Development and application of multiplex PCR method for simultaneous detection of seven viruses in ducks

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Abstract
Background: Major viruses, including duck-origin avian influenza virus, duck-origin Newcastle disease virus, novel duck parovirus, duck hepatitis A virus, duck Tembusu virus, fowl adenovirus, and duck enteritis virus, pose great harm to ducks and cause enormous economic losses to duck industry. This study aims to establish a multiplex polymerase chain reaction (m-PCR) method for simultaneous detection of these seven viruses.

Results: Specific primers were designed and synthesized according to the conserved region of seven viral gene sequences. Then, seven recombinant plasmids, as the positive controls, were reconstructed in this study. Within the study, D-optimal design was adopted to optimize PCR parameters. The optimum parameters for m-PCR were annealing temperature at 57 °C, Mg²⁺ concentration at 4 mM, Taq DNA polymerase concentration at 0.05 U/μL, and dNTP concentration at 0.32 mM. With these optimal parameters, the m-PCR method produced neither cross-reactions among these seven viruses nor nonspecific reactions with other common waterfowl pathogens. The detection limit of m-PCR for each virus was 1 × 10⁴ viral DNA copies/μL. In addition, the m-PCR method could detect a combination of several random viruses in co-infection analysis. Finally, the m-PCR method was successfully applied to clinical samples, and the detection results were consistent with uniplex PCR.

Conclusion: Given its rapidity, specificity, sensitivity, and convenience, the established m-PCR method is feasible for simultaneous detection of seven duck-infecting viruses and can be applied to clinical diagnosis of viral infection in ducks.

Keywords: M-PCR, Duck, Virus, Clinical detection

Background
The Food and Agriculture Organization (FAO) suggests that approximately 818 million ducks were raised annually in China, accounting for 65.96% of the World’s stock in 2016 (http://www.fao.org/faostat/en/#data/QA). However, along with the development of China’s duck industry, many problems emerged. Among them, viral infection is one of the most important problems endangering the waterfowls [1]. The major viruses that cause enormous economic losses to duck industry include duck-origin avian influenza virus (AIV), duck-origin Newcastle disease virus (NDV), novel duck parovirus (NDPV), duck hepatitis A virus (DHAV), duck Tembusu virus (DTMUV), fowl adenovirus (FAdV), and duck enteritis virus (DEV) [2].

AIV has been isolated from numerous avian species. Normally, waterfowls are the primary reservoir hosts. However, highly pathogenic avian influenza could cause high mortality in ducks and geese [3]. Therefore, determining AIV in ducks is important in epidemiology research [4]. Similar to AIV, waterfowls are generally considered potential reservoirs for NDV, which has been occasionally reported in China in ducks. In addition, since 1997, Newcastle disease occurred frequently in geese throughout China, causing devastating economic losses [5]. NDPV, a new variant of goose parvovirus, is a
novel duck parvovirus. The virus is currently infecting ducks across China, the morbidity of which is approximately 20–40%, thereby causing serious economic loss for the duck industry in China [6]. DHAV is a highly contagious, acute, fatal disease of ducklings, causing more than 80% of mortality in ducklings under three weeks old. Therefore, DHAV has jeopardized the duck industry throughout the world [7, 8]. DTMUV is another newly emerged infectious disease mainly affecting laying ducks. When the ducklings are infected with DTMUV, the morbidity can reach 90%, and the mortality rate can vary from 5 to 30% [9]. FAdV is another major concern in the poultry industry; it may cause immunodeficiency or vaccination failure and result in serious economic loss [10]. DEV could also cause considerable economic losses to the commercial duck industry due to the high mortality and decreased egg production rate. Therefore, it poses a continuous threat to wild and migratory waterfowl populations [11].

The development of rapid and convenient methods to determine these viruses and accordingly implementing preventive measures to reduce economic losses as soon as possible [1, 12] are important given the great danger of these viruses on the duck industry. Thus far, many methods, such as virus isolation and identification, serological detection, immuno-electron microscopy, enzyme-linked immune sorbent assay (ELISA), lateral flow assay (LFA), and Polymerase Chain Reaction (PCR) techniques, have been applied for virus detection [13–18]. Virus isolation and identification is a confirmation method for virus detection. However, the method is time consuming, greatly hindering its application in rapid clinical detection [19]. Immunoassay-based methods, such as ELISA, have been widely used. The problem of such technique is the requirements of special antibodies, whose production is time consuming and exhausting [20]. Immuno-electron microscopy requires sophisticated instrumentation and high amount of virus; thus, it is not feasible in clinical diagnosis [21]. Different from these methods, PCR is a common used technique in molecular biology [22]. It can exponentially amplify a single copy or few copies of a specific DNA segment. It has been widely used in clinical laboratory research for a broad variety of pathogen detection because of its high sensitivity, non-strict detection conditions, strong specificity, high speed, and safety [20, 23].

The m-PCR refers to the PCR reactions, in which two or more primer pairs are used in a PCR reaction tube, and multiple nucleic acid fragments are amplified simultaneously [24]. Compared with uniplex PCR, m-PCR shows unparalleled advantages, including high amplification efficiency, time-saving, and high throughput [25, 26]. More importantly, this approach can differentially diagnose various viruses at the same time; it is an effective method for rapid diagnosis of mixed-virus infection in clinical detection [19, 27]. We aimed to develop and optimize a single-step m-PCR method capable of detecting and differentiating seven major duck viruses, including AIV, NDV, NDPV, DHAV, DTMUV, FAdV, and DEV.

Results
Optimization and establishment of the m-PCR method
A D-optimal design was used to optimize the m-PCR method with 22 runs performed in one randomized batch (in duplicate measurements). As an example, the three-dimension response surface curves of DHAV are shown in Fig. 1a. Similarly, 4D plots indicated the interaction between the four factors (Fig. 1b). We obtained the ultimate optimum parameters, considering the economic perspective, with the necessary compromise, with annealing temperature at 57 °C, Mg²⁺ concentration at 4 mM, Taq DNA Polymerase concentration at 0.05 U/μL, and dNTP concentration at 0.32 mM. With the ultimate optimum primers and parameters (Table 1), we successfully established the m-PCR method, which could also effectively amplify genes of duplex, triplex, and even septuplet (Fig. 2).

Specificity of the m-PCR method
The specificity of the m-PCR method was evaluated with these seven viruses and DRV, APMV-4, APMV-8, DHAV-3, Escherichia coli, Salmonella, Riemerella anatipestifer, Pasteurella multocida, and Clostridium perfringens that may infect ducks. As shown in Fig. 3a, the band for each virus was clear for m-PCR analysis, similar to that of uniplex PCR. Furthermore, although other viruses or bacteria genomes were mixed in the sample pool, only the genes of these seven viruses were specifically amplified; no amplification occurred with those interfering genomes (Fig. 3b). The sequencing results further demonstrated the good specificity of the m-PCR.

Sensitivity of the m-PCR method
Each of the seven recombinant plasmids and the mixed plasmds were diluted from 1 × 10⁸ to 1 × 10⁰ copies/μL by 10-fold gradient dilution. Then, m-PCR was performed. As shown in Fig. 4, when single template was applied, the detection limits were 1 × 10² copies/μL for DEV, DTMUV, and NDPV, 1 × 10³ copies/μL for FAdV and DHAV, and 1 × 10⁴ copies/μL for NDV and AIV. When seven DNA templates were mixed, the detection limit of each virus was 1 × 10⁴ copies/μL.

Reproducibility of the m-PCR method
The results of reproducibility of the uniplex PCR and m-PCR method are shown in Additional file 1: Figure S1. Uniplex PCR was conducted with plasmids of concentrations of 10⁵, 10⁴, 10³, and 10² copies/μL. and 10⁵, 10⁴,
Fig. 1 Response surface plots for DHAV. Different response plots for DHAV for various combinations of the investigated parameters. 

**a** Different combinations of four factors to form the three-dimension response surface curves; Mg: Mg$^{2+}$ concentration, Tm: the annealing temperature, DNA pol: Taq DNA polymerase concentration, dNTP: dNTP concentration. 

**b** 4D plots indicated the interaction between the four factors.
10^6 copies/μL for m-PCR method, confirming that the proposed method was reliable.

Co-infection model and clinical sample detection
As shown in Fig. 5a and Additional file 2: Figure S2, we simulated duplex infections of different combinations of viruses in the same concentration. Simultaneously, triple, quadruple, and quintuple infections of different combinations of viruses in the different concentrations were detected (Fig. 5b). In addition, a total of 60 clinical samples were examined using the developed m-PCR method and uniplex PCR. The result showed that 11 samples were virus positive, including 1 positive sample for FAdV, 4 positive samples for DHAV, 2 positive samples for DEV, 2 positive samples for DTMUV, 2 positive samples for AIV, and 1 positive sample for NDPV. The positive rate was 18%. Among these 11 positive samples, sample 8 was co-infected with DHAV and AIV (Table 2). Moreover, these results were further confirmed by the primers published previously.

Discussion
Because of the development of mixed culture models, enhanced mobility of humans and animals, and environment pollution, viral infection in ducks is getting serious in recent years [1]. Among the popular viruses that infected ducks, AIV, NDV, NDPV, DHAV, DTMUV, FAdV, and DEV were very common, seriously endangering ducks’ health and causing great economic losses [5, 7, 10, 11, 28–30]. Although virus isolation is considered to be the gold standard for viral detection, it is unsuitable for clinical rapid detection [31]. Other methods, such as serological detection, immuno-electron microscopy, ELISA, and real-time PCR, have been applied for detecting these viruses [32–34]. However, rapid detection of the seven viruses simultaneously in ducks has not been reported. Therefore, we aim to develop a specific, sensitive, and rapid m-PCR method for clinical diagnosis of co-infection of duck virus. In establishing a successful multiplex PCR method, primer design is the key factor [35]. Mutual interference, such as mismatch and dimer

| Viruses | Primer sequences (5’-3’) | Targeted genes | Position | Accession no. | Product size (bp) | Working concentration (μM) |
|---------|--------------------------|----------------|----------|---------------|------------------|---------------------------|
| FAdV   | F: CAACAGCCCTCTGTACCCAG  | Hexon          | 839–858  | AJ459805      | 102              | 0.12                      |
|         | R: CCGATGTTGGGCTGAGGAG    |                | 921–940  |               |                  |                           |
| DHAV   | F: CTTCCACCTCCTGCTCCCAAC | VP1            | 510–528  | KJ606043      | 140              | 0.12                      |
|         | R: TTGCTCCCACATCCTCCTCA   |                | 629–649  |               |                  |                           |
| DEV    | F: ATCGCATGTTGAGCTTGGTTG  | UL2            | 726–745  | EU885419      | 172              | 0.16                      |
|         | R: AGACACCGGTAGTGAGTGG    |                | 879–897  |               |                  |                           |
| DTMUV  | F: AATCCGCTTAGTGCTTGG    | Envelope       | 432–449  | AB110495      | 288              | 0.28                      |
|         | R: AGTCTGCGCCATGATGATAT   |                | 701–719  |               |                  |                           |
| NDV    | F: CACCGGCAACCTTATCTGCA   | Fusion         | 853–872  | M24701        | 330              | 0.28                      |
|         | R: AGTGCAGCCTCAGTCTTATGTA |                | 1163–1182|               |                  |                           |
| AIV    | F: GCGCACTACTACAAACCA    | Matrix          | 521–539  | DQ064401      | 435              | 0.24                      |
|         | R: CTGCTGTTCATGCTCGTAAT   |                | 937–955  |               |                  |                           |
| NDPV   | F: TATGTCCTGGGCGTCCGCTAC | VP3            | 448–467  | EF014903      | 516              | 0.24                      |
|         | R: AGCTGACACAGGTCAGGTT    |                | 944–963  |               |                  |                           |
between different primers is common when different pairs of primers are mixed [37]. Within this experiment, a 5-plex PCR was first developed. When we enlarged the system into 7-plex PCR, with the addition of two pairs of new primers, interfering bands appeared, indicating that the original primers used in 5-plex PCR is unsuitable for 7-plex PCR. After careful optimization, seven pairs of primers for the viruses were found at last, with which seven viruses could be amplified separately. Moreover, these amplification products were consistent with the target gene fragments. The results proved that the m-PCR method could detect the seven viruses flexibly and that a usable m-PCR method has been established successfully.

Many other factors may affect the amplification efficiency. For example, even the smallest deviation of annealing temperature could lead to the nonspecific amplification [38]. Similarly, if Mg$^{2+}$ in the reaction is extremely high, then the method lacks specificity, whereas if it is extremely low, then bad amplification can be expected [39]. Hence, by using the design of experiments, we optimized four pivotal parameters including annealing temperature, Mg$^{2+}$ concentration, Taq DNA Polymerase concentration, and dNTP concentration that may influence the PCR [39]. A D-optimal design with 22 runs performed in one randomized batch (in duplicate measurements) was used for this optimization. The peak area was carefully investigated. In general, if a factor was considered critical, then the level, at which a factor gave the highest response, was used in the final experimental protocol [40]. In cases of conflicting results, the favorable level of a factor was decided in terms of the number of parameters from which the highest response was obtained. Therefore, a compromise was made to obtain the optimum parameters.

AIV, NDV, NDPV, DHAV, DTMUV, FAdV, DEV, and other viruses and bacteria that may infect ducks were utilized to evaluate the specificity of the m-PCR method. The result indicated that the m-PCR method produced neither cross reactions among these seven viruses nor nonspecific reactions with other common duck pathogens when all the DNA templates existed in the sample pool [41], thereby further demonstrating that the proposed primers were highly specific. The circulating NDPV is originated from the GPV lineage [30], thus the developed m-PCR method can also identify GPV at the same time. In addition, mix infection of different pathogens is common in clinical practice [42]. When ducks are infected with one virus, they are susceptible to others [9, 43], which may have a more powerful pathogenicity and bring about more serious economic losses [21]. Therefore, co-infection was performed to determine the practicality of the m-PCR method. The assay showed that the result of co-infection experiment was consistent with the grouping of seven viruses, which manifested that the m-PCR method could flexibly and specially detect several random virus combinations. The detection limit of the m-PCR method was 1 × 10^4 copies/μL, which was higher than that of the uniplex PCR. The possible reason might be the mutual interference between multiple templates and primers [44]. However, considering its convenience and high throughput in sample analysis, the m-PCR method is more applicable. In addition, the m-PCR method showed highly consistent results with uniplex PCR, demonstrating the good accuracy of this method. The assay was performed to assess the repeatability and reproducibility of the method. This result suggested that the m-PCR method was highly reliable and stable [45].
Fig. 4 Sensitivity of the m-PCR method. Templates of pMD-FAdV (a), pMD-DHAV (b), pMD-DEV (c), pMD-DTMUV (d), pMD-NDV (e), pMD-AIV (f), pMD-NDPV (g), and mixture of seven virus plasmids (h). M, DL600 marker; NC, negative control. Plasmids were diluted from $10^8$ to $10^0$ DNA copies/μL.
The result of the field analysis with the developed m-PCR method was consistent with that of the uniplex PCR and standard PCR. Furthermore, all clinical positive samples were confirmed by sequencing. The samples collected from agricultural markets did not produce any specific band, and these positive samples were from different duck farms. Thus, the samples collected from the markets were healthy and safe. Furthermore, this result demonstrated that the m-PCR method was specific, sensitive, rapid, and practical in laboratory and clinical diagnoses.

Conclusion

We established a multiplex PCR method with high specificity, good sensitivity, and reproducibility that could detect and differentiate seven major viruses causing duck diseases, including AIV, NDV, NDPV, DHAV, DTMUV, FAdV, and DEV. Thus, preventive measures can be implemented as early as possible to reduce economic losses.

Methods

Pathogen

DHAV (attenuated vaccine strain A66, purchased from Chengdu Tecbond Biological Products Co., Ltd. Cat. no. 220012214), DTMUV (attenuated vaccine strain WF100, purchased from QiLu Animal Health Products Co., Ltd. Cat. no. 1502522), and DEV (attenuated vaccine strain C-KCE, purchased from Guangxi Liyuan Biological Co., Ltd. Cat. no. 200352023) were isolated from the vaccine strain; AIV (H9N2), NDV, NDPV, goose parvovirus (GPV) and FAdV (serotype 4) were isolated from the clinical samples. Duck reovirus (DRV), Avian paramyxoviruses (APMV-4) Escherichia coli (wild-type, serotype

Table 2 Result of the clinical positive samples detected by m-PCR and uniplex PCR

| Positive sample | Viruses   |
|----------------|----------|
|                | FAdV | DHAV | DEV | DTMUV | NDV | AIV | NDPV |
| 1               | (-)   | (+)  | (-) | (-)   | (-) | (-) | (-)  |
| 2               | (-)   | (-)  | (-) | (+)   | (-) | (-) | (-)  |
| 3               | (-)   | (-)  | (+) | (-)   | (-) | (-) | (-)  |
| 4               | (-)   | (-)  | (-) | (-)   | (-) | (+) | (-)  |
| 5               | (-)   | (-)  | (-) | (-)   | (-) | (+) | (-)  |
| 6               | (-)   | (+)  | (-) | (-)   | (-) | (+) | (-)  |
| 7               | (+)   | (-)  | (-) | (-)   | (-) | (+) | (-)  |
| 8               | (+)   | (-)  | (-) | (-)   | (-) | (+) | (-)  |
| 9               | (-)   | (-)  | (+) | (-)   | (-) | (+) | (-)  |
| 10              | (-)   | (+)  | (+) | (-)   | (-) | (+) | (-)  |
| 11              | (-)   | (+)  | (+) | (-)   | (-) | (+) | (-)  |
| Positive rates (%) | 1/60 (1.7) | 4/60 (6.7) | 2/60 (3.4) | 2/60 (3.4) | 0/60 (0) | 2/60 (3.4) | 1/60 (1.7) |

Note: Data in the racket showed the result detected by uniplex PCR
The nucleic acid of the seven types of viruses was synthesized using the Viral RNA/DNA Extraction Kit (Sangon Biotech, China) and dissolved with nuclease-free water. The RNAs of DHAV, DTMUV, AIV, and NDV were reverse-transcribed into cDNA by the Reverse Transcription Kit (Thermo Scientific, USA). The concentration and purity of each genome were determined by spectrophotometry (Thermo Scientific, USA). The DNA/cDNA was stored at −20 °C.

**Nucleic acid extraction**

The nucleic acid of the seven types of viruses was extracted using the Viral RNA/DNA Extraction Kit (Sangon Biotech, China) and dissolved with nuclease-free water. The RNAs of DHAV, DTMUV, AIV, and NDV were reverse-transcribed into cDNA by the Reverse Transcription Kit (Thermo Scientific, USA). The concentration and purity of each genome were determined by spectrophotometry (Thermo Scientific, USA). The DNA/cDNA was stored at −20 °C.

**Primer design**

The complete gene sequences of the DHAV, DTMUV, DEV, NDV, AIV, FAdV, and NDPV strains were acquired from the GenBank, and the conserved region of viral genes sequences was aligned by DNAMAN (LynnonBiosoft, USA). We designed seven pairs of specific primers for each virus by Primer Premier 5 (Premier, Canada), according to the results of the sequence alignment (Additional file 3: Figure S3). Primers that were listed in Table 1 (BLAST results of primers are shown in Additional file 4: Table S1) and synthesized by Sangon Biotech. The target genes include Matrix (M) gene for AIV, Fusion (F) gene for NDV, viral structural protein (VP3) gene for NDPV, viral structural protein (VP1) gene for DHAV, Envelope (E) gene for DTMUV, Hexon (H) gene for FAdV, and unique long region 2 (UL2) gene for DEV. These genes were the highly conserved region of these viruses (Table 1).

**Standard plasmid preparation**

Seven recombinant plasmids were used as the positive controls. These were reconstructed in this study. Briefly, specific target fragments were amplified first with the primers (Table 1), and then these fragments were cloned into the pMD-18T vector (TaKaRa, China) to obtain the recombinant plasmids pMD-AIV, pMD-NDV, pMD-NDPV, pMD-DHAV, pMD-DTMUV, pMD-FAdV, and pMD-DEV. The concentration of the recombinant plasmids was determined by NanoDrop-1000 (Thermo Fisher NanoDrop, USA), and the plasmid copy number was calculated using the following formula: copy number (copies/μL) = NA (copies/mol) × concentration (g/μL) / MW (g/mol), where NA is Avogadro’s number, and MW is the base number times 340 [21].

**Uniplex PCR**

The total volume of each reaction system was 25 μL, containing 2.5 μL 10× Taq Buffer (Mg²⁺ free), 4 μL 25 mM MgCl₂, 0.75 μL dNTP Mix (10 mM each), 0.25 μL 5 U/μL Taq DNA Polymerase (Vazyme, China), 1 μL of the forward primer, 1 μL of the reverse primer, 1 μL of the single-virus template, and 14.5 μL of ddH₂O. The PCR procedure was performed as follows: pre-denaturation at 95 °C for 5 min, followed by denaturation at 95 °C for 30 s, annealing at 57 °C for 30 s, extension at 72 °C for 40 s, 35 cycles, with a final extension at 72 °C for 10 min. The PCR products were analyzed by 1.5% agarose gel electrophoresis. Double distilled water was used as the blank control.

**Experimental design for multiplex PCR method**

The m-PCR method was optimized using a D-optimal design consisting of 22 experiments. Four factors were considered including annealing temperatures (48 °C–62 °C), Mg²⁺ concentrations (1–6 mM), Taq DNA Polymerase concentrations (0.02–0.06 U/μL), and dNTP concentrations (0.08–0.48 mM). As a response, intensity of the PCR amplified bands (quantitated using the Image J software) was used for statistical analysis [39, 46]. All analyses were performed using MODDE 12.1 software (Umetrics, Sweden). The relationship between the response Y and the variables Xᵢ, Xⱼ, Xᵢ, Xⱼ was expressed as $Y = \beta₀ + \beta_iX_i + \beta_jX_j + \beta_0X_iX_j + \beta_0X_i^2 + \beta_0X_j^2 + ... + \epsilon$, where $\betaₖ$ represents the regression coefficients, and $\epsilon$ was the experimental error. The linear coefficients $\beta_i$ and $\beta_j$ were the quantitative effect of the respective variables. The cross coefficient $\beta_{ij}$ measured the interaction between the variables, and the square terms of $\beta_iX_i^2$ and $\beta_jX_j^2$ described the non-linear effects on the response [47].

**Sensitivity and specificity of the m-PCR method**

The mixed plasmids were diluted from 1 × 10⁸ copies/μL to 1 × 10⁶ by 10-fold gradient dilution and were used to detect the sensitivity of the m-PCR method. The specificity of m-PCR was tested using other common duck viruses and bacteria (including DRV, APMV-4, APMV-8, DHAV-3, Escherichia coli, Salmonella, Riemerella anatipestifer, Pasteurella multocida, and Clostridium perfringens).

**Repeatability and reproducibility of the m-PCR method**

The repeatability of the method was evaluated at three concentrations on the same day, whereas the reproducibility study was performed at three concentration levels on three successive days. The concentrations of seven recombinant plasmids were 1 × 10⁴, 1 × 10⁵, and 1 × 10⁶ copies/μL [44]. In addition, the repeatability and reproducibility of the uniplex PCR were carried out with the plasmid concentration of 1 × 10⁵, 1 × 10⁶, 1 × 10⁷, and 1 × 10⁷ copies/μL.
Co-infection model and clinical specimen analysis

Co-infection analysis was designed to determine the practicality of the m-PCR method [45]. Furthermore, 60 clinical samples (40 tissue samples and 20 throat swabs) were collected from the duck farms and live poultry markets in Jiangsu province (These samples collection were permitted by the owner of the animals and suffering were minimized to these animals). The established m-PCR method was used for PCR amplification. Results were further confirmed by uniplex PCR and standard or published PCR methods [1, 48–52]. Then, the positive products of PCR amplification were sequenced to confirm the detection results.

Additional files

Additional file 1: Figure S1. Repeatability and reproducibility of the uniplex PCR and m-PCR methods. The plasmid concentration for reproducibility analysis of the uniplex PCR for FAdV (A) and DHAV (B) was set at 10^3, 10^4, and 10^5 copies/μL, and that for DEV (C), DTMUV (D), NDV (E), AIV (F), and NDPV (G) were set at 10^3, 10^4, and 10^5 copies/μL. The plasmid concentration for reproducibility analysis of the m-PCR (H) was set at 10^3, 10^4, and 10^5 copies/μL. (JPG 229 kb)

Additional file 2: Figure S2. Co-infection analysis by m-PCR method. Detection result of co-infection of two viruses with plasmid concentration of 1 × 10^4 copies/μL. Lanes 1 and 9, positive control; Lanes 2 and 8, NDV + DTMUV; Lanes 3 and 7, AIV + DEV; Lanes 4 and 6, DHAV+NDPV; Lane 5, AIV + FAdV; M, DL600 marker; NC, negative control. (JPG 155 kb)

Additional file 3: Figure S3. Alignment results of the seven viruses’ primers. (DOXX 791 kb)

Additional file 4: Table S1. BLAST results of the seven viruses’ primers. (DOXX 15 kb)

Abbreviations

AIV: Avian influenza virus; C. perfringens: Clostridium perfringens; ddH2O: Double distilled water; DEV: Duck enteritis virus; DHAV: Duck hepatitis A virus; dNTP: Deoxy-ribonucleotide triphosphate; DRV: Duck adenovirus; FAO: Food and Agriculture Organization; GPV: Goose parvovirus; H. hexon; LFA: Lateral flow assay; M: DL600 marker; M. Matrix; Mg2+: Magnesium ion; m-PCR: Multiplex PCR; MW: Base number times; NA: Avogadro’s number; NC: Negative control; NDPV: Novel duck parvovirus; NDV: Newcastle disease virus; P. multocida: Pasteurella multocida; PCR: Polymerase Chain Reaction; RA: Remmerella anaplasitfer; SE: Salmonella; UL2: Unique long region 2; VP1: Viral structural protein; VP3: Viral structural protein

Acknowledgements

Not applicable.

Funding

The design of the m-PCR method was supported by the National Key Research and Development Program (2016YFD0500800). The clinical samples collection was supported by Key Program of Science and Technology Planning of Guangdong Province (2017B020202010) and the Natural Science Foundation of Jiangsu Province (BK20161452). The analysis and interpretation of data was supported by Forestry Science and Technology Innovation and Promotion Project of Jiangsu Province (LYKJ201822) and Fundamental Research Funds for the Central Universities (Y2016000146, KY201848). The writing of the manuscript was supported by Priority Academic Program Development of Jiangsu Higher Education Institutions, and NAU International Cooperation and Cultivation Project (2018-AF-20).

Availability of data and materials

All data generated or analyzed during this study are included in this published article and its additional files.

Authors’ contributions

LPY and SQS designed the study and approved the manuscript. MY, XYZ and DNX developed the multiplex PCR method, analyzed data and drafted the manuscript. MY and YFG collected clinical samples. All authors read and approved the final manuscript.

Ethics approval and consent to participate

This study was performed in accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the Ministry of Health, China. The protocol of the current study was reviewed and approved by the Institutional Animal Care and Use Committee of Nanjing Agricultural University (approval no. SYXK 2017–0007). Written informed consent to use 60 clinical samples, which were collected from the duck farms and live poultry markets, were obtained from the owner of the animals. All efforts were made to minimize animal suffering during sample collection.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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Received: 15 October 2018 Accepted: 25 February 2019

Published online: 01 April 2019

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