Mucosal Mast Cells Response in the Jejunum of Ascaridia galli-Infected Laying Hens

Darmawi*a, U. Balqis*a, M. Hambal*a, R. Tiuria*a, Frengki*a, & B. P. Priosoeryanto*b
*aFaculty of Veterinary Medicine, Syiah Kuala University
Jln. Tgk. H. Hasan Krueng Kale No. 8 Darussalam, Banda Aceh 23111, Indonesia
*bFaculty of Veterinary Medicine, Bogor Agricultural University
Jln. Agatis, Kampus IPB Dramaga Bogor 16680, Indonesia
(Received 26-06-2013; Reviewed 06-09-2013; Accepted 17-09-2013)

ABSTRACT

Intestinal defense mechanism against helminthes parasitic nematode to be associated with mucosal mast cells reaction. The aim of this research was to examine the effect of infection by Ascaridia galli parasite to trigger mucosal defense based on mucosal mast cells response in laying hens. Amount of ten head laying hens 12-wk old were divided into two groups containing five chickens of each. The first group, chickens were left as un-infected controls. The second group, chickens were infected orally with 1,000 embryonated eggs of A. galli. Mucosal mast cell responses were assayed by in situ jejunal mast cell counts in stained serial histological sections with Alcian blue (pH 0.3) and Safranin-O (pH 0.1) of the jejunum. Mucosal mast cells response were observed and counted on days post infection. The result showed that A. galli infection was able to increase significantly (P<0.05) mast cells response. This research concluded that the A. galli infection can trigger the involvement of mucosal mast cells response in jejunal defense of laying hens against parasitic diseases caused by A. galli.

Key words: Ascaridia galli, laying hen, mucosal mast cell

INTRODUCTION

Ascaridia galli is one of the major nematode parasites causing substantial economic losses in domesticated chickens farming worldwide. The normal habitat of the parasitic stages of A. galli is in the small intestine of poultry. Sexual reproduction of A. galli nematodes leads to release of eggs through feces into the environment. These eggs to become embryonated and resulting larval transmission stages reinfect the host by oral ingestion. Although A. galli adult worms survive in the lumen of intestine, Luna-Olivares et al. (2012) found that A. galli larvae had penetrated in the epithelium and were positioned in the lamina propria. To undertake “histotrophic phase”, the A. galli larvae were localized within the epithelium or in the lumen of the crypts at 3 d post...
infection. Thus, *A. galli* was capable of developing in the lumen but could also enter the lining intestinal with migration to the tissue.

Characteristic immune responses occur during parasite infection in the small intestine. It has long been known that the mast cells are contributed in defense mechanism against parasite infection, particularly in locations that are in close contact with the external environment such as intestines (Urb & Sheppard, 2012), with these responses peaking at the time of parasite expulsion from the host (McDermott et al., 2003), but the mucosal mast cells precise mechanisms involved have remained obscure. Infection induces mucosal mast cells degranulation in the intestinal that is considered to be a host defense mechanism against the parasite. In support of this hypothesis, various authors described that mast cells involved in mucosal defense mechanism. Li et al. (2004) showed that mast cells are important for rapidly controlling murine infection with the protozoan parasite *Giardia lamblia*. Okayama & Kawakami (2006) described that the number of mast cells in inflamed tissue can be regulated by proliferation, migration, and survival (apoptosis). Anthony et al. (2007) explained that many of effector cells are activated in response to most helminth infections including mast cells. Mucosal mast cells contribute to expulsion of a number of gastrointestinal nematode parasites (Afferson et al., 2012).

Mast cells differentiate from multipotent hematopoietic stem cells in the bone marrow that give rise to committed mast cell progenitors in the blood and are recruited to tissues, where they mature. Franco et al. (2010) suggested that mast cell development is most closely associated with the megakaryocyte/erythocyte lineage. The lambs infected with *Haemonchus contortus* had significantly greater numbers of mucosal mast cells in abomasal mucosa of lambs (Shakya et al., 2009). Ortolani et al. (2013) investigated the greater number of mucosal mast cells was able to decrease worm burden in the abomasums of sheep infected with *H. contortus*. Ohan & Nawa (2004) observed that mastocytosis occurred in jejunal sections of mice challenged with *Strongyloides venezuelensis*.

Currently, little information is available describing the effects of *A. galli* infection on the mucosal defense regarding mast cells response of small jejunal of laying hens. Reference to mucosal mast cells mediate *Trichinella spiralis*, the nematode resides within enterocytes of the jejunum, expulsion from the intestine of mice has been reported (Ierna et al., 2005; Suzuki et al., 2008; Afferson et al., 2012); however, De-yuan et al. (2003) reported that the mucosal mast cells in jejunum of chickens infected with *A. galli* were increased with no significantly difference but a remarkable decrease of mast cells in the thymic medulla. Thus, the aim of the current study was to investigate the effects of *A. galli* infection on the mucosal defense based on mucosal mast cells response in jejunal of laying hens. Therefore, we investigated the distribution of mucosal mast cells in laying hens whether these mucosal mast cells are strong associated for mucosal defense in the jejunal against *A. galli* infection.

### MATERIALS AND METHODS

#### Chickens

*Ascaridia galli* adult worms were procured from the intestine of freshly slaughtered chickens. They were brought to the laboratory from local restaurant. Worms were washed with phosphate buffered saline (PBS). For collecting the egg worms, the selected *A. galli* female adult worms were wounded in half of body length using a needle with sharp tip under stereo microscope. The eggs obtained from uteri female adult worms were incubated in sterile aquadestilata at room temperature for 20-31 d till to develop embryonated eggs. The eggs were counted under stereomicroscope for preparing doses of 1,000 embryonated *A. galli* eggs (Darmawi et al., 2007; Balqis et al., 2009).

#### Ascaridia galli-Infected Laying Hens

The laying hens devided into two groups contained five chickens of each. First groups, the chickens were introduced with 0.5 mL PBS. Second group, the chickens were orally infected with doses of 1,000 embryonated *A. galli* eggs contained in 0.5 mL PBS and introduced directly into the oesophagus using a needle with blunt oval tip (Darmawi et al., 2007).

#### Tissue Preparation for Mast Cells Protocol

Intestines were devided into 3 segments, namely duodenum, jejunum, and ileum. Jejunum of laying hens were taken because the normal habitat of the parasitic stages of *A. galli* infection is manly located in the jejunum as described by Luna-Oliveres et al. (2012). Jejunum’s segment was dissected, flushed with cold sterile saline solution, opened longitudinally, and placed, mucosa side up, onto small pieces of blotting paper. The segments were then fixed in 10% buffered normal formalin. This process was performed for each laying hen using sterile instruments for each dissection. Fixed samples were dehydrated in the ascending concentrations of ethanol (50%, 60%, 70%, 80%, 96% (1), 96% (2), and 100%). The samples were cleared in xylol and were embedded in paraffin wax as described by Darmawi et al. (2012) with certain modifications. Three of each histological sections (3-5 μm of thickness) were stained with Alcian blue (pH 0.3) and Safranin-O (pH 0.1)
(Sigma). After washing, sections were counterstained with eosin and mounted. The number of mast cells per 10 villus crypt units (VCUs) was counted on each section. Mast cell counts were performed under light microscopy using an eyepiece square graticule (eyepiece ×10, objective ×40), and data expressed as mean number of mucosal mast cells (MMCs) per VCU as described by previous authors (McDermott et al., 2003; Noviana et al., 2004; Li et al., 2004; Königová et al., 2008) with certain modifications.

Statistical Analysis

The MMC responses in different groups of laying hens were analyzed by the Student t test, where t test was used for comparisons of mast cell numbers. P value of <0.05 was taken to indicate a significant difference.

RESULTS AND DISCUSSION

The simplest interpretation of the finding as seen in Figure 1, is that at least some mast cells go through mucosae in close contact with the external environment, jejunum, mediating the expulsion of A. galli from the intestine. Staining with Alcian Blue–Safranin O revealed mast cells in all the organs examined. Mast cells were identified as blue granules against a pale brown background. Here, we regarded them as mucosal mast cell. Two major subtypes of mast cells have been identified in dogs: connective tissue type, particularly localized in skin, around blood vessels, and in the peritoneal cavity; and mucosal type, which is associated with mucosal surfaces such as those in the gut or airways (Noviana et al., 2004). Regarding the distribution of mast cells within the various locations similar for and support those of Königová et al. (2008), who observed mucosal in the lamina propria mucosae, meanwhile connective tissue mast cells were found in the tela submucosa in the stomach of Mongolian gerbils. In the present study, we recorded degranulated mucosal mast cells in jejunum of laying hens. The present study clearly demonstrate that the embryonated eggs of A. galli stimulated the immune mechanism particularly in mucosal defense by mean of mucosal mast cells response in the jejunal of infected laying hens.

Embryonated eggs of A. galli were hatched in the small intestinal of chickens. The previous study demonstrated that the A. galli larvae were successfully isolated from intestinal of Isa brown laying hens infected with the ascending dose embryonated eggs of A. galli (Darmawi et al., 2007). In this study, we agree with and support those of Luna-Olivares et al. (2012) who found that the normal habitat of the parasitic stages of A. galli is in the profound crypt zone of the mucosa and in the tissue of the jejunum in layer pullets. However, the young larvae grown and survived in the lumen to achieved adult worm. The worm parasitic established in definitive host released antigenic materials in relationship between host-parasite interaction. Previously, investigators reported that the proteins were secreted by females and males adult worm of Syngamus trachea throughout amphial glands, excretory/secretory gland cells, pharyngeal glands (Rica et al., 2005).

The common antigenic substances in many parasites were found in both somatic and excretory/secretory products. This hypothesis supported by many previously reports exist about the role of somatic and or excretory/secretory released by nematode. Our previous investigation showed that excretory/secretory protein released by A. galli could be applied for generating the immune response by mean of immunoglobulin yolk (IgY) antibody formation in egg yolks (Darmawi et al., 2008; 2010) and serum (Darmawi et al., 2013) of immunized laying hens. Karimi et al. (2008) reported that in excretory/secretory and somatic of Ornithobilarzia turkestanicum contained material antigenic substances, similar to the findings of Prasad et al. (2008) successfully purified the fraction of excretory/secretory antigen of H. contortus in sheep. In the study of Smith et al. (2009) analyzed of excretory/secretory products released by Teladorsagia circumcincta. In another study using excretory/secretory antigen of Toxocara vitulorum infective larvae, Hassan & Aziz (2010) noticed that the antigen was able to detect toxocariasis in buffalo calves. Previously, Rokni & Kia (2005) have been reported the excretory/secretory and somatic antigen of Strongyloides stercoralis in human

Figure 1. Mast cells were identified as blue granules. The section was stained with Alcian Blue–Safranin O, m: mucosa, sm: submucosa, ml: muscularis, bar = 200 μm. A= Mast cells in jejunal of uninfected chicken (arrow); B= Mast cells in jejunal of infected with dose 1,000 embryonated of A. galli (arrow).
intestinal nematode infection. More previously, Choi et al. (2003) reported that the excretory/secretory antigen to be a better antigen for a serodiagnosis of clonorchiasis.

There are numerous studies regarding the secretory products of parasites involved in the stimulating of immune response. Excretory/secretory product of T. circumcincta were potentially involved in immunity so targets of local immunoglobulin A (IgA) responses in mucus from sheep rendered immune to infection (Smith et al., 2009). Venom allergen-like (VAL) proteins from gastrointestinal nematode Heligmosomoides polygyrus allow functional testing of the various potentially immunomodulatory (Hewitson et al., 2011). In confirmation of our previous study, we found that IgY antibody formation in egg yolk of laying hens stimulated by excretory/secretory was able to recognized the antigen in the tissue of A. galli (Darmawi et al., 2012).

In this study, the jejunal of both normal and infected chicken groups, mast cells were found in three tissue layers. Large numbers of mast cells were observed in the mucosa. Fewer mast cells were apparent in the submucosa and tunica muscularis/serosa, respectively. We recorded that the mast cells degranulated in lamina propria mucosae. The result showed that in laying hens infected with embryonated eggs of A. galli, significantly more mucosal mast cells were found in the lamina propria mucosae in comparison with mucosal mast cell numbers in laying hens uninfected with the A. galli within the first 14 days p.i. as seen in Table 1. This reflect that the mast cell progenitors are released from the bone marrow into the blood from where they localize to different tissues including in the mucosae throughout the body. Various authors described that multipotent hematopoietic stem cells give rise to committed mast cell progenitors under the influence of growth factors. Once in the tissues, mast cell maturation proceeds, with local factors determining the mature phenotype appropriate for the particular location (Okayama & Kawakami, 2006; Franco et al., 2010).

In normal and infected laying hens, staining with Alcian Blue–Safranin O revealed mast cells in all the organs examined. However, their numbers varied widely and they distributed within the layer of jejunum (Figure 1). In the jejunal tract of both normal and infected laying hens, mast cells were found in lamina propria mucosae tissue layers. Large numbers of mast cells were observed in the jejunum of infected laying hens (Table 1). We described that laying hens infected with embryonated eggs of A. galli accumulated mast cells in the jejunum. Increased numbers of mucosal mast cells are often observed in affected tissues during helminth infections. On the other hand, the number of mucosal mast cells in healthy individuals is stable, but their numbers increase in helminth infection. This phenomenon support that the mast cells play an important role for controlling of A.galli infection. Similarly with many previous reports exist about the role of parasite in attracting mast cells in the tissue. Under various experimental conditions, there were shown that mast cell is important in the immune response in mice against S. venezuelensis (Onah & Nawa, 2004), G. lambia (Li et al., 2004), Fasciola hepatica (Vukman et al., 2013), Acanthocheilonema vitaeae (Ball et al., 2013). The similar phenomenon observed by De-yuan et al. (2003) in chickens against A. galli, Suzuki et al. (2008) in rats against T. spiralis, Königová et al. (2008) in Mongolian gerbils against H. contortus, and the data presented here argue that mast cells are also involved in laying hens against A. galli infection.

In the gastrointestinal defense literature, it is well known that mast cells are key effector cells in mediating worms expulsion from the small intestine, and the increase in parasite loss may therefore be explained by the correlation with the enhanced mastocytosis. Importantly, mast cells can regulate both innate and adaptive immune responses of host defense against helminth infection. Galli & Tsai (2010) explained that mast cells can participate in direct killing of organisms by phagocytosis and reactive oxygen species production. Mast cells can modulate host innate immune responses through the release of granular and secreted mediators. However, Urb & Sheppard (2012) described that mast cells contribute to host defense by mean of to serve as immune sentinel cells to both respond directly to pathogens and send signals to other tissues to modulate both innate and adaptive immune responses. Moreover, our results in this study agree with and support those of Galli et al. (2008) who showed that mast cells are also able to influence disease directly via the release of pro-inflammatory mediators. Therefore, those data suggest that mast cells might be responsible for the gastrointestinal helminth expulsion. Indeed, Suzuki et al. (2008) showed that, in rats as well as in mice infected with T. spiralis, the mast cells other than the IgE antibody are an important effector for worm expulsion.

Upon helminthes stimuli cause synchronous development of new population of mast cells, antibodies-producing plasma cells, and plasma cells synthesizing antibodies prosesing an anti-worms effect. The immuno-

---

Table 1. Mucosal mast cell number/10 villus crypt unit (mean ± SD) in the jejunum from laying hens infected orally with 1,000 embryonated eggs of A. galli

<table>
<thead>
<tr>
<th>Groups</th>
<th>Jejunum layer</th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mucosa</td>
<td>Submucosa</td>
<td>Muscularis</td>
<td>Total</td>
<td></td>
</tr>
<tr>
<td>Uninfected</td>
<td>73.40±16.59*</td>
<td>160.60±20.83*</td>
<td>58.60±10.45</td>
<td>292.60±35.85*</td>
<td></td>
</tr>
<tr>
<td>Infected</td>
<td>140.00±24.78*</td>
<td>212.40±16.29*</td>
<td>64.60±10.19</td>
<td>417.00±21.83*</td>
<td></td>
</tr>
</tbody>
</table>

Note: *Significantly different from uninfected and infected chickens (P<0.05). Results shown are representative of two independent experiments.

MMC/10VCU ± SD

August 2013
globulin recognizes, and binds on worm surface antigen by mean of fragment antibody (Fab) and therefore can potentially respond to opsonized organisms. Meanwhile, fragment crystalline (Fc) of immunoglobulin plays a role in stimulating for mast cell migration. Mast cells can be activated by aggregation of surface Fc receptors, including the cell-surface expression of the high-affinity Fc receptor (FcR) for IgE (FcεRI) (Anthony et al., 2007). The presence of chickens IgY antibody homologous to mammalian IgE. The Fc region of IgY mediates most biological effector functions in the chicken, such as complement fixation, opsonization, and anaphylactic reactions, a function that is attributed to IgE in mammals. In many ways IgY combines the functions associated with mammalian IgG and IgE in the chicken (Hau & Hendriksen, 2005; Kazimierczuk et al., 2005; Lee et al., 2009; Chalghoumi et al., 2009; Diravigam et al., 2011; Darmawi et al., 2012).

Mast cells can be activated by directly interacting with pathogens through pattern recognition receptors. Selective engagement of pattern recognition receptors is also an important mechanism in governing the type of mast cell response. Regarding migration of mast cells in the tissue, Okayama & Kawakami (2006) explained that critical signals for homing and recruitment of mast cells to various tissues are also provided by stem cell factor (SCF). The membrane bound SCF and/or its soluble isomer is chemotactic for mast cells and their progenitors; SCF not only elicits adhesion of mast cells, but also facilitates their proliferation and sustains their survival, differentiation, and maturation. Vukman et al. (2013) reported that the F. hepatica tegumental coat antigen indirectly induces mast cell migration by dendritic cell-derived chemokines. Urb & Sheppard (2012) described that the binding of an antigen by FcεRI-bound specific IgE leads to FcεRI clustering, which in turn induces downstream signalling events and ultimately the release of mediators. Monomeric IgE binding to FcεRI enhances mast cell survival mainly by an autocrine production of IL-3 (Okayama & Kawakami, 2006). The release of these mediators is induced by mast cell degranulation, which in turn is induced by mast cell activation triggered by cross-linking of the FcεRI with an antigen-IgE immune complex. Recently, Ball et al. (2013) described that the product excretory/secretory-62 secreted by filarial nematodes, A. viteae, was an immunomodulator at least in part by inducing the desensitisation of FcεRI-mediated mast cell responses.

Mast cells undergo a degranulation process, release histamine, and proteases, and give rise to globule leukocytes. Various authors explained that histamine and other vasoactive mediators increases vascular permeability and local blood flow, and can act on smooth muscle to increase the expulsion of mucosal parasites (Okayama & Kawakami, 2006; Anthony et al., 2007; Urb & Sheppard, 2012). In addition, Urb & Sheppard (2012) described that mast cell production of chemotactic factors can enhance the recruitment of multiple inflammatory cells including eosinophils (eotaxin), natural killer (NK) cells (interleukin namely IL-8), and neutrophils (IL-8 and tumor necrosing factor namely TNF-α). Mast cell granules contain an array of mediators such as biogenic amines (typically histamine), proteoglycans, and neutral proteases (Noviana et al., 2004). Weller et al. (2005) suggested that leukotriene (LT) B4 released from activated mature mast cells may also have an important autocrine role in regulating the release of mast cell progenitors from the bone marrow and/or their recruitment into tissues before maturation. Interestingly, because of the various mediators they produce, mast cells are potent immune effector cells involved in cuticle degradation and worm expulsion, important modulatory cells that help link innate and adaptive immunity in the fight against helminthes.

**CONCLUSION**

The *A. galli* infection can trigger the involvement of mucosal mast cells response in mucosal defense of jejunum in laying hens against parasitic diseases caused by *A. galli*.

**ACKNOWLEDGMENT**

We wish to thank the Ministry of Research and Technology, Republic of Indonesia for funding the work from the Riset Unggulan Terpadu (No. 47/H11.2/PL/RUT-L/l/2007). We also thank Mr. Sulaeman, Mr. Kosaisih, and Mr. Kasnadi, Faculty of Veterinary Medicine of Bogor Agricultural University, for their expert technical help in the materials preparation.

**REFERENCES**


August 2013
