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Original article

The *irp2* and *fyuA* genes in High Pathogenicity Islands are involved in the pathogenesis of infections caused by avian pathogenic *Escherichia coli* (APEC)

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Abstract

Avian pathogenic *Escherichia coli* (APEC) is a major bacterial infectious disease that may lead to local or systemic infections in chickens with clinical manifestations. The *irp2-fyuA* gene cluster has been confirmed to be the main genes involved in the synthesis of HPI. The objective of this study was to determine the influence of the *irp2* and *fyuA* genes in the high pathogenicity island (HPI) of avian pathogenic *Escherichia coli* (APEC) on its pathogenicity by knocking out these genes. The Δ AE17 (lacking *irp2*) and $\Delta\Delta$ AE17 (lacking *irp2* and *fyuA*) strains of APEC were constructed. The Δ AE17 and $\Delta\Delta$ AE17 strains showed significantly impaired capacity to adhere onto DF-1 cells. The LD₅₀ results indicated that the virulence of the Δ AE17 and $\Delta\Delta$ AE17 strains was decreased in comparison with that of the AE17 strain. We concluded that the knock-out of the core HPI genes weakened APEC adhesion onto DF-1 cells, inhibited transcription of virulence genes, and reduced pathogenicity in chicks. The effects of genetic deletion of *irp2* and *fyuA* on APEC were more severe than those produced by deletion of *irp2* only, indicating that *irp2* and *fyuA* co-regulate APEC pathogenicity.

Key words: avian pathogenic *Escherichia coli*, *irp2*, *fyuA*, mutant strain, pathogenicity

Introduction

Avian pathogenic *Escherichia coli* (APEC) is a major bacterial infectious disease that may lead to local or systemic infections in birds such as chickens, ducks, and geese, with clinical manifestations including air sacculitis, perihepatitis, pericarditis, omphalitis, peritonitis, encephalitis, and salpingitis. APEC has seriously restricted the development of the poultry industry (Schouler et al. 2012). The pathogenicity

of APEC is manifested as adhesion to host cells and enhanced transcription of virulence factors including adhesion, iron-binding proteins, antiserum factors, outer membrane proteins, and hemolysis (Han et al. 2013).

The *Yersinia* high pathogenicity island (HPI) was first identified by its association with pathogenic phenotypes in mice (Carniel et al. 1996). The HPI is a long chromosomal fragment containing several regulatory genes and the genes related to synthesis,

uptake, and transport of siderophores such as yersiniabactin (Ybt), which is involved in the virulence and pathogenicity of *Yersinia* species (Rakin et al. 1999, Jacobi et al. 2001).

According to the gene sequence analysis, HPI could be divided into two evolutionary lines. The *Yersinia pestis* HPI and the *Yersinia pseudotuberculosis* HPI have a higher similarity in the nucleotide sequences, so the two bacterial HPI is an evolutionary system, referred to as YPs HPI. There are obvious differences in the HPI variable region sequences of *Yersinia enterocolitica*, so it is classified as another evolutionary system, referred to as Yen HPI. YPs HPI and Yen HPI are composed of two parts including the functional core area (about 30.5kb) and the variable region. The functional core area in HPI is highly conserved and up to 98-99% sequence homology. But the variable region in HPI is completely different between YPs HPI and Yen HPI (Carniel et al. 1996).

According to previous studies, HPI in *Yersinia* mediated iron uptake and regulation. At the same time, it is essential for the expression of mouse lethal protein. Although the gene names of the functional core area in YPs HPI or Yen HPI are different, the structure and location of these genes is similar. The gene location of the functional core area in YPs HPI is: 5'-asn-intB-irp9-irp8-irp7-irp6-ybtA-irp2-irp1-irp3-irp4-irp5-fyuA-3'. And the gene location of functional core area in Yen HPI is: 5'-asn-int-ybtS-ybtX-ybtQ-ybtP-ybtA-irp2-irp1-ybtU-ybtT-ybtE-psn-3'. In HPI, the irp2-irp1-irp3-irp4-irp5-fyuA gene cluster has been confirmed to be the main genes involved in the synthesis of Ybt components, called the irp2-fyuA gene cluster. At present, we have explored the function of irp1, irp2 and fyua. The irp1 and irp2 as the ferric uptake regulator gene, are expressed only in pathogenic *Yersinia* and closely related to the virulence. The two genes express the HMWP1(240KD) and HMWP2 (280KD) proteins, respectively. The HMWP1 protein is a multi enzyme complex and mediates the synthesis of no-ribosomal proteins or proteinoid. It is essential to the Ybt positive phenotype. The HMWP2 protein is mainly involved in siderophore and induces the expression of pesticin receptor and ironophore. The fyua gene encodes a 73.7 kD iron-inhibition outer membrane protein which is the co-receptor between Bacterium bacteriocin and Bacillus yersini bacteriocin. The irp3 gene is 1098 bp in size and contrary to the direction of transcription of irp1 and irp2 genes. But the function of the gene is not clear. The irp4 and irp5 genes are the same to direction of transcription of irp1 and irp2 genes and may be related to Ybt synthesis. Studies have shown that the HPI is widely

distributed in various intestinal pathogens. Clinical isolates of 93% of enteroaggregative *E. coli*, 27% of enteroinvasive *E. coli*, 5% of enteropathogenic *E. coli*, and 5% of enterotoxigenic *E. coli* carry the irp2-fyuA gene cluster of the HPI. The sequences of the irp2 and fyua gene in *E. coli* are almost identical to those of *Yersinia* spp. (Schouler et al. 2005). The structural characteristics of the HPI in intestinal pathogens and *Yersinia* species are consistent and show high homology (Bach et al. 2000, Karch et al. 1999), indicating horizontal transmission of the HPI between species of bacteria. The irp2 and fyua genes are the main structural genes in the irp2-fyuA gene cluster, which is the core region of the HPI. The irp2 gene is the primary marker gene for HPI detection and an iron-regulating gene closely related to virulence (Carniel et al. 1996). The fyua gene influences iron absorption and encodes an outer membrane receptor for Ybt, bacteriocin, and *Pasteurella multocida* toxin (Fetherston et al. 1994, Fetherston et al. 1995).

In this study, we used Red homologous recombination to knock out the irp2 and fyua genes in HPI-positive APEC strains to construct the Δ AE17 (lacking irp2) and $\Delta\Delta$ AE17 (lacking irp2 and fyua) strains to assess the influence of key HPI genes on the expression of APEC virulence genes, cell adhesion capacity, and pathogenicity in chicks. The study revealed that the Δ AE17 and $\Delta\Delta$ AE17 strains showed significantly impaired capacity to adhere onto DF-1 cells and transcript levels of luxs, pfs, tsh, iss, ompA, and fimC were significantly decreased compared to those of the AE17 strain. And the virulence of the Δ AE17 and $\Delta\Delta$ AE17 strains was decreased in comparison with that of the AE17 strain which was certified by chick LD₅₀ results. This study laid the foundation for further investigation of the role of genes in the core region of the HPI in the pathogenesis of APEC.

Materials and Methods

Bacterial strains and culture conditions

AE17 was isolated from ducks in An'hui Province, China in 2008. AE17 was identified as *E. coli* stains by biochemical identification and 16S rDNA analysis and then identified as the serotype O2 strain of *E. coli* by serological identification. AE17 was grown on liquid LB medium or on 1.2% solid LB medium at 37°C. Ampicillin (100 mg/ml) and chloramphenicol (34 mg/ml) were added to the medium for the antibiotics group.

Table 1. Primers used in the gene deletion.

Primer	Primer sequence	Size (bp)
irp2-UF	GACTCCCACTGCTGTTGC	760
irp2-UR	GTCAAGCAACCCGGAATTCAGCGAATCCTGAGATGGT	
irp2-DF	GGATTCGCTGGAATTCGGGTTGCTTGACCTCTTCAC	1114
irp2-DR	ATAGCCGACCTTTCTGTT	
pkD3- EcoRI- F	CCGGAATTCTGTAGGCTGGAGCTGCTTCGA	1013
pkD3- EcoRI- R	CCGGAATTCATATGAATATCCTCCTTAG	
irp2-inF	ACAACCATTCGTCCTACTC	
irp2-inR	GTTATCCAGACACAGCCA	
irp2-outF	TTTAAACGCGGGATGTA	
irp2-outR	GAAGGGTGGGTGAAGAG	
fyuAUF	TCAGCCAACAACGTCTCG	551
fyuA-UR	CCCTGGTCGCCGCGTCGACCGCAGTTTCACATTCCCTAT	
fyuA-DF	TGAAACTGCGGTCGACGCGGCCAGGAGGTAAGAGCA	651
fyuA-DR	GGTTATCGACATAGACGGAAAT	
pkD3- Sall – F	CGCGTTCGACTGTAGGCTGGAGCTGCTTCGA	1013
pkD3- Sall – R	CGCGTTCGACCATATGAATATCCTCCTTAG	
fyuA-inF	GCCTATGTGGGATGGAATG	
fyuA-inR	TGCCAGGTCAGGTCCTGTATG	
fyuA -outF	AATGCCCAGACTTCACAGC	
fyuA -outR	CGACGGTCGAACAGGTTA	

Construction and identification of the *irp2* mutant strain Δ AE17

Red homologous recombination was used to construct the Δ AE17 strain lacking *irp2*. The homology arms upstream and downstream of the *irp2* gene were amplified and performed to amplify the *irp2*-Up-Down fragment with *irp2*-Up and *irp2*-Down as templates. The *irp2*-Up-Down fragment was cloned to the zero background vector pZero-back (Tiangen, China) to form recombinant plasmid pZero-Up-Down. The fragment was amplified from the pKD3 plasmid (Yale University Preservation Center, New Haven, CT, USA) flanked by EcoRI restriction enzyme digestion sites. The recovered product and the plasmid were simultaneously digested by EcoRI restriction enzyme (Promega, Madison, WI, USA) and connected to form pZero-Up-cat-Down. Taking pZero-Up-cat-Down as a template, the *irp2*-UF and *irp2*-DR primers were used to amplify the large fragment *irp2*-Up-cat-Down for Red homologous recombination. The purified product was added to competent cells of strain AE17 containing the pKD46 plasmid (Yale University Preservation Center, New Haven, CT, USA). Electroporation was performed using a Bio-Rad Gene Pulser Xcell Electroporation System (200 Ω , 25 μ F, 2.5 kV) at 37°C overnight (Bio-Rad Laboratories, Hercules, CA, USA). Then the cells were inoculated in 100 ml LB medium (Sangon Biotech, China) containing 100 mg/ml ampicillin at a rate of 1:100. After electroporation and culture, the cells were spread on a chloramphenicol-coated plate. Two pairs of primers

irp2-inF/*irp2*-inR and *irp2*-outF/*irp2*-outR were used to identify chloramphenicol-resistant cells to detect whether the *irp2* gene of AE17 was knocked out.

The plasmid (Yale University Preservation Center, New Haven, CT, USA) was electroporated into the chloramphenicol-resistant recombinant cells. The LB plate showing ampicillin/chloramphenicol double resistance was used to culture the recombinants at 28°C. The positive transformants were picked, inoculated in antibiotic-free LB medium, and cultured at 42°C overnight. FLP recombinase expression was induced thermally. By recombination, the sequence between the FRT sites was removed, preserving only one FRT site. PCR using the *irp2*-outF/*irp2*-outR primers was carried out for verification.

Construction and identification of the *irp2/fyuA* mutant strain $\Delta\Delta$ AE17

The $\Delta\Delta$ AE17 strain lacking *fyuA* and *irp2* was created from the Δ AE17 strain by the same method used for knocking out the *irp2* gene. The homology arms upstream and downstream (*fyuA*-Up, 551 bp; *fyuA*-Down, 651 bp) of the *fyuA* gene were amplified with the *fyuA*-UF/*fyuA*-UR and *fyuA*-DF/*fyuA*-DR primer pairs, respectively. Overlap PCR was performed to amplify the *fyuA*-Up-Down fragment with the *fyuA*-UF and *fyuA*-DR primers. The PCR product was cloned into the zero background vector pZero-back (Tiangen, China) to form the pZero-Up-Down plasmid. The chloramphenicol-resistant fragment was amplified from the pKD3 plasmid

Table 2. Primers used in the Real time PCR.

Gene	Primer	Primer sequence (5' to 3')	Size (bp)
dnaE	dnaE-F	GATTGAGCGTTATGTCCGAGGC	80
dnaE	dnaE-R	GCCCCGAGCCGTGAT	
luxs	luxs-F	ACGCCATTACCGTTAAGATG	81
luxs	luxs-R	AGTGATGCCAGAAAGAGGGA	
pfs	pfs-F	CGGCAACAGCCAGGAACTCA	169
pfs	pfs-R	GCGAAAATCCGCCACAATT	
tsh	tsh-F	GCACGAACTGGGAAGTATGGA	118
tsh	tsh-R	GGCATAGAAACCACCACCCC	
ibeA	ibeA-F	TTGTTTTGGCGGAATGATG	118
ibeA	ibeA-R	CATTGATTTTGCCGTTTCTTCT	
stx2f	stx2f-F	GTGTAAAACACTACGCCATCCG	186
stx2f	stx2f-R	AAGCCCAGAACCAGACTCCC	
iss	iss-F	CCGACAGCAGTAACACCAAAGG	105
iss	iss-R	TTCTGCACCGCCAACAAATT	
ompa	ompa-F	TCCAGAGCAGCCTGACCTTC	152
ompa	ompa-R	GCTGAGCCTGGGTGTTTCTCT	
vat	vat-F	TAACGGTATCCGACTTCTGC	171
vat	vat-R	ATGGGGTGGTGGTTTCTATG	
fimc	fimc-F	GCCGATGGTGTAAGGATGG	127
fimc	fimc-R	AACTTTCCCGATCCTGTGGC	

using the primers pkD3-SalI-F/pkD3-SalI-R flanked by SalI restriction enzyme digestion sites. The recovered product and the recombinant plasmid pZero-Up-Down were simultaneously digested by the SalI enzyme (Promega, Madison, WI, USA). The digested chloramphenicol-resistant fragment was connected with pZero-Up-Down to form pZero-Up-cat-Down. Using pZero-Up-cat-Down as a template, the primers *fyuA*-UF and *fyuA*-DR were used to amplify the fragment *fyuA*-Up-cat-Down (2215 bp) for Red homologous recombination. The sequences of the primers are shown in Table 1.

The constructed fragment *fyuA*-Up-cat-Down was electroporated into Δ AE17 cells containing pKD46. Two pairs of primers (*fyuA*-inF/*fyuA*-inR and *fyuA*-outF/*fyuA*-outR) were used for the identification of chloramphenicol-resistant recombinants.

Assay for bacterial adherence

The AE17, Δ AE17, and $\Delta\Delta$ AE17 strains were cultured in liquid LB medium at 37°C until reaching the logarithmic phase. The bacterial solution was centrifuged at 6000 \times g for 5 min at 4°C. The cells obtained were rinsed 3 times with sterile PBS, antibiotic-free DMEM was added (Thermo Scientific, Pittsburgh, PA, USA), and they were shaken well. DF-1 cells (DF-1 is an immortalized cell line of chicken embryo fibroblasts) were cultured in DMEM nutrient solution

containing 10% fetal calf serum and antibiotics. The cells were added to a 24-well plate and cultured at 37°C in an incubator with 5% CO₂ and saturated humidity. When the well was overgrown with cells, the medium was removed and the cells were rinsed 3 times with sterile PBS. Each of the experimental groups with three replicates.

After rinsing, the AE17, Δ AE17, and $\Delta\Delta$ AE17 strains (multiplicity of infection (MOI) = 200) were added to the DF-1 cells. The control group of DF-1 cells was exposed to an equal volume of antibiotic-free DMEM. Each group was assessed in 3 replicates wells. Low-velocity centrifugation was performed to bind the bacteria to the cells, and the cells were cultured in an incubator at 37°C for 1.5 hr. The cells were rinsed 5 times with sterile PBS and 200 μ L 0.1% pancreatin was added to each well to disrupt the cells at room temperature for 10 min. The plate count method was used to count the number of bacteria in the cytosol. The colonies were counted after the cells were cultured on LB plates at 37°C overnight.

RNA isolation, reverse transcription and quantitative real-time PCR

The virulence genes *luxs*, *pfs*, *tsh*, *ibeA*, *stx2f*, *iss*, *ompa*, *vat*, and *fimc* of avian pathogenic *E. coli* were selected for quantitative real-time PCR (qPCR). The housekeeping gene *dnaE* was used to normalize the

amount of RNA across the samples and to assess the reverse transcription efficiency of cDNA (Table 2). Total RNA of AE17, Δ AE17, and $\Delta\Delta$ AE17 strains was extracted from the 3 groups of bacteria separately. The RNA concentration and purity were detected by a Nucleic Acid Protein Detector (ZHD-3, Nanjing Chishun Science & Technology Co., Ltd., Nanjing, China). The DNA-freeTM kit (Ambion[®], Life Technologies, Carlsbad, CA, USA) was used to digest the DNA in the RNA solution. The RNA was uniformly mixed with 0.1 V DNase I buffer and 1 μ L of DNase I, reacted in a 37°C water bath for 30 min, and 0.1 V DNase inactivation reagent was added. The solution was centrifuged at $1000 \times g$ for 2 min, and the supernatant containing the purified RNA was collected. Next, 1 μ L of random primer was added to 2 μ g RNA. The RNA was degenerated and exposed to an ice bath for 1 min, after which 1 μ L of M-MLV, 5 μ L of 5x M-MLV reaction buffer, 4 μ L dNTP mixture, and 1 μ L of RNase inhibitor were added. RNase-free water was added. The mixture was reacted at 42°C for 1 h and stored at -40°C. The cDNA was diluted stepwise and used to plot a 5-point relative standard curve. Reaction system (20 μ L): 10 μ L SYBR Premix Ex Taq (Promega, Madison, WI, USA), 0.5 μ L Primer 1 (10 pmol·L⁻¹), 0.5 μ L Primer 2 (10 pmol·L⁻¹), 1 μ L template, 8 μ L ddH₂O. Each sample had 3 replicates. A melting curve was obtained to analyze the specificity of the primers. Virulence analysis was performed with the 2- $\Delta\Delta$ CT method, and the transcript levels of virulence genes were calculated.

Animals

One-day-old Roman chicks, purchased from Changfeng County, Hefei City, Anhui Province, were reared at 28-30°C with free access to food and water (complete diet without antibiotics) and 12-hr illumination every day. Healthy 7-day-old chicks were selected for the experiment.

Determination of the lethal dose of the bacteria (LD₅₀)

The AE17, Δ AE17, and $\Delta\Delta$ AE17 strains were separately added to liquid LB medium and cultured at 37°C until reaching the logarithmic phase. The cells were collected, rinsed 3 times with sterile PBS, and resuspended. The number of bacteria was regulated and multiple proportion dilution was performed. Each strain was inoculated into 5 groups (n = 8 each) of 7-day-old Roman chicks. The dosages of the AE17 and Δ AE17 strains in the 5 groups were 2×10^8

CFU/chick, 2×10^7 CFU/chick, 2×10^6 CFU/chick, 2×10^5 CFU/chick, and 2×10^4 CFU/chick in the 5 groups. The dosages of the $\Delta\Delta$ AE17 strain in the 5 groups were 2×10^{10} CFU/chick, 2×10^9 CFU/chick, 2×10^8 CFU/chick, 2×10^7 CFU/chick, and 2×10^6 CFU/chick. The control group was treated with sterile PBS. The chicks were infected by intramuscular injection. After injection, the chicks were observed for 7 days and the survival status was recorded. The survival rate was used to calculate the 50% lethal dose (LD₅₀) of the 3 strains based on the Reed-Muench method.

Results

Identification of the mutant strain Δ AE17 and $\Delta\Delta$ AE17

PCR confirmed that the *irp2* knockout strain Δ AE17 was constructed successfully (Fig. 1). The pKD46 plasmid and the target fragment *fyuA*-up-cat-down were successively electroporated into the Δ AE17 strain. PCR was performed to verify the construction of the *irp2/fyuA* knockout strain $\Delta\Delta$ AE17 (Fig. 2).

Comparison of AE17, Δ AE17, and $\Delta\Delta$ AE17 adhesion to DF-1 cells

The adhesion capacity of the AE17, Δ AE17 and $\Delta\Delta$ AE17 strains was 9.04×10^4 , 5.97×10^4 , and 4.84×10^4 CFU/well, respectively. As shown in Figure 3, in comparison with the AE17 strain, the adhesion capacity of the Δ AE17 and $\Delta\Delta$ AE17 strains was significantly decreased (P<0.01). The adhesion capacity of $\Delta\Delta$ AE17 strain was significantly lower than that of the Δ AE17 strain (P<0.05).

Transcriptional analysis of virulence genes in the AE17, Δ AE17, and $\Delta\Delta$ AE17 strains

The melting curve analysis showed that the primers for the reference gene *dnaE* and the virulence genes *luxs*, *pfs*, *tsh*, *ibeA*, *stx2f*, *iss*, *ompA*, *vat*, and *fimC* had high specificity. RT real-time PCR results indicated that the Δ AE17 strain had significantly decreased levels of the *luxs*, *iss*, *ompA*, and *fimC* transcripts (P<0.01), which were 0.32, 0.27, 0.33, and 0.25 times as abundant as those of the AE17 strain, respectively. In the Δ AE17 strain, *pfs* mRNA abundance was significantly decreased to 0.79 times than that of the AE17 strain. However, the abundance of

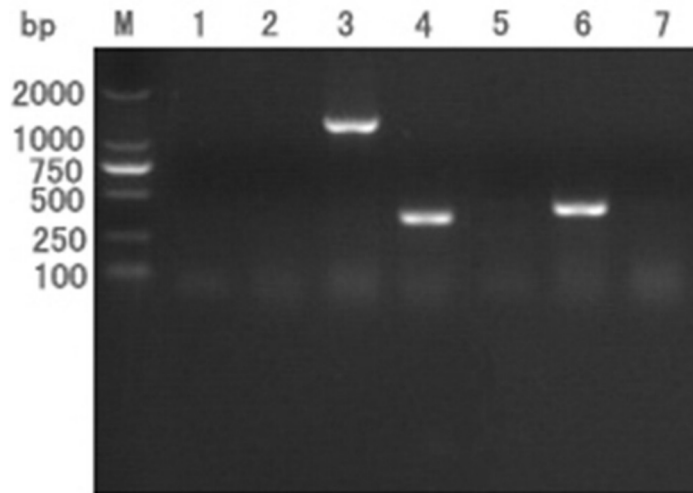


Fig. 1. PCR analysis of the mutant strain Δ AE17. Lane 1: Negative control. Lane 2: The mutant strain Δ AE17, mutant strain showed no PCR product using primers *irp2-outF/irp2-outR*. Lane 3: The Δ AE17, mutant strain Δ AE17 showed a 1282 bp PCR product (Containing the chloramphenicol resistance gene) using primers *irp2-outF/irp2-outR*. Lane 4: The Δ AE17 mutant strain, Δ AE17 showed a 281 bp PCR product (Does not containing the chloramphenicol resistance gene) using primers *irp2-outF/irp2-outR*. Lane 5: Negative control. Lane 6: The mutant strain Δ AE17, mutant strain showed a 418 bp PCR product using primers *irp2-inF/irp2-inR*. Lane 7: The mutant strain Δ AE17, mutant strain showed no PCR product using primers *irp2-inF/irp2-inR*.

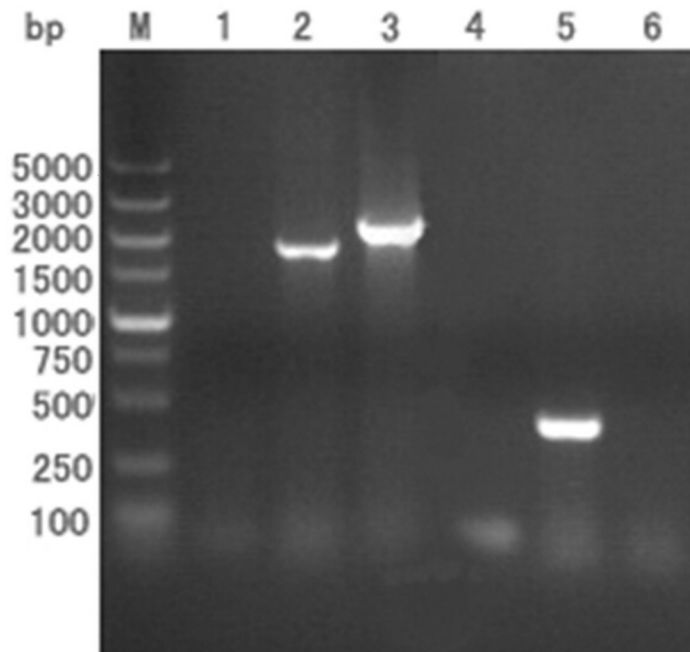


Fig. 2. PCR analysis of the mutant strain Δ Δ AE17. Lane 1: Negative control. Lane 2: The mutant strain Δ Δ AE17 mutant strain showed a 1865 bp PCR product using primers *fyuA-outF/fyuA-outR*. Lane 3: The mutant strain Δ Δ AE17 mutant strain showed a 2316 bp PCR product (Containing the chloramphenicol resistance gene) using primers *fyuA-outF/fyuA-outR*. Lane 4: Negative control. Lane 5: The mutant strain Δ Δ AE17 mutant strain showed a 410 bp PCR product using primers *fyuA-inF/fyuA-inR*. Lane 6: The mutant strain Δ Δ AE17 mutant strain showed no PCR product using primers *fyuA-inF/fyuA-inR*.

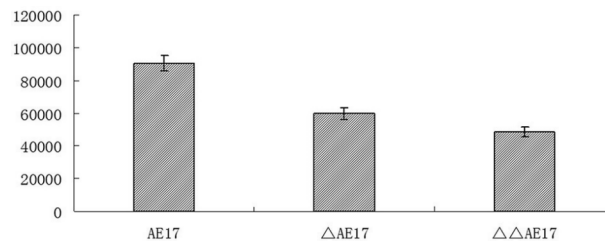


Fig. 3. Adherence assay. Adherence to DF-1 cells by the Δ AE17 and $\Delta\Delta$ AE17 mutant strains were significantly reduced compared with AE17 strain. The ability of Δ AE17 adhesion of the DF-1 cells was significantly reduced decreased, in comparison with that of the $\Delta\Delta$ AE17 strain. The columns represent the means and standard deviations.

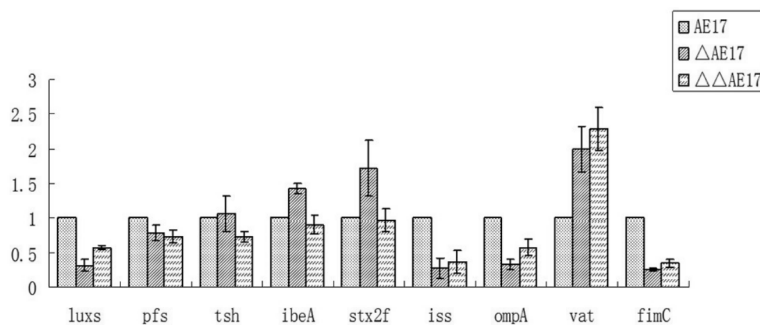


Fig. 4. Differences were found in virulence gene expression in the Δ AE17 and $\Delta\Delta$ AE17 strains. The three columns show the virulence gene mRNA transcript levels of the AE17, Δ AE17, and $\Delta\Delta$ AE17 strains. The relative expression level of each gene was normalized to that of *dnaE*. Results are shown as relative expression ratios compared to the expression of each transcript in the wild-type strain AE17. Each value represents the mean and standard deviation.

stx2f mRNA in the Δ AE17 strain was significantly increased to 1.72 times than that of the AE17 strain. In the Δ AE17 strain, abundance of *ibeA* and *vat* mRNA was increased significantly to 1.43 and 2.00 times than that of the AE17 strain, respectively. There was no significant change in the transcript levels of *tsh*. The $\Delta\Delta$ AE17 strain showed significant decreases in the levels of *luxs*, *pfs*, *tsh*, *iss*, *ompA*, and *fimC* transcripts, which were 0.57, 0.73, 0.73, 0.37, 0.57, and 0.35 times as abundant as those of the AE17 strain, respectively ($P < 0.01$). As shown in Figure 4, there were no significant differences in the levels of *ibeA* and *stx2f* mRNA transcripts in the $\Delta\Delta$ AE17 and AE17 strains. The results shown that the *irp2* and *fyuA* genes in high pathogenicity island may induce transcript levels of *luxs*, *pfs*, *tsh*, *iss*, *ompA*, and *fimC* significantly decreased in avian *Escherichia coli*.

AE17, Δ AE17, and $\Delta\Delta$ AE17 median lethal dose (LD_{50}) determination

After the Roman chicks were injected with the bacteria for 24 hr, they began to show lassitude, inappetence, and hypothermia. After 36 hr, there was some mortality in the APEC-infected groups, but not in the control group. Karber's method was modified

to determine the LD_{50} values for the APEC strains. The LD_{50} values of the AE17, Δ AE17, and $\Delta\Delta$ AE17 strains were 1.5×10^6 , 5.1×10^7 , and 7.5×10^8 CFU/chick, respectively. In comparison with the AE17 strain, the virulence of the Δ AE17 and $\Delta\Delta$ AE17 strains reduced by 34- and 500-fold. The $\Delta\Delta$ AE17 strain was 14.7-fold less virulent than the Δ AE17 strain (Tables 3 and 4). The results shown that the *irp2* and *fyuA* genes in high pathogenicity island may induce the virulence of avian *Escherichia coli* decreased which was certified by chick LD_{50} results.

Discussion

By using Red homologous recombination, the *irp2* mutant and the *irp2/fyuA* mutant strains were constructed successfully in this study. These mutants served as the foundation for further study of the relationship between the *Yersinia* HPI and APEC pathogenicity. Red recombination technology is a genetic engineering technology based on the Red recombinase of the bacteriophage λ and *in vivo* homologous recombination. Using 3' recombinant bacteriophage λ proteases, this technology realizes homologous recombination between DNA fragments and the chromosome *in vitro* (Muyrers et al. 2001, Poteete et al. 2001).

Table 3. Mortality of *E. coli*-infected chicks.

Group	Number of experimental animals	Number of dead animals	Mortality rate (%)
Positive	20	14	70
AE-a1	20	8	40
AE-t	20	0	0
Control	20	0	0

Table 4. The comparison of virulence of AE17 and $\Delta\Delta$ AE17.

Group	Dilution						LD ₅₀ (CFU)
	AE17	2×10 ⁸ 8/8	2×10 ⁷ 7/8	2×10 ⁶ 5/8	2×10 ⁵ 1/8	2×10 ⁴ 0/8	Saline 0/8
$\Delta\Delta$ AE17	10 ¹⁰ 8/8	10 ⁹ 5/8	10 ⁸ 0/8	10 ⁷ 0/8	10 ⁶ 0/8	Saline 0/8	7.5×10 ⁸

As an essential nutrient for the microbial growth, iron is an indispensable bacterial growth factor and a regulator of bacterial virulence factor expression. Many bacteria have the ability to acquire iron from host species, and this ability is a major influence factor on their pathogenesis. The HPI encodes genes that uptake iron from the host through Ybt and then affects virulence of APEC. As important structural genes in the core region of the HPI, *irp2* and *fyuA* play key roles in the iron uptake system encoded by the HPI of *E. coli* (Kurmanova et al. 2007). Therefore, it was speculated that deletion of *irp2* and *fyuA* may influence the iron uptake capacity of APEC strains and the expressions of some virulence factors. This study showed that, in the Δ AE17 strain, the transcription of *luxs*, *iss*, *ompA*, *fimC*, and *pfs* was down-regulated, and the transcription of *stx2f* and *ibeA* genes was up-regulated. In the $\Delta\Delta$ AE17 strain, the transcription of *luxs*, *pfs*, *tsh*, *iss*, *ompA*, and *fimC* was down-regulated. Notably, compared with the single knockout Δ AE17 strain, the double knockout $\Delta\Delta$ AE17 strain showed more down-regulated virulence genes. Although transcription of *stx2f* and *ibeA* was up-regulated in the Δ AE17 strain, there was no significant difference in the expression of these genes between the Δ AE17 and AE17 strains. The decreased transcription of virulence gene indicated weakened AE17 pathogenicity, and thus the double HPI gene knockout impaired pathogenicity more strongly than the single HPI gene knockout.

The capacity of bacteria to adhere to host cells or the surfaces of other organs is vital for their colonization and survival (Schouler et al. 2005). The analysis of adhesion capacity indicated that, compared with the AE17 strain, the capacity to adhere to the cells of the Δ AE17 and the $\Delta\Delta$ AE17 was reduced. Adhesion, an important virulence factor of pathogens, is local-

ed on the bacterial surface. Adhesin can bind to the surface receptors of the host cell to escape scavenging, and it is thus very important in the pathogenic process (Antao et al. 2009). The *fimC* and *ompA* genes encode adherence factors that influence *E. coli* adherence to host cells (Jones et al. 1993). These results confirmed that decreased *fimC* and *ompA* transcription may reduce the adhesion capacity of APEC.

LD₅₀ values intuitively and accurately reflect bacterial virulence. LD₅₀ is inversely proportional to the bacterial virulence; larger LD₅₀ values indicate less virulence. Our LD₅₀ results showed that the AE17 strain showed the strongest virulent in the tested strains. Gene deletion significantly decreased virulence, and double gene-deleted strain showed the weakest virulent. After the determination of LD₅₀ values of the AE17, Δ AE17, and $\Delta\Delta$ AE17 strains, the difference in the pathogenicity of these strains in chicks was studied.

In this work, we report the successful construction of the *irp2* knockout APEC strain and the *irp2/fyuA* knockout APEC strain. Deletion of the HPI core gene *irp2* reduced transcription of virulence genes, capacity to adhere to DF-1 cells, and pathogenicity in chicks, and the deletion of *irp2* and *fyuA* resulted magnified these effects. This study might lay the foundation for further investigation of the genes in the core region of the HPI in the pathogenesis of APEC

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