Risk factors for sporadic *Giardia* infection in the USA: a case-control study in Colorado and Minnesota

H. E. Reses1, J. W. Gargano1, J. L. Liang1, A. Cronquist2, K. Smith3, S. A. Collier1, S. L. Roy1, J. Vanden Eng1, A. Bogard3, B. Lee3, M. C. Hlavsa1, E. S. Rosenberg4, K. E. Fullerton1, M. J. Beach1 and J. S. Yoder1

1Division of Foodborne, Waterborne, and Environmental Diseases, National Center for Emerging and Zoonotic Infectious Diseases, Centers for Disease Control and Prevention, Atlanta, Georgia, USA; 2Colorado Department of Public Health and Environment, Denver, Colorado, USA; 3Minnesota Department of Health, Saint Paul, Minnesota, USA and 4Department of Epidemiology and Biostatistics, University at Albany School of Public Health, SUNY, Rensselaer, New York, USA

**Abstract**

*Giardia duodenalis* is the most common intestinal parasite of humans in the USA, but the risk factors for sporadic (non-outbreak) giardiasis are not well described. The Centers for Disease Control and Prevention and the Colorado and Minnesota public health departments conducted a case-control study to assess risk factors for sporadic giardiasis in the USA. Cases (*N* = 199) were patients with non-outbreak-associated laboratory-confirmed *Giardia* infection in Colorado and Minnesota, and controls (*N* = 381) were matched by age and site. Identified risk factors included international travel (aOR = 13.9; 95% CI 4.9–39.8), drinking water from a river, lake, stream, or spring (aOR = 6.5; 95% CI 2.0–20.6), swimming in a natural body of water (aOR = 3.3; 95% CI 1.5–7.0), male–male sexual behaviour (aOR = 45.7; 95% CI 5.8–362.0), having contact with children in diapers (aOR = 1.6; 95% CI 1.01–2.6), taking antibiotics (aOR = 2.5; 95% CI 1.2–5.0) and having a chronic gastrointestinal condition (aOR = 1.8; 95% CI 1.1–3.0). Eating raw produce was inversely associated with infection (aOR = 0.2; 95% CI 0.1–0.7). Our results highlight the diversity of risk factors for sporadic giardiasis and the importance of non-international-travel-associated risk factors, particularly those involving person-to-person transmission. Prevention measures should focus on reducing risks associated with diaper handling, sexual contact, swimming in untreated water, and drinking untreated water.

**Introduction**

*Giardia duodenalis*, a flagellated protozoan, is a major cause of diarrhoeal illness worldwide [1]. In the USA, *Giardia* is the most commonly identified enteric parasite of humans [2], with an estimated 1.2 × 10⁶ episodes of illness occurring annually [3] and *Giardia* identified in 4–7% of stool specimens from patients with diarrhoeal illness tested by clinical diagnostic laboratories [2]. *Giardia* infection typically results in self-limited gastrointestinal (GI) illness characterised by diarrhoea, abdominal cramps, bloating, malaise, nausea, weight loss and malabsorption [1, 4]. Asymptomatic infection also occurs frequently [1, 4]. Additionally, giardiasis has been associated with the development of chronic diarrhoea or irritable bowel syndrome [5], fatigue, reactive arthritis and allergies [4, 6, 7]. Cases of giardiasis are estimated to result in 1918 emergency department visits, 3581 hospitalisations and $34 million in hospitalisation costs annually in the USA [8, 9].

The infectious dose of *Giardia* is low, with ingestion of as few as ten cysts needed to cause infection [10]. Cysts are immediately infectious upon being excreted in feces [10] and infected individuals can shed 1 × 10⁸ to 1 × 10⁹ cysts in their stool each day for several months [10–12]. *Giardia* cysts are environmentally hardy and moderately chlorine tolerant and can, therefore, survive in water, food, or on surfaces for several weeks to several months [1, 13]. *Giardia* infection is transmitted through the faecal-oral route and results from the ingestion of *Giardia* cysts. Transmission occurs indirectly through the consumption of faecally contaminated water or food and directly through animal-to-person transmission and person-to-person transmission. Additional host-level factors such as illnesses and medications might impact individuals’ susceptibility to infection [14].

Most information on risk factors for domestically-acquired giardiasis in the USA has come from outbreak investigations [15]. From 1971–2011, 242 US giardiasis outbreaks were reported to the Centers for Disease Control and Prevention (CDC). These outbreaks resulted from waterborne (74.8%), foodborne (15.7%), person-to-person (2.5%) and zoonotic (1.2%)
transmission [16]. However, >99% of giardiasis cases are sporadic (i.e., not associated with a recognised outbreak) [17] and only a few case-control studies in the USA have investigated risk factors for sporadic giardiasis [18–20]. The CDC collaborated with the Colorado and Minnesota public health departments to conduct a case-control study of sporadic giardiasis in Colorado and Minnesota aimed at evaluating a wider range of risk factors among more participants across an expanded geographic area compared with the relatively small, single-state studies that preceded it.

Methods

Data collection

Persons with laboratory-confirmed Giardia infection were identified by the Colorado and Minnesota public health departments in 2003 and 2004. Under a protocol approved by the CDC and the states’ Institutional Review Boards, public health departments identified case-patients over a 12-month period by monitoring reports of laboratory-confirmed Giardia infections not associated with known outbreaks under investigation by the state or local health departments. Colorado monitored the Colorado Electronic Disease Reporting System for laboratory-confirmed cases in their catchment area, consisting of seven counties within the Denver metro area. Minnesota received reports from the entire state through a passive reporting system. Due to the higher incidence of giardiasis and larger catchment area in Minnesota, every second laboratory-confirmed case was eligible for the study.

Cases were defined as individuals with symptomatic, non-outbreak-associated, laboratory-confirmed giardiasis and an onset of illness within 6 weeks of their public health interview date. For each case-patient, up to two matched controls were enrolled. Controls were matched to case-patients by site (Colorado and Minnesota) and age group (≤11 months, 1–4 years, 5–11 years, 12–17 years, 18–44 years, 45–64 years and ≥65 years). Controls were recruited using progressive digit dialing anchored on the case-patient’s phone number, where one number was added to the case-patient’s phone number until someone who met the inclusion criteria was reached. Only one control was recruited per household.

Investigators contacted case-patients and controls by telephone, obtained informed consent (from parents or guardians for participants aged <18 years) and enrolled them in the study. A structured telephone questionnaire was administered by trained staff to participants ≥12 years of age or to the parent or guardian of participants <12 years of age. Participants were asked about demographic information, medical conditions, illness symptoms and possible exposures occurring in the 2 weeks prior to the onset of GI symptoms in the matched case-patient. Risk factor questions addressed health status and medications, drinking-water consumption, recreational water exposure, travel and outdoor activity, food and drink consumption, contact with young children and persons with diarrhoea, contact with animals, and sexual practices (only among persons ≥18 years of age).

Case-patients and controls were excluded if they lived outside of the states’ surveillance catchment areas, were not reached after 15 telephone attempts, had unavailable telephone numbers, were physically or emotionally unable to answer questions, were non-English speakers, or were recent immigrants, adoptees, or refugees. Additional exclusion criteria that applied only to case-patients included asymptomatic infection, unknown illness onset date, previous Giardia infection within 3 months, and not having the earliest symptom onset date within a household cluster of laboratory-confirmed giardiasis. Additional exclusion criteria that applied only to controls included diarrhoea during the 4 weeks before the illness onset date of the matched case-patient, or diarrhoea in a household contact during the 4 weeks before the illness onset date of the matched case-patient.

Statistical analysis

Over 500 exposure questions were included in the questionnaire; to prioritise potential risk factors for model building, we developed a framework to organise host factors and transmission pathways for Giardia infection (Fig. 1). We subdivided transmission pathways into direct faecal contact and indirect faecal contact. We defined direct faecal contact as risk factors for animal-to-person transmission and person-to-person transmission. We defined indirect faecal contact as activities that could lead to faecal ingestion without contact with the human or animal source, including drinking-water consumption, recreational water exposures, food consumption, travel and outdoor activity. Host factors included illnesses and medications that might impact susceptibility to infection. Several analysis exposures were derived by combining multiple items from the questionnaire and recoding free text responses. We selected potential giardiasis risk factors from the framework for analysis based on biological plausibility, previously reported associations and those that contained at least 25 exposed individuals.

Data were analysed using SAS software (Version 9.3; SAS Institute, Cary, NC) with multivariable logistic regression. Crude odds ratios (OR) and 95% confidence intervals (CI) were estimated for each exposure variable of interest. ORs were also calculated for each exposure, controlling for the matching variables age and site; these values were essentially equivalent to the crude estimates, so results from the unadjusted models were considered in further analyses. Exposures associated with disease status with P < 0.20 in the crude models or those that had a previously reported association with giardiasis were included in the multivariable logistic regression model. This model estimated adjusted ORs (aOR) and 95% CIs, controlling for age, site and sex. The interaction between male or female sex and having a sex partner of the same sex was assessed in this model. Population attributable fractions (PAF) of international travel during the 2-week exposure period. This multivariable model, referred to as the domestic-only model, also controlled for sex, age and site.

Statistical analysis: semi-Bayes shrinkage methods

Given the large number of potential risk factors assessed in the questionnaire, a separate analysis applied semi-Bayes shrinkage methods to the distribution of ORs for 70 exposures that have not previously been shown to have an association with giardiasis, or those that had too few cases and controls exposed to result in stable estimates in the conventional multivariable regression model that evaluated 18 potential risk factors. After excluding all associations that we expected a priori, we applied semi-Bayes
shrinkage methods to the distribution of the remaining ORs [22, 23]. Semi-Bayes shrinkage narrows the distribution of observed ORs and improves their precision by applying a regression-derived shrinkage estimator [24]. When this method is applied, imprecisely measured associations that are well above or below the null are drawn towards the center of the distribution, thereby reducing the potential to overestimate true associations and allowing us to identify new associations with improved validity [24]. To implement this method, it was assumed that 95% of the true ORs would fall between 0.25 and 4. Each OR was adjusted for age, site and sex. We also adjusted each OR for international travel and drinking water from a river, lake, stream, or spring because these exposures are previously reported strong risk factors and therefore could be confounders [25].

**Results**

**Study population**

During the 12-month study period, 653 cases of laboratory-confirmed *Giardia* infection were identified in Colorado and Minnesota. A total of 213 case-patients were enrolled, of which 199 were matched with 381 controls; 14 case-patients were excluded because no matched controls were found (Fig. 2).

**Demographic characteristics**

Case-patients were mostly white (97.0%) and non-Hispanic (92.9%) (Table 1). The median age of enrolled case-patients and controls was 37 years. Case-patients and controls were generally similar with regard to the distribution of race, ethnicity, income, site and age, but more case-patients were male compared with controls (56.3% of case-patients vs. 39.1% of controls; OR = 2.0; 95% CI 1.4–2.8; *P* < 0.0001).

**Risk factor analysis**

In the bivariate and multivariable analyses, international travel, having contact with children in diapers, drinking water from a river, lake, stream, or spring, swimming in a natural body of water, taking antibiotics and having a chronic GI condition or taking medication indicated for such a condition (hereafter referred to as chronic GI condition), were associated with a statistically significant increased odds of giardiasis, and eating raw vegetables or fruit was associated with a decreased odds of giardiasis (Table 2). In addition to these main effects, an interaction between having a sex partner of the same sex and being male/female was detected (Wald $\chi^2 = 7.3$; *P* = 0.007). Male–male sexual behaviour was associated with significantly increased odds of giardiasis (aOR = 45.7; 95% CI 5.8–362.0), but female–female sexual behaviour was not
Factors that were associated with giardiasis in the bivariate analysis but not after multivariable adjustment included camping, animal contact, swimming in a manmade aquatic venue, and taking psychoactive medication. Results were similar after excluding individuals with a history of international travel, except that an additional significant crude association between giardiasis and travelling in the USA was found (Supplementary Table S1; Supplementary Material is available on the Cambridge Core website).

PAFs

Among all case-patients, the proportion whose illness could be attributed to foreign travel was 11.2% (Table 2). The largest PAFs resulted from contact with children wearing diapers (15.1%), having a chronic GI condition (12.5%), swimming in a natural body of water (10.5%), drinking water from a river, lake, stream, or spring (9.4%) and male–male sexual behaviour (8.4%).

Semi-Bayes shrinkage methods

Supplementary Table S2 shows the conventional and semi-Bayes-adjusted ORs and 95% CIs for 70 exposures that do not have a previously reported association with giardiasis, or those that had too few cases and controls exposed to result in stable estimates in the conventional multivariable regression model. Prior to semi-Bayes adjustment, the conventional ORs ranged from 0.1 (95% CI 0.01–0.5) for swimming in or entering a recreational waterpark to 35.8 (95% CI 4.6–276.2) for male–male sexual behaviour. After semi-Bayes adjustment, the ORs ranged from 0.5 (95% CI 0.2–1.2) for taking pain medication to 3.9 (95% CI 1.2–13.1) for same-gender sexual behaviour. Of the 70 exposures considered, 11 were associated with an increased odds of giardiasis and four were associated with a decreased odds of giardiasis prior to semi-Bayes adjustment. After the adjustment, the only exposure that remained significantly associated with giardiasis was same-gender sexual behaviour.

Discussion

This case-control study represents the largest and most extensive assessment of risk factors for sporadic Giardia infection in the USA to date. The results from this study demonstrate the importance of domestic transmission and the diversity of risk factors for giardiasis across multiple transmission pathways. Among these pathways, person-to-person transmission was responsible for the largest proportion (32.9%) of illness among case-patients, representing an opportunity for intervention through modifiable, individual-level changes in behaviour.

Giardia infection was strongly associated with international travel (aOR = 13.9), supporting the results of previous case-control studies in the USA [18] and other developed countries including the UK [26, 27], Canada [28] and New Zealand [29]. Despite this strong association, only 11.2% of the illness in this group of case-patients was attributed to international travel, underscoring the importance of domestic risk factors for giardiasis.

(aOR = 0.6; 95% CI 0.1–6.4). Factors that were associated with giardiasis in the bivariate analysis but not after multivariable adjustment included camping, animal contact, swimming in a manmade aquatic venue, and taking psychoactive medication.

Results were similar after excluding individuals with a history of international travel, except that an additional significant crude association between giardiasis and travelling in the USA was found (Supplementary Table S1; Supplementary Material is available on the Cambridge Core website).
The actual transmission pathways that result in infection associated with international travel likely include drinking local water, eating undercooked food and direct faecal contact due to inadequate sanitation. Therefore, individuals should practice extra caution when traveling and avoid consuming undercooked food and inadequately treated water and ice [30]. In the unadjusted analyses, travelling in the USA (in the domestic-only model) and camping were associated with giardiasis. Camping has been frequently reported as a risk factor for giardiasis [18, 20, 26, 27] and one study in New Zealand reported domestic travel as a risk factor [31]. Like international travel, domestic travel and camping might confer an elevated risk through behaviours across multiple transmission pathways including eating undercooked food, practicing poor hygiene, drinking untreated surface water and drinking water from new sources with different pathogens to which individuals lack immunity.

Our study also provides evidence for transmission through consumption of contaminated water and food. Drinking water from a river, lake, stream, or spring was identified as a strong risk factor for giardiasis. Waterborne outbreaks and sporadic cases of giardiasis have been previously associated with consuming untreated water from natural bodies of water including rivers and streams [16, 19, 20, 32–34]. Studies in Colorado [20], Minnesota [34] and New Hampshire [19] identified consumption of untreated water as an important cause of sporadic giardiasis in the USA. In addition to direct consumption of contaminated water, recreational activities in lakes, rivers and swimming pools have been implicated as risk factors in numerous outbreak investigations and studies of sporadic giardiasis [15, 16, 29, 35, 36]. In our study, swimming in a natural body of water was associated with *Giardia* infection. Natural recreational waters are not treated with disinfectants, such as chlorine or bromine, the primary barrier to the transmission of infectious pathogens in aquatic venues [37]. Swimming in a manmade aquatic venue (i.e., swimming pool, waterpark, or hot tub/spa) was a risk factor in the unadjusted analysis (although not in the multivariable analysis), emphasising the importance of public health efforts to promote healthy swimming behaviours (e.g., staying out of the water if one has diarrhoea) to improve swimmer hygiene and prevent water contamination in all recreational water venues.

As with water, food can become contaminated by *Giardia* cysts and links between giardiasis and various types of produce have been reported [15, 31]. We found that eating raw vegetables or fruit was inversely associated with giardiasis. A similar protective effect was reported in a sporadic giardiasis case-control study in the UK [27]. It is possible that repeated exposure via contaminated raw produce could provide protective immunity [38]. Fruits and vegetables can become contaminated during irrigation and fertilisation activities, by water used during processing or packing, or by unwashed hands during harvesting or any point in the production chain [39]. Since fruits and vegetables are often eaten raw, preventing produce contamination and washing produce before use are particularly important [40]. The addition of serologic approaches to future studies could provide more data on the possible role of fresh produce for protective immunity. However, it is possible that this result might reflect increased healthful behaviours among controls compared with case-patients; individuals who frequently consume fruits and vegetables might have better general health than non-produce-consumers and could be less likely to contract giardiasis or develop a symptomatic infection.

Direct person-to-person transmission represented the most important transmission pathway in our study, as the risk factors within this pathway were responsible for the largest proportion of illness among case-patients. Contact with children in diapers had a stronger association with infection among men who have sex with men (MSM), was quantified in our study to document a statistically significant association between male sexual behaviour [42, 43], but this is the first epidemiological study to document a statistically significant association between giardiasis and male–male sexual behaviour. Further, same-gender

### Table 1. Demographic characteristics for giardiasis cases and controls in the Colorado and Minnesota giardiasis case-control study

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Cases (N = 199)</th>
<th>Controls (N = 381)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gendera</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>87/199 (43.7)</td>
<td>232/381 (60.9)</td>
</tr>
<tr>
<td>Male</td>
<td>112/199 (56.3)</td>
<td>149/381 (39.1)</td>
</tr>
<tr>
<td>Race</td>
<td></td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>191/197 (97.0)</td>
<td>336/364 (92.3)</td>
</tr>
<tr>
<td>Black</td>
<td>1/197 (0.5)</td>
<td>10/364 (2.8)</td>
</tr>
<tr>
<td>Otherb</td>
<td>5/197 (2.5)</td>
<td>18/364 (5.0)</td>
</tr>
<tr>
<td>Ethnicity</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hispanic</td>
<td>14/198 (7.1)</td>
<td>23/378 (6.1)</td>
</tr>
<tr>
<td>Non-Hispanic</td>
<td>184/198 (92.9)</td>
<td>355/378 (93.9)</td>
</tr>
<tr>
<td>Age</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0–12 months</td>
<td>1/199 (0.5)</td>
<td>1/375 (0.3)</td>
</tr>
<tr>
<td>1–5 years</td>
<td>16/199 (8.0)</td>
<td>29/375 (7.7)</td>
</tr>
<tr>
<td>5–12 years</td>
<td>17/199 (8.5)</td>
<td>30/375 (8.0)</td>
</tr>
<tr>
<td>12–&lt;18 years</td>
<td>4/199 (2.0)</td>
<td>8/375 (2.1)</td>
</tr>
<tr>
<td>18–&lt;45 years</td>
<td>92/199 (46.2)</td>
<td>172/375 (45.9)</td>
</tr>
<tr>
<td>45–&lt;65 years</td>
<td>58/199 (29.2)</td>
<td>116/375 (30.9)</td>
</tr>
<tr>
<td>65+ years</td>
<td>11/199 (5.5)</td>
<td>19/375 (5.1)</td>
</tr>
<tr>
<td>Household income</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;$25 000</td>
<td>25/171 (14.6)</td>
<td>50/318 (15.7)</td>
</tr>
<tr>
<td>$25 000–$55 000</td>
<td>53/171 (31.0)</td>
<td>107/318 (33.7)</td>
</tr>
<tr>
<td>$55 000–$75 000</td>
<td>36/171 (21.1)</td>
<td>59/318 (18.6)</td>
</tr>
<tr>
<td>&gt;$75 000</td>
<td>57/171 (33.3)</td>
<td>102/318 (32.1)</td>
</tr>
<tr>
<td>State of residence</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Colorado</td>
<td>100/199 (50.3)</td>
<td>200/381 (52.5)</td>
</tr>
<tr>
<td>Minnesota</td>
<td>99/199 (49.8)</td>
<td>181/381 (47.5)</td>
</tr>
</tbody>
</table>

aA statistically significant association between gender and giardiasis was identified (odds ratio = 2.0; 95% confidence interval = 1.4, 2.8; *P* = 0.0001).
bOther race includes American Indian/Alaskan Native, Asian/Pacific Islander, mixed, other.
sexual behaviour remained associated with giardiasis after semi-Bayes adjustment, which strengthened the evidence of risk. The relatively high PAFs associated with person-to-person transmission representing the largest risk factors within this pathway were underestimated as case-patients were excluded from the study if they did not have the earliest symptom onset date within a household cluster. Accordingly, it is likely that the estimates for risk factors in our study underestimated the risk associated with childcare centres, as transmission to close contacts of persons attending or working in childcare centres has been frequently reported [18, 19]. The relatively high PAFs associated with person-to-person transmission (32.9%), in spite of the exclusion of ill household

<table>
<thead>
<tr>
<th>Exposure</th>
<th>Cases (N = 199)</th>
<th>Controls (N = 381)</th>
<th>Unadjusted ORa</th>
<th>Adjusted ORb</th>
<th>Population-Attributable Fraction (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>International travel</td>
<td>24 (12.1)</td>
<td>5 (1.3)</td>
<td>10.3 (3.9, 27.5)</td>
<td>13.9 (4.9, 39.8)</td>
<td>11.2</td>
</tr>
<tr>
<td>Travel in USAf</td>
<td>78 (39.2)</td>
<td>123 (32.3)</td>
<td>1.4 (0.9, 1.9)</td>
<td>1.0 (0.7, 1.6)</td>
<td>0.0</td>
</tr>
<tr>
<td>Camp</td>
<td>29 (14.6)</td>
<td>26 (6.8)</td>
<td>2.3 (1.3, 4.1)</td>
<td>1.2 (0.6, 2.6)</td>
<td>2.4</td>
</tr>
<tr>
<td>Animal contacti</td>
<td>129 (64.8)</td>
<td>278 (73.0)</td>
<td>0.7 (0.5, 0.99)</td>
<td>0.7 (0.4, 1.1)</td>
<td>–</td>
</tr>
<tr>
<td>Contact with children wearing diapers</td>
<td>80 (40.2)</td>
<td>138 (36.2)</td>
<td>1.2 (0.8, 1.7)</td>
<td>1.6 (1.01, 2.6)</td>
<td>15.1</td>
</tr>
<tr>
<td>Contact with person with diarrhoea1</td>
<td>24 (12.1)</td>
<td>31 (8.1)</td>
<td>1.5 (0.9, 2.7)</td>
<td>1.5 (0.8, 2.8)</td>
<td>4.0</td>
</tr>
<tr>
<td>Contact with children in childcareg</td>
<td>32 (16.1)</td>
<td>54 (14.2)</td>
<td>1.2 (0.7, 1.9)</td>
<td>1.5 (0.8, 2.8)</td>
<td>5.4</td>
</tr>
<tr>
<td>Male–female sexual behaviourh</td>
<td>17 (15.2)</td>
<td>1 (0.7)</td>
<td>26.5 (3.5, 202.0)</td>
<td>45.7 (5.8, 362.0)</td>
<td>8.4</td>
</tr>
<tr>
<td>Female–female sexual behaviourh</td>
<td>1 (1.2)</td>
<td>5 (2.2)</td>
<td>0.5 (0.1, 4.6)</td>
<td>0.6 (0.1, 6.4)</td>
<td>–</td>
</tr>
<tr>
<td>Drink water from wellj</td>
<td>48 (24.1)</td>
<td>87 (22.8)</td>
<td>1.1 (0.7, 1.6)</td>
<td>1.1 (0.7, 1.9)</td>
<td>2.2</td>
</tr>
<tr>
<td>Drink water from lake, river, stream, or spring</td>
<td>22 (11.1)</td>
<td>5 (1.3)</td>
<td>9.3 (3.5, 25.1)</td>
<td>6.5 (2.0, 20.6)</td>
<td>9.4</td>
</tr>
<tr>
<td>Swim in natural body of wateri</td>
<td>30 (15.1)</td>
<td>16 (4.2)</td>
<td>4.1 (2.1, 7.6)</td>
<td>3.3 (1.5, 7.0)</td>
<td>10.5</td>
</tr>
<tr>
<td>Swim in manmade aquatic venuek</td>
<td>48 (24.1)</td>
<td>61 (16.0)</td>
<td>1.7 (1.1, 2.6)</td>
<td>1.2 (0.7, 2.1)</td>
<td>4.0</td>
</tr>
<tr>
<td>Eat raw vegetables or fruits1</td>
<td>187 (94.0)</td>
<td>374 (98.2)</td>
<td>0.3 (0.1, 0.8)</td>
<td>0.2 (0.1, 0.7)</td>
<td>–</td>
</tr>
<tr>
<td>Take antibiotics</td>
<td>21 (10.6)</td>
<td>22 (5.6)</td>
<td>1.9 (1.03, 3.6)</td>
<td>2.5 (1.2, 5.0)</td>
<td>6.3</td>
</tr>
<tr>
<td>Take psychoactive medication</td>
<td>21 (10.6)</td>
<td>15 (3.9)</td>
<td>2.9 (1.4, 5.7)</td>
<td>1.7 (0.7, 4.1)</td>
<td>4.3</td>
</tr>
<tr>
<td>Chronic gastrointestinal condition or medicationi</td>
<td>56 (28.1)</td>
<td>68 (17.9)</td>
<td>1.8 (1.2, 2.7)</td>
<td>1.8 (1.1, 3.0)</td>
<td>12.5</td>
</tr>
<tr>
<td>Immunosuppressing condition, medication, or treatmenti</td>
<td>21 (10.6)</td>
<td>41 (10.8)</td>
<td>1.0 (0.6, 1.7)</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

aUnadjusted odds ratio. 
bAdjusted odds ratio from full multivariable model adjusted for age, site, sex and all exposures that have been previously reported to be associated with giardiasis and showed an association with disease status with P < 0.2 in the bivariate models. Given these criteria, immunosuppressing condition, medication, or treatment was excluded from the multivariable analysis.

cThe variable ‘travel in the United States’ was derived by grouping travel within home state more than 100 miles and travel outside of home state within the USA.

deThe variable ‘animal contact’ was derived by grouping any contact with domestic pets (i.e., kitten, cat, puppy, dog) and contact with farm mammals (i.e., calf, cow, goat, sheep, horse, pig).

fThe variable ‘contact with children wearing diapers’ was derived by grouping children in home wear diapers, children at child’s childcare location wear diapers, contact with children in diapers and change diapers.

gThe variable ‘contact with person with diarrhoea’ was derived by grouping direct contact with person with diarrhoea and provide care for person with diarrhoea.

hThe variable ‘contact with children in childcare’ was derived by grouping attends childcare, children in home attend out-of-home childcare, provide childcare outside of home, provide childcare in home.

iStatistically significant interaction was identified between sex (male vs. female) and having a same-gender sex partner; this is presented stratified by individual sex.

jThe variable ‘drink water from well’ was derived by grouping well as source of drinking water at home and well as source of drinking water outside of home.

kThe variable ‘swim in natural body of water’ was derived by grouping swim, wade, or enter an ocean, lake, pond, river, stream, or hot spring.

lThe variable ‘swim in manmade aquatic venue’ was derived by grouping swim, wade, or enter a swimming pool, hot tub/spa, or waterpark.

mThe variable ‘eat raw vegetables or fruits’ was derived by grouping eat lettuce, garden salad, raw vegetables, raw berries, raw fruits with skin/peel and any organic produce.

nThe variable ‘chronic gastrointestineal (GI) condition’ was derived by grouping chronic lower GI conditions (i.e., chronic diarrhoea, Crohn’s disease, irritable bowel syndrome), chronic lower GI medications (i.e., laxatives, enemas, antidiarrieal medications) and chronic upper GI medications (i.e., stomach-acid reducing medications).

oThe variable ‘immunosuppressing condition, medication, or treatment’ was derived by grouping immunosuppressing conditions (i.e., diabetes, cancer, HIV/AIDS, other immunodeficiency), medications (i.e., steroids, cyclosporine) and treatments (i.e., organ transplant, chemotherapy, radiation).
members, suggest that person-to-person transmission represents an important pathway for sporadic giardiasis transmission in the USA and prevention efforts should target behaviours associated with the direct faecal contact.

Our study was novel in its consideration of the role of host factors (e.g., illnesses and medications that might influence host susceptibility) in *Giardia* transmission. We identified a previously unreported risk for giardiasis associated with taking antibiotics during the 2 weeks prior to symptom onset. Disruption of the microbiota with antibiotics is known to precede the emergence of several enteric pathogens including *Salmonella* spp. and *Clostridium difficile* [46–48]. Antibiotic use may lead to prolonged alterations in the microbiota and decreased colonisation resistance, lowering the dose of pathogens needed for colonisation and infection and therefore increasing the risk of illness [48]. While antibiotics are beneficial when clearly indicated, our result underscores the importance of avoiding unnecessary use. In addition to antibiotics, we found a previously unreported host-level risk associated with taking psychoactive medication in the bivariate analysis. Psychoactive medications, including medications for anxiety, depression and psychosis, have antimicrobial activity and might influence gut motility and sensation; this could also change the composition of gut microbiota and increase infection susceptibility [49, 50]. Further, we identified an additional novel host-level association between giardiasis and having a chronic GI condition. This was an important risk factor, with a PAF of 12.5%. Elevated giardiasis prevalence has been reported in patients with GI conditions including Crohn’s disease [51], irritable bowel syndrome [27], celiac disease [52] and other GI diseases [51]. It has been suggested that GI conditions might influence *Giardia* infection susceptibility by disturbing the protective function of the mucosa in inflammatory states [51]. However, it is not clear whether individuals with these conditions are more susceptible to infection or whether they are more likely to develop symptomatic disease. In addition, conditions with symptoms resembling those of giardiasis (e.g., chronic GI conditions, co-infection with other enteric pathogens) might increase the probability of a *Giardia* test being performed, thereby increasing the probability of classifying asymptomatic infection as symptomatic giardiasis. While diarrhea caused by other conditions is possible, the finding of this association warrants further investigation in studies with more extensive clinical data [6].

Our study has several limitations. The study design was a retrospective case-control in which exposure information was ascertained via subject interview after disease status was known, potentially resulting in recall bias. The study design might have also led to a selection bias in favour of controls who were more likely to be at home and, thus, perhaps not representative of the general population. In addition, persons with asymptomatic infections might have been misclassified as controls since stool samples were not collected to confirm the absence of infection in the controls. Further, our data were collected in two states over 1 year, thereby limiting generalisability. Finally, PAFs from case-control studies should be interpreted with caution because ORs might not represent causal effects. Although the data for this study were collected in 2003 and 2004, the main giardiasis transmission pathways have likely remained commonplace in the USA. Rates in the overall population have declined, but it is unclear if this is related to changes in disease transmission or changes in surveillance, given the 2011 modification of the case definition clarifying that clinical symptoms are necessary for categorising giardiasis cases as confirmed [45]. Changes in surveillance capacity and a decreased emphasis on giardiasis surveillance might have also occurred [45].

In summary, this study represents the largest, most recent and most comprehensive case-control study of sporadic giardiasis in the USA to date. Our results highlight the diversity of risk factors for sporadic giardiasis and the importance of domestic risk factors, particularly those that are related to direct transmission from person-to-person. We also identified novel risk factors and host-level factors associated with infection, including male–male sexual behaviour, taking antibiotics and having a chronic GI condition. Educational efforts to prevent direct faecal contamination and transmission and to decrease exposure to unsafe drinking and recreational water have the potential to considerably reduce sporadic giardiasis transmission in the USA. In addition, the low percentage of illnesses associated with international travel (11%) emphasises the importance of domestic giardiasis transmission and the need for healthcare providers and the general public to have improved awareness of non-international-travel-associated infection through multiple transmission pathways.

**Supplementary material.** The supplementary material for this article can be found at [https://doi.org/10.1017/S0950268818001073](https://doi.org/10.1017/S0950268818001073)

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**Declaration of interest.** None.

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