Genetic characterization and phylogeny of pigeon paramyxovirus isolate (PPMV-1) from Pakistan

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Abstract

Background: Knowing the genome characteristics of circulating Newcastle disease viruses [avian paramyxoviruses (APMV-1) and pigeon paramyxoviruses (PPMV-1)] is important to devise appropriate diagnostics and control strategies. APMVs originating from chicken and wildlife in Pakistan are well-elucidated; nevertheless, molecular characterization for the circulating PPMV-1 is largely unknown.

Findings: Here, we have performed fusion (F) and hemagglutinin (HN) gene based characterization of PPMV-1 isolated from an outbreak in a pigeon flock. With F0 proteolytic cleavage site (112RRQKR↓F117), characteristic of velogenic/mesogenic serotype, the complete F and HN gene based sequence analysis of the isolate revealed evolutionary relationship to genotype VI. Further analysis of hyper-variable region of F-gene demonstrated clustering of the study isolate with genotype VIb. The deduced residue analysis for both F and HN protein showed a number of substitution mutations in the functional domains distinct from representative strains of each genotype including the vaccine strains; some of them were found exclusive to the study isolate.

Conclusions: Though limited and preliminary data, the findings enhance our knowledge towards circulating strains of PPMVs in Pakistan. Further studies are needed to ascertain its potential for transmission in the wild birds, commercial and backyard poultry and its subsequent shedding into the environment.

Keywords: Pigeon paramyxovirus type 1 (PPMV-1), Newcastle disease virus, Pigeon, Pakistan

Background

Pigeon Paramyxovirus type 1 (PPMV-1) is an antigenic and host variant of classical Newcastle Disease virus (NDV) or Avian Paramyxovirus type 1 (APMV-1) that causes Newcastle Disease (ND)-like infection and pathology in pigeons (Ujvari et al. 2003). Including the earliest known PPMV-1 (Iraq78) originating from the Middle-East (ME), and the strains responsible for third panzootic in 80 s involving a pigeon-adopted APMV-1 are considered likely to be derived from these ME viruses (Kaleta et al. 1985; Ujvari et al. 2003) and as yet continue to circulate around the globe. The virus has an approximate genomic length of 15192 nt and, despite characteristic F protein cleavage site for velogenic strains (112GRQKRF117 or 112RRKKRF117 or 112RRQKRF117), differences in pathogenicity index range from moderate to no virulence for chicken (Collins et al. 1994; Dortmans et al. 2010). However in recent years, some PPMV-1 have been reported to be highly pathogenic for chicken after passage either in chicken or chicken embryo indicating their potential to cause ND outbreaks (Dortmans et al. 2011). Today, ND caused by either virulent APMV-1 or PPMV-1 is considered endemic to feral and domestic pigeon (Columbiformes) worldwide (Aldous et al. 2004). Clinical symptoms differ depending upon host immune titre and virulence of isolate involved (Dortmans et al. 2011). Most of the times, the observed clinical symptoms relate

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to neurotropic form of ND including tremors, torticollis and disturbed equilibrium, however naturally infected pigeon also exhibit respiratory symptoms such as sneezing, coughing and tracheal rales (Marlier and Vindevogel 2006).

Since the establishment of well-organized poultry sector in Pakistan to-date, there have been a number of epidemics of ND in commercial poultry, wild birds and domestic pigeons. Although, APMV-1 from commercial poultry and wild birds have been isolated and well-characterized as genotype VI and VII (Munir et al. 2010; Shabbir et al. 2012, 2013a, b), despite a number of clinically suspected outbreaks in pigeons including vaccinated ones (Lab data, not published), there is absolute paucity of information pertaining to circulating lineage of PPMV-1 and its nucleotide and amino acid profile to reference and vaccinal strains. We analysed complete F and HN genes of an isolate recovered from ND outbreak in a pigeon flock. The obtained sequences were processed for phylogeny and amino acid residue analyses giving an insight towards genetic diversity of the indigenous strain and its comparative evolution to those reported earlier.

Methods

A Newcastle disease outbreak occurred in a flock of racing pigeons (n = 56) in district Lahore, Pakistan (31.5790°N, 74.3096°E) during December, 2014. Four days post appearance of clinical symptoms, 13 died and 29 were morbid. The clinical symptoms observed in affected pigeons were circling movement and tremors while some birds were also exhibiting mild respiratory symptoms such as sneezing, coughing and nasal discharge. Necropsy was performed and samples (trachea, lungs, and brain) were processed for isolation of NDV in 9 day old embryonated chicken eggs (ECE). Presence of agglutinating virus was confirmed in harvested allantoic fluid by spot agglutination assay using 10 % chicken red blood cells (RBCs). Later, identity of the isolate was confirmed as NDV through standard haemagglutination inhibition assay using specific antisera.

Total RNA was extracted from allantoic fluid using commercially available RNA extraction kit as per manufacturer’s instructions (QIAamp Viral RNA Mini Kit, Qiagen, USA). Quantity (NanoDrop, USA) and quality (QubitFlourometer, USA) of extracted RNA was measured. The extracted RNA was subjected to amplification of complete F and HN genes spanning the genomic region from 4498 to 6330 nucleotide through reverse transcription polymerase chain reaction (RT-PCR) using the primers and protocols as described previously (Munir et al. 2010). The amplified products were purified by 1.0 % gel electrophoresis, using the Wizard® SV Gel and PCR Clean-Up System (Promega, Co., Madison, WI, USA) as per manufacturer’s instructions. Using the same primers (as used previously for F and HN gene amplification) and ABI PRISM BigDye Terminator version 3.1 (Applied Biosystems, Foster City, CA, USA), the purified genomic DNAs were processed for sequencing reaction on a 3100 DNA Analyzer (Applied Biosystems, Foster city, CA, USA). Each genomic fragment was sequenced twice in both forward and reverse directions to generate a reliable consensus sequence. The consensus sequences of F and HN genes of the study isolate (Pi/MZS1-UVAS/2014) has been submitted to GenBank under the accession number KU644586 and KU644587, respectively.

The obtained sequences and sequences reported earlier (GenBank) were aligned in BioEdit version 5.0.6 (Hall, 1999) using ClustalW and cut to equal lengths. Phylogenetic relationships of complete F, HN gene and hyper-variable region of F gene of study isolate were elucidated to the corresponding region of previously characterized viruses around the globe (http://www.ncbi.nlm.nih.gov/nuccore/?term=partial+F+gene+of+Newcastle+disease+virus) at the level of genotype and sub-genotype using the MEGA version 6.0 software (Tamura et al. 2013). The evolutionary distances were inferred and expressed based on the number of nucleotide substitutions per site. The codon positions included in the analysis were the 1st, 2nd, 3rd, and noncoding. All positions containing gaps and missing data were eliminated from the data set (the “complete deletion” option). Furthermore, comparative residue analysis of the representative strains of each known genotype was analysed through BioEdit.

Results and discussion

Agglutination of harvested virus from allantoic fluid with chicken RBCs (1:128) and subsequent inhibition with specific antisera (1:128) confirmed the presence of NDV. Since a particular emphasis is given to F and HN gene-based molecular characterization of NDV strains in the previous studies (Toyoda et al. 1988; Yussof and Tan, 2001; Aldous et al. 2003; Ujvari et al. 2003; Kim et al. 2008; Munir et al. 2012; Shabbir et al. 2012), we also used complete F and HN gene as well as hyper-variable region of F-gene (47–421 nt, 375 bp) to determine phylogenetic relationship of the study isolate to previously known representative strains of APMV-1 and PPMV-1 around the globe at the level of genotype and sub-genotype. Based upon complete F and HN gene phylogeny, the study isolate clustered within genotype VI close to an isolate reported previously from Russia (Pi/Rus/Vladimir/687/05) (Fig. 1a, b). Further analysis of hyper-variable region of F gene revealed clustering of the study isolate to sub-genotype VIb (Fig. 1c). The clustering of study isolate close to the ones from Russia highlights potential ancestor or origin of the virus that may involve.
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**Fig. 1** Phylogenetic consensus tree for the pigeon-originated NDV isolate for fusion gene (a), hemagglutinin gene (b) and hypervariable region of F gene (c). The nucleotide sequences of study isolate for each gene were compared with corresponding genes of representative strains reported previously to public database, the GenBank. The evolutionary history was inferred using the Neighbor-Joining method with 1000 bootstrap value in MEGA version 6.0.
migratory route of birds from North to South. Since the first introduction of genotype VI, various strains such as V1a, V1b, V1c and V1d responsible for pigeon-origin panzootics have been reported from time to time with widespread geographic movement (Kaleta et al. 1985, Ujvari et al. 2003; Aldous et al. 2004; Lee et al. 2004). The most recent example is the identification of an isolate of genetic clade V1a from China (pi/GX/1015/13) that is supposed to have ancestor similar to isolate (W4/2005) from Europe (Wang et al. 2015). It is noteworthy that NDVs belonging to genotype VI was identified from chicken in district Karachi back in 2005 (Khan et al. 2010) and, more recently, genotype V1c was reported from a commercial poultry farm in northern areas of Pakistan (Shabbir et al. 2013a). As the presence of cleavage motif in mesogenic strains does not necessarily correlate with the pathogenicity for chickens (Kommers et al. 2002; Ujvari et al. 2003; Kim et al. 2008), studies ascertaining transmission of pigeon-originated NDVs in vaccinated as well as immunologically naïve flocks and their subsequent shedding are essential in understanding their potential to cause disease in wild, commercial and backyard poultry in Pakistan.

Nucleotide and amino acid residues of the isolate were identified in the complete coding region of the F (4550–6211, 1662 bp, 571 aa) and HN (6418–8133, 1716 bp, 571 aa) genes. Although, differences in amino acid length of HN protein have been reported previously from different strains of NDV (Romer-Oberdorfer et al. 2006), the study isolates had open reading frame that encoded 571 residues, a feature common to most of the virulent NDVs (Triumurugan et al. 2011). While comparing nucleotide and residues of the study isolate to representative strains of different genotypes, the percent similarity for F gene was found to be maximum with genotype VI (92.4 and 94.2) while it was observed to vary from 84 to 90 for genotype representing vaccine strain used in Pakistan and other representative strains, we observed a number of substitution mutations in residue sequences for fusion peptide (117–142 aa) at position 121 (V → I), 124 (S → G) and 132 (A → S), hydrophilic region-a (143–185 aa) at position 179 (V → I), hydrophilic region-b (268–299 aa) at position 272 (N → H), 288 (T → N) and hydrophilic region-c (471–500 aa) at position 482 (E → A), 487 (K → R), 492 (N → D). Mutation at position 506 (V → I) and 516 (I/M → S) of the major transmembrane region (501–521 aa) further indicates lack of conserveness of this particular part of genome (Fig. 2).

On the other hand, for HN gene, analysis of predicted amino acid sequence revealed conserveness of cysteine residues at positions 123, 172, 186, 196, 238, 247, 251, 344, 455, 461, 465, 531 and 542; key residues for receptor (sialic acid) binding at positions R174, E401, Y526; residues for the hydrophobic core of the stalk 4HB at positions Y85, V88, S92, L96, T99, I103, I107, L110 and L114; and stalk residues involved in direct interaction with F protein at positions R83, A89, L90, L94 and L97 (Ke et al. 2010; Yuan et al. 2011). Six N-glycosylation sites at positions 112NNS121, 341NNT434, 431NKT435, 483NHT483, 531NNT543 and 538NKT540 were identified. When compared to most common vaccine strain used in Pakistan (genotype II), beside substitutions at several point in total length of HN gene, we found substitutions in the
Fig. 2 Alignment of deduced amino acid sequence of complete F gene of pigeon isolate. The residue profile of study isolate is compared with strains of NDVs representing different genotypes including the vaccine strains. Structurally and functionally important residues are boxed and highlighted.
Alignment of deduced amino acid sequence of complete HN gene of pigeon isolate.

The residue profile of study isolate is compared with highlighted strains of NDVs representing different genotypes including the vaccine strains. Structurally and functionally important residues are boxed and highlighted.

**Fig. 3**
transmembrane domain (I29V, T33I, V35M, S43V, L45V, Y46H, G49R), antigenic sites (K98N, G494D, I514V) and neutralization epitope (N263R) (Fig. 3). It is interesting to note that some substitutions were found to be exclusive to the study isolate notably at positions V179I, K321R, I448V, K487R, K/N492D, V506I, L537M for F protein and Q/R7K, V8I, T61A, S/G75N, S/R269P, I/T289A for HN protein (Figs. 2, 3). This is important as some geographically conserved mutation have been reported recently; K4I is conserved for strains reported from Japan (Umali et al. 2013) while I52V, K78R, R101K are considered conserved for isolates originating from the Far East Asia (Japan, China and Taiwan). Taken together, variation in nucleotide and subsequent substitutions/alternations in amino acid profile such as observed in this study is consistent with previously described theories of evolution of RNA viruses particularly NDV (Yu et al. 2001; Umali et al. 2013).

As the pigeon flock had a history of vaccination with lentogenic strain (LaSota), identification of cleavage motif similar to velogenic raises concerns for the type of vaccine used to vaccinate the flock and the need for post-vaccine evaluation. Substitution and subsequent mutations at fusion peptide, HR regions, trans-membrane domain and antigen neutralization sites could affect the fusion activity of NDV and, alteration in antigenic epitopes particularly those that are involved in virus attachment, could result in escape variants and subsequent vaccine failure (Cho et al. 2007; Umali et al. 2014; Wang et al. 2015). Beside potential compromise in procedures used in vaccine storage and administration, the expected genetic distance between vaccine strains and the study isolate seems to be well-explained by Wang et al. (2015) through cross-HI assay. While evaluating the antigenic diversity of different strains through cross-HI assay, they reported a lower R-value (0.13–0.18) for interaction of PPMVs to LaSota than between PPMVs (VIa and VIb, 0.7) indicating an obvious antigenic difference with vaccine strain.

**Conclusion**

We characterized the circulating genotype of pigeon (PPMV-1) closely related to previously known clades of VIb. Since outbreaks of NDV in commercial poultry has been reported in one of the province of Pakistan from a closely related genotype (VIC) to the study isolate (VIb), future studies relating to disease surveillance coupled with experiments involving potential to transmit and shedding from either vaccinates or non-vaccinates are essential. Moreover, immunity barrier provided by LaSota and other vaccinal strains needs to be evaluated in the face of newly emerging NDV strains.

**Authors’ contributions**

Conceived and designed the experiment: SA, MAM, MR and MZS. Sampling and laboratory work: SA, MYT, AR and MZS. Analyzed the data: SA, KM, AR and MZS. Wrote the manuscript: SA, AR and MZS. All authors read and approved the final manuscript.

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**Competing interests**

The authors declare that they have no competing interests.

**Ethical approval**

This article does not contain any experiment or research with animals performed by any of the authors.

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