OPEN LETTER

Risk assessment for the implementation of controlled human Schistosoma mansoni infection trials in Uganda [version 1; peer review: 1 approved, 1 approved with reservations]

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Abstract

Schistosomiasis is a parasitic infection highly prevalent in sub-Saharan Africa, and a significant cause of morbidity; it is a priority for vaccine development. A controlled human infection model for Schistosoma mansoni (CHI-S) with potential to accelerate vaccine development has been developed among naïve volunteers in the Netherlands. Because responses both to infections and candidate vaccines are likely to differ between endemic and non-endemic settings, we propose to establish a CHI-S in Uganda where Schistosoma mansoni is endemic. As part of a “road-map” to this goal, we have undertaken a risk assessment. We identified risks related to importing of laboratory vector snails and schistosome strains from the Netherlands to Uganda; exposure to natural infection in endemic settings concurrently with CHI-S studies, and unfamiliarity of the community with the nature, risks and rationale for CHI. Mitigating strategies are proposed. With careful implementation of the latter, we believe that CHI-S can be implemented safely in Uganda. Our reflections are presented here to promote feedback and discussion.
Keywords
Schistosoma mansoni, Controlled Human Infection Studies, Uganda, risk assessment

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Competing interests: The authors have declared no personal financial competing interests. However, we are working collaboratively to develop the CHI-S for implementation in Uganda, and therefore have research goals with potential to influence our approach to this risk assessment. This is, in part, our motivation for publishing it on an open peer review platform.

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Background

Schistosomiasis is a parasitic infection affecting approximately 230 million people worldwide\(^1\). Infection is caused by trematodes (flukes) of the genus *Schistosoma*. Because the infection is responsible for considerable morbidity worldwide, particularly in Africa, schistosomiasis was recently listed among the top 10 infections for which a vaccine should urgently be developed\(^2\).

Controlled human infection (CHI) studies are an important tool for vaccine development. They provide a platform to safely and swiftly test vaccine candidates for the pathogen in question. Furthermore, they can contribute to understanding host-pathogen interactions and help to unravel the nature of protective immunity. They have been used successfully for a substantial number of infectious diseases, including malaria, dengue, and influenza\(^3\). A CHI model has now been developed for schistosomiasis at Leiden University Medical Center, where Dutch volunteers with no previous exposure to schistosomiasis participated\(^4\). However, the response to schistosome infection, and to candidate vaccines, is likely to be different in endemic countries. In such settings multiple differences in environmental exposures, as well as prior exposure to schistosomes, drive differences in both the innate and adaptive immune responses which determine infection susceptibility and vaccine responses\(^4,5\).

We are therefore working towards the establishment of a controlled human infection model for schistosomiasis in Uganda, where *Schistosoma mansoni* is highly endemic. Almost 30% of the population is estimated to be infected\(^6\), with half the population at risk\(^7\). As a first step we held a stakeholders’ meeting in Uganda in November 2017, and we published the meeting report and resultant road-map for the implementation process\(^8\). A key element of the road-map was to undertake a risk assessment. This document therefore aims to provide an assessment of risks that may arise before, during and after start of a controlled human infection model with *Schistosoma mansoni* (CHI-S) in Uganda.

Male and female schistosomes live in the mesenteric or perivesical veins of their human host, where they mate and produce eggs. These eggs are either released into the environment through faeces and urine or stay within the host tissue where they induce inflammation. When the excreted eggs reach fresh water, they hatch and release miracidia that can then infect a suitable snail host. Infected snails are able to shed larvae, called cercariae, which infect humans. The Leiden University Medical Center (LUMC) CHI-S exposed healthy naïve volunteers to increasing doses of male cercariae to study the tolerability of such a controlled human infection model. This male-only model avoids the risk of pathology caused by schistosome eggs. To generate the infectious cercariae for a male-only CHI-S, individual laboratory-reared freshwater snails are infected, each with a single miracidium. Clonal replication follows, such that thousands of single-sex cercariae are subsequently shed by the snail. The sex of the cercariae can be determined by PCR, and the appropriate number of cercariae can be prepared for dermal infection. Because snails shed thousands of cercariae over a period of weeks, every time they are exposed to light, it is possible to first perform quality control (QC) testing on every batch (e.g., to assess the viability, sex and bioburden of the cercariae). Following principles set forward in good manufacturing practices (GMP) guidelines, the cercariae and their excipients are produced and tested for consistent quality according to predefined criteria. Only when compliant, is the cercariae batch released for clinical use. To this date, 17 people have been exposed to *S. mansoni* cercariae during CHI-S studies in Leiden.

In terms of the technical aspects of shipping infectious material to Uganda, culturing the infectious material in Uganda and preparing the infectious cercariae, we have considered three options.

**Option 1: Shipping of parasites and snails from the Netherlands to Uganda.** In this scenario, *S. mansoni* parasites and snails would be shipped from Leiden (The Netherlands) for preparation of the cercariae for human infection in Uganda. From a technical perspective, the easiest approach to rapid implementation of CHI-S in Uganda would be to produce and release the infectious snails in Leiden and subsequently ship them to Uganda. In Uganda, a further snail shedding would be used to generate the infectious cercariae. Alternatively, *S. mansoni* parasites (for example in the form of *S. mansoni* eggs contained in a rodent liver) could be shipped separately from uninfected snails, which would mitigate shipment risks.

The CHI-S model in Leiden uses a schistosome strain which has been genotyped and has been mapped to be of Puerto Rican origin\(^8\). Because this strain has been laboratory adapted and kept in the Leiden facility since 1955, it has the advantage of its known virulence in animals, experience of its effects in the Dutch human volunteers, and its sensitivity to praziquantel. As well, the Leiden model uses *Biomphalaria glabrata* snails which are not indigenous to Uganda (Appendix 1 [Extended data\(^9\)]). Therefore, the ecological risks of accidental release of schistosomes or snails into the environment have to be considered.

**Option 2: Shipping of parasites from The Netherlands followed by use of local Ugandan snails.** This scenario would involve transporting only *S. mansoni* parasites (Puerto-Rican strain), then using local snail species such as *B. choanomphala* (from Lake Victoria) or *B. stanleyi* (from Lake Albert) to produce cercariae in Uganda\(^10\). Advantages, as in option 1, would be the fact that the parasite strain has been characterized in both animals and humans, which decreases its potential risk for the volunteers. Disadvantages would be possible technical hurdles to be overcome to establish a local snail colony and achieve successful infection with release of infectious *S. mansoni* cercariae. However, expertise in these processes already exists in Uganda\(^10\), subject to laboratory renovations and staff training to ensure compliance with GMP principles. This option would also be relatively simple to implement.

**Option 3: Using local Ugandan parasites and local Ugandan snails.** In this scenario the full *S. mansoni* laboratory life
cycle would be established in Uganda, using a local snail species and starting with a new S. mansoni strain, and a rodent mammalian host. Although the risk of clinically unexpected, unwanted side effects, or of relative resistance to praziquantel treatment, might be higher when using the local strain of S. mansoni, the ecological risk would be lowest.

All options require preparation of the cercariae for human infection under strict Quality Assurance and controlled conditions in Uganda with adherence to Good Manufacturing Guidelines. In Leiden, procedures were developed based on GMP principles contained in the European Commission directive 2003/94/EC, with the infectious cercariae considered as an ‘auxiliary medicinal product’. Details of the procedures have been published1. These include production in a biosafety level 3 facility, governed by stringent standard operating procedures including for quality control, logging and monitoring; production and counting of infectious cercariae by two independent technologists; and antibiotic treatment and microbiological bioburden testing to ensure that the cercarial product is free of pathogens with potential to harm CHI volunteers. Equivalent procedures and quality control will be needed in Uganda in order to implement CHI-S.

In this document we address risks associated with CHI-S in Uganda on three different levels: i) the introduction of new species (the transport of snails, the snail culture facilities, the potential for ecological harm as a result of importing snails), ii) the introduction of a new schistosome strain into Uganda, and iii) clinical trial risks common to all options (natural infection during the trial period, and the risks to volunteers resulting from the controlled infection).

**Risk assessment methods**

We identified risks and potential approaches to mitigation based on relevant literature, experience from the Leiden CHI-S model, stakeholder discussions, and discussion with experts. The level of risk and effectiveness of proposed controls was determined by consensus between the authors. The inherent risk was defined as the risk before putting controls in place, calculated by consensus between the authors. The level of risk and effectiveness of proposed controls was determined by consensus between the authors. The residual risk was similarly calculated, based on likelihood and impact scores after controls have been put in place. Mitigating controls could reduce the residual risk score by reducing the likelihood of an event occurring, or by reducing the impact if it should occur. Likelihood was scored as almost certain/common, 5; likely, 4; possible, 3; unlikely, 2; rare, 1. Impact was scored as critical, 5; major, 4; moderate, 3; minor, 2; insignificant 1. Resulting risk scores of 18–25 were considered high, and unacceptable. Resulting risk scores in the range of 9–17 were considered moderate, with further controls desirable if possible, and caution required if implemented at this risk level. Resulting scores of 0–8 were considered low, and usually acceptable.

**Option 1: Shipping of parasites and snails from the Netherlands to Uganda**

According to our first idea, infected snails would be shipped. The WHO report ‘Guidance on regulations for the Transport of Infectious Substances 2017–2018’11 provides information on how to adequately transport infectious substances. In accordance with these guidelines, shipment of S. mansoni infected snails falls under ‘CATEGORY B, INFECTIOUS SUBSTANCES’ (UN3373). Shipment of live snails is a time-sensitive undertaking and therefore can only be facilitated by air shipment. Infectious substances cannot be carried on as hand-luggage. Transport of infectious substances are subjected to International Air Transport Association (IATA) requirements. Packaging of Category B substances need to comply with rules set out in the P650 packaging instruction11. This involves triple packaging and proper marking and documentation. Upon arrival in Uganda, it would be crucial for the package to clear customs as quickly as possible so that snails arrive in good condition. In order to achieve this, the customs office should be notified about the arrival of the shipment. In collaboration with the customs officer, all required documentation should be prepared in advance and approval for import of the products should be sought.

Alternatively, snails and *Schistosoma* parasites would be shipped separately. Uninfected snails can be shipped more easily because this shipment does not have to comply with the regulations for the transport of infectious substances. Similar to the previous option, shipment should clear customs as soon as possible. These snails could be kept to reproduce in the Ugandan laboratory to sustain their life cycle.

A second shipment would contain *Schistosoma* parasites. There are two ways in which this material can be transported (still under the ‘CATEGORY B, INFECTIOUS SUBSTANCES’ (UN3373)):

1) Within a living host such as a *Schistosoma*-infected hamster. These animals can shed *Schistosoma* eggs that can be used to infect the snails.

2) Within a preserved liver sample kept on medium from a *Schistosoma*-infected hamster. This liver sample contains *Schistosoma* eggs. Upon arrival in Uganda, further processing of the sample provides miracidia which can be used to infect the snails. Test shipments should be scheduled to determine the feasibility of such transports and the conditions in which the liver sample should be shipped. From previous experiments in Leiden, the preserved liver sample can be used to infect snails for up to one week after being harvested.

Risks associated with shipping of parasites and snails from the Netherlands to Uganda, and mitigating strategies, are summarized in Table 1.

**Option 1: Snail culture facilities; potential ecological harm**

To house the *Biomphalaria glabrata* snails in Uganda, they would need to be kept in strict quarantine. *B. glabrata* are not a naturally occurring snail host in Uganda, and should therefore not spread to the environment. In order to house snails, an incubator, or room temperature, set and monitored at 28°C is needed. The incubator (if used) door should be fully closed when the laboratory is not in use. Precautionary measures to contain the snails to the facility should be taken and include physical
Table 1. Risks associated with shipping of Schistosoma mansoni parasites and Biomphalaria glabrata snails.

<table>
<thead>
<tr>
<th>Risk</th>
<th>Inherent risk score</th>
<th>Controls</th>
<th>Residual risk score</th>
<th>Total risk post control</th>
</tr>
</thead>
<tbody>
<tr>
<td>Death of snails in transport</td>
<td>Likely</td>
<td>Pilot transport with low numbers of snails to optimize transport conditions</td>
<td>Possible</td>
<td>Critical</td>
</tr>
<tr>
<td>Delays in customs clearance</td>
<td>Likely</td>
<td>Contacting customs officials to discuss required documentations and preparing documents prior to shipment</td>
<td>Possible</td>
<td>Major</td>
</tr>
<tr>
<td>Spill of infectious materials and non-indigenous snail species</td>
<td>Possible</td>
<td>Proper packaging</td>
<td>Unlikely</td>
<td>Moderate</td>
</tr>
<tr>
<td>Establishment of a B. glabrata colony outside laboratory facility</td>
<td>Possible</td>
<td>Proper packaging</td>
<td>Rare</td>
<td>Critical</td>
</tr>
</tbody>
</table>

Risks associated with culture of B. glabrata in Uganda, and mitigating strategies, are summarised in Table 2.

Option 2: transport of S. mansoni infectious material and use of local snail species for cercarial production

This approach only requires transport of S. mansoni infectious material. This would use the second transport approach described in option 1, within a preserved liver sample from a schistosomiasis-infected hamster. The same regulatory guidelines for transporting infectious material apply. A major advantage of this approach is that the potential ecological and genetic risks related to introduction of a non-endemic snail species can be avoided.

Option 3: re-establishing the full S. mansoni laboratory life cycle in Uganda, using a local snail species and S. mansoni strain

The alternative to shipping infectious material and snails from The Netherlands is to re-establish the full laboratory life cycle of S. mansoni using Ugandan snail species and Ugandan isolates of S. mansoni. The life-cycle has been maintained in the past at the Vector Control Division of the Ministry of Health, but is not currently available. The advantages of using a Ugandan life cycle include reducing the environmental risk associated with non-endemic snail species and schistosome strains. In addition, this model would be most representative of the field infections in Uganda. There are however several challenges with using Ugandan snails and isolates. With regard to Ugandan snail species, there is variability between snail species in susceptibility to S. mansoni infection; however, there is experience of conducting infection of local species at the Vector Control Division, so this is expected to be feasible. With regard to the new schistosome laboratory strain, the characteristics of this would be unknown in terms of virulence and susceptibility to praziquantel treatment. Determining these characteristics would not be simple, since validated tests for schistosome

barriers, such as rooms with closed doors and windows. The snail culture basins and water drainage system should be covered with fine mesh to prevent escape (appendix 1 [Extended data]). In addition, access to the laboratory should be restricted to the research team. The incubator (if used) should preferably be positioned away from the door. Additional security measures could be a double door to create a sluice. Appendix 2 (Extended data) lists precautionary measures that should be taken when working with schistosomes. Standard operating procedures (SOPs) will be exchanged with LUMC and reviewed to fit the Ugandan facility. These SOPs deal with culture processes as well as the disposal of infectious material.

In case a single snail would accidentally be released into the environment, it is capable of reproducing in the absence of an opposite-sex snail using self-insemination\(^2\). This ability poses an ecological hazard where a single snail could develop into a colony. In addition, snails can be transported over large distances attached to birds and can survive dry conditions for up to two months. This snail itself is not endemic in Uganda, although previously this species has been held at the Vector Control Division of the Ministry of Health for a different project. The consequences of accidental introduction of this new species are difficult to predict, however it may result in the following (Appendix 1 [Extended data]):

1) Interspecific hybridization between B. glabrata and local Biomphalaria species
2) Uncontrolled spread due to lack of natural enemies, competitors or pathogens
3) Altered S. mansoni dynamics, because of potentially higher susceptibility of B. glabrata for S. mansoni infection

Spread to the environment of B. glabrata may go unnoticed, because of its similar morphology to endemic snail species.
resistance are currently not available. In addition, the new isolate would not be clonal and variability within the newly collected schistosome population might result in variable responses in the host, and to drug treatment. In addition, dose-finding studies would start from scratch to find the balance between tolerability and attack rate.

Options 1, 2 and 3 all require the establishment of facilities in Uganda for production of the infectious cercariae under GMP principles, in order to ensure high quality, reproducible infectious doses. Option 3 requires also the establishment of suitable, specific pathogen free animal facilities to house the rodents (hamsters or mice) that will provide the mammalian hosts in the laboratory life cycle. Risks associated with these elements are also considered here (Table 3).

### Natural infection during trial period

The single-sex *S. mansoni* challenge has been designed to prevent the occurrence of egg-associated morbidity. In the current model, volunteers participating in the trial will be infected using only male cercariae which penetrate the skin and result in potent infection. In future, female cercariae may also be used to infect volunteers. The sex of the male cercariae can be determined using a specifically designed multiplex real-time PCR which has been described elsewhere. Once infected, individuals should avoid any exposure to contaminated water. If a subject were to be naturally infected over the course of the study, this might lead to mixed, male and female, infections, with mating of the schistosomes resulting in egg production that causes morbidity. If the Puerto Rican strain used in Leiden is imported for use in Uganda, mating and excretion of eggs into the environment could alter the genetic make-up of Ugandan schistosome populations, with unknown consequences. However, given that the Puerto-Rican strain is relatively inbred after prolonged passage in the laboratory, and was shown to be praziquantel-sensitive in the CHI-S, hybridisation with Ugandan schistosome populations is unlikely to result in increased praziquantel resistance or virulence.

The chance of natural infection can be limited by choosing a study population which does not come into contact with freshwater. However, this would over-restrict recruitment from the true target population, which is people at risk of *S. mansoni* infection. Options to minimise this risk among volunteers from the preferred target population include the following:

1. **The feasibility of avoiding fresh water may be surveyed using questionnaires in a pilot study at the field site and the information used to select volunteers least at risk of re-exposure, and to make provisions to support volunteers to avoid re-exposure.**

2. **While selecting subjects, the investigator may ask whether the subject is likely to spend time in, or to travel to, areas where the risk of contracting a natural infection is high. If so, once again it should be stressed that contact with fresh water should be avoided; volunteers unlikely to achieve this would be excluded.**

3. **Apart from providing information to the volunteer and raising awareness of this issue, frequent testing for eggs in stool and urine samples may be performed by microscopy (and PCR). Eggs can be found 5–7 weeks after mixed male and female infection.** *S. mansoni* eggs in stool would indicate a concomitant natural infection, which would necessitate immediate treatment of the volunteer with praziquantel. However, stool microscopy and PCR is likely to be unreliable given variable egg excretion and the low sensitivity of stool examination for eggs.

4. **In those trials in which natural infection may be a considerable risk, testing using plasma circulating anodic antigen (CAA) may be conducted weekly from the outset of the trial. Both natural and experimental infections may then be terminated as soon as patent infection has been detected (e.g. at ~7 weeks post controlled human infection, when CAA levels > 1pg/mL). Early abrogation of the infection will prevent mating and egg

| Table 2. Risks associated with snail culture facilities. |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|
| **Risk**                        | **Inherent risk score** | **Total inherent risk** | **Controls**                        | **Residual risk score** | **Total risk post control** |
| **Likelihood**                  | **Impact**       |                  |                                | **Likelihood** | **Impact** |
| **Spread of Biomphalaria**      |                  |                  |                                | **Rare**       | **Critical** |
| **glabrata snail to environment** |                  |                  | 1) Precautionary measures for snail housing facility including physical barriers and restricted access |                  |                  |
|                                 |                  |                  | 2) Use of SOPs regarding disposal of infectious material and non-indigenous snail species |                  |                  |
| **Establishment of a B. glabrata** |                  |                  | 1) Development of containment strategies |                  |                  |
| **colony outside laboratory**   |                  |                  |                                | **Rare**       | **Critical** |
| **facility**                    |                  |                  |                                |                  |                  |
laying. There would be modest drawbacks to the resulting data, because it would not be possible to study the dynamics of antigen excretion over time and quantitation of infection would be less accurate.

5) Alternatively, volunteers may be displaced to a non-endemic region for the study duration. However, the prolonged, seven to 12-week “admission” required for the CHI-S would be a major burden and inconvenience, as opposed to the relatively short-duration (24 days) for malaria CHI studies where such approach has been employed.11 The possibility of volunteers absconding during the study, given the long duration, might be significant, abrogating the value of such an approach. Additionally, this would have cost implications, in terms of providing suitable accommodation and compensation for loss of income.

Risks associated with natural infection during the CHI-S, and mitigating strategies, are summarised in Table 4.

**Risks to volunteers resulting from the controlled human infection**

Controlled infection with *S. mansoni* has been successfully performed in 17 Dutch volunteers. Although the single sex infection does not cause egg-related morbidity in volunteers, it may cause symptoms in response to the infection. These include dermatitis due to the percutaneous penetration of the cercariae and an acute schistosomiasis as a consequence of a systemic hypersensitivity response. Severe acute schistosomiasis syndrome (Katayama fever) may present with symptoms such as fever, fatigue, myalgia, malaise, non-productive cough, eosinophilia and patchy infiltrates on chest radiography. In Leiden, several volunteers reported with systemic symptoms which seemed to be an acute schistosomiasis syndrome, with one volunteer presenting with prolonged symptoms of Katayama fever. In addition, one volunteer presented with peri-orbital oedema which lasted one day, and may have been related to the infection. Such symptoms can be treated symptomatically and all recovered. Both these volunteers had received the highest dose of cercariae (30 cercariae) used in Leiden. The risk of severe symptoms can be minimised by dose escalation in modest increments. The impact can be reduced by careful monitoring, provision of symptomatic relief and abrogation of infection by treatment if necessary. Frequent follow up visits need to be scheduled throughout the trial to discuss adverse events and conduct clinical assessments of the study volunteers. Safety laboratory tests need to be routinely performed. Volunteers can also experience side effects related to the praziquantel treatment. Common side effects include nausea, dizziness, and fatigue. Volunteers can be reassured that these symptoms are well recognised and transient. Their severity can be reduced by taking praziquantel after food. Symptomatic relief can be provided when required.

The 2017 stakeholders’ meeting identified community engagement to ensure proper understanding of the CHI-S as an essential basis for ethical conduct of a CHI study. CHI is a novel concept in Uganda, where CHI have not been undertaken in the past and understanding of medical research, in general, is at a low level. The idea of a “medical” procedure being undertaken should be explained to volunteers, who can identify, and help to address, misinformation; effective education of volunteers to a full understanding of the expected effects of the CHI (and reasons for undertaking it) will all be essential to the smooth and safe running of these projects.

Risks related to volunteers and communities during the CHI-S, and mitigating strategies, are summarised in Table 5.

### Table 3. Risks associated with re-establishing Uganda *Schistosoma mansoni* life cycle.

<table>
<thead>
<tr>
<th>Risk</th>
<th>Inherent risk score (Likelihood</th>
<th>Total inherent risk</th>
<th>Controls</th>
<th>Residual risk score (Likelihood</th>
<th>Total risk post control</th>
</tr>
</thead>
<tbody>
<tr>
<td>New isolates of <em>S. mansoni</em> from the Ugandan population might exhibit variable praziquantel susceptibility, or praziquantel resistance</td>
<td>Possible Critical 15</td>
<td>1) Test new isolates for praziquantel susceptibility <em>in vitro</em> and in an animal model before use in CHI</td>
<td>Unlikely Critical 10</td>
<td></td>
<td></td>
</tr>
<tr>
<td>New isolates of <em>S. mansoni</em> from the Ugandan population might exhibit unexpected virulence</td>
<td>Possible Critical 15</td>
<td>1) Test new isolates for relative virulence in an animal model before use in CHI</td>
<td>Unlikely Critical 10</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Production processes based on GMP principles for single-sex infectious cercariae not established in Uganda</td>
<td>Possible Critical 15</td>
<td>1) Development of appropriate animal and snail facilities 2) Training of Ugandan staff 3) Monitoring and review by experienced LUMC collaborators 4) Monitoring and review by Ugandan regulators</td>
<td>Rare Critical 5</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Table 4. Risks associated with natural infection during trial period.

<table>
<thead>
<tr>
<th>Risk</th>
<th>Inherent risk score</th>
<th>Total inherent risk</th>
<th>Controls</th>
<th>Residual risk score</th>
<th>Total risk post control</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mixed sex infection in trial volunteers</td>
<td>Likely</td>
<td>Moderate</td>
<td>1) Avoidance of fresh water bodies during trial period</td>
<td>Rare</td>
<td>Moderate</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>2) Pilot survey to establish feasibility of fresh water avoidance</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>3) Selection of trial volunteers with low risk of contracting natural infection</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>4) Abrogation of infection as soon as the trial endpoint has been reached (e.g. CAA &gt; 1 pg/mL)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>5) Displacement of volunteers to non-endemic setting with excellent water and sanitation facilities</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mixed sex infection in trial volunteers</td>
<td>Likely</td>
<td>Moderate</td>
<td>1) Full clearance of infections before trial starts</td>
<td>Rare</td>
<td>Moderate</td>
</tr>
<tr>
<td>leading to release of Puerto Rican strain into environment</td>
<td></td>
<td></td>
<td>2) Continuous screening for egg production</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>3) Abrogation of infection as soon as the trial endpoint has been reached (e.g. CAA &gt; 1 pg/mL)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>4) Displacement of volunteers to non-endemic setting with excellent water and sanitation facilities</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* The impact of natural co-infection on morbidity is classed as moderate (rather than major or critical) since volunteers who acquire such an infection would presumably be at risk of mixed-sex natural infections as a result of their usual behaviours and occupation. The risk of egg-related morbidity due to the presence of male worms from the CHI-S would therefore add little to the risk resulting from exposure to natural infection. CAA - circulating anodic antigen

### Table 5. Risks associated with controlled human infection with Schistosoma mansoni.

<table>
<thead>
<tr>
<th>Risk</th>
<th>Inherent risk score</th>
<th>Total inherent risk</th>
<th>Controls</th>
<th>Residual risk score</th>
<th>Total risk post control</th>
</tr>
</thead>
<tbody>
<tr>
<td>Symptoms related to infection</td>
<td>Common</td>
<td>Major</td>
<td>1) Slow dose escalation in modest increments</td>
<td>Common</td>
<td>Moderate</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>2) Frequent follow up visits and collection of adverse events.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>3) Clinical assessment and routine safety lab.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>4) Symptomatic treatment with corticosteroids or abrogating infection with praziquantel (which kills adult worms) if needed.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>5) Abrogate infection with artesunate (which kills immature forms)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Symptoms related to treatment with praziquantel</td>
<td>Common</td>
<td>Moderate</td>
<td>1) Take praziquantel with food</td>
<td>Common</td>
<td>Minor</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>2) Clinical assessment, reassurance, symptomatic relief if needed</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Misunderstanding of the nature of CHI-S studies</td>
<td>Likely</td>
<td>Critical</td>
<td>1) Education of community leaders, opinion makers and regulators</td>
<td>Possible</td>
<td>Major</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>2) Work with community advisory board</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>3) Education of potential volunteers using tested materials</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>4) Informed consent verified with tests of comprehension</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Discussion
In this document we have reflected on the potential risks involved in establishing a controlled human infection model for schistosomiasis in Uganda. The opinions expressed and risk scores allocated have been arrived at by discussion between the authors and are therefore subjective. In submitting this document to open peer review through the African Academy of Sciences Open Research Platform we welcome discussion of these issues.

Based on the assessments made, our own reflections and proposed plans are as follows.

First, we have decided not to pursue the option of importing B. glabrata snails from the Netherlands to Uganda. Although the proposed controls were estimated to reduce the risk or establishing a colony outside the laboratory to low, it seems unnecessary to incur them. Since snail species endemic to Uganda are susceptible to S. mansoni infection we expect that option 2 will work.

Second, we propose to further pursue the option of using the Puerto Rican laboratory strain of S. mansoni in the CHI-S in Uganda. We consider that the recognised virulence and praziquantel susceptibility profile of this strain makes it a safe option for CHI-S. The long-term in-breeding of the laboratory strain is an asset in this regard, making the characteristics of each clone of male cercariae reasonably predictable. The potential variability of a newly isolated schistosome population from Uganda would be a concern – even if generally of modest virulence and good praziquantel susceptibility, an individual clone might exhibit undesirable properties.

To generate infectious cercariae for human infection and challenge studies following the principles of GMP it will be essential to establish a suitably controlled snail facility in Uganda. For sustainability (to avoid the need of repeated shipping of infectious material from the Netherlands) it will also be necessary to establish a specific pathogen free animal facility to house the mammalian host and complete the laboratory life cycle.

With regard to the selection of volunteers, and avoidance of natural infection during the CHI-S, current activities include engagement with relevant Ugandan communities which are potential settings for recruitment of volunteers. As part of the engagement, options for avoidance are being explored. Our current view is that careful volunteer selection, close follow up and immediate abrogation of infection (on detection of CAA) will be preferable to 12-week “admissions”; but views from the communities will influence our future approach.

Controlled human infections with known pathogens inevitably involve risks and possibly the burden of symptoms. Available mitigations in several examples reduced our risk scores only to moderate, rather than low: for example, symptomatic treatment and early abrogation of infection cannot reduce the likelihood of symptoms below common, but can reduce the impact of the symptoms. Such areas emphasise the need for caution – for example, small group sizes and carefully monitored dose-escalation approaches.

We realize that symptoms may be different among Ugandan volunteers than among Dutch volunteers. Particularly, Katayama fever is considered less likely to occur in subjects from endemic, compared to subjects from non-endemic settings1. Nevertheless, we shall provide full information to potential volunteers about symptoms predicted from the literature, and those which occurred previously in the Dutch volunteers. We are currently piloting educational materials, volunteer information sheets, and tests of comprehension in order to ensure that Ugandan volunteers can be enrolled with genuine understanding and fully informed consent. As well, we shall work with community leaders and advisors to ensure optimal understanding of the work, and to mitigate the impact of rumours about the work which are likely to arise.

We conclude that, with careful risk management, CHI-S can be safely implemented in Uganda with a view to accelerating vaccine development against this important communicable disease.

Disclaimer
Data availability
Underlying data
No data are associated with this article.

Extended data
Open Science Framework: Controlled Human Infection Model – Schistosomiasis. https://doi.org/10.17605/OSF.IO/53GT9

This project contains the following extended data:

• Appendix 1.docx (risk assessment report addressing the intended transfer to and culturing of the snail Biomphalaria glabrata in Uganda)

• Appendix 2.docx (Summary of safety precautions for working with Schistosoma)

Data are available under the terms of the Creative Commons Zero “No rights reserved” data waiver (CC0 1.0 Public domain dedication).

Grant information
The work was supported by a pump-priming grant from the HIV-Vac network. The HIC-Vac network is supported by the GCRF Networks in Vaccines Research & Development, which is co-funded by the Medical Research Council (MRC) and the Biotechnology and Biological Sciences Research Council (BBSRC). This UK funded award is part of the EDCTP2 programme supported by the European Union.

AME is a fellow of the African Academy of Sciences.

The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Acknowledgements
We thank Dr A. J. de Winter of the Naturalis Biodiversity Center, Leiden, The Netherlands for his expert contribution on snail biology.


Open Peer Review

Current Peer Review Status:

Version 1

Reviewer Report 26 June 2019

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Donald Harn
Department of Infectious Diseases, College of Veterinary Medicine, Center for Tropical and Emerging Global Diseases, University of Georgia, Athens, GA, USA

Overall, the authors have put together 3 different scenarios that would eventually provide for controlled human infections with male schistosomes in Uganda. The authors present good rationale for why such CHI may be valuable for vaccine trials, and I generally agree with their rationale.

The authors produced a document that attempts to summarize the risks associated with each scenario and how they would be reduced if proper interventions were imposed. Overall, a nicely written paper that concludes that “Option 2” introducing the Puerto Rican strain of Schistosoma mansoni into Ugandan labs, and possibly the ecology, is the best way forward. I disagree with this assessment as outlined here and feel that the more costly (initially) “Option 3” is the best way forward to minimize unknown potential ecological risks.

A few things concern me. One, the authors state “The level of risk and effectiveness of proposed controls was determined by consensus between the authors.” Some of these authors were involved in establishing CHI in Dutch volunteers in Leiden. However, I am concerned with the level of expertise the authors have in ecology or ecologic modeling to accurately assess the likelihood of introduction of new parasite or snail into the environment, and the impact of such new species into the Ugandan environment?

For Option 3 the authors state “There are however several challenges with using Ugandan snails and isolates. With regard to Ugandan snail species, there is variability between snail species in susceptibility to S. mansoni infection;” Wouldn't this concern be the same for option 2.?

Also under Option 3, the authors state “With regard to the new schistosome laboratory strain, the characteristics of this would be unknown in terms of virulence and susceptibility to praziquantel treatment. Determining these characteristics would not be simple, since validated tests for schistosome resistance are currently not available. In addition, the new isolate would not be clonal and variability within the newly collected schistosome population might result in variable
responses in the host, and to drug treatment. In addition, dose-finding studies would start from scratch to find the balance between tolerability and attack rate.”

I find all of these arguments not justifiable. Determining susceptibility to PZQ in their animal model is straightforward and they can do this. They can easily produce a clone or clones of schistosomes to initiate these studies. Yes, infectious dose studies will have to start from scratch, but in reality, as this is an endemic population, not Dutch volunteers, this will have to be done with Ugandans anyway.

In the “Natural Infection during trial period” the authors note that at some point they may introduce female cercariae infections. Are they implying single-sex female cercariae infections? Later in this section the authors note that infected individuals should avoid any contact with schistosome contaminated water. How feasible is this? Much of the risk here will depend on the residence of the cohorts for the Ugandan CHI trials. If the volunteers are urban, with little to no chance of encountering contaminated water, this point is moot and perhaps the authors have considered this as a likely way to mitigate this potential problem.

General, the Tables are informative. I may have missed this but, it is not clear to me how the authors determined the reduced “Total risk post control” score? We can see large swings in score from initial “Total inherent risk” to the score for “Total risk post control” but do not have a numerical rationale mentioned or discussed for why this lower score.

Minor:
Potent should be patent

Hybridisation with local schistosome population unlikely to result in praziquantel resistance? Maybe true, but you still have created a hybrid parasite.

Under “Option 1: snail culture facilities, potential ecological harm” page 5, first paragraph, line 4 should be corrected to read “access to the laboratory should be controlled...”

Is the rationale for the Open Letter provided in sufficient detail?
Yes

Does the article adequately reference differing views and opinions?
Partly

Are all factual statements correct, and are statements and arguments made adequately supported by citations?
Partly

Is the Open Letter written in accessible language?
Yes

Where applicable, are recommendations and next steps explained clearly for others to follow?
Partly
**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Vaccine design, development, and delivery. Field studies. Immunology. Tropical Medicine/Parasitology.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

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**Author Response 05 Aug 2019**

**Alison Elliott**, Medical Research Council/Uganda Virus Research Institute and London School of Hygiene & Tropical Medicine Uganda Research Unit, Entebbe, Uganda

We thank Dr Harn for his review. Our response is as follows.

Comment. The authors produced a document that attempts to summarize the risks associated with each scenario and how they would be reduced if proper interventions were imposed. Overall, a nicely written paper that concludes that “Option 2” introducing the Puerto Rican strain of Schistosoma mansoni into Ugandan labs, and possibly the ecology, is the best way forward. I disagree with this assessment as outlined here and feel that the more costly (initially) “Option 3” is the best way forward to minimize unknown potential ecological risks.

Response. We agree that the choice between the safety concerns on the one hand and the ecological risk on the other, is a challenging one. We also understand from this comment that we have not been clear about why, on balance, we prefer option 2. We have extended the paragraph in the discussion on this decision to be more specific about why this is our current preference.

Comment. A few things concern me. One, the authors state “The level of risk and effectiveness of proposed controls was determined by consensus between the authors.” Some of these authors were involved in establishing CHI in Dutch volunteers in Leiden. However, I am concerned with the level of expertise the authors have in ecology or ecologic modeling to accurately assess the likelihood of introduction of new parasite or snail into the environment, and the impact of such new species into the Ugandan environment?

Response. We agree that we, the authors, have little experience in assessment of ecological risks. For these parts of the risk assessment, we therefore consulted relevant ecologists and geneticists including Dr A. J. de Winter (Naturalis Biodiversity Center, Leiden, The Netherlands) regarding the snails and Dr. M. Berriman (Wellcome Trust Sanger Institute, UK) regarding the parasite. To make this clear, we have better outlined the considerations. With his permission, we have added Dr. Berriman to the acknowledgements.

Comment. For Option 3 the authors state “There are however several challenges with using Ugandan snails and isolates. With regard to Ugandan snail species, there is variability between snail species in susceptibility to S. mansoni infection;” Wouldn't this concern be the
same for option 2.?

Response. This is correct. We have adjusted accordingly and mention this concern for option 2 as well.

Comment. Also under Option 3, the authors state “With regard to the new schistosome laboratory strain, the characteristics of this would be unknown in terms of virulence and susceptibility to praziquantel treatment. Determining these characteristics would not be simple, since validated tests for schistosome resistance are currently not available. In addition, the new isolate would not be clonal and variability within the newly collected schistosome population might result in variable responses in the host, and to drug treatment. In addition, dose-finding studies would start from scratch to find the balance between tolerability and attack rate.”

I find all of these arguments not justifiable. Determining susceptibility to PZQ in their animal model is straightforward and they can do this. They can easily produce a clone or clones of schistosomes to initiate these studies. Yes, infectious dose studies will have to start from scratch, but in reality, as this is an endemic population, not Dutch volunteers, this will have to be done with Ugandans anyway.

Response. We agree that an inbred population of Ugandan schistosomes could be established in a rodent model, but are not confident that it would be straightforward. Many generations of crossing following initial infection with a clone of males and a clone of females would be required to produce a monomorphic strain. Initiating the strain with a single clone of males and a single clone of females would minimise variability, but might also result in quite atypical parasites, not necessarily representative of the Ugandan population of schistosomes in general. Starting with a more diverse selection of cercariae would generate a more representative laboratory population of Ugandan schistosomes, but would mean that the characteristics of any particular clone (notably pathogenicity or praziquantel resistance) selected for CHI-S would be unpredictable.

Indeed our greatest concern is the potential for praziquantel resistance. Ugandan populations have been exposed to regular praziquantel treatment for over a decade, so there is a risk that the initial isolates would include individuals with relative praziquantel resistance which could not be established with certainty in the initial stages of the above process. While we agree that some testing could be done in animals the pharmacokinetics and pharmacodynamics of praziquantel are complex and different between rodents and humans.

With regards to the dose-finding study, we agree that this would be the case for any of the three options and therefore removed this sentence.

Comment. In the “Natural Infection during trial period” the authors note that at some point they may introduce female cercariae infections. Are they implying single-sex female cercariae infections? Later in this section the authors note that infected individuals should avoid any contact with schistosome contaminated water. How feasible is this? Much of the risk here will depend on the residence of the cohorts for the Ugandan CHI trials. If the volunteers are urban, with little to no chance of encountering contaminated water, this
point is moot and perhaps the authors have considered this as a likely way to mitigate this potential problem.

Response. We have clarified the sentence on female cercariae – we indeed plan on developing a single-sex female model in the future.

With regard to avoiding contact with contaminated water, the location we are considering to recruit volunteers for this work is a peri-urban fishing village close to Entebbe, well known to the research team, where many communal taps are available to provide an alternative water source. In addition, it will be possible to select volunteers who have access to adequate sanitation. We thus envision that it will be feasible to avoid any lake contact for these inhabitants. Through surveys and group discussions, we are currently assessing possible strategies to incentivise the use of tap water – for example, making it freely available for study participants. We hope that this will allow participation from the true target population i.e. people with previous exposure to *Schistosoma mansoni*.

Comment. General, the Tables are informative. I may have missed this but, it is not clear to me how the authors determined the reduced “Total risk post control” score? We can see large swings in score from initial “Total inherent risk” to the score for “Total risk post control” but do not have a numerical rationale mentioned or discussed for why this lower score.

Response. The inherent risk was defined as the risk before putting controls in place, calculated as the product of the likelihood and impact scores. The residual risk was similarly calculated, based on likelihood and impact scores after controls have been put in place. The difference in the calculated scores is based on the degree to which the proposed controls are expected to alter either the likelihood of the event happening, or the impact of the event, should it happen. We have added the scores to the legend of the tables for clarity.

Comment. Potent should be patent

Response. We have changed this.

Comment. Hybridisation with local schistosome population unlikely to result in praziquantel resistance? Maybe true, but you still have created a hybrid parasite.

Response. This is true. We have added a sentence to include this concern.

Comment. Under “Option 1: snail culture facilities, potential ecological harm” page 5, first paragraph, line 4 should be corrected to read “access to the laboratory should be controlled...”

Response. We have corrected this.

**References**

Reduced Efficacy of Praziquantel Against Schistosoma mansoni Is Associated With Multiple Rounds of Mass Drug Administration. Clin Infect Dis. 2016 Nov 1;63(9):1151-9

Competing Interests: No competing interests were disclosed.

Reviewer Report 17 June 2019

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James E. Meiring
1 Malawi-Liverpool Wellcome Trust Clinical Research Programme, Blantyre, Malawi
2 Oxford Vaccine Group, Oxford University, Oxford, UK

This is a well written, and referenced letter outlining the risk assessments for setting up CHI-S in Uganda. The authors should be commended for a very thorough approach to the topic, highlighting all areas of risk, from shipment and storage of infectious diseases and animal hosts through to the community-based responses to CHI-S and possible impact on other research activity.

In addition to what is presented here, I think it might be helpful for the authors to outline their thinking on expectations for financial reimbursement for participants who go through the trial in Uganda, and if there are any risks attached to this, as discussed in Gordon, S et al. 2018 published in Wellcome Open Research1.

It may also be beneficial to consult the literature for other examples of CHI studies that have been moved from non-endemic, high income countries to endemic sites. Although the individual risks may be different, the general lessons may be applicable and helpful.

References

Is the rationale for the Open Letter provided in sufficient detail?
Yes

Does the article adequately reference differing views and opinions?
Yes

Are all factual statements correct, and are statements and arguments made adequately supported by citations?
Yes

Is the Open Letter written in accessible language?
Yes

Where applicable, are recommendations and next steps explained clearly for others to follow?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Typhoid Fever epidemiology and vaccine trials. I was a clinical research fellow in CHI for typhoid in Oxford, UK, with clinical responsibility for patients, and am currently study clinician and co-PI for a typhoid vaccine trial in Blantyre, Malawi.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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Author Response 05 Aug 2019

**Alison Elliott**, Medical Research Council/Uganda Virus Research Institute and London School of Hygiene & Tropical Medicine Uganda Research Unit, Entebbe, Uganda

Comment: In addition to what is presented here, I think it might be helpful for the authors to outline their thinking on expectations for financial reimbursement for participants who go through the trial in Uganda, and if there are any risks attached to this, as discussed in Gordon, S et al. 2018 published in Wellcome Open Research.

Response: We agree that the financial reimbursement is an important topic for debate. We have thus added a paragraph on this matter and included it in the table with risks related to the controlled human infection with Sm.

Comment: It may also be beneficial to consult the literature for other examples of CHI studies that have been moved from non-endemic, high income countries to endemic sites. Although the individual risks may be different, the general lessons may be applicable and helpful.

Response: We agree that it will be essential to interact with experienced researchers who have implemented CHI studies in LMIC. We added reference to this important topic by addressing a paper on the first malaria CHI in Kenya that describes issues that are also applicable for our CHI-S.
Competing Interests: No competing interests were disclosed.