Viral infections in goose flocks in Poland

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Abstract

The aim of this study was to determine the infectious agents isolated from infection – suspected geese sent for the diagnostic examination to National Veterinary Research Institute. The birds were sent from goose flocks localized in different parts of Poland. Totally, 1,013 birds from 122 flocks were examined. The presence of goose parvovirus (GPV), goose haemorrhagic polyomavirus (GHPV), and goose circovirus (GoCV) was detected by triplex PCR. The presence of GPV DNA was shown in 36 flocks. The disease was most frequently diagnosed in goslings aging 3.5 weeks (ten flocks), and 2.5 weeks (six flocks). The analysis of the nucleotide sequence of VP1 encoding region has shown close similarity of Polish GPV strains within the group which ranged from 92% to 100%. Moreover, the similarity level of these strains with GPV isolated in Europe was from 91.3% to 100%. The occurrence of GoCV DNA was shown in 25 goose flocks. The presence of GoCV DNA was found among geese aged from 2 to 6 weeks, but predominantly in those aging 3.5 (three flocks) and 5 weeks (five flocks). The sequence analysis of PCR products from the sequenced region of ORFC1 capsid protein of GoCV has shown that Polish isolates share from 85% to 91% similarity with the sequences of GoCV strains isolated in other countries. The presence of DNA of GHPV was found in 3-week-old geese. During the last 2 years the presence of GHPV was confirmed in three flocks of goslings at the age from 3 to 3.5 weeks. During the last 12 years the occurrence of coinfection with GPV and GoCV was detected in six flocks aging from 5 to 6 weeks.

Key words: geese, parvovirus, circovirus, polyomavirus, infection

Introduction

Infectious diseases of viral aetiology present an epidemic threat in goose production. Derzsy's disease is the most important viral disease of geese but infections with polyoma- and circoviruses are frequent as well.

Derzsy's disease (DD) is caused by Derzsy's disease virus (DDV) also called goose parvovirus (GPV). The birds susceptible to the infection are domestic and wild geese as well as Mullard and Barbarie ducks. The course of the infection and specific clinical symptoms depend on the age of birds, their immune status, and breeding conditions (Brown et al. 1995).

In spite of protective vaccination that limited the occurrence of Derzsy's disease, it is still a cause of considerable losses in goose flocks. The morbidity due to DD reach up to 80% of birds in a flock while the mortality is frequently about 60%.

The first cases of the disease were reported in Poland in the 1970s. Due to this fact, protective vaccinations of breeder flocks were introduced in 1982, what remarkably limited the occurrence of the disease. However, in the middle of the 1990s, the number
of clinical cases of DD increased again. Actually, the disease progresses as a chronic form with lower mortality than before. In Poland the disease is responsible for 16.5% to 19% of overall goose mortality. These data indicate the importance of this problem. Another important infectious disease of geese is syndrome of hemorrhagic nephritis and enteritis (HNEG) caused by goose haemorrhagic polyomavirus (GHPV). The disease is frequent in goslings of the age from 4th to 10th weeks (Guerin et al. 2000). The course of HNEG is acute, the mortality is high and the observed lesions in affected birds may give false evidence of DD. The third factor of viral infections of waterfowl is represented by goose circovirus (GoCV). These infections are frequent in goslings older than 5 weeks. The clinical symptoms are non-specific and include weight losses and feathering disorders, which are also observed in case of DD. The main role of this virus is associated with immunosuppression (Todd 2000).

Therefore, in case of mortality occurring in goose flocks, these three viral diseases of geese should be taken into account during differential diagnosis.

The aim of this study was to determine the infectious agents isolated from geese sent for the diagnostic examination to National Veterinary Research Institute during the years 1998-2010.

Materials and Methods

**Birds.** The infection-suspected birds were sent to the laboratory during 1998-2010 from goose farms localised in different parts of Poland. Totally, 1,013 birds from 122 flocks were examined. The number of birds in the flocks ranged from 1,200 to 12,000, most frequently from 3,000 to 5,000 birds. The age of the examined geese ranged from 1 to 9 weeks, but predominantly between 2 and 5 weeks. The clinical symptoms and pathological lesions suggested the occurrence of Derzsy’s disease.

**Standard viruses.** The vaccine strain MFP from Palmivax vaccine (Mérial, France) was used as the standard strain of GPV. The GoCV standard strain and GHPV strain were obtained thanks to courtesy of Dr Vilmos Palya (Ceva-Phylaxia, Ceva Sainte Ani male, Budapest, Hungary).

**DNA extraction.** DNA from standard strains was extracted from 200 μl of viral stock according to recommended procedure. Sections of the liver, heart, and kidneys were collected from the examined birds during post-mortem examination. Next, the sections were homogenised and total DNA was extracted according the procedure recommended by the manufacturer of QIAmp DNA Mini Kit (Qiagen, Germany).

**PCR primers.** Three pairs of specific primers were used. The sequences of primers used for GPV detection were: VPD1 – 5’ CCG GGT TGC AGG AGG TAC 3’; VPD1 – 5’ AGC TAC AAC AAC CAC ATC 3’. The expected PCR product size was 806 bp. For GoCV the primer sequences were: GCVP1 – 5’ TAA ATG CGT TGA TGT GTC T 3’; GCVP2 – 5’ CAT TTA ACC CCT TCC AAA GAG T 3’, and the expected product size was 571 bp, while for GHPV the sequences were as follows: GPOL1 – 5’ GAG GTT GGT GGA GTG ACC ACA ATG 3’; GPOL2 – 5’ ACA ACG GTG CAA GGG TTC 3’ with the expected product about 180 bp long.

**Triplex PCR assay.** Amplification was carried out in 25 μl of reaction mixture that contained: 2.5 μl of 10X concentrated PCR buffer (EurX, Poland), 1 μl of dNTP mixture (0.2 mM of each dNTP), 1 μl of MgSO4 (5 mM), 2 μl of each primer (VPD, GCV, GPOL) (10 mM), 2 μl of template DNA, 0.5 μl of Taq polymerase (5 U/ml), and PCR-grade deionised water. The reaction conditions were as follows: 30 cycles: 94°C/30 s (initial denaturation), 52°C/30 s (primers annealing), 72°C/1 minutes (elongation of products), 72°C/10 minutes (final elongation of products). Triplex PCR products were separated in 2% agarose gel with addition of ethidium bromide (0.5 μg/ml). To each well in gel, 5 μl of triplex PCR products and 1 μl of loading buffer (Fermentas, Lithuania) were loaded.

**Electrophoresis of PCR products.** The electrode buffer used for electrophoresis was TBE (Tris – Base 10.8 g, boric acid 5.5 g, 4 ml 0.5 M EDTA, pH 8.0). Electrophoresis was carried out for 1 h under the voltage of 10V/cm agarose gel. The results were documented using UV-light transiluminator (Vilber-Lourmat, France). Each product was compared with the molecular mass marker of 80-10 000 bp (Mass ruller DNA ladder 100 bp plus, Fermentas, Lithuania).

**Sequence analysis.** After gel electrophoresis the PCR products were cut from the gel and extracted from agarose slices using Gel Extraction Kit (Qiagen, Germany). The purified products were sequenced using the primers used for PCR by the Institute of Biochemistry and Biophysics of the Polish Academy of Science in Warsaw, Poland. The raw sequences were manually edited and compared with sequences accessible in the NCBI GeneBank database using BLAST algorithm.

**Results.**

During the twelve years (1998-2010) totally 1,013 birds from 122 flocks were examined. According to interviews and our observations, the clinical symptoms
occurred predominantly in the 1st and 2nd week of life of the birds and were manifested by a decrease in consumption of water and food, locomotor dysfunction, gathering, inhibition of growth, and feathering disorders. Meanwhile, in some of birds breathing difficulties, watery diarrhoea and neural disorders visible as paralysis of legs and neck were observed. The morbidity of the birds in the flock ranged from 30% to 60%.

The analysis of mortality of the birds in the examined flocks has shown that the mean percentage of dead birds was 13.95% to 42.45% and the range was from 9.9% to 61.7%. The highest mortality of birds was noted in 1999 (42.45%) and the lowest in 2003 (13.95%). Our personal observations have shown that after introduction to the market of the inactivated bivalent vaccine containing goose parvovirus and Muscovy duck parvovirus, the proportion of dead birds affected by Derzsy’s disease has considerably decreased. Meanwhile, the mortality observed in 2008 was 17.40% while in 2010 was only 5.70% (Fig. 1).

The clinical symptoms in geese were predominantly observed as weight losses (33%). Other clinical symptoms and anatomopathological lesions of the minor significance are presented in the Fig. 2. The most frequently occurring lesions were observed in the liver and they included congestion (41%) or paleness (16%), enlargement of the organ (22%) and fibrinous perihepatitis (19%). Additionally, the occurrence of small necrotic focuses was found in 2% of birds.

The lesions were also observed in other visceral organs. Rounded heart apex was found in 46% of the examined birds, 7% of the birds showed fibrinous pericarditis, while in 14% of the geese congestion of the intestines was found. Interestingly, both fibrinous pericarditis and congestion of the intestines were observed in birds at the age over 3.5 week.

The study conducted by triplex PCR has shown the presence of GPV DNA in 36 examined flocks (Fig. 3). The number of GPV-positive flocks was constant during the studied period and concerned three – four flocks per year. These flocks originated from areas with the highest density of goose breeding, predominantly from the west and east part of Poland. The disease was most frequently diagnosed in goslings at the age of 3.5 weeks (ten flocks) and 2.5 weeks (six flocks) (Fig. 4). The analysis of the nucleotide sequence of VP1 encoding region has shown a close similarity of isolated Polish GPV strains which ranged from 92% to 100%. Moreover, the similarity level of these strains with GPV strains isolated in Europe was from 91.3% to 100%. These data suggest that Polish GPV strains have the common European origin.

The study on the occurrence of GoCV has shown the presence of the virus in 25 goose flocks. The first case was diagnosed in 2002 and the positive flocks were localised in the west and east part of Poland similarly to GPV localisation. The presence of GoCV DNA was found among geese at the age from 2 to 6 weeks, but predominantly 3.5 (three flocks) and 5 weeks (five flocks) of age (Fig. 5). The sequence analysis of PCR products from the sequenced region of ORFC1 capsid protein of GoCV has shown that Polish isolates share from 85% to 91% similarity with the sequences of GoCV strains isolated in other countries.

During the third stage of this study, the presence of DNA of GHPV was found for the first time in 2007 in 3-week-old geese. During 1998–2010 the presence of GHPV was confirmed in three flocks of goslings aging from 3 to 3.5 weeks. The sequencing of PCR products of GHPV was not conducted in this study; however, so far the occurrence and analysis of GHPV was done in Germany and Hungary, what indicates its importance in epidemiology of viral infection of geese.

The applied triplex PCR allowed for easy DNA detection of three waterfowl viruses. This technique
is useful in differential diagnosis of viral infection of geese, which are frequently manifested by similar clinical symptoms and pathological lesions.

This aspect has a major importance when taking into account that during the last 12 years, in six examined flocks with geese aging from 5 to 6 weeks, the occurrence of co-infection with GPV and GoCV was detected (Fig. 6). This may suggest the possible role of these viruses in immunosuppression of the birds, which needs to be studied more comprehensively.
Discussion

The present results are the first analysis of viral infection in flocks of geese in Poland.

The most important viral infection is the goose parvovirus infection. Despite the prophylactic vaccination it is still the cause of deaths of young birds. As reported here, parvovirus infection were found in 36 goose flocks. Goose parvovirus was isolated in many countries of Europe and Asia. In Japan most GPV was isolated in geese at about 2 weeks, and mortality among birds was up 80% – 100%. GPV infection has also been reported in England and Wales and clinical symptoms and pathological changes were identical to the changes described in geese in Poland. In Sweden mortality of birds due to infection with GPV was from 40% to 90% (Takehara et al. 1995, Jansson et al. 2007, Irvine et al. 2008).

The goose circovirus infection was detected in 25 examined goose flocks. The results of phylogenetic analysis indicated the similarity of Polish isolates with GoCV strains isolated worldwide, but also suggest the variable nature of circoviruses which will be studies in the future (Chen et al. 2003, Ball et al. 2004, Yu et al. 2007).

The goose polyomavirus was detected in 3 examined goose flocks. The sequencing of PCR products of GHPV was not conducted in this study, however, so far the occurrence and analysis of GHPV was done in Germany and Hungary, what indicates its importance in epidemiology of viral infection of geese (Miksch et al. 2002, Palya et al. 2004).

The obtained results pointed out on the important role of viral disease of geese, which are still an epi-

zootic problem. As we presented in our study, in spite of annually applied prophylaxis of GPV infection, the coexistent infectious with GoCV and GHPV make this problem more complexive and indirectly influence the health status of geese.

References


