DISCOVERY OF HUMAN-SPECIFIC IMMUNODOMINANT CHLAMYDIA TRACHOMATIS B CELL EPITOPES

K. Shamsur Rahman,1,2 Toni Darville,1,2 Ali N. Russell,1,2 Catherine M. O’Connell,1,2 Harold C. Wiesenfeld,3 Sharon L. Hillier,3 Erfan U. Chowdhury,1,2 Yen-Chen Juan,1,2 Bernhard Kaltenboeck1

1 Department of Pathobiology, College of Veterinary Medicine, Auburn University, Auburn, Alabama, USA
2 Department of Pediatrics, University of North Carolina at Chapel Hill, Chapel Hill, North Carolina, USA
3 Department of Obstetrics, Gynaecology and Reproductive Sciences, the University of Pittsburgh School of Medicine and the Magee-Womens Research Institute Pittsburgh, Pittsburgh, Pennsylvania, USA

ABSTRACT

Chlamydia species-specific serology is compromised by cross-reactivity of the gold standard microimmunofluorescence (MIF) or commercial enzyme-linked immunosorbent assays (ELISAs). This study was conducted to discover novel C. trachomatis-specific peptide antigens that were recognized only by the antibody response of the natural human host. We evaluated a library of 271 peptide antigens from immunodominant C. trachomatis proteins by reactivity with 125 C. trachomatis antibody-positive sera from women with PCR-confirmed C. trachomatis infection and 17 C. trachomatis antibody-negative sera from low-risk women never diagnosed with C. trachomatis infection. These C. trachomatis peptide antigens had been predicted in silico to contain B cell epitopes but had been nonreactive with mouse hyperimmune sera against C. trachomatis. We discovered 38 novel human host-dependent antigens from 20 immunodominant C. trachomatis proteins (PmpD, IncE, IncG, CT529, CT618, CT442, TarP, CT143, CT813, CT795, CT223, PmpC, CT875, CT579, LcrE, IncA, CT226, CT694, Hsp60, and pGP3). Using these human sera, we also confirmed 10 C. trachomatis B cell epitopes from 6 immunodominant C. trachomatis proteins (OmpA, PmpD, IncE, IncG, CT529, and CT618) as host species-independent epitopes that had been previously identified by their reactivity with mouse hyperimmune sera against C. trachomatis. ELISA reactivities against these peptides correlated strongly with the C. trachomatis microimmunofluorescence (MIF) text results (Pearson’s correlation coefficient [R] = 0.80; P < 10⁻⁶). These C. trachomatis peptide antigens do not cross-react with antibodies against other Chlamydia species and are therefore suitable for species-specific detection of antibodies against C. trachomatis. This study identified an extended set of peptide antigens for simple C. trachomatis-specific ELISA serology.

IMPORTANCE

Current serological assays for species-specific detection of anti-Chlamydia species antibodies suffer from well-known shortcomings in specificity and ease of use. Due to the high prevalences of both anti-C. trachomatis and anti-C. pneumoniae antibodies in human populations, species-specific serology is unreliable. Therefore, novel specific and simple assays for chlamydial serology are urgently needed. Conventional antigens are problematic due to extensive cross-reactivity within Chlamydia spp. Using accurate B cell epitope prediction and a robust peptide ELISA methodology developed in our laboratory, we identified immunodominant C. trachomatis B cell epitopes by screening performed with sera from C. trachomatis-infected women. We discovered 38 novel human host-dependent antigens from 20 immunodominant C. trachomatis proteins, in addition to confirming 10 host-independent mouse serum peptide antigens that had been identified previously. This extended set of highly specific C. trachomatis peptide antigens can be used in
Obligate intracellular bacteria of the genus *Chlamydia* infect virtually all vertebrates and cause largely chronic and asymptomatic disease conditions (1, 2). The principal human chlamydial pathogens are *C. trachomatis* and *C. pneumoniae* (1, 2). *C. trachomatis* serovars A to C cause ocular infection and are the leading causes of preventable blindness, affecting tens of millions of people in developing countries (3). *C. trachomatis* serovars D to K cause genitourinary tract infections, and serovars L1 to L3 cause lymphogranuloma venereum (1, 3). Infections with *C. trachomatis* genital serovars remain clinically silent in most men and women, but in women, they can ascend to the upper genital tract, leading to pelvic inflammatory disease, infertility, and ectopic pregnancy (4). The single human serovar of *C. pneumoniae* is a common cause of respiratory infection; such infections lead to pharyngitis, bronchitis, and community-acquired pneumonia (1, 2, 5) and have been associated with atherosclerosis (1, 2, 6, 7).

The remaining 9 chlamydial species have animal hosts (8, 9). *C. psittaci* infects birds, and *C. abortus* causes abortion in ruminants; both occasionally cause severe zoonotic human infection. *C. felis* transmitted from cats is thought to sporadically cause human follicular conjunctivitis or atypical pneumonitis (9). Other chlamydial species are endemic in swine (*C. suis*), ruminants (*C. pecorum*), and poultry (*C. gallinacea*), with poorly understood public health impact (8, 9). *C. caviae*, *C. avium*, and *C. muridarum* are found in guinea pigs, birds, and rodents, respectively, but their significance with respect to epidemiology and public health is largely unknown (8, 9).

Nucleic acid amplification tests (NAAT) are most commonly used for diagnosis of chlamydial infections and for DNA sequence-based differentiation of chlamydiae (3, 6, 9–12), but they provide information only at a single point in time. In contrast, serological assays (13–20) have the power to indicate the history of exposure to an infectious agent and are generally preferable to antigen detection for epidemiological or retrospective analyses. The microimmunofluorescence (MIF) test for detection of antichlamydial antibodies has remained the gold standard since its introduction (21–26). MIF is performed as an indirect fluorescent antibody technique that enables microscopic observation of captured antibody on fixed whole chlamydial elementary bodies (EBs) (21, 23, 24, 26). This is a painstaking technique that requires extensive technical expertise, imposing a risk of high interlaboratory variation in results. The high prevalence of *C. pneumoniae* respiratory infection in children (27–29) complicates results of serological studies of *C. trachomatis* due to the possibility of seropositivity arising from a remote *C. pneumoniae* infection.

Several studies evaluated the suitability of *C. trachomatis*- or *C. pneumoniae*-specific ELISAs for analysis of immunodominant antigens (13–20). The majority of such antigens are highly conserved within the *Chlamydia* genus, and determination of suitable antigens for species-specific and sensitive ELISAs is difficult (13, 16–18). ELISAs based on whole elementary bodies (EBs), lipopolysaccharide (LPS), major outer membrane protein (MOMP), Omp2, or Hsp60 suffer from lack of specificity due to cross-reactivity of ELISA antigens (13, 16–18). The Pgp3 protein expressed by the chlamydial plasmid has been extensively studied as a candidate *C. trachomatis*-specific antigen (13, 18, 30–34). This plasmid protein offers the advantage that it is rarely found in human *C. pneumoniae* isolates (35); thus, *C. pneumoniae* infections would not confound the specificity of human Pgp3 *C. trachomatis* serology. Recently, Horner et al. (34) remedied suboptimal Pgp3 ELISA sensitivity with a double-antigen sandwich *C. trachomatis* Pgp3 ELISA, a method that is, however, cumbersome and labor intensive. In addition, the Pgp3 protein is present and highly conserved in most other *Chlamydia* spp. infecting animal hosts (35, 36) and thus cannot be used to resolve
cross-reactivity concerns after human exposure to these animal chlamydiae, while it also may be absent in certain *C. trachomatis* strains following the loss of the plasmid (37, 38). Thus, assays that determine antibody responses against a wide spectrum of *C. trachomatis*-specific antigens are still needed to improve the specificity and sensitivity of *C. trachomatis* serology.

We previously identified *Chlamydia* species-specific immunodominant B cell epitopes using mouse hyperimmune sera generated by three high-dose intranasal inoculations of mice with live chlamydial organisms (39). These B cell epitopes were used as peptide antigens in ELISAs for detection of species-specific anti-*Chlamydia* antibodies. Reactivities of *C. pecorum*-specific peptide antigens were confirmed with sera from cattle, the natural host of *C. pecorum* (39). However, reactivity of *C. trachomatis* and *C. pneumoniae* peptide antigens has not been confirmed with sera of naturally infected humans.

In the present study, we first expanded our bank of *Chlamydia* species-specific mouse sera to all 11 chlamydial species and used it to identify additional immunodominant B cell epitopes. We then confirmed that the mouse-identified *C. trachomatis* B cell epitopes (referred to as “host-independent” epitopes) are similarly immunodominant in humans, using sera from young women actively infected with *C. trachomatis*, many of whom also had documented a prior chlamydial genital infection(s). Since antibody responses can be influenced by host-dependent expression of protein antigens in natural versus natural hosts (40), we also sought to identify “host-dependent” *C. trachomatis* B cell epitopes that might be immunodominant in human infection but fail to elicit antibodies in the heterologous murine host. Since *C. pneumoniae* is a common infectious pathogen in children and young adults, we also utilized the human sera to probe for host-independent and -dependent B cell epitopes specific for this respiratory pathogen. Using this approach, we have identified highly species-specific *C. trachomatis* and *C. pneumoniae* peptide antigens.

**RESULTS**

**Mouse-reactive Chlamydia species-specific peptide antigens.** Using monospecies-specific mouse antiserum pools, species-specific reactivities of *Chlamydia* peptide antigens were first confirmed or identified for *C. avium* and *C. gallinacea* (Fig. 1 and Table S1). The majority of peptide antigens (49 of 60; Fig. 1 and Table S1) reacted with high specificity only with homologous sera. Only 11 peptides (18%) showed cross-reactivity with 1 or more of the 10 heterologous serum pools, in addition to reacting with the corresponding homologous serum pools (Fig. 1 and Table S1). However, these cross-reactivities were observed among closely related chlamydial species, and the majority of the observed cross-reactivities were weak. These results clearly show that these peptide antigens can be used for *Chlamydia* species-specific detection of anti-*Chlamydia* antibodies.

**Antibodies against Chlamydia spp. in women with *C. trachomatis* infections.** To ascertain the suitability of the sera for identification of immunodominant *C. trachomatis* B cell epitopes, the set of 60 *Chlamydia* species peptide antigens (Fig. 1) was tested with polyclonal anti-IgG and monoclonal anti-IgG1 and anti-IgG3 conjugates for reactivity with the *C. trachomatis*-positive serum pools (Fig. 2). All 10 *C. trachomatis* peptides showed consistently strong IgG, IgG1, and IgG3 reactivity with the *C. trachomatis*-positive pool (Fig. 2). This result confirmed that both long-lived IgG1 and short-lived IgG3 were present in the *C. trachomatis*-positive serum pool, and this finding was in agreement with the recent *C. trachomatis* infection history of the study subjects. None of the *C. trachomatis* peptide antigens showed reactivity with the *C. trachomatis*-negative serum pool (Fig. 2), confirming the specificity of these antigens. Further evidence for the overall specificity of peptide antigens is the fact that all 10 peptides of *C. muridarum*, a chlamydial species closely related to *C. trachomatis*, showed no cross-reactivity with the anti-*C. trachomatis* sera.

*C. pneumoniae* peptides from non-OmpA source proteins (PmpD, IncA, CT529, and CT618) showed substantial reactivities with the *C. trachomatis*-positive serum pools, particularly for long-lived total IgG and IgG1 antibodies (Fig. 2). These data suggest that
the study human sera, particularly the sera from 125 women with active *C. trachomatis* infection, also retained moderate amounts of anti-*C. pneumoniae* antibodies. Combined with the reactivity of only a single CT529 peptide with short-lived IgG3 antibodies, these results mainly represent past *C. pneumoniae* infections. Importantly, weak reac-

| Peptide antigens | Anti-*Chlamydia* monospecies-specific sera |
|------------------|------------------------------------------|
| Strain of *Chlamydia* spp., source protein, and peptide position | C. trachomatis (Ctr) C. muridarum (Cmu) C. suis (Csu) C. pneumoniae (Cpn) C. psittaci (Cps) C. abortus (Cab) C. caviae (Cca) C. felis (Cfe) C. pecorum (Cpe) C. avium (Cav) C. gallinacea (Cga) |
| D/UW-3/CX_OmpA_82-105 | [Green] [Red] [White] [White] |
| D/UW-3/CX_OmpA_152-167 | [Green] [Red] [White] [White] |
| D/UW-3/CX_OmpA_226-265 | [Green] [Red] [White] [White] |
| D/UW-3/CX_OmpA_313-339 | [Green] [Red] [White] [White] |
| D/UW-3/CX_PmpD_336-365 | [Green] [Red] [White] [White] |
| D/UW-3/CX_IncE_106-120 | [Green] [Red] [White] [White] |
| D/UW-3/CX_IncE_108-147 | [Green] [Red] [White] [White] |
| D/UW-3/CX_CT529_200-239 | [Green] [Red] [White] [White] |
| D/UW-3/CX_CT618_58-73 | [Green] [Red] [White] [White] |
| Nigg_OmpA_91-104 | [Green] [Red] [White] [White] |
| Nigg_OmpA_160-170 | [Green] [Red] [White] [White] |
| Nigg_OmpA_240-257 | [Green] [Red] [White] [White] |
| Nigg_OmpA_320-335 | [Green] [Red] [White] [White] |
| Nigg_PmpD_724-739 | [Green] [Red] [White] [White] |
| Nigg_PmpD_1038-53 | [Green] [Red] [White] [White] |
| Nigg_IncG_124-139 | [Green] [Red] [White] [White] |
| Nigg_CT529_207-222 | [Green] [Red] [White] [White] |
| Nigg_CT618_100-205 | [Green] [Red] [White] [White] |
| MD6_S_OmpA_139-154 | [Green] [Red] [White] [White] |
| MD6_S_OmpA_300-315 | [Green] [Red] [White] [White] |
| 99DC1_IncA_319-298 | [Green] [Red] [White] [White] |
| 99DC1_CT529_207-236 | [Green] [Red] [White] [White] |
| CWL029_OmpA_89-104 | [Green] [Red] [White] [White] |
| CWL029_OmpA_154-173 | [Green] [Red] [White] [White] |
| CWL029_OmpA_242-257 | [Green] [Red] [White] [White] |
| CWL029_PmpD_654-693 | [Green] [Red] [White] [White] |
| CWL029_PmpD_1131-70 | [Green] [Red] [White] [White] |
| CWL029_IncA_331-370 | [Green] [Red] [White] [White] |
| CWL029_CT529_236-275 | [Green] [Red] [White] [White] |
| CWL029_CT618_201-216 | [Green] [Red] [White] [White] |
| 02DC15_OmpA_159-181 | [Green] [Red] [White] [White] |
| 02DC15_OmpA_250-265 | [Green] [Red] [White] [White] |
| 02DC15_OmpA_333-348 | [Green] [Red] [White] [White] |
| 02DC15_PmpD_49-86 | [Green] [Red] [White] [White] |
| 02DC15_PmpD_1039-92 | [Green] [Red] [White] [White] |
| 02DC15_IncA_331-360 | [Green] [Red] [White] [White] |
| 02DC15_CT529_105-134 | [Green] [Red] [White] [White] |
| 02DC15_CT618_189-228 | [Green] [Red] [White] [White] |
| 526/3_OmpA_89-104 | [Green] [Red] [White] [White] |
| 526/3_PmpD_576-605 | [Green] [Red] [White] [White] |
| 526/3_IncA_124-139 | [Green] [Red] [White] [White] |
| 526/3_CT529_207-222 | [Green] [Red] [White] [White] |
| QFIC_OmpA_159-168 | [Green] [Red] [White] [White] |
| QFIC_IncA_316-355 | [Green] [Red] [White] [White] |
| QFIC_CT618_134-163 | [Green] [Red] [White] [White] |
| QFIC_CT618_224-249 | [Green] [Red] [White] [White] |
| Fe/C-56_OmpA_160-175 | [Green] [Red] [White] [White] |
| Fe/C-56_PmpD_1005-1024 | [Green] [Red] [White] [White] |
| Fe/C-56_IncA_108-137 | [Green] [Red] [White] [White] |
| ESB_IncA_311-326 | [Green] [Red] [White] [White] |
| ESB_CT529_200-248 | [Green] [Red] [White] [White] |
| 10DC88_IncA_319-348 | [Green] [Red] [White] [White] |
| 02DC15_OmpA_309-324 | [Green] [Red] [White] [White] |
| 02DC15_IncA_331-360 | [Green] [Red] [White] [White] |
| 02DC15_CT529_207-236 | [Green] [Red] [White] [White] |
| 02DC15_CT618_105-134 | [Green] [Red] [White] [White] |
| 02DC15_CT618_189-228 | [Green] [Red] [White] [White] |

**FIG 1** Reactivities of 60 peptide antigens from 11 *Chlamydia* species with *Chlamydia* species-specific mouse sera. Each peptide was ELISA tested with 11 pools of 9 to 50 hyperimmune mouse sera obtained by 3× intranasal inoculation with live inocula of a single chlamydial species (39). Green cells represent the reactivity of peptide antigens with their corresponding homologous antisera pools. Red cells indicate peptide antigen cross-reactivity with nonhomologous antiserum (ELISA signals > background + 2 standard deviations [SD]). Green and red color intensities indicate signal strength, and white cells indicate nonreactivity. Peptide designations consist of three-letter *Chlamydia* species acronyms (defined in the headings of columns 3 to 13) followed by strain, source protein, and the amino acid positions of the peptide in the protein. RLU indicates relative light units per second.
The reactivity of only a few peptide antigens of *C. abortus*, *C. felis*, *C. caviae*, *C. pecorum*, and *C. suis* suggests sporadic exposure of human hosts to *Chlamydia* spp. from nonhuman hosts (Fig. 2), given that these peptide antigens showed highly species-specific reactivity with mouse sera (Fig. 1).

**FIG 2** Reactivities of the 60 mouse antiserum-reactive chlamydial peptide antigens with *C. trachomatis*-positive (Ctr-Pos) and -negative (Ctr-Neg) human serum pools. The reactivity of each of the 60 *Chlamydia* species-specific peptide antigens (Fig. 1) was tested with the human serum pools. The *C. trachomatis*-positive pool consisted of sera from 125 women with *C. trachomatis* infection, and the *C. trachomatis*-negative pool consisted of sera from 17 women never diagnosed with *C. trachomatis* infection who were EB ELISA negative for anti-*C. trachomatis* antibodies. Polyclonal anti-human IgG HRP conjugate was used for detection of bound total IgG, and monoclonal antibody conjugates were used to detect bound long-lived IgG1 or short-lived IgG3 isotypes. Peptide reactivities are shown in Log2 RLU signal bars. Different colors are used for the chlamydial species for convenient visualization.
Reactivities of host-independent peptide antigens with anti-*C. trachomatis* MIF titer-ranked subpools. To determine anti-*C. trachomatis* antibody-dependent signal intensity, the set of 60 mouse-reactive *Chlamydia* species peptides was also tested for reactivity with the four human serum subpools ranked by the *C. trachomatis* MIF titers of the constituent individual sera (Fig. 3). The reactivities of 10 *C. trachomatis* peptide antigens correlated strongly with the *C. trachomatis* MIF titers of 4 *C. trachomatis*-positive serum subpools (R = 0.80; P < 10⁻⁶). For instance, the serum subpool with the highest anti-*C. trachomatis* MIF titers (1:256 to 1:2,048) showed the highest reactivity with all 10 individual *C. trachomatis*-specific peptides (Fig. 3). Signals above background were observed for 3 *C. pneumoniae* peptides and six peptides from other *Chlamydia* species showed a signal above background that did not correlate with MIF titers (P = 0.78).

**FIG 3** Anti-*C. trachomatis* MIF titer-dependent IgG reaction intensity of the 60 chlamydial peptide antigens. *C. trachomatis*-positive sera from 108 women with known anti-*C. trachomatis* microimmunofluorescence (MIF) test results were combined by MIF titer into 4 subpools of 19 to 35 sera. Reactivities of the 4 subpools, indicated by color intensity, with peptides from all 11 chlamydial species (Fig. 1) are shown in Log₂ RLU IgG signal bars on the scale as described for Fig. 2. All 10 *C. trachomatis* peptides showed a signal above background, and the signals correlated highly significantly with MIF titers (R = 0.80; P < 10⁻⁶). Three *C. pneumoniae* peptides and six peptides from other *Chlamydia* species showed a signal above background that did not correlate with MIF titers (P = 0.78).
matis, but not those of other chlamydial species, showed anti-\textit{C. trachomatis} MIF titer-dependent reactivity.

Overall, these results confirmed the specific reactivity of the \textit{C. trachomatis} peptide antigens (Fig. 1 to 3). Additionally, the presence of high-titer anti-\textit{C. trachomatis} antibodies in the serum pools from infected women (Fig. 2 and 3) indicated that these sera were suitable for discovering additional immunodominant \textit{C. trachomatis} B cell epitopes that had not previously been recognized by mouse hyperimmune sera (39). Antibodies against \textit{C. pneumoniae} were the second most prevalent antichlamydial antibodies in the human sera (Fig. 2 and 3), in agreement with the high prevalence of \textit{C. pneumoniae} infections in humans (1, 2, 5–7). This opened the possibility to also identify, by the use of these sera, \textit{C. pneumoniae} B cell epitopes that were not recognized by anti-\textit{C. pneumoniae} sera from the heterologous murine host.

Identification of host-dependent B cell epitopes of \textit{C. trachomatis} and \textit{C. pneumoniae}. In the preceding study that identified the set of \textit{Chlamydia} species-specific peptide antigens (39), many more predicted peptides did not react with hyperimmune mouse sera. Therefore, in the present study, we rescreened the \textit{C. trachomatis} and \textit{C. pneumoniae} peptides of the library of mouse nonreactive antigens with the human \textit{C. trachomatis}-positive and -negative serum pools. Of the 271 \textit{C. trachomatis} and 153 \textit{C. pneumoniae} peptides tested, the set of 38 reactive \textit{C. trachomatis} and 8 reactive \textit{C. pneumoniae} peptides is shown in Fig. 4. These \textit{C. trachomatis} peptides reacted, almost uniformly, with the \textit{C. trachomatis}-positive serum pool but not with the \textit{C. trachomatis}-negative pool, indicating high specificity (Fig. 4). As expected, mouse nonreactive \textit{C. pneumoniae} peptide antigens also reacted with these human serum pools (Fig. 4).

Reactivities of host-dependent peptide antigens with anti-\textit{C. trachomatis} MIF titer-ranked subpools. To determine the \textit{C. trachomatis} MIF titer-dependent reactivity of the 46 host-dependent peptide antigens (Fig. 4), they were tested with the four \textit{C. trachomatis} MIF titer-ranked human serum subpools (Fig. 5). Similarly to the host-independent peptides (Fig. 3), the \textit{C. trachomatis} peptides, except for the Hsp60 and Pgp3 peptides, reacted most intensely with the subpool with the highest MIF titer (Fig. 5). The reactivities of \textit{C. trachomatis} peptides also correlated strongly with the MIF titers of the serum subpools ($R = 0.79; P < 10^{-6}$), except for Hsp60 and Pgp3 peptides. However, the reactivities of \textit{C. pneumoniae} peptides did not correlate with the \textit{C. trachomatis} MIF titers ($P = 0.61$). Thus, host-dependent peptide antigens of \textit{C. trachomatis}, but not of \textit{C. pneumoniae}, showed anti-\textit{C. trachomatis} MIF titer-dependent reactivity.

The overall comparison of murine and human host responses to \textit{C. trachomatis} proteins indicates that a much wider spectrum of proteins is recognized by the antibodies of natural human hosts of \textit{C. trachomatis} (Fig. 2 to 5: OmpA, PmpD, IncE, IncG, CT529, CT618, CT442, TarP, CT143, CT813, CT795, CT223, PmpC, CT875, CT579, LcrE, IncA, CT226, CT694, Hsp60, and pGP3) than by those of natural murine hosts (Fig. 1: OmpA, PmpD, IncE, IncG, CT529, and CT618). Similarly, humans also produce antibodies against a wider range of \textit{C. pneumoniae} proteins (Fig. 2 to 5: OmpA, PmpD, IncA, CT529, CT618, PmpG/I, Cpn0525, and YscC) than the natural murine hosts (Fig. 1: OmpA, PmpD, IncA, CT529, and CT618).

Ranking of candidate \textit{C. trachomatis} and \textit{C. pneumoniae} peptide antigens for serological assays. Table 1 presents the summary of the serological screening results for \textit{C. trachomatis} and \textit{C. pneumoniae} peptide antigens that may be suitable for development of species-specific serology. Ten human host-independent and 38 host-dependent peptide antigens of \textit{C. trachomatis} were ranked by an overall reactivity score (Table 1; see also Table S2 in the supplemental material). Similarly, 10 human host-independent and 8 host-dependent peptide antigens of \textit{C. pneumoniae} were ranked (Table 1). Only 10 \textit{C. trachomatis} peptides and 10 \textit{C. pneumoniae} peptides had been previously identified as B cell epitopes by their reactivity with murine hyperimmune sera (39).
Sequence conservation and probability of peptide antigen cross-reactivity. The intended use of these antigens for species-specific detection of antichlamydial antibodies requires maximum divergence from all other chlamydial species but highest sequence conservation within the target species. By these criteria ($\text{SeqID} \leq 50\%$, $P_{\text{cross}} \leq 0.04$), the vast majority of these peptide antigens were highly specific for *C. trachomatis* and were conserved within all *C. trachomatis* serovars but not in the remaining *Chlamydia* spp. (Table 1; see also Table S2). The 30 *C. trachomatis* peptides from the following 16 proteins fit these criteria: CT442, IncE, PmpD, TarP, CT618, CT529, CT813, CT875, CT795, PmpC, IncG, LcrE, IncA, CT694, and CT226. The exceptions with a high likelihood of cross-reactivity with other chlamydial species were two *C. trachomatis* Hsp60 peptides (74% to 85% SeqID) and two pGP3 peptides (40% to 80% SeqID). In addition, 9 peptides of 7 *C. trachomatis* proteins (OmpA, CT618, CT143, CT529, CT579, LcrE, CT875) had $P_{\text{cross}}$ values of 0.16 to 0.89 only with *C. suis* or *C. muridarum* (40% to 93% SeqID), which are closely related to *C. trachomatis*.

Five B cell epitope regions of the strongly reactive *C. trachomatis* OmpA (64% to 76% SeqID) and TarP (77% to 90% SeqID) antigens are polymorphic within *C. trachomatis*. As
indicated in Table S2, the OmpA peptides (variable domain 1 [VD1] to VD4) showed high levels of sequence polymorphism within 22 strains that represent the major clades of *C. trachomatis*. The highest levels of sequence polymorphisms (Table 1; see also Table S2) were observed within the VD1 region, with only 64% average sequence identity across these 22 *C. trachomatis* strains (average *P*<sub>cross</sub> value, 0.18), followed by 74% identity for VD4 (*P*<sub>cross</sub> = 0.42), 76% for VD2 (*P*<sub>cross</sub> = 0.48), and 94% for VD3 (*P*<sub>cross</sub> = 0.89). Therefore, these reactive regions with high levels of sequence polymorphism have the potential for use in the design of peptide antigens that discriminate among antibodies against different *C. trachomatis* serovariants.

In contrast, the 13 *C. pneumoniae* peptides of the following 9 proteins were conserved within *C. pneumoniae* strains but highly divergent from those of other chlamydial species: IncA, PmpD, PmpG/I, CT618, PmpG, OmpA, Cpn0525, YscC, and CT529 (Table 1; see also Table S2). The exceptions with a high likelihood of reactivity with other chlamydial species were 5 peptides from *C. pneumoniae* proteins PmpG/I (40% to 69% SeqID), OmpA (40% to 81% SeqID), and Cpn0525 (40% to 75% SeqID).

**Reactivities of *C. trachomatis*- and *C. pneumoniae*-specific peptides with individual sera.** To further characterize peptide antigens by testing with individual rather than pooled sera, the most highly ranked 38 *C. trachomatis* and 18 *C. pneumoniae* peptides (Table 1) were tested with a panel of 48 individual serum samples from women with *C. trachomatis* infection (Fig. 6). Results showed that the newly identified host-dependent epitopes had reaction strengths and frequencies similar to those seen with the host-independent epitopes identified previously with mouse sera (Fig. 6; see also Table 1).
| Peptidea | Sequenceb | Pooled human sera | Mouse serum statusc | Peptide sequence % identity | Probability of cross-reactivity (Pcross)d |
|---------|-----------|-------------------|--------------------|--------------------------|-------------------------------------|
| CtrOmpA_313-339 | IFDITLNPITIAAGDVKTAEGQLGSD | 10.3 | 1 | + | 74 | <40 | 0.55 | 0.01 |
| CtrOmpA_226-265 | NVLCNAAEFITNKPKGYYQKEFPLDLTATGDAATGTDAS | 8.0 | 5 | + | 94 | 55 | 0.89 | 0.09 |
| CtrOmpA_152_167 | SFNLVGLFNDENQKT | 2.4 | 27 | + | 76 | 50 | 0.29 | 0.04 |
| CtrOmpA_82-105 | FQMGAKPITTDGTNSAAPSTLTARE | 3.9 | 39 | + | 64 | 40 | 0.06 | 0.01 |
| CtrCT44_135-150 | YVESLRRNLSVLQDTQ | 8.4 | 22 | 3 | - | 89 | <40 | 0.01 | 0.01 |
| CtrIncE_81-120 | LFAISLDDLVHEGLGVCPFLCKSSPAEMPVTQVFQKGG | 8.4 | 3 | 3 | - | 97 | <40 | 0.01 | 0.01 |
| CtrIncE_117-132 | FKNGKSGAKVLIYTO | 12.7 | 6.5 | 17 | - | 97 | <40 | 0.01 | 0.01 |
| CtrPmpD_727-742 | EKVVEPPAPAEQKDNN | 8.2 | 4 | 4 | - | 100 | <40 | 0.01 | 0.01 |
| CtrPmpD_536-555 | ARPAQOLPQTEPFLKSEKPRGLSSYSG | 7.7 | 6 | 6 | - | 100 | <40 | 0.01 | 0.01 |
| CtrPmpD_1036-65 | SGTPQGQHISKPEAIESSSEPEGAHSL | 6.8 | 14 | + | 98 | <40 | 0.01 | 0.01 |
| CtrPmpD_760-775 | QALFASEGDGLPSSESS | 6.6 | 16 | - | 95 | <40 | 0.01 | 0.01 |
| CtrTarP_151-180 | SSNYYDAADYEPRTTENIYEGGSRTS | 7.4 | 7 | - | 95 | <40 | 0.01 | 0.01 |
| CtrTarP_116-145 | TSSSHIPSYYDDGVSNSGDINYDDVGS | 6.3 | 21 | - | 77 | <40 | 0.01 | 0.01 |
| CtrTarP_63-78 | TVVNITYNSASPNTVT | 4.4 | 33 | - | 94 | <40 | 0.01 | 0.01 |
| CtrTarP_406-421 | FSFXGDIWDLSVEPMV | 3.7 | 40 | - | 90 | <40 | 0.07 | 0.01 |
| CtrCT618_185-206 | GNKQKNKPTGTEGSNFGMARL | 7.4 | 8 | - | 99 | <40 | 0.01 | 0.01 |
| CtrCT618_58-73 | TVSETQQQQLTETTI | 5.0 | 31 | + | 100 | <40 | 0.04 | 0.01 |
| CtrCT618_101-122 | KTPNPGDFQLPDVPSQRTTLSP | 4.2 | 37 | - | 98 | <40 | 0.39 | 0.01 |
| CtrCT143_2-27 | KRPVFTGQPGRPGSTEEGTGVKQDNQ | 7.3 | 9 | - | 100 | <40 | 0.20 | 0.01 |
| CtrCT529_200-239 | SAERADCEACRARIARELLELVQGENACCKVAGEKAK | 7.3 | 10 | + | 96 | <40 | 0.01 | 0.01 |
| CtrCT529_197-132 | KAFFTQPNKMARVNN | 4.0 | 38 | - | 44 | <40 | 0.16 | 0.02 |
| CtrCT529_105-120 | SHMAKASQKTQGEDG | 3.4 | 42 | - | 96 | <40 | 0.01 | 0.01 |
| CtrCT83_235-284 | AENLDLMAYAMEEFEKEKHPGKRGS | 7.2 | 11 | + | 100 | <40 | 0.01 | 0.01 |
| CtrCT83_203-232 | TVYTDLEAQKOLEEKYVTDLESEKOEELREL | 6.0 | 23 | 100 | <40 | 0.01 | 0.01 |
| CtrCT875_398-427 | KGTHYRAPRDLLEPSGASLAELTARFADD | 5.9 | 24 | - | 100 | <40 | 0.03 | 0.01 |
| CtrCT795_148-163 | IMDIFTEIPSNPVEFVE | 7.0 | 12 | - | 99 | <40 | 0.02 | 0.01 |
| CtrCT223_5-34 | ALGTSNGVEAINGNDLSPAPEAKKTTGSL | 7.0 | 13 | - | 97 | <40 | 0.01 | 0.01 |
| CtrPmpC_483-498 | APSLEAESDQDOTQTE | 6.8 | 15 | - | 97 | <40 | 0.01 | 0.01 |
| CtrPmpC_608-637 | AIVESTPEAEIPYVEGEAETATPDNSN | 6.4 | 20 | - | 99 | <40 | 0.01 | 0.01 |
| CtrPmpC_222-301 | ETEQTESNGQDGSSSETDQVESSEPSPT | 4.5 | 32 | - | 99 | <40 | 0.01 | 0.01 |
| CtrPmpC_1299-14 | EQQNNDASNGESAN | 3.7 | 41 | - | 98 | <40 | 0.01 | 0.01 |
| CtrIncG_108-147 | RPSQOEFGSRLSEASAPQSAPSTSFTGLESALRSGDS | 6.5 | 18 | + | 98 | <40 | 0.01 | 0.01 |
| CtrIncG_97-112 | KRSPFEEIGGNAPQDO | 6.4 | 19 | - | 100 | <40 | 0.02 | 0.01 |
| CtrIncG_149-164 | SGAFFDINDKNSR5S | 5.3 | 40 | + | 97 | <40 | 0.02 | 0.01 |
| CtrCT757_256-286 | ALDDVAGTATAVAGAKTSAGAASASAAATK | 5.6 | 25 | - | 100 | 42 | 0.52 | 0.03 |
| CtrCT759_100-129 | AQAVHGRDSGNSGDSLPSPTGTEVG | 2.1 | 45 | - | 100 | <40 | 0.69 | 0.01 |
| CtrLcE_392-421 | RSFSSTPSPHVPQPQSEIPTSPTSTOPSP | 5.4 | 26 | - | 100 | 42 | 0.06 | 0.01 |
| CtrLcE_242-271 | ATWEDINHHRLPWCWEDEETKTNKPPFLIQML | 2.9 | 43 | - | 100 | <40 | 0.16 | 0.01 |
| CtrCT875_228-257 | NDPILGRTPNQSKNPQETYGQNSMFYDGP | 5.4 | 28 | - | 100 | <40 | 0.52 | 0.01 |
| CtrCT787_486-515 | OQHYQDPRSPDFYDLPALSYDYLPRSPYPTP | 5.4 | 29 | - | 100 | <40 | 0.24 | 0.02 |
| CtrIncA_234-273 | SKTLTSQALOREKSSDLCSQRELSPRSPKASPTKSS | 4.3 | 34 | - | 100 | <40 | 0.01 | 0.01 |
| CtrCT694_64-93 | NRGTTTPSRVTQAINHPISTISQGAQOP | 4.2 | 36 | - | 93 | <40 | 0.06 | 0.01 |
| CtrCT694_267-296 | ENEMNQIILQDGQDQPDQVHQDQDNKELQK | 2.5 | 44 | - | 98 | <40 | 0.05 | 0.01 |
| CtrCT226_147-176 | AQVSDKDELAQKKEIQEQLQGSLKVCLEESLI | 4.2 | 35 | - | 97 | <40 | 0.01 | 0.02 |

(Continued on next page)
| Peptide Sequence | Pooled human sera | Peptide sequence % identity | Probability of cross-reactivity (P<sub>cross</sub>) | Mouse serum status | C. trachomatis strain<sup>f</sup> | C. pneumoniae strain<sup>g</sup> | C. suis or C. muridarum<sup>h</sup> | Chlamydia spp<sup>i</sup> |
|------------------|-------------------|-----------------------------|-----------------------------------------------|-------------------|--------------------------|--------------------------|--------------------------|--------------------------|
| CtrHsp60_519-544 | TEALIAEIPEEKPAAPAMPGAGMDY | 6.3 22 - | 100 81 | 0.95 0.74 | - | - | - | 0.74 |
| CtrHsp60_338-367 | EKEALEARCEIKKOQEDSSDYEDEKQLE | 0.2 48 - | 100 70 | 0.89 0.69 | - | - | - | 0.69 |
| CtrpGP3_37-66   | GTKSTPVAAMTDGSLTVSNNSSTNAS | 2.0 46 - | 96 &lt;40 | 0.24 0.07 | - | - | - | 0.07 |
| CtrpGP3_198-227 | SSGVPNCLSRLTSINTGLPTPTSLRGVGG | 0.9 47 - | 97 &lt;40 | 0.61 0.61 | - | - | - | 0.61 |
| CnpiAC_331-370  | QKAES3FIAVCVRDFRGGRETPTPVVEGDESEQEDEG | 3.5 1 + | &lt;40 95 | 0.01 0.01 | - | - | - | 0.01 |
| CnpPmpG/I_1223-38 | HGQVSYGRNHNNMITTK | 4.6 2 - | &lt;40 100 | 0.01 0.04 | - | - | - | 0.04 |
| CnpPmpG/I_947-62 | AGTTLETTTNNTDGS | 5.2 3 - | &lt;40 100 | 0.01 0.04 | - | - | - | 0.04 |
| CnpPmpG/I_714-729 | INNTARSGGSGYAPK | 6.5 9 - | 63 100 | 0.20 0.29 | - | - | - | 0.29 |
| CnpPmpG/I_232-237 | TADKGGYGGYSEKDS | 5.7 14 - | 44 98 | 0.01 0.02 | - | - | - | 0.02 |
| CnpPmpD_1131-70 | NKEETLVASAVQWMSSTPNKDADVPTVLDLSISITVD | 1.7 4 + | &lt;40 100 | 0.01 0.01 | - | - | - | 0.01 |
| CnpPmpD_521-536 | RSNPKLEQKDSGNNIN | 1.5 5 - | &lt;40 100 | 0.01 0.01 | - | - | - | 0.01 |
| CnpPmpD_654-693 | EKSLNACHDHYPKPTVEEVPSLLEEHVPSSTDIRG | 1.3 10 + | &lt;40 100 | 0.01 0.01 | - | - | - | 0.01 |
| CnpPmpD_147-186 | EKISSDTREKDKLEDEPSKSKGLKVEVSDLPKSTAV | 0.2 12 + | &lt;40 100 | 0.01 0.01 | - | - | - | 0.01 |
| CnpCT618_201-216 | PEISIDPMNKPSAE | 0.2 6 + | &lt;40 100 | 0.01 0.01 | - | - | - | 0.01 |
| CnpOmpA_158-173 | FGKVTGVANELPVW | 0.2 7 + | &lt;40 96 | 0.01 0.07 | - | - | - | 0.07 |
| CnpOmpA_309-324 | AVLNLTAWSPLLGNAA | 0.2 15 + | &lt;40 98 | 0.01 0.64 | - | - | - | 0.64 |
| CnpOmpA_89-104 | PTGAAANNYYTAVDRP | 0.0 16 + | &lt;40 98 | 0.01 0.46 | - | - | - | 0.46 |
| CnpOmpA_242-257 | VAFPLPDAGVATAGT | 0.0 17 + | &lt;40 100 | 0.01 0.16 | - | - | - | 0.16 |
| CnpPmpG_88-903 | LLRGLNVNYVNSCEL | 3.3 8 - | &lt;40 100 | 0.01 0.01 | - | - | - | 0.01 |
| Cnp0525_239-254 | AQNSTAKRRRRRAV | 4.8 11 - | 63 100 | 0.29 0.46 | - | - | - | 0.46 |
| CnpOmpA_77-92 | HTKXTPPGSPKSVS | 2.1 13 - | &lt;40 100 | 0.01 0.01 | - | - | - | 0.01 |
| CnpCT529_236-275 | RASKYNLRELANILONGQGDAVLSAERALRKHEVATL | 1.5 18 + | &lt;40 100 | 0.01 0.01 | - | - | - | 0.01 |

*a* A set of 48 C. trachomatis (Ctr) and 18 C. pneumoniae (Cpn) peptide antigens with highest reactivity is shown. Included are 10 each of previously identified C. trachomatis and C. pneumoniae mouse serum-reactive host-independent peptides (39) and 46 top-ranked host-dependent peptides identified in this investigation by screening 424 mouse nonreactive peptides.

*b* Only the actual chlamydial sequence of peptide antigens is shown, without the N-terminal biotin or the serine-glycine-serine-glycine spacer that is attached to each peptide (39).

*c* Reactivity scores are weighted averages of seven values corresponding to the reactivities of total and subpooled sera as described in Materials and Methods.

*d* The rank of each peptide within the list of 48 C. trachomatis or 18 C. pneumoniae B cell epitopes is shown. C. trachomatis rank is based on reactivity score, and C. pneumoniae rank is based on reaction frequency with 48 individual sera shown in Fig. 6.

*e* Reactivity (+) or lack of reactivity (−) with C. trachomatis-specific or C. pneumoniae-specific hyperimmune mouse sera is shown (39).

*f* Average percent amino acid sequence identity with 22 strains representing all major clades of C. trachomatis (Table S2) is shown. Sequences with identities below 40% typically cannot be aligned correctly, and the probability of peptide cross-reactivity is less than 1% (39).

*g* Average percent amino acid sequence identity with 6 major strains of C. pneumoniae (Table S2) is shown.

*h* Maximum probability of cross-reactivity (39) with C. muridarum- or C. suis-specific sera based on sequence identity (Table S2) is shown.

*i* Maximum probability of cross-reactivity (39) with sera specific for the remaining 8 Chlamydia spp. (Table S2) is shown.
Peptide/MIF Test specificity

| Serum # | C. trachomatis (Ctr) MIF | C. pneumoniae (Cpn) MIF |
|---------|--------------------------|------------------------|
|         |                          |                        |

**FIG 6** IgG1 and IgG3 antibodies against *C. trachomatis* and *C. pneumoniae* peptides in 48 individual sera. Sera from 48 *C. trachomatis*-positive women that covered the spectrum of *C. trachomatis* and *C. pneumoniae* MIF titers (rows 2 and 3), as well as the racial origin of the study subject, were selected. A total of 56 unique *C. trachomatis* and *C. pneumoniae* peptide antigens with potential utility for human population surveys were tested with these individual sera. These peptides are divided into 6 sets based on *C. trachomatis* and *C. pneumoniae* species, serovar specificity, source proteins, reaction
Correlation of *C. trachomatis* and *C. pneumoniae* peptide reactivities and species-specific MIF titers. Reactivity of the 48 individual sera against *C. trachomatis* peptide antigens (peptide sets 1 to 3 in Fig. 6) correlated significantly with the anti-*C. trachomatis* MIF titers of the individual sera (*R* = 0.56 to 0.33; *P* = 10^-4 to 0.02; Table 2). However, the reactivities of *C. trachomatis* peptide set 4 (Hsp60, pGP3; Fig. 6) did not correlate with anti-*C. trachomatis* MIF titers (*P* = 0.42; Table 2). As expected, the signal intensities of all of the sets of *C. trachomatis* peptides (sets 1 to 4) failed to correlate with anti-*C. pneumoniae* MIF titers. Similarly, signals of *C. pneumoniae* peptides (set 5) did not correlate with anti-*C. trachomatis* MIF titers (*P* = 0.66). Surprisingly, the reactivities of *C. pneumoniae* peptides (set 5) did not correlate significantly with anti-*C. pneumoniae* MIF titers (*P* = 0.39). This may have been due to strong human serum reactivities with the non-OmpA proteins of *C. pneumoniae* but weak reactivities with OmpA antigens (major component of MIF antigens). Overall, the results in Table 2 indicate that *C. trachomatis*, but not *C. pneumoniae*, peptide reactivities correlated with the respective MIF titers (Fig. 6). As expected, the ELISA signal intensities of the *C. trachomatis* peptides in set 1 (Fig. 6) showed strong correlation with other *C. trachomatis* peptides (set 2) (*R* = 0.81; *P* = 10^-6; Table 2). In contrast, the reactivities of the *C. pneumoniae* peptides (set 5) showed only weak correlation with the *C. trachomatis* peptides in set 1 (*R* = 0.42; *P* = 0.003) (Table 2).

Reactivities of serovariant peptides from *C. trachomatis* OmpA variable domain 1 (VD1) to VD4. As shown in Fig. 6, the OmpA VD1 peptide of *C. trachomatis* serovar D/UW-3/CX showed IgG1 reactivity with about 44% of the individual sera (21 of 48 sera) and the VD2, VD3, and VD4 peptides with 38%, 50%, and 79% of the sera, respectively. Testing of 5 additional serovariant peptides from OmpA VD1 increased the frequency of individual serum IgG1 reactivity by 20% (65% with six OmpA VD1 serovariant peptides compared to 44% with a single serovar D VD1 peptide). The use of 3 additional serovariant peptides from OmpA VD4 increased the reactivity frequency by only 4% (83% for four VD4 serovariant peptides compared to 79% with one serovar D VD4 peptide). Reactivities of short-lived IgG3 were almost identical to those of long-lived IgG1, except for distinct differences in some individuals (Fig. 6). These exceptions suggest a potential to distinguish current/recent infections indicated by the presence of IgG3 antibodies from past infections indicated by the presence of IgG1 antibodies. Interestingly, the six serovariant peptides from *C. trachomatis* OmpA VD1 generated infrequent but distinct signals (Fig. 6). In contrast, the OmpA VD4 peptides showed frequent and common reactivities. Combined with sequence polymorphism data (Table 1; see also Table S2), this result suggests that the use of several serovariant peptides from the highly polymorphic OmpA VD1 region may identify *C. trachomatis* serovar-specific antibodies and that assaying just a few variant peptides from OmpA VD3 and VD4 may allow detection of all anti-*C. trachomatis* antibodies.

*C. pneumoniae* OmpA and non-OmpA peptide reactivities. In contrast to the highly reactive *C. trachomatis* OmpA peptide antigens (Fig. 6), *C. pneumoniae* OmpA peptides 53 to 56 showed rare and weak reactivity that was much weaker and less frequent than the reactivity seen with dominant peptides 39 to 47 from other *C. pneumoniae* proteins. All four OmpA variable-domain peptides and the non-OmpA peptides (InCA, PmpD, PmpG/I, CT618, PmpG, Cpn0525, and YscC) (Fig. 6) were highly conserved across all strains of *C. pneumoniae* (95% to 100% sequence identity) (*P* = 0.91 to 0.95) (Table 1; see also Table S2). Several of those *C. pneumoniae* non-OmpA peptides (InCA, PmpD, PmpG/I, and CT618; Fig. 6) showed frequent detection of anti-*C. pneumoniae* antibodies.

**FIG 6** Legend (Continued)

frequency, and the immunoglobulin subclasses detected. The host-independent peptides shown in black font had previously reacted with mouse anti-*C. trachomatis* sera (39), while the host-dependent peptides shown in red font were not recognized by mouse sera but showed reactivity in this study with human sera. Blue font indicates eight non-serovar D peptides from OmpA variable domains 1 and 2. The column following the peptide designations shows *C. trachomatis* or *C. pneumoniae* specificity. Individual sera are numerically identified in the top row and are arranged in order of their average level of IgG1 reactivity with the first set of 11 *C. trachomatis* peptides. Peptide reactivities with the individual 48 sera are indicated by color intensities (legend at bottom), and absence of reactivity is shown by white cells.

July/August 2018 Volume 3 Issue 4 e00246-18
TABLE 2 Correlation of *C. trachomatis* and *C. pneumoniae* peptide reactivities with species-specific MIF titers

| Peptide set<sup>a</sup> | Chlamydia species specificity | Source proteins | Antibody isotype | Correlation to<sup>b</sup>: |  |
|-------------------------|------------------------------|-----------------|-----------------|-----------------------------|-----------------|
|                         |                              |                 | C. *trachomatis* | MIF titer      | C. *pneumoniae* | MIF titer      | Peptide set 1 |
|                         |                              |                 |                | P   | R   | P   | R   | P   | R   |
| 1 (peptides 1–11)       | *C. trachomatis*             | OmpA, TarP, CT143, PmpD, IncE, CT529, CT442, CT618 | IgG1            | 10<sup>-4</sup> | 0.56 | 0.91 | –   | –   | –   | –   |
| 2 (peptides 12–22)      | *C. trachomatis*             | CT813, LcrE, CT226, IncE, IncG, CT223, CT618, CT795, PmpC | IgG1            | 10<sup>-3</sup> | 0.47 | 0.20 | –   | –   | <10<sup>-6</sup> | 0.81 |
| 3 (peptides 23–34)      | *C. trachomatis*             | OmpA, variable domains (VD) 1, 2, 3, 4 | IgG1 | 0.02 | 0.33 | 0.32 | –   | 10<sup>-5</sup> | 0.65 |
| 3 (peptides 23–34)      | *C. trachomatis*             | OmpA (VD1 to VD4) | IgG3 | 0.01 | 0.36 | 0.12 | –   | 10<sup>-5</sup> | 0.60 |
| 4 (peptides 35–38)      | *C. trachomatis*             | Hsp60, pGP3     | IgG1 | 0.42 | –    | 0.92  | –   | –   | 0.002 | 0.44 |
| 5 (peptides 39–50)      | *C. pneumoniae*              | IncA, PmpD, PmpG/I, CT618, PmpG, Cpn0525, YscC | IgG1 | 0.66 | –    | 0.39  | –   | 0.003 | 0.42 |

<sup>a</sup>Peptide sets and peptide numbers are identical to those of the peptides shown in Fig. 6.

<sup>b</sup>Pearson’s correlation coefficient (R) and P values were calculated in linear regression analyses. The correlations are based on the mean reactivity of a peptide set with each of 48 individual sera (Fig. 6). –, absence of significant correlation or self-comparison.
DISCUSSION

In this study, we first reconfirmed the specificities of 60 peptide antigens from the 11 *Chlamydia* spp. (39), using *Chlamydia* monospecies-specific hyperimmune mouse sera (Fig. 1 and Table S1). Subsequent testing of these peptides with the human *C. trachomatis*-positive and -negative sera confirmed the *C. trachomatis*-specific reactivity of these peptides with human sera and established the mouse/human host-independent nature of the chlamydial B cell epitopes (Fig. 1, 2, and 3). Next, by screening large numbers of additional *C. trachomatis* candidate peptide antigens with human sera (Table 1), we identified novel human host-dependent B cell epitope peptide antigens of *C. trachomatis* that had not previously reacted with mouse anti-*C. trachomatis* hyperimmune sera (Fig. 4 and 5). Additionally, using the previously identified mouse species-specific peptide antigens (39), we confirmed the presence of both anti-*C. pneumoniae* and anti-*C. trachomatis* antibodies in these human sera (Fig. 6).

Again, by screening a large number of additional *C. pneumoniae* candidate peptide antigens with the human sera (Table 1), we identified novel human host-dependent *C. pneumoniae* B cell epitope peptide antigens (Fig. 4 and 5).

These experiments conclusively established the concept of human host-dependent antibody responses against certain immunodominant B cell epitopes of *C. trachomatis* and *C. pneumoniae*. Overall, the identification of many human host-dependent B cell epitopes from two natural human chlamydial pathogens confirmed the concept that the range of antibody responses to chlamydiae in the natural host species is wider than in the experimental murine host (40).

Compared to the frequent reactivities of *C. trachomatis* and *C. pneumoniae* peptide antigens (Fig. 2 to 5), only few antigens from *C. abortus*, *C. felis*, *C. pecorum*, and *C. psittaci* produced ELISA signals, and the signals that those few produced were very weak (Fig. 2 and 3), suggesting sporadic exposure of the human hosts to those animal *Chlamydia* spp. Additionally, all 10 peptide antigens from *C. muridarum* were nonreactive with the study sera (Fig. 2 and 3), despite the high anti-*C. trachomatis* antibody levels in the *C. trachomatis*-positive serum pool. This indicates that these *C. muridarum* peptide antigens do not cross-react with anti-*C. trachomatis* antibodies, despite minimal evolutionary divergence between these two chlamydial species. Similarly, reactivity of peptides of *C. suis*, another species closely related to *C. trachomatis*, was essentially absent compared to that seen with *C. trachomatis* (Fig. 2 and 3). These results clearly indicate that these peptides are highly species-specific antigens (Fig. 2).

With the completion of this study, we report now an expanded set of peptide antigens that may be valuable in serodiagnosis of human chlamydial infections (Table 1). These peptide antigens are sufficiently divergent to eliminate virtually any cross-reactivity among antibodies against other *Chlamydia* spp. (see Table S2 in the supplemental material). Derived from the most strongly reactive regions of immunodominant proteins identified in the chlamydial proteome (40–43), they produce very strong ELISA signals under conditions of detection with anti-human antibody conjugates (Fig. 6). The distinguishing characteristic of these peptide antigens is that they react highly specifically with their cognate antibodies and can therefore be used in assays for species- and even serovar-specific detection of antibodies against *C. trachomatis* and *C. pneumoniae*. This fills a distinctive void among *Chlamydia* antibody assays for effective species/type-specific detection of antichlamydial antibodies (13–20, 44–48).

A clear absence of reactivity to certain peptides in individual serum samples (“hole in antibody repertoire”), but strong reactivity to others (Fig. 6), points to the molecular uncertainty principle of the antibody response. We attribute this to the stochastic “roll of the dice” nature of immunoglobulin gene recombination (49), which may have failed to create even a single B cell receptor molecule that bound to a B cell epitope, preventing epitope-specific B cell expansion and antibody affinity maturation. These data strongly argue that combined results from multiple peptide antigens, preferably from several proteins, will be required for accurate anti-*C. trachomatis* antibody detec-
tion that approximates the quantitative accuracy of methods using complex multi-
epitope antigens.

*C. trachomatis*-seropositive individuals show holes in the antibody repertoire not
only against individual peptide antigens but also against whole-protein antigens
(40–43, 50–53). Thus, the molecular uncertainty principle of the antibody response
applies also to proteins. The use of single antigens in serology therefore inherently
compromises sensitivity as well as quantitative accuracy. These disadvantageous per-
formance characteristics of the use of single protein or peptide antigens may inherently
impede their suggested use in programs for control and monitoring for trachoma,
sexually transmitted infections (STI), or community-acquired pneumonia (52–56).

Interestingly, *C. pneumoniae* OmpA peptides were strongly reactive with anti-
*C. pneumoniae* mouse sera (Fig. 1), but not with human sera (Fig. 6), while *C. trachomatis* OmpA peptides reacted strongly with anti-*C. trachomatis* mouse sera as well as
human sera (Fig. 1 and 6). In agreement with these findings, Campbell et al. reported
that rabbit anti-*C. pneumoniae* immune sera recognized *C. pneumoniae* OmpA poorly
but that *C. trachomatis* immune sera dominantly recognized the respective OmpA
proteins (57). These results may explain the finding that anti-*C. trachomatis* but not
anti-*C. pneumoniae* MIF titers correlated significantly with the corresponding species-
specific peptide reactivities (Table 2), since OmpA is the dominant MIF antigen. In
agreement with the previous findings (29, 57), these results suggest that the *C. trachomatis* but not the *C. pneumoniae* OmpA peptides would be suitable for sensitive antibody detection in serological assays.

Synthetic OmpA peptides have been tested before for *Chlamydia* species-specific
serology, and OmpA peptide-based assays provided specificity but lacked sensitivity
(58–61). However, those studies used short, spacerless peptide antigens, resulting in
low sensitivity that was most likely due to weak antibody binding to the short peptides
(62) and/or to steric hindrance of antibody binding (63). Recently, several groups have
attempted to identify non-OmpA peptide antigens for *C. trachomatis* and *C. pneu-
moniae* serology and to use levels of antibody against 1 to 3 of these peptides as
markers for disease (64, 65). However, those assays had low sensitivity, possibly because
the peptides do not represent immunodominant B cell epitopes. Therefore, given that
only a fraction of *Chlamydia*-infected individuals respond, those assays are inherently of
limited use as disease markers. And even if the epitopes were immunodominant, use
of multiple antigens is required for sensitive detection of a wide repertoire of anti-
*C. trachomatis* antibodies, as results determined by us and others (40, 41) indicate
(Fig. 6).

These synthetic peptide antigens can be readily commercialized in the standard
ELISA format as well as in the microarray format (63, 66, 67), and such assays may
become widely available tools for chlamydial serology. In a recent study (66), we
detected species- and serovar-specific antibodies against *Chlamydia* spp. by a peptide
microarray that used several of the peptide antigens described here (Fig. 1; see also
Table 1). With the possibility of spotting 196 to 784 peptides on a single chip (67), such
microarrays offer unprecedented opportunities for *Chlamydia* species serology by
reducing the need for labor-intensive multipeptide ELISAs. Averaged microarray signals
of multiple peptide antigens may reliably quantify the overall antichlamydial antibody
response, and specific seroreactivity patterns of individual peptides may be associated
with a risk of sequelae following chlamydial infection. In conclusion, these peptide
antigens will improve *C. trachomatis* and *C. pneumoniae* serology and will thereby help
to more accurately diagnose infections and sequelae and aid in *C. trachomatis* surveil-
lance and control programs (30–34, 50, 59, 68).

**MATERIALS AND METHODS**

**Mouse sera.** *Chlamydia* monospecies-specific mouse sera for each of the 11 *Chlamydia* spp. were
used to confirm the specificity of peptide antigens for detection of species-specific antibody. Nine of such
hyperimmune sera had been raised in our previous study (39), but additional *C. avium*- and *C. gallinacea-
specific sera were raised in the current study. Briefly, EB suspensions of *C. avium* strain 10DC88 and
*C. gallinacea* strain 08-1274/3 were intranasally inoculated in A/J mice as described previously (39). Sera
of 9 to 50 animals infected with *C. avium* or *C. gallinacea* were collected into a monospecies-specific serum pool. All animal experimental protocols were approved by the Institutional Animal Care and Use Committees at Auburn University.

**Human sera.** Sera were collected from 125 women with *C. trachomatis* infection confirmed by NAAT (69) and from 18 healthy, low-risk women who were never diagnosed with *C. trachomatis* infection. *Chlamydia trachomatis* MIF titers (A to I, K, and L1 to L3 antigens) were determined for 108 of the 125 *C. trachomatis* NAAT-positive women at the University of Washington, and 106 were also tested for *C. pneumoniae* (TW183 antigen) (21–24, 26). Serum samples from 18 healthy women at low risk for *C. trachomatis* exposure were tested for antibodies against *C. trachomatis* by an ELISA using a *C. trachomatis* EB antigen (GenWay Biotech, Inc., San Diego, CA). Seventeen anti-*C. trachomatis* EB antibody-negative sera were included in this study as *C. trachomatis*-negative-control sera, and a single positive serum sample was excluded. These samples from the 17 women at low risk for *C. trachomatis* infection were not tested by MIF. The chlamydial antibody status of all human sera was also characterized by the reactivity with mouse serum-reactive peptide antigens of all 11 chlamydial species.

Among the 125 *C. trachomatis*-positive women, 74 were African American, 28 were Caucasian, and 23 were Hispanic, Asian, or mixed race. The 17 *C. trachomatis*-negative healthy women were all Caucasian but were of similar age to the *C. trachomatis*-positive serum donors. The study protocol was approved by both the Institutional Review Boards for Human Research of the University of Pittsburgh and the University of North Carolina. All participants provided written informed consent at the time of enrollment.

**Pooling of human sera.** Aliquots of serum from the 125 women diagnosed with *C. trachomatis* were pooled and are referred to here as the *C. trachomatis*-positive pooled sera. Serum aliquots from the 17 women never diagnosed with *C. trachomatis* were similarly pooled and are referred to here as the *C. trachomatis*-negative pooled sera. The *C. trachomatis* MIF titer for 108 *C. trachomatis*-infected women was known at study onset, and these 108 samples were used to prepare four serum subpools ranked by *C. trachomatis* MIF titer. The serum subpools were prepared by mixing equal proportions of 19, 21, 33, and 35 serum samples from women with *C. trachomatis* MIF titers of 1:4 to 1:8, 1:16 to 1:32, 1:64 to 1:128, and 1:128 to 1:1,048, respectively. To identify additional peptide antigens that showed host-dependent reactivity with human sera, we used the human serum pool or subpools of 125 *C. trachomatis*-positive and 17 *C. trachomatis*-negative women to screen a large panel of 271 *C. trachomatis* and 153 *C. pneumoniae* peptides that had been nonreactive with mouse sera.

**Peptide antigens.** The B cell epitope prediction approach has been described in detail in preceding studies (39, 62). In brief, we first identified and ranked 72 immunodominant proteins among all chlamydial proteomes on the basis of published data (40–43). On the basis of alignments of each of these 72 individual proteins, suitable regions for identification of genus-, species-, and serovar-specific epitope candidates were further subjected to *in silico* B cell epitope analyses (62). The majority of the peptide antigens were selected from polymorphic protein regions for species-specific reactivity and smaller numbers from highly polymorphic regions (serovar-specific reactivity) or conserved regions (genus-specific reactivity). Conserved B cell epitopes that corresponded to a given *Chlamydia* species but were divergent from those of other *Chlamydia* spp. were selected for analyses of the species specificity of the peptide antigens. Divergent B cell epitopes within a *Chlamydia* species were selected for serovar-specific reactivity.

Peptide antigens were chemically synthesized with N-terminal biotin followed by a serine-glycine-serine-glycine spacer and captured on streptavidin-coated white microtiter plates (Fisher Scientific, Roskilde, Denmark). For antigen optimization of some B cell epitope regions, they were scanned by the use of 16- to 40-mer peptides overlapping by 4 to 12 amino acids (aa), and peptides of different lengths corresponding to regions showing peak reactivity were tested to maximize reactivity.

**ELISA.** Primary antibodies were detected with horseradish peroxidase (HRP)-conjugated secondary antibodies in chemiluminescent ELISAs. Overall methods were employed as described in detail by Rahman et al. (39) with the following modifications. The wash buffer consisted of 0.15 M NaCl, 20 mM Tris-HCl (pH 7.5), 0.025% Tween 20, and 0.001% benzalkonium chloride. The assay diluent consisted of 0.125 M NaCl, 20 mM Tris-HCl (pH 7.5), 0.025% Tween 20, 2% rabbit serum, 0.2% bovine serum albumin, 0.2% casein, 0.2% polyethylene glycol, and 0.005% benzalkonium chloride. The blocking buffer consisted of 0.125 M NaCl, 20 mM Tris-HCl (pH 7.5), 2% rabbit serum, 0.2% bovine serum albumin, 0.2% casein, 0.2% polyethylene glycol, and 0.005% benzalkonium chloride. A polyclonal rabbit anti-human IgG·h+1 cross-adsorbed antibody-HRP conjugate was obtained from Bethyl Laboratories (Montgomery, TX) (catalog no. A80-218P). The following monoclonal mouse anti-human antibody conjugates were purchased from Southern Biotech, Birmingham, AL: IgG1 hinge-HRP (catalog no. 9052-05) and IgG3 hinge-HRP (catalog no. 9210-05).

Chemiluminescent detection was chosen because of the signal linearity over 4 to 5 orders of magnitude, and light emission was measured as relative light units (RLU) per second. Background signals were determined for wells without peptide antigen and wells coated with nonspecific peptide antigens (39). The differences between all background signals and all peptide signals were analyzed by one-tailed paired Student’s *t* test. This analysis established that peptide signal correction by subtraction of the background mean value plus 3 standard deviations allowed unequivocal identification of reactive peptides. The polyclonal signal value was expressed as the base value × 10^3 for comparative displays, since polyclonal anti-human Ig conjugates produced higher chemiluminescent signals than monoclonal Ig conjugates.
Confirmation of peptide antigens with monospecies-specific anti-Chlamydia mouse serum pools. A set of previously reported (39) and new peptide antigens from each of the 11 Chlamydia spp. was first tested with monospecies-specific anti-Chlamydia hyperimmune mouse serum pools. Peptide antigens that reacted with the homologous anti-Chlamydia serum pool were further tested with the remaining 10 heterologous species-specific mouse serum pools to determine specificity and cross-reactivity.

Testing of mouse-reactive peptide antigens of 11 Chlamydia spp. with human serum pools. A set of 60 mouse-reactive peptide antigens of all 11 Chlamydia spp. was further tested with the pools of C. trachomatis-positive and -negative human sera by using polyclonal anti-human IgG. For confirmation of the presence of and determination of the levels of bound antibody isotypes, reactive peptide antigens were also tested by use of monoclonal conjugates against long-lived IgG1 and short-lived IgG3 antibodies. To determine anti-C. trachomatis MIF titer-dependent signal intensities, these peptide antigens were further tested with the 4 C. trachomatis MIF titer-dependent human serum subpools by using the human anti-IgG conjugate.

Testing of mouse reactive C. trachomatis peptide antigens with human serum pools. A large panel of 271 mouse serum-reactive C. trachomatis peptides, predicted as high-scoring B cell epitopes, was first tested with the human serum pool of C. trachomatis-positive sera by using the anti-human IgG polyclonal conjugate. Reactive peptides were further tested as described above for mouse reactive peptides.

Ranking of C. trachomatis peptide antigens. C. trachomatis peptides that reacted with the C. trachomatis-positive serum pool but not with the C. trachomatis-negative pool were ranked by an arbitrary reactivity score corresponding to the trachomatis-positive serum pool. These scores were determined by combining 7 weighted signal intensities of serum samples as follows—(i) 1/6 weight of the level of polyclonal anti-IgG conjugate reactivity with the pool of 125 serum samples from C. trachomatis-infected women; (ii) 1/12 weight for each polyclonal anti-IgG conjugate reactivity of the 4 MIF titer subpools; and (iii) 1/4 weight each for monoclonal anti-IgG1 and anti-IgG3 conjugate reactivities with the positive serum pool. This approach balanced all reactivities of serum total and subpools and weighted the data from the polyclonal and monoclonal conjugates equally.

Reactivity of C. trachomatis peptide antigens with 48 individual human sera. As a final confirmation of the identifications of the novel peptide antigen reagents, the top-ranked peptide antigens of C. trachomatis were tested for IgG1 and IgG3 reactivity with individual serum samples from 48 C. trachomatis-infected women. The serum samples were selected based on their distribution across the complete spectrum of C. trachomatis and C. pneumoniae MIF titers, proportional representation of racial origin of the study subjects, and available volume. All tests were performed by using single peptide antigens per well with a single serum sample and a single conjugate.

Testing of mouse reactive C. pneumoniae peptide antigens with human sera. To determine and contrast the reactivities of C. pneumoniae peptide antigens and C. trachomatis peptide reactivities, a panel of 153 C. pneumoniae peptide antigens was similarly tested with both C. trachomatis-positive human serum pools and C. trachomatis-negative pools as described above for the C. trachomatis peptide antigens. All 153 C. pneumoniae peptides were initially ranked by combining 7 weighted signal intensities of serum samples as described for C. trachomatis. A set of 18 top-ranked C. pneumoniae peptides was then tested with a panel of 48 individual sera, and the final rank was determined based on the frequency of positive reactivities with these 48 individual sera.

Statistical analyses. All statistical analyses were performed and graphical outputs were generated by the use of Microsoft Excel 2016 or the software package Statistica 7.1 (StatSoft, Tulsa, OK). To determine the relationships between continuous variables, Pearson’s correlation coefficient (R) values were calculated in linear and polynomial regression analyses from the square root of R². R values of 0.01 to 0.30 were considered to represent very weak correlations, 0.30 to 0.50 weak correlations, 0.50 to 0.70 moderately strong correlations, 0.70 to 0.90 strong correlations, and 0.90 to 1.00 very strong correlations. The probability of peptide cross-reactivity \( P_{cross} \) was calculated as follows: \( P_{cross} = e^{0.718 - 9.4153 \times 0.123223 \times \text{percent sequence identity} / (1 + 0.84153 \times 0.123223 \times \text{percent sequence identity})} \). At 45%, 60%, 75%, and 90% sequence identity (%SeqID), this translates to \( P_{cross} \) values of 0.02, 0.12, 0.46, and 0.84, respectively. This probability of cross-reactivity of peptide antigens with antibodies against a heterologous B cell epitope had been described earlier based on a large experimental data set (39). It is derived by logistic regression analysis of the observed cross-reactivity of chlamydial peptides with monospecies-specific serum samples against other chlamydial species and the percentage of sequence identity with the corresponding peptide antigens of the respective other chlamydial species (39).

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/mSphere.00246-18.

TABLE S1, XLSX file, 0.03 MB.

TABLE S2, XLSX file, 0.1 MB.

ACKNOWLEDGMENTS

This work was supported by grant 103 from the Auburn University Molecular Diagnostics Laboratory to K.S.R. and by National Institute of Allergy and Infectious Diseases grants U19 AI084024 and R01 AI119164.
pgp3 protein in animal and human chlamydial infections. Vet Microbiol 135:181–185. https://doi.org/10.1016/j.vetmic.2008.09.037.

37. Stothard DR, Williams JA, Van Der Pol B, Jones RB. 1998. Identification of a Chlamydia trachomatis serovar E urogenital isolate which lacks the cryptic plasmid. Infect Immun 66:6010–6013.

38. Schachter J. 2007. The Chlamydia trachomatis plasmid deletion mutant—which does it mean to us? Sex Transm Dis 34:257. https://doi.org/10.1097/OLQ.0b013e3180f52099.

39. Rahman KS, Chowdhury EU, Poudel A, Rouettger A, Sachse K, Kaltenboeck B. 2015. Defining species-specific immunodominant B cell epitopes for molecular serology of Chlamydia species. Clin Vaccine Immunol 22: 539–552. https://doi.org/10.1128/CVI.0102-15.

40. Wang J, Zhang Y, Lu C, Lei L, Yu P, Zhong G. 2010. A genome-wide profiling of the humoral immune response to Chlamydia trachomatis infection reveals vaccine candidate antigens expressed in humans. J Immunol 185:1670–1680. https://doi.org/10.4049/jimmunol.1001246.

41. Bunk S, Susnea I, Rupp J, Summersgill JT, Maass M, Stegmann W, Schrattenholz A, Wendel A, Przybyski M, Hermann C. 2008. Immunoprofiling and characterization of antibodies to novel Chlamydia pneumoniae antigens that are associated with persistent C. pneumoniae infections. J Immunol 180:5490–5498. https://doi.org/10.4049/jimmunol.1080849.

42. Li Z, Chen C, Chen D, Wu Y, Zhong Y, Zhong G. 2008. Characterization of fifty putative inclusion membrane proteins encoded in the Chlamydia trachomatis genome. Infect Immun 76:2746–2757. https://doi.org/10.1128/IAI.00010-08.

43. Yasui Y, Yanatori I, Kawai Y, Miura K, Suminami Y, Hirot A, Tamari M, Ouchi K, Kishi F. 2012. Genomic screening for Chlamydiaphila pneumoniae-specific antigens using serum samples from patients with primary infection. FEMS Microbiol Lett 329:168–176. https://doi.org/10.1111/j.1574-6968.2012.02520.x.

44. Maass M, Gieffers J, Krause E, Engel PM, Bartels C, Solbach W. 1998. Poor immunoprofiling and serological responses to novel Chlamydia pneumoniae antigens that are associated with persistent C. pneumoniae infections. J Immunol 180:5490–5498. https://doi.org/10.4049/jimmunol.1080849.

45. Rahman et al. 2008. Characterization of C. trachomatis pgp3 antibody prevalence in young women in England, 1992–2010. PLoS One 8:e72001. https://doi.org/10.1371/journal.pone.0072001.

46. Martin DL, Bid R, Sandi F, Goodnew EB, Massae PA, Laswasy A, Philippin H, Makupa W, Molina S, Holland MJ, Mabey DC, Drakeley C, Lammie PJ, Solomon AW. 2015. Serology for trachoma surveillance after cessation of mass drug administration. PLoS Negl Trop Dis 9:e0003555. https://doi.org/10.1371/journal.pntd.0003555.

47. Gwyn S, Mitchel A, Dean D, McKoch H, Handali S, Martin DL. 2016. Lateral flow-based antibody testing for Chlamydia trachomatis. J Immunol Methods 435:27–31. https://doi.org/10.1016/j.jim.2016.05.008.

48. Campbell LA, Kuo CC, Grayston JT. 1990. Structural and antigenic analysis of Chlamydia pneumoniae. Infect Immun 58:93–97.

49. Närävänä A, Puolakkainen M, Hao W, Kino S, Junii J. 1997. Detection of antibodies to Chlamydia trachomatis with peptide-based species-specific enzyme immunoassay. Infect Dis Obstet Gynecol 5:339–354. https://doi.org/10.1155/10167449970000616.

50. Bas S, Genevay S, Schenk MC, Vischer TL. 2002. Importance of species-specific antigens in the serodiagnosis of Chlamydia trachomatis reactive arthritis. Rheumatology 41:1017–1020. https://doi.org/10.1093/rheumatology/41.10.1017.

51. Morre SA, Munk C, Persson K, Krüger-Kjær S, van Dijk R, Meijer CJ, van den Brule AJ. 2002. Comparison of three commercially available peptide-based immunoglobulin G (IgG) and IgA assays to microminouflow fluorescence assay for detection of Chlamydia trachomatis antibodies. J Clin Microbiol 40:584–587. https://doi.org/10.1128/JCM.40.2.584-587.2002.

52. Klein M, Kötz A, Bernardo K, Krönke M. 2003. Detection of Chlamydia pneumoniae-species specific antibodies binding to the VD2 and VD3 regions of the major outer membrane protein. J Clin Microbiol 41:1957–1962. https://doi.org/10.1128/JCM.41.5.1957-1962.2003.

53. Rahman KS, Chowdhury EU, Sachse K, Kaltenboeck B. 2016. Inadequate reference datasets biased toward short non-epitopes confound B-cell epitope prediction. J Biol Chem 291:14585–14599. https://doi.org/10.1074/jbc.M116.729020.

54. Andresen H, Grötzinger C, Zarse K, Kreuter OJ, Ehrenreich-Förster E, Bier FF. 2006. Functional peptide microarrays for specific and sensitive antibody diagnostics. Proteomics 6:1376–1384. https://doi.org/10.1002/pmc.200500343.

55. Stansfield SH, Patel P, Debattista J, Armitage CW, Cunningham K, Timms P, Allen J, Mittal A, Huston WM. 2013. Proof of concept: a bioinformatic and serological screening method for identifying new peptide antigens for chlamydia trachomatis related sequelae in women. Results Immunol 3:33–39. https://doi.org/10.1016/j.jirin.2013.05.001.

56. Menon S, Stansfield SH, Logan B, Hocking JS, Timms P, Rombouts L, Allen JA, Huston WM. 2016. Development and evaluation of a multi-antigen peptide ELISA for the diagnosis of Chlamydia trachomatis-related infertility in women. J Med Microbiol 65:915–922. https://doi.org/10.1099/jmm.0.000311.

57. Sachse K, Rahman KS, Schnee C, Müller E, Peisker M, Schumacher T, Schubert E, Rouettger A, Kaltenboeck B, Ehrlich R. 2018. A novel synthetic peptide microarray assay detects Chlamydia species-specific antibodies in animal and human sera. Sci Rep 8:4701. https://doi.org/10.1038/s41598-018-23117-8.

58. Ehrlich R, Adelhelm K, Monecke S, Huelseweh B. 2009. Application of peptide-based species-specific antibodies to Chlamydia trachomatis diagnostics: importance of methodology in relation to timing of sampling. Clin Microbiol Infect 15:42–49. https://doi.org/10.1111/j.1469-0691.2008.02075.x.

59. Solomon AW. 2006. World Health Organization and International Trachoma Initiative. Trachoma control: a guide for programme managers. http://apps.who.int/iris/bitstream/handle/10665/34305/9241546095_eng.pdf;jsessionid=E324284246710189FF786C38EFA558E7F?sequence=1.

60. Russell AN, Zheng X, O’Connell CM, Wiesenfeld HC, Hillier SL, Taylor BD, Picard MD, Flechtner JB, Zhong W, Frazer LC, Danville T. 2016. Identification of Chlamydia trachomatis antigens recognized by T cells from highly exposed women who limit or resist genital tract infection. J Infect Dis 214:1884–1892. https://doi.org/10.1093/infdis/jiw485.