Effects of Dietary Additives and Early Feeding on Performance, Gut Development and Immune Status of Broiler Chickens Challenged with Clostridium perfringens

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ABSTRACT : The effects of dietary additives and holding time on resistance and resilience of broiler chickens to Clostridium perfringens challenge were investigated by offering four dietary treatments. These were a negative control (basal), a positive control (Zn-bacitracin) and two dietary additives, mannanoligosaccharides (MOS), and acidifier. Two holding times included (a) immediate access to feed and water post hatch (FED) and (b) access to both feed and water 48 h post hatch (HELD). Chicks fed Zn-bacitracin had no intestinal lesions attributed to necrotic enteritis (NE), whereas chicks fed both MOS or acidifier showed signs of NE related lesions. All dietary treatments were effective in reducing the numbers of C. perfringens in the ileum post challenge. The FED chicks had heavier body weight and numerically lower mortality. The FED chicks also showed stronger immune responses to NE challenge, showing enhanced (p<0.05) proliferation of T-cells. Early feeding of the MOS supplemented diet increased (p<0.05) IL-6 production. The relative bursa weight of the FED chicks was heavier at d 21 (p<0.05). All the additives increased the relative spleen weight of the HELD chicks at d 14 (p<0.05). The FED chicks had increased villus height and reduced crypt depth, and hence an increased villus/crypt ratio, especially in the jejunum at d 14 (p<0.05). The same was true for the HELD chicks given dietary additives (p<0.05). It may be concluded that the chicks with early access to dietary additives showed enhanced immune response and gut development, under C. perfringens challenge. The findings of this study shed light on managerial and nutritional strategies that could be used to prevent NE in the broiler industry without the use of in-feed antibiotics. (Key Words : Necrotic Enteritis, Early Feeding, Gut Morphology, Alternative to Antibiotics, Mannanoligosaccharides (MOS))

INTRODUCTION

Necrotic enteritis (NE) is a costly poultry disease caused by the bacterium Clostridium perfringens (Parish, 1961; Hofacre, 2001). The C. perfringens is found in the intestine of healthy chickens and the environment (Ficken and Wages, 1997), and it only causes NE when it grows unchecked and produces toxins (Al-Sheikhly and Truscott, 1977). Dietary and husbandry factors contribute to the outbreak of this disease, including a high level of viscous cereal grains (for example wheat, barley and oat) and animal protein (i.e., fish meal and meat meal) in the diets, and factors that cause damage to the intestinal mucosa (i.e., coccidiosis) (Branton et al., 1987; Kaldhusdal and Skjerve, 1996; Kaldhusdal, 2000). Stress factors, such as stocking density may also contribute the pathogenesis of the disease (McDevitt et al., 2006). However, the single most important factor believed to be responsible for the disease is a toxin, netB, produced by some isolates of C. perfringens (Keyburn et al., 2006).

Traditionally, antibiotics such as bacitracin, virginiamycin, avoparcin, lincomycin, tylosin, and penicillin, have been used as a successful means to control NE in broilers (Ficken and Wages, 1997; Williams, 2005). However, there is a worldwide trend that more and more chickens will be produced under an antibiotic-free environment in the future. Such a demand will require not only nutriceutical alternatives (organic acids, enzymes, probiotics, prebiotics, nucleotides, functional carbohydrates, betaine, spice and plant extracts, and so on) but also novel managerial and environmental measures. A hygienic environment will prevent the exposure of animals to...
potential pathogens, thus alleviating the pressure on their immune system. During the last two decades, early nutrition has been studied intensively. Early access to nutrients can improve the immunity and enhance the health status of chicks (Dibner et al., 1998). However, in practice, newly hatched chicks are held for a long time, sometimes more than 48 h without access to feed and water, due to processing at hatchery and transportation to farms (Noy and Sklan, 1999). Under some circumstances, chicks are held without access to nutrients deliberately in an attempt to minimise possible bacterial contamination of incompletely healed navel of young chicks and initiate a vaccine response (Dibner et al., 1998). Early feeding, either given access to feed and water to newly hatched birds, or injected in ovo has been demonstrated to be beneficial (Hornasio et al., 2011). When antibiotics are not included in the diet, a sound immune competence is of paramount importance in maintaining the health of animals. Chicken cytokines have been considered as another candidate for replacing in-feed antibiotics (Lowenthal et al., 1994) However, it is not known whether the use of feed additives either under normal feeding practices or in an early feeding regime will impact on IL-6 production.

This study investigated the effects of early nutrition and dietary additives, used solely or in combination, on the performance, immune competence and gut morphology in broilers under NE challenge.

**MATERIALS AND METHODS**

**Experimental diets**

A 4 × 2 factorial arrangement of treatments was used in this study. Four dietary treatments consisted of a negative control (basal diet, designated as “control”), positive control (Zn-bacitracin, designated as “Antibiotic”), mannan-oligosaccharides (MOS) (Bio-Mos®, Alltech Inc., USA), and MOS+acidifier (Acid-Pak 4-way, Alltech Inc., USA, designated as “acidifier”). The chicks were fed under two holding times, immediate access to both feed and water after hatch (FED) and 48 h delayed access to both feed and water post hatch (HELD). Bio-MOS and Acid-Pak 4-way were supplemented at a rate of 2.0 and 3.0 kg per tonne of feed, respectively.

All commercial products used in this study were kindly supplied by the manufacturers. Zn-bacitracin was purchased from a local feedstuff supplier (Ridley AgriProducts, Tamworth, NSW, Australia). The chicks were fed starter diets for 0 to 21 d and then finisher diets until d 42. All diets were wheat-SBM based and were cold-pelleted (60-75°C). The chicks were fed a high-protein diet containing 40% fish meal from d 8 to d 14 prior to the inoculation with *C. perfringens*, to trigger the outbreak of NE (Table 1).

| Ingredient (g/kg) | Starter | Hi-protein starter | Grower/finisher |
|------------------|---------|--------------------|-----------------|
| Wheat            | 474.5   | 441.0              | 499.5           |
| Oats             | 100.0   | 100.0              | 100.0           |
| Wheat offal      | -       | 40.0               | -               |
| Rice pollard     | 26.5    | -                  | 26.0            |
| Tallow           | 35.0    | -                  | 62.0            |
| Soybean meal 48% | 190.0   | -                  | 150.0           |
| Meat-bone meal 50% | 80.0      | -                  | 75.0            |
| Peas             | 75.0    | -                  | 72.0            |
| Fish meal        | -       | 400.0              | -               |
| Limestone 38%    | 5.0     | 5.0                | 4.0             |
| Sodium bicarbonate | 3.4    | 3.4                | 2.0             |
| Salt             | 1.0     | 1.0                | 1.5             |
| Lysine-HCl       | 2.2     | 2.2                | 2.3             |
| DL-methionine    | 3.3     | 3.3                | 2.5             |
| L-threonine      | 1.0     | 1.0                | 0.3             |
| Choline chloride | 0.6     | 0.6                | 0.4             |
| Vitamin/mineral premix$^1$ | 2.0 | 2.0 | 2.0 |
| Xylanase$^2$     | 0.5     | 0.5                | 0.5             |
| **Nutrients**    |         |                    |                 |
| **Protein**      | 225.0   | 330.0              | 205.0           |
| **ME (MJ/kg)**   | 12.6    | 12.1               | 13.4            |
| **Lysine**       | 12.0    | 24.1               | 10.9            |
| **Met+cys**      | 9.3     | 14.7               | 8.6             |
| **Calcium**      | 9.8     | 18.5               | 9.0             |
| **Available P**  | 4.8     | 13.4               | 4.6             |

$^1$ Provided per kg of diet: vitamin A (as all-trans retinol) 3.03 mg, cholecalciferol 0.09 mg; vitamin E (as d-α-tocopherol) 20 mg; vitamin K$_2$ 2 mg; thiamine 2 mg; riboflavin 6 mg; pyridoxine hydrochloride 5 mg; vitamin B$_2$; 0.2 mg; biotin 0.1 mg; niacin 50 mg; D-calcium pantothenate 12 mg; folic acid 2 mg; Mn 80 mg; Fe 60 mg; Zn 80 mg; Cu 8 mg; I 1 mg; Co 0.3 mg; Se 0.1 mg; Mo 1 mg (obtained from Ridley AgriProducts, Tamworth, New South Wales, Australia).

$^2$ Allzyme PT, Alltech Inc., USA.

**Bird management**

Four hundred (400) male Cobb 500 broiler chicks were obtained from a local hatchery, with a special arrangement to take chicks out of hatchery within 2 h of hatch. The chicks were reared in floor pens in groups of 50 chicks per pen in the same environmentally controlled facility. All the chicks were wing-tagged and weighed. Half the chicks were given access to feed and water immediately after they were weighed, while the other half was held in chicken boxes at 33°C for 48 h to mimic the condition in incubator prior to giving access to feed and water.

On d 9 (five days before *C. perfringens* inoculation), all chicks were given a one-off dose of 1 ml Eimeria inoculation containing 4,500 *E. maxima*, 6,000 *E. tenella* and 10,000 *E. acervulina* (obtained from the Animal
Research Institute, Queensland Department of Primary Industries and Fisheries, Brisbane, Australia). On days 14, 15 and 16 all the chicks were orally infected with 2 ml C. perfringens culture (approx. 10^8 CFU/ml) in a thioglycolate broth (Oxoid, UK). The C. perfringens inoculant was prepared using a subculture of a C. perfringens field isolate of C. perfringens type A (obtained from the CSIRO, Geelong, Victoria, Australia). Necropsies of all dead chicks from d 14 onwards were conducted to determine the cause of death. Mortality associated with lesions of confluent necrosis or sloughing of the epithelial lining of the intestinal tract was considered to be caused by NE (Helmboldt and Bryant, 1971). For chicks died as a result of NE, intestinal swab samples from the infected area were taken and plated onto C. perfringens agar (Oxoid, UK). Total mortality and mortality due to NE were recorded daily.

The experiment was approved by the Animal Ethics Committee of the University of New England. Health and animal husbandry practices complied with the “Australian Code of Practice for the Care and Use of Animals for Scientific Purposes” issued by the Australian Government (National Health and Medical Research Council, 2004).

Live weight, feed intake and mortality

Body weight and feed intake on a cage basis were recorded weekly for calculation of weight gain during the entire experimental period. Mortality was recorded daily.

C. perfringens counts

On days 17 and 20, four chicks from each treatment group were randomly selected, weighed, and killed by cervical dislocation. The numbers of C. perfringens (CFU/g digesta) in the ileum were measured by plating onto C. perfringens agar plate.

NE lesion scores

On days 17 and 20, the small intestine from each killed bird was incised longitudinally and examined for evidence of gross NE lesion (0 = none, 1 = mild, 2 = moderate, 3 = marked/severe).

Lymphoid organ development

On sample collection days, the bursa of Fabricius and spleen were removed and weighed. The weights were expressed as a percentage of body weight.

Chicken interleukin-6 bioassay

Production of Chicken interleukin-6 (ChIL-6) was investigated using a murine IL-6 7TD1 bioassay as described by van Snick et al. (1986) with some minor modifications. The bioassay of ChIL-6 activity was conducted at the Australian Animal Health Laboratory (AAHL), CSIRO Livestock Industries, Geelong, Victoria, Australia.

Approximately 5 ml of blood were taken from each bird by wing bleeding on d 26, 10 days post infection (p.i.), using 5 ml vacutainers containing Lithium Heparin (Becton Dickinson, NJ 07417, USA). The blood samples were stored in room temperature and analysed within 24 h. The IL-6 secretion was expressed by titres (U/ml). A titre is defined as the reciprocal of the lowest dilution of a sample that gives half the maximum response. Maximum response equals the maximum count minus the background count.

Chicken T-cell proliferation assay

The same blood samples used for ChIL-6 bioassay were used for T-cell proliferation assay as described by Lowenthal et al. (1994). This assay was also conducted at the AAHL, CSIRO Livestock Industries, Geelong, Victoria, Australia. A mitogen called Concanavalin A (ConA, Sigma, Israel) was used to stimulate the proliferation of the T-cells. Two concentrations of ConA, 50 μg/ml and 10 μg/ml, were used in the assay and made by adding ConA into DMEM medium. DMEM medium was designated as 0μg/ml and used as a negative control. For each sample the assay was conducted in triplicate. An aliquot of 200 μl of each medium, 0 μg/ml, 10 μg/ml and 50 μg/ml, were pipetted into pre-labelled 96-well plates (Nunc, Kamstrup, Denmark) and 5 μl of whole blood was added to each well. The plates were gently mixed using a Dynatech Micro Shaker (IKA® Works, Selangor, Malaysia) and then placed in a humidified cell incubator at 37°C with 5% CO2 (Forma Scientific, USA) for three days. On day three, the wells were pulsed with 25 μl of 1:50 dilution of 3H thymidine and mixed gently. The plates were incubated for 18 h. On day four, the plates were removed and shaken again on a micro shaker for approximately three min to ensure that the cells were suspended in solution. The cells were harvested onto glass fibre filter mats (Wallac, Finland) using a plate harvester (Wallac Harvester 96 Mach III M, Tomtec, USA), saturated with scintillant fluid (Wallac Betaplate Scint, Wallac, Finland), and the radioactivity was measured (MicroBeta 1450 Trilux Liquid Scintillation and Luminescence Counter, Wallac, Finland). The ability of T-cells to proliferate is expressed by Stimulation Index (SI) relative to cells cultured in the absence of mitogen.

Gut morphology

On sampling days, four chicks from each treatment group were killed by cervical dislocation. The body cavity was opened, and 20 mm segments from the jejunum (20 mm distal to the point of entry of the bile ducts) and the ileum (20 mm proximal to the ileo-caecal junction) were removed for morphometric analyses. The intestinal samples
were flushed clean with phosphate buffered saline (PBS, pH 7.4) and fixed in 10% neutral buffered formalin solution for at least 24 h. The fixed tissue samples were processed in an automatic tissue processor (TOSCO, Thomas Optical and Scientific Co., Melbourne and Sydney, Australia) and embedded in paraffin using a Histo Embedding Centre (Leica EG 1160, Leica Microsystems, Bensheim, D-64625, Germany). Embedded samples were subsequently sectioned sagittally with a Rotary Microtome (Leitz 1516, Leica Microsystems, Bensheim, D-64625, Germany) at 5 µm. The tissue sections on the slides were stained using Harris’s haematoxylin (Gurr’s, UK) and eosin Y (Chroma, Germany) stains and mounted with DPX Mountant for histology (Aldrich Chemical Company, Inc., Milwaukee, WI 53255, USA). Morphometric indices were determined using computer-aided light microscope image analyses (SPOT 3.1, Diagnostic Instruments, Inc., Sterling Heights, MI 48314, USA) as described by Bird et al. (1994). The morphometric variables analysed included: villus length (from the tip of the villus to muscularis mucosa), villus height using the lamina propria as the base (from the tip of the villus to the villus crypt junction), villus width at half height, crypt depth as the depth of the invagination between adjacent villi, and thickness of the muscularis (the external muscle layer) and muscularis mucosa. Villus area was calculated from the villus height and width at half height. Values are means from 12 different villi and only vertically oriented villi and crypts were measured (Uni et al., 1999).

**Statistical analysis**

All data were analysed using Statgraphics software (Manugistics, Inc., 2000). ANOVA (by the general linear model procedure) was used to determine the significance of the main effects and interactions. Duncan’s multiple-range test was used to separate means when significant effects (p<0.05) were detected by multifactorial analyses of variance (Duncan, 1955).

**RESULTS**

**Growth performance**

Growth performance of chicks before and after *C. perfringens* challenge is presented in Table 2. FED chicks had higher (p<0.001) body weight throughout the experiment. Chicks given the MOS and acidifier both had heavier (p<0.001) body weight at d 14 than the antibiotic-supplemented chicks and the control chicks (before *C. perfringens* challenge). Furthermore, the HELD chicks given antibiotic had lighter (p<0.001) body weight than the control chicks at d 14.

The HELD chicks lost 12.3% of their initial body weight during the first 48 h post hatch, while the FED chicks gained 25.6% of the initial body weight during the same period. A higher relative growth occurred in HELD chicks immediately after the chicks were given access to nutrients from d 3 to d 28 (p<0.001) (Table 3). However, the growth rate of FED chicks became greater during week 5 (p<0.001). Chicks given dietary supplements had higher body growth during the first two weeks before *C. perfringens* infection (p<0.001). The chicks given the antibiotic exhibited higher body growth from second week post hatch (p<0.001). During wk 3, when clinical signs of NE occurred, only the chicks given the antibiotic-supplemented diet had a greater growth rate, compared to the other treatment groups (p<0.001).

The NE-related and non-NE related mortality figures for the HELD chicks were 14.6 and 6.9%, respectively, and both were numerically higher than that of the FED chicks, which were 12.2 and 2.8%, respectively.

**C. perfringens counts and necrotic enteritis lesion scores**

Only the antibiotic offered a complete protection against NE (Table 4). There seemed to have an interaction between early nutrition and dietary treatments on the *C. perfringens* counts for chicks at d 17 (3 d after challenge), with the FED chicks given MOS and acidifier having numerically lower *C. perfringens* counts in the ileum.

**Lymphoid organs and immune responses**

As shown in Table 5, the FED chicks had a significantly

| Table 2. Effects of post hatch holding time and dietary supplements on body weight at d 7, 14 and 21
| --- | --- | --- |
| Diet | Body weight (g) |   |
|   | d 7 | d 14 | d 21 |
| --- | --- | --- | --- |
| HELD |   |   |   |
| Control | 114.2<sup>ab</sup> | 302.4<sup>b</sup> | 662.4<sup>b</sup> |
| Antibiotic | 111.2<sup>b</sup> | 305.9<sup>b</sup> | 804.9<sup>a</sup> |
| MOS | 117.0<sup>a</sup> | 321.6<sup>a</sup> | 685.2<sup>b</sup> |
| Acidifier | 116.3<sup>ab</sup> | 317.3<sup>ab</sup> | 707.5<sup>b</sup> |
| FED |   |   |   |
| Control | 87.5<sup>d</sup> | 254.0<sup>d</sup> | 583.3<sup>c</sup> |
| Antibiotic | 77.4<sup>c</sup> | 233.7<sup>d</sup> | 654.6<sup>b</sup> |
| MOS | 87.6<sup>d</sup> | 259.0<sup>d</sup> | 537.3<sup>c</sup> |
| Acidifier | 93.7<sup>c</sup> | 276.5<sup>c</sup> | 586.6<sup>c</sup> |
| SEM | 1.8 | 6.7 | 18.9 |

<sup>1</sup>Values are means of 20 replicates.

<sup>2</sup> NS = Not significant; * p<0.05; ** p<0.01; *** p<0.001.

<sup>a,b,c,d</sup> Means within a column without a common superscript differ significantly.
### Table 3. Effects of post hatch holding time and dietary supplements on relative growth rate of chicks at d 3, 7, 14, 21, 28 and 35

| Treatment | Relative growth (%)<sup>a</sup> | 0-2 d | 3-7 d | 8-14 d | 15-21 d | 22-28 d | 29-35 d |
|-----------|-------------------------------|-------|-------|--------|---------|---------|---------|
| FED       |                               |       |       |        |         |         |         |
| Control   | 26.5<sup>b</sup>              | 92<sup>c</sup> | 160<sup>d</sup> | 127<sup>b</sup> | 76<sup>b</sup> | 55<sup>a</sup> |
| Antibiotic| 19.5<sup>b</sup>              | 96<sup>c</sup> | 176<sup>c</sup> | 166<sup>a</sup> | 70<sup>c</sup> | 52<sup>ab</sup> |
| MOS       | 30.1<sup>a</sup>              | 90<sup>c</sup> | 173<sup>c</sup> | 122<sup>b</sup> | 76<sup>b</sup> | 53<sup>ab</sup> |
| Acidifier | 28.4<sup>a</sup>              | 93<sup>c</sup> | 175<sup>c</sup> | 125<sup>b</sup> | 74<sup>bc</sup> | 53<sup>ab</sup> |
| HELD      |                               |       |       |        |         |         |         |
| Control   | -12.2<sup>2</sup>             | 115<sup>b</sup> | 184<sup>b</sup> | 128<sup>b</sup> | 84<sup>a</sup> | 47<sup>b</sup> |
| Antibiotic| 95<sup>c</sup>                | 204<sup>a</sup> | 190<sup>a</sup> | 84<sup>a</sup> | 43<sup>c</sup> |         |
| MOS       | 115<sup>b</sup>              | 197<sup>ab</sup> | 119<sup>b</sup> | 83<sup>a</sup> | 46<sup>b</sup> |         |
| Acidifier | 132<sup>a</sup>              | 194<sup>ab</sup> | 114<sup>b</sup> | 77<sup>b</sup> | 33<sup>a</sup> |         |
| SEM       | 1.3                           | 3     | 5     | 7      | 3       | 2       |

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| Diet       |         |       |       |       |       |
|------------|---------|-------|-------|-------|-------|
| ***        | ***     | **    | ***   | **    | ***   |

| Holding time |         |       |       |       |       |
|--------------|---------|-------|-------|-------|-------|
| ***          | ***     | ***   | NS    | ***   | ***   |

| Diet-holding time |         |       |       |       |       |
|-------------------|---------|-------|-------|-------|-------|
| NS                | NS      | NS    | NS    | NS    | NS    |

<sup>a</sup> Values are means of 20 replicates.<sup>b</sup> Value is mean of HELD chicks 48 h post hatch.<sup>c</sup> NS = Not significant; * p<0.05; ** p<0.01; *** p<0.001.<sup>d</sup> Means within a column without a common superscript differ significantly.<sup>e</sup> Weight gain relative to initial body weight at d 0.

### Table 4. Effects of post hatch holding time and dietary supplements on NE lesion scores and *C. perfringens* counts 3 and 6 days after challenge

|               | d-17 (3 d after challenge) | d-20 (6 d after challenge) |
|---------------|-----------------------------|----------------------------|
|               | NE lesion scores (10<sup>5</sup> CFU/g) | *C. perfringens* counts (10<sup>5</sup> CFU/g) | NE lesion scores (10<sup>5</sup> CFU/g) | *C. perfringens* counts (10<sup>5</sup> CFU/g) |
| FED           |                             |                            |                             |                            |
| Control       | 2.17<sup>a</sup>           | 6.38<sup>a</sup>          | 1.38<sup>a</sup>           | 0.98<sup>a</sup>          |
| Antibiotic    | 0.00<sup>c</sup>           | 0.00<sup>b</sup>         | 0.13<sup>bc</sup>         | 0.00<sup>b</sup>         |
| MOS           | 1.88<sup>a</sup>           | 1.20<sup>ab</sup>        | 0.75<sup>ab</sup>         | 0.68<sup>ab</sup>        |
| Acidifier     | 1.88<sup>a</sup>           | 1.44<sup>ab</sup>        | 0.75<sup>ab</sup>         | 0.33<sup>ab</sup>        |
| HELD          |                             |                            |                             |                            |
| Control       | 2.00<sup>ab</sup>          | 1.24<sup>ab</sup>        | 0.75<sup>ab</sup>         | 0.03<sup>b</sup>         |
| Antibiotic    | 0.25<sup>bc</sup>          | 0.00<sup>b</sup>         | 0.00<sup>b</sup>         | 0.00<sup>b</sup>         |
| MOS           | 1.63<sup>ab</sup>          | 2.58<sup>ab</sup>        | 1.75<sup>c</sup>         | 1.10<sup>a</sup>         |
| Acidifier     | 2.38<sup>a</sup>           | 2.56<sup>ab</sup>        | 1.00<sup>ab</sup>         | 0.30<sup>ab</sup>        |
| SEM           | 0.53                        | 1.84                      | 0.30                        | 0.31                      |

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| Diet        |         |       |       |       |       |
|-------------|---------|-------|-------|-------|-------|
| ***         | NS      |       |       |       |       |

| Holding time |         |       |       |       |       |
|--------------|---------|-------|-------|-------|-------|
| NS           | NS      | NS    | NS    | NS    | NS    |

| Diet-holding time |         |       |       |       |       |
|-------------------|---------|-------|-------|-------|-------|
| NS                | NS      | NS    | NS    | NS    | NS    |

<sup>a</sup> Values are means of 4 replicates for each treatment group.<sup>b</sup> NS = Not significant; * p<0.05; ** p<0.01; *** p<0.001.<sup>c</sup> Means within a column without a common superscript differ significantly.
heavier relative bursa weight at d 21 (p<0.05). MOS supplementation tended to increase the spleen weight at d 14 (p = 0.056). There was an interaction between early nutrition and dietary treatments on IL-6 production (p<0.05), with the FED chicks given MOS producing more IL-6 than other treatments (Figure 1). Early nutrition had no effect on IL-6 production in birds given the control, antibiotic and acidifier diets. T-cells proliferation was enhanced by early access to nutrients (p<0.001) (Figure 2). In the FED chicks, both MOS and acidifier reduced T-cell proliferation, compared to antibiotic (p<0.05).

**Gut morphology**

The morphology of intestinal segments was studied before and after the *C. perfringens* challenge, the FED chicks regardless of dietary treatment and the HELD chicks given dietary supplements had shallower crypts and greater villus/crypt ratio in the jejunum (p<0.01). Dietary supplements also increased villus height in the jejunum (p<0.001), but not for FED chicks given the MOS or the HELD bird given the acidifier. Early feeding and dietary supplements had a similar effect on the ileal segment, albeit less apparent. The HELD chicks had thicker submucosa in both the jejunum and ileum (p<0.001). Early feeding increased villus height in the jejunum (p<0.05), but not in the ileum.

At d 21, only the FED chicks given dietary supplements had thinner crypts and greater villus/crypt ratio in the jejunum.

**Figure 1.** Effects of post hatch holding time and dietary supplements on Chicken IL-6 production 10 days post infection (d.p.i.) (Mean values, n = 6; error bars indicate SEM; bars without a common superscript differ significantly).

**Figure 2.** Effects of post hatch holding time and dietary supplements on T-cell proliferation of broilers 10 days post infection (d.p.i.) when stimulated with different levels of Con A, expressed by Stimulation Index (Mean values, n = 6; error bars indicate SEM; bars without a common superscript differ significantly).
Table 6. Effects of post hatch holding time and dietary supplements on the morphology of small intestine at d 14

|                | Jejunum | Ileum |
|----------------|---------|-------|
|                | Submucosa | Villus height | Crypt depth | Villus/crypt ratio | Submucosa | Villus height | Crypt depth | Villus/crypt ratio |
| **FED**       |          |                 |            |                  |          |                 |            |                  |
| Control       | 16.9<bc  | 1,240.2<bc     | 195.6<ab   | 6.7<            | 20.6<abc | 555.6<bc      | 164.9<ab   | 3.5<             |
| Antibiotic    | 15.2<bc  | 1,214.1<bc     | 182.6<bc   | 7.1<            | 22.3<abc | 685.4<bc      | 186.8<     | 4.0<             |
| MOS           | 16.5<bc  | 1,069.6<bc     | 158.0<bc   | 7.3<            | 18.5<d   | 577.8<bc      | 147.3<b    | 4.1<             |
| Acidifier     | 17.7<ab  | 1,366.7<bc     | 187.8<bc   | 7.4<            | 23.0<abc | 684.2<bc      | 162.6<ab   | 4.4<             |
| **HELD**      |          |                 |            |                  |          |                 |            |                  |
| Control       | 20.2<a   | 1,059.5<c      | 225.0<b    | 4.9<b           | 25.2<a   | 605.0<b       | 190.4<     | 3.5<             |
| Antibiotic    | 19.5<a   | 1,241.9<b      | 203.4<bc   | 6.4<            | 23.7<ab  | 684.9<bc      | 189.7<     | 3.7<             |
| MOS           | 17.9<ab  | 1,219.2<b      | 180.5<bc   | 7.2<            | 23.6<ab  | 686.2<bc      | 178.9<     | 3.9<             |
| Acidifier     | 18.9<ab  | 1,166.8<bc     | 178.9<bc   | 7.1<            | 19.3<cd  | 592.7<b       | 166.8<bc   | 3.9<             |
| SEM           | 0.8      | 36.8            | 10.3       | 0.4             | 1.2      | 18.5           | 10.1       | 0.3              |
| **Probability of greater F value in analysis of variance** |     |      |      |     |     |      |      |     |

Diet: NS *** *** ** NS *** * NS
Holding time: *** * * * *** *** *** ***
Diet-holding time: NS NS NS NS *** *** *** ***

1 Values are means of 12 replicates. 2 NS = Not significant; * p<0.05; ** p<0.01; *** p<0.001.

jejunum and ileum (p<0.001). In the HELD chicks, the antibiotic and acidifier gave greater villus height, but only the antibiotic gave shallower crypts and higher villus/crypt ratio in the jejunum (p<0.001). Again in the HELD chicks, the acidifier increased villus height, while the antibiotic reduced it in the ileum (p<0.001). The FED chicks given dietary supplements had shallower crypts in the ileum (p<0.001), while in the HELD chicks, only the antibiotic had such an effect (p<0.001).

**DISCUSSION**

The current NE model was designed to mimic real-life situation where mortality from NE should be moderate and

Table 7. Effects of post hatch holding time and dietary supplements on the morphology of small intestine at d 21

|                | Jejunum | Ileum |
|----------------|---------|-------|
|                | Submucosa | Villus height | Crypt depth | Villus/crypt ratio | Submucosa | Villus height | Crypt depth | Villus/crypt ratio |
| **FED**       |          |                 |            |                  |          |                 |            |                  |
| Control       | 30.0<ab  | 1,417.9<ab     | 280.5<ab   | 5.14<c           | 28.8<a   | 893.2<a        | 198.3<     | 4.81<            |
| Antibiotic    | 23.2<abc | 1,542.8<c      | 200.4<cd   | 8.93<            | 25.8<abc | 772.1<bc       | 102.0<     | 8.04<            |
| MOS           | 21.2<de  | 1,239.6<bc     | 174.9<d    | 7.39<            | 24.4<abc | 782.4<bc       | 103.6<     | 8.23<            |
| Acidifier     | 26.6<ab  | 1,418.7<bc     | 227.1<bc   | 6.61<cd          | 28.9<a   | 890.4<a        | 116.7<     | 7.65<            |
| **HELD**      |          |                 |            |                  |          |                 |            |                  |
| Control       | 19.0<bc  | 1,321.3<bc     | 189.8<bc   | 7.14<c           | 26.4<ab  | 803.5<bc       | 108.3<     | 7.73<            |
| Antibiotic    | 18.6<abc | 1,435.7<bc     | 135.0<     | 10.88<           | 21.7<c   | 694.5<bc       | 79.1<      | 9.33<            |
| MOS           | 19.7<de  | 1,062.9<bc     | 264.1<b    | 5.32<de          | 21.8<bc  | 835.1<bc       | 115.9<     | 7.34<            |
| Acidifier     | 22.8<cd  | 1,566.2<bc     | 262.1<b    | 6.12<de          | 23.9<bc  | 893.3<bc       | 114.6<     | 8.05<            |
| SEM           | 1.3      | 59.6            | 24.6       | 0.45             | 1.4      | 29.2           | 6.4        | 0.49             |
| **Probability of greater F value in analysis of variance** |     |      |      |     |     |      |      |     |

Diet: *** * *** *** ** *** *** ***
Holding time: *** NS NS NS *** NS *** **
Diet-holding time: NS NS NS NS *** *** *** ***

1 Values are means of 12 replicates. 2 NS = Not significant; * p<0.05; ** p<0.01; *** p<0.001.

*a,b* Means within a column without a common superscript differ significantly.
performance of unaffected chicks relevant to commercial standard (Wu et al., 2010). In the current study, only the antibiotic, Zn-bacitracin, offered a complete protection against the outbreak of the disease. The birds used in the study performed to commercial standard, suggesting that there were no lingering health problems with the chicks. Some of the feed additives as well as early nutrition were helpful in preventing performance loss through effects on gut development and immunity. These effects are detailed in the following sections.

Growth performance

As growth rate increases and market age decreases, the percentage of time a broiler bird spends as a neonate increases (De Beer et al., 2011). The impact of nutrition at an early age on the chicks has been studied intensively. It is now well accepted that early access to nutrients post hatch has a major impact on chicks’ both immediate and long-term development. Early growth is directly correlated with the final body weight and feed efficiency. Heavier body weight on placement usually results in heavier body weight at marketing (Wilson, 1991). Research in early life of chickens has led to the conclusion that the immediate post hatch period is critical for the development of all major organs in chicks (Uni, 1998; Hornasio et al., 2011). Early feeding has been reported to enhance the body growth, uniformity and health status of the chicks (Moran, 1990; Casteel et al., 1994; Sklan et al., 2000).

In this study, body weight of the FED chicks increased by 25.6% compared with their initial body weight during the first 48 h post hatch, while the HELD chicks lost 12.3% of the initial body weight during the same period. The HELD chicks demonstrated superior growth relative to body weight from d 3 to d 14, before the \textit{C. perfringens} infection. However, after the chicks were given \textit{C. perfringens} challenge, the FED chicks seemed to be better able to grow than the HELD chicks. This was not the case in a previous study (Ao and Choc, 2003) where the chicks were reared under a higher standard of hygiene without NE challenge, suggesting an interaction between early feeding and disease challenge. Thus delayed access to nutrients up to 48h post hatch may have caused a long-term compromise on the competence of chicks to resist disease challenge and to gain compensatory growth. Both the MOS and acidifier used in the current study failed to prevent the outbreak of NE. This is of practical importance because under commercial production systems, broiler chickens face many stressors like delayed access to feed, dirty litter and a frequent change of diets. Therefore it might be worthwhile to consider early feeding strategies, in combination with good environmental management, where antibiotics are not allowed in feed.

NE lesion scores and \textit{C. perfringens} counts

Feed additives such as acidifiers, enzymes, probiotics and prebiotics have all been reported to inhibit the pathogen colonisation of the intestine and subsequently reduce the incidence of some diseases, such as NE, with varying levels of success (Corrier et al., 1995; Thompson and Hinton, 1997; Finuance et al., 1999; Kaldhusdal, 2000). Whilst some of these additives are claimed to have a direct effect on the organisms themselves, others are deemed to have a more indirect impact, such as enhancing immunity. It is reported frequently that early access to nutrients post hatch benefit the health status of chicks through aiding the immune system. But it was not known whether birds with a stronger immune system would be better able to resist disease, for example NE. In the current study, the HELD chicks had numerically higher NE lesion scores and \textit{C. perfringens} counts in the intestine, especially three days after \textit{C. perfringens} challenge. These results appeared to indicate that under the condition of this study, early feeding offered a degree of protection even though the reduction of NE lesion scores and \textit{C. perfringens} counts was not statistically significant.

Lymphoid organs

The protective immunity of chicks plays an important role in preventing infectious diseases, especially when in-feed antibiotics are not used. The first week post hatch is a critical period and the nutritional status of chicks can impact the immune system (Klasing, 1998). The bursa of Fabricius, a primary immune organ, plays a major role in creating antibody diversity (Uni, 1998), and inhibition of bursal development was found to result in the failure of normal development of the spleen, the secondary immune organ (Glick, 1967). The bursa and spleen were both found to be responsive to environmental stress of chicks at an early age (Wyatt et al., 1986). Dibner et al. (1998) found that chicks had early access to feed and water show a higher relative bursa weight, earlier appearance of germinal centres, and better disease resistance than their HELD hatch mates. This is also supported by the data of Wyatt et al. (1986) who found that broiler chicks held in incubators for 30 h without access to feed and water before placement showed significantly lower bursa and spleen weights. This suggests that early stress to young broiler chicks, such as holding without feed and water post hatch, alters the immune capabilities and growth rate later in their life. In the current study, under a challenged environment, the FED chicks given the MOS diet had heavier bursas. This appears to suggest that when chicks are exposed to certain environmental stressors, early feeding and certain dietary additives might have a greater ability to modulate the development of these lymphoid organs and enhance the immune competence of the bird.
Immune responses

The immune response to infection is controlled by a complex interplay between various cytokines (Kelso and Metcalf, 1990). IL-6 is a multifunctional cytokine that plays a major role in regulating immune responses, acute phase reactions and haemopoiesis (Hirano et al., 1986). IL-6 is produced by many different cell types and acts on B-cells, T-cells, hepatocytes, haematoipoietic progenitor cells and cells of the central nervous system, immediately following infection or vaccination (Hirano et al., 1986; Gauldie et al., 1987; Ikehuchi et al., 1987).

Since T-cells are the major source of cytokines, the ability of these cells to proliferate in response to their mitogens has been used to determine the development of the immune responses of chickens (Lowenthal et al., 1994). Studies conducted by these authors suggested that susceptibility of newly hatched chicks to infection is due to a period of transient T-cell unresponsiveness to immune stimulation, a deficiency that may be amenable to cytokine therapy. In this study, only the FED chicks given MOS diet had elevated IL-6 production. This indicates that there is an interaction between early nutrition and dietary supplements. Early administration of MOS might augment the production of IL-6 under challenged conditions. In the current study, early feeding significantly increased T-cell proliferation. Interestingly, the T-cells of the chicks given the control or the antibiotic diet had a significantly enhanced ability to proliferate, while all the other additives had the opposite effects. Such an effect has not been demonstrated elsewhere. The mechanism by which this occurs is not understood. This result, together with the data on IL-6 production, suggests that early access to nutrients post hatch might enhance the immune responses to infectious challenge.

Gut morphology

Hydrolysis of macromolecules in the small intestine is achieved, to a large extent, by pancreatic enzyme activities, which are correlated with body weight and intestinal weight (Sklan and Noy, 2000). Undoubtedly early access to nutrients stimulates the gastro-intestinal tract (GIT) activity (Uni et al., 1998; Noy and Sklan, 1999). It was reported that the GIT increases in size and weight more rapidly in relation to body weight, during early post hatching period, than other organs and tissues of chickens (Lilja, 1983; Katanbaf et al., 1988). The preferential early growth of the small intestine occurs both in the presence and absence of the feed, although in the absence of exogenous feed both the absolute and the relative growth rates are lower (Noy and Sklan, 1999). The intestine is also the largest immunological organ and contains a large number of lymphocytes, most of them are T-cells located predominantly in the Lamina propria (Perdigon et al., 1991).

The interaction between the microflora and the morphology of the intestinal wall is clearly shown by alteration in the structure and morphology of the GIT of germ-free animals compared with that in conventional animals (Heneghan, 1965). The small intestine villi of germ-free species are usually slender and uniform in shape, whereas crypts are shorter (Gordon and Bruckner-Kardoss, 1961). Cook and Bird (1973) associated the presence of pathogenic micro-organisms with a change in the intestinal wall and in the surface area for nutrient absorption. They demonstrated a shortening of the villi and a decrease in their epithelial layers when counts of pathogenic bacteria increased, in company with a deeper crypt. Schneeman (1982) suggests that shorter villi relative to crypt depths result in less absorptive and more secretory cells. This change in the intestinal morphology due to pathogenic organisms is more pronounced in the upper than in the lower parts of the intestinal tract. It was reported that Coliform bacteria (Truscott and Al-Sheikhly, 1977) and Clostridia (Kaldhusdal and Hofshagen, 1992) in the intestinal tract cause damage to the mucosal layer of the intestine of broiler chicks. Based on these results it is possible that morphological changes in the intestinal wall are indicative of a disturbance in the balance between non-pathogenic and pathogenic micro-organisms. In addition, research showed that chicks with immediate access to feed and water post hatch have increased villus height and surface area of the small intestine, and enhanced crypt development (Uni et al., 1998). The length of villi within the jejunum and ileum of chicks receiving an amylase-supplemented corn-soy based diet was significantly increased, with a concomitant improvement in growth rate (Ritz et al., 1995). This might suggest an increased absorptive area capable of greater absorption of available nutrients (Caspary, 1992).

In the current study, when the chicks were 14 days old, just before the C. perfringens infection, all the dietary supplements tended to increase villus height and reduce crypt depth in the jejunum, thus increasing the villus height/crypt depth ratio. This was particularly obvious in the HELD chicks. On d 21, seven days after the initial C. perfringens challenge, the FED chicks given the supplements still showed reduced jejunal crypt depth and increased villus/crypt ratio; while in the HELD chicks, the supplements increased the crypt depth and reduced the villus/crypt ratio. The morphology of the ileum followed a similar trend, although less pronounced. The results on the intestinal morphology suggest that under unchallenged conditions, both dietary supplements and early access to nutrients may enhance the absorptive capacity and reduce the secretion in the small intestine. But under pathogenic challenge, early feeding and dietary supplements gave the FED chicks reduced crypt depth and thus increased villus/crypt ratio, but not the HELD chicks, indicated the
gut microflora community might be disturbed for HELD chicks given the supplements.

The present study indicated that the supplementation of MOS or acidifier to broiler diets enhanced the absorptive surface area of the small intestine, and improved the growth performance of chicks before *C. perfringens* challenge. However, neither of these additives gave the chicks the same degree of protection against *C. perfringens* infection as an antibiotic did.

Early access to certain supplements post hatch, such as MOS and acidifier, improved the development of intestinal morphology and immune competency of the chicks, thus enhanced the absorptive potential and the immune response of chicks under pathogenic infection.

In conclusion, under challenged condition, chicks with early access to feed and water exhibit superior disease resistance and growth performance through enhanced immunity and intestinal development.

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