Effects of Probiotic on the Expression of IL-7 Gene and Immune Response to Newcastle Disease Vaccine in Broilers

Research Article


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Abstract

Probiotics have potential as prevention strategies or therapies for multiple diseases in chickens, such as Newcastle Disease (ND), heat stress, and Coccidiosis, but the mechanism still partially understood. It is an important and warranted issue for further investigation that how probiotics modulated gene expression in disease. In this study, 80 Specific-pathogen-free (SPF) broilers were treated with probiotics, and inoculated La Sota vaccine, then detected the changes of the number of antibody-producing cells (IgA+, IgM+, and IgG+) and the expression of IL-7 mRNA in mucosa associated lymphoid tissue (MALT: Harder’s gland, cecal tonsils, duodenum and ileum Peyer’s patch), and the content of Immunoglobulins (IgA, IgM, and IgG) in local humor (tear, trachea fluid, bile and intestinal fluid) by immunohistochemistry staining, RT-qPCR, and IELISA, respectively. Results showed that the contents of IgA, IgM and IgG in local humor, the number of IgA+, IgM+, and IgG+ and the expression of IL-7 mRNA in MALT of inoculated birds were significantly higher than control birds (P < 0.05 or P < 0.01) after 7 days post-inoculation (PI), and chickens showed higher immunity response after treated with probiotics. These results indicated that probiotics could enhance the humoral immunity in the respiratory and digestive tracts of chickens, and the immunological response to ND vaccine, and ND vaccine in combining with probiotics will get a better effect on humoral immunity. Both the probiotics and ND vaccine could increase the expression of IL-7 mRNA, and then improve the humoral immunity in the chickens, and the IL-7 maybe one of the ways by which the probiotics improve the humoral immunity.

Keywords: Newcastle Disease; Probiotics; Chickens; Humoral Immunity; IL-7.

Introduction

Newcastle Disease Virus (NDV) is an economically important and frequently isolated worldwide pathogen whose listed status with OIE marks its importance to both commercial poultry producers and poultry trading [2], as it causes the highly contagious and acute septic infectious diseases of poultry with the typical lesions for respiratory and gastrointestinal mucosal bleeding. As the number of livestock units lost for poultry species, Newcastle Disease (ND) has been ranked as the fourth most important disease [33], and the continuous variation of the ND pandemic strain contribute to the hard-work in prevention and treatment. Now, vaccination for NDV is primarily by mass application of live-virus vaccines in commercial poultry, but immune protection involves responses that are presently incompletely defined [28]. In terms of ND vaccine, failure reports on the ability of classical ND vaccines to stop spread of disease from many countries, renewed interest in a more complete understanding of the global immune response of poultry to NDV will be critical in developing new control strategies and intervention programs in the future [14].

In the last century, many studies have reported that probiotic play an important role in maintaining the balance of the digestive tract flora, improving the utilization of feed, promoting the absorption of nutrients, and improving the performance of poultry production [11, 22], and also in the modulation of immunological, respiratory, inflammation, and gastrointestinal functions [6]. It is available to consumers mainly in the form of dietary supplements and foods. Unlike antibiotics, the probiotics are the green and safe feed additives as for the no residues, non-toxic, and no resistance [20]. Research found that probiotics have effects on the modulation of humoral, cellular and nonspecific
Interleukin 7 (IL-7) is a pluripotent cytokine that plays a critical role in lymphocyte development and homeostasis, it has been suggested as potential immunomodulatory for use in HIV infection [17]. Reported claimed that IL-7 is indispensable to the mature of surface immunoglobulin (sIgM+) in the conventional cells culture, and cell culture with IL-7 balance the generated from B cells by controlling cell death. IL-7 receptor-deficient mice are always along with the absence of y8TCR and Intestinal intraepithelial lymphocytes (IEL), and their IgA+ plasma cells was significantly decreased in intestinal mucosa, the secretion of the IgA and the response to antigen's stimulation of IgA also declined [18]. In normal mice, IL-7 have a stimulating effect on the B cells of spleen, lymph nodes and bone marrow, so it often used to induce B cell in vitro [21]. People also found the IL-7 pathway is essential to immune system development and maintenance in physiological conditions [19]. Recently, more and more reports on the IL-7 application and clinical treatment, IL-7 can effectively elicit an immune reconstruction in a variety of lymphopenia's model, and injected IL-7 can accelerate recovering the number of CD4+ and CD8+ T cells in the spleen and lymph node [10], and it reported that autologous tumor vaccine modified with NDV strain LX/(IL-7) could promote the antitumor immune responses mediated by CD8+ T cells and significantly improve the efficacy of the Autologous tumor vaccine modified with nonlytic Newcastle disease virus (ATV-NDV) [36]. However, limited studies have been conducted to the relationship between IL-7 and probiotics, and evaluate the potential of IL-7 expressing ND vaccine to induce anti-NDV immune response in chickens with probiotics protection efficacy. Consequently, a rapid, sensitive, and specific RT-qPCR assay was designed for the detection of IL-7 gene expression of IL-7 mRNA and immune response in mucosa with SYBR Green, try to clarify the effects of probiotic on the protection efficacy. Consequently, a rapid, sensitive, and specific RT-qPCR assay was designed for the detection of IL-7 gene expression of IL-7 mRNA and immune response in mucosa with SYBR Green, try to clarify the effects of probiotic on the

Materials and Methods

Chickens, probiotics, and ND vaccine

Specific pathogen-free (SPF) Avian broilers (n=80, one-day-old) were purchased from and fed in SPF Laboratory Animal Center of Harbin Veterinary Research Institute, Chinese Academy of Agricultural Sciences (CAAS), Harbin, China. Probiotics were provided by HUAERKANG Biotechnology Co. Ltd, Jinan, China, which is mainly contains of Bacillus subtilis, Bacillus licheniformis, Lactobacillus, Bifidobacterium, yeast, etc., as well as some enzymes like cellulase, hemicellulase and pectinase. Numbers of total and viable bacteria is more than 20 billion per gram, and (Total) enzyme activity is more than 1,000 u/g. LaSota vaccine (IV strain) were from First Biological Factory of Harbin Pharmaceutical Group, Harbin, China.

Experimental design

Random allocation was taken to averagely divide the broilers into four groups named as Control group (C), Probiotics treated group (P), ND vaccine inoculated group (N), and Probiotics treated and ND vaccine inoculated group (S). The schedule of all treatment was presented in Table 1.

The MALT (Harder's gland, cecal tonsils, duodenum Peyer's patch, and ileum Peyer's patch) and local humor (tear, trachea fluid, bile and intestinal fluid) were collected from three birds per group on days 0, 7, 14, 28 and 42 post-inoculation (PI).

Indirect Enzyme-linked Immunosorbent assay (IELISA)

Collected tear, trachea fluid, bile and intestinal fluid were diluted by PBS (0.05M pH9.6) according to 1:10, but tear according to 1:50, and then add to the 96-well polystrene microtiter plate, 200µl per hole. Here, the dilution for rabbit anti-chicken whole serum was 1:200 as primary antibodies, and the dilution for HRP-labeled Goat Anti-Rabbit IgG conjugate was 1:500 as second antibodies. However, 100µl TMB (tetrathylbenzidine, Sigma,
USA) as the color-developing agent was added to each hole. As soon as the chromogenic reaction was terminated, the OD (optical density) value was counted on 450nm by Universal Microplate Spectrophotometer (Molecular Devices, USA), and calculated the average of three holes of each sample.

Immunohistochemistry streptavidin-peroxidase (IHC - SP) staining

The procedure of IHC - SP staining was performed by the immunohistochemical kit (ZSGB-Bio, Beijing, China). The dilution for rabbit anti-chicken polyclonal antibody (Prepared in our laboratory) was 1:100 as primary antibodies, and the dilution for HRP-labeled Goat Anti-Rabbit IgG (H+L) was 1:500 as second antibodies. PBS replaced antibody in the negative control group. In addition, the TMB is for color reactions, and the apparent brown color observed under a light microscope indicated positive staining of the bound toxin. Finally, the number of IgA+, IgM+, and IgG+ in MALT were enumerated with point-counting method by H600L Microscopic pathological imaging system (Nikon, Japan).

Real-time Quantitative PCR (RT-qPCR)

Totals RNA of MALT were extracted by using a Trizol Reagent (Invitrogen, USA) as recommended by the manufacturer. The qualities of RNA were analyzed by the electrophoresis 1.0% agarose gel. Moreover, the concentrations of RNA were quantified by UV Spectrophotometer (APEL, Japan) at 260 nm and 280 nm (OD260/OD280 = 1.8). According to the specific sequences of chicken IL-7 and β-actin in Genbank, primers were synthesized by Invitrogen (USA). Details of the primers are given in Table 2. Complementary DNA (cDNA) was synthesized in a volume of 25μl with Moloney Murine Leukemia Virus (M-MLV) Reverse transcriptase (Promega, USA) according to the manufacturer's protocol. IL-7 and β-actin was amplified using Taq DNA Polymerase (Promega, USA) according to the manufacturer's protocol. The resulting PCR products were cloned into pMD18-T Vector (TakaRa, Dalian, China) and propagated in competent Escherichia coli DH5α cells (Life Technologies, USA), according to the manufacturer's instructions. Plasmid DNA was purified on columns with a StarPrep Plasmid Miniprep Kit (GenStar Bio Solutions, Beijing, China) and the concentrations were measured by UV spectrophotometer. 10 fold dilutions of plasmid DNA were used to construct the standard curve. RT-qPCR reactions were performed in Light Cycler 2.0 (Roche Diagnostics, USA) with a total reaction volume of 20μl, containing 10μl Platinum® SYBR® Green qPCR SuperMix-UDG (Invitrogen, USA), 2.0μl plasmid DNA, 0.8μl (10 μmoL/L) of each primers, and 1.0μl Bovine Serum Albumin (BSA; Thermo Fisher, USA). Each amplification was performed in triplicate wells. The temperature-time profile was pre-denaturation at 95°C for 30s for 45 cycles of 95°C for 5s, 60°C for 20s, and a final extension cycle at 72°C for 10s.

Statistical analysis

The data of RT-qPCR were recorded and analyzed by the Light Cycler software 4.0. Copy number of target genes (relative to β-actin) was defined by 2ΔΔCt [7], where ∆∆Ct=∆Ct [sample] - ∆Ct [calibrator] = (Ct_IL-7(sample) - Ct_β-actin(sample)) - (Ct_IL-7(calibrator) - Ct_β-actin(calibrator)). All of the statistical analyses were performed using SPSS 17.0. The Student’s t-test was used to determine significant differences between change values of the six time points post infection. A value of p < 0.05 was considered to be significant. Statistical analysis was carried out using Prism 5.0 (GraphPad Software).

Results

Changes of IgA, IgM, and IgG in local humor by IELISA

The contents of IgA, IgM, IgG in the tear, trachea fluid, bile and intestinal fluid of P, N and S showed higher than C at the same time point PI, and S were higher than N, N were higher than P (Figure 1, 2). And the contents of IgA, IgM, and IgG in these humors are increased before 7 day PI, and then recovered to normal. There was no significant difference (P > 0.05) in the contents of IgA, IgM, and IgG in the tear and bile between P and C among 7 to 42 day PI, while the contents of IgA, IgM, and IgG in intestinal fluid, and the contents of IgA in trachea fluid of P showed a very significantly higher (P < 0.01) than C among 7 to 14 day PI. In the comparison of P and S, S was higher than P, the significant difference occurred in the tear (IgA) at 14 day PI (P<0.01), in the trachea fluid (IgM) at 7 and 28 day PI (P<0.01 or P<0.05), in bile (IgA, IgG) at 7 to 14 day PI (P<0.01), and in the intestinal fluid (IgA, IgM, IgG) at 7 to 14 day PI (P<0.01). As for P and N, the contents of IgA, IgM, and IgG in all detected humors in N were very higher than P at 0 day PI, and the same results still found in the bile (IgG) and intestinal fluid (IgA) at 14 day PI. In addition, there was a significantly difference between N and S in the contents of IgA in the tear, trachea fluid, bile and intestinal fluid at 0 day, and then also showed the same results in tear and bile at 14 day, 7 and 42 day PI, respectively. The contents of IgM in intestinal fluid of S were significantly higher than N at 0 to 14 day PI. The contents of IgG in bile and intestinal fluid showed a significant difference between N and S at 0 to 14 day PI (P < 0.01).

Changes of IgA’, IgM’, and IgG’ in MALT by IHC - SP staining

From the Figure 3 and 4, the numbers of IgA’, IgM’, and IgG’ in Harder's gland, cecal tonsils, duodenum Peyer's patch and ileum

Table 2. Sequence of primers used in this study.

<table>
<thead>
<tr>
<th>Primers</th>
<th>Sequences</th>
<th>Accession No./ Amplicon size</th>
</tr>
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<tbody>
<tr>
<td>IL-7(F)</td>
<td>TCTATCTTTCGAGGTCTCTGCCA</td>
<td>AM931037 / 135bp</td>
</tr>
<tr>
<td>IL-7(B)</td>
<td>CATGACATCGAAGGCTGGAAG</td>
<td></td>
</tr>
<tr>
<td>β-actin(F)</td>
<td>TGAAGGCCGACAGCCTAAAGAGGTAT</td>
<td>AF294323 / 135bp</td>
</tr>
<tr>
<td>β-actin(B)</td>
<td>TGCTCCTCAGGCCGCTACTCTC</td>
<td></td>
</tr>
</tbody>
</table>
Figure 1. Contents of IgA, IgM, and IgG in the tear (A, B, C) and trachea fluid (D, E, F) of experimental chickens, respectively. Data are mean ± SD of three chickens per group. Adjacent letters indicates significantly different between any two groups at the same time point (P < 0.05), with interval letter indicates very significantly difference at P < 0.01; the same letters and no letters means that no significant different (P > 0.05).

Figure 2. Contents of IgA, IgM, and IgG in the bile (G, H, I) and intestinal fluid (J, K, L) of experimental chickens, respectively. Data are mean ± SD of three chickens per group. Adjacent letters indicates significantly different between any two groups at the same time point (P < 0.05), with interval letter indicates very significantly difference at P < 0.01; the same letters and no letters means that no significant different (P > 0.05).
Figure 3. Numbers of IgA⁺, IgM⁺, and IgG⁺ in Harder's gland (A, B, C) and cecal tonsils (D, E, F) of experimental chickens, respectively. Data are mean ± SD of three chickens per group. Adjacent letters indicates significantly different between any two groups at the same time point (P< 0.05), with interval letter indicates very significantly difference at P< 0.01; the same letters and no letters means that no significant different (P> 0.05).

Figure 4. Numbers of IgA⁺, IgM⁺, and IgG⁺ in duodenum Peyer's patch (G, H, I) and ileum Peyer's patch (J, K, L) of experimental chickens, respectively. Data are mean ± SD of three chickens per group Adjacent letters indicates significantly different between any two groups at the same time point (P< 0.05), with interval letter indicates very significantly difference at P< 0.01; the same letters and no letters means that no significant different (P> 0.05).
Figure 5. Establishment of RT-qPCR to detect the expression of IL-7 mRNA in MALT of experimental chickens. (A), (B), (C) were represent the amplification, standard, and melting curve of pMD18-T-IL-7 plasmid, respectively. (D), (E), (F) were represent the amplification, standard, and melting curve of pMD18-T-β-actin plasmid, respectively. R2 values for IL-7 and β-actin were 0.999 and 0.998, respectively. And the slopes of the standard curve for IL-7 and β-actin were -3.217 and -3.358, respectively.

Figure 6. Expression of IL-7 mRNA in Harder’s gland (A), cecal tonsils (B), duodenum Peyer’s patch (C), and ileum Peyer’s patch (D) of experimental chickens. Data are mean ± SD of three chickens per group. Adjacent letters indicates significantly different between any two groups at the same time point (P< 0.05), with interval letter indicates very significantly difference at P< 0.01; the same letters and no letters means that no significant different (P> 0.05).
PL, and there was a significant different the two group birds in the IgM^+ and IgG^+ of duodenum and ileum Peyer's patch (P < 0.01 or P < 0.05). Additionally, the number of IgA^+ in Harder's gland of S were higher than that of N from 0 to 7 day PI, and it showed same results in cecal tonsils and Peyer's patch at 42 day PI (P < 0.01); there was a significantly different between S and N for the numbers of IgM^+ in Harder's gland and cecal tonsils at 42 day PI (P < 0.01), and the same results found in duodenum and ileum Peyer's patch at 1 to 28 day, 7 and 42 day PI, respectively. As for the number of IgG^+, S had a significant higher than P in duodenum and ileum Peyer's patch after 28 day PI.

Expression of IL-7 mRNA in MALT by RT-qPCR

The comparative analysis of nucleotide homology between IL-7 and β-actin sequences which has been determined by Takara (Beijing, China) and results are 100% and 99.5% in comparison of the sequence that has been published in GenBank, respectively. Tenfold IL-7 and β-actin standard dilutions (10^6, 10^5, 10^4, 10^3, 10^2, 10^1, 10^0) amplified by RT-qPCR (Figure 5A and 5D). In our standard curves (Figure 5B and 5D), R2 values for IL-7 and β-actin were 0.999 and 0.998, respectively, which were greater than 0.99. And the slopes of the standard curve for IL-7 and β-actin were -3.217 and -3.358, respectively, which also showed a high credibility. The melting curves of IL-7 and β-actin formed a single peak at 79.6°C and 81.7°C, respectively (Figure 5C and 5F). In addition, all negative control were flat, indicated that the primers of IL-7 and β-actin were specificity for this study.

As results illustrated in Figure 6, the IL-7 mRNA expression in Harder's gland, cecal tonsil, duodenum Peyer's patch, and ileum Peyer's patch of all experimental group (P, N, and S) are higher than C at the same time point, and it showed the low expression in these immune tissues of all experimental groups at 14 day PI, in comparison with other days PI. In the view of C and P, the IL-7 mRNA expression of P in these immune tissues had a significantly higher than P at 0 to 42 day PI (P<0.01 or P<0.05). In the comparison of P and N, the IL-7 mRNA expression in these immune tissues showed significant difference between two groups among 0 to 42 day PI(P<0.01 or P<0.05), excluding 28 day PI in Harder's gland (compare P and N), 42 day PI in ileum Peyer's patch (compare P and N), and 28 day PI in ileum Peyer's patch (compare N and S), that the three comparison data showed no significantly deference (P>0.05).

Discussion

Mucosal immune system (MIS), as the first barrier to protect from pathogenic microorganisms for poultry, can produce both humor immunity and cell immunity by antigen, but the former is dominated. It's reported that more than 95 % of the poultry's infections occur in or invaded through the mucosal [25]. IgA, IgM, and IgG antibodies are produced as part of the immune response in chickens [10]. It was found that Peyer's patches are a highly enriched source of cells which have the potential to proliferate and differentiate into IgA-producing immunocytes. Cecal tonsil as an important source of antibodies, not only does it involve in the production of antibodies, but also in cell-mediated immune (CMI) response, and has a local immunity effects on intestinal bacteria and other antigenic material. Harder's gland is a specific immune tissue in chicken's eye ground, which plays a significant role on local immunity. When chickens infected with vaccine, Harder's gland generated and improved immune response to the vaccine with the independent of maternal antibody interference [31].

In this study, probiotic contains of variety of living bacteria and enzymes, it was found that probiotic can not only improve the humor immunity of chickens, but also enhanced the humor immunity response to ND vaccine, when used in combination with ND vaccine it is recognized as the more effective treatment for improving humor immunity. These results are in consistent with the previously reports. People found that probiotics initially planting on the gut play a pivotal on stimulating the development of immunity system. It have a wide range of enhancing immune function, such as activating the immune cells, increasing the number of T and B cells, promoting the proliferation and maturation of T and B cells, enhancing the capability of immune recognition, and inducing cytokines production, moreover, it can strengthen the immune defense by lymphocyte recirculation to activate the whole immune system in host [8, 24]. Lactobacillus-based probiotics have been shown to improve humor immunity in chicken's response to live ND vaccines during heat stress [30]. In weaned rabbits study, Fed-bacillus licheniformis had an increasing of humor immunity and the effect of vaccine vaccinated with Bacillus licheniformis was better than only vaccine in weaned rabbits [13].

As a heterologous antigen, the vaccine recognized and processed to Th cells by antigen presenting cell (APC), and activated Th cells to secrete various cytokines, and then these cytokines combine with B-cell receptor (BCR) to promote the production of antigen for humor immunity. Furthermore, B cells can recognize the antigen, and differentiate and mature by Th cells’ assist for humor immunity. It was reported that probiotics can significantly improve the level of Igs in serum and gut in chickens inoculated with tetanus toxoid and Clostridium perfringens α-type vaccines [8]. Moreover, probiotics can increase the level of IgG in serum of baby for β-type haemophilus influenza vaccine [15]. However, the dominant antibody isotype of the mucosal immune system is IgA, especially sIgA, so we also found that the contents of IgA increased more than IgM and IgG, even the three Igs all increased after chicken fed with probiotics and inoculated with ND vaccine. Study showed that the contents of IgA increased rapidly in the tear, Harder's gland and tracheal liquid after chickens vaccinated with ND vaccine by nasal drop, especially in the tear at 2 weeks after vaccinated [34]. Moreover, sIgA expressed high level in intranasal, trachea, bronchi and bronchiolo mucus after inoculated with a lethal dose of influenza virus for mice, which demonstrated that sIgA participate in the defense against influenza virus in the upper respiratory tract [1]. Study on the cold stress cause the change of the change of immune function in chicken intestinal, demonstrated that the expression levels of IgM, IgA, IgG, and IL-7 had an increased tendency in the small intestine of broilers following acute and chronic cold [35]. After CD4+ T-deficient mice inoculated with Formalin inactivated influenza virus A/PR8/34, the contents of IgG and IgA in serum and sIgA in mucosal secretions were high increased. However, recently research claimed that antibodies remain the primary mechanism contribute to the protection against virulent NDV in the CMI detection following live NDV, and CMI are still an important factor to be considered in the face of field challenge [14].
Here we showed that the expression of IL-7 mRNA in Harder's gland, cecal tonsil, duodenum and ileum Peyer's patch of probiotics protected chickens were higher than control birds, and it also increased after probiotic protected chickens following the inoculation of ND vaccine, these results indicated that probiotic could upregulate the expression of IL-7 mRNA in MALT, which improve the MIS response to ND vaccine, and IL-7 may one of ways for probiotics improve the humoral immunity. Humoral and cell mediated immune response involve in the protection against NDV infection [26]. IL-7 is a tissue-derived cytokine secreted from stromal and epithelial cells in various locations, as MALT, bone marrow, thymic stromal cells and keratinocyte, and there are many tumor cells and cords found to secret IL-7, as well [9, 32]. It was reported CD8+ T cell play a role in protection against NDV, and the expression of IL-7 and their receptors have been shown be crucial for CD8+ T cell memory generation and maintenance, moreover, as the only cytokine that has been identified can stimulate memory CD4+ T cell homeostasis and survival, IL-7 play a decisive role in the homeostasis and survival of naïve CD4+ and CD8+ T cell in the immune restoration [12, 27]. Researchers also found that IL-7 can promote the growth of B cell, the differentiation and proliferation of pre-B cell, and then B cells transform into plasma cells after stimulating by antigen to synthesize and secret the Igs [4, 29]. Last but not the least, IL-7 could improve significantly the number of cells which can express typical dendritic cells' mark, and enhance the activity of macrophage cytotoxic factors, and inducing the secretion of cytokines, like interferon (IFN), Tumor necrosis factor (TNF) [3]. IL-7 must be bind to its receptor (IL-7R) that can produce biological effect, and after the combination of IL-7 and IL-7R, there are several signaling pathway be activated, mainly include the Janus Kinase/Signal Transducer and Activator of Transcription (JAK/ STAT) pathway and the Phosphatidylinositol-3-kinase (PI3K)/ AKT pathway, but the mechanism underlying IL-7 mediated signaling pathway in chickens is only partially understood [23].

To sum up, our results demonstrated that probiotics could improve the humoral immunity in respiratory and digestive tracts of chickens, and enhance the immunological response to ND vaccine. The expression of IL-7 mRNA, which probably reflects the humoral immunity of MALT in chickens, and it should have a better synergistic promoting effect on the MIS response to ND vaccine, it may serve as an indicator for the mechanism of probiotics research.

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References


