Molecular Detection, Virus Isolation and Pathotyping of a Newcastle Disease Virus Field Strain from Backyard Chickens in Qatar

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Abstract: A first trial of detection, isolation and pathotyping of a Newcastle Disease Virus (NDV) field strain was conducted in Qatar. NDV RNA was isolated, amplified and detected from oropharyngeal and cloacal swabs from clinically diseased backyard chickens and Amnioallantoic fluids (AAF) of infected chicken embryos using primary RT-PCR, nested RT-PCR and gel electrophoresing techniques. The 356 and 216 bp RNA fragments were retrieved in primary and nested RT-PCR, respectively. NDV was isolated, identified and pathotyped in commercial chicken embryos using Haemagglutination (HA), Haemagglutination Inhibition (HI) tests, Mean Death Time (MDT) and Intracerebral Pathogenicity Index (ICPI). The causative agent was found to be a highly virulent strain.

Key words: Newcastle diseases, newcastle disease virus, chicken, RT-PCR, nested RT-PCR, virus isolation, pathotyping

INTRODUCTION

Newcastle Disease (ND) is a highly contagious viral malady of avian species known to cause significant worldwide afflictions to poultry industry. The virus belongs to Avian Paramyxovirus type 1 (APMV-1) serotype, genus Avulavirus that relates to the family Paramyxoviridae (Mayo, 2002). Up to date, over 250 species were estimated to be susceptible (Alexander, 2008). Although, ND was diagnosed earlier in Saudi Arabia neighboring Qatar (El-Zein, 1986) and frequent virulent isolates had been reported in the middle east (Alexander et al., 1987), only clinical reports of presence of the disease in Qatar were known. Information about the history, incidence and prevalence of the disease is limited. Because of lack of laboratory facilities at that time, no detection and isolation trials were conducted to substantiate the presumptive diagnosis describing the disease. The newly establishment of such means has paved the way to carry definitive laboratory diagnosis.

In order to investigate the presence of Newcastle Disease Virus (NDV) in samples from backyard chickens during ND-suspected outbreak in January 2008, Reverse Transcription Polymerase Chain Reaction (RT-PCR), nested RT-PCR, virus isolation and pathotyping techniques were used.

MATERIALS AND METHODS

Samples: Oropharyngeal and cloacal swabs, spleens, kidneys, tracheae and livers were collected from apparently diseased chickens. Swabs were preserved individually into 1 mL virus transport medium and were refrigerated for future processing. Necropsies were stored frozen at -40°C till used. Amnioallantoic Fluids (AAF) from dead organ tissue-inoculated chicken embryos were either used freshly or as frozen stocks.

Viral RNA extraction: Separate pools of oropharyngeal and cloacal swab fluids and AAF from dead organ tissue inoculated chicken embryos were used for isolation of viral RNA using AM 1929 MagMax™ AL/ND Viral RNA Isolation Kit (Ambion ABI, USA) following the manufacture’s instructions. LaSota B, type, clone N79 and HN79 mass (Schering-Plough Animal Health, Millsboro, Delaware, USA) vaccine strains used for vaccination in the country were included as positive controls. Briefly, 400 µL from each pool was added to 802 µL viral lysis/binding solution, vortexed gently for 30 sec followed by brief centrifugation. The lysate was mixed with 20 µL RNA magnetic bead resuspension, vortexed gently for 4 min and centrifuged briefly for 2 sec. Then, the mixture was captured to a magnetic stand for 3 min. The supernate was removed and the trapped viral RNA was washed

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3 times with two washing solutions. The beads were dried for 2 min before elution of the isolated RNAs in 100 µL elution buffer. All RNAs were collected after magnetically trapping the beads and were stored at -20°C for amplification.

Reverse Transcription Polymerase Chain Reaction (RT-PCR): The forward (5'-GCA GCT GCA GGG ATT GTG GT-3', nucleotide position 158-177) and reverse (5'-TCT TTGAGCAGGAGATGTTG-3', nucleotide position 513-493) oligonucleotide primer sets were used for amplification of 356 bp amplicons corresponding to the cleavage activation site of NDV-F gene (Nanthalakumar et al., 2000). Reverse transcription and PCR amplification followed QiaAmp one step RT-PCR procedure (Qiagen). A 50 µL total reaction mix was prepared using 10 µL 5X QiaIgen RT-PCR buffer, 2 µL dNTPs, 10 µL 5X Q-solution, 6 µL 5 mM F and R primers each, 2 µL enzyme mix, 0.5 µL 20 µM L-1 RNase inhibitor, 9 µL MQ H2O and 100 µg viral RNA. A three-step amplification method of 35-cycle-programmed PCR machine (9800 Fast Thermal Cycler) was used for cDNA amplification as follows: 50°C for 30 min (reverse transcription), 95°C for 15 min (initial PCR activation), 3-step cycling 94°C for 45 sec (denaturation), 58°C for 45 sec (annealing), 72°C for 45 sec (extension) and 72°C for 5 min (final extension).

Nested RT-PCR: Nested RT-PCR was used for confirmation. About 1 µL of each 10% diluted primary amplicon was amplified with the F-primer (5'-CCC CTG TGG AGG CAT AC-3', nucleotide position 282-298) and R-primer (5'-TGT TGG CAG CAT TTT GAT TG-3', nucleotide position 497-478) targeting the 216 bp internal sequence of the cleavage activation site of the NDV-F gene (Nanthalakumar et al., 2000). GeneAmp Gold PCR Reagent kit was used for amplification in a total volume of 50 µL reaction mix containing 5 µL 10X PCR buffer, 15 pmol each of the F and R primers, 200 µM each dNTP, 2.5 mM MgCl2, and 3 units AmpliTaq Gold DNA polymerase. MQ H2O was included as negative control in both primary and nested RT-PCR techniques. Amplification phases were as that in RT-PCR and cycles were adjusted following 95°C for 5 min (initial activation), 3-step cycling 94°C for 45 sec (denaturation), 56°C for 45 sec (annealing), 72°C for 45 sec (extension) for 35 cycles followed by 72°C for 5 min (final extension). All amplicons were determined by electrophoresis, visualization and documentation following standard procedures.

Virus isolation: Tissue homogenates were used for virus isolation. The 10% pools were prepared in sterile Hank’s-Balanced Salt Solution (SHBSS) (Euro C-ione) followed by centrifugation for 10 min at 1000 rpm at 20°C and filtration through 0.45 µm millipore membranes. Each of the final inocula was supplemented with 50 µL of 200 IU mL-1 cell culture grade penicillin, 200 µg mL-1 streptomycin and 20 mg mL-1 gentamycin. About 200 µL of each homogenate was inoculated into the allantoic sac of five 10 days old commercial chicken embryos (Loghman breed). Controls received the same dose of SHBSS. All eggs were incubated at 37°C for 8 days observation. About 100 µL of AAF obtained from the first passage was inoculated successively for another two passages using the same inoculation conditions.

Haemagglutination (HA) and Haemagglutination Inhibition (HI) test: HA and HI were used for virus identification follows standard procedure (OIE, 2008). The 10%, 4 weeks old washed chicken RBC was used for spot HA testing individual AAF from the dead embryos prior to harvesting. Four HAU were used in HI test using reference anti-NDV serum (PA0155, VLA, UK).

Determination of the 50 Embryo Lethal Dose (ELD50) and Mean Death Time (MDT): A single-step determination of both ELD50 and MDT was conducted. About 100 µL of 10^-1 to 10^-4 AAF dilutions prepared from the second passage was inoculated each into the allantoic sac of five 10 days old Loghman breed commercial chicken embryos while controls were inoculated with SHBSS (Euro C-ione). All eggs were incubated at 37°C and observed for 8 days at 12 h intervals. ELD50 was calculated according to Spearman-Karber method (Villegas and Purchase, 1980).

Determination of the Intracerebral Pathogenicity Index (ICPI): The 50 µL of 10% (10^3 ELD50 mL^-1) third passage AAF was inoculated intracerebrally to each of the ten 30 h old Loghman breed commercial chicks. Twenty chicks were used as controls, 10 were inoculated with the same dose and technique using antibiotic-free sterile phosphate-buffered saline and the other 10 were kept untreated. Normal, sick and dead birds were recorded daily at the same time of inoculation for successive 8 days. The test followed standard procedures (OIE, 2008). ICPI was determined according to Allan et al. (1987).

RESULTS AND DISCUSSION

Epidemiological and clinical observations: The disease was of a rapid pattern (1-3 days). All the infected chickens
were non-vaccinated. Mortality rate exceeded 60%. Nervous signs were predominant. Necropsized birds showed ecchymotic haemorrhages in the mucosae of the proventriculus and intestines.

Molecular detection of viral RNA in oropharyngeal and cloacal swabs and Aminioallatoic Fluids (AAF): Newcastle disease virus was detected in the swabs, AAF of infected chicken embryos and the vaccine strains. cDNA coinciding with the positive controls measuring 356 and 216 bp were detected in the primary RT-PCR (Fig. 1) and nested RT-PCR (Fig. 2), respectively. Of the positive samples, cDNA from AAF had highly distinct 356 bp band followed by the oropharyngeal swabs. No band was detected from the cloacal swabs. Positive bands of 216 bp were noticed in all tested samples using nested RT-PCR. No bands were obtained from the negative controls in both techniques.

Recovery of NDV from tissues: NDV virulent field strain was successfully recovered into 10 days old commercial chicken embryos inoculated with the NDV-suspected tissue homogenates. Three successive viral passages were attained. The mean HA and HI titres are shown in Table 1.

ELD50, MDT and ICPI of the isolate: The ELD50 and MDT of the isolate are shown in Table 1. All the AAF of the dead chicken embryos showed positive HA on spot testing indicative of viral infection. The MDT value was found to be equivalent to the 10^7 dilution (10^6 ELD50 mL^-1). ICPI of the isolate was shown in Table 1.

The least definitive diagnosis of ND necessitates virus isolation, identification and pathotyping of the strain. In this investigation, a high virulent NDV field strain was detected, isolated and pathotyped based on clinical epidemiology of the disease, RT-PCR, nested RT-PCR, virus isolation, MDT and ICPI results. The successful use of RT-PCR to rapidly diagnose ND indicates the efficacy of the technique as a primary diagnostic tool especially where standard virus isolation facilities are not available. The application of the nested RT-PCR for confirmation showing almost similar band intensity in all samples has further reflected the sensitivity and identity of the primary RT-PCR results (Jestin et al., 1993; Kho et al., 2000; Yousof et al., 2005). In addition, it proved in this study the suitability of the kit and the single step amplification technique followed. The use of the highly conserved F genes for detection of the viral RNA amplicons could also further help future studies aiming at probing the biology, ecology and molecular epidemiology of the isolate.

![Fig. 1: Agarose gel electrophoresis of RT PCR products of NDV field isolate and reference vaccine strains using 356 primer sets. Lane M: pGem DNA molecular size marker. Lane V1: LaSota c1N79 vaccine strain. Lane V2: LaSota c1HN79 mass vaccine strain. Lane AAF: Aminioallatoic Fluids. Lane O: Oropharyngeal swabs. Lane C: Cloacal swabs. Lane N: Negative control](image1)

![Fig. 2: Agarose gel electrophoresis of nested RT PCR products of NDV field isolate and reference vaccine strains using 216 bp primer sets. Lane M: pGem DNA molecular size marker. Lane V1: LaSota c1N79 vaccine strain. Lane V2: LaSota c1HN79 mass vaccine strain. Lane AAF: Aminioallatoic Fluids. Lane O: Oropharyngeal swabs. Lane C: Cloacal swabs. Lane N: Negative control](image2)

Table 1: Some properties of the isolated NDV field strain

<table>
<thead>
<tr>
<th>HA (log2)</th>
<th>HI (log2)</th>
<th>ELD50 (log10)</th>
<th>MDT (h)</th>
<th>ICPI</th>
</tr>
</thead>
<tbody>
<tr>
<td>9</td>
<td>7</td>
<td>8.2</td>
<td>60</td>
<td>1.59</td>
</tr>
</tbody>
</table>

Although, virus isolation is a gold standard measure for diagnosis yet time consuming and laborious, the investigation successfully used commercial eggs for isolation. Owing to less sensitivity of this tool compared to specific pathogen free eggs (Wakamatsu et al., 2006), and because of diversity in tissue affinity among variant NDV virulent strains (Brown et al., 1999; Hofstad, 1951; Alexander, 2008), pooling of samples from different organ tissues was followed. This process was recommended to attain productive results (Smietanka et al., 2006).
Many people in Qatar have backyard chickens stocked with variety of birds including turkeys, pigeons, geese, ducks and peafowl. Although, the study did not include the sources of infection, it suggests incrimination of imported birds, an appreciable source of birds in the country. Live market birds and to a lesser extent, the wild species could be involved in spreading the disease. This assumption is based on absence of active surveillance programmes and lack of consistent vaccination schedules against ND in most rearing communities. Only casual immunization at the will of the owner is practiced.

**CONCLUSION**

In this study, it is concluded that NDV is a wide host range virus and it could aggravates the sequence of other infective pathogens (Bano et al., 2003; Monne et al., 2006; Aamir et al., 2007), isolation of a high virulent strain from non-vaccinated flocks with such rearing system warrants attention to the growing poultry industry in the country.

**RECOMMENDATIONS**

Efficient public awareness and empowerment of bird trade legislations are needed. Further, epidemiological and molecular studies to give insights on the candidate vaccines and the recommended vaccination policy against the disease are needed.

**ACKNOWLEDGEMENTS**

This investigation was conducted at the premises of the Department of Animal Resources, state of Qatar. The researchers are grateful to the director of the Department of Animal Resources and the Head of Veterinary Research Laboratories for the help, encouragement and permission to use those facilities. Due thanks also to the technical staff for the assistance they afforded. Thanks also go to the Arabian-Qatari company for poultry production for provision of the fertile eggs and chicks.

**REFERENCES**


