Clinicopathological evaluation of Newcastle disease virus vaccination using gums from Cedrela odorata and Khaya senegalensis as delivery agents in challenged chickens

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ABSTRACT

Following previous studies on delivery potential and immune response of chickens given Newcastle disease vaccine with gums, this study was conducted to evaluate the protective ability of vaccines delivered with plant gums against clinicopathological features of Newcastle disease (ND). Processed gums from incised trunks of Cedrela odorata and Khaya senegalensis trees were combined with ND vaccine in ratio 2:2:1 and administered at 21 days to white leghorn cockerels after weaning of maternal antibodies. The birds were grouped into gum-vaccine-oral (GVOR), vaccine-oral (VOR), gum-vaccine-ocular (GVOC), vaccine-ocular (VOC), gum-ocular (GOC), gum-ocular (GOR), no-gum-no-vaccine/challenged (NGNV/C), no-gum-no-vaccine/unchallenged (NGNV/U). Vaccination was boosted with the same preparation at day 42 while birds were challenged with live ND virus (KUDU strain) at day 84. Clinical signs (Dullness, Diarrhoea, Paralysis, Torticollis) Post infection (Pi), terminal weakness, gross and histology lesions were scored on a severity scale from absent (-), mild (+1) to moderate (+2) and severe (+3). Scores were assigned a quantitative score of 0, 10, 20, 30 respectively. Clinical signs scores for the 5 week Pi were subjected to Friedman test to assess the significance of severity among the groups. The test was significant at 1% significance level which implies that the clinical signs ranked highest in the NGNV/C, followed by the Gum alone groups, the vaccine alone groups and the gum-vaccine groups irrespective of route. Moribund birds subsequently euthanized were seen in the GOR and GOC group at 21% each and at 57% in NGNV/C group alone. No signs were seen in the NVNG/U group. Grossly, mild to moderate lesions were seen in all groups except GVOR and NGNV/U. At histology, pulmonary congestion, acute pneumonia, cecal tonsilar haemorrhages, gliosis and neuronophagia were present at different proportions in all groups except the GVOR and NGNV/U. Overall, lesion severity was least in the gum-vaccine groups while the oral groups had less lesion score compared to the ocular. From this study, phytogenic mucoadhesives polymers used hold immense potential as a delivery agent capable of improving protection against clinicopathologic features of Newcastle disease in previously vaccinated birds.

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1. Introduction

Newcastle disease is an infectious often fatal viral disease of poultry. It is endemic in most countries of the world with seasonal or sporadic epizootic spikes since first reported in England [1,2]. It is caused by Avian Paramyxovirus 1 (APMV-1) of the genus Avulavirus in the family Paramyxoviridae which was formerly alongside grouped with other nine serotypes (APMV 2–9) into
genus Rubulavirus until recently [1,3]. It is also responsible for major economic losses in both commercial and backyard flocks because outbreaks often reach 100% mortality in virulent strains [4].

Broadly, there are 3 classifications of APMV-1 strains namely Velogenic (high virulence), Mesogenic (intermediate virulence) and Lentogenic (low virulence) although an overlap of these strain classes exists in the field [4]. Severity of the disease often follows the virulence of the infecting strain amongst other factors such as host immune status, co-infections with other agents, age of host, environmental stress and endemicity [5,6].

Generally, non-specific signs of the disease could include depression, greenish diarrhoea, prostration and oedema of the head including the wattles, paralysis, torticollis and other respiratory signs. In layer flocks, drop to complete stoppage of egg production, small, spotted or soft shelled eggs have been reported to indicate the presence of the virus [7]. However grossly, post mortem lesions could range from severe eyelid oedema, haemorrhages in the intestinal tracts (duodenum, jejunum and ileum), caeca tonsils, proventriculus, perithymic areas to necrohemorrhagic plagues in the ceca and small intestine. Spleenic congestion, enlargement, stippled appearance extending from the capsule into the parenchyma could also be noticed in other strains [4]. Catarrhal or caseous exudates may be present from associated secondary bacterial infection [8]. Birds in lay could develop egg yolk peritonitis. The ovarian follicles may be flaccid and degenerative in growing pullets [8,9]. Histologically, depletion of mononuclear cells from the spleen, extensive fibrin deposits around the periarteriolar lymphoid sheaths, considerable destruction of lymphoid areas in the ceca and intestine with deposit of necrotic materials are often observed. In some cases, lymphoid depletion of the bursa and thymus might be present as well as ulcers in the small intestine. In the central nervous system, focal neuronal degeneration, gliosis, presenting with a non-suppurative encephalomyelitis are the main reported lesions [10,11].

Major control measures aside proper biosecurity involves the use of vaccines with recommended schedules. However, major constraint with existing vaccine protocols in developing nations with endemic ND burden involves short duration and inconsistent protective humoral response. The latter is associated with live attenuated vaccines administered orally or intraocular, while invasiveness/ease of administration and unwanted reactions such as egg drop are seen with parenteral oil based vaccines. Newer recombinant vaccines also have low coverage and are cost prohibitive. Hence these shortfalls have resulted in disharmonized or immune responses along mucosa surfaces with reduction of clinical signs and associated mortality resulting in economic losses influenced the purpose of this research. This study was aimed at evaluating the clinicopathologic features of challenged chickens previously vaccinated with ND vaccine using natural mucoadhesive biopolymers such as gums from plants. This was followed recent reports by Emikpe et al. [13] who reported strong mucoadhesive property of gums obtained from Cedrela odorata and Khaya senegalensis trees when used as vaccine vehicles with a suspected immunopotentiating action when investigated using ex-vivo means. Further, Oyebanji et al. [14] reported higher, faster maturation and recorded as mortality), gross and histopathology lesions were followed for this in vivo study.

2. Materials and methods

2.1. Ethical approval

All international protocols concerning animal studies were followed for this in vivo study.

2.2. Vaccine reconstitution and experimental groupings

Following recommended dilution for the gum delivery agent [14], NDV Lasota (BN:183888) vaccine was combined with 1 in 8 bench dilution of 400 mg of purified and lyophilized Cedrela odorata and Khaya senegalensis gums (1:1) dissolved in 8 mL distilled water. This above mixture was the dilution needed to vaccinate exactly 200 birds giving 2 drops of constituted gum-vaccine preparation using 3 mL and 1 mL rubber pipettes per bird through the oral and ocular routes respectively as follows:

- **Group A**: Gum vaccine oral (GVOR)
- **Group B**: Vaccine oral (VOR)
- **Group C**: Gum vaccine ocular (GVOC)
- **Group D**: Vaccine ocular (VOC)
- **Group E1**: Gum alone oral (GOR)
- **E2**: Gum alone ocular (GOC)
- **Group F1**: No Gum No Vaccine/Challenged (NGNV/C)
- **F2**: No Gum No Vaccine/Un-challenged (NGNV/U).

Each vaccine vial was reconstituted as stated for the treatment groups. Birds were randomly grouped along the treatments into a number of 36 birds each and the NGNV/C and NGNV/U had 18 birds each.

2.3. Live virus for challenge test

Live Newcastle disease viral stock (KUDU strain) was acquired from the National Veterinary Research Institute, Vom Jos, Nigeria. The virus was inoculated in specific pathogen free (SPF) 9–11 day old embryonated chicken eggs to determine Embryo Lethal Dose (ELD) 50 dose as well as the titre. Allantoic fluid from the eggs with known titre per mL (10^5.5 per ELD) at 3.5 ELD_{50} was used for the challenge experiments at day 42 post booster vaccination.

2.4. Clinical features, gross, histopathology and scoring

Post infection, clinical signs; terminal weakness (euthanized and recorded as mortality), gross and histopathology lesions were recorded and scored as described by Emikpe et al. [18] with modifications. Daily, presence or absence of clinical signs (dullness, diarrhoea, prostration, and torticollis) with degree of severity was scaled as absent (0–); mild (1+), moderate (2+) and severe (3+). These scales were then assigned a score of 10 degrees such that 0– is assigned score of 0; 1+ assigned score of 10; 2+ assigned score of 20 and 3+ assigned score of 30. Scores for each clinical parameter were summed up as the collective clinical score for each group on a weekly basis. Gross lesions and histopathological were also scored as described above for lesions observed in main organs at the termination of experiment. Terminal weakness (euthanized......
and recorded as mortality) observed were recorded and percentage determined on a weekly basis. Post-infection, tissues were harvested and routinely processed for histologic examination. Briefly, samples were fixed in 10% formalin before trimming and sectioning for histology. Formalinized samples were dehydrated in grades of alcohol concentrations, cleared in xylene, and embedded in molten wax. On solidifying, sections of 5 μm thick were made with microtome (Leica RM2125 RTS), floated in water bath and incubated at 60°C for 30 min. The sectioned tissue was again cleared in grades of xylene and dehydrated in alcohol graded concentrations while sections were stained with routine hematoxylin and eosin stains. Examination of stained sections was done using light microscope (Olympus CH, XSZ107BN; 072351) while photomicrographs were taken with Ucmos series microscope camera Mu900 and Toupview 3.2 image software.

2.5. Statistical analysis

Friedman test (MINITAB version 16), a nonparametric alternative to the randomized complete block experiment was used to assess the significance of the clinical signs from the eight experimental groups for the five week observatory period post infection.

3. Results

The scored clinical signs, gross and histopathological lesion severity were presented in the bar charts (Figs. 1–4). Post infection clinical signs observed were dullness, greenish watery diarrhoea, leg paralysis, torticollis, and terminal weakness recorded as mortality. From the clinical score in the treatment groups as presented in the Friedman test results (Table 1) and illustrated in Fig. 1, severity of clinical signs ranked highest in the No-gum-No-Vaccine/Challenged group, followed by the vaccine alone groups irrespective of route, then the vaccine alone groups and finally, the gum-vaccine groups irrespective of route. The No-gum-No-Vaccine/Unchallenged group showed least severity being the control uninfected.

Further, in detail, first week post infection (PI), greenish diarrhoea was observed in all groups except the no gum-no-vaccine/unchallenged (NGNV/U) group. These became moderately severe in the vaccine alone group by the 2nd week and more severe in the gum alone and no-gum-no-vaccine/challenged (NGNV/C) groups throughout the PI period. By the day 9, diarrhoea sign had subsided in the gum vaccine oral group (GVOR), day 15 in the vaccine oral group (VOR), and in the gum vaccine ocular groups (GVOC), however, it continued in the vaccine ocular (VOC) group till 3rd week before it subsided while it was present throughout the 5 weeks PI in gum alone and unvaccinated infected groups. Consequently, dullness was observed in the gum alone flock irrespective of the route as well as in the NGNV/C group throughout PI period beginning from 2nd week PI. Also, torticollis and leg paralysis followed a similar pattern in these groups.

Grossly, at termination of experiment, main organs that had visible lesions grossly across the groups with the exception of NGNV/U were the lungs, ceca tonsils, intestines (jejunum and ileum) and spleen. In the lungs, congestion and focal areas of consolidation were most severe in the vaccine ocular (VOC), gum oral (GOR), gum ocular (GOC), and no-gum-no-vaccine/challenged (NGNV/C), moderately severe in the vaccine oral (VOR) and gum-vaccine ocular (GVOC) groups while absent in the gum-vaccine oral (GVOR) and no-gum-no-vaccine/Unchallenged (NGNV/U) group. In the ceca tonsils, mild petechial haemorrhages were seen in all groups except the GVOR and NGNV/U groups. In the intestine, mild petechial to echymotic haemorrhages were seen only in the NGNV/C group. In the spleen, No lesions were observed for the GVOR, VOR, GVOC and NGNV/U groups. However, moderate splenomegaly with hyperplastic follicles were seen in the VOC, GOR, GOC and NVNG/C groups.

![Clinical severity score for each group post-infection](image-url)

**Fig. 1.** Bar chart showing clinical signs (dullness, diarrhoea, paralysis, and torticollis) severity scores for each group post-infection. Absent (0–) = 0; Mild (1+) = 10; Moderate (2+) = 20; Severe (3+) = 30. A: GVOR = Gum-Vaccine Oral; B: VOR = Vaccine oral; C: GVOC = Gum-Vaccine Ocular; D: VOC = Vaccine ocular; E1: GOR = Gum alone Oral; E2 = GOC: Gum alone Ocular; F1: NGNV/C = No gum No Vaccine Challenged; F2: NGNV/U = No gum No Vaccine Unchallenged.
Fig. 2. Bar chart showing mortality pattern per week as observed in different groups. GVOR: Gum-Vaccine Oral; VOR: Vaccine oral; GVOC: Gum-Vaccine Ocular; VOC: Vaccine ocular; GOR: Gum alone Oral; GOC: Gum alone Ocular; NGNV/C: No gum No Vaccine Challenged; NGNV/U: No gum No Vaccine Unchallenged.

Fig. 3. Bar chart showing Gross lesion scoring of harvested organs following euthanasia of selected birds at week 5: Lungs: Congestion, Focal areas of consolidation; Ceca tonsils, Intestine (jejunum): Haemorrhages (petechiae to echymotic); Spleen: Hyperplastic splenomegaly (follicular). Severity scores for each group. Absent (0–) = 0; Mild (1+) = 10; Moderate (2+) = 20; Severe (3+) = 30. GVOR: Gum-Vaccine Oral; VOR: Vaccine oral; GVOC: Gum-Vaccine Ocular; VOC: Vaccine ocular; GOR: Gum alone Oral; GOC: Gum alone Ocular; NGNV/C: No gum No Vaccine Challenged; NGNV/U: No gum No Vaccine Unchallenged.
At histology, acute interstitial pneumonia, parabronchial infiltration of heterophils and mild pulmonary congestion were observed in all groups except the GVOR and NGNV/U groups (Fig. 5). In the cerebrum, severe gliosis, neuronal cell body necrosis and lymphocytic perivascular cuffing were observed in the VOC, GOR, GOC and NGNV/C groups, moderate lesions in VOR and GVOC while these lesions were absent in the GVOR and NGNV/U groups. In the spleen, moderate necrosis and depletion of lymphoid follicles were observed in the VOC, GOR, GOC and NGNV/C groups while these lesions were moderate in VOR, GVOC, GOC and mild in the GOR groups. Bursa of Fabricius presented with mild lymphocytic depletion of follicles and disruption of follicular architecture only in the NGNV/C group.

In summary, the severity of clinical signs, gross and histopathological changes could be graded from least severe to the most severe as follows:

- GVOR < NGNV/U < GVOC < GOR < VOR < VOC < GOC < NGNV/C.

### 4. Discussion

This study evaluates clinical signs and pathological changes observed in chickens challenged with a virulent field strain of Newcastle disease virus after prior and booster vaccinations with Newcastle disease vaccine delivered using gums from *Cedrela odorata* and *Khaya senegalensis* through the oral and ocular routes. From this study, it was evident that the vaccine used gave 100% protection with regards to terminal weakness leading to mortality among vaccinated groups irrespective of the route or delivery agent used. This validates the effectiveness of the vaccine used as well as the protection offered by both routes against mortality induced by Newcastle disease in birds. Protective capacities of vaccination through oral and ocular routes in chickens have already been reported in literature [19]. This is owing to the large surface area of the mucosa surface exposed to constant irritation and adapted for antigenic sampling and effector mechanisms [14,20].

However, significant difference exists in severity of clinical signs and post mortem lesions observed as the oral groups fared better than the ocular groups while the gum-vaccine groups showed lesser degree of clinical signs/post mortem lesions compared to the vaccine alone groups. This is especially true in the GVOR group where mild greenish diarrhoea was only observed for a duration of nine days post-infection (PI) without any other

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**Table 1**: Results of Friedman test for severity of observed clinical signs.

| Treatment | Number | Estimated median | Sum of Ranks |
|-----------|--------|------------------|--------------|
| A: GVOR   | 5      | 0.00             | 15.0         |
| B: VOR    | 5      | 0.00             | 17.0         |
| C: GVOC   | 5      | 0.00             | 15.0         |
| D: VOC    | 5      | 0.00             | 17.0         |
| E1: GOR   | 5      | 81.25            | 32.5         |
| E2: GOC   | 5      | 81.25            | 32.5         |
| F1: NGNV/C| 5      | 107.50           | 40.0         |
| F2: NGNV/U| 5      | 0.00             | 11.0         |

Grand median = 33.75.

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**Table 4.** Bar chart showing histopathologic changes score of harvested organs following euthanasia of selected birds at week 5 post-infection: Lungs: Congestion, Focal areas of consolidation; Cecal tonsils, Intestine (jejenum): Haemorrhages (petechial to ecchymotic); Spleen: Hyperplastic splenomegaly (follicular). Severity scores for each group. Absent (0−) = 0; Mild (1+) = 10; Moderate (2+) = 20; Severe (3+) = 30. GVOR: Gum-Vaccine Oral; VOR: Vaccine oral; GVOC: Gum-Vaccine Ocular; VOC: Vaccine ocular; GOR: Gum alone Oral; GOC: Gum alone Ocular; NGNV/C: No gum No Vaccine Challenged; NGNV/U: No gum No Vaccine Unchallenged.
sign or post mortem lesion compared to GVOC, vaccine alone or gum alone groups where moderate to severe diarrhoea extended for longer duration. This observed degree of severity of clinical signs and absence/lesser degree of post mortem lesions in oral groups (GVOR, VOR and GOR) as against the ocular (GVOC, VOC, GOC) highlights the effect of route of administration. This was evident by moderate to mild lungs lesions such as congestion and acute pneumonia marked by presence of inflammatory cells present in the ocular groups but absent in the oral groups. This disparity might not be unconnected with larger surface area for antigen induction, processing and more diffusely associated lymphoreticular structures with effector sites in the oral groups (gastrointestinal tract) compared to the ocular groups [21].

The larger surface area of the oral route with more inductive and effector sites in the gastrointestinal mucosa i.e. more follicles with specialized M-cells, Peyer’s’ patches, ceca tonsils may help to elicit higher immune response than the ocular route with fewer surface area for antigen induction/effector. This might account for the preferential protection offered by the oral route compared to the ocular one. This report is however at variance with reports...
by Okwor et al. [19] which favoured the intraocular route to the oral route mostly citing possible destruction of vaccine antigens by gastrointestinal secretions as probable reasons for reduced optimization of uptake and subsequent immune response. Hence, phyto- topical vaccine used in this study might have mitigated the destruction of vaccine antigens thereby optimizing uptake, response and subsequent reduced clinicopathologic feature following challenge by field virus.

Further, the better protection offered by the addition of the gum to the vaccine given through the ocular route as against the vaccine alone along the same route supports the suspected immunopotentiating property of the gum [14]. This could be mediated either through a slow antigen release ensuring prolonged antigenic exposure and stimulation or through an induction of a faster recall-memory response. This could ensure faster destruction of challenge live viruses since the birds were infected through the ocular route. Additionally, if the gum enhances immunologic memory induction, the principle of common mucosal immunity would assist in ensuring fairly even protection along mucosal effector sites thus reducing viral replication. Thus, under field conditions, protection along the ocular route could be maximized, by the addition of gums although the precise mechanism still remains to be evaluated.

Generally, the observed reduction in clinico-pathological features observed in the gum-vaccine groups might be related to the humoral systemic and mucosal responses to previous booster vaccination as suggested and demonstrated in our previous studies [13,14]. These studies showed a higher and faster peak humoral (mucosal and systemic) response (using HI and ELISA) [14] in challenged chickens after prior and booster oral NDV vaccinations when gum vehicle was employed. Thus, this further highlights the importance of the gum vehicle in infection immune response when previously included in vaccine preparation.

Darabighane and Nahashon [22] reported similar reduction of clinico-pathologic changes from other studies using plant parts. The report highlighted several studies about improved humoral, cellular responses as well as reduced clinical signs and lesions associated with different diseases as attributable to polysaccharides either present in plants used or given alone e.g. acemannan. In this study, as earlier stated, Cedrela odorata and Khaya senegalensis contain abundant polysaccharides. Thus, these polysaccharides may play a pivotal role in immunomodulation mechanisms observed leading to reduced clinical signs and pathologic changes in the gum-vaccine groups. Studies to validate this are presently underway.

Notably, presence of lung lesions and absence of marked lesions in the central nervous system, proventriculus and intestines of groups showing lesions might be attributed to the tropism of the infecting virus [KUDU] strain. Such reported preferential predilection of strains was reported by Wambura et al. [23] who reported preferential tropism of I-2 NDV vaccine strain in lungs than V4 strain which favoured the intestine. Either way, the inclusion of gums in the oral delivery of Newcastle disease vaccine ensured protection against the challenge virus which seems to show more preference for the lungs.

5. Conclusions

In conclusions, the use of phyto-genic mucoadhesives as delivery agents for Newcastle disease vaccines in chickens holds novel potentials in reduction of clinical and subsequent pathologic features associated with ND thus reducing economic losses usually observed with the disease. Further research studies are underway to evaluate the mechanism employed by gum vehicle in proffering better protection from clinical and pathologic changes of Newcastle disease observed in the study and immune response pattern observed from previous studies.

Competing interests

The authors of this manuscript have no competing interest.

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References

[1] Alexander DJ. Newcastle disease and other avian Paramyxoviridae infections. In: Diseases of poultry, 10th Ed. BW. Calnek with HC, Barnes, CW, Beard, LR. McHugh propulsion & YM. Saif, eds. Mosby-Wolfe, London 1997, p. 541–70.

[2] Health BC, Lindsey MS, McManus RF, Clixton PD. Newcastle disease vaccine for village chicken. Spadbrow P.D. (Ed). Australian Center for International Agricultural Research, Canberra, Australia 1991, p. 103.

[3] De Leeuw O, Peeters B. Complete nucleotide sequence of Newcastle disease virus: evidence for the existence of a new genus within the subfamily Paramyxovirinae. J Gen Virol 1999;80:131–6.

[4] Alexander DJ. Newcastle disease and other avian paramyxoviruses. Rev Sci Technol 2000;19:443–62.

[5] Okwor EC, Eze DC. Newcastle disease in layers: preliminary studies on the preference for the lungs.

[6] Okwor EC, Eze DC. Newcastle disease observed in the study and immune response pattern observed from previous studies.

[7] Okwor EC, Eze DC. The annual prevalence of Newcastle disease in commercial chickens rear in South Eastern Savanna zone of Nigeria. Res J Poult Sci 2010:3.23–6.

[8] Spadbrow PD. Epidemiology of Newcastle disease and the economics of its control. Proceeding of the workshop on poultry as a tool in poverty eradication and promotion of gender equality, March 23–26, Tune Landboskole, Denmark; 1999. p. 1–6.

[9] Oladele SB, Abdu PA, Nok AJ, Esevo KAN, Useh NM. Preliminary reports on neuraminidase, enzytocyte surface and free serum siaic acid concentrations in serum of healthy and Newcastle infected Chickens. Med Vet Paps Trop 2002;55:265–8.

[10] Oluwole OE, Emikpe BO, Ologasa BO. Attitude of poultry farmers towards vaccination against Newcastle disease and Avian influenza in Ibadan, Nigeria. Sokoto J Vet Sci 2012;10:5–12.

[11] Emikpe BO, Oyebanji VO, Odeniyy MA, Salaam AM, Okelede AE, Jarikre TA, et al. Ex vivo evaluation of the mucoadhesive properties of Cedrela odorata and Khaya senegalensis gums with possible applications for veterinary vaccine delivery. SpringerPlus 2016;5:1289.

[12] Oyebanji VO, Emikpe BO, Oladele AO, Odeniyi MA, Salam A, Osowole OJ, et al. Evaluation of immune response in challenged chickens vaccinated with Newcastle disease vaccine using gums from Cedrela odorata and Khaya senegalensis as delivery agents. J Immunol Med Immunol 2016;23:1–11.

[13] de Troconis GN, Martinez M, de Pinto LG, Bhasas A. Estudio químico y spectroscopico del polisacárido rido de la goma de Cedrela odorata. Ciencia Tecnol 2000;19:443–62.

[14] Okwor EC, Eze DC. Newcastle disease observed in the study and immune response pattern observed from previous studies.
[17] Waihenya R, Mtambo M, Nkwengulila G. Evaluation of the efficacy of the crude extract of Aloe secundiflora in chickens experimentally infected with Newcastle disease virus. J Ethnopharmacol 2002;79:299–304.

[18] Emikpe BO, Tanko PN, Onilude OM, Sabri MY. The influence of dexamethasone treatment and successive road transport stress on the occurrence of caprine pneumonia in a hot humid tropical environment. Vet World 2013;6:497–501.

[19] Okwor EC, Eze DC, Uzuegbu OM. Comparative studies on the oral and intraocular routes of administration of Newcastle disease vaccine, La Sota in adult chickens. J Agric Vet Sci 2013;3:48–51.

[20] Peralta MF, Danelli MGM, Vivas A. Rediscovering the importance of Mucosal Immune System (MIS) in poultry. Acad J Biotechnol 2016;4:91–5.

[21] Fix A, Arp LH. Morphologic characterization of conjunctiva-associated lymphoid tissue in chickens. J Vet Res 1991;52:1852–8.

[22] Darabighane B, Nahashon SN. A review on effects of Aloe vera as a feed additive in broiler chicken diets. Ann Anim Sci 2014;14:491–500.

[23] Wambura P, Meers J, Spradbrow P. Determination of organ tropism of Newcastle disease virus (Strain I-2) by virus isolation and reverse transcription-polymerase chain reaction. Vet Res Commun 2006;30:697–706.