of 0.4). Cultures of the donor and recipient were mixed and centrifuged for 15 min in a prewarmed centrifuge at $4,500 \times g$ to form a cell pellet. The pellet was resuspended in 1 ml of BHI broth with or without 50 μ g of DNase I ml⁻¹ (35), and 150- μ l aliquots were spread on 0.45-µm-pore-size nitrocellulose filters (Sartorius) on tryptic soy agar plates, which were then incubated for 15 h at 37°C. The filters were removed from the agar plates and placed in 50-ml tubes containing 10 ml of sterile BHI broth, and bacteria were recovered by vortexing for 30 s. Various dilutions were spread on agar plates supplemented with the appropriate antibiotics, and the plates were incubated for 24 h in order to count the CFUs of the donor, the recipient, and the transconjugants

DNA sequencing and sequence analysis. DNA sequencing was performed on recombinant plasmids and PCR products by Beckman Coulter genomics. BLASTN, BLASTX, BLASTP, and PSI-BLAST (36) were used to search similarities to sequences in the GenBank database. Sequences were aligned using Vector NTI advance 11 (Invitrogen).

Statistical analysis. Statistical analysis was performed as described by Georgin and Mouet (37) and Cumming et al. (38). The means and standard errors from at least three independent experiments were determined.

RESULTS

Excision of the putative ICEs integrated in the 3' end of a tRNA Iysine gene in *S. agalactiae.* We previously reported that eight sequenced genomes of *S. agalactiae* display a genomic island integrated into the 3' end of the tRNA^{Lys CTT} gene (14). Among these elements, four could be ICEs (in strains 18RS21, 515, COH1, and 2603V/R). The name that was given to the elements is composed of the nature of the element followed by the name of the strain and the integration locus. One of these putative ICEs, ICE_2603_tRNA^{Lys}, is a composite element that carries an additional recombination module, an additional putative relaxase gene and an internal *attI* site that probably results from the accretion of a putative ICE (left side from *attL* to *attI*) and a putative IME (right side from *attI* to *attR*) (Fig. 1).

We looked for excised circular forms of these four putative ICEs: one for ICE_515_tRNA^{Lys}, ICE_COH1_tRNA^{Lys}, and ICE_18RS21_tRNA^{Lys} (Fig. 2A) and three different possible circular forms for ICE_2603V/R_tRNA^{Lys} since this element carries an internal *att1* site (Fig. 2C). By nested PCR, we detected an excised circular form for ICE_515_tRNA^{Lys} and ICE_18RS21_tRNA^{Lys} but not for ICE_COH1_ tRNA^{Lys} (Fig. 2B). An empty *attB* site was, however, detected for this latter strain (data not shown). Furthermore, two different excised forms were observed for ICE_2603_tRNA^{Lys} (Fig. 2D), indicating that the IME integrated beside the ICE in this strain is able to excise alone or together with the ICE. All of the fragments obtained by nested PCR were confirmed by sequencing.

Test of intraspecies conjugative transfer of the putative ICEs integrated in the 3' end of a tRNA lysine gene in *S. agalactiae*. To test whether the four putative ICEs described above are able to transfer by conjugation, they were tagged at their left extremity by an erythromycin resistance cassette after allelic replacement of a gene encoding a membrane subunit of a putative ABC transporter (SAL_2079/SAG2026 gene) (Fig. 1).

The strains harboring these tagged elements were then used as donor cells in filter mating experiments with three different recipient strains that all harbor elements flanked by *attL* and *attR* sites: (i) strain COH1, which harbors a putative ICE as mentioned previously; (ii) strain A909, which harbors a putative IME (i.e., an



FIG 2 Schematic representation of the site-specific excision of the ICEs integrated in the 3' end of the *S. agalactiae* tRNA^{Lys} gene. Primers used to detect DNA molecules resulting from site-specific recombination events are indicated by letters. (A) Model of ICE_515_tRNA^{Lys}, ICE_COH1_tRNA^{Lys}, and ICE_18RS21_tRNA^{Lys} excision. (B) Electrophoresis of nested PCR product obtained for the 515, COH1, and 18RS21 strains of *S. agalactiae*. (C) Model of ICE_603V/R_tRNA^{Lys} excision where three different circular forms could be obtained. (D) Electrophoresis of nested PCR products obtained for the 2603V/R strain.

element with a recombination module, a gene encoding the relaxase, but lacks genes required for mating pore assembly); and (iii) strain Nem316, which carries an element devoid of recombination and conjugation genes (CIME). We were not able to test a recipient strain devoid of element integrated in the tRNA^{Lys CTT} gene since all our efforts to obtain such a strain (by screening of a large collection of strains or by artificial curing of a strain carrying an element) were unsuccessful (see Materials and Methods for more details).

ICE_515_tRNA^{Lys} was found to transfer by conjugation to the Nem316 strain and to the COH1 strain at frequencies of $4.0 \times 10^{-7} \pm 0.8 \times 10^{-7}$ and $0.7 \times 10^{-7} \pm 0.1 \times 10^{-7}$ CFU of transconjugants per CFU of donor cells, respectively. As expected, PCR on the SAL_2079/SAG2026 gene gave two fragments for the transconjugants (one for the gene present on the resident element and another of higher size corresponding to the gene that is carried by the incoming ICE and is interrupted by the erythromycin cassette) (data not shown). No transconjugant was obtained when cell-cell contacts were prevented by placing a filter between donor and recipient cells. Furthermore, the addition of DNase I in the growth medium had no impact on the number of transconjugants, indicating that DNA transfer occurred by conjugation. No transconjugant was obtained when the A909 strain was used as recipient cells.

ICE was found to retransfer if transconjugants (carrying a CIME *cat*-ICE *ery* element) were used as donor cells and rifampinstreptomycin mutants of the Nem316 strain were used as recipient cells. The frequency of retransfer was higher $(2.2 \times 10^{-5} \pm 0.9 \times 10^{-5}$ CFU of transconjugants per CFU of donor cells) than the one obtained for transfer from strain 515 to strain Nem316. This indicates that ICE_515_tRNA^{Lys} is able to self-transfer by conjugation and therefore is an ICE. In contrast, no conjugative transfer was observed for the three other putative ICEs tested (ICE_2603_ *tRNA^{Lys}*, ICE_18RS21_ *tRNA^{Lys}*, and ICE_COH1_*tRNA^{Lys}*).

Test of interspecies conjugative transfer of ICE_515_ tRNA^{Lys}. Since ICE_515_tRNA^{Lys} was shown to transfer by conjugation to a S. agalactiae recipient strain, we also tested whether it can transfer to other bacterial species. Seven different species belonging to the Firmicutes were tested as recipient cells: S. salivarius (CIP102503 and JIM8777), S. thermophilus (LMG18311), S. pyogenes (ATCC 12202), S. uberis (20388 and 21458), S. dysgalactiae subsp. dysgalactiae (14998 and 16192), S. mutans (UA159), and E. faecalis (JH2-2). Spontaneous rifampin and streptomycin mutants were first selected for these strains in order to use them as recipient cells in the mating experiments. Transconjugants were obtained only when using S. pyogenes as recipient cells. However, even for this species, transconjugants were obtained only in three experiments out of nine experiments performed, and only 20 different clones of transconjugants were isolated. The acquired ICE is integrated into the 3' end of the tRNA^{Lys CTT} gene of the transconjugants, as shown by PCR using primers located in this tRNA^{Lys} gene and in the integrase gene of ICE_515_tRNA^{Lys} (Fig. 3A). In addition, they have the same genetic background than the recipient S. pyogenes strain, as shown by pulsed-field gel electrophoresis (Fig. 3B).

Site-specific accretion between ICE_515_tRNA^{Lys} and related elements. Since the Nem316 strain used as recipient cells for the mating experiments carries an internal recombination site in its element integrated in the tRNA^{Lys CTT} gene, the incoming ICE could integrate into three different positions: the left (*attL*), the internal (*attI*), or the right (*attR*) recombination sites of the resident CIME (Fig. 4). Testing of 100 transconjugants by PCR indicated that integration occurs preferentially in the *attR* recombination site (96% versus 4% in the *attL* site and 0% in the *attI* site),



FIG 3 Analysis of eight clones of transconjugants obtained after filter-mating experiments between donor strain 515 (carrying ICE_515_tRNA^{Lys}) and recipient strain *S. pyogenes* ATCC 12202. (A) Amplification of the right junction of the ICEs in these clones as for donor cells, using primers specific of the integrase genes and of the tRNA^{Lys} gene (lane D). The negative control is *S. pyogenes* ATCC 12202, used as recipient cells (lane R). Lane M shows DNA molecular weight marker. (B) Analysis of the transconjugants by pulsed-field gel electrophoresis in parallel to the donor strain (lane D) and the recipient strain (lane R).

giving rise to a CIME-ICE *ery* tandem (Fig. 4). Mating experiments were also performed with a Nem316 recipient strain carrying a CIME tagged with a chloramphenicol resistance gene, giving rise to CIME *cat*-ICE *ery* tandem. PCRs were performed to amplify the two other tRNA^{Lys} genes to determine whether they can be used as secondary integration sites. None of the 100 transconjugants tested carried an ICE in the two other tRNA^{Lys} genes, indicating that integration is specific to the tRNA^{Lys} cTT gene.

Strain COH1 also gave transconjugants, although it already carries a putative resident ICE that is integrated into the same site and harbors very closely related conjugation and recombination modules. The incoming ICE can thus theoretically integrate at two different positions: in the left (*attL*) or in the right (*attR*) recombination site of the resident ICE. Analysis of 12 transconjugants indicated that they all integrated in the *attR* recombination site giving rise to the formation of a tandem of ICEs.

Conjugative mobilization of CIME 316 tRNA^{Lys} in *cis* by ICE_515_tRNA^{Lys}. The empty *attB* site (Fig. 5B) and the *attI* site (Fig. 5C) resulting from the excision of the whole CIME cat-ICE ery tandem were detected by nested PCR in transconjugants, showing the intracellular mobilization in *cis* of the CIME by the ICE. The conjugative transfer of this composite element was then investigated by filter matings using, as donor cells, the transconjugant deriving from Nem316 strain carrying this tandem and, as recipient cells, rifampin-streptomycin spontaneous mutant of the Nem316 strain. Putative transconjugants deriving from Nem316 Rif^r Sm^r were recovered on agar plates containing rifampin, streptomycin, and either erythromycin or chloramphenicol. Among 900 erythromycin-resistant transconjugants analyzed, only one carried both ICE and CIME; the others carried only the ICE alone. The only transconjugant recovered by plating on chloramphenicol-containing agar was shown to carry both ICE and CIME. These two transconjugants were analyzed by pulsed-field gel electrophoresis. Their electrophoresis pattern reflects the acquisition of a CIME-ICE tandem (one or two copies depending on the transconjugant) (Fig. 6). Therefore, the accretion has led not only to the intracellular mobilization in cis of the CIME by the ICE but also to its conjugative mobilization in cis.



FIG 4 Schematic representation of the site-specific integration of ICE_ 515_tRNA^{Lys} in the genome of strain Nem316 and schematic localization of primers used to detect DNA molecules resulting from the site-specific recombination events. ICE_ 515_tRNA^{Lys} appears in black, and CIME_ Nem_tRNA^{Lys} appears in gray.

DISCUSSION

A circular form was detected for three out of the four ICEs integrated in a tRNA^{Lys} gene of S. agalactiae tested. In strain 2603V/R, the whole composite ICE and the IME corresponding to its right end are able to excise. The lack of excision of the putative ICE corresponding to the left part of the composite ICE suggests that the integrase requires sequences present in the tRNA^{Lys} gene which are absent in the internal attI site. Alternatively, it is possible that integrase binding to the *attL* site and to *attR* moiety of *attI*_{in} is not optimal for recombination, explaining why the distal attR site is used for recombination instead of internal attR moiety of attI_{in}. Since the *attR* and *attL* sites of the ICE and of the IME are very different, the excision of the whole composite element ICE-IME requires that the integrase (of the ICE or of the IME) recognizes the other type of sequence (attL of the ICE if excision is mediated by the integrase of the IME or *attR* of the IME if the excision is ensured by the integrase of the ICE). This suggests that this IME is functional and could thus be trans- or cis-mobilized by an ICE. In both cases, it would need a functional helper conjugation pore to be transferred to a recipient cell. Although putative IMEs are frequent in genomes in particular in S. agalactiae (14) and the presence of a high number of isolated relaxase genes in bacterial chromosomes suggests that these elements are very frequent in bacteria (6), very few IMEs have actually been characterized until now in bacteria (5, 39). No circular form was detected for the fourth ICE (ICE_COH1_tRNA^{Lys}), but an empty integration site was detectable. This could indicate a loss of circular form in this strain in the extraction conditions tested due to less quantity or higher degradation of circular form than in the other strains. Alternatively, the absence of amplification can also be due to the longer expected size of the amplified fragment in this strain that carries two additional genes at the left-hand side of its ICE.

Screening of a large collection of strains of *S. agalactiae* did not enable us to find a strain that does not carry a genetic element integrated in the tRNA^{Lys CTT} gene in order to be used as recipient in the mating experiments. Even a strain isolated from fish carries



FIG 5 Analysis of the site-specific excision of the CIME *cat*-ICE *ery* composite element integrated in the 3' end of the tRNA^{Lys} gene of transconjugants of *S. agalactiae* Nem316. (A) Schematic representation of the circular form (with its *attI* site) and the empty *attB* integration site resulting from excision of the composite element. Primers used to detect DNA molecules resulting from site-specific recombination events are indicated by letters. (B) Amplification of an *attB* site in 10 transconjugants of *S. agalactiae* Nem316. (C) Amplification of an *attI* site in the same 10 clones. ICE_515_tRNA^{Lys} appears in black, and CIME_Nem_tRNA^{Lys} appears in gray.

an element at this locus (40), which thus appears as a hot spot of integration in this species. All curing attempts failed, although we tested different strategies that proved to be efficient for distantly related ICEs (26, 41). This could be due to the presence of a toxinantitoxin system that kills the cells that have lost the ICE. However, interruption of the toxin gene of the putative toxin-antitoxin system identified on ICE_515_tRNA^{Lys} did not allow curing the strain even after a mitomycin C treatment. Our hypothesis is that the ICE replicates in the cell as reported for two ICEs belonging to the ICESt3 family (26, 41), leading to a high frequency of reintegration in the chromosome even if excision is artificially increased. This hypothesis would also explain why the numerous attempts to interrupt the excisionase, integrase, relaxase, and other genes of



FIG 6 Pulsed-field gel electrophoresis analysis of the two transconjugants (M1 and M2) obtained after conjugative mobilization in *cis* of the CIME *cat* by ICE *ery*. A strain deriving from Nem316 and harboring a tandem CIME *cat*–ICE *ery* (indicated by "D" on the schema) and a mutant strain resistant to rifampin and streptomycin deriving from Nem316 already harboring a resident CIME (indicated as recipient "R") were used as donor and recipient strains, respectively.

the ICE failed (data not shown). Since we did not find a *S. agalactiae* strain with an empty tRNA^{Lys} gene site, we used in the mating experiments a recipient strain that already carries a genetic element (CIME, IME, or ICE) integrated at this locus.

In this study, we demonstrated the functionality of an ICE carried by a human isolate of S. agalactiae. This is only the second description of conjugative transfer of an ICE in this species (the first one is TnGBS2 which uses a DDE transposase for its integration instead of a tyrosine recombinase [42] and carries an unrelated conjugation module). This ICE is related to two other ICEs whose conjugative transfer has been demonstrated in other streptococcal species: (i) RD2 of S. pyogenes (67 to 96% of identity for the proteins involved in conjugation transfer but different recombination module enabling integration into the 3'end of tRNA^{Thr} gene) (41) and (ii) ICESt3 of S. thermophilus (35 to 67% of protein identity for the conjugation proteins, unrelated recombination module providing specific integration into the 3'end of *fda* gene) (26). ICE_515_tRNA^{Lys} was successfully transferred by conjugation to two other S. agalactiae strains: Nem316 strain (carrying a CIME) and COH1 strain (carrying a defective ICE). This led in both cases to an accretion of two elements (CIME-ICE or ICE-ICE) with a preferential integration of the incoming ICE at the attR recombination site (that includes the end of the tRNA^{Lys} gene). This attR preference was also observed for ICESt3 of S. thermophilus, although it encodes an unrelated integrase that catalyzes the integration in the *fda* gene instead of the tRNA^{Lys} gene (13). This suggests that the integrase or a cofactor needs to bind sequences present in the tRNA^{Lys} gene but absent in *attL* and attI sites for efficient integration of the ICE.

The frequency of conjugative transfer was low compared to the frequency of intraspecies transfer reported for ICEs of other *Firmicutes* (10-fold less than ICESt3 of *S. thermophilus* [26] and RD2 of *S. pyogenes* [41]). One hypothesis is that the occupation of the recipient *attB* site by an element flanked by *attR* and *attL* recombination sites reduces the frequency of ICE_515_tRNA^{Lys} transfer, as was previously found for ICESt3 (13). The efficiency of ICE

integration in recombination sites flanking the resident element is likely reduced compared to an empty integration site. In addition, the frequency of transfer is even lower when the resident element of the recipient strain is an ICE (i.e., in strain COH1). This could be due to an exclusion mechanism as described for other ICEs (7). Furthermore, the *attR* recombination site of IME_*A909_tRNA*^{Lys} and CIME_*Nem316_tRNA*^{Lys} resident elements greatly differs from the *attR* site of ICE_*COH1_tRNA*^{Lys}.

The genetic background of the recipient cells also likely plays a role since the frequency of transfer was 100-fold higher when transferring the ICE between strains with the same genetic background (experiments of retransfer using Nem316 strain as donor and recipient strain). This could be due to restriction-modification systems that degrade foreign DNA, including incoming ICE. Furthermore, no conjugative transfer of ICE 515 tRNA^{Lys} was obtained when the A909 strain (carrying an IME) was used. A CRISPR system has been described recently in this strain (43). Analysis of the spacers present in this CRISPR system indicate that three of them match a sequence found in ICE 515 tRNA^{Lys} (the coupling protein gene, the intergenic region between coupling protein and relaxase genes, and the gene of ICE_515_tRNA^{Lys} encoding a putative DNA adenine methylase). Strain A909 thus likely carry a functional CRISPR system which targets foreign DNA including ICEs belonging to the same family than ICE_515_tRNA^{Lys} and protect cell from invasion by these MGEs.

No conjugative transfer was observed for the three other putative ICEs tested (ICE_*COH1_tRNA^{Lys}*, ICE_*2603_tRNA^{Lys}*, and ICE_*18RS21_tRNA^{Lys}*). These three elements all carry an *orfD* pseudogene. This conjugation gene encodes a putative ATPase belonging to the FtsK superfamily with characteristic Walker A and Walker B domains. It is thus a pTi VirB4 homolog like ConE (YddE) of ICEBs1, TcpF of pCW3, or Orf16 of Tn916. This protein could play a role in energizing the conjugation machinery and as such is likely crucial for conjugative transfer of the element. This would explain why we did not observe conjugative transfer of ICE_ *COH1_tRNA^{Lys}*, ICE_*2603_tRNA^{Lys}*, and ICE_*18RS21_tRNA^{Lys}*.

Interspecies transfer was obtained only with S. pyogenes as recipient cells. In this species, the tRNALys CTT gene displays one difference in the 11 terminal base pairs compared to S. agalactiae. Other species tested in the present study (Streptococcus salivarius, *Streptococcus uberis*, and *S. thermophilus*) carry a tRNA^{Lys CTT} gene with a 3' end identical to the gene of S. agalactiae. Other factors thus impact the compatibility of donor-recipient cells. At least two of the strains tested as recipient already carry a putative IME integrated in the tRNA^{Lys CTT} gene. This could limit the integration of the incoming ICE. Little is known about the cell-cell contacts that need to be established between donor and recipient cells during conjugative transfer. These contacts could be more difficult between bacteria of different species. Furthermore, restriction-modification systems probably interfere with the maintenance of ICE foreign DNA in the recipient cell. Even if we did not find in vitro conditions propitious to interspecies conjugative transfer of ICE_515_tRNA^{Lys}, there are evidences of a spread of this type of ICE in other streptococcal species. For example, S. urinalis 2285-97 carries an almost identical ICE, and we recently identified an ICE integrated in a tRNA^{Lys} gene and carrying a related conjugation module in a strain of S. uberis (unpublished results). We are currently characterizing the functions encoded by the accessory genes carried by ICE_515_tRNA^{Lys} since they could confer an adaptive advantage to the recipient strains. We showed in particular that this ICE encodes a functional CAMP factor toxin (a gene also found on *S. urinalis* putative ICE) (44). Transfer of ICEs carrying such accessory gene to other bacterial species would thus disseminate this toxin.

Conjugative transfer of ICE_515_tRNA^{Lys} to the Nem316 that carries a CIME led to a CIME-ICE tandem in the recipient strains, thus offering us the opportunity to examine if ICE 515 tRNA^{Lys} is able to *cis*-mobilize genes of CIME_Nem_tRNA^{Lys}. The attB and attI sites resulting from the coexcision of both elements were detected by PCR, showing the intracellular mobilization in cis of the CIME by the ICE. Although quite infrequently (0.1% of the events of retransfer), the ICE-CIME tandem was also shown to transfer by conjugation. Such phenomenon has been described recently for ICESt3 of S. thermophilus (13) and could be a shared characteristic of ICEs using site-specific recombinases to excise and integrate. ICE_COH1_tRNA^{Lys}, ICE_2603_tRNA^{Lys}, and ICE_ 18RS21_tRNA^{Lys}, although not self-transferable, could also be transferred by conjugation after accretion of an ICE and cis-mobilization. Since ICE_2603_tRNA^{Lys} is able to excise, it could also be transferred by conjugation by using the mating apparatus encoded by a related ICE (mobilization in *trans*).

ICEs likely play an even higher role in horizontal gene transfer since it was reported recently that ICEBs1 of B. subtilis is able to mobilize in trans plasmids lacking dedicated mobilization proteins (45). Furthermore, Tn916 was found to mobilize in trans MTnSag1, an element from S agalactiae which encodes a DDE transposase and carries its own oriT unrelated to the one of Tn916 but lacks conjugation or mobilization genes (46). Mobile genomic islands (MGI) carrying their own tyrosine integrase and an their own oriT but no conjugation or mobilization gene were also reported to be mobilized in trans by an ICE harboring a related oriT in Vibrio (47). Mobilization in cis can also lead to conjugative transfer of chromosomal sequences (up to 1 Mb) located at the 5' side of the MGI (47). Conjugative transfer of chromosomal DNA (up to 334 kb) by an Hfr-type mobilization initiated from the oriT of a genomic island was also reported in S. agalactiae (48). cis- and trans-mobilization of DNA by ICEs are thus probably very common and likely contribute to the evolution of genomic islands and bacterial genomes.

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