Actinobacillus suis isolated from diseased pigs are phylogenetically related but harbour different number of toxin gene copies in their genomes

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Abstract
Objective: The Gram-negative bacterium Actinobacillus suis is an agent of global importance to the swine industry and the cause of lethal respiratory or septicaemic disease in pigs of different ages. Between 2018 and 2019, seven commercial farms in western Canada experienced episodes of increased mortality due to A. suis infection in grower pigs. The goal of this work was to profile, with molecular methods, A. suis isolated from diseased pigs and to compare them to other isolates.

Design: This inferential observational study used nine western Canadian strains obtained from diseased lungs (n = 6), heart (n = 2) and brain (n = 1) and whole genome sequencing was performed. Comparative genomic analyses were performed to characterise the genetic variability, antimicrobial resistance and the virulence genes present.

Results: Compared to the reference strain (ATCC 33415), an increased number of RTX (repeats in the structural toxin) gene copies were identified in strains isolated from organs without a mucosal surface, thus theoretically harder to invade. Western Canadian strains did not harbour genes associated with resistance to antimicrobial agents used in swine production. Novel regions were also identified in the genomes of five of nine strains demonstrating recombination and emergence of novel strains.

Conclusions: The results obtained in this study were associated with the emergence of new lineages. An increased number of RTX toxin gene copies is suggested to be associated with increased virulence. This study will contribute to improve our understanding regarding A. suis and may help guide vaccine development and agent control measures.

INTRODUCTION

Actinobacillus suis is a facultative anaerobic, non-motile, Gram-negative bacillus. It is an important pathogen of swine, leading to respiratory disease or septicaemia in pigs of different ages and stages of production. The organism was first identified and described in 1955, when it was associated with acute septicaemia and sudden death in domestic swine. Actinobacillus suis colonises the nasopharynx and palatine tonsils of healthy pigs; it is a commensal. It can also colonise the vaginal mucosa of healthy sows. Disease is often associated with stressful conditions such as weaning, farrowing, transportation or suckling. Weaned pigs often do not develop clinical signs and are found dead following fulminant septicaemia, while grower-finisher or older pigs in high-health status (HHS) herds develop respiratory signs due to necro-haemorrhagic pleuropneumonia. A third syndrome observed in HHS reproductive-aged pigs is characterised by acute septicaemia, lethargy, anorexia, fever, abortion and erysipelas-like lesions on the skin. Treatment strategies include parenteral antimicrobial therapy with cefetiofur, penicillin, ampicillin, neomycin, gentamicin, sulphadimethoxine, tiamulin, tetracycline or tylosin. Variation in sensitivity to antimicrobial therapy has been reported and antimicrobial susceptibility testing is suggested when clinical cases are identified. Autogenous bacterins have been used to prevent disease and shown variable results. These evidence a gap in our understanding of A. suis pathogenesis, contributing to our inability in developing efficient prevention, treatment and control tools.
Actinobacillus suis can be readily isolated using blood and McConkey media, producing β-haemolytic smooth, translucent colonies. Genes belonging to different functional classes have been proposed to contribute to virulence and facilitate colonisation of mucosal sites. A previous study described 13 virulence genes in A. suis, identified using genome-wide screening by PCR-based signature-tagged transposon mutagenesis. Noteworthy is the RTX (repeats in the structural toxin) gene family. It encodes a pore-producing protein and is proposed as a disease determinant associated with A. suis. The RTX genes identified in A. suis is immunologically and genetically related to the RTX genes described in other pathogenic bacteria such as Escherichia coli, Pasteurella haemolytica and Actinobacillus pleuropneumoniae.

Sporadic outbreaks of A. suis were previously reported in Canada, the United States, UK, India, New Zealand, Hungary, Croatia and Australia. Since 1990, an increasing number of clinical disease outbreaks have been reported in pigs of varying ages in North America and considered as an emerging disease. Therefore, the purpose of this study was to investigate the molecular epidemiology, virulence and antimicrobial gene profile in A. suis isolated from diseased pigs in western Canada.

**MATERIAL AND METHODS**

**Bacterial strains**

Between 2017 and 2019, clinical samples (n = 9) were collected from pigs with signs suggestive of A. suis infection in seven commercial farms in western Canada. Isolates were obtained from lungs (n = 6), heart (n = 2) and brain (n = 1) (Table 1). Samples were streaked on 5% blood agar and incubated at 37°C under aerobic conditions and plates were examined at 24 and 48 h for growth. Matrix-assisted laser desorption/ionisation-time of flight was performed for initial identification; colonies identified as A. suis were selected and saved for further genomic analysis.

**Genome sequencing and assembly**

Genomic DNA from isolates was extracted using DNeasy ultraclean microbial kit (QIAGEN, Toronto, Ontario, Canada). Sequencing libraries for isolates 19_419, 19_418, 18_466, 19_298, 18_292, 4286 and 17_417 were prepared using the Nextra XT DNA library (Illumina, San Diego, CA, USA) and sequencing was performed using an Illumina MiSeq Platform (Illumina Inc., 500 cycles, 2 × 250) and MiSeq V3 reagent kits (Illumina Inc.). Isolates 20_377_1a and 20_277_1a had DNA libraries prepared using a Rapid Sequencing Kit (SQK-RAD0003, Oxford Nanopore Technologies, Oxford, UK). Libraries were sequenced on a MinION sequenced (Oxford Nanopore Technologies) using a Flow Cell model FLO-MIN 106 v R9.4 running the software MinKNOW v1.10 (Oxford Nanopore Technologies) with default parameters. Genomes were trimmed, assembled, annotated and analysed using the comprehensive genomic analysis tool in Pathosystem Resource Integration Center (PATRIC).

**Phylogenetic tree of Actinobacillosis suis and whole genome comparisons**

Three complete and four draft A. suis genomes isolated from pigs with clinical disease suggestive of A. suis infection were included in these analyses, as well as A. pleuropneumoniae strain NCTC 10976, which was included in the analysis as an outgroup. These were initially identified between 1963 and 1991 in North America; they were retrieved from National Center for Biotechnology Information (www.ncbi.nlm.nih.gov) (Table 2). Randomised accelerated maximum likelihood for high-performance computing (RAxML-VI-HPC) in PATRIC using 500 genes and allowing neither deletions nor duplications was used to build the phylogenetic tree. Orthologous average nucleotide identity was performed to measure the overall similarity between genome sequences using OrthoANI V0.93.1 available at EzBioCloud. Protein sequences of all western Canadian strains were compared to the reference genome ATCC 33415 using bidirectional BLASTP with parameters set to minimum coverage = 30%, minimum identity = 10% and BLAST E-value = 1e-5 and a circular genomic map was constructed. Progressive Mauve was used to reorder contigs and align query genomes to ATCC 33415 to identify homologous regions distribution.

| Strain ID | Source organ | Postmortem diagnosis | Collection location | Collection date |
|-----------|--------------|----------------------|---------------------|-----------------|
| 20_277_1a | Heart        | Septicaemia          | Farm 6              | June 2020       |
| 20_377_1a | Brain        | Septicaemia          | Farm 7              | April 2020      |
| 17_471    | Lung         | Bronchopneumonia     | Farm 1              | September 2017  |
| 4286      | Lung         | Bronchopneumonia     | Farm 2              | October 2018    |
| 19_419    | Lung         | Bronchopneumonia     | Farm 3              | September 2019  |
| 19_418    | Lung         | Bronchopneumonia     | Farm 3              | September 2019  |
| 19_298    | Heart        | Bronchopneumonia/septicaemia | Farm 4     | June 2019       |
| 18_466    | Lung         | Bronchointerstitial pneumonia | Farm 5    | October 2018    |
| 18_292    | Lung         | Bronchopneumonia     | Farm 6              | June 2019       |
TABLE 2  Publicly available *Actinobacillus suis* genomes used for comparison with western Canadian strains

| Strain ID | Serovar | Country, province of origin | Isolation source | Collection year | Reference |
|-----------|---------|-----------------------------|------------------|----------------|-----------|
| ATCC 33415 | O1:K1 | USA | Pig with septicemia | Not available | 19 |
| ATCC 15557 | O1:K1 | USA, Washington DC | Blood of irradiated adult pig | 1963 | 20 |
| NCTC 12996 | Not available | Not available | Pig with septicemia | 1980 | NCBI |
| AS108/13 | Not available | Brazil, Minas Gerais | Pig with pneumonia | 2013 | 21 |
| H89-1173 | O2:K3 | Canada, Ontario | Lung of pig with pneumonia | 1989 | 22 |
| H91-0380 | O2:K2 | Canada, Ontario | Pig with septicemia | 1991 | 23 |
| H91-0406 | O2:K3 | Canada, Ontario | Lung of pig with pneumonia | 1991 | 22 |

Abbreviation: NCBI, National Center for Biotechnology Information.

| Strain ID | Source | Genome accession | Contigs | GC content (%) | Sequence length (bp) | # Genes identified |
|-----------|--------|------------------|---------|----------------|---------------------|-------------------|
| 20_277_1a | This study | CP090556 | 2 | 40.26 | 2549182 | 2775 |
| 20_377_1a | This study | JAKEZJ0000000000 | 1 | 40.23 | 2551416 | 2772 |
| 17_471 | This study | SAMN24451496 | 37 | 40.08 | 2533834 | 2416 |
| 4286 | This study | JAKEZJ0000000000 | 29 | 40.16 | 2456581 | 2320 |
| 19_419 | This study | JAJU0Y0000000000 | 29 | 40.09 | 2534652 | 2414 |
| 19_418 | This study | SAMN24451500 | 30 | 40.09 | 2534921 | 2412 |
| 19_298 | This study | JAJU0Z0000000000 | 29 | 40.16 | 2453999 | 2399 |
| 18_466 | This study | SAMN24451498 | 30 | 40.14 | 2569370 | 2467 |
| 18_292 | This study | JAJU0PA0000000000 | 28 | 40.09 | 2536093 | 2416 |
| ATCC 33415 Public | | CP009159 | 1 | 40.22 | 2501598 | 2341 |
| ATCC 15557 Public | NZ_MTBV0000000000 | 80 | 40.22 | 2405979 | 2179 |
| NCTC 12996 | | NZ_LT906456.1 | 1 | 40.22 | 2501595 | 2338 |
| H91-0380 Public | | CP003875 | 2 | 40.24 | 2484940 | 2249 |
| AS108/13 | | QCXP01 | 44 | 40.17 | 2617798 | 2528 |
| H89-1173 Public | NZ_MTBW0000000000 | 74 | 40.23 | 2450899 | 2345 |
| H91-0460 Public | NZ_MTBX0000000000 | 116 | 40.35 | 2423500 | 2318 |

*BioProject ID PRJNA792721.*

Functional characterisation

Genomes were screened for the presence of virulence factors using BLASTP.10,11,24 Antimicrobial resistance genes (ARG) in all *A. suis* genomes were identified using the Resistance Gene Identifier platform and the Comprehensive Antimicrobial Resistance Database, only genes with greater than 80% identity were reported.15,25

RESULTS

Whole genome comparisons and phylogenetic analysis

A summary of the *A. suis* genomes used in this study is listed in Table 3. Average genomic length and number of genes identified were numerically higher in the seven genomes from this study (2.52 ± 0.4 Mbp and 2478 ± 173 genes) when compared to publicly available genomes (2.48 ± 0.6 Mbp and 2328 ± 107 genes). Average nucleotide identity (ANI) scores for all pairwise comparisons, including previously published genomes, were greater than 99% (Table 4). Noteworthy, isolates 18_466, 18_292, 19_419, 19_418, 17_471, H91-0406 and H91-0380 came from different barns and were 99.9% similar to H89-1173, which was isolated from a diseased pig in Ontario (in 1989). Multiple genome alignment identified a genomic region present in strains ATCC 33415, 17_471 and AS108/13; this was absent in strains 4286, ATCC 15557, H89-1173, H91-0380, H91-0406, 19_298, 20_277_1a and 20_377_1a (Figure 1, yellow region at approximately 1 Mbp). Strain AS108/13 was also noted to have two strain-specific regions at approximately 0.5 and 1.2 Mbp (Figure 1).

Phylogenetic analysis (Figure 2a) revealed two groups, one composed of the *A. suis* strains, markedly separated from the *A. pleuropneumoniae* NCTC 10976 strain. When investigating the *A. suis* strains only (Figure 2b), western Canadian isolates obtained from lungs clustered together; they were found closely related to the three reference isolates recovered two decades ago (H91-0406, H89-1173, H91-0380) from Ontario, Canada, two of which were also obtained from lungs samples. While European and USA isolates clustered separately from Canadian *A. suis* isolates, the position of the South American strain AS108/13 is unclear. This strain clustered separately or within the Canadian cluster depending on the phylogenetic reconstruction (Figure 2a,b), sometimes with low bootstrap
values. Genome sequences from other South American strains will be necessary to clarify the interrelationship between A. suis of the different geographic regions.

**Functional characterisation**

Proteome analysis revealed that most proteins encoded were greater than 99% similar to the reference genomes. However, a few sections with only 20%–70% identity were also identified in all genomes (Figure 3). Genomic gaps in four of nine and seven of nine western Canadian strains, when compared to the reference strain, were identified at 1.0 and 2.5 Mbp regions, respectively. The complete list of protein sequences identified with less than 70% similarity: long-chain fatty acid lead, cadmium, zinc and mercury-transporting ATPase, cell division proteins FtsP and CopG (copper reductase). Strains on lyclide in ATCC33415a provide in Supporting Information S1. In all genomes, five predicted proteins were identified with less than 70% similarity: long-chain fatty acid transport protein, Cu(I)-responsive transcriptional regulator, lead, cadmium, zinc and mercury-transporting ATPase, cell division proteins FtsP and CopG (copper reductase). Strains 20_277_1a, 20_377_1a and 4286 were the only strains with less than 70% similarity to ATCC33415 KpsT gene (capsular polysaccharide ABC transporter). This region also contained a tyrosine recombinase, usually involved in chromosomal integration of mobile genetic elements.

Putative virulence genes were identified in all western Canadian strains (Table 5). Noteworthy, strains 20_277_1a and 20_377_1a, which were isolated from heart and brain samples, carried more RTX-associated genes (n = 9) than all other strains (including ATCC33415 and all other strains isolated from lung samples). Antimicrobial resistance genes were identified only in western Canadian strains 20_277_1a and 20_377_1a, which had an ARG profile similar to the publicly available genomes (Table 6). The exception was strain AS108/13 that harboured more ARG genes than all other strains, in a unique profile that may be related to the presence of the unique genome regions mentioned above.

**DISCUSSION**

In this study, we showed that A. suis isolates obtained from organs without a mucosal surface harboured an increased number of RTX toxin genes, a potential disease determinant in A. suis. We also observed that clinical A. suis strains circulating in Canadian commercial herds between 2018 and 2019 were phylogenetically similar to those characterised in the late 1900s. Taken together, these findings reiterate the importance of A. suis surveillance to prevent the introduction of new strains and sheds a light on potential markers of virulence.

RTX toxins are widespread in the *Actinobacillus* genus. These toxins have haemolytic, cytotoxic and cytolytic activity; they can induce the production of inflammatory mediators, thereby leading to necrosis in host tissue. In this study, we found an increased number of RTX-coding genes in isolates obtained from theoretically sterile organs. In yeast, an increased number of gene copies resulted in increased protein production. When yeasts were grown under stress, such as glucose-deprived environment, cells with multiple gene copies associated with glucose metabolism were selected and thrived in that environment. In addition, *E. coli* and *Streptococcus pneumoniae* strains, with increased antimicrobial resistance gene copies, show increased phenotypic resistance to specific drugs. Gene copy number change is a phenomenon established as part of adaptation and evolution of bacteria. RTX toxins are suggested as disease determinants in A. suis. Previous reports investigating the A. suis genomes have reported a maximum of seven RTX gene copies per genome, but no correlation with clinical findings was performed. We postulate that strains with increased RTX copies may more efficiently colonise non-mucosal tissues (such as blood) reaching immune-privileged organs, leading to increased pathogenicity. Given the small number of strains with an increased number of RTX gene copies identified in this study, it remains to be verified if this change truly is associated with increased virulence. Further research focused on evaluating this change using animal models may
### TABLE 5

| Virulence factors | Reference | Source organ | ATCC | NCTC | H91-418 | 18_292 | H89-1173 | H91-0380 | H91-0406 | 19_290 | D20_277_1a | D20_377_1a | 4296 | AS108_13 | ATCC 15557 | NCTC 12996 |
|------------------|-----------|--------------|------|------|---------|--------|----------|----------|----------|--------|----------|----------|-----|--------|------------|-----------|
| Source organ     |           |              |      |      |         |        |          |          |          |        |          |          |     |        |            |           |
| Autotransporters | 42        | Blood        | 8    | 11   | 16      | 24     | 32       | 32       | 24       | 12     | 12       | 12       | 12  | 12     | 12         | 8         |
| Outer membrane   | 42        | Heart        | 13   | 11   | 17      | 24     | 32       | 32       | 24       | 12     | 12       | 12       | 12  | 12     | 12         | 8         |
| Filamentous      | 42        | Brain        | 12   | 12   | 13      | 24     | 32       | 32       | 24       | 12     | 12       | 12       | 12  | 12     | 12         | 8         |
| Haemagglutinin   | 42        | Lung         | 2    | 2    | 1       | 2      | 2        | 2        | 2        | 1      | 1        | 1        | 1   | 1      | 1           | 1         |
| Sialic acid      | 44        | Heart        | 7    | 9    | 7       | 7      | 7        | 7        | 7        | 7      | 7        | 7        | 7   | 7      | 7           | 7         |
| Lipo polysaccharides | 44     | Heart        | 5    | 6    | 4       | 4      | 4        | 4        | 4        | 4      | 4        | 4        | 4   | 4      | 4           | 4         |
| Biofilm          | 19        | Lung         | 2    | 2    | 2       | 2      | 2        | 2        | 2        | 2      | 2        | 2        | 2   | 2      | 2           | 2         |
| Iron acquisition | 45        | Lung         | 1    | 2    | 1       | 1      | 1        | 1        | 1        | 1      | 1        | 1        | 1   | 1      | 1           | 1         |
| Repeat in toxins | 10        | Blood        | 9    | 8    | 4       | 4      | 4        | 4        | 4        | 4      | 4        | 4        | 4   | 4      | 4           | 4         |

**FIGURE 1** Multiple genome alignment across all strains used in this study. Blocks with the same colours depict homologous regions.

help clarify this question. Several other virulence factors, such as lipopolysaccharides (LPS), capsular polysaccharides, proteins associated with iron chelation and cellular adhesion also play a role in *A. suis* pathogenesis. Changes in the LPS structure also appear to be associated with the range of virulence observed in different *A. suis* strains. Adhesins facilitate bacterial colonisation and biofilm formation, including the adhesion of other bacteria. Finally, this bacterium also carries receptors that recognise and bind to haemoglobin-bound iron, transferring it to the bacterial cytoplasm.

Despite decades between the isolation of the strains obtained in this study and some of those publicly available, withstanding the exposure to antibiotics used for pork production, we found limited evidence of major changes to the gene content of *A. suis* isolated from diseased pigs. In fact, some of the current strains were phylogenetically very similar to those isolated in the last century, with a high nucleotide
Phylogenetic relationship of *Actinobacillus suis* strains identified in this study (n = 9) and publicly available strains (n = 7) in relation to an *Actinobacillus pleuropneumoniae* reference strain (a) and the *A. suis* strains only (b). Trees were constructed using RAxML program by comparing 500 randomly selected genes. Organ from which isolates were obtained are coded by colour (lung: blue; heart: orange; blood: red; brain: green). Shapes indicated geographical origin of each isolate (circle: Manitoba, Canada; square: Ontario, Canada; triangle: USA; diamond: Brazil; rectangle: UK).

Similiarity score. Limited genomic fragment insertions were identified in the western Canadian strains and the gene order identified was the same. It has been established that the genome structure of bacteria can change slowly with time due to local or global mutations.5-37

Treatment and control of *A. suis* is usually achieved through antimicrobial therapy, often based on ceftiofur, gentamicin and trimethoprim/sulphadiazine, ampicillin, sulphadimethoxine or tiamulin.38 Interestingly, we did not identify ARG related to any drugs used in pork production.
in the western Canadian strains. The only strain harbouring ARG for β-lactams, sulphadiazine and sulphadimethoxine was ASI08/13. One caveat in this analysis was the lack of phenotypic antimicrobial susceptibility data. While future investigations should focus on evaluating the efficacy of *A. suis* whole genome sequencing in predicting resistance patterns, it has been shown to be highly correlated to phenotypical resistance results for other bacterial species.\(^4\) Despite this, we did identify nucleotides changes in the gene coding for the copper resistance protein CopG present in the western Canadian strains, when compared to the ATCC 33415 reference strain. CopG encodes a 14 kDa periplasmic protein that is suggested to contribute to copper resistance by reducing free copper ions.\(^4\) This protein is widely spread across Gram-negative bacterial species; genes encoding CopG homologues are often found associated with other known metalloproteins.\(^4\) One study\(^1\) proposed that CopG is the main copper detoxification mechanism employed by *Vibrio cholerae* when under anaerobic conditions. Future research may further explore how the changes identified in the western Canadian strains contribute to the biological activity of CopG.

*A. suis* remains a challenge for swine production. As the livestock industry moves towards a reduction in overall antimicrobial usage, it is likely that this agent will become more relevant in the coming years. Here, we have shown that isolates obtained from brain and heart had more RTX toxin gene copies in their genomes than those isolated from lungs. We have shown that *A. suis* recently circulating in commercial farms in Canada are similar to isolates from 1990 to 2000s. Antimicrobial resistance genes detected in clinical isolates suggest the potential for resistance to drugs of importance to human medicine.

### Author Contributions
Matheus de Oliveira Costa acquired funding and conceptualized the study. Matheus de Oliveira Costa, Dharmasiri Gamage Ruwini Sulochana Kulathunga and Alaa Abou Fakher performed the analyses. Dharmasiri Gamage Ruwini Sulochana Kulathunga drafted the first version of the manuscript. Matheus de Oliveira Costa reviewed the manuscript and the data analyses.

### Acknowledgements
We thank the Veterinary Diagnostic Service, Manitoba, Canada and Gallant Custom Laboratories (Cambridge, Ontario, NIR 6J9, Canada) for their assistance with clinical sample handling. This work was supported by a Natural Sciences and Engineering Research Council of Canada grant to Matheus de Oliveira Costa (RGPIN06353-2020).

### Conflicts of Interest
The authors declare they have no conflicts of interest.

### Data Availability Statement
Supporting information data are available at https://doi.org/10.23644/uu.19346372

### Ethics Statement
No ethical approval was required because the work did not directly use any live pigs.

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### Table 6

| Strain ID   | Resistance genes (>80% identity) |
|-------------|----------------------------------|
| 20_277_1a  | EF-Tu, Sul2, APH(3′)-Ia, EF-Tu   |
| 20_377_1a  | EF-Tu, Sul2, APH(3′)-Ia, EF-Tu   |
| 19_419     | EF-Tu, Sul2, APH(3′)-Ia, EF-Tu   |
| 19_418     | EF-Tu, Sul2, APH(3′)-Ia, EF-Tu   |
| 17_471     | EF-Tu, Sul2, APH(3′)-Ia, EF-Tu   |
| 4286       | EF-Tu, Sul2, APH(3′)-Ia, EF-Tu   |
| 18_292     | EF-Tu, Sul2, APH(3′)-Ia, EF-Tu   |
| 18_466     | EF-Tu, Sul2, APH(3′)-Ia, EF-Tu   |
| 19_298     | EF-Tu, Sul2, APH(3′)-Ia, EF-Tu   |
| ATCC 33415 | EF-Tu, Sul2, APH(3′)-Ia, EF-Tu   |
| ATCC 15557 | EF-Tu, Sul2, APH(3′)-Ia, EF-Tu   |
| H89-1173   | EF-Tu, Sul2, APH(3′)-Ia, EF-Tu   |
| H91-0380   | EF-Tu, Sul2, APH(3′)-Ia, EF-Tu   |
| H91-0406   | EF-Tu, Sul2, APH(3′)-Ia, EF-Tu   |
| NCTC 12996 | EF-Tu, Sul2, APH(3′)-Ia, EF-Tu   |
| ASI08/13   | ROB-10, Tet(H), APH(3′)-Ia, EF-Tu, Sul2, APH(3′)-Ia, EF-Tu |

Abbreviations: APH(3′)-Ia, aminoglycoside; APH(6)-Id, aminoglycoside; EF-Tu, *Escherichia coli* EF-Tu mutants conferring resistance to pulvomycin/elfamycin class; ROB-10, cephalosporin, penam; Sul2, sulphonamide; Tet(H), tetracyclines.
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**SUPPORTING INFORMATION**

Additional supporting information can be found online in the Supporting Information section at the end of this article.