Chitosan nanoparticle vaccine loaded with crude extracellular proteins of *C. perfringens* and *Salmonella* flagella is partially protective against necrotic enteritis in broiler chickens.

Gabriel Akerele\textsuperscript{a}, Nour Ramadan\textsuperscript{a}, Muhammed Mortada, Revathi Shanmugasundaram\textsuperscript{a}, Sankar Renu\textsuperscript{b}, Gourapura J. Renukaradhya\textsuperscript{b}, Ramesh K Selvaraj\textsuperscript{a*}

\textsuperscript{a}Department of Poultry Science, The University of Georgia, Athens, GA 30602.

\textsuperscript{b}Food Animal Health Research Program, Ohio Agricultural Research and Development Center, The Ohio State University, Wooster-44691, Ohio.

*Corresponding author at: The University of Georgia, Athens, GA 30602. E-mail address: selvaraj@uga.edu

**Contact Information**

Ramesh Selvaraj

Professor

Department of Poultry Science

University of Georgia

110 Cedar Street

Athens, GA 30602

Ph: 706-542 3706

Fax: 706-542-1827

selvaraj@uga.edu
Abstract
Necrotic enteritis (NE) causes significant economic losses and food shortages world-wide. There are currently no licensed commercial vaccines against NE in broilers. Chitosan nanoparticles were formulated with extracellular proteins of *C. perfringens* surface-tagged with *Salmonella* flagellar proteins. One-day-old male broiler chicks were completely randomized to 3 treatments: Non-vaccinated non-challenged as negative control, Vaccinated-challenged, and non-vaccinated challenge as positive control. On day of hatch, d7, and d14 post-hatch, vaccinated-challenged birds were orally gavage with 50µg vaccine in 0.5ml PBS while positive control birds were gavage with 0.5ml PBS only. Birds in the vaccinated-challenged and positive control groups were orally infected on d14 post-hatch, with 5,000 oocysts/bird of *E. maxima*, followed by log 8 CFU of a virulent strain of *C. perfringens* on d19, d20, and d21 post-hatch. From d14 to 21 and d14 to 28 post-hatches, mortality in the vaccinated-challenged group was higher than that in the positive control group, approaching statistical significance (p=0.07). On d21 post-hatch, the mean lesion score of 3 birds/cage in the vaccinated-challenged group was higher than the positive control group, approaching statistical significance (p = 0.05). From d14 to 28 post-hatch, the feed intake was higher and feed conversion ratio lower in the vaccinated-challenged group compared to the positive control group (p<0.05). On d21 post-hatch, antigen specific recall proliferation in the vaccinated-challenged group was higher than that in the negative and positive control groups (p<0.05). On d21 post-hatch, cecal tonsils CD8+ T lymphocytes expression in the vaccinated-challenged group was similar to the negative control group (p>0.05) but higher than that in the positive control group (p<0.05). Finally, vaccination resulted in an increase in ileal mRNA levels of zonula occluding on d21 post-hatch. In conclusion, there were numerical but not statistically significant decrease in NE lesions and mortality in vaccinated and challenged
broilers. Further studies are needed to improve the efficacy of the vaccine and understand the
mechanism underlying protection in vaccinated birds.

Introduction

Necrotic enteritis (NE) is a disease of *Clostridium perfringens* that affects poultry gut
health. Losses are estimated to be at least 2 billion USD annually from clinical and subclinical
infections. The resurgence in infection is due to the voluntary or regulatory withdrawal in the use
of antibiotics as growth promoters [1]. Poultry NE is a multifactorial disease with a complex
etiology. However, the overgrowth of commensal or virulent *C. perfringens* strains is the
primary cause of infection [2]. Current intervention strategies include the use of feed additives,
improving biosecurity/management practices, anticoccidial drugs and vaccinations [3, 4]. There
are currently no commercial vaccines against NE in broilers. Most commercial vaccine products
are anti coccidia vaccines because coccidiosis is one of the most important predisposing
conditions for NE [5].

Two important considerations for any vaccine design strategy are antigen selection and
route of delivery. Subunit antigens are generally safer than live antigens for vaccines. Protein
subunits in the extracellular secretion of *C. perfringens* (ECP) have been explored as antigens for
candidate vaccines against NE with mixed results, with the most promising results involving
parenteral routes of administration [6, 7, 8]. Parenteral antigen delivery is neither economical nor
practical for large commercial operations. However, unencapsulated, subunit antigens such as
proteins are denatured or broken down by the low pH conditions and gut enzymes when
administered by oral gavage. Unencapsulated, subunit antigens are therefore inefficiently
delivery to inductive sites, compromising their immunogenicity and protective efficacy [9].
Therefore, there is a need to develop immunogenic, safe and efficacious vaccines based on *C.
perfringens ECP that can be administered by orally to help mitigate production losses to NE in the poultry industry.

The safety and immunogenicity of nanoparticles as encapsulating and antigen delivery vehicles have been well documented [10, 11, 12]. Chitosan has emerged as a promising candidate for effective antigen delivery [13, 14]. A previous study conducted in our laboratory indicates that chitosan nanoparticles entrapped with C. perfringens ECP and surface-tagged with Salmonella flagella proteins is safe for oral vaccination and immunogenic in broiler chickens [15]. However, immunogenicity does not always correlate with protection. Therefore, further studies are required to characterize the protective efficacy of the previously synthesized chitosan nanoparticle vaccine against NE in broiler chickens.

The specific objectives of this study were to determine the immunogenicity and protective efficacy of an oral chitosan nanoparticle vaccine against experimentally induced NE. It is hypothesized that chitosan loaded with extracellular proteins of C. perfringens and delivered orally to broilers will stimulate antigen-specific immune responses and will reduce the severity of NE by improving boiler production performance, decreasing gut lesions.

Materials and methods

Experimental animals: All animal protocols were approved by the Institutional Animal Care and Use Committee of the Southern Poultry Research Group. Birds were monitored daily by trained veterinary care staff for diarrhea, bloody feces, lethargy, ruffled feathers, refusal to eat and dehydration in the course of the experiment. Birds that could not move or refused to eat were immediately humanely euthanized by carbon dioxide. A total of 2 animals reached the humane
end point during the course of this study. All remaining birds were also humanely euthanized by carbon dioxide on the last day of sampling (day 28).

A total of 144 one-day-old male broiler chicks obtained from a commercial hatchery were raised in Petersime battery cages for 28 days at the Southern Poultry Research facility (Athens, GA). Chicks were weighed by pen and then randomly distributed to one of three treatments groups in a completely randomized design: Negative control group consisted of chicks that were not vaccinated and not challenged. The vaccinated-challenge group consisted of chicks that were both vaccinated and challenged. The positive control group consisted of chicks that were mock vaccinated and challenged. A cage was treated as a replicate. Each treatment was replicated in six cages (n = 6) of eight chicks per cage. Birds were raised under standard management practices and had *ad libitum* access to food and water. Feed intake, body weight and mortality were measured weekly from day of hatch. Mortality figures were compiled from birds that met the criteria for euthanasia as well as birds that died suddenly without meeting the criteria for euthanasia, after confirming NE by necropsy.

*Preparation and administration of vaccine, necrotic enteritis challenge and lesion scoring:* A field strain of *C. perfringens* (a gift from Dr. C. Hofacre, Southern Poultry Research Group) was obtained. Chitosan nanoparticle vaccine was synthesized with native extracellular proteins of a virulent field strain of *C. perfringens* and surface tagged with flagella proteins according to methods detailed in a previous publication [15]. Synthesized nanoparticles were resuspended to 0.1 mg/ml loaded protein in PBS at pH 7.2. On d 0 (day of hatch), d 4 and d 14 post-hatch, each bird in the vaccinated-challenged and positive control group was orally gavage with 50 μg loaded protein in nanoparticle suspension in 0.5 ml PBS and 0.5 ml PBS only, respectively.
Each bird in the vaccinated-challenged and positive control group was infected by oral gavage with 5,000 oocysts of *E. maxima* on d 14 post-hatch and then orally gavage with $1.0 \times 10^8$ colony forming units of the same isolate of *C. perfringens* used to prepare the vaccine on d 19, 20 and 21 post-hatches.

Three randomly selected birds per pen were humanely euthanized by cervical dislocation and scored for NE lesions on d 21 post-hatch. The scoring system was based on a score of 0-3 as follows: 0 for a normal intestine, 1 for the presence of a slight mucus covering and loss of tone, 2 for severe NE and 3 for severe NE with the presence of blood in the lumen.

**Ex vivo recall response of splenic and cecal tonsil mononuclear cells of chickens**

Cecal tonsil and spleen samples were collected from one bird per cage on d 18, and two birds per cage on d 21 and d 28 post-hatch after cervical dislocation. Single cell suspensions of cecal tonsils and spleen mononuclear cells were obtained as described earlier [16].

Approximately $5 \times 10^4$ mononuclear cells/well were plated in duplicates per sample in 100µl of RPMI-1640 culture media (Sigma Aldrich, St. Louis, MO) supplemented with 10 % fetal bovine serum and 1 % Penicillin and Streptomycin. 100µl of RPMI-1640 culture media only (negative control) or 25µg, or 50 µg or 100µg of native ECP or 0.1 µg of Concanavin A (Con A) in 100 µl of RPMI-1640 culture media was added to each well and incubated for 5 d at 37 °C in the presence of 5 % CO$_2$. Lymphocyte proliferation was measured using [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide] (MTT) assay as described earlier [13]. Values are reported as Optical density measured at 570 nm using a spectrophotometer.
**CD4+ and CD8+ cell percentages in cecal tonsils of vaccinated chickens**

On d 18, 21 and 28 post-hatches, single cell suspensions of cecal tonsils and spleen mononuclear cells were obtained as described earlier [16]. The cell suspensions were concentrated for lymphocytes by density centrifugation over 1.077 g/mL histopaque (Sigma–Aldrich, St. Louis, MO). For CD4+ and CD8+ T cell analysis, single-cell suspensions of the cecal tonsils (1 × 10^6 cells) were incubated with FITC-conjugated mouse anti-chicken CD4, PE-conjugated mouse anti-chicken CD8 (Southern Biotech, Birmingham, AL) at 1:200 dilution, and unlabeled mouse IgG at 1:200 dilution in a 96-well plate for 20 minutes. After incubation, cells were washed twice by centrifugation at 400 x g for 5 minutes using wash buffer comprising 2 mM EDTA and 1.5% FBS in 1× PBS to remove unbound antibodies. After washing, cells were analyzed using cytosoft software (Guava EasyCyte, Millipore, Billerica, MA). The percentage of live CD4+ and CD8+ T-cells was analyzed after gating on live cells based on forward-scatter and side-scatter.

**Nitrite production from adherent splenic and cecal tonsil mononuclear cells of chickens**

On d 18 and d 28 post-hatches, single cell suspensions of cecal tonsils and spleen mononuclear cells were obtained as described earlier [16]. Nitrite production from adherent cells was carried out according to a previous method [17] with some modifications. Briefly, mononuclear cells were seeded at 1 × 10^6 splenocytes/well in 96-well plates incubated for 24 hours at 37 °C in the presence of 5 % CO₂, to allow them attach. Unattached cells were removed with media and replaced with RPMI freshly prepared as stated above. Adherent cells were stimulated with approximately 0 µg (200 µl RPMI-1640 culture media only) or 25 µg, 50 µg or 100 µg of ECP or 0.1 µg of *Salmonella Enteritidis* lipopolysaccharide (LPS) in 200 µl of RPMI-1640 culture media per well and incubated for 5 d at 37 °C in the presence of 5 % CO₂. Samples
were centrifuged at 630 X g for 5 min at 10°C and 100 μL of the supernatants were removed. The nitrite content of the supernatant was determined using a sulfanilamide/N-(1-naphthyl) ethylenediamine dihydrochloride solution (Ricca Chemical Co., Arlington, TX) following the manufacturer’s instructions. Nitrite concentrations were determined from a standard curve with sodium nitrite standards.

Anti-ECP- and anti-flagellar-specific IgG and IgA antibodies in serum and bile of chickens

Serum from the and bile were collected from one bird per cage on d 18, and 2 birds per cage on d 21 and d 28 post-hatch. The amounts of anti-ECP- and anti-flagellar- specific IgG and IgA antibodies in serum and bile were determined by ELISA as previously described [11] with modifications. Native ECP was coated at 10 µg/ml (IgA) or 20µg/ml (IgG) on ELISA plates (Nunc Maxisorp™, ThermoFisher Scientific, Waltham, MA). Bile was diluted to 1:200 and serum was diluted to 1:20 in PBS containing 2.5 %, non-fat dry milk and 0.1% Tween 20 (VWR, Radnor, PA). Horseradish peroxidase (HRP) conjugated polyclonal goat anti-chicken IgG (Bethyl, Montgomery, TX) at 1:20,000 dilution or HRP-conjugated polyclonal goat anti-chicken IgA (SouthernBiotech, Birmingham, AL) at 1:10,000 was used as a secondary antibody. Optical density was measured as absorbance at 450nm using a spectrophotometer (Biochek, Scarborough, ME).

Effect of serum antibodies from chickens on ECP neutralization

Liver male hepatoma cells (LMH) were seeded in 96-well plates at 4 x 10⁴ cells per well for 30 minutes in 100µl of Dulbecco’s modified eagle medium (DMEM). The DMEM was supplemented with 10% Fetal bovine serum and 1% Penicillin and Streptomycin. Antigen-
antibody reaction was prepared in duplicates by incubating 15µl of \textit{C. perfringens} supernatant containing 825µg protein with 15µl serum from each bird (n=6) for 60 minutes. The antigen-antibody mixture was added to the growing LMH culture. Cells were incubated in 5 % CO\textsubscript{2} at 37°C for 8 hours. LMH cytotoxicity was measured according to the protocol outlined by OPS diagnostics (OPS diagnostics Lebanon, NJ) to measure the release of lactate dehydrogenase (LDH) from dying cells. Briefly, 50µl of cell-free supernatant of LMH culture was incubated with 50µl of 50mM Lithium Lactate and 50µl of a mixture composed of 100µl of 9mg/ml Phenazine methosulfate (PMS), 100µl of 33mg/ml iodonitrotetrazolium 2-(4-iodophenyl)-3-(4-nitrophenyl)-5-phenyl-2H-tetrazolium) (INT) and 2.3ml of 3.7mg/ml Nicotinamide adenine dinucleotide (NAD) solution. Color development was allowed for at least 5 minutes. Absorbance was measured at 490nm using a spectrophotometer (Biochek, Scarborough, ME).

\textit{Effect of vaccination and challenge on the cecal tonsil mRNA levels of tight junction proteins}

Total RNA from jejunum and ileum tonsils were collected from 2 birds per cage on d 21 and d 28 post-hatch. The mRNA was reverse transcribed into cDNA using methods described earlier [18]. The mRNA was then analyzed for the tight junction proteins claudin-2 and zonula occluden-1 by real-time PCR using SyBr green, after normalizing for β-actin mRNA. Primer sequences and annealing temperatures are described in Table 1. The fold change with respect to the reference gene was calculated using the method $2^{(\text{Ct Sample} - \text{Housekeeping})/(\text{Ct Reference} - \text{Housekeeping})}$ with Ct being the threshold cycle [19]. The Ct value was determined by CFX Maestro software when the fluorescence rises exponentially 2-fold above background (Biorad, Hercules, CA.). The reference group was the negative control birds.
Table 1. Primers and PCR Conditions for RT-qPCR.

| Target gene | Sequence          | Annealing temperature | Reference |
|-------------|-------------------|-----------------------|-----------|
| CLAU-2-F    | CCTGCTCACCCCTCATTGGAG | 57.5°C               | [20]      |
| CLAU-2-R    | GCTGAACCTCACTCTTGGCT |                      |           |
| ZO-1-F      | TGTAGCCACAGCAAGAGGTG | 57.4°C               | [21]      |
| ZO-1-R      | CTGGAATGGCTCCTTGTGGT |                      |           |
| β-actin-F   | ACCGGACTGTTACCAACACC | 57.0°C               | [22]      |
| β-actin-R   | GACTGCTGCTGACACCTTCA |                      |           |

Statistical Analysis: All statistical analyses were carried out with statistical software (SAS, v. 9.4, SAS Institute Inc., Cary, NC, USA). Analysis of body weight gain (WG), feed conversion ratio (FCR), feed intake (FI), toxin neutralization, nitrite production, antibody and T lymphocyte responses was carried out using a one-way analysis of variance (ANOVA) and pre-planned orthogonal contrasts to compare negative control vs vaccinated-challenged, and vaccinated-challenged vs negative control. Analysis of mortality and gut lesions was carried using Kruskal-Wallis chi-square test. Significance was determined at P < 0.05 and/or at P < 0.01. P values less than 0.1 were determined as approaching significance.

Results

**Effect of vaccination and challenge on mortality and jejunum lesions of broiler birds.**

Birds in the vaccinated-challenged group had comparable mortality to that in the negative control group, from d 14 to d 21 (one-week post challenge) and d 14 to d 28 post-hatch (two weeks
post challenge), but had lower mortality than that in the positive control group which approached significance (P = 0.07, Table 1).

Birds in the vaccinated-challenged group had higher lesion scores (Table 2) than that in the negative control group but had numerically lower lesion scores than that in the positive control group which approached significance (P = 0.05, Table 2) on d 21 post-hatch.

**Effect of vaccination and challenge on feed intake, average body weight gain and feed conversion ratio of broiler birds**

All three treatment groups had comparable FI, WG and FCR from d 0 to d 14 post-hatch (Table 3). However, the vaccinated-challenged group had numerically lower feed intake than that in the negative control group which approached significance (P = 0.08, Table 3), but had comparable feed intake to the positive control group from d 14 to d 21 post-hatch (Table 3).

The vaccinated-challenged group also had comparable FI to the negative control group (Table 3) but had statistically significantly higher FI than that in the positive control group from d 14 to d 28 post-hatch (p < 0.05, Table 3).

The vaccinated-challenged group had comparable FCR to the negative control group (Table 3) but had statistically significantly lower FCR than that in the positive control group from d 14 to d 21 post-hatch and from d 14 to d 28 post-hatch (p < 0.01, Table 3).

**Effect of vaccination and challenge on the proliferation of splenic PBMCs of broilers birds**

Splenic mononuclear cells obtained from all 3 treatment groups on d 18 post-hatch and stimulated with 0 µg ECP, 50 µg ECP and 0.1 µg Con A had comparable proliferation (Fig 1A).

Splenic mononuclear cells obtained from the vaccinated-challenged group on d 18 post-hatch
and stimulated with 25 μg ECP had comparable proliferation to that in the negative control group (Fig 1A) but had lower proliferation than that in the positive control birds approaching significance (p = 0.09, Fig 1A). Splenic mononuclear cells obtained from the vaccinated-challenged group on d 18 post-hatch and stimulated with 100μg ECP had comparable proliferation to that in the negative control group (Fig 1A), but statistically significantly lower proliferation than that in the positive control group (p < 0.01, Fig 1A).

Splenic mononuclear cells obtained from all 3 treatment groups on d 21 post-hatch, and stimulated with 0μg ECP, 50μg ECP, 100μg ECP, and 0.1μg Con A had comparable proliferation (Fig 1B). Splenic mononuclear cells obtained from the vaccinated-challenged group on d 21 post-hatch and stimulated with 25μg ECP had statistically significantly higher proliferation than that in the negative control and positive control groups (p < 0.05, Fig 1B).

Splenic mononuclear cells obtained from all 3 treatment groups on d 28 post-hatch and stimulated with 0μg ECP, 25μg ECP, and 50μg ECP had comparable proliferation (Fig 1C). Splenic mononuclear cells obtained from chickens in the vaccinated-challenged group on d 28 post-hatch and stimulated with 100μg ECP had statistically significantly lower proliferation than that in the negative control group (p < 0.05, Fig 1C) but had comparable proliferation to the positive control group (Fig 1C). Splenic mononuclear cells obtained from chickens in the vaccinated-challenged group on d 28 post-hatch, and stimulated with 0.1μg ConA had comparable proliferation to that in the negative control group but had statistically significantly lower proliferation than that in the positive control group (p < 0.05, Fig 1C).
Fig 1. Effect of broiler vaccination and challenge on the proliferation of splenocyte mononuclear cells. Broiler chickens were orally vaccinated on day 0, day 4 and day 14 post-hatch with Chitosan-nanoparticles entrapped with *C. perfringens* ECP and *Salmonella* Enteritidis flagellar proteins, and challenged with *Eimeria* on day 14 and *C. perfringens* on day 19, 20, and 21 post-hatch. (A) On day 18, (B) day 21 and (C) day 28 post-hatch, splenic mononuclear cells were stimulated with 0.05, 0.1, 0.25 or 0.5 μg ECP for 5 days. Lymphocyte proliferation was measured using MTT assay and values reported as Optical Density (OD) values. Mean ± SEM of 6 replicates (n = 6). P < 0.05.

Effect of broiler vaccination and challenge on the percentages of splenic and cecal tonsils

*CD4*, *CD8*+ cells, double positive *CD4*+ - *CD8*+ cells and *CD4*+:*CD8*+ ratio

Splenic mononuclear cells obtained from all treatment groups on d 18, 21 and 28 post-hatch had comparable proportion of *CD4*+ cells (Fig 2A). Cecal tonsil mononuclear cells obtained from all treatment groups on d 18 and 28 post hatch also had comparable proportion of *CD4*+ cells (Fig 2A). Cecal tonsil mononuclear cells obtained from the vaccinated-challenge group on d 21 post-hatch had a comparable proportion of *CD4*+ cells to that in the negative control group (Fig 2A), but had a lower proportion of *CD4*+ cells than that in the positive control group, approaching statistical significance (p = 0.07, Fig 2A).

Splenic mononuclear cells obtained from all treatment groups on d 18 and 28 post-hatch had a comparable proportion of *CD8*+ cells (Fig 2B). Splenic mononuclear cells obtained from the vaccinated-challenged group on d 21 post-hatch had a numerically lower proportion of *CD8*+ cells than that in the negative control group approaching statistical significance (p = 0.06, Fig 2B) but had a comparable proportion of *CD8*+ cells to that in the positive control group (Fig 2B).
Cecal tonsil mononuclear cells obtained from the vaccinated-challenge group on d 18 post-hatch had a comparable proportion of CD8\(^+\) cells to that in the negative control group (Fig 2B), but had a statistically significantly lower proportion of CD8\(^+\) cells than that in the positive control group (p < 0.01, Fig 2B). Cecal tonsil mononuclear cells obtained from the vaccinated-challenge group on d 21 post-hatch had a comparable proportion of CD8\(^+\) cells to that in the negative control group but had a numerically lower proportion of CD8\(^+\) cells than that in the positive control group, approaching statistical significance (p = 0.07, Fig 2B). Cecal tonsil mononuclear cells obtained from all treatment groups on d 28 post-hatch had comparable percentages of CD8\(^+\) cells (Fig 2B).

Splenic mononuclear cells obtained from all treatment groups on d 18, 21 and 28 post-hatch had comparable percentages of double positive CD4\(^+\)CD8\(^+\) cells (Fig 2C). Cecal tonsil mononuclear cells obtained from the vaccinated-challenge group on d 18 post-hatch had a comparable proportion of double positive CD4\(^+\)CD8\(^+\) cells to that in the negative control group but had a statistically significantly lower proportion of double positive CD4\(^+\)CD8\(^+\) cells than that in the positive control group (p < 0.05, Fig 2C). Cecal tonsil mononuclear cells obtained from the vaccinated-challenge group on d 21 post-hatch had a comparable proportion of double positive CD4\(^+\)CD8\(^+\) cells to that in the negative control group but had a numerically lower proportion of double positive CD4\(^+\)CD8\(^+\) cells than that in the positive control group, approaching statistical significance (p = 0.07, Fig 2C). Cecal tonsil mononuclear cells obtained from all treatment groups on d 28 post-hatch had comparable proportion of double positive CD4\(^+\)CD8\(^+\) cells (Fig 2C).

Splenic mononuclear cells obtained from the vaccinated-challenged group on d 18 post-hatch had a numerically higher CD4\(^+\):CD8\(^+\) ratio than that in the negative control group.
approaching significance (p = 0.05, Fig 2D) but had a comparable CD4+:CD8+ ratio to that in positive control group (Fig 2D). Splenic mononuclear cells obtained from all treatment groups on d 21 and d 28 post-hatch had comparable CD4+:CD8+ ratios (Fig 2D). Cecal tonsil mononuclear cells obtained from all three treatment groups on d 18 and 21 post-hatch had comparable CD4+:CD8+ ratios (Fig 2D). Cecal tonsil mononuclear cells obtained from the vaccinated-challenge group on d 28 post-hatch had comparable CD4+:CD8+ ratio to that in the negative control group but had a statistically significantly lower CD4+:CD8+ ratio than that in the positive control group (p < 0.01, Fig 2D).

**Fig 2. Effect of broiler vaccination and challenge on their CD4+ and CD8+ cell populations.**

Broiler chickens were orally vaccinated on d 0, 3 and 14 post-hatch with Chitosan-nanoparticles entrapped with *C. perfringens* ECP and *Salmonella* Enteritidis flagellar proteins, and challenged with *Eimeria* on day 14 and *C. perfringens* on day 19, 20, and 21 post-hatch.

(A) Percentages of live cells that are CD4+. (B) Percentages of live cells that are CD8+. (C) Percentage of live cells that double positive CD4+ and CD8+. (D) The ratio of CD4+:CD8+ cells. Mean ± SEM.

n = 6 replicates. P < 0.05.

**Effect of vaccination and challenge on nitrite production from cecal tonsil mononuclear cells of broiler birds on day 18 and 28 post-hatch**

Cecal tonsil mononuclear cells obtained from chickens in all 3 treatment groups on d 18 post-hatch and stimulated with 0μg ECP, 25μg ECP, 50μg ECP, and 100μg ECP had comparable nitrite production (Fig 3A). Cecal tonsil mononuclear cells obtained from the vaccinated-challenged group on d 18 post-hatch, and stimulated with 0.1μg LPS had comparable nitrite
production to that in the negative control group (Fig 3A), but had statistically significantly lower nitrite production than that in the positive control group (p < 0.05, Fig 3A).

Cecal tonsil mononuclear cells obtained from all 3 treatment groups on d 28 post-hatch and stimulated with 0µg ECP, 100µg ECP and 0.1µg LPS had comparable proliferation (Fig 3B). Cecal tonsil mononuclear cells obtained from chickens in the vaccinated-challenged group on d 28 post-hatch and stimulated with 25µg ECP had statistically significantly lower nitrite production than that in the negative control group (p < 0.05, Fig 3B), but had comparable nitrite production to that in the positive control group (Fig 3B). Cecal tonsil mononuclear cells obtained from chickens in the vaccinated-challenged group on d 28 post-hatch, and stimulated with 50µg ECP had comparable nitrite production to that in the negative control group (Fig 3B) but had statistically significantly higher nitrite production than that in the positive control group (p < 0.05, Fig 3B).

Fig 3. Effect of broiler vaccination and challenge on nitrite production from adherent cecal tonsil mononuclear cells. Broiler chickens were orally vaccinated on day 0, day 4 and day 14 post-hatch with Chitosan-nanoparticles entrapped with *C. perfringens* ECP and *Salmonella Enteritidis* flagellar proteins, and challenged with *Eimeria* on day 14 and *C. perfringens* on day 19, 20, and 21 post-hatches. On d 18 (Panel A), and d 28 (Panel B) nitrite production was measured using sulfanilamide/N-(1-naphthyl) ethylenediamine dihydrochloride solution. Nitrite concentrations were determined from a standard curve with sodium nitrite standards. Mean ± SEM of 6 replicates (n = 6). P < 0.05.
**Effect of vaccination and challenge on nitrite production from splenic mononuclear cells of broiler birds on d 18 and d 28 post-hatch**

Splenic mononuclear cells obtained from all 3 treatment groups on d 18 post-hatch and stimulated with 0μg ECP, 50μg ECP, 100μg ECP, and 0.1μg LPS had comparable nitrite production (Fig 4A).

Splenic mononuclear cells obtained from all 3 treatment groups and stimulated with 0μg ECP, 50μg ECP, 100μg ECP, and 0.1μg LPS had comparable proliferation (Fig 4B). Splenic mononuclear cells obtained from chickens in the vaccinated-challenged group on d 28 post-hatch and stimulated with 25μg ECP had comparable nitrite production to that in the negative control group (Fig 4B), but had statistically significantly lower nitrite production than that in the positive control group (p < 0.05, Fig 4B).

**Fig 4. Effect of broiler vaccination and challenge on nitrite production from splenocyte mononuclear cells.** Broiler chickens were orally vaccinated on day 0, day 4 and day 14 post-hatch with Chitosan-nanoparticles entrapped with *C. perfringens* ECP and *Salmonella* Enteritidis flagellar proteins, and challenged with *Eimeria* on day 14 and *C. perfringens* on day 19, 20, and 21 post-hatches. On d 18 and d 28 post-hatch, splenocyte mononuclear cells were stimulated with 0, 0.05, 0.1, 0.25 or 0.5 μg ECP for 48 hrs. (A) on day 18 and (B) on day 28 post-hatch, nitrite production was measured using sulfanilamide/N-(1-naphthyl) ethylenediamine dihydrochloride solution. Nitrite concentrations were determined from a standard curve with sodium nitrite standards. Mean ± SEM of 6 replicates. P < 0.05.
Effect of vaccination on anti-ECP-specific serum IgG and bile IgA

Serum and bile obtained on d 18, 21, and 28 post-hatches from all treatments had comparable anti-ECP antibodies in (Fig 5).

Fig 5. Anti-ECP-specific serum IgG and bile IgA. Broiler chickens were orally vaccinated on day 0, day 4 and day 14 post-hatch with Chitosan-nanoparticles entrapped with C. perfringens ECP and Salmonella Enteritidis flagellar proteins, and challenged with Eimeria on day 14 and C. perfringens on day 19, 20, and 21 post-hatch. Indirect ELISA was used to measure anti-ECP specific antibodies in serum and bile. Mean ± SEM of 6 replicates. P < 0.05.

Effect of vaccination and challenge on ECP toxin neutralization by serum and bile antibodies

Serum obtained from all 3 treatments groups on d 18 and 28 post-hatch, and incubated with ECP and LMH cells had comparable LDH release (Fig 6A). Serum obtained from the vaccinated-challenged group on d 21 post-hatch and incubated with ECP and LMH cells had lower LDH release than that in the negative control group approaching statistical significance (p = 0.08, Fig 6A) but had statistically significantly higher LDH release than that in the positive control group (p < 0.05, Fig 6A).

Bile obtained from the vaccinated-challenged group on d 18 post-hatch and incubated with ECP and LMH cells had comparable LDH release to that in the negative control group but had higher LDH release than that in the positive control group, approaching statistical significance (p = 0.07, Fig 6B). Bile obtained from all three treatment groups on d 21 post-hatch and incubated with ECP and LMH cells had comparable LDH release (Fig 6B). Bile obtained from the vaccinated-challenged group on d 28 post-hatch and incubated with ECP and LMH cells had
comparable LDH release to that in the negative control group but had statistically significantly lower LDH release than that in the positive control group (p < 0.01, Fig 6B).

**Fig 6. Effect of broiler vaccination and challenge on ECP toxin neutralization.** Broiler chickens were orally vaccinated on day 0, day 4 and day 14 post-hatch with Chitosan-nanoparticles entrapped with *C. perfringens* ECP and *Salmonella* Enteritidis flagellar proteins, and challenged with *Eimeria* on day 14 and *C. perfringens* on day 19, 20, and 21 post-hatch. (A) serum and (B) bile were collected on day 18, 21, and d 28 post-hatch and incubated with 825 µg ECP to neutralize ECP. LMH cells were incubated with the above neutralizing solution for 8 hours. Cytotoxicity was measured by LDH release and values reported as Optical Density (OD) values. Mean ± SEM. n = 4 replicates. P < 0.05.

**Effect of vaccination and challenge on the jejunum and ileum mRNA levels of claudin-2, zonula occluden-1, muc-2**

Jejunum samples obtained from the vaccinated-challenged group on d 21 post-hatch had statistically significantly higher zonula occluden-1 mRNA levels than that in the negative control and positive control group (P < 0.05, Fig 7). Ileum samples obtained from all treatment groups on d 21 post-hatch had comparable claudin-2, and zonula occluden-1 mRNA levels (Fig 7).

Jejunum samples obtained from all treatment groups on d 28 post-hatch had comparable claudin-2 and zonula occluden-1 (Fig 7). Ileum samples obtained from the vaccinated-challenged group on d 28 post-hatch had statistically significantly lower claudin-2 mRNA level
than that in the negative control group (P < 0.05, Fig 7) but had comparable claudin-2 to that in
the positive control group (Fig 7).

Fig 7. Effect of broiler vaccination and challenge on tight junction mRNA level. Broiler
chickens were orally vaccinated on day 0, day 4 and day 14 post-hatch with Chitosan-
nanoparticles entrapped with *C. perfringens* ECP and *Salmonella* Enteritidis flagellar proteins,
and challenged with *Eimeria* on day 14 and *C. perfringens* on day 19, 20, and 21 post-hatch
(dph). On day 21 and day 28 post-hatch, relative claudin-2 and zonula occuloden-1 mRNA
content were analyzed after correcting for β-actin mRNA and normalizing to the mRNA content
of the negative control group. Mean ± SEM. n = 6 replicates. P < 0.05.

Discussion

To the best of our knowledge, this study is the first to identify the immunogenicity and
protective efficacy of nanoparticles loaded with *C. perfringens* ECP and *Salmonella* flagella
proteins. The study was designed on the back of promising results from a previous study carried
out by our lab that demonstrated the safety and immunogenicity of the same chitosan
nanoparticle vaccine when administered by oral gavage to broilers, in the absence of an
experimental challenge [15]. The experimental NE challenge model employed was successful in
reproducing clinical NE as shown by 33% mortality and 0.72 average NE gut lesion in the
unvaccinated challenged group. Over the 2-week period post challenge, the vaccinated-
challenged group had numerically higher mortality and intestinal lesions than the negative
control group but had numerically lower mortality and intestinal lesions than positive control
birds. Vaccination also improved the feed conversion ratio and feed intake of challenged birds.
Increased mortality, feed intake, lower feed conversion, body weight gain and the presence of intestinal lesions have been associated with clinical NE in broilers [23]. Vaccination did not significantly improve body weight gain of challenged birds at the intervals examined. However, the improvement in the other performance parameters suggests that the vaccine is partially protective against NE in broilers. In support of this, Hu et al. in 2013 [24] demonstrated that nanoparticle detained toxins are immunogenic and protective in mice. In addition, Zhao et al. in 2012 [13] demonstrated that chitosan nanoparticles vaccines loaded with Newcastle disease virus antigens offered protection against an oral challenge of Newcastle disease.

It has been shown that a cell-mediated immune response is more important for immunity to Eimeria infections [25] and chitosan has shown potential as an adjuvant to drive cell-mediated immunity [26]. It is not clear why splenic mononuclear cells from positive control birds had higher recall response than splenocytes from vaccinated-challenged and negative control birds on day 18 post-hatch, although higher proliferation was found in vaccinated-challenged birds compared to controls at the peak of infection, on day 21 post-hatch. A study by Zhao et al. in 2012 [13] also found increased lymphoproliferation of splenocytes of chickens vaccinated with chitosan nanoparticles encapsulating Newcastle disease virus compared to non-vaccinated controls. ConA is a non-specific T cell mitogen from plants that can be used to measure T cell proliferation in response to vaccination [27]. However, there was no increase in T cell proliferation in vaccinated-challenged birds suggesting either a different dose of ConA may be appropriate for ex vivo stimulation or vaccination was inducing a diminished T cell response at the assay time points. Furthermore, the phenotype of proliferating T cells can variably be dominated by effector or regulatory cells [28]. Eimeria has previously been demonstrated to induce IL-10 which can increase regulatory T cells [29]. Overall, there appears to be an antigen
concentration-dependent effect on splenocyte proliferation. The decrease in proliferation of splenic mononuclear cells at 100μg ECP concentration on d 28 post-hatch may be due to the cytotoxicity of the ECP to a population of cells present at this time point [15].

The overall higher proportion of CD4+, CD8+, and double positive CD4+ CD8+ cells in the positive control birds compared to vaccinated challenge and negative control birds may explain the increased proliferation found in positive control birds discussed previously. According to Luhtala et al. in 1997 [30], unlike thymic CD4+ CD8+ double positive T-cells which are immature, peripheral CD4+ CD8+ double positive T-cells can respond to in vitro stimulation with antigens. Necrotic enteritis was also found to increase CD4+ and CD8+ and double positive CD4+ CD8+ cell population in chickens by Runhke et al. in 2017 [31]. A study by Hong et al. in 2006 [32] also demonstrated the persistence of relatively higher numbers of CD4+ T-cells during a secondary or challenge infection with *Eimeria*. The enhanced cell-mediated immune response associated with the positive control group may also be associated with increased disease pathology associated with susceptible chickens [9]. The numerical increase in the CD4:CD8 ratio on day 18 may be indicative of a Th2 response that generally results in antibody production [33] although there were no differences between treatments in the specific antibodies against ECP. There was an increase in the CD4:CD8 T cell ratio on day 28 post-hatch, in the cecal tonsils in positive control birds compared to vaccinated-challenged birds. The implication for this is not clear as the mechanisms underlying the temporal and spatial CD4 and CD8 response during NE is not yet fully understood.

Ex vivo stimulation of cecal tonsil mononuclear cells for nitrite production was carried out to assess innate priming. Vaccination resulted in an antigen-dependent modulation of nitrite production from ex-vivo stimulated cecal tonsil cells. LPS stimulated increased NO production
in the positive control birds on day 18 post-hatch in the cecal tonsils, but down regulated NO production in the cecal tonsils by day 28 post-hatch. Nitric oxide is usually induced early during an infection and then rapidly cleared; else it can have a negative impact on bird performance [34]. iNOS has been demonstrated to be upregulated in chicken intestinal cells in response to *Eimeria* infection [35] but unchanged or down-regulated in chickens when *Eimeria* was co-infected with *C. perfringens*. The positive control birds compared to vaccinated and control birds, had increased NO production from ex vivo stimulated spleen mononuclear cells on d 28 post-hatch.

Indirect ELISA carried out to detect specific antibodies against ECP in serum or bile did not detect any differences between treatments. This may be because a crude antigen was used. Metabolic enzymes such as fructose 1,6 biphosphate aldolase and glyceraldehyde-3-phosphate dehydrogenase [36, 37] which are found in commensal organisms can generate antibodies against Clostridial infections which can interfere with ELISA results. Antigen neutralization assay was therefore carried out to determine whether undetected antibodies in serum or bile can neutralize toxins in ECP, thereby protecting LMH cells from lactose dehydrogenase release. LMH cells have been demonstrated to be vulnerable to LDH release by toxins such as NetB in the supernatant of virulent *C. perfringens* [38] and the strain of *C. perfringens* used for this study is *netB* positive (data not shown). Contrary to expectations however, increased cytotoxicity or decreased neutralization was observed in the serum and bile of vaccinated-challenged birds, compared to positive control birds on d 18 and d 21 post-hatch. The reason for this trend is not immediately clear, although for this study, endogenous complement was not inactivated and exogenous complemented was not added to serum or bile. Antibody-dependent and independent complement system has been characterized in poultry such as duck [39] and demonstrated to
play a role *in vitro* in pathogen neutralization [40]. Also, non-specific complement cytotoxicity to certain cell lines is possible [41].

Quantitative PCR was carried out to determine the effect of vaccination on the mRNA levels of tight junction proteins such as claudin-2 and zonula occulodens-1. Claudin-2 is normally a pore forming protein associated with paracellular water and ion channels [42] while ZO-1 is positively associated with the formation of epithelial tight junctions by acting as tether for claudins [43]. A study by Bortoluzzi et al. in 2019 [44] showed that necrotic enteritis upregulates claudin-2 and downregulates ZO-1 at 21 days post-hatch. The decreased expression of claudin-2 in vaccinated-challenged and positive control birds compared to negative control birds on d 28 post hatch may therefore be associated with recovery. However, there was no difference between vaccinated-challenged and positive control birds claudin-2 levels. Vaccination improved the expression of ZO-1 compared to negative control and positive control birds and this may have contributed to the improved performance of vaccinated-challenged birds compared to positive control birds.

**Conclusion**

Chitosan nanoparticles loaded with ECP of *C. perfringens* and *Salmonella* flagellar proteins were immunogenic and partially protective against experimentally induce NE. The vaccine produced cell mediated and humoral immune response in the birds. In order to understand the mechanisms underlying protection, further studies are required to identify the different populations of proliferating cells in vaccinated challenged, and non vaccinated challenged birds. Further studies are also needed to understand at the proteomic level, the expression of Th1 and Th2 cytokines. The vaccine was designed with only *C. perfringens*
proteins with a homologous challenge. Further studies may be needed to determine heterologous protection, and if protection can be improved by adding *Eimeria* proteins, varying the route of administration, and refining the antigen selection.

Acknowledgements

We acknowledge the efforts of the staff of Southern Poultry Research farms at which the field study was conducted. We also appreciate the support of members of the Selvaraj lab: Dr Theros Ng, Dr Jarred Oxford, Keila Acevedo, and Ragini Reddyvary for their contributing efforts during sampling collections and laboratory analysis.
References

1. M’Sadeq SA, Wu S, Swick RA, Choct M. Towards the control of necrotic enteritis in broiler chickens with in-feed antibiotics phasing-out worldwide. Anim Nutr. 2015; 1(1): 1–11. https://doi.org/10.1016/j.aninu.2015.02.004

2. Cooper KK, Trinh HT, Songer JG. Immunization with recombinant alpha toxin partially protects broiler chicks against experimental challenge with Clostridium perfringens. Vet. Microbiol. 2009;133: 92–7.

3. Peek HW, Landman WJM. Coccidiosis in poultry: anticoccidial products, vaccines and other prevention strategies. Vet Q. 2011;31(3): 143–161. https://doi.org/10.1080/01652176.2011.605247

4. Hofacre CL, Reynolds DJ, Mathis GF, Lumpkins BS, Ollis N, Smith JA, Demey V. Effect of a Competitive Exclusion Culture in a Necrotic Enteritis Challenge Model in Broilers. J Appl Poult Res. 2019;28(2): 350–355. https://doi.org/10.3382/japr/pfy078

5. Collier CT, Hofacre CL, Payne AM, Anderson DB, Kaiser P, Mackie RI, Gaskins HR. Coccidia-induced mucogenesis promotes the onset of necrotic enteritis by supporting Clostridium perfringens growth. Vet Immunol Immunopathol. 2008; 122(1–2): 104–115. https://doi.org/10.1016/j.vetimm.2007.10.014

6. Saleh N, Fathalla, SI, Nabil R, Mosaad, AA. Clinicopathological and immunological studies on toxoids vaccine as a successful alternative in controlling clostridial infection in broilers. Anaerobe. 2011;17: 426–430.

7. Lillehoj HS, Jang SI, Panebra A, Lillehoj EP, Dupuis L, Arous JB, et al. NetB proteins in Montanide IMS adjuvant increases protective immunity against...
experimentally-induced necrotic enteritis. Asian-Australas J Anim Sci. 2017;30(10): 1478–1485.

8. Mot D, Timbermont L, Haesebrouck F, Ducatelle R, Van Immerseel, F. Progress and problems in vaccination against necrotic enteritis in broiler chickens. Avian Pathol. 2014; 43(4): 290–300. https://doi.org/10.1080/03079457.2014.939942

9. Kim, S, Jung, D, Yang, I., Jang S, Kim J, Truong TT, et al. Application of an M-cell-targeting ligand for oral vaccination induces efficient systemic and mucosal immune responses against a viral antigen. 2017;25(11): 623–632. https://doi.org/10.1093/intimm/dxt029

10. Nel A, Xia T, Madler L, Li N. Toxic potential of materials at the nano-level. Science. 2006;311: 622-627.

11. Thomas C, Rawat A, Hope-Weeks L, Ahsan F. Aerosolized PLA and PLGA Nanoparticles Enhance Humoral, Mucosal and Cytokine Responses to Hepatitis B Vaccine Mol Pharm. 2011;8: 405-415.

12. Carrillo-Conde B, Song EH., Chavez-Santoscoy A, Phanse Y, Ramer-Tait AE, Pohl NL, et al. Mannose-functionalized "pathogen-like" polyanhydride nanoparticles target C-type lectin receptors on dendritic cells. Mol. Pharmacol. 2011;8: 1877-1886.

13. Zhao K, Chen G, Shi XM, Gao TT, Li W, Zhao Y. et al. (2012). Preparation and Efficacy of a Live Newcastle Disease Virus Vaccine Encapsulated in Chitosan Nanoparticles. PLoS ONE. 2012;7(12): 1–11. https://doi.org/10.1371/journal.pone.0053314
14. Renu S, Markazi AD, Dhakal S, Lakshmanappa YS, Shanmugasundaram R,
Selvaraj RK, Renukaradhyya GJ. Oral Deliverable Mucoadhesive Chitosan-
Salmonella Subunit Nanovaccine for Layer Chickens. International journal of
nanomedicine, 2020;15: 761–777. https://doi.org/10.2147/IJN.S238445

15. Akerele G, Ramadana N, Renu S, Renukaradhyab GJ, Shanmugasundarama R,
Selvaraj RK. In vitro characterization and immunogenicity of chitosan
nanoparticles loaded with native and inactivated extracellular proteins from a field
strain of CLostridium perfringens associated with necrotic enteritis. Vet Immunol
Immunopathol. 2020;224: 110059.

16. Shanmugasundaram R, Kogut MH, Arsenault RJ, Swaggerty CL, Cole K,
Reddish JM, Selvaraj RK. Effect of Salmonella infection on cecal tonsil
regulatory T cell properties in chickens. 2015: 1828–1835.
http://dx.doi.org/10.3382/ps/pev161.

17. Acevedo-Villanueva KY, Lester B, Renu S, Han Y, Shanmugasundaram R,
Gourapura R, Selvaraj R. Efficacy of chitosan-based nanoparticle vaccine
administered to broiler birds challenged with Salmonella. PloS one. 2020;15(4):
e0231998. https://doi.org/10.1371/journal.pone.0231998

18. Selvaraj RK, Klasing KC. Lutein and eicosapentaenoic acid interact to modify
iNOS mRNA levels through the PPARγ/RXR pathway in chickens and HD11 cell
lines. J. Nutr. 2006;136: 1610-1616.

19. Schmittgen TD, Livak, KJ. Analyzing real-time PCR data by the comparative
C(T) method. Nat Protoc. 2008;3: 1101-1108.
20. Chen YP, Cheng YF, Li XH, Yang WL, Wen C, Zhuang S, Zhou YM. Effects of threonine supplementation on the growth performance, immunity, oxidative status, intestinal integrity, and barrier function of broilers at the early age. Poult Sci. 2017;96: 405-413.

21. Yi GF, Allee GL, Knight CD, Dibner JJ. Impact of glutamine and Oasis hatchling supplement on growth performance, small intestinal morphology, and immune response of broilers vaccinated and challenged with *Eimeria maxima*. Poult Sci. 2005;84: 283-293.

22. Shanmugasundaram R, Selvaraj RK. Effect of killed whole yeast cell prebiotic supplementation on broiler performance and intestinal immune cell parameters. Poult. Sci. 2012;91: 107-111

23. Latorre JD, Adhikari B, Park SH, Teague KD, Graham LE, Mahaffey BD, et al. Evaluation of the epithelial barrier function and ileal microbiome in an established necrotic enteritis challenge model in broiler chickens. Front Vet Sci. 2018;5(AUG): 1–11. https://doi.org/10.3389/fvets.2018.00199

24. Hu, CMJ, Fang RH, Luk BT, Zhang L. Nanoparticle-detained toxins for safe and effective vaccination. Nat Nanotechnol. 2013. https://doi.org/10.1038/nnano.2013.254

25. Rose ME, Davison TF, Morris TR, Payne LN. Immunity to coccidia Poultry immunology. Darfax Publishing Company, Abington. 1996.

26. Mori A, Oleszycka E, Sharp FA, Coleman M, Ozasa Y, Singh M, O'Hagan DT, et al. The vaccine adjuvant alum inhibits IL-12 by promoting PI3 kinase signaling while chitosan does not inhibit IL-12 and enhances Th1 and Th17
27. Linghua Z, Xingshan T, Fengzhen Z. Vaccination with Newcastle disease vaccine and CpG oligodeoxynucleotides induces specific immunity and protection against Newcastle disease virus in SPF chicken. Vet. Immunol. Immunopathol. 2007;115(3–4): 216–222. https://doi.org/10.1016/j.vetimm.2006.10.017

28. Yuan F, Li Z. Induction of immunological tolerance in chickens inoculated with xenogeneic antigens at a late stage of embryonic development. Int Immunol. 2012;24(4): 267–272. https://doi.org/10.1093/intimm/dxr112

29. Rothwell L, Young JR, Zoorob R, Whittaker CA, Hesketh P, Archer A, et al. Kaiser, P. Cloning and Characterization of Chicken IL-10 and Its Role in the Immune Response to *Eimeria maxima*. J Immunol. 2004;173(4): 2675–2682. https://doi.org/10.4049/jimmunol.173.4.2675

30. Luhtala M, Lassila O, Toivanen P, Vainio O. A novel peripheral CD4+ CD8+ T cell population: inheritance of CD8+ alpha expression on CD4+ T cells. Eur. J. Immunol. 1997;27: 189-193

31. Ruhnke I, Andronicos NM, Swick RA, Hine B, Sharma N, Kheravii SK, Wu S, Hunt P. Immune responses following experimental infection with *Ascaridia galli* and necrotic enteritis in broiler chickens. Av Pathol. 2017;46(6): 602-609.

32. Hong YH, Lillehoj H., Lillehoj EP, Lee SH. Changes in immune-related gene expression and intestinal lymphocyte subpopulations following *Eimeria maxima* infection of chickens. Vet Immunol Immunopathol. 2006;114: 259–272.
33. Alam, SI, Dwivedi P. Putative function of hypothetical proteins expressed by *Clostridium perfringens* type A strains and their protective efficacy in mouse model. Infect Genet Evol. 2016;44: 147–156.

https://doi.org/10.1016/j.meegid.2016.06.040

34. Lillehoj HS, Lee SH, Park SS, Jeong M, Lim Y, Mathis GF, et al. Calcium Montmorillonite-Based Dietary Supplement Attenuates Necrotic Enteritis Induced by *Eimeria* maxima and *Clostridium perfringens* in Broilers. Av dis. 2016;44(2): 408-425

35. Allen PC. Nitric oxide production during *Eimeria* tenella infections in chickens. Poult Sci. 1997;76(6): 810-3. doi: 10.1093/ps/76.6.810. PMID: 9181612.

36. Kulkarni R.R, Parreira VR, Sharif S, Prescott JF. *Clostridium perfringens* antigens recognized by broiler chickens immune to necrotic enteritis. Clin Vaccine Immunol. 2006;13(12): 1358-62.

37. Matsunaga N, Shimizu H, Fujimoto K, Watanabe K, Yamasaki T, Hatano N, et al. Expression of glyceraldehyde-3-phosphate dehydrogenase on the surface of *Clostridium perfringens* cells. Anaerobe. 2018;51: 124-130

38. Keyburn AL, Boyce JD, Vaz P, Bannam TL, Ford ME, Parker D, et al. NetB, a new toxin that is associated with avian necrotic enteritis caused by *Clostridium perfringens*. PLoS Pathog. 2008;4(2).

https://doi.org/10.1371/journal.ppat.0040026

39. Koppenheffer TL, Chan SWS, Higgins DA. The complement system of the duck. Av Pathol. 1999;28(1): 17–25. https://doi.org/10.1080/03079459995000
40. Sahin O, Luo N, Huang S, Zhang Q. Effect of Campylobacter-specific maternal antibodies on *Campylobacter jejuni* colonization in young chickens. J Appl Environ Microbiol. 2003;69(9): 5372–5379. https://doi.org/10.1128/AEM.69.9.5372-5379.2003

41. Sugimoto C, Kodama H, Mikami T. Complement-dependent antibody cytotoxicity test of chicken antibody with duck complement used against T-cells of a Marek's disease lymphoma-derived cell line (MSB-1). Av dis. 1979;23(1): 229–234.

42. Van Itallie C, Rahner C, Anderson JM. Regulated expression of Claudin-4 decreases paracellular conductance through a selective decrease in sodium permeability. J. Clin. Invest. 2001; 107:1319–1327. doi: 10.1172/JCI12464.

43. Stevenson BR, Siliciano JD, Mooseker MS, Goodanough DA. Identification of ZO-1: A high molecular weight polypeptide associated with tight junction (zonula occludens) in a variety of epithelia. J. Cell Biol. 1986; 103:755–766. doi: 10.1083/jcb.103.3.755.

44. Bortoluzzi C, Lumpkins B, Mathis GF, França M, King WD, Graugnard DE, Dawson KA, Applegate TJ. Zinc source modulates intestinal inflammation and intestinal integrity of broiler chickens challenged with coccidia and *Clostridium perfringens*. Poult. Sc. 2019;98: 2211-2219.
Figure
Figure
A
- Negative control
- Vaccinated + Challenged
- Positive control

B

NO production (µM)

0µg antigen  25µg antigen  50µg antigen  100µg antigen  0.1µg LPS

Stimulation

< 0.01

0µg antigen  25µg antigen  50µg antigen  100µg antigen  0.1µg LPS

Stimulation
Figure
