The Impact of Intensive Versus Standard Anthelminthic Treatment on Allergy-related Outcomes, Helminth Infection Intensity, and Helminth-related Morbidity in Lake Victoria Fishing Communities, Uganda: Results From the LaVIISWA Cluster-randomized Trial

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Background. The prevalence of allergy-related diseases is increasing in low-income countries. Parasitic helminths, common in these settings, may be protective. We hypothesized that intensive, community-wide, anthelminthic mass drug administration (MDA) would increase allergy-related diseases, while reducing helminth-related morbidity.

Methods. In an open, cluster-randomized trial (ISRCTN47196031), we randomized 26 high-schistosomiasis-transmission fishing villages in Lake Victoria, Uganda, in a 1:1 ratio to receive community-wide intensive (quarterly single-dose praziquantel plus albendazole daily for 3 days) or standard (annual praziquantel plus 6 monthly single-dose albendazole) MDA. Primary outcomes were recent wheezing, skin prick test positivity (SPT), and allergen-specific immunoglobulin E (asIgE) after 3 years of intervention. Secondary outcomes included helminths, haemoglobin, and hepatosplenomegaly.

Results. The outcome survey comprised 3350 individuals. Intensive MDA had no effect on wheezing (risk ratio [RR] 1.11, 95% confidence interval [CI] 0.64–1.93), SPT (RR 1.10, 95% CI 0.85–1.42), or asIgE (RR 0.96, 95% CI 0.82–1.12). Intensive MDA reduced Schistosoma mansoni infection intensity: the prevalence from Kato Katz examinations of single stool samples from each patient was 23% versus 39% (RR 0.70, 95% CI 0.55–0.88), but the urine circulating cathodic antigen test remained positive in 85% participants in both trial arms. Hookworm prevalence was 8% versus 11% (RR 0.55, 95% CI 0.31–1.00). There were no differences in anemia or hepatosplenomegaly between trial arms.

Conclusions. Despite reductions in S. mansoni intensity and hookworm prevalence, intensive MDA had no effect on atopy, allergy-related diseases, or helminth-related pathology. This could be due to sustained low-intensity infections; thus, a causal link between helminths and allergy outcomes cannot be discounted. Intensive community-based MDA has a limited impact in high-schistosomiasis-transmission fishing communities, in the absence of other interventions.

Clinical Trials Registration. ISRCTN47196031.

Keywords. helminths; Schistosoma mansoni; mass drug administration; allergy-related disease; Africa.

The prevalence of allergy-related diseases (ARD), such as eczema, rhinitis, and asthma, increased rapidly in high-income countries in the twentieth century [1] and is now increasing in tropical, low-income countries (LICs) [2]. Nevertheless, populations in LICs, particularly in rural settings, remain relatively protected [3]. Understanding this phenomenon is crucial to elucidating the causes and improving prevention of ARD.

By contrast, LICs carry the largest burden of parasitic helminth infections; these are associated with some severe and much subtle morbidity [4, 5]. Major anthelminthic mass drug administration (MDA) has taken place in the last decade but, although prevention of severe helminth-induced morbidity is important, wider benefits [6] and the sustainability of helminth control by MDA [7, 8] have been questioned.

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Certain helminth antigens are highly homologous to allergens; immunoglobulin (Ig)-E and the atopic pathway are presumed to have evolved to protect mammals against such organisms [9]. Parasitic helminths must modulate such responses to survive within mammalian hosts. Animal and human epidemiological and in vitro studies indicate that, through bystander effects of such immunomodulation, chronic helminth infection protects against atopy and ARD [10]. If helminths protect against ARD, MDA programs may adversely affect these outcomes. Observational studies, many of which indicate an inverse association between helminths and ARD, are subject to confounding and reverse causation; therefore, several groups have investigated the effects of anthelminthic treatment on ARD in clinical trials. Some studies show increased atopy after anthelminthic intervention, but 2 large, school-based, individually-randomized intervention trials focusing on soil-transmitted helminths (STH) reported no effect on atopy or ARD [11, 12]. A recent household-randomized trial of intensive albendazole for STH showed no effect on ARD, but upregulated pro-inflammatory responses and reduced immunoregulatory molecules [13].

East African fishing communities bear an intense schistosomiasis burden [14]. During a *Schistosoma mansoni* infection, adult worms reside in mesenteric blood vessels and eggs are excreted through intestinal mucosa, causing intestinal and tissue (notably liver) pathology [5]. *Schistosoma* infection has shown even stronger inverse associations with atopy than STH [15] and there is evidence of increased SPT reactivity with treatment [16], but no large-scale randomized trial on the allergy-related effects of intensively treating schistosomiasis has been conducted.

We undertook the Lake Victoria Island Intervention Study on Worms and Allergy-related diseases (LaVIISWA; ISRCTN47196031) [17], a cluster-randomized trial of extended (3-year) intensive versus standard anthelmintic intervention, to assess the causal role of helminths in allergy-related outcomes and the benefits of intensive intervention for helminth-related morbidity in a schistosomiasis hot spot.

**METHODS**

**Design and Setting**

This was a 2-arm, open, cluster-randomized trial of intensive versus standard anthelmintic treatment conducted among fishing villages in the Koome islands, Lake Victoria, Uganda, between September 2012 and August 2016. The protocol has been published previously [17]. We randomized 26 villages 1:1 to either intensive or standard intervention. Village-level cluster-randomization aimed to minimize contamination from reinfection by untreated neighbours. Before the study, annual praziquantel treatment was offered to these communities, but hampered by logistics. In our baseline survey, 17% participants reported treatment in the prior year [17].

**Interventions**

Standard intervention, based on the Uganda Ministry of Health guidelines, was annual single dose praziquantel at 40 mg/kg (Cipla; CSPC OUYI Pharmaceuticals, India; AGOG Pharma, India) to community members ≥94 cm, as estimated by a height pole, plus 6 monthly single dose albendazole at 400 mg (CSPC OUYI Pharmaceuticals, India; AGOG Pharma, India; Medreich, India) to all aged ≥1 year. Intensive intervention was quarterly single dose praziquantel at 40 mg/kg to individuals ≥60 cm (to allow treatment of younger children) [18], as estimated by an extended height pole, plus quarterly triple dose albendazole (400 mg daily for 3 days) to all aged ≥1 year. Pregnant women were included in both arms, receiving single dose albendazole [19, 20].

Treatment, distributed house-to-house in collaboration with the Uganda Ministry of Health Vector Control Division, was directly observed and documented against household registers, with the exception of post–day 1 albendazole in the intensive arm.

**Participants and Surveys**

Leaders of all 27 Koome fishing villages gave written consent for their village’s participation. Allocated interventions were given to all community members (of eligible age and height) unless they were absent, sick, or refused.

Household-based surveys were conducted at baseline [21] and after 3 years of intervention. All primary and most secondary outcomes were assessed in both baseline and outcome surveys. Smaller surveys were conducted at years 1 and 2 to assess helminth trends (Supplementary Figure). Separate random household samples were selected for each survey (overlap was possible). There was no individual participant follow-up. Surveys were conducted immediately prior to respective quarterly treatments.

Household registers were updated before each survey. Villages generally comprised an intensely-populated center and a scattered periphery. Peripheral households were excluded from surveys to avoid contamination from neighboring villages, but received allocated interventions.

Baseline survey methods (previously reported) were similar to the 3-year outcome survey described below [21]. For interim surveys, stool and blood samples were collected from community members selected using a 2-stage method: 1 person was randomly selected from each of 15 randomly-selected households per village. For the 3-year survey, 70 households per village were randomly selected using a Stata program (StataCorp, College Station). In the selected households, all members ≥1 year were invited to participate. Household heads gave permission for household participation and provided the demographic details (age, sex) of all household members. Written informed consent was obtained from all adults and emancipated minors and from parents/guardians for children, with
additional assent from children ≥8 years. For each participant, a questionnaire was completed; an examination and SPT were performed; and blood, urine, and 1 stool sample were obtained. Abdominal ultrasonography was performed on children.

Outcomes
Primary outcomes were recent (last 12 months) self-reported wheezing, stratified by age (<5 years, ≥5 years); SPT positivity to mites (Dermatophagoides mix, Blomia tropicalis) and German cockroaches (Blattella germanica); and allergen-specific IgE (asIgE) to Dermatophagoides and German cockroaches (common allergens in Uganda [22]). Secondary outcomes were visible flexural dermatitis (assessed using standardized procedures); helminth infections; haemoglobin; growth (height-for-age, <20 years; weight-for-age, <11 years; and weight-for-height, <6 years, z-scores); and hepatosplenomegaly (by palpation). An additional secondary outcome, schistosomiasis-related liver and spleen morbidity assessed by abdominal ultrasonography (<18 years), was included after trial interventions commenced, when additional funding became available. Exploratory outcomes were recent urticaria and rhinitis. For logistical reasons, we could not provide infant vaccines ourselves or obtain post-immunization samples at consistent timepoints, so planned vaccine response secondary outcomes are not reported. Details on outcome ascertainment are provided (Supplementary Methods).

Randomization
At a public ceremony, 1 village was randomly selected for piloting while 26 were randomized 1:1, using restricted randomization to balance village size, prior praziquantel treatment, and distance from the sub-county health center [17] (Supplementary Methods).

Statistical Methods
For the outcome survey, we planned to sample 1540 individuals per arm (Supplementary Methods). Data were analyzed using Stata v14.0. Baseline characteristics were tabulated. Characteristics of survey participants were compared with those of non-participants by chi-squared tests. Treatment uptake was calculated both by village and treatment round as the number of people receiving treatment divided by the total number of residents.

Trial analyses were done at the cluster level. Crude and adjusted analyses (adjusting for sex, age, and the corresponding baseline summary measure of the outcome, where available) were performed. For binary outcomes, risk ratios (RRs) were calculated as the mean of the intensive-arm cluster proportions divided by the mean in the standard arm, with 95% confidence intervals (CI) calculated using a Taylor series approximation for the standard error and \( P \) values from unpaired \( t \)-tests. Where the distribution of cluster proportions was skewed, the log-cluster proportions were compared and the results were back-transformed. A 2-stage approach was used for adjusted analyses [23] (Supplementary Methods).

For continuous outcomes, intervention effects were quantified as the differences in mean outcome between trial arms, with 95% CIs calculated using the \( t \)-distribution. Non-normally distributed continuous outcomes were log-transformed and the results were back-transformed to obtain geometric mean ratios. For ordered categorical outcomes, a proportional-odds model was used.

Trial analyses were conducted in 2 populations: the primary analysis population (intention-to-treat population) included all individuals. The secondary analysis population comprised all individuals who had lived in their village throughout (or were born into their village during) the intervention period (per-protocol analysis).

Using a cluster-level approach [24], we conducted post hoc subgroup analyses by age group (<4 years, ≥4 years) for the primary outcomes to assess whether intervention effects differed among those exposed to differential anthelminthic interventions from birth.

Ethics Statement
Ethical approval was given by the Uganda Virus Research Institute (GC127), Uganda National Council for Science and Technology (HS 1183), and London School of Hygiene and Tropical Medicine (6187).

RESULTS
Participants and Intervention Uptake
Characteristics assessed in the baseline survey (October 2012–July 2013) were balanced between trial arms, with the exception that, compared to villages in the intensive arm (intensive villages), villages in the standard arm (standard villages) had fewer public toilets but contained more households with private toilets [17].

Figure 1 summarizes the treatment uptake. Both the praziquantel and albendazole uptake increased during the trial. The mean uptake per round was 63% for praziquantel and 64% for albendazole (intensive villages), compared to 56% and 73% (standard villages). In standard villages, the albendazole uptake was lower in the treatment rounds where praziquantel treatment was also given. Reported receipt of ≥1 dose of praziquantel in the preceding year was higher in intensive, compared to standard, villages (93% versus 75%, respectively). Reported receipt of ≥1 dose of albendazole was universally high (99% versus 98%, respectively).

Between September 2015 and August 2016, 70 households from each village were randomly selected for the outcome survey (Figure 2): 84 (5%) households refused to participate, 17 (1%) consented but no demographic data were captured, and 300 (17%) had no members that could be contacted. The remaining 1419 participating households contained 3566 residents aged ≥1 year. Overall, 3350 (94%) household members provided data for at least 1 primary outcome (recent wheezing...
Impact of Intensive Versus Standard Anthelminthic Treatment on Primary Outcomes

The prevalence of reported wheezing among ≥5-year-olds was 3%, with little difference between trial arms (Table 2). There were 9 individuals <5 years old who reported wheezing; no formal analysis was done for this outcome. Regarding atopy, 19% participants had a positive SPT to ≥1 allergen. Of those tested using ImmunoCAP, 54% were positive (IgE > 0.35kUa/L) for either cockroach or dust mite allergens. Enzyme-linked immunosorbent assay (ELISA) and ImmunoCAP results were positively correlated for both dust mites and cockroaches (Spearman’s correlation coefficient 0.32 and 0.29, respectively). There was no effect of intensive versus standard treatment on atopy (by SPT or IgE; Table 2). For all primary outcomes, there remained little evidence of a difference between trial arms in the per-protocol analysis (Supplementary Table 1) or among age groups (Supplementary Table 2), although RRs for SPT responses to individual allergens increased in both the per-protocol analysis and in children <4 years old.

Impact of Intensive Versus Standard Anthelminthic Treatment on Secondary and Exploratory Outcomes

The Schistosoma mansoni infection prevalence was lower in the intensive-treatment villages when assessed by stool Kato
### Table 1. Characteristics of Outcome Survey Participants

| Cluster-level Characteristics | Standard Arm (n = 13) | Intensive Arm (n = 13) |
|-------------------------------|----------------------|-----------------------|
| Mean no. of households per village (range) | 307 (124–882) | 289 (87–544) |
| Mean no. of participating households (range) | 55 (48–63) | 54 (48–64) |
| Mean no. of individuals resident in participating households (range) | 137 (69–161) | 137 (65–177) |
| Mean no. of individuals included in analysis (range) | 129 (84–150) | 129 (79–169) |
| Villages with any public toilets | 5 | 6 |
| Median no. of public toilets (range) | 0 (0–16) | 0 (0–20) |
| Median no. of private toilets (range) | 8 (0–59) | 3 (1–29) |
| Water supply other than lake | 3 | 5 |
| Piped water | 2 | 2 |
| River or open spring | 1 | 2 |
| Open well | 0 | 1 |
| Household-level characteristics (n = 714) | 881 | 857 |
| Individual-level characteristics (n = 1675) | 53% | 51% |

| Age in years, grouped |
|-----------------------|
| 0–4 | 283 | 264 |
| 5–9 | 173 | 219 |
| 10–14 | 66 | 115 |
| 15–19 | 102 | 79 |
| 20–24 | 212 | 179 |
| 25–29 | 239 | 216 |
| 30–34 | 198 | 211 |
| 35–39 | 175 | 140 |
| 40–44 | 86 | 106 |
| 45+ | 141 | 146 |

| Place of birth (mv 9, 19) |
|---------------------------|
| This fishing village | 439 | 477 |
| Other fishing village | 48 | 20 |
| Other rural village | 1021 | 1002 |
| Town | 127 | 127 |
| City | 31 | 30 |

| Maternal history of allergic diseases (mv 9, 19) |
|-----------------------------------------------|
| No history | 1193 | 1204 |
| History of asthma, eczema or allergies | 258 | 266 |
| Don’t know | 215 | 186 |

| Paternal history of allergic diseases (mv 9, 19) |
|-----------------------------------------------|
| No history | 1248 | 1244 |
| History of asthma, eczema or allergies | 145 | 155 |
| Don’t know | 273 | 257 |

| Occupation, grouped by type (mv 8, 19) |
|---------------------------------------|
| Child, not at school | 289 | 275 |
| Student | 257 | 345 |
| Housewife | 120 | 101 |
| Fishing or lake related | 564 | 467 |
| Shops, saloons, artisans, service providers | 118 | 102 |
| Bars, restaurants, food providers, entertainment | 114 | 103 |
| Agricultural, lumbering, charcoal | 157 | 201 |
| Professional | 11 | 19 |
| Unemployed | 37 | 43 |

| Malaria treatment with coartem (mv 190, 167) |
|--------------------------------------------|
| Malaria positivity by blood smear (Plasmodium falciparum; mv 213, 214) | 50 | 52 |
Katz (23% versus 39%, respectively; adjusted RR 0.70, 95% CI 0.55–0.88; Table 3) and stool polymerase chain reactions (39% versus 60%, respectively; adjusted RR 0.76, 95% CI 0.65–0.88), but urine circulating cathodic antigen positivity remained high and similar across trial arms (both 85%; Table 3), indicating that the intensive treatment was more effective than the standard treatment in reducing heavy-intensity *Schistosoma* infections, particularly in younger age groups, but had little impact on the light-infection prevalence (Figure 3). The incidence of *Schistosoma* infections was lower in both trial arms, compared to baseline, with 49% and 23% pre- and post-intervention in the intensive arm, respectively, and 56% and 39% in the standard arm, respectively. Our interim survey data suggested a greater initial reduction in intensive villages, which then plateaued, and

Table 1. Impact of Intensive Versus Standard Anthelminthic Treatment on Primary Outcomes

| Outcome | Standard Arm | Intensive Arm |
|---------|--------------|---------------|
| Wheeze (age ≥5 years)<sup>b</sup> | 44/1384 (3.2%) | 43/1392 (3.1%) |
| Wheeze (age <5 years) | 6/284 (2.1%) | 3/264 (1.1%) |

Table 2. Impact of Intensive Versus Standard Anthelminthic Treatment on Primary Outcomes

| Outcome | n/N (%)/Geometric Mean | Unadjusted | Adjusted for Outcome at Baseline by Age and Sex<sup>a</sup> |
|---------|------------------------|------------|----------------------------------------------------------|
| wheeze (age ≥5 years)<sup>b</sup> | 44/1384 (3.2%) | 1.06 (0.61–1.87) | 1.11 (0.64–1.93) |
| SPT positivity to any allergen | 273/1514 (18.0%) | 1.09 (0.83–1.44) | 1.10 (0.85–1.42) |
| Dermatophagoides | 273/1514 (18.0%) | 1.09 (0.83–1.44) | 1.10 (0.85–1.42) |
| Blomia tropicalis | 102/1514 (6.7%) | 1.26 (0.83–1.90) | 1.27 (0.85–1.91) |
| German cockroach | 156/1513 (10.3%) | 1.24 (0.87–1.77) | 1.22 (0.87–1.71) |
| Dermatophagoides or cockroach positivity (>0.35 kUa/L) | 214/390 (54.9%) | 0.97 (0.83–1.13) | 0.96 (0.82–1.12) |
| Dermatophagoides positivity (asIgE > 0.35 kUa/L) | 134/390 (34.4%) | 0.95 (0.76–1.20) | 0.96 (0.77–1.19) |
| German cockroach positivity (asIgE > 0.35 kUa/L) | 201/390 (51.5%) | 0.94 (0.79–1.11) | 0.94 (0.79–1.11) |
| Concentration of asIgE to *Dermatophagoides* (kUa/L)<sup>f</sup> | GM: 0.158 | 0.78 (0.51–1.17) | 0.77 (0.52–1.3) |
| Concentration of asIgE to *German cockroach* (kUa/L)<sup>f</sup> | GM: 0.342 | 0.82 (0.55–1.22) | 0.81 (0.55–1.20) |

Abbreviations: ART, antiretroviral therapy; HIV, human immunodeficiency virus; IQR, interquartile range; mv, missing values; no., number.
A gradual reduction in standard villages (Figure 3). The STH prevalence was relatively low. The intensive treatment reduced hookworm prevalence; no significant reductions were seen for other nematodes (among all participants; Table 3). There was no impact of the intensive versus standard treatment on anthropometric or clinical outcomes, including hepatosplenomegaly.

Table 3. Impact of Intensive Versus Standard Anthelminthic Treatment on Helminths, Clinical Outcomes, Hepatosplenomegaly by Palpation, and Anthropometry

| Outcome | Standard | Intensive | RR/ Mean Difference (95% CI) | P Value | Adjusted for Outcome at Baseline by Age and Sex | RR/ Mean Difference (95% CI) | P Value |
|---------|----------|-----------|-----------------------------|---------|---------------------------------------------|-----------------------------|---------|
| Helminth infections | | | | | | | |
| Schistosoma mansoni, stool Kato Katz | 523/1355 (38.6%) | 323/1396 (23.1%) | 0.64 (0.43–0.94) | .02 | 0.70 (0.55–0.88) | .003 |
| Schistosoma mansoni, stool PCR | 797/1353 (59.9%) | 541/1394 (38.8%) | 0.68 (0.52–0.89) | .007 | 0.76 (0.65–0.88) | .001 |
| Schistosoma mansoni, urine CCA | 1229/1444 (85.1%) | 1216/1435 (84.7%) | 0.99 (0.91–1.08) | .85 | 1.00 (0.93–1.08) | .93 |
| Hookworm, stool PCR* | 147/1353 (10.9%) | 112/1394 (8.0%) | 0.54 (0.28–1.02) | .06 | 0.55 (0.31–1.00) | .05 |
| Strongyloides stercoralis, stool PCR | 112/1353 (8.3%) | 78/1394 (5.6%) | 0.34 (0.50–1.11) | .14 | 0.78 (0.54–1.44) | .21 |
| Trichuris trichiura, stool Kato Katz | 137/1355 (10.1%) | 108/1396 (7.7%) | 0.91 (0.40–2.09) | .82 | 0.85 (0.48–1.50) | .55 |
| Ascaris lumbricoides, stool Kato Katz | 11/1355 (0.8%) | 3/1396 (0.2%) | | | | |
| Clinical outcomes | | | | | | | |
| Visible flexural dermatitis | | 1/1558 (0.1%) | 4/1553 (0.3%) | | | | |
| Haemoglobin | 14.0 | 13.9 | -0.06 (-0.37–0.25) | .70 | 0.00 (-0.24–0.25) | .97 |
| Anthropometry | | | | | | | |
| Height-for-age z-score, age 1–19 years | -0.48 | -0.49 | -0.01 (-0.20–0.19) | .95 | 0.02 (-0.16–0.20) | .83 |
| Weight-for-age z-score, age 1–10 years | -0.06 | -0.17 | -0.11 (-0.31–0.09) | .27 | -0.05 (-0.23–0.12) | .52 |
| Weight-for-height z-score, age 1–5 years | 0.15 | 0.19 | -0.09 (-0.43–0.26) | .62 | -0.06 (-0.40–0.28) | .72 |
| Hepatosplenomegaly, palpation | | | | | | | |
| Hepatomegaly, palpation | 100/1546 (6.5%) | 98/1546 (6.3%) | 0.97 (0.71–1.32) | .83 | 0.96 (0.70–1.32) | .80 |
| Splenomegaly, palpation | 87/1549 (5.6%) | 63/1547 (4.1%) | 0.73 (0.43–1.25) | .20 | 0.70 (0.43–1.15) | .13 |
| Hepatosplenomegaly, palpation* | 22/1548 (1.4%) | 14/1548 (0.9%) | 0.85 (0.52–1.39) | .49 | 0.78 (0.47–1.30) | .33 |
| Reported clinical outcomes (exploratory) | | | | | | | |
| Urticaria, last 12 months | 162/1667 (9.7%) | 172/1656 (10.4%) | 1.06 (0.86–1.30) | .59 | 1.06 (0.88–1.27) | .51 |
| Rhinitis, last 12 months | 78/1667 (4.7%) | 74/1656 (4.5%) | 1.02 (0.73–1.42) | .92 | 1.00 (0.74–1.36) | .99 |

*For this outcome, a natural log transformation was applied to village-level proportions to correct skewed distributions.

Figure 3. (A) Intensity of schistosomiasis infection in the outcome survey, by age group and trial arm, with prevalence assessed by KK examination of a single stool sample, PCR, and urine CCA. (B) Prevalence of Schistosoma mansoni infection over time (pre-intervention baseline survey, interim survey at 1 year, interim survey at 2 years, outcome survey at 3 years), by trial arm. Data are shown as the mean of village prevalences over time +/- 95% confidence intervals, assessed using KK analysis of a single stool sample (with duplicate slides) at each time point. Abbreviations: CCA, circulating cathodic antigen; KK, Kato Katz; PCR, polymerase chain reaction.
assessed either by palpation (Table 3) or ultrasound (among children; Supplementary Table 3). The per-protocol analysis did not yield any hitherto-unseen differences (Supplementary Table 4).

**Serious Adverse Events**

In total, 77,739 praziquantel treatments and 102,219 albendazole treatments were given. There were 4 serious adverse events reported (2 in each trial arm), all among adults and within 2 days of treatment: gastrointestinal symptoms leading to hospitalization (1) or requiring intravenous fluids (1); abdominal pain and vaginal bleeding in a non-pregnant woman (1); and vaginal bleeding 1 day after treatment in a pregnant woman, followed by delivery 3 days later (probably premature) and subsequent neonatal death (1). Clinic records suggested that this last woman had concurrent malaria, but this remained unconfirmed.

**DISCUSSION**

We report the first trial to address the community-level effects of intensive anthelminthic MDA in a high-Schistosoma mansoni-transmission setting. After 3 years, we found no effect of an intensive, compared to standard, intervention on allergy-related or helminth-associated disease outcomes. Intensive, compared to standard, treatment with praziquantel achieved a substantial reduction in S. mansoni intensity, most marked after 1 year, but infection remained almost universal. Intensive, compared to standard, treatment with albendazole achieved a modest reduction in hookworm prevalence, but had little impact on Trichuris or Strongyloides.

The prevalence of wheezing was lower than anticipated based on previous reports [25], limiting power for this outcome. However, the understanding of “wheeze” in our study communities was poor: there are no words for wheeze or asthma in the vernacular and asthma is rare. That said, there was no effect of the intensive intervention on wheezing, and no increase in wheezing during the intervention (5% at baseline [13], 3% after 3 years). These results provide reassurance that anthelminthic MDA is unlikely to have an immediate adverse effect on asthma among high-schistosomiasis-transmission communities, although no conclusions can be drawn on the impact of effective, universal S. mansoni removal.

SPT positivity was common. There was no increase in SPT positivity during the intervention (19% at baseline [13], 18% and 20% in the standard and intensive arms, respectively, after 3 years). There was a suggestion, especially in the per-protocol analysis and in under-4 year olds, that SPT responses increased with intensive treatment. This could be a chance finding, since a substantial number of (planned) statistical tests were conducted. This warrants more detailed investigation, as it may presage the emergence of increased atopy and ARD when helminth infections are more completely cleared. The effect of treatment may have differed based on pre-treatment infection intensity [26]. We could not assess this hypothesis, because our study was not a cohort of individual subjects.

Despite our emphasis on schistosomiasis and on long-term, community-based intervention, our results accord with previous, shorter-term trials focussing on STH [10]. However, it seems premature to conclude that high helminth prevalence has no causal link with the low ARD prevalence in LICs, given the strong effects and demonstrated mechanisms in animal models and experiments using human samples in vitro [27].

The most obvious explanation for a lack of impact on allergy-related (or helminth-associated) diseases is a failure to clear helminth infections. All villages were continuously exposed to S. mansoni-infested lake water because of a lack of alternative safe water, an involvement in fishing, and open defecation due to a scarcity of latrines. Although single- and first-dose treatments were directly observed, compliance was imperfect; albendazole uptake in the standard arm was lower in the rounds where praziquantel was given, indicating that villagers were averse to the praziquantel side effects. Furthermore, we cannot rule out the possible role of reduced drug efficacy [28, 29]. However, as a differential effect on helminth intensity was achieved, particularly for schistosomiasis, our results cast doubt on the extent to which intensity reduction (without elimination) substantially modifies the overall immunological or pathological effects in high-schistosomiasis-transmission settings.

Other factors contributing to the lack of impact on allergy-related outcomes may include the long-term immunological effects of helminth exposure through a persistence of antigen or through epigenetic changes in immunological pathways [30]. Also, in tropical, low-income settings, numerous other exposures—including immunomodulating infections such as malaria, exposure to dirt and domestic animals, or the microbiome profile—may impact allergy-related outcomes, such that modifying helminth exposure alone may have a limited impact [31].

A recent meta-analysis examined the effects of schistosomiasis treatments on related morbidity [32]. The results indicated wide-ranging benefits, with an increased impact when egg reduction rates were greatest and, for anaemia and chronic morbidities, when treatments were repeated over periods of greater than 24 months. Thus, we were disappointed that, despite a differential reduction in schistosome intensity, we found no evidence that 3 years of intensive (compared to standard) intervention achieved improvement in any morbidity measure. This adds to the evidence base showing the limited effects of MDA on such outcomes at the community-level. We identified surprisingly little severe Schistosoma-related morbidity in this community, despite intense infections, consistent with earlier work from Lake Victoria communities; it is possible that intensive interventions would have a greater benefit in settings (such as Lake Albert) where severe pathology is more common [33].
Our experience emphasizes that MDA may struggle to eliminate helminths as a public health problem, especially in high-transmission environments. The substantial decline in *S. mansoni* infections (as measured by Kato Katz) achieved in year 1 led us to hope that intensive intervention could make an important contribution to schistosomiasis control in these challenging hot spots. The subsequent plateau and persistent infection (as measured by circulating cathodic antigen) were disheartening. This phenomenon (a large drop in prevalence, followed by a subsequent plateau) has also been reported in Kenyan districts bordering Lake Victoria [34]. Besides reinfection, the possibility of selection for praziquantel-resistant or -tolerant strains is of concern [35]. A radically different approach is needed, with complementary interventions, including improved water supplies and sanitation, behavior changes, and vector controls, as well as an effective vaccine against schistosomiasis [36].

Observational analyses addressing the effects of helminths remain limited by confounding by poverty and environment. Our strategy aimed to pinpoint helminth effects by randomizing their treatment, but was constrained by difficulties in achieving removal. Trials designed so that helminths are cleared in settings where reinfection can be avoided, and with substantial follow-up, are needed for a full understanding of the risks and benefits of deworming.

**Supplementary Data**

Supplementary materials are available at *Clinical Infectious Diseases* online. Consisting of data provided by the authors to benefit the reader, the posted materials are not copyedited and are the sole responsibility of the authors, so questions or comments should be addressed to the corresponding author.

**Notes**

The LaVISSWA trial project leaders, physicians, and post-doctoral scientists are Richard Sanya, Margaret Nampijja, Harriet Mpairwe, Geraldine O’Hara, and Barbara Nereumia; the statisticians and data managers are Emily Webb, Remy Hoek Spaans, Lawrence Musungu, Lawrence Lubayi, Helen Akurut, Fatuma Nalukenga, Beatrice Mirembe, Justin Okello, Sebastian Owilla, Jonathan Levin, and Stephen Nash; the clinical officers are Milly Namutebi and Christopher Ziwa; the nurses are Esther Nakazibwe, Josephine Tumusiime, Caroline Ninsiima, Susan Amongi, Grace Kamukama, Susan Iwala, and Florence Akello; the internal monitor is Mirrion Akelo; the field workers are Robert Kizindo, Moses Sewankambo, and Denis Nsubuga; the laboratory staff and collaborators are Stephen Cosse, Linda Wammes, Prossy Kabuubi Nakawungu, Emmanuel Niwagaba, Gloria Oduru, Grace Kabani, Elson Ayabo, Eric Ssebagala, Fred Muwonge Kakooza, Joyce Kabagama, Gayiira Nkurungi, Angela Nabwoga, Denisson Kizito, John Vianney Tushabe, Jacent Nassuna, Jaco Verweij, Serge Versteeg, and Ronald van Ree; the social scientist is Edward Nalwoga, Dennison Kizito, John Vianney Tushabe, Jacent Nassuna, Jaco Verweij, Serge Versteeg, and Ronald van Ree; the internal monitor is Mirrion Akelo; the field workers are Robert Kizindo, Moses Sewankambo, and Denis Nsubuga; the laboratory staff and collaborators are Stephen Cosse, Linda Wammes, Prossy Kabuubi Nakawungu, Emmanuel Niwagaba, Gloria Oduru, Grace Kabani, Elson Ayabo, Eric Ssebagala, Fred Muwonge Kakooza, Joyce Kabagama, Gayiira Nkurungi, Angela Nabwoga, Denisson Kizito, John Vianney Tushabe, Jacent Nassuna, Jaco Verweij, Serge Versteeg, and Ronald van Ree; the scientific officer is Elly Tumushabe and the clerk for Koome sub-county (Asuman Muwumuza), who are members of the Trial Steering Committee, for their support. They also thank the other members of the Trial Steering Committee: Heiner Grosskurth (chair), Edridah Tukahebwa, Narcis Kabaterine, Neil Pearce, Anatoli Kamali, and Monica Kuteesa. They thank the Makerere University Joint AIDS Programme and the Calvary Chapel Island Mission for providing voluntary counselling and human immunodeficiency virus testing for community members, including study participants, in collaboration with their research program. They thank colleagues, especially Richard Hayes, Narcis Kabaterine, and Heiner Grosskurth, for their comments on the manuscript.

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