

Characterization of Pigeon Paramyxoviruses (Newcastle disease virus) Isolated in Kazakhstan in 2005*

Andrey Bogoyavlenskiy^{1**}, Vladimir Berezin¹, Alexey Prilipov², Eugeni Usachev²,
Ilya Korotetskiy¹, Irina Zaitceva¹, Aydyn Kydyrmanov¹ and Marat Sayatov¹

(1. Institute of Microbiology & Virology, Almaty 050010, Kazakhstan; 2. D.I. Ivanovsky Institute of Virology, Russian Academy of medical sciences, Moscow 123098, Russia)

Abstract: Isolates of Newcastle disease virus (NDV) from deceased wild and domestic pigeons in Kazakhstan were obtained from the Almaty region during 2005 and were genotypically analyzed by using reverse transcription polymerase chain reaction (RT-PCR) with primers specific to the viral fusion (F) protein gene. Part of the amplified F protein DNA product (nucleotide sequence 47–422) and the deduced amino acid sequences were compared phylogenetically with those from strains previously reported in other geographic regions. Phylogenetic analysis indicated that the Kazakhstani pigeon paramyxovirus type 1 (PPMV-1) isolates belong to genotype VI or 4bii. To our knowledge, this is the first reported VI isolates that possess the sequences of ¹¹²GKRQKR¹¹⁶* F¹¹⁷ within the F0 protein. The information is fundamental to improving the efficiency of control strategies and vaccine development for NDV.

Key words: Newcastle disease virus; Paramyxovirus; Phylogenetic characterization; Pigeon

Avian paramyxovirus type I of pigeons (PPMV-1) is an antigenic variant of avian paramyxovirus type 1 (APMV-1; Newcastle disease virus) of chickens that is responsible for an autonomous Newcastle disease (ND)-like infectious disease of pigeons. Clinical symptoms pigeons and doves infection include

paralysis wings, legs or stiff neck and excessive drinking, watery to haemorrhagic diarrhoea^[3]. The APMV-I viruses, including PPMV-1, are members of the genus *Avulavirus* in the family *Paramyxoviridae*, order *Mononegavirales*^[10]. NDV isolates have been divided into nine genotypes by phylogenetic analysis of the part of gene, including the F protein cleavage site, and were reclassified into six distinct lineages (1 to 6) by the using of monoclonal antibody binding method (mAb group P)^[2, 16] PPMV-I viruses associated with the permanent panzootic in pigeons and doves were placed into sublineage 4b(VIb) of lineage 4(VI)

Received: 2011-11-18, Accepted: 2012-02-27

* Foundation items: USDA-ISTC partner project (K-747p, Institute of Microbiology and Virology funding, and USDA CRIS (6612-32000-038-00 D)

** Corresponding author.

Phone: +727-291-8248, Fax: +727-291-3355,

E-mail: anpav_63@mail.ru

which later were divided into two 4bi and 4bii groups. Each of subgroups, 4bi and 4bii were subdivided into some clades, a, b, c and d, e, f, g respectively^[2,4,8,10,16,17]. In chickens, intracerebral pathogenicity (ICPI) values for PPMV-1 are typical of mesogenic Newcastle disease viruses but in most cases, PPMV-1 isolates have increased their virulence for chickens after passage, and therefore represent a threat to poultry production^[2, 9]. Besides pigeons, doves and chickens, PPMV-1 viruses have also been isolated from kestrels, falcons, cockatoos, budgerigars, pheasants, swans and a robin^[1, 2, 9, 10, 12].

Large die-offs in doves and pigeons have occasionally been reported in the Almaty region of Kazakhstan in 2005. In the present study, the phylogenetic relationships between 3 Kazakhstani PPMV-1 viruses isolated from pigeons were investigated. We decided to describe these three strains of NDV due to the fact that since 1995, only a few strains of this monophyletic group of viruses isolated in Poland, Austria and Croatia were reported, and suddenly the same viruses were isolated in Kazakhstan which was previously characterized by another group of these viruses.

MATERIALS AND METHODS

Viruses

Three NDV isolates were recovered from pigeons during the 2005 NDV outbreaks in Almaty, Kazakhstan. The observed clinical signs and high mortality were evidence of the severity of this Newcastle disease outbreak (NDV). NDV isolates were recovered from samples taken from dead birds. Initial isolation of the virus was performed in 9–10 day old embryonated chicken eggs (ECE). The type of

the virus was determined in standard haemagglutination inhibition and neuraminidase inhibition tests using specific antisera to the reference strains of paramyxovirus and influenza virus. Allantoic fluids were harvested from ECE inoculated with the viruses and used as a stock for sequence analysis. Methods for biological and molecular characterization were as described previously^[5, 6].

Sequencing and Phylogenetic Analysis

Sequence analysis of PCR products was completed using an “fmol DNA Sequencing System” (Promega, USA) or ABI PRISM BigDyeTM Terminator cycle sequencing reaction kit (Applied Biosystem, USA).

After sequencing, assembly of sequences, removal of low quality sequence data and nucleotide sequence translation into protein sequence, additional multiple sequence alignments and processing were performed with the Lasergene package and the BioEdit software version 7.1.3 with an engine based on the Clustal W version 2.0 algorithm. Phylogenetic analysis, based on the comparison of the nucleotide sequences of F gene fragments (from 47 to 421), was conducted with the Molecular Evolutionary Genetics Analysis (*MEGA*, version 5.0) software package using maximum likelihood method to infer evolutionary trees and conduct the bootstrap test for nucleotide and amino acid alignments^[13-15].

In addition to the 3 NDV strains isolated in Kazakhstan, 136 previously described NDV sequences representing different NDV genotype groups were used for comparison (on-line Supplementary Table S1). The nucleotide sequences determined in this study are available in GenBank under accession numbers JN806235, JN806236, JN806237.

Supplementary Table S1. List of representative NDV strains taken from GenBank database for phylogenetic analysis

Isolate identification	Genotype or subgroup	Accession number	Isolate identification	Genotype or subgroup	Accession number
PUKPI86239	VI	AY471854	PDKPI93202	VI	AY471761
PUKPI183299	VI	AY471852	Pi-Japan-Tochigi-95	VI	Ab070419
PUKPI183264	VI	AY471851	Pi-Japan-Fukushima-96	VI	Ab070423
PITPI84354	VI	AY471850	Pi-Japan-Saitama-97	VI	Ef030957
PUKCK84263	VI	AY471858	Pi-Japan-Shiga-96	VI	Ab070422
PHKPI86358	VI	AY471856	PFRPI98372	VI	AY471765
PBEPI84352	VI	AY471849	Pi-Japan-FK-1-84	VI	AB070434
PUKPI84260	VI	AY471848	Pi-Japan-Niigata-88	VI	AB070434
PUKPI84259	VI	AY471847	FR-99299	VI	AB070434
PITDO00289	VI	AY471846	STP96	VI	AB070434
PUKPI84342	VI	AY471855	Vh 357-06	VI	EU240576
EG-3-87	VI	AY150111	HR-3-02	VI	AY150165
EG-6-87	VI	AY150112	HR-111-01	VI	AY150162
NO-1-85	VI	AY150105	DOZA05AM68313	VI	EF030952
GB-1168-84(Vib)	VI	AF109885	DOZA05N240	VI	EF030953
SE-2-83	VI	AY150100	PATPI00323	VI	AY471789
Pi-Japan-Ehime-93	VI	AB070416	WX-10-07-Pi	VI	GQ281086
SE-8-00	VI	AY150158	DOZA06N591	VI	EF030959
Pi-Japan-Kumamoto-95	VI	AB070417	Vh 536-06	VI	EU240582
Pi-Japan-Tokachi-91	VI	AB070413	Vh 529-06	VI	EU240580
China-Sh-2-98	VI	AF458017	Vh 538-06	VI	EU240583
China-XJ-1-97	VI	AF458021	Gxp40	VI	AY635815
SZb9803	VI	AY390293	SD-15-08-Pi	VI	GQ245801
SZA9803	VI	AY390292	Pi-Dnipro-2007	VI	EF030957
NP9904	VI	AY390290	PI-PL-H2-10	VI	HM627541
YZ9712	VI	AY390288	BG-99-82	VI	AF402131
NC9701	VI	AY390289	CA-28-89	VI	AY150119
China-XJ-3-97	VI	AF458019	Chicken-Japan-Chiba-69	VI	AB070387
China-ZhJ-2-86	VI	AF458016	DE-60-93	VI	AY150134
SE-3-90	VI	AY150126	DE-61-93	VI	AY150135
ZC94	VI	AY390295	HR-65-95	VI	AY150144
Z32	VI	AY390296	HR-155-01	VI	AY150163
Pi-Japan-Gunna-2000	VI	AB070434	HU-61-83	VI	AY150099
Astr-74	VI	NDI243391	HU-238-84	VI	AY150103
Iraq AG68	VI	AF001108	JAEFA96038	VI	AY175738
Israel 70	VI	AF001111	JN08	VI	HM776583
Kuwait 256	VI	AF001109	Lebanon 70	VI	AF001110
Vh448-06	VI	EU240578	NDV05-027	VI	DQ439884
JS-35-07PI	VI	GQ281085	Pigeon-NY-US-1984	VI	EF520716
PIQPI178442	VI	AY471857	SD-2-75	VI	AY151384
PUKPI02434	VI	EF030957	BG-29-86	VI	AF402135
PUKPI02431	VI	AY471755	BG-72-74	VI	AF402116
PUKPI01426	VI	AY471753	BG-48695	VI	AF402138
PUKPI02435	VI	AY471754	BG-99-82	VI	AF402131
PAEKE99364	VI	AY471785	GR-12-68	VI	EU604251
YZ-21-07-Pi	VI	AB070434	KR-5-99	VI	EU665683
PTRBU95211	VI	AY471773	CH-1-95	VI	AF001132
Warwick	VI	NDVWAWRFG	DK-1-95	VI	AF001129
PDEPI94216	VI	AY471763	PB9601	VI	AY390291
ES-3-92	VI	AY150132	China-98-1	VI	AF358785
DOZA05N247	VI	EF030954	China-JC-2-98-Go(VIb)	VI	AF456439
DOZA05N417	VI	EF030955	TW-06-223	VI	DQ898521
OZA06N549	VI	EF030957	Chicken-Japan-Chiba-81	VI	AB070388

Table 1 (continue)

Kr-12A-8	VI	AY630414	NL-1-93	VII	AF001124
Kr-102-89	VI	GQ507801	TR-8-97	VII	AF136785
Kr-163-90	VI	AY630416	ZA-5-68	VIII	AF136762
Kr-M-88	VI	AY630413	ZA-10-74	VIII	AF136763
SNU88139	VI	AF257749	B-14-93	VII	AF001121
BOR74	I	Y16049	BG-109-84	VII	AF402133
Ulstr	I	NDVULSTFG	China-T02	VII	AY390307
Komarov	II	AY170137	China-YLF3	VII	AY390312
ZA-405-01	II	AF532748	D-83-95	VII	AF001118
H-Ph-02	III	AY170136	MZ 46-95	VII	AF136778
Muktesvak1	III	AY117022	FIN-1-96	VII	AF091623
Pok-70	IV	NDI243388	RI-1-88	VII	AF001134
Simf-64	IV	NDV19017	ZA-32-93	VII	AF136771
YU(VO)-C-91	V	AY117003	ZA-549-99	VII	AY210500
IT-129-88	V	AF218131			

RESULTS AND DISCUSSION

Estimation of NDV Isolates Pathotype

The biological properties of pigeon NDV strains isolated in Kazakhstan during 2005 are presented in Table 2. NDV isolates were characterized as mesogenic based on their mean death time (MDT), which ranged from 60,8 to 73 h and intracerebral pathogenicity index (ICPI) that ranged from 0,51 to 1.06. The deduced amino acid sequence of F protein cleavage site for all of these strains was determined to be ¹¹²GKRQKR¹¹⁶*F¹¹⁷ and was characteristic of mesogenic NDV strains.

Phylogenetic Relationships among NDV Isolates

Phylogenetic analysis of pigeon NDV strains isolated in Kazakhstan and study of their phylogenetic relationships with other NDV worldwide isolates was completed based on sequence analysis of the variable region of the F gene (47-421) (Fig. 1 and Fig. 2). Although the bootstrap values supporting the phylogenetic relationships within lineages 4bi and 4bii are low^[1,2], amino acid sequences support the phylogenetic relationships: lineage 4bi clade (c) is distinguished from clades (a) and (b) by T3, L10 and R27 substitutions, and lineage 4bii clade (d) differs

from clade (e) by a P36→S substitution. The low level of bootstrap analysis is connected with continuing evolution of this group of viruses from regular outbreaks

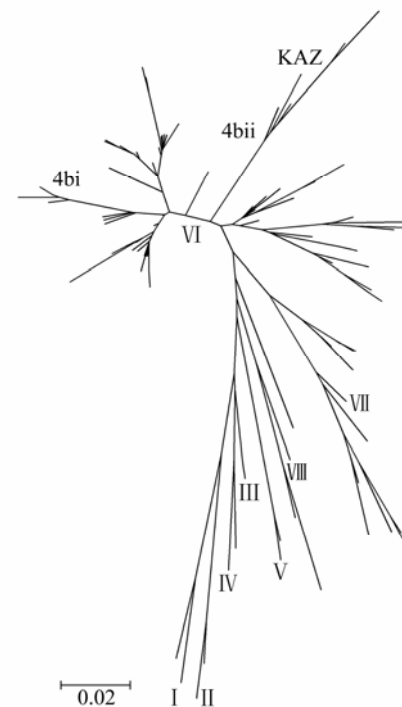


Fig. 1. Unrooted maximum likelihood radial phylogram based on nucleotide sequence data from 139 APMV-1 isolates, including 115 PPMV-1 isolates and 24 representative of the other genetic lineages^[2]. The region analysed was a 374 base pair fragment (47 to 422) at the 3' end of the fusion protein gene. Branch lengths represent the predicted number of substitutions and are proportional to the differences between the isolates. The individual names of each PPMV-1 isolate included in this phylogram are not included. Bar=0.02 μ m.

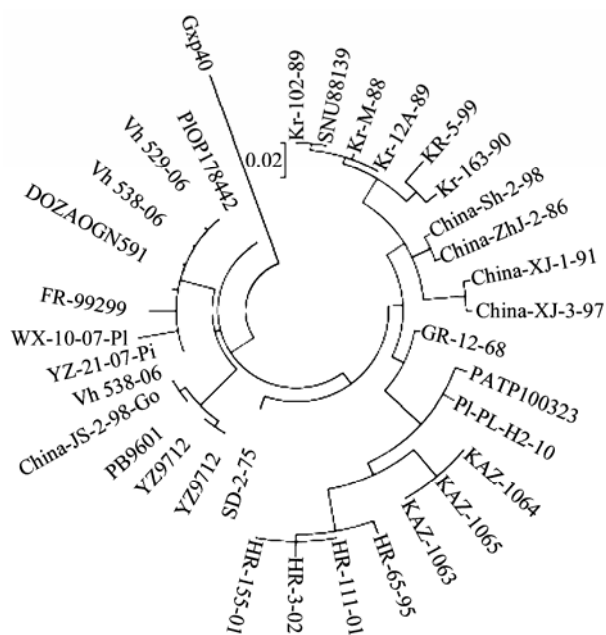


Fig. 2. Unrooted maximum likelihood circular phylogram based on nucleotide sequence data from 39 PPMV-1 4bii isolates. The evolutionary history was inferred by using the Maximum Likelihood method based on the Tamura-Nei model [14]. The tree with the highest log likelihood (-987.0619) is shown. Initial tree(s) for the heuristic search were obtained automatically as follows. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. The analysis involved 35 nucleotide sequences. Evolutionary analyses were conducted in MEGA5 [15].

of pigeons in the various regions of the globe. The phylogenetic analysis indicated that the 3 Kazakhstania PPMV-1 strains cluster together with viral isolates from Poland, Austria and the Croatia. The nucleotide substitutions characteristic for this group of Kazakhstan viruses were found in positions 18 (T changed to C), 32 (T changed to C), 43 (C

changed to T), 111 (T changed to C), and 187 (G changed to A). Calculation of the synonymous and non-synonymous substitution rate demonstrate that all isolates have a rate ranging between 0.081 - 0.264; a value less than one demonstrates the presence of only purifying (negative) selection, despite the most variable portion of the F gene between nucleotide positions 47-421 was used in the analysis. Phylogenetic analysis based on the amino acid NDV fusion protein exhibited similar properties (Fig. 3). According to the phylogenetic tree, all the isolates belong to the genotype to the genotype VI or 4bii—of sublineage 4b. A further analysis on the amino acid sequence of the protease cleavage site reveals that the F1 protein of all the recent Kazakhstania isolates contain a phenylalanine (F) on the residue 117 on the N-terminus and four basic amino acids in the motif. ¹¹²GKRQKR¹¹⁶*F¹¹⁷, which indicates they are velogenic strains^[2, 3]. Based on amino acids analysis, unique amino acid substitutions (V₁₁ to A₁₁, C₂₇ to R₂₇, V₆₃ to I₆₃, T₉₀ to A₉₀) in all strains of this monophyletic group were found.

Although these results demonstrated some sequence similarity between the isolated strains and strains responsible for outbreaks of Newcastle disease in Europe, it is the region of isolation of this monophyletic group of viruses that is unusual. The results indicate that the exchange of breeding birds by owners of pigeon lofts may become a serious factor in

Table 2. Biological and molecular characteristics of NDV isolates recovered from pigeons in Kazakhstan during 2005

Isolate identification	Strain abbreviation	ICPI	MDT	Fusion protein cleavage site (molecular pathotyping)
PPMV- 1/pigeon/Алматы/1063/05	KAZ-1063-05	0,51	60,8	¹¹² GKRQKR ¹¹⁶ *F ¹¹⁷
PPMV- 1/pigeon/Алматы/1064/05	KAZ-1064-05	1,06	73,0	¹¹² GKRQKR ¹¹⁶ *F ¹¹⁷
PPMV- 1/pigeon/Алматы/1065/05	KAZ-1065-05	0,81	60,8	¹¹² GKRQKR ¹¹⁶ *F ¹¹⁷

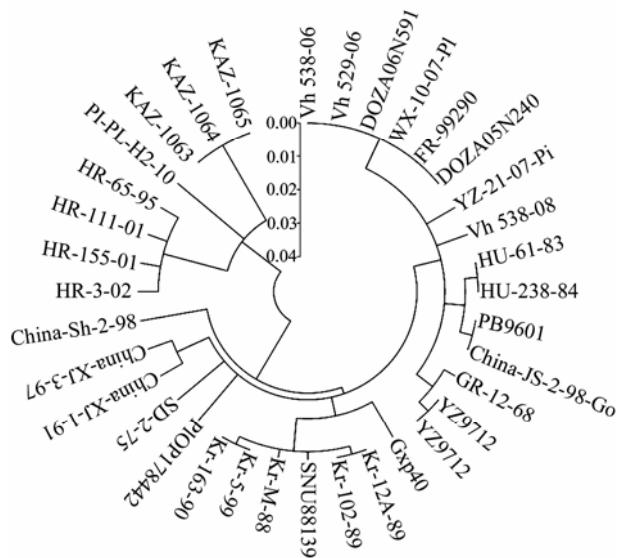


Fig. 3. Unrooted maximum likelihood circular phylogram based on amino acids sequence data from 39 PPMV-1 4bii isolates. The evolutionary history was inferred by using the Maximum Likelihood method based on the JTT matrix-based model [7]. The tree with the highest log likelihood (-445.5714) is shown. Initial tree(s) for the heuristic search were obtained automatically as follows. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. Evolutionary analyses were conducted in MEGA5^[15].

the spread of Newcastle disease virus because, at the time of this outbreak, the National Exhibition of Racing Pigeons took place, organized by the Polish Association of Racing Pigeon Breeders, the main institution which deals with pigeon breeding. Another possibility may be introduction of new NDV strains from migrating birds during their direct contact with chickens in primarily open poultry farms in Kazakhstan. Further characterization of NDV strains circulating in Kazakhstan and other Central Asia countries is important for better control and understanding of NDV epizootics in the Central Asia region.

Acknowledgments

The study was supported by USDA-ISTC partner

project (K-747p, Institute of Microbiology and Virology funding, and USDA CRIS (6612-32000-038-00 D).

References

- 1 **Abolnik C, Gerdes G H, Kitching J, et al.** 2008. Characterization of pigeon paramyxoviruses (Newcastle disease virus) isolated in South Africa from 2001 to 2006. *Onderstepoort J Vet*, 75:147-152.
- 2 **Aldous E W, Fuller C M, Mynn J K.** 2004. A molecular epidemiological investigation of isolates of the variant avian paramyxovirus type 1 virus (PPMV-1) responsible for the 1978 to present panzootic in pigeons. *Avian Pathol*, 33(2): 258-269.
- 3 **Alexander D J, Parsons G.** 1986. Pathogenicity for chickens of avian paramyxovirus type I isolates obtained from pigeons in Great Britain during 1983-1985. *Avian Pathol*, 15: 487-493.
- 4 **Biancifiore F, Fioroni A.** 1983. An occurrence of Newcastle disease in pigeons: virological and serological studies on the isolates. *Comp Immunol Microbiol Infect Dis*, 6: 247-252.
- 5 **Bogoyavlenskiy A, Berezin V, Prilipov A, et al.** 2009. Newcastle disease outbreaks in Kazakhstan and Kyrgyzstan during 1998, 2000, 2001, 2003, 2004, and 2005 were caused by viruses of the genotypes VIIb and VIId. *Virus Genes*, 39(1): 94-101.
- 6 **Bogoyavlenskiy A, Berezin V, Prilipov A, et al.** 2005. Molecular characterization of virulent Newcastle disease virus isolates from chickens during the 1998 NDV outbreak in Kazakhstan. *Virus Genes*, 31(1):13-20.
- 7 **Jones D T, Taylor W R, Thornton J M.** 1992. The rapid generation of mutation data matrices from protein sequences. *Comput Appl Biosci*, 8: 275-282.
- 8 **Kaleta E F, Alexander D J, Russell P H,** 1985 The first isolation of the avian PMV-1 virus responsible for the current panzootic in pigeons? *Avian Pathology*, 14: 553-557.
- 9 **Krapez U, Racnik J, Slavec B, et al.** 2010. Detection and molecular characterization of a pigeon variant of avian paramyxovirus type 1 virus (PPMV-1) from a blackbird (*Turdus merula*). *Slov Vet Res*, 47(3): 83-90.

- 10 **Lee Y J, Sung H W, Choi J G, et al.** 2004. Molecular epidemiology of Newcastle disease viruses isolated in South Korea using sequencing of the fusion protein cleavage site region and phylogenetic relationships. **Avian Pathol**, 33(5): 482-491.
- 11 **Mayo M A.** 2002. Virus Taxonomy – Houston. 2002. **Arch Virol**, 147: 1071-1076.
- 12 **Rui Z, Juan P, Jingliang S, et al.** 2010. Phylogenetic characterization of Newcastle Disease virus isolated in the mainland of China during 2001-2009. **Vet Microbiol**, 141: 246-257.
- 13 **Saitou N, Nei M.** 1987. The neighbor-joining method: A new method for reconstructing phylogenetic trees. **Mol Biol Evol**, 4: 406-425.
- 14 **Tamura K, Nei M, Kumar S.** 2004. Prospects for inferring very large phylogenies by using the neighbor-joining method. **P Natl Acad Sci USA**, 101:11030-11035.
- 15 **Tamura K, Peterson D, Peterson N, et al.** 2011. MEGA5: Molecular Evolutionary Genetics Analysis using Maximum Likelihood, Evolutionary Distance, and Maximum Parsimony Methods. **Mol Biol Evol**, 28(10): 2731-2739.
- 16 **Ujvari D, Wehmann E, Kaleta E F, et al.** 2003. Phylogenetic analysis reveals extensive evolution of avian paramyxovirus type 1 strains of pigeons (*Columba livia*) and suggests multiple species transmission. **Virus Res**, 96: 63-73.
- 17 **Wilson G W.** 1986. Newcastle disease and paramyxovirus 1 of pigeons in the European Community, **World Poultry Sci J**, 42: 143-153.
- 18 **Zuckerandl E, Pauling L.** 1965. Evolutionary divergence and convergence in proteins. In: **Evolving Genes and Proteins**, Bryson V, Vogel H J, ed. New York: Academic Press, pp97-166.