

~~Shared Expression of Mucin12 contribute in both common antigenicity of host-parasite relationship between *Ascaris Lumbricoides* and Human Small Intestine~~

Authors

Affiliations

\*Corresponding author:

Conflict of interest:

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**Commented [SC ED2]:** The title of the paper should be concise and yet present the main research focus of the paper. We have therefore, revised it to make it impressive and informative for the readers.

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**Commented [SE3]:** Please mention the names of the authors who were involved in conducting this research. Please follow the below format for the names: First Author<sup>1</sup>, Second Author<sup>2\*</sup>, Third Author<sup>3</sup>

If available, please provide the e-mail address of each author.

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An example of the format is given below:

<sup>1</sup>First Department, First University, Address, City, Country Name

**Commented [SE5]:** Please provide the details of corresponding author from among the authors that you have listed above. The corresponding author will be the one who will communicate with the journal after submission, on behalf of remaining authors in the paper.

**Commented [SE6]:** Please provide a declaration of any conflict of interest. If there are no conflict of interest to declare, the authors can mention "There are no conflict of interests to declare."

## Summary/Abstract

*Ascaris lumbricoides* is one of the most common parasites in the world. The purpose of this research study is to focus on the host specificity of human *Ascaris lumbricoides*, which is a parasitic parasite of the small intestine and is also one of the commonest parasites worldwide. As part of this investigation, we examined, at a genetic level, we examined the common antigenicity existing in *A. lumbricoides* and human small intestinal mucosa to unravel the this host-parasite relationship. We obtained three DNA clones after by screening analysis for common antigenicity of using a human colon cDNA library on common antigenicity using and anti-*A. lumbricoides* polyclonal antibodies. After sequencing analysis, we identified one of them is the transmembrane mucin12 gene was identified as a gene of interest. The specific signals of immunostaining with polyclonal anti-mucin12 antibodies were observed in the mucous secretory organs, epidermis, and intestinal canal of *A. lumbricoides*. These signals were disappeared when immunohistochemistry was performed using pre-absorbed polyclonal antibodies with a specific peptide. These results suggested that mucin12 mucin12 was is localized in the mucous secretory organs to in the epidermis of *A. lumbricoides*. Furthermore, we examined the site of mucin12 mucin12 localization on in the host side; the specific mucin12 signals of mucin12 were observed on the mucosal epithelia present around intestinal crypts and villi of the small intestine. Therefore, it is suggested that mucin12 mucin12 is one of the proteins that show the common antigenicity in both parasites, *A. lumbricoides* and its host. It is presumed that adult *A. lumbricoides* live in its their ideal preferred environment, which is the small intestine, by secreting mucin12 mucin12, which is the common antigenicity in the small intestine, to avoid being attacked by the host immune system.

## Key Words:

Human *Ascaris Lumbricoides*, Human human Small small Intestinal intestinal Mucosa mucosa, Mucin12 mucin12, Host host-parasite relationship, sequencing.

## Abbreviations

### 1. Introduction

Infection with the parasite *A. lumbricoides* infection is a disease caused by parasitizing of *A. lumbricoides* and is widespread throughout the world. Many Several cases-infections develop in tropical, subtropical, and temperate regions, although a few

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**Commented [SC ED10]:** Although you have concluded the Abstract by stating the defensive mechanism of the parasite in human small intestine, please note that this is quite a known fact reported in several papers. Hence, the final statements here should reflect the potential of your findings from a diagnostic and prognostic view to appeal the reader's attention.

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also develop in few may also occur in cool regions. In Japan, the there were many prevalence of *A. lumbricoides* patients infection increased to such an extent after World War II and that it was called deemed a national affliction; but however, the number of patients prominently has decreased considerably, thanks to group disinfection, usage of chemical fertilizers, and the improved conditions of living environment. However, increased international travel ing has been causing new problems in recent years because infection sources have increased due to of infected travelers from overseas countries entering Japan.

Infection begins when mature Looking at the life history of *A. lumbricoides* , at first mature eggs were are ingested orally ingested. Once they reach the , and reached to the small intestine, they and hatches, and Hatching the larvae invade the intestinal walls, of the small intestine, and They are then able to enter the systemic circulation the circulatory system via the portal vein and reach the lungs from the heart. They break rupture the alveoli and are is swallowed once more by through the pharynx via the, bronchus, and the trachea, and enabling them to returns to the small intestine again. Finally, they become adults reach maturity in the small intestine, where they mature and remain and stay. The human small intestine is considered to be the ideal habitat for *A. lumbricoides* can be the human small intestine, but although its immunological escape mechanism has not yet been sufficiently clearly elucidated yet. Few gene level studies of the genetics of *A. lumbricoides* exist, and no research has been conducted regarding concerning the common antigenicity between *A. lumbricoides* and the human small intestine has been conducted.

However, some studies have investigated the biochemistry of the intestinal mucosa and its relevance in *A. lumbricoides* infection. As for research concerning *A. lumbricoides* and human intestinal mucosa, Scientists have proved reported the existence of an antibody against nematodes in the blood serum of ulcerative colitis patients through using the Ouchterlony method. The researcher reported the existence of common antigen substances in the cortical laminae and basal laminae of the cuticles on of *A. lumbricoides* side. Antigens common to *A. lumbricoides* , local existence are exist of common antigens with *A. lumbricoides* around the basement membrane in normal human intestinal mucosa, and that a protein with a molecular weight of 41.38 kDa in a crude antigen of normal intestinal mucosa antigen is the substance that owns shares a common antigen with *A. lumbricoides*.

## 2. Materials and mMethods

### 2.1 Preparation of *A. lumbricoides* crude antigen and polyclonal antibody

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~~The~~ adult ~~worm~~ of *A. lumbricoides* were homogenized and ~~the~~ crude antigen was extracted using ~~a~~ homogenizer with PBS at 4°C. ~~The~~ rabbit polyclonal antibody specific ~~for~~ to *A. lumbricoides* ~~were~~ ~~was~~ kindly donated ~~from~~ by Dr. Ishida [5].

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## 2.2 Screening ~~using~~ against a human colon cDNA library

~~In order to obtain~~ We screened for ~~the~~ a cDNA clone that ~~show~~ ~~determines~~ the ~~common~~ antigenicity between ~~host~~ and *A. lumbricoides* and its host. ~~we used~~ the screening method of using a  $\lambda$ -phage method using and polyclonal anti-*A. lumbricoides* antibodies. Briefly, ~~the~~ plaques ~~that~~ ~~were~~ produced by using  $\lambda$ -phage human colon cDNA libraries were transferred to ~~the~~ a nitrate cellulose membrane, which was saturated with 10 mM IPTG. The membranes were screened with polyclonal anti-*A. lumbricoides* antibody-antibodies and ~~some~~ of positive phage clones were ~~obtained~~ ~~identified~~ by developing color using development using 3,3'-diaminobenzidine-4-hydrochlorides (DAB).

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## 3. 2.3 Sequence analysis of the sequence

The positive plaques ~~for~~ reactive against *A. lumbricoides* antibody-antibodies were ~~picked up and~~ transformed into ~~an~~ the *E. coli* host. These clones were constructed as a plasmid, pExcell. The sequences of positive clones were analyzed using the BigDye Terminator v3.1 cycle sequencing kit (Becton Dickinson Biosciences). The plasmid sequences were then compared with ~~the~~ sequences in the GenBank database (National Institute for Biotechnology Information) to determine the identity of genes.

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## 4.2.4 Identifying the site of mucin12 localization in *A. lumbricoides*

The ~~A~~ 14 amino acid synthetic peptide of 14 amino acid (HREQYDVPQEWKRK) from amino acid 396 to 409 of ~~mucin12~~ mucin12 ~~were~~ ~~was~~ synthesized and administered ~~it~~ to rabbits to prepare anti-~~mucin12~~ mucin12 polyclonal antibody-antibodies by Sigma Ggenosys (Hokkaido, Japan). On day 0, ~~The~~ first administration to the rabbit ~~on~~ day 0 was given a 200  $\mu$ g dose of the peptide, ~~then~~ ~~boosted~~ ~~by~~ ~~followed~~ ~~by~~ an additional 100  $\mu$ g in 5 subsequent administrations on days 7, 14, 21, 27, and 42 with incomplete Freund's adjuvant. On day 49, ~~exsanguination~~ was conducted. ~~Furthermore,~~ ~~the~~ serum was purified by ammonium sulfate precipitation.

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**Commented [SE28]:** The "Ethical approval" aspect of this paper is blank. Approval for the use of rabbits for generation of polyclonal serum and human tissue for staining of mucin isoforms is not described in the materials and methods. No major journal will accept these studies without statement of ethical approval.

To identify the localization of mucin12 in *A. lumbricoides*, we conducted immunostaining with sections of *A. lumbricoides*. Three frozen sections were used: ~~one~~ from the

head, the quarter point ~~off from~~ the head, and the quarter point ~~from of~~ the tail of *A. lumbricoides*. The sections of *A. lumbricoides* were ~~cut into sliced with~~ ~~of~~ ~~thickness~~ ~~slices~~ and stained with HE. ~~In order to identify the localization of mucin12 in A. lumbricoides, we conducted immuno-staining with sections of A. lumbricoides.~~ After blocking, the sections were stained with the polyclonal anti-~~mucin12~~ ~~mucin12~~ antibody ~~iesy~~ ( $\times 150$  dilution) and mucin12 ~~antibody~~ ~~antibodies~~ after the absorption treatment with the synthetic peptide described ~~above~~ ~~earlier in this section~~ to identify the specificity of the primary antibody. After washing with PBS with Tween 20, FITC anti-rabbit IgG (Sigma-Aldrich,  $\times 150$  dilution) ~~were was used for as~~ the secondary antibody. ~~The~~ sections were observed ~~by using~~ a laser confocal microscope LSM510 (Carl Zeiss).

To confirm the mucin12 existing in *A. lumbricoides* crude antigen, SDS-PAGE was performed using a ~~crude extract of A. lumbricoides~~. For western ~~blotting~~, crude extract was transferred onto nitrocellulose membrane by a ~~semidry~~ transfer system (ATTO, Japan) and incubated with anti-mucin12 polyclonal antibody ~~iesy~~ ( $\times 1,000$  dilution) for primary antigen after blocking. ~~Furthermore, the~~ The membrane was incubated with HRP ~~a~~ Anti-rabbit IgG for the secondary antibody ~~iesy~~ and the reacted bands were visualized ~~by using the~~ ECL method (GE healthcare) ~~using and~~ X-ray films.

~~In order to~~ To confirm that the results of immunoe-staining were not affected by ~~mucin12~~ ~~mucin12~~ ~~that were derived from of~~ human ~~origin~~, ~~w~~estern ~~blotting~~ was performed with protein extracts of *A. lumbricoides* crude antigen and mouse Embryonic Stem (ES) cell lines using ~~Ra~~ Anti-GAPDH polyclonal antibody ~~iesy~~ that could cross-react with ~~some~~ mammalian GAPDH (human, mouse, and rat, etc.) (Cat: ab9485-25, Abcam,  $\times 1,000$ ) for the primary antibody ~~iesy~~ and HRP anti-rabbit IgG (Sigma-Aldrich, as ~~described~~ above) for the secondary antibody ~~iesy~~.

### 3. Results

#### 1-3.1 Screening ~~with against~~ a human cDNA library

~~As the results of homology search, t~~he fragments of 389 bp ~~was identified~~ ~~matched with~~ ~~as from~~ base 756 bases to 1144 base of human transmembrane ~~mucin12~~ ~~mucin12~~ (AF147790) ~~with and its consistent was~~ 99.5% ~~consistency~~ (Fig. 1).

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#### 2-3.2 Identifying the site of mucin12 localization in *A. lumbricoides*

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In order to identify the localization of ~~mucin12~~ ~~in~~ *A. lumbricoides*, immunohistochemistry was performed with the anti-~~mucin12~~ polyclonal antibody. ~~As the results, the~~ FITC-labeled signals were detected in the mucous secretory organs, epidermis, and intestinal canal (Fig. 2b and 2c). ~~In order to~~ To confirm the specificity of the antibody, the ~~mucin12~~ antibody ~~was~~ ~~were~~ pre-absorbed with a synthetic peptide that was used for the preparation of polyclonal antibody and ~~done the staining as same~~ stained as described ~~above~~ in section 2. As shown in Fig. 2d and 2e, the signals ~~were clearly~~ disappeared. These results ~~were suggested~~ that the protein ~~which that~~ reacts with anti-human ~~mucin12~~ is localized in the mucous secretory organs, epidermis, and intestinal canal ~~in~~ of *A. lumbricoides*.

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#### 4. Discussion

~~It was discussed~~ has been suggested that ~~one of an~~ the parasite's immunological escape mechanism ~~utilized by~~ *A. lumbricoides* is the expression of ~~the matter~~ an antigen very similar ~~to to its~~ a host antigen [6]. However, the parasite's immunological escape mechanisms ~~have have~~ not yet been ~~sufficiently clearly~~ unraveled ~~yet~~ elucidated, so this hypothesis has not been definitively confirmed. Therefore, the ~~researchers~~ authors of the present report decided to ~~reveal~~ examine ~~the~~ this potential immunological escape mechanism using a molecular ~~biological-biology~~ tool. After ~~conducting~~ a screening against a ~~human colon~~ cDNA library ~~of human colon~~ using anti-*A. lumbricoides* polyclonal ~~antibody~~ antibodies and checking for a common antigen, ~~3~~ three positive clones were obtained. ~~One of the~~ Results of the analysis of each sequence showed that one clone possessed high ~~homologous~~ homology with transmembrane ~~mucin12~~ ~~mucin12~~.

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Mucins are ~~mucous~~ glycosylated proteins that are important components of ~~mucous that~~ covering inner cavities such as the trachea, the digestive tract including the stomach and intestines, and the gonads [11].

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~~At the time of writing~~ Currently, 18 different mucin ~~genes~~ proteins have been reported to exist in humans. ~~There are secreted mucins, which are secreted from epithelial cells, and membrane-associated mucins, which have transmembrane sections and exist under cellular membrane-bound conditions.~~

~~We examined transmembrane mucin12 In order to observe their the context~~

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association of transmembrane mucin12 in of the rest of the mucin family. Genomic mucin12 DNA can be found on chromosome 7, along with that of mucin12 including in mucin\_3 and mucin\_17 are existed on same chromosome 7 [20], and mucin12 mucin12 protein is comprised of comprises a total of 588 amino acid sequences.

Mucin 12 Mucin12 was localized in the mucous secretory organs and the epidermis of hypodermis of *A. lumbricoides* by immunostaining with m Mucin12 mucin12 polyclonal antibody (Fig. →). The mucous secretory organs are connected to the epidermis of hypodermis and the lateral cord, and are involved in excretion and form the cluster-like [21]. There is an excretory canal in the lateral cord, and it is connected to the surface of the worm body.

Then, Western blotting was conducted to examine the existence of common antigenicity of mucin12 mucin12 in *A. lumbricoides* crude antigen. A band was detected around 37 kDa. Additionally, a specific band was detected in the protein extracted from cultured mouse cell but not in human *A. lumbricoides* crude antigen.

Researchers The authors would like to further examine the protein, which is cross-reactive with mucin12 mucin12 in the *A. lumbricoides* crude antigen, and the matter substance similar to actin and beta-casein-like protein detected as noted to have common antigenicity, both detected in this study. We would also like And to research which investigate whether any *A. lumbricoides* hosts are immune to avoidance by mucin 12 mucin12, of the mechanism and the other mechanisms of immune avoidance utilized by *A. lumbricoides*.

## 5. Conclusion

In this study, analysis of common antigenicity between *A. lumbricoides* and intestinal mucosa obtained three DNA clones. After analyzing Analysis of each clone sequence indicated, it was clarified that one of them has high homology with transmembrane mucin12 mucin12.

Localization of mucin12 mucin12 was confirmed in the mucous mucosal epithelia present around the intestinal crypts and villi of the human small intestine. These data suggest that expression of mucin proteins by helminths may be one mechanism by through which the helminth parasite evades immunological detection within the mammalian host.

## 6. Acknowledgments

### Ethical approval

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#### Figure Legends

**Figure 1.** Sequence analysis of a newly found-identified clone after screening of of the *A. lumbricoides* cDNA library. -TM12: human transmembrane mucin 12-mucin12.-

**Figure 2.** Identification of the localization for of mMucin12-ucin12 in human-the *A. Lumbricoides-lumbricoides* adult worm. The sSections of *A. Lumbricoides-lumbricoides* were stained with HE (a) and anti-mMucin12-ucin12 polyclonal antibodies. The sections of *A. lumbricoides* were stained with anti-Mucin12 polyclonal antibody that was pre-treatedpretreated with synthetic mMucin12-ucin12 peptide (d and e). The scale bar indicates represents 200 μm (b and d), and 100 μm (a, c and e), respectively.

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