Ribose operon repressor (RbsR) contributes to the adhesion of Aeromonas hydrophila to Anguilla japonica mucus

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Abstract
The characterization of adhesion between pathogenic bacteria and the host is critical. Pathogenic Aeromonas hydrophila was shown to adhere in vitro to the mucus of Anguilla japonica. To further investigate the adhesion mechanisms of A. hydrophila, a mini-Tn10 transposon mutagenesis system was used to generate an insertion mutant library by cell conjugation. Seven mutants that were impaired in adhesion to mucus were selected out of 332 individual colonies, and mutant M196 was the most severely impaired strain. National Center for Biotechnology Information (NCBI) blast analysis showed that mutant M196 was inserted by mini-Tn10 with an ORF of approximately 1,005 bp (GenBank accession numbers KP280172). This ORF is predicted to encode a protein consist of 334 amino acid, which displays the highest identity (98%) with the RbsR of A. hydrophila ATCC 7966. Random inactivation of rbsR gene affected the pleiotropic phenotypes of A. hydrophila. The adhesion ability and the survival level of the rbsR gene mutant (M196) were attenuated compared with the wild-type and rbsR complementary type. The findings of this study indicated that RbsR plays roles in the bacterial adhesion and intracellular survival of A. hydrophila.

KEYWORDS
A. hydrophila, adhesion, intracellular survival, Mini-Tn10Km, rbsR

1 | INTRODUCTION

Adhesion to the host external surfaces and tissues by pathogenic bacteria represents the first crucial step in most infections, and it had been regarded as an important virulence factor in colonizing the host surface for proliferation and for the release of virulence factors of many pathogenic bacteria (Grześkowiak, Collado, Vesterlund, Mazurkiewicz, & Salminen, 2011; Huang et al., 2016; Wang et al., 2015). The ability of pathogens to attach to the host mucus, cells, and other tissues is mediated by adhesions that can cause many infectious diseases or other harm (Hori & Matsumoto, 2010). Therefore, the inhibition of bacterial adhesion to the host could be useful therapeutically (Acord, Maskell, & Sefton, 2005). It is important to understand the mechanisms of bacterial adhesion as well as the major factors controlling this attachment. Recently, the adhesion of many clinical pathogens to their hosts has been elucidated, and bacterial adhesion was shown to be mostly influenced by motility, cell hydrophobicity, and bacterial surface structures (Chen, Yan, Wang, Zhuang, & Wang, 2008; He et al., 2015; Losensky, Vidakovic, Klingl, Pfeifer, & Fröss, 2015). Experiments with the pathogens Pseudomonas aeruginosa, Vibrio alginolyticus; Vibrio vulnificus, and Legionella pneumophila (Huang, Hu et al., 2015; Kim & Rhee, 2003; Luo et al., 2016; Olga, Jens, & Michael, 2011; Shao et al., 2013) demonstrated the important role of flagella or lipopolysaccharides in establishing the initial interaction with mucosal surfaces or cells and that...
this defect can reduce the ability of adhesion. In addition, the genetic basis of colonization has been studied in V. vulnificus, L. pneumophila, and Streptococcus suis (Chang et al., 2005; Kim & Rhee, 2003; Zhang et al., 2015) and in the genes that impact bacterial adhesion that can also contribute to bacterial pathogenesis. In the pathogenic bacteria V. vulnificus, FlgC was shown to be involved in adhesion to HeLa cells, and the flgC mutation could decrease the level of adhesion, cytotoxicity, and lethality in mice (Kim & Rhee, 2003). Intracellular bacteria L. pneumophila was shown to be involved in adhesion to the human lung alveolar epithelial cell line A549, and the laIA gene mutation could cause reduced mortality in A/J mice (Chang et al., 2005). The dnaJ of S. suis Type 2 was important for adhesion to host cells, and the dnaJ mutation could result in reduced adhesion to cultured cell monolayers (Zhang et al., 2015). These pathogenic genes were all considered to be the key factors for bacterial adhesion and pathogenicity.

Aeromonas hydrophila has been recognized as a gram-negative bacillus of the Aeromonadaceae family, and it can survive in a wide variety of aquatic systems (Cao et al., 2012). It is an opportunistic pathogen and can infect a great variety of poikilothermic and homeothermic animals, including humans (Krovacek et al., 1994; Mccoy et al., 2010; Sime-Ngando, 2015). Several structures have previously been demonstrated to be related to the pathogenicity of A. hydrophila, including adhesion (Grześkowiak et al., 2011; Hori & Matsumoto, 2010; Van Der Marel, Schroers, Neuhaus, & Steinhagen, 2008). A. hydrophila attached strongly to the host surface using flagella-promoted motility in the optimum environment, and the environmental factors obviously affected bacterial adhesion (Benhamed, Guardiola, Mars, & Esteban, 2014). Several genes essential for A. hydrophila adhesion have been identified. The A. hydrophila flgE gene mutation can cause the absence of flagella, and the mutant bacteria can exhibit inadequate motility, adhesion, invasion, and survival in host macrophages when compared with the wild-type B11 and the complementary strain (Qin et al., 2014). Single gene mutations of minD can lead to flagella deficiency and adhesion reduction (Huang, Qin et al., 2015).

It has been extensively reported that some genes associated with flagella play an important role in A. hydrophila adhesion. However, few studies have focused on fully exploring the adhesion-related genes of the pathogenic bacteria A. hydrophila, especially genes related to nutrient metabolism or enzymes. The aim of this study was to identify more adhesion-related genes of pathogenic A. hydrophila with random gene inactivation method and to probe other pleiotropic phenotypes of A. hydrophila by constructing mutant and complementary strains to study the mechanisms of these genes.

2 | EXPERIMENTAL PROCEDURES

2.1 Bacterial strains and growth conditions

The bacterial strains and plasmids used in this study are listed in Table 1. Pathogenic A. hydrophila was isolated from the naturally infected Anguilla japonica. Matrix-Assisted Laser Desorption/Ionization Time of Flight Mass Spectrometry (MALDI-TOF-MS) and sequencing of 16S rRNA, and confirmed as a pathogen by artificial infection. A. hydrophila and Escherichia coli were grown in Luria-Bertani Miller broth (LB) at 28°C and 37°C, respectively. The bacterial sample was centrifuged at 4,000 g at 4°C for 5 min, and bacterial cells were resuspended in phosphate-buffered saline (PBS, pH 7.4) after incubation for 18 hr. The medium was supplemented with the appropriate antibiotics at the following concentrations: 600 μg/ml kanamycin (Km),

| Strain or plasmid | Characteristic(s) | Source or reference |
|-------------------|-------------------|---------------------|
| **Strains**       |                   |                     |
| A. hydrophila     |                   |                     |
| W1                | wild-type strain  |                     |
| M01 ~ M332       | mini-Tn10Km insertion mutant (SmR KmR) | This study |
| M196              | rbsR: mini-Tn10Km (SmR KmR) | This study |
| MC196             | WM196 complemented with pACYC184-rbsR (CmR) | This study |
| **E. coli**       |                   |                     |
| SM10              | thi-1 thr-1 leu-1 tonA lacY supE recA RP4-2-Tc::Mu::Km (λpir) F-, 80dlacZ ΔM15, Δ(lacZYA-argF) U169, deoR, recA1 | (Herrero et al., 1990) |
| E. coli DH5α      | endA1, hsdR17 [K−, mK+], phoA, supE44, λ−, thi-1, gyrA96, relA1 |                     |
| **Plasmid**       |                   |                     |
| pMD18-T           | Cloning vector (ApR) | Takara |
| pLOF/Km           | Tn10-based delivery plasmid with (KmR ApR); | (Herrero et al., 1990) |
| pACYC184          | (CmR TcR) | provided by Prof. Nie |
| pACYC184-rbsR     | pACYC184 derivative containing 1,005 bp fragment of rbsR putative promoter and ORF(CmR) | This study |

*TABLE 1* Strains and plasmids used in this study
100 μg/ml ampicillin (Amp), 100 μg/ml streptomycin (Sm), and 34 μg/ml chloramphenicol (Cm).

2.2 | Preparation of *A. japonica* mucus and *A. hydrophila* adhesion in mucus in vitro

A group of five healthy *A. japonica* were purchased from an aquatic product market in Xiamen of Fujian Province. The skin mucus was prepared using a modification of a previously described technique (Huang, Qin et al., 2015). Briefly, the skin mucus gel was collected by scraping the skin surfaces with a rubber spatula, and the gel was homogenized in PBS and centrifuged twice at 20,000 g at 4°C for 30 min to remove particulates. The supernatant was filtered sequentially through .45 and .2 μm pore size filters. The resulting mucus sample was adjusted to 1 mg of protein/ml with sterile aged seawater, and the protein concentration was determined using the method that was described by Bradford (Bradford, 1976).

Gill mucus was prepared using a technology we described before (Chen et al., 2008). Blood was completely removed from the caudal vessel, and the gill arches were excised and soaked in PBS for 2 hr at 4°C, with occasional shaking. The mucus preparation was centrifuged twice at 20,000 g for 30 min at 4°C to remove particles and cellular material, followed by filtration of the final supernatant through .45 and .2 μm pore size filters. The protein concentration of mucus preparation was adjusted to 1 mg/ml in PBS. The protein concentration was determined using the method of Bradford (1976).

Intestinal mucus was prepared using a technique we described before (Yan, Chen, Ma, Zhuang, & Wang, 2007). The fish were starved for 48 hr without feeding and the intestines were then removed and divided into foregut and hindgut according to intestinal morphology. The guts were transferred to sterile petri dishes and washed by sterile .01 mol/L PBS (pH 7.2). Next, they were split open with a scalpel, and the foregut and hindgut mucus were collected by scraping of the inner surface of the intestines with a rubber spatula to remove the layer of mucus gel covering the intestinal lumen, respectively. The mucus was homogenized in .01 mol/L PBS (pH 7.2). The mucus preparations were centrifuged twice at 20,000 g for 30 min at 4°C to remove particulate and cellular material, followed by filtration of the final supernatant through .45- and .2-μm pore size filters. The mucus samples were adjusted to 1 mg protein/ml with PBS. The protein concentration was determined using the method of Bradford (1976).

An in vitro adhesion assay was performed using an indirect enzyme-linked immunosorbsent assay (ELISA) method with 96-well microtiter polystyrene plates, as previously described and modified (Qin et al., 2014). Briefly, 100 μl of the mucus samples were added to each well, the mucus was fixed by incubation overnight at 4°C, and unbound mucus was removed by washing the wells twice with 200 μl of PBS. Equal volume aliquots of bacteria were added to the wells and incubated at 28°C for 2.5 hr. Then, adhered bacterial cells to mucus were removed by washing the wells twice with 200 μl of PBS, and the bacteria were dried at 60°C. The wells were washed three times with 200 μl of phosphate-buffered saline Tween-20 (PBST), and all wells were blocked with fetal bovine serum (1:10). Subsequently, a sample (100 μl) of the indicated dilution of rabbit antibacterial serum in PBS was added to each well after washing the wells three times with 280 μl of PBST, incubating for 1 hr at 37°C, and washing three times with PBST. Horseradish peroxidase (HRP) goat anti-rabbit immunoglobulin G diluted (1:1,000) in PBS was then added to each well and incubated for 1 hr at 37°C. After washing four times with PBST, 100 μl of a fresh OPD-H₂O₂ was added to each well. The color was allowed to develop for 30 min and was then measured at 492 nm with a microtiter plate reader (Bio-Tek). Controls consisted of wells without bacteria to ensure that the primary antibody did not cross-react with the endogenous bacteria or with the host cells and wells without the immobilized cells to determine the amount of nonspecific binding of the bacteria to the plastic.

To further confirm the result of bacterial adhesion assay, a total of 50 μl of mucus was evenly spread on a 22 mm × 22 mm glass slide area and fixed with methanol for 20 min. Then, 1 ml of bacterial suspension (10⁸ CFU/ml) was placed on the mucus-coated glass slides, incubated for 2 hr at 25°C in a humidified chamber, and washed with PBS five times. Lastly, the bacteria were fixed using 4% methanol for 30 min, dyed via crystal violet for 3 min, and counted under a microscope (×1,000). Each group was conducted in three trials, and 20 fields of view were selected. Negative control was performed using PBS instead of bacterial suspension.

2.3 | Mini-Tn10 mutagenesis of *A. hydrophila* strain W1 and genetic screening for adhesion-deficient mutants

The mini-Tn10 Km transposon was transformed into *A. hydrophila* wild strain W1 on the suicide vector pLOF/Km carried by *E. coli* Sm10 through a filter mating technique that had been previously developed (Qin et al., 2014). Briefly, the suicide vector mini-Tn10 Km (KmR and AmpR) was extracted from *E. coli* CC118 and transferred into *E. coli* Sm10. Then, the *A. hydrophila* wild-type (W1) and *E. coli* Sm10 pLOF/Km (KmR and AmpR) were combined with Isopropyl β-D-1-thiogalactopyranoside (IPTG) induction. Serial dilutions of the mating mixture were plated on LB supplemented with kanamycin to select the mutants. The mutants in the genes were identified by gene walking polymerase chain reaction (PCR, Takara).

2.4 | Southern blotting

A single insertion event in the rbsR::Km mutant was confirmed. A single chromosomal transposon insertion in the rbsR::Km mutant was confirmed with genomic DNA prepared from the rbsR::Km mutant using the standard method (Rock & Nelson, 2006). Briefly, DNA was extracted from the bacterial cells by proteinase K-sodium dodecyl sulfate digestion followed by phenol-chloroform purification. Approximately 4 g of genomic DNA of wild type and rbsR::Km mutant were digested with Sal I. A total of 10 μg of DNA was digested to completion with Sal I and run on agarose gels. Southern blots were prepared on Hybond nylon filters. Oligonucleotide probes end-labeled with a Dig-High Prime DNA Labeling and Detection Starter Kit II.
were used for detection according to the manufacturer’s protocol. A 176-bp STM PCR product was amplified using primers Tn10kan (CGGGGATCCGAGTGG) and Tn10kan (TGGAAGCCGCGATGC) and labeled using a Dig-High Prime DNA Labeling and Detection Starter Kit II. The probe was hybridized to the genomic DNA on the membrane and detected using the Dig-Highreagents in accordance with the manufacturer’s instructions.

2.5 | Sequence analysis of the mini-Tn10 insertion mutants

Thermal asymmetric interlaced polymerase chain reaction (TAIL-PCR) (Singer & Burke, 2003) was used to obtain the mutant DNA sequences flanking the transposon to find the insertion sites in *A. hydrophila*. The arbitrary primers were provided by a genomic walking kit (Takara, Bio). The specific transposon nested primers are listed in Table 2. LSP4, LSP5, and LSP6 were used to amplify upstream sequences flanking the transposon, and RSP4, RSP5, and RSP6 were used to amplify downstream sequences flanking the transposon. TAIL-PCR products were purified with a Gel Extraction Kit (Omega, Bio-Tek), cloned into a pMD18-T vector (Takara, Bio) and sequenced. DNA sequences were analyzed by BLASTN (National Center for Biotechnology Information, Bethesda, MD, USA) and other software such as DNA Star.

2.6 | Plasmid complementation

For complementation studies, *A. hydrophila* wild-type DNA fragments with *rbsR* were PCR amplified using primers *rbsR*-EcoRV-For (5′-GAT ATC CTA ACC AGA GAG GGT GGA ATA GTG C-3′) and *rbsR*-2HA-Sphl–Rev (5′-GCA TGC TTA CTA GAG GCT AGC ATA ATC AGG AAC ATC ATA CGG ATA CCG CAG TGA CTT GAC GAC GC-3′) (enzyme restriction sites are underlined, the HA-tag sequence is in italic and the stop codon is underlined and in italic). Amplified fragments of 1,005-bp were ligated into pMD18-T (Takara). The DNA insert was recovered by BamH I and Sphl restriction digestion and was ligated into BamH I and Sphl phosphatase-treated pACYC184 to generate plasmid pACYC-rbsR. This vector was transferred into *E. coli* DH5α, and recombinant plasmids surviving on chloramphenicol plates were selected by tetracycline sensitivity. Then, the recombinant plasmids were electroporated into a *rbsR* mutant strain, and bacteria complementary strains surviving on chloramphenicol plates were selected from tetracycline sensitivity.

Whole-cell proteins were obtained from Aeromonas strains grown at 28°C. Equivalent numbers of cells were harvested by centrifugation, and each cell pellet was suspended in 50–200 μl of sodium dodecyl sulfate-polyacrylamide gel electrophoresis loading buffer and boiled for 10 min. The expression of *rbsR* was detected by western blot according to the supplier’s instructions with an anti-HA tag antibody that was appropriate for the method (Green & Sambrook, 2012). Following sodium dodecyl sulfate-polyacrylamide gel electrophoresis and transfer to nitrocellulose membranes, the membranes were blocked with skimmed milk powder (5 mg/ml) and probed with hemagglutinin (HA)-tag antibodies (1:1,000). The unbound antibody was removed by three washes in PBST, and a goat anti-rabbit HRP-conjugated secondary antibody (1:100) was added. The unbound secondary antibody was removed by three washes in PBST. The bound conjugate was then measured using a Super Signal West Pico Trial kit (Thermo, USA) and electrochemiluminescence (ECL) western blot system (LAS-4000 mini, Fuji, Japan) according to the manufacturer’s instructions.

2.7 | Preparation of macrophages, and in vitro invasion and survival

Tilapia macrophages were prepared, and in vitro invasion and survival of *A. hydrophila* in tilapia macrophages were conducted. Healthy tilapia (*Oreochromis niloticus*) individuals were obtained from an aquaculture farm. *O. niloticus* macrophages were prepared as Qin et al. (Qin et al., 2014) described with some modification. Briefly, *O. niloticus* was bled by syringe as completely as possible from the caudal circulation to reduce the blood volume in the pronephros, and then the surface of the fish was sterilized with 70% alcohol. The tissue overlaying the pronephros was removed and then placed in 2 ml of ice-cold L-15 medium (HyClone Laboratories, Inc., Utah, USA) supplemented with 100 IU S/P ml⁻¹ and 2% fetal bovine serum. The tissues of pronephros were sheared and pushed through a 100 μm nylon mesh. The cell suspension was layered onto a 34%/51% discontinuous Percoll (GE Healthcare Bio-Sciences AB, Sweden) density gradient with a syringe and centrifuged at 400 x g for 30 min at 4°C. The band of cells lying at the interface of the 34%/51% Percoll gradient was collected, washed twice and the living and dead cells were counted by trypan blue stainning. Then, the cells were counted with hemocytometers and adjusted to 2.4 x 10⁷ cells ml⁻¹ in a L-15 medium with 5% FBS, 100 IU S/P ml⁻¹ and 10 IU heparin ml⁻¹ and transferred to 6-well plates, 1 ml/well.

In vitro bacterial invasion tests were performed as previously described (Martin & Mohr, 2000) with some modification. Briefly, this assay was performed according to the following steps. First, 1 ml of the macrophage suspension was added to each well of 6-well culture plates and incubated for 0.5 hr. Then, an equal volume of bacterial suspension with a multiplicity of infection of 100 (100 bacteria per macrophage added) was added to each well and incubated at 28°C. After 1 hr of bacterial invasion, the cell suspensions were treated with 2,500 μg of gentamicin ml⁻¹ for 20 min at 28°C to kill the extracellular bacteria, and the supernatant was carefully removed without disturbing the packed cells. Then, fresh L-15 medium with 10 IU heparin ml⁻¹, 10% FBS and 100 units S/P was added to the packed cells and allowed

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**TABLE 2** The primer specific primers used in TAIL-PCR

| Primer | Sequence |
|-------|----------|
| LSP4  | 5′ ATGCTTATGTCGAGAAGGGA 3′ |
| LSP5  | 5′ CATGCGGCTTCACATACAGG 3′ |
| LSP6  | 5′ ATTATCGGAGGCCATTATACCC 3′ |
| RSP4  | 5′ CCTGTGGAAACAGTGTCGGAAAGA 3′ |
| RSP5  | 5′ GATTTTGGCCTTCATTGACCTG 3′ |
| RSP6  | 5′ TTACGCTACCTGAGCGGCAGG 3′ |
to incubate at 28°C in 5% CO₂ with the time point denoted as 0 hr. After incubation for 1 hr, the cells were centrifuged for 5 min at 400 g at 28°C, and the supernatant was aspirated. Then 1 ml of sterile distilled water was added for 30 min to lyse the cells. The CFU number of the cell lysate was determined by plate counting.

2.8 | Statistical analyses

All data were statistically analyzed with SPSS16.0 (SPSS, Chicago, IL, USA). The mean ± standard deviation was calculated for each sample. Unless otherwise stated, all experiments were performed at least three times in triplicate assays.

3 | RESULTS

3.1 | Adhesion-reduced mutant isolation

A mini-Tn10 transposon mutagenesis system was used to generate an insertion mutant library by cell conjugation. A mutagenesis library with 332 random insertion mutants of A. hydrophila was constructed, and each mutant in the library was subjected to the adhesion assay. Seven mutants were found to be reduced in adhesion to A. japonica skin mucus compared with W1, and the mutant strain (M196) was the most significantly inadequate regarding adhesion to A. japonica mucus (Figure 1).

3.2 | Southern blotting

Southern blotting analysis showed that a single band was present in all mutants and the plasmid positive control pLOF/Km. No signal was detected in the negative control wild-type strain W1 (Figure 2), which confirmed that the mutations were caused by the insertion of transposon mini-Tn10Km and that only a single transposon insertion was present in the chromosome for each mutant.

3.3 | Mini-Tn10 insertion site in the mutant strain (M196)

Sequence analysis showed that the mini-Tn10 insertion site was a 1,005 bp ORF that shared the highest identity (98%) with the rbsR gene of A. hydrophila ATCC 7966 (accession no: CP000462.1). This rbsR gene encoded a ribose operon, which is a repressor protein that was predicted to contain 334 amino acid residues with a predicted 33 kDa molecular weight. The Tn10 insertion site is indicated with a bold and enlarged letter (Figure 3).

3.4 | RbsR protein expression in the complementary strain

After the recombinant expression plasmid pACYC184-rbsR was introduced into the mutant (M196), the expression of protein RbsR in the complementary strain (MC196) was detected by western blotting. Western blotting analysis showed that a positive signal of approximately 33 kDa was detected, which confirmed that the fusion protein RbsR-HA was expressed in the complementary strain (MC196) but not in the mutant strain (M196; Figure 4).

3.5 | Adhesion assay

Adhesion of the wild-type (W1), the rbsR mutant strain (M196), and the complementary strain (MC196) to the host skin, intestinal and gill mucus was detected. The results showed that the adhesion of the M196 bacteria to skin, intestinal and gill mucus was approximately 15%, 46% and 14%, respectively, compared with that of the wild-type (W1) strain. The adhesion of bacteria to the skin, intestinal and gill mucus with the complementary strain recovered to more than 43%, 59%, and 35% of that of the wild-type strain, respectively (Figure 5). These data suggested that the mutation in rbsR significantly affected bacterial adhesion.

3.6 | Intracellular survival in macrophages

Given the importance of rbsR in A. hydrophila virulence, the number of invasive and surviving bacteria of the wild-type (W1), rbsR mutant (M196), and complementary strains (MC196) in macrophages was determined in vitro. The results at 0 hr show the number of invasive bacteria into macrophages, and the results at 1 hr represent the number of surviving bacteria after entry into the macrophages for 1 hr. The data showed that there was no difference between the number of invasive wild-type bacteria, mutant strain and that of the
complementary strain, while the survival number of the mutant strain (M196) was significantly lower than that of the wild-type (W1) and the complementary strain (MC196) after 1 hr (Figure 6). Furthermore, there was only approximately 17% of invasive mutant bacteria that survived in macrophages 1 hr after entry into the cells, while the survival rates of wild-type (W1) and the complementary strain (MC196) were ~26% and ~31%, respectively, which suggested that the survival of the mutant bacteria in host macrophages were seriously impaired.

4 | DISCUSSION

Recently, adhesion has been found to be associated with the virulence of many types of pathogens (Benhamed et al., 2014; Luo et al., 2016). Some studies have suggested that bacterial adhesion genes, such as flrA, flrB, and flrC can mostly be related to flagella. Bacterial adhesion is an extremely complicated process and can be controlled by one or more genes. To understand the genetic basis of adhesion in the A. hydrophila prototype, transposon insertion mutants that were inadequate in adhesion isolated were characterized. In this study, pathogenic A. hydrophila W1 could adhere to A. japonica mucus well, and a gene corresponding to a repressor protein was identified: rbsR (M196).

Previous studies have found a few genes, such as CtbP, which is not traditionally considered as an adhesion factor, that can impact adhesion (Grooteclaes & Frisch, 2000). The results confirmed the screening data in this study.

In this study, the mutant strain (M196) was significantly inadequate regarding adhesion to A. japonica mucus and the complementary strain (MC196) recovered the adhesion ability to some extent. rbsR deletion can immediately affect the mRNA level of the whole rbsRACBD (rbs) operon, and the gene products of the rbs operon of Corynebacterium glutamicum encode a ribose-specific ATP-binding cassette (ABC) transport system with corresponding regulatory proteins (i.e., rbs). ABC transporters are membrane proteins that couple the energy from ATP hydrolysis to transport substrates, including nutrients and toxins (Chen, Lu, Lin, Davidson, & Quirocho, 2003; Nentwich et al., 2009). Previous reports have shown that NCAM-mediated adhesions may be modulated by extracellular ATP (Skladchikova, Ronn, Berezin, & Bock, 1999). Potassium requires ATP to stimulate the adhesion of P. aeruginosa; ATP synthesis may play an important role in the adhesion process (Espinosa-Urgel, Salido, & Ramos, 2000). Mostly, nutrients influence the expression of adhesion and invasion proteins in Listeria monocytogenes, and a starvation assay that inhibited V. alginolyticus adhesion to the fish mucus showed that the nutrients were important to bacteria adhesion (Chen et al., 2008; Jaradat & Bhunia, 2002). It can be deduced that RbsR may affect the expression of ABC transporters that couple the energy from ATP hydrolysis to transport nutrients and that the nutrients can affect the bacterial adhesion. Although specific

FIGURE 2 Detection of a mini-Tn10 insertion in the wild-type strain (W1) and the mutants by Southern blotting; a single band was present in all mutants and in the plasmid positive control pLOF/Km.
mechanisms on how RbsR functions in ATP and some adhesion processes remain uncertain, our findings provide an important new understanding of this process.

Once bacteria have irreversibly attached to a surface, the process of biofilm maturation begins (Dunne, 2002). Cells within the biofilm can modulate cytokine synthesis and can remain hidden from antibodies and complement factor recognition and from subsequent host cell phagocytosis (Cortés, Consuegra, Sinisterra, & Mendez-Vilas, 2011; McCann, Gilmore, & Gorman, 2008). Brucella suis releases compounds, such as adenine or GMP, that can inhibit phagocyte functions and contribute to the intracellular survival of the bacteria (Foulongne, Bourg, Cazevieille, Michaux-Charachon, & O’Callaghan, 2000). RbsR protein has been believed to be an ordinary LacI-type repressor for a single target rbsDACBK operon, which may be involved in the production of GMP in nucleotide synthesis (Shimada, Kori, & Ishihama, 2013). Mostly, the E. coli rbsR gene was found to be a transcriptional repressor for the ribose operon that is involved in the regulation of purine nucleotide synthesis and nutrient synthesis and transport (Daigle, Graham, & Curtiss, 2001; Shimada et al., 2013). In this study, the data suggested that the survival ability of the rbsR mutant in host macrophages was severely impaired. This impairment could be related to the result that Salmonella typhi rbsR mutations could decrease the level of survival within human macrophages at both 2 and 24 hr after bacterial infection (O’Callaghan, Maskell, Liew, Easmon, & Dougan, 1988). We hypothesized that the A. hydrophila rbsR mutation could affect nutrient synthesis and transportation causing poor survival ability. However, further research is still necessary. Our future research efforts will focus on the strategies that permit A. hydrophila intracellular survival within phagocytes.

In summary, our study showed that rbsR mutations affected A. hydrophila behavior related to virulence, such as adhesion and intracellular survival ability.

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CONFLICT OF INTEREST

The authors declare no potential conflicts of interest.

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