Identifying human encounters that shape the transmission of *Streptococcus pneumoniae* and other acute respiratory infections

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ABSTRACT

Although patterns of social contacts are believed to be an important determinant of infectious disease transmission, it remains unclear how the frequency and nature of human interactions shape an individual’s risk of infection. We analysed data on daily social encounters individually matched to data on *S. pneumoniae* carriage and acute respiratory symptoms (ARS), from 566 individuals who took part in a survey in South-West Uganda. We found that the frequency of physical (i.e. skin-to-skin), long (≥1 h) and household contacts – which capture some measure of close (i.e. relatively intimate) contact – was higher among pneumococcal carriers than non-carriers, and among people with ARS compared to those without, irrespective of their age. With each additional physical encounter the age-adjusted risk of carriage and ARS increased by 6% (95%CI 2–9%) and 7% (2–13%) respectively. In contrast, the number of casual contacts (<5 min long) was not associated with either pneumococcal carriage or ARS. A detailed analysis by age of contacts showed that the number of close contacts with young children (<5 years) was particularly higher among older children and adult carriers than non-carriers, while the higher number of contacts among people suffering from ARS was more homogeneous across contacts of all ages.

Our findings provide key evidence that the frequency of close interpersonal contact is important for transmission of respiratory infections, but not that of casual contacts. Those results are essential for both improving disease prevention and control efforts as well as informing research on infectious disease dynamics and transmission models, and more studies should be undertaken to further validate our results.

1. Introduction

The transmission of respiratory infections is likely to depend on the frequency and age structure of human social contacts, as well as other factors including pre-existing immunity from prior infection or vaccination (Smieszek et al., 2012; Mossong et al., 2008; Eubank et al., 2004). To understand the dynamics of such infections, studies to quantify social mixing patterns have been conducted in various settings, under the assumption that self-reported encounters reflect transmission probabilities of pathogens transmitted through close contact (Mossong et al., 2008; Wallinga et al., 2006; Horby et al., 2011; Grijalva et al., 2015; Johnstone-Robertson et al., 2011; Kiti et al., 2014; Fu et al., 2012; Read et al., 2014; Edmunds et al., 1997). Combined with disease transmission models, these data are increasingly being used to inform infection control policies (Halloran et al., 2008; Baguelin et al., 2013). There is evidence from population-based models that using self-reported social mixing patterns in such models can reproduce observed aggregated data for chickenpox (Santermans et al., 2015), mumps (Wallinga et al., 2006), parvovirus (Melegaro et al., 2011), influenza (Wallinga et al., 2006; Kwok et al., 2014; Kucharski et al., 2014) and whooping cough (Rohani et al., 2010). Moreover, it has been suggested that age-stratified social mixing patterns can capture individual
influence risk, as measured by a four-fold rise in neutralization titres over the course of an epidemic (Kucharski et al., 2014; Rohani et al., 2010). However, as yet no study has directly linked individual’s contact patterns with their risk of infection at the same point in time.

To establish how social behaviour shapes individual-level infection, we explored whether the frequency and duration of different types of social encounters were associated with an individual’s risk of respiratory infection, using nasopharyngeal (NP) carriage of Streptococcus pneumoniae (the pneumococcus) and self-reported acute respiratory symptoms (ARS) as endpoints.

Streptococcus pneumoniae is one of the main causes of pneumonia and sepsis globally (O’Brien et al., 2009), disproportionally so in low-income settings (O’Brien et al., 2009; Black et al., 2010; Cohen et al., 2012). Colonization of the nasopharynx is a precondition to disease, and the main source of human-to-human transmission. Given that most episodes of carriage remain asymptomatic, social behaviour is unlikely to change as a result of carriage, making pneumococcal carriage a more suitable endpoint than symptomatic illness to explore the association between social behaviour and infection risk, given that people tend to limit their contacts during symptomatic illness (Van Kerckhove et al., 2010). However, as yet no study has directly linked individual contact lasting for 20 minutes (NP swabs) was taken from each participant. NP samples were collected, transported and analysed as per WHO guidelines (Satzke et al., 2013). NP swabs (flocked nylon swabs, COPAN, Italy) were inoculated in a skim milk tryptone-glucose-glycerol (STGG) medium, transported in cool boxes and frozen at the research laboratory at −20 °C within 8 h of collection. Specimens were inoculated onto a selective agar plate of 5 mg/L gentamicin-Columbia agar with 5% sheep blood and incubated at 37 °C in 5% CO2 atmosphere overnight. Pneumococcal identification was based on optochin susceptibility testing of all alpha-hemolytic colonies and bile solubility testing in case of intermediate susceptibility to optochin. Further details are described elsewhere (Nackers et al., 2017).

2.2. Statistical analysis

We first performed descriptive analyses, with age-specific probability weights to account for different inclusion probabilities by age at the design stage, and adjusted for the clustering by village through the use of clustered ‘sandwich’ variance estimators to account for possible correlation within each of the sixty clusters (Rogers, 1993). We explored social contact patterns within and between age groups through contact matrices. In such matrices we report the mean number of ordinary contacts of participants in age group i with contacts in age group j (mi,j), adjusted for reciprocity, as in Melegaro et al. (Melegaro et al., 2011).

We then analysed whether and how the frequency distribution of contacts was associated with pneumococcal carriage or self-reported ARS.

We modelled the effect of ‘ordinary’ contacts (defined as contacts ≥5 min long) on carriage or respiratory symptoms as a function of contact frequency, through a log-binomial model with a robust variance estimator, and inclusion probability weights by age group. We treated contacts as continuous variables, but assessed departure from linearity through likelihood ratio tests and model comparisons of Bayesian Information Criterion (BIC). In multivariable analysis, we considered for inclusion any covariate significantly associated with the outcome at P < 0.10 in univariable analysis. Model improvement was considered for any decrease in the BIC. Further details are provided in the Supporting Information.

Next, we used the same analytical approach to assess whether carriage and ARS were associated with the level of self-reported ‘casual’ contacts (i.e. < 5 min long).

Finally, we explored differences in the social mixing matrices by status of pneumococcal carriage and ARS. To do so, we computed the ratio of the mean number of reported contacts by participants j with contacts i among carriers compared to non-carriers (Rij C) or symptomatic compared to asymptomatic individuals (Rij S), such that $R_{ij}^{C} = \frac{m_{ij}^{C}}{N_{i}^{C}}$ and $R_{ij}^{S} = \frac{m_{ij}^{S}}{N_{i}^{S}}$, where $C = \text{carriers}, R = \text{participants with ARS}$ and $N_{i}^{C}$ is the total number of participants. Estimates were not adjusted for reciprocity given that subpopulations were not closed (e.g. contacts of carriers may not be carriers), and that our aim was to compare reported number of contacts rather than calculate a contact matrix. The uncertainty in reported values and ratios was obtained through resampling techniques, drawing random samples from each $m_{ij}$, with the number of draws equal to the study population in each age group j. Ratios and uncertainty around them was obtained from the ratio of bootstrapped matrices.

The same approach as described above was used to compute the differences in $m_{ij}$ with $D_{ij}^{C} = m_{ij}^{C} - m_{ij}^{C-\text{C}}$ and $D_{ij}^{S} = m_{ij}^{S} - m_{ij}^{S-\text{C}}$. All analyses were performed in Stata 13.1 IC and R version 3.2.

Next, participants were asked about respiratory symptoms experienced in the two weeks prior to the survey, including any of the following: cough, runny nose, sneezing, sore throat, difficulty breathing.

Finally, after the interview was completed, a nasopharyngeal swab was taken from each participant. NP samples were collected, transported and analysed as per WHO guidelines (Satzke et al., 2013). NP swabs (flocked nylon swabs, COPAN, Italy) were inoculated in a skim milk tryptone-glucose-glycerol (STGG) medium, transported in cool boxes and frozen at the research laboratory at −20 °C within 8 h of collection. Specimens were inoculated onto a selective agar plate of 5 mg/L gentamicin-Columbia agar with 5% sheep blood and incubated at 37 °C in 5% CO2 atmosphere overnight. Pneumococcal identification was based on optochin susceptibility testing of all alpha-hemolytic colonies and bile solubility testing in case of intermediate susceptibility to optochin. Further details are described elsewhere (Nackers et al., 2017).

2. Materials and methods

2.1. Data collection

The study was conducted in Sheema North Sub-District (Sheema district, South-West Uganda) between January and March 2014. Sixty clusters were randomly selected from the 215 villages and two small district towns (Kabwohe and Tendero) in the study area, proportionally to the population size of each village and town. In each cluster, from 29 or 30 individuals randomly sampled from different households for inclusion in a nasopharyngeal carriage study (Nackers et al., 2017), a subset of 11 or 12 individuals were selected to answer questions about their social contacts and their history of respiratory illness in the last two weeks, in addition to having a nasopharyngeal swab taken. A household was defined as the group of individuals living under the same roof and sharing the same kitchen on a daily basis. The full descriptive results from the social contact study are available elsewhere (le Polain de Waroux et al., 2018).

For the social contact questionnaire, participants were first asked to list all the individuals with whom they had a two-way conversational contact lasting for ≥5 min during a period of approximately 24 h prior to the survey day (from wake up the previous day until wake up on the survey day). Such encounters were defined as ‘ordinary contacts’. For each reported ordinary contact, participants (or their parent/guardian) were asked to estimate the contact’s age (or estimated age), how long the encounter lasted for and whether it involved skin-to-skin touch or utensils passed from mouth to mouth (either of those defined as casual contacts). For very short social encounters (< 5 min), which were defined as ‘casual contacts’ (e.g. seeing someone on the way, encounter in a shop etc.), participants were asked to estimate the number of encounters based on pre-defined categories (< 10 contacts, 10–19 contacts, 20–29 contacts, ≥30 contacts), but not to provide further details about each contact.
2.3. Ethics

Approval was obtained from the Ethical review boards of the London School of Hygiene and Tropical Medicine, Médecins Sans Frontières, the Faculty of Medicine Research & Ethics Committee of the Mbarara University of Science and Technology, the Institutional Ethical Review Board of the MUST, and the Uganda National Council for Science and Technology.

3. Results

3.1. Study population

Data on social contacts, carriage and ARS were available from 566 participants. Full descriptive results from the nasopharyngeal carriage study and the social contact study are available elsewhere (Nackers et al., 2017; le Polain de Waroux et al., 2018).

Briefly, participant’s age spanned across age groups and the sex distribution was reasonably balanced (58% female). The majority (98%) of children aged 6–15 years attended school or college. More than a third of all adults (36%) worked in agriculture and 22% were homemakers/housewives.

On average, people made seven ‘ordinary’ contacts (defined as contacts ≥5 min long), ranging from 0 to 25, the majority of which were physical (i.e. ‘skin-to-skin’ ordinary contact or indirect physical contact through utensils passed from mouth-to-mouth). There was no evidence that the average number of contacts differed by weekday or between weekdays and weekends (P = 0.623). Children aged 5–9 years reported most contacts and children < 5 years the fewest (Fig. 1). The majority of contacts made by children were physical. The proportion of non-physical contacts, contacts outside the household and contacts of short (< 1 h) duration was higher among teenagers and adults than younger children.

The most intense mixing tended to be between individuals of the same age group (i.e. assortative mixing), but there was also substantial mixing between age groups. Contact from and with children aged < 10 years involved proportionally more physical touch than contacts between older children and adults (Fig. 2A and B). There was no difference in contact patterns by sex, for all types of contacts considered.

Four hundred and ninety (87%) participants estimated how many casual contacts (i.e. < 5 min in duration) they had the day before the survey. Over a third (36%) of these reported 10 ore more casual contacts, and 11% reported more than 20 casual contacts. Among the 13% who did not know how many casual contacts they had, there were proportionally more children, with over half (56%) under 10 years.

The mean number of reported ‘ordinary’ contacts was higher among individuals reporting ≥10 casual contacts than those reporting fewer than 10 contacts (mean 8.9 vs 5.9, P < 0.001), as well as among the 78 individuals who did not know how many casual contacts they had (mean 8.8, P < 0.001).

The prevalence of pneumococcal carriage was strongly age dependent, decreasing from 75% in children < 5 years to 46% among 5–9 year olds, 17% in 10–19 year olds, and further decreasing to 8% and 7% among 20–39 years and ≥40 years old respectively (Fig. 2C).

Fig. 1. Number of reported contacts by age group and type of contact.
Legend: Boxplots showing the median (horizontal line), interquartile range (boxes) and 95% confidence interval around the mean (whiskers). The outliers are shown as bubbles above or under the whiskers.
Overall, 72 (13%) people reported having suffered from ARS in the two weeks prior to the survey. The prevalence of ARS varied much less with age than that of carriage (Fig. 2D), ranging from 20% among 5 to 9 year olds to 8% among ≥ 40 years old. There was no sex difference in the prevalence of carriage (age-adjusted RR for males 0.97, P = 0.814) or ARS (age-adjusted RR for males 1.40, P = 0.134).

The Table 1 below provides the crude risk ratios by age for prevalence of carriage, and acute respiratory symptoms (ARS).

<table>
<thead>
<tr>
<th>Age group</th>
<th>Prevalence</th>
<th>Risk ratio (95%CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NP Carriage</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 5 years</td>
<td>75.1%</td>
<td>1.00</td>
</tr>
<tr>
<td>5 – 9 years</td>
<td>45.9%</td>
<td>0.61 (0.48 ; 0.77)</td>
</tr>
<tr>
<td>10 – 19 years</td>
<td>19.3%</td>
<td>0.24 (0.16 ; 0.37)</td>
</tr>
<tr>
<td>20 – 39 years</td>
<td>6.1%</td>
<td>0.08 (0.04 ; 0.15)</td>
</tr>
<tr>
<td>≥40 years</td>
<td>4.8%</td>
<td>0.06 (0.03 ; 0.13)</td>
</tr>
<tr>
<td>ARS</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 5 years</td>
<td>17.4%</td>
<td>1.00</td>
</tr>
<tr>
<td>5 – 9 years</td>
<td>20.3%</td>
<td>1.16 (0.63 ; 2.12)</td>
</tr>
<tr>
<td>10 – 19 years</td>
<td>10.3%</td>
<td>0.68 (0.32 ; 1.46)</td>
</tr>
<tr>
<td>20 – 39 years</td>
<td>9.9%</td>
<td>0.56 (0.28 ; 1.15)</td>
</tr>
<tr>
<td>≥40 years</td>
<td>7.5%</td>
<td>0.43 (0.22 ; 0.85)</td>
</tr>
</tbody>
</table>

3.2. Social contacts as a risk for pneumococcal carriage or ARS

Overall, the mean number of contacts among carriers was significantly higher than non-carriers, and this observation was consistent across age groups, although most differences were not statistically significant due to small numbers (Fig. 3). In particular, carriers had more physical contacts (Fig. 3A). This pattern was also consistent for ARS, with the exception of 5–9 year olds in whom the mean number of contacts among individuals with ARS was lower for physical contacts (Fig. 3B).

In univariable analysis, the risk of carriage increased with all contacts, household contacts, contacts ≥ 1 h long and physical contacts. The latter had the largest effect size, with a 13% increased risk for each additional contact reported by participants (Table 2). Physical contacts and contacts ≥ 1 h were strongly correlated (R = 0.76), particularly among children < 5 years (R = 0.85) and children aged 5–9 years (R = 0.87), hence their effect could not be disentangled, whereas the correlation between physical contacts and household contacts was moderate (R = 0.61, ranging from 0.47 to 0.70 between age groups).

After age adjustment, physical contacts or contacts ≥ 1 h remained most significantly associated with carriage, with a 6% increased risk (95%CI 2–10%) for each unit increase in the number of reported contacts (Table 2). We found good evidence that the number of household contacts increased the risk as well (Table 2). There was no confounding effect by other covariates and models were therefore only adjusted for age (Supporting Information).

An increase in physical, household and long (≥ 1 h) contacts, were also associated with an increased risk of ARS in univariable analysis. There was little or no evidence of a confounding effect of age, given the more constant prevalence of ARS across age groups (Table 2). Unlike pneumococcal carriage, however, the relative risk was more constant across types of contacts, and household contacts rather than physical contacts were most strongly associated with a risk for ARS, with a risk increase of 9% (1–18%) for each additional reported contact.

Next, we analysed whether the number of casual contacts (i.e. contacts lasting < 5 min) was associated with either pneumococcal carriage or ARS. We found no evidence that the prevalence of
pneumococcal carriage or the risk of ARS were associated with reporting higher levels of casual contacts, as shown in Table 3. Given the small numbers of individuals reporting ≥30 casual contacts, we pooled the 56 individuals reporting ≥20 casual contacts into one category.

In univariable analysis the risk of pneumococcal carriage was higher in the group of 78 participants who did not know how many casual contacts they may have had. This was due to the higher proportion of children < 5 years and aged 5–9 years in that group, however, after age-adjustment, there was no evidence that the risk of pneumococcal carriage was higher in that group. Similarly, a higher number of casual contacts was associated with increased risk of ARS, and with no confounding effect of age or other variables on the estimates.

We found no confounding effect on casual contacts on the relative risk of carriage or ARS as a function of the frequency of reported ‘ordinary’ contacts. Finally, we explored the characteristics of age-specific mixing patterns by ARS or carriage status in greater detail, with a focus on physical contacts and household contacts.

We computed the ratio of the mean number of contacts within and between age groups among carriers compared to non-carriers, and among individuals with ARS compared to those without. The average number of physical encounters within and between age groups tended to be higher for carriers than non-carriers in most instances, albeit with substantial uncertainty owing to small numbers (Fig. 4).

Similar findings were seen for ARS, however, unlike for pneumococcal carriage, symptomatic adults did not have more contacts with

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**Table 2**

Relative risk of *S. pneumoniae* carriage and ARS by frequency of contact.

<table>
<thead>
<tr>
<th>Contact types</th>
<th>Mean contacts</th>
<th>Crude RR (95%CI)</th>
<th>Age-adjusted RR (95%CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NP carriage</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All contacts</td>
<td>8.09</td>
<td>1.04 (1.00; 1.07)</td>
<td>1.03 (1.00; 1.07)</td>
</tr>
<tr>
<td>Physical contacts</td>
<td>6.83</td>
<td>1.13 (1.09; 1.17)</td>
<td>1.06 (1.02; 1.09)</td>
</tr>
<tr>
<td>Non-physical contacts</td>
<td>1.25</td>
<td>0.78 (0.72; 0.86)</td>
<td>0.97 (0.91; 1.05)</td>
</tr>
<tr>
<td>Household contacts</td>
<td>5.64</td>
<td>1.09 (1.03; 1.14)</td>
<td>1.04 (0.99; 1.10)</td>
</tr>
<tr>
<td>Non-household contacts</td>
<td>2.22</td>
<td>0.98 (0.93; 1.04)</td>
<td>1.03 (0.98; 1.09)</td>
</tr>
<tr>
<td>Contacts ≥ 1 h</td>
<td>7.56</td>
<td>1.08 (1.05; 1.12)</td>
<td>1.06 (1.02; 1.10)</td>
</tr>
<tr>
<td>Contacts &lt; 1 h</td>
<td>2.38</td>
<td>0.81 (0.73; 0.92)</td>
<td>0.94 (0.85; 1.03)</td>
</tr>
<tr>
<td>ARS</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All contacts</td>
<td>8.18</td>
<td>1.07 (1.02; 1.12)</td>
<td>1.07 (1.02; 1.13)</td>
</tr>
<tr>
<td>Physical contacts</td>
<td>6.10</td>
<td>1.09 (1.03; 1.15)</td>
<td>1.07 (1.00; 1.14)</td>
</tr>
<tr>
<td>Non-physical contacts</td>
<td>2.07</td>
<td>1.00 (0.89; 1.10)</td>
<td>1.06 (0.95; 1.18)</td>
</tr>
<tr>
<td>Household contacts</td>
<td>5.70</td>
<td>1.10 (1.02; 1.19)</td>
<td>1.09 (1.01; 1.18)</td>
</tr>
<tr>
<td>Non-household contacts</td>
<td>2.47</td>
<td>1.04 (0.98; 1.11)</td>
<td>1.05 (0.98; 1.13)</td>
</tr>
<tr>
<td>Contacts ≥ 1 h</td>
<td>7.23</td>
<td>1.07 (1.01; 1.13)</td>
<td>1.07 (1.00; 1.14)</td>
</tr>
<tr>
<td>Contacts &lt; 1 h</td>
<td>3.14</td>
<td>1.05 (0.96; 1.15)</td>
<td>1.08 (0.99; 1.18)</td>
</tr>
</tbody>
</table>

**RR in bold**: Associations significant at P < 0.05. RR = Risk Ratio. CI = Confidence Interval. NP+ = pneumococcal carrier, NP- = non-carrier, ARS+ = with ARS, ARS- = without ARS.
young children than asymptomatic ones. Results based on absolute differences in the mean number of contacts rather than relative means are displayed in Fig. S1 (Supporting Information), showing the similar associations than the reported ratios, but providing a quantified difference in mean number of contacts instead.

4. Discussion

Our study provided a unique opportunity to explore whether and how social contact patterns are associated with someone’s risk of acute respiratory infection in a mostly rural East African setting. Our results show that people who tend to have more frequent close contacts are more likely to be pneumococcal carriers or to report acute respiratory symptoms, irrespective of their age. In contrast, we found that less intimate or short casual contacts were not associated with someone’s infection risk, suggesting that social contacts important for transmission are close interpersonal encounters.

Existing evidence on the association between contact patterns and risk of infection is mostly ‘ecological’, with very few studies based on individual-level data. Using data of influenza A/H1N1 seroconversion in Hong Kong coupled to social contact data in the same population, Kwok et al. (Kwok et al., 2014) showed that age rather than social contact patterns were the main driver of the individual risk of infection in that setting, and further work by Kucharski et al. (Kucharski et al., 2014) on the same data further supported the finding that someone’s risk of infection is related to the average mixing pattern within their age group rather than their reported number of contacts. However, the validation of the ‘social contact hypothesis’ with such data remains difficult, due to challenges in accounting for pre-existing levels of protective immunity in the population, assumptions around stability of individual behaviour is unlikely to account for a large proportion of ARS cases (Berkeley et al., 2010), the deflation was used as a non-specific proxy for acute respiratory infection and may have captured other infectious and non-infectious conditions, including chronic conditions. Notwithstanding such caveats, results for ARS were similar to those for nasopharyngeal carriage, and under the hypothesis that a significant proportion of individuals presenting with acute respiratory symptoms have an acute respiratory infection, our findings support the hypothesis that the number of close interpersonal contacts, and particularly household contacts, play a role in the transmission of acute respiratory pathogens.

One of the striking features of our analysis of the relative number of mean contacts between age groups is that adults colonised with *S. pneumoniae* reported more than twice as many close contacts with children under five than non-carriers, and to a lesser extent also more proportionally more contact with adults. The higher number of contacts with children is in agreement with observational and modelling studies showing that pneumococcal carriage risk increases with household size and with the number of children < 5 years in the household (Farida et al., 2014). It also supports the finding that carriage acquisition in adults occurs mostly within the household as a result of contact with young children who are drivers of infection (Melegaro et al., 2004). In contrast, however, we found that adults with ARS reported an equal or lower number of contacts with children < 5 years than asymptomatic adults. This likely reflects the epidemiological differences between

![Fig. 4. Ratio of the mean reported number of physical contacts between pneumococcal carriers and non-carriers (panel A) and individuals with ARS compared to those without ARS (Panel B).](image-url)
carriage and ARS, given the very high prevalence of carriage among < 5 year olds and the much less marked difference in age-specific prevalence of ARS across age groups, in addition to other potential factors such as acquired immunity among adults more frequently exposed to very young children. Some of the specific details of the contact patterns and differences between S. pneumoniae and ARS are harder to interpret, due in part to wide statistical uncertainty, particularly among adults in whom the number of carriers or symptomatic individuals is small, as well as the multiple underlying causes of ARS as mentioned earlier. However, overall the results from our analysis support the general finding that increased close contacts are associated with higher risk of ARS or pneumococcal carriage.

There is still much debate about how respiratory pathogens are transmitted from person to person; whether through close direct or indirect physical contact, through large droplet transmission at close range (< 1 m), or through aerosolized particles floating over longer distances, particularly in poorly ventilated indoor settings (Brankston et al., 2007; Larson and Liverman, 2011). It is likely that many pathogens can be transmitted through a combination of these routes. Yet the contribution of each mechanism remains uncertain (Brankston et al., 2007; Larson and Liverman, 2011). It is generally assumed that the main transmission route of S. pneumoniae is through direct contact (Ferreira et al., 2013), as well as indirectly through shared glasses or bottles (Levine et al., 2012). Analogously, for other colonising bacteria such as N. meningitidis, close contact and intimate kissing are known risk factors among teenagers and young adults (Maclennan et al., 2006). Similarly, it is believed that influenza and other respiratory viruses are primarily transmitted through direct contact or contact with large droplet transmission at close range rather than aerosolized particles (Brankston et al., 2007).

Although our objective was not to demonstrate transmission, our findings strongly support that direct close interpersonal contact is an important mode of transmission for pathogenic bacteria of the nasopharynx as well as respiratory viruses, and strengthens the scientific evidence for public health measures based upon these assumptions, such as hand washing campaigns or chemoprophylaxis of close contacts of cases of meningococcal meningitis to name a few. The strong association of ARS with household and other close contacts, independently from physical touch, might suggest that other mechanisms such as indirect transmission through fomites or aerosol transmission may play a role. Elucidating the contribution of such factors in this context would be an important question for future research. In addition, given that our results are based on a single study, more work should be undertaken to corroborate our findings in other settings.

Our findings have also several implications for infectious disease research. Contact structures are central to transmission models, and appropriate assumptions about what type of contact drives infectious disease transmission are essential. Our results suggest that the parameterisation of transmission models of S. pneumoniae and similar pathogens using mixing matrices based on physical or another measure of close interpersonal contact would more likely capture relevant contact patterns than those based on any type of social encounter. This has also implications for the design of contact studies, particularly in low-income settings given the scarcity of published data currently available and the need to collect additional data from many more settings (Johnstone-Robertson et al., 2011; Kiti et al., 2014). In contrast to diary-based approaches that have been used by many (Mossong et al., 2008; Wallinga et al., 2006; Johnstone-Robertson et al., 2011; Kiti et al., 2014), the study design here was relatively simple and only involved a single face-to-face interview. A drawback of such retrospective approach is the lack of detailed information about very short (i.e. ‘casual’) contacts, as such information was deemed unreliable (Le Polain de Waroux et al., 2018) – and this is further supported by evidence that short contacts tend to be inconsistently recorded even in prospective diary-based approaches (Smieszek et al., 2012). However, assuming that short contacts do not account for much of the transmission, as our study suggests, a more simple retrospective design is a potential attractive option in other settings where contact data are lacking, and in which data collection through more comprehensive diary-based approaches may be difficult to implement.

There are some additional limitations to our study. We were unable to explore other potential confounding factors, such as bed share, ventilation, indoor smoke or hand washing (Tumwesigire and Barton, 1995), and the contribution of such factors should be explored further. Among other important factors, it is worth mentioning that our results were not impacted by pneumococcal vaccination, as the vaccine had not been introduced in the study area at the time. Moreover, it remains unclear to what extent our results can be generalised to any acute respiratory infection. For example, factors such as ventilation and airflow may be of particular importance for aerosolized transmission of pathogens such as mycobacteria (Johnstone-Robertson et al., 2011), compared to influenza or S. pneumoniae (Brankston et al., 2007; Larson and Liverman, 2011), and whether household or physical contacts reflect contact patterns important for aerosolized transmission remains uncertain (Johnstone-Robertson et al., 2011). Finally, while we found a strong association at the individual level, our study does not demonstrate causality.

However, in the absence of detailed longitudinal data on acquisition events between every single contact—which would be challenging and possibly unrealistic to obtain—our findings provide consistent evidence of a ‘dose-response’ association at the individual level between close social encounters and acquisition risk for respiratory pathogens, and therefore provides support both for the social contact hypothesis, and for research and policy work based upon this hypothesis. Additional studies in other settings should be undertaken to further validate our results.

Competing interests

The authors declare they have no conflict of interest.

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Appendix A. Supplementary data

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References
