INTRODUCTION

Intestinal parasitic infections have a significant impact on health outcomes and morbidity in adults and children worldwide, ranging from diarrhea and stunting in children to impaired cognitive development from iron deficiency anemia.1 Globally, approximately 430 million people are infected with hookworm (Ancylostoma duodenale and Necator americanus).2 Among those infected with intestinal nematodes, hookworm infections are associated with the greatest years lived with disability (YLDs), with recent estimates indicating that human hookworm disease is associated with 4.1 million disability-adjusted life years.3 This large contribution of hookworm infections to the YLDs among those with nematode infections is due to the link between hookworm disease and anemia in children.4-6 Beyond its health impact, the anemia it induces is linked to a moderate economic burden ranging up to $139 billion annually.3 Necator americanus is the major hookworm species in the Americas. Transmission requires fecal contamination of soil and dermal penetration of human hosts. Conditions for larval survival, include moist and temperate environments. These soil-transmitted helminths are mainly found in areas where sanitation and hygiene are poor.7 most commonly in resource-limited countries. However, in the early 1930s, the southern United States had a high prevalence of hookworm infections caused by N. americanus. The initial surveys found 53.6% of the population to be infected with hookworm; in some areas, the prevalence rose as high as 76%.8 To control the disease, thousands of individuals were treated, decreasing prevalence to 39%.9 After these interventions, there were increases in school enrolment, attendance, and literacy, and those within the treated cohort had substantial gains in long-term incomes.10 However, because of posttreatment reinfection and widespread transmission, hookworm infection and disease continued to persist in the southern United States, especially in areas of extreme poverty.11 According to a study in the 1950s, rural Alabama still suffered from a high prevalence of hookworm infection in schoolchildren, with some counties having 60% infection.12 With improved sanitation and waste disposal infrastructure, in association with aggressive economic development in the southern United States, the prevalence of hookworm infection decreased. In the 1990s, surveillance studies for enteropathogens in southern Alabama show a 30% prevalence for all soil-transmitted helminths combined, including Ascaris lumbricoides, N. americanus, and Enterobius vermicularis.13 This part of the United States was identified as one at high risk for intestinal helmint infections.14 A more recent systematic review, however, found that few surveys for intestinal helmint infections have been conducted in recent decades, with limited information about these diseases, especially in poor rural and southern United States.

According to the Alabama Center for Rural Enterprise (ACRE), an organization that addresses poverty and economic development in one of the poorest areas of the nation,16 there continue to be residences without adequate sanitation systems, increasing exposure to open sewage near dwellings. The “Black-Belt” soil native to this area is composed of a firm sedimentary limestone bed overlain with a layer of dark, rich soils,17 which requires expensive septic systems for proper waste disposal. In Lowndes County, Alabama, where the per capita income is $18,046, and 31.4% of the population lives below the poverty line,18 sanitation systems are unaffordable. For rural, impoverished individuals, the main form of waste removal involves use of “straight piping,” a method involving a series of ditches or crudely constructed piping systems to...
guide human waste away from the residence. Most pipes never reach more than 10 meters in length, and during rainstorms or flooding, the residents report visible stool entering their homes (reported by ACRE, unpublished data).

While the prevalence of intestinal parasites has decreased in the past 85 years, conditions amenable to the parasite life-cycle and transmission persist. Because of these conditions, a more current investigation using modern molecular diagnostics is warranted to determine the burden of intestinal parasitic infection. The goal of this study seeks to address the current prevalence of helminthic and protozoan infections and determines correlation with sewage exposure in an economically-challenged region of the southern United States. Given continued poverty in these areas such as Lowndes County, lack of access to contained septic systems, and potential exposure to raw sewage, intestinal helminth infection likely persists despite previous interventions. The results of this study will provide insight into disease burden within this community and a basis for further study on possible risk factors, intervention, and effects on the community.

**METHODS**

**Sample population.** The study was performed in Lowndes County, an area in Alabama known to have low per capita income. According to the 2010 US Census, Lowndes County, has 5,270 houses with a 11,299 total population. ACRE reports approximately 50% of households have a failing or no sewage system (ACRE, unpublished data). Dwellings and individuals were preselected by ACRE organizers from areas with poor sanitation and used open sewage systems as a means of waste management. Enrolment goals of 100 subjects were planned, all subjects enrolled were African American. A total of 66 subjects were enrolled in the study from a total of 24 homes. There were single individuals from seven homes, four homes with two subjects, seven homes with three subjects, three homes with four subjects, one home each with five, six, and eight subjects, respectively. The research team and those assisting from ACRE visited the selected dwellings, interviewed residents using a questionnaire to obtain data regarding their dwelling and health, and collected soil from the surrounding areas and stool samples from participants. The questionnaire included information regarding sex, age, previous travel outside the United States, prior history of parasitic infections, exposure to sewage within the home, asthma, and diarrheal symptoms. Before participating in the study, each participant or guardian signed a written consent form. Participants were excluded if they have ever traveled outside the United States or were under 2 years of age, pregnant, HIV-positive, or immunocompromised. This study was approved by the Baylor College of Medicine Institutional Review Board and the Alabama Department of Public Health.

**Sample collection.** Stool samples were self-collected by the individual participants after instructions were provided verbally and in writing, and soil samples were obtained near areas of sewage collections or run-off by research team members. Participants were instructed to fill stool containers with approximately 5 grams of stool and store in provided opaque double-sealed biohazard bags in the refrigerator overnight, or if possible, the same day as the scheduled interview and collection. All samples were stored in dry ice coolers after collection and transported to the Laboratory of Clinical Parasitology and Diagnostics, Baylor College of Medicine within 5 days of collection, and stored in −20°C freezers.

**Molecular methods.** DNA was extracted from the frozen stool samples the following week after collection using MP FastPrep® spin kits for soil (MP Biomedicals, Santa Ana, CA) after a modified method was developed by the primary investigator, as previously described, for *Trichuris trichiura* DNA. Parasite DNA was detected using a multiparallel quantitative real-time PCR (qPCR) protocol. Species-specific primers and 6-carboxyfluorescein-labeled Minor groove binder probes (Applied Biosystems, Foster City, CA) were selected for each of the eight parasites tested, including helminths (*Ascaris lumbricoides*, *Trichuris trichiura*, *Strongyloides stercoralis*, *Ancylostoma duodenale*, and *N. americanus*), and protozoa (*Giardia lamblia*, *Cryptosporidium* species, and *Entamoeba histolytica*) (Supplemental Table 1). Samples were analyzed on an ABI ViA 7 Real-Time PCR System (Applied Biosystems) using default parameters for fast chemistry and 40 cycles. DNA concentrations were calculated using parasite plasmid standard curves. All controls (positive, negative, and a pBR322 plasmid internal control) were performed in triplicate with subject samples in duplicate. Samples were repeated for discordant results. The threshold for positivity was set at 38 cycle thresholds (Ct), which was the limit of detection for our dynamic range of positive standard curves. Fifty grams of top soil was collected at sites close to the opening of “straight pipes,” and the same protocol used for stool was implemented for extracting DNA from soil samples.

**Additional analysis.** A subsequent serological evaluation using an enzyme-linked immunosorbent assay (ELISA) protocol for the detection of *Toxocara* species was performed on 11 patients who were found to be positive for *N. americanus* and had serum available for study. Because of logistical restraints, blood from only those positive for any parasite was requested, and only 11 subjects consented. A commercially available *Toxocara* ELISA was used per manufacturer protocol (Abnova, Taiwan) with positive absorbance readings accepted as greater than or equal to 0.3 optical density units. For the detection of *Strongyloides stercoralis*, a previously described NIE-ELISA protocol was implemented using serum from the same 11 participants who were tested for *Toxocara* species. Unfortunately, subject’s stool positive for *Strongyloides* by qPCR declined further serum work up. Any positive laboratory result (qPCR, microscopy, ELISA) was communicated to the subject and their primary care provider, who decided what clinical course to take.

Gel electrophoresis with 4% agarose gel was also used in this study as a confirmatory tool for positive results. Sample standards were used, as well as positive and negative controls. Bands were visualized by ultraviolet fluorescent detection. Those testing positive for *N. americanus* infection had their stool sent to the Centers for Disease Control (CDC) for ova and parasites, and concentration microscopic evaluation.

**Data analysis.** All statistical analyses were performed using Prism software version 5.0d (GraphPad, La Jolla, CA). The level of significance was set at *P* < 0.05 for all analyses, and all statistical tests were two-sided. Descriptive statistics were computed to describe data for sample characteristics. Frequency distribution and measures of central tendency and dispersion were expressed by geometric means, medians, standard deviations, and proportions. Fisher’s exact test was
used to compare categorical variables and proportions. All qPCR positive samples were reported as parasite DNA concentrations measured in femtograms per microliter (fg/μL). Data were generated by detecting which parasites were present in each person. Linear regression was used to calculate fg/μL to hookworm eggs per gram (epg) of stool by microscopy in a previous study, which provides a good representation of parasitic infections within a population.\(^{21}\)

RESULTS

The questionnaires were collected for 67 individuals, but given exclusion criteria, only 66 were included in this study. Not all questions were answered, therefore, the questionnaire results were evaluated separately in a subgroup analysis (Figure 1). Age ranged from 2 to 85 years, with a mean of 34.2 years of age. The participants included 36 females and 25 males. No one reported a previous diagnosis of a parasitic infection. Only six individuals reported having exposure to raw sewage inside their domicile (42.4% of those answering the questionnaire within this study population). Twenty-eight individuals reported having exposure to raw sewage inside their domicile (42.4% of those answering the questionnaire within this study population).

Stool samples were collected for 55 individuals. Of these, 19 (34.5%) were positive for \(N\). \(americanus\), four (7.3%) for \(Strongyloides stercoralis\), and one (1.8%) for \(Entamoeba histolytica\) (Figure 2). Polyparasitism was identified in two stool samples, both of which were positive for \(N\). \(americanus\) and \(Strongyloides stercoralis\). All 55 samples were negative for \(Giardia lamblia\), \(Cryptosporidium\) species, \(Ascaris lumbricoides\), and \(Trichinella triichiura\). Stool samples positive for \(N\). \(americanus\) contained low levels of parasite DNA (range of 0.013–0.059 fg/μL), and the estimated epg, based on calculations from a previous study,\(^{21}\) ranged from 0.474 to 2.14 epg of stool. Microscopy performed on qPCR positive stool samples were negative for ova or parasites. The 19 \(N\). \(americanus\) positive samples detected by qPCR were run on 4% agarose gel electrophoresis with sample standards, and all 19 samples had bands corresponding to the appropriate size using a DNA ladder (New England Biolabs, Ipswich, MA) (Supplemental Figure 1). None of the hookworm samples were positive for \(Ancylostoma duodenale\). Eleven of the 23 participants with stool samples positive for any parasite consented to further serology assays. Of these 11 participants, one (5.2%) tested positive for \(Toxocara\) species. None tested positive for \(Strongyloides stercoralis\) IgG antibodies based on ELISA. Of the 34 soil samples collected, one (2.9%) tested positive for the protozoa, \(Cryptosporidium\) species.

Further subanalysis was performed on 26 individuals with fully completed questionnaires and qPCR stool testing. Those meeting these criteria included 10 males and 16 females. Stool samples tested negative for 15 patients, \(N\). \(americanus\) positive for 11 patients, and \(N\). \(americanus\) and \(Strongyloides stercoralis\) coinfection positive for two patients. Females comprised 61.5% of this subgroup analysis. There was no statistically significant difference between males and females with or without \(N\). \(americanus\) infection \((P = 0.68)\) (Table 1). Raw sewage was reported inside the home for 73.1% of individuals included within this subanalysis, but there was no statistically significant difference between those testing positive and negative for \(Necator americanus\) \((P = 0.61)\). Three individuals reported a diagnosis of asthma, and one reported an episode of diarrhea within the past 7 days, but were not statistically different between the uninfected and those infected with \(N\). \(americanus\) \((P \geq 0.99)\) (Table 1).

DISCUSSION

Using field-tested molecular assays, 34.5% of individuals living in a high-risk environment within a region of the southern United States were found to have stool samples testing positive for \(N\). \(americanus\) (hookworm). Even though the parasite burden was low, with up to 0.059 fg/μL of parasite DNA and an estimated egg burden up to 2.1 epg of stool, there is evidence of endemic and autochthonous infection within this population. In addition, 73% of Lowndes County residents who were tested and completed the questionnaire reported exposure to raw sewage inside their home, and even though not statistically significant (likely due to small sample size), 72.7% with this exposure tested positive for parasitic infection. By using qPCR, gastrointestinal parasites known to be endemic to developing countries have now been identified...
in a resource-limited county in Alabama, among people who have never traveled abroad, likely from autochthonous infection.

Despite decreased hookworm prevalence reported by previous soil helminth eradication programs, infection is still present within this population. During the last century, repeat surveys were performed to track the improvement of helminth infections among local populations, but these studies used microscopy as the diagnostic modality, which is limited by time requirements, manpower, and need for skilled microscopists. Even among those with ample training and experience within the field of microscopy, sensitivity for ova and parasite detection is 50–85%. In addition, detection of ova and parasites is severely limited by the burden of disease—those with less parasite burden are less likely to test positive via microscopy. Given these limitations, this study was unable to provide a correlation between the detection of parasite DNA and the microscopy analysis of qPCR positive stool samples. This was most likely due to the low N. americanus egg burden (calculated 1–2 epg of stool), which is below the microscopy limits of detection (12 epg). These limitations further justify the utility of qPCR as a method for detection of intestinal parasite infections within these resource-limited communities, where the burden of disease might be too low to accurately detect via microscopy. A new method for parasite detection could provide a more sensitive diagnostic approach for those within these communities with a continued risk of low hookworm burden and other parasitic infections. These constraints explain why qPCR is a desirable method to use with the potential of multiplexing samples for varied infections. Moreover, qPCR requires less skilled operators, and has a significantly higher sensitivity for detection of parasite DNA, especially those with a low burden of disease, detecting levels as low as 100 attograms [1 × 10^-18] of DNA. The limit of detection for qPCR and its specificity effectively rule out A. duodenale hookworm DNA in the samples, together with a historical record showing an absence of A. duodenale infections in the southern United States. Together with the positive gel bands (Supplemental Figure 1), strengthens the likelihood that the positive DNA signals for N. americanus hookworm in this study do not reflect false positives and that transmission continues in the modern era.

As shown in the 1950s Alabama study, hookworm infection rates were as high as 60% in some of the more poverty-stricken communities. In association with overall economic development in the years during and after Franklin Delano Roosevelt’s New Deal legislation, together with improvements in sanitation, urbanization, and industrialization, these pockets of infection were thought to have resolved. However, given continued poor sanitation and advancement of detection methods (improved sensitivity with PCR compared with microscopy), low burdens of infection have now been discovered in the United States among populations with autochthonous transmission. Further testing is necessary as hookworm continues to be a problem in areas with poor sanitation, allowing recurrent infections due to repeat exposure. Other areas in the United States have also been found to harbor significant pockets of parasitic infections thought only endemic to developing countries. Strongyloides stercoralis has also been found in some Appalachian mountain communities, as well as in Kentucky and rural Tennessee. Interestingly, we detected a 5.2% Toxocara IgG serological prevalence, which coincides with predicted national prevalence for this parasite among African Americans. While the soil sampling did not reveal the presence of helminths, this was likely because of the limitations of random sampling; however a single Cryptosporidium species sample was detected and reinforces the perception that human or animal fecal contamination occurs near these houses. Another limitation of soil was that no concentration steps were performed, only 50 mg of soil per 50 gm soil sample was processed for DNA extraction. This small amount of soil and low sample numbers likely contributed to low-parasite positive samples. The discovery of these parasitic diseases within the United States begins to shift the idea behind global health. One concept is blue marble health, which reveals that many of the world’s neglected tropical diseases are paradoxically found in some of the wealthiest countries, especially in these regions of extreme poverty. With the introduction of more advanced diagnostic techniques, emergence of rare, endemic infections may eventually become less defined by geographic location, but more by economic status.

Unfortunately, because of the mistrust stemming from the illegals of the self-constructed “straight pipe” waste

![](image.png)

**Figure 2.** Parasite DNA fg/µL in positive stool samples from rural Alabama.

### Table 1

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Uninfected [N = 15 (%)*]</th>
<th>Necator only infection [N = 9 (%)*]</th>
<th>Any infection [N = 11 (%)*]</th>
<th>P value†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex (N = 26)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male 10 (38.5%)</td>
<td>5 (33.3%)</td>
<td>4 (44.4%)</td>
<td>5 (45.5%)</td>
<td>0.679</td>
</tr>
<tr>
<td>Female 16 (61.5%)</td>
<td>10 (86.7%)</td>
<td>5 (55.6%)</td>
<td>6 (54.5%)</td>
<td></td>
</tr>
<tr>
<td>Raw sewage in home</td>
<td>11 (73.3%)</td>
<td>8 (88.9%)</td>
<td>8 (72.7%)</td>
<td>0.615</td>
</tr>
<tr>
<td>Asthma or other lung disease</td>
<td>3 (20.0%)</td>
<td>2 (22.2%)</td>
<td>2 (18.2%)</td>
<td>&gt; 0.999</td>
</tr>
<tr>
<td>Diarrhea in past week</td>
<td>1 (6.7%)</td>
<td>1 (11.1%)</td>
<td>1 (9.1%)</td>
<td>&gt; 0.999</td>
</tr>
</tbody>
</table>

*Percentage of individuals within each infection group.
†P values derived from Fisher’s exact test for association between uninfected and Necator infection. All P values were above 0.05 comparing uninfected to any infection groups.
disposal systems, as well as toward the medical community, the number of individuals included in this study was much smaller than expected. By working with the ACRE organization, which has fostered trust and worked with several members within the community, the research team was able to include more participants, but some of the data obtained from the questionnaires were incomplete because of different individuals performing the interviews. This should be considered when interpreting the results of this study. Performing subgroup analyses with the information provided some baseline data that could be assessed and compared among those being tested for infection. These incomplete forms were still included in data analyses given the difficulty of obtaining this information by other means and to provide further insight into the local population.

This preliminary study demonstrates that gastrointestinal parasites are present in > 30% of this at-risk population in Lowndes County, Alabama. Further testing in this community is vital to better comprehend the parasitic burden in this population, and additional studies should be initiated to further understand the implications and effects on health and the community. Parasitic testing needs to be expanded to include more households to determine the prevalence and quantitative parasitic burden. Hemoglobin monitoring should be performed among those testing positive for infection, against a negative control group, to determine if there is an impact on health with low parasitic burdens or a correlation between parasitic burden and hemoglobin levels. Hookworm (*N. americanus*) has also been shown to impair immune recovery (CD4+ cells) in HIV-infected individuals by an average of 102 cells/mL fewer CD4+ cells in those infected with *Necator.* Interestingly, the participants of this study also had low *Necator* burden of infection (0.025 fg/mL, 0.72 epg) and similar to the Alabama findings. According to 2014 data, the rates of HIV infection in Lowndes county is 758 per 100,000 African Americans and is an area with some of the highest rates of HIV infections in the United States. The combination of HIV and subclinical hookworm infections in Alabama could have an impact on immune recovery. Also, previous studies found a correlation between treatment and substantial gains in long-term income, as well as improvement of school enrollment, attendance, and literacy after hookworm eradication programs. By monitoring for changes in median income, attendance, and literacy after hookworm eradication, found a correlation between treatment and substantial gains in health outcomes, with the overall goal of eradicating this neglected tropical disease in the United States and worldwide. 

*INTESTINAL PARASITES AND POOR SANITATION IN ALABAMA 1627*

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Authors’ addresses: Megan L. McKenna, Shannan McAtee, Patricia E. Bryan, Rebecca Jeun, Tabitha Ward, Jacob Kraus, Maria E. Bottazzi, Peter J. Hotez, and Rojelio Mejia, National School of Tropical Medicine, Baylor College of Medicine, Houston, TX, E-mails: mmckenna@bcm.edu, shannon.mcatee@gmail.com, patricia.e.bryan@uth.tmc.edu, rjeun1@jhmi.edu, tabitha.ward@bcm.edu, jckraus@gmail.com, bottazzi@bcm.edu, hotez@bcm.edu, rojelio.mejia@bcm.edu. Catherine C. Flowers, Alabama Center for Rural Enterprise, Montgomery, AL, E-mail: cathnerflowers@aol.com.

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