

ISSN 0215-8930

MEDIA

Kedokteran Hewan

Veterinary Medicine Journal



MKH (Vet.Med.J.)	Vol. 25	No. 2	Pp. 67-144	Surabaya, May 2009	ISSN 0215-8930
------------------	---------	-------	------------	--------------------	----------------

Accreditation by Director General of Higher Education, No. 108/Dikti/Kep/2007, August 23, 2007

Veterinary Medicine Journal

Vol. 25, No. 2, May 2009

Veterinary Medicine Journal contains scientific literature in the field of
Veterinary Science and Animal Husbandry
it was first published in 1985 with a frequency published three times a year in the month:
January, May and September

Composition of the Board Editor

Chief Editor:

Kusnoto

Secretary:

Erma Safitri

Treasurer:

Lilik Maslachah

Ads and subscriptions:

Boedi Setiawan

Executive Editor:

Sri Subekti Bendryman Soedjoko

Sri Agus Sudjarwo

Suwarno

Epy Muhammad Luqman

Translation Team:

Mas'ud Hariadi

Suzanita Utama

Muchammad Yunus

Mustofa Helmi Efendi

Editor supervisor:

Susilowati

Address: Faculty of Veterinary Medicine, Airlangga University
Kampus "C" Unair, Jl. Mulyorejo Surabaya 60115
Telp. (031) 5992785; 5993016; Fax. (031) 5993015
e-mail: mkh_ua@yahoo.com

Account: Tabungan Bisnis Mandiri No. 14100-07144132 (a.n. Media Kedokteran Hewan)

Veterinary Medicine Journal published by the Association of Indonesia Veterinarians (PDHI)
and the Faculty of Veterinary Medicine, Airlangga University, Surabaya

Accreditation by Director General of Higher Education, No.108/Dikti/Kep/2007.

Veterinary Medicine Journal

Vol. 25, No. 2, May 2009

AUTHOR GUIDELINES

1. General Regulation
 - a. Veterinary Medicine Media/MKH publishes research articles, reviews and case reports in the scopes of veterinary medicine and animal husbandry.
 - b. Manuscript/articles must be original and may not have been published. Manuscripts accepted for publication in Vet. Med. J. should not be submitted for any other publication.
2. Writing Standard
 - a. Manuscripts are typed in double-line spacing except for title, abstract, figure/table legends, table content, references and appendices.
 - b. New paragraphs use first line indentation by 0.3".
 - c. Standard Font for manuscript writing is 11 point Book Antiqua.
 - d. Manuscript should be printed on A4 HVS paper.
 - e. Manuscript could be submitted in Indonesian or English.
 - f. Tables / Figures must be very contrast and scanning files (photograph) must be enclosed separately in **JPG format**. Table/Figure Legends or other notes in Appendices should be typed in single-line spacing.
3. Manuscript writing guidelines
 - a. The whole manuscript should be composed of 12–14 pages.
 - b. Topics (Title, Author Names, Abstract, Introduction, Methods, etc.) should be written in Title Case, not Uppercase (all capitals) and are to be left-justified.
 - c. Manuscript should be written following this order: Title, Author names and Identity, Corresponding Address, Abstract with Keywords, Introduction, Methods, Results and Discussion, Conclusion, Acknowledgement (if any), References and Appendices (if any).
 - d. Title must be concise, specific, written in full (not abbreviated), informative, written in Indonesian and English.
 - e. Author names are placed beneath the title, author identity and corresponding address should be clear, written in full and placed after the author names.
 - f. Abstract is written in Indonesian and English, single-spaced and not exceed 200 (two hundred) words.
 - g. Keywords is placed after the abstract, list up to 5 (five) words.
 - h. Methods include equipments/materials which were used (particularly the specific ones), research procedure and statistical analysis (if used).
 - i. References should be listed in alphabetical order without numbering. Journal names should be abbreviated according to the system used by each journal. References should be single spaced with hanging indentation by 0.3" and before spacing by 3.6 pt. References should be composed of 60 % scientific articles and 40 % scientific text books. Below are the examples of references style to journals and books.

Roitt I, Brostoff J, and Male D. 1996. Immunology. 4th Ed. Blackwell Scientific Pub. Oxford.

Staropoli I, Clement JM, Frenkiel MP, Hofnung M, and Deuble V. 1996. Dengue-1 virus envelope glycoprotein gene expressed in recombinant baculovirus elicits virus neutralization antibody in mice and protects them from virus challenge. Am.J. Trop. Med. Hygi. 45: 159-167.
4. Manuscripts submission could be performed at any time by sending 3 print copies and one soft copy in a CD to Media Kedokteran Hewan, Faculty of Veterinary Medicine, Airlangga University, Kampus C Unair, Jalan Mulyorejo, Surabaya 60115, Telephone 031-5992785; 5993016; Fax. 031-5993015; e-mail: **mkh_ua@yahoo.com**
5. The editors reserve the right to
 - a. Publish manuscript/article without revision
 - b. Publish manuscript/article with revision
 - c. Reject manuscript/article
6. The editors are not responsible for all statements in published manuscripts.
7. Accepted manuscript for publication requires author to cover publishing and delivery cost.
8. Authors / subscribers may send publication or subscription cost by bank-transfer to **Media Kedokteran Hewan FKH UNAIR, on Tabungan Bisnis Mandiri, account no. 1410007144132.**
9. All editorial decisions are final; no further correspondence will be responded.

Veterinary Medicine Journal

Vol. 25, No. 2, May 2009

Published every 4 months, in January, May and September

Acknowledgments

Editors, writers and readers of Veterinary Medicine Journal award and thanks the highest to the experts below, as contribution partners who have examined all the good writing that was published or rejected according to the recommendations submitted to the editor of Volume 25 No.2, May 2009 edition.

Prof.Dr. Aulani'am, drh.,DEA (Faculty of Mathematic and Natural Science, Brawijaya University)

Prof. Dr. Bambang Sektiari, drh., DEA (Faculty of Veterinary Medicine, Airlangga University)

Prof. Dr. Fedik Abdul Rantam, drh. (Faculty of Veterinary Medicine, Airlangga University)

Prof. Dr. Setiawan Koesdarto, drh., MSc. (Faculty of Veterinary Medicine, Airlangga University)

Prof. Dr. Imam Mustofa, drh., M.Kes. (Faculty of Veterinary Medicine, Airlangga University)

Veterinary Medicine Journal

Vol. 25, No. 2, May 2009

Published every 4 months, in January, May and September

CONTENS

		Page
13	The Role of TLR 2,4 and Characteristics as Signaling of Immuns System by Chickens Infected Avian Influenza Virus (Rantam et al.)	74 - 81
14	Characterization and the Increase of Chicken Interferon-gamma Production as a Measure of T-cell Responses to <i>Eimeria tenella</i> Antigens (Yunus et al.)	82 - 89
15	Characterization of Specific Protein of <i>Toxocara canis</i> for the Development of Diacnostic by Antibodies Examination of Case Toxocariasis (Kusnoto)	90 - 95
16	Study of Skin Premature Aging Induced by Ultraviolet B Light Irradiation in <i>Balb/C</i> Mouse Model (Purwandari and Widyarini)	96 - 103
17	Pharmacokinetic Profile and Pharmacokinetic/Pharmacodynamic (PK/PD) Parameter of Doxycycline in Broiler Plasma and Tissues after Single Dose Intravenous Administration (Wijayanti et al.)	104 - 109
18	Goat's Immune Response for Prolactine from Isolation in Moulting Duck (Safitri et al.)	110 - 115
19	Detection of Enterotoxin Genes of <i>Staphylococcus aureus</i> Isolated from Waste Slaughtered House and Carcass of Chickens (Khusnan et al.)	116 - 121
20	Reaction of Thyroxin, Hematocrit, Haemoglobin on Reducing Feedstuff and Drinking Water (Achmad)	122 - 125
21	<i>In vitro</i> Anticestode Activity of Painted Nettle Leaves Extract to <i>Hymenolepis microstoma</i> : Observation Using SEM (Ridwan et al.)	126 - 133
22	Suckling Behaviour of Bali Calves with Dams Different in Milk Production (Henderiana et al.)	134 - 139
23	Comparison between Calipers and Ultrasonography for Measurement of Testicular Volume and Mass in Dogs (Junaidi)	140 - 146

Characterization and the Increase of Chicken Interferon-gamma Production as a Measure of T-cell Responses to *Eimeria tenella* Antigens

Yunus, M.¹, Suwarno², Suwanti, L.T.¹ and Mufasirin¹

¹Department of Veterinary Parasitology, ²Department of Veterinary Microbiology, Faculty of Veterinary Medicine, Airlangga University, e-mail: muhyunus_99@yahoo.com

Abstract

Characterization and the increase of chicken interferon-gamma (ChIFN- γ) production were investigated in *Eimeria tenella* (*E. tenella*) infected chicken. Twenty poultry broilers at three weeks old were divided into two groups. The first group was control and those chickens were not infected anything. The second group was infected with 4×10^3 of *E. tenella*. Spleen cells of both the groups were cultured and induced mitogen. This study was to characterize ChIFN- γ protein through extraction technique of supernatant protein of the culture medium of chicken spleen cell that was induced Con-A by SDS-PAGE. The result of characterization showed that molecule weight of ChIFN- γ protein was 17 kDa. Then, Optical Density (OD) of ChIFN- γ protein was detected ELISA reader 450 nm of both *E. tenella* uninfected and infected chickens was 1.29 and 1.95, respectively, whereas, concentration of ChIFN- γ protein of both those groups was 10.312 pg/ml and 15.569 pg/ml, respectively.

Key words: *E. tenella*, chicken interferon-gamma.

Introduction

IFN- γ is a cytokine with a wide range of biological effects including antiviral and macrophage activating capacities and inhibition of the intracellular development of protozoan parasites (Houglum, 1983; Dijkmans and Billiau, 1988; Murray, 1988; Liesenfeld *et al.*, 1996). In mice, elevation in the production of IFN- γ correlates with a genetically-determined disease phenotype; an earlier increase in IFN- γ occurred following primary infection (pi) with *E. vermiformis* in a resistant mouse strain compared to relatively susceptible mice (Wakelin *et al.*, 1993). Although IFN- γ has been shown to play an important role in host defense against *Eimeria* (Rose *et al.*, 1989, 1991; Lillehoj and Choi, 1998), *Leishmania* (Scott, 1991), *Plasmodium* (Schofield *et al.*, 1987) and *Toxoplasma* (Suzuki *et al.*, 1988), the mechanisms of its action have yet to be clarified.

Cytokines and lymphokines have been shown to influence the course of coccidial infections. Cell culture supernatant from concanavalin A-stimulated lymphocytes inhibited the replication of *Eimeria* parasites in MDBK cell cultures (Lillehoj *et al.*, 1989). The same supernatant, when administered to chickens, reduced oocyst production following both *E. acervulina* and *E. tenella* infections (Lillehoj *et al.*, 1989). Supernatant from concanavalin A-stimulated lymphocytes also inhibited the growth of *E. bovis* and

E. papillata in bovine monocyte cultures and activated murine macrophages and a bovine monocyte cell line to kill *E. bovis* parasites (Hughes *et al.*, 1987).

IFN- γ production in chickens has been used as a measure of T-cell responses to coccidial antigens (Byrnes *et al.*, 1993; Martin *et al.*, 1994; Prowse and Pallister, 1989). Lymphocytes from *Eimeria*-infected chickens produced a higher level of IFN- γ when induced with concanavalin A than did lymphocytes from uninfected chickens (Martin *et al.*, 1994). Chicken IFN- γ regulates acquired immunity by activating lymphocytes and enhancing expression of MHC class II antigens (Kaspers *et al.*, 1994). Treatment of MDBK, fibroblast, and epithelial cell cultures with recombinant bovine IFN- γ inhibited *E. tenella* and *E. vermiformis* development (Kogut and Lange, 1989; Rose *et al.*, 1991). Pretreatment of sporozoites with IFN- γ did not affect growth, indicating that IFN- γ alters some aspects of the host cells but not those of the parasites. However, until recombinant chicken IFN- γ become available, the role of IFN- γ in avian coccidiosis remains to be determined.

Recently, the availability of recombinant chicken IFN- γ has led to a better understanding of its physiologic and immunologic roles in chicken coccidiosis (Lillehoj and Choi, 1998; Lowenthal *et al.*, 1997; Song *et al.*, 1997). Chicken recombinant IFN- γ was capable of protecting chick fibroblasts from virus

mediated lysis, induced nitrite secretion from macrophages *in vitro*, and enhanced MHC class II antigen expression on macrophages (Lowenthal *et al.*, 1997). Administration of exogenous recombinant IFN- γ to chickens significantly hindered intracellular development of *Eimeria* parasites and reduced body weight loss (Lillehoj and Choi, 1998). When chicken fibroblast cells transfected with the IFN- γ gene were infected with *E. tenella* sporozoites, significant reductions in parasite intracellular development occurred although the ability of parasites to bind and to invade host cells was not affected (Lillehoj and Choi, 1998). Briefly, the purpose of this study was to characterize and to know the influence of *E. tenella* infection on the increase of chicken interferon gamma as a measure of T-cell responses to coccidial antigens.

Materials and Methods

Animals: Male CP707 broiler, 3-5 weeks old with around 1000 g body weight (BW) were purchased from Poultry Supplier Co. Surabaya, housed in clean cages and fed with a standard diet without coccidiostat and tap water *ad libitum* in room temperature ($24 \pm 1^\circ\text{C}$), under conventional conditions with a 12:12 hr, light: dark cycle. They were kept as outlined in the guide for the care and use of laboratory animals by the Faculty of Veterinary Medicine, Airlangga University.

Parasites: The pathogenic agent used in this study was *E. tenella* obtained from field and routinely maintained in our laboratory by oral passage through CP707 broiler.

Experimental procedures: Twenty CP707 broilers were divided into two groups, each group composed ten chickens. The first group was control group and to those chickens were not infected anything. The second group was infected with *E. tenella* sporulated oocyst. All infective doses of *E. tenella* sporulated oocysts were orally given by 1 ml spuite, as 4×10^3 oocysts/chicken in 1 ml of distilled water. The second infection with the same doses as the first infection was given at the second group to know protective immunity due to the first infection by oocysts production. Fecal pellets were collected from the infected chicken between days 6 until 12 post infection (pi). Spleen cells of both the groups were cultured at 1.7×10^9 cells/ml and placed in 16 well plastic tissue culture test plates, for 48 hr in RPMI-1640 medium containing 2 mM L-glutamine, 10% fetal calf serum, 100 U/ml of penicillin, 100 U/ml of streptomycin and supplemented with 1.2 $\mu\text{g/ml}$ Concanavalin A (Con-A, Sigma, St. Louis, MO), incubated at 37°C and 5% CO_2 . Supernatant of tissue culture was predicted to contain ChIFN- γ isolated and characterized by SDS-PAGE and confirmed with ELISA and Dot Blot.

Data Analysis: Oocysts output was analyzed descriptive. The character of ChIFN- γ expressed by molecule weight and described with comparison analysis, concentration and OD of ChIFN- γ were statistically analyzed using student *t*-test and a *p* value below 0.05 was considered significant (Steel and Torrie, 1995).

Results and Discussion

The pattern and total of oocyst output of *E. tenella* infection: The temporal pattern of oocyst output per day confirms those previously reported (Stiff and Bafundo, 1993) with this isolate of *E. tenella*. Oocyst first appeared on the seven days pi, then reached peak on the 10 days pi before numbers declined rapidly and the fewest oocysts were detected on 12 days pi. Basically, the pattern of daily oocyst output was clearly seen in the first infection of infected chickens groups, but in the second infection, oocyst output per day as well as totally were significantly very lower and very few than the first infection (Figs. 1 and 2).

The total numbers of oocysts output of the 1st *E. tenella* infected chickens in this study was $[23.5 \pm 5.3] \times 10^6/\text{chicken}$ and the 2nd *E. tenella* infected chickens $[1.1 \pm 0.2] \times 10^6/\text{chicken}$, the period of patency was $[12.4 \pm 0.7]$ days (Fig. 2). Total of oocysts output in the 2nd *E. tenella* infected chickens were significantly decreased ($p < 0.01$) about 95 % compared with the 1st *E. tenella* infected chickens (Fig. 2). Clinical signs (such as anemia, anorexia) of the 2nd *E. tenella* infected chickens were slighter than the 1st *E. tenella* infected chickens. Consistency and colour of feces appeared normal in the 2nd *E. tenella* infected chickens compared the 1st *E. tenella* infected chickens. Pathological changes and lesion score of cecum in the 2nd *E. tenella* infected chickens more slight compared than the 1st *E. tenella* infected chickens (Unpublished data). Endogenous development of *E. tenella* (schizogony and gametogony) in the 2nd *E. tenella* infected chickens was suppressed and/or incompleated undergone. Several generations of schizont appeared degenerated consequently unbreak schizont, damaged cecal mucosa epithelial cell was not occurred and automatically there were no bleeding in cecum. Many abnormal endogenous developments of parasites such as gametogony result in disturbing syngamy of microgamete and macrogamete. Thus, oocyst formation was not perfectly continued. In contrast, endogenous development of parasites in the 1st *E. tenella* infected chickens occurred well and no inhibition. Infection with one species of *Eimeria* induces protective immunity in the host that is long lasting and exquisitely specific to that particular parasite (Yun *et al.*, 2000). While a large number of

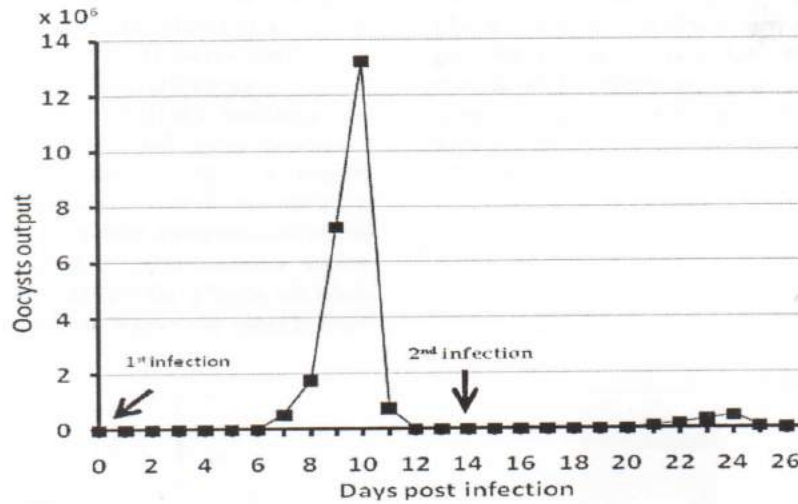


Figure 1. The pattern of oocyst output per day of 1st and challenge infection of *E. tenella* infected chickens. The initially oocyst output on the 7th day, then to peak level the 10th day and for limit around 12 days post infection. Each value of oocyst output per day represents mean of 10 chickens.

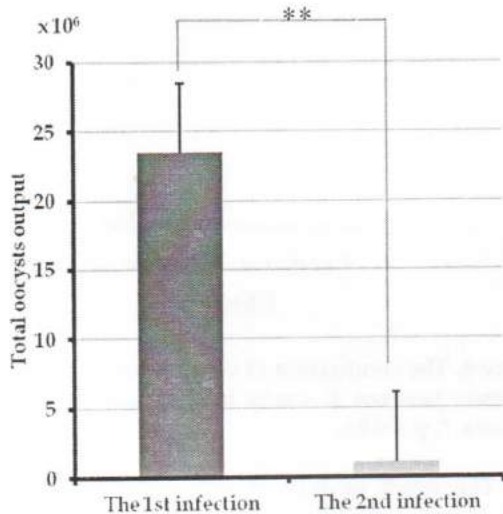


Figure 2. The differences among 1st and challenge infection on total of oocysts output from *E. tenella* infected chickens. From left to right, each column represents 1st and challenge infection status. Each value represents mean \pm SD of 10 chickens. ** $p < 0.01$.

inoculating oocysts is generally required to generate an immune response against *Eimeria*, some exceptions have been noted, e.g. *E. maxima* is highly immunogenic and requires only a small number of oocysts to induce almost complete immunity. The early endogenous stages of the parasite life cycle are considered to be more immunogenic than the later sexual stages

(Yun *et al.*, 2000) although Wallach *et al.* (1990 and 1995) showed that immunization with recombinant gamete associated antigen induced partial protection against challenge infection. Studies using oocysts irradiated to prevent intracellular development, but not invasion, demonstrated partial protection against challenge infection, thereby suggesting that sporozoites may also be immunogenic (Jenkins *et al.*, 1991).

Spleen cell culture of uninfected and infected chickens in induction of Con-A for isolation of ChIFN- γ protein. Splens of *E. tenella* infected chickens and *E. tenella* uninfected chickens were cultured in 16 well plastic tissue culture test plates, for 48 hr in RPMI-1640 medium containing 2 mM L-glutamine, 10% fetal calf serum, 100 U/ml of penicillin, 100 U/ml of streptomycin and supplemented with 1.2 μ g/ml Concanavalin A (Con-A, Sigma, St. Louis, MO), incubated at 37°C and 5% CO₂.

The daily check up on the growth of spleen cell cultured appeared the good growth and proliferation of both spleen cells infected and uninfected chickens (Fig. 5). Additional mitogen (Con-A) be able to induce release and production of IFN- γ by lymphocytes. Lymphocytes from *Eimeria*-infected chickens produced a higher level of IFN- γ when induced with Con-A than did lymphocytes from uninfected chickens (Martin *et al.*, 1994).

The examination of concentration of Ch-IFN- γ that expressed in supernatant of spleen cell cultured with Con-A induction of *E. tenella* infected chickens compared *E. tenella* uninfected chickens using indirect ELISA showed significantly differences ($p < 0.05$)

(Fig. 4) which concentration of Ch-IFN- γ *E. tenella* infected chickens increased 46% higher than *E. tenella* uninfected chickens that was 15.569 pg/ml and 10.312 pg/ml, respectively. Pattern of optical density (OD) of both *E. tenella* infected chickens and *E. tenella* uninfected chickens was also same as pattern of concentration that was 1.95 and 1.29, respectively (Fig. 5).

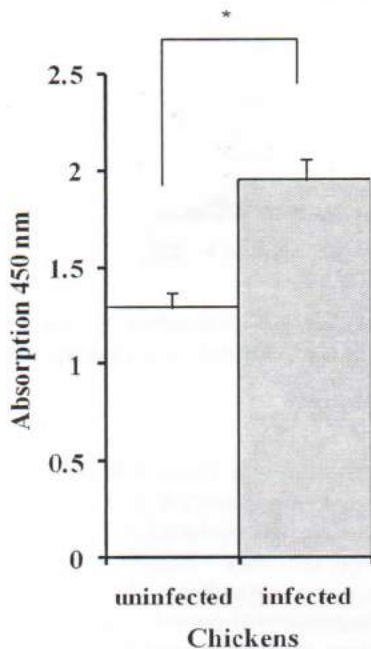


Figure 3. The comparison of optical density (OD) of ChIFN- γ protein between *E. tenella* infected and uninfected chickens. *, $p < 0.05$.

Cytokines (i.e. IFN- γ) are proteins that are naturally produced by the body's immune system immediately following infection or vaccination, resulting in protection from disease. Cytokine activities during avian coccidiosis are major roles (Choi *et al.*, 1999; Lillehoj and Choi, 1998). Chicken IFN- γ regulates acquired immunity on *Eimeria* infection by activating lymphocytes and enhancing expression of MHC class II antigens (Kaspers *et al.*, 1994; Lillehoj, 1989). IFN- γ production in mice (Rose *et al.*, 1991) and chickens (Martin *et al.*, 1994; Yun *et al.*, 2000) has been used as a measure of T cell responses to coccidial antigens. Study by Yun *et al.*, (2000) showed that production of IFN- γ was high in intestine tissue of coccidia development.

IFN- γ mRNA expression is significantly increase in infected chickens compared uninfected chickens. Correlation of immunity on disease with local IFN- γ production early indicates important roles of IFN- γ in protective immunity. Level of IFN- γ increased in SC

compared TK chickens and this cytokine appeared in intestine particularly in circulation (Yun *et al.*, 2000).

Characterization of Ch-IFN- γ Protein by SDS-PAGE. In supernatant of chicken spleen cell cultured was identified Ch-IFN- γ protein by SDS-PAGE and confirmed using Dot Blot. Identification Ch-IFN- γ protein of *E. tenella* infected as well as uninfected chickens was shown several bands in variety several bands in conformity with marker in certain molecule weight standard (Fig. 7). Based on measurement of molecule weight, all supernatants showed the same band of molecule weight of 17 kDa.

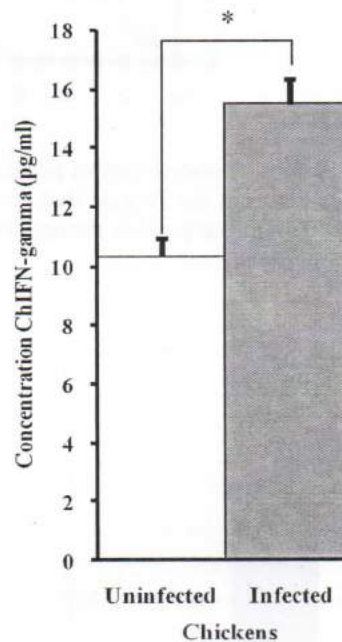


Figure 4. The comparison of concentration of ChIFN- γ protein between *E. tenella* infected and uninfected chickens. *, $p < 0.05$.

The result analysis of SDS-PAGE to be sure more specific of Ch-IFN- γ protein confronted with immunoblotting (Dot Blot) (Fig. 8). Characterization of Dot Blot showed protein was specific for monoclonal antibody anti Ch-IFN- γ , except panel A (antigen, mouse IFN- γ). Panel B and C expressed Ch-IFN- γ protein that known by monoclonal antibody anti Ch-IFN- γ . In *E. tenella* infected chickens was shown clear spot which intensity was stronger than *E. tenella* uninfected chickens. Briefly, concentration of Ch-IFN- γ protein of *E. tenella* infected chickens higher than *E. tenella* uninfected chickens, however both those groups had strong antigenicity which proved the result of examination both showed positive reaction.

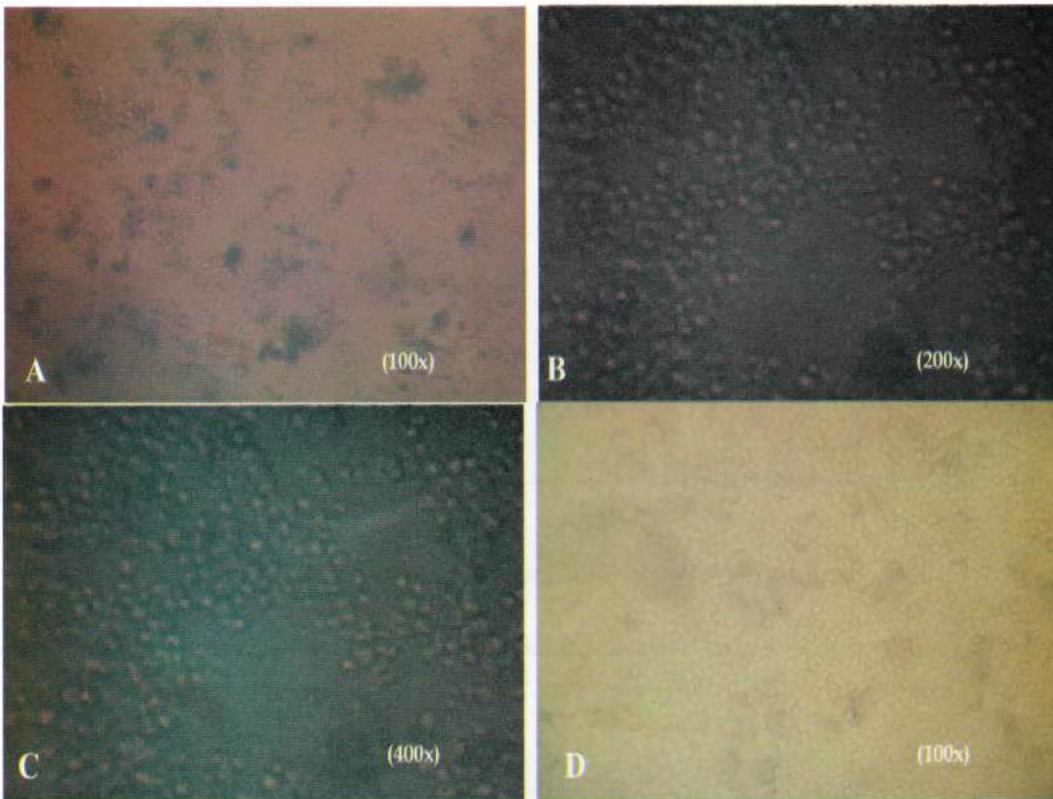


Figure 5. Spleen cell cultured and cell number in initial culture $\sim 1.7 \times 10^9$ /ml (panel A, B, C), 48 hours after undergoing development (panel D).

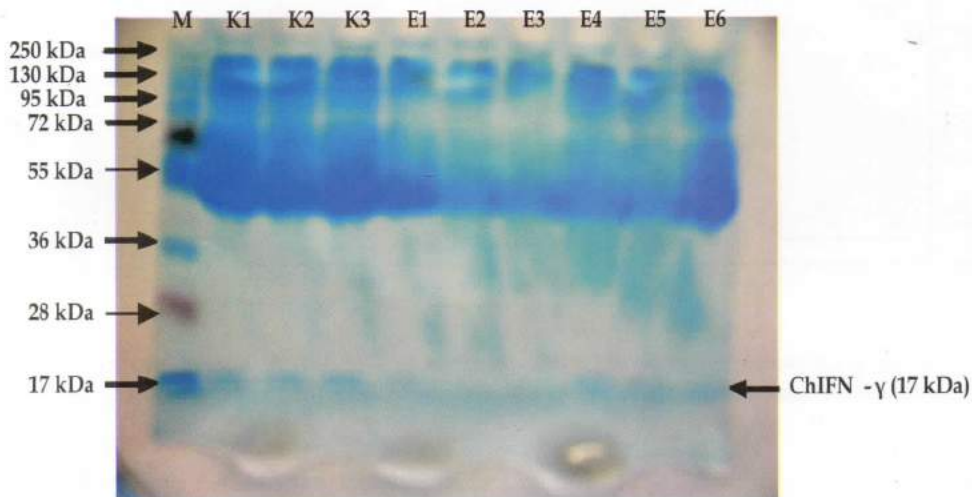


Figure 6. Molecule weight of ChIFN- γ protein. M: marker, K1-3: *E. tenella* uninfected chickens, E1-6: *E. tenella* uninfected chickens (Song, *et al.*, 2007, molecule weight of ChIFN- γ protein recombinant is around 17-18 kDa).

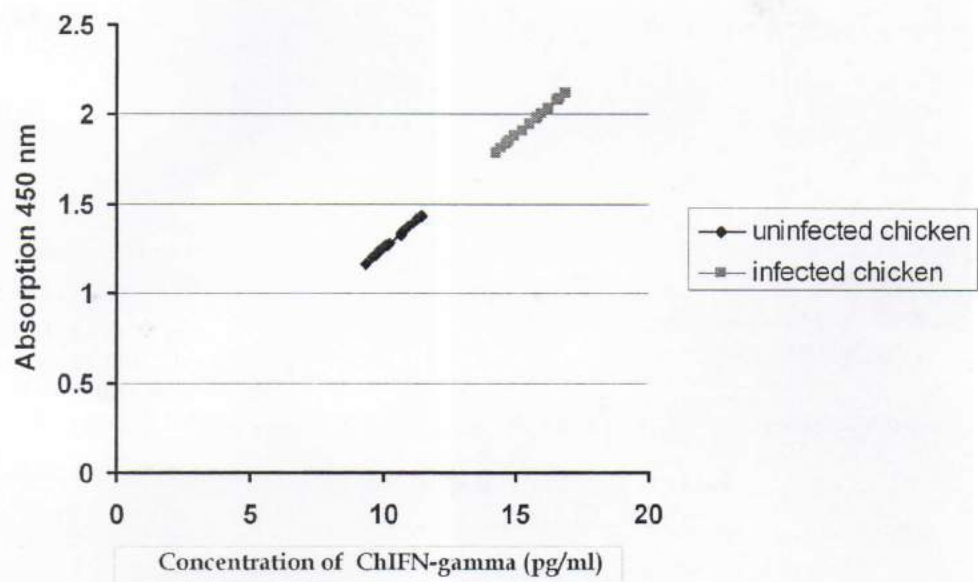


Figure 7. Comparison optical density and concentration ChIFN- γ between *E. tenella* infected and uninfected chickens.

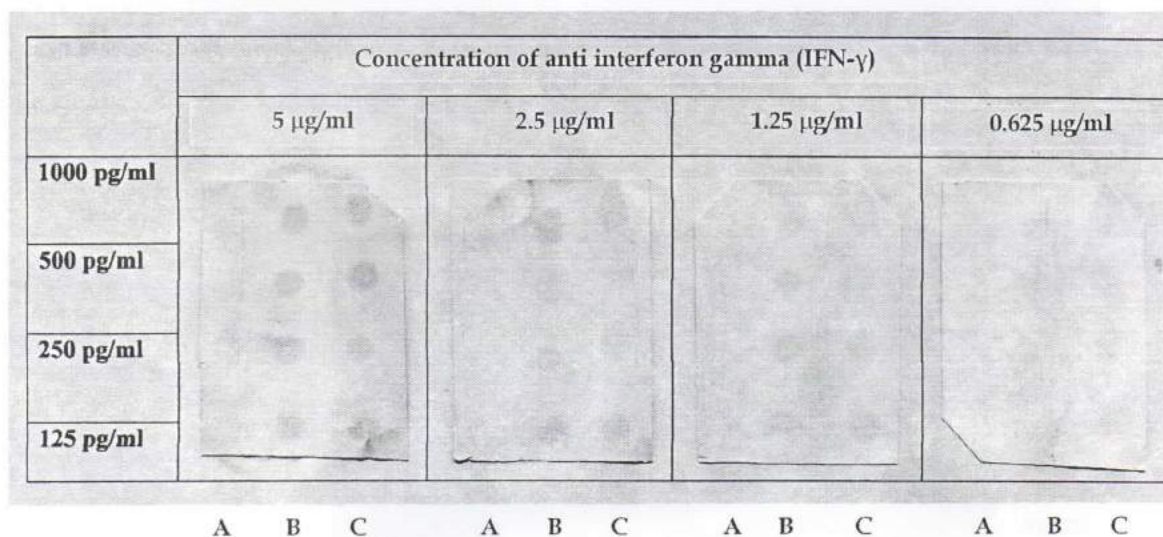


Figure 8. Immunoblotting for characterization of ChIFN- γ protein by *Dot Blot*. A, mouse interferon-gamma; B, *E. tenella* infected chickens interferon-gamma; C, *E. tenella* uninfected chickens interferon-gamma.

Conclusions

Measurement of molecule weight of chicken interferon gamma protein of *E. tenella* uninfected and infected chickens showed the same band of molecule weight of 17 kDa. Lymphocytes from *Eimeria*-infected chickens produced a higher level of IFN- γ

when induced with Con-A than lymphocytes from uninfected chickens. Moreover, the concentration of Ch-IFN- γ that expressed in supernatant of spleen cell cultured with Con-A induction of *E. tenella* infected chickens higher compared *E. tenella* uninfected chickens.

References

- Byrnes S, Eaton R, and Kogut M. 1993. In vitro interleukin-1 and tumor necrosis factor-alpha production by macrophages from chickens infected with either *Eimeria maxima* or *Eimeria tenella*. *Int. J. Parasitol.* 23: 639-645.
- Choi KD, Lillehoj HS, and Zalenga DS. 1999. Changes in local IFN- γ and TGF- β 4 mRNA expression and intraepithelial lymphocytes following *E. acervulina* infection. *Vet. Immunol. Immunopathol.* 71: 263-275.
- Dijkmans R, and Billiau A. 1988. Interferon gamma: a master key in the immune system. *Curr. Opin. Immunol.* 1: 269-274.
- Houglum JE. 1983. Interferon: mechanisms of action and clinical value. *Clin. Pharm.* 2: 20-28.
- Hughes HPA, Speer CA, Kyle JE, and Dubey JP. 1987. Activation of murine macrophages and a bovine monocyte cell line by bovine lymphokines to kill the intracellular pathogens *Eimeria bovis* and *Toxoplasma gondii*. *Infect. Immun.* 55: 784-791.
- Jenkins MC, Castle MD, and Danforth HD. 1991. Protective immunization against the intestinal parasite *Eimeria acervulina* with recombinant coccidial antigen. *Poult. Sci.* 70: 539-547.
- Kaspers B, Lillehoj HS, Jenkins MC, and Pharr GT. 1994. Chicken interferon-mediated induction of major histocompatibility complex class II antigens on peripheral blood monocytes. *Vet. Immunol. Immunopathol.* 44: 71-84.
- Kogut M, and Lange C. 1989. Interferon- γ mediated inhibition of the development of *Eimeria tenella* in cultured cell. *J. Parasitol.* 75: 313-317.
- Liesenfeld O, Kosek J, Remington JS, and Suzuki Y. 1996. Association of CD4+ T-cell-dependent, interferon-gamma-mediated necrosis of the small intestine with genetic susceptibility of mice to peroral infection with *Toxoplasma gondii*. *J. Exp. Med.* 184: 597-607.
- Lowenthal JW, York JJ, and O'Neil TE. 1997. In vivo effect of chicken interferon- γ during infection with *Eimeria*. *J. Int. Cytol. Res.* 17: 551-558.
- Martin A, Lillehoj HS, Kaspers B, and Bacon LD. 1994. Mitogen-induced lymphocyte proliferation and interferon production induced by coccidian infection. *Avian Dis.* 38: 262-268.
- Murray HW. 1988. Interferon-gamma, the activated macrophages, and host defense against microbial challenge. *Ann. Intern. Med.* 108: 595-608.
- Prowse SJ, and Pallister J. 1989. Interferon release as a measure of the T-cell response to coccidial antigens in chickens. *Avian Pathol.* 18: 619-630.
- Rose ME, Wakelin D, and Hesketh P. 1989. Gamma interferon controls *Eimeria vermiformis* primary infection in BALB/c mice. *Infect. Immun.* 57: 1599-1603.
- Rose ME, Smith AL, and Wakelin D. 1991. Gamma interferon-mediated inhibition of *Eimeria vermiformis* growth in cultured fibroblasts and epithelial cells. *Infect. Immun.* 59: 580-586.
- Schofield L, Villaquiran J, Ferreira A, Schellekens H, Nussenzweig R, and Nussenzweig V. 1987. Gamma interferon/CD8+ T-cells, CD8+ T-cells and antibodies required for immunity to malaria sporozoites. *Nature.* 330: 664-666.
- Scott P. 1991. IFN-gamma modulates the early development of Th1 and Th2 responses in a murine model of cutaneous leishmaniasis. *J. Immunol.* 47: 3149-3155.
- Song KD, Lillehoj HS, Choi KD, Zalenga D, and Han JY. 1997. Expression and functional characterization of recombinant chicken interferon- γ . *Vet. Immunol. Immunopathol.* 58: 321-333.
- Steel, R. G. D. and Torrie. 1995. Prinsip dan Prosedur Statistika. Penerbit P.T. Gramedia Pustaka Utama Jakarta.
- Stiff MI, and Bafundo KW. 1993. Development of immunity in broilers continuously exposed to *Eimeria* sp. *Avian Dis.* 37: 295-301.
- Suzuki Y, Orellano MA, Schreiber RD, and Remington JS. 1988. Interferon- γ : the major mediator of resistance against *Toxoplasma gondii*. *Science.* 240: 516-518.
- Wakelin D, Rose ME, Hesketh P, Else KJ, and Grecis, RK. 1993. Immunity to coccidiosis: genetic influences on lymphocyte and cytokine responses to infection with *Eimeria vermiformis* in inbred mice. *Parasite Immunol.* 15: 11-19.
- Wallach M, Pillemer G, Yarus S, Halabi A, Pugatsch T, and Mencher D. 1990. Passive immunization of chickens against *Eimeria maxima* infection with a monoclonal antibody developed against a gametocyte antigen. *Infect. Immun.* 58: 557-562.

Wallach M, Smith NC, Petracca M, Miller CM, Eckert J, and Braun R. 1995. *Eimeria maxima* gametocyte antigens: potential use in a subunit maternal vaccine against coccidiosis in chickens. *Vaccine*. 13: 347-354.

Yun CH, Lillehoj HS, and Lillehoj EP. 2000. Intestinal immune responses to coccidiosis. *Develop. Comp. Immunol.* 24: 303-324.