Neurocysticercosis (NCC) is a disease caused by central nervous system infection by the larval stage of the pork tapeworm, *Taenia solium*. In developing countries, NCC is a leading cause of adult-onset epilepsy. Case reports of NCC are increasing among refugees resettled to the United States and other nations, but the underlying prevalence among refugee groups is unknown. We tested stored serum samples from the Centers for Disease Control and Prevention Migrant Serum Bank for antibodies against *T. solium* cysts by using the enzyme-linked immunoelectrotransfer blot. Seroprevalence was high among all 4 populations tested: refugees from Burma (23.2%), Lao People’s Democratic Republic (18.3%), Bhutan (22.8%), and Burundi (25.8%). Clinicians caring for refugee populations should suspect NCC in patients with seizure, chronic headache, or unexplained neurologic manifestations. Improved understanding of the prevalence of epilepsy and other associated diseases among refugees could guide recommendations for their evaluation and treatment before, during, and after resettlement.

Cysticercosis is a disease caused by infection with the larval stage of the pork tapeworm, *Taenia solium*. Humans and pigs acquire cysticercosis by ingesting *T. solium* eggs shed in the feces of humans with taeniasis (i.e., infected with an adult intestinal tapeworm). Upon ingestion, tapeworm eggs release oncospheres, which invade the intestinal wall and disseminate through the bloodstream to form cysts throughout the body. The natural lifecycle of *T. solium* tapeworms completes when a human eats pork contaminated by *T. solium* larval cysts because these can then develop into adult egg-producing intestinal tapeworms. This endemic lifecycle occurs primarily in regions where sanitation is poor and where pigs are allowed to roam and access raw human sewage.

Neurocysticercosis (NCC) occurs when cysts develop within the central nervous system (CNS); NCC is the primary cause of illness in *T. solium* infection. The clinical features of NCC cover a diverse range of neurologic manifestations, including seizures, headache, intracranial hypertension, hydrocephalus, encephalitis, stroke, cognitive impairment, and psychiatric disturbances (1,2). In areas in which *T. solium* infection is endemic, it is a major cause of epilepsy, with 30% of seizure disorder attributable to NCC (3–5).

Numerous reports document that cysticercosis in the United States occurs primarily among migrants and travelers who are presumed to have acquired their infection in another country (6–9). Refugees represent a large group of migrants in which the frequency of *T. solium* infection has not been described. Approximately 690,000 refugees resettled in the United States during 2000–2010 (10). Resettlement from regions with known pockets of *T. solium* tapeworm endemicity, including Southeast Asia, central Asia, and sub-Saharan Africa, is common. Cysticercosis among resettled refugees has been reported, but the underlying prevalence in refugee populations is unknown.

Seroprevalence of Antibodies against *Taenia solium* Cysticerci among Refugees Resettled in United States

Seth E. O’Neal, John M. Townes, Patricia P. Wilkins, John C. Noh, Deborah Lee, Silvia Rodriguez, Hector H. Garcia, and William M. Stauffer
Understanding the prevalence of *T. solium* infection could guide recommendations on evaluating and treating refugees before, during, and after resettlement.

During 2010, we used the classic enzyme-linked immunoelectrotransfer blot for lentil-lectin purified glycoprotein (EITB LLGP) to measure the seroprevalence of antibodies against *T. solium* cysts among several refugee populations resettled to the United States in previous years. We present the results, discuss clinical and public health implications, and suggest topics for further research.

**Methods**

**Study Populations**

Refugees who apply for resettlement to the United States are required to undergo a predeparture medical screening examination that includes collection of a peripheral blood sample from persons ≥15 years of age. The Migrant Serum Bank, established by the Division of Global Migration and Quarantine at the Centers for Disease Control and Prevention (CDC, Atlanta, GA, USA) in 2002, retains a convenience sample of de-identified serum samples from these examinations. Each sample has associated demographic information, including refugee group, age, birth country, refugee camp, and site and date of specimen collection. At the time of this study, ≈31,000 serum samples were available that represented resettled refugee populations from the Middle East, Southeast Asia, and Africa. We identified refugee populations represented in the Migrant Serum Bank in which cases of human cysticercosis or NCC have been reported in the countries of origin (16). We then randomly selected serum samples from each of these identified populations to test by EITB LLGP for antibodies against *T. solium* cysts. Populations with limited numbers of samples were excluded because lack of statistical power could impede prevalence estimations. Our final sample comprised 2,001 serum samples from resettled refugees from Laos, Burma (renamed Myanmar in 1989), Bhutan, and Burundi (Figure 1). The institutional review boards at CDC and at Oregon Health & Science University reviewed and approved this study.

**Laboratory Methods**

Individual 100-μL aliquots of each sample were separated at the CDC Central Repository, stored in microtubes, and shipped on dry ice to the CNS Parasitic Diseases Research Unit, Universidad Peruana Cayetano Heredia (Lima, Peru), for processing. Serum samples were analyzed by EITB for the presence of antibodies against *T. solium* cysts (EITB LLGP) as described (17). The EITB LLGP uses a semipurified fraction of homogenized *T. solium* cysts containing 7 *T. solium* glycoprotein antigens named after the Kda molecular weights of the corresponding reactive bands (GP50, GP42, GP24, GP21, GP18, GP14, GP13). Reaction to any of these 7 glycoprotein antigens is considered positive. When applied in community settings, the EITB LLGP provides an estimate of population exposure to *T. solium* cyst antigens. A positive EITB LLGP result alone does not definitely establish active infection.

Figure 1. Geographic location and background of refugee populations sampled for antibodies against *Taenia solium* cysticerci by using the classic enzyme-linked immunoelectrotransfer blot for lentil-lectin purified glycoprotein. Countries of origin are shaded dark grey (Burundi, Bhutan, Burma [Myanmar], Laos). Host countries are shaded light grey (Tanzania, Nepal, Thailand). Burundi: ≈14,000 Burundian refugees who lived in camps in Tanzania since 1972 were resettled during 2006–2008. Resettled refugees were primarily ethnic Hutu. Bhutan: ethnic Lhotshampa Bhutanese refugees arrived in Nepal ≈1990. Resettlement began in 2008 and is ongoing, with ≈40,000 resettled thus far. Burma: there has been intermittent influx of refugees into Thailand from Burma since 1984. Resettlement began in 2004 and is ongoing, with ≈90,000 resettled thus far. Resettled refugees in this group are primarily ethnic Karen and Karenni. Laos: refugees from Laos arrived in Thailand as early as 1975, and many resettled soon thereafter. The most recent round of resettlement from the Wat Tham Krabok camp occurred during 2004–2006 with resettlement of ≈16,000 ethnic Hmong refugees.
because antibodies can persist even after parasite clearance. The clinical significance of specific glycoprotein bands or combinations of bands in community studies has not been described. Although a highly sensitive and specific EITB is available to detect serum antibodies against adult *T. solium* intestinal infection, the unknown duration of antibody persistence after parasite clearance and the large sample size required for reasonable confidence intervals precluded our use of this assay in this study (18).

**Data Analysis**

Data were analyzed by using Stata version 10 (StataCorp LP, College Station, TX, USA). Direct standardization was used to facilitate comparison across refugee groups, with age–sex standardized seroprevalence calculated as the weighted average of stratum-specific seroprevalence. Continuous variables were assessed by using Kruskal-Wallis for differences among groups of interest. Pearson χ² and Fisher exact tests were used to compare distributions of proportions or to examine association between pairs of categoric measures. Logistic regression models were constructed for each refugee population to calculate odds ratios for seropositivity among strata (refugee camp or birth country) while controlling for age and sex. All tests are 2-sided, and significance was set at 0.05.

**Results**

A total of 2,001 samples were distributed approximately equally among refugees from Burma (499 [24.9%] refugees), Laos (502 [25.1%]), Bhutan (500 [25.0%]), and Burundi (500 [25.0%]). The median age of refugees sampled was 26 years (interquartile range 20–40 years, range 15–99 years). No significant difference existed between the proportions of samples from male (984 [49.2%]) and female (1,017 [50.8%]) refugees (p = 0.30). Of the 2,001 samples, 22.5% (95% CI 20.7%–24.4%) were EITB LLGP–positive for antibodies against *T. solium* cysts.

The aggregate seroprevalence was statistically homogenous across categories of age and sex. However, within individual refugee groups, seroprevalence differed across strata of age (Figure 2) and sex (Table 1). Male refugees from Burma were 2× more likely than female refugees from Burma to be seropositive (odds ratio [OR] 2.0, 95% CI 1.3–3.1). This association between male sex and positive serologic test results was not present in the other refugee groups. The proportion of seropositive results also varied by age category in refugees from Laos (p = 0.04) and Bhutan (p = 0.12).

Refugees from Burundi were significantly younger than those from the other countries (p<0.01), but Burma had a higher proportion of male refugees (p = 0.04) (Table 2). The crude seroprevalence (25.8%, 95% CI 22.0–29.6) and age–sex standardized seroprevalence (27.4%, 95% CI 22.8–32.0) were highest among refugees from Burundi. Samples from Burundian refugees were collected during 2006–2007 from persons in a single camp (Kibondo, Tanzania).

Of the 499 samples from refugees from Burma, 459 (92.0%) were collected during 2006–2007, a period of increased resettlement; the remaining 40 were collected during 2004–2005. The refugees came from 5 refugee camps and 2 urban populations, with most samples from
Mae La Camp (326 [65.3%]) and Tham Hin Camp (130 [26.1%]). The proportion of seropositive refugees was not equal between camps (p<0.01). The crude seroprevalence was significantly higher in Mae La (28.5%, 95% CI 23.6%–33.4%) than in Tham Hin (12.3%, 95% CI 6.6%–18.0%). After controlling for age and sex, we found that refugees from Mae La were >3× more likely than refugees from Tham Hin to be seropositive (OR = 6.2, 95% CI 2.6–14.6). The proportional distribution of an atypical reaction occurring in a sample from Burundi (6 [5.2%]), Laos (2 [2.2%]), or Bhutan (0). The odds of an atypical reaction occurring in a sample from Burundi were 6× greater than for all of the other groups combined (OR 6.2, 95% CI 2.6–14.6). The proportional distribution of atypical reactions did not differ with respect to age (p = 0.94) or sex (p = 0.54).

Discussion

We demonstrated that exposure to *T. solium* parasitic infection is common among refugees from Burma, Laos, Burundi, and Bhutan who resettled to the United States. All 4 populations had seroprevalence of antibodies against *T. solium* cysts comparable to or higher than the seroprevalence in well-characterized *T. solium*–endemic communities in Latin America where illness attributable to NCC is common (4,19,20). The widespread exposure among these groups has clinical and public health implications because these populations are resettling to the United States, where the infection is not endemic and where many clinical providers are not familiar with the disease manifestations, diagnosis, or treatment.

### Table 1. Relationship between sex and seroprevalence of antibodies against *Taenia solium* cysts among refugees resettled in the United States

<table>
<thead>
<tr>
<th>Male refugees</th>
<th>Female refugees</th>
<th>Odds ratio* (95% CI)</th>
<th>p value†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Country of origin</td>
<td>No. positive/total no.</td>
<td>% (95% CI)</td>
<td>No. positive/total no.</td>
</tr>
<tr>
<td>Burma (Myanmar)</td>
<td>78/273</td>
<td>28.6 (23.2–34.0)</td>
<td>38/226</td>
</tr>
<tr>
<td>Laos</td>
<td>41/240</td>
<td>17.1 (12.3–21.9)</td>
<td>51/262</td>
</tr>
<tr>
<td>Burundi</td>
<td>57/234</td>
<td>24.4 (18.8–29.9)</td>
<td>72/266</td>
</tr>
<tr>
<td>Bhutan</td>
<td>56/237</td>
<td>23.6 (18.2–29.6)</td>
<td>58/263</td>
</tr>
<tr>
<td>Total</td>
<td>232/984</td>
<td>23.6 (20.9–26.2)</td>
<td>219/1,017</td>
</tr>
</tbody>
</table>

*Odds of positive result for enzyme-linked immunoelectrotransfer blot for lentil-lectin purified glycoprotein (EITB LLGP) testing of samples from male refugees compared with samples from female refugees.

†Pearson χ².

### Table 2. Crude and age–sex standardized seroprevalence of antibodies against *Taenia solium* cysts among refugees resettled in the United States

<table>
<thead>
<tr>
<th>Variable</th>
<th>Burma, n = 499</th>
<th>Laot, n = 502</th>
<th>Bhutan, n = 500</th>
<th>Burundi, n = 500</th>
<th>p value*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, y, median (interquartile range)</td>
<td>29 (22–40)</td>
<td>28 (20–47)</td>
<td>30 (22–45)</td>
<td>21 (18–25)</td>
<td>&lt;0.01†</td>
</tr>
<tr>
<td>Male, no. (%)</td>
<td>273 (54.7)</td>
<td>240 (47.8)</td>
<td>237 (47.4)</td>
<td>234 (46.8)</td>
<td>0.04</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Seroprevalence, % (95% CI)</th>
<th>Age–sex standardized†</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Burma, n = 499</td>
</tr>
<tr>
<td>Crude</td>
<td>23.2 (19.5–27.0)</td>
</tr>
<tr>
<td>Age–sex standardized†</td>
<td>23.0 (19.1–26.8)</td>
</tr>
</tbody>
</table>

*Pearson χ² unless otherwise noted.
†Kruskall–Wallis χ².
‡Direct standardization method.
Epilepsy and other neurologic diseases associated with *T. solium* infection may be prevalent among certain populations of resettled refugees. We were unable to characterize the prevalence of *T. solium*–related disease because of the retrospective nature of this study and the lack of clinical data accompanying the anonymous samples. However, case reports of symptomatic NCC among resettled refugees are now being reported in the literature (11–15). Clinicians who care for migrants from countries where *T. solium* infection is endemic should have a high index of suspicion for NCC when encountering seizure disorder, chronic headache, or other neurologic deficits of unknown cause. Up-to-date information on diagnosis and treatment of NCC is available in recent reviews (21–23).

Although human cysticercosis is considered a dead end in the *T. solium* life cycle, a person with taeniasis can transmit infection to others by shedding infective eggs in feces. An adult-stage tapeworm can live for several years within the human intestine and intermittently releasing proglottids containing tens of thousands of potentially infective eggs. The infrequent reports of *Taenia* spp. infections in fecal samples of resettled refugees may not reliably indicate the true prevalence of taeniasis (24). Routine screening of fecal samples is done by light microscopy, which has low sensitivity (<40%) for *Taenia* spp. (25,26). The number of imported *T. solium* taeniasis infections can be estimated among the populations we tested by extrapolation from other communities with similar seroprevalence in areas in which it is endemic. Multiple studies using the highly sensitive (99%) coproantigen ELISA in Latin America have shown the prevalence of *Taenia* spp. taeniasis in *T. solium*–endemic communities to be 2.0%–3.5% (19,27–30). These estimates include both *T. solium* and *T. saginata* tapeworms because the coproantigen ELISA used in those studies does not differentiate between these species (26). Approximately 87,000 refugees resettled to the United States from the 4 refugee populations we sampled during 2004–2009 (10). By using a conservative estimate of 1% prevalence, ≈870 refugees with *T. solium* taeniasis may have entered the United States from these populations alone.

Identifying and treating taeniasis among resettling refugees could prevent further transmission of cysticercosis in destination countries. The current approach for controlling intestinal helminthes among resettling refugees involves presumptive treatment before resettlement rather than routine fecal screening (www.cdc.gov/immigrantrefugeehealth/guidelines/overseas/intestinal-parasites-overseas.html). *Taenia* spp. tapeworms are not specifically targeted, although refugees from *Schistosoma* spp.–endemic areas in Africa may receive presumptive treatment with praziquantel, which is an effective treatment for *T. solium* taeniasis. Household screening for taeniasis when cysticercosis is diagnosed in an area to which it is not endemic is an alternate approach that can identify persons with *T. solium* intestinal tapeworm infection (6,31–35). Clinicians who diagnose cysticercosis should consider screening the index case-patient for taeniasis and household members for taeniasis and NCC. A combination of clinical history, laboratory analysis of feces and serum, and neuroimaging may be required.

The demonstration of widespread exposure to *T. solium* tapeworms among certain refugee populations is of concern because of the potential for severe adverse events related to presumptive treatment for intestinal helminthes. Refugees resettling to the United States from Africa and Asia receive presumptive treatment for intestinal roundworms. All
refugees without contraindication receive a single dose of albendazole, and refugees originating in sub-Saharan Africa receive additional treatment for schistosomiasis with praziquantel before departure. These guidelines are consistent with program strategies of presumptive treatment in parasite-endemic areas used by the World Health Organization for soil helminth infections (www.who.int/intestinal_worms/strategy/en/) and schistosomiasis (www.who.int/schistosomiasis/strategy/en/). Both medications are used in the treatment of NCC because of their ability to penetrate the CNS and to damage *T. solium* cysts. Corticosteroids are typically administered simultaneously in treatment of NCC to control resulting inflammation and to prevent neurologic complications (22). The long latency between CNS infection and development of symptoms means that some persons with NCC will have occult viable brain cysts. Inadvertent damage to these occult brain cysts during presumptive treatment for intestinal helminthiasis could precipitate an inflammatory CNS reaction in patients for whom presumptive treatment would otherwise have been contraindicated had their infection been known. Multiple case reports describe new-onset seizures in persons with underlying NCC who receive treatment with these agents (12,35–39). The Food and Drug Administration recently updated label precautions for albendazole and praziquantel to inform clinicians about this potential adverse event (www.accessdata.fda.gov/drugsatfda_docs/appletter/2009/020666s005,s006ltr.pdf and www.accessdata.fda.gov/drugsatfda_docs/appletter/2010/018714s012ltr.pdf).

The current CDC refugee predeparture health guidelines advise avoiding presumptive treatment for intestinal helminthiasis in patients with known history of cysticercosis or previous seizure (www.cdc.gov/immigrantrefugeehealth/guidelines/refugee-guidelines.html). However, refugees from *T. solium* tapeworm–endemic regions may harbor occult viable CNS cysts that could increase their risk for severe adverse events. Refugees are observed for 1–3 days after drug administration before departure; however, enhanced surveillance with systematic data collection would help inform risk–benefit analyses of these programs. Prospective studies that monitor adverse neurologic reactions after mass treatment with albendazole and/or praziquantel in *T. solium* tapeworm–endemic areas are needed to quantify the actual risk. Clinicians should be aware of potential adverse treatment events when evaluating refugees who develop neurologic symptoms after presumptive therapy and should report suspected cases by email to CDC (RefGuidelines@cdc.gov) or telephone (1-404-498-1600) and to the Food and Drug Administration though the Adverse Events Reporting System (www.fda.gov/Drugs/GuidanceComplianceRegulatoryInformation/Surveillance/AdverseDrugEffects/ucm115894.htm).

Although we cannot confirm that active transmission is occurring within the refugee camps and/or within surrounding communities on the basis of seroprevalence alone, we do suspect that active transmission is occurring for 2 reasons: 1) the populations we sampled lived in the refugee camps for years to decades before resettlement, and 2) the antibody response to *T. solium* cysts detected on EITB LLGP has been demonstrated to be transient, with ≈40% reversion from seropositive to seronegative after 1 year in serial community studies (40). Although we were unable to adequately explore risk factors for positive serologic findings in this study, we did detect significant differences in the odds of exposure between refugee camps. Further investigation within the camps and surrounding communities may clarify reasons for the variation observed. Areas for further study include characterizing 1) the prevalence of epilepsy and other neurologic disease associated with NCC, 2) the prevalence of and risk factors for taeniasis, 3) the prevalence of and risk factors for porcine cysticercosis, and 4) animal husbandry practices and market structure for sale of pork. Interventions, such as screening for and treatment of taeniasis, use of corrals for raising pigs, and improved sanitation infrastructure and education, may reduce transmission among refugees and ultimately prevent disease.

This study has limitations. The EITB LLGP is known to have low sensitivity for detecting single parenchymal cysts and calcified cysts alone. On the other hand, the 100% specificity to the larval stage of *T. solium* means that false-positive reactions are unlikely. Seroprevalence estimates based on the EITB LLGP are therefore likely to underestimate the overall prevalence of exposure to *T. solium* eggs in a community. Although *T. asiatica* tapeworms are co-endemic in Southeast Asia, there is no evidence for or against potential cross-reactivity of this related species on the EITB LLGP. However, *T. asiatica* cysticercosis has not been reported among humans. We preselected our sample to include refugee populations in which we expected to find evidence of endemic *T. solium* transmission. Our seroprevalence estimates should not be generalized to all resettling refugee populations, particularly those from Middle Eastern or northern African countries, to which *T. solium* tapeworms are not thought to be endemic. Our seroprevalence estimates also may not be generalizable to the broader population in the refugees’ countries of origin. Refugee populations often include ethnic minority groups whose compilation of risk factors may not represent those of the majority population in their countries of origin. Nevertheless, our study does provide seroprevalence estimates for regions in which little to no data were previously available.

Exposure to *T. solium* parasitic infection is widespread among specific refugee groups resettled to the United States. Clinicians should suspect NCC in patients from these
regions who have seizure, headache, or other unexplained neurologic manifestations and should consider screening household members for additional cases. Systematic screening and treatment for taeniasis among refugees may prevent additional cases of NCC. Further investigation is needed to characterize illness and risk factors associated with *T. solium* infection in refugee populations. Additional serologic testing of stored samples from different regions of the world with the EITB LLGP can improve understanding of the global distribution of *T. solium* infection and may highlight regions that could benefit from control or elimination interventions.

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**References**


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