Visceral Marek’s disease in white-peafowl (*Pavo cristatus*)

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ABSTRACT

Marek’s disease (MD) is a lymphoproliferative disorder caused by *Gallid herpesvirus* 2 (MDV) that infects mainly domestic gallinaceous birds although wild birds may occasionally be affected. The current report describes the anatomopathological and molecular findings of a case of MD in a white-peafowl (*Pavo cristatus*). The signs included apathy, hyporexia, and diarrhea. Grossly, 0.5 to 1.5cm in diameter, yellow, soft nodules were observed in the skeletal muscle, lung, kidney, air sacs, small intestine, heart, ovary, ventriculus, and proventriculus. Microscopically, numerous atypical round neoplastic cells were noted. The molecular detection of MDV DNA was implemented to amplify part of the meq gene and products were sequenced for the phylogenetic analysis. Template DNA was obtained from tissues of the affected bird and from blood of all the gallinaceous birds of the Zoo. The expected amplicon for the partial amplification of MDV *meq* gene was obtained and the amplicons were sequenced. Sequences obtained enabled grouping the strain (accession no. KT768121) with MDV serotype 1 strains from the GenBank. Based on the anatomopathological and molecular findings, the diagnosis of MD in a white-peafowl was reached, and to the authors’ knowledge, no previous report regarding MD was published in *Pavo cristatus*.

Keywords: Marek’s disease, lymphoma, oncogenic virus, MDV, herpesvirus

RESUMO

Doença de Marek (MD) é uma desordem linfoproliferativa causada pelo Gallid herpesvirus 2 (MDV), que infecta principalmente galiáceos domésticos, porém aves silvestres podem ser ocasionalmente afetadas. O presente relato descreve os achados anatomopatológicos e moleculares de um caso de MD em um pavão-branco (*Pavo cristatus*). Os sinais clínicos incluíram apatia, hiporexia e diarreia. Grossamente, foram observados nódulos macios, de 0,5 a 1,5cm de diâmetro, no músculo esquelético, no pulmão, nos rins, nos sacos aéreos, no intestino delgado, no coração, no ovário, no ventrículo e no proventrículo. Microscopicamente, numerosas células redondas neoplásicas atípicas foram notadas. A detecção molecular do DNA do MDV foi implementada para amplificar parte do gene meq, e os produtos foram sequenciados para análise filogenética. DNA foi obtido de tecidos de aves afetadas e do sangue de todos os galiáceos do zoológico. A esperada amplificação de parte do gene meq de MDV amplificado foi ampliada e sequenciada. As sequências obtidas permitiram o agrupamento da cepa (acesso KT768121) com cepas do sorotipo 1 de MDV do GenBank. O diagnóstico de MD em pavão-branco foi obtido com base nos achados anatomopatológicos e moleculares e, pelo conhecimento dos autores, não há relatos anteriores publicados de MD em *Pavo cristatus*.

Palavras-chave: doença de Marek, linfoma, vírus oncogênico, MDV, herpesvírus

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INTRODUCTION

Marek’s disease (MD) is a lymphoproliferative disorder caused by Marek’s disease virus (MDV) serotype 1 of the Family Herpesviridae, subfamily Alphaherpesvirinae, genus Mardivirus and species Gallid herpesvirus 2 (Murata et al., 2012). MDV replicates in the feather follicle epithelium and natural desquamation (dander) is readily disseminated in premises and inhaled by susceptible birds of genus Gallus (Schat and Nair, 2008). The disease occurs in the domestic chicken and related birds of order Galliformes (Schat and Nair, 2008). However, little is known regarding the disease in wild species of other genera (Cho and Kenzy, 1975).

The transmission to migratory birds which share the environment with chickens is considered (Schat and Nair, 2008). In wild species, MD is more prevalent in captive birds, as compared to free living, although the actual prevalence is unknown (Murata et al., 2012). In peafowl, previous reports describe a lymphoproliferative disease, one case of reticuloendotheliosis (Miller et al., 1998) and two others without the determination of the etiology (Kaliner and Miringa, 1972; Sah et al., 1973). In San Diego Zoological Garden (California, USA), the virological and serological evaluation of 111 birds, of 49 species and 14 orders, resulted in the detection of MDV in ten birds, all of genus Gallus (Cho and Kenzy, 1975).

Clinical signs may vary with the MDV strain and the degree of effect in organs, but may be mostly unspecific, including apathy, weight loss, diarrhea, and dyspnea, although the type of paralysis of limbs would be indicative. Most birds would not develop clinical signs and function as lifelong reservoirs and virus source, expressing MDV in cycles (Payne and Venugopal, 2000). At necropsy, whitish soft metastases may be disseminated in viscera, mainly liver, spleen, thymus, heart, kidneys, gonads, lungs, proventriculus, mesentry, intestines, cloacal bursa, adrenal, pancreas, skeletal muscle, iris, and skin (Schat and Nair, 2008). Microscopically, tumors consist of pleomorphic T lymphocytes (Payne and Venugopal, 2000). Virus isolation, immunohistochemistry or DNA detection by PCR are required for the definitive diagnosis and to differentiate from other lymphoproliferative diseases such as avian leukosis, reticuloendotheliosis, and lymphoproliferative disease of turkeys (Payne and Venugopal, 2000; Schat and Nair, 2008). The aim of this article is to describe the clinical, pathological, and molecular findings of a natural MDV occurrence in white peafowl (Pavo cristatus).

CASUISTIC

An adult female peafowl of the Jardim Zoológico de Brasília (FJZB) (DF, Brazil) died after a month-long clinical disease which included apathy, hyporexia, and diarrhea, despite being medicated with enrofloxacin (Baytril®, Bayer S.A., São Paulo, Brazil), sulfamethoxazol plus trimethoprim (Trisulfin®, Ouro Fino, Cravinhos, Brazil), and metronidazole (Flagyl®, Sanofi-Aventis Farmacêutica Ltda, Suzano, Brazil). The bird was found dead in the premises and after a preliminary examination, all the gallinaceous (Galliformes; n=36) (Tab. 1) birds of the premises were examined, taking a blood sample for PCR. Blood samples (1mL) were collected at the ulnar or at the brachial vein in anticoagulant (EDTA).

Table 1. Species of order Galliformes evaluated by PCR for Marek’s disease virus

<table>
<thead>
<tr>
<th>Species</th>
<th>Number sampled</th>
</tr>
</thead>
<tbody>
<tr>
<td>Red-throated piping guan (Aburria cujubi)</td>
<td>1</td>
</tr>
<tr>
<td>Black-fronted piping guan (Aburria jucatoga)</td>
<td>4</td>
</tr>
<tr>
<td>Black curassow (Crax alector)</td>
<td>1</td>
</tr>
<tr>
<td>Red-billed curassow (Crax blumenbachii)</td>
<td>1</td>
</tr>
<tr>
<td>Bare-faced curassow (Crax fasciolata)</td>
<td>3</td>
</tr>
<tr>
<td>Razor-billed curassow (Pauxi tuberosum)</td>
<td>5</td>
</tr>
<tr>
<td>Nocturnal curassow (Nothocrax urutumutum)</td>
<td>2</td>
</tr>
<tr>
<td>Indian peafowl (Pavo cristatus)</td>
<td>8</td>
</tr>
<tr>
<td>Dusky-legged guan (Penelope obscura)</td>
<td>1</td>
</tr>
<tr>
<td>Rusty-margined guan (Penelope superciliiaris)</td>
<td>4</td>
</tr>
<tr>
<td>Chestnut-bellied guan (Penelope ochrogaster)</td>
<td>5</td>
</tr>
<tr>
<td>Silver pheasant (Lophura nycthemera)</td>
<td>2</td>
</tr>
</tbody>
</table>

The hemogram revealed non-regenerative anemia and leukopenia, with approximately 98% of lymphocytes reactive, piknotic and with a few in mitosis. At necropsy, poorly delimited, soft, 0.5 to 1.5cm in diameter yellow nodules were noted in the pectoral muscle (Fig. 1A), lungs, kidneys, air sacs, small intestine, heart (Fig. 1B), ovary, gizzard (ventriculus), and proventriculus.
All samples were fixed in buffered formaldehyde 10%, processed routinely for histopathology, and stained with hematoxylin-eosin (HE). Tissue samples from kidneys and breast muscle were also collected and stored frozen for PCR. The histopathology revealed densely cellular neoplasia without capsule and infiltrating into the affected organs. The neoplasma was composed of atypical lymphocytes arranged in layers maintained by a scarce fibrovascular stroma (Fig. 1C and 1D). Neoplastic cells presented scarce, with distinct limits, eosinophilic, and homogenous cytoplasm. The nuclei were central, rounded, single, basophilic with loose chromatin and with one to four separate nucleoli. Discrete anisocytosis and moderate anisokaryosis with a 4:1 nucleus-cytoplasm ratio were evident. Within ten high magnification (1000x) fields, 27 mitoses were observed.

DNA extractions were performed by silica adsorption. Three volumes of NaI (6M) were added to one volume of tissue or blood (approximately 200µl of tissue or blood and 600µl of NaI), macerated and incubated at 55°C/15min. in an end-over-end mixer. The mixture was centrifuged (2,000xg/30s), the supernatant discarded and the pellet washed in NaI, centrifuged (2,000xg/30s) and the pellet washed twice in washing buffer (ethanol 50% 50mM Tris-HCl pH 8.0, 10mM EDTA pH 8.0) by centrifugation (2,000xg/30s),
with the supernatant discarded. One ml of acetone was added to the pellet, vortexed, centrifuged (2,000xg/30s), the supernatant discarded and any residue evaporated (56°C/10min). The adsorbed DNA to silicon microbeads was eluted with 50µl TE (5mM Tris-HCl pH 8.0, 0.5mM EDTA pH 8.0), incubated at 50°C/5 min in the end-over-end mixer and centrifuged (2,000xg/30s). The supernatant was removed with pipette and stored (-20°C) until used as template for PCR. The meq-specific oligonucleotide primers used 5'-ATGTCCTCAAGAGCCAGGCGGCT-3', and 5'-GGGGCACTAGGCATGTCGCTAGG-3' amplify a sequence of the meq gene with a 1,062bp product. PCR was performed in 50µl volumes containing 1.5mM MgCl2, 0.1µM of each primer, 500ng of template DNA, 0.2mM of dNTP mix and 0.2 unit of the Taq DNA polymerase (Phoneutria, Belo Horizonte, Brazil). After initial 94°C/5min incubation, the 40-cycle PCR incubation protocol was run at 94°C/60s, 57°C/60s and 72°C/90s, with a final incubation at 72°C/7min.

The proviral DNA of avian leukosis virus was investigated using previously described primers for fowl glioma-inducing virus (FGV) (Hatai et al., 2005). A nested PCR for FGV with external primers EnvFwd3 (59-ACGTTCGCGGGAATGTGTTGT-39) and L7Krev (59-CTATACACACCTAGTCCC-39) and internal primers 5FGOG (59-TGTAGGCGGCTTTGTATT-39) and 3FGOG (59-TTGTCGGTCAAGCCTTGCCTT-39), was used to amplify a FGV non-coding region with a 135bp product. PCR products were purified for sequencing with polyethylene glycol (PEG). Single band PCR products, as determined by electrophoresis, were purified. Equal volumes of PEG 8000 20% and amplicon were stirred for 15s and incubated for 15min at 37°C. The mix was centrifuged at 12,000xg for 15min and the supernatant carefully removed with pipette and discarded. The sediment was washed twice with 100µl of 80% ethanol added slowly without disturbing the pellet and centrifuged for 5min at 12,000xg. The supernatant was discarded by inversion, the tube was incubated at 37°C for ethanol evaporation and 10-15µl of ultrapure water were added. The sediment was suspended by pipetting and the purified DNA fragment size was confirmed by electrophoresis on 1.5% agarose gel in 0.53 TBE and quantified (NanoVue, GE Healthcare, Little Chalfont, United Kingdom).

The purified product was sequenced by the dideoxynucleotide method using a sequencing kit (BigDyeH Terminator Mix, Applied Biosystems, Carlsbad, USA) according to the manufacturer instructions. One microliter of each purified product was used. The reaction was performed using 1µl of each primer (forward and reverse) at 10pmol, 1µl of BigDye, 1µl of SaveMoney™ buffer (Applied Biosystems) and ultrapure water 18.2Ω q.s.p. to 10µl. Reactions were performed in a thermocycler (PTC-100, MJ Research, Inc.) with denaturation at 96°C for 15 sec, annealing of primer at 50°C for 15 sec, and extension at 60°C for 4 min, repeated 30 times. The sequencing PCR product was purified by isopropanol and ethanol precipitation and homogenized in Hi-Di™ formamide (Invitrogen, São Paulo, Brazil) and rapidly denatured at 95°C / 2 min and immediately placed on ice. The analyses were made in an automated capillary electrophoresis system (ABI Prism 310 Genetctic Analyzer, Perkin Elmer, Waltham, USA) in the Laboratory of Genetics, Department of Animal Sciences (Escola de Veterinária UFMG, Belo Horizonte, Brazil). The generated sequences compositions were compared with sequences available in the database of the National Center for Biotechnology Information (NCBI, 2015). The Basic Local Alignment Search Tool (BLAST 2.0), BLASTn, and BLASTx (BLAST, 2015) software, developed by the NCBI, were used to determine similarity of nucleotide and amino acid compositions, respectively. The electropherograms of sequences of sense and antisense strands were analyzed using BioEdit.

All final sequences generated were compared with sequences available in the database of the NCBI. The nucleotide and amino acid sequences were aligned with sequences obtained in the GenBank using Clustal X software, in the Molecular Evolutionary Genetics Analysis MEGA version 5.0 for Windows. The confidence on the topology of the phylogenetic tree was insured by 1,000 bootstrap oversampling. The nucleotide and the amino acid substitutions were assessed using the Kimura 2 parameter or the JTT for amino acid substitution, respectively. The expected molecular mass product (1,062bp) was obtained as revealed by electrophoresis. No amplicon was obtained for avian leukosis virus.
Comparing the sequenced PCR products obtained from tumors with sequences on GenBank revealed identity with *Gallid herpesvirus 2* (MDV) (Fig. 2). However, MDV DNA was not detected in any of the other birds (n=36) examined. The phylogenetic analysis of the MDV is based on the *meq* gene partial DNA sequence, performed by the neighbor-joining method using MEGA version 5.0 software packages and bootstrap confidence limits for 1000 replicates. The isolate *Gallid herpesvirus 2 Marek’s disease virus type 1 Brasilia 1* was identified using the nomenclature of strain or isolate name referring to the geographic location. The partial sequence of *meq* gene was deposited in the GenBank with the Accession number KT768121.

![Figure 2](image)

**DISCUSSION**

The anatomopathological evaluation and mainly the molecular detection were conclusive for the diagnosis of MD in peafowl. No previous report of MD was found for the white peafowl. Some reports, however, describe lymphomatosis in peafowl (Kenzy *et al.*, 1964; Kaliner and Miringa, 1972; Sah *et al.*, 1973; Miller *et al.*, 1998). MD is usually diagnosed in domestic gallinaceous birds and descriptions in other species of birds are scarce.

The clinical signs, gross and microscopic lesions were compatible with the literature (Schat and Nair, 2008). MDV is replicated initially in
B-lymphocytes, then in the feather follicle epithelium and subsequently integrates genomically into T lymphocytes. Except for the lymphoproliferative disease, the atrophy of lymphoid organs and immunodepression (Payne and Venugopal, 2000), were not detected in the present study. Lymphoproliferative diseases must be differentiated microscopically at histopathology and the etiology may be determined through the genomic detection and characterization. Lymphoid leukosis presents homogenous lymphoblast proliferations which are distinguishable from the MD that usually has discrete anisocytosis and moderate anisokaryosis as found in the current study (Payne and Venugopal, 2000). MD is an infectious disease of worldwide occurrence, mostly in immature chickens (Schat and Nair, 2008). In the present report, the white peafowl was an adult bird. The reasons for differences between chicken and peafowl regarding the susceptibility to infection and to the behavior to the MD lymphoproliferative disease are not known.

All other gallinaceous birds captive at the FJZB were evaluated for MD and ALV genome and resulted negative. The source of MDV remains to be determined. Occasional local migratory birds visit the FJZB and the possibility of such birds previously visiting free-range chickens. However, the local geographical range migratory birds, such as the cattle egret (Bubulcus ibis) and the white-faced whistling duck (Dendrocygna viduata) are not considered a source of MDV. Free-range chickens may act as natural reservoir of MDV and a few individuals from flocks regularly succumb to MD in the rural areas of Minas Gerais, Brazil, (unpublished findings) and possibly other Brazilian states.

MD has been one of the most important viral diseases of the poultry industry since the prosperity of the industry has been enabled through its control by vaccination (Payne and Venugopal, 2000). However, although chickens are by far the most important host for MDV, quail, turkey, and pheasant are also susceptible species to infection and disease. In addition to chicken, gamefowl (Kenzy et al., 1964), native gallinaceous breeds (Grewal et al., 1977), and jungle fowl (Cho and Kenzy, 1975) are susceptible to MDV infection and tumor formation. Most other domestic avian species including duck, partridge, peafowl, pigeon, and sparrow (Baxendale, 1969; Kenzy and Cho, 1969) are considered refractory to MDV infection.

A novel GaHV-3 has been described in the mountain peacock pheasant (Polyplectron inopinatum), the Malayan peacock pheasant (Polyplectron malacense), and the Congo peafowl (Afropavo congensis), with hepatocyte necrosis and nuclear eosinophilic inclusions, which showed 92% identity to GaHV-3, Marek’s disease virus serotype 2, despite serotype 2 strains are considered not pathogenic in chickens. However, no description of MDV tumors was found in the literature for peacock (Seimon et al., 2012).

Based on the clinical, anatomopathological, and molecular results, the diagnosis of MD was confirmed in a peafowl of the zoological garden of Brasilia, Midwestern Brazil. The evaluation of all other gallinaceous birds resulted negative for MDV and the source of infection, with the possibility of a role for free-range chickens, which remains to be determined.

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