Outbreak of Viscerotropic Velogenic form of Newcastle disease in vaccinated six weeks old pullets

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Abstract

An outbreak of very virulent form of Newcastle disease in 6 week old pullets is reported. The flock was vaccinated against Newcastle disease with Newcastle disease vaccine intra ocular at the hatchery and Newcastle disease vaccine Lasota at 4 weeks of age at the farm. The signs noticed by the farm Manager were yellowish and greenish watery faeces, weakness and decrease in feed consumption. Response to treatment with Bioxin® (oxytetracycline) was poor. The sick birds never recovered. The gross lesions observed were: Necrosis of the gastrointestinal tract (Payers patches), caecal tonsils and bursa of fabricius. The disease lasted for 11 days and the mortality rate was 99.7%. The high mortality rate suggested that the vaccine had failed to immunize the birds against Newcastle disease and the birds were highly susceptible and had been infected with a very pathogenic stain of Newcastle disease virus.

Key words: Velogenic Newcastle Disease, Vaccine, Pullets and Mortality

Introduction

Newcastle disease (ND) is an acute contagious and pathogenic disease of poultry (Heath et al., 1991). The mortality rate can be up to 100% and it is considered to be among the most important viral disease s of poultry in the World (Heath et al., 1991; Alders and Spradbrow, 2001). The disease is characterized by respiratory sings, Nervous manifestation and diarrhoea (Alexander, 1990). Different isolates and strains of the virus may produce enormous variations in the severity of the disease in a given host even the chicken (Alexander, 1997). The first documented outbreak of the disease occurred between December 1952 and February 1953 in and around Ibadan (Hill et al., 1953). Since then the disease has been reported to endemic in both Local and Commercial poultry with annual epidemics being recorded in highly susceptible poultry flocks (Adu et al., 1986; Sa’idu et al., 1994; Halle et al., 1999; Orajaka et al., 1999). ND was also described as one of the most economically important disease of poultry in Nigeria (Okeke and Lamorde, 1988). The disease was also reported to be more common during the dry harmattan period (Sa’idu et al., 1994; Halle et al., 1999) and the stress of the harmattan period is known to worsen the out come of the disease (Abdu et al., 1992).

Newcastle disease vaccines have been of considerable value in reducing losses from the disease in general (Philip, 1973; Allan et al., 1978.; Shamaki et al., 1989). However the disease continues to occur in both vaccinated and unvaccinated flocks (Halle et al., 1999; Sa’idu et al., 2006). This paper reports an outbreak of the viscerotropic velogenic form of Newcastle disease in a vaccinated flock in due to vaccine failure.

Materials and Methods

On presentation, the flock history was obtained, the farm was visited and the clinical signs observed were recorded, Post mortem (P M) examination was conducted on dead birds. The flock was monitored on daily basis throughout the course of the disease and the daily morbidity and mortality were recorded. Ten birds were selected randomly and bled through the wing vein at the onset of the disease and three weeks after on set of the disease. Extra vials of the Newcastle disease vaccine and Gumboro disease vaccines used were obtained from the farm manager. Five Local chickens and five 6 week old chickens were used for vaccine trials .The birds were bled before vaccination and 2
weeks post vaccination. Serum was separated from the blood by centrifugation at 2000 rpm for 10 minutes. The sera were used for haemagglutination inhibition (HI) test as described by Allan and Gough (1974) Allan et al. (1978). The antigen used for the HI test contain 4 HA units. The sera were also tested for antibodies to Infectious bursal disease by the Agar gel precipitin test (AGPT) as described by Hirai et al. (1972) and Wood et al. (1979). Bursa of fabricius of birds that suffered from typical Infectious bursal disease (IBD) was used as positive antigens for the AGPT. The sera were also tested for antibodies to Salmonella and Mycoplasma by the rapid plate test. Sections of the intestine and caeca were taken for parasitological examination.

Results

History
A total of 5,054 day old Lohman brown chicks were bought from S and D farms Abeokuta. They were divided into 2 and housed in two different pens (1 and 4). Pen 1 contains 2542 and pen 4 contains 2512 chicks. The following drugs were used before the outbreak, Tetroxy® (oxytetracycline) within the first week, Vitaperos® (vitamins and minerals) for 5 days at 2 weeks, Furaprol® (Furaltadone 20% and Amprol 20%) for 5 days at 3 weeks Vitaperos® for 3days at 4 and 5 weeks of age. In addition the birds were vaccinated against ND with ND vaccine intra ocular (i/o) at the hatchery and on the farm, NDV Lasota at 4 weeks and Gumboro disease vaccine at 2 and 5 weeks of age. The vaccines were purchased directly from the manufacturer. The birds were fed with Silver® feed ad libitum.

The signs observed by the farm manager were yellowish and greenish watery faeces, Weakness, Sleepiness and decrease in feed consumption. Response to administration of Bioxin® (oxytetracycline) in drinking water was poor and sick birds never recovered. The disease was first noticed in house 1 and by the sixth day after onset of the disease it spread to house 4.

Clinical Examination
Yellowish or greenish diarrhea, anorexia, weakness, somnolence, dropped beak prostration, severe depression, dehydration, abnormal respiration and trembling were notices at the onset of the outbreak. Later tracheal rales, sneezing, gasping, coughing and clonic spasms were also noticed. Towards the end of the outbreak the remaining birds showed circling, trembling, torticollis and star gazing. The mortality pattern is shown in table 1. The total mortality was 99.7% and the disease lasted for 11 days and most of the death (80%) was recorded on the 5th day (house 4) and on the 9th day (house 1) after onset of the disease.

Post Mortem Examination
The trachea, lungs, pancreas, thymus and muscles were congested. There were haemorrhages in the trachea, proventriculus, duodenum, jejunum, ilium, caeca, caecal tonsils and bursa of fabricius. Necrosis was observed in the gastrointestinal tract (Peyers patches areas), caecal tonsils and bursa of fabricius, other gross lesions seen were ascites, caecal cores, regression of the bursa and enlargement of the spleen, liver and kidneys.

Laboratory Findings
Antibodies to Salmonella and Mycoplasma were not detected. However 82% of the birds tested had antibodies to Gumboro disease. At the onset of the outbreak the mean HI antibody titre to ND was low (≤ log2, 2.6). The mean antibody titre to ND however increased three (3) fold, 3 weeks after the onset of the disease. (≥ log2, 9.0). Very few coccidian oocyst were seen in the intestinal content and Histomonas specie were present in the intestinal and caecal segments. The antibody titre to Gumboro disease in experimentally vaccinated birds was low (≤ log2, 2.0) at 2 weeks post vaccination and the HI titre for ND was equally low (≤ log2, 4.0) (Table 2).

Treatment
Furaprol® at the dose rate of 5 gram per 5 litres of drinking water for 7 days

Advice
The client was advised to ensure day old vaccination against ND and 3 weeks with Lasota rather than 4 weeks of age. He was also advised to sero-monitor the birds before and 2 weeks post vaccination to ensure that they have produced enough anti-bodies that will ensure they are well protected when exposed to pathogenic ND.
Table 1:
Course and mortality rate of the Newcastle disease in vaccinated pullets.

<table>
<thead>
<tr>
<th>House</th>
<th>Course of the disease in days</th>
<th>Mortality rate ( % )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>11</td>
<td>99.7</td>
</tr>
<tr>
<td>4</td>
<td>10</td>
<td>99.7</td>
</tr>
</tbody>
</table>

Table 2: Comparision of antibody response of chickens vaccinated with Newcastle disease Vaccine (Lasota), Gumboro vaccine and chickens suffering from Newcastle disease

<table>
<thead>
<tr>
<th>Type of birds</th>
<th>Mean HI antibody titre log2</th>
<th>Mean antibody titre to Gumboro disease log2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Pre-vaccination 2weeks post vaccination</td>
<td>Pre-vaccination 2 weeks post vaccination</td>
</tr>
<tr>
<td>Improved chickens</td>
<td>2.0 3.3</td>
<td>- 2.0</td>
</tr>
<tr>
<td>Local chickens</td>
<td>2.1 3.8</td>
<td>- 0.4</td>
</tr>
<tr>
<td>Sick chickens</td>
<td>At onset of the disease 3 weeks Post on set</td>
<td>At onset of the disease 3 weeks Post on set</td>
</tr>
<tr>
<td></td>
<td>2.6 10.2</td>
<td>- 0.4</td>
</tr>
</tbody>
</table>

Discussion

The chicks were susceptible to ND at the onset of the outbreak despite prior vaccination 11 days earlier, because the antibody level in the sera of the chicks were similar to those of the birds that have not been vaccinated against ND or exposed to ND previously (OIE, 1996). It was either that the vaccines used contain very low viral titres or the birds failed to respond to the vaccination. The most likely reason was the former based on the antibody response of the vaccinated birds as the mean HI titre expected in vaccinated birds ranges from log2, 4.0 – log2, 6.0 (Alexander, 1990; OIE, 1996 ). The clinical signs, mortality rate and post mortem lesions are similar to what was reported in birds infected by the Viscerotrophic Velogenic ND virus ( Alexander, 1990 ). The histomonas and the coccidian oocyst seen in intestinal sections in this case might have been complications following infection with the viscerotropic velogenic ND virus. It is well known that Histomonas meleagridis is a primary cause of disease in turkeys and occasionally in chickens and other species of birds (Tress, 1990). Treatment was not reviewed because the furaprol contain drugs active against coccidian, histomonas, gram negative and positive bacteria. This report highlights the need to conduct pre and post vaccination serology.

References


