Presence and Characterization of a Novel cfr-Carrying Tn558 Transposon Derivative in *Staphylococcus delphini* Isolated From Retail Food

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Antimicrobial resistance has become a major public health threat. Food-related *Staphylococcus* species have received much attention due to their multidrug resistance. The *cfr* gene associated with multidrug resistance has been consistently detected in food-derived *Staphylococcus* species. In this retrospective study, we examined the prevalence of *cfr*-positive *Staphylococcus* strains isolated from poultry meat in different geographical areas of China from 2011 to 2016. Two *cfr*-positive *Staphylococcus delphini* strains were identified from poultry meat in China. Comparative and whole-genome analyses were performed to characterize the genetic features and overall antimicrobial resistance genes in the two *S. delphini* isolates 245-1 and 2794-1. Whole-genome sequencing showed that they both harbored a novel 20,258-bp *cfr*-carrying Tn558 transposon derivative on their chromosomes. The Tn558 derivative harbors multiple antimicrobial resistance genes, including the transferable multiresistance gene *cfr*, chloramphenicol resistance gene *fexA*, aminoglycoside resistance genes *aacA-aphD* and *aadD*, and bleomycin resistance gene *ble*. Surprisingly, within the Tn558 derivative, an active unconventional circularizable structure containing various resistance genes and a copy of a direct repeat sequence was identified by two-step PCR. Furthermore, core genome phylogenetic analysis revealed that the *cfr*-positive *S. delphini* strains were most closely related to *S. delphini* 14S03313-1 isolated from Japan in 2017 and 14S03319-1 isolated from Switzerland in 2017. This study is the first report of *S. delphini* harboring a novel *cfr*-carrying Tn558 derivative isolated from retail food. This finding raises further concerns regarding the potential threat to food safety and public health safety. The occurrence and dissemination of similar *cfr*-carrying transposons from diverse *Staphylococcus* species need further surveillance.

Keywords: Tn558, cfr, *Staphylococcus delphini*, unconventional circularizable structure, multidrug resistance
INTRODUCTION

In recent years, resistance in bacteria has spread worldwide and presents a serious threat to human health. Linezolid is an oxazolidinone antibiotic and is considered as the last-resort antibiotic for the treatment of infections caused by multidrug-resistant (MDR) Gram-positive pathogens, including Staphylococcus species (Wilson et al., 2008). The antibiotic targets the P site in the peptidyl transferase center of the 23S ribosomal RNA of the 50S ribosomal subunit, acting on this target and blocking protein synthesis (Aoki et al., 2002). In fact, due to the synthetic nature of the drug, resistance to this antibiotic is rare. However, the cfr gene could mediate resistance to linezolid (Long et al., 2006). This gene encodes a methyltransferase that catalyzes the posttranscriptional methylation of adenosine at nucleotide position 2503 (Escherichia coli numbering) in 23S rRNA, which replaced the target of binding for linezolid (Corinna et al., 2005; Giessing et al., 2009; Anna et al., 2016). However, due to overlapping binding sites, cfr methylation also confers resistance to four other classes of antimicrobial agents and results in the PhLOPSA multiresistance phenotype, including resistance to phenicols, lincosamides, oxazolidinones, pleuromutilins, and streptogramin A compounds (Long et al., 2006; Anna et al., 2016). Interestingly, cfr is often associated with erm, fexA, lsa(B), and ter(L), which can assist in co-selecting the cfr gene and in its spread (Shen et al., 2013; Mendes et al., 2014).

Generally, the cfr gene is often associated with mobile genetic elements (MGEs) (plasmids, integrative, and conjugative elements or transposons), which have great potential for dissemination (Shen et al., 2013). Tn558 is one of these bacterial transposons and was first identified on the plasmid pSCFS2 harboring the antimicrobial resistance gene (ARG) fexA from Staphylococcus lentus (Kehrenberg and Schwarz, 2005). Currently, this transposon is often harbored with cfr, and derivatives of Tn558 usually carry other acquired ARGs (Kehrenberg et al., 2007; Li et al., 2018). Therefore, this transposon plays an important role as vectors in the spread of transposon-borne ARGs.

Members of the genus Staphylococcus are widespread in nature and play vital roles in disease causation in humans and animals (McGavin and Heinrichs, 2012; Vrbovská et al., 2020). Among these species, Staphylococcus delphini is a pathogen that causes animal and human infections (Magleby et al., 2019; Ruiz-Ripa et al., 2019). It belongs to the Staphylococcus intermedius group and was first described in purulent skin lesions of dolphins (Varaldo et al., 1988). S. delphini is further separated into two subgroups, groups A and B, based on the phylogenetic analysis of the sodA, hsp60, and nuc genes and DNA–DNA hybridization (Sasaki et al., 2007). Although this staphylococcal species is poorly documented due to misidentification with S. intermedius, it has been isolated from humans and a wide range of diseased animals, including domestic pigeons, camels, horses, magpies, cinereous vultures, and mustelids, which serve as the natural hosts of S. delphini group A (Devriese et al., 2005; Sasaki et al., 2007; Sledge et al., 2010; Guardabassi et al., 2012; Sudagidan and Aydin, 2012; Stull et al., 2014; Magleby et al., 2019; Ruiz-Ripa et al., 2019).

In this retrospective study, we examined the prevalence of cfr-positive Staphylococcus isolates in poultry meat from 2011 to 2016. We determined the complete genome sequence of cfr-positive S. delphini and described their phenotypic and genotypic profiles. This is the first report of a Tn558 derivative-embedded cfr in S. delphini isolated from retail food.

MATERIALS AND METHODS

Bacterial Isolation

From July 2011 to June 2016, we collected 4,300 retail food samples from supermarkets, fairs, and farmer markets, covering most of the provincial capitals of China (Supplementary Figure 1), and isolated 1,581 Staphylococcus strains, including Staphylococcus aureus, Staphylococcus argenteus, S. delphini, Staphylococcus epidermidis, and other staphylococci from 1,063 positive samples from all the sampling sites (Wu et al., 2018a,b). During the retrospective study of cfr-positive Staphylococcus species among these isolates, the cfr-positive strains 2451 and 2794-1 were isolated from frozen duck wings in Guangzhou 2013 and frozen duck legs in Kunming 2014, respectively. The isolates were further identified as S. delphini by the MALDI-TOF/MS system (Bruker, Bremen, Germany) (Decristophoris et al., 2011).

PCR Detection

The presence of the resistance gene cfr was identified by PCR and Sanger sequencing (Kehrenberg et al., 2009). The presence of the two direct repeats (DRs) and circular intermediate translocatable units (TU) was detected by PCR and inverse PCR (the primers and conditions are shown in Table 2). To minimize the detection of artificial products, a high-fidelity polymerase (PrimeSTAR GXL DNA Polymerase, Takara, Dalian, China) and an 8-min elongation step were used (Tansirichaiya et al., 2016). The amplicons obtained by PCR and inverse PCR experiments were subjected to Sanger sequencing.

Antimicrobial Susceptibility Testing

Minimum inhibitory concentrations (MICs) were determined using a standard broth dilution method according to the CLSI guidelines with S. aureus ATCC 29213 as a quality control strain (Weinstein and Clinical and Laboratory Standards Institute, 2018). The MICs for all of the following antimicrobials were determined: FFC, florfenicol; CHL, chloramphenicol; CLI, clindamycin; TIA, tiamulin; LZD, linezolid; K, kanamycin; ERY, erythromycin; FOX, cefoxitin; VAN, vancomycin; RIP, rifampicin; and DAP, daptomycin. The MIC breakpoints of each antibiotic, except florfenicol, were used as recommended by the current CLSI guidance (Weinstein and Clinical and Laboratory Standards Institute, 2018). For florfenicol, the results were interpreted according to the Veterinary CLSI (VET01-A5).

Whole-Genome Sequence and Analysis

Genomic DNA for whole-genome sequencing was extracted from the cfr-positive strains using a genomic extraction...
kit (Magen Biotech, Guangzhou, China) according to the manufacturer’s instructions. Whole-genome sequencing of the cfr-positive strains was performed using the Illumina HiSeq Xten platform (800-bp paired-end reads with 100-fold average coverage) and a PacBio Sequel II sequencing instrument (100-fold average read depth). The chromosome sequences were assembled into one scaffold using the software SMRT Portal, version 3.2.0. The genomic DNA annotation was performed in Prokka NCBI-BLASTP/BLASTX (Torsten, 2014). The single-nucleotide polymorphisms (SNPs) between strains 245-1 and 2794-1 were identified with Snippy software1.

The acquired antibiotic resistance genes were identified by ResFinder 3.02 and were further verified through a BLAST search against the Comprehensive Antibiotic Resistance Database (EA et al., 2012). The genetic environment of the cfr gene was analyzed using BLAST3, followed by visualization of the comparative cfr multiresistance region (MRR) with Easyfig, v2.2.2 (Sullivan et al., 2011).

Phylogenetic Analysis
All publicly available draft genome sequences of S. delphini strains were acquired (22 strains with at least 50 × read coverage), and core SNP alignments were produced via Snippy using the S. delphini 8086 complete genome sequence (ASMO011v1) as a reference (see text footnote 1). The maximum-likelihood (ML) phylogenetic tree was constructed with RAxML-NG based on the ML optimality criterion (Kozlov et al., 2019). The locations of recombined regions on each branch were detected, and this tree was reconstructed by ClonalFrameML (Didelot and Wilson, 2015). FigTree, v1.4.3, was used to finalize the tree visualization (Morariu et al., 2008).

Nucleotide Sequence Accession Numbers
The complete genomic sequences of 245-1 and 2794-1 have been deposited in GenBank: 245-1 (GenBank ID: CP063368) and 2794-1 (GenBank ID: CP063367).

RESULTS

Phenotypic Characteristics of cfr-Positive S. delphini
In this study, 245-1 and 2794-1 displayed the same MDR profiles. Antimicrobial susceptibility testing showed that these strains were resistant to chloramphenicol, florfenicol, tiamulin, clindamycin, and linezolid, exhibiting a high level of resistance to florfenicol (MIC = 256 µg/ml), chloramphenicol (MIC > 128 µg/ml), and tiamulin (MIC > 128 µg/ml). Moreover, the isolates were susceptible to vancomycin, daptomycin, and rifampicin (Table 1).

Basic Genomic Information for cfr-Positive S. delphini
To understand the molecular characteristics and resistomes of the two strains of S. delphini, they were submitted for whole-genome sequencing. Basic information related to the complete genome sequence of cfr-positive S. delphini is shown in Figure 1. The chromosomal DNA of 245-1 and 2794-1 consisted of 2,708,646 bp with 2,486 predicted ORFs along with 102 RNAs and 2,707,963 bp with 2,486 predicted ORFs along with 102 RNAs, respectively. The genome analysis of the complete chromosomal DNA revealed that there were 166 variants between the chromosomes of 245-1 and 2794-1, and there were multiple ARGs located on their chromosomes, including fexA (conferring resistance to chloramphenicol), aacA-aphD and aadD (resistance to aminoglycosides), ble (resistance to bleomycin), and the multiresistance gene cfr (resistance to phenicols, lincosamides, oxazolidinones, pleuromutilins, and streptogramin A).

Core Genome Phylogenetic Analysis of cfr-Positive S. delphini
To further investigate the potential sources of cfr-positive S. delphini 245-1 and 2794-1, we performed a core genome phylogenetic analysis of all publicly available draft genome sequences of S. delphini strains. The phylogenetic analysis shows that 245-1 and 2794-1 are most closely related to S. delphini 14S03331-1 (GCA_0002374125.1) isolated from Japan in 2017 and 14S03319-1 (GCA_002369675.1) isolated from Switzerland in 2017 (Figure 2). This phylogenetic analysis did not reveal the origin of 245-1 and 2794-1, indicating that the scarcity of genomic sequences may be the constraint, and further genomic sequencing is needed to identify the source of the cfr-positive strains.

Genetic Environment of cfr Located on a Novel Tn558 Transposon Derivative
Genomic mining revealed that the cfr gene, along with four other ARGs, namely, fexA, aacA-aphD, aadD, and ble, was located on a 20,258-bp (62,847–83,104 nt on the chromosomes of 245-1 and 2794-1 in Figure 1) MRR on the chromosomes. Further BLAST analysis showed that the ARGs aacA-aphD, aadD, ble, and cfr were flanked by two DRs oriented in the same direction within the MRR and that the two DRs both belonged to Tn558 (Figure 3). The presence of the two DRs was further identified by PCR assays followed by sequencing of the amplicons (primers shown in Table 2). Both DRs were 1,326 bp in size, except for 18-bp exchanges in DRB compared to DRA. DRB contained partial fexA (430 bp) and orf38 sequences, while DRB comprised partial orf1 (430 bp) and orf2 sequences. Further analysis revealed that the single-nucleotide exchange TAG (orf138) → TAC (orf2) caused the termination codon to mutate to a Tyr codon, resulting in an extension of the open reading frame that transformed orf138 to orf2.

To further determine whether these unknown DRs in 245-1 and 2794-1 could mediate the formation of circular intermediate TUs, inverse PCR (P3, P4) was performed, followed by sequencing of the amplicons. Two identical PCR products

1https://github.com/tsseemann/snippy
2https://cge.cbs.dtu.dk/services/ResFinder/
3http://blast.ncbi.nlm.nih.gov/Blast.cgi
TABLE 1 | Phenotypic and genotypic characteristics of *Staphylococcus delphini*.

| Bacterial isolate | MIC (µg/mL) | Resistance genes |
|-------------------|-------------|------------------|
|                   | FFC  | CHL  | CLI  | TIA  | Lzd  | K    | Ery  | Fox  | Van  | Rfp  | Dap  |
| 245-1             | 256  | 128  | 4    | 128  | 8    | 16   | 0.25 | 0.5  | <0.015 | 0.25 | cfr, fexA, ble, ascA-aphD, aadD |
| 2794-1            | 256  | 128  | 4    | 128  | 8    | 16   | 0.25 | 0.5  | <0.015 | 0.5  | cfr, fexA, ble, ascA-aphD, aadD |
| 29213             | 8    | 8    | 0.0625 | 0.25 | 4    | 1    | 0.125 | 4    | 0.5  | 0.0078 | 0.5  | NONE |

FFC, florfenicol; CHL, chloramphenicol; CLI, clindamycin; TIA, tiamulin; Lzd, linezolid; K, kanamycin; Ery, erythromycin; FOX, cefoxitin; VAN, vancomycin; Rfp, rifampicin; Dap, daptomycin.

FIGURE 1 | Circular representation of the cfr-positive *Staphylococcus delphini* 245-1 and 2794-1 genomes. From the outer to the inner circles in the chromosome circular map: slot 1 (ARGs) and slots 2–9 (slot 2, genome size; slot 3, forward strand gene, colored according to the cluster of orthologous groups classification; slot 4, reverse strand gene, colored according to the cluster of orthologous groups classification; slot 5, forward strand ncRNA; slot 6, reverse strand ncRNA; slot 7, repeat; slot 8, GC content; and slot 9, GC skew).

(1,824 bp) were acquired from 245-1 and 2794-1, including a copy of DR₈, orf138, and part of fexA, as determined by sequencing (Figure 3). The TUs (13,613 bp) resulted from the recombination between DR₈ and DR₉, including multi-ARGs and one copy of DR₈. The PCRs (P1, P2) containing one copy of DR₈ detected the remaining structures after the excision of unconventional circularizable structures (UCSs) on chromosomes, and the results were consistent with the inverse PCR results (Figure 3). Importantly, the remaining structures were Tn558. These results confirmed the excision and cyclization of the structure (Figure 3). Further BLAST analysis revealed that the left ΔfexA-UCS exhibited 99.88% nucleotide identity to the corresponding region of the plasmids pWo28-1 (KX982171.1) and pWo28-3 (KY601170.1) from *S. sciuri* and plasmid pJP2 (KC989517.1) from *S. rostri* lacking DR₈ (Figure 3).

The sequence alignment analysis showed that the cfr MRR consisted of a Tn558 homologous region (6,644 bp) and a 13,613-bp region (Figure 3). This arrangement is a novel derivative of the Tn558 transposon. Compared to the fexA, orf138, tnpC, tnpB, and tnpA genes in Tn558 (Kehrenberg and Schwarz, 2005), a closer inspection of the Tn558 derivative showed that several nucleotide exchanges were identified in fexA (14 bp), tnpB (20 bp), and tnpA (14 bp), except for orf138. To further explain the genetic environment of the cfr MRR in this study, the plasmids pWo28-1 (KX982171.1) and pWo28-3 (KY601170.1) from *S. sciuri* and plasmid pJP2 (KC989517.1) from *S. rostri* are also shown in Figure 3. Analysis of the regions flanking the Tn558 derivative insertion in the chromosome identified a reading frame encoding a putative protein of 114 aa (62,844–62,846 and 83,105–83,443 nt on chromosomes 245-1 and 2794-1) that shared 98.54% nucleotide identity with a 148-aa DNA repair protein from *Macrococcus canis* (CP021059.1) (Gobeli et al., 2017). Additionally, a minicircle of Tn558, an indication of Tn558 having transposition activity, was identified via PCR (primers shown in Table 2) and sequencing of the derivative.

DISCUSSION

Naturally, *S. delphini* is widely susceptible to clinically relevant classes of antibiotics. In a previous study from Denmark,
among 55 *S. delphini* isolates recovered from mink, only some isolates were resistant to tetracycline (51%), penicillin (47%), and erythromycin (20%), whereas all the isolates tested susceptible to a vast majority of the antimicrobials assayed, including cefoxitin (Nikolaisen et al., 2017). In 2019, Magleby et al. also reported the first human case of *S. delphini* infection and found that...
the isolated exhibited low MIC values for all the antimicrobials assayed, including oxacillin (Mageley et al., 2019). Remarkably, the multiresistance gene *cfr* was shown to encode Cfr, an RNA methyltransferase that affects the binding of at least five chemically unrelated antimicrobial classes, namely, phenicols, lincosamides, oxazolidinones, pleuromutilins, and streptogramin A antibiotics, ultimately leading to a multidrug resistance phenotype (Long et al., 2006). Thus, the emergence and the global spread of the multiresistance gene *cfr* reduce the efficacy of a number of antibiotics in the control of Gram-positive bacteria. In this study, we identified the *cfr* gene in two food-related *S. delphini* strains. To the best of our knowledge, this study is the first report of the *cfr* gene existing in *S. delphini*. Furthermore, the *cfr* gene was located in an MRR with a number of antibiotic genes (*fexA*, *aacA-aphD*, *aadD*, and *ble*). The coexistence of *cfr* and other ARGs limits the choice of antibiotic therapy and may lead to the co-selection of these genes even without direct selection pressure, thereby increasing the retention and dissemination of these ARGs in *Staphylococcus*.

In this study, MRRs, including *cfr* and other ARGs, were confirmed as novel derivatives of the *Tn558* transposon. *Tn558* is a 6.6-kb bacterial transposon. It was first identified on the plasmid pSCFS2 harboring ARG *fexA* from *S. lentus*, and then numerous derivatives harboring numerous ARGs were found (Kehrenberg and Schwarz, 2005; Kehrenberg et al., 2007; Li et al., 2018). With a few exceptions, *cfr* is often harbored in the *Tn558* transposon as coexisting with other ARGs, such as *fexA*, *meca*, *erm(A/B/C)*, *tet(K/L/M)*, and *drf(K/G)* in the plasmids pSCFS3, pSCFS6, and pSCFS7 in previous studies (Witte and Cuny, 2011), but in this study, the derivative of the *Tn558* transposon harbored *cfr*, *fexA*, *aacA-aphD*, *aadD*, and *ble* on the chromosomes. In addition, the additional DRs within the *Tn558* derivative further confirm the particularity of this transposon. As previously reported for *Tn558* derivatives, there are no inverted repeats at the ends and no duplication of the target sequence at the integration site of the *Tn558* derivative. The typical 6-bp core sequences 5′-GATGTA-3′ at the left-end junction and 5′-GATCCA-3′ at the right-end junction were replaced by 5′-CATCCT-3′ and 5′- TAAAGCT-3′ in the novel derivative. The disappearance of target duplication and the alteration of the typical core sequences may have occurred during the transposition process (Diaz-Arco et al., 1987; Murphy, 1990). Moreover, the reading frame, including the insertion site of the *Tn558* derivative, is similar to the protein containing the *Tn558* site, and the excision of TUs in this *Tn558* derivative could lead to the formation of *Tn558*, indicating that the *DA* and *DB* in this study may be involved in the evolution of *Tn558* and that this derivative may be the ancestor of *Tn558* (Kehrenberg and Schwarz, 2005). Although multiple conjugation assays failed, the presence of a circular *Tn558* structure is indicative of the functional activity, suggesting that this novel *Tn558* derivative is a transposable element and may mediate the transfer of the *cfr* gene in the process of transposition (Kehrenberg and Schwarz, 2005).

Generally, the *cfr* gene often coexists with other ARGs on transposons or plasmids and is often in close proximity to insertion sequences (ISs), such as IS21-558, IS256, or ISEnu4, which play a crucial role in the mobility of *cfr* (Witte and Cuny, 2011; Wendlandt et al., 2015). These mobile structures have been detected among several Gram-positive bacteria, such as *staphylococci, Enterococcus faecalis, Macroccocus caseolyticus, Jeotgalicoccus pinnipedialis, Bacillus spp.*, and *Streptococcus suis*, as well as in Gram-negative bacteria, such as *E. coli* and *Proteus vulgaris* (Shen et al., 2013). However, mobile structures can form UCSs (Palmieri et al., 2013). UCSs lack recombinase genes.

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**TABLE 2 | Primers used for detecting antibiotics resistance genes, the circular forms and the structures not included in the corresponding region of the unconventional circularizable structures.**

| Primer | Sequence (5′ to 3′) | Product size (bp) | Annealing temperature (°C) | Purpose |
|--------|---------------------|-------------------|---------------------------|---------|
| Tn-F   | CGGTGCGTATACATCGGTGTC | 872               | 55                        | Detection of minicircle form of *Tn558* |
| Tn-R   | CGTTAACCGGCTTCTATGAC | 1824              | 65                        | Detection of the formation of translocatable units (TUs) between DR*DA* and DR*DB* |
| P1     | GAAAAACGTTGGCAGGTA   | 1817              | 65                        | Detection of remaining structures after the excision of UCSs on chromosomes |
| P2     | CTTCACTTCTCAAAGGCTCTGT | 746               | 58                        | Detection of of UCSs on chromosomes |
| P3     | GGCAQAATCGTGAAGCA    | 1880              | 55                        | Detection of DR*DA* |
| P4     | ACCATGATAGACAGGCTATAT | 1683              | 55                        | Detection of DR*DB* |
| A-F    | TGGTCCGATTGCTAGTT    | 1680              | 55                        | Detection of remaining structures after the excision of UCSs on chromosomes |
| A-R    | AAAAACTCATCCTGCAAGCTT | 1824              | 65                        | Detection of the formation of translocatable units (TUs) between DR*DA* and DR*DB* |
| B-F    | TGCGTGGATCGAAGACCG   | 746               | 58                        | Detection of of UCSs on chromosomes |
| B-R    | CCCTCGTTCAAGGAGCAGTTT | 1824              | 65                        | Detection of the formation of translocatable units (TUs) between DR*DA* and DR*DB* |
| cfr-F  | TGAAGTATAAACCAGGTTGGGAGCTCA | 746               | 58                        | Detection of *cfr* |
| cfr-R  | ACCATATACATCGACACACAGC | 1817              | 65                        | Detection of the formation of translocatable units (TUs) between DR*DA* and DR*DB* |

*Primer Tn-F/Tn-R were used for detecting the formation of minicircle of *Tn558* derivative; primers P1/P2 were used for detecting the formation of translocatable units (TUs) between DR*DA* and DR*DB*; primers P3/P4 were used for detecting the remaining structures after the excision of UCSs on chromosomes; primers A-F/A-R, B-F/B-R were used for detecting DR*DA* and DR*DB*; primers cfr-F/cfr-R were used for detecting resistance gene *cfr*. |
and can be excised in circular form due to the extensive DRs flanking the DNA segment undergoing excision (Locke et al., 2012; Palmieri et al., 2012, 2013). Thus, they are very important for the horizontal transmission of ARGs. In this study, the ARGs aacA-aphD, aadD, ble, and cfr, bracketed by DRs, formed a novel genetically mobile structure. The particular genetic structures identified by the analysis were referred to as UCSs. Two-step PCR results indicated that this structure can be looped out and excised from the chromosome, leading to the formation of Tn558 (Figure 3), which suggests that the DR is active and involved in the mobility of the Tn558-carried cfr gene in this study. Further BLAST analysis revealed that the left AβexA-UCS exhibited 99.88% nucleotide identity to the corresponding region of the plasmids pWo28-1 (KX982171.1) and pWo28-3 (KY601170.1) from S. sciuri and plasmid pJP2 (KC989517.1) from S. rostri lacking DR (Figure 3). Therefore, the DR and DR in this study, similar to ISs, might facilitate the dissemination and accumulation of ARGs in Tn558 (Palmieri et al., 2013; Harmer et al., 2014). Of course, the functions of these two unknown DRs still need to be further studied and explored in the future.

Unconventional circularizable structures are widely distributed in Gram-negative and Gram-positive bacteria and play an important role in the dissemination of ARGs (Palmieri et al., 2013; Chanchaithong et al., 2019). The DRs in UCSs are usually long and are more than 100 times longer than the att sites functioning in traditional MGEs (Frost et al., 2005). The DRs may contain genes, such as erm(B), mef, (macrolide efflux), and ofr138 in this study, but they are not involved in transposition (Locke et al., 2012; Hao et al., 2019). The exact mechanism of mobilization has not been determined, although hypotheses have been proposed (Azpiroz et al., 2011). This transfer mechanism may be similar to that of IS26 via site-specific recombination, including a multistep process that requires the formation of a TU, precise excision of the TU, and integration targeting the preexisting DR (Harmer et al., 2014; Harmer and Hall, 2015). The endogenous instability of UCSs endows the encompassed niche adaptation determinants with the ability to be transferred. Moreover, they are often carried by MGEs, which prompts the updating of MGEs (such as the derivative of Tn558) and further accelerates the spread of UCSs. Furthermore, the presence of DRs on this novel cfr-carrying Tn558 derivative may accelerate the spread and persistence of ARGs among staphylococci and exacerbate the threat of superbugs, such as methicillin-resistant S. aureus. The proliferation of the transferable ARG cfr kidnapped by transposons or other MGEs has impaired the efficiency of oxazolidinones in clinical settings and threatens public health (Li et al., 2018).

CONCLUSION

To the best of our knowledge, this study is the first report of S. delphini harboring a novel cfr-carrying Tn558 derivative. The constant occurrence of the cfr gene in new staphylococcal host species underlines its strong transmissibility and wide distribution. This finding raises further concerns regarding the potential threat to food safety and public health safety. The occurrence and the dissemination of similar cfr-carrying transposons from diverse Staphylococcus species need further surveillance.

DATA AVAILABILITY STATEMENT

The complete genomic sequences of 245-1 and 2794-1 have been deposited in GenBank: 245-1 (GenBank ID: CP063368) and 2794-1 (GenBank ID: CP063367).

AUTHOR CONTRIBUTIONS

QW, JZ, SW, and TL conceived and designed the experiments. FZ, JH, and JD performed the experiments. FZ, SW, and RY analyzed the data. YD, LX, MC, and JW contributed reagents, materials, and analysis tools. FZ, SW, and JW contributed to the writing of the manuscript. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2020.598990/full#supplementary-material

Supplementary Figure 1 | Sample collection locations for this study in China.

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FZ, SW, and RY performed the experiments. FZ, JH, and JD performed the experiments. FZ, SW, and RY contributed reagents, materials, and analysis tools. FZ, SW, and JW contributed to the writing of the manuscript. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2020.598990/full#supplementary-material

Supplementary Figure 1 | Sample collection locations for this study in China.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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