Outbreak of cutaneous form of avian poxvirus disease in previously pox-vaccinated commercial turkeys

Bruna C. Ferreira, Roselene Ecco*, Rodrigo M. Couto, Humberto E. Coelho, Daise A. Rossi, Marcelo E. Beletti and Paulo L. Silva

ABSTRACT - Ferreira B.C., Ecco R., Couto R.M., Coelho H.E., Rossi D.A., Beletti M.E. & Silva P.L.
E-mail: ecco@vet.ufmg.br

This study describes an outbreak of avian poxvirus disease in previously pox-vaccinated turkeys in Brazil. The turkeys had suggestive gross lesions of cutaneous avian poxvirus in the skin of the head and cervical area without changes in the flock mortality rates. In the slaughterhouse, 30 carcasses were removed from the slaughter line to collect tissue from cutaneous lesions for histological analyses and characterization of the virus. The virus was identified by conventional polymerase chain reaction (PCR) and subsequent gene sequencing. Acanthosis, hyperkeratosis, and hydropic degeneration were seen on skin histopathology. Eosinophilic intracytoplasmic inclusion bodies (Bollinger) on keratinocytes were observed in 46.6% of the samples. Avian poxvirus DNA was detected on PCR in 93.3% of the total samples. PCR associated with histopathology had 93.3% of positivity for avian poxvirus. In the phylogenetic study, samples show 100% matching suggesting that the outbreak occurred by a single viral strain and was different from those strains affecting other wild birds such as canaries and sparrows. A single mutation (Adenine for Guanine) was detected in our study’s strain and in the strains of turkey, chickens, and vaccine strains published in GenBank. When the sequence strain of the present study and sequences from GenBank of canarypox and sparrowpox strains were aligned, a Thymine was found replacing the Adenine or Guanine. The in ovo vaccination method as single-use in turkeys of this study apparently did not provide adequate protection against avianpox disease, but additional vaccination administered by wing-web when turkeys were 45-60 days old in the new flocks controlled the disease. In the subsequent year, new cases of this disease were not found. It was not possible to confirm the source of the virus strain, but infection with a field strain derived from chickens is one possibility, considering the poultry farm population in the area and biosecurity aspects. For wide characterization of avipoxvirus and differentiation among strains, the complete sequence of the viral genome is required.

INDEX TERMS: Avian poxvirus disease, avipoxvirus, fowlpox, pox-vaccinated, turkeys, pathology, fpv167 gene, sequencing, vaccine.
Although it is not a highly lethal disease and shows an atypical occurrence, avianpox disease is being frequently reported in avian wildlife in Brazil (Catroxo et al. 2009, Vargas et al. 2011, Pereira et al. 2014). In broilers, skin lesions caused by avian poxvirus were detected in carcasses of slaughtered birds from unvaccinated flocks (Fallavena et al. 2000). There are no recent studies using molecular characterization of poxvirus in Brazil, particularly in turkeys.

Avianpox reports are common in many countries and phylogenetic studies were described by Luschow et al. (2004), Jarmin et al. (2006), Manarolla et al. (2010), and Gyurancz et al. (2013) with samples from the United States, Italy, Germany, and other countries. Molecular research correlated with clinical manifestations allow the identification and comparison of genetic strains by consulting a database of strains identified by infected species, country of origin, year of isolation, isolated type (cutaneous form, diphtheric, or mixed) and form of isolation (cultivation in embryonated eggs).

Live attenuated vaccines have been used to prevent disease in susceptible birds. The strains come from fowlpox of chickens or pigeons (Tripathy & Reed 1997). However, the efficacy of these vaccines has been questioned worldwide, because in recent years, outbreaks have been reported in previously vaccinated turkeys (Odoya et al. 2006, Estrella-Tec et al. 2013).

An experimental study in vaccinated and unvaccinated turkeys for fowlpox revealed that vaccination did not influence the incidence of the disease but reduced the severity of clinical signs and mortality in young birds in the studied populations (Estrella-Tec et al. 2013).

Monitoring avianpox disease at slaughter was responsible for the increase in carcass condemnation related to this condition in turkeys from state of Minas Gerais, Brazil. In this study, we describe an avian outbreak of avianpox disease in previously vaccinated commercial turkeys.

MATERIALS AND METHODS

To obtain information related to the occurrence of avianpox disease in turkeys in the state of Minas Gerais, Brazil, data from official nosological maps constructed by the regional Federal Inspection Service were accessed. Turkeys (Meleagris gallopavo) from one farm only were examined and and samples from these turkeys were collected for histopathology and molecular study.

Sample collection. In June 2013, 115 day-old turkeys with an average of 15.44 kg, from a farm in Indianapolis, Minas Gerais, were slaughtered. We selected 30 carcasses from the slaughter line of this flock to collect skin fragments with suggestive lesions of cutaneous fowlpox. A small piece of skin was pinched and pulled out using a scalpel and sterile tweezers. These fragments were sectioned to contain areas of the skin lesion and normal skin margin. Each section was divided in two portions: one was fixed in 10% buffered formalin for histological evaluation and another portion was stored at -20°C until further molecular analysis.

Histopathological analysis. The fragments were fixed in 10% buffered formalin for 48h and then transferred to 70% ethanol. The tissues were dehydrated in increasing ethanol series, diaphanized in xylene, embedded in paraffin to obtain sections of 5 μm thickness, stained with hematoxylin and eosin, and evaluated using light microscopy. Histopathological diagnosis of avian fowlpox was based on the observation of typical lesions associated with intracytoplasmic eosinophilic inclusion bodies (Tripathy & Reed 2008). The inflammatory infiltration was graded according intensity in mild, moderate and marked.
Molecular analysis. For DNA extraction, the modified protocol of Vogelstein & Gillespie (1979) and Boom et al. (1990) was used.

Tissue samples were scraped using a 2.0 scalpel blade in a glass petri dish, and then added to a 1.5-mL microtube and macerated with a pestle. Thereafter, the homogenate was added to 300μL of sodium iodide (NaI). For DNA adsorption, 40μL of silica (silicon dioxide, Sigma-Aldrich, St. Louis, MO, USA) was used. Then, the mixture was washed thrice in 500μL of ethanol and finally rinsed with 1 mL acetone. Extracted DNA was eluted in 40 μL tris EDTA (TE) and then analyzed and quantified by atomic absorption at 260nm in a spectrophotometer (Nanodrop ND- 1000, Thermo Scientific, Wilmington, USA).

PCR for amplification of the gene fragment fpv167 (P4b locus) was performed using the following specific primers: 5’-CAGCAGGTGCTAAACAACAA-3’ and 5’-CGGTAGCTTAAGCGGAATA-3’ (Lawson et al. 2012); the product size was 578bp. Each reaction was performed in a 25μL volume (PCR Master mix Promega) containing 200ng of DNA sample and 0.02nmol of each primer. Amplification was performed in a thermocycler under the following conditions: Initial denaturation at 94°C for 5min followed by 45 cycles at 94°C for 1min, 60°C for 1 min and 72°C for 1min, and a final extension at 72°C for 7min. The amplified PCR products were separated by electrophoresis on 1.5% agarose gel and visualized after staining with ethidium bromide under UV transillumination.

The amplified DNA products were extracted and purified from the agarose using the Invisorb® DNA Extraction Spin Kit, (B-Bridge, Santa Clara, CA, USA) according to manufacturer’s recommendations. Each sample was bidirectionally sequenced in triplicate as per Sanger’s method with capillary electrophoresis (ABI Taq DyeDeoxy Terminator Cycle Sequencing Kit version 3.1) on an automatic sequencer (ABI 3130 Genetic Analyzer).

For sequencing, four samples (P1, P6, P10, and P26) that yielded a single stronger band of correct size in the electrophoresis were selected. The sequencing reactions were performed with an initial denaturation step at 96°C for 1 min; 30 cycles at 96°C for 15 s, 50°C for 15 s, and 60°C for 4 min; ending in a final step at 8°C indefinitely (SeqDNA Program). Next, this reaction product was precipitated with 40μL of 65% isopropanol, incubated at room temperature in the dark for 30 min, and centrifuged at 14,000 rpm for 45min. The supernatant was removed, added to 500μL of 60% ethanol (Merck KGaA, Darmstadt, Germany), centrifuged at 14,000rpm for 10min, and the precipitate was dried in an oven at 95°C for 5min.

We analyzed the obtained sequences and created consensus sequences by using the SeqScap® software program (version 2.5). Mega program (version 5.0) was used to align sequences with the GenBank database (NCBI) using the BLASTn platform. Phylogenetic trees were generated by the neighbor-joining method with bootstrap 1000, Kimura two parameters.

The published sequences in Genbank used for comparison with the strain isolated from turkeys with avian fowlpox are shown in the results part.

RESULTS

The monitoring for avianpox disease occurrence started in January 2013 and continued for a year. In 2013, 16,830 turkeys were identified with suggestive lesions of avianpox disease at the slaughter. Turkeys from several farms on the region presented avianpox lesions, despite in ovo vaccination with live fowl pox virus. The monthly percentages of turkeys slaughtered with avianpox lesions ranged from 1.0% to 40.38%, with high occurrence especially on the summer.

The examined turkeys presented lesions characteristic of the cutaneous form of fowlpox. Crusted and nodular coalescing lesions were found disseminated in the skin of the head (Fig.1A and B) and cranial cervical region. No additional clinical signs or increasing mortality in the flock were recorded by the owner.

Histopathology of all sampled skin lesions showed hyperkeratosis, hyperplasia with hydropic degeneration of keratinocytes, and desquamation (Fig.2A). Intracytoplasmic eosinophilic inclusion bodies (Bollinger) in keratinocytes (Fig.2B) were found in 46.6% of samples. Skin lesions with predominance of necrosis (ulceration) of the entire epithelial layer with replacement by necrotic and keratin debris were observed in samples where inclusion bodies were not found.

In the dermis, there was multifocal to coalescing inflammatory infiltration of lymphocytes, forming aggregates (Fig.3). The classification of these inflammatory infiltrates was performed in three degrees considering the intensity...
and distribution on each skin section with lesions. In 56.6% of the samples, there was diffuse inflammatory infiltrate, 30% multifocal inflammatory infiltrate, 10% focal inflammatory infiltrate, and 3.3% did not present inflammatory infiltrates (Table 1). In the skin of some birds, there were also erosions and ulcerations associated with heterophilic infiltration and bacterial colonies.

Histopathological examination associated with PCR allowed the confirmation of the disease caused by avian fowlpox in 93.3% of the samples analyzed in this study. Histopathology alone allows establishment of a definitive diagnosis for the disease in 46.6% of cases. Avipoxvirus was detected by PCR in 83% (25/30) of the samples. Negative results for avipoxvirus by PCR were detected in 17% (5/30) of the samples.

Sequencing of samples P1, P6, P10, and P26 showed that the region amplified was identical to the complete virus genome (complete fowlpox 4b). The Turkeypox sequences, Turkeypox GB134/01, Turkeypox PA213/07, fowlpox IPVD/LSA/2012/02, fowlpox IPVD/LSA/2012/03, fowlpox IPVD/LSA/2012/04 (subtype turkeys), Cow Pox Chickenpox Mild Fort Dodge, Cow Pox Chickenpox Diftosec Merial, fowlpox Nobilis variol MDS (vaccines), fowlpox Hungary, Chickenpox CVL 174/4/04 and fowlpox HP (subtype chickens) also had 100% identity with the sequence obtained (Fig. 4). These strains belong to the subgroup A. This subgroup identification allows us to infer that the turkey samples from this study, which are separate from subgroup B that include pigeon, turkey and ostrich strains and subgroup C, which include canary and sparrow strains, are not related to canary strains or strains found in other wild birds.

All samples evaluated in this study had Adenine (A) to Guanine (G) mutation at nucleotide 549 (Table 2).

### Table 1. Samples with presence (P)/absence (A) of intracytoplasmic eosinophilic inclusion bodies, classification of lymphocytic and plasma cells infiltration and PCR results

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IB = Intracytoplasmic eosinophilic inclusion bodies, PCR = polymerase chain reaction, P = positive, N = negative; - absent, + mild, ++ moderate, +++ marked.
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Table 2. Nucleotide sequence of positions 544 to 552 in the fpv167 gene of strains from Minas Gerais and other published avipoxvirus strains

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Fig.4. Phylogenetic tree generated from the concatenated sequence grouped and aligned to P4b nucleotide fragment of the fpv167 gene divided into three subgroups. Method: Neighbor-joining analysis with 1000 bootstrap replicates by using the Kimura 2-parameter model. (A) Chicken strain, (B) Pigeon strain, (C) Canary strain.
DISCUSSION

Histopathological lesions were consistent with the cutaneous form of avianpox in turkeys, confirming the clinical and macroscopic findings. Molecular analysis by PCR and sequencing confirmed the diagnosis of infection with avipoxvirus.

The result of monitoring carried out in the region in conjunction with the Federal Inspection Service, showed that the disease occurred frequently in 2013. Nevertheless, most cases occurred in the summer (data not shown). The prophylactic vaccination adopted in the region apparently did not prevent new cases of the disease in these birds during the monitoring year. Avianpox disease is widely distributed and has cyclical occurrence in endemic areas, especially in high-density areas of poultry production (Tripathy & Reed 2008, Bernardino 2009) such as the outbreak area.

Some factors may be relevant in identifying the source of an outbreak. Scratches produced by the claws of other infected birds in the flock, presence of blood-sucking insects at the time of the outbreak (summer), and the possibility of virulence reversion of vaccine strain or type of vaccine strain should be considered (Fallavena et al. 1993, Tripathy & Reed 2008, Bernardino 2009). The presence of mosquitoes is an important factor in the virus introduction in the region. In addition to the 11 species cited by Akey et al. (1981), Zylberberg et al. (2013) reported that Aedes aegypti and Culex quinquefasciatus, frequently found in Brazil, are also possible vectors of transmission of avipoxvirus. Singh et al. (2000) and Ramos et al. (2002) theorized a hypothesis about fowlpox outbreaks in previously vaccinated flocks concerning the emergence of strains containing reticuloendotheliosis virus genes integrated in the genome, which could possibly increase the virulence of these strains.

Usually the fowlpox is a self-limiting disease and has milder manifestations, as reported by Silva et al. (2009) in Brazilian free-range chickens. In the present case, a challenger viral strain might have disseminated to turkey flocks of the Triângulo Mineiro region. Bloodsucking insects and/or insufficient biosecurity measures and immunization (Tripathy & Reed 2008) were possibly involved in the introduction and dissemination of the virus to these turkey flocks.

The main gross lesions in this episode were nodular crusted lesions mainly found in the skin of head and neck, including caruncle, snood, wattles, eyelids, and beak commissure, consistent with previous avian poxvirus cutaneous disease reported in turkeys (Bernardino 2009, Hess et al. 2011). The clinical and gross presumptive diagnoses were confirmed by histopathological examination of lesions characterized by proliferative dermatitis containing eosinophilic intracytoplasmic inclusion bodies inside keratinocytes. These findings are consistent with the descriptions of Tripathy & Reed (2008) and Hess et al. (2011). The eosinophilic intracytoplasmic inclusion bodies are pathognomonic for avianpox disease and could be present at 72 h post-infection in the epithelial tissue (Arhelger et al. 1962). Because of the absence of the inclusion bodies, the remaining 53.4% samples were diagnosed based on suggestive lesions of avianpox disease in association with PCR results. The absence of the characteristic findings such as intracytoplasmic inclusion bodies may be related to chronic injuries and extensive necrosis of the epidermis, aggravated by secondary bacterial infection.

The use of specific techniques such as PCR combined with histopathology allows a definitive diagnosis of various diseases. In this study, three samples (3/5) with PCR results negative for avipoxvirus had intracytoplasmic eosinophilic inclusion bodies, which is a lesion characteristic of the disease (Tripathy & Reed 2008). Two remaining samples were negative in both histopathology and PCR tests. Probably, the sample collected had DNA virus lower than the detection threshold. Very small and extensively ulcerated samples with primary viral lesions replaced by secondary bacterial and cells debris could produce negative results (Tripathy & Reed 2008, Parker et al. 2011). These results highlight the importance of tissue selection at moment of sampling to obtain reliable results.

In this study, considering the sequence of the fpv167 gene fragment, four samples showed 100% identity to each other suggesting that the outbreak occurred by a single virus strain. Phylogenetic studies by Jarmin et al. (2006) and Manarolla et al.
(2010) found that the vast majority of avipoxvirus strains are grouped into three major subtypes, represented by chicken, psittacines, and canary strains.

The virus strain of this study was positioned near the turkeys’ field strains identified in Brazil, called fowlpox IPVDF/LSA/2012/01, fowlpox IPVDF/LSA/2012/02, fowlpox IPVDF/LSA/2012/03, and fowlpox IPVDF/LSA/2012/04; all these were also grouped near the 4b gene fragment of the complete genome of the chicken viruses, indicating 100% identity. The relationship with field or vaccine strains using the gene fragment sequenced was not possible because the fragment analyzed was short and similar for both strains.

The alignment of sequences of avipoxvirus gene in our study demonstrated that the virus origin is not from wild birds or passerines. The phylogenetic characterization established that the fragment /pv167 of the gene used was not sufficient for differentiating the strain in this study from field and vaccine strains from chickens and turkeys. Nevertheless, a single mutation (Adenine for Guanine) was detected in the present strain and in the strains of turkeys, chickens, and vaccine strains submitted to GenBank database. More studies are required to ascertain whether this mutation could influence some aspects of strain virulence and/or whether they belong to specific-host strains. Also, when the sequence strain of the present study and sequences from GenBank of canarypox and sparrowpox strains were aligned, a Thymine was found replacing the Adenine or Guanine. Although day-old turkey poults had been vaccinated with fowlpox vaccine at the hatchery (Sarma et al. 2015), the in ovo vaccination method as single-use in turkeys of this study apparently did not provide adequate protection against avianpox disease. According Tripathy & Reed (2008), avianpox vaccination in turkeys should be performed intradermally about midway on the thigh when they are 2-3 months old. The wing-web method may create dry cutaneous pox lesions on the turkey’s head (Tripathy & Reed 2008). In the subsequent year to this outbreak, new episodes of this disease were not recorded. According to the veterinarian, improvement of the biosecurity measures and additional fowlpox vaccine administered by wing-web method when turkeys were 45-60 days old was adopted in an attempt to control the disease. Despite recommendations aforementioned for vaccine injection site, no avianpox lesions were reported again. The in ovo vaccination was routinely adopted in the region because the company obtained an automated egg injection machine for injecting vaccine into a plurality of eggs. Nevertheless, in farms or regions with high viral challenge, a second vaccine dose apparently is necessary to protect turkeys against avianpox disease. It was not possible to confirm the source of the virus strain, but infection with a field strain derived from chickens is one possibility, considering the farm poultry population in the area and biosecurity aspects. Nevertheless, the possibility of reversion to virulence of the live attenuated poxvirus strain cannot be excluded and complete sequence of the viral genome is required for complete characterization of avipoxvirus and differentiation among strains.

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