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Prevalence of *Chlamydia trachomatis*-Specific Antibodies before and after Mass Drug Administration for Trachoma in Community-Wide Surveys of Four Communities in Nepal

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Abstract. The target end date for the global elimination of trachoma as a public health problem is 2020. As countries begin the process for submitting their dossier for the validation of elimination of trachoma as a public health problem, strategies for post-validation surveillance must be considered. Seroprevalence of antibodies against antigens from the causative bacteria *Chlamydia trachomatis* (*Ct*) in young children has been shown to reflect trachomatous inflammation–follicular (TF) rates in both endemic and previously endemic settings. However, none of these studies has directly compared age seroprevalence in the same communities before and after mass drug administration (MDA) for trachoma. Here we report a marked shift in age seroprevalence curves in four villages in Kapilvastu District, Nepal, before and after MDA. Clinical examinations were performed and blood was taken before (N = 659) and 5 years after (N = 646) MDA. Rates of TF decreased from 17.6% in ≤ 9-year-olds before MDA (N = 52) to 0% in ≤ 9-year-olds (N = 73) after MDA. Positive antibody responses to *Ct* in the entire population decreased from 82.1% pre-MDA to 35.8% post-MDA, whereas those among ≤ 9-year-olds decreased from 59.6% to 4.1%. These data show that the postintervention decrease in TF was reflected in a drop in anti-*Ct* antibody responses, suggesting that antibody responses could be useful indicators for post-validation surveillance.

INTRODUCTION

Trachoma is caused by repeated ocular infection with *Chlamydia trachomatis* (*Ct*) and is the leading infectious cause of blindness in the world today. Over 200 million people are at risk of infection with an estimated 1.9 million people blind or visually impaired due to trachoma (www.trachomacoalition.org/GET2020). Efforts by the World Health Organization (WHO) and other partners are underway to eliminate trachoma.1–4 The current WHO endpoint for MDA is a prevalence of trachomatous inflammation–follicular (TF) rates in both endemic and previously endemic settings. However, none of these studies has directly compared age seroprevalence in the same communities before and after mass drug administration (MDA) for trachoma. Here we report a marked shift in age seroprevalence curves in four villages in Kapilvastu District, Nepal, before and after MDA. Clinical examinations were performed and blood was taken before (N = 659) and 5 years after (N = 646) MDA. Rates of TF decreased from 17.6% in ≤ 9-year-olds before MDA (N = 52) to 0% in ≤ 9-year-olds (N = 73) after MDA. Positive antibody responses to *Ct* in the entire population decreased from 82.1% pre-MDA to 35.8% post-MDA, whereas those among ≤ 9-year-olds decreased from 59.6% to 4.1%. These data show that the postintervention decrease in TF was reflected in a drop in anti-*Ct* antibody responses, suggesting that antibody responses could be useful indicators for post-validation surveillance.

MATERIALS AND METHODS

Study population. The study was conducted in four villages in the Kapilvastu District of the Lumbini Zone in southwestern Nepal. The three annual rounds of MDA were completed in 2009 (given in 2007, 2008, and 2009). All community members aged 1 year and over were invited to participate in the study. Participation rates were 81–87% across the four villages. The same communities were sampled before MDA (2000 and 2002) and after MDA (2014). Demographic data on villages are presented in Supplemental Table 1; two of these villages were contiguous and so treated as a single site in this table.

Ethics statement. Institutional Review Board (IRB) approvals were obtained for both data collections from Children’s Hospital and Research Center Oakland (IRB number 2013-043) and by Nepal Netra Jhoti Sangh (Nepali Prevention of Blindness Program), and data were not anonymized for response to elimination program interventions. Monitoring conjunctival infection by commercial nucleic acid amplification tests is an option but is costly and time consuming.5 Age-specific prevalence of antibody responses to *Ct* at the community level could provide an informative proxy measure of the intensity of transmission. Serological tests for measuring antibodies represents a means for monitoring cumulative exposure to ocular infection in children.6–10 Serological testing would also facilitate integration of trachoma surveillance with other health program activities in which blood collection is occurring, such as demographic and health surveys, malaria indicator surveys, or vaccine-coverage surveys.

Here, we examined the use of serological tools for monitoring and evaluation in post-MDA settings by assessing the age-specific prevalence of clinical signs and antibody responses of trachoma within four communities that received MDA for trachoma 5 years before the study.
these researchers. Centers for Disease Control and Prevention researchers were non-engaged (i.e., did not have access to patient-identifying information) in the study. All study participants gave written informed consent, or written parental consent was obtained for participants under the age of 18.

Clinical examination for trachoma. Grading of the upper tarsus of each study participant was performed according to the WHO simplified trachoma grading system by experienced graders (D.D. and R.P.K.).

Blood collection and preparation. Serum or finger prick blood was obtained from all participants; 2000/2002 samples were all serum and 2014 samples were a mix of finger prick blood from primarily < 15-year-olds and serum from ≥ 15-year-olds. Finger prick blood was collected onto filter papers with six circular extensions calibrated to absorb 10 μL of whole blood (TropBio Pty Ltd, Townsville, Queensland, Australia) and stored at −20°C before use. One bloodspot extension for each participant was eluted overnight at 4°C with 500 μL of phosphate-buffered saline (PBS) containing 0.5% bovine serum albumin (BSA), 0.05% Tween 20, 0.02% sodium azide, 0.5% polyvinyl alcohol, and 0.8% polyvinylpyrrolidone, designated as PBN1. This elution was equivalent to a serum dilution of approximately 1:100. Eluates (100 μL) were diluted to a final volume of 400 μL in PBN1 with 0.5% w/v of *Escherichia coli* crude extract to block nonspecific binding [8] for a final serum dilution of approximately 1:400. Sera were diluted 1:400 in 400 μL PBN1 with 0.5% w/v of *E. coli* crude extract and stored overnight at 4°C.

Multiplex bead array assay. Bloodspot eluates and serum dilutions were screened in duplicate with Pgp3 and CT694 antigen-coupled beads in a multiplex bead assay as previously described. Briefly, Pgp3- and CT694-coupled beads (2,500 each) were added to each well of a prewet filter papers with six circular extensions calibrated to absorb 10 μL of whole blood (TropBio Pty Ltd, Townsville, Queensland, Australia) and stored at −20°C before use. One bloodspot extension for each participant was eluted overnight at 4°C with 500 μL of phosphate-buffered saline (PBS) containing 0.5% bovine serum albumin (BSA), 0.05% Tween 20, 0.02% sodium azide, 0.5% polyvinyl alcohol, and 0.8% polyvinylpyrrolidone, designated as PBN1. This elution was equivalent to a serum dilution of approximately 1:100. Eluates (100 μL) were diluted to a final volume of 400 μL in PBN1 with 0.5% w/v of *Escherichia coli* crude extract to block nonspecific binding [8] for a final serum dilution of approximately 1:400. Sera were diluted 1:400 in 400 μL PBN1 with 0.5% w/v of *E. coli* crude extract and stored overnight at 4°C.

Table 1

| TF prevalence and seroprevalence pre- and post-MDA, Nepal, 2000/2 and 2014 |
|---------------------------|--------------------------|---------------------------|---------------------------|---------------------------|
| Pre-MDA % (95% CI) | Post-MDA % (95% CI) | All ages Pre-MDA % (95% CI) | All ages Post-MDA % (95% CI) |
| TF | 17.6 (7.25–27.9) | 0 (0–5.6) | 6.5 (4.6–8.4) | 1.24 (0.39–2.09) |
| Pgp3+ | 59.6 (46.3–72.9) | 2.7 (0.0–6.4) | 35.8 (73.6–80.0) | 13.2 (28.4–35.6) |
| CT694+ | 53.8 (40.3–67.4) | 1.3 (0.0–3.9) | 71.1 (67.6–74.6) | 34.9 (31.2–38.58) |
| Ab+* | 58.0 (46.3–72.9) | 4.0 (0.0–8.4) | 82.5 (78.6–85.4) | 38.6 (34.85–42.35) |
| Pgp3 < CT694- | 53.8 (40.3–67.4) | 0.0 (0–8.4) | 56.5 (61.9–69.1) | 24.6 (23.19–30.01) |

* Data shown are percent positive responses for given indicator, with 95% CI in parentheses for samples taken in 2000/2 (pre-MDA, N = 52 for < 9-year-olds and 659 for all ages) and 2014 (post-MDA, N = 73 for 1–9-year-olds and 646 for all ages). CI = confidence interval; MDA = mass drug administration; TF = trachomatous inflammation-follicular.

* Denotes positive antibody responses to Pgp3 alone, CT694, alone or both Pgp3 and CT694. Pgp3 < CT694 denotes positive antibody responses to both antigens.
to Pgp3 and 1.3% (1/73, 95% CI: 0–3.9) recognizing CT694 (Table 1). None of the ≤9-year-olds had antibody responses to both Pgp3 and CT694 post-MDA (Table 1).

The MFI of the three ≤9-year-olds with positive antibody responses post-MDA was 1,513 and 2,485 for Pgp3 reactivity and 490 for CT694 reactivity (data not shown). By contrast, the specimens from ≤ 9-year-olds taken pre-MDA had median MFI value of 25,160 for positive Pgp3 responses (range 1,200–31,534) and median MFI value of 3,347 for positive CT694 responses (range 460–30,445, data not shown).

When evaluating all ages, pre-MDA TF was 6.5% (43/659, 95% CI: 4.6–8.4) and post-MDA was 1.24% (8/646, 95% CI: 0.39–2.69) (Table 1). The percent of all participants with any positive anti-Ct response pre-MDA was 82.5% (544/659, 95% CI: 79.6–85.4), with 77.5% (511/659, 95% CI: 73.6–80.0) having antibodies recognizing Pgp3 and 71.1% (468/659, 95% CI: 67.6–74.6) with antibodies recognizing CT694 (Table 1). After MDA, 40.4% (261/646, 95% CI: 36.7–44.2) of individuals had antibody responses to either Ct antigen, with 34.9% (226/646, 95% CI: 31.2–38.5) recognizing Pgp3 and 32.0% (207/646, 95% CI: 28.4–35.6) recognizing CT694 (Table 1).

**Shift in age-specific seroprevalence after MDA.** Line graphs of seroprevalence grouped by decade and age-stratified MFI of the antibody responses differed in the pre- and post-MDA phases (Figures 1 and 2). Pre-MDA, the proportion antibody-positive in all age groups was greater than 60%, and the mean MFI values for the different age groups were statistically similar (Pgp3: P = 0.6984; CT694: P = 0.5629, Figure 2). Post-MDA, there was a marked age-dependent increase in both the proportion antibody-positive (Figure 1) and the mean MFI values among age groups (Pgp3: P < 0.001; CT694: P < 0.001, Figure 2).

**DISCUSSION**

The data presented here show marked shifts in the Ct-specific age-seroprevalence curve in trachoma-endemic communities before and after cessation of MDA. Rates of TF in ≤9-year-olds dropped from 17.6% before MDA to 0% after MDA in these communities. Antibody positivity in ≤9-year-olds before MDA was approximately 59%, whereas the TF rates in this age group were only 17.6%. This confirms previously reported data that antibody positivity rates exceed TF rates in trachoma-endemic communities. In multiple studies in trachoma-endemic communities, the antibody positivity rate is approximately 2.5 to 5× that of the TF rate. This suggests that antibody responses are longer-lived than TF responses, which themselves can last for weeks or months. The relationship between TF and antibody-positive rates in trachoma-endemic communities will be critical for threshold determination for potential programmatic use of seroprevalence assays.

Agreement between antibody responses against Pgp3 and CT694 among ≤9-year-olds differed between the pre- and post-MDA setting; the pre-MDA agreement was 90%, whereas the post-MDA agreement was 0%. For the post-MDA samples collected 5 years after cessation of MDA, no TF was seen in ≤9-year-olds, and antibody positivity rates were 2.7% for Pgp3, 1.3% for CT694, and 4.1% for either antigen. This lack of concordance between Pgp3 and CT694 reactivity was seen previously in communities in which trachoma has been controlled or eliminated. Pgp3 is a well-defined immunodominant Ct antigen that is encoded in the multi-copy Ct plasmid and, therefore, should be highly specific for Ct, but not other chlamydial infection. Available data suggest that CT694 is recognized by the immune system in a smaller proportion of individuals with active urogenital Ct infection than Pgp3. The intensities of the responses against Pgp3 and CT694 were also much lower post-MDA than before MDA when transmission was active, possibly related to less exposure to ocular Ct in the post-MDA setting. Although Pgp3-positive individuals had low-positive responses, the CT694-positive response post-MDA were in the indeterminate range of the assay, similar to previous observations in Tanzania. It may be useful to focus operational research on Pgp3, and in fact we have selected Pgp3 as the antigen for use in rapid antibody tests for trachoma.

One explanation for this shift in the age-seroprevalence data in 2014 is that other factors during this time frame ca. 1999 caused a decrease in transmission before the initiation of MDA in 2007. One potential factor is the improved socioeconomic status of the community that occurred as young Nepali men worked abroad and sent funds home to support their families. If the transmission rates were changing in 1999, those changes might not be reflected in the antibody data from the earlier surveys in 2000 and 2002 as it would take time for any socioeconomic effect to occur. An alternative explanation is that after 5–8 years of low to no transmission after MDA, ocular Ct infection in the community antibody responses...
were waning and that individuals were seroreverting. The only published data on the longevity of the antibody response for trachoma is from a single Tanzanian hyperendemic community (46% TF in 1–6-year-olds) 6 months after MDA, in which no seroreversion was observed. Despite this, it is not unreasonable to predict that in a setting with a lower initial TF prevalence (17.6% in 1–9-year-olds) that some individuals of all ages would lose their antibody response over time and that repeated exposure may be required to maintain long-lived antibody production for trachoma. There may be some threshold level or number of infections that results in long-lived, or possibly irreversible, anti-Ct antibody responses. The longevity of the antibody response will determine how restrictive the age range for testing will need to be for post-validation surveillance. In the case of very long-lived antibody responses at a population level, the focus for monitoring the impact of MDA would need to be on children born after the cessation of MDA. If the responses are shorter-lived (months to a few years, rather than decades) and seroreversion occurs in most or all of the population, a broader age range could be considered for monitoring for reemergence of transmission.

A major limitation of this study was the small number of ≤9-year-olds enrolled and the lack of 1-year-olds. The WHO indicator age range for TF is 1–9-years and data from children in that range are, therefore, of key importance in studies evaluating the utility of serological testing for trachoma surveillance. One-year-olds routinely have the lowest antibody positivity rates in studies from trachoma-endemic villages, so the lack of 1-year-olds here may slightly inflate the pre-MDA seroprevalence estimates but likely have no effect on the post-MDA estimates. Despite the small number of samples from ≤9-year-olds in this study (52 pre-MDA and 73 post-MDA), we observed a patently steep drop in antibody positivity among this group post-MDA. However, the number of samples may limit the generalizability of these data in helping to calculate a threshold for antibody positivity for trachoma post-MDA surveillance. In addition, although the setting is characterized as “post-MDA,” there has been considerable socioeconomic improvement in these communities since the “pre-MDA” time frame. Therefore, the stark change in the age-seroprevalence curve may not be attributable solely to MDA. The cause of the decline in TF in these communities may be less important than the fact that the drop in TF prevalence was accompanied by a decrease in antibody prevalence at the 2014 data collection. Finally, although we collected data from four villages, two of these were contiguous and could not be considered as separate clusters, and we therefore could not perform cluster-level analysis without the variance calculations being overwhelmed by the limited number of clusters. This study took advantage of

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**Figure 2.** Intensity of anti-Chlamydia trachomatis antibody responses by age pre- and post-MDA, Nepal, 2000/2 and 2014. Left: the median fluorescent intensity of anti-Pgp3 antibodies with background subtracted out (Pgp3 MFI-BG) are shown pre-MDA (top) and post-MDA (bottom) for all ages grouped by decade. Right: the MFI of anti-CT694 antibodies with background subtracted out (CT694 MFI-BG) are shown pre-MDA (top) and post-MDA (bottom) for all ages grouped by decade. Y-axes are shown on a logarithmic scale. Boxes show the 25–75% quartile of data with solid horizontal lines in the middle representing median proportion-positive for each grouping. The upper and lower whiskers represent the minimum and maximum range. MDI = mass drug administration; MFI = median fluorescence intensity.
the presence of stored samples from a pre-MDA study in these villages; prospective studies evaluating the pre- and post-MDA seroprevalence in a population-based cluster survey are needed.

These data are the first to directly compare age-seroprevalence data before and after completion of MDA for trachoma from the same communities. We observed a clear and dramatic shift in age-seroprevalence data after intervention, reflecting the TF prevalence before and after MDA. Therefore, serosurveillance for trachoma has the potential as a useful tool for evaluating transmission status in settings that have moved from programs focused on eliminating trachoma as a public health problem to programs with the aim of control.

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