Genotypic characterisation of Avian paramyxovirus type-1 viruses isolated from aquatic birds in Uganda

Avian paramyxovirus type-1 (APMV-1) viruses of the lentogenic pathotypes are often isolated from wild aquatic birds and may mutate to high pathogenicity when they cross into poultry and cause debilitating Newcastle disease. This study characterised APMV-1 isolated from fresh faecal droppings from wild aquatic birds roosting sites in Uganda. Fresh faecal samples from wild aquatic birds at several waterbodies in Uganda were collected and inoculated into 9–10-day-old embryonated chicken eggs. After isolation, the viruses were confirmed as APMV-1 by APMV-1-specific polymerase chain reaction (PCR). The cleavage site of the fusion protein gene for 24 representative isolates was sequenced and phylogenetically analysed and compared with representative isolates of the different APMV-1 genotypes in the GenBank database. In total, 711 samples were collected from different regions in the country from which 72 isolates were recovered, giving a prevalence of 10.1%. Sequence analysis of 24 isolates revealed that the isolates were all lentogenic, with the typical GGRQGL’L avirulent motif. Twenty-two isolates had similar amino acid sequences at the cleavage site, which were different from the LaSota vaccine strain by a silent nucleotide substitution T357C. Two isolates, NDV/waterfowl/Uganda/MU150/2011 and NDV/waterfowl/Uganda/MU186/2011, were different from the rest of the isolates in a single amino acid, with aspartate and alanine at positions 124 and 129, respectively. The results of this study revealed that Ugandan aquatic birds indeed harbour APMV-1 that clustered with class II genotype II strains and had limited genetic diversity.

Introduction

Avian paramyxovirus type 1 (APMV-1) belongs to the genus *Avulavirus*, in the family Paramyxoviridae, and is responsible for causing Newcastle disease (ND), a highly infectious disease for poultry (Alexander 2003). These viruses are usually grouped according to the pathotype based on the severity of the disease they may cause: velogenic (high virulence), mesogenic (mild virulence) and lentogenic (low virulence) viruses. The genetic basis of virulence is defined by the amino acid residues at the cleavage site of the fusion protein precursor (F₁) (Aldous & Alexander 2001). The more virulent pathotypes that cause ND in poultry have multiple basic amino acids (at least three arginine or lysine residues between residues 113 and 116) with a phenylalanine at position 117 of the C-terminus of the F₂ protein (also the N-terminus of the F₁ protein) that make them cleavable by many proteases within the host tissues (Aldous & Alexander 2001; Aldous et al. 2003). The avirulent strains have fewer basic amino acid residues, with a leucine residue at position 117 of the F-protein. This portion of the cleavage site of the F-protein has been used not only for pathotyping APMV-1 viruses but also for genotyping the different strains (Aldous et al. 2003) among other typing methods that classify these strains into genotypes and lineages (Kim et al. 2007; Snoeck et al. 2009). Most of the genotyping methods classify these viruses into two classes, I and II, and further subdivide them into different genotypes and clades. The most recent classification has utilised the full F gene to classify these viruses further (Diel et al. 2012; Snoeck et al. 2013). The class II viruses are the most commonly reported and studied viruses and are associated with disease in poultry, pet and wild birds.

The role of APMV-1 recovered from wild birds has been alluded to in the epidemiology of ND in domestic poultry and aquatic birds in particular are thought to be the natural reservoirs (Jorgensen et al. 2004). Avirulent strains have particularly been consistently recovered from aquatic birds and their potential to mutate to virulent form upon passage in poultry has been confirmed (Shengqing et al. 1997; Takakuwa et al. 1998).
The migration of wild bird populations along various migration pathways across the world constitutes a serious threat to possible spread of these viruses and pose a risk of transmission to domestic poultry (Zarkov et al. 2005). Studies have already shown similarities between strains recovered from aquatic birds and shorebirds with those isolated from live-bird markets in some parts of the world (Kim et al. 2007), further confirming this threat. We have previously demonstrated that virulent APMV-1 strains circulate in live-bird markets in Uganda in apparently healthy birds (Byarugaba et al. 2014). However, despite the massive number of migratory and resident birds that rest along several waterbodies in Uganda, no studies have been undertaken to understand whether these birds harbour APMV-1. With the biggest proportion of the poultry production sector in Uganda being backyard, there is a high risk of transmission of the APMV-1 from aquatic birds into the poultry population that may result in economic losses to both the small-scale poultry farmers and the country. This study sought to establish if APMV-1 strains circulate in aquatic birds in Uganda and how they compare genetically to others elsewhere.

**Materials and methods**

**Sample collection**

Fresh faecal samples were collected from aquatic bird (which includes all aquatic birds and waterfowl) roosting sites along various waterbodies across the country using sterile dacron swabs into cryovials containing virus transport medium supplemented with antibiotics (isotonic phosphate buffered saline, 2000 U/mL penicillin, 2 mg/mL streptomycin, 50 μg/mL gentamycin, 50 U/mL nystatin and 0.5% bovine serum albumin). The samples were stored and transported in dry shippers until delivered to the laboratory where they were stored at -80°C until further use. A total of 711 samples were collected from various sites including Musambwa Island, Makanaga Bay, Lutembe Bay, Makamba Bay, Nakivogo landing site, Samuka Island, Macdonald Bay, Doho Rice Scheme, Lake Bisma Island, Murchison Falls National Park, Queen Elizabeth National Park, and Kibimba Dam Rice Scheme.

**Virus isolation**

The samples were inoculated (in triplicate) by the allantoic route into 9–10-day embryonated chicken eggs for virus isolation according to the World Organisation for Animal Health (OIE) Manual of Standards for Diagnostic Tests and Vaccines (OIE 2008). Allantoic fluid was harvested 3 days post-inoculation and subsequently tested for haemagglutination (HA) using 1% chicken erythrocytes and haemagglutination inhibition (HI) with in-house-generated polyclonal anti-APMV-1 sera as described (OIE 2008). The HI-positive samples were subsequently confirmed by polymerase chain reaction (PCR).

**Confirmation by reverse transcription-polymerase chain reaction**

Viral ribonucleic acid (RNA) was extracted from all the HI-positive samples using the QIAamp Viral RNA mini kit (Qiagen, Germantown, MD, USA) according to the manufacturer’s instructions. Polymerase chain reaction was performed with a Qiangen one-step reverse transcription-polymerase chain reaction (RT-PCR) kit (Qiagen, USA) according to the manufacturer’s instructions, with the following APMV-1 primers FOP1: 5’TACACCTCATCCCCAGA CAGGAGTC 3’ (nucleotide position, 158–177) and FOP2: 5’AGGCCAGGGAAGTGATTGTGGC 3’ (nucleotide position, 493–513) according to Kho et al. (2000). The primers were used for the amplification of a 356 bp region corresponding to the cleavage activation site of F gene of APMV-1. The RT-PCR was performed in a 25 μL reaction volume containing 5 μL of 5X RT-PCR buffer, 11 μL of RNase-free H2O, 1 μL of 10 mmol/L dNTPs, 1.5 μL of 10 mmol/L of each primer, 2 μL of 50 mM MgCl2, 1 μL of enzyme mix (Taq DNA polymerase and reverse transcriptase) and 2 μL of viral RNA extract. Amplification was carried out in an Applied Biosystems Veriti 96-well thermocycler with a single reverse transcription (RT) step of 50 ºC for 30 min, a denaturation step of the RT (95 ºC) for 15 min, followed by 40 cycles with 30 s denaturation at 95 ºC, 30 s of primer annealing at 58 ºC, 1 min of extension at 72 ºC and a final extension for 10 min at 72 ºC. The samples (including a known positive control) were then separated on a 1% agarose gel with a 100-bp marker.

**Sequencing**

A total of 24 representative isolates were selected for sequencing of the partial cleavage site of the fusion gene. Out of the 72 isolates, we selected 24 isolates chosen proportionally from each site including 7/23 from Musambwa, 8/22 from Lutembe, 5/8 from Makanaga, 1/6 from Samuka, 1/7 from Nakivogo and 1/6 from Queen Elizabeth National Park. The fragments were run on a 1% agarose gel, excised from the gel and purified with QIAquick PCR Purification Kits (Qiagen, USA) according to the manufacturer’s recommendations. Sanger sequencing was carried out on the purified PCR products using the same primers that were used for the PCR. Sequencing was performed on a 3130XL Applied Biosystems capillary sequencer at the Plateau de Génomique GeT-Purpan, UDEAR UMR 5165 CNRS/UPS, CHU PURPAN, Toulouse, France.

**Phylogenetic analysis**

The Basic Local Alignment Search Tool (BLAST) was used to find similar F gene sequences for APMV-1 in the Genbank. Sequences covering the cleavage site of the fusion gene representing all 18 genotypes of APMV-1 recently described by Diel et al. (2012), including all the Ugandan and East African sequences, were retrieved. The sequences were aligned together with the Ugandan sequences generated in this study using Clustal W and edited using Bioedit Software version 5.0.9 (Hall 1999). Phylogenetic analysis was performed using the MEGA version 5.05 program (Tamura et al. 2011) with the neighbour-joining (NJ) Kimura 2-parameter method and 1000 bootstrap replicates. The amino acids around the fusion protein cleavage site were compared to representative sequences from each of the genotypes. Vaccine strains – LaSota, accession number: JF950510; Hitchner BI, accession
number: JN872151 and I-2, accession number AY935499 – were also included in the analysis.

Availability of data and materials

The sequences of the cleavage site of the isolates analysed in this study were deposited in GenBank with accession numbers LT549451, LT549452 and LT549453. These include NDV138/aquatic birds/Uganda/2011 representing the 22 isolates with identical sequences and NDV150/aquatic birds/Uganda/2011 and NDV186/aquatic birds/Uganda/2011 as indicated in the phylogenetic tree legend in Figure 2.

Ethics

This study was approved by the College of Veterinary Medicine Animal Resources and Biosecurity Higher Degrees Research Committee and Uganda National Council of Science and Technology (Approval # HS 776).

Results

Occurrence of Avian paramyxovirus type-1

From the 711 samples collected, 72 isolates were recovered. The prevalence at each site ranged from 0% to 36% and the average was estimated as 10.1% by HI (Table 1). No isolates were recovered from Doho Rice Scheme, Lake Bisina, Mabamba, Murchison Falls and Kibimba Dam Rice Scheme. Six isolates were obtained from Queen Elizabeth National Park, 22 from Lutembe, 8 from Makanaga, 23 from Musambwa, 7 from Nakivogo and 6 from Samuka (Table 1). Musambwa Island and Lutembe Bay provided the highest number of isolates, with Queen Elizabeth and Nakivogo providing the lowest number of isolates. Prevalence was highest in Musambwa, Lutembe and Makanaga. At the sites, the most common bird species were the grey-headed gull, white-winged tern, gull-billed tern, long-tailed cormorants, great cormorant, Egyptian geese and others as indicated in Table 1.

Pathotypes of Avian paramyxovirus type-1 in Ugandan aquatic birds

The deduced amino acid sequences of the F gene cleavage site were used to determine the pathotypes and are shown in Table 2. All 24 isolates sequenced had a lentogenic motif of $^{111}$GGRQGR'T$^{117}$ characteristic of the avirulent strains. However, isolates NDV150/waterfowl/Uganda/2011 and NDV186/waterfowl/Uganda/2011 were different from the rest of the 22 isolates in a single amino acid; aspartate and alanine at positions 124 and 129, respectively. The rest of the 22 isolates were identical at all positions.

Phylogenetic analysis: To determine the phylogenetic relationships between Ugandan aquatic birds’ isolates, other Ugandan strains from poultry and the rest of the world, the sequences of the 201-bp hypervariable region of the F gene were compared to the corresponding region of viruses available in GenBank. Results from the phylogenetic analysis clustered our isolates with genotype II strains which had also been historically described as genotype II or lineage 2. They were different from the recently isolated strains from poultry in Uganda which belonged to genotype V as shown in Figure 2.

Discussion

This study is the first to isolate and characterise APMV-1 from aquatic birds in Uganda. Most work on APMV-1 in many parts of the world has focused on poultry, where occurrence of both virulent strains and lentogenic strains of class II has been reported with limited studies in wild birds (de Almeida et al. 2013; Miller, Decanini & Afonso 2010). Virulent APMV-1 is a common cause of infections in birds and more than 230 bird species have been reported to be susceptible in experimental infections (USDA/APHIS/WS 2016) including those we found at the sites where we collected our samples, such as the cormorants. The economic impact of ND on the poultry industry in Uganda and elsewhere is significant both in the backyard and commercial flocks. Little is known about the APMV-1 strains circulating in wild birds, their evolution and their role in the epidemiology of the disease. Czeglédi et al. (2006) speculated that class I and class II genotype I are ancestral representatives of APMV-1 maintained by their natural hosts, the wild waterfowl.

In the present study, we demonstrated the occurrence of APMV-1 among aquatic birds in Uganda with a 10.1% prevalence, which was higher than that reported (2.1%) in other studies in Africa (de Almeida et al. 2013). A few studies in Uganda have shown the occurrence of virulent APMV-1 strains (genotype V) that circulate among Ugandan poultry and live-bird markets (Byarugaba et al. 2014; Otim et al. 2004). Unlike in our study where the 24 sequenced isolates were clustered with genotype II, other studies of APMV-1 in wild birds in Africa and elsewhere have demonstrated the presence.
TABLE 1: Occurrence of Avian paramyxovirus type-1 by site among migratory aquatic birds.

| Sampling site                      | GPS coordinates (longitude, latitude) | Estimates of total bird | Most predominant bird species at site | Occurrence |
|------------------------------------|--------------------------------------|-------------------------|--------------------------------------|------------|
| Samuka Island                      | 33.275185, 0.399174                  | 420                     | Long-tailed Cormorant, Little Egret, African Open-billed Stork, S. longirostris, Gull, S. heuglini | 9.2        |
| Makenaga wetland                   | 32.655487, 0.091581                   | 250                     | Slender-billed Gull, Gull-billed Tern, A. jamaicensis, Long-tailed Cormorant, Yellow-billed Duck, G. atriceps | 17.4       |
| Macdonald Bay                     | 33.762360, 0.143766                   | 760                     | Long-tailed Cormorant, Grey-headed Gull, White-winged Tern, A. excavata, Greater Cormorant, E. tobata | 0.0        |
| Doho Rice Scheme                  | 33.704681, 0.135127                   | 560                     | Little Egret, African Open-billed Stork, Black Crane, S. longirostris, Black-headed Heron, G. atriceps | 0.0        |
| Musambwa Island                    | 32.439880, -1.750159                  | 650                     | Grey-headed Gull, Long-tailed Cormorant, Greater Cormorant, S. longirostris, Sacred Ibis, P. phalacrocorax, G. atriceps | 35.9       |
| Lutembe wetland                   | 32.581716, 0.155726                   | 1670                    | White-winged Tern, G. atriceps, Yellow-billed Duck, S. longirostris, Long-toed Plover, G. atriceps | 22.0       |
| Murchison Falls NP                | 31.482525, 2.201648                   | 1560                    | Pied Kingfisher, A. jamaicensis, African Darter, C. albus, Hadada Ibis, E. tobata | 0.0        |
| Nakiwogo –L. Victoria             | 32.365894, -0.047604                  | 8900                    | White-winged Tern, G. atriceps, Grey-headed Gull, Slender-billed Gull, L. longirostris, Egyptian Goose | 10.7       |
| Queen Elizabeth NP                | 29.911222, -0.222585                  | 4200                    | Pied Kingfisher, Greater Cormorant, E. tobata, Egyptian Goose, Hamerkop, A. jamaicensis | 4.4        |
| L. Bisina                         | 34.050064, 1.342055                   | 50                      | Long-tailed Cormorant, African Open-billed Stork, S. longirostris, White-faced Whistling Duck, E. tobata | 0.0        |
| Mabamba wetland                   | 32.430611, -0.027916                  | 220                     | Long-toed Plover, A. jamaicensis, Pied Kingfisher, White-faced Whistling Duck, Malachite Kingfisher, Common Squacco, Heron, L. longirostris | 0.0        |
| Kibimba Dam Rice Scheme           | 33.776093, -0.012789                  | 1800                    | Egrets, A. jamaicensis, T. sandvicensis, Long-tailed Cormorant, S. longirostris | 0.0        |

GPS, global positioning system.
†, number of positive over total samples collected.

TABLE 2: F-Protein cleavage site motif of the Newcastle disease viruses sequenced in this study.

| No. | Isolate name            | Date of isolation | Site/location | F gene cleavage site (111-117) sequence | Pathogenicity |
|-----|-------------------------|-------------------|---------------|-----------------------------------------|---------------|
| 1   | APMV-1/waterfowl/Uganda/MU122/2011 | 17/07/2011        | Queen Elizabeth NP/Kasese | GGRQGRFL | LENTOGENIC |
| 2   | APMV-1/waterfowl/Uganda/MU125/2011 | 11/07/2011        | Lutembe Bay/Entebbe | GGRQGRFL | LENTOGENIC |
| 3   | APMV-1/waterfowl/Uganda/MU126/2011 | 11/07/2011        | Lutembe Bay/Entebbe | GGRQGRFL | LENTOGENIC |
| 4   | APMV-1/waterfowl/Uganda/MU129/2011 | 11/07/2011        | Lutembe Bay/Entebbe | GGRQGRFL | LENTOGENIC |
| 5   | APMV-1/waterfowl/Uganda/MU130/2011 | 11/07/2011        | Lutembe Bay/Entebbe | GGRQGRFL | LENTOGENIC |
| 6   | APMV-1/waterfowl/Uganda/MU131/2011 | 11/07/2011        | Lutembe Bay/Entebbe | GGRQGRFL | LENTOGENIC |
| 7   | APMV-1/waterfowl/Uganda/MU132/2011 | 11/07/2011        | Lutembe Bay/Entebbe | GGRQGRFL | LENTOGENIC |
| 8   | APMV-1/waterfowl/Uganda/MU137/2011 | 11/07/2011        | Lutembe Bay/Entebbe | GGRQGRFL | LENTOGENIC |
| 9   | APMV-1/waterfowl/Uganda/MU138/2011 | 11/07/2011        | Lutembe Bay/Entebbe | GGRQGRFL | LENTOGENIC |
| 10  | APMV-1/waterfowl/Uganda/MU149/2011 | 25/09/2011        | Makanaga Bay/Mpigi | GGRQGRFL | LENTOGENIC |
| 11  | APMV-1/waterfowl/Uganda/MU150/2011 | 25/09/2011        | Makanaga Bay/Mpigi | GGRQGRFL | LENTOGENIC |
| 12  | APMV-1/waterfowl/Uganda/MU151/2011 | 25/09/2011        | Makanaga Bay/Mpigi | GGRQGRFL | LENTOGENIC |
| 13  | APMV-1/waterfowl/Uganda/MU152/2011 | 25/09/2011        | Makanaga Bay/Mpigi | GGRQGRFL | LENTOGENIC |
| 14  | APMV-1/waterfowl/Uganda/MU154/2011 | 25/09/2011        | Makanaga Bay/Mpigi | GGRQGRFL | LENTOGENIC |
| 15  | APMV-1/waterfowl/Uganda/MU159/2011 | 12/09/2011        | Musambwa Island/Rakai | GGRQGRFL | LENTOGENIC |
| 16  | APMV-1/waterfowl/Uganda/MU162/2011 | 12/09/2011        | Musambwa Island/Rakai | GGRQGRFL | LENTOGENIC |
| 17  | APMV-1/waterfowl/Uganda/MU165/2011 | 12/09/2011        | Musambwa Island/Rakai | GGRQGRFL | LENTOGENIC |
| 18  | APMV-1/waterfowl/Uganda/MU167/2011 | 12/09/2011        | Musambwa Island/Rakai | GGRQGRFL | LENTOGENIC |
| 19  | APMV-1/waterfowl/Uganda/MU170/2011 | 12/09/2011        | Musambwa Island/Rakai | GGRQGRFL | LENTOGENIC |
| 20  | APMV-1/waterfowl/Uganda/MU171/2011 | 12/09/2011        | Musambwa Island/Rakai | GGRQGRFL | LENTOGENIC |
| 21  | APMV-1/waterfowl/Uganda/MU172/2011 | 12/09/2011        | Musambwa Island/Rakai | GGRQGRFL | LENTOGENIC |
| 22  | APMV-1/waterfowl/Uganda/MU173/2011 | 12/09/2011        | Musambwa Island/Rakai | GGRQGRFL | LENTOGENIC |
| 23  | APMV-1/waterfowl/Uganda/MU181/2011 | 12/03/2012        | Nakiwogo/Entebbe | GGRQGRFL | LENTOGENIC |
| 24  | APMV-1/waterfowl/Uganda/MU186/2011 | 23/09/2011        | Samuka Island/ Jinja | GGRQGRFL | LENTOGENIC |

Note: Twenty-four representative APMV-1 isolates were sequenced. All the isolates showed the lentogenic amino acid motif GGRQGRFL111. APMV-1, Avian paramyxovirus type-1.
of both genotype I and II strains in wild birds (Miguel et al. 2013; Snoeck et al. 2009), with a possible involvement of inter- or intracontinental bird migration. Others have reported lentogenic APMV-1 in wild birds outside Africa (Banura et al. 2013; Krapez et al. 2010; Lindh et al. 2012; Stanislawek et al. 2000; Takakuwa et al. 1998). Some of these studies indicate

**FIGURE 2:** Phylogenetic analysis of partial F nucleic acid sequences of Avian paramyxovirus type-1. The tree was generated by a neighbour-joining algorithm, and alignments were bootstrapped 1000 times (only bootstraps > 50 are shown). The strain names were edited to include origin of isolates where needed. Genotypes are marked on the right. Representative isolates from the 18 recently described genotypes (according to the new proposed classification by Diel et al. (2012) are included and collapsed together, except for the genotype II where the aquatic birds isolates from this study belong.

NDV, Newcastle disease virus.
that wild birds may play a role as a potential source of virulent APMV-1 for poultry. Moreover, it is suggested that velogenic APMV-1 might arise from lentogenic APMV-1 in nature through point mutations in the F-protein cleavage site, making them virulent for poultry (de Leeuw et al. 2003; Takakuwa et al. 1998). Evolution of APMV-1 has continuously posed threats for emergence of new virulent strains and challenges for diagnosis of ND (Cattoli et al. 2010; Miller et al. 2010). Toyoda et al. (1989) suggested that different strains of APMV-1 evolve through various degrees of accumulation of point mutations rather than gene exchange by recombination. Such emerging virulent strains related to lentogenic strains (antigenically and genetically) have been suspected to have caused outbreaks in Ireland in 1990 (Toyoda et al. 1989; Alexander et al. 1992) and in Australia in 1998–2000 (Gould et al. 2001; Westbury 2001).

Phylogenetic analysis of APMV-1 isolates isolated in the current study showed that the partial F gene sequences clustered with those of genotype II viruses. This is consistent with previous reports of a predominance of genotype II viruses in wild birds (Hoque et al. 2012). The earlier revelation that the highly virulent strains could evolve from viruses of low virulence by mutation (Gould et al. 2001; Westbury 2001) underscores the significance of more detailed genomic studies to ascertain possible epidemiological linkages of strains circulating in wild birds and poultry. Although velogenic strains of APMV-1 have been isolated from wild birds suggesting an epidemiological link with strains in poultry (Huovilainen et al. 2001; Jorgensen et al. 2004; Liu et al. 2008; Snoeck et al. 2013; Zarkov et al. 2005; Zhu et al. 2010), the current study did not recover any virulent pathotype. Our recent studies on APMV-1 in domestic poultry revealed a separate genotype (V) (Byarugaba et al. 2014). This does not mean virulent strains may not occur in aquatic birds in the country and therefore more extensive molecular epidemiological and routine monitoring for APMV-1 in aquatic birds is important for early detection to prevent any possible spillover into domestic poultry. Such detailed genomic studies will elucidate the exact role of wild birds in the ecology and epidemiology of APMV-1 in poultry and inform control strategies.

Acknowledgements

The sample collection, virus isolation and sequencing of the isolates were supported by the World Bank Millennium Science Initiative Project (Grant # MSl/03/32/2010) to Denis K. Byarugaba through the Uganda National Council of Science and Technology. We thank Mathilde Paul and Agnès Waret-Szkuta (IHAP, Université de Toulouse, INRA, ENVT, Toulouse, France) for their help with the cartography.

Competing interests

The authors declare that they have no financial or personal relationships that may have inappropriately influenced them in writing this article.

Authors’ contributions

D.K.B., A.W., H.K. and J.L.N. designed the study. A.W., D.K.B. and M.F.D. analysed the data and drafted the manuscript. A.W., J.B.O. and K.K.M. collected the samples, isolated APMV-1 strains and identified the viruses. A.T. and M.F.D. carried out the molecular characterisation and sequence analysis of the isolates. All authors read and approved the final manuscript.

References

Aldous, E.W. & Alexander, D.J., 2001, ‘Detection and differentiation of Newcastle disease virus (avian paramyxovirus type 1)’, Avian Pathology 30(2), 117–128. https://doi.org/10.1080/03079450120044515

Aldous, E. W., Mynn, J.K., Banks, J. & Alexander, D.J., 2003, ‘A molecular epidemiological study of avian paramyxovirus type 1 (Newcastle disease virus) isolates by phylogenetic analysis of a partial nucleotide sequence of the fusion protein gene’, Avian Pathology 32(3), 239–256. https://doi.org/10.1080/03074501000059763

Alexander, D.J., 2003, ‘Newcastle disease, other paramyxoviruses and Pneumovirus infections’, in Y.M. Saif, H.J. Barnes, J.R. Glisson, A.M. Fadly, L.R. McDougall & D.E. Swaye (eds.), Diseases of poultry, 11th edn., pp. 63–99, Iowa State University Press, Ames.

Alexander, D.J., Campbell, G., Manvell, R.J., Collins, M.S., Parsons, G. & McNulty, M.S., 1992, ‘Characterization of an antigenically unusual virus responsible for two outbreaks of Newcastle disease in the Republic of Ireland in 1990’, Veterinary Record 130(4), 65–68. https://doi.org/10.1136/vr.130.4.65

Banura, B., Timur, G., Fethiye, Ç., Ilhami, K. & Atilla, D., 2013, ‘Determination of Newcastle disease virus among wild bird populations in Lake Van basin turkey’, Turkish Journal of Veterinary and Animal Sciences 37, 86–93.

Byarugaba, D.K., Mugomba, K.K., Omori I.B., Okitiwi M., Wanyana, A., Otim, M. et al., 2014, ‘High pathogenicity and low genetic evolution of avian paramyxovirus type 1 (Newcastle disease virus) isolated from live bird markets in Uganda’, Virology Journal 11, 173. https://doi.org/10.1186/1475-282X-11-173

Cattoli, G., Fusaro, A., Monne, I., Molia, S., Le Menach, A., Maregoya, B. et al., 2010, ‘Emergence of a new genetic lineage of Newcastle disease virus in West and Central Africa – implications for diagnosis and control’, Veterinary Microbiology 142, 168–176. https://doi.org/10.1016/j.vetmic.2009.09.063

Czeglédi, A., Ujvári, D., Somogyi, E., Wehmann, E., Werner, O. & Lomniczi, B., 2006, ‘Third genome size category of avian paramyxovirus serotype 1 (Newcastle disease virus) and evolutionary implications’, Virus Research 120(1–2), 36–48. https://doi.org/10.1016/j.virusres.2005.11.009

De Almeida, R.S., Hamburger, S., Goh, B., Briend, F.X., Molia, S., Gaidet, N. et al., 2013, ‘New avian paramyxovirus type I strains identified in Africa provide new outcomes for phylogeny reconstruction and genotype classification’, PLoS One 8(10):e76413. https://doi.org/10.1371/journal.pone.0076413

De Leeuw, O.S., Koch, G., Hartog, L. & Peeters, B.P., 2003, ‘Effect of fusion protein cleavage site mutations on virulence of Newcastle disease virus: Non-virulent cleavage site mutants revert to virulence after one passage in chicken brain’, Journal of General Virology 84, 475–484. https://doi.org/10.1099/vir.0.1878.0

Diez, P.G., De Silva, L.H., Liu, H., Wang, Z., Miller, P.J. & Afonso, C.L., 2012, ‘Genetic diversity of avian paramyxovirus type 1: Proposal for a unified nomenclature and classification system of Newcastle disease virus genotypes’, Infection, Genetics and Evolution 12(8), 1770–1779. https://doi.org/10.1016/j.meegid.2012.07.012

Gould, A.R., Kattenbelt, J.A., Selleck, P., Hansma, S., Gaidet, N. et al., 2004, ‘Avian paramyxovirus type 1 (Newcastle disease virus) and evolutionary implications’, Journal of Virology 77, 51–60. https://doi.org/10.1128/JVI.03167-1700.20040265-9

Hall, T.A., 1999, ‘BioEdit: A user-friendly biological sequence alignment editor and analysis for Windows 95/98/NT’, Nucleic Acids Symposium Series 41, 95–98.

Hoque, M.A., Burgos, G.W., Karo-Karo, D., Cheam, A.L. & Skerratt, L.F., 1992, ‘Characterization of an antigenically unusual virus responsible for two outbreaks of Newcastle disease in the Republic of Ireland in 1990’, Veterinary Record 130(4), 65–68. https://doi.org/10.1136/vr.130.4.65

Huovilainen, A., Komonen, E.K., Marvell, C. & Kinnunen, R., 2001, ‘Phylogenetic analysis of avian paramyxovirus 1 strains isolated in Finland’, Archives of Virology 146, 1775–1785. https://doi.org/10.1007/s0070501007063

Jorgensen, P.H., Handberg, K.J., Ahrens, P., Therkildsen, O.R., Manvell, R.J. & Alexander, D.J., 2004, ‘Strains of avian paramyxovirus type 1 of low pathogenicity for chickens isolated from poultry and wild birds in Denmark’, Preventive Veterinary Medicine 63(1), 49–62.

Kho, C.L., Mohd-Azmi, M.L., Arshad, S.S. & Yusoff, K., 2000, ‘Performance of an RT-nested PCR ELISA for detection of Newcastle disease virus’, Journal of Virological Methods 86(1), 71–83. https://doi.org/10.1016/S0166-0934(99)00185-8

Kim, L.M., King, D.J., Curry, P.E., Suarez, D.L., Swaye, D.E. et al., 2007, ‘Phylogenetic diversity among low virulence Newcastle disease viruses from waterfowl and shorebirds and comparison of genotype distributions to poultry-origin isolates’, Journal of Virology 81, 12641–12653. https://doi.org/10.1128/JVI.00843-07
Krapez, U., Steyer, A.F., Slavec, B., Barlic-Maganja, D., Dovc A., Racnik, J. et al., 2010, 'Molecular characterization of avian paramyxovirus type 1 (Newcastle disease) viruses isolated from pigeons between 2000 and 2008 in Slovenia', Avian Diseases 54(3), 1075–1080. https://doi.org/10.1637/9161-111709-ResNote.1

Lindh, E., Ek-Kommonen, C., Vaananen, V.M., Alasaari, J., Vaheri, A, Vapalahti, O. et al., 2012, 'Molecular epidemiology of outbreak-associated and wild waterfowl-derived Newcastle disease virus strains in Finland, including a novel class I genotype', Journal of Clinical Microbiology 50, 3664–3673. https://doi.org/10.1128/JCM.01427-12

Liu, H., Wang, Z., Wang, Y., Sun, Ch., Zheng, D. & Wu, Y., 2008, 'Characterization of Newcastle disease virus isolated from waterfowl in China', Avian Diseases 52, 150–155. https://doi.org/10.1637/8030-061507-Reg

Miguel, E., Grosbois, V., Berthouly-Salazar., C., Caron, A., Cappelle, J. & Roger, F., 2013, 'A meta-analysis of observational epidemiological studies of Newcastle disease in African agro-systems, 1980–2009', Epidemiology and Infection 141(8), 1117–1133. https://doi.org/10.1017/S0950268812002610

Miller, P.J., Decanini, E.L. & Afonso, C.L., 2010, 'Newcastle disease: Evolution of genotypes and the related diagnostic challenges', Infection, Genetics and Evolution 10(1), 26–35. https://doi.org/10.1016/j.meegid.2009.09.012

OIE, 2008, Manual of diagnostic tests and vaccines for terrestrial animal, 6th edn., World Organisation for Animal Health, Paris, France.

Otim, M.O., Christensen, H., Jørgensen, P.H., Handberg, K.J. & Bisgaard, M., 2004, ‘Molecular characterization and phylogenetic study of Newcastle disease virus isolates from recent outbreaks in eastern Uganda’, Journal of Clinical Microbiology 42(6), 2802–2805. https://doi.org/10.1128/JCM.42.6.2802-2805.2004

Shengqing, Y., Shinya, K., Otsuki, K., Ito, H. & Ito, T., 1997, ‘Isolation of myxoviruses from migratory waterfowls in San-in district, western Japan in winters of 1997–2000’, The Journal of Veterinary Medical Science 64, 1049–1052. https://doi.org/10.1292/jvms.64.1049

Snoeck, C.J., Ducatez, M.F., Owoade, A.A., Faleke, O.O., Alkali, B.R., Tahita, M.C. et al., 2009, 'Newcastle disease virus in West Africa: New virulent strains identified in noncommercial farms', Archives of Virology 154, 47–54. https://doi.org/10.1007/s00705-008-0269-5

Snoeck, C.J., Owoade, A.A., Couacy-Hymann, E., Alkali, B.R., Okwen, M.P., Adeyanju, A.T et al., 2013, 'High genetic diversity of Newcastle disease virus in poultry in West and Central Africa: Circorrelation of genotype and newly defined genotypes XVII and XVIII', Journal of Clinical Microbiology 51(7), 2250–2260. https://doi.org/10.1128/JCM.00684-13

Stanislawlek, W.L., Wilks, C.R., Meers, J., Horner, G.W., Alexander, D.J., Manvell, R.J. et al., 2000, ‘Avian paramyxoviruses and influenza viruses isolated from mallard ducks [Anasplatyrhynchos] in New Zealand’, Archives of Virology 147, 1287–1302. https://doi.org/10.1007/s00705-002-0818-2

Takakuwa, H., Ito, T., Takada, A., Okazaki, K. & Kida, H., 1998, ‘Potentially virulent Newcastle disease viruses are maintained in migratory waterfowl populations’, Japanese Journal of Veterinary Research 46(4), 207–215.

Tamura, K.P.D., Peterson, N., Stecher, G., Nei, M. & Kumar, S., 2011, ‘MEGAN: Molecular evolutionary genetics analysis using maximum likelihood, evolutionary distance, and maximum parsimony methods’, Molecular Biology and Evolution 28, 2731–2739. https://doi.org/10.1093/molbev/msr121

Toyoda, T., Sakaguchi, T., Hirota, H., Gotoh, B., Kuma, K., Miyata, T. et al., 1989, ‘Newcastle disease virus evolution. II. Lack of gene recombination in generating virulent and avirulent strains’, Virology 169, 273–282. https://doi.org/10.1016/0042-6822(89)90152-9

USDA/APHIS/WS, 2016, Avian Health, viewed 12 October 2016, from https://www.aphis.usda.gov/wcm/connect/aphis_content_library/sa_our_focus/sa_wildlife_damage/sa_programs/sa_nwrc/sa_nwdp/ct_avian_health

Westbury, H., 2001, ‘Newcastle disease virus: An evolving pathogen?’, Avian Pathology 30(1), 5–11. https://doi.org/10.1080/03079450020023131

Zarkov, I., Bochev, I., Oreshkova, N. & Manvell, R., 2005, ‘Isolation of avian paramyxovirus type 1 (APMV-1) from free-living mallards [Anasplatyrhynchos] (preliminary communication)’, Bulgarian Journal of Veterinary Medicine 8, 173–181.

Zhu, W., Dong, J., Xie, Z., Liu, Q. & Khan, M.I., 2010, ‘Phylogenetic and pathogenic analysis of Newcastle disease virus isolated from house sparrow (Passer domesticus) living around poultry farm in southern China’, Virus Genes 40(2), 231–235. https://doi.org/10.1007/s11262-009-0436-0