# Mosaic tetracycline resistance genes encoding ribosomal protection proteins

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First reported in 2003, mosaic tetracycline resistance genes are a subgroup of the genes encoding ribosomal protection proteins (RPPs). They are formed when two or more RPP-encoding genes recombine resulting in a functional chimera. To date, the majority of mosaic genes are derived from sections of three RPP genes, tet(O), tet(W) and tet(32), with others comprising tet(M) and tet(S). In this first review of mosaic genes, we report on their structure, diversity and prevalence, and suggest that these genes may be responsible for an underreported contribution to tetracycline resistance in bacteria.

## Introduction

Tetracyclines bind to the A-site on the bacterial ribosome, resulting in steric blocking of the aminoacyl-tRNA binding site, which prevents protein synthesis. They are effective against both Gram-positive and Gram-negative bacteria and, due to the relative lack of major side effects and cheap cost, have been used extensively in the treatment of infections as well as growth promoters in animal husbandry.

Bacterial resistance to tetracycline is often mediated through the acquisition of DNA encoding proteins that confer resistance by one of three main mechanisms: ATP-dependent efflux, enzymatic inactivation of tetracycline, or ribosomal protection.<sup>2</sup> To date, a total of 60 different classes of tetracycline resistance gene, including oxytetracycline resistance genes, have been reported. These include 33 predicted or proven to encode active efflux pumps, 12 encoding ribosomal protection proteins (RPPs), 13 encoding inactivating enzymes and 1 reported to confer resistance via an as yet undetermined mechanism, designated tet(U) (a full list is periodically updated by Roberts<sup>4</sup>). Although it has vet to be assigned a mechanistic class, tet(U) has been identified in Enterococcus and Staphylococcus isolates. 5,6 However, a study by Caryl et al. reported that when tet(U) was cloned and expressed in Escherichia coli, the transformants were not resistant to tetracycline.

To be considered a new class of tetracycline resistance gene, it must encode a protein <80% identical to known tetracycline resistance proteins. Determinants representing new classes were originally assigned a letter from the English alphabet. However, as all letters are used, they are now assigned an

Arabic numeral,<sup>8</sup> with new determinants referred to the Levy group (bonnie.marshall@tufts.edu) in order to obtain a designation prior to publication to avoid duplication and ensure taxonomic consistency.

## **RPPs**

RPPs are a related group of proteins that, when bound to the ribosome, result in the release of tetracycline from the ribosome, enabling protein synthesis to proceed<sup>10</sup> (reviewed by Thaker et al.<sup>11</sup>). Of the 12 classes of RPP gene currently reported [tet(M), (O), (Q), (S), (T), (W), (32), (36), (44), B(P), otr(A) and tet], tet(M) is considered the most prevalent due to its association with the broad host range Tn916/Tn1545 family of conjugative transposons.<sup>12</sup> However, a subgroup of RPP genes has been identified that consist of regions of different, already characterized RPP genes that appear to have undergone recombination forming a mosaic gene. It must be stressed here that the progenitors of mosaic genes are assumed based purely on the order in which they were discovered and we cannot be sure of the directionality of mosaic gene formation.

# Mosaic RPP genes

In 2003, Stanton and Humphery<sup>13</sup> reported two RPP genes in *Megasphaera elsdenii* that encoded predicted proteins showing 89.1% and 91.9% identity to Tet(W) (accession number AJ222769) from *Butyrivibrio fibrisolvens*. As this was above the <80% cut-off, they did not qualify as a new resistance class

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under the nomenclature system. However, further analysis of the amino acid sequence revealed variability in the percentage identity to Tet(W) across its length. The large central section in both sequences showed 98.1% identity to Tet(W), while small sections at the N- and C-terminal ends were found to have a lower amino acid sequence identity to Tet(W) [between 66.6% and 75.3%]. However, these same N- and C-terminal sections were shown to have between 99.3% and 100% amino acid identity to Tet(O) (accession number M18896), despite the central section showing identity to Tet(W). Given the evidence, this suggested recombination had occurred, creating a mosaic determinant with a central Tet(W) region flanked by two Tet(O) regions. Although never before observed between two different RPP classes, recombination resulting in functional genes has previously been reported between different phylotypes of  $tet(M)^{14}$  as well as in other antibiotic resistance genes, such as penA and pbp2x, which confer resistance to penicillin. 15,16 Furthermore, in vitro experiments have successfully recombined tet(A) and tet(C) to create mosaics that confer resistance to tetracycline at levels comparable to the nonmosaic tet(C).

The guideline for determining a new resistance gene class was established prior to the discovery of these mosaic RPP genes and none of the mosaic genes qualified as a new class when analysed as one single continuous sequence. It was clear, however, that these mosaic genes were different from their non-mosaic counterparts and that the current classification did not adequately reflect the true evolutionary background of these genes. Therefore, an expansion of the nomenclature system was suggested whereby the mosaic gene would receive a designation that reflected the structural order and class of the genes they comprised, better reflecting their variable nature. <sup>18,19</sup> For example, the two resistance genes reported in *M. elsdenii*, which comprised a central *tet*(W) region flanked by two *tet*(O) regions, were designated *tet*(O/W/O). <sup>13</sup>

Although Stanton and Humphrey<sup>13</sup> were the first to report mosaic RPP genes, Melville et al. 20 had unknowingly reported a mosaic gene 2 years previously. This resistance gene, found in Clostridium saccharolyticum K10, encoded a predicted protein that showed 76% amino acid identity to Tet(O) (accession number Y07780). As per the original nomenclature guidelines, it was given the new designation Tet(32). However, subsequent re-examination of the sequence found that only the central section showed <80% identity to known proteins, while the Nand C-terminal regions flanking the central section shared 100% and 97.7% identity, respectively, to Tet(O) (accession number M18896). The central region was still thought to represent a section of a new Tet(32) class and therefore the determinant was reclassified Tet(0/32/0). 18 Subsequently, the proposed full, non-mosaic sequences of Tet(32) have been reported in several isolates identified from the human oral cavity, 21,22 with the Tet(0/32/0) mosaic determinant now showing 89% amino acid identity to these.

Similarly, the previously reported tet(S) allele (accession number AY534326) on the conjugative transposon Tn916S<sup>23</sup> has subsequently been reclassified as a result of *in silico* analysis. The amino acid sequence shows identity to Tet(S) across 595 amino acids (1–595 inclusive), with the final 61 amino acids at the C-terminus end identical to Tet(M) (accession number U09422), resulting in it being reclassified as Tet(S/M).<sup>24</sup>

## Mosaic gene diversity

To date, a total of 30 mosaic genes have been reported in the literature, of which 26 currently have sequences deposited in GenBank (Table 1). Some studies have reported multiple occurrences of known genes; however, many of these have been characterized by PCR amplification only. Structurally, these chimeric genes currently comprise either two [e.g. tet(O/W)], three [e.g. tet(O/W/O)], four [e.g. tet(O/W/32/O)] or six [e.g. tet(O/W/32/O/ W/O)1 different regions (Figure 1), with tet(0), tet(W) and tet(32)being the predominant RPP genes reported to form mosaic genes, comprising all but two of the reported variants, and tet(M) and tet(S) forming the remaining two. 24,25 Given the prevalence of tet(M) in certain samples, and the previous reports of selfrecombination, <sup>14,26</sup> it is surprising that there are so few reports of mosaic genes containing tet(M). Furthermore, alignment of 12 representative RPP gene sequences shows tet(M) sharing 75% and 70% identity, respectively, to tet(0) and tet(44), which is higher than the percentage identity observed between the more commonly reported RPP mosaic genes comprising tet(0), (W) and (32) (Table 2). However, mosaic genes comprising tet(M) and any other gene, with the exception of tet(S), have yet to be reported. It is entirely possible that this may be due to a lack of investigation rather than an absence of recombination followed by fixation of the recombinant allele in the bacterial population. Alternatively, it is possible that there is little selective pressure for tet(M)-based mosaic genes if the resultant protein is no more efficient than Tet(M) itself and/or there is no indirect selective pressure for mosaicism. A similar situation may exist for other proteins, such as Tet(S). Stanton et al.<sup>27</sup> reported that the protein encoded by the tet(O/W/O) mosaic genes in M. elsdenii conferred a higher level of resistance to tetracycline than their non-mosaic counterparts, but similar investigations are still to be reported for other RPP genes. Therefore, the prevalence of certain mosaic gene variants could suggest that they are in some way more beneficial to the host than the non-mosaic genes they comprise.

## **PCR-based** analysis

PCR-based assays have been developed to help researchers detect specific mosaic genes. Stanton and Humphrey  $^{13}$  describe an assay that distinguished between the non-mosaic genes tet(0) and tet(W) and the mosaic tet(0/W/0) from Megasphaera strains, enabling them to detect tet(0/W/0) variants in six additional M. elsdenii strains. Patterson  $etal.^{21}$  investigated the presence of mosaic genes using various specific oligonucleotide sets that either bound within the resistance genes or flanked them. Amplicons specific to tet(0/W), tet(0/32) and tet(W/32) were detected in faecal samples, with tet(0/32) being the most common of these mosaic amplicons; it was amplified in all 12 pig faecal samples and 6 of 7 human faecal samples tested. In contrast, the faecal samples from cows and sheep, as well as human saliva samples, failed to produce any amplicons for these mosaic genes, suggesting they were not present at detectable levels.

Chen et al. <sup>28</sup> also used an oligonucleotide primer set that annealed outside tet(0) to determine the presence of tetracycline resistance genes in two *Streptococcus suis* isolates. Although no amplicon was produced using internal, tet(0)-specific primers, the primers binding to flanking DNA yielded an amplicon, indicating the presence of mosaic genes [identified as tet(0/32/0) and

**Table 1.** A summary of the mosaic tetracycline genes reported to date

Gene	Organism	Source(s)	Accession number	Reference(s)	
tet(O/W)	Bifidobacterium thermophilum B0219	environmental (pig slaughterhouse) sample	AM889118	32	
tet(O/W)	B. thermophilum B0241	pig faeces	AM889119	32	
tet(O/W)	B. thermophilum B0242	pig faeces	AM889120	32	
tet(O/W)	B. thermophilum B0253	pig faeces	AM889121	32	
tet(O/W)	B. thermophilum B0256	pig faeces	AM889122	32	
tet(O/W)-2	Megasphaera elsdenii 25-51	swine faeces	AY485122	18,27	
$tet(O/W)-1 [n=15^{a}]$	M. elsdenii 27-51	swine faeces	AY485126	27,33	
tet(O/W/O)-4	uncultured bacterial clone	pig faeces	no accession number	21	
tet(O/W/O)-3 [n=9]	uncultured bacterial clone	pig faeces	EF065524	21	
$tet(O/W/O)-2 [n=28^b]$	M. elsdenii 14-14	swine caecum	AY196920	13,18,27,33	
tet(O/W/O)-1 [n=2]	M. elsdenii 7-11	swine caecum	AY196921	13,18,27	
tet(0/W/32/0) [n=32]	uncultured bacterial clone	pig faeces	EF065523	21	
$tet(O/W/32/O) [n=7^{c}]$	Streptococcus suis Ss1303	pig (brain, lung and spleen) and human (CSF) samples	FM164392	34	
tet(O/W/32/O)	S. suis 32457	diseased pig lung	FR823304	34,35	
tet(O/W/32/O)	Streptococcus gallolyticus subsp. gallolyticus ATCC 2069 plasmid pSGG1	human blood	FR824044	36	
tet(O/W/32/O)	S. suis	diseased pig (blood, brain, heart, joint and lung) samples	JQ740053	28	
tet(O/W/32/O/W/O)	Lactobacillus johnsonii G41	human faeces	DQ525023	32	
tet(O/W/32/O/W/O)	uncultured bacterial clone	pig faeces	DQ679926	21	
tet(0/32/0)	S. suis	diseased pig (blood, brain, heart, joint and lung) samples	JQ740052	28	
tet(0/32/0)	Clostridium saccharolyticum K10	human colon	AJ295238	18	
tet(0/32/0)-2 [n=3]	uncultured bacterial clone	human and animal faecal samples	no accession number	21	
tet(O/32/O)-3	uncultured bacterial clone	human and animal faecal samples	no accession number	21	
tet(0/32/0)-4	uncultured bacterial clone	human and animal faecal samples	no accession number	21	
tet(0/32/0)-5	uncultured bacterial clone	human and animal faecal samples	no accession number	21	
tet(O/32/O)	Dorea longicatena AGR2136	rumen microbiome	NZ_AUJS01000017 (41626-43545 bp)	direct submission, analysed in this study	
tet(O/32/O)	Campylobacter coli 202/04	human faeces	AINH01000038 (2361-4280 bp)	direct submission, analysed in this study	
tet(O/32/O)	C. coli 317/04	human faeces	NZ_AINJ01000054 (2094-4013 bp)	direct submission, analysed in this study	
tet(O/32/O)	Campylobacter jejuni subspecies jejuni 2008-894	human	AIOQ01000025 (14515-16434 bp)	direct submission, analysed in this study	
tet(O/32/O)	Roseburia intestinalis XB6B4	human intestinal tract	FP929050 (2873814-2875733 bp)	direct submission, analysed in this study	
tet(S/M)	Streptococcus equinus 1357	food	HM367711	25	
tet(S/M)	Streptococcus intermedius	human isolate	AY534326	23,24	

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Gene	Organism	Source(s)	Accession number	Reference(s)
tet(W/32/O)	B. thermophilum B0219	environmental (pig slaughterhouse) AM710601 sample	AM710601	32
tet(W/32/0)	B. thermophilum B0241	pig faeces	AM710602	32
tet(W/32/0)	B. thermophilum B0242	pig faeces	AM710603	32
tet(W/32/0)	B. thermophilum B0253	pig faeces	AM710604	32
tet(W/32/0)	B. thermophilum B0256	pig faeces	AM710605	32

Fourteen of the 15 tet(O/W)-1 variants were only determined by PCR analysis and so could be either tet(O/W)-1 or tet(O/W)-2. PEleven of the 28 tet(O/W/O)-2 variants were only determined by PCR analysis and so could be either tet(O/W/O)-2 or tet(O/W/O)-1. he number given in square brackets indicates the instances of that mosaic gene variant reported, if more than one. All S. suis isolates, but not the same strain. tet(O/W/32/O)]. This full-length oligonucleotide primer set does aid the identification of mosaic genes; however, it is only specific for those with regions homologous to tet(O) flanking sequences. Since PCR strategies aimed at identifying resistance genes require knowledge of the sequence of the target, mosaic RPP genes are likely to be largely undetected and under-reported by PCR-based studies.

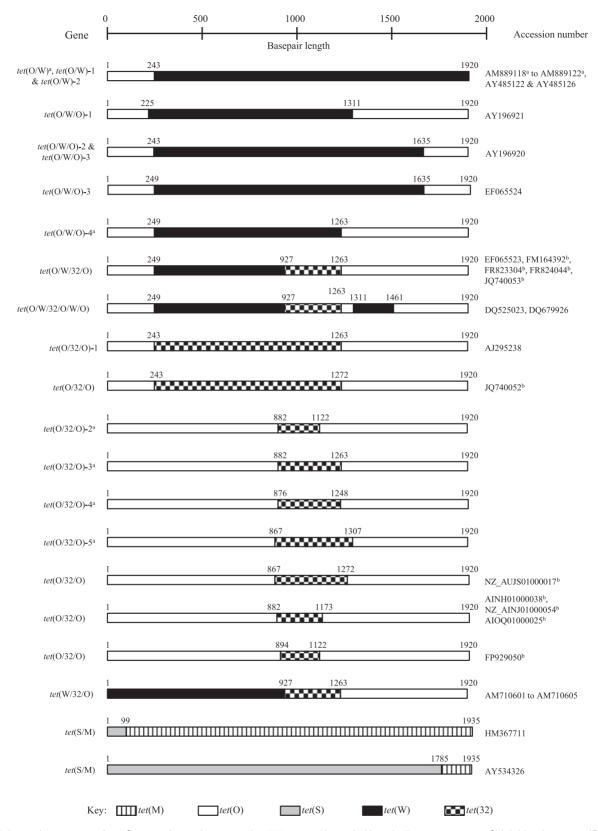
Reflecting the findings by Patterson et al., 21 almost all the mosaic genes reported to date have originated from faecal samples, with the majority identified from a porcine origin and less commonly from humans (Table 1). The gut houses a complex and diverse bacterial community with potential for widespread horizontal gene transfer, and the mosaic genes found in faecal samples are likely to reflect the pool of non-mosaic genes present within the gut microbiota. Genes such as tet(W) and tet(O) are commonly reported from these types of samples, <sup>29</sup> but the prevalence of tet(32)-containing mosaic genes suggests that tet(32) may be more common than initially thought. In fact, tet(0/32/0) was found to be the most common mosaic gene in both the human and pig faecal samples tested and was present in almost as many samples tested as the non-mosaic tet(0) and tet(W) genes.<sup>21</sup> In contrast, mosaic genes have not vet been reported in faecal samples from bovine and ovine origin or in human saliva.<sup>21</sup> Why they are predominantly found in pigs while as yet unreported in other animals is not immediately clear, though the extensive use of tetracyclines in the swine industry<sup>3,30,31</sup> may have contributed to their selection.

# Draft genome analysis

The advent of high-throughput genomic sequencing has led to an increase in the number of genomes being deposited in sequence databases. Many contain tetracycline resistance genes that are generically labelled simply as 'tetracycline resistance protein' or as 'tet(M)-like', the designation of which may be a result of automated annotation pipelines. A preliminary search of the NCBI nucleotide database, using tet(0) (accession number Y07780) as the guery, found that some of these generically labelled tetracycline resistance genes gave a partial match to tet(0). Further examination indicates that some are as yet uncharacterized and unreported mosaic genes, which have been further defined for this review using the nucleotide sequence to determine the crossover points. For example, the tet(M)-like gene (accession number NZ\_AUJS01000017, location  $41626-43\overline{5}45$  bp) in the draft genome of Dorea longicatena AGR2136 from a human faecal sample appears to be a previously unreported variant of tet(O/32/O)(Figure 1).

Furthermore, the tetracycline resistance genes present in *Campylobacter jejuni* subspecies *jejuni* 2008-894, *Campylobacter coli* 202/04, *C. coli* 317/04 (accession numbers AIOQ01000025, AINH01000038 and NZ\_AINJ01000054, respectively) and *Roseburia intestinalis* XB6B4 (accession number FP929050) are also structurally novel variants of *tet*(O/32/O) (Figure 1). The three mosaic genes present in the *Campylobacter* spp. are identical to each other, while that in *R. intestinalis* is different. Taking into account these newly defined genes, the total number of mosaic genes reported increases from 30 to 35 (not including those identified via PCR amplification only; Table 1) and suggests that other generically labelled tetracycline resistance genes

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**Figure 1.** Schematic representation of reported mosaic tetracycline RPP genes. The coded bars indicate sequences of high identity to specific RPP genes: vertical line bars for tet(M), white bars for tet(M), grey bars for tet(S), black bars for tet(W) and checked bars for tet(32). The number above the bar indicates the reported crossover point. <sup>a</sup>Indicates those sequences that are incomplete or absent in GenBank, with the crossover points taken from the publication. <sup>b</sup>Indicates sequences that have been analysed in this study due to no specific crossover point(s) reported.

**Table 2.** Sequence identity matrix showing the percentage nucleotide identity between representatives of all 12 RPP gene classes, in descending order, compared with tet(M)

RPP gene	tet(M)	tet(S)	tet(O)	tet(44)	tet(32)	tet(W)	tet(T)	tet(36)	tet(Q)	tetB(P)	otr(A)	tet
tet(M)	100	78	75	70	69	64	57	49	46	23	11	11
tet(S)		100	70	69	67	62	56	56	48	11	11	10
tet(O)			100	69	69	65	56	49	48	15	12	11
tet(44)				100	71	64	50	58	46	15	11	10
tet(32)					100	67	55	49	47	11	12	10
tet(W)						100	12	45	15	5	14	12
tet(T)							100	57	56	18	8	2
tet(36)								100	64	9	11	11
tet(Q)									100	13	12	12
tetB(P)										100	1	1
otr(A)											100	63
tet												100

Accession numbers of representative genes included in the matrix: tet(M), U09422; tet(O), Y07780; tetB(P), AE001437; tet(Q), X58717; tet(S), X92946; tet(T), L42544; tet(W), AJ222769; tet(32), DQ647324; tet(36), AJ514254; tet(44), FN594949; otr(A), X53401; tet, AL939106. Shaded boxes represent those genes currently reported to comprise mosaic genes.

present in the database [e.g. those labelled as tet(M)-like] could be further classified, helping to understand mosaic gene proliferation and diversity.

#### **Conclusions**

Our knowledge of the mosaic RPP gene group is steadily increasing since their discovery in 2003, with the majority derived from tet(0), tet(W) and tet(32) and others deriving from tet(M) and tet(S). It is clear that these genes are being under-reported both in terms of experimental detection and also within genomic data. Further work and increased attention on mosaic RPP genes is important if we are to understand the evolutionary selective pressures driving their fixation in bacterial populations and the subsequent effects on resistance and mobile genetic element evolution within their host.

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## **Transparency declarations**

None to declare.

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