A Probiotic Adjuvant *Lactobacillus rhamnosus* Enhances Specific Immune Responses after Ocular Mucosal Immunization with Chlamydial Polymorphic Membrane Protein C

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Abstract

Recent advances in the development of chlamydia vaccines, using live-attenuated or ultraviolet light-inactivated chlamydia, are paving the way for new possibilities to oppose the societal challenges posed by chlamydia-related diseases, such as blinding trachoma. An effective subunit vaccine would mitigate the risks associated with the use of a whole-cell vaccine. Our rationale for the design of an efficient subunit vaccine against *Chlamydia trachomatis* (Ct) is based on the membrane proteins involved in the initial Ct-host cell contact and on the route of immunization that mimics the natural infection process (i.e., via the ocular mucosa). The first aim of our study was to characterize the specific conjunctival and vaginal immune responses following eye drop immunization in BALB/c mice, using the N-terminal portion of the Ct serovar E polymorphic membrane protein C (N-PmpC) as the subunit vaccine antigen. Second, we aimed to examine the adjuvant properties of the probiotic *Lactobacillus rhamnosus* (LB) when formulated with N-PmpC. N-PmpC applied alone stimulated the production of N-PmpC- and Ct serovar B-specific antibodies in serum, tears and vaginal washes, whereas the combination with LB significantly enhanced these responses. The N-PmpC/LB combination initiated a T cell response characterized by an elevated percentage of CD25+ T cells and CD8+ effector T cells, enhanced CD4+ T-helper 1 skewing, and increased regulatory T cell responses. Together, these results show that eye drop vaccination with combined use of N-PmpC and a live probiotic LB stimulates specific cellular and humoral immune responses, not only locally in the conjunctiva but also in the vaginal mucosa, which could be a promising approach in Ct vaccine development.
Introduction

Eye drop vaccination via the ocular mucosa constitutes an attractive immunization approach, particularly for the prevention/treatment of ocular infections and their blinding sequelae [1, 2]. Furthermore, ocular topical immunization is safer than nasal immunization because there is no risk of antigen redirection towards the nervous system [1] and is needle-free, which is more comfortable and safer than parenteral vaccination [3].

*Chlamydia trachomatis* (Ct) is a human pathogen causing chronic conjunctivitis and is also the most common cause of sexually transmitted disease. Ct infections can be asymptomatic and, if left untreated, can result in blinding trachoma (the leading cause of preventable blindness worldwide) and pelvic inflammatory disease, which can lead to infertility and ectopic pregnancy [4–6]. Regardless of the infection site, there is a consensus that a vaccine is needed [7–11]. Even a partially effective vaccine would contribute in reducing the global disease burden caused by Ct [12].

Polymorphic membrane proteins (Pmps) are essential in early *Chlamydia*-host cell interactions [13–15] and are recognized as potential vaccine antigens to inhibit both contact and infection [15]. Furthermore, a relatively high portion of the Ct genome encodes for Pmp superfamily proteins [16], suggesting their importance in the *Chlamydia* life cycle [17]. Epitope mapping of the PmpC region encompassing the amino acid residues 605–840 demonstrated a broad B cell recognition range. In addition, the full-length protein was shown to react with the serum from Ct-infected minipigs [18]. We have previously shown that N-PmpC can trigger heterologous immunity as well as the positive influence of particulate Gram-negative bacterial adjuvants on the development of antigen-specific immune responses after topical ocular immunization [19].

Available data suggest a hyporesponsiveness of conjunctiva-associated immune cells to lipopolysaccharide stimulation [20] and an important role for TLR2 agonists in the abrogation of the immunosuppressive mechanisms naturally occurring within the conjunctiva [21]. These findings led to the hypothesis that Gram-positive bacteria, e.g., probiotic *Lactobacillus rhamnosus* (LB), which are also corpuscular in nature, may be employed as an efficient adjuvant when immunizing via the conjunctiva. Furthermore, *Lactobacillus* spp. (*Lactobacillus acidophilus*) was previously used in ocular surface applications without signs of toxicity or safety risks [22].

The aim of our study was to characterize the local and systemic immune responses produced by the chlamydia-specific subunit antigen N-PmpC by analysing serum and mucosal washes from *Chlamydia*-relevant surfaces (i.e., conjunctiva and vagina). In addition, we sought to assess the modulatory effect of the probiotic bacterial adjuvant LB on the post-immunization immune response.

Materials and Methods

Ethics statement

All experiments were approved by the "Ethics Committee for the Welfare of Experimental Animals" and by the committee section at the Institute of Virology, Vaccines and Sera – TORLAK. All experiments conformed to the Serbian laws and European regulations on animal welfare (Approval No. 011-00-00510/2011-05). Every effort was made to minimize animal suffering. Mice that were immunized were anesthetized by intraperitoneal (i.p.) administration of a mixture of xylazine (Sigma-Aldrich, Kansas, KS, USA) and ketamine (Richter Pharma AG, Wels, Austria). The method for mice euthanasia was cervical dislocation. We did not observe any unexpected deaths of animals during this study.
Animals

Ten-week-old BALB/c female mice (six mice per group) were housed at the Animal Facility of the Institute of Virology, Vaccines and Sera (TORLAK) and kept at a temperature of 21°C under a 12:12 h light: dark cycle with ad libitum access to water and food.

Antigens and adjuvants

A recombinant N-PmpC protein fragment (1–565 amino acids) from Ct serovar E produced in E. coli [15, 23] was used as the antigen. LB (characterized by 16S rRNA sequencing, NCIMB Ltd, Aberdeen, UK, ref.no. NCSQ 18723; TORLAK) was used as the adjuvant. During preliminary validation experiments, LB was administered via the conjunctiva (5 μl/eye) at a concentration of 1 x 10^8 CFU/ml, resulting in an LB immunization dose of 1 x 10^6 CFU/dose. Additional preliminary results with different LB doses (1x10^9 CFU/ml, 1x10^8 CFU/ml, 1x10^7 CFU/ml, 1x10^6 CFU/ml) on days 0, 14 and 28 applied topically revealed no visible signs of inflammation or infection at the ocular surface, where signs of ocular irritation were monitored in all mice on a daily basis (during the course of immunization). Two “blinded” ophthalmologists, using magnifying loupes, assessed conjunctival hyperemia, edema and corneal clarity.

Immunization schedules

Ten-week-old BALB/c female mice (n = 6 mice per group) were immunized with N-PmpC on days 0, 14 and 28, and the resulting immune responses were evaluated two weeks after the last immunization. The mice were immunized via the conjunctiva (conj//) with N-PmpC alone in PBS (conj//N-PmpC) or N-PmpC combined with LB at 1 x 10^8 CFU/ml in PBS (conj//N-PmpC/LB). The concentration of N-PmpC in all vaccines was 1.5 mg/ml. Each mouse was immunized with 15 μg N-PmpC, with or without a 10^6 CFU dose of LB, in a total volume of 10 μl (5 μl/eye, both eyes were treated). A group of age-matched non-immunized mice was used as the normal control group (nc). Our experimental procedure did not cause any visual impairment (including blindness) in animals, which was examined by an ophthalmologist during the whole immunization period. Furthermore, we did not observe any changes in behavioural pattern between treated animals and their respective controls.

Sample collection

Blood serum samples were collected from the mouse tail vein (6 individual sera from 6 animals per group) two weeks after completion of the indicated immunization protocol. Wash samples were obtained two weeks after completion of the indicated immunization protocol by lavage with 15 μl of PBS for each eye for tear-wash samples, and with 150 μl for vaginal-wash samples.

Detection of PmpC-specific and CtB-specific immunoglobulins

Quantification of N-PmpC- and Ct serovar B (CtB)-specific antibodies in sera, tear and vaginal washes was performed as previously described [19], slightly modified. Briefly, ELISA plates (MaxiSorp; Nunc, Roskilde, Denmark) were coated (50 μl/well) with N-PmpC (10 μg/ml N-PmpC in PBS) or CtB (1 x 10^6 IFU/ml in PBS) by overnight adsorption at 4°C. Appropriately diluted non-pooled sera (1:100), tear or vaginal washes (1:10) were used as samples. Specific antibody levels in wash samples were expressed as a relative amount, calculated as the concentration of specific antibody in a particular sample divided by the lowest concentration of the same antibody in the corresponding nc sample.
Lymphocyte phenotyping by FACS analysis

Submandibular lymph nodes (SMLN) from mice immunized via the ocular conjunctivae or control mice were aseptically isolated as previously described [19].

SMLN cells (1 x 10^6 cells/sample) were immunostained using fluorochrome-conjugated antibodies specific for murine CD3 (FITC-conjugated, eBioscience, San Diego, CA), CD4 (PE-conjugated, Biolegend, San Diego, CA), CD8 (PerCP-conjugated, Biolegend), CD19 (PECy5-conjugated, eBioscience), CD25 (PE-conjugated, Biolegend or PECy5-conjugated, eBioscience) and Foxp3 (Alexa488-conjugated, Biolegend). Before staining, the cells were washed in a cold 2% BSA/0.1% NaN_3/PBS solution (2 x centrifugation at 300 g, 5 min, 4°C). Fluorochrome-conjugated antibodies were added to the resuspended cell pellets and incubated in the dark for 30 min at 4°C. Discrete aliquots of each analysed cell suspension were incubated with the corresponding isotype control antibodies and used as the unstained reference for setting the FACS analysis staining thresholds. Unbound antibodies were removed by washing in cold 2% BSA/0.1% NaN_3/PBS solution (3 x centrifugation at 300 g, 5 min, 4°C).

For the intracellular Foxp3 staining, the cells were first stained with anti-mouse CD4-PE and anti-mouse CD25-PECy5 and then fixed and permeabilized using the BD Cytofix/Cytoperm Buffer and BD Cytoperm Permeabilization Buffer Plus (BD Biosciences, San Jose, CA, USA), respectively, according to the manufacturer’s instructions. Washes between all steps were performed using the Perm/Wash Buffer (BD Biosciences; 3 x centrifugation at 300 g, 5 min, 4°C). An anti-mouse Foxp3-Alexa488 monoclonal antibody was added to the resuspended cell pellets and incubated in the dark for 30 min at 4°C. Unbound antibodies were removed by centrifugation (washing).

Stained cells were analysed using the BD FACScan™ flow cytometer (BD Biosciences). BD CellQuest™ software was used for analysis.

Proliferation assay

Evaluation of proliferative response of SMLN cells upon stimulation with N-PmpC (10 μg/ml) and CtB (1 x 10^6 IFU/ml) was performed as previously described [19]. Briefly, SMLN cells were plated into 96-well plates (100 μl/well, 2 x 10^6 cell/ml in 10% FCS/50 μM β-mercaptoethanol/RPMI 1640) and incubated for 48 h (5% CO_2, 37°C) without additional stimulation or in the presence of the stimulators. Cell Counting Kit-8 reagent (10 μl/well, Sigma Aldrich) was added upon 48 h long incubation, and the cells were incubated for additional 4h. Reactions were stopped by the addition of 1% (w/v) sodium dodecyl sulfate (10 μl/well), and absorbance values were measured at 450/650 nm (A_450/650) using a spectrophotometer (Ascent 6–384 [Suomi], MTX Lab Systems Inc., Vienna, VA, USA).

The number of viable cells per well was calculated using a standard curve A_450/650 = f(number of cells). Discrete pool of non-stimulated cells was used as standard after counting in the presence of trypan blue (Countess Automated Cell Counter, Invitrogen). Standard suspension was plated in serial dilutions prior to centrifugation and further treated identically as the experimental wells.

A proliferation index (PI) for each specifically stimulated cell suspension was calculated per individual animal. The PI index was defined as the ratio of number of viable cells per well present in stimulated (S) to number of viable cells per well present in corresponding non-stimulated (So) cultures, such that PI = Ss/So.

SMLN cell cytokine profiling

Production of IFN-γ, IL-4, IL-17A and IL-10 was analysed by measuring their concentrations in the supernatants of non-stimulated, N-PmpC- (10 μg/ml) and CtB-stimulated (1 x 10^6 IFU/ml)
SMLN cultured cells (2 x 10^6 cell/ml in 10% FCS/50 μM β-mercaptoethanol/RPMI 1640; 37°C, 5% CO₂, 48 h) using sandwich ELISA with commercially available monoclonal antibodies (eBioscience) [19].

Statistical analyses

The statistical significance of the observed differences was evaluated using Kruskal-Wallis test followed by Dunn’s multiple comparisons test to compare between groups. All statistical analyses were performed with the GraphPad 6.0 software. A probability (P) value of 0.05 was set as the significance threshold.

Results

Ocular mucosal immunization increased anti-PmpC IgA levels in sera

Ocular immunization with N-PmpC, either alone or formulated with LB, promoted a systemic rise in the level of N-PmpC-specific IgA vs the nc group (Fig 1A; P < 0.005 for conj//N-PmpC, P < 0.05 for conj//N-PmpC/LB). However, no significant difference was found between conj// groups in the levels of all antibodies analysed. The mean serum anti-N-PmpC IgA concentrations recorded in the nc, conj//N-PmpC and conj//N-PmpC/LB groups were 4.25±0.30 μg/ml, 11.14±1.71 μg/ml and 14.91±4.62 μg/ml, respectively.

Analysis of anti-N-PmpC IgG showed no significant difference between the conj// groups (Fig 1B). However, the mean levels of anti-N-PmpC IgG were higher in the serum of the conj//N-PmpC and conj//N-PmpC/LB groups (165.68±47.42 μg/ml and 171.93±45.56 μg/ml, respectively) vs the nc group (80.67±4.76 μg/ml). In addition, subclass level analyses showed no significant difference in the contribution of N-PmpC-specific IgG1 and IgG2a antibodies in both conj// groups vs the nc group (Fig 1C).

N-PmpC-specific SIgA levels in mucosal washes were elevated significantly following immunization

An evaluation of N-PmpC-specific antibodies in the tears and vaginal washes generally showed a rise in the local concentrations of both N-PmpC-specific IgG and IgA after N-PmpC immunization via the ocular mucosa (Fig 2). The concentration of N-PmpC-specific
SIgA was significantly higher in the wash samples from both conj// groups vs the nc group (Fig 2A and 2B). In addition, an elevated level of specific SIgA was found in the vaginal wash samples from the conj//N-PmpC/LB group compared with the conj//N-PmpC group (Fig 2B; *P* < 0.05).

The N-PmpC-specific IgG concentration was also elevated in the wash samples collected from conj// mice compared with nc mice (Fig 2C and 2D). A significant rise in anti-N-PmpC IgG level was observed in the tears of conj//N-PmpC mice (Fig 2C; *P* <0.05 vs nc) and in the vaginal washes collected from both conj// groups vs the nc group (Fig 2D; *P* <0.05 for both groups). Comparison of anti-N-PmpC IgG levels in the washes collected at the same mucosal surface from the conj//N-PmpC and conj//N-PmpC/LB mice revealed no significant differences.
Anti-CtB antibody levels in sera, tears and vaginal washes were increased significantly following immunization

IgG and IgA analysis of sera and washes collected from mucosal surfaces revealed that conj// immunization with N-PmpC promoted the production of both CtB-specific immunoglobulins (Fig 3).

The mean concentration of anti-CtB antibodies in the sera of the nc, conj//N-PmpC and conj//N-PmpC/LB mice were 0.24±0.01 μg/ml, 0.34±0.07 μg/ml and 0.26±0.01 μg/ml, respectively, for IgA (Fig 3A), and 0.80±0.05 μg/ml, 1.48±0.44 μg/ml and 1.01±0.14 μg/ml, respectively, for IgG (Fig 3D).

The most prominent increase in anti-CtB antibody levels after either formulation of conjunctival N-PmpC immunization was recorded in the vaginal washes. Indeed, the conj//N-PmpC group produced more vaginal anti-CtB IgG (Fig 3F; $P < 0.005$ vs nc), whereas the conj//N-PmpC/LB group produced more of both vaginal anti-CtB IgA (Fig 3C; $P < 0.0009$ vs nc) and anti-CtB IgG (Fig 3D; $P < 0.05$ vs nc). Compared with the group immunized with

![Fig 3. Anti-CtB antibody levels in serum and on mucosal surfaces.](image-url)
N-PmpC alone, LB co-administration exerted a positive effect on vaginal CtB-specific SIgA levels (Fig 3C; \(P < 0.005\)).

LB co-administration exerted a significant positive effect on CtB-specific SIgA level in tears as well (Fig 3B; \(P < 0.05\) vs nc).

**N-PmpC immunization promoted effector T cell differentiation in SMLNs**

Analysis of the SMLN T cell pool (i.e., CD3+ cells within the lymphocyte gate, S1 Fig) revealed that conjunctival immunization with N-PmpC in the presence of LB promoted the expansion of CD8+ T cells over the CD4+ (Fig 4A, percentage of CD8+ T cells in conj/N-PmpC/LB, \(P < 0.05\) vs nc). In addition, the percentage of CD25+ T cells in conj/N-PmpC LB was markedly increased compared with the nc group (Fig 4B, \(P < 0.005\)). The increased percentage of CD25+ T lymphocytes was mainly due to the higher abundance of CD3+CD4+CD25+ lymphocytes (Fig 4C, conj/N-PmpC/LB, \(P < 0.005\) vs nc).

Simultaneous analysis of CD4, CD25 and Foxp3 expression on SMLN lymphocytes (S2 Fig) revealed a significantly lower percentage of CD4+CD25+Foxp3+ cells (Treg) in the nc group than in the conj/N-PmpC/LB mice (Fig 4D, \(P < 0.005\)).

**Chlamydia antigens stimulated in vitro proliferation of SMLN cells**

With the aim of evaluating local immune responses in mice immunized with N-PmpC via the conjunctiva, SMLN cells were stimulated *in vitro* with the N-PmpC and CtB antigens (Fig 5). Irrespective of the stimulator (N-PmpC or CtB), the proliferation of SMLN cells isolated from the conj/N-PmpC/LB group was significantly higher than that of SMLN cells from both corresponding nc and conj/N-PmpC groups (N-PmpC stimulation: \(P < 0.005\); CtB stimulation: \(P < 0.05\)).

**LB adjuvant affected the cytokine pattern of SMLN cells and their response to Ct antigen stimulation**

The basal *in vitro* production of effector cytokines (IFN-\(\gamma\) as Th1 marker, IL-4 as Th2 marker, IL-17A as Th17 marker) and the regulatory cytokine IL-10, recorded in non-stimulated SMLN
cultures, differed among conjunctively immunized groups and was altered compared with the 
culture (Fig 6).

Compared with the nc non-stimulated SMLN cells, the conj//N-PmpC non-stimulated 
SMLN cells produced slightly more IFN-γ and IL-10, whereas the secretion of IL-17A was 
significantly reduced ($P < 0.05$). The basal IFN-γ and IL-10 production levels were highest in the 
SMLN cultures from the conj//N-PmpC/LB group (IFN-γ: $P < 0.005$ vs nc; IL-10: $P < 0.005$ vs 
nc) and were accompanied by a significant reduction in IL-4 secretion ($P < 0.05$ vs nc). SMLN 
cells from the conj//N-PmpC/LB group without any stimulation secreted IL-17A in amounts 
comparable with those recorded in non-stimulated nc SMLN cells.

In nc SMLN cultures, N-PmpC stimulation enhanced the secretion of IFN-γ ($P < 0.05$ vs 
non-stimulated nc), IL-10 and IL-17A but did not affect the secretion of IL-4. Compared with 
the corresponding non-stimulated cultures, N-PmpC stimulation enhanced the production of 
IFN-γ ($P < 0.005$), IL-17A and IL-10, and inhibited IL-4 secretion in conj//N-PmpC SMLN 
cells. In contrast, N-PmpC stimulation decreased IL-17A, IFN-γ ($P < 0.005$ vs non-stimulated) 
and IL-10 ($P < 0.05$ vs non-stimulated) production while enhanced IL-4 secretion ($P < 0.05$ vs 
non-stimulated) in conj//N-PmpC/LB SMLN cells.

CtB stimulation had less impact on the overall production of all tested cytokines. Compared 
with the corresponding non-stimulated cultures, CtB stimulation did not significantly affect 
the production of the tested cytokines. However, in the CtB-stimulated conj//N-PmpC/LB 
cultures compared with the CtB-stimulated nc SMLN cultures, the productions of IFNγ 
($P < 0.005$) and IL-10 ($P < 0.05$) were significantly higher while IL-4 production was decreased 
($P < 0.05$).
Discussion

The administration of the N-PmpC chlamydia-specific subunit antigen via the ocular mucosa elicited antigen specific humoral and cell-mediated immune responses.

The importance of the humoral immune response in the protection against Ct has been debated extensively [24–26]. Compared with immunization with N-PmpC alone, LB co-administration had no significant impact on specific serum antibodies. Interestingly, the levels of chlamydial specific SIgAs were significantly higher in the vaginal washes of mice immunized with N-PmpC/LB vs mice immunized with N-PmpC alone. This finding shows that ocular mucosal immunization could increase antigen-specific antibodies in distant mucosal compartments, such as the vaginal mucosa, when antigen is presented to the immune system within an

Fig 6. Effect of N-PmpC and CtB stimulation on the in vitro cytokine pattern in SMLN cells isolated after ocular mucosal immunization with N-PmpC and N-PmpC/LB. Bar graphs representing the levels of IFNγ, IL-4, IL-10 and IL-17A in the supernatants of non-stimulated, N-PmpC- or CtB-stimulated SMLN cells isolated from BALB/c mice immunized via the conjunctiva (bars). The corresponding measurements in SMLN cells isolated from age-matched controls (nc) are presented on the graphs as a solid line (mean value) and dotted lines (upper and lower standard error values). SMLN cells were cultured at 37°C under a 5% CO2 atmosphere for 48 h in 10% FCS/50 μM β-mercaptoethanol/RPMI 1640 medium supplemented or not with the indicated stimulator (10 μg/ml for N-PmpC or 1x10^6 CFU/ml for CtB). The results are presented as the mean concentrations ± SE (n = 6). The statistical significance of the observed differences was evaluated using Kruskal-Wallis test followed by Dunn’s multiple comparisons test to compare between groups (immunized vs nc stimulated in a same way P < 0.05*, P<0.005**; vs non-stimulated samples of the same group P < 0.05#, P<0.005##). doi:10.1371/journal.pone.0157875.g006

Probiotic Live Adjuvant and Its Influence on the Immune Response

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adequate adjuvant delivery system. This might be important because subunit formulations with different adjuvants delivered into the vagina were shown to be poorly immunogenic due to immunological properties of the female reproductive tract during the menstrual cycle [27, 28]. The mechanism behind this warrants further attention and will be explored in our further experiments.

The fact that anti-N-PmpC antibody levels changed in parallel with corresponding (same isotype and location) anti-CtB antibodies levels implies that the observed anti-CtB reactivity is likely due to cross-reactivity between anti-N-PmpC and anti-CtB antibodies. This suggests that the antibodies secreted upon N-recombinant PmpC stimulation might 1) recognize naturally occurring antigens and 2) cross-react with the PmpC fragment of Ct serovars that were not the source the recombinant antigen.

It is known that the cellular immune response is important for protection against Ct infection [29–31]. IFN-γ, known as the key molecule in the protection against Ct invasion, can be produced by both Th1 (CD4+) and Tc1 (CD8+) cells [24]. However, CD4+ T cell depletion was shown to abrogate the protection against Ct-infection more profoundly than the depletion of CD8+ T cells [32]. In our study, LB promoted the expansion of CD8+ T cells and Th effector (CD3+CD4+CD25+) cells compared with the conj//N-PmpC group. The elevated percentage of CD8+ cells within the SMLN T cells of conj//N-PmpC/LB mice is in line with the results of MHC I/proteasome cleavage prediction testing, which suggested that N-PmpC-derived peptides could be presented in the context of MHC I [32]. The abovementioned findings, together with the strong proliferative response observed in conj//N-PmpC/LB SMLN cells upon CtB stimulation, suggest that LB or similar probiotic adjuvants may be promising tools for the development of efficient anti-Ct vaccines. Moreover, cytokine production analysis in ex vivo SMLN cultures suggests that N-PmpC administration via the conjunctiva promoted a slight Th1 immune response, which was prominently enhanced by LB co-administration. Further, the comparison of SMLN cell responses to in vitro N-PmpC and CtB stimulations confirms that the applied immunization protocol influences not only the intensity but also the quality of the response to subsequent antigen stimulation. Results we got in vitro imply that in N-PmpC/LB-immunized mice subsequent stimulation with N-PmpC alone would not be able to enhance Th1 immune response. Even more, it is likely that Th1 specific immune response would be inhibited. This result suggests that for an eventual booster immunization N-PmpC also has to be formulated with some adjuvant capable to stimulate innate immunity, in order to tackle natural Ct infection. However, during naturally occurring infection PmpC is not alone i.e. it is in the context of chlamydia elementary bodies. Accordingly, we consider the results gotten upon CtB stimulation more relevant for the prediction of susceptibility to infection.

We also demonstrated that the increased percentage of effector T cells was accompanied by an expansion of the Treg cell population, and that, in all cases, enhanced IFN-γ secretion was paralleled by enhanced IL-10 production. However, the activation of regulatory mechanisms following vaccination may be a double-edged sword [25]. This response may represent a beneficial self-limiting mechanism preventing a strong inflammatory immune response, thereby diminishing pathological sequelae and prevent potentially harmful autoreactive immune responses through molecular mimicry [33]. However, an increase in the number of Foxp3+ Tregs has been shown to prevent the development of an efficient immune response upon immunization [24]. Besides, one of the inherent characteristics of Foxp3+ T cells is “plasticity” i.e. they can undergone reprogramming in the sense of acquisition of an effector phenotype [34]. It was shown by others that stimulation of rabbit Tregs through TLR2 decreased their immunosuppressive potential and promoted the concomitant expansion of T effector cells [21]. Similar findings were reported in mice [35, 36]. Although Foxp3 expression is considered a marker of the Treg population and implies on cells capable of exerting suppressive impact,
the real suppressive potential of Tregs in this particular situation (i.e. their functional status) cannot be discerned by Foxp3 expression only. For all these reasons, the actual immunosuppressive potential of Tregs during our immunization protocol should be evaluated in more detail.

Although the precise mechanisms by which probiotics act as adjuvants remain unknown, an immunoenhancing strain of *Lactobacillus* (*L. rhamnosus* HN001) identified by Gill et al and isolated originally from Cheddar cheese, compared with placebo, was shown to significantly induce phagocytic cell activity and anti-cholera specific antibody production in gut mucosa when delivered simultaneously with a cholera toxin vaccine [37]. The proposed mechanisms of action for the adjuvant activity of probiotics include an influence on innate immune cells, such as intestinal macrophages or dendritic cells, which may in turn enhance antigen presentation and promote the preferential differentiation of mucosal lymphocytes towards the production of protective antibodies particularly at sites where a multitude of antigens are encountered constantly, such as in the conjunctiva- and vagina-associated lymphoid tissues [38].

In conclusion, we provided evidence that ocular mucosal immunization with combined use of a specific antigen and a probiotic stimulates specific immune responses not only locally but also in the vaginal mucosa.

**Supporting Information**

**S1 Fig. Phenotyping of SMLN lymphocytes by FACS analysis.** Lymphocytes were gated according to their position within the forward scatter (FSC) vs side scatter (SSC) plots and analysed for the percentage of T (CD3+CD19-) and B (CD3-CD19+) cells. T cells (gated CD3+ lymphocytes) were further analysed for the expression of CD4, CD8 and CD25. Cells were analysed using a BD FACScan "flow cytometer (BD Biosciences) and BD CellQuest" software. Representative dot plots and histograms are presented. (TIF)

**S2 Fig. Phenotyping of SMLN Treg lymphocytes by FACS analysis.** The co-expression of CD4, CD25 and Foxp3 on lymphocytes from SMLN of BALB/c mice immunized via the conjunctiva and age-matched controls (nc) was analysed. Lymphocytes were gated according to their position within the FSC vs SSC plots. Then, CD4+ lymphocytes were gated and further analysed for the expression of CD25 and Foxp3. Cells were analysed using a BD FACScan "flow cytometer (BD Biosciences) and BD CellQuest" software. Representative dot plots and histograms are presented. (TIF)

**Author Contributions**

Conceived and designed the experiments: AIK MS TBA. Performed the experiments: AIK MS EM EB ES IL RD NS. Analyzed the data: AIK MS. Contributed reagents/materials/analysis tools: JHH TBA. Wrote the paper: MS AIK TBA.

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