Concordance of Helicobacter pylori infection among children in extended-family homes

P. K. GARG¹, S. PERRY*, L. SANCHEZ² AND J. PARSONNET²

¹ Northwestern University Feinberg School of Medicine, Chicago, IL, USA
² Division of Geographic Medicine and Infectious Disease, Stanford University School of Medicine, Stanford, CA, USA

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SUMMARY

Helicobacter pylori is transmitted within households and high concordance is observed among siblings. To better understand the contributions of close interpersonal contact and family relatedness to transmission, we compared concordance of H. pylori infection among 241 sibling and non-sibling children aged 2–18 years in 68, predominantly low-income, Hispanic households with at least two nuclear families. Prevalence of H. pylori infection was 24%. Compared to children with no infected siblings or non-siblings and adjusting for age, odds of H. pylori infection were 1.2 (95% CI 0.52–2.9), 3.2 (95% CI 1.14–9.1), and 9.4 (95% CI 3.1–28.5) for children residing with at least one infected non-sibling, one infected sibling, and with at least one infected sibling and non-sibling, respectively. The study further implicates intersibling transmission as a pathway for H. pylori infection in childhood. In addition, living with a non-sibling in extended-family homes may contribute to infection risk but only in households with prevalent H. pylori infection within all family groups.

INTRODUCTION

Helicobacter pylori, one of the most common chronic infections of the developing world and a known cause of gastric ulcers and cancer, appears to be acquired predominantly in childhood [1]. In industrialized countries, transmission has decreased in recent decades and only 10–20% of adults less than 30 years of age are infected [2] compared to 50% of the population above 60 years of age [3, 4]. The different prevalence by age reflects a ‘cohort’ phenomenon, i.e. decreasing risk of acquiring new infection over time. This decreased risk, in turn, is thought to reflect improvements in household sanitation and hygiene throughout the 20th century. Although H. pylori acquisition has been clearly associated with hygienic factors – such as bed sharing, household crowding, and lack of indoor plumbing – the exact route of transmission remains unclear [5–7].

Epidemiological data have suggested mechanisms consistent with person-to-person transmission. The organism has been cultured from vomitus [8], human faeces and saliva [9–13]. In addition, the many studies documenting clustering of fingerprinted organisms have supported transmission within families, finding infection rates among children to be significantly higher when their mothers were infected [14–20]. In addition, children may also have different strains from their parents [20]. H. pylori infection in young
children has also been strongly linked to having infected siblings close in age [14, 17, 19, 21–24], and, particularly having infected older siblings [21].

The majority of evidence studying concordance of infection status among family members, particularly among children in the same age group, is consistent with person-to-person transmission, although exposure to common environmental sources has not been clearly eliminated [25, 26]. Concordance among family members, however, has also introduced the possible role of genetic susceptibility in influencing such transmission. Because studies of familial transmission include only participants sharing a strong genetic similarity, they are unable to clearly discriminate between the importance of close contact and that of genetics in person-to-person transmission. The only study to date that strongly suggested a genetic contribution to transmission identified higher concordance rates within the monozygotic than among dizygotic twin pairs [27].

A large ongoing trial of gastroenteritis transmission within Bay area households has provided us with an opportunity to explore concordance of \textit{H. pylori} infection within complex households containing related and unrelated individuals. In these households, we examined concordance of infection among children of extended families (defined as at least two nuclear families) living together. By comparing infection rates among siblings and non-siblings living in the same home, we hope to better dissect the contributions of shared living and familial relatedness to \textit{H. pylori} transmission.

\section*{METHODS}

\subsection*{Study design and population}

The Stanford Infection and Familial Transmission (SIFT) study is a prospective cohort study initiated in 1999 to assess \textit{H. pylori} infection in association with household episodes of gastroenteritis [28]. Households in the Santa Clara and San Mateo counties of Northern California were recruited by community outreach as well as through 15 cooperating community health-care programmes, including general medicine outpatient clinics, emergency rooms, and paediatric clinics, and two county environmental health surveillance programmes that receive reports of gastroenteritis from the community.

Cases of diarrhoea and/or vomiting of suspected infective aetiology (‘index case’) who presented at cooperating clinics were asked for permission to be contacted by study personnel. Those who consented to referral received a brief telephone interview to elicit study eligibility criteria, and, if appropriate, to schedule a home visit for which all interested household members were asked to be present. At each home visit informed consent was administered individually to all interested household members (‘participants’) and, for those consenting, a structured questionnaire regarding demographic characteristics, socioeconomic markers, risk factors for \textit{H. pylori} infection, household composition and family relationships was administered. Serum samples were collected for determination of \textit{H. pylori} infection status. All visits were conducted by research staff with phlebotomy certification and fluent in the preferred language of the household. On average, \(\sim 87\%\) of all known household members have been available for interviews and \(\sim 62\%\) of those over age 2 years have participated in laboratory testing as well as interviews.

A household member was defined as someone who spent at least 20 h per week in the home and shared kitchen and bathroom facilities. As part of the home interview, participants were asked to identify distinct biological and/or economic family units within the home and the relationship of each household member to the index case (parent/spouse, child/sibling, aunt/uncle, niece/nephew, grandparent, child-care worker, unknown). Within family units, individuals were further identified as parent, offspring, other relative, or unknown. Households contained from 2–21 members and up to eight distinct family units. For the purposes of our current analysis, only households containing at least two distinct family units, each with at least one child between the ages of 2 and 18 years tested for \textit{H. pylori}, were selected.

\subsection*{Laboratory methods}

\textit{H. pylori} infection was diagnosed by enzyme-linked immunosorbent assay (ELISA) for IgG using an assay validated in our laboratory, as previously described [29]. This assay uses high-molecular-weight, cell-associated proteins for five strains as antigen, including two Mexican strains. The sensitivity and specificity of this assay based on 77 persons from different ethnic groups were 94 and 91\% respectively. Borderline results (\(\sim 4\%\) of serology runs) were considered negative. Because serology may be less reliable in children <2 years of age, these children were excluded from the analysis.
The social structure of these 68 households is summarized in the Figure, with the family of the index case denoted as the index family, and other family units defined as related or unrelated to the index unit. Of a total 241 children included in the analysis, 100 (41%) were members of the index unit, and 141 were members of other family units within the home, including 115 (82%) who were related to the index case, and 26 (18%) who were unrelated. Of these 26 children, only four were unrelated to each other. The 241 children resided with a median of three other children (range 1–9), including a median of one sibling (range 0–4) and two non-siblings (range 1–9). A total of 96 children (40%) resided with other (predominately related) non-sibling children only. Compared to SIFT children residing in single-family homes with at least two children, SIFT children living in extended-family households (Table 1) were more likely to be of Hispanic ethnicity but did not differ significantly in age or gender.

Compared to all other households participating in the SIFT study (Table 2), households of the children included in the analysis were larger (median 10 vs. 6 members, \( P < 0.0001 \)), including twice as many children, of greater sleeping density (median 3 vs. 2 persons/bedroom, \( P < 0.0001 \)), of lower educational attainment (12 vs. 11 years, \( P = 0.02 \)), and more likely to report Spanish as the primary language of the household (93% vs. 71%, \( P < 0.0001 \)). In addition, 88% of these households had at least one adult (\( \geq 18 \) years of age) member who tested positive for infection, vs. 65% of other households (\( P < 0.0001 \)). Compared to all other extended-family homes, households included in the analysis were larger (median 10 vs. 7 members, \( P < 0.0001 \)), with greater sleeping density (median 2.5 vs. 2 persons/bedroom, \( P < 0.01 \)), and more likely to have at least one adult infection (88% vs. 73%, \( P < 0.01 \) respectively).
**H. pylori** prevalence was 24% overall among study children, compared with 15 and 19% in predominantly sibling-only households, extended-family homes containing at least two children in one family unit and single-family homes with at least two children respectively (Table 1, \(P = 0.06\)). Without accounting for infection status of other children, factors associated with **H. pylori** infection included age of the child (OR 2.0, 95% CI 1.4–2.8, per 5-year difference, \(P < 0.0001\)), and total number of other siblings in the home (OR 1.4, 95% CI 1.1–1.8, per one sibling, \(P = 0.02\)). The total number of other children in the household, total number of other non-siblings, gender, household educational attainment,

![Figure](https://doi.org/10.1017/S0950268805005352)

**Table 1.** Characteristics of children aged 2–18 years residing with at least one other child in extended-family and single-family homes

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Extended-family home</th>
<th>Single-family home</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Children in multiple family units* ((n=241))</td>
<td>Children in one-family unit ((n=284))</td>
</tr>
<tr>
<td>Age of children (years)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median (range)</td>
<td>7 (2–18)</td>
<td>7 (2–18)</td>
</tr>
<tr>
<td>Race</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hispanic</td>
<td>235 (98%)</td>
<td>271 (95%)</td>
</tr>
<tr>
<td>Gender</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>121 (50%)</td>
<td>140 (49%)</td>
</tr>
<tr>
<td>No. children</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median (range)</td>
<td>4 (2–10)</td>
<td>2 (2–4)†</td>
</tr>
<tr>
<td><strong>H. pylori</strong> infection</td>
<td>57 (24%)</td>
<td>44 (15%)</td>
</tr>
</tbody>
</table>

* Children included in the analysis.
† \(P < 0.05\) compared to 241 children included in the analysis.
‡ \(P = 0.06\) for comparison of infection rate across three groups.

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[https://doi.org/10.1017/S0950268805005352](https://doi.org/10.1017/S0950268805005352)
and sleeping density of members within a household were not associated with *H. pylori* infection.

Prevalence of infection rose significantly with number of infected other children in the household (Table 3), including 14, 20, 47 and 70% of children residing with no infected children, with one, two, or three or more infected children respectively ($\chi^2$ test for trend, *P* < 0.0001). Twelve ‘high transmission’ households – those homes with at least two infected children – accounted for nearly 65% of all *H. pylori* infections. Compared to children residing with no infected other children and controlling for age as well as household membership (Table 4), children residing with an infected sibling were significantly more likely to be infected themselves [adjusted odds ratio (AOR) 3·2, 95% CI 1·1–9·1], and children residing with at least one infected sibling and non-sibling were nine times more likely to be infected themselves (AOR 9·4, 95% CI 3·1–28·5). By contrast, children who resided only with infected non-siblings were not more likely

Table 3. *Frequency of H. pylori infection among 241 children in extended-family homes*

<table>
<thead>
<tr>
<th>No. of other children positive for <em>H. pylori</em></th>
<th>Total no.</th>
<th>Infected no. (%)</th>
<th>AOR (95% CI)*</th>
<th><em>P</em> value</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>139</td>
<td>20 (14·4%)</td>
<td>1·0</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>60</td>
<td>12 (20·0%)</td>
<td>1·4 (0·65–3·3)</td>
<td>0·37</td>
</tr>
<tr>
<td>2</td>
<td>19</td>
<td>9 (47·3%)</td>
<td>4·8 (1·7–13·8)</td>
<td>0·004</td>
</tr>
<tr>
<td>$\geq 3$</td>
<td>23</td>
<td>16 (70·0%)</td>
<td>11·3 (4·0–31·4)</td>
<td>&lt;0·0001</td>
</tr>
<tr>
<td>Total</td>
<td>241</td>
<td>57 (24%)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

AOR, Adjusted odds ratio; CI, confidence interval.

* Adjusted for child’s age.

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**Table 2. Household characteristics**

<table>
<thead>
<tr>
<th>Household characteristic</th>
<th>All households (n=1186)</th>
<th>Children in one family only (n=301)</th>
<th>Households included in analysis (n=68)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Household size</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median (range)</td>
<td>6 (2–21)</td>
<td>7 (3–21)</td>
<td>10 (5–20)*†</td>
</tr>
<tr>
<td>Adults</td>
<td>2 (0–15)</td>
<td>4 (1–12)</td>
<td>4 (1–15)</td>
</tr>
<tr>
<td>Children</td>
<td>2 (0–11)</td>
<td>3 (1–9)</td>
<td>4 (2–10)</td>
</tr>
<tr>
<td>No. members tested for <em>H. pylori</em> infection</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median (range)</td>
<td>3 (2–17)</td>
<td>4 (2–10)</td>
<td>7 (3–20)</td>
</tr>
<tr>
<td>No. family units‡</td>
<td>1 (1–8)</td>
<td>2 (2–8)</td>
<td>3 (2–8)</td>
</tr>
<tr>
<td>Sleeping density (persons/bedroom)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>2</td>
<td>2·5</td>
<td>3*†</td>
</tr>
<tr>
<td>Household income§</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;$30000/year (%)</td>
<td>59%</td>
<td>70%</td>
<td>69%</td>
</tr>
<tr>
<td>Highest educational level (years)</td>
<td>12</td>
<td>11</td>
<td>11*</td>
</tr>
<tr>
<td>Spanish-speaking household</td>
<td>856 (72%)</td>
<td>268 (89%)</td>
<td>63 (93%)*</td>
</tr>
<tr>
<td>$\geq 1$ adult <em>H. pylori</em> infection</td>
<td>771 (65%)</td>
<td>220 (73%)</td>
<td>59 (87%)*†</td>
</tr>
</tbody>
</table>

* *P* <0·05 vs. 1186 households not included in the analysis.
† *P* <0·05 vs. 301 households not included in the analysis.
‡ A family unit was defined as a separate biological and/or economic unit within the household.
§ Based on 71% response rate.
than children residing with no infected children to be infected (AOR 1·22, 95% CI 0·52–2·9). Thus, the effect of living with infected non-siblings was significant only in concert with the presence of an infected sibling.

Excluding the 96 children who did not reside with a sibling (and thus had no chance of sibling transmission), the findings remained significant (Table 5). The odds of infection given at least one infected sibling was, adjusting for age (AOR 4·4, 95% CI 1·4–13·9), and for residing with at least one infected sibling and one infected non-sibling (AOR 14·3, 95% CI 4·5–45·8). In the 12 high-transmission households, 70% of children had at least one infected sibling.

Results of the bootstrap simulations (Tables 4 and 5) were largely corroborative of these results; however, as expected, CIs were wider, and only the category of children residing with both an infected sibling and non-sibling remained significant after adjustment for age.

### DISCUSSION

*H. pylori* is one of the most common bacterial pathogens of humans, with considerable public health importance. Although evidence suggests infection is acquired primarily in early childhood [1], the epidemiology of childhood transmission has yet to be clearly elucidated. Despite well-established associations between *H. pylori* infection and crowded living conditions and strong evidence of familial concordance, knowledge of specific transmission pathways remains largely circumstantial. Studies delineating the social structures of high-risk households can help to clarify the social and familial pathways involved in *H. pylori* transmission.

In this study, we evaluated concordance of infection among children living in extended-family US immigrant homes. These predominately Hispanic households were crowded and complex, containing up to 10 related and unrelated children in up to eight different family units. Prevalence of *H. pylori* transmission among children

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**Table 4. Odds of *H. pylori* infection accounting for infection status and familial relationship of other children in the household**

<table>
<thead>
<tr>
<th>No. of infected siblings</th>
<th>No. of infected non-siblings</th>
<th>Total no.</th>
<th>Infected no. (%)</th>
<th>AOR (95% CI)*</th>
<th>P value</th>
<th>Bootstrap AOR (95% CI)†</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>139</td>
<td>0 (14%)</td>
<td>1·0</td>
<td></td>
<td>1·0</td>
</tr>
<tr>
<td>0</td>
<td>1+</td>
<td>56</td>
<td>11 (20%)</td>
<td>1·2 (0·52–2·9)</td>
<td>0·65</td>
<td>1·6 (0·7–3·7)</td>
</tr>
<tr>
<td>1+</td>
<td>0</td>
<td>23</td>
<td>10 (43%)</td>
<td>3·2 (1·14–9·12)</td>
<td>0·03</td>
<td>3·7 (0·8–16·4)</td>
</tr>
<tr>
<td>1+</td>
<td>1+</td>
<td>23</td>
<td>16 (70%)</td>
<td>9·4 (3·1–28·5)</td>
<td>&lt;0·0001</td>
<td>8·2 (1·4–30·0)</td>
</tr>
</tbody>
</table>

AOR, Adjusted odds ratio; CI, confidence interval.

* Random effects model accounting for household membership (adjusted for age).

† 1500 replicates sampling one child per household (adjusted for age).

**Table 5. Odds of *H. pylori* infection for only children living with at least one sibling and one non-sibling, accounting for infection status and familial relationship of other children in the household**

<table>
<thead>
<tr>
<th>No. of infected siblings</th>
<th>No. of infected non-siblings</th>
<th>Total no.</th>
<th>Infected no. (%)</th>
<th>AOR (95% CI)*</th>
<th>P value</th>
<th>Bootstrap AOR (95% CI)†</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>77</td>
<td>9 (12%)</td>
<td>1·0</td>
<td></td>
<td>1·0</td>
</tr>
<tr>
<td>0</td>
<td>1+</td>
<td>22</td>
<td>2 (9%)</td>
<td>0·68 (0·12–3·6)</td>
<td>0·64</td>
<td>1·5 (0·5–5·6)</td>
</tr>
<tr>
<td>1+</td>
<td>0</td>
<td>23</td>
<td>10 (43%)</td>
<td>4·4 (1·4–13·9)</td>
<td>0·01</td>
<td>5·9 (1·7–22·9)</td>
</tr>
<tr>
<td>1+</td>
<td>1+</td>
<td>23</td>
<td>16 (70%)</td>
<td>14·3 (4·5–45·8)</td>
<td>&lt;0·0001</td>
<td>16·4 (3·5–73·7)</td>
</tr>
</tbody>
</table>

AOR, Adjusted odds ratio; CI, confidence interval.

* Random effects model accounting for household membership (adjusted for age).

† 1000 replicates sampling one child per 45 extended-family households with at least one sibling and at least one non-sibling (adjusted for age).
infection among children was 24%, and higher than infection rates estimated for the general population of US children. While household characteristics, including total number of other children in a household, were not associated with *H. pylori* infection; residing with two or more other children who were infected was highly associated with infection. In addition, when evaluated separately, residing with an infected sibling was more influential than residing with an infected non-sibling. At the same time, a significant additive effect of sibling and non-sibling status was also observed, i.e. children residing in households including at least one infected sibling and non-sibling were three times more likely than children residing with an infected sibling only to be concordant for infection. This suggests that the presence of infected non-siblings can play an important role in transmission in households with multiple infections among children. Transmission from infected non-siblings may occur less frequently than inter-sibling transmission, but in concert with infected siblings, risk of transmission may be modified importantly in crowded homes.

The importance of intra-familial transmission in *H. pylori* infection has been previously established [5, 30, 31]. Several studies have reported strong vertical (parent–child) associations, particularly between mother and child, probably reflecting the mother's role as the primary caretaker in early childhood [14–20]. In crowded homes, however, it is not uncommon for children to share beds or for older children to participate in child-care responsibilities. Although not consistently as strong as vertical (parent–child) associations, more recent studies have suggested that close contact among siblings is also associated with household prevalence of infection among children [14, 17, 19, 21–24]. Among a paediatric Taiwanese population, children with an older seropositive sibling were nearly four times more likely than children with a seronegative older sibling to be infected [24]. A Brazilian study found that preschool children with a positive sibling had a nearly two-fold greater risk of infection [14]. In a study of rural Andean children aged 2–9 years, Goodman et al. reported a gradient effect, ranging from 1.7 to 7.1, for children with 1–4 positive siblings when compared to children with no infected siblings. The pattern of transmission was found strongest in those siblings close in age, particularly from older siblings to younger ones [21]. Our results expand on this research and also suggest that delineating familial relationships among cohabitating children in high-risk homes may yield further insight into the interaction of social and genetic risk factors for transmission.

*H. pylori* DNA strain analysis in families has also illustrated *H. pylori* transmission patterns among siblings as a predominant pathway [20, 30–33]. Over 80% of 36 families in a Swedish-based study had at least two siblings sharing *H. pylori* strain concordance. By comparison, mother and child concordance was only 50% in 18 families analysed [20]. A smaller Taiwanese study found strain concordance among the siblings in each of the five families studied whereas only two families revealed the same children sharing a strain of *H. pylori* with at least one of the parents [33]. Inter-generational strain segregation is consistent with the hypothesis of secular cohort effects in prevalence of *H. pylori* infection, including an important role for child-to-child transmission within households.

Siblings may play more closely together than non-siblings and may also have longer periods of cohabitation. Since non-siblings were predominately second-degree relatives and we did not collect information on duration of cohabitation, the higher concordance of infection among siblings than among non-siblings cannot be deemed in this study to implicate genetics in *H. pylori* transmission (although neither can we rule out this possibility). The lower risk conferred by living with infected non-siblings could suggest that higher thresholds of contact with infected, non-sibling children might be required for their presence to impact on incidence of infection. Of 42 children living in households with at least two other infected children, 83% were in an environment where at least half of the infected children were non-siblings. Therefore, our result was not simply masking the known significant effect infected siblings’ play in transmission.

Although intra-familial transmission has been accepted as a major factor for *H. pylori* infection the relative contributions of close interpersonal contact and genetic similarity to this pathway are not well understood. Our study is unique in that other studies directly comparing the effects of siblings and non-siblings in child-to-child transmission of *H. pylori* are not available. The impact on transmission by unrelated children is still unmeasured as *H. pylori* concordance studies of children in day-care centres are largely lacking. A study of Swedish schoolchildren aged 10–12 years found patterns of infection in children more closely correlated among their family
members rather than their peers [34]. The study had limitations as there were obvious differences in degree of contact between the two groups and that family members, not solely siblings, was one comparison group. This study provides an initial analysis towards better understanding underlying factors for intrafamilial transmission.

Some caveats about our study should be noted. The sampling design elicited households that are not typical of US homes, and even somewhat atypical of settled immigrant homes. Even within the context of our underlying cohort study, which includes many low-income Hispanic families, these homes were larger, with higher sleeping densities, of lower income, and of lower educational attainment – all factors linked with increased risk of *H. pylori* infection [5–7, 35–38]. Nonetheless, prevalence of *H. pylori* infection among children (24%) was not dissimilar to estimates for other US immigrant children [39–42]. Further, extended-family living arrangements are often a matter of convenience, and the composition of these homes can change frequently. Some arrangements in our study may have been temporary, and others more permanent, thus affecting the amount of time children are in close living situations with each other.

Although exposure to infected siblings appeared to play a significant role in explaining infection rates among children in these crowded households, 35 of the 68 households included in the analysis had no infected children and, ultimately, did not contribute to distinguishing the risk of transmission between sibling and non-sibling household contacts. Non-siblings in this study were usually second-degree relatives such as cousins, and the number of unrelated non-siblings was too small to distinguish these sources of non-sibling exposure. Only four children in 12 ‘high-transmission’ homes were found to be unrelated to the index child. Thus, it is unlikely that exposure to infected, unrelated non-siblings played a significant role in the results. While we cannot distinguish genetic from social explanations, sibling exposure was an independent predictor of infection when adjusted for sheer number of children, age, and other environmental factors such as household size or sleeping density.

Household concordance analysis involves complex clustering as well as redundancy effects. For this reason, we tested our results in a bootstrap simulation using replicate random samples of one child per household. While this analysis did widen CIs, conclusions were similar to the basic analysis for all households and those only restricted to having both siblings and non-siblings. Bootstrapping does not compensate for limitations of sample size. Although observed effect sizes, particularly for exposure to an infected sibling, were substantial, the modelling of interactions is a cautionary exercise with small numbers of children in some categories. In general, more complex statistical models are needed to delineate transmission problems where multiple interactions are of interest. Emerging methodological work in the area of network analysis may prove useful.

Our study further confirms child–child transmission as a likely pathway for *H. pylori* infection. That concordance was more common between siblings than non-siblings sharing a household may suggest a role for genetic similarity in transmission or that the contact required to transmit *H. pylori* must be at a high threshold to induce transmission. Further studies that are able to more fully account for duration of cohabitation and degree of contact can offer further clarification.

**ACKNOWLEDGEMENTS**

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**DECLARATION OF INTEREST**

None.

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