Identifying HLA \textit{DRB1-DQB1} alleles associated with \textit{Chlamydia trachomatis} infection and in silico prediction of potentially-related peptides

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HLA class II (HLA-II) genes’ polymorphism influences the immune response to \textit{Chlamydia trachomatis} (Ct), it is considered a sexually transmitted infection. However, associations between HLA-II alleles and Ct-infection have been little explored in humans; this study was thus aimed at determining HLA-\textit{DRB1-DQB1} alleles/haplotypes’ effect on Ct-infection outcome in a cohort of Colombian women. Cervical sample DNA was used as template for detecting Ct by PCR and typing HLA-\textit{DRB1-DQB1} alleles/haplotypes by Illumina MiSeq sequencing. Survival models were adjusted for identifying the alleles/haplotypes’ effect on Ct-outcome; bioinformatics tools were used for predicting secreted bacterial protein T- and B-cell epitopes. Sixteen HLA-\textit{DRB1} alleles having a significant effect on Ct-outcome were identified in the 262 women analysed. \textit{DRB1}\textsuperscript{*}08:02:01G and \textit{DRB1}\textsuperscript{*}12:01:01G were related to infection-promoting events. Only the \textit{DQB1}\textsuperscript{*}05:03:01G allele related to clearance/persistence events was found for HLA-\textit{DQB1}. HLA-\textit{DRB1} allele homozygous women were associated with events having a lower probability of clearance and/or early occurrence of persistence. Twenty-seven peptides predicted in silico were associated with protective immunity against Ct; outer membrane and polymorphic membrane protein-derived peptides had regions having dual potential for being T- or B-cell epitopes. This article describes HLA-\textit{DRB1-DQB1} alleles/haplotypes related to Ct-infection resolution and the peptides predicted in silico which might probably be involved in host immune response. The data provides base information for developing future studies leading to the development of effective prevention measures against Ct-infection.

\textit{Chlamydia trachomatis} (Ct) is the commonest bacteria-related, sexually-transmitted infection (STI) worldwide\textsuperscript{1}; the WHO’s Report on global sexually transmitted infection surveillance estimated that there are 127 million new cases annually\textsuperscript{2}. Most Ct infections have an asymptomatic clinical course, but some might lead to severe complications, such as pelvic inflammatory disease (PID) and recurrent abortions\textsuperscript{1}.

Ct mainly affects 18–20 year-old women and 20–24 year-old men, asymptomatic cases mainly occurring in women (close to 90%)\textsuperscript{3}; however, symptomatic infections usually occur weeks or months after exposure. Abundant secretion, dysuria and postcoital bleeding are amongst the commonest symptoms\textsuperscript{3}.

Previous studies have revealed that late detection of Ct asymptomatic infection (together with other agents such HPV) could lead to conditions such as squamous cell carcinoma and cervical cancer (CC)\textsuperscript{4}. It has been

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shown that a fourth of \textit{Ct} infections become spontaneously resolved and without treatment; factors, such as host immune response, thus significantly contribute towards eliminating genital tract infection\cite{5}.

The major histocompatibility complex (MHC) is a mechanism greatly determining infections’ clinical course; establishing its role in \textit{Ct} infection dynamics would thus explain their outcome\cite{5}. Some studies have suggested the relationship between class II HLA alleles, \textit{Ct} and tubal factor infertility (TFI)\cite{04:02} as DQA\textsuperscript{*}03:01, DQA\textsuperscript{*}05:01 and DQB\textsuperscript{*}04:02 have been described as being associated with \textit{Ct} and DRB1\textsuperscript{*}17:03 and DRB5\textsuperscript{*}01:01 with TFI\cite{04:02}.

An efficient immune response against intracellular pathogens like \textit{Ct} requires cell-mediated immunity, this being stimulated by bacterial peptides presented to T-cells by the MHC\cite{7}. Predicting these molecules and establishing their potential role in the natural history of \textit{Ct} infection could contribute towards understanding its dynamics\cite{8}.

Studies have searched for prophylactic interventions; most have evaluated the immune response in mice experimentally infected by \textit{Ct} or \textit{C. muridarum}\cite{9,10}. Murine model studies have demonstrated the importance of an HLA class II molecule-mediated immune response regarding bacterial infection resolution\cite{11}. However, few studies have comprehensively evaluated HLA alleles’ role regarding human \textit{Ct} infection outcome and the peptides probably related to immune response\cite{04:02,05:01}. This study was thus aimed at determining HLA-DRB1-DQB1 alleles/haplotypes’ effect on \textit{Ct} infection, persistence, clearance and redetection in a cohort of Colombian women. The results provided information for developing suitable prevention measures for managing and controlling \textit{Ct} infection.

**Results**

**HLA-DRB1-DQB1 allele effects on \textit{Ct} outcome.** The demographic characteristics of 262 women complying with the retrospective study’s inclusion criteria were analysed; mean age was 41.7 years-old (23.1 IQR) and median age at onset of sexual life 18.0 years-old (4.0 IQR) (Table 1). \textit{Ct} was detected retrospectively; study results gave the highest rate for persistence events (26.0 per 100 women/month), followed by clearance (16.1 per 100 women/month) and redetection events (15.6 per 100 women/month) (Supplementary Fig. S1). Survival data was estimated (using Kaplan Meier survival functions) for each event; Fig. 1 shows the probability of \textit{Ct} infection, clearance, persistence and redetection risk throughout the follow-up period.

| Variable                                      | Years |
|-----------------------------------------------|-------|
| Mean age (SD)                                 | 41.7 (23.1) |
| Median age at first intercourse (IQR)         | 18.0 (4.0) |
| Years of active sex life (SD)                 | 23.1 (10.6) |
| **n (%)**                                     |       |
| Ethnicity—not mestizo\textsuperscript{a}     | 7 (2.6) |
| More than three lifetime sexual partners       | 55 (21.8) |
| Marital status—married/cohabiting             | 231 (88.1) |
| Having had more than two pregnancies          | 37 (14.5) |
| A history of abortions                         | 93 (35.5) |
| A history of sexually-transmitted diseases    | 60 (22.9) |
| Contraceptive method—hormonal                 | 29 (11.9) |
| Educational level—illiterate/primary          | 122 (47.8) |
| Living outside Bogotá                         | 180 (68.7) |
| Average monthly income—less than minimum wage\textsuperscript{b} | 323 (89.6) |
| HPV positive                                  | 218 (83.2) |
| Cytological findings—abnormal                 | 23 (8.9) |

Table 1. Base line demographic characteristics and risk factors for the 262 women included in the study. *This category included Indigenous and Afro Colombian ethnicities. *The Colombian minimum average monthly income would be roughly US$ 250.
found that HLA-DRB1 homozygous women had a LP effect on clearance and an EO effect related to persistence events, suggesting that this characteristic could represent a genetic disadvantage for its carriers as it makes them more susceptible to Ct infection. However, more experimental evidence is required to confirm such hypothesis.

**HLA-DRB1-DQB1 haplotype effect on Ct outcome.** Forty-seven of the 142 DRB1-DQB1 haplotypes had statistically significant values when analysing the effect of each haplotype on Ct outcome (Table 3). Twenty-seven associations were found regarding infection events; 16 related to the event (GP EO) (e.g. DRB1*01:02:01G-DQB1*03:03:02G) whilst 11 associations did not (LO) (e.g. DRB1*12:01:01G-DQB1*03:03:02:01G). DRB1*09:01:02G-DQB1*03:01:01G and DRB1*12:01:01G-DQB1*03:02:01G had an EO effect on clearance whilst DRB1*04:05:01-DQB1*03:01:01G was associated with LP (Table 3 and Supplementary Table S7).

Eighteen associations were found regarding persistence; sixteen of them related to the event (GP/EO), DRB1*04:05:01-DQB1*03:01:01G and DRB1*11:01:01G-DQB1*03:02:01G having the greatest effect. By contrast, DRB1*10:01:01G-DQB1*05:01:01G and DRB1*14:02:01G-DQB1*05:01:01G reduced persistence (LP/LO). Eight haplotypes were related to redetection events (Table 3 and Supplementary Table S8).

Nine haplotypes were related to more than one event when analysed; 6 were related to some infection events, such as EO for DRB1*04:05:01-DQB1*02:01:01G on infection and redetection, LO for DRB1*12:01:01G-DQB1*03:02:01G on infection and GP on clearance;

whilst others had an opposite effect on events, LO of DRB1*01:01:01G-DQB1*03:01:01G infection and EO on persistence (Table 3).

**In silico predicted peptides associated with Ct events.** Peptides derived from proteins predicted as being secreted (Supplementary Table S9) were predicted in silico since it has been shown that they could be associated with protective immunity or susceptibility to Ct infection. Fifteen out of 24 proteins had peptides having T-cell epitopes binding strongly to HLA-II-DRB1 but not to -DQB1 molecules (Supplementary Tables S10 to S24). Some peptides might have been related to increased susceptibility against Ct as they were associated with GP and EO effects related to persistence or redetection events. Twenty-seven peptides were related to protection (of which 7 were OMP-derived and 17 from PMP) since they were associated with effects related to Ct elimination (LO of infection, GP of clearance and LP of persistence) (Supplementary Tables S10 to S24). Interestingly, 11
of them had potential B-cell epitope regions, thus highlighting their possible role as cell and/or humoral immune response mediators (Table 4).

### Discussion
*Ct* is the commonest sexually-transmitted bacterial pathogen worldwide; it can provoke serious consequences regarding reproductive sexual health once it becomes a chronic infection. Despite significant advances having been made regarding its control, clear and effective tools for reducing its impact on public health are still not

| Locus | Allele | Event | Coeff. | 95%CI | Pc | Effect |
|-------|--------|-------|--------|-------|----|--------|
| 01:02:01G | Infection | 8.36 | 1.71–4.08 | 0.000 | Greater probability |
| 01:03:01 | Redetection | 0.01<sup>c</sup> | 0.00–0.05 | 0.000 | Earlier occurrence |
| 03:02:01 | Infection | −0.98<sup>b</sup> | −1.29–(−0.67) | 0.000 | Earlier occurrence |
| 03:02:02 | Infection | −0.55<sup>b</sup> | −0.86–(−0.25) | 0.000 | Earlier occurrence |
| 04:10:01G | Infection | 2.91<sup>b</sup> | 2.49–3.36 | 0.000 | Later occurrence |
| 08:01:01G | Infection | −0.87<sup>b</sup> | −1.20–(−0.54) | 0.000 | Earlier occurrence |
| 08:02:01G | Persistence | 12.71 | 4.35–37.09 | 0.000 | Greater probability |
| Redetection | −0.32<sup>a</sup> | −0.51–(−0.13) | 0.030 | Earlier occurrence |
| 09:01:02G | Clearance | 5.53 | 2.91–10.49 | 0.000 | Greater probability |
| 10:01:01G | Persistence | 0.33 | 0.19–0.57 | 0.000 | Lower probability |
| 11:01:01G | Persistence | 22.20 | 5.70–86.49 | 0.000 | Greater probability |
| 11:01:02 | Infection | 3.00<sup>b</sup> | 2.48–3.51 | 0.000 | Later occurrence |
| 12:01:01G | Infection | 3.23<sup>b</sup> | 2.73–3.73 | 0.000 | Later occurrence |
| Persistence | 5.53 | 2.91–10.49 | 0.000 | Greater probability |
| 13:05:01 | Persistence | 3.67 | 1.74–7.75 | 0.032 | Greater probability |
| 14:01:01G | Persistence | 3.92 | 1.86–8.25 | 0.000 | Greater probability |
| 14:02:01G | Infection | 2.40<sup>b</sup> | 1.87–3.10 | 0.000 | Later occurrence |
| Persistence | 27.79 | 5.30–145.59 | 0.000 | Greater probability |
| 16:01:01 | Redetection | −0.17<sup>a</sup> | −0.27–(−0.08) | 0.000 | Earlier occurrence |
| 05:03:01G | Clearance | 4.97 | 1.79–13.74 | 0.024 | Greater probability |
| Persistence | 3.92 | 1.86–8.25 | 0.000 | Greater probability |

Table 2. HLA *DRB1-DQB1* alleles associated with *Ct* infection, clearance, persistence and redetection. Alleles affecting an event are shown as green associations [i.e. greater probability (GP) or earlier occurrence (EO)] and those hindering them [lower probability (LP) or later occurrence (LO)] as red associations. *Pc* corrected *p* value, 95%CI 95% confidence interval, Coeff regression coefficient, *DRB1* DR beta 1, *DQB1* DQ beta 1. Values in bold indicate statistical significance based on 95%CI, *p* < 0.05. The Cox proportional hazards model did not fulfil the assumption of proportionality; a logistic parametric model was constructed. cThe Cox proportional hazards model did not fulfil the assumption of proportionality; a lognormal parametric model was constructed.

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of them had potential B-cell epitope regions, thus highlighting their possible role as cell and/or humoral immune response mediators (Table 4).

**Discussion**
*Ct* is the commonest sexually-transmitted bacterial pathogen worldwide; it can provoke serious consequences regarding reproductive sexual health once it becomes a chronic infection. Despite significant advances having been made regarding its control, clear and effective tools for reducing its impact on public health are still not
available. Therefore, comprehensively evaluating a Ct infection dynamics-related immune response represents
an alternative approach for developing effective control tools15,16.

This study was focused on comprehensively investigating (the first time) the effect of HLA-DRB1-DQB1 alleles on Ct infection outcome, given that it has been shown that HLA molecules could be related to Ct-induced diseases, such as trachoma17, PID6,18 and infertility19,20, or be associated with infection prevalence and bacterial reinfection8,21,22. It was found that HLA-DRB1 alleles were associated with cervical-related Ct infection outcome (Tables 2 and 3). Some alleles were less common for this locus and occurred at lower frequency in the target population. MHC-pathogen coevolution models indicate that less commonly occurring alleles provide greater protection against pathogens than more commonly occurring ones to which pathogens may have become adapted23,24.

DQB1*05:03:01G was only associated with Ct clearance and persistence events (Table 2); however, previous studies have reported HLA-DQB1 (DQB1*06 and DQB1*04:02) alleles’ association with Ct infection and reinfection and increased bacterial persistence marker cHSP608,21,22. Such discrepancy could be explained by the genetic background of the particular population being studied (African compared to South-American in this study) thereby contributing to modulating an immune response to bacterial infection6,25. However, these alleles only had similar associations to those reported in previous studies when configured as haplotypes, i.e. when they have been combined with a DRB1 allele (Table 3).

| Haplotype DRB1 DQB1 | Event       | Coeff. | 95%CI     | Pcd | Effect          |
|----------------------|-------------|--------|-----------|-----|----------------|
| 01:01:01G 03:01:01G  | Infection   | 2.73a  | 2.21–3.25 | 0.000 | Later occurrence |
| 01:01:01G 03:02:01G  | Persistence | 0.48a  | 0.31–0.66 | 0.000 | Earlier occurrence |
| 01:01:01G 03:02:01G  | Infection   | 4.76   | 2.03–11.14 | 0.000 | Greater probability |
| 01:01:01G 03:03:02G  | Infection   | 3.10a  | 2.61–3.59 | 0.000 | Later occurrence |
| 01:02:01G 03:03:02G  | Infection   | 14.82  | 4.54–48.31 | 0.000 | Greater probability |
| 01:03:01 02:01:01G  | Persistence | 9.52   | 2.77–32.71 | 0.000 | Greater probability |
| 01:03:01 02:01:01G  | Redetection | 0.01c  | 0.00–0.05 | 0.000 | Earlier occurrence |
| 01:03:01 03:01:01G  | Infection   | −1.27b | −1.64(−0.90) | 0.000 | Earlier occurrence |
| 01:03:01 03:02:01G  | Infection   | 2.97a  | 2.44–3.49 | 0.000 | Later occurrence |
| 01:03:01 05:01:01G  | Persistence | 9.52   | 2.77–32.71 | 0.000 | Greater probability |
| 01:03:01 05:01:01G  | Redetection | 0.01c  | 0.00–0.05 | 0.000 | Earlier occurrence |
| 03:01:01G 05:01:01G  | Infection   | 14.08  | 5.36–36.96 | 0.000 | Greater probability |
| 03:02:01 04:02:01G  | Infection   | −0.98b | −1.29(−0.67) | 0.000 | Earlier occurrence |
| 03:02:01 04:02:01G  | Infection   | −0.55b | −0.86(−0.25) | 0.000 | Earlier occurrence |
| 04:02:01 03:01:01G  | Infection   | 0.83a  | 0.41–1.26 | 0.000 | Earlier occurrence |
| 04:03:01G 02:01:01G  | Persistence | 5.25   | 2.10–13.15 | 0.000 | Greater probability |
| 04:03:01G 02:01:01G  | Infection   | 5.32   | 2.58–10.95 | 0.000 | Greater probability |
| 04:03:01G 03:02:01G  | Persistence | 6.89   | 2.82–16.83 | 0.000 | Greater probability |
| 04:05:01 02:01:01G  | Infection   | −0.80b | −1.30(−0.48) | 0.000 | Earlier occurrence |
| 04:05:01 02:01:01G  | Redetection | −0.47a | −0.61(−0.34) | 0.000 | Earlier occurrence |
| 04:05:01 03:01:01G  | Clearance   | 0.23   | 0.11–0.47 | 0.000 | Lower probability |
| 04:05:01 03:01:01G  | Persistence | 51.72  | 10.70–249.94 | 0.000 | Greater probability |
| 04:05:04 03:02:01G  | Infection   | 3.08a  | 2.57–3.59 | 0.000 | Later occurrence |
| 04:08:01 02:01:01G  | Infection   | 3.10a  | 2.60–3.61 | 0.000 | Later occurrence |
| 04:08:01 03:01:01G  | Infection   | 3.10a  | 2.60–3.61 | 0.000 | Later occurrence |
| 07:01:01G 02:01:01G  | Redetection | 0.01c  | 0.00–0.05 | 0.000 | Earlier occurrence |
Table 3. HLA DRB1-DQB1 haplotypes associated with Ct infection, clearance, persistence and redetection. Analysis was adjusted for age, age at onset of sexual life, lifetime amount of sexual partners, planning method, abortions, history of other STI and HPV infections. Alleles affecting an event are shown as green associations [i.e. greater probability (GP) or earlier occurrence (EO)] and those hindering them [lower probability (LP) or later occurrence (LO)] as red associations. Values in bold indicate statistical significance based on 95%CI, p < 0.05. Pc corrected p value, 95%CI 95% confidence interval, Coeff regression coefficient, DRB1 DR beta 1, DQB1 DQ beta 1. a The Cox proportional hazards model did not fulfil the assumption of proportionality, the logistic parametric model was constructed. b The Cox proportional hazards model did not fulfil the assumption of proportionality, the lognormal parametric model was constructed. c The Cox proportional hazards model did not fulfil the assumption of proportionality, the Gompertz parametric model was constructed. d The Bonferroni method was used for correcting all models’ p values.
LP of clearance and EO of persistence were found for homozygous HLA-DRB1 (Supplementary Tables S3 and S4). It has been reported that homozygosity is related to susceptibility to infection whilst heterozygosity is associated with a higher probability of eliminating it, possibly due to a greater immune response, given the broader amount of HLA-II restricted epitopes that can be presented to T-cells. It is worth noting that findings regarding alleles causing effects on events promoting or reducing infection (Table 2) are useful when designing infection control strategies; for example, considering peptides presented by DRB1*12:01 could represent a good strategy since it is associated with events related to infection resolution whilst peptides presented by DRB1*08:02 should be avoided as it is related to events associated with infection, such as persistence and redetection.

It has been suggested that Ag presentation during adaptative response could be an important mechanism for controlling Ct infection; identifying T-cell antigens able to stimulate protection-inducing immunity is thus the key for developing anti-Ct vaccines. Analysing Ct molecules’ T-cell epitopes (Supplementary Tables S10 to S24) whose role in protection-inducing immunity was experimentally evaluated revealed that 3-oxoacyl-[acyl-carrier protein] reductase had a peptide related to a GP of Ct clearance (Table 4). This peptide had been found in an immunoproteomics study demonstrating that inoculating dendritic cells previously pulsed with a peptide mixture (including the peptide discussed here) triggered a response partially protecting mice from intranasal and genital tract Chlamydia infection. It has been demonstrated that CPAF-derived peptides could be related to a protection-inducing effect in a HLA-DR4 (HLA-DRB1*04:01) transgenic mouse model, however, no allele/peptide association was found in this study (Supplementary Table S12), possibly due to the allele’s low frequency in the studied population (less than 0.2%), suggesting an allele-specific effect.

The OMPs and PMPs had various regions containing T-cell epitopes, the most important ones being related to events associated with Ct infection elimination (Table 4). Interestingly, there was discrepancy amongst several events which could have been explained by binding core mutations for some PMP-derived peptides (PMP-B and PMP-F); these enabled discriminating between invasive (L2 and LGV) and non-invasive variants (A, B, C, D, E, F, H, J, Ja and La) (Supplementary Table S16 and S18).

It has been reported recently that the Ct OMP (CTH522) protein being evaluated in phase 1 trials was able to trigger a more consistent cell-mediated immune response profile after its immunisation using CAF01 liposomes compared to the placebo group, thus highlighting its potential usefulness as a vaccine candidate. Such result, added to this study’s findings, supports the idea that T-cell epitopes derived from the antigens analysed here (mainly surface-derived molecules, particularly totally conserved ones) could be regions of interest for the future design of novel interventions aimed at controlling Ct infection.

| Protein name | Event | Locus | Allele | Effect | Peptide |
|--------------|-------|-------|--------|--------|---------|
| 3-OXO-RED   | Clearance | *09:01 | Major probability | RGSPGQTNYAAKAGIFGS |
| OMP-C       | Infection | *04:10 | Later occurrence | MAEGLSTEVSLADTKADN |
|             | Clearance | *09:01 | Major probability | SYVCKPVASYSNSPGDLV |
| PMP-A       | Infection | *04:10 | Later occurrence | ANKKGGAAYAQYVNEONQD |
|             |                  | *12:01 | Major probability | SGDFAGSRLFLNQQTTE |
| PMP-B       | Infection | *04:10 | Later occurrence | TCENSHRLQFLKNSSDKGG |
|             | Persistence | *10:01 | Lower probability | RNQNVCKVRVLSSNAGQV |
| PMP-D       | Infection | *04:10 | Major probability | GAIAAQEIVSNSNQAGSF |
| PMP-F       | *04:10 | Later occurrence | RGKAEQPILEITEIINQDGQL |
| PMP-G       | *12:01 | VAGRGGGIAAYQDQGQYVS |
|             | *12:01 | EIGVQLPVITPSKVLNNEL |
|             | *12:01 | GTIYKSTDLLNNEKF SY |
| LSU         | Clearance | *09:01 | Major probability | IAVKGNFVPPAAPVYVDRP |
| OMP-B       | Infection | *04:10 | Later occurrence | RHASCPIDYIANSQANPEV |
| OMP-H       | Infection | *10:01 | Major probability | AAEELRKKFEDLSAEYNTAQG |
| VPP         | clearance | *09:01 | Major probability | EPLKDQQIIIITSTPVKA |

Table 4. Protein regions having the potential to be T- or B-cell epitopes. T-(green) or B-cell (red) epitope prediction is shown; regions having dual prediction are shown in purple. Peptides which could be related to GP of clearance when they are presented by the same allele. Peptides related to LP of persistence when presented by DRB1*10:01 allele.
In silico analysis suggested that some protection-related predicted peptides would specifically stimulate T-cells whilst others would stimulate both T- and B-cells (Table 4). Vaccination with various OMP serovars (D, E and F) has elicited an antibody (Ab) response neutralising bacteria in vitro17. Furthermore, PMPs can trigger an immune response against genital34 and ocular35 Chlamydia infection and a serological response in humans46. Mice vaccinated with DC/PMP-derived peptides (G, E and F) or with just immunogens in formulation have developed immunity against genital tract and pulmonary Chlamydia infection, significantly reducing bacteria in UFI assays2,35. Interestingly, phase 1 clinical trial vaccination using the Ct-OMP version showed accelerated seroconversion, increased IgG titres and enhanced mucosal profile, thus making CTH522 a promising candidate for further clinical development42.

As CD4 T-cells are essential for resolving primary genital infection48 and CD8 T-cells are important for eliminating Ct-infected cells by effector mechanisms49, then peptides stimulating both types of effector cells (B- and T), as predicted here, could be considered most suitable for controlling Ct infection and therefore as promising candidates for future studies (Table 4).

Antigenic peptides recognised by both CD4 and CD8 T-cells could be promising diagnostic and therapeutic tool candidates since one of the main limitations for developing an effective vaccine lies in identifying Ct epitopes capable of being recognised by both cell types30; an Ab-mediated immune response would reduce bacterial load, thereby facilitating further elimination of infection via a cell-mediated immune response3,40.

Considering this and given the in silico analysis performed here, it can thus be suggested that a universal anti-Ct vaccine should contain peptides having the following characteristics: they should induce immunity and be protein-derived, have a high degree of conservation, be associated with protection-related events (such as GP of clearance and LP of infection, redetection and persistence) and be able to stimulate T- and B-cell responses.

HLA-DRB1-DQB1 alleles/haplotypes having an effect on Ct resolution have thus been reported here, along with in silico predicted epitopes derived from predicted protection-related peptides. However, functional read-outs for demonstrating the effect of presentation regarding the predicted antigens (i.e. wet-laboratory assays) were not performed here, thus failing to obtain a complete panorama of anti-Ct immune responses constitutes a limitation of this study.

Addressing new prophylactic and therapeutic targets must become a high priority as the tools used to date for Ct control have not had a significant impact on reducing bacterial infection load. Future analysis should be aimed at validating predicted epitopes’ immunogenic and immunological in vitro and in vivo properties and their safe and efficacy regarding humans. Such data will provide relevant knowledge for understanding the usefulness of peptides as a vaccine component and the influence of host factors on the clinical course of Ct infection.

Materials and methods

Study design and participants. A cohort was studied between 2007 and 2010; that previous study was aimed at determining the natural history of HPV infection in women from the Colombian cities of Bogotá, Girardot and Chaparral; all the women were attending hospital clinics as outpatients. The study’s objective was explained to them and they voluntarily accepted participating in the study by signing an informed consent form, as described previously13. Retrospective analysis inclusion criteria consisted of having available cervical samples for typing HLA-DRB1 and DQB1 and Ct detection, women having attended at least four follow-up sessions (one base line and three visits) and 6-monthly periods between visits (±3 months).

The women filled in a survey form during each visit for compiling data regarding sociodemographic information and risk factors. Such information included data regarding whether they had received/used any type of treatment between visits; none of the women reported using antibiotics during follow-up. The women did not receive antibiotic treatment for Ct infections detected during the study, given the retrospective nature of Ct detection. The Universidad del Rosario’s School of Medicine and Health Sciences Research Ethics Committee approved the study (CEI-ABN026-000135). All procedures were performed in accordance with Helsinki Declaration guidelines.

Ct detection and HLA-DRB1-DQB1 typing. Previously obtained genomic DNA (gDNA)13,41 was used as template for detecting Ct by conventional PCR, amplifying a cryptic plasmid ORF2 segment from KL5/KL6 and KL1/KL2 primers43, An Illumina MiSeq (San Diego, CA, USA; Histogenetics, Ossining, NY, USA) sequencer was used for typing HLA alleles from DRB1-DQB1 loci exons 2 and 3; the IPD-IMGT/HLA database (https://www.ebi.ac.uk/ipd/imgt/hla) published in January 2018 (3.31.0) was used for assigning alleles13.

Statistical analysis. Qualitative variables were expressed as percentages. The Chi2 or Fisher’s exact tests were used for evaluating association/concordance amongst categorical variables. Continuous variables were expressed as means [with standard deviations (SD) for measure of dispersion] or medians [interquartile ranges (IQR)].

Ct infection was defined for this study’s purposes as PCR detection of bacterial DNA at any point during follow-up (2 years). Clearance was understood as the elimination of infection via a previous positive Ct result. A percentage of infections persisted before becoming eliminated; this event was evaluated in the study and was defined as Ct being detected during two or more consecutive follow-ups. Redetection was defined as bacterial detection after not having detected bacterial DNA during a previous follow-up.

Incidence rates for events were reported along with 95% confidence intervals (95%CI). The Kaplan–Meier estimator was used for estimating the probability of subjects continuing event-free. Cox proportional hazards models were constructed for evaluating outcome probability; such models’ coefficients were expressed as hazard rate (HR) and used for identifying alleles/haplotypes related to the events being evaluated. Schoenfeld residuals were plotted to test the proportional hazard assumption; the covariables considered for the plot were those
having \( p < 0.200 \) in univariate analysis (Supplementary Table S2). Variance inflation factor (VIF) and tolerance values were used for evaluating multicollinearity between covariables\(^3\).

Different parametric survival models were constructed when proportional hazards assumptions were not met. Akaike (AIC) and Bayesian (BIC) information criteria were used for selecting the models having the best fit. The Bonferroni method was used for correcting \( p \)-values for each model\(^3\). All two-tailed hypothesis tests (except those involved in constructing the models) were run with 0.05 significance. STATA14 software was used for analysis.

**T- and B-cell epitope prediction.** Ct variant-derived protein amino acid (aa) sequences (Supplementary Table S25) were downloaded from the PATRIC 3.5.11 database (https://www.patricbrc.org) and analysed first using classical pathway secretion predictor (SignalP 5.0)\(^42\) and those not assigned by this predictor were analysed by the non-classical one, SecretomeP 2.0\(^43\).

The Technical University of Denmark’s Systems Biology Department’s Center for Biological Sequence Analysis’ NetMHCIIpan 3.2 server was used for assessing peptides having high predicted HLA-DRB1-DQB1 allele binding activity, i.e. proteins predicted as secreted\(^44\). Peptides having <2.0% rank were considered to have strong binding. The BepiPred 2.0 tool (0.6 epitope threshold) was used for calculating B-cell epitopes derived from proteins whose peptides were associated with protection\(^45\).

**Data availability**
The datasets produced and/or analysed during this study are available from the corresponding author on reasonable request.

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Author contributions

L.P., M.C. and D.A.M.P.: designed the study, performed the experiments, analysed the data, wrote the first draft and revised the final draft of the manuscript. R.S. and L.D.R.O.: designed the study, contributed to data analysis and critically reviewed the first and final drafts of the manuscript. I.M.B.M.: contributed to the experiments and revised the final draft of the manuscript. R.S. and L.D.R.O.: designed the study, contributed to data analysis and critically reviewed the first and final drafts of the manuscript. R.S. and L.D.R.O.: designed the study, contributed to data analysis and critically reviewed the first and final drafts of the manuscript. M.E.P. and M.A.P.: critically reviewed the first and final drafts of the manuscript. All authors approved the final draft.

Competing interests

The authors declare no competing interests.

Additional information

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