Identification and characterization of the chromosomal yefM-yoeB toxin-antitoxin system of Streptococcus suis

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Toxin-antitoxin (TA) systems are widely prevalent in the genomes of bacteria and archaea. These modules have been identified in Escherichia coli and various other bacteria. However, their presence in the genome of Streptococcus suis, an important zoonotic pathogen, has received little attention. In this study, we describe the identification and characterization of a type II TA system, comprising the chromosomal yefM-yoeB locus of S. suis. The yefM-yoeB locus is present in the genome of most serotypes of S. suis. Overproduction of S. suis YoeB toxin inhibited the growth of E. coli, and the toxicity of S. suis YoeB could be alleviated by the antitoxin YefM from S. suis and Streptococcus pneumoniae, but not by E. coli YefM. More importantly, introduction of the S. suis yefM-yoeB system into E. coli could affect cell growth. In a murine infection model, deletion of the yefM-yoeB locus had no effect on the virulence of S. suis serotype 2. Collectively, our data suggested that the yefM-yoeB locus of S. suis is an active TA system without the involvement of virulence.

Toxin-antitoxin (TA) systems are widely distributed and highly abundant in the genomes of bacteria and archaea1,2. Typically, they consist of two genes: one encoding the antitoxin (A) and the other encoding the toxin (T)3. The two genes are usually organized as operons3. TA systems were initially identified on low copy number plasmids, where they help to maintain plasmid stability by a mechanism called post-segregational killing or genetic addiction5,6. Currently, TA systems are classified into five types, according to the nature and mode of action of the antitoxin2,7. In the type I TA system, the antitoxin is an antisense RNA transcribed from the toxin gene, but in the reverse orientation, thus it binds directly to the toxin mRNA and inhibits production of the toxin1. The antitoxin of type III TA system is also an RNA, which interacts directly with the toxin and inhibits its toxicity8. Type IV and V are two recently reported TA systems, in which the antitoxin either binds to the toxin targets and functions as an antagonist for the toxin toxicity (type IV)9 or inhibits the toxin by cleaving its mRNA specifically (type V)10. Type II TA systems are pretty prevalent in bacterial chromosomes and have been most extensively studied1,11. In the type II TA system, the antitoxin and toxin are both proteins. By forming a complex with its cognate toxin, the antitoxin blocks the toxicity of the toxin1.

Various roles are proposed for TA systems, including maintenance of plasmid stability, coping with stress, as mediators of programmed cell death and antiphage activity2,12–14. However, the possibility that TA systems could contribute to bacterial pathogenicity has been largely neglected. Mycobacterium

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tuberculosis possesses a total of 79 TA systems, whereas its nonpathogenic counterpart, Mycobacterium smegmatis contains only four. This led to speculation that TA systems may be related to the pathogenicity of pathogens. A recent study reported that TA systems can promote colonization and stress resistance of uropathogenic E. coli. In addition, TA modules have been shown to be involved in the virulence of Salmonella and Leptospira interrogans. Thus, evidence is increasing that some TA systems are implicated in bacterial pathogenesis and host-pathogen interactions.

Streptococcus suis (S. suis) is a major swine pathogen causing severe economic losses in the swine industry worldwide. It is also the causative agent in humans in close contact with infected swine or pork by-products. S. suis infections in swine lead to a variety of diseases, including meningitis, septicemia, endocarditis, arthritis and even sudden death. In humans, the infection causes meningitis, septicemia and streptococcal toxic shock syndrome (STSS). To date, 33 serotypes (types 1 to 31, 33, and 1/2) of S. suis have been described based on the composition of the capsular polysaccharide (CPS). Among them, S. suis serotype 2 (S. suis 2 or SS2) is the most virulent and the most commonly isolated serotype in association with diseases in most countries. In 1998 and 2005, two large outbreaks of human S. suis 2 infections occurred in China, resulting in 25 cases with 14 deaths and 215 cases with 38 deaths, respectively. Despite numerous studies having been carried out over the past few years, knowledge of the pathogenesis of S. suis remains very limited. In the genome of S. suis 2 strain SC84, nine type II TA systems have been predicted by bioinformatics analysis (unpublished data). Among them, only SezAT has been identified as an active TA system, yet its functions remain elusive. To gain further insight into the pathogenesis of S. suis infection, other TA systems should be identified and their roles should be explored.

The yefM-yoeB module belongs to one of the well-characterized TA systems in E. coli and S. pneumoniae. This TA module was first identified in E. coli, as the homolog of the Enterococcus faecium ace-tx TA system. Since then, the yefM-yoeB TA system has been described in many bacterial species, including major pathogens such as S. pneumoniae, M. tuberculosis and Staphylococcus aureus. Moreover, the yefM-yoeB locus was identified as the first TA system in Streptococcus. Recently, the expression pattern and structural peculiarities of the yefM-yoeB TA system were investigated in Lactobacillus rhamnosus. E. coli YefM and YoeB form a heterotrimeric complex (YefM-YoeB) that blocks the effect of the toxin.

When liberated from the complex, YoeB binds to the 50 S ribosomal subunit and cleaves mRNA at the A site, thereby inhibiting translation initiation in E. coli and S. aureus. Overproduction of the Lon protease specifically activates YoeB-dependent mRNAs cleavage, suggesting that Lon might be responsible for YefM degradation. In addition, the yefM-yoeB TA system is transcriptionally autoregulated. YefM can bind to the promoter and repress transcription, while YoeB acts as a repression enhancer. Bioinformatics analysis revealed that the yefM-yoeB locus is also present in the genome of most serotypes of S. suis. However, whether this locus encodes an active TA system is still unknown.

In this study, a selective expression vector was constructed to characterize the TA systems in E. coli. We showed that the yefM-yoeB locus is an active TA system of S. suis, and introduction of this system into E. coli could affect cell growth. We also explored the involvement of yefM-yoeB in the virulence of S. suis 2 and found that the module has no role in virulence.

**Results**

**Identification of the yefM-yoeB locus in S. suis.** To identify the YoeB toxin homologs, a BlastP search against the proteins annotated in the genome of S. suis SC84 was performed, using the YoeB sequences of S. pneumoniae R6 and E. coli MG1655 as query sequences. Both searches revealed an open reading frame (SSUSC84_1817) encoding an 85 amino acid protein sharing 75% and 50% identity with the S. pneumoniae and E. coli YoeB toxins, respectively. Hence, the protein was termed YoeB. The YefM antitoxin encoded by SSUSC84_1818 was identified by the same method. BlastP analysis showed that the S. suis YefM has 79% and 29% amino acid sequence identity with YefM from S. pneumoniae and E. coli, respectively. Multiple sequence alignments further revealed that 1) the S. suis YefM-YoeB system shares high level of homology with that from S. pneumoniae and E. coli; 2) YoeB processes several conserved residues required for its activity (Glu46, Arg65, His83 and Tyr84) (Fig. 1). Protein homology modeling using CPHmodels predicted the structures of S. suis YefM and YoeB. The secondary structure of YefM is proposed to consist of four α-helices and three β-sheets (Fig. 1a), while that of YoeB contains two α-helices, five β-sheets and a coil (Fig. 1b). To determine whether the yefM-yoeB locus was universally present in the genomes of S. suis, BlastN analysis was performed using the 21 complete genomes of S. suis available in the National Centre for Biotechnology Information database as of 31 May 2015. The results confirmed that all strains harbour the yefM-yoeB locus, except for strain D12, a serotype 9 S. suis (see Supplementary Fig. S1 online).

A genetic structure analysis revealed that yefM is located upstream of yoeB, and the two genes are separated by one nucleotide, apparently arranged in a bicistronic operon (Fig. 2a). BPROM analysis of the upstream region of the yefM gene (about 300 bp) identified the putative −35 and −10 regions, which are located in the intergenic region between the yefM gene and its upstream gene (Fig. 2a). To assess whether yefM and yoeB are co-transcribed in S. suis, a reverse transcription polymerase chain reaction (RT-PCR) analysis was performed. Reverse transcriptase was used to synthesize cDNAs and the resulting cDNAs were PCR amplified using primer pair A1/T2. The individual yefM and yoeB genes were also amplified using primer pairs A1/A2 and T1/T2, respectively. As shown in Fig. 2b, the PCR products...
were of the expected sizes for yefM (261 bp), yoeB (258 bp) and yefM-yoeB (520 bp), all consistent with that of the genomic DNA. No PCR products were evident in the negative controls, in which the reverse transcription was performed without the enzyme, therefore eliminating possible DNA contamination. These results demonstrated that in S. suis, yefM and yoeB are actively co-transcribed, thus forming a bicistronic operon.

Construction of a selective expression vector to characterize the toxin-antitoxin systems in E. coli. To characterize the toxin-antitoxin systems in E. coli, a selective expression vector was constructed as previously described. A DNA fragment containing the araC gene and the promoter PBAD was amplified from the pBADhisA plasmid, digested with the XhoI and HindIII enzymes, and then cloned into pET-30a, an expression vector induced by isopropyl β-D-thiogalactopyranoside (IPTG), to generate the selective expression vector, designated pETBAD (see Supplementary Fig. S2 online). Plasmid pETBAD has five unique restriction sites for cloning and possesses the IPTG-inducible promoter Plac and the arabinose-inducible promoter PBAD, thus expression can be induced by IPTG and/or arabinose (see Supplementary Fig. S2 online).

Overproduction of YoeB inhibits cell growth in E. coli which can be alleviated by YefM. To determine whether the yefM-yoeB locus is indeed an active TA system, the yefM and yoeB genes were cloned separately as well as together into the pBADhisA expression vector. The plasmids were introduced into E. coli Top10 cells, and the transformsants were selected in LB agar plates with 0.2% D-glucose (repressed conditions of PBAD). E. coli Top10 cells harbouring the corresponding plasmids were grown in LB medium, and 0.2% D-glucose or L-arabinose was added at time zero. In the presence of 0.2%
D-glucose, Top10 cells harbouring the pBADhisA-yefM and pBADhisA plasmids showed no major difference in growth, while cells carrying the pBADhisA-yoeB plasmid showed a moderate growth defect (Fig. 3a). In the case of inductive conditions (0.2% L-arabinose), E. coli Top10 cells harbouring the pBADhisA-yoeB plasmid exhibited drastic growth inhibition, while cells harbouring other two plasmids showed only moderate reductions in their OD<sub>600</sub> value (Fig. 3b). Surprisingly, under both repressed and inductive conditions, E. coli Top10 cells harbouring the pBADhisA-yefM-yoeB plasmid exhibited obvious growth inhibition (Fig. 3), even so, Top10 cells carrying the pBADhisA-yefM-yoeB plasmid showed much better growth than that carrying pBADhisA-yoeB (Fig. 3b), indicating that YoeB-induced growth inhibition could be alleviated by YefM.

We further investigated the toxic and antitoxic effect of the TA components using the selective expression system constructed here. In the selective expression plasmid, the IPTG-inducible promoter <i>P<sub>lac</sub></i> and the arabinose-inducible promoter <i>P<sub>BAD</sub></i> control the expression of YefM and YoeB, respectively (Fig. 4a). Thus, E. coli BL21 (DE3) cells harbouring the pETBAD-<i>yefM</i><sub>Ssu</sub>-yoeB plasmid could express the <i>S. suis</i> YefM and/or YoeB upon induction with IPTG and/or L-arabinose. As shown in Fig. 4b, the E. coli BL21 (DE3) cells exhibited considerable growth inhibition in the presence of L-arabinose. In contrast, only moderate growth inhibition was observed in the presence of IPTG or IPTG and L-arabinose together.

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**Figure 2.** The yefM-yoeB locus is organized as an operon. (a) Genetic organization of the yefM-yoeB locus in <i>S. suis</i> strain SC84. The putative −35 and −10 regions are boxed. The primers used for RT-PCR or PCR are drawn as arrows, and the expected sizes of the corresponding PCR products are shown below. (b) Co-transcription analysis. Total RNAs extracted from <i>S. suis</i> SC84 were used to synthesize cDNAs. PCR was carried out with primer pairs A1/A2, T1/T2 and A1/T2, respectively. Lanes 1, 4 and 7 represent the amplification using cDNAs as the template; Lanes 2, 5 and 8 represent the amplification using genomic DNA (gDNA) as the template; Lanes 3, 6 and 9 represent the amplification using cDNA- (cDNA reaction without reverse transcriptase) as the template. The DL 2000 DNA Marker is shown on the left (lane M).

**Figure 3.** Effect of induction of YoeB, YefM or YefM-YoeB on growth of <i>E. coli</i>. Overnight cultures of <i>E. coli</i> Top10 cells harbouring the plasmid pBADhisA-yefM, pBADhisA-yoeB, pBADhisA-yefM-yoeB and pBADhisA were diluted 1:1000 in LB-ampicillin. Each culture was then divided into two equal volumes. The first half served as the control, to which 0.2% D-glucose was added (a), 0.2% L-arabinose was added to the second half to induce expression of the target gene (b). Culture growth was evaluated by measuring the OD<sub>600</sub> every hour. The data shown are averages with standard deviations for the results from three independent experiments.
These results indicated that the protein encoded by the yoeB gene is a toxin against *E. coli* and that the protein encoded by the yefM gene could counteract the toxicity. Therefore, the *S. suis* yefM-yoeB locus works as a typical TA system.

**YoeB<sub>ssu</sub>-induced growth inhibition in *E. coli* that could be alleviated by YefM<sub>ssu</sub> but not by YefM<sub>eco</sub>.** In *S. pneumoniae* and *E. coli*, the toxicity of YoeB could be counteracted only by its cognate antitoxin<sup>28</sup>. To test whether there was cross-complementation between non-cognate YefM and *S. suis* YoeB, *E. coli* BL21 (DE3) cells were transformed with plasmid pETBAD-yefM<sub>ssu</sub>-yoeB and pETBAD-yefM<sub>eco</sub>-yoeB (Fig. 4a). As seen in Fig. 4c, induction of YoeB<sub>ssu</sub> resulted in a drastic reduction in OD<sub>600</sub> value, whereas coinduction of YefM<sub>ssu</sub> and YoeB<sub>ssu</sub> alleviated the growth inhibition in *E. coli*, indicating that the toxic effect of YoeB<sub>ssu</sub> was counteracted by coexpression of YefM<sub>ssu</sub>. However, coinduction of YefM<sub>eco</sub> and YoeB<sub>ssu</sub> did not neutralize the YoeB<sub>ssu</sub> toxicity (Fig. 4d). In addition, induction of YefM<sub>ssu</sub>, YefM<sub>ssu</sub>, and YefM<sub>eco</sub> also displayed an effect on growth inhibition in *E. coli* (Fig. 4b–d).

**Introduction of the *S. suis* yefM-yoeB system into *E. coli* could affect cell growth.** It seemed likely that introduction of the *S. suis* yefM-yoeB system into *E. coli* affects cell growth, since *E. coli* Top 10 cells carrying the pBADhisA-yefM-yoeB plasmid showed considerable growth inhibition under both repressed and inductive conditions. To test the hypothesis, the yefM and yoeB genes were cloned together into the pET-30a expression plasmid. When introduction of the pET30a-yefM-yoeB and pET-30a plasmids into *E. coli* Trans5α and Top10 strains, cells carrying pET30a-yefM-yoeB showed an obvious growth defect compared with cells carrying pET-30a (Fig. 5a). However, when introduction of the two plasmids into *E. coli* BL21(DE3) strain, no major difference in growth was found (Fig. 5a).

The same experiments were carried out with the pSET2-yefM-yoeB plasmid. In this plasmid, the *S. suis* yefM-yoeB locus is under the control of its own promoter. As shown in Fig. 5b, *E. coli* Trans5α cells harbouring the pSET2-yefM-yoeB plasmid exhibited a remarkable growth defect compared with that harbouring the empty plasmid. The growth inhibition effect was even more severe when the plasmid was transformed into Top10 strain, as cells carrying the pSET2-yefM-yoeB plasmid showed growth arrest.
over a period of 12 hours (Fig. 5b). However, BL21(DE3) strain harbouring pSET2-yefM-yoeB showed only a slight defect in growth.

Next, the pBADhisA-yefM-yoeB and pBADhisA plasmids were transformed into E. coli Trans5α and BL21(DE3) strains. Both strains carrying the pBADhisA-yefM-yoeB plasmid exhibited an obvious defect in growth compared with that carrying the pBADhisA plasmid (Fig. 5c).

Finally, growth of E. coli BL21(DE3) strain harbouring the pET30a-yefM-yoeB and pET-30a plasmids was determined in the presence of 1 mM IPTG. Under inductive conditions, BL21(DE3) cells harbouring pET30a-yefM-yoeB also displayed an obvious growth inhibition effect (Fig. 5d).

Taken together, the results clearly demonstrated that introduction of the S. suis yefM-yoeB system into E. coli could affect cell growth.

Construction and microbiological characterization of the ΔyefM-yoeB mutant. To investigate the functions of the yefM-yoeB locus in S. suis 2, an isogenic yefM-yoeB knockout mutant of S. suis 2 strain SC19, termed ΔyefM-yoeB, was constructed through homologous recombination (Fig. 6a). To rule out the possible polar effect and introduction of a second mutation during the construction of ΔyefM-yoeB, we generated a complementation strain, designated CΔyefM-yoeB using the E. coli-S. suis shuttle vector pSET241. The resulting mutant and complementation strains were confirmed by PCR (Fig. 6b), RT-PCR (Fig. 6c) and direct DNA sequencing (data not shown).

The effects of yefM-yoeB deletion on the basic microbiological properties of S. suis 2 were investigated in terms of morphology, haemolytic activity and in vitro growth. The cell morphologies of the ΔyefM-yoeB mutant, WT and CΔyefM-yoeB strains were examined under light microscope using Gram staining. However, no obvious differences were found (see Supplementary Fig. S3a online). When inoculated on sheep blood agar plates, the three strains showed similar haemolytic activity (see Supplementary Fig. S3b online). The growth kinetics of the mutant strain were compared with those of the WT and complementation strains by measuring the optical density at 600 nm (OD600) every hour. We found that the growth kinetics of ΔyefM-yoeB were essentially identical to those of the WT and CΔyefM-yoeB strains (Fig. 7), indicating that the yefM-yoeB locus of S. suis 2 plays no role in growth in vitro.

Deletion of the yefM-yoeB locus has no effect on S. suis 2 virulence in mice. To assess the role of the yefM-yoeB locus in the pathogenesis of S. suis 2, we performed an experimental infection model in CD1 mice. As an initial comparison of virulence, groups of ten CD1 mice were inoculated intraperitoneally with 6×10⁵ CFU of the WT, ΔyefM-yoeB, CΔyefM-yoeB strains or PBS. Most mice infected with S. suis strains developed typical clinical signs of S. suis 2 infection, including depression, lethargy,
weakness, prostration and rough coat hair during the first 72 h post infection. Ultimately, the survival rates of mice in the WT, ΔyefM-yoeB and CΔyefM-yoeB groups were 50%, 60% and 30%, respectively (Fig. 8a). By contrast, all mice inoculated with PBS remained healthy and survived. No significant difference in survival rates was observed between the ΔyefM-yoeB group and the WT group (P = 0.6793),
or the CΔyeFM-yoeB group (P = 0.1924). Thus, it seemed likely that the yeFM-yoeB locus has no role in S. suis 2 virulence.

A competitive-infection assay was adopted to further compare the abilities of the WT strain and the ΔyeFM-yoeB mutant to establish infection. Four mice were inoculated intraperitoneally with a 1:1 mixture of the WT and mutant bacteria. Mice were sacrificed to collect blood, brain, heart, liver, spleen, lung and kidney samples 24 h after inoculation. Bacterial cells recovered from various tissue samples were analysed by colony PCR to determine the competitive index (CI). The result showed that for each tissue, the mean CI values were approximately 1 (Fig. 8b), suggesting that the mutant and WT strains have similar abilities to colonize the tissues.

Taken together, these results indicated that the yeFM-yoeB locus is not involved in the virulence of S. suis 2.

**Discussion**

TA systems have attracted an increasing concern in recent years because of their abundance in the genomes of bacteria and archaea on the one hand, and the limited of knowledge of their physiological functions on the other. In E. coli K12, at least 33 TA systems have been identified, with several being well characterized. However, only one TA module, SezAT, has been described in S. suis, yet its function has not been demonstrated. The yeFM-yoeB module is one of the best studied TA systems and has been described in various bacteria, including E. coli, S. pneumoniae, M. tuberculosis, S. aureus, Staphylococcus equorum and Streptomyces.
In this study, we showed that the chromosomally encoded yefM-yoeB locus of *S. suis* is an active TA system with yoeB encoding the toxin and yefM encoding the cognate antitoxin. This is not surprising, as this system shows considerable similarity to the YefM-YoeB system from *S. pneumoniae* and *E. coli*. Like most TA systems, the yefM and yoeB genes are co-transcribed. Upstream of the yefM gene, there is an intergenic region of 75 nucleotides, which may act as the promoter region. Overproduction of the YoeB toxin in *E. coli* Top10 and BL21 (DE3) cells both resulted in toxic effects commonly linked to toxin activity. YoeB homologs were identified as endonucleases that inhibit translation by cleaving mRNA, either in a ribosome-dependent or -independent manner. We therefore reasoned that *S. suis* YoeB inhibits cell growth via a similar mechanism.

The toxic effect of *S. suis* YoeB toxin could be neutralized by both the cognate YefM<sub>Spn</sub> and the heterologous YefM<sub>Spn</sub> but not by the *E. coli* counterpart, consistent with the fact that YefM<sub>Spn</sub> shares higher levels of identity with YefM<sub>Spn</sub> (79% identity versus 29% for YefM<sub>Eco</sub>). As YefM<sub>Spn</sub> and YefM<sub>Spn</sub> display a high level of sequence homology, we speculated that YefM<sub>Spn</sub> could bind to and neutralize the YoeB<sub>Spn</sub>. The lack of cross-complementation between YefM<sub>Eco</sub> and YoeB<sub>Spn</sub> suggested that there is no favourable interaction between the two heterologous proteins. Similar behaviour has also been reported for YoeB<sub>Eco</sub>, whose activity could be alleviated by both the cognate YefM<sub>Eco</sub> and the antitoxin Axe of *E. faecium*, but not by YefM<sub>Spn</sub>. A previous study showed that overexpression of YefM<sub>Eco</sub> displayed toxicity in *E. coli* at high expression levels, consistent with our observation that YefM<sub>Spn</sub> YefM<sub>Spn</sub> and YefM<sub>Spn</sub> had an effect on growth inhibition in *E. coli*.

An interesting observation was that *E. coli* Top10 cells harbouring the pBADhisA-yefM-yoeB plasmid showed an obvious growth defect under both repressed and inductive conditions. Similar experiments were then performed with other plasmids and strains. Plasmids containing the *S. suis* yefM-yoeB system were introduced into different strains of *E. coli*. Except for BL21(DE3) strain carrying the pET30a-yefM-yoeB plasmid, all tested strains harbouring the plasmids containing the *S. suis* yefM-yoeB system showed growth inhibition. A previous study revealed that the YefM-YoeB complex forms a 2:1 heterotrimer. We speculated that the yefM-yoeB system could be expressed even under repressed conditions and that YefM expression was not enough to counteract YoeB, thus leading to growth inhibition in *E. coli*. Since pET-30a and *E. coli* BL21(DE3) strain constitute a precise inducible expression system, it is possible that the yefM-yoeB locus on pET-30a was not expressed without inducer, therefore growth inhibition was not observed for BL21(DE3) carrying pET30a-yefM-yoeB under normal growth conditions. The speculation agrees with the result that BL21(DE3) carrying the pET30a-yefM-yoeB plasmid exhibited a growth defect under inductive conditions.

To investigate the functions of the yefM-yoeB locus in *S. suis* 2, a knockout mutant and the corresponding complementation strain were constructed. No obvious differences between the WT and the yefM-yoeB deletion mutant were found in terms of their cell morphology, haemolytic activity on blood agar plates, and *in vitro* growth. The potential role of TA systems in bacterial pathogenesis has been neglected for a long time. More and more studies have revealed that TA systems are involved in bacterial pathogenicity and host-pathogen interactions. Therefore, we evaluated the effect of this TA system on the pathogenesis of *S. suis* 2 using a murine infection model. Survival curves of mice and the competitive-infection assay both demonstrated that deletion of the yefM-yoeB locus had no role in the pathogenicity of *S. suis* 2. It was reported that *Yersinia pestis* lacking the hicB3 antitoxin is virulence-attenuated; however, the mutant lacking the whole hicA3B3 locus is fully virulent. In contrast, the toxins ChpK and MazF but not the antitoxins ChpI and MazE are involved in the virulence of *L. interrogans* during infection. Future experiments should evaluate the involvement of the individual genes of the yefM-yoeB locus in the virulence of *S. suis* 2.

It should be noted that TA systems play important roles in the physiology of cells, including biofilm formation and multidrug resistance. The effect of the yefM-yoeB module on biofilm formation, stress tolerance and formation of persistor cells will be explored in future studies. It is proposed that TA systems are potential targets for antibiotics. Given the fact that *S. suis* YoeB can inhibit the growth of *E. coli* considerably, we are planning to examine the effect of YoeB on *S. suis*. If a similar toxic effect is observed, a multivalent strategy to synthesize an inhibitor that interacts with the YefM antitoxin and frees the toxin YoeB to inhibit bacterial growth could be promising for the development of new antibiotics.

In conclusion, the yefM-yoeB locus was identified as a new TA system of *S. suis*. The present study clearly demonstrated that the yoeB gene encodes a toxin that can inhibit the growth of *E. coli*. Specifically, the toxicity of *S. suis* YoeB could be alleviated by the cognate *S. suis* YefM and heterologous *S. pneumoniae* YefM. More importantly, we reported that introduction of the yefM-yoeB TA system into *E. coli* could affect cell growth. In addition, deletion of the yefM-yoeB locus had no effect on the virulence of *S. suis* 2.

**Methods**

**Bacterial strains and growth conditions.** Bacterial strains and plasmids used in this study are listed in Supplementary Table S1. *S. suis* strains were maintained on Tryptic Soy Broth (TSB) or Tryptic Soy Agar (TSA; Difco Laboratories, Detroit, MI, USA) with 10% (vol/vol) newborn bovine serum at 37°C, unless otherwise specified. *E. coli* strains were cultured in Luria-Bertani (LB) broth or on LB agar at 37°C. When necessary, antibiotics (purchased from Sigma) were added at the following concentrations:
for *E. coli*, ampicillin, 75 μg/ml; kanamycin, 25 μg/ml and spectinomycin, 50 μg/ml; for *S. suis*, spectinomycin, 100 μg/ml.

**RNA isolation and RT-PCR analysis.** Total RNA samples were prepared from *S. suis* cultures using an SV total RNA isolation system (Promega), according to the manufacturer's protocol. RNA concentrations and integrity were determined by UV spectrophotometry and agarose gel electrophoresis, respectively. RT-PCR was carried out using a QuantiTect Reverse Transcription Kit (Qiagen), according to the manufacturer's instructions. For the co-transcription assay, the gene specific primers A1, A2, T1 and T2 were used for RT-PCR analysis (see Supplementary Table S2 online). To identify the mutant and complementation strains, primer pair ATin1/ATin2 was used.

**Plasmid Construction.** Plasmids were constructed as follows using the primers listed in Supplementary Table S2.

1. pETBAD. Primer pair BAD1/BAD2 amplified the DNA fragment containing the *araC* gene and the promoter *Pbad* from the pBADhisA plasmid. The DNA fragment was digested with the *Xho* I and *Hind* III enzymes, and cloned into pET-30a, to generate the selective expression plasmid pETBAD.
2. pBADhisA-*yefM* and pBADhisA-*yoeB*. The *yefM* gene was amplified from the *S. suis* 2 genome using primer pair yeM1/yeM2. The PCR product was digested with the *Xho* I and *Hind* III enzymes, and then cloned into pBADhisA, to generate plasmid pBADhisA-*yefM*. Plasmid pBADhisA-*yoeB* was constructed in a similar manner.
3. pBADhisA-*yefM*-yoeB and pET30a-*yefM*-yoeB. The *yefM* and *yoeB* genes were amplified from the *S. suis* 2 genome using primer pairs yeM1/R1 and R2/yoeB2, respectively. The two DNA fragments were fused into one fragment using overlap extension PCR. This DNA fragment was digested with the *Xho* I and *Hind* III enzymes, and then ligated into pBADhisA, to generate pBADhisA-*yefM*-yoeB. Plasmid pET30a-*yefM*-yoeB was constructed in a similar manner, except that the *yefM* gene was amplified using primer pair yeM3/R1 and the fused DNA fragment was digested with the *Bam* HI and *Hind* III enzymes.
4. pETBAD-*yefM*spa-yoeB, pETBAD-*yefM*spa-yoeB and pETBAD-*yefM*spa-yoeB. The *yefM* gene was amplified from the *S. suis* 2 genome using primer pairs Spa1/Spa2. The DNA fragment was digested with the *Kpn* I and *Eco* R I enzymes, and then cloned into plasmid pETBAD to yield pETBAD-*yefM*. The *yoeB* gene was amplified from the *S. suis* 2 genome using primer pair Spa1/Spa2. After digestion with the *Hind* III and *Sac* I enzymes, the fragment was cloned into pETBAD-*yefM*, to generate pETBAD-*yefM*-yoeB. This construct placed the *yefM* and *yoeB* genes under the control of the IPTG-inducible promoter *Plac* and the arabinose-inducible promoter *Pbad*, respectively. Thus, IPTG could induce the expression of *yefM* and arabinose could induce *yoeB*. The two other plasmids were constructed using the same procedure, except that the *yefM* and *yoeB* genes were amplified from the *S. pneumoniae* R6 and *E. coli* K12 genomes, respectively.
5. pSET2-*yefM*-yoeB. A DNA fragment containing the *yefM*-yoeB locus and its predicted promoter was amplified from the *S. suis* 2 genome using primer pair CAT1/CAT2. After digestion with the *Pst* I and *Eco* R I enzymes, the fragment was cloned into pSET2, to generate the plasmid pSET2-*yefM*-yoeB.
6. pSET4s-*ΔyefM*-yoeB. Two flanking fragments (LA and RA) of the *yefM*-yoeB locus were amplified from the *S. suis* 2 genome using primer pairs LA1/LA2 and RA1/RA2, respectively. After digestion with the appropriate restriction enzymes, the two fragments were simultaneously cloned into pSET4s to generate the knockout plasmid pSET4s-*ΔyefM*-yoeB.

**E. coli growth analysis.** *E. coli* Top10 cells transformed with pBADhisA-*yefM*, pBADhisA-*yoeB*, pBADhisA-*yefM*-yoeB and pBADhisA were cultured overnight in LB broth supplemented with 75 μg/ml ampicillin and 0.2% D-glucose. The next day, the four cultures were diluted 1:1000 in LB-ampicillin. Each culture was then divided into two equal volumes. The first half served as the control, to which 0.2% D-glucose was added, while 0.2% L-arabinose was added to the second half to induce expression of the target gene. Culture growth was evaluated by measuring the OD600 every hour.

*E. coli* BL21(DE3) cells harbouring the respective selective expression plasmids were incubated in LB broth supplemented with 25 μg/ml kanamycin to an OD600 of about 0.3. Each culture was divided into four equal parts, to three of which was individually added 0.2% L-arabinose, 1 mM IPTG or both, respectively. The fourth part had nothing added to it and served as a control. These cultures were further incubated and samples were taken every hour to determine the OD600.

*E. coli* strains were transformed with plasmids containing the *S. suis* *yefM*-yoeB system or the corresponding empty plasmids. Cells were cultured overnight in LB broth supplemented with antibiotics and diluted 1:1000 in fresh medium. Culture growth was monitored by measuring the OD600. For those cells showing growth arrest in LB broth, such as Top10 carrying pSET2-*yefM*-yoeB, isolated colonies were used as inocula. In parallel, colonies of the same size for strains carrying the corresponding empty plasmids were used.
Deletion of the yefM-yoeB locus and functional complementation. Gene knockout mutant of the yefM-yoeB locus was constructed using plasmid pSET4s as described previously. The knockout plasmid pSET4s-ΔyefM-yoeB was introduced into the competent cells of S. suis SC19 by electroporation. After two steps of allelic exchange at 28 °C, spectinomycin-sensitive clones were selected to identify the mutants by PCR using primers listed in Supplementary Table S2. The mutants were further confirmed by RT-PCR analysis and direct DNA sequencing.

For complementation assays, the recombinant plasmid pSET2-ΔyefM-yoeB was introduced into the ΔyefM-yoeB mutant by electroporation. The complementation strain CΔyefM-yoeB was selected with spectinomycin and confirmed using PCR, RT-PCR and DNA sequencing.

Mouse infections. The Laboratory Animal Monitoring Committee of Huazhong Agricultural University approved all the animal experiments, which were performed in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of Hubei Province, China. Forty female CD1 mice (5-weeks-old) were randomly divided into four groups with 10 mice per group. Mice in Groups 1, 2 and 3 were inoculated intraperitoneally with 6 × 10⁵ CFU in 200 μL PBS of the WT, ΔyefM-yoeB and CΔyefM-yoeB strains, respectively. Group 4 was injected with 200 μL PBS, and served as the control group. Mice were monitored daily over 14 days for clinical signs and survival rates.

For the competitive-infection assay, four mice were inoculated intraperitoneally with a mixture of the WT and mutant strains at a ratio of 1:1 (1 × 10⁸ CFU). The actual ratio in the inoculum was determined by plating the suspension of each strain before mixing. Mice were sacrificed 24 h after inoculation and brain, heart, liver, spleen, lung and kidney samples were collected, homogenized and diluted for plating. Blood samples were directly diluted for plating. The ΔyefM-yoeB:WT ratios in these samples were determined by analysing 70–90 colonies using colony PCR with primer pair ATout1/ATout2, which yielded PCR products of 597 bp and 1117 bp for the ΔyefM-yoeB and WT strains, respectively. The competitive index (CI) was calculated as the ΔyefM-yoeB:WT ratio in each sample divided by the ratio in the inoculum.

Bioinformatics and statistical analysis. Multiple sequence alignments were processed using ClustalW2 (http://www.ebi.ac.uk/Tools/msa/clustalw2/), and the modeled structures of S. suis YefM and YoeB were generated using CPHmodels 3.2 Server (http://www.cbs.dtu.dk/services/CPHmodels), which searches for templates from known structures. The promoter of the yefM-yoeB locus was predicted by BPROM (http://linux1.softberry.com/berry.phtml).

Statistical analyses were carried out using GraphPad Prism 5 (San Diego, USA). Log-rank test was used to analyse the mice survival curves. Two-tailed paired t test was used to analyse the data in the competitive-infection assay. A P value of <0.05 was considered statistically significant.

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**Author Contributions**
C.Z., J.X., H.C. and W.B. conceived and designed this project and experiments. C.Z., J.X., S.R., J.L. and M.X. performed the experiments. C.Z. and J.X. analysed the data and contributed to the development of the figures and tables. C.Z. wrote this manuscript.

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