A Critical View of Helminthic Therapy: Is It a Viable Form of Treatment for Immune Disorders Under the Category of Inflammatory Bowel Disease?

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A Critical View of Helminthic Therapy: Is it a viable form of treatment for immune disorders under the category of inflammatory bowel disease?

Abstract

In the developed world, Crohn’s disease and colitis affects the lives of many individuals. Recently, a new form of treatment for these autoimmune diseases has gained recognition. This treatment uses helminths (Trichuris trichiura and Trichuris suis) as an immunomodulant to a human immune system. Various studies (Summers et. al., 2004; Dige et. al., 2016; Lopes et. al., 2016) have shown the safety and viability of this form of treatment, but many believe this area of research leaves much to be desired. Through this paper, the topic of helminthic therapy and its viability as a form of treatment for autoimmune diseases under the category of inflammatory bowel disease (IBD) will be discussed. A conclusion will be reached by reviewing these three key studies: Helminth Regulation of Immunity: A Three-pronged Approach to Treat Colitis by Lopes et. al., Trichuris suis therapy in Crohn’s disease by Summers et. al., and Mucosal and systemic immune modulation by Trichuris trichiura in a self-infected individual by Dige et. al.

Part 1: Helminths

Understanding the following research includes a knowledge of certain key terms. The terms “helminth” or “helminthic therapy” are casting an incredibly broad net. Helminth is a vague term meaning “worm”. Pre-fixes are used to distinguish the different types of helminths; the main two being platy-helminth (flatworms) and nemat-helminth (roundworms) (Weinstock, 2012). Both species of helminths commonly used in helminthic therapy are nemat-helminth, and specifically whipworms. Trichuris trichiura (T. trichiura) is the human whipworm, and Trichuris suis (T. suis) is the pig whipworm (Cross, 1996). Focusing on the evolutionary paths and interactions with a human host for these two species of helminths will shed light on the viability of helminthic therapy for inflammatory bowel disease (IBD).

Parasites and parasitic roundworms have history that dates back to the beginning of man.
Written record of parasitic infection is seen in Egyptian medicine, and recently a new section of science has developed called paleoparasitology (Cox, 2002). The discovery of helminth eggs in fossilized feces spurred the development of paleoparasitology, and has furthered understanding of the history of parasitic helminths. Through this look at the history of human-parasite interaction, various theories have arisen. Many believe we have “heirloom” parasites, which were inherited from our primate ancestors (Cox, 2002). Along with these passed down parasites, humans have picked up “souvenir” parasites while we spread across the globe. With the rise in global trading throughout human history, many parasites have been passed from one population of humans to another (Hawash et. al., 2016). This trading resulted in a wide spread of human parasites (about 300 species). Although humans are the hosts for these many parasites, with the rise of technology, many parts of the world rarely experience infection. The hygiene hypothesis is a theory that looks at the effect of this loss of parasitism on humans, and is part of the basis for researching helminthic therapy (Robinson and Bradley, 2010).

Its complex life cycle consists of six distinct stages (CDC, 2013). Beginning with unembryonated eggs being released through the feces (usually human), they then develop into a two cell organism and advance to a cleavage stage outside of their human host (CDC, 2013). Following these stages, the eggs become embryonated, and are able to start infecting their human host (CDC, 2013). The eggs are then ingested (through contact with infected soil), and once inside the host, the eggs will hatch in the small intestine and release larvae (CDC,2013). Once mature, the adult whipworm will attach itself in the ascending colon and cecum, and the females will start to produce eggs (CDC, 2013). Adults are around four centimeters in length, and can shed anywhere from 3,000 to 20,000 eggs per day, thus starting the cycle again (CDC, 2013).
In order to complete its life cycle, *T. trichiura* must advert the host's immune system. All helminths secrete large amounts of antigenic material, which many believe overloads the human immune system and renders it useless (Wakelin, 1996). Since *T. trichiura* (like many other helminths) is large and motile, it is physically able to withstand attacks from its host (from both the immune system and the high acidity of the stomach). Another explanation as to why *T. trichiura* is able to survive the human immune system is through the secretion of lymphocyte suppressor factors, which will reduce immune responsiveness (Wakelin, 1996). It is this explanation that is a major part of the basis for research of helminthic therapy, and why an infection of helminths could combat the symptoms of IBD.
The helminth *T. suis* is also playing a part in helminthic therapy research. The life cycle of *T. suis* is almost identical to that of *T. trichiura*, except that it has co-evolved with the primary host as a pig (Pittman *et al.*, 2010). Because of this coevolution with pigs, *T. suis* is not as adept at avoiding a human immune attack as *T. trichiura*. However, like many other helminths, *T. suis* still secretes certain suppressants that lower immune responsiveness even in humans (Summers *et al.*, 2004). Since *T. suis* shares these immunomodulatory aspects with *T. trichiura*, but usually are unable to withstand a human environment for long, they make an optimal candidate for helminthic therapy research (Summers *et al.*, 2003).

**Part 2: The Immune System**

Another important part of this research is the role of the human immune system. Dissecting how the mucosal immune system in the digestive tract fends off non-self attackers, like *T. suis* or *T. trichiura*, and how is this process interacts with autoimmune disorders. The mucosal immune system is made of two distinct parts, the innate immune system and the adaptive immune system (NIH, 2003). For the purpose of this paper we will focus on the adaptive immune systems T-lymphocytes, also known as T-cells, due to the important part they play in the mucosal immune system (McGhee *et al.*, 1992).

The adaptive immune system is further divided into two sub-categories (humoral and cell-mediated). Within the humoral immune system, lymphocytes, specifically B-lymphocytes, are the main tracker cells. B-cells mature in the bone marrow, where they undergo rigorous trials to insure they are ready to produce the proper antibodies (making sure they will only produce antibodies that will tag non-self antigens) (Janeway *et al.*, 2001). Once matured they will travel to secondary lymphoid organs and await activation. Each B-cell is highly specific, meaning each
will produce a specific antibody for a specific antigen (Janeway et. al., 2001). This works for the checks and balances part of the humoral immune system, allowing little error for over-reaction to other, similar, antigens. When a B-cell is alerted to a specific antigen, it begins to proliferate. Soon it will have numerous copies of itself, some being memory B-cells and others being effector B-cell (Janeway et. al., 2001). The memory B-cell will survive for decades to allow your immune system to have a faster and stronger defense towards this specific antigen, preventing symptoms from developing during secondary exposure to the pathogen (Janeway et. al., 2001).

On the other side of this system are the effector B-cells. These B-cells will be the ones to poison the water supply, so to speak, becoming antibody factories and producing free-flowing antibodies that will mark antigens for phagocytosis, complement activation, neutralization or precipitation (Janeway et. al., 2001). These antibodies will allow macrophages, or other attacker cells to identify the pathogenic antigens.

T-cells fall into four categories, T-helper cells, T-regulatory cells, cytotoxic cells and memory cells. T-helper cells play an integral role in both cytotoxic (CD-8) T cells and activating B lymphocytes for antibody production. T lymphocytes gain immunocompetence in the thymus through a process of clonal selection (Janeway et. al., 2001). In the immune system, T-helper cells act as an activator for the immune system (calling the troops to take up their arms). In a similar fashion to a B-cell, T-helper cells are incredibly specific and usually each T-helper cell is tuned to one specific antigen. This is again to help the immune system prevent cross-reactivity with other (possibly self) antigens. When a T-helper cell is presented its specific antigen by an antigen presenting cell (i.e. a B-cell, or dendritic cell, etc.), the T-helper cell will begin to sound the alarm (Janeway et. al., 2001). This includes proliferation of this specific T-helper cell, and
some will become memory T-cells and others will become effector T-helper cells. Memory T-helper cells will stick around in the immune system (as was the case for memory B-cells), but the effector T-cells will rush into action. Effector T-cells will start to release cytokines, which signal other activated immune cells to ramp up production and get ready for battle (Janeway et al., 2001). Cytokines are essentially proteins or polypeptides that act as a signaling switch for other activated immune cells (i.e. effector B-cells). Earlier in this discussion, it was mentioned that when B-cells become activated they proliferate. This activation is made possible and monitored by effector T-helper cells (Janeway et al., 2001). This is another way the immune system is making sure it actually needs this large response to occur. With all of these ways that the immune system has to monitor itself, it seems there is little room for error or self-harm. However, in science there are usually exceptions to a rule, and for the immune system, autoimmune disorders are the exception.

Autoimmune diseases are still an elusive topic of research. Many health professionals and research scientists are not quite sure what causes certain autoimmune disorders, and have trouble coming up with cures or treatments for these diseases. When it comes to the umbrella term of inflammatory bowel disease (IBD), it is usually marked by a rise in the $T_{h1}$ and $T_{h2}$ immune response (Jackson et al., 2008). $T_{h1}$ and $T_{h2}$ are sub-classifications of the T-helper cells discussed earlier in this research, and both give rise to the signature symptom of IBD which is inflammation (Jackson et al., 2008). When a patient is diagnosed with an autoimmune disease, their immune system works in a slightly different way than most. Where a healthy immune system would only mark and attack non-self antigens, an immune system with autoimmune tendency marks and attacks self-cells. In a simplified explanation, $T_{h1}$ activate cytotoxic cells and
macrophages to attack self cells and Tₕ,2 cells activate B lymphocytes to produce antibodies against our own healthy cells (Jackson et. al., 2008). Normally, the body will release a Tₕ,1 immune response to an antigen for initial attack, and if it is able to kill and eliminate said antigen the body will return to equilibrium. If the antigen is not eliminated by the body’s Tₕ,1 response, then the body will move to a Tₕ,2 response. A Tₕ,2 immune response will lead to widespread antibody production and a much more powerful response. This cycle between Tₕ,1 and Tₕ,2 immune responses is what causes the cyclic symptoms of IBD (Jackson et. al., 2008). The Tₕ,1 will start to attack self-cells, causing inflammation and other painful symptoms. If the “antigen” is not killed the body cycles into a Tₕ,2 immune response and starts to produce self-marking antibodies, which only proliferate the attack on self-cells (Jackson et. al., 2008). It was this knowledge of the Tₕ,1 and Tₕ,2 immune response that lead researchers to believe that certain helminths may be helpful in combating IBD.

The way helminths, like T. suis and T. trichiura, avoid the immune system is by secreting suppressor factors as stated earlier. These proteins depress the body’s natural Tₕ,1 and Tₕ,2 immune responses, and therefore decrease the severity of these responses (Jackson et. al., 2008). Helminthic therapy research is based on the idea that a body with a heightened Tₕ,1 and Tₕ,2 immune responses could be regulated through the use of helminth infection, due to the fact that these lymphocyte suppressor factors down regulate the Tₕ,1/Tₕ,2 immune response (Jackson et. al., 2008).

Part 3A: Article Review

In the article Helminth Regulation of Immunity: A Three-pronged Approach to Treat Colitis, Lopes et. al. brought forward three distinct forms of helminthic therapy to treat patients
with ulcerative colitis. Lopes et. al. began their research with a few hypotheses that have driven helminthically therapy research. The “red queen hypothesis” states that as the helminth evolves better methods to evade its host’s immune system, said host evolves more efficient ways of ejecting the parasite from its system (Lopes et. al., 2016). Another hypothesis Lopes et. al. reference was the “hygiene hypothesis”, stating that a helminth could provide some type of protection against concomitant disease. These two hypotheses combined could explain why the helminth has been able to coevolve with the human species (Lopes et. al., 2016). As the two species (human and helminth) kept cycling through evolutionary tactics, the helminth presented protection against concomitant diseases. This could be seen as a health benefit to the host, and lead to a type of symbiotic lifestyle between host and parasite.

With these two hypotheses as a basis for research, Lopes et.al. presented these three forms of treatment for colitis: ingestion of viable ova or larvae of a helminth, use of crude helminthic extracts, and cellular immunotherapy. The most common type of helminthic therapy is through ingestion of viable ova or larvae, and through comparison of animal and human models it proves to be successful (Lopes et. al., 2016). Lopes et. al. discussed that in human models using T. suis ova or larvae there has been little to no side effects reported. Referencing studies that use both open label and placebo methods, Lopes et. al. provided adequate evidence of successful studies. However, Lopes et. al. state that there is a gap between animal (rodent) studies and human studies. This gap shows the need for further research of helminthic therapy using human patients.

Lopes et. al. also reviewed the use of helminth-derived extracts in treatment for colitis patients. Extracts could include things such as: soluble antigens from adult worms, egg antigens,
or simply adult worm extracts (Lopes et al., 2016). The use of these extracts negates the concerns of using live parasites for treatment of a patient with ulcerative colitis (Lopes et al., 2016). Furthermore, these extracts could be used as blueprints for the development of new anti-inflammatory drugs (Lopes et al., 2016). By reviewing a study featuring the use of Schistosoma mansoni (S. mansoni) egg extracts, Lopes et al. showcased the success these extracts can have. However, in this same study, not all results were positive as certain types of colitis were not affected by the egg extracts (Lopes et al., 2016). This lead the researchers to propose that helminthic therapy research should be conducted with defined patient cohorts. Having a defined patient cohort, in this sense, would mean the patients with colitis being studied would all have similar types of colitis and similar symptoms in response to the colitis (Lopes et al., 2016).

Thirdly, Lopes, et al. proposed the use of cellular immunotherapy as a treatment for patients with colitis. Traditionally, cellular immunotherapy is used by transplanting cells into the patient in order to combat the disease. Lopes et al. proposed transplanting cells from an individual infected with helminths to a patient diagnosed with colitis. The viability of this treatment has been shown through successful mice models, however there is a feasibility limitation when it comes to human patients (Lopes et al., 2016). Even though there is this feasibility issue, it is shown that a helminth infection (or the antigens helminths produce) induce certain cell phenotypes that could be used for treatment of IBD (Lopes et al., 2016).

Lopes et al. concluded that research on treatments for IBD is urgently needed, due to the terrible toll colitis can take on patient’s lives. Furthermore, through these three treatment methods Lopes et al. determined a trend towards a personalized approach in patient care and
treatment. Finally, Lopes et. al. concluded that there is definitely a link between host and parasite that may be beneficial to the treatment of IBD.

*Helminth Regulation of Immunity: A Three-pronged Approach to Treat Colitis* is a useful review to examine, because of its overview of three distinct treatment options. Many other studies focus their attention on the ingesting of live ova or larvae, and leave out the other options of cellular immunotherapy or helminthic extracts. Including these two other methods is inventive, and strengthens helminthic therapy research in various ways and broadens the field of research for helminthic therapy. Including results from various studies, and comparing those results between different treatment options gives a deeper insight into the gap in helminthic therapy research. Lopes, *et. al.*’s conclusions support helminthic therapy, especially in the sense of developing varying forms of treatment options. Some of the drawbacks of this article are based in the gap between human and animal trials. Throughout *Helminth Regulation of Immunity: A Three-pronged Approach to Treat Colitis*, Lopes, *et. al.* related the human trials to rodent trials that have had more conclusive results. Lopes, *et. al.* discussed that the less conclusive results in human trials are due to the lack in extensive research in this area. Although it seems likely that more research would close this gap in results, there may be other explanations as to why the human trials tend to be non-conclusive. One explanation could be the way trials are being conducted. Many of the studies using human trials do not have a defined patient cohort, which could be skewing their results. Lopes, *et. al.* mentioned the great necessity for defined patient cohorts in helminthic therapy research. Especially in the realm of cellular immunotherapy, the use of a more personalized care plan could drastically change the results of some of these studies. Because many autoimmune diseases vary from patient to patient, even
when they have the same diagnoses, it makes more sense to vary the treatment method to fit each individual.

Defining a patient cohort for a study of helminthic therapy would involve finding patients with similar responses to a specific autoimmune disease. For example, when creating a human trial studying colitis patients, the researcher would need to decide on the type of response they would be looking into. If they decide to look at an immune response marked by heightened $T_{h2}$ response, then they would need to test patient candidates for this response before including them in the study. This way, researchers can look at several different methods of helminthic therapy, and compare the results against one another to determine the best course of treatment. This extra setup step would allow researchers to tailor design a treatment method for patients with a specific immune response to colitis. Even though this would take more time, effort and money, it has the potential to change the lives of those living with IBD.

Part 3B: Article Review

*Trichuris suis therapy in Crohn’s disease* is a 24-week open-label study on the safety and efficacy of ingesting *T. suis* ova as a treatment for Crohn’s disease. Summers, *et. al.* (2004) conducted the study on patients with ages ranging from 18 to 72 years old, with a Crohn’s disease activity index (CDAI) from 220 to 450. The CDAI is a form of measuring symptoms of Crohn’s diseases, where 220 is moderately ill and 450 is severe disease (Summers *et. al.*, 2004). A score is calculated by taking in various symptoms (i.e. number of liquid stool per day, abdominal pain, number of antidiarrheal drugs used) summed over the course of seven days, after adjustment by a weighting factor, and relating that score to an average (Summers *et. al.*, 2004) (see table below).
<table>
<thead>
<tr>
<th>Clinical or laboratory variable</th>
<th>Weighting factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of liquid or soft stools each day for seven days</td>
<td>x 2</td>
</tr>
<tr>
<td><strong>Abdominal pain</strong> (graded from 0-3 on severity) each day for seven days</td>
<td>x 5</td>
</tr>
<tr>
<td>General well-being, subjectively assessed from 0 (well) to 4 (terrible) each day for seven days</td>
<td>x 7</td>
</tr>
<tr>
<td>Presence of complications^</td>
<td>x 20</td>
</tr>
<tr>
<td>Taking Lomotil or opiates for diarrhea</td>
<td>x 30</td>
</tr>
<tr>
<td>Presence of an abdominal mass (0 as none, 2 as questionable, 5 as definite)</td>
<td>x 10</td>
</tr>
<tr>
<td><strong>Hematocrit</strong> of &lt;0.47 in men and &lt;0.42 in women</td>
<td>x 6</td>
</tr>
<tr>
<td>Percentage deviation from standard weight</td>
<td>x 1</td>
</tr>
</tbody>
</table>

Figure 2: CDAI data table. (Summers et al., 2004)

Along with this age and CDAI range, the patients could be taking immunosuppressant drugs as long as it was kept constant. Patients were not allowed in the study if they filled any of the following criteria: they were taking immunomodulatory drugs; had previous treatments with antibiotic, antifungal, or antiparasitic medications; were diagnosed with diseases other than Crohn’s disease; had undergone an ileostomy, colostomy, resection of greater than 50 cm, or had obstructive symptoms; or were anticipating the need for surgery.

The method used to obtain *T. suis* ova included using pathogen free pigs, and ensuring the ova remained uncontaminated. First the pathogen free pigs were infected with *T. suis* ova, which were allowed to come to maturation. Once maturation was reached the eggs were collected from the colon, and cultured in vitro. The ova were embryonated in a phosphate buffered saline solution at 25°C, and kept bacteria free through the use of 0.2% K₂Cr₂O₇ solution.
and washed with sterile saline solution. Storing the ova at 5°C also ensured that there was no contamination by bacteria or other pathogens (Summers et al., 2004). Through this process of obtaining *T. suis* ova, Summers et al. showed that *T. suis* ova can stay viable, when stored properly, for up to nine months. This conclusion lends support to the potential for *T. suis* ova becoming a treatment which could be done by the patient at home with monthly checkups with a physician.

Through this 24-week open-label study, the following procedures were in use. The patients were given a dose of 2500 *T. suis* ova, suspended in liquid, every three weeks. At the time of entry and every six weeks, the following data points were collected: medical history, physical examination, pregnancy test, complete blood count, liver profile, stool examination checking for ova, pathogens, and *Clostridium difficile* toxin. Patients also kept diaries of their symptoms, and if a patient was taking immunosuppressant medication the dosage was kept constant. Summers *et al.* also used the two-tail Fisher’s exact test to examine characteristics that may indicate response or remission.

Results of this study support the viability and safety of using *T. suis* ova as a form of treatment for Crohn’s disease. With 75.9% of patients experiencing response (decrease of CDAI>100 or CDAI<150) and 65.5% of patients experiencing remission (CDAI<150) by week 12, this lends hard evidence to helminthic therapy being a viable form of treatment. Then by week 24, 79.3% of patients had experienced response and 72.4% of patients experienced remission. These results partnered with the mean CDAI decrease being 195.1 by week 12 and 187.2 by week 24, showing substantial evidence for the viability of helminthic therapy.

With these results, Summers, et. al. concluded that the use of *T. suis* ova was a safe and
effective treatment for Crohn’s disease. Since *T. suis* was not a natural parasite for humans, it lowered the risk of over-colonization of the patient (Summers *et. al.*, 2004). This paired with the ability to control the source of the ova (pathogen-free pigs), ensured the patient's safety throughout the course of treatment. With an 80% response rate and a 73% remission rate, even in patients with refractory disease, the outcome of this study clearly shows significant evidence towards the viability of helminthic therapy (Summers *et. al.*, 2004). However, with this study being an open-label study, it is impossible to rule out a placebo effect. This means that the results from this study justify further research (i.e. a blind clinical study), but cannot have truly conclusive results on its own.

The Summers *et. al.* study is important to look at, because of its unique use of the Crohn’s disease activity index. Many other studies focus on a change from a cellular level, and do not look at patient’s results holistically. By using the CDAI, Summers *et. al.* were able to show the change helminthic therapy was eliciting in their patient’s lives. Also, this holistic approach resulted in a greater change over time than other studies focusing on change in certain interleukins or $T_{h1}/T_{h2}$ response. While it is valuable to mark the clinical changes from the cellular level of a patient, it can be just as valuable to also look at the overall change in symptoms a patient is experiencing.

*Trichuris suis therapy in Crohn’s disease* did support the viability and safety of helminthic therapy, and had substantial evidence behind this conclusion. Summers, *et. al.* produced viable results that were even better than predicted. Even with the possible placebo effect, Summers *et. al.* created a new way of viewing helminthic therapy research. With studies like these it is easy to see the need for further research, particularly blind studies, but it also
showcases the real possibility for helminthic therapy to make a difference in the lives of those living with Crohn’s disease.

Part 3C: Article Review

*Mucosal and systemic immune modulation by Trichuris trichiura in a self-infected individual* is an open label study that consists of one self-infected patient (Dige et. al., 2016). The purpose of this study was to determine if the helminth *T. trichiura* was able to immunomodulate a human immune system. Through this study Dige et. al. could single out the immunomodulant aspect of helminthic infect, and rule out the possible interaction of an intestinal disease.

Because this study consisted of a single self-infected patient with no prior intestinal disease, the methods used were slightly different than many other helminthic therapy studies. The patient was a 38 year-old with untreated psoriasis, but with no prior medical history of intestinal disease. Being 1.92 m tall and maintaining a constant body weight of 71 kg, Dige et. al. determined the patient’s daily fecal production to be 250g. This data point is necessary to determine the amount of *T. trichiura* ova needed to produce a moderate infection. After factoring in the amount of eggs that would actually hatch in the intestine (80%) and the sex ratio (1:1), Dige et. al. decided that a dosage of 600 eggs would produce the infection needed. After nine weeks with no eggs in the patient’s feces, the patient voluntarily took an extra three small doses of *T. trichiura* eggs. Throughout this infection period, the following data points were collected before and after infection: blood and faecal samples (biweekly), biopsies and ileo-colonoscopies at four locations (the terminal ileum, the cecum/ascending colon, the transverse colon and the sigmoid colon), and monitoring of the patient for signs of diarrhea, anemia, and weight loss. At
the end of the study, the helminthic infection was terminated through the use of a mebendazole treatment.

The results Dige et. al. produced support the immunomodulatory effects of *T. trichiura* on a human immune system. This support was shown through a marked rise in T-cell cytokines that are directly linked to autoimmune diseases under the umbrella of IBD (i.e. T_{h1}/T_{h2}). With a rise in T-cell cytokines, the patient’s immune system lowered its T_{h1} response and raised its T_{h2} response. Dige et. al. conclude that this marked switch in T-cell response shows the ability of *T. trichiura* to be an immunomodulant for a human immune system. Another interesting result from this study was the patient developing *Campylobacter* colitis after the infection with *T. trichiura*. Dige et. al. concluded that an infection with helminths may leave the human immune system open to infection with other bacteria. Through this finding, Dige et. al. suggest that future studies ensure the patients have no prior infection of *Campylobacter*, and that the ova being ingested are free of contaminants.

*Mucosal and systemic immune modulation by Trichuris trichiura in a self-infected individual* is an article that brings an entirely new perspective to the study of helminthic therapy. Through the use of a patient with no prior intestinal disease, Dige et. al. were able to produce results that negate the coincidental changes in disease activity. Focusing on only the interaction between *T. trichiura* and the human immune system allowed Dige et. al. to produce results that support the immunomodulatory effects helminths can have on humans. Having conclusive evidence of this interaction could further helminthic therapy research, and supports the hygiene hypothesis. Through this study Dige et. al. also found that helminths may leave the human immune system vulnerable to co-infection. Even though this result seems negative for the use of
helminthic therapy, it was able to provide instruction for future studies. Through this study conducted by Dige et. al., helminthic therapy research was furthered by the knowledge of the immunomodulatory effects of T. trichiura has on a human immune system, and given valuable instruction to avoid co-infection of Campylobacter.

Part 4: Conclusion

In conclusion, treatment of autoimmune diseases categorized as inflammatory bowel disease (IBD) with infection of helminths (T. suis) is both safe and effective. Though more research is needed, evidence of the viability of helminthic therapy is clearly shown in the studies discussed in this paper. Researchers Lopes et. al. in Helminth Regulation of Immunity: A Three-pronged Approach to Treat Colitis support the use of helminths as a treatment for IBD in various forms. These forms include helminth extracts, cellular immunotherapy, and the most common method of ingesting live ova or larvae (Lopes et. al., 2016). Presenting evidence that each of these three forms have produced successful results (patients gaining remission status, or positively responding to treatment) in clinical trials, expands helminthic therapy research to include these innovative methods.

In Summers et. al.’s research, Trichuris suis therapy in Crohn’s disease, supporting evidence was shown through tracking patient progress using the Crohn’s disease activity index (CDAI). Using the CDAI to determine the viability of helminthic therapy was a unique way of examining results, and provided a holistic approach to Summers et. al.’s research. Their results were positive, with a majority of their patients gaining remission or at least responding positively to the treatment (CDAI decreasing by at least 100 points) (Summers et. al., 2004). Through this supporting evidence, Summers et. al. were able to provide a new way of examining results of
helminthic therapy studies.

Dige et. al., authors of *Mucosal and systemic immune modulation by Trichuris trichiura in a self-infected individual*, also presented concrete results supporting the immunomodulatory effects helminths have on a human immune system. By studying a self-infected individual with no prior intestinal disease, Dige et. al. were able to rule out the possibility of coincidental changes in disease activity in their results. Having results like these presents concrete evidence of the immunomodulatory influence helminths can have on a patient’s immune system (Dige et. al., 2016). This study solidifies the basis of helminthic therapy research, and warns future research of things to avoid.

After reviewing these three studies, helminthic therapy seems to be a safe and effective form of treatment. However, in science, nothing is ever truly proven and research must always be followed by more research. For future helminthic therapy research, the following points should be taken into consideration. When creating a clinical study, patients should be chosen from a defined cohort. A defined patient cohort would limit the influencing variables in a study, therefore the results would be more conclusive (Lopes et. al., 2016). Future studies should also use the CDAI to examine results, and pair this with determining results from a cellular level. This would allow studies to view their data comprehensively, thus solidifying their results. Finally, future research should ensure a lack of contamination when using live ova or larvae. This is important, because helminths can leave the human immune system vulnerable to bacterial infection.


K. Robinson and J. E. Bradley, The allergy epidemic: can helminths supply the antidote?, 2010 (40) 1586-1589.


