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Published in:
Immunology and Cell Biology

Link to article, DOI:
[10.1038/icb.2015.79](https://doi.org/10.1038/icb.2015.79)

Publication date:
2016

Document Version
Publisher's PDF, also known as Version of record

[Link back to DTU Orbit](#)

Citation (APA):
Bøje, S., Olsen, A. W., Erneholm, K., Agerholm, J. S., Jungersen, G., Andersen, P., & Follmann, F. (2016). A multi-subunit Chlamydia vaccine inducing neutralizing antibodies and strong IFN-(+) CMI responses protects against a genital infection in minipigs. *Immunology and Cell Biology*, 94(2), 185-195.
<https://doi.org/10.1038/icb.2015.79>

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ORIGINAL ARTICLE

A multi-subunit *Chlamydia* vaccine inducing neutralizing antibodies and strong IFN- γ ⁺ CMI responses protects against a genital infection in minipigs

Sarah Bøje^{1,2}, Anja Weinreich Olsen², Karin Erneholm^{1,2}, Jørgen Steen Agerholm¹, Gregers Jungersen³, Peter Andersen² and Frank Follmann²

Chlamydia is the most widespread sexually transmitted bacterial disease and a prophylactic vaccine is highly needed. Ideally, this vaccine is required to induce a combined response of Th1 cell-mediated immune (CMI) response in concert with neutralizing antibodies. Using a novel Göttingen minipig animal model, we evaluated the immunogenicity and efficacy of a multi-subunit vaccine formulated in the strong Th1-inducing adjuvant CAF01. We evaluated a mixture of two fusion proteins (Hirep1 and CTH93) designed to promote either neutralizing antibodies or cell-mediated immunity, respectively. Hirep1 is a novel immunogen based on the variant domain (VD) 4 region from major outer membrane protein (MOMP) serovar (Sv) D, SvE and SvF, and CTH93 is a fusion molecule of three antigens (CT043, CT414 and MOMP). Pigs were immunized twice intramuscularly with either Hirep1+CTH93/CAF01, UV-inactivated *Chlamydia trachomatis* SvD bacteria (UV-SvD/CAF01) or CAF01. The Hirep1+CTH93/CAF01 vaccine induced a strong CMI response against the vaccine antigens and high titers of antibodies, particularly against the VD4 region of MOMP. Sera from Hirep1+CTH93/CAF01 immunized pigs neutralized *C. trachomatis* SvD and SvF infectivity *in vitro*. Both Hirep1+CTH93/CAF01 and UV-SvD/CAF01 vaccination protected pigs against a vaginal *C. trachomatis* SvD infection. In conclusion, the Hirep1+CTH93/CAF01 vaccine proved highly immunogenic and equally protective as UV-SvD/CAF01 showing promise for the development of a subunit vaccine against *Chlamydia*.

Immunology and Cell Biology (2016) 94, 185–195; doi:10.1038/icb.2015.79

Chlamydia is the most common bacterial sexually transmitted disease, and worldwide, more than 100 million people are infected each year with the urogenital serovars (Svs) of *Chlamydia trachomatis*.¹ The infection is often without symptoms and therefore left undiagnosed and untreated. This increases the risk of developing severe complications, such as pelvic inflammatory disease and infertility.² At present, the main public intervention to reduce the incidence of genital chlamydia is routine screening programs, which until now has been largely unsuccessful,¹ and development of a vaccine against chlamydia is therefore an international priority.³

In humans, repeated chlamydial infections are common, although natural immunity can develop.⁴ The exact immune correlates of protection are unknown, however, it is generally accepted that both humoral and cell-mediated immune (CMI) responses are important. The presence of antibodies in cervical secretions of women correlates with lower bacterial shedding,⁵ and interferon (IFN)- γ -positive responses in peripheral blood mononuclear cells (PBMCs) stimulated

with cHsp60 correlates with a decreased risk of acquiring a subsequent chlamydial infection.⁶ Studies in mice consistently identify a key protective role for CD4⁺ Th1 cells and the cytokine IFN- γ .^{7–9} The protective role of antibodies is more multifaceted and ranges from direct bacterial neutralization¹⁰ to accelerating CMI through Fc-mediated uptake in antigen-presenting cells.¹¹

Vaccines based on intact *Chlamydia* bacteria demonstrate that protection can be achieved both in non-human primates and in clinical trials.^{12,13} However, protection seems to be Sv-specific¹³ and a whole-cell vaccine may potentially generate unwanted responses and lead to pathology,¹⁴ emphasizing the need for a broadly protective subunit vaccine. Vaccine research efforts have led to the identification of a large number of antigens with significant protective efficacy.^{15–18} Efforts to identify a target for neutralizing antibodies have mainly focused on major outer membrane protein (MOMP), but the complex structure of the protein has complicated its use as a vaccine antigen.^{19,20}

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Received 22 April 2015; revised 14 July 2015; accepted 29 July 2015; accepted article preview online 13 August 2015; advance online publication, 22 September 2015

Table 1 Design of study

No of pigs	Immunization groups	Timepoint			
		Week 0	Week 3	Week 7 Day 0 post infection	Day 12 pi
5	CAF01	First immunization	Second immunization	Infection	Killing
5	10 µg Hirep1+CTH93/CAF01				
5	100 µg Hirep1 +CTH93/CAF01				
4	125 µg UV-SvD/CAF01				

Abbreviation: pi, post infection.

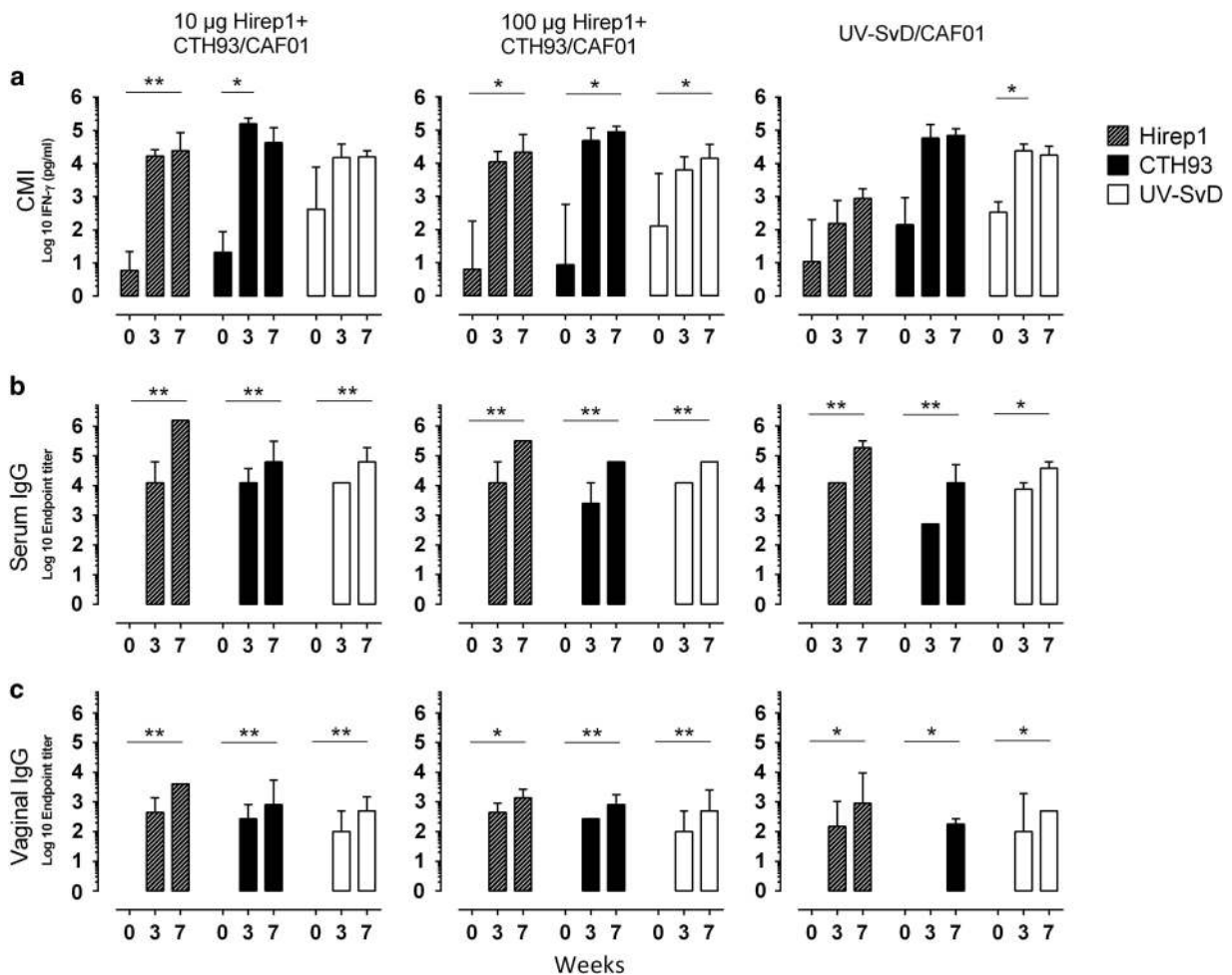


Figure 1 Characterization of vaccine-induced immune responses. Pigs were intramuscularly immunized at week 0 and 3 with 10 µg ($n=5$) or 100 µg ($n=5$) Hirep1+CTH93/CAF01 or UV-SvD/CAF01 ($n=4$). Blood samples and vaginal swabs were taken at weeks 0, 3 and 7 to evaluate CMI and humoral responses. (a) PBMCs were restimulated *in vitro* with Hirep1, CTH93 and UV-SvD. For each pig, the mean IFN- γ of triplicate culture wells was calculated and the bars represent the median \pm interquartile range within a group. (b) Serum IgG and (c) vaginal IgG titers specific for Hirep1, CTH93 and UV-SvD were analyzed in duplicates and the mean were calculated. Bars represent medians \pm interquartile range within a group. Statistical difference was calculated using the Friedman test followed by Dunn's multiple comparison test, * $P < 0.05$, ** $P < 0.01$.

In addition to identifying the right antigens, vaccine-induced immunity to *Chlamydia* highlights the challenge of identifying a clinically relevant delivery system that would induce a strong Th1 response, antibodies and long lived memory.²¹ The CAF01 adjuvant has demonstrated potent induction of CMI and humoral immunity with several chlamydial antigens in mouse models.^{17,22} Importantly, the adjuvant has also been tested in phase I trials^{23,24} with an excellent

safety profile and is a promising adjuvant for a recombinant vaccine against *Chlamydia*.

So far, most *Chlamydia* vaccine research has been carried out in mice.^{15–17} To move promising vaccine candidates closer to a clinical trial, there is a need for testing in animal models with closer resemblance to humans than rodents. Pigs are immunologically and physiologically in many ways comparable with humans,^{25–27}

facilitating the evaluation of vaccine immunogenicity in this animal species. Importantly, a previous study has demonstrated that pigs are susceptible to genital *C. trachomatis* infection.²⁸

We recently constructed a novel recombinant version of MOMP, based on the variant domain (VD)4 regions of SvD, SvE and SvF (Hirep1) that induces high titered neutralizing antibodies, covering SvS causing up to 90% of all infections and showing for the first time that vaccine-induced antibodies can control *Chlamydia* infections.²⁹ In the present study, we combine Hirep1 with a recombinant fusion molecule CTH93, consisting of previously identified antigens; CT043,^{17,30,31} MOMP (CT681)_{amino acid (aa)34-371}^{15-17,31} and CT414_{aa605-840}. CTH93 represents a broad epitope repertoire covering both T- and B-cell epitopes, but lacks the ability to induce neutralizing antibodies. In the current study, we evaluate the immunogenicity and efficacy of Hirep1+CTH93 formulated in CAF01 in Göttingen minipigs.

RESULTS

The Hirep1+CTH93/CAF01 vaccine is highly immunogenic

Göttingen minipigs were immunized with a mixture of Hirep1 (comprising the VD4 domain of MOMP SvD, SvE and SvF) and CTH93 (consisting of MOMP SvD_{aa34-371}, CT043 and CT414_{aa605-840})

(Supplementary Figure 1) formulated in CAF01, with the purpose of inducing neutralizing antibodies and strong T-cell responses. Throughout the study, the Hirep1+CTH93/CAF01-induced responses were compared with immune responses in UV-SvD/CAF01-immunized pigs. Nineteen sexually mature minipigs were randomly allocated into four immunization groups receiving 10 µg Hirep1+CTH93/CAF01, 100 µg Hirep1+CTH93/CAF01, 125 µg UV-SvD/CAF01 or CAF01 alone (Table 1). The pigs were immunized twice in the neck muscles spaced with an interval of 3 weeks.

At week 0, 3 and 7, PBMCs were isolated from blood samples and restimulated with the recombinant, vaccine antigens (Hirep1 and CTH93) and with UV-SvD. After immunizations, Hirep1+CTH93/CAF01-immunized groups showed significant higher levels of IFN-γ against Hirep1 and CTH93 compared with week 0 (Figure 1a). Furthermore, the Hirep1+CTH93/CAF01 vaccine-induced CMI responses were able to recognize UV-SvD, comparable with the levels in UV-SvD/CAF01-immunized pigs.

To examine humoral immune responses, we collected sera and vaginal swabs at week 0, 3 and 7. Both groups of pigs immunized with Hirep1+CTH93/CAF01 had significant levels of specific IgG in serum (Figure 1b). The Hirep1+CTH93/CAF01-generated antibodies were able to recognize the surface of UV-SvD, comparable with the levels

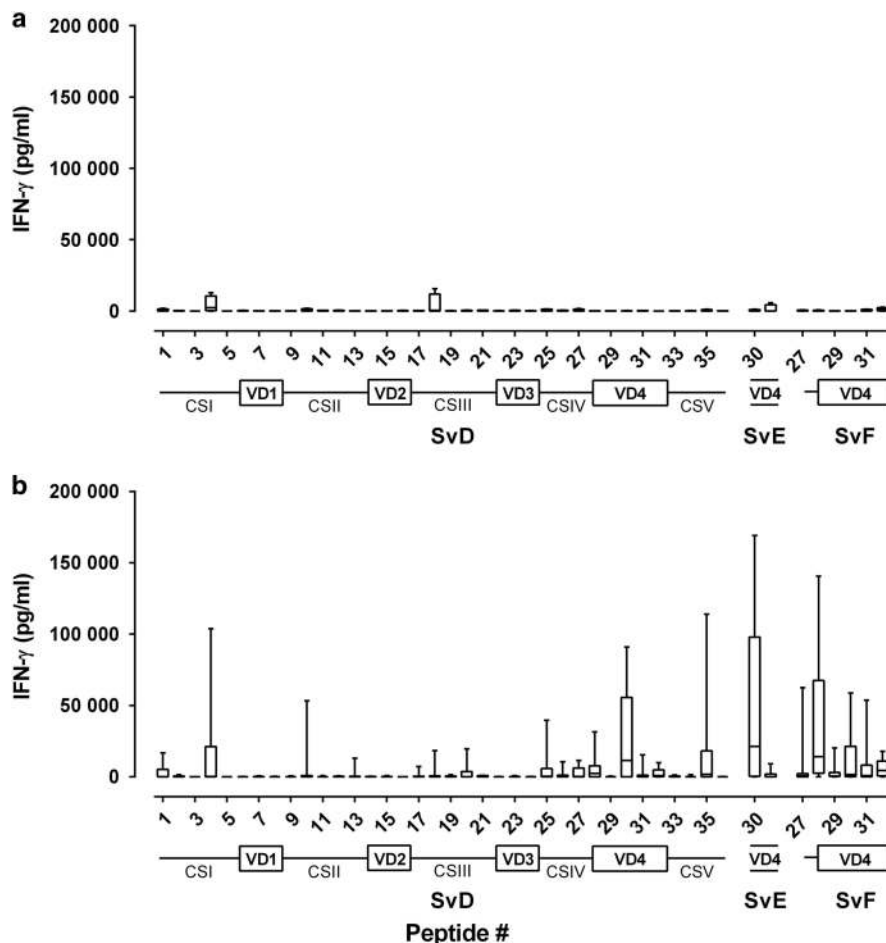


Figure 2 Mapping of T-cell epitopes after two immunizations in peptides (20-mer with 10 aa overlap) covering MOMP. PBMCs from (a) UV-SvD/CAF01 ($n=4$)- and (b) Hirep1+CTH93/CAF01 ($n=10$)-immunized pigs were restimulated *in vitro* with individual peptides (see Supplementary Table 1 for sequences), spanning the MOMP SvD protein including SvE- and SvF-specific peptides in VD4. The graph is based on the mean IFN-γ value of duplicate culture wells for each pig. The box indicates the 25th and the 75th percentile within the group, the horizontal line within a box represents the median, and whiskers indicate min to max values. The sequence of constant sequences (CS) and VDs of MOMP is depicted below the graphs.

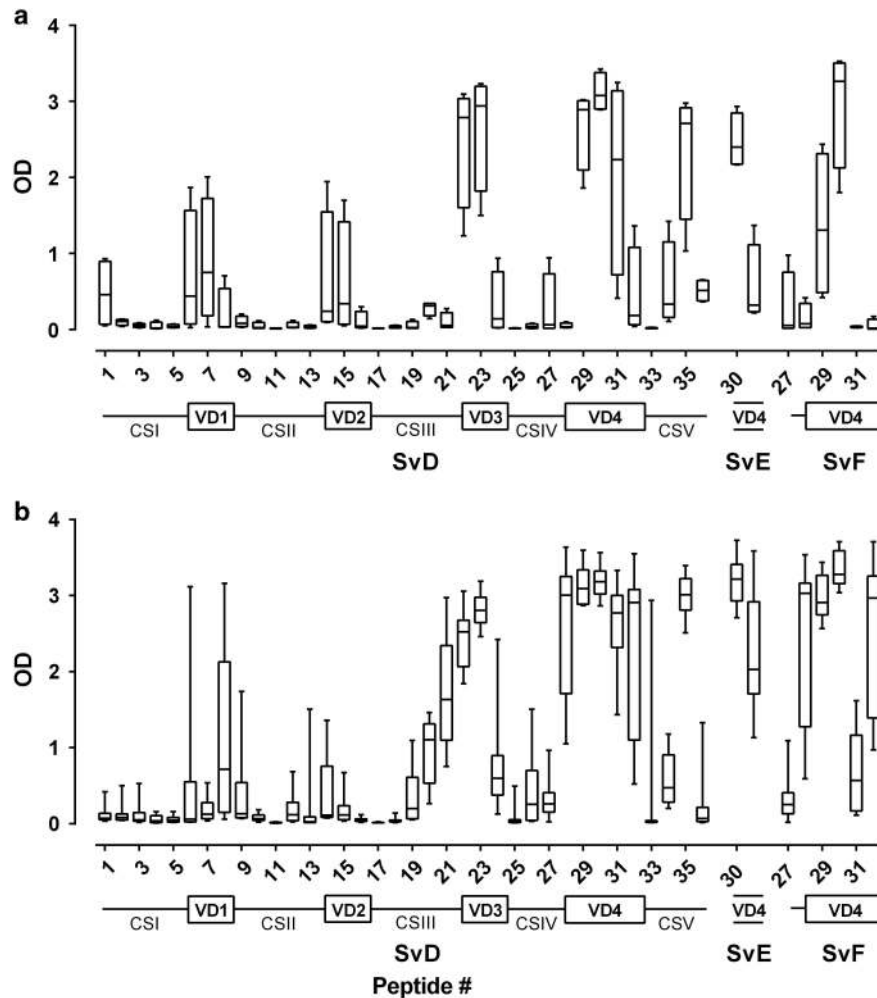


Figure 3 Mapping of B-cell epitopes after two immunizations in peptides (20-mer with 10 aa overlap) covering MOMP. The mapping was performed with sera from (a) UV-SvD/CAF01 ($n=4$)- and (b) Hirep1+CTH93/CAF01 ($n=10$)-immunized pigs. Serum antibody recognition of overlapping peptides (see Supplementary Table 1 for sequences) spanning MOMP SvD, and SvE- and SvF-specific sequences in VD4. The graph is based on the mean OD value of duplicates for each pig. The box indicates the 25th and the 75th percentile within the group, the horizontal line within a box represents the median, and whiskers indicate min to max values. The sequence of CSs and VDs of MOMP is depicted below the graphs.

induced in UV-SvD/CAF01-immunized pigs. Importantly, high titers of vaccine-specific antibodies were also detected on the vaginal mucosa (Figure 1c), reflecting transudation of serum into the genital tract in the vaccinated pigs.

Hirep1+CTH93/CAF01 vaccination primarily induces T-cell epitopes in the VD4 regions of MOMP from SvD, SvE and SvF

As there were no significant differences between immune responses in the groups receiving 10 and 100 μ g Hirep1+CTH93/CAF01 (shown in Figure 1), we decided to pool results from the two hybrid vaccine groups for further analysis.

T-cell responses against the individual vaccine components were characterized in more detail by stimulating with peptide pools representing the individual vaccine components (Hirep1, MOMP SvD, CT043 and CT414_{aa605-840}) (Supplementary Figure 2). In general, peptide pools representing the individual vaccine components, Hirep1, MOMP SvD and CT043, were frequently and strongly recognized in Hirep1+CTH93/CAF01-vaccinated pigs with levels of IFN- γ significantly different from naïve controls. In contrast, recognition of CT414_{aa605-840} was low. UV-SvD/CAF01-immunized pigs did not respond to peptide pools representing any of the vaccine antigens.

To localize T-cell epitopes in the individual components, PBMCs were stimulated with individual 20-mer peptides with 10 aa overlap (Supplementary Tables 1–3) covering the vaccine antigen components (Hirep1 and MOMP SvD (Figure 2), CT043 and CT414_{aa605-840} (Supplementary Figure 3A)). For MOMP, peptides covering full length MOMP SvD were used. To cover the Hirep1 fusion construct, peptides covering VD4 of SvE and SvF were included (SvE peptide 30–31, SvF peptide 27–32), when they differed in sequence from VD4 of MOMP SvD. In agreement with the lower recognition of the MOMP SvD peptide pools in the UV-SvD/CAF01-vaccinated group, only few T-cell epitopes were detected (Figure 2a). In Hirep1+CTH93/CAF01-immunized pigs, multiple T-cell epitopes were identified throughout MOMP, both in constant and variable domains (Figure 2b). T-cell epitopes were predominantly identified in a region of MOMP (peptide 25–35) comprising conserved sequences and VD4 of SvD, SvE and SvF. Especially, peptide 30 of SvD and SvE, and peptide 28 and 32 of SvF were frequently recognized. The recognition of the Sv-specific peptides, especially SvF peptide 27–32, may indicate a Hirep1-induced T-cell response. In Hirep1+CTH93/CAF01-immunized pigs, the epitopes in CT043 and CT414_{aa605-840} (Supplementary Figure 3A) were recognized by most pigs in peptides 10–13 and 18–23,

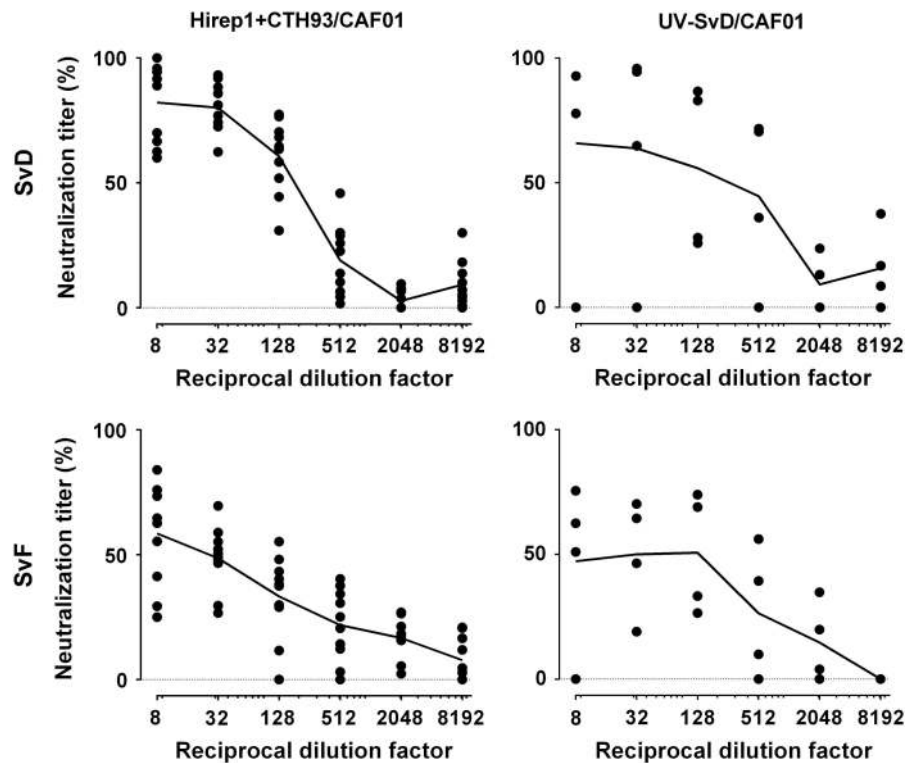


Figure 4 *In vitro* neutralization of *C. trachomatis* SvD and SvF. Serial dilutions of sera from Hirep1+CTH93/CAF01 ($n=10$)- and UV-SvD/CAF01 ($n=4$)-immunized pigs were incubated with SvD or SvF. The sera-bacteria solutions were transferred to HaK cells, and IFUs were fixed, stained and counted. Results are given as the neutralization titer of each immunized pig in relation to pooled sera from naïve controls ($n=5$). Each dot represents the mean of duplicate wells, and a connecting line between the dilutions is shown.

respectively, whereas the CT043 and CT414_{aa605-840} peptide recognition in UV-SvD/CAF01-immunized pigs was low (data not shown).

Strong antibody recognition of surface-exposed and Sv-specific regions of MOMP

To characterize antibody recognition of vaccine components, epitope mapping of linear B-cell epitopes was performed post immunizations (week 7) by evaluating serum antibody recognition of overlapping peptides (20-mer with 10 aa overlap) (Supplementary Tables 1–3) covering the individual vaccine components (MOMP SvD and Hirep1 (Figure 3), CT043 and CT414_{aa605-840} (Supplementary Figure 3B). In UV-SvD/CAF01-immunized pigs, the antibody recognition of MOMP was clustered, largely consistent with the surface-exposed VD sequences (Figure 3a), especially VD3 and VD4. The pattern was similar in Hirep1+CTH93/CAF01-immunized pigs (Figure 3b), but additionally, transmembrane and periplasmic regions adjacent to VD3 and VD4 were recognized. This demonstrates broader antibody recognition owing to presentation of more B-cell epitopes in the hybrid vaccine. Recognition of non-Hirep1 peptides in MOMP SvD (peptide # < 25) indicates a CTH93-induced response. In both vaccine groups, peptide 30 of SvD, SvE and SvF was strongly recognized. In addition, peptides of CT043 (peptide 1–2, 6–7 and 14–16) and CT414_{aa605-840} (peptide 1, 4, 14–16 and 22) were strongly recognized in Hirep1+CTH93/CAF01-immunized pigs (Supplementary Figure 3B) compared with low responses in the UV-SvD/CAF01-vaccinated group (data not shown).

Immune sera neutralizes SvD and SvF infectivity *in vitro*

Peptide 30 in MOMP contains the TLNPITAG sequence (Supplementary Table 1) associated with induction of neutralizing

antibodies. As shown in Figure 3, peptide 30 in MOMP SvD, SvE and SvF was strongly recognized by serum antibodies from both Hirep1+CTH93/CAF01 and UV-SvD/CAF01 immunized pigs. Further analysis of sera recognition of peptide 30 in MOMP SvD demonstrated high end point titers in both Hirep+CTH93/CAF01 (median: 125.000, interquartile range: 125.000–125.000)- and UV-SvD/CAF01 (median: 125.000, interquartile range: 35.000–500.000)-immunized pigs. To evaluate the neutralizing abilities of the antibodies, an *in vitro* neutralization assay³² was performed using sera collected post immunizations (week 7) and tested against SvD and SvF. We observed strong and broadly neutralizing capacity against SvD and SvF of sera from all individual animals in the Hirep1+CTH93/CAF01 group (Figure 4). Sera from pigs immunized with UV-SvD/CAF01 also had the capability to neutralize SvD and SvF, however, within this group, we observed great variability in responses ranging from no neutralization to as high as observed in the Hirep1+CTH93/CAF01 group.

Hirep1+CTH93/CAF01 vaccination protects against vaginal infection

To evaluate the protective efficacy of the vaccines, the pigs were estrus cycle synchronized and intravaginally inoculated with 3.9×10^9 inclusion forming units (IFUs) during estrus at week 7, and vaginal swabs were taken at days 3, 5, 7 and 12 pi. The levels of chlamydial 16S rRNA were quantified by qPCR, and for comparison at day 3 pi, vaginal swabs were also cultured on McCoy cells to detect the number of IFUs (Figure 5). At day 3 pi, levels of chlamydial DNA and IFUs were significantly decreased in Hirep1+CTH93/CAF01-immunized pigs compared with controls. However, as control pigs rapidly cleared

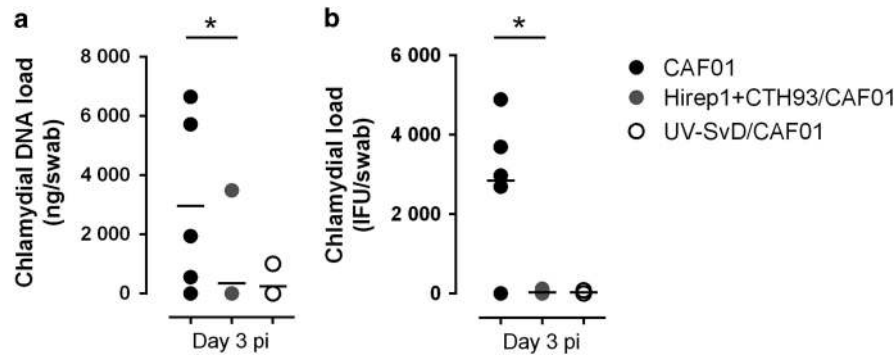


Figure 5 Vaginal loads of *C. trachomatis*. Chlamydial load in CAF01 controls ($n=5$), Hirep1+CTH93/CAF01 ($n=10$)- and UV-SvD/CAF01 ($n=4$)-immunized pigs at day 3 pi was measured in vaginal swabs by (a) qPCR (16S rRNA) and (b) swab culturing in McCoy cells. In qPCR, each dot represents the mean of triplicates for each pig. In swab culturing, each dot represents one well for each pig. Lines indicate the medians. Statistical difference was calculated using the Kruskal-Wallis test followed by Dunn's multiple comparisons test, $*P<0.05$.

the infection, this protection was only detected at day 3 pi, after which the levels were comparable (data not shown).

Immune responses post infection demonstrate significant CMI recall in vaccinated pigs

Evaluation of antibody responses after the infection was carried out in sera and vaginal swabs against Hirep1, CTH93 and UV-SvD, and results revealed no significant sera and vaginal antibody recall responses in vaccinated pigs (data not shown). Evaluation of CMI responses post challenge was carried out in PBMCs collected at days 0 and 7 pi. CMI responses to the infection alone, represented by pigs immunized with adjuvant alone (CAF01 controls) (Figure 6a), demonstrated UV-SvD-specific IFN- γ responses at day 7 pi ($P=0.0625$), and responses against Hirep1 were also increased ($P=0.0625$). In Hirep1+CTH93/CAF01-vaccinated pigs (Figure 6b), the CMI recall responses against both UV-SvD, Hirep1 and CT414_{aa605-840} was significantly increased, whereas the responses against peptide pools of CT043 and MOMP SvD_{aa34-371} were only modestly elevated. In contrast, the UV-SvD/CAF01 group only responded against UV-SvD (Figure 6c).

DISCUSSION

Considerable data support the concept that an ideal *Chlamydia* vaccine would need to elicit both humoral and CMI-based immune responses.^{33,34} Ideally, efficient mucosal neutralizing antibodies reduce initial infectious bacterial load and once intracellular, the bacteria are eliminated by a strong CMI-driven response. Trachoma studies in NHPs and clinical trials in the 1960s using whole-cell vaccines were at least partially successful in reducing conjunctival infection, but protection was short-lived¹⁴ and Sv-specific;¹³ suggesting that the infection can be controlled by a vaccine if long-term protection and broad Sv coverage are established.

In the present study, we evaluate a multi-subunit vaccine containing Hirep1+CTH93/CAF01. CTH93 is a fusion molecule consisting of three antigens identified in our discovery program: CT414_{aa605-840} (unpublished), CT043 and MOMP SvD_{aa34-371}.^{17,31,35} Hirep1 is a molecularly designed version of MOMP that induces broadly neutralizing antibodies targeting the most prevalent SvS.²⁹ To induce a combined humoral and CMI response, we used the CAF01 adjuvant, which has previously been shown to induce both antibody and Th1 cell responses³⁶ and has passed clinical safety evaluation.²⁴ The vaccines were administered intramuscularly twice, and, regardless of dose (10 or 100 μ g), induced highly specific serum and vaginal IgG

levels and strong CMI responses (Figure 1). Individual antigens were recognized with a broad spectrum of B- and T-cell epitopes presented.

Epitope mapping of CT043 and CT414_{aa605-840} demonstrated broad B-cell recognition for both, whereas especially CT043 induced a strong and broad T-cell epitope recognition pattern. CT414_{aa605-840} was initially identified in our laboratory in a phage display assay probed with sera from human patients diagnosed with a genital infection (unpublished data) and the full length protein has been shown to react with sera from infected patients,³⁵ indicating that the antigen is a B-cell target in infected humans. CT043 was originally identified as a frequent human T-cell antigen^{30,31,37} and has later been shown to elicit CD4 T-cell-mediated protection against genital infections in mice, and CT043-specific antibodies recognize the chlamydial surface.¹⁷

MOMP contains numerous human T-cell epitopes,^{38,39} consequently aa34–371 of MOMP SvD was included in CTH93 to expand the overall T-cell repertoire. T-cell epitope mapping in MOMP (Figure 2) demonstrated that in pigs, T-cell epitopes are present throughout MOMP with frequently recognized peptides in peptide 25–35 including constant sequence IV, which is known to contain several human T-cell epitopes.³⁸ In mice, Hirep1 has been shown to induce both antibody and T-cell responses to Sv-specific and conserved epitopes.²⁹ Hirep1-vaccinated mice are protected against genital challenge with *C. trachomatis* SvD and SvF, and this protection is mediated by broadly neutralizing antibodies in concert with a CMI response. Interestingly, we observe multiple vaccine-induced T-cell epitopes recognized in the VD4 region. In particular, peptide 30 in MOMP SvD and SvE containing the conserved neutralizing antibody epitope TLNPTIAG was strongly and frequently recognized. Noticeably, also Sv-specific regions covered in peptides 28, 31 and 32 were recognized. This broad and significant T-cell response covering all SvS represented in the vaccine was unique for the Hirep1+CTH93/CAF01-immunized pigs and not found in the UV-SvD/CAF01 group. Importantly, this Hirep1-specific T-cell response was significantly boosted following the infection (Figure 6), indicating a strong T-cell recall response targeting the VD4 region in MOMP during infection. This broadens the immunological importance of the VD4 region and suggests that beside a clear target for antibodies, this region is also a possible target for T cells during infection. Karunakaran *et al.*⁴⁰ recently suggested that surface proteins, including MOMP, are enriched among the *Chlamydia* T-cell immunoproteome. In that study, mouse MHC class II-bound

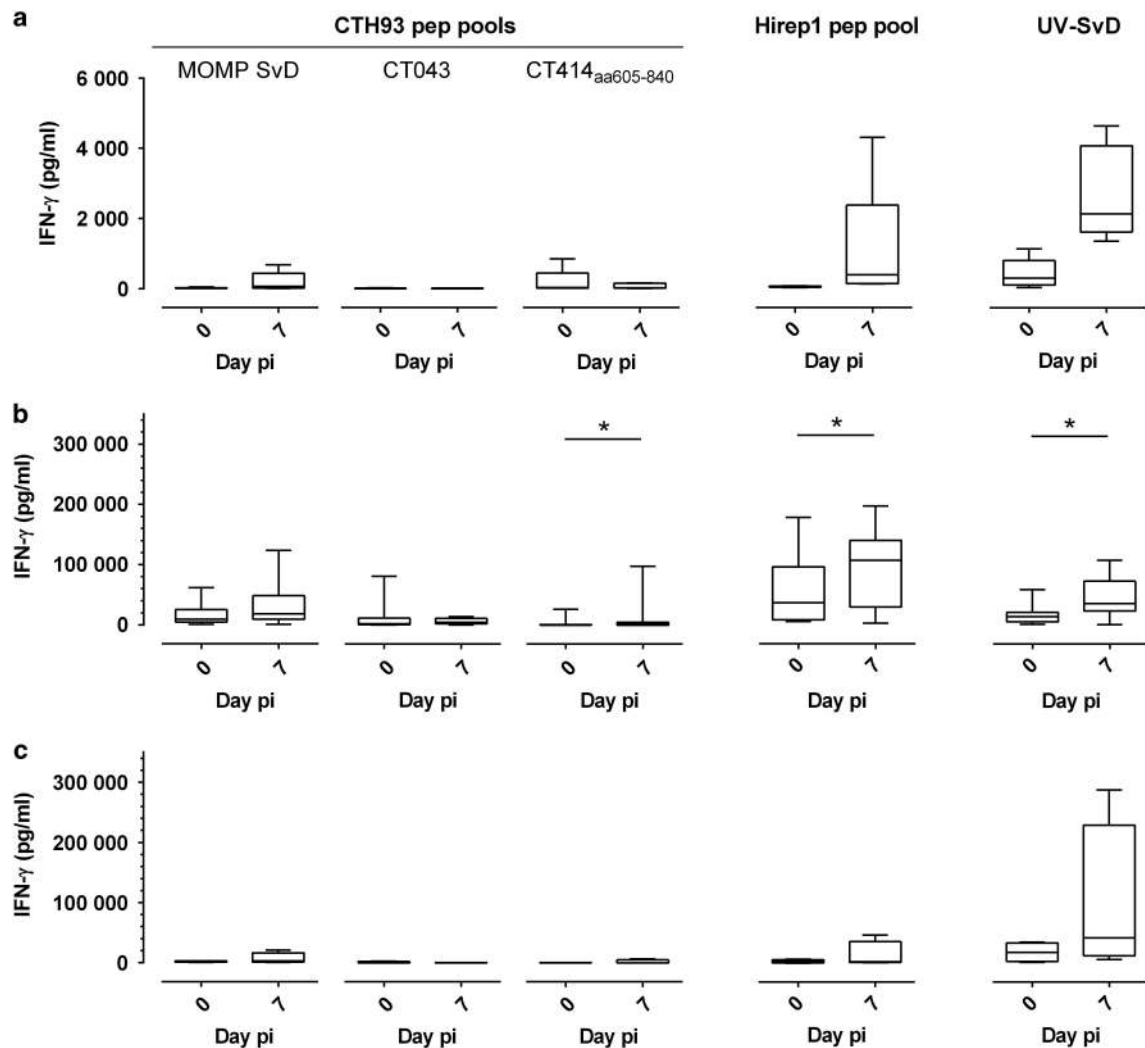


Figure 6 CMI responses post infection. PBMC were stimulated (days 0 and 7 pi) with CTH93 peptide pools (MOMP SvD, CT043 and CT414_{aa605-840}), Hirep1 peptide pool and UV-SvD. (a) Infected controls ($n=5$), (b) Hirep1+CTH93/CAF01-immunized pigs ($n=10$) and (c) UV-SvD/CAF01-immunized pigs ($n=4$). For each pig, the mean IFN- γ value of triplicate wells is shown. The box indicates the 25th and the 75th percentile within the group, the horizontal line within a box represents the median, whiskers indicate min to max values. Statistical difference was calculated using the Wilcoxon's matched-pairs rank test, * $P < 0.05$.

C. trachomatis peptides were identified by proteomics and they identified a peptide, IFDFTTLNPTIAGAGDVK, which is covered in the highly recognized peptide 30 from SvD and SvE used in the present study. Consequently, in line with our observations, this confirms that this region is an important target for vaccine-induced T cells.

Very few MOMP T-cell epitopes were induced in the UV-SvD/CAF01 group, and epitopes recognized were primarily situated within the constant sequences. This is in line with epitope mapping in *C. trachomatis*-infected humans, in which T-cell epitopes of MOMP are mainly located in the CSs,^{38,41} although also few T-cell epitopes exist in the VD_s.⁴²

In Hirep1+CTH93/CAF01-immunized pigs, B-cell epitope mapping in MOMP revealed high and frequent antibody recognition of the variable domains and adjacent sequences; in particular, VD3 and VD4 of SvD, and SvE- and SvF-specific sequences of VD4. Sera from Hirep1+CTH93/CAF01-immunized pigs recognized Sv-specific peptides of SvD, SvE and SvF, all surrounding peptide 30 containing the conserved neutralizing epitope TLNPTIAG.^{10,43} In comparison,

the B-cell recognition in UV-SvD/CAF01-immunized pigs showed a narrower epitope pattern, with fewer epitopes recognized, however, still including peptide 30.

The Hirep1+CTH93/CAF01 vaccine induced strong neutralizing antibodies consistently in all animals and sera neutralized both SvD and SvF infection, which confirms our previous findings in mice²⁹ and validates the vaccine for future testing in clinical trials. Also, the UV-SvD/CAF01 vaccine induced neutralizing antibodies in line with previously published studies,^{19,20} but the levels were varying with ranges from not neutralizing to highly neutralizing within the group. Although the UV-SvD/CAF01 group did also recognize peptide 30 encompassing the neutralizing TLNPTIAG epitope, we cannot exclude that other surface proteins may also be involved in induction of neutralizing antibodies, such as PorB⁴⁴ and PmpD⁴⁵ or even conformational epitopes in MOMP. Although with neutralizing potential, antibodies induced by whole-cell vaccines¹³ and derivatives thereof, that is, purified native MOMP, have been shown to promote homotypic immunity in NHPs,⁴⁶ an obstacle that is overcome by the use of recombinant Hirep1 subunit vaccine.

Deep vaginal challenge established an infection that resolved relatively fast and although we observed shedding up to day 12 pi, the levels were very low and no differences between groups were found after day 3 pi (data not shown). Nevertheless, at the very onset on infection (day 3 pi), we did observe protection in both the Hirep1+CTH93 and UV-SvD groups (Figure 5). Given the time point and the neutralizing capacity of sera in both groups, this protection is likely mediated by, at least in part, antibodies. This would also be in line with our previous studies in mice where we observe significant control of bacteria at the early time point (day 3 pi).²⁹ Interestingly, we find that Hirep1+CTH93/CAF01-vaccinated pigs mount a significant T-cell recall response directed at peptides representing CT414 and especially Hirep1 indicating that this region is not only a target for antibodies but also for T cells during the infection.

In non-vaccinated controls, the infection induced a T-cell response against UV-SvD. The UV-SvD/CAF01 group had a recall T-cell response against UV-SvD, whereas responses against Hirep1 and CTH93 components were not boosted. In Hirep1+CTH93/CAF01-immunized pigs, CMI responses against UV-SvD, Hirep1 and CT414_{aa605-840} were significantly increased, demonstrating a recall response against both Hirep1 and CTH93 components (Figure 6). This indicates that although the protective efficacy of the Hirep1+CTH93/CAF01 vaccine and the UV-SvD/CAF01 is equal, the antigen specificity behind this protection might be different.

In conclusion, evaluation of Hirep1+CTH93/CAF01 vaccination in pigs demonstrates a highly immunogenic vaccine with uniform patterns of T- and B-cell epitopes against vaccine components, including high and frequent recognition of SvE- and SvF-specific sequences. Hirep1+CTH93/CAF01 vaccination generates both neutralizing antibodies against SvD and SvF and protective immune responses comparable with the UV-SvD/CAF01 vaccine, but the hybrid vaccine would be a safer alternative, excluding antigens associated with immunopathology.

METHODS

Ethics statement

The experimental procedures were approved by the Danish Animal Experiments Inspectorate of the Danish Veterinary and Food Administration, license no: 2008/561 – 1581.

Chlamydia trachomatis

C. trachomatis SvD (Trachoma type D strain UW-3/Cx, ATCC VR-885) was propagated in Hela cells (Human epithelial cell line, Hela 229, ATCC CCL-2.1), and chlamydial elementary bodies (EBs) were harvested and purified as described by Olsen *et al.*³⁷ with few modifications: Hela cells were cultured in six-well plates and were infected with 1.5 *C. trachomatis* SvD IFUs per Hela cell. Instead of diethylaminoethyl-dextran treatment, the cells were centrifuged at 750 g for 1 h. For use as vaccine antigen, the chlamydial EBs were inactivated by exposure to ultraviolet light emission as described,⁴⁷ and the inactivation step was confirmed by culture in Hela cells. Finally, the content of protein was quantified by the bicinchoninic acid assay, as described in the Micro BCA Protein Assay Kit (cat no 23235, Thermo Scientific, Rockford, IL, USA). For use as the challenge infection, the purified chlamydial EBs were IFU quantified in Hela cells.

Chlamydia trachomatis antigens

Recombinant fusion proteins CTH93 and Hirep1 were produced as follows; CTH93 was designed as a fusion protein of CT043 (full length), CT414_{aa605-840} and MOMP SvD_{aa34-371} (Supplementary Figure 1) with an added N-terminal histidine tag, all sequences obtained from the original annotation published in the study by Stephens *et al.*⁴⁸ The Hirep1 hybrid vaccine was composed of VD4 regions from SvD, SvE and SvF, including residues in the constant regions adjacent to VD4.²⁹ Synthetic DNA constructs were codon-optimized for

expression in *Escherichia coli* followed by insertion into the pJexpress 411 vector (DNA2.0). To avoid disulfide bridge formation during recombinant production, cysteines in MOMP were exchanged systematically with serines. Purification was carried out essentially as described by Olsen *et al.*⁴⁹

Vaccines

The hybrid antigens were diluted in 1 ml sterile Tris buffer (10 mM, pH 7.4) and were mixed by vortexing with 1 ml cationic liposomal adjuvant CAF01 (500 µg of glycolipid trehalose 6,6'-dibehenate and 2500 µg of dimethyldioctadecylammonium bromide) and sucrose into a total volume of 2 ml isotonic (9% sucrose) vaccine.

Animals

Nineteen, sexually mature, female Göttingen minipigs were purchased from Ellegaard Göttingen Minipigs A/S, Dalmose, Denmark, which is breeding microbiologically well-defined minipigs in a barrier unit, in which the herds are health-monitored twice a year, following FELASA recommendations. At delivery, the minipigs were 5–6 months old. The pigs were housed in groups of four to six, fed with standard minipig diet twice a day and water *ad libitum*. The study was carried out in a laboratory animal isolation facility at University of Copenhagen.

Design of study

The 19 female minipigs were randomly divided into four groups immunized with either CAF01, 10 µg Hirep1+CTH93/CAF01, 100 µg Hirep1+CTH93/CAF01 or 125 µg UV-SvD/CAF01; see Table 1. The vaccines were administered twice intramuscularly in the neck muscles, week 0 and week 3. Pigs immunized with CAF01 and UV-SvD/CAF01 are also included in another study focusing on immunohistochemical analysis supported by immunological data and chlamydial shedding. To estrus synchronize the pigs, they were treated orally with 20 mg progestagens (Regumate Equine, MSD Animal Health, Ballerup, Denmark) for 18 days and 6 or 8 days after cessation of treatment, they went into estrus with proximity of a teaser boar. The pigs were sedated with Zoletil mixture (2.5 mg ml⁻¹ tiletamine, 2.5 mg ml⁻¹ zolazepam, 20 mg ml⁻¹ xylazine, 100 mg ml⁻¹ ketamin, 10 mg ml⁻¹ butorphanol) and inoculated intravaginally at the most cranial part of the vagina with 3.9 × 10⁹ IFUs of *C. trachomatis* SvD diluted in 5 ml SPG (0.2 M sucrose, 20 mM sodium phosphate and 5 mM glutamic acid buffer) with an insemination catheter (Osiris, cat no 902011, E-Vet, Haderslev, Denmark). The pigs were inoculated 30 or 32 days after the second immunization and were killed at day 12 pi by deep anesthesia followed by exsanguination. Rectal temperature and clinical signs were recorded after each vaccination day and on a daily basis after the challenge. Blood samples and vaginal swabs were taken at week 0 and 3 and at days 0, 7 and 12 pi; vaginal swabs were additionally taken at days 3 and 5 pi. For collection of sera, unstabilized blood was centrifuged at 2400 g for 15 min, and serum was transferred into new tubes and kept at 20 °C until analysis.

Vaginal sampling

Before vaginal sampling, the pigs were anesthetized in Zoletil mixture, and the outer vulva was rinsed in water and 70% ethanol. By using a speculum, a swab was introduced into the deeper part of the vagina, close to the cervix, where the swab was rolled clockwise twice along the vaginal wall. The swabs were kept in tubes containing 1 ml SPG. Three autoclaved glass beads were added to each swab sample, followed by whirl mixing for 2 min. The samples were kept at –80 °C until further analysis. On day 3 pi, the swabs were cultured fresh as described below.

Detection of chlamydial load by culturing

The protocol for detection of chlamydial load in vaginal swabs was modified from the protocol of Hansen *et al.*⁵⁰ Briefly, McCoy cells (murine fibroblast cell line, ATCC CRL-1696) were cultured in infection media (RPMI 1640 supplemented with 5% fetal calf serum, 10 mM HEPES, 1% (vol/vol) non-essential amino acids for MEM Eagle, 2 mM L-glutamine, 1 mM pyrovalate, 10 µg ml⁻¹ gentamicin, 70 µM mercaptoethanol) in 48-well plates for 24 h. *C. trachomatis* bacteria were detected with sera from MOMP and *C. trachomatis*

heat shock protein 60-immunized rabbits followed by staining with 4 $\mu\text{g ml}^{-1}$ Alexa Flour 488-labeled goat-anti rabbit antibody (Cat no: A11008, Life Technologies, Paisley, UK), and McCoy cells were stained with propidium iodide. At least 20 ($\times 40$) fields were counted for IFU in each well using an Olympus IX71 fluorescence microscope (Ballerup, Denmark), and the no of IFUs per swab was calculated.

Detection of chlamydial load by quantitative PCR

DNA extraction from the vaginal swab samples was performed with Chelex 100 (Bio-Rad, Life Science, Copenhagen, Denmark). One hundred microliters of the swab material were mixed with 300 μl of a 20% Chelex solution in TE buffer (T9285, Sigma Aldrich, Copenhagen, Denmark), vortexed for 60 s and incubated at 96 °C for 10 min. The sample was then centrifuged for 10 min at 17 500 g and 4 °C and hereafter triplicates of 5 μl of the supernatant was used for PCR. Real-time qPCR detection of *C. trachomatis* in the vaginal swab samples was performed by detection of the 16S rRNA gene. An internal control (IC) was run in all samples to detect possible inhibition of the PCR. The IC primers were bought from TAG Copenhagen A/S (Copenhagen, Denmark): IC-F 5'ACCGCTCAGGCATTGCT-3' and IC-R 5'CCGGGACGTATCATGCT3'. The remaining primers and probes were bought from Applied Biosystems: Ct 16s-F GGATCTTCGGACCTTTCGGT; Ct 16s-R AATCTCTCAATCCGCCTAGACA; Ct 16s-probe FAM-AAGGGAGAGTCTATGTGATAT-MGBNFQ, IC-probe NED-CCTTCGTGATATCGGACGTTGGCTG-MGBNFQ. The assay was performed with: Perfecta qPCR SuperMix (UNG, low ROX, 95066-02K (2000 rx), Quantum Biosciences). The samples were run on a StepOne Real-time PCR instrument (Applied Biosystems) and the instrument was programmed to run 2 min at 95 °C and 40 cycles of denaturation at 95 °C for 15 s and annealing/extension at 60 °C for 1 min. The C_t cutoff was determined to be 37, hence C_t values greater than 37 were considered as nonsense.

Measurement of antigen-specific antibodies in serum and in vaginal swabs

Determination of antigen-specific antibodies in serum and vaginal swabs was carried out by indirect ELISA. Ninety-six-well plates were coated overnight with 0.4 μg of UV-SvD (using Polysorp plates, NUNC, Roskilde, Denmark) and 0.1 μg Hirep1, 0.1 μg CTH93 or 0.1 μg MOMP SvD peptide 30 (using Maxisorp plates, NUNC) in bicarbonate/carbonate buffer (pH 9.6) per well. Study samples were added in duplicates in serial dilutions and incubated overnight. An internal standard was used on each plate to correct for plate-to-plate variation. Sample antibodies were detected with horse radish peroxidase-conjugated goat-anti pig IgG secondary antibodies (1:10,000, AAI41P, Serotec, Oxford, UK). The wells were visualized with TMB PLUS substrate (Kem-En-Tec, Taastrup, Denmark) and 0.5 M sulfuric acid. The plates were read at the wavelength of 450 nm with correction at 650 nm. To compensate for plate-to-plate variation, the log-transformed OD mean of the titrated internal standard on each plate was correlated with the log-transformed mean of the internal standard on all plates analyzed simultaneously, resulting in a regression equation for each plate. For each plate, the sample OD values were transformed according to the regression equation. After that, the end point titers were calculated as the highest sample dilution before reaching the cutoff. The cutoff value was calculated for each individual antigen as the mean of the lowest dilution from all week 0 samples added to two times the standard deviation.

B-cell epitope mapping was performed as described for detection of antibodies with few modifications: 96-well Maxisorp plates were coated overnight with duplicates of overlapping peptides (20-mer with 10 aa overlap) (5 $\mu\text{g ml}^{-1}$) covering vaccine antigens (MOMP, Hirep1, CT043 and CT414_{aa605-840}). The peptides are listed in Supplementary Tables 1. After the blocking and washing steps, sera drawn after two immunizations (week 7) were diluted (1:200) and added to the wells and serial dilutions of an internal standard were used to minimize plate-to-plate variation, and the plates were incubated overnight and developed as described. The results were given as OD values.

Measurement of neutralizing antibodies *in vitro*

The neutralizing capacity of the serum antibodies were investigated by an *in vitro* assay, as described.³² HaK cells (Syrian Golden hamster epithelial cell line, ATCC CCL-15) were cultured in infection media (as described for culturing swabs in McCoy cells) in Nunclon sterile 96-well plates (Cat no: 167008, Thermo Scientific). *C. trachomatis* SvD or SvF EBs were incubated for 30 min at 37 °C with titrations of heat-inactivated sera from immunized and naïve pigs. EB-sera mixtures were added to the confluent HaK cell layers in duplicates, and SPG and pooled naïve sera from controls were added on each plate as controls. After 2 h of incubation at 35 °C, the EB-sera mix was removed from the wells, and new infection media with 0.5% glucose and 1 $\mu\text{g ml}^{-1}$ cycloheximide were added, followed by incubation at 37 °C for 42 h. *Chlamydia* bacteria were detected with sera from CT043 and cHsp60-immunized rabbits followed by staining with 4 $\mu\text{g ml}^{-1}$ Alexa Flour 488-labeled goat-anti rabbit antibody (Cat no: A11008, Life Technologies), and HaK cells were stained with propidium iodide. At least 20 fields ($\times 40$) were counted for IFUs in each well using an Olympus IX71 fluorescence microscope, and for each dilution, the % inhibition of the sample related to the pooled naïve control sera on the same plate was calculated by the formula: Neutralization % = 100—((Mean IFUs of duplicate wells/mean IFUs of pooled sera from naïve controls) $\times 100$).

In vitro stimulation of PBMCs

PBMCs were isolated by density gradient centrifugation on Lympholyte-Mammal medium (1.086 g cm^{-3} at 22 °C, Cedarlane Labs, Skanderborg, Denmark) and cultured in RPMI-1640 with 5×10^{-5} M 2-mercaptoethanol, 1 mM glutamine, 1% pyruvate, 1% penicillin-streptomycin, 1% HEPES and 10% fetal calf serum (Invitrogen, Taastrup, Denmark), with 2×10^5 cells/well were added in Nunclon microtiter plates (NUNC). The cells were stimulated in triplicates with UV-SvD, Hirep1, CTH93, CT043, CT414_{aa605-840} and MOMP in 10 $\mu\text{g ml}^{-1}$. Synthetic peptides (20-mer) with 10 aa overlap covering CT043, CT414_{aa605-840}, MOMP and Hirep1 were purchased from (GenScript, Piscataway, NJ, USA) and used for stimulation as pools (5 $\mu\text{g ml}^{-1}$ of each peptide in the peptide pools) or as individual peptides (5 $\mu\text{g ml}^{-1}$). The peptides are listed in Supplementary Tables 1. Wells with medium alone and with Staphylococcal enterotoxin B (1 $\mu\text{g ml}^{-1}$) were included as negative and positive control, respectively. After 3 days of incubation at 37 °C with 5% CO₂, cell culture supernatants were harvested and stored at -20 °C.

Determination of IFN- γ in cell culture supernatants by ELISA

The levels of IFN- γ in cell culture supernatants were determined by a monoclonal ELISA using anti-porcine IFN- γ monoclonal antibody clone P2F6 (MP700, Thermo Scientific) and biotin-conjugated mouse anti-pig IFN- γ monoclonal antibody clone P2C11 (559958, BD Pharmingen, San Diego, CA, USA), as described by Riber *et al*.⁵¹ The plates were developed with TMB Plus (Kem-En-Tec) in the dark, and the reaction was stopped by adding 0.5 M sulfuric acid. The OD was read at wavelength 450 nm with correction at 650 nm. From the standards, a double logarithmic curve was constructed to quantify the levels of IFN- γ in the samples.

Statistics

All statistical analyses were performed using Graph Pad Prism software. To compare time-dependent differences within a group, we used the paired, non-parametric Friedman test followed by Dunn multiple comparisons test. Comparison of difference between selected groups at the same time point was performed by the non-parametric Kruskal-Wallis test followed by Dunn multiple comparisons test. Evaluation of variance within a group at two timepoints was carried out by Wilcoxon's matched-pairs rank test. Significant difference was defined as $P < 0.05$ (* $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$).

CONFLICT OF INTEREST

P Andersen, AW Olsen and F Follmann are co-inventors on a patent application relating to Chlamydia vaccines. All rights have been assigned to Statens Serum Institut, a Danish not-for-profit governmental institute.

ACKNOWLEDGEMENTS

We thank Vita Skov for technical assistance, and the staff of Laboratory Animal Unit at the University of Copenhagen for taking care of the pigs and for assistance during procedures in the animal facilities. We further thank Maria Rathmann Sørensen for development of a *C. trachomatis*-specific qPCR. The study was supported by Innovation Fund Denmark [069-2011-1], the European Commission through the ADITEC consortium contract [FP7-HEALTH-2011.1.4-4-280873], Aase og Ejnar Danielsen's fond and Danish Council for Independent Research, Medical Sciences [09-067407].

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