“At first, I was very afraid”—a qualitative description of participants’ views and experiences in the first Human Infection Study in Malawi [version 2; peer review: 2 approved]

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Abstract

Background: Human infection studies (HIS) involve deliberately infecting healthy volunteers with a pathogen in a controlled environment to understand infection and support the development of effective vaccines or treatments. HIS research is expanding to many low and middle-income settings to accelerate vaccine development. Given the implementation of the first HIS research to establish the experimental human pneumococcal carriage model’s feasibility, we sought to understand the participant’s opinions and experiences.

Methods: We used a qualitative, descriptive approach to understand participants perceptions and experiences on HIS participation. Sixteen healthy adult participants were invited to participate in in-depth exit interviews to discuss their experiences, motivations and concerns.

Results: Our findings showed that the likelihood of participation in HIS research rests on three essential conditions: motivation to participate, compensation and advocacy. The motivation and decision to participate was based on reasons including altruism, patriotism, monetary and material incentives, and while compensation was deemed appropriate, concerns about unanticipated research-related risks were raised. Participant advocate groups were recommended for increasing awareness and educating others in the broader community about HIS research.

Conclusions: Participants’ experiences of HIS in Malawi provide the basis of what can be acceptable in HIS research in lower-income countries and areas where study procedures could be adjusted.

Keywords
Human infection studies, ethics, acceptability, Malawi

Open Peer Review

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Introduction

Human infection studies (HIS) or controlled human infection models (CHIM) often involve deliberately infecting healthy adult student volunteers with a pathogen in a controlled environment to better understand infection and support the development of effective vaccines or treatments1. CHIM have been conducted in different participant groups over hundreds of years and have contributed vital scientific knowledge leading to advances in the development of drugs and vaccines. Recently, CHIM have received renewed interest, particularly in disease-endemic settings, because they offer an efficient and cost-effective approach for selecting the most promising vaccines for further development in populations that bear the more significant burden of infection. CHIM allows efficacy data to be generated quickly and promotes the identification of good immune correlations, the down-selection of vaccine candidates and early vaccine formulation decisions, and therefore provide an opportunity to circumvent the large-scale field efficacy studies to deselect intervention candidates2-3.

CHIM have been conducted in low and middle-income countries (LMICs) such as Columbia, Kenya, Tanzania and Thailand, Gabon, Equatorial Guinea, and Mali. Zambian scientists have conducted a pilot using a live attenuated Rotavirus vaccine in Zambia (Chilengi, 2020)5. Plans to conduct CHIM in other settings such as India, Vietnam, Uganda, Malawi, to name a few, are underway to explore the acceleration of vaccine development relevant to LMIC populations. While CHIM offer scientific opportunities, there are some ethical issues concerning CHIM, especially in LMICs. Some authors raise concerns around benefit-sharing, limits to risk, the right to withdraw, informed consent, compensation for participants, and research with children or other vulnerable participants, for example, pregnant women6. Other concerns relate to compensation, particularly if participant exposure to risks or harm is perceived as high, and potential burdens of participation are perceived to be impactful7. Yet other concerns relate to issues of governance of CHIM research, particularly in LMICs. The lack of specific ethical frameworks and guidelines for ethics committees raises issues about a lack of review procedures that will guide the evaluation of human infection research8. Other authors argue that these concerns are not unique to LMICs. Although the ethical issues may be more amplified in LMICs, they do not differ significantly from the ethical issues raised by CHIM in high-income countries9. Despite the concerns, CHIM offers potential opportunities to quickly identify and develop vaccines and provide them to those who need them.

Evidence on experiences and decision making among CHIM participants remains limited, but a small number of articles have pointed to a range of motivations for taking part. Research with malaria CHIM participants in Kenya found that monetary compensation was a primary motivation10. In the US, alongside financial incentives, participants reported altruism, and experiential motivations, such as curiosity, as reasons for participating in CHIM research11. Beyond CHIM, there is limited evidence on motivation among healthy research volunteers.
Only a few studies have examined why they join research, how they evaluate risks, and how well they understand these risks\(^9\). A review of the literature on motivation among healthy volunteers across a range of clinical research studies (though almost all in high-income countries) found that participation was motivated by financial reward, contributing to science or the health of others, accessing ancillary healthcare benefits, interest in the science or goals of the study, meeting people and curiosity\(^7\). Exploring participants’ perceptions and experiences in LMICs is essential for understanding what is appropriate, acceptable, and practically feasible to provide ethical guidance for the conduct of CHIM research.

CHIM in Malawi

The Malawi-Liverpool Wellcome Trust Clinical Research Programme (MLW) has recently completed a feasibility study of experimental human pneumococcal carriage (EHPC) in Malawi\(^1\) under an umbrella programme titled: Malawi Accelerated Research in Vaccines using Experimental and Laboratory Systems (MARVELS). MARVELS is a clinical research programme conducting CHIM for vaccine development – targeting both transmission and infection. The current research question that the MARVELS group of researchers seek to address is why pneumococcal conjugate vaccination in Malawi does not result in the decrease in vaccine-type pneumococcal carriage seen in other parts of the world. This question is highly tractable to the EHPC model. Before implementing the feasibility study, the MARVELS group conducted a formative qualitative study interviewing research staff, clinicians, district health officials, ethics committee members, medical students, and community representatives from rural and urban Blantyre to gather views on the acceptability of CHIM in Malawi. The study’s findings revealed that acceptability depended on various factors related to informed consent procedures, inclusion criteria, medical care and support, compensation, regulation and robust community engagement. The interviewed stakeholders also expressed concerns about the safety of study volunteers and distrust or confrontation from community members if a participant were to become ill\(^1\). The findings informed the implementation of the pneumococcal feasibility study. In particular, information on stakeholder views helped MARVELS researchers understand how to design the study so that recruitment procedures addressed stakeholder concerns yet still maintained the scientific procedures based on an established pneumococcal carriage model from the Liverpool School of Tropical Medicine (LSTM), UK\(^2\).

Aim of the study

Building on the formative qualitative study of stakeholder views, this study aimed to describe the experiences, motivations and concerns of the enrolled healthy adult volunteers who had completed the MARVELS feasibility study. Specifically, we wanted to explore their opinions on study recruitment and consent procedures, medical care and support, compensation, and community engagement. We needed feedback from participants to help establish some degree of acceptability, identify areas where CHIM researchers in Malawi could improve the study design to ensure participants have a positive experience, minimise risk perceptions, and maximise their comfort. These findings will help inform the design of a pneumococcal conjugate vaccine (PCV13) trial in Malawi similar to that carried out in the UK\(^2\).

Methods

Overview of the MARVELS pneumococcal feasibility study

In the MARVELS pneumococcal feasibility study, 24 healthy adult student volunteers were inoculated with viable Streptococcus pneumoniae serotype 6B or 0.9% saline (sham inoculation) to the inside of each nostril. Blood, throat swabs, saliva, nasal scrapes, and nasal wash samples were obtained at days 2, 7, and 14 post-inoculation following the pneumococcal challenge and participants were discharged from the study on day 21. Of the 24 participants enrolled in the study, some participants received a dose between 20,000 or 80,000 bacterial colony-forming units to each nostril (naris) in 100 µl of saline. A pre-specified randomisation list determined the randomisation of participants. Participants were monitored for safety and the establishment of the pneumococcal carriage during their enrollment. Participants were provided with a thermometer and antibiotics following inoculation. They were advised to monitor their temperature daily and report any signs and symptoms during the enrollment period. Any reported symptoms were characterised as mild, moderate or severe by the study doctor and treated according to a standard operating procedure. Participants were required to contact (text message/phone) a specified member of the research team before 12:00 hours every day for seven days post-inoculation irrespective of whether they had experienced symptoms or not. Participants were provided with hotel accommodation for the first three nights. A field nurse was stationed at the hotel during all participant overnight stays to monitor any adverse reactions that may have occurred within 24 hours of receiving the inoculum. Participants were then checked out from the hotel after their stay and reminded to return to the clinic on the scheduled visits. Participants received MWK 8,400 (~11 USD) per study visit as reimbursement for out-of-pocket expenses such as travel and compensating for time spent and burdens incurred while participating in the study. In total, participants received MWK 67,200 (~91 USD). The compensation offered in this study was consistent with the remuneration guidelines published in Malawi\(^1\), paid pro-rata (per activity and not dependent on the completion of the study).

Recruitment

Before recruitment, enrolment and consent procedures for the feasibility study, the MLW Science Communication department organised the public engagement activity at the University of Malawi’s Polytechnic college through the Polytechnic dean of students office and the student’s union (PSU). The Polytechnic campus is approximately 3km from QECH. After the public engagement activity, interested participants were given contact information and invited to a one-on-one visit to the research clinic. There they were briefed about the research and study procedures, including their participation in an exit interview. They were provided with an opportunity to ask questions and
seek further clarifications. During this information visit, the research team did not ask participants to consent to the interview on the spot to give them a cooling-off period to make a decision, and subsequent non-engagement was taken as a decision not to participate. These two processes (the public engagement activity and the one-on-one research clinic informational visit) allowed participants to meet the research team, ask questions and have time for contemplation and reflection. These processes were designed to improve their understanding of the research and help their decision-making. Only on the day of screening and enrolment was consent for participation obtained.

### Study site

Interviews for the qualitative study were conducted in a private meeting room in the MLW research institution building. MLW is based in Blantyre and is situated adjacent to Queen Elizabeth Central Hospital (QECH). QECH is the largest government referral hospital in the country, with an official bed capacity of 1350. The hospital serves as the College of Medicine’s teaching and research hospital.

On their last scheduled study visit (day 21), we invited participants to participate in an exit interview. Study participants consented to participate in the exit interviews in the MLW research institution building. The sample size was not defined because the MARVELS pneumococcal feasibility study was powered to recruit only 36 participants to establish carriage. By the time we experienced our first wave of COVID-19 in April 2020, a total of 24 participants had completed the study. At that point, we interviewed 16 participants and observed through ongoing data analysis that no new information was being gathered or learned. At which point, we determined to have achieved saturation.

Biases within qualitative research are well-known, and although the exit interview was held on MLW premises, specific attention was given to avoid biases. Particular, attention was given to address interviewee biases knowing that interviewees may choose to withhold detailed descriptions or embellish them, mainly if the ‘truth’ is inconsistent with their preferred self-image, experience, opinion or wish to impress the interviewer. To address this concern, the interviewers made sure to probe, seek clarification, and continually refer to what the participant had said, mainly if there appeared to be contradictions. Attention was also given to reflexivity threats because while researchers wish to adopt a relatively neutral role, they may inadvertently demonstrate a preference for a particular perspective, and in the process, bias their findings. To this end, the two main social science researchers (NT and LMT) worked independently from the clinical team with a clear objective for reporting on the participants’ ethical concerns and experiences and continually discussed and agreed on codes, categories, and themes.

### Interviews

We used a semi-structured exit interview-guide that was piloted and tested in March 2019. The interview-guide (Table 1) was piloted on two health workers to check for clarity, relevancy, comprehensiveness, and question flow. Questions that we identified as ambiguous were amended. Two social science researchers (NT and LMT) from the MARVELS project conducted exit interviews together or separately. Most interviews were conducted primarily in English, with one conducted in Chichewa, based on participant preferences. All interviews were conducted face-to-face. Interviews lasted about 60 minutes each, and the open-ended questions covered topics about reasons and decisions to participate, understanding of the purpose of the study, procedures and risks, views of information provision, satisfaction with study experiences, compensation and any social impacts on participation, for example, education, family or home environment. The semi-structured interview topics were developed based on common concerns about human challenge studies raised in ethics literature and the formative research by Kapumba et al.

### Data analysis procedures

Interviews were audio-recorded and transcribed by NT. Transcripts were de-identified and uploaded onto REDCap version 10, a web-based application used to capture data in a secure environment so that research teams can collect and store highly sensitive information for data management and analysis. Data was coded systematically and manually by two researchers (NT and LMT) in Microsoft Word version 16. NT and LMT developed and iteratively refined the codebook beginning with a priori codes from the interview guide. We used the research questions to group the data and then look for similarities and differences. We developed a hierarchical coding framework to analyze texts based on participants feelings, opinions and experiences, with broader higher-order codes providing an overview and detailed lower order codes allowing for distinctions to be made within and between cases. NT and LMT each coded transcripts separately, discussed the codes, reconciled any discrepancies, and then summarised each code’s content. The codes were sorted into themes. These summaries formed the basis of our thematic content analysis. We achieved saturation from the 16 participants interviewed. No new codes occurred from the data, suggesting that no further or new information was being gathered or learned about the volunteers’ perceptions and experiences.

### Ethical approval

The EHP C study, consent forms and interview guides were approved by the National Health Sciences Research Committee (NHSRC) ethics committee (protocol number 19/08/2246) and the Liverpool School of Tropical Medicine (LSTM) ethics committee (protocol number 19-017). Participants were provided with an information sheet and verbal explanations, and they provided oral and written consent.

### Results

We interviewed 16 participants that were recruited between December 2019 and March 2020. Of the 16 participants, 15 were interviewed after all study procedures and after the final study visit and one participant was interviewed on a subsequent date rescheduled for participant convenience. The sample of participants is displayed in Table 2.
| Table 1. Participants Exit Interviews Topic Guide. |

| Question | Probe on the following |
|----------|------------------------|
| **General views on the study** | |
| How did you hear about this study? | Motivation to participation? |
| What made you decide to take part in this study? | |
| What are your general experiences of participating in this kind of study | Anything that made you unsure about joining the study? |
| How does that compare to how you feel about it now? | |
| **Recruitment** | |
| What do you think about our study participant recruitment approaches? | Access to target population, use of flyers? |
| What did you find the most important part of the recruitment process? | Participant information sheet? Opportunity to provide further details? |
| What other recruitment approaches need to be considered for similar studies in the future? | |
| **Screening and consenting** | |
| What can you tell me about your experiences with the screening procedures which included checking your eligibility to participate in the study? | Sample collection procedures, screening questions, and HIV testing? |
| What was your experience with nasal washing and with nasal scraping? | Preference and why? |
| **Pneumococci Inoculation** | |
| What can you tell me about your experiences with the procedures of being infected with the pneumococcal bacteria? | General feeling, fears, anticipated and unanticipated symptoms and AEs, disease expectations, and laboratory results for the collected samples? |
| Did you have any thoughts on the risks of possible unexpected and (unconsented harm) from being infected with pneumococcal bacteria? | Harm to self, including possible harms to others and the environment? |
| **Safety monitoring** | |
| What symptoms did you experience during the study period? | |
| What is your general overview of the safety monitoring processes put in place? | Whether the safety monitoring procedures put in place satisfactory? |
| Which approach worked for monitoring your safety? Calling and or SMS systems? | |
| Was it enough? What need to be considered for future CHIM? | |
| Follow-up visits – how convenient were they to you? | Time, frequency of the visits in days, facilities, travel, sample collection procedures? |
| Were you given antibiotics to take at home? How important was it for you to have antibiotics at home? What need to be considered on the issue of antibiotics from your experience? | Information given on when to take the antibiotics? |
| **Residential stay** | |
| What do you think about your experience of staying at Grace Bandawe for 3 days during the study? | Concerns, challenges, family or relations, work, or school demands? Effect on daily life experiences? |
| What if you stayed at home? Would you consider that important? | |
| What would need to be considered to ensure the residential stay is okay for future CHIM studies in Malawi? | Accommodation, food, time? |
As displayed in Table 2, most of our participants were male (n=11), and the remaining were female (n=5). Most were 3rd and 4th-year university students, while 1 participant had completed their secondary school education (O-level equivalent). All participants were 18 years or above, and all indicated that this was the first time they had ever participated in a clinical research study.

We present our findings under four main themes. First, we present participants’ decision making around participation in the pneumococcal feasibility study, including their views on the public engagement event at the Polytechnic campus and study consenting procedures described above in the recruitment procedures section, and motivations for participating. Second, we describe their experiences with study procedures and methods, including safety monitoring plans and the three-night hotel stay. Third, we discuss views on compensation. Finally, we present the participants’ suggestions for engagement and recruitment for future CHIM.

Participants’ decision-making and motivations for participating in the EHPC feasibility study
Most participants informed us that they decided to join the study after attending a public engagement event organised by the MLW research team.

“...the very first time they came to school [the Polytechnic], they presented the study, but there were some areas, which I did not understand, and there were questions, which I could not ask in the presence of others, you could like to ask in person with the team doing the research. I had that time to ask, and they told me all the relevant information about what is it all about for the research. So that helped me, that was a good move that I had that ample time to speak out anything I was afraid of, and they could answer me.” (CHIM 1028)

Participants particularly liked the private setting at the research clinic, which enabled them to ask questions they would not have done in their peers’ presence after the public engagement event and before enrollment.
Since most of the volunteers were students studying at a university, it was common for them to use the Internet to research the study independently. The participants informed us that they conducted Internet searches on the pneumococcal carriage model from the LSTM to understand the pneumococci bacteria they would be inoculated with and the inoculation procedures to assess the study risks.

“To know more about bacteria, I researched...I just Googled pneumococci, and the Internet explained what this bacteria, pneumococci was... They said that this is mostly or is mostly in kids, if I am not wrong. Those affected most are children or those whose immunity is low, like people with HIV and AIDS and the elderly whose immunity system is low that bacteria can affect them, not healthy adults... I wanted to know more about this study... It helped me to have confidence enough that I join the study.” (CHIM 1036)

One participant consulted their family in the process of deciding to participate in the study. While the relatives did not offer a direct opinion, the participant felt that this was their tacit way of giving permission and reassurance that a sensible decision was being made.

“Maybe the part I can explain about, which made me reach the point of deciding to participate was, it was when I came to explain to my relatives at home after I gave them the form after they read it through, I felt to say: ah I think I can participate (CHIM 1184).

The motivation to participate was varied and based on perceived individual benefits and societal value. As one participant put it,

“My interest is to focus much on research, so the most important reason is to participate and gain more knowledge” (CHIM 1069).

A couple of participants were motivated to participate as they were very keen to know their general health status, which included an HIV antigen test.

“I was so motivated when they said we would have your health check-up. In most cases, I have been looking for that, but then if you look at the money going to the hospital and then having the check-up, it would cost me a lot. So, I was partly motivated because of that that I should know how I am. Even though the study did not have that fully checking up of the entire body, you had just some special areas that you were supposed to look at, but still more it has been good” (CHIM1119).

Only one participant mentioned money as the primary motivating factor for participating in the EHPC study.

“Besides money [laughs], but otherwise no, I just wanted to take part... Anyway, okay, generally, money is a basic need, it is a necessity, so with sixty-seven thousand, it was at least an attractive package” (CHIM1010).

The social value of participating in the research was expressed in the form of altruism and patriotism. Almost all participants spoke of the desire to help humanity and Malawians impacted by pneumonia and the scientific community develop a better vaccine.

“I just felt like at least I should be one of the people that could contribute to something good to the entire nation because I know this could be. It is something that has been recommended in Malawi. Therefore, it would carry a certain value. So, I thought of being part of it” (CHIM 1119)

Patriotism, expressed in the form of self-sacrifice or selflessness, was articulated when one participant described himself as a “risk-taker”.

“Ah, here I can say that I had no worry considering that I am a risk-taker. So joining it was just like that. I think I made up my general mind that I am joining this, and if they are telling me that this is the way it is handled. You will experience this and that, and nothing else can harm me as they said that it worked 100% in Liverpool. I was, like ah. I think this is just okay after so my first days when I experienced nothing, I did not have a fear that anything can happen” (CHIM 1028)

While this participant expressed the view that he was a “risk-taker”, his motivation and ultimate decision to participate were balanced by assessing the risks and the potential benefits.

“My interest was that I would be the one involved in helping the community because I know that when a vaccine is found, and I was involved in the process of making the vaccine, you will be reaching to many people that I cannot personally reach” (CHIM 1028).

Experiences with study procedures and methods

Participants described positive and negative aspects of the study procedures. Positive factors included staff attitudes, safety monitoring and support for health care.

Several participants commented on the research staff’s friendly nature, which helped participants feel comfortable with the study follow-up visits and clinical procedures.

“The nurses were friendly, the way they talked to us, the way they were handling us, it was just okay, and I could feel that sense of closeness, sure. I think, for me, I can say it was just okay, because even they were flexible may be telling them that we will meet at such a time or that maybe you have changed time due to some changes, they would understand. I think it is just okay” (CHIM, 1028).

Participants also appreciated the safety monitoring procedures, including the hotel accommodation for three nights after receiving the inoculum. The Grace Bandawe Conference Centre [GBCC] is situated 3.4 km (four-minute drive) from a private medical facility (Mwaiwathu Private Hospital) to provide study participants with medical care should they experience a severe adverse event.
“My experience was great. Yeah, everything about food, the place that we were sleeping, and it was a comfortable place and a very conducive environment for research like this” (CHIM 1085).

“You provided a person with all the options which he or she can follow if at all is feeling unwell. For example, you have given us an allowance, let’s find somebody gets sick while he is maybe somewhere very far, so he could travel using that money to go to the hospital. You also gave us a card whereby you can go to the Mwaiwathu hospital at any time where we were feeling unwell. The third was being accommodated at Grace Bandawe, which is very close to the Mwaiwathu hospital. Now, that thing itself is very good and has caused my experience to be excellent as well” (CHIM 1143).

Participants also appreciated receiving other forms of immediate medical care and support as part of the study safety monitoring. This included being escorted by a nurse/fieldworker for the nights spent at the hotel, and the ability to access the medical team at all hours via cellphone, and a safety information sheet listing possible side-effects, such as a temperature of >37.5°C, shivering, headache, new rash, drowsiness, cough, earache, and or new eye infection. Participants were also provided with a thermometer to monitor their temperature personally and provided antibiotics to take if their temperature was too high.

Because having a thermometer that’s, because one of the symptoms of you know these bacteria is a rise in body temperature. So, having that in mind, I think that is enough. Furthermore, that card, like the information sheet, was good. There were just so many points that you cannot miss out on them, you see, maybe you are feeling something, it was listed on the card, so, that was enough for me” (CHIM 1010).

There were mixed views on the actual inoculation procedures. Most participants expressed initial feelings of fear and apprehension about receiving the inoculum.

“At first, I was very afraid. Actually, I was so afraid of, okay, I that running nose, that’s exactly one thing that I hate the most. I fail to study when I have a runny nose, so I hate it a lot” (CHIM 1093).

Another participant also felt unwell after receiving the inoculum and reported experiencing fatigue.

“Somehow I was feeling like just tired, somehow a little headache and that general feeling that I am not well, but I cannot explain it the way it was, but just feeling that I am not okay” (CHIM 1051).

Other concerns about inoculation related to the potential harm that the bacteria can cause to others.

“Yes, you know the social world we live in we always associate with people. So whenever you are with a friend you have met, certain family and they have a kid, you feel like carrying the kid, but when you think of oh, I am going under this, so you are somehow refraining, avoiding meeting people, avoiding going to family members, like family friends to chat, like there are kids there they will need me to carry. I have a sister, and she has a kid and that day, the kid was seven months old, so I never held her. It was just like I was avoiding her (CHIM 1093).

But many also talked about feeling reassured because of the antibiotics the study provided, the independent research they had done before joining the study and the fact they did not experience any untoward event.

“I was nervous. A bacteria is a bacteria; it is a microorganism. Sometimes it can decide to misbehave. There can be something that would alter the body, like the chemical reactions and the likes, so, I was afraid, of course, I was nervous. However, I knew, like from education-wise, we know what bacteria do, and there are treatments for bacteria. Yea, so, if anything, then take the antibiotics” (CHIM 1010).

Despite the worries, one study participant reported feeling at ease with receiving the inoculum.

“My experience to say that after inoculation as I said at first that I was like somehow worried but in the process, it was found that the worry disappeared because I did not experience anything unusual or that I was feeling sick. I was just okay the way I was even before joining the research. Hence I had the confidence that aah, I think these things are all right because the way they explained before the research that there is nothing harmful that can happen after inoculation, after being inoculated the bacteria there was like no harm that happened in line with what they said” (CHIM 1028).

Participants were also asked about their experiences with sample procedures, including the throat swab, nasal scraps, nasal washes, blood and saliva samples. Most participants were comfortable having their blood and saliva samples taken. However, many found the nasal scraps, nasal washes and throat swabbing procedures uncomfortable. Holding water in the nose and then expelling it created discomfort. Similarly, having a throat swab sample taken was awkward because of the gag reflex.

Compensation

Questions regarding the appropriate compensation levels for CHIM research in Malawi required evaluation. Participant views on the monetary payments they received varied. Some participants felt that the amount of compensation provided was reasonable.

“The study was not that demanding, and comparing the risk and the compensation that we are getting, it is fair, yes. So, the study was not demanding that much, and the clinic was just in a convenient place. It was easy to
get to the clinic. So, mainly I would say the compensation was mainly for the airtime and the transport I would use to get to the clinic” (CHIM 1150).

However, others felt the compensation was inadequate compared to the perceived health issues that might appear after study participation is complete.

“What we are dealing with here is the human health of which if somebody has been inoculated with that bacteria, we never know some further reactions in their bodies that could happen even after the study maybe one year after now or two years from now. You know some things may happen after a very long period. Maybe this will have an adverse effect in the future. We never know. So, if we are to look at the health status of that participant, then the money is not enough” (CHIM 1119).

Another participant believed that monetary compensation for research burdens could not be equated with the value of life. Therefore, money would not be able to compensate for the loss of life.

“It is just a little money… I told you that it is about health hazard when you are making a decision you are sure that you are putting your life at risk, you never know what may come out of the study” (CHIM 1127).

Apart from monetary compensation, participants were asked to reflect on what other payment forms would be suitable to offer in CHIM. One notable view was to provide participants with health insurance to help compensate an individual should they fall ill during and long after the study is completed. Other ideas suggested were to offer participants risk allowances to cover unexpected and unpredicted costs, including t-shirts and certificates of appreciation to acknowledge their role and contribution to research.

Suggestions for engagement and recruitment in future CHIM research in Malawi

Participants in the study offered a mixture of suggestions on using community and public engagement platforms to communicate and promote understanding about CHIM research, particularly if future studies plan to recruit individuals from the broader community who are not as educated or literate as university students. Key among the suggestions offered were participant advocates in the public and community engagement activities to provide first-hand testimonials about their experiences in participating in CHIM research.

“For them to understand, they need people who have participated in this study, not just telling them that you have to do this but the experienced one” (CHIM 1127).

Another participant echoed the above sentiment and explained how involving former participants as advocates would help dispel possible misconceptions about this type of research that deliberately infects an individual with a disease-causing agent.

“some people might have misconceptions about the study. So, to hear from somebody who has gone through the study, their experience might be a bit more, how I can put this? They might feel a bit more reassured that somebody who is like me in some sort went through the study and is testifying. So, I can put it that way that the study is not as bad as I think it is or it is not what I think that the study is all about. So, to say for participant advocacy groups are very important” (CHIM 1150).

Social media platforms such as WhatsApp was also proposed as a more effective way to reach people.

“I feel like the approach is good, but maybe you can also employ social media, especially WhatsApp, because nowadays people are so active on WhatsApp. Sometimes just posting posters around, most people do not like looking at posters around, maybe with our generation. However, when something circulates on WhatsApp, it is easy to reach people. So, maybe you can include the spreading of the message using WhatsApp” (CHIM 1135).

Other recommendations included offering study participants full-time residency for the entire study period and home visits as another way to bolster CHIM safety and monitoring procedures.

Discussion

This is the first study in Malawi to examine participants’ perceptions and experiences of participating in a CHIM. Participants reported initial fears and concerns before they enrolled. Their anxiety levels dissipated after inoculation, primarily because of the strong communication with the nursing team and assurances by the study safety monitoring procedures, including a three-night residential stay at the GBCC after inoculation. From this, we can infer that CHIM research is acceptable in Malawi. However, participation in CHIM research rests on three essential conditions: motivation to participate, compensation and advocacy. In this section, we compare our findings to existing literature and map out recommendations for future studies.

Our findings on the factors influencing the decision and motivation to participate in the study included altruism, patriotism, financial benefits and medical health checks. This mirrors previously reported results in Malawi, where access to healthcare, monetary and material incentives given were the main reported reasons for participation in biomedical research. Furthermore, the decision to participate was also influenced by the methods used for recruitment. For example, the public engagement event held at the Polytechnic college, where the Liverpool EHPC research was presented and demonstrated to be safe, helped establish integrity and trust. Moreover, the research team’s attitude, which was described as friendly and flexible during screening and enrolment, further helped influence participants’ decisions to participate because they gained trust in the research team.
However, these views and experiences, which informed decisions to participate, can be partly attributed to cognitive dissonance theory. Cognitive dissonance suggests that individuals tend to seek consistency among their cognitions (i.e., beliefs and opinions) that can influence behaviours and actions. In turn, these beliefs and views can help consolidate the perception that researchers have their best interests at heart. Cognitive dissonance tends to occur in situations where an individual must choose between two incompatible beliefs. For example, believing that research participation may have risks versus feeling that the researchers would not have suggested participation if the activity was risky. Participants in the MARVELS study may have decided to participate and stay in the study to counter their beliefs around risk because of the study’s material and monetary incentives. This kind of reasoned behaviour must be approached with some degree of caution, particularly for the future of CHIM research in Malawi, as it relates to evaluating whether participants are making an informed decision to participate in the study. Future research on decision-making in CHIM needs to interrogate whether participants understand the differences between the study’s goals and personal goals.

While financial compensation for participation in research was a motivational factor, for the most part, our findings show that the notions of altruism, patriotism, perceived individual benefits, societal value, and volunteerism were primary reasons. Altruism, patriotism, and volunteerism were important aspects of achieving informed consent. This showed us that participants appreciated the risks, benefits and burdens. The issue of risk-allowances and health insurance pointed to participants’ concern about how researchers would take care of unforeseen or unanticipated research-related illnesses. Similar problems were noted in a study done in India that reported on public perceptions of CHIM. The MARVELS feasibility study paid for no-fault insurance for participants to compensate for those who would sustain research-related injuries. This is a regulatory requirement for high-risk research in Malawi.

Nevertheless, risk-allowance and health insurance require some consideration, mainly these concerns expose CHIM researchers to additional ethical obligations beyond simply paying for time and inconvenience. It requires scientists, regulatory and ethics experts to consider paying for risk in medical research, thus incorporating compensating for actual or possible harm that may occur due to a study. In essence, Grimwald et al. propose a “payment for risk model” that involves paying for time, pain, inconvenience, and risks associated with participation. The payment for risk and pain would cover unforeseen impacts participating in CHIM on participants mental health and physical well-being. The payment for risks and pain would be similar to the type of compensation provided to those taking on risky jobs, especially when those jobs are a direct service to society. Risk payment supports the legal obligation of responsibility in research. It reflects the ethical commitment to establish justice and fairness in research. It serves as a reminder to CHIM researchers in LMICs like Malawi that “an injury to one is an injury to all.” The payment of risk model would help build public trust by demonstrating a commitment to the value of life, which is consistent with the duty of care, especially in CHIM research, where the possibility to cause harm could have severe implications for public support and willingness to participate. Providing participants with clinical trial insurance, health insurance, and risk-allowances can be interpreted as the researcher and participant working together to share the risks and burdens inherent in CHIM research.

Participant advocate groups (PAGs) for CHIM in Malawi could be an essential tool for community and public engagement. This study’s findings highlight an opportunity to establish PAGs to help researchers conduct CHIM with public awareness. PAG members would comprise previous research participants who volunteer their time, knowledge and experience in research. Their role would not be to convince people to join in CHIM or facilitate recruitment, but rather to learn about the public’s fears, concerns and expectations and to respond to the questions and concerns that communities and the general public may raise, ideally before the implementation of the study and their participation. PAGs, because of their experience, would help dispel misconceptions, increase awareness of health and medical research and improve engagement between researchers and the public. Moreover, PAGs could be a means to engender the ethical conduct of CHIM research further. Like community advisory groups (CAGs), PAGs would also be the eyes and ears of researchers and the community and, in so doing, be able to protect the public’s interest and help them gain trust in the research process.

Our study has some limitations. Before starting the MARVELS study, we did not collect qualitative data on initial feelings and questions or concerns that participants may have had before joining the research. This is because we anticipated that potential participants who chose to enrol in the study would be open about their opinions at the end of their participation in the study. However, future qualitative research work must consider including this approach to help deepen our understanding of the underlying motivation to participate in CHIM research in Malawi. Moreover, the participants were interviewed at the clinical research institution after exiting the study. This may have affected openness because of the perceived links between the interviewers and the research team. Also, we may have ignored others’ views; for a more rounded assessment on the experiences, additional insights on experiences and acceptability with frontline research staff would have been helpful. Wider opinions from family members would also be necessary for understanding acceptability beyond immediate participants’ views. This would have given us a better understanding of informed consent, acceptable levels of risk and compensation. A varied view of perspectives and experiences in implementing and participating in CHIM research will help deepen our understanding of acceptability in Malawi and contribute to the growing discourse and literature on the ethically acceptable approaches to CHIM, and develop ethical frameworks, guidelines and principles on how to conduct CHIM in LMICs.
Conclusion
Though participants were, at first afraid of participating in the first CHIM human pneumococcal carriage research, participants’ experiences of CHIM in Malawi provide the basis of what can be acceptable and areas where study procedures could be adjusted. MARVELS will use these to design the next steps of their research. Additionally, findings will also inform regulatory thinking on guidelines, frameworks and principles on how ethics review committees in Malawi should handle the governance of CHIMs. Finally, future research ought to consider issues around the theory of cognitive dissonance and the role that it can play in explaining participant experiences and decision-making to bring greater transparency about the acceptability of CHIM in LMICs.

Data availability
Underlying data
The data generated and analysed are not publicly available because consent was not obtained for these to be made public even if anonymized but are available from the corresponding author on reasonable request.

Acknowledgements
We want to thank our study participants for their valuable time because we would not be able to share our findings without them. We would also like to thank MLW’s Science Communication team, the Polytechnic dean of students and the student’s union for facilitating the public engagement event.

Author’s contributions
KG developed the protocol, conceived the study design and topic guide, and these were adapted by BK. NT and LMT collected the data. NT translated and transcribed the data. LMT initiated the development of the codes and the codebook. NT completed coding the data. LMT reviewed the data, codes and developed the themes. LMT conceived the manuscript and wrote the first draft. NT contributed to the draft manuscript. KJ, JR, SB, and BM provided input on HIS plans supporting social science work. SG secured the funding and supported the social science work. All authors were involved in reviewing and providing comments to the first draft and revisions to the paper. All authors read and approved the final manuscript.

MARVELS consortium
• Clinical team: Clara Ngoliwa, Edward Mangani, Modesta Reuben and Helen Thomson
• Laboratory team: Chris Mkandawire, Simon Sichone, Raphael Kamng’ona, Mphatso Mayuni and Percy Mwenechanya
• Data managers: Joel Gondwe and Clemens Masesa
• Administration: Sandra Antoine
• Collaborators: Kate Gooding, Markus Gmeiner, Mike Parker, Andrew Pollard, Rob Heyderman, Jason Hinds, Mark Alderson, Chris Bailey, Marien de Jonge, Robert Kneller, Jeremy Brown, Jane Mallewa, David Goldblatt, Richard Malley, Jeff Weiser, Jonathon Grigg, Henry Mwandumba, and Debby Bogaert

Ethical approval
We obtained ethics approval from the National health Sciences Reseach Committee (NHSRC) ethics committee (protocol number 19/08/2246) and the Liverpool School of Tropical Medicine (LSTM) ethics committee (protocol number 19-017). We sought permission from the Blantyre District health Office (DHO) and the Queen Elizabeth Central Hospital’s Director to conduct this study. Participants were provided with an information sheet and verbal explanations, and they provided oral and written consent.

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Open Peer Review

Current Peer Review Status:  

Version 2

Reviewer Report 12 October 2021

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Evelyne Kestelyn
Oxford University Clinical Research Unit, Ho Chi Minh City, Vietnam

I have no further comments. The authors addressed all the comments thoroughly.

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Clinical trials, Infectious Diseases, Human challenge ethical and governance frameworks

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.

Version 1

Reviewer Report 25 May 2021

https://doi.org/10.21956/wellcomeopenres.18279.r43585

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Evelyn Muleba Kunda-Ng’andu
Research Department, The Centre for Infectious Disease Research in Zambia, Lusaka, Zambia

Toto et al. explored the experiences, motivations and concerns of HIS participants in Malawi. The reasons for participation included altruism, patriotism, monetary and material incentives. Some participants expressed concerns about the risks involved vis-a-vis the value of compensation they
received from the study. Participants recommended the use of advocate groups to engage the wider community.

This article provides very relevant and timely information on HIS for those planning to undertake HIS studies, especially in LMICs. This is because understanding the experiences of actual HIS participants provides insights that may be applied for better results in a HIS in similar settings. The paper makes interesting reading and provides clear methods and steps used making it appropriate for replication.

Just a few notes for the authors.

**Abbreviations**
It is important for the authors to be consistent in the presentation of these. Some have been presented in lower case while others have been presented by Capitalizing each word.

**Quotes**
Authors should re-look some of the quotes to ensure that they read clearly to reflect the theme in the discussion.

The quote below was repeated on page 9 of the paper. It is presented on page 9 and repeated immediately on page 9. Please try and find another quote to replace ‘it’, on either page or better still delete it from page 9.

“At first, I was very afraid. Actually, I was so afraid of, okay, I that running nose, that's exactly one thing that I hate the most. I fail to study when I have a runny nose, so I hate it a lot” (CHIM 1093).

**Abstract/Reporting qualitative findings/Discussion**
Authors should avoid the use of the word ‘likelihood’, as it is used to show probability. The first sentence of the results in the abstract should be revised to sound more qualitatively appropriate and clear.

**Introduction**
Where the authors describe the HIS situation in LMICs, it would be imperative to note that countries like Zambia conducted a pilot HIS and have since begun conducting a full study. Furthermore, South Africa has also started conducting a HIS.

**Aim of the study**
From the paper, I gather that the aim of the research was to explore and describe the experiences; motivations and concerns of enrolled HIS participants. In the section of the aim of the study, authors say they aimed ‘to assess acceptability among the healthy adult volunteers who had completed the feasibility study, including their opinions on study recruitment and consent procedures, medical care and support, compensation, and community engagement’. There is a need to revise this for consistency.

**Discussion**
The authors discuss the decision process using the theory of cognitive dissonance and that this could affect the behaviours and actions of the participants. This may have been implied in the statements provided, but the authors may consider discussing further that participants of ‘their’ study may have decided to participate and stay in the study as a counter reaction to the disbeliefs
and conflicting feelings. The participants may have decided not to show this for various reasons including material and monetary incentives.

I must state that it is a very important theory for consideration in a HIS as discussed in the paper. Well-done.

Did the study have a psychologist to assess the mental health aspect of the study participants?

**Conclusion**

The authors should consider expanding on the conclusion to state what they would suggest given what they found in the study; this may include for future research.

Thank you for the opportunity to review this interesting piece.

**References**

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**Is the work clearly and accurately presented and does it cite the current literature?**

Yes

**Is the study design appropriate and is the work technically sound?**

Yes

**Are sufficient details of methods and analysis provided to allow replication by others?**

Yes

**If applicable, is the statistical analysis and its interpretation appropriate?**

Not applicable

**Are all the source data underlying the results available to ensure full reproducibility?**

Yes

**Are the conclusions drawn adequately supported by the results?**

Partly

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

**Reviewer Expertise:** Bioethics, Ethics of HIS, Public Health, Health Research, Qualitative research methods

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.

**Author Response 22 Jul 2021**

Neema Mtunthama Toto, Malawi-Liverpool Wellcome Trust Clinical Research Programme,
Blantyre, Malawi

23rd June 2021

To: The Editors, Wellcome Open

Re: “At first I was very afraid” – A qualitative description of participants’ views and experiences in the first Human Infection Study in Malawi [Version 1; peer review: 1 approved, 1 approved with reservations]

Firstly, appreciate the encouraging words following the review of our manuscript. Second, we thank the reviewers for their significant observations and contributions to improving clarity in our findings' write-up. Below we present our point-by-point responses to the reviewers.

Reviewer's Comments

Reviewer 1

Abbreviations the authors need to be consistent in the presentation of these. For example, some have been presented in lower case, while others have been presented by Capitalizing each word.

Authors' response
Thank you very much. We have correctly presented the abbreviations.

Action taken
Please refer to the highlighted text on the abbreviations list on page 2.

Quotes Authors should re-look some of the quotes to ensure that they read clearly to reflect the theme in the discussion. The quote below was repeated on page 9 of the paper. It is presented on page 9 and repeated immediately on page 9. Please try and find another quote to replace ‘it’ on either page or, better still, delete it from page 9.
"At first, I was very afraid. Actually, I was so afraid of, okay, I that running nose, that's exactly one thing that I hate the most. I fail to study when I have a runny nose, so I hate it a lot" (CHIM 1093).

Authors' response
Thanks for that helpful observation. We have removed redundant or irrelevant quotes throughout the manuscript results section and deleted the duplicate quote.

Action taken
Please refer to the highlighted text on page 17.

Abstract/Reporting qualitative findings/Discussion Authors should avoid using the word 'likelihood', as it is used to show probability. The first sentence of the results in the abstract should be revised to sound more qualitatively appropriate and clear.
Authors' response
Thank you very much. We have removed the word "likelihood" of participation altogether.

Action taken
Please refer to the highlighted text on pages 1 and 21, respectively.

Introduction
Where the authors describe the HIS situation in LMICs, it would be imperative to note that countries like Zambia conducted a pilot HIS and have since begun conducting a full study. Furthermore, South Africa has also started conducting a HIS.

Authors' response
We have included information on Zambia and South Africa.

Action taken
Please refer to the highlighted text on page 3.

Aim of the study
From the paper, I gather that the objective of the research was to explore and describe the experiences, motivations and concerns of enrolled HIS participants. In the section of the aim of the study, authors say they aimed 'to assess acceptability' among the healthy adult volunteers who had completed the feasibility study, including their opinions on study recruitment and consent procedures, medical care and support, compensation, and community engagement. There is a need to revise this for consistency.

Authors' response
Thank you very much. While the overall objective was to describe participants' experiences, motivations, and concerns enrolled in the study. Our specific objectives explore their opinions on study recruitment and consent procedures, medical care and support, compensation, and community engagement. This was done to establish some acceptability and inform the design of future HIS studies in Malawi. We hope that this rephrasing of our statements accurately conveys our objectives, specific aims and goals.

Action taken
Please refer to the highlighted text on page 5 under the heading, Aim of the study.

Discussion,
The authors, discuss the decision process using the theory of cognitive dissonance and that this could affect the behaviours and actions of the participants. This may have been implied in the statements provided. Still, the authors may consider discussing further that participants of 'their' study may have decided to participate and stay in the study as a counter-reaction to the disbeliefs and conflicting feelings. The participants may have decided not to show this for various reasons, including material and monetary incentives. I must state that it is a very important theory for consideration in a HIS, as discussed in the paper. Well-done.

Authors' response
Thank you for your appreciation of the theory used to explain participant behaviours and actions. We have considered your suggestion on discussing the concept further.

Action taken
Please refer to the highlighted text on page 22.

Did the study have a psychologist to assess the mental health aspect of the study
authors' response
No, the study did not conduct any psychometric tests on study participants. It was not an objective or included as part of the eligibility criteria.

Action taken
N/A

Conclusion The authors should consider expanding on the conclusion to state what they would suggest given what they found in the study; this may include for future research.

Authors' response
Thank you for your suggestion. We have included a statement on what future research may need to be considered to expand our understanding of cognitive dissonance and the role that it can play in explaining participant experiences and decision-making to bring greater transparency about the acceptability of HIS in LMICs.

Action taken
Please refer to the highlighted text on page 24.

We hope that we have appropriately and adequately addressed all the reviewer comments described above and highlighted (in yellow) in the revised manuscript.

Please do not hesitate to contact me should you require further amendments.

Kind regards,

Lucinda Manda-Taylor, PhD

Competing Interests: We do not have any competing interests to disclose

Reviewer Report 17 May 2021
https://doi.org/10.21956/wellcomeopenres.18279.r43583

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Evelyne Kestelyn
Oxford University Clinical Research Unit, Ho Chi Minh City, Vietnam

Thank you for the opportunity to review this manuscript. This is clear and well written,
contributing to the growing field of knowledge about the conduct of CHIM in LMICs.

**Introduction:**
It would be beneficial to define from the start the study population not only as healthy adult volunteers but as students. I assume the literacy level but potentially also the socio-demographic status of these students might not be fully representative for the population in Malawi.

Paragraph 2, In the text: '...there are ethical issues concerning HIS, especially in LMICs' Some authors argue the ethical issues are not very different across settings. See: Kaewkungwal J, Adams P, Sattabongkot J, Lie RK, Wendler D (2019) Conducting human challenge studies in LMICs: A survey of researchers and ethics committee members in Thailand. PLoS ONE 14(10): e0223619.¹ 'results suggest that, in the view of important stakeholders in Thailand, the ethical issues raised by human challenge studies in LMICS do not differ significantly from the ethical issues raised by human challenge studies in high income countries'. I know this is contested but I would nuance this statement or provide further argumentation/references.

**Interviews:**
The interview guide was piloted on two health workers. Again I assume health workers might have a better grasp of medical terminology or research terminology than the general population. Did you consider piloting this guide with the students?

The interview guide mentions:‘How does that compare to how you feel about it now?’, "it" referring to decision making or general experiences? My main issue here is that the interviews are all exit interviews and that issues such as desirability or recall bias are not addressed. Most research shows a difference between initial worries and doubts about a study intervention and experiences after study completion where these feelings of worry fade away as the participants get more familiarized with the study staff and experience positive situations. Did you collect any data before the study start on initial feelings, questions?

Experience with study procedures and methods.
The quote about the runny nose is used twice, are there other quotes that you might use instead?

**Compensation:**
I would encourage the authors to refer to the literature around compensation for harm. The idea of compensation after study completion or to pay for insurance is interesting. Given the current debate around payment for harm versus payment for risk in CHIM, this is certainly worth elaborating on. See: Grimwade O, Savulescu J, Giubilini A, et al. Payment in challenge studies: ethics, attitudes and a new payment for risk model. Journal of Medical Ethics 2020;46:815-826.²

**Discussion:**
The discussion mentions initial fears dissipating but again this is at study end and it would have been very interesting to have initial (pre-enrolment data) versus end of study data. Regarding wider opinions; not only family members etc... could give valuable input but were any screen-failures or students who refused to participate interviewed?

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**Is the work clearly and accurately presented and does it cite the current literature?**
Yes

**Is the study design appropriate and is the work technically sound?**
Yes

**Are sufficient details of methods and analysis provided to allow replication by others?**
Yes

**If applicable, is the statistical analysis and its interpretation appropriate?**
Not applicable

**Are all the source data underlying the results available to ensure full reproducibility?**
Partly

**Are the conclusions drawn adequately supported by the results?**
Partly

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

**Reviewer Expertise:** Clinical trials, Infectious Diseases, Human challenge ethical and governance frameworks

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.

---

**Author Response 22 Jul 2021**

**Neema Mtunthama Toto**, Malawi-Liverpool Wellcome Trust Clinical Research Programme, Blantyre, Malawi

**23rd June 2021**

**To:** The Editors, Wellcome Open

**Re:** "At first I was very afraid" – A qualitative description of participants' views and experiences in the first Human Infection Study in Malawi [Version 1; peer review: 1 approved, 1 approved with reservations]
Firstly, appreciate the encouraging words following the review of our manuscript. Second, we thank the reviewers for their significant observations and contributions to improving clarity in our findings' write-up. Below we present our point-by-point responses to the reviewers.

**Reviewer's Comments**

**Reviewer 2**

**Authors' response**

**Action taken**

**Introduction:** It would be beneficial to define from the start the study population not only as healthy adult volunteers but as students.

**Authors response**

Thank you, we have added healthy adult student volunteers so that it remains clear that these characteristics formed part of the eligibility criteria and recruitment plan.

**Action taken**

Please refer to the highlighted text in the abstract and on pages 2 and 5.

I assume the literacy level, but potentially also the socio-demographic status of these students, might not be fully representative for the population in Malawi.

**Authors response**

Yes, it is correct that the literacy level is not fully representative of the population in Malawi. However, their socio-demographic status may well represent the general population of Malawi because most students enrolled in public universities in Malawi are from backgrounds that are not necessarily affluent.

**Action taken**

N/A

Paragraph 2, In the text: ‘...there are ethical issues concerning HIS, especially in LMICs’

Some authors argue the ethical issues are not very different across settings. See Kaewkungwal J, Adams P, Sattabongkot J, Lie RK, Wendler D (2019) Conducting human challenge studies in LMICs: A survey of researchers and ethics committee members in Thailand. PLoS ONE 14(10): e0223619.

‘results suggest that, in the view of important stakeholders in Thailand, the ethical issues raised by human challenge studies in LMICS do not differ significantly from the ethical issues raised by human challenge studies in high-income countries. I know this is contested, but I would nuance this statement or provide further argumentation/references.

**Authors response**

Thank you for this helpful suggestion. We have included this statement.

**Action taken**

Please refer to the highlighted text on page 3.

**Interviews:** The interview guide was piloted on two health workers. Again I assume health
workers might have a better grasp of medical terminology or research terminology than the general population. Did you consider piloting this guide with the students?

**Authors response**
Thank you for your excellent questions. We wanted to pilot the interview guide on health workers first because we wanted to avoid any risk of misconception about the study, given that this was the first study we would conduct in Malawi. We decided to test the questions on a health worker to ensure clarity and establish some relevance. While a health worker may better grasp medical terminology, we wanted to a) establish relevancy in the questions we were asking and b) we needed to obtain feedback where questions were ambiguous. Besides, our questions were broad and general enough for University students to understand once we had established clarity and relevance.

**Action taken**
N/A

The interview guide mentions: 'How does that compare to how you feel about it now?', "it" referring to decision making or general experiences? My main issue here is that the interviews are all exit interviews and that issues such as desirability or recall bias are not addressed. Most research shows a difference between initial worries and doubts about a study intervention and experiences after study completion, where these feelings of worry fade away as the participants get more familiarized with the study staff and experience positive situations. Did you collect any data before the study start on initial feelings, questions?

**Authors response**
The question, "How does that compare to how you feel about it now" was asking the participant to reflect on how they felt after joining the study to the point of exiting the study. Since the interviews took place immediately after the participant concluded all study procedures and had formally exited the trial, we believe that recall bias was not affected much. We also mention in the manuscript that as social scientists within the MARVELS consortium, we tried to be as reflexive and as neutral as possible during the data collection and analysis process by paying careful attention to our positionality. We also ensured that where participants offered conflicting views, we probed around contradictory points during questioning in an encouraging and supportive way.

**Action taken**
N/A because on page 7 of the manuscript, under the heading "Study site", we did mention how we carefully considered our position.

**Authors response**
Finally, we did not collect any data before starting the study on initial feelings and questions. This is because we felt that participants would be open enough with their opinions even at the end of the study. However, it is a valid point that you have raised. We have noted it as a possible limitation of our research to contextualize our understanding of the underlying motivation to participate in the first HIS study in Malawi.

**Action taken**
Please refer to the highlighted text on page 23.
Experience with study procedures and methods.
The quote about the runny nose is used twice, are there other quotes that you might use
instead?

**Authors response**
Thank you for noting this. We have addressed the concern by deleting the duplicate quote.
We have opted not to include another quote because other participants echoed similar
experiences.

**Action taken**
Please refer to the highlighted text on page 17.

**Compensation:** I would encourage the authors to refer to the literature around
compensation for harm. The idea of compensation after study completion or to pay for
insurance is interesting. Given the current debate around payment for harm versus
payment for risk in CHIM, this is certainly worth elaborating on. See:
Grimwade O, Savulescu J, Giubilini A, et al.
Payment in challenge studies: ethics, attitudes and a new payment for risk model. Journal of
Medical Ethics 2020;46:815-826.

**Authors response**
Thank you for pointing us to this important piece of literature. We have included the idea of
payment for risk as part of the overall discourse on compensation in HIS research.

**Action taken**
Please refer to the highlighted text on page 23.

**Discussion:** The discussion mentions initial fears dissipating but again, this is at the study
end, and it would have been very interesting to have initial (pre-enrolment data) versus the
end of study data.

**Authors response**
Thank you for your observation. Indeed, it would have been helpful to conduct pre-and
post-interviews, and future qualitative research on participant opinions about HIS should
consider this. We have taken note of this point and included it as a recommendation for
future research

**Action taken**
Please refer to the highlighted text on page 23.

Regarding broader opinions, not only family members, etc., could give valuable input, but
were any screen failures or students who refused to participate interviewed?

**References**
1. Kaewkungwal J, Adams P, Sattabongkot J, Lie RK, et al.: Conducting human challenge
   studies in LMICs: A survey of researchers and ethics committee members in Thailand. *PLoS One.* 2019; 14 (10): e0223619 PubMed Abstract | Publisher Full Text
2. Grimwade O, Savulescu J, Giubilini A, Oakley J, et al.: Payment in challenge studies: ethics,
   attitudes and a new payment for risk model. *J Med Ethics.* 46 (12): 815-826 PubMed Abstract
   Publisher Full Text
Authors response
As with any clinical trial protocol, the eligibility and exclusion criteria is clearly defined. For the feasibility study, those participants with the natural serotype identified 6B were not included in the study. We only had one participant who was excluded based on these criteria. We did not experience any refusals from students who we invited to participate in the exit interview.

Action taken
N/A

We hope that we have appropriately and adequately addressed all the reviewer comments described above and highlighted (in yellow) in the revised manuscript.

Please do not hesitate to contact me should you require further amendments.

Kind regards,

Lucinda Manda-Taylor, PhD

Competing Interests: We do not have any competing interests to disclose