Deletion of AS87_03730 gene changed the bacterial virulence and gene expression of *Riemerella anatipestifer*

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*Riemerella anatipestifer* is an important pathogen of waterfowl, which causes septicemia anserum exsudativa in ducks. In this study, an AS87_03730 gene deletion *R. anatipestifer* mutant Yb2ΔAS87_03730 was constructed to investigate the role of AS87_03730 on *R. anatipestifer* virulence and gene regulation. By deleting a 708-bp fragment from AS87_03730, the mutant Yb2ΔAS87_03730 showed a significant decreased growth rate in TSB and invasion capacity in Vero cells, compared to wild-type strain Yb2. Moreover, the median lethal dose (LD50) of Yb2ΔAS87_03730 was 1.24 × 10^7 colony forming units (CFU), which is about 80-fold attenuated than that of Yb2 (LD50 = 1.53 × 10^5 CFU). Furthermore, RNA-Seq analysis and Real-time PCR indicated 19 up-regulated and two down-regulated genes in Yb2ΔAS87_03730. Functional analysis revealed that 12 up-regulated genes were related to "Translation, ribosomal structure and biogenesis", two were classified into "Cell envelope biogenesis, outer membrane", one was involved in "Amino acid transport and metabolism", and the other four had unknown functions. Polymerase chain reaction and sequence analysis indicated that the AS87_03730 gene is highly conserved among *R. anatipestifer* strains, as the percent sequence identity was over 93.5%. This study presents evidence that AS87_03730 gene is involved in bacterial virulence and gene regulation of *R. anatipestifer*.

*Riemerella anatipestifer* is a significant pathogen of waterfowl, turkey and other birds\(^1,2\). It causes epizootic infectious polyserositis in ducks characterized by lethargy, diarrhea, respiratory and nervous symptoms, which led to high mortality and consequently to great economic losses\(^3\). In view of its veterinary importance, many investigations have been carried out on the virulence factors of *R. anatipestifer*, including CAMP cohemolysin, OmpA, nucleoside-diphosphate-sugar epimerase, and glycosyl transferase etc.\(^4-7\). Recently, 49 virulence associated genes were identified by random transponson mutagenesis\(^8\).

The *luxE* gene from a number of bioluminescent bacteria had been well defined by genetic and biochemical analysis, including *Vibrio harveyi*, *Vibrio fischeri*, *Photobacterium phosphoreum*, etc\(^9-11\). The *luxE* gene encodes acyl-protein synthetase (LuxE) which activates the fatty acid, results in the formation of a fatty acyl-AMP intermediate and functions as the second step in the bioluminescent fatty acid reduction system\(^12\). Up to date, most studies focus on the role of *luxE* in bacterial bioluminescence reaction, and the role of *luxE* in *R. anatipestifer* has remained unknown. In this study, a *luxE* homology gene deletion mutant strain Yb2ΔAS87_03730 was constructed by allelic exchange, and the roles of AS87_03730 gene on bacterial growth, adherence and invasion capability, as well as colonization and development during infection were investigated. Furthermore, the function of AS87_03730 on the gene regulation at genome level of *R. anatipestifer* was investigated using RNA-Seq, the differentially expressed genes between mutant strain Yb2ΔAS87_03730 and wild-type strain Yb2 were analyzed. The distribution of AS87_03730 gene in 36 *R. anatipestifer* strains with different serotypes and the gene homology were also analyzed.

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Results

Characterization of mutant strain Yb2ΔAS87_03730. The AS87_03730 gene was deleted from the chromosome of *R. anatipestifer* Yb2 by allelic exchange, which was replaced by a Spec<sup>R</sup> cassette. The mutant strain was screened on TSA plates containing kanamycin (50 μg/ml) and spectinomycin (80 μg/ml), and confirmed by PCR amplification of the AS87_03730 and 16S rRNA fragments from transconjugants (Fig. 1A). Real-time PCR analysis further confirmed that AS87_03730 transcription was abolished in the mutant strain; however, inactivation of the AS87_03730 gene had no effect on the transcription of chromosomally upstream AS87_03725 and downstream AS87_03735 genes (Fig. 1B). The AS87_03730 deletion mutant strain was designated as Yb2ΔAS87_03730.

Mutant strain Yb2ΔAS87_03730 was viable when grown on TSB medium supplemented with kanamycin (50 μg/ml) and spectinomycin (80 μg/ml), however, the growth rate was significantly lower than that of wild-type strain Yb2 (Fig. 1C). When grown on TSA, mutant strain Yb2ΔAS87_03730 formed smooth, slightly raised and non-pigmented colony, which is similar to that of wild-type strain Yb2. Biochemical analysis demonstrated that both mutant strain Yb2ΔAS87_03730 and wild-type strain Yb2 were unable to ferment saccharides (glucose, fructose, galactose, lactose, maltose, sucrose), but produced urease and liquefied gelatin (data not shown). The drug sensitivity of mutant strain Yb2ΔAS87_03730 was also determined, no differences were observed when compared to wild-type strain Yb2 (data not shown). These data suggest that the AS87_03730 gene has no influence on biochemical property and drug sensitivity of *R. anatipestifer*.

Deletion of AS87_03730 gene decreased bacterial adherence and invasion abilities. The adherence and invasion abilities of mutant strain Yb2ΔAS87_03730 to Vero cells were compared with wild-type strain Yb2 to investigate the role of AS87_03730 gene on the attachment and invasion of *R. anatipestifer* to host cells. When infection was performed at a multiplicity of infection (MOI) of 50, the adhered bacteria of mutant strain Yb2ΔAS87_03730 was 1,090 CFU/well, which was slightly decreased in comparison with that of wild-type strain Yb2 (1,456 CFU/well). After additional 1 h of incubation with 100 μg/ml gentamicin, the invaded bacterial counts of mutant strain Yb2ΔAS87_03730 was 140 CFU/well, about 3-fold lower than that of wild-type strain Yb2 (410 CFU/well) (p = 0.0144). The results demonstrated that deletion of AS87_03730 gene has no influence on the bacterial adherence, but decreased bacterial invasion significantly (Fig. 2).

Determination of the bacterial virulence. Bacterial virulence was evaluated by the median lethal dose (LD<sub>50</sub>) using 18-day-old Cherry Valley ducks. The LD<sub>50</sub> for mutant strain Yb2ΔAS87_03730 was 1.24 × 10<sup>7</sup> CFU, which was 80-fold attenuated virulence than that of wild-type strain Yb2 (1.53 × 10<sup>8</sup> CFU). To further investigate the role of AS87_03730 gene on systemic infection *in vivo*, the bacterial loading in blood of infected ducks was quantified. The bacterial recovery of mutant strain Yb2ΔAS87_03730 was 1,255 CFU/ml,
1,845 CFU/ml, and 2,980 CFU/ml at 6, 12 and 24 h post infection (h.p.i.), respectively, whereas the bacterial recovery of wild-type strain Yb2 was 1,580 CFU/ml at 6 h.p.i., $2.36 \times 10^4$ CFU/ml at 12 h.p.i., and $3.80 \times 10^4$ CFU/ml at 24 h.p.i. respectively, indicating that mutant strain Yb2ΔAS87_03730 was drastically attenuated (Fig. 3).

Determination and functional categories of the differentially expressed genes. Differentially expressed genes between mutant strain Yb2ΔAS87_03730 and wild-type strain Yb2 was investigated using strand-specific Illumina RNA-Seq analysis. In total, 31 genes were up-regulated and three genes were down-regulated in mutant strain Yb2ΔAS87_03730, compared to wild-type strain Yb2 (see Supplementary Table S1). Of them, real-time PCR further verified that 19 genes in Yb2ΔAS87_03730 were up-regulated by >2-fold at the transcriptional levels (Table 1). The proteins encoded by these genes were categorized on the basis of their putative functions. The products of up-regulated gene were mostly (63.16%, 12/19) related to “Translation, ribosomal structure and biogenesis”, and two proteins, encoded by AS87_03970 and AS87_06085, were classified into “Cell envelope biogenesis, outer membrane”. The protein encoded by AS87_02350 was involved in “Amino acid transport and metabolism”. There are only two down-regulated genes for hypothetical proteins, and one is involved in “Defense mechanisms”, the other has unknown function. Pathway analysis for the differentially expressed genes was performed according to the latest KEGG database (http://www.genome.jp/kegg/), and the differentially expressed genes were distributed in clusters. Furthermore, the up-regulated genes mostly (63.16%, 12/19) occurred within ribosome pathway (see Supplementary Figure S1), indicating AS87_03730 regulates the genes mainly responsible for translation process.

AS87_03730 gene is highly conserved in R. anatipestifer strains. The AS87_03730 gene was successfully amplified from all 36 R. anatipestifer strains tested. Eight of the PCR products were further subjected to sequence analysis, indicating that Yb2 AS87_03730 gene shares a sequence identity of more than 93.5% with other 7 strains (Fig. 4). In addition, BLAST analysis showed that Yb2 AS87_03730 gene exhibits 100%, 100%, 97%, 97% and 91% identity compared to RA-CH-2 (GenBank accession no. CP004020.1), DSM15868 (GenBank accession no. CP002346.1), CH3 (GenBank accession no. CP006649.1), RA-CH-1 (GenBank accession no. CP003787.1).
and RA-GD (GenBank accession no. CP002562.1), respectively. These data indicated that AS87_03730 gene was highly conserved in *R. anatipestifer*. Moreover, AS87_03730 gene shares 71% similarity to the BD94_0721 gene encoding putative acyl protein synthase/acyl-CoA reductase of *Elizabethkingia anophelis* NUHP1 (GenBank accession no. CP007547.1). The predicted LuxE exhibits sequence identity to acyl transferase from other bacteria, including *Chryseobacterium formosense* (GenBank accession no. KFE99559.1; 72%), *Epilithonimonas* sp. FH1 (GenBank accession no. KFC20122.1; 71%), *Elizabethkingia menigoseptica* (GenBank accession no. WP_026149215.1; 69%) and *Bergeyella zoohelcum* (GenBank accession no. WP_002687639.1; 61%).

| Gene locus | Description of genes | Subcellular location | Function group (COGs) | 2ΔΔCt |
|------------|----------------------|---------------------|-----------------------|-------|
| AS87_05650 | hypothetical protein | Cytoplasmic         | —                     | 4.31  |
| AS87_04530 | 50S ribosomal protein L10 | Unknown               | COG02443             | 4.30  |
| AS87_06160 | 50S ribosomal protein L28 | Cytoplasmic         | COG0227            | 4.25  |
| AS87_03970 | dTDP-4-dehydrohamnose 3,5-epimerase | Cytoplasmic | COG1898M             | 4.19  |
| AS87_07745 | 50S ribosomal protein L23 | Cytoplasmic         | COG0809J             | 3.88  |
| AS87_07845 | 50S ribosomal protein L36 | Unknown               | –                     | 3.76  |
| AS87_06455 | 50S ribosomal protein L21 | Cytoplasmic         | COG0261J             | 3.54  |
| AS87_06085 | dolichyl-phosphate beta-D-mannosyltransferase | Cytoplasmic | COG0463M             | 3.45  |
| AS87_07850 | 30S ribosomal protein S13 | Cytoplasmic         | COG0909J             | 3.31  |
| AS87_07760 | 50S ribosomal protein L22 | Cytoplasmic         | COG0901J             | 3.22  |
| AS87_02350 | cystathionine beta-synthase | Cytoplasmic        | COG0301E             | 2.82  |
| AS87_07855 | 30S ribosomal protein S11 | Cytoplasmic         | COG0100J             | 2.53  |
| AS87_07860 | 30S ribosomal protein S4 | Cytoplasmic         | COG0522J             | 2.24  |
| AS87_07775 | 50S ribosomal protein L29 | Unknown               | –                     | 2.13  |
| AS87_00900 | hypothetical protein | Cytoplasmic         | –                     | 2.08  |
| AS87_07790 | 50S ribosomal protein L24 | Cytoplasmic         | COG0198J             | 2.04  |
| AS87_07830 | 50S ribosomal protein L15 | Cytoplasmic         | COG0200J             | 2.01  |
| AS87_07815 | 50S ribosomal protein L18 | Cytoplasmic         | COG0256J             | 2.01  |
| AS87_07805 | 30S ribosomal protein S8 | Cytoplasmic         | COG0906J             | 2.00  |
| AS87_05175 | 30S ribosomal protein S12 | Cytoplasmic         | COG0408J             | 1.97  |
| AS87_04540 | 50S ribosomal protein L11 | Cytoplasmic         | COG0808J             | 1.88  |
| AS87_05170 | 30S ribosomal protein S7 | Cytoplasmic         | COG0409J             | 1.78  |
| AS87_02865 | hypothetical protein | Cytoplasmic         | –                     | 1.75  |
| AS87_04500 | hypothetical protein | Unknown               | –                     | 1.72  |
| AS87_07455 | membrane protein | Unknown               | –                     | 1.65  |
| AS87_07825 | 50S ribosomal protein L30 | Cytoplasmic         | COG1841J             | 1.65  |
| AS87_07870 | 50S ribosomal protein L17 | Cytoplasmic         | COG0203J             | 1.57  |
| AS87_01795 | membrane-binding protein | Unknown               | –                     | 1.55  |
| AS87_05160 | 30S ribosomal protein S10 | Cytoplasmic         | COG0015J             | 1.55  |
| AS87_04930 | glycosyltransferase | Cytoplasmic         | –                     | 1.29  |
| AS87_02265 | S-adenosylmethionine tRNA ribosyltransferase | Cytoplasmic | COG0809J             | 1.09  |
| AS87_03335 | hypothetical protein | Unknown               | –                     | 0.61  |
| AS87_05990 | hypothetical protein | Unknown               | –                     | 0.51  |
| AS87_03730 (mutated gene) | acyl-protein synthetase | Cytoplasmic         | –                     | 0.01  |

Table 1. Real-time PCR verification of differentially expressed genes in mutant strain Yb2ΔAS87_03730. Functional categories: (1) Information storage and processing: (J: Translation, ribosomal structure and biogenesis; K: Transcription; L: DNA replication, recombination and repair); (2) Cellular processes and signaling: (D: Cell division and chromosome partitioning; V: Defense mechanisms; O: Posttranslational modification, protein turnover, chaperones; M: Cell envelope biogenesis, outer membrane; N: Cell motility; T: Signal transduction mechanisms); (3) Metabolism: (C: Energy production and conversion; G: Carbohydrate transport and metabolism; H: Coenzyme transport and metabolism; P: Inorganic ion transport and metabolism; E: Amino acid transport and metabolism; F: Nucleotide transport and metabolism; Q: Secondary metabolites biosynthesis, transport and catabolism); (4) Poorly characterized: (R: General function prediction only; S: Function unknown).

*Based on *R. anatipestifer* Yb2 genome (accession number: CP007204). Subcellular locations were predicted by the PSORTb Version 3.0 server (http://www.psort.org/). Functional characterization of the proteins was predicted by searching against eggNOG (Version 3) database using BLASTP. No related COG. Results are presented as 2ΔΔCt. Figures = 1 indicated that the gene is expressed similarly in both mutant strain Yb2ΔAS87_03730 and wild-type strain Yb2, figures > 1 indicated that the gene is over expressed in mutant strain Yb2ΔAS87_03730, and figure < 1 indicated that the gene is expressed less in mutant strain Yb2ΔAS87_03730.
Cofactor S-adenosyl-L-methionine. In addition, CBS deficiency leads to changes in protein unfolding. Due to the cofactor pyridoxal-phosphate (PLP) and can be allosterically regulated by effectors such as the ubiquitous component 3-deoxy-L-glycerol-tetronic acid. In R. anatipestifer, the homology gene of luxE is AS87_03730, the prevalence of the gene in R. anatipestifer that do not produce detectable light has led to the question of what role, if any, is there for the activity encoded by the AS87_03730 gene in non-luminescent bacteria. To better evaluate the role of AS87_03730 in R. anatipestifer, we constructed a mutant strain Yb2ΔAS87_03730, in which the AS87_03730 gene was inactivated by deleting a 708-bp fragment from the gene in wild-type strain Yb2. The mutant strain Yb2ΔAS87_03730 showed different growth characteristics in TSB, decreased capacity of invasion, and attenuated virulence in ducks.

R. anatipestifer infection is characterized as septicemia, hence, this species of bacteria have ability to colonize and develop in the tissues of their host. We found that the bacterial loadings in the blood of ducks infected with mutant strain Yb2ΔAS87_03730 was significantly decreased than that of Yb2 infected ducks. In addition, infection with mutant strain Yb2ΔAS87_03730 resulted in a marked delay in mortality, and more than 80-fold attenuated virulence. Our results indicated that the AS87_03730 gene is important for R. anatipestifer to establish a systemic infection. A previous study has revealed that the lux gene loci (luxA, luxI, and luxR) of Vibrio fischeri play an important role in colonization and development of host light organ.

A report has suggested that the products of luxI and luxR control the expression of several non-lux loci, including one required for V. fischeri to remain competitive in mixed symbiotic infections. To investigate whether AS87_03730 gene regulates the expression of other genes in the genome, RNA-Seq analysis was performed to identify differentially expressed genes in mutant strain Yb2ΔAS87_03730. As a result, 31 genes were up-regulated and three genes were down-regulated in mutant strain Yb2ΔAS87_03730. Real-time PCR verification further confirmed 19 genes were up-regulated by over 2-fold. Among them, 12 were classified into "Translation, ribosomal structure and biogenesis". The ribosome is responsible and essential for translation process. The bacterial ribosome consists of three RNA molecules and approximately 55 proteins, components that are put together in an intricate and tightly regulated way. Regulation of gene expression on transcriptional level relies on signals transferred to the RNA polymerase (RNAP) that alter the enzyme's activity or specificity. Normally, rRNA transcription was affected by the stringent response and growth rate regulation. Stringent control leads to a reduction of stable RNA synthesis in response to amino acid starvation, while the growth rate control leads to adjustment of stable RNA synthesis in response to changes in the nutritional quality of growth medium.

In this study, the expressions of 12 rRNA proteins were up-regulated in mutant strain Yb2ΔAS87_03730, indicating that it is attempting to adapt to changing condition as a result of inactivation of AS87_03730 gene. Two up-regulated genes (AS87_03970 and AS87_06085) were involved in "Cell envelope biogenesis, outer membrane". The predicted product of AS87_03970 is a dTDP-4-dehydrodolichomannose 3, 5-epimerase, which is involved in the biosynthesis of dolichyl-phosphate beta-D-mannosyltransferase, which transfers mannose from Dol-P-mannose to Ser or Thr residues on proteins. In Saccharomyces cerevisiae, dolichyl-phosphate beta-D-mannosyltransferase is required in vivo for glycosyl phosphatidylinositol membrane anchoring, O mannosylation, and N glycosylation. In glycoprotein biosynthesis of Candida albicans, dolichyl phosphate mannose synthase is activated by CAMP-mediated protein phosphorylation. Similarly, our results showed that AS87_03730 gene defects in mutant strain Yb2ΔAS87_03730 up-regulated the expression of dolichyl-phosphate beta-D-mannosyltransferase.

One up-regulated gene AS87_02350 encoding cystathionine beta-synthase (CBS) was related to "Amino acid transport and metabolism". CBS, primarily found in eukaryotes, is the first enzyme in the transsulfuration pathway, catalyzing the conversion of serine and homocysteine to cystathionine and cysteine. The enzyme uses the cofactor pyridoxal-phosphate (PLP) and can be allosterically regulated by effectors such as the ubiquitous cofactor S-adenosyl-L-methionine. In addition, CBS deficiency leads to changes in protein unfolding. Due to AS87_03730 gene deletion, the biosynthesis of proteins involved in fatty acid metabolism was stalled in mutant...
Disturbance of fatty acid metabolism consequently resulted in up-regulated expression of AS87_02350. In conclusion, for the first time, we demonstrated that the AS87_03730 gene is involved in bacterial virulence and gene regulation in R. anatipestifer. Further investigation is necessary to clarify the regulation mechanisms of the AS87_03730 gene in R. anatipestifer.

### Methods

#### Bacterial strains, plasmids and culture conditions.

The bacterial strains and plasmids used in this study, and their relevant characteristics, are described in Table 2. R. anatipestifer strains were grown on tryptic soy agar (TSA, Difco, USA) at 37 °C in 5% CO2 or tryptic soy broth (TSB, Difco). Escherichia coli strains were grown at 37 °C on Luria-Bertani (LB) plates or in LB broth. For selective growth of bacterial strains, antibiotics were added at following concentrations: ampicillin (100 μg/ml), kanamycin (50 μg/ml), spectinomycin (80 μg/ml) and chloramphenicol (30 μg/ml). R. anatipestifer serotype 2 strain Yb2 was selected to construct an AS87_03730 gene deletion mutant because serotype 2 is one of the most prevalent serotypes in China and Yb2 is a kanamycin resistant (KanR) and spectinomycin sensitive (SpecS) strain which is good for the further mutant selection.

#### Ethics Statement.

One-day-old Cherry Valley ducks were purchased from Zhuanghang duck farm (Shanghai, China) and kept under a controlled temperature (28 to 30 °C). The ducks were housed in cages with a 12-h light/dark cycle and free access to food and water during this study. This study was carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the Institutional Animal Care and Use Committee (IACUC) guidelines set by Shanghai Veterinary Research Institute, the Chinese Academy of Agricultural Sciences (CAAS). The protocol was approved by the Committee on the Ethics of Animal Experiments of Shanghai Veterinary Research Institute, CAAS (Permit Number: 13–11). All surgeries were performed under sodium pentobarbital anesthesia, and all efforts were made to minimize suffering.

#### Construction of mutant strain Yb2ΔAS87_03730.

An AS87_03730 gene deletion mutant was constructed by allelic exchange through the recombinant suicide plasmid pDS132 as described previously. Briefly,
a Spec cassette (1,119 bp) was amplified from plasmid pFW5 using primers Spec-F and Spec-R. A 1,076 bp left flanking region of AS87_03730 gene was amplified from R. anatipestifer Yb2 genomic DNA using primers AS87_03730 Left-F and AS87_03730 Left-R, and cloned into pGEM®-T easy vector (Promega, Madison, WI, USA). A 1,132 bp of right flanking region of AS87_03730 gene was amplified using primers AS87_03730 Right-F and AS87_03730 Right-R, and cloned into the above-mentioned plasmid; and then a Spec cassette was inserted at the PstI site. The recombinant DNA was moved to SphI- and SalI-digested suicide plasmid pDS132 to produce pDS132-AS87_03730-LSR (Spec cassette). Subsequently, the recombinant plasmid pDS132-AS87_03730-LSR was transformed into R. anatipestifer strain Yb2 by conjugation. The transconjugants were selected by plating on TSA plates containing 50 μg/ml kanamycin and 80 μg/ml spectinomycin. Deletion of AS87_03730 gene in the mutant was confirmed by PCR analysis and resulting mutant strain was designated as Yb2ΔAS87_03730.

**Phenotype characteristics of Yb2ΔAS87_03730.** To determine whether AS87_03730 gene deletion would influence the bacterial growth, the growth curves of mutant strain Yb2ΔAS87_03730 and wild-type strain Yb2 were measured as previously described. Briefly, equal amount of each bacterial culture in mid-exponential phase was transferred into fresh TSB medium at a ratio of 1:100 (v/v) for growing at 37°C with shaking at 200 rpm. Optical density at 600 nm (OD600) was measured at 1 h intervals for 15 h using a spectrophotometer (BIO-RAD, USA). Bacterial growth rate was expressed as OD600 values.

The biochemical test for mutant strain Yb2ΔAS87_03730 and wild-type strain Yb2 was carried out using bacterial biochemical tube (Hangwei, Hangzhou, China) following the manufacturer’s instruction. Both strains were also tested against ten commonly used antibiotic disc (ampicillin, amoxicillin, gentamicin, tetracycline, erythromycin, chloramphenicol, ciprofloxacin, rifampicin, polymyxin B and trimethoprim-sulfamethoxazole). Inoculums from the tested bacteria were prepared depending on Kirby-Bauer antibiotic testing.

**Adhesion and invasion assays.** Adhesion and invasion assays were performed on Vero cells (ATCC CCL-81) in 24-well plates (Corning, NY, USA). The mutant strain Yb2ΔAS87_03730 and wild-type strain Yb2 were grown to mid-logarithmic phase and washed three times with phosphate-buffered saline (PBS) prior to use. Vero cells (approximately 2.5 × 10⁵ cells) were inoculated with treated bacteria at a multiplicity of infection (MOI) of 50, and incubated at 37°C with 5% CO₂ for 1.5 h. The infected cells were rinsed three times with PBS to remove unbound bacteria, and then incubated with 0.1% trypsin (100 μl/well) to lyse the eukaryotic cells. The number of cell-adherent bacteria was counted after dilution and plating. For the invasion assay, the infected cells were further incubated for an additional 1 h with DMEM medium supplemented with 100 μg/ml gentamicin to kill extracellular bacteria. After washed three times with PBS, the infected cells were lysed and the amount of intracellular bacteria was determined. All samples were tested in triplicate, and experiments were repeated three times.

**Determination of the bacterial virulence and survival in vivo.** To determine whether AS87_03730 gene deletion had an influence on virulence of R. anatipestifer, the bacterial median lethal dose (LD₅₀) of mutant Yb2ΔAS87_03730 was determined using 18-day-old Cherry Valley ducks as described and compared with that of wild-type strain Yb2. A total of 64 ducks were divided randomly into eight groups (8 ducks per group) and injected intramuscularly with respective bacterial strain at a dose of 10⁴–10⁸ CFU. Ducks in groups 1–4 were injected with 10⁴, 10⁵, 10⁶ or 10⁷ CFU of Yb2 bacteria, and ducks in groups 5–8 were injected with 10⁵, 10⁶, 10⁷ or 10⁸ CFU of Yb2ΔAS87_03730 bacteria, respectively. The infected ducks which appeared clinical signs (clinical signs include lethargy, diarrhea, rough hair coat, frequent seizure activity, paralysis and no eating or drinking, etc.) were sacrificed humanely with an intravenous injection of sodium pentobarbital at a dose of 120 mg/kg and counted as dead. The infected ducks were clinically observed three times at 8-h intervals daily for a period of 7 days post-challenge. The LD₅₀ value was calculated by improved Karber’s method.

The bacterial loadings in the blood of infected ducks were counted to evaluate bacterial survival in vivo. Twelve 18-day-old Cherry Valley ducks were divided randomly into two groups, and infected intramuscularly with mutant strain Yb2ΔAS87_03730 or wild-type strain Yb2 at a dose of 10⁶ CFU in 0.5 ml PBS. The blood samples were collected at 6 h, 12 h and 24 h post infection, diluted appropriately and plated on TSA for bacterial counting.

**Bacterial RNA isolation.** The mutant strain Yb2ΔAS87_03730 and wild-type strain Yb2 were grown to mid-logarithmic phase, harvested by centrifugation at 10,000 × g for 1 min, and then washed once with PBS for RNA extraction. Total RNA was extracted from bacteria (2.5 × 10⁹ CFU) using Trizol reagent (Invitrogen, Carlsbad, CA, USA), according to the manufacturer’s instructions. All RNA samples were treated with TURBO DNA-free™ kit (Ambion, Grand Island, NY, USA) to remove DNA contamination. Quantification and quality assessment of RNA samples were performed with NanoDrop ND-100 instrument (NanoDrop Technologies, Inc., Wilmington, DE, USA). RNA integrity was checked using 1% agarose gel electrophoresis.

**Illumina sequencing for RNA-Seq.** Total RNA quantification and quality were assessed by spectrophotometer, and illumina RNA-Seq libraries were generated as following steps: 1) RNA fragmentation; 2) reverse transcription; 3) double-strand cDNA synthesis with dUTP; 4) cDNA fragmentation and end-repair; 5) 3’end “da” base addition and illumina adapter ligation; 6) cleavage of the uridine containing strand with Uracil DNA Glycosylase; 7) PCR amplification; 8) size selection and purification. The complete libraries were then quantified with Agilent 2100 Bioanalyzer using DNA 1000 Kit. The DNA fragments in these libraries were denatured with 0.1M NaOH to generate single-strand DNA molecules, loaded onto channels of Illumina flow cell at 8 pM concentration, amplified in situ using TrueSeq SR Cluster Kit v3-cBot HS (Illumina), and subsequently sequenced for 100 cycles on Illumina HiSeq 2000 according to the manufacturer’s instruction.
Differential expression analysis. Image analysis and base calling was performed using Solexa pipeline Version 1.8 (Off-Line Base Caller software, Version 1.8)\(^{35}\). Trimmed reads were aligned to \textit{R. anatipestifer} Yb2 genome using TopHat2 software (Version 2.0.9)\(^{35}\). Transcript levels were calculated as FPKM (fragments per kilobase DNA per million fragments mapped). Differential expressed genes were analyzed using Cufflinks software (Version 2.1.1) with fold change (cutoff = 2.0)\(^{36}\), and considered statistically significant if the fold change was >2.0 and the \(p\)-value was <0.05.

Real-time PCR analysis. Transcriptional levels of differently expressed genes obtained in the transcriptome analysis were further confirmed by real-time PCR. Primers were designed from \textit{R. anatipestifer} Yb2 using primer3 online software Version 0.4.0 (http://bioinfo.ut.ee/primer3-0.4.0/) and described in Supplementary Table 2. RNA samples were extracted and purified as mentioned above. cDNA was synthesized using PrimeScript RT Master Mix (Takara) according to the manufacturer’s protocol, and the resulting cDNAs were diluted 3-fold served as templates. Real-time PCR reaction was carried out in Go Taq qPCR Master Mix (Promega, Fitchburg, WI, USA), and subjected to the Mastercycler ep realplex4 apparatus (Eppendorf, Germany). For each gene, reactions were performed for three RNA samples and each reaction was performed in triplicate. Quantification of transcriptional level was calculated according to the \(2^{-\Delta\Delta C_t}\) method using RA 16S rRNA gene for normalization\(^{37}\).

Distribution and sequence analyses of \textit{AS87_03730} gene in \textit{R. anatipestifer} strains. \textit{AS87_03730} gene in \textit{R. anatipestifer} strains was amplified using PCR. Genomic DNAs of 36 \textit{R. anatipestifer} strains with different serotypes were extracted using TIANamp Bacteria DNA kit (Tiangen, Beijing, China). The primers \textit{AS87_03730} orf-F and \textit{AS87_03730} orf-R were used to amplify the \textit{AS87_03730} sequence using Premix LA Taq\(^{\text{TM}}\) (loading dye mix) (Takara, Dalian, China) according to the following cycle parameters: 94 °C for 4 min, 30 cycles of 94 °C for 40 s, 52 °C for 40 s and 72 °C for 1 min, followed by one cycle of 72 °C for 10 min. To investigate \textit{AS87_03730} homology among different \textit{R. anatipestifer} strains, the PCR products from 8 strains were cloned into pGEM®-T easy vector (Promega, Madison, WI, USA), respectively. DNA sequencing was performed on an Applied Biosystems DNA sequencer (ABI 3730XL) by Thermo Fisher Scientific Inc. (Thermo Fisher Scientific, Shanghai, China). The \textit{AS87_03730} gene sequences from these \textit{R. anatipestifer} strains have been submitted to the GenBank database under the accession numbers KM676074 ~ KM076081. Homology of the \textit{AS87_03730} gene sequences was analyzed with DNASTAR software (DNASTAR Inc., Madison, WI, USA).

Statistical analysis. The statistical significance of the data was determined by the Student’s \(t\)-test in the SPSS Version 17.0 software (SPSS Inc., Chicago, IL, USA). A \(p\) value of < 0.05 was considered to be statistically significant.

Nucleotide sequence accession numbers. The accession number for \textit{R. anatipestifer} strain Yb2 in the GenBank is CP007204

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Author Contributions
Conceived and designed the experiments: S.Y. and X.W. Performed the experiments: X.W., J.Y. and S.W. Analyzed the data: X.W., J.Y., S.Y. and C.D. Contributed reagents/materials/analysis tools: X.W., B.L. and M.T. Wrote the paper: X.W. and S.Y. All authors received the manuscript.

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