Molecular Identification of a Velogenic Newcastle Disease Virus Strain Isolated from Egypt

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ABSTRACT

Newcastle Disease Virus (NDV) is still a major concern for the Egyptian poultry industry in spite of the mass vaccination programs implemented from a long years ago. The current study aimed to carry out the molecular identification of surface glycoprotein genes of NDV field strain isolated from the Giza governorate, Egypt. Tracheae were collected from 10 broilers NDV-vaccinated chicken flocks (at least three samples from each flock) suffering from mild to moderate respiratory symptoms; with mortalities varying from 10-40% during October 2019. Only five samples showed HA positive activity after propagation in specific pathogen-free embryonated chicken eggs and only one sample was positive for Avian avulavirus 1 by real-time reverse transcription-PCR. Sequencing for the cleavage site of the F protein gene of the positive isolate showed the typical known sequence of velogenic NDV strains (1RRQKRF117). Phylogenetic analysis of both F and HN genes showed high similarity and close relation to Chinese strains of Genotype VII and more specifically subtype VId, suggesting the role of migratory wild birds in NDV evolution in Egypt. In conclusion, further epidemiological and surveillance studies are strongly recommended to define the exact role of migratory wild birds in NDV evolution in Egypt.

Keywords: Broilers, Newcastle Disease, Poultry industry, Velogenic

INTRODUCTION

Newcastle Disease (ND) is one of the most important viral diseases affecting poultry which is caused by Avian avulavirus 1 (APMV-1) (Abd El Aziz et al., 2016). The natural hosts of ND virus (NDV) include chickens, turkeys, ducks, geese, pigeons, quail, pheasants, guinea fowl, ostriches, and several species of wild birds (Wang et al., 2015).

ND as an acute viral infectious disease affects domestic poultry regardless of gender and age (Saad et al., 2017) and causes great economic losses, especially in developing countries (Westbury, 2001). Production inefficiencies are considered as a greater concern compared to mortality losses in breeders and layers flocks while mortalities usually reported to be more significant in broilers (Shahid Mahboob et al., 2020). Many ND outbreaks were reported in the past years around the world, as in Japan (Mase et al., 2002), in Brazil (Marks et al., 2014), in China (Kang et al., 2014), in South America (Diel et al., 2012) and in Malaysia (Jaganathan et al., 2015). NDV was recorded in Egypt since 1942 (Daubeny and Mancy, 1947) and has been reported ever since (Hussein et al., 2000; Mohamed et al., 2011; Selim et al., 2018). Recently in Egypt, NDV outbreaks have been reported in both vaccinated and non-vaccinated flocks (Abd El Aziz et al., 2016; Ewies et al., 2017). A subclinical infection manifested by respiratory, intestinal, and nervous symptoms, with mortalities up to 100% may be a result of NDV infection according to virus strain pathogenicity in infected birds. Based on the pathogenicity of the virus, NDV strains can be categorized into three main types; lentogenic, mesogenic, and velogenic strains. NDV can be classified into two classes; class I and class II. NDV isolates of class I are grouped into one genotype, whereas NDV isolates of class II are grouped into at least eighteen genotypes, some with subgenotypes. Genotype VII viruses are responsible for the fourth panzootic that has spread from Asia, Africa, Europe, and has even been isolated in South America, which continues today (Dimitrov et al., 2016). The NDV is an enveloped virus that has a linear, single-stranded RNA genome of negative
polarity; with a genome length of about 15.2 kb (Aldous et al., 2003; Ashraf et al., 2016). The genome of NDV consists of 15,186, 15,192 nucleotides or 15,198 nucleotides that contains six genes coding six structural and two non-structural proteins including an RNA-directed RNA polymerase (L), hemagglutinin-neuraminidase protein (HN), fusion protein (F), matrix protein (M), phosphoprotein (P), and nucleoprotein (N). Both F with HN proteins play a collective role in NDV infection processes. The fusion protein is the most important key in the NDV virulence determining process (Peeters et al., 1999). Mutations affecting NDV viral genome which alter its biological properties and virulence, in addition to altered immunity, and improper vaccination processes can increase the incidence of NDV outbreaks in vaccinated flocks (Kattenbelt et al., 2006). Virulence of ND can be distinguished on the basis of the cleavage site sequence of the F protein (Selim et al., 2018). HN is one of the membrane glycoproteins, through its neuraminidase (NA) activity it mediates attachment to sialic acid-containing receptors (Wang et al., 2015). Recently, the molecular identification and phylogenetic analysis of any new NDV isolates become an important and usual approach to find out which of the applied control measures needs to be improved (Fringe et al., 2012; Hassan et al., 2016). Sequence analysis of mainly F and of HN proteins genes - two surface glycoproteins- is wildly used for molecular identification of NDV isolates. Brevity, the current applied NDV vaccination programs consist of live and/or inactivated genotype I or II NDV or genetically modified vaccines depending on flock age and type.

In the present study, analysis of nucleotides sequences of F and HN genes were done for a recently isolated NDV strain obtained from samples collected from different chicken flocks showing mild to severe respiratory symptoms with variable mortality rates in Giza governorate, Egypt.

MATERIALS AND METHODS

Ethical approval
Institutional, national, and international animal care guidelines were followed.

Sampling and samples history
Tracheae (at least three samples from each flock) from 10 freshly dead broilers, NDV vaccinated chicken flocks suffering from mild to moderate respiratory symptoms; with mortalities varying from 10 to 40 % as well as a range of NDV indicative postmortem lesions at Giza governorate during October of 2019. Tracheae from the same flock kept together for isolation. Samples history mentioned in Table 1.

Table 1. History of flock sampled in the present study

<table>
<thead>
<tr>
<th>Sample</th>
<th>Birds No/Flock</th>
<th>Mortalities (%)</th>
<th>Age of birds (day)</th>
<th>NDV vaccination</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>10,000</td>
<td>12.5 %</td>
<td>35</td>
<td>Twice, live LaSota</td>
</tr>
<tr>
<td>S2</td>
<td>8,000</td>
<td>14 %</td>
<td>55</td>
<td></td>
</tr>
<tr>
<td>S3</td>
<td>11,000</td>
<td>37.6 %</td>
<td>24</td>
<td></td>
</tr>
<tr>
<td>S4</td>
<td>4,000</td>
<td>35.8 %</td>
<td>26</td>
<td></td>
</tr>
<tr>
<td>S5</td>
<td>15,000</td>
<td>22.4 %</td>
<td>22</td>
<td></td>
</tr>
<tr>
<td>S6</td>
<td>2,000</td>
<td>10.7 %</td>
<td>23</td>
<td></td>
</tr>
<tr>
<td>S7</td>
<td>1,000</td>
<td>34.5 %</td>
<td>35</td>
<td>Once, live LaSota</td>
</tr>
<tr>
<td>S8</td>
<td>3,000</td>
<td>21.3 %</td>
<td>40</td>
<td>Twice, live LaSota</td>
</tr>
<tr>
<td>S9</td>
<td>4,000</td>
<td>39.1 %</td>
<td>41</td>
<td></td>
</tr>
<tr>
<td>S10</td>
<td>12,000</td>
<td>35.7 %</td>
<td>33</td>
<td></td>
</tr>
</tbody>
</table>

Isolation
Virus isolation was done from tracheal swabs after immersion in Phosphate-Buffered Saline (PBS) mixed with gentamycin antibiotic (50 μg/ml) and mycostatin (1000 units/mL). Swabs from different birds from the same flock were immersed in the same PBS solutions. Samples were named numerically as sample 1 (S1): sample 10 (S10).

PBS-containing samples were clarified by centrifugation at 5000 rpm for 15 minutes. A 200 μl of supernatant fluid from each sample was inoculated into the allantoic cavity of five 10-day-old Specific Pathogen Free Embryonated Chicken Eggs (SPF-ECE). Allantoic fluid from each egg was harvested 3 to 5 days post-inoculation and was tested for hemagglutination (HA) activity by rapid slide HA test. HA negative samples were submitted for two blind passages of SPF-ECE. Collectively, samples that showed HA positive activity were kept for further molecular identification (OIE manual, 2018).

Viral RNA extraction
Viral RNA from HA positive allantoic fluid was extracted using Pure Link® (Invitrogen, USA) RNA Mini Kit following the manual’s instruction.

Real-time reverse transcription-polymerase chain reaction
Real-time Reverse Transcription PCR (RT-qPCR) was performed in one step. Using TOPreal™ One-step - SYBR Green with low ROX - RT qPCR Kit (Enzymomics, Korea) according to the manufacturer’s instructions and
using the CFX96 Touch real-time PCR detection system (Bio-Rad Laboratories, USA). Primers used were designed according to Wise et al. (2004) which are specific for the matrix protein gene of APMV-1 viruses selected from a conserved region of the M gene (Table 2).

The thermal conditions were as follows; reverse transcription at 50 °C for 30 mins followed by 10 mins at 95 °C for reverse transcriptase inactivation and initial denaturation. Then, followed by 40 amplification cycles of 95 °C denaturation for 5 s, 52 °C annealing for 10 s, and 60 °C extension for 30 s.

Melting curve analysis was performed to determine the specificity of amplification as follows: 95°C denaturation for 10 s, 65°C annealing for 5 s, and heating to 95 °C with an increment 0.5 °C for 0.05 s.

The melting temperature (Tm) of melting curves and Cp values were calculated using the Bio-Rad CFX manager 3.1 software (Figure 1).

Table 2. Primers used for Newcastle disease virus detection using RT-qPCR.

<table>
<thead>
<tr>
<th>Primer Name</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>F Primer</td>
<td>5’-ATGGGCTCCAAACCTTCT-3’</td>
</tr>
<tr>
<td>R primer</td>
<td>5’-CCTGAGGAGGCAATTGCTA-3’</td>
</tr>
</tbody>
</table>

**F and HN genes amplification**

Positive NDV RNA samples (by RT-qPCR) samples were subjected to one-step RT-PCR using SuperScript™ III One-Step RT-PCR System with Platinum™ Taq DNA Polymerase according to the manufacturer’s instructions to amplify full-length F protein gene and HN protein gene using two sets of primes kindly provided by Dr. Mohammed Rohaim, Virology Department, Cairo University, Egypt (Table 3) and using the ProFlex PCR thermal cycler (Applied biosystem, USA).

Thermal amplification conditions were as follows; reverse transcription at 50 °C for 30 min followed by initial denaturation for 2 min at 94 °C. Then followed by 40 cycles of denaturation at 94 °C for 15 s, annealing at 65 °C for 30 s for F gene while 51 °C for 30 s for HN gene, and extension at 68 °C for 120 s followed by one cycle of final extension at 68 °C for 5 min.

PCR products were analyzed by agarose gel electrophoresis (1%) and then purified using a QIAquick Gel Extraction Kit (Qiagen) following the manufacturer’s instructions.

<table>
<thead>
<tr>
<th>Target gene</th>
<th>Primer Name</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>F protein</td>
<td>Fus-F</td>
<td>5’-ATGGGCTCCAAACCTTCT-3’</td>
</tr>
<tr>
<td>HN protein</td>
<td>Hae-F</td>
<td>5’-CATGACGCCGCGGTTAAC-3’</td>
</tr>
<tr>
<td>HN protein</td>
<td>Hae-R</td>
<td>5’-CTAAACTCTATCATCCTTG-3’</td>
</tr>
</tbody>
</table>

**Sequencing**

Sequencing of the purified RT-PCR products was done by the Bigdye Terminator V3.1 cycle sequencing kit (Perkin-Elmer, Foster City, CA) and Applied Biosystems 3130 genetic analyzer machine (ABI, USA).

**Genetic alignment**

The quality of obtained F and HN genes sequences were checked, assembled, edited using Bioedit software version 7.0.4.1 (Hall, 1999), and submitted to GenBank using BankIt tool of the GenBank (http://www.ncbi.nlm.nih.gov/WebSub/?tool=genbank), with accession numbers MN905162 and MN905163, respectively.

**Phylogenetic analysis**

The tree was constructed using the neighbor-joining method; bootstrapping at 500 repeats using Mega 6 software version 7.0.26 (Tamura et al., 2013).

**RESULTS**

**Hemagglutination activity**

After three blind passages only S3, S4, S7, S9, and S10 samples were positive for hemagglutination activity. S7 and S10 were positive for HA after the 1st egg passage, S3, and S4 were positive for HA after the 2nd egg passage, and S9 was positive for HA after the 3rd egg passage. RNA from 5 positive HA samples were sent for one-step RT-qPCR.

**NDV detection by RT-qPCR**

Only S4 was positive for *Avian avulavirus* 1 by RT-qPCR with a threshold cycle (CT) of 29.34 with a starting quantity of 3.033 log10 in comparison with a standard curve (Figure 2) with melting peak at 79 °C (Figure 3 and 4).
Amplification of full F and full HN proteins genes by RT-PCR

RT PCR products gel electrophoresis revealed the expected and correct size bands for full-length F and HN proteins genes.

Genetic and phylogenetic analysis

F protein gene

Blasting of sequence results obtained for the full F protein gene showed similarities with Chinese genotype VII strains with similarities varying from 95.5% to 97.28% and with many Egyptian isolates varying from 94.5% to 95.5%. The phylogenetic tree of the full F protein gene showed that S4 isolate is closely related to genotype VII subtype D (Figure 5).

HN protein gene

Blasting of sequence results obtained for the full HN protein gene showed similarities with Chinese genotype VII strains with similarities varying from 95.69% to 98.72% and with some Egyptian isolates varying from 94.9% to 95.45%. The phylogenetic tree of the full HN protein gene showed that S4 isolate is closely related to the Chinese genotype VII (Figure 6). Three-dimensional structure of F and HN monomer for S4 isolate was created by SWISS-Model modeling online server and visualized by PyMOL program version 2.3.4 (Figure 7 and 8).

Figure 1. Thermal conditions applied at RT-qPCR and for the melting curve.

Figure 2. Threshold cycles of tested samples, green lines represent positive control samples (standard curve samples), the red line represents positive for Avian avulavirus sample (S4) appeared after 29.34 CT, and pink lines represent the negative for Avian avulavirus samples.
Figure 3. Melting curve of tested samples, green lines represent positive control samples (standard curve samples), the red line represents positive for *Avian avulavirus* sample (S4), and pink lines represent the negative for *Avian avulavirus* samples.

Figure 4. Melting peak of tested samples, green lines represent positive control samples (standard curve samples), the red line represents positive for *Avian avulavirus* sample (S4) showed a different melting peak, and pink lines represent the negative for *Avian avulavirus* samples.
Figure 5. Neighbor-joining phylogenetic tree of the full-length F gene of Egyptian isolate of Newcastle disease virus (NDV) (S4) in comparison to other NDV strains from GenBank. Bootstrap values are shown above the branches. S4 isolate is indicated by a solid green circle.

Figure 6. Neighbor-joining phylogenetic tree of the full-length HN gene of Egyptian isolate of Newcastle disease virus (NDV) (S4) in comparison to other NDV strains from GenBank. Bootstrap values are shown above the branches. S4 isolate is indicated by a solid green circle.

Figure 7. 3D structure for F protein of Newcastle disease virus (S4 isolate) created by SWISS-Model modeling online server and visualized by PyMOL program version 2.3.4. Red color represents the cleavage site.

Figure 8. 3D structure for HN protein of Newcastle disease virus (S4 isolate) created by SWISS-Model modeling online server and visualized by PyMOL program version 2.3.4.
DISCUSSION

In the current study, only five samples (50% samples) showed HA positive activity indicating infection with a hemagglutinating virus. To confirm NDV infection, RT-qPCR was performed using the HA positive samples.

Only S4 isolate was positive for NDV using universal primers for APMV-1. Negative RT-qPCR results for S3, S7, S9, and S10 may indicate an infection with another hemagglutinating virus-like avian influenza H9 or H5; however, history of mortalities and symptoms severity indicated H9 infection mixed with other respiratory pathogens other than H5 (Hussein et al., 2014; Sedeik et al., 2018). The most important pathogenicity indicator for NDV is the F protein gene sequence analysis mainly for cleavage site in which velogenic strains have polybasic amino acid sequences; therefore, molecular identification and phylogenetic analysis of the F gene is a major determinant of NDV virulence instead of conventional methods (Mohamed et al., 2011; Damena et al., 2016). Also, it can be considered as a reliable way for NDV virulence evaluation when compared to traditional ways of evaluation (Ganar et al., 2014).

Results of F protein gene sequencing revealed that the cleavage site motif of S4 isolate has the sequence of S4 isolate has the sequence of the phenylalanine (F) residue at position 117 (Collins et al., 1993). The full sequence of both F and HN protein genes were submitted to the GenBank database with accession number MN905162 for the full F protein gene sequence and MN905163 for the full HN protein gene sequence.

F and HN proteins genes genetic and phylogenetic analysis in the present study revealed high similarity of S4 isolate with Chinese isolates and relatively fewer similarities with the Egyptian isolates which may strongly refer to the role of migratory wild birds in NDV evolution in Egypt.

CONCLUSION

Newcastle disease still occurs in sporadic cases despite massive vaccination programs implemented in the Egyptian poultry field. Migratory wild birds are supposed to have a big role in the continuous evolution of NDV in Egypt. Further epidemiological and surveillance work is strongly recommended to define the exact role of migratory wild birds in NDV evolution in Egypt with defining the main causes of the inability of currently used vaccines to protect chickens against infection with Newcastle disease virus.

DECLARATION

Authors’ contributions

All authors reviewed the final manuscript. This work is a part of Mira Maher, and Abdulrahman S. Metwally thesis under the supervision of Shakal M, and Gehan Safwat. Shakal M. designed, supervised the experiments, and co-wrote the paper. Gehan Safwat co-designed the experiment and co-wrote the paper. Mohammed A. Abdel Sabour conducted samples pooling, virus isolation, and co-wrote the paper. Mira Maher and Abdulrahman S. Metwally conducted RNA extraction, genes amplification by PCR, and conducted genetic alignment. Yahia M. Madbouly conducted RNA extraction, real-time reverse transcription PCR, GenBank submission, phylogenetic analysis, and co-wrote the paper.

Competing interests

The authors declare that they have no competing interests.

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