Ultrastructural Assessment of the Integrity of the enteric Mucosa of Commercial Turkeys Vaccinated against Coccidiosis

ABSTRACT

The objective of this study was evaluated the villus integrity of commercial turkeys submitted to coccidiosis prevention methods and challenged with *Eimeria* field oocysts, using scanning electron microscopy. Sixty BUT 9 female commercial turkeys were distributed in a completely randomized block design split with two treatments: T1-control diet without vaccinations against coccidiosis and anticoccidial drug, and T2- vaccinated against coccidiosis (commercial vaccine). On d 21 of life, all birds of all treatments were challenged with a mixed-species containing *E. meleagrimitis* and *E. gallopavonis*, via crop intubation with 1 mL of ~20,000 sporulated oocysts/bird. The size of the inoculum was determined in previous experiments. At 27 and 70 days of age, five birds per treatment were randomly removed to evaluate the intestinal integrity. Duodenum, jejunum and ileum segments were collected and processed according to routine scanning electron microscope. The enteric mucosa integrity of the commercial turkeys subjected to coccidiosis vaccination was reduced when compared to the birds of the control treatment. There was no difference in the performance results, therefore these results cannot be attributed to the vaccination as well as to the ineffectiveness against the challenge that was administered.

INTRODUCTION

The most relevant enteric diseases of infectious origin in turkeys are hemorrhagic enteritis, necrotic enteritis, histomoniasis, poult enteritis and mortality syndrome (PEMS), salmonellosis, and coccidiosis (Hafez, 2004; Hafez, 2013).

Coccidiosis or eimeriosis caused by protozoan parasites of the genus *Eimeria*, and it is considered one of the most important diseases in the poultry industry. Eimeria not only causes enteritis and diarrhea, impairing the intestinal absorption of nutrients, but also has a synergistic effect with other diseases, which become more severe in the presence of coccidiosis (Allen & Fetterer, 2002). Low infection levels of coccidia may reduce the birds’ metabolic and immune efficiency, causing significant economic losses in the turkey industry (Lillehoj & Lillehoj, 2000).

Disease prevention methods, associated with adequate management practices, are necessary to proper control and reduce disease outbreaks. When disinfection and cleaning alone are not sufficient to control the disease, anticoccidials need to be added to the feed or birds need to be vaccinated, because *Eimeria* oocysts remain viable for more than a year in the environment (Allen & Fetterer, 2002). Live vaccines are still not frequently used in turkeys, particularly in commercial production. The economic impact, the production benefits, and the limitations of vaccination and control strategies of coccidiosis is improving over time (Allen & Fetterer, 2002).
The objective of this study was to assess the integrity of the enteric mucosa of commercial turkeys vaccinated against coccidiosis and challenged with oocysts of a field species of *Eimeria*, using scanning electron microscopy.

**MATERIAL AND METHODS**

In this experiment, seventy-one-day-old female commercial turkey poult (British United Turkeys – BUT Big 9) were housed in the experimental aviary of UNESP (São Paulo State University), Veterinary Medicine and Animal Science School (FMVZ), Botucatu, Brazil. The poult were vaccinated at the hatchery against Newcastle disease and infectious rhinotracheitis. All birds were submitted to the same housing, feed management, and daily management practices, according to the recommendations of the BUT manual [12].

The birds were distributed in a completely randomized block design into two treatments. In Treatment 1, birds were not vaccinated against coccidiosis, and in Treatment 2, birds received a vaccine against coccidiosis (commercial vaccine, with a pool of oocysts of *E. adenoeides*, *E. dispersa*, *E. meleagrimitis*, and *E. gallopavonis*).

Birds were housed in experimental pens covered with sawdust litter. Birds were housed at densities of 21 birds/m² in the first week, 9.8 birds/m² between the second and sixth week, and 4.2 birds/m² from the seventh week up to the slaughter age (BUT, 2013).

Water and feed were supplied *ad libitum*. The diets were iso-nutritious and the feeding program was divided into four phases: starter (0 to 3 wk), grower (4 to 6 wk), developer (7 to 8 wk), and finisher (9 to 10 wk). The experimental diets were formulated to meet or exceed the minimum nutritional requirements of turkeys of turkeys according to the NRC (1994).

On d 21, each bird of both treatments was challenged with 1 mL of a mixed-species inoculum containing ~20,000 sporulated oocysts of *E. meleagrimitis* and *E. gallopavonis*. The concentration of the inoculum was determined in previous experiments, and it was administered directly into the esophagus of the birds, via gavage.

On d 27 and d 70, five birds per treatment were randomly selected, sacrificed, and necropsied to evaluate the integrity of the intestinal mucosa. The ileum, duodenum and jejunum were longitudinally opened, stretched, and washed with 0.1 M phosphate buffer (pH 7.3) to remove the intestinal contents. Tissues were fixed in 2.5% glutaraldehyde in phosphate buffer (pH 7.3, 0.1 M) for 24 hours at 4°C. After fixation, the samples were rinsed in distilled water and fixed for 30 minutes in 0.5% osmium tetroxide in distilled water. The material was dehydrated in an ascending ethanol series (7.5% to 100%) and then dried with liquid carbon dioxide in a Balzers CPD-020 critical point drying apparatus, assembled on “stubs”, and metalized with 10 nm gold using a sputter coater Bal-Tec SCD 050. The samples were examined and photographed using a scanning electron microscope FEI Quanta 200 from 10 to 15 kV.

Each bird represented an experimental unit. The results were subjected to analysis of variance (ANOVA) and the means were compared by the t-test (p<0.05), using the SAS statistical program (SAS, 2004).

The experimental protocol was approved by the Commission of Ethics in Research (n°188/2007 CEEA) of the Universidade Estadual Paulista “Júlio de Mesquita Filho”, Botucatu Campus, Brazil.

**RESULTS AND DISCUSSION**

The vaccinated turkeys were significantly heavier (p<0.05) than those not vaccinated at slaughter age. In the initial stage (21 days) the mean weights were T1 (control) = 843 g (± 42 g) and T2 (vaccination against coccidiosis) = 984 g (± 50 g). The values obtained for transitional (35 d) were T1 = 1390 g (± 79 g) and T2 = 1534g (± 82 g), for fattening (53 d) were T1 = 3773 g (± 148 g) and T2 = 3965 g (± 158 g) and final stage (70 days) were T1 = 5730 g (± 308 g) and T2 = 5870 g (± 311 g). During the growth phase (d 0-28), after coccidiosis challenge, only the birds belonging to the control group presented clinical signs of diseases such as diarrhea and prostration; there was no disease-provoked mortality.

Tests with the same commercial vaccine used in this study were previously conducted in commercial turkey flocks (Lee & White, 2001; Clark & Augustine, 2003; Linckersdorff-Sietz et al., 2004; Lee et al., 2006), and no typical clinical signs of “vaccine reaction” or significant impairment of live performance were reported. In chickens, according Peek & Landman (2011), the best method for coccidiosis control in the field is the use of live attenuated vaccines, as these promote good protection and do not cause severe injury to the intestinal mucosa, which may happen with the use of live vaccines. However, there are no attenuated vaccines for turkeys.
The enteric morphology evaluation in the growth phase (0-28d) confirmed better villus integrity in the duodenum, jejunum and ileum of the vaccinated birds (T2) compared with the same segments of the control birds (T1), indicating the protection of the intestinal cells provided by the vaccine against the injury caused by the coccidial challenge (Figure 1). According to Chapman et al. (2005), because live anti-coccidial vaccines are administered only once during the grow-out, re-infection, by ingesting small doses of vaccine oocysts from the feces or even from non-vaccine oocysts present in the litter, is essential for the establishment of immunity. The results of the present study indicate that the vaccinated birds made better use of the energy and of the chemical compounds, which besides ensuring adequate structural conditions of the mucosal cells of the digestive tract. They also ensured a better balance of the processes that determine the enteric mucosa turnover, resulting in a better absorption and digestion of the products.

Irrespective of the treatments, changes in the shape and organization of the villi were observed at different ages. At 27d of age, duodenal villi were dispersed and leaf-shaped, whereas the jejunum and the ileum presented high villus density and finger-shaped villi. At 70 days of age, duodenal villi were very similar to those observed at 21 days of age, i.e., dispersed and leaf-shaped. On the other hand, the jejunum and the ileum presented high villus density, and tongue-shaped villi arranged in a zigzag manner (Figure 2). These observations are consistent with literature reports on the mucosa of broiler chickens (Yamauchi & Isshiki, 1991; Pelicano et al., 2003; Van Leeuwen et al., 2004). According to those authors, the zigzag organization of the villi allows more efficient nutrient uptake when compared with randomly organized villi, since the distance traveled by the food bolus is higher in a zigzag flow, thereby enabling greater contact between the nutrients and the surface of the intestinal epithelium.

**CONCLUSION**

The vaccination used as a method to prevent and control coccidiosis preserved the intestinal morphology integrity of turkeys, besides proving to be efficient in the initial stage.

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