Characterization of chicken IFI35 and its antiviral activity against Newcastle disease virus

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ABSTRACT. Interferon-induced protein-35 kDa (IFI35) was an antiviral protein induced by interferon (IFN)-γ, which plays an important role in the IFN-mediated antiviral signaling pathway. Here, we cloned and identified IFI35 in the chicken for the first time. Chicken IFI35 (chIFI35) contains an open reading frame (ORF) of 1,152 bp encoding a protein of 384 amino acids containing two conserved Nmi/IFI35 domain (NID) motifs. Tissue distribution analysis of chIFI35 in healthy and Newcastle disease (ND) virus-infected chickens indicated a positive correlation between chIFI35 mRNA transcription and ND viral loads in various tissues. The role of chIFI35 in regulation NDV replication were further assessed by up- or down-regulated chIFI35 expression in DF-1 cells transfected with plasmid harboring chIFI35, pCMV-3HA-chIFI35 or shRNA targeting chIFI35, pshRNA-chIFI35 plasmids. NDV replications in DF-1 cells were significantly reduced or slightly increased by over- or under-expression of the chIFI35 protein, respectively, indicating the role of chIFI35 in anti-NDV infection. Moreover, chIFI35 also involved in regulation of viral gene transcription and IFNs expression. The collected data were meaningful for research of chicken antiviral immunity and shed light on the pleiotropic antiviral effect of chIFI35 during NDV infection.

KEY WORDS: antivirus, chicken, interferon, interferon-induced protein-35 kDa, Newcastle disease virus

Interferon-induced protein-35 kDa (IFI35), also known as IFP35, was one of the antiviral proteins induced by both type I and type II interferons (IFNs) [1]. IFI35 was originally identified in 1994 as an IFN-induced protein by screening a cDNA library from HeLa cells treated with IFN-γ [1]. It was widely expressed in various cells, such as fibroblasts, mononuclear macrophages, and epithelial cells, and its expression in the mitochondrial/lysosomal or microsomal fractions was increased by IFN treatment [1, 4]. This protein could translocate to the nucleus from the cytoplasm upon IFN stimulation. The protein interacts with N-Myc interacting protein (Nmi) and casein kinase 2 interacting protein-1 (CKIP-1), and which regulate the stability of IFI35 [17, 28, 30]. Moreover, over-expression of IFI35 could efficiently inhibit the replication of bovine foamy virus, prototype foamy virus [21], and foot-and-mouth disease virus [29], however, IFI35 promoted vesicular stomatitis virus and Senda virus proliferation through degradation of retinoic acid inducible-gene I (RIG-I) suppressing the expression of IFNs [5]. IFI35 has also been reported involving in various diseases, such as Sezary syndrome, which it was significantly down-regulated in patients, suggesting a role of it in suppression tumor [19]. To our knowledge, the pleiotropic effects of IFI35 have not been examined in birds, besides limited reports about mammals and fish [5, 18, 29].

Newcastle disease (ND), one of the most highly contagious and infectious diseases, was causing serious death and economic losses in both domestic and wild avian species [20]. Newcastle disease virus (NDV), the causative agent of ND, belongs to the order Mononegavirales, family Paramyxoviridae and genus Avulavirus [24]. The etiology of the disease was the velogenic strains of NDV, including viscerotropic and neurotropic velogenic NDV strains [11]. NDVs were identified to be excellent inducer of IFNs, while they gain potent immunosuppressive ability though evolution [9, 16]. NDVs regulated IFNs expression have been identified through impact key elements, like MDAS, STAT, in IFN pathway [13, 25]. While, there will be a series of steps before we completely reveal these mechanism involving in the host-virus interactive stages during NDV infection. Especially, the functions

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of most downstream molecules in chicken IFN pathways, including IFI35, OASL, and Mx, are still not clear. In our previous study, IFI35 was significantly up-regulated in velogenic NDV-infected chicken visceral tissues using transcriptome sequencing compared with lentogenic NDV or control-infected chicken tissues. Here, the molecular characterization of chicken IFI35 (chIFI35) and its antiviral role in against NDV infection were analyzed in the current study.

MATERIALS AND METHODS

Animals, cells, and virus

The specific pathogen free (SPF) chicken embryonated eggs were purchased from the Merial-vital Laboratory Animal Technology (Beijing, China). Nine-day-old SPF chicken embryos were incubated at 37°C for viral amplification or making chick embryo fibroblast (CEF) cells. The SPF chickens were fed in isolator until 21 days old for experiments.

All incubations and reactions were performed under 5% CO₂ and 37°C in flat-bottom plates. DF-1, CEF, and HEK293T cells were maintained and cultured in Dulbecco’s modified Eagle’s medium (DMEM, Gibco, Grand Island, NY, USA) with 10% fetal bovine serum (FBS, Hyclone, Logan, UT, USA).

NDV strains, velogenic strain F48E9, and lentogenic strain rLa Sota-GFP were obtained from Northwest A&F University Infectious Diseases Lab. Viruses were propagated in 9-day-old SPF chicken embryonated eggs, titrated with the hemagglutination (HA) test, and then stored at −80°C for further use.

RNA extraction and cDNA preparation

Total RNA was extracted using Trizol (TaKaRa, Dalian, China) according to the instructions. The cDNA was synthesized using StarScript II First-Strand cDNA Synthesis Mix (GenStar, Beijing, China) according to the instructions. In brief, the reaction was performed in a volume of 20 μl, containing 500 ng total RNA, 10 μl 2 × Reaction mix, 1 μl oligo (d)₁₈ (50 μM), 1 μl StarScript II RT Mix, and RNase-free ddH₂O up to 20 μl. Each reaction used the following conditions: 25°C for 10 min, 42°C for 30 min, and 85°C for 5 min. The resultant cDNA was stored at −20°C for further use.

Cloning, sequencing, and bioinformatic analysis of chIFI35 sequences

The primers of the partial sequence of chIFI35 including the full-length ORF were designed based on the computer-predicted sequence of Gallus gallus IFI35 and our transcriptome data of chicken visceral tissues (data not shown). Primers are listed in Table 1. The amplification was performed as follows: initial denaturation at 95°C for 5 min followed by 35 cycles at 95°C for 30 sec, annealing at 55°C for 30 sec and extension at 72°C for 30 sec, and a further extension at 72 for 10 min following the last cycle. The PCR products were purified and cloned into the pMD19-T vector (TaKaRa, Dalian, China), and then sent to Genweizhi (Suzhou, China) for sequencing.

The sequence and protein were analyzed as previous described [18, 25–27]. In brief, the potential open reading frame (ORF) and homology search of chIFI35 were analyzed by the ORF finder (http://www.ncbi.nlm.nih.gov/orf/gorf/gorf.html) and BLAST program (http://blast.ncbi.nlm.nih.gov/Blast.cgi) in NCBI. Protein molecular weight, isoelectric point, and instability index of chIFI35 were predicted by the online server site (http://www.expasy.org/protparam/), while the domains and motifs of chIFI35 were determined using SMART (http://smart.embl-heidelberg.de/) and MotifScan (http://myhits.isb-sib.ch/cgi-bin/motif_scan). The MultiLoc tool (http://abi.inf.uniba.it/services/MultiLoc/) predicted the cellular distribution of chIFI35, and SignalIP (http://www.cbs.dtu.dk/services/SignalP/) and the SACS MEMSAT2 tool (http://www.sacs.ucsf.edu/cgi-bin/memsat2) were used to analyze the signal peptide and transmembrane segments of chIFI35. Multiple sequence alignment was conducted using Clustal W 1.83, and the sequence diversity and phylogenetic analyzes were predicted using the MEGA 5.2 program.

Real-time quantitative PCR (RT-qPCR)

RT-qPCR was used to detect the expression of mRNA of target genes using the 2 × RealStar Green Power kit (GenStar, Beijing, China) according to the manufacturer’s instructions. Primers used in this study are listed in Table 1 [7, 12, 14]. The relative expression levels of the target genes were calculated using the 2^ΔΔCT method [15], and 28S was used to normalize the fold-changes in expression.

Tissue distribution profile of chIFI35 expression in vivo

In order to investigate the tissue distribution of chIFI35, various tissues, including the blood, brain, bursa of fabricius, liver, spleen, heart, lung, intestines, pancreas, kidney, proventriculus, gizzard, trachea, skin, and muscle were collected from 21-day-old SPF chickens for RT-qPCR.

To evaluate the mRNA expression level of chIFI35 in the tissues in NDV-infected chickens, the tissues that highly expressed chIFP35 were collected from NDV-infected chickens and analyzed using RT-qPCR. In brief, 21-day-old SPF chickens were injected with 100 μl F48E9 (2^8 HA) and control birds were injected with 100 μl PBS (three chicken). At 24 hr post-infection, all chickens were euthanized, and the organs were collected and analyzed by RT-qPCR as above.

Expression plasmid construction and cell transfection

To demonstrate the molecular function of chIFI35 in vitro, the full-length chIFI35 sequence was cloned into the pCMV-3HA or
Table 1. The primers used in this study

| Primers       | Sequences (5’-3’)                                                                 | Methods      | Size (bp) | References |
|---------------|------------------------------------------------------------------------------------|--------------|-----------|------------|
| IRF1          | F: GCTACACCCGCTCACGA R: TCAGCCATGGCGGATTTC                                          | RT-qPCR      | 133       | [12]       |
| MAVS          | F: CCTGACTCAAACAAAGGGAAG R: AATCAGAGGGACTGCAACAG                                    | RT-qPCR      | 123       | [7]        |
| STING         | F: TGACCCAGAGCTCCTAAAGG R: CTTGGGCAAGAATCTTCCAAGC                                   | RT-qPCR      | 63        | [7]        |
| TBK1          | F: AAGAAGGCAACATCCGGAGA R: GTGATGGCTGAAAATAGCAG                                     | RT-qPCR      | 152       | [7]        |
| NF-κB         | F: CATTGCGAGAGGCTCCTAT R: TCCAGTTCCCGTTTCCTCA                                    | RT-qPCR      | 102       | [7]        |
| IRF7          | F: CAGTGCTTCTCCAGACAAAA R: TGCATGTGGATTAGCTCGAT                                       | RT-qPCR      | 169       | The present study |
| IFI35         | F: AGGGGATGCTTCTGAGTAC R: GCTCTCCAGACGCACT                                          | RT-qPCR      | 229       | The present study |
| LGP2          | F: CCAGAATGCGAGCAGCGAG R: AATGTGGTCAACTGAGGATGT                                     | RT-qPCR      | 109       | The present study |
| MDA5          | F: TTGTGCAGAGAGCTGTTAAGG R: GAATCTACGTCGTGACTCTGTC                                   | RT-qPCR      | 109       | The present study |
| IFN-α         | F: GACATGGGCTCTCCAAACTCC R: AGGCGCTGAATCTTGCATT                                       | RT-qPCR      | 330       | [14]       |
| IFN-β         | F: TTCAGCTCCTCGAAATCG R: TCCATCATGACGATCAG                                          | RT-qPCR      | 192       | The present study |
| IFN-γ         | F: TGGACCAGAGTTGTCGAT R: CTTGGCCAGGTCCATGATA                                         | RT-qPCR      | 134       | [14]       |
| 28S           | F: GGTATGCGCCGCGAGCCT R: CCAGTGCCGAGCTCCAT                                          | RT-qPCR      | 160       | The present study |
| NP            | F: TCGGATGAAAGGAGAATAATCG R: GTCTCCAGAATGATGCTCA                                    | RT-qPCR      | 182       | The present study |
| P             | F: CAAATAATCGTCAAATGAATTA R: CTCATTCAAGACATCAG                                      | RT-qPCR      | 322       | The present study |
| M             | F: CCGATGCTCCTACAGACAG R: GGAGGCTCGTACAGGAGCAT                                       | RT-qPCR      | 223       | The present study |
| F             | F: GACGGATTATGCTGCACTACG R: GCCGCTACGATGAGGTCG                                        | RT-qPCR      | 294       | The present study |
| RNAS          | F: CTACAGGATGGTTGCAGCT R: CCTGTGCGATGTCGGAGGCA                                       | RT-qPCR      | 151       | The present study |
| L             | F: CCATTTGCTGACCAACATACT R: GCCCTTGACAGCTCCAT                                          | RT-qPCR      | 325       | The present study |
| cPAGGS-Flag-chIIF35 | F: GGATTTCATTTATGCCTGAGGAGGAGGACTC R: GAAGATCTCTTACATTGCTGCTGCATCTGTAATCGGCTGAGCGGCTCCATCACCC | IFA         | 1,191     | The present study |
| cCMV-3HA-chIIF35 | F: GGTTATCATTTATGCCTGAGGAGGAGGACTC R: GAAGATCTCTTACATTGCTGCTGCATCTGTAATCGGCTGAGCGGCTCCATCACCC | Amplication   | 1,167     | The present study |
| IFI35-shRNA1  | F: GATCCGCTGAGAGGATGAGAAGGTTTACAAATCCAGAGGTTAAGCTGCTACAGCTGTTTTCGTGATGTTGAGGGTTTAA AGGAGGTTTACAAATCCAGAGGTTAAGCTGCTACAGCTGTTTTCGTGATGTTGAGGGTTTAA | Interference | 63        | The present study |
| IFI35-shRNA2  | F: GATCCGCTGAGAGGATGAGAAGGTTTACAAATCCAGAGGTTAAGCTGCTACAGCTGTTTTCGTGATGTTGAGGGTTTAA AGGAGGTTTACAAATCCAGAGGTTAAGCTGCTACAGCTGTTTTCGTGATGTTGAGGGTTTAA | Interference | 63        | The present study |
| IFI35-NC      | F: GATCCGCTGAGAGGATGAGAAGGTTTACAAATCCAGAGGTTAAGCTGCTACAGCTGTTTTCGTGATGTTGAGGGTTTAA AGGAGGTTTACAAATCCAGAGGTTAAGCTGCTACAGCTGTTTTCGTGATGTTGAGGGTTTAA | Interference | 63        | The present study |

pCAGGS vector. Primers are listed in Table 1. Recombinant plasmids, including pCMV-3HA-chIIF35 and pCAGGS-Flag-chIF35, were confirmed by DNA sequencing. In order to test the effect of chIIF35 on NDV proliferation, we designed shRNA primers (IFI35-shRNA1, IFI35-shRNA2, and IFI35-NC) using the BLOCK-it™ RNAi Designer (https://rnadesigner.thermofisher.com/rnaexpress/design.do), and the primer sequences are listed in Table 1. Next, double-stranded of shRNA was synthesized and cloned into the lentiviral expression plasmid pCD513B-U6-neo, which we named pCD513B-chIIF35-shRNA1 (shRNA1), pCD513B-
chIFI35-shRNA1 (shRNA2), and pCD513B-chIFI35-negative control (NC). All the sequences of the plasmids were confirmed by sequencing. Other plasmids, including pCAGGS-Flag-chIFN-α, pCAGGS-Flag-chIFN-β, and pCAGGS-HA-P were previously constructed in our laboratory.

Cell transfection was carried out using Turbofect transfection reagent (Thermo Scientific, Waltham, MA, USA) according to the manufacturer’s instructions. In brief, DF-1 cells were seeded in 24-well plates, and a mixture of Turbofect and plasmids was added, and then cultured at 5% CO2 and 37°C.

**Virus titration**

Viral multiplications were quantified by the 50% tissue culture infection dose (TCID50) as previous described [6]. Briefly, DF-1 cells were seeded into 96-well plates for 24 hr. Virus supernatants from cell cultures were 10-fold serially diluted and then 100 µl/well were respective added (three replicates) for infection. Five days after infection, the TCID50 was calculated using Reed–Muench method.

**Western blotting**

Western blotting was used to analyze *IFI35* expression in DF-1 cells post-treatment. After transfected for 48 hr, cells were washed twice with PBS and lysed for 10 min by radio immunoprecipitation assay (RIPA) buffer with protease inhibitor Phenylmethanesulfonyl fluoride (PMSF) on ice. Cell lysate proteins were separated by 10% SDS-PAGE, blotted onto polyvinylidene difluoride (PVDF) membranes (Millipore, Boston, MA, USA), and subsequently probed with mouse anti-β-actin monoclonal antibody and mouse anti-β-actin monoclonal antibody (CoWin Biosciences, Beijing, China) for 2 hr at room temperature to detect *IFI35* and GAPDH, respectively. HRP-conjugated goat anti-mouse IgG (Sangon Biotech, Shanghai, China) was used as a secondary antibody. Immuno-stained proteins were visualized using ECL peroxidase substrate (Millipore, Boston, MA, USA).

**Immunofluorescence staining**

DF-1 cells were grown to 50–60% confluence on coverslips and were co-transfected with the plasmids pCAGGS-HA-P and pCAGGS-Flag-chIFI35 for 36 hr. Cells were fixed in 4% paraformaldehyde for 30 min, permeabilized with 0.25% Triton X-100 for 15 min, and blocked with 30% bovine serum albumin (BSA) for 30 min. Next, cells were stained with anti-Flag and anti-HA mAbs, followed by staining with secondary antibodies conjugated to Alexa Fluor 594 or fluorescein isothiocyanate (FITC; Sangon Biotech, Shanghai, China). Nuclei were stained with 4,6-diamidino-2-phenylindole (DAPI), and the co-localization of *P* protein and chIFI35 were visualized using a Nikon C1-si confocal fluorescence microscope (Nikon Instruments Inc., Melville, NY, USA).

**Statistical analysis**

Experiments were repeated independently at least twice. GraphPad prism was used to statistical analysis. All data are showed as the mean ± standard error of the mean (SEM). The significant differences between two groups were performed using Student’s *t*-tests in GraphPad software. The results were considered to be statistically significant as follows: ***P*<0.001, **P*<0.01, *P*<0.05, and ns mean not significant.

**RESULTS**

**Molecular and genetic character of chIFI35**

In our previous transcriptome study, we discovered that *IFI35* was up-regulated in both velogenic (F48E9) and lentogenic (La Sota) NDV infected groups, but the expression in F48E9 infected (5.34 fold-change) was much higher than La Sota infected (1.25 fold-change) using next-generation sequencing [10]. In this study, an 1,152 bp full-length ORF of *IFI35* was first amplified and cloned from chicken visceral tissue (chIFI35). The obtained sequence (GenBank accession number: KY931454) has 99% similarity with computer-predicted *Gallus gallus* *IFI35* sequence (XM_418132.5). Phylogenetic analysis based on the deduced amino acid sequences certified that *IFI35* was conserved among similar species but significant difference among different species (Fig. 1). ChIFI35 was composed of 383 amino acids (Fig. 1A), with the theoretical isoelectric point of 5.14 and calculated molecular mass of 46.3 kDa. It was predicted unstable due to 51.59% instability probability, which indicated that this protein was unstable.

ChIFI35 protein mainly distributed in the cytoplasm with cellular location prediction by MultiLoc tool. No putative signal peptide and trans-membrane region were found based on signal peptide analysis and trans-membrane domain prediction program. According to analysis with SMART software, the domain structure of the obtained chIFI35 was similar to the predicted sequence (XM_418132.5) and both composed of two characteristic Nmi/IFI35 domains (NIDs) at positions 166–263 and 274–361.

ChIFI35 sequence showed 89.2% identity to that from *Meleagris gallopavo* (avian), 47.9% from *Xenopus tropicalis* (amphibian), 44.5% from *Cynoglossus semilaevis* (fish), 52.5% from *Anolis carolinensis* (reptile), and 40.2% from *Mus musculus* (mammalian) (Table 2). Five clusters present in the Phylogenetic tree constructed based on the *IFI35* amino acid sequences from different species (Fig. 1B), and interestingly, the different taxa-derived *IFI35* counterparts formed separate sub-clusters that agreed with their orthodox taxonomy. ChIFI35 belongs to the I/FI35 group of birds and has a close relationship with *Meleagris gallopavo.*
Tissue distribution profile of chIFI35 and transcriptional response against immune stimulation by NDV infection in SPF chicken

In the previous study of F48E9-infected chicken embryo (Fig. 2A, the allantoic cavity infection) or bursa of fabricius (Fig. 2B, Intramural injection) RNA sequencing, it was found that the expression of IFI35 was significantly increased [10, 22]. In healthy SPF chickens, chIFI35 mRNA had high level transcription in the lungs, glandular stomach, brain, bursa of fabricius, gizzard, kidney, heart, intestines, liver, and muscle, whereas, lower level expressed in the trachea, spleen, pancreas, and skin (Fig. 2C).

Velogenic NDV, F48E9, infection increased chIFI35 expression in various immune-associated tissues of chickens (Fig. 2D). The highest level of chIFI35 expression was detected in intestine, which also was one of the major target of NDV. The expressions of chIFI35 in trachea, bursa of fabricius, and spleen were also significantly unregulated comparing with control group. Interestingly, high viral loads were detected in intestine, which also was one of the major target of NDV. The expressions of chIFI35 were highest level in kidney, heart, intestines, liver, and muscle, whereas, lower level expressed in the trachea, spleen, pancreas, and skin (Fig. 2C).

Table 2. Estimates of evolutionary divergence between chicken interferon-induced protein-35 kDa (chIFI35) and other species sequences

| Divergence | Percent identity |
|------------|------------------|
|            | 1    | 2    | 3    | 4    | 5    | 6    | 7    | 8    |
| 1          | 47.9 | 89.2 | 44.5 | 46.2 | 39.9 | 40.2 | 52.5 |
| 2          | 86.9 | 47.4 | 45.7 | 46.3 | 39.8 | 41.1 | 49.0 |
| 3          | 5.2  | 85.2 | 45.9 | 46.9 | 46.5 | 46.2 | 58.8 |
| 4          | 92.4 | 100  | 88.3 | 62.3 | 36.6 | 36.4 | 44.9 |
| 5          | 85.9 | 98.7 | 81.8 | 55.1 | 40.6 | 39.4 | 45.8 |
| 6          | 75.9 | 92.2 | 74.3 | 103.1| 79.5 | 84.0 | 45.9 |
| 7          | 75.6 | 87.5 | 76.5 | 107.0| 86.9 | 26.3 | 45.5 |
| 8          | 65.1 | 81.5 | 63.5 | 99.3 | 92.5 | 79.8 | 82.8 |

1–8 represent the species of Gallus gallus, Xenopus tropicalis, Melocys galepavo, Cynoglossus semilaevis, Kryptolebias marmoratus, Homo sapiens, Mus musculus, and Anolis carolinensis, respectively.

ChIFI35 plays an antiviral role against NDV infection in DF-1 cells

Over-expressed or down-expressed chIFI35 in DF-1 cells transfected with plasmids were confirmed by western blotting assay at 24 hr post transfection (hpt) (Fig. 3A and 3D). NDV strains, F48E9 or rLa Sota-GFP, were used to infect the cells at 24 hpt. The proliferations of the virus in supernatants of different treated cells were quantified by testing viral TCID50. Comparing with the untreated cells, over-expressed chIFI35 could significantly decrease F48E9 and rLa Sota-GFP propagation (Fig. 3B and 3C). However, viral proliferations in IFI35-down-regulated DF1 cells were slight enhanced with no significantly different (Fig. 3E and 3F). These data suggested that chIFI35 has a certain antiviral ability.

ChIFI35 affects the transcription of RNA viral proteins

IFI35 affect DNA virus transcription by interacts with one or more viral proteins. Whether it also plays a role in RNA virus replication has not been clearly Illuminated. The distribution of chIFI35 and NDV protein P were identified by co-transfected plasmids, pCAGGS-Flag-chIFI35 and pCAGGS-HA-P, and the proteins of Flag-chIFI35 and HA-P in cells were co-localization under laser confocal scanning microscopy (Fig. 4A). Furthermore, the transcription of different NDV genes in cell transfected empty vector or plasmids overexpressing of chIFI35 were detected. The CT value of P gene was significant increased when over-expression of chIFI35, which indicated that P gene transcription was reduced at 12 hr after F48E9 infection. The transcription of NP, P, F and HN genes were also down regulated at 24 hr (Fig. 4B). Hence, these data indicated that chIFI35 may inhibit NDV proliferation by affect NDV gene transcription, especially for P and F.

ChIFI35 enhances IFN expression

Since IFI35 involved in IFN pathway, the correlation between IFI35 and IFNs expression were further analysis with RT-qPCR. Cells were collected at different time points after F48E9 infection, and then the expression of chIFI35, IFNa, IFNb, and IFNg were detected. ChIFI35 expression was keep increasing after NDV infection, which was similar to the IFNa/β/γ expression (Fig. 5A–D). The expressions of IFNa/β/γ in cells over-expressed chIFI35 were also quantified. Over-expressed chIFI35 could significantly enhanced type I and type II IFNs expression in cells (Fig. 5E). And interestingly, chIFI35 was also significantly up regulated when IFNa/β were over-expressed in cells (Fig. 5F).

Furthermore, the expression of IFNs signaling pathway molecules, including MDA5, LGP2, MAVS, STING, TBK1, IRF1, IRF7, and NF-κB, were detected and compared in cells over- or normal- expressing chIFI35 after F48E9 infection. The transcriptions of MAVS, IRF7, and NF-κB were not significant differently both in CEF and DF-1 cells, whereas MDA5, LGP2, STING, TBK1, and IRF-1 were significantly increased (Fig. 6) when chIFI35 was overexpressed, indicating that chIFI35 play a role in promoting the expression of IFNs.
Fig. 1. Multiple alignment analysis and phylogenetic tree construction based on the interferon-induced protein-35 kDa (IFI35) amino acid sequences. (A) Multiple alignment analysis of IFI35 family amino acid sequences using Clustal W in DNAstar. These sequences were downloaded from GenBank with accession numbers as follows: XM_418132 (Gallus gallus IFI35, ggaIFI35), XM_010724650 (Meleagris gallopavo IFI35, mgpIFI35), XM_002932512 (Xenopus tropicalis IFI35, xtpIFI35), XM_00829193 (Cynoglossus semilaevis IFI35, cysIFI35). Amino acids conserved among species were indicated as identical “•”. (B) Evolution analyzes were conducted using MEGA6.0. Numbers in branches indicate evolutionary distances. The evolutionary history was inferred using the neighbor-joining (NJ) method. The evolutionary distances were computed using the Poisson correction method.
Fig. 2. Tissue distribution profile of chicken interferon-induced protein-35 kDa (chIF35). The relative expression of chIF35 of F48E9 infected chicken embryo using RNA-Seq and RT-qPCR (A). The relative expression of chIF35 at 24 hr and 48 hr after F48E9 infected the bursa of fabricius (B). The tissue distribution profiles of chIF35 in various tissues in healthy chicken (C) and F48E9-infected chickens (D). (E) The viral loads of Newcastle disease virus (NDV) in different tissues.

Fig. 3. Viral titers of Newcastle disease virus (NDV) in DF-1 cells with interferon-induced protein-35 kDa (IF35) over- or under-expression. DF-1 cells were transfected with pCMV-3HA-IF35, empty vector plasmids (A), or interference plasmids (D) for 24 hr, and infected with La Sota (1 moi) or F48E9 (0.1 moi). The viral titer of La Sota (B, E) and F48E9 (C, F) at different time points were detected using the TCID_{50} method. **p<0.01, *p<0.05, and ns, not significant.
Fig. 4. Chicken interferon-induced protein-35 kDa (ChIF35) affects viral gene transcription. (A) The plasmids pCAGGS-HA-P and pCAGGS-Flag-chIF35 were co-transfected for 36 hr in DF-1 cells, and observed the cell sublocalization of chIF35 and NDV P protein under the laser confocal microscopy (400×). (B) CT values of each viral gene transcript after F48E9 infection. Data are shown as the mean ± SEM (n=3). The statistical analysis was performed in GraphPad Prism using unpaired two-tailed t-tests: *P<0.05, **P<0.01.

Fig. 5. Relative expression of chicken interferon-induced protein-35 kDa (chIF35) and interferons (IFNs) during F48E9 infection. DF-1 cells were infected with F48E9 (0.1 moi, A–D), and the cells were collected at various timepoints to detect the expression changes using RT-qPCR of chIF35 (A), IFNa (B), IFNβ (C), and IFNγ (D). (E) ChIF35 was increased by IFNa/β over-expression. (F) IFNa/β/γ expression using RT-qPCR following transfection of DF-1 cells with chIF35 and empty vector plasmids for 24 hr. Data are shown as the mean ± SEM (n=3). The statistical analysis was performed in GraphPad Prism using unpaired two-tailed t-tests: **P<0.01, ***P<0.001.
**DISCUSSION**

*IFI35* was first found in HeLa cells after interferon-γ treatment in 1994 [1]. *IFI35* sequences have been reported in mouse, cattle, and fish [5, 18, 21]. To date, the pleiotropic effects of *IFI35* have not been examined in birds, with only limited reports in mammals and teleosts. In this study, the full-length ch*IFI35* cDNA was successfully cloned and sequenced for the first time. The ch*IFI35* protein consists two NID domains in the N-terminal, similar to human *IFI35* (hu*IFI35*) and rockfish *IFI35* (Rf*IFI35*) [18], suggesting that it interacts with Nmi protein to maintain stable and subcellular localization similar to hu*IFI35* or Rf*IFI35* [3]. In order to identify the evolutionary relationship between ch*IFI35* and other species, a neighbor-joining (NJ) tree was constructed, and which demonstrated that the amino acids sequences of *IFI35* were with high identity in birds, but with low identities to species in other four classes, which consistent with the amino acid sequences analysis. The pairwise alignment and phylogenetic analysis results showed that ch*IFI35* was homology among birds, suggesting similar structures and functions of it in birds.

Tissue distribution analysis revealed a broad tissue distribution of ch*IFI35* in healthy chickens. High lever expressions of ch*IFI35* were detected in the lung, proventriculus, brain, bursa of fabricius, kidney, and intestine. The rule of tissue distribution was similar to *IFI35* of fish and human [18]. Analysis ch*IFI35* expression in immune-associated tissues after F48E9 infection found that ch*IFI35* mRNA was high level expression in intestines, which was one of the major targets of velogenic NDV. Moreover, ch*IFI35* highly expressed in respiratory tract, lung and trache, as well as in the digestive tract, intestines, proventriculus, and gizzard. These tissues are the main targets of NDV and also present typical lesions of NDV infection. The viral loads of the trachea and kidney were slightly low, with slightly different from the expression of ch*IFI35*, which may be caused by the difference in inoculation method or sample collection time. Basically synchronized tissues distribution of ch*IFI35* and NDV suggests that ch*IFI35* involves in NDV infection process.

The current understanding of the role of *IFI35* in virus infections was limited. A previous study suggested that *IFI35* negatively regulates the innate immune signaling, and could promote VSV replication via keeping RIG-I in phosphorylated form and mediating proteasomal ubiquitination of RIG-1 [5]. However, in this study, over-expression *IFI35* increased NDV, F48E9 and La Sota viral replication in DF-1 cells. This suggested that an anti-NDV infection role of ch*IFI35*, consistent with its reported role in against bovine foamy virus (BFV) [21] and foot-and-mouth disease virus (FMDV) [29]. The different roles of *IFI35* in VSV and NDV infection may cause by RIG-1. As well known, RIG-1 was deficient in chicken [11]. Though MDA5 replaced parts function of RIG-1, they are difference in response to viral infection and their final destiny (the way of degradation) [25].

Previous reports also demonstrated that ch*IFI35* could inhibit virus replication by interacting with viral structural proteins, such as the Btas protein of BFV [21]. Beside, *IFI35* could transfer into the nucleus from the cytoplasm and where it affects the virus
genome transcription or replication [21]. However, limited knowledge about IFI35 in anti RNA virus infection has been revealed. In the current study, we identified a relationship between chIF35 and the transcription of NDV genes. Compared with the CT values of each viral gene between chIF35 over-expressing cells and vector-transfected cells, the CT values of NP, P, F and HN were significant increased, especially P gene, which indicates that chIF35 might affect the transcription of viral proteins in the viral genome. NP and P proteins are two important structural proteins and they compose the RNA-dependent RNA polymerase (RdRp) with the large (L) protein [8, 23]. The F and HN proteins are also two significant structural proteins that are mainly involved in viral invasion [23]. The subcellular location—determined using laser confocal microscopy—also showed that chIF35 co-localized with P protein in the cytoplasm. Those results suggested that chIF35 could affect the viral gene transcription, but the further research was needed to investigate the specific mechanism. Moreover, the nonstructural protein V was transcripted from P protein, which was considered as an antagonistic protein of IFN [2]. Here, the less transcripts of P gene could affect the expression of V protein, so as to result in decreasing the suppression of IFN from V protein.

IFI35 was an interferon-induced protein and so we investigated the relationship between chIF35 expression and IFNs, and they have similar expression trends at different time points in F48E9-infected DF-1 cells, which was similar to RIIIF-γ and RDIIF35 mRNA expression in fish [18]. However, we observed that IFN-γ expression induced by IFI35 was apparently greater than that of type I IFNs, which indicated that the anti-NDV function of chIF35 may also depend on IFN-γ. NDV was considered an interferon inducer, especially type I, and we detected several upstream molecules of the interferon signaling pathways in uninfected or infected cells during chIF35 over-expression. We found that chIF35 could induce increased of the expression of IFN upstream molecules, including MDA5, LGP2, STING, TBK1, and IRF1, but have no significant changed in MAVS, IRF7 and NF-kB. These data showed that chIF35 could induce IFNs and its regulator, which play an antiviral role in NDV replication. These results also indicated that chIF35 may play a significant role in the signal transmission of IFN production, similar to the result of IFI35 in FMDV infection [29]. Unfortunately, the lack of chicken antibodies meant that we were unable to study the antiviral mechanism and protein levels in depth.

In summary, the present study identified and characterized IFI35 in chicken and showed that this protein was conserved in birds. The highest expression of chIF35 mRNA was observed in the lung, followed by the brain and bursa of fabricius. chIF35 plays an important role in the antiviral NDV infection. The antiviral function of chIF35 may relate with that chIF35 expression and protein levels in depth.

CONFLICT OF INTEREST. The authors declare no conflict of interest.

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