Comparative Study on Some Characteristics of Newcastle Disease Virus Field Strains Isolated From Captivated Avian Species in Qatar

Abstract
Six Newcastle disease virus (NDV) field strains (NDV-QF10, NDV-QF13, NDV-QH10-1, NDV-QH10-2, NDV-QQ09 and NDV-QP09) were isolated and pathotyped from samples collected from captivated falcons, houbara bustard, quaïl and pigeon, respectively, during the study period of 2012 and 2013 in Qatar. Virus cultivation in specific pathogen free embryonating chicken eggs (SPF ECE), haemagglutination (HA), haemagglutination inhibition (HI) and mean death time (MDT) assays used in the investigation displayed virus identity and pathogenicity. The six NDV isolates were classified velogenic strains. Sequence analysis of 150bp of the amplified 216bp nucleotides of the functional region of the F-gene encoding the F2-F1 cleavage site, revealed one amino acid motif confirming velogenicity, and that the three strains clustered under NDV genotype VII either subgroup d or f.

Keywords
Newcastle disease virus; Captivated avian species; Virus isolation; Pathotyping; Phylogenetic analysis; Genotyping

Abbreviations
ND: Newcastle Disease; NDV: Newcastle Disease Virus; SPF-ECE: Specific Pathogen-Free Embryonating Chicken Eggs; AAF: Amnio Allantoic Fluid; HA: Haemagglutination; HAU: Haemagglutinating Units; HI: Haemagglutination Inhibition; MDT: Mean Death Time; ICPI: Intracerebral Pathogenicity Index; SPF-ECE: Specific Pathogen-Free Embryonating Chicken Eggs; HA: Haemagglutination; HAU: Haemagglutinating Units; HI: Haemagglutination Inhibition; MDT: Mean Death Time; ICPI: Intracerebral Pathogenicity Index; SPF-ECE: Specific Pathogen-Free Embryonating Chicken Eggs; SPF: Specific Pathogen-Free

Introduction
Newcastle disease (ND) is a known highly contagious disease of a significant worldwide impact on avian species. The virus belongs to avian paramyxovirus type 1 (APMV-1) serotype, genus Avulavirus that relates to the family Paramyxoviridae and Mononegavirales order [1,2]. Hundreds species were estimated to be susceptible [3] rendering serious losses among poultry rearing and industry economy. Beside, the virus possess zoonotic characteristics that causing mild conjunctivitis to mankind. Since the first informative detection, isolation and conventional pathotyping of Newcastle disease virus (NDV) in Qatar [4], continuous reporting of the disease was reported. Focal annual outbreaks of 120, primarily among chickens, pigeons and other domesticated birds were observed [5]. This situation signals public health hazards to tendency to rear feral birds, the estimated 3.5 million traditional poultry system that represents 23% of the total bird population, and the 11 million bird of the growing industrial poultry business that valued to 77% of the domesticated birds were observed [5]. This situation signals public health hazards to tendency to rear feral birds, the estimated 3.5 million traditional poultry system that represents 23% of the total bird population, and the 11 million bird of the growing industrial poultry business that valued to 77% of the state poultry population [6].

Aiming to have a preliminary insight on the nature of Newcastle disease virus (NDV) strains circulating in captivated avid species reared in Qatar, conventional virological methods coupled with late molecular assays were used to reveal some properties of the strains including virulence, genotype and phylogenetic relatedness to other known worldwide NDV strains.

Materials and Methods

Samples
Tracheal and cloacal swabs, spleen and liver samples collected from 51 apparently healthy and ND-suspected captivated falcons (Falco peregrines peregrinus), diseased Houbara (Chlamydotis undulata), quaïl (Colinus virginanus) and pigeons (Columbia livia) were used for the study. Swabs were immediately transported in cool chamber condition and frozen at -40°C till used. Positive amnio allantoic fluids (AAF) from specific pathogen free (SPF) embryonating chicken eggs (ECE) inoculated with the collected samples were either used freshly or as frozen stocks.

Virus isolation
Tracheal and cloacal Swabs’ elution and tissue homogenates were used for virus isolation. Ten per cent of each sample was prepared in sterile phosphate-buffered saline (SPBS), followed by centrifugation for 10 min at 2500 rpm at 20°C. Each of the final inoculum was supplemented with 100 μL 200 IU mL⁻¹ cell culture grade penicillin, 100 mg mL⁻¹ streptomycin and 20 mg mL⁻¹ gentamycin suspensions. Two hundred μL of each swab elution and tissue homogenate was inoculated into the allantoic sac of five 10-day-old SPF ECE, Loghman breed (VALO, Germany). Controls received the same dose of SPBS. All eggs were incubated at 37°C for 8 days observation. Two hundred microlitre of each neat AAF obtained from the first passage was inoculated for a successive second passage using the same inoculation conditions.
Haemagglutination (HA) and Haemagglutination Inhibition (HI) test

AAF harvest of passage level two was used for virus identification applying HA and HI assay following standard procedures [7]. Ten percent 4-week-old washed chicken RBC was used for rapid spot HA testing individual AAF of each dead embryo prior to harvesting and pooling. Four HAU determined by HA assay of the positive AAF pools of each isolate was used in HI to determine the virus strains specificity to the reference anti-NDV serum (PA0155, VLA, UK).

Determination of the 50 Embryo Lethal Dose (ELD50), Minimum Embryo Lethal Dose (MELD) and Mean Death Time (MDT)

MDT determination follows [8] standard procedure with minor modifications. A single-step determination of ELD50, MELD and MDT was conducted. Two hundred microlitre of 10^5-10^6 AAF dilutions prepared from the second passage was inoculated each into the allantoic sac of five 10-day-old SPF ECE (Loghman breed; VALO, Germany). All eggs including uninfected controls were incubated at 37°C and observed for 8 days at 12 h intervals. ELD50 was calculated according to [9].

Viral RNA extraction

Pools of positive haemagglutination (HA)/haemagglutination inhibition (HI) NDV AAF were used for viral RNA extraction using QIAamp (QIAamp Viral RNA Mini Handbook, 12/2005). Briefly, 140µL of each pool was added to 560µL buffer AVL RNA carrier, pulse-vortexed for 15 sec and incubated at room temperature (15-25°C) for 10 min. 560µL 96-100% ethanol was used for precipitation purposes. QIAamp Mini spin column contained in a 2 mL collection tube was used for loading of the lysed material. 500µL Buffer AW1 and AW2 were used successively to maintain RNA binding and washing. Finally, 80 µL AVE Buffer was used to elude the RNA extract after 1 min room temperature incubation and 1 min 6000 x g (8000rpm) centrifugation. All viral RNA extracts were stored at -20°C till amplification.

Reverse transcription polymerase chain reaction (RT-PCR)

The forward (5′- GCA GCTGCA GGG ATT GTG GTG-3′, nucleotide position 158-177) and reverse (5′- TCT TTG ACCAGG AGG ATG TTG-3′, nucleotide position 513-493) oligo nucleotide primer sets were used for amplification of 356bp amplicons corresponding the cleavage activation site of NDV-F gene [10,11]. RT-PCR amplification followed Access Quick RT-PCR Kit (Cat. no. A1700). 25.5µL total reaction mix was prepared using 12.5µL 2X Access Quick Master Mix, 2.5µL 1µM upstream F-primer and control R-primer each, 2.5µL Nuclease-free water, 0.5 µL 5 U/µ AMV RT enzyme. A three-step amplification method of 35-cycle programmed PCR machine (9800 Fast Thermal Cycler) was used for cDNA amplification as follows: 45°C for 45 min and 95°C for 2 min (RT-step), 3-step cycled of 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).

Sequence analysis of the complementary deoxynucleotide acid (cDNA) fragments

cDNA RT-PCR products were measured for concentration and run on 1.5% agarose gel according to standard procedures. Fragments purification was done according to ExoSAP-IT instructions. Amplification was performed with the 216bp NDV F-gene nested primer sets (F-primer: 5′-CCC CGT TGGGAG CAT AC-3′, nucleotide positions 282-298; R-primer: 5′-TGT TGG CAG CAT TTG GATGG-3′, nucleotide positions 497-478) according to ExoSAP-IT instructions (PCR: 78200/01/02/05/50-Affymetrix, USA) using 16.2µL ddH2O, 0.8µL each F and R primers, 1µL purified cDNA product. The deduced nucleotide sequences were generated, constructed and aligned using MEGA (Version 5).

Results and Discussion

The epidemiological and clinical findings of this investigation demonstrated that all of the birds’ populations involved in these focal outbreaks were susceptible to NDV and not vaccinated. ND-infected birds demonstrated a rapid disease course pattern of about 1-3 days. All birds showed nervous signs. Beside, the houbara and quails showed diarrhea. The quail necropsies exhibited ecythymosis in the mucosa of the proventriculus and intestines. Based on successful recovery of the strains on inoculation of the suspected NDV swabs’ elution samples and tissue homogenates into AAF of the 10-day-old SPF ECE, exhibition of the biological HA characteristics, HI of the isolates on testing against the reference anti-NDV serum with a minimum HI values of 4log2 (Table 1), retrieval of the specific 356bp NDV F-gene products at the cleavage site of the NDV F protein using RT-PCR; six NDV field strains were isolated.

The fact that the isolated strains are originating from samples collected from four different avian type species (falcon, houbara, quail and pigeon) signifies the circulation of ND among Qatari birds. Further, it also indicates the endemic situation of the disease in the country. However, not necessary genetically related, the dose range of the ELD50 of the isolated strains (10^10 to 10^12.3 ml^-1) and the similarity of the MDT (48h) indicate the close pathogenic properties among the isolated NDV strains. MDT records classify the strains pathologically velogenic. Further, the sequence analysis of the corresponding 216bp nested NDV F-gene segments of NDV-QF13, NDV-QH10-2 and NDV-QQ09 strains Figure 1 showed "R-R-K-R-F" amino acid sequence motif indicative of velogenicity of these isolates. These findings simulate that of the first observations of [4] when a velogenic NDV strain (NDV-QC08) was isolated from ND-suspected backyard chickens using ICPI. Although the investigation did not include viral tropism, the necropsy findings of 2 of the houbara and the quail were typical viscerotropic [12,13].

Despite all strains were successfully propagated at passage level one and two; only at dilution 10^6 could the study retain the biological HA ability of the NDV-QH10-1 isolate with the highest scored ELD50 value of 10^12.3. This finding might further reinforce the virulence potentiality of this isolate.

Even so, the study did not apply intracerebral pathogenic
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In the present investigation, it was clear that none of NDV-QF13, NDV-QH10-2 or NDV-QQ09 has 100% nucleotide homology with the GenBank-derived NDV vaccines; Hitchner B1-47, Lasota, VGGA-87 and Clone30 and that they were clustered under NDV genotype VII group whereas the vaccines are under genotype II (Figure 3). These facts reflect the basic finding that the isolated

Table 1: Some properties of the isolated NDV field strains tested at egg passage level 2 and compared to NDV-QC08 (the first NDV isolate derived from chicken in Qatar [4]).

| NDV Strain | Species | HA (Log 2) | HI (> 4Log 2) | ELD50 (Log 10) (1mL) | MELD (Log 10) (1mL) | MDT (Hour) | ICPI |
|------------|---------|------------|--------------|---------------------|--------------------|------------|------|
| NDV-QF10   | Falcon  | Positive   | 9.8          | 5                   | 48                 | ND         |
| NDV-QF13   | Falcon  | Positive   | 9.7          | 6                   | 48                 | ND         |
| NDV-QH10-1 | Houbara | Positive   | 12.3         | 11                  | 48                 | ND         |
| NDV-QH10-2 | Houbara | Positive   | 9.8          | 8                   | 48                 | ND         |
| NDV-QP09   | Quail   | Positive   | 10.2         | 7                   | 48                 | ND         |
| NDV-QQ09   | Chicken | Positive   | 9.8          | 6                   | 48                 | ND         |
| NDV-QC08   | Chicken | Positive   | 8.2          | 7                   | 60                 | 1.58       |

NDV-QF10: Newcastle Disease Virus Qatar Falcon10; NDV-QF13: Newcastle Disease Virus Qatar Falcon13; NDV-QH10-1: Newcastle Disease Virus Qatar Houbara-1; NDV-QH10-2: Newcastle Disease Virus Qatar Houbara-2; NDV-QQ09: Newcastle Disease Virus Qatar Quail09; NDV-QP09: Newcastle Disease Virus Qatar Pigeon09; NDV-QQ09: Newcastle Disease Virus Qatar Chicken09; HA: Haemagglutination; HI: Haemagglutination Inhibition; ELD50: 50% Embryo Lethal Dose; MELD: Minimum Embryo Lethal Dose; MDT: Mean Death Time; ICPI: Intracerebral Pathogenicity Index; ND: Not Done

Figure 1: Sequence analysis of the amplified F gene of NDV-QF13, NDV-QH10-2 and NDV-QQ09 showing the gene cleavage site and its deduced amino acid sequences compared to NDV-QC08 and other Genbank worldwide strains (MEGA, Tamura-Nei model 1993).
field strains are of unique virulence to that of these known vaccines. The falcon (NDV-QF13) and the quail (NDV-QQ09) strains showed 100% homology in their nucleotide sequence (Figure 1 and 2), and that they were within the subgroup VII-f (Figure 3). However, both were different in nucleotide sequence correlated to NDV-QC08 [17], the first known NDV field isolated in Qatar [4]. Although the nucleotide sequence and homology of the houbara isolate (NDV-QH10-2) was different from that of NDV-QC08 (Figure 1 and 2), both were within the subgroup VII-d (Figure 3), which was known to be responsible for most of the global ND outbreaks since the 20th century. Regional wise, NDV-QF13, the Qatari falcon strain, was found to be genetically different from the United Arab Emirates NDV field isolates (falcon strain AV 1196/96 18, genotype IVa, Genbank accession No. AY175738, peregrine falcon strain AV 1573/92 804/90, genotype IV-c, Genbank accession no. AY175677, domesticated fowl strain AV 51/9640-1084-95, genotype IV-c, Genbank accession no. AY175663 and ostrich strain AV1009/98 169.1668-98, genotype IVc, Genbank accession no. AY175666). Conclusively, all these findings might reflect the variation in the epidemiology of the field NDV strains circulating in Qatar than that of the regional reference Genbank-derived strains used in the study. However, further molecular investigations with a wide-based alignment to known GenBank-derived NDV strains are recommended to have more perception about the molecular epidemiology of the Qatari NDV field strains (Table 2).

Generally, it is acceptable that sequence of as little as 250bp always give meaningful convenient phylogenetic analysis compared to much longer sequences [18,19]. Nonetheless, in this study, only 150bp of the totally amplified cDNA of the F-gene was successfully used to reveal results for molecular pathotyping and phylogenetic analysis of the 216bp-amplified cDNA of the isolated NDV strains. The reverse is true for a convenient and precise genetic characterization studies [18,19]. Previous investigations

Figure 2: Homology percentage of the nucleotide sequence analysis (MEGA) of NDV-QH10-2, NDV-QF13 and NDV-QQ09 compared to NDV-QC08 and other NDV sequences retrieved from the Genbank (Modern Evolutionary Genetic Analysis “MEGA”), Tamura-Nei model 1993.)

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have established that, the fusion protein is likely one of the major proteins that play an important role in determination of the genetic variations among different NDV strains, and that, it is more sensitive than the internal genes demonstrating genetic characteristics [20,21]. For that, further future molecularly established investigations using F-gene primer sets of a convenient length is recommended to cast light on the biology, ecology, and epidemiology of the isolated NDV strains and the newly expected emerging strains as well.

**Conclusion**

The fact that falcons play a significant role in the traditions of Gulf citizens is very observable in Qatar State. Since, houbara, quails and pigeons are the common birds used as a main raw meat source for feeding falcons, and that they are highly ND-susceptible and reservoir birds, reared in mixed communal backyards and traditional farms, perpetuation of NDV in the country is anticipated. In addition, increased possibilities of emergence of new virulent virus mutants are also expected. Accordingly, much attention and more strict health measures are recommended to guard the growing investment in the country's poultry industry. The cross-boundary nature of falcons as a hunting bird will also warrants additional health measures. Control measures based on identity of the nature of the candidate vaccine to combat the incriminated virus in the region should be escorted carefully.

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Table 2: Genbank accession data of the reference NDV strains aligned to the NDV field strains isolated from the four avid species (NCBI Blast-BLASTN 2.2.29+).

| Isolate | Host | Protein/Gene | Accession No. |
|---------|------|--------------|---------------|
| NDV isolate IS/375/01 | - | Fusion protein (F) gene- partial cds | JF795626.1 |
| NDV isolate IS/374/01 | - | Fusion protein (F) gene- partial cds | JF795622.1 |
| NDV isolate IS/242/01 | - | Fusion protein (F) gene- partial cds | JF795621.1 |
| NDV isolate Chicken/China/Guangxi9/2003 | Chicken | Complete genome | F343539.1 |
| NDV isolate Chicken/China/Guangxi9/2003 | Chicken | Complete genome | DQ485230.1 |
| NDV isolate Duck/H/GD/YP-09 | Duck | Complete cds | KJ750156.1 |
| NDV isolate Chicken/China/Hebei/01/2006 | Chicken | Complete genome | KC542895.1 |
| NDV strain WF00D | - | Complete genome | FJ754272.1 |
| NDV strain IR-HGT2012.2 | - | Fusion protein (F) gene- partial cds | JX131350.1 |
| NDV strain IR-Ostrich-HGT2012 | Ostrich | Fusion (F) gene- partial cds | JX131358.1 |
| NDV isolate NR-29 | - | Fusion (F) gene- partial cds | KC161990 |
| NDV isolate IR-HGT2010.2 | - | Fusion protein (F) gene- partial cds | JX131353.1 |
| NDV isolate NR_91 | - | Fusion (F) gene- partial cds | JX129804.1 |
| NDV isolate IR-HGT2012.2 | - | Fusion protein (F) gene- partial cds | JX131350.1 |
| NDV strain IR-Ostrich-HGT2012 | Ostrich | Fusion protein (F) gene- partial cds | JX131358.1 |
| NDV strain IR-HGT2012.1 | - | Fusion (F) gene- partial cds | JX131357.1 |
| NDV isolate NR_98 | - | Fusion (F) gene- partial cds | JX129809.1 |
| NDV isolate NR_97 | - | Fusion (F) gene- partial cds | JX129808.1 |
| NDV isolate NR_95 | - | Fusion (F) gene- partial cds | JX129807.1 |
| NDV isolate NR_94 | - | Fusion (F) gene- partial cds | JX129806.1 |
| NDV isolate NR_94 | - | Fusion (F) gene- partial cds | JX129805.1 |
| NDV/FALCON/RL/IS/2012/911 | Falcon | Fusion protein (F) gene- partial cds | KF650624.1 |
| NDV/CANARY/Je/IS/2012/875 | Canary | Fusion protein (F) gene- partial cds | KF650623.1 |
| NDV/CHICKEN/Hol/IS/2012/870 | Chicken | Fusion protein (F) gene- partial cds | KF650622.1 |
| NDV/PARROT/BS/IS/2012/84 | Parrot | Fusion protein (F) gene- partial cds | KF650619.1 |
| NDV-Ck-Pakistan-NARC-13N20-2013 | Chicken | Fusion protein (F) gene- partial cds | KC811835.1 |
| NDV-Ck-Pakistan-NARC-13N94-2013 | Chicken | Fusion protein (F) gene- partial cds | KC811831.1 |
| NDV-Ck-Pakistan-NARC-12N152-2012 | Chicken | Fusion protein (F) gene- partial cds | KC811830.1 |
| NDV-Ck-Pakistan-NARC-12N148-2012 | Chicken | Fusion protein (F) gene- partial cds | KC811841.1 |
| NDV-Ck-Pakistan-NARC-12N80-2012 | Chicken | Fusion protein (F) gene- partial cds | KC811813.1 |
| NDV-Ck-Pakistan-NARC-28991-2012 | Chicken | Fusion protein (F) gene- partial cds | KC811821.1 |
| NDV/PARROT/BS/IS/2012/84 | Parrot | Fusion protein (F) gene- partial cds | KF650619.1 |
| NDV-Ck-Pakistan-NARC-13N20-2013 | Chicken | Fusion protein (F) gene- partial cds | KC811835.1 |
| NDV-Ck-Pakistan-NARC-13N94-2013 | Chicken | Fusion protein (F) gene- partial cds | KC811831.1 |
| NDV-Ck-Pakistan-NARC-12N152-2012 | Chicken | Fusion protein (F) gene- partial cds | KC811830.1 |
| NDV-Ck-Pakistan-NARC-12N148-2012 | Chicken | Fusion protein (F) gene- partial cds | KC811841.1 |
| NDV-Ck-Pakistan-NARC-12N80-2012 | Chicken | Fusion protein (F) gene- partial cds | KC811813.1 |
| NDV-Ck-Pakistan-NARC-28991-2012 | Chicken | Fusion protein (F) gene- partial cds | KC811821.1 |
| NDV-Ck-Pakistan-NARC-12N126-2012 | Chicken | Fusion protein (F) gene- partial cds | KC811811.1 |
| NDV-Ck-Pakistan-NARC-28789-2012 | Chicken | Fusion protein (F) gene- partial cds | KC811810.1 |
| NDV-Ck-Pakistan-NARC-28859-2011 | Chicken | Fusion protein (F) gene- partial cds | KC811809.1 |

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