Naturally Occurring Egg Drop Syndrome Infection in Turkeys

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Abstract


A decrease in the egg quality, production, fertility and hatchability without serious clinical signs of illness was recorded in turkey flocks in Croatia at the beginning of 2002. It was assumed that the egg drop syndrome virus might be one of the etiological agents responsible for the abnormalities in the egg production. The systematic serological monitoring, using a haemagglutination inhibition test, showed that the antibodies to the egg drop syndrome virus existed in 94.4 and 55.1% of the sera analysed in 2002 and 2003, respectively. The haemagglutination inhibition titres ranged from 16 to 128. The sera samples were randomly collected from 11 - to 46-week-old hens from the affected flocks. The serological evidence of the egg drop syndrome virus infection was confirmed by detection of the presence of the virus genome in the turkey sera by the polymerase chain reaction. Vaccination of the 18- and 25-week-old turkey hens against the egg drop syndrome virus started in March 2003. After this period, the presence of antibodies to the egg drop syndrome virus (the haemagglutination inhibition titres between 16 and 256) was found in 96.7% of the analysed sera, while the egg production reached normal or higher values for the Nicholas hybrid line of turkeys.

Croatia, egg drop syndrome virus, haemagglutination inhibition, Meleagris gallopavo

Egg drop syndrome (EDS) affecting the flocks of laying fowl was described for the first time in 1976 (Van Eck et al. 1976). The EDS virus (EDSV), an etiological agent of the disease, designated as a duck adenovirus type 1, belonging to Atadenovirus genus (Harrach et al. 1997; Hess et al. 1997; Đan et al. 1998; Benkő et al. 2005), was isolated from the chickens during the 1980s and 1990s worldwide (McFerran and Adair 2003). The EDSV or the antibodies against the virus have been detected not only in hens but also in wild birds (Malkinson and Weisman 1980), wild waterfowl (Schlör 1980; Gulka et al. 1984), and pigeons (Durojaiye et al. 1992). In spite of the fact that the disease outbreaks were recorded only in laying hens, it has been demonstrated that ducks and geese were natural EDSV hosts (Schlör 1980; Zsak et al. 1982; Bartha and Mézáros 1984; Brugh et al. 1984). The receptivity to the infection and its transmission by contact were observed in pheasants, guinea fowls and quails (Zanella et al. 1980). The EDS outbreaks observed in the quail flocks, maintained together with infected chickens, resulted in the fall of the egg production, the increase of the number of the soft-shelled eggs, as well as the development of the HI-antibodies to virus (Das and Pradhan 1992). The involvement of the EDSV in a severe respiratory disease of the young goslings was also reported (Ivanics et al. 2001).

Optimal growth of the EDSV has been observed in the duck cells, but it was poor in the turkey cells. However, the experimental infection confirmed that turkeys can be infected by a direct contact, the eye-drop and the combined intranasal and oral routes (Parsons et al. 1980; Zanella et al. 1980; Kaleta et al. 2003), without any clinical signs of the infection. The virus could be detected in cloacal swabs during the period between 3 and 10 days after...
the infection. A rapid humoral response has been generated one week post infection and the detectable antibody titres were found 28 weeks later. In spite of this fact, no data concerning natural infection of the turkey flocks with the EDSV have been reported as yet.

A significant decrease in both egg quality and production was recorded in several turkey flocks in Croatia at the beginning of 2002. The affected flocks were known to have avian pneumovirus (APV) infection, but the abnormalities in the egg production were not recorded prior to the year 2002. At the same time, the turkeys were negative to the avian influenza virus (AIV), *Mycoplasma gallisepticum* (MG), and *Mycoplasma synoviae* (MS), while the Newcastle disease virus (NDV) infection was controlled by a systematic vaccination. These facts suggested that the decline in egg production might be caused by some other agent, possibly a naturally-occurring EDSV infection. For this reason, a serological and histopathological monitoring, as well as polymerase chain reaction (PCR) analysis of the sera samples collected from the birds suspected to be infected with the EDSV, were carried out. A differential diagnosis including the APV, NDV, AIV, MS and MG, was performed.

**Materials and Methods**

**Turkeys**

Nicholas hybrid line breeders were placed into production units with about 1100 birds each. During the whole period they were housed on litter in a controlled environmental housing. The birds were fed nutritionally adequate diets and treated according to the producer’s recommendations. At the age of 2 weeks, the hens were vaccinated against NDV by oculo/nasal (o/n) administration of a live vaccine. At the age of 5 weeks, they received a live NDV vaccine by spray. The sixteen-, 18- and 25-week-old breeders were boostered intramuscularly (i/m) with an inactivated NDV oil-vaccine. A modified vaccination schedule was applied from March 2003. The eighteen- and 25-week-old hens received i/m 1.0 ml of a two-component oil-vaccine containing inactivated ND and EDS viruses, instead of a one-component inactivated oil-vaccine against the NDV. The egg production was expressed as the average number of the eggs/turkey/week. Morbidity and mortality were monitored daily. The dead birds were necropsied and the organs were submitted for pathological analysis. The blood samples were taken by brachial vein puncture.

**Serological analyses**

The seroprevalence of antibodies to EDSV was determined in the sera collected from 30 breeder flocks (322 samples) prior to the vaccination, and 13 breeder flocks (151 samples) after the vaccination. The turkeys were between 11 and 46 weeks old. The levels of the antibodies against the EDSV were determined by haemagglutination inhibition test (HI-test). The two-fold serial dilution test was performed with 4 haemagglutination (HA) units of the EDS’76 strain BC14 and 0.8% chicken erythrocytes (Adair et al. 1979). The HI-titres ≥ 16 were considered positive.

The presence of the antibodies against the NDV was determined by the HI-test using a β-micrititration procedure (Allan and Gough 1974). The HI-tests were performed with a constant volume of 50 μl of serially double-diluted sera in multiwell plastic plates with V-shaped wells in the presence of 1% suspension of the chicken erythrocytes and the 4 HA units NDV La Sota strain as an antigen. The HI-titre of an antiserum corresponds to the highest serum dilution that inhibits haemagglutination. The HI-titres ≥ 16 were considered positive.

The sera samples were analysed for the antiAPV antibodies using a Flockscreen™ (Guildhay Ltd, Guildford, England) ELISA kit. Flockscreen ELISA titres ≥ 1210 were considered positive. The titres ≤ 876 were considered negative, while the sera titres between 877 and 1209 were in the suspect range.

Antibodies against AIV were determined by immunodiffusion test in 0.8% agar gel (AGID) supplemented with 0.15 M NaCl and 0.1% NaN₃, by using the AIV strain A/Shearwater/Australia/70 (H6N5) as an antigen (kindly provided by prof. B. Sinković, Central Veterinary Laboratory, Glenfield, N.S.W. 2167, Australia). The antisera that formed immunoprecipitates with the antigen after 24-hour incubation at 37 °C in a wet chamber were considered positive.

The presence of antibodies against *Mycoplasmae* was detected by a rapid serum agglutination test using the Nobilis® MG strain S6 of Adler and MS strain WVU-1853 antigens (Intervet International B. V., Boxmeer, The Netherlands). The test procedure and the evaluation of the results were given according to the prescriptions of the antigen producer. A blue-coloured flocculation occurring within 2 minutes, characterized a positive reaction. The reaction was considered negative for the fail of agglutination after an observation period of 2 min. The flocculation which occurred after 2 min was considered a doubtful reaction.

**PCR analysis for determination of EDSV**

Sera samples were collected on 4 breeding farms during the period between 2002 and 2003. A trivalent oil-vaccine Nobilis IB+ND+EDS (Intervet, Boxmeer, The Netherlands) containing the EDS’76 strain BC14, was used as a positive control. A haemorrhagic enteritis virus of turkey antigen for agar gel immunodiffusion test (Poultry Centre, Zagreb) was used as negative control.
From the sera and the vaccine, 200 μL was submitted to DNA extraction using a Machery-Nagel Nucleospin® Blood extraction kit (Düren, Germany) according to the manufacturer’s instructions. The concentration of the DNA extracted was determined spectrophotometrically at 260 nm.

Primers H5F and H6R specific for the EDSV hexon gene were used according to the data given by Rauwe and Hess (1998). For PCR, 50 ng of DNA were mixed together with 0.2 mM of each primer in a total volume of 25 μL using an AccuPrime™ Taq DNA Polymerase System (Invitrogen, Carlsbad, CA, USA). The PCR was carried out using a GenAmp PCR System 2400 (Applied Biosystems, Foster City, CA, USA) under the following conditions for 30 cycles, 94 ºC for 45 s, 54 ºC for 45 s, 68 ºC for 1.5 min. The amount of 10 μL of each PCR product was analysed by electrophoresis in a 1% agarose gel stained with ethidium bromide.

Results and Discussion

Rough, thin, soft-shelled or irregularly shaped eggs accompanied by a fall in the egg production, were recorded in turkey laying flocks for the first time during the first quarter of 2002. During the period from the 3rd to the 5th week of laying, the drop in the egg production in three flocks varied between 3.7 and 14.5% (Fig. 1, 2002a). At the same time, a decrease in fertility was observed in 8.7 to 19.0% of the eggs, while the 1st class hatchability was low in 1% of the eggs during 5 weeks. The HI-titre values of all the sera collected from 36-week-old hens at the 5th week of lay (Fig. 1, 2002a, denoted by an arrow) were positive to the antiEDSV antibodies and varied between 16 and 64.

The fall in egg quality and quantity continued in 2002 and at the beginning of 2003. Two diagrams characteristic for the egg production in 2002 (Fig. 1, 2002a and 2002b) and one for the beginning of 2003 (Fig. 1, 2003a) are presented. The egg shell changes and the reduced egg quality preceded or were concurrent with the decline of the egg production. A total decrease of the hatchability rate amounting to 7.7% accompanied the drop of the egg production and quality in 2002.

During the period of the decreased egg quantity and quality, 322 sera samples (233 in 2002, and 89 at the beginning of 2003), were analysed for the presence of the antiEDSV antibodies. It was found that 94.4 and 55.1% of the sera samples collected during 2002 and at the beginning of 2003, respectively, were antiEDSV positive (Fig. 2). The antibody HI-titres ranged from 16 to 128. The presence of the antiEDSV antibodies was detected in the sera collected at the beginning and during the laying period at the hen age of 30, 31, 36, 38, 39, 43, 46 weeks, as well as prior to the laying period in the sera of 11-, 15-, 21-, 24- and 27-week-old pullets. During the period of a decrease in the egg production, no serious clinical signs of the illness were observed in the birds. Only a few birds showed very mild respiratory difficulties. The food intake was not reduced. No specific pathological changes were found during the histopathological examinations. Mortality was within a normal range for the Nicholas hybrid line.

Hen vaccination against the EDSV began in March 2003. The egg production reverted to the normal values or to the values higher than normal for the Nicholas hybrid line (Fig. 1, 2003b). As a result of the vaccination, 96.7% of a total of 151 analysed sera were antiEDSV positive (Fig. 2). Higher average values of the antiEDSV antibody levels were generated by a vaccination rather than by a naturally-occurring infection (Fig. 2). The vaccination against the EDSV was carried out using a vaccine dose twice as high as that used for the chickens. We assumed that, for this reason, the naturally-occurring infection resulted in a lower humoral immune response than that elicited by a high EDSV vaccine dose.

The PCR analysis confirmed the presence of the EDSV in the serum sample of a 36-week-old antiEDSV positive non-vaccinated hen (Fig. 3, lane 5). The serum sample was collected during the period of the first observation of abnormal eggs (Fig. 3, lane 5). The results of PCR analyses of 7 randomly chosen antiEDSV negative sera samples, collected from 20- to 30-week-old turkey hens during the period of a suspected EDSV infection prior to the vaccination, are presented in lanes 6 - 12 (Fig. 3). The presence of EDSV was detected in 4 serum samples (lanes 6, 8, 10 and 11). The
EDSV was also detected in a serum sample of a hen twice vaccinated against the EDSV (Fig. 3, lane 4).

Primers H5/H6 (Raue and Hess 1998) hybridized in two variable regions (L1 and L4) of the EDS hexon gene where the lowest sequence identities between the EDS and other fowl adenoviruses (FAVs) were found which enable a clear differentiation between these viruses. This was verified by use of a commercially-available EDSV vaccine as positive control, and antigen for HEV detection as negative control for the PCR analysis. The PCR confirmed the presence of the EDSV in the antiEDSV positive sera samples of a non-vaccinated and a vaccinated hen (Fig. 3, lanes 5 and 4). On the other hand, the EDSV genome was also detected in the antiEDSV negative sera (Fig. 3, lanes 6, 8, 10 and 11). This result can be explained by higher sensitivity of PCR than that of HI-test. It is also possible that the antiEDSV negative turkeys were still infected by the EDSV, but
there was not enough time for the development of measurable levels of specific antibodies.

The antiNDV antibodies were detected in more than 95% of a total of 195 analysed sera collected prior to and after the vaccination against the EDSV. The antibody HI-titres varied between 16 and 128. The presence of antiNDV antibodies in almost all of analysed sera can be explained as a consequence of a systematically performed turkey vaccination against NDV.

All analysed sera, collected prior to and after the vaccination against the EDSV, were negative to antibodies against AIV (197 samples), MS (192 samples) and MG (192 samples).

A total of 433 sera were analysed for the presence of the antiAPV antibodies. The ELISA tests showed that 72.1 and 45.5% of the sera analysed were antiAPV positive prior to and after the vaccination against the EDSV, respectively. The ELISA titres of the antiAPV positive sera varied from 940 to 11 400. In some of the sera samples, both the antibodies...
against the EDSV and the APV were found simultaneously. The antibodies against the APV were detected in 74.4% of the antiEDSV positive sera prior to the vaccination against the EDSV. After the vaccination, 43.3% of the sera positive to the antiEDSV, were also positive to the antiAPV. According to these results, the decrease of the egg production and quality might be influenced by the APV infection. On the other hand, the APV infection was permanently present in turkey flocks prior to 2002 and no abnormalities in the egg production were detected during this period. After the vaccination against the EDSV, the egg quality and the production reached the normal values (Fig. 1, 2003b). The levels of the antiAPV antibodies in the breeder sera collected during this laying period were still high. These results and the absence of the AIV, MS and MG infections suggest the possibility that the egg production prior to the vaccination against the EDSV was influenced solely by a naturally-occurring infection. The potential source of the outbreak of the EDS in turkeys could be a horizontal spread of the infection from the non-vaccinated chickens in the neighbourhood of the turkey farms. The EDS, recorded in the turkey flocks not earlier than 2002, is permanently present in the domestic chickens in Croatia. A systematic vaccination by an inactivated vaccine prior to the laying period (Bidin et al. 1997) is applied for the prevention of the infection in the breeding and commercial egg producer chicken farms but not in the small back-yard flocks.

In the presented examination we showed that after the vaccination programme was introduced, the egg quality and the production reached normal values. This data, together with the serological and PCR analyses of the samples collected under the field conditions proved the existence of a naturally-occurring EDSV turkey infection.

Presented results provide a background for further investigation of the possible new EDSV outbreaks in other turkey farms. Further research will be completed with field virus isolation and characterization.

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