Does infection with *Chlamydia trachomatis* induce long-lasting partial immunity? Insights from mathematical modelling

Ryosuke Omori,1,2,3 Hiam Chemaitelly,3 Christian L Althaus,4 Laith J Abu-Raddad3,5,6

ABSTRACT

**Objectives** To explore whether existence of long-lasting partial immunity against reinfection with *Chlamydia trachomatis* is necessary to explain *C. trachomatis* prevalence patterns by age and sexual risk, and to provide a plausible estimate for the effect size, defined here as a reduction in susceptibility to reinfection.

**Methods** A population-based mathematical model was constructed to describe *C. trachomatis* natural history and transmission dynamics by age and sexual risk. The model was parameterised using natural history, and epidemiological and sexual behaviour data, and applied for UK and US data. Sensitivity analyses were conducted to assess the robustness of predictions to variations in model structure and to examine the impact of alternative assumptions for the mechanism underlying partial immunity.

**Results** Partial immunity against reinfection was found necessary to explain observed *C. trachomatis* prevalence patterns by age and sexual risk. The reduction in susceptibility to reinfection was estimated at 93% using UK data (95% uncertainty interval (UI)=88%–97%) and at 67% using US data (95% UI=24%–88%). The model-structure sensitivity analyses affirmed model predictions. The immunity-mechanism sensitivity analyses suggested a mechanism of susceptibility reduction against reinfection or a mechanism of infectious-period duration reduction upon reinfection.

**Conclusions** A strong long-lasting partial immunity against *C. trachomatis* reinfection should be present to explain observed prevalence patterns. The mechanism of immunity could be either a reduction in susceptibility to reinfection or a reduction in duration of infection on reinfection. *C. trachomatis* infection appears to naturally elicit a strong long-lasting immune response, supporting the concept of vaccine development.

INTRODUCTION

*Chlamydia trachomatis* is a common bacterial STI.1,2 Untreated *C. trachomatis* infection is associated with pelvic inflammatory disease, infertility and ectopic pregnancy among women, and causes urethritis among men.3,4 Most genital *C. trachomatis* infections are asymptomatic.2 The WHO estimates that 4.2% of the population aged 15–49 years is infected.1

An understanding of *C. trachomatis* natural history and its implications on transmission dynamics is needed to guide public health strategies for its control.3 Its infection is characterised by a prolonged interval between infection acquisition and spontaneous cessation of shedding.4 In a study among 82 women followed for >5 years, 46% of infections were persistent at 1 year, 18% at 2 years and 6% at 4 years.5 *C. trachomatis* immunity appears to take at least months to acquire.6,7

*C. trachomatis* control has focused on early detection and treatment,8,9 but the effectiveness of such strategy is subject to debate.9,10 Despite widespread testing of young asymptomatic adults and treatment of infected individuals,10–13 *C. trachomatis* prevalence in the targeted age group has not changed markedly.14,15 It is argued that *C. trachomatis* early detection and treatment hinder the development of an adequate immune response, thereby increasing susceptibility to reinfection at the individual level, reducing herd immunity at the population level and counteracting reductions in prevalence.8,16

Since *C. trachomatis* infection is curable, there are ethical constraints to conducting studies that directly assess existence/strength of a protective immunity against reinfection.17 There is, nonetheless, evidence from animal models suggesting short-term complete immunity and long-term partial immunity.18 Partial immunity can take different forms, such as reduction in susceptibility to reinfection, reduction in infectious-period duration on reinfection and/or reduction in infectiousness upon reinfection (lower organism load). Indirect evidence from human studies supports the concept of protective partial immunity.17,19 The evidence includes rapidly declining prevalence with age, similarity in prevalence despite high variability in sexual risk, lower organism load with age and repeat infection, reduction in concordance rate in couples with age, and an apparent treatment attenuation of protective immunity.17,20,21

Against this background, we present a novel hypothesis-generation approach to assess the existence, and to provide a plausible effect-size estimate, of the partial immunity against reinfection. The approach rests on the concept that natural history effects at the individual level have manifestations at the population level. Starting from population-level measures, we used mathematical modelling to provide the link between these measures and natural history effects.
Although our approach uses an indirect method, mathematical modelling, its strength lies in that it provides an independent assessment of the effect that capitalises on existing quality population-based data. Specifically, we used two population-level distributions, prevalence by age and by sexual risk, to explore a population-level ‘signature’ of partial immunity. We also factored in the model different biological (eg, various forms of immunity and impact of treatment on immunity development) and behavioural (eg, impact of age on sexual behaviour and various forms of sexual mixing by age and risk behaviour) mechanisms that potentially may explain the observed patterns, irrespective of inclusion of an immunity effect. Accordingly, we aimed to answer two questions: (1) Is the existence of partial immunity against reinfection necessary to explain prevalence patterns by age and sexual risk? (2) How strong is the effect of partial immunity likely to be?

METHODS
We constructed a population-based deterministic compartmental model to assess the role of long-lasting partial immunity in C. trachomatis epidemiology. The model was developed based on a review of existing mathematical models for this infection.5 21–26 The model stratified the population according to infection status, immune status, age and sexual risk behaviour, and to reduce complexity, sex was not included explicitly in the model, nor did the model explicitly distinguish between different forms of sexual transmission.

Mathematical model
Model description and C. trachomatis natural history
Online supplementary figure S1 shows a schematic diagram of the model. The model assumes full susceptibility to infection at birth. Following the first C. trachomatis exposure, infected individuals enter a non-infectious latent period of 14 days,5 followed by an infectious period that can be asymptomatic or symptomatic. We assumed that asymptomatic infections comprise 62.5% of all infections.5 The infectious period lasts for 300 days for asymptomatic infection,5 but only 35 days for symptomatic infection, on account of treatment-seeking behaviour.5

We assumed that, by the time of infection clearance, asymptomatic individuals acquire short-term temporary (90 days) but full immunity against reinfection, per the recent Althaus et al model5 and as informed by animal model studies.17 This is followed by long-lasting partial immunity that reduces susceptibility to reinfection by a fraction α17. The latter mechanism is informed by a review of evidence from human studies that examined both biological (eg, markers of protective immunity) as well as epidemiological (eg, patterns in different populations) data.17

Meanwhile, symptomatic individuals do not acquire immunity and revert back to the fully susceptible state after clearance by treatment—treatment shortens infection duration in symptomatic individuals, thus reducing their chance to develop an adequate immune response.5

Details on model structure, equations and parameterisation are in the online supplementary materials.

Demography and sexual risk behaviour
The model assumes a stable population with a balance between births and deaths. It further incorporates 20 age groups, each of which describes a 5-year age band in the population. Demographics for the UK and the USA were drawn from the United Nations Population Division databases.27 Sexual activity lifespan extends in the model from ages 15 to 74, with sexual activity declining with older age. For each 5-year age category, the model incorporates six sexual risk groups describing a hierarchy of sexual risk behaviour varying from low to high levels. Accordingly, the model accommodates for the broad behavioural heterogeneity that typically exists in a given population.38–40

Distribution of the population across risk groups is informed by data for the number of heterosexual partners during the last 12 months, as reported in the second UK National Survey of Sexual Attitudes and Lifestyles (Natsal-2).21 22 Distribution of the level of risk behaviour across risk groups follows a power-law function as informed by sexual partnership data30 and analyses of complex weighted networks.33–36 Partnership acquisition rate in each risk group varies with age as informed by the Natsal-2 data.27 Coital frequency also varies with age as informed by the US data.39

The pattern of sexual mixing between sexually active individuals is determined by two mixing matrices describing the likelihood of a partnership to be formed based on age group or risk group.30 39 40 Each matrix describes a mixing continuum varying between proportionate mixing (no preferential bias based on age or risk group) and full assortativity (partnerships formed exclusively within the same age or risk group).

Details on the inclusion of demography and sexual behaviour in the model are in the online supplementary materials.

Data sources
The model was parameterised using available data for C. trachomatis natural history (online supplementary table S1) and sexual behaviour (online supplementary table S1 and S2). The key model parameters were based on the median value of the range that was used in published models.5 21–26 Empirical data describing the distribution of C. trachomatis prevalence by age were obtained from (1) a systematic review of studies assessing C. trachomatis prevalence in the UK41 and (2) the US Centers for Disease Control and Prevention (CDC) STI database.42 The distribution of C. trachomatis prevalence by sexual risk group was obtained through Natsal-2 data analysis.21 31

Plan of analysis
We conducted analyses to explore the role of partial immunity in explaining observed C. trachomatis epidemiological patterns.

Partial immunity and age-specific and sexual risk-specific distributions of C. trachomatis prevalence
We generated model predictions for the age-specific and sexual risk-specific distributions of C. trachomatis prevalence in the UK at different strengths for the partial immunity, and compared these distributions with empirical data. The comparison was conducted to assess whether the existence of immunity yields enhanced agreement between model predictions and empirical data.

Estimation of the strength of partial immunity against reinfection
The strength of the long-lasting partial immunity against reinfection (α) was estimated by fitting model predictions to data from the following surveys: (1) UK age-specific41 and sexual risk-specific21 31 C. trachomatis prevalence distributions, and (2) US age-specific39 C. trachomatis prevalence distribution. No fitting was done for the US sexual risk-specific C. trachomatis distribution, as such distribution was not available for the used CDC data. The fitting was implemented by minimising the residual

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sum of squares between data points and model predictions. Analyses were conducted assuming endemic equilibrium.

The uncertainty intervals (UIs) for estimates were calculated through multivariate uncertainty analyses with respect to variations in the model’s sexual behaviour structure. This was done using Monte Carlo sampling from (conservatively) uniform probability distributions, assuming 20% uncertainty around the parameters’ point estimates, as informed by the range of available data and previous modelling studies.\textsuperscript{29–30,41} Each new set of parameters was used to refit \emph{C. trachomatis} prevalence distributions, and accordingly estimate $\alpha$. We implemented 500 uncertainty runs for each estimation and determined the median and mean and associated 95% UI.

Sensitivity analyses with respect to variations in model structure
Several sensitivity analyses were conducted with respect to variations in model structure. These are summarised in table 1.

<table>
<thead>
<tr>
<th>Sensitivity analysis</th>
<th>Description</th>
<th>Result</th>
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<tbody>
<tr>
<td>1. Variation in the distribution of risk behaviour across risk groups.*</td>
<td>Explored the impact of variation in the distribution of risk behaviour across risk groups by varying (in univariate analysis) the parameter $\sigma$ of the distribution of risk behaviour (online supplementary materials), but fixing $\alpha$ at its model-predicted baseline value.</td>
<td>The predicted age-specific \emph{Chlamydia trachomatis} prevalence distribution was largely invariable despite the variation in the distribution of risk behaviour across risk groups (online supplementary figure S2A).</td>
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<tr>
<td>2. Variation in the sexual mixing by age.*</td>
<td>Explored the impact of variation in sexual mixing by age (in univariate analysis) across the full spectrum starting from proportionate mixing-up to fully assortative mixing. This was done by varying $\epsilon_1$, the parameter describing the degree of assortativity in mixing by age (online supplementary materials).</td>
<td>The predicted age-specific \emph{C. trachomatis} prevalence distribution was largely invariable despite the variation in the sexual mixing by age (online supplementary figure S2B).</td>
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<tr>
<td>3. Variation in the sexual mixing by risk.*</td>
<td>Explored the impact of variation in sexual mixing by risk (in univariate analysis) across the full spectrum starting from proportionate mixing-up to fully assortative mixing. This was done by varying $\epsilon_2$, the parameter describing the degree of assortativity in mixing by risk (online supplementary materials).</td>
<td>The predicted age-specific \emph{C. trachomatis} prevalence distribution was largely invariable despite the variation in the sexual mixing by risk (online supplementary figure S2C).</td>
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<td>4. Temporal variation in risk behaviour.</td>
<td>Explored the impact of temporal variation in risk behaviour on our estimated partial immunity strength by assuming that 10% of individuals change their risk group every year.</td>
<td>$\alpha$ for the UK data was estimated at 93% (95% UI: 89%–95%) with an uncertainty analysis median of 93%—similar to the original estimate.</td>
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<tr>
<td>5. Removal of latent period in \emph{C. trachomatis} natural history.</td>
<td>Explored the impact of removing the latent period in \emph{C. trachomatis} natural history.</td>
<td>$\alpha$ for the UK data was estimated at 93% (95% UI: 88%–97%) with an uncertainty analysis median of 93%— similar to the original estimate.</td>
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<tr>
<td>6. Inclusion of partial immunity for the symptomatically infected individuals.</td>
<td>Explored the impact of inclusion of partial immunity for the symptomatically infected individuals.</td>
<td>$\alpha$ for the UK data was estimated at 93% (95% UI: 89%–96%) with an uncertainty analysis median of 93%— similar to the original estimate.</td>
</tr>
<tr>
<td>7. Variation in the duration of the short-term temporary but full immunity.</td>
<td>Explored the impact of varying the duration of the short-term temporary but full immunity over a range of 0–100 days.</td>
<td>Variation in the short-term temporary immunity had limited impact on the estimated effect size of partial immunity (online supplementary figure S3).</td>
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</tbody>
</table>

All sensitivity analyses were applied to the model fit of the UK data.

* Conducted in view of the fundamental ambiguity in defining ‘sexual risk’,\textsuperscript{26–46} and done on the prediction for the age-specific \emph{C. trachomatis} prevalence distribution, since this distribution is the most prototypical pattern in \emph{C. trachomatis} epidemiology. Ul, uncertainty interval.

RESULTS
Figure 1A shows the predicted age-specific \emph{C. trachomatis} prevalence distribution for the UK data at different $\alpha$ (the fractional reduction in susceptibility to reinfection). With no immunity ($\alpha = 0\%$), it was not possible to generate the typically observed distribution where the prevalence is highest in those aged 15–24 years and declines to <1% by age 30. $\alpha$ must be large enough, in the range of 70%–90%, to reproduce this prototypical \emph{C. trachomatis} empirical pattern. Moreover, it was not possible to generate the distribution without incorporating the age dependence of sexual behaviour.

Figure 1B shows the predicted age-specific \emph{C. trachomatis} prevalence distribution at different $\alpha$, but assuming no variation in sexual behaviour with age—absence of this variation did not lead to a good fit of empirical data.

Figure 1C shows the predicted risk-specific \emph{C. trachomatis} prevalence distribution for the UK data at different $\alpha$. With $\alpha = 0\%$, the prevalence in high-risk populations was much higher than that in low-risk populations, in contrast to empirical data that suggest a more even distribution by risk.\textsuperscript{21} In order to generate such more even distribution, $\alpha$ must be large enough to substantially lower \emph{C. trachomatis} prevalence in high-risk populations, who are more likely to experience reinfections. Of notice that the outlier for risk group 4 may be due to sampling variation or to the ambiguity of defining ‘sexual risk’.\textsuperscript{26–46}—a definition based on simply reported number of partners may not capture the true risk of infection exposure in the sexual network.

To derive a plausible estimate for the effect size of $\alpha$, we fitted \emph{C. trachomatis} prevalence distribution by age and by risk for the UK data (figure 2A,C and online supplementary tables S3, S4) and by age for the US data (figure 2B and online supplementary materials).
Figure 1  
*Chlamydia trachomatis* (CT) prevalence in the UK by age and by sexual risk under different assumptions for the effect size of partial immunity against reinfection (α). Model predictions for (A) age-specific distribution of CT prevalence at variable levels of α and assuming age dependence of sexual behaviour, (B) age-specific distribution of CT prevalence at variable levels of α but with no age dependence of sexual behaviour, and (C) sexual risk-specific distribution of CT prevalence at variable levels of α for those 18–44 years of age. Empirical data (illustrated by ‘*’) were provided from reference 41 for CT prevalence by age and from references 21 31 for CT prevalence by sexual risk. Distribution of the population across risk groups was based on the reported number of heterosexual sex partners during the last 12 months in the second National Survey of Sexual Attitudes and Lifestyles.21,31

Figure 2  
Model fits for *Chlamydia trachomatis* (CT) prevalence in the UK by age and by sexual risk, and in the USA by age. Model fits for (A) age-specific distribution of CT prevalence in the UK, (B) age-specific distribution of CT prevalence in the USA and (C) sexual risk-specific distribution of CT prevalence in the UK for those 18–44 years of age. Empirical data (illustrated by ‘**’) were provided for the UK from references 21 31 41 and for the USA from reference 42.

table S5). The model produced robust fits of these distributions. α for the UK data was estimated at 93% (95% UI=88%–97%) with an uncertainty analysis median of 93%. α for the US data was estimated at 67% (95% UI=24%–88%) with an uncertainty analysis median of 69%.

Table 1  shows the results of the sensitivity analyses with respect to variations in model structure. The analyses indicated
that the estimated $\alpha$ is not likely to be affected appreciably by the ambiguity in defining sexual risk nor by lack of detailed knowledge of sexual networking in a population.

Figure 3 shows the predicted age-specific C. trachomatis prevalence distribution under three distinct assumptions for the mechanism of partial immunity: susceptibility reduction (our baseline assumption; figure 3A), infectious-period duration reduction with reinfection (figure 3B) and infectiousness reduction with reinfection (figure 3C). The results indicated that a mechanism of infectiousness reduction has a limited effect on the age-specific prevalence distribution, particularly in older populations, regardless of the immunity effect size. Meanwhile, a mechanism of infectious-period duration reduction affects the age-specific prevalence distribution in a similar manner to that of susceptibility reduction.

**DISCUSSION**

We explored the role of immunity in C. trachomatis epidemiology using a hypothesis-generation mathematical modelling approach. Our results indicated that a strong effect of partial immunity against reinfection should be present to explain observed prevalence patterns. The immunity effect size, reduction in susceptibility to reinfection, may also exceed 65%. The results further indicated that the mechanism of immunity could be either a reduction in susceptibility to reinfection or a reduction in duration of infection upon reinfection. Observed patterns could not be explained by merely an immunity effect on infectiousness (organism load).

The presented analyses highlighted how a natural history effect that occurs at the individual level (ie, development of partial immunity) expresses itself indirectly as an observed effect on prevalence patterns at the population level—thereby facilitating a derivation of a plausible estimate for the effect size using a population-level mathematical model. The immunity effect led to high prevalence in youths, with rapid decline in prevalence with age, testifying to age being a strongly predictive risk factor for C. trachomatis infection. The immunity effect led also to a ‘dispersion’ effect of more even prevalence distribution by risk group—affirming the empirically observed smaller Gini coefficient for C. trachomatis, compared with gonorrhoea and syphilis.

The observed strong immunity effect is consistent with existing direct evidence from animal studies and indirect evidence from human studies. In the context of previous studies on C. trachomatis control, the observed effect suggests that control programmes focused on early detection and treatment may not be effective, if they hinder the development of an adequate immune response. The roll-out of screening in several countries was paralleled by increases in reinfection rates that remain unexplained by advances in diagnostic tests, screening coverage and sexual behaviour. This outcome demonstrates how it is difficult to achieve C. trachomatis control without probably a vaccine—the observed strong immunity effect supports the concept, and may suggest the feasibility, of vaccine development. A vaccine that is administered to adolescents, in similar fashion to human papillomavirus vaccination, may have a larger impact in controlling C. trachomatis transmission, and be more cost-effective, than a screening programme.

The sensitivity analyses of alternative mechanisms for the immunity effect indicated that infectiousness reduction cannot explain observed patterns (figure 3). Even if this mechanism is present biologically, it can affect the overall intensity of C. trachomatis transmission in a population, but not as much the observed age-specific prevalence distribution. Meanwhile, a mechanism of infectious-period duration reduction affects this distribution just as susceptibility reduction, and therefore provides an alternative plausible explanation for the effect of partial immunity. Further work is warranted to investigate which of these two mechanisms, or a combination, underlie immunity. Of notice that these three mechanisms, to one degree or another, could eventually be determined to be in action and simultaneously. If so, C. trachomatis immunity may be proven to be more influential in the epidemiology of this infection than previously thought.

**Figure 3** Sensitivity analyses of the impact of alternative biological mechanisms, for the effect of partial immunity, on the model-predicted age-specific Chlamydia trachomatis (CT) prevalence in the UK. Model predictions for the age-specific CT prevalence, under different levels for the effect size of partial immunity ($\alpha$), assuming a mechanism of (A) susceptibility reduction to reinfection, (B) infectious-period duration reduction with reinfection or (C) infectiousness reduction with reinfection. Empirical data (illustrated by ‘*’) were provided from reference.21
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Limitations may have affected this study. Although an elaborately developed mathematical model was used to capture the complexity of C. trachomatis transmission dynamics, the results depend on the used model. We used a frequency-dependent model, but this type of model may overestimate immunity effects. The model depends on availability and representativeness of input data—the estimate for the USA had a wide UI, as the sexual risk-specific prevalence distribution was not available for the used CDC data. The sexual risk-specific prevalence distribution for the UK data was based on the number of sexual partners, which alone may not sufficiently capture sexual risk. This distribution, however, was influential in estimating such a strong risk for the UK data. If the shape of this distribution was less even, we would still have estimated strong immunity, but with a smaller α (Figure 1). We assumed that symptomatically infected persons do not develop partial immunity because of treatment-seeking behaviour—a reasonable assumption but with undetermined validity. Lastly, for sexual mixing, we assumed a specific mix of proportionate and assortative mixing since the actual mixing in the population is not known.

Despite these limitations, the model was sufficiently complex to incorporate the main factors that can affect our research questions, and produced robust fits for C. trachomatis prevalence by age and by risk. None of the biological and behavioural factors suspected to affect the study outcome affected the study conclusions, and the different sensitivity analyses affirmed our findings. We conducted multivariate uncertainty analyses with respect to variations in model structure, rather than using maximum likelihood methods, to account for broader uncertainty in the estimated effect size. Although the exact effect size of C. trachomatis immunity is yet to be determined with precision, and there was a difference between the UK-based and US-based estimates, conclusively the results affirmed a strong effect of partial immunity. With sampling variation affecting C. trachomatis prevalence distributions in population-based surveys, and in the context of fundamental ambiguity in defining ‘sexual risk’, the presented approach is not best suited to provide an exact effect size, but is sufficient to demonstrate the likely existence of a strong partial immunity effect.

In conclusion, we explored the existence of partial immunity against C. trachomatis reinfection using an indirect, but novel hypothesis-generation approach. Our results indicated that a strong immunity effect should be present to explain observed C. trachomatis prevalence patterns. These findings are suggestive of a strong natural immune response against C. trachomatis infection, which upon elucidation of its detailed biological mechanisms may have implications for vaccine development.

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Contributors RO conceived the study, developed the mathematical models and conducted the analyses. HC contributed to data analysis and wrote the first draft of the article. LIA-R provided technical input. LIA-R led the study design and analyses. All authors contributed to results generation and interpretation and to writing of the article.

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Key messages

• Partial immunity against Chlamydia trachomatis reinfection can explain the observed prevalence patterns by age and sexual risk.

• C. trachomatis infection appears to elicit a strong natural immune response that reduces substantially the susceptibility to reinfection.

• Immunity mechanism can be either a reduction in susceptibility to reinfection or a reduction in infectious-period duration upon reinfection, or a combination of both.

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