Do Hookworms Elicit Protective Immunity in Man?

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The two main species of human hookworm, Ancylostoma duodenale and Necator americanus (Table 1), are together believed to infect about 900 million people—mainly in tropical countries where adequate sanitary facilities may be lacking. But interactions between the two species, and their relative contributions to observed age-related infection patterns and seasonal cycles of transmission, continue to engender controversy. People tend to remain susceptible to infection throughout life, even with constant exposure to the infective stages. So what role does human immunity or resistance play in the epidemiology and control of infection?

In this article, Jerzy Behnke reviews the epidemiology of hookworm infection in the light of current understanding of mechanisms involved in host responses to infection and hookworm evasion of those responses. As he stresses, much further work is required.

It is often stated that A. duodenale is a northern species—the principal agent of human hookworm disease north of 20° latitude. In contrast, N. americanus is considered a southern species. However, this distinction is not so simple because foci of A. duodenale have been found in many parts of Africa. Eggs of the two species are not easily distinguishable, so estimates of the relative intensity of infection in mixed infections depend either on the recovery of adult worms from the faeces after chemotherapy or on culture of eggs to the infective stage, when the larvae can be individually identified. Both are labour intensive exercises and it is not surprising that many workers have not confirmed the identity of the parasites beyond their recognition as hookworms. This is a continuing problem in hookworm research, but nevertheless it is quite clear that there are important differences in the biology and host-parasite relationships of N. americanus and A. duodenale. These differences may account for some of the inconsistencies in epidemiological findings from different parts of the World, and there is an urgent need to define accurately the extent of infection with both organisms in future studies.

Longevity and Egg Output

Hookworms are considered to be long-lived parasites. Few longitudinal studies have been carried out but age-intensity profiles of affected communities reveal that children tend to acquire infection within the first 4 years of life (Fig. 1). This is followed by increasing intensity of infection monitored through faecal egg counts and expressed as the mean number of eggs per gram of faeces—EPG—which peaks or plateaus in the 20–30 age group. Thereafter EPGs tend to remain stable into late life or even rise further in old age. One study in Nigeria reported a decline in mean EPG in the 45+ age group to a level comparable to that of 1–4 year old children. The identity of the hookworm was not given but other studies in Nigeria report that N. americanus accounts for around 88% of the hookworm infections. No other survey to date has yielded quite comparable results, and it is more common for the age-intensity curve to stabilize in adult life—particularly in regions where only Necator occurs, such as Gambia and Puerto Rico. A study in Taiwan, where A. duodenale is dominant, found that the mean EPG fell in females aged 45+ but continued to rise in men among whom the heaviest infections were in 55–60 year age group. Similar observations have also been made in India.

Collectively these studies do not lend support for the existence of immunity to hookworms under field conditions. However, longitudinal studies across 1–2 transmission seasons report fluctuations in EPG.
which suggest that parasites are regularly lost and subsequently reacquired in the following transmission season. It is exceptional for hookworm larvae to be continuously available throughout the entire year, but even then the density of infective larvae in contaminated soil oscillates with the seasons. In Nigeria\textsuperscript{14} the density of hookworm larvae increased over 10-fold from very few in the dry months (January-February) to a maximum in July-October when rainfall was frequent. In Tanzania\textsuperscript{3} and West Bengal\textsuperscript{13} transmission seasons are sharply delimited by the hot dry climate between monsoons.

Early studies in India\textsuperscript{15} first suggested that faecal egg counts rose during the early part of the rainy season, peaked, and declined in the dry season (Fig. 2). This feature of hookworm epidemiology has been recently confirmed\textsuperscript{16} and attributed to population changes in \textit{A. duodenale} rather than \textit{N. americanus}. In this work\textsuperscript{16}, the population studied was about equally affected by both species, and the authors concluded that \textit{A. duodenale} was a relatively short-lived species whilst \textit{N. americanus} gave rise to more stable, long-lasting infections.

This conclusion is supported by several other studies which reveal that \textit{N. americanus} infections are more stable in the field under endemic conditions. Longitudinal surveys in Brazil\textsuperscript{17}, Gambia\textsuperscript{9} and East

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\includegraphics[width=\textwidth]{Fig. 1. Age-intensity curves for hookworm infection in four different regions (open points – females, closed points – males). a – Gambia\textsuperscript{9} (circles – Marakissa community; squares – Mandinari community), b – West Bengal, India\textsuperscript{15}, c – Taiwan\textsuperscript{8}, d – Nigeria\textsuperscript{11},}
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Hookworms are long-lived parasites. \textit{Necator americanus} can survive in the gut for up to 15 years, while \textit{Ancylostoma duodenale} lives for more than 12 months—probably up to 5 years.
Africa all suggest that where *N. americanus* is the dominant species, there is no obvious pattern of regular worm loss and reacquisition, despite short term fluctuations in EPG. The exception is amongst children still acquiring worms, where an upward trend is usually discernible in the transmission season. Further evidence for the stability of *N. americanus* infections comes from monitoring individuals who have been isolated by hospitalization or imprisonment or have been self-infected. The worms can survive for up to 15 years and loss is gradual over this period. Nevertheless there is still some disagreement. Nwosu and Anya found seasonal oscillations in EPG coinciding with rainfall, but their data were based on a population in which there was 88% prevalence with *N. americanus*. At the low point in the dry season, EPGs were still quite high (4000) and the fluctuation in egg counts above this baseline may therefore represent loss and gain of *A. duodenale*, which although in a minority in this community, is known to be considerably more fecund than *N. americanus*. In areas where *A. duodenale* is the dominant species, EPGs fluctuate in regular annual cycles in relation to rainfall and transmission season. The interpretation of annual cycles in EPG is not straightforward. It is widely assumed that egg counts reflect the number of parasites harboured by the host, but this is not a reliable indicator of parasite burden. Infections of only male worms would not be recognized through faecal egg counts, and larval burdens would also be missed. Furthermore there is evidence for a density-dependent reduction in female fecundity, and the possibility that female worms reduce or even cease egg output in the dry season when transmission is virtually abolished. The relationship between EPG and worm burdens (differential counts for both species) urgently requires examination, and a comparison of worm burdens...
during dry season and transmission season, following chemotherapy, would go a long way to providing relevant answers.

**Arrested development**

Another factor which may contribute to fluctuating egg counts is arrested development (hypobiosis) in *A. duodenale*. Individuals infected with the West Bengal strain took 22–38 weeks to develop mature worms – considerably longer than the normal pre-patent period\(^20\). These subjects were not exposed to further infection and it is conceivable that the parasites had lain dormant in the intervening period. Reinterpretation of Maplestone's earlier work\(^22\) reinforces the concept that in West Bengal, *A. duodenale* may have the capacity for arrested development. Egg counts on prisoners who commenced their sentences just after the monsoon season showed that EPGs fell steadily until February, but from March onwards increasing quantities of eggs were passed, peaking in May despite the absence of reinfection. Field observations provide further support\(^20\). The increase in egg production usually associated with the monsoon season actually began to be apparent in West Bengal in late April when ground conditions were still inappropriate for transmission. A temporary reduction in the fecundity of female worms could explain these observations, but it seems more likely that arrested larvae from the previous season began to mature in advance of the new transmission season, synchronizing their reproduction to environmental conditions most likely to ensure further infections. *A. duodenale* in the field may therefore have a relatively short survival time, adult worms declining in the dry season leaving newly acquired larvae to survive the months least suitable for transmission in an arrested state. In contrast, *N. americanus* continue to produce eggs steadily throughout the year and so, in regions where both species coexist, faecal cultures in the dry season yield a greater proportion of *N. americanus* larvae\(^16\).

**Worm Loss and Immunity**

Does the regular loss of adult hookworms imply immune mediated expulsion of worms and acquired immunity? Evidence for the induction of protective immune responses by hookworms in man is still scarce, although recent work supports the idea that some people are predisposed to reinfection following treatment with anthelminitics\(^23\)\(^,\)\(^24\). It may be that *A. duodenale* burdens decline each year through natural senility, but such a short life span is not compatible with Kendrick's study on experimentally infected prisoners in Madras penitentiary\(^25\). All the prisoners monitored for 12 months or more remained infected, and in some, egg output was maintained for over 5 years. *A. duodenale* therefore can live for longer than 12 months under conditions free from reinfection. In endemic regions the life span of adult worms is shorter – Nawalinski *et al.*\(^16\) calculated a 60% annual turnover for *A. duodenale* in West Bengal.

These observations imply that although a significant proportion of adult *A. duodenale* are lost in the post monsoon season, some would survive together with arrested larvae (and *N. americanus*) to continue the cycle into the next season. Perhaps worm loss is selective – acting on individual worms rather than on the entire worm burden. Possibly there are highly localized responses to individual parasites in the intestine, or alternatively age-related changes within the parasites may increase their susceptibility to host effector mechanisms.

It is likely that humans express a range of response levels to hookworm infection, as do sheep to infection with *Haemonchus concor- tortus*\(^30\), and inbred mouse strains to intestinal nematodes\(^26\)–\(^29\). Studies in Tanzania for example, where *A. duodenale* and *N. americanus* coexist in almost equal proportions, show that in some children, egg production was rapidly curtailed after the transmission season – while in others, more stable EPG patterns emerged with varying degrees of fluctuation. Individuals who do not pass hookworm eggs have been recognized in all studies, despite the generally high prevalence in affected communities. Moreover, there is good evidence from studies of chemotherapy followed by natural reinfection, that certain individuals are predisposed to heavier infections than others. But it is very difficult to distinguish between predisposition due to particular levels of exposure to infection, and predisposition through a genetically determined capacity for a particular level of immunological response.
Fig. 3. Reinfection with hookworms after treatment with anthelmintic. (a) India [all ages and both sexes]; (b) Puerto Rico [● – males, ○ – females]. Both figures show mean faecal egg counts before and after treatment. The arrow indicates the point at which anthelmintic was administered. (c) Ceylon, showing the number of worms estimated from faecal egg counts to have been harboured by tea plantation labourers and village residents in north and central Ceylon. Individuals were grouped according to the year in which they had previously been treated for hookworms. The first point shows the mean worm recovery from 3825 persons who had never before received treatment. The second point is based on 1728 persons re-examined within 15 days of treatment, and the remaining points refer to groups of 154-635 individuals who were previously given anthelmintic in the year indicated. The data for 1916 are based on only 29 persons. (A group of 18 individuals previously treated in 1923 had a mean worm burden of 101. This point is not shown.)

Son found that only 30% of the precontrol level was restored 630 days after chemotherapy – in agreement with earlier work. This slow reacquisition of adult hookworms contrasts with the relatively more rapid reinfection rates observed after elimination of Ascaris lumbricoides, where pretreatment intensity has been reported within 12 months of chemotherapy.

Two recent studies found significant correlation between worm burdens before, and many months after, treatment. Individuals heavily infected before treatment, subsequently reacquired above average worm burdens – supporting the idea that they are predisposed to heavy infections. But although it is tempting to suggest that predisposition may be linked to genetically determined susceptibility and resistance to infection, this hypothesis has not yet been specifically addressed in appropriately designed epidemiological studies.

Antibody Responses to Hookworms

Antibody responses to hookworms have been documented by a variety of assays using L3 and adult worm homogenates. Naturally infected patients and volunteers infected with N. americanus show elevated haemagglutinating antibody, complement fixing antibody, raised IgE and anti-parasite acetylcholinesterase responses. Yet they remain susceptible to infection. There is no evidence for resistance to human hookworms involving systemically circulating antibody.

Among this array of responses there may be host protective responses which are masked by non-protective responses to the numerous antigens in the parasite preparations. Local intestinal responses have generally been ignored but there is some evidence that IgA may figure prominently in the host-parasite relationship. Kumar et al. reported that IgA in the intestinal aspirates of hookworm patients was depressed, but normal levels were restored after chemotherapy. It may be that the presence of hookworms led to altered IgA secretion or reduced IgA half-life in the gut lumen, or the depressed IgA may reflect inactivation of this immunoglobulin by worm products. Hookworms are known to secrete proteases which are believed to aid in digesting the bolus of host tissues taken in during feeding. Some of these products may be directed at IgA, as occurs among other microorganisms living on mucosal surfaces.

Experiments with A. ceylanicum in dogs lend support for the importance of IgA in hookworm infection. A vigorous parasite-specific IgA response was detected in dogs which resisted a challenge infection administered 28 weeks after primary exposure to infective larvae. In contrast, no such response was evident in dogs vaccinated with adult worm antigen, even though they...
were equally resistant to challenge. Specific IgG levels however, were considerably higher in both groups and this may have reflected responses to the somatic migratory stages: the IgA response detected in reinjected dogs may have been the consequence of enhanced secondary responses in the intestine. Vaccinated animals which did not show an elevated IgA response to challenge would not have experienced local mucosal stimulation during vaccination and therefore would not be primed for an anamnestic response to challenge.

**Evasion of Immunity**

If IgA is involved in protective immunity to hookworms, it appears that the parasites have evolved a counter-strategy to inactivate this immunoglobulin class in the gut. Hookworms may also secrete immunomodulatory components to inactivate local immune responses. Other species causing chronic infections are known to do this, but there is yet no evidence that comparable strategies have been evolved by hookworms. Antigens presented orally can induce life-long tolerance of cell-mediated responses to dermal challenge and long-lasting tolerance of antibody responses, whilst initiating local immunity in the form of specific IgA. Hookworms reside in the gut lumen but their antigenic presentation is not directly akin to the oral route. Hookworm mouthparts penetrate deep into the gut mucosa and their excretory products probably enter the circulation – bypassing the normal mucosal-Peyer's patch route of antigen entry from the intestine. The possibility that hookworms interfere with the immunoregulatory circuits in the intestine has not yet been examined experimentally but the hamster models use of hamster-adapted strains of *N. americanus* and *A. ceylanicum* now offers the possibility of exploring these aspects of the hookworm-host-parasite relationship.

Observations on the feeding activity of the dog hookworm, *A. caninum*, have shown that parasite attachment to the gut mucosa is temporary, lasting for 4–6 hours at most. The lesions created during feeding are rapidly repaired by the host but areas of infiltration by inflammatory cells have been observed around parasites embedded in host mucosa. The parasites regularly change their feeding sites and may thus avoid potential damage which could be inflicted by inflammatory cells. It may be that the ability of worms to relocate is diminished with age so that, for example, older *A. duodenale* may be more readily caught in lesions around feeding sites than younger worms or *N. americanus*.

There is some evidence that hookworms have evolved strategies for evading antibody responses. Adult *A. caninum* shed immunoglobulin from their cuticle, whilst dead parasites bind antibody which can be detected using immunofluorescent techniques. Studies with *N. americanus* surface-labelled with 125I suggest that they also shed surface antigens. This may be an evasive mechanism preventing antibody from adhering to the parasite, although it is difficult to envisage how antibody bound to cuticle could affect hookworm viability.

A specific and continuous intestinal antibody response may explain epidemiological and experimental observations for the slow loss of hookworms. Such a response may cause the gradual deterioration of older worms but not affect young parasites, both of which are recruited from the environment which still have potent evasive mechanisms. It would also explain why *A. duodenale* can be lost in the post-monsoon season leaving *N. americanus* unscathed.

**Strategies for Vaccination**

Whatever the nature of immune involvement in hookworm infection, it is clear that the parasites have the upper hand in most people living in endemic areas. A vaccine against hookworms must therefore be considered improvement on natural human responses to these parasites. However, it is quite possible that antigens will be identified which are not normally made available to the host during natural infection. Two such possibilities are apparent – high affinity antibodies against key components of the various nematode sensory receptors may block sensory input and so inhibit parasite attachments, and, because hookworms pass large quantities of serum through their intestines, a serum antibody response against the parasite intestine may cause sufficient damage for the worms to be lost. During natural infection, such antigens may never be exposed or may present insufficient antigenic challenge to the host, but their isolation may be feasible using fresh parasites grown in hamsters – enabling vaccine production through gene cloning and antigen synthesis by recombinant DNA methods.

There is considerable scope for future
research on hookworms, using recently developed animal models and by epidemiological projects linking immunological assays with resistance and worm loss in the field. Understanding the mechanisms involved in worm loss will be an important step in potential vaccine production, and I hope that a concerted effort to apply the tools of modern experimental medicine will help to elucidate aspects of hookworm biology which have eluded experimenters in the past.

But even in areas of high malaria transmission, only a small proportion of malaria infections reach a sufficiently high parasitaemia to cause an acute illness (Fig. 1). The remainder produce a low-level, asymptomatic parasitaemia which may last for weeks or months. Are these asymptomatic infections of any clinical significance? In non-immune patients, most P. falciparum infections result in high parasitaemias and severe symptoms unless treated early. Fortunately, however, in areas where malaria is endemic and where most of the population have some immunity, this is not the case and severe, clinical infections are relatively uncommon.

What proportion of new malaria infections causes symptoms in semi-immunes is uncertain. In The Gambia, we have found that children who live in a rural area experience about 1 clinical attack of malaria per year and a similar incidence has been noted in Liberia. In our survey, children were visited only once a week so that mild attacks may have gone unrecorded between visits. But even in areas of high malaria transmission, it is unlikely that children experience more than 2 clinical attacks of malaria per year. In our study area in The Gambia, children are exposed to about 50 bites from infected mosquitoes per year (S. Lindsay et al. unpublished) but in other parts of Africa, where transmission is more intense, inoculation rates 2 or 3 times higher have been recorded. Not all sporozoite inoculations lead to infection but these entomological observations suggest that in endemic areas, only a small proportion of malaria infections reach a sufficiently high parasitaemia to cause an acute illness (Fig. 1). The remainder produce a low-level, asymptomatic parasitaemia which may last for weeks or months. Are these asymptomatic infections of any clinical significance?