Development of a vaccine for Chlamydia trachomatis: challenges and current progress

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Abstract: Chlamydia trachomatis remains an enigmatic bacterial pathogen with no vaccine yet available to treat human ocular and genital tract infections caused by tissue-tropic serovars of the organism. Globally, it is the leading cause of preventable blindness as well as the leading cause of bacterial sexually transmitted infections. The pathogen has a range of virulence factors that enable it to successfully evade both the innate and adaptive immune system of the host. The host immune system, although protective, paradoxically is also associated closely with the pathologies of trachoma and pelvic inflammatory disease – disease sequelae of some chlamydial infections and reinfections in some genetically susceptible hosts. In this review, we focus on what is known currently about the pathogenesis of ocular and genital infections caused by this mucosal pathogen. We also discuss novel insights into the pathogenesis of infections caused by the genital and ocular serovars of C. trachomatis, including a discussion of both pathogen and host factors, such as the human microbiota at these mucosal sites as well as the current immunological challenges facing vaccine development. Finally, we discuss the current progress toward development of a vaccine against C. trachomatis. A wide range of recombinant protein antigens are being identified and, hence, are available for vaccine trials. A plasmid-free live strain has recently been produced and evaluated in the mouse (Chlamydia muridarum) and monkey (C. trachomatis) models. The data for ocular infections in the monkey model was particularly encouraging, although the path to regulatory approval of a live vaccine is still uncertain. While still a major challenge, vaccines for ocular and genital C. trachomatis infections are looking more promising.

Keywords: C. trachomatis, ocular and genital infections

Chlamydial epidemiology: global burden of disease

Chlamydia trachomatis is an obligate intracellular bacterial pathogen primarily of human conjunctival and urogenital columnar epithelial cells.1,2 Infections of cells at these mucosal surfaces with tissue-tropic ocugenital serovars of C. trachomatis can result in trachoma (ocular serovars A, B, Ba, and C),3 the leading cause of infectious blindness, and can also result in the leading cause of bacterial sexually transmitted infection (STI) (genital serovars D–K, Da, Ia, and Ja).4 The STIs can lead to tubal infertility in infected, untreated (or treatment failure) women as a result of ascension of the bacteria into the endometrium and fallopian tubes.5 The global burden of these infectious diseases remains a significant public health problem as many primary infections remain undiagnosed and untreated due to the largely asymptomatic nature of each disease. It is estimated that 85% of women with STIs caused by C. trachomatis are asymptomatic.6 If left untreated, primary chlamydial infections may also persist...
for years and cause disease sequelae in some infected hosts. Indeed, it is estimated that up to 40% of women with an untreated chlamydial genital tract infection develop pelvic inflammatory disease (PID) and also that one in four women with PID will develop infertility. Repeat detection of \textit{C. trachomatis} commonly occurs in up to 20% of patients, and if the same strain is detected by genotyping but not by culture, this may in fact represent a persistent, rather than a recurrent, infection.

Globally, the World Health Organization (WHO) estimates that annually over 105 million adults have newly acquired STIs caused by \textit{C. trachomatis}. In the USA, between 2007 and 2012, the overall chlamydial prevalence was 1.7%, suggesting that approximately 1.8 million infections were nationally prevalent among sexually active females aged 14–39 years in this developed country. In those sexually active females aged 14–24 years, chlamydial prevalence was reportedly 4.7% overall, with 13.5% prevalence among non-Hispanic blacks. Notably, however, \textit{C. trachomatis} STIs are much more important in developing countries, with estimates of 8.3 million new cases in the WHO African Region and 7.2 million new cases in the WHO South-East Asia Region.

Trachoma is responsible for the global visual impairment of approximately 2.2 million people, 1.2 million of whom are irreversibly blind from the disease. Current reports are that 51 countries are known or suspected to be endemic for blinding trachoma, with an additional seven previously endemic countries (Gambia, Ghana, Iran, Morocco, Myanmar, Oman, and Vietnam) reporting that they have now achieved the outcome indicator target for “elimination of blinding trachoma as a public health problem”. Despite this encouraging outcome, the 2014 estimate for the global population at risk of trachoma is 232 million people.

Chlamydial disease pathogenesis both for ocular and for genital infections is immune-mediated, with bacterial infections producing inflammatory reactions at both mucosal sites. The immune responses to chlamydial infections are partially protective; however, these local inflammatory reactions often result in asymptomatic infections that can covertly spread to cause tissue damage and scarring complications that can lead to blindness and infertility in a subset of those infected.

Public health programs currently are implemented worldwide in attempts to control for both ocular and genital infections caused by \textit{C. trachomatis}. The SAFE (Surgery, Antibiotic treatment, Facial cleanliness, and Environmental improvements) strategy is recommended by the WHO for trachoma control primarily to interrupt bacterial transmission. Worldwide, by the end of 2013, 31 countries reported that they were actively implementing the SAFE strategy. In addition, national screening programs have been used in many countries for over 20 years along with early treatment interventions for controlling \textit{C. trachomatis} STIs. However, both of these public health control programs, although effective at reducing active \textit{C. trachomatis} infections and producing declined rates of disease sequelae, are also problematic. For the STI programs, there are two main problems: first, the greatest areas affected worldwide with these infections, ie, developing countries, have virtually no control programs in place; and second, it is hypothesized (the “arrested immunity” hypothesis) that early antibiotic treatment interventions may in fact disrupt the acquisition of protective immunity in these populations. Trachoma elimination efforts also confront major obstacles for the successful implementation of the SAFE program due to conflict and insecurity in several areas of the 46 countries comprising the WHO African Region. A further problem is that the SAFE program involves mass antibiotic treatments of populations which in itself could potentially induce antibiotic resistance.

Obviously the current control programs for ocular and genital infections caused by \textit{C. trachomatis} are suboptimal, with rising reinfection rates still occurring despite, or perhaps even attributed to, some aspects of these control efforts. Thus vaccines that are designed to prevent the acquisition and transmission of chlamydial ocular and genital infections, as well as to prevent the development of immunopathologies following these infections, clearly would be advantageous for controlling trachoma and STIs caused by \textit{C. trachomatis}.

**New insights into \textit{C. trachomatis} pathogenesis**

Infections of ocular and genital mucosae with the genetically similar serovars of \textit{C. trachomatis} damage epithelial cells eliciting complex immune events to control disease outcome. However, chronic inflammatory responses involving both innate and adaptive immunity can have pathological consequences for the host with submucosal tissue remodeling and scarring of the conjunctival surface and fallopian tubes causing the symptomatic diseases of trachoma and PID in some patients.

Disease severity following chronic inflammatory responses to persistent or repeated reinfections is associated with pathogen factors including virulence gene variants and the \textit{C. trachomatis} plasmid. Infection susceptibility and disease pathogenesis is also associated with
a myriad of host factors including genetic polymorphisms and related immune responses. Changes in bacterial community structures and diversities at the ocular and genital mucosal sites are also important host factors that influence chlamydial disease outcomes. For example, it is becoming increasingly apparent that changes in the conjunctival and vaginal microbiomes of the host occur in trachomatous disease and also influence genital tract infections caused by *C. trachomatis*.20,21

**Pathogen factors**

Although the ocular and genital strains of *C. trachomatis* are strictly differentiated, with ocular strains lacking tryptophan synthase function,22 the genomes of *C. trachomatis* serovars are more than 99% identical,23,24 and the host–pathogen interactions in chlamydial infections are known to be serovar-specific. A recent report from a Dutch STI clinic showed that in comparison with the other urogenital isolates, serovars D and E, two of the most prevalent urogenital serovars worldwide,25,26 induced the highest serological immunoglobulin G responses in women with urogenital *C. trachomatis* infections.27 In addition, it has also recently been shown that not only is the host–*Chlamydia* interaction serovar-specific, but that it is also able to induce different levels of immune responses in disparate host cells. When *C. trachomatis* serovars Ba and D infected monocytes and dendritic cells (DCs), lower levels of inflammatory cytokines were produced in the monocytes allowing the serovars to persist while higher cytokine levels were induced in the DCs causing degradation of serovars in these cells.28

**Chlamydial plasmid**

Important virulence factors of *C. trachomatis* that link to disease severity include the translocated actin-recruiting phosphoprotein (Tarp)29 and the Inc. proteins found on the surface of the parasitophorous compartments (inclusions) inside infected host cells.30 Another key *C. trachomatis* virulence factor is the highly conserved 7.5 kB cryptic plasmid that regulates the transcription of chlamydial plasmid-responsive chromosomal loci31 and expression of pgp3,32 the plasmid gene encoding the virulence factor Pgp3.33

Studies using live attenuated chlamydial strains cured of the plasmid reveal that the plasmid contributes both to STI and to blinding trachoma pathogenesis. Genital tract infection of mice with the inflammation-inducing, plasmid-containing wild-type *C. trachomatis* Lymphogranuloma venereum (LGV) serovar L2 caused infertility in these animals, but infection with the non-inflammation-inducing, plasmid-free strain did not.34 The plasmid is also a crucial virulence factor in the nonhuman primate infection model of trachoma, where it has been shown that infections of macaque eyes with *C. trachomatis* organisms lacking the cryptic plasmid are highly attenuated and did not produce measurable ocular clinical pathology in these nonhuman primates,17 with recent confirmation that loss of pgp3 alone is enough to attenuate the pathogenesis of acute chlamydial conjunctivitis in macaques.35 It is also known that the plasmid is not naturally transferred readily between clinical isolates supporting a host–plasmid tropism relationship.36 This has been confirmed by recent experiments aimed at developing a transformation system for non-LGV isolates of the ocular tropic *C. trachomatis* serovar A where it was observed that stable plasmid transformants were only obtained when the transforming plasmid shuttle vector was matched with the parental strain.37 More recently, transcriptional profiling of cultured human epithelial cells infected with plasmid-deficient and -proficient *C. trachomatis* organisms has revealed statistically significant increases in the levels of expression of host genes coding for proinflammatory, immune suppression, and cell growth and fibrosis proteins in plasmid-bearing strains.38 This latter observation further supports the vital role that the chlamydial plasmid plays in immune avoidance and host cell inflammatory responses to *C. trachomatis* infections. Overall, it is interesting to note that the vast majority of *C. trachomatis* isolates have the plasmid. Naturally occurring plasmid-free strains are very rare. This confirms the central role that the plasmid has for chlamydial virulence and survival. While not proven, the plasmid products may also enhance transmissibility between hosts in some way.

**Host factors**

Host factors contributing to *C. trachomatis* disease severity include genetic variation in epithelial cells (eg, interferon [IFN] gamma receptors) and immune cells (eg, human leukocyte antigen [HLA] alleles on monocytes and interleukin [IL]-10 promoter polymorphisms). Indeed, host polymorphisms in immune response genes – immunogenetic risk factors – are known to be significantly associated with susceptibility to, and severity of, *C. trachomatis*-induced inflammatory fibrotic diseases of trachoma and tubal factor infertility (TFI).16

In trachoma, several host inflammatory response genes have been identified as contributing to the severity of fibrotic disease leading to conjunctival scarring and trichiasis (turning-in of the eyelashes).39–41 For example, part of the host immune system’s initial defense against *C. trachomatis* infection is the activation and cytotoxic response of natural
killer (NK) cells which has been shown to be an important determinant of the severity of active trachoma.\textsuperscript{42,43} Recently, it has been reported that variants of the HLA gene C (HLA-C), in certain combinations with killer-cell immunoglobulin-like receptor (KIR) genes (eg, the HLA-C2 homozygous KIR2DL2\textsuperscript{a}, KIR2DL3\textsuperscript{a} genotype), are associated with a sixfold increase in the relative risk of scarring damage in trachoma, and these authors suggest a model of NK-mediated scarring in trachoma in which HLA-C2 genotype favors chronic infection while KIR2DL2/L3 heterozygosity favors chronic inflammation.\textsuperscript{44}

Immunogenetic associations are also widely reported in many published studies for \textit{C. trachomatis}-induced genital tract pathology such as TFI. These studies have investigated both immunological regulation of infection focused on genes that code cytokines and HLAs as well as genes related to fibrosis and tissue remodeling such as the matrix metalloproteinases (MMPs). The IL-12-related single nucleotide polymorphism \textit{IL12B} 1188 A/C was reportedly associated with TFI and the severity of tubal damage in a study of 164 TFI women with various degrees of tubal damage, 137 of whom had evidence of a past \textit{C. trachomatis} infection.\textsuperscript{45} The chemokine CXCL13 (the ligand for CXCR5) was induced in fallopian tube tissue following \textit{C. trachomatis} infection of the genital tract\textsuperscript{46} and the frequency of CXCR5 single nucleotide polymorphism +10950T>C (rs 3922) was found to decrease in women who developed tubal pathology after a \textit{C. trachomatis} infection.\textsuperscript{47}

In both ocular and sexually transmitted \textit{C. trachomatis} infections, a specific genotype of the IL-10-coding \textit{IL10} gene has been associated with disease. In scarring trachoma, the \textit{IL10}-1082 GG genotype is associated with an increased risk of trachoma\textsuperscript{48} while the \textit{IL10}-1082/-819/-592 GCC haplotype has been associated with a lower risk of recurrent infection in STIs.\textsuperscript{49} The tumor necrosis factor TNF-308A allele has also been associated with both ocular and genital \textit{C. trachomatis} disease severity. This allele has been correlated with increased TNF production and increased risk of pathological sequelae in trachoma\textsuperscript{50} and moderate-to-severe adhesions in TFI.\textsuperscript{44}

**Conjunctival and vaginal microbiomes**

The human microbiome and particularly the diversity of commensal bacteria at healthy conjunctiva\textsuperscript{50} and vagina\textsuperscript{51} is increasingly being investigated for its impact on ocular and reproductive health in humans following infections with the tryptophan auxotroph \textit{C. trachomatis}.

Of relevance to note here is that: 1) chlamydial growth and development in epithelial cells is inhibited by the cytokine IFN-\textgamma that induces indoleamine-2,3-dioxygenase that can catabolize tryptophan\textsuperscript{52-55} and 2) genital (D–K), but not ocular (A–C), serovars have retained a functional tryptophan synthase that enables them to salvage any indole from their environment and synthesize tryptophan for growth.\textsuperscript{22} Also of note is that tryptophan deficit causes \textit{Chlamydia} to enter the persistent (viable but non-cultivable\textsuperscript{56}) growth form (reviewed in\textsuperscript{57}). The ability of \textit{C. trachomatis} to develop into a persistent form has been suggested as key pathogenetic mechanism underlying ocular and genital infections and the subsequent chronic disease sequelae of blinding trachoma, PID, ectopic pregnancy, and infertility. This is likely in part due to that fact that the persistent form of the organism can evade the host immune response and is more difficult to eliminate with antibiotics.\textsuperscript{59} Subfertile women with tubal pathology also have serological markers of persistent \textit{C. trachomatis} infections and these markers are significantly more common when compared to women without tubal pathology.\textsuperscript{59}

The abundance and composition of bacterial communities at the conjunctival site in healthy (or asymptomatic) humans have been characterized using 16S deep sequencing of the bacterial ribosomal gene in two studies. The first study was from four healthy American subjects and revealed two prominent non-indole-producing genera, \textit{Pseudomonas} and \textit{Bradyrhizobium}, in the ocular microbiomes of the 69.3% of sequence reads categorized to genus level, in addition to \textit{Propionibacterium} and \textit{Corynebacterium}.\textsuperscript{58} However, in contrast to these findings, a more recent study of 105 Gambian participants with normal healthy conjunctivae has reported six genera found in at least 80% of participants, and these included \textit{Corynebacterium}, \textit{Streptococcus}, \textit{Propionibacterium}, \textit{Staphylococcus}, \textit{Bacillus}, and \textit{Ralstonia}, with \textit{Corynebacterium} and \textit{Streptococcus} higher in 115 participants with conjunctival scarring, and with \textit{Corynebacterium} in even higher abundance in adults with clinical signs of scarring and trichiasis.\textsuperscript{20} Despite the fact that indole-producing \textit{Corynebacterium} is a predominant component of healthy eyes, this genus has also been reported in trachomatous eyes,\textsuperscript{60} suggesting that changes in the normal conjunctival microbiome occur during trachomatous disease and also suggesting that non-chlamydial bacteria are associated with clinical signs of trachoma.\textsuperscript{20} Interestingly, genital tract serovars of \textit{C. trachomatis} can also cause inclusion conjunctivitis, a sexually transmitted disease that generally follows autoinoculation from contaminated...
genital secretions. Classically, chlamydial conjunctivitis initially affects one eye and this was reported in a recent case study. It is therefore tempting to speculate that perhaps the indole produced by Corynebacterium is scavenged by the genital strains of C. trachomatis that are able to use indole for growth using the functional tryptophan synthase produced by these genital serovars.

In the vaginal microbiome of apparently healthy reproductive-age women, sequencing of ribosomal RNA genes revealed that non-indole-producing Lactobacillus species such as L. iners, L. crispatus, L. gasseri, and L. jensenii comprise 73% of the bacterial community while the remaining consisted of a large heterogeneous group of strictly anaerobic microorganisms including Prevotella sp. that may express tryptophanase to produce indole. This latter bacterial community is the common and apparently normal vaginal community found in black and Hispanic women.

Increased risks of acquiring and transmitting STIs such as HIV have been shown to be associated with decreased Lactobacillus species in the vaginal microbiota. The interactions of vaginal microbiota and C. trachomatis were recently studied by analyzing the effects of Lactobacillus strains (L. brevis and L. salivarius) on the different phases of C. trachomatis developmental cycle. Lactobacilli had an adverse effect on chlamydial elementary bodies, on chlamydial adsorption to epithelial cells, and on intracellular phases of chlamydial replication. However, lactobacilli had protective effects toward persistent forms of C. trachomatis that were induced by HSV-2 coinfection. The role of vaginal coinfections on human C. trachomatis genital tract infections has recently been reviewed with particularly focus on the suggestion that an altered vaginal microbiome (as occurs in bacterial vaginosis) may provide a source of indole in vivo and that genital chlamydial strains can use indole to grow in this environment. Lactobacillus strains have also recently been reported to inactivate C. trachomatis serovar L2 (the LGV pathogen) primarily through maintaining acidity by producing lactic acid and not through the production of hydrogen peroxide. Finally, it is also known that the vaginal microbiota is also impacted by sex hormones although the precise mechanisms of the estrogen/Lactobacillus association are yet to be determined.

These novel insights into the interactions of chlamydiae with their ocular and genital host cells, including host genetic polymorphisms and microbiota, need to figure prominently in the deliberations for designing vaccines to treat C. trachomatis infections (Figure 1) and are opined in the current literature.

The need for chlamydial vaccines

Given the significant impact of C. trachomatis STI and disease worldwide, there continues to be major efforts to control
it. These efforts can be divided into primary, secondary, and tertiary strategies. Primary prevention strategies traditionally have focused on education and changing people’s behavior and attitudes, and they have been successful in increasing condom use; however, they have had negligible impact on other sexual risk practices. Despite widespread knowledge to the contrary, recent data suggests an increase in the risk factors strongly associated with Chlamydia, such as having multiple sexual partners and overlapping partnerships. While education should continue to be a major control strategy, it is unlikely, on its own, to curb the increasing chlamydial STI rates. Secondary prevention involves screening, or at least testing, and then antibiotic treatment of infected individuals. Prevalence data from the UK and the USA, where over 25% of young women are screened annually, unfortunately does not show any reduction in prevalence of Chlamydia over time.

It is not clear why there is no observed reduction in chlamydial STI rates, when mathematical models suggest that if 20% of women under 30 years of age are screened each year, the prevalence should be reduced by at least 40%. One explanation may be the “arrested immunity” hypothesis, which suggests that early treatment actually prevents the development of any “natural” immunity, making these individuals as equally susceptible as naive individuals to subsequent chlamydial exposure. The other weakness of secondary prevention is the perceived effectiveness of the most commonly used antibiotic, azithromycin. While azithromycin is very effective in vitro, in practice, there are many more “treatment failures” than originally thought. Treatment failure rates range from 8% in women to 20% in men with “men who have sex with men” infections being particularly challenging.

The reasons for treatment failure are several, including: 1) true antibiotic resistance – unlikely; 2) failure of the individual to take the antibiotic – more likely for tetracyclines requiring multiple days of administration, but less likely for single-dose azithromycin; 3) different mucosal absorption rates between individuals; and 4) heterotypic resistance in relation to populations of the organism. Tertiary prevention strategies involve targeting the downstream pathology caused by Chlamydia infections, such as PID, infertility, and ectopic pregnancy. The issues here are that these pathologies are often: 1) asymptomatic, especially for women; 2) poorly diagnosed clinically; and 3) multifactorial. Biomarkers that are predictive of disease are, therefore, urgently needed. For all of these reasons, a vaccine for genital tract infections caused by Chlamydia is considered the only feasible solution.

The greatest demand for control of ocular chlamydial infections leading to trachoma is in the developing world. While the SAFE strategy does work, the challenge is to deliver it to the poor regions where it is needed. If a vaccine was available for ocular Chlamydia infections, then this could be utilized in conjunction with the SAFE program and elimination of trachoma may be one step closer.

The immunological challenges
Chlamydial immune responses can be partially protective but also can cause pathology

Protective immune responses to human ocular Chlamydia infections are not well understood; however, what is known is that protective immunity does develop following natural infections and is associated with reduced incidence and duration of infections. Acquisition of natural immunity is also known to occur following human genital tract Chlamydia infections; however, the time to clear infection can take many months with a long-term follow-up study of the natural course of infection in asymptomatic women reporting that approximately 54% of the infections were cleared at 1-year follow-up. The risk for PID is also increased following additional exposures to human genital chlamydial infections. The immune responses to natural human ocular and genital tract infections with Chlamydia thus provide incomplete protection and, in addition, are important in the development of tissue damage and the disease sequelae of blindness and TFI.

The pathogenesis and scarring caused by chlamydial infections has been proposed by Stephens as encompassing two paradigms: the cellular paradigm (epithelial cell layer is sufficient to explain pathology) and the immunological paradigm (in which adaptive immunity explains pathology).

Innate immunity: protection and pathology

Protection from chlamydial ocular and genital infections involves both innate and adaptive immunity following the initial induction of proinflammatory and chemotactroc cytokines such as IL-1α, IL-1β, IL-12, and granulocyte-macrophage colony-stimulating factor from infected epithelial cells. A subsequent upregulation of these chemokines in Chlamydia-infected epithelial cells recruits innate immune cells to the infection site. For example, recruited to the site of infection following induction of CCL2 and CCL5 are macrophages and DCs; neutrophils...
following chemokine (C-X-C motif) ligand 11 and CXCL8 induction; and also NK cells following CXCL9, CXCL10, and CXCL16 induction. Immune-mediated control of chlamydial growth occurs through the contribution of TNF-α and IFN-γ that are released from these immune cells, noting that IL-12 levels were seen to be increased in the cervical monocytes from women infected with *C. trachomatis*. Innate immune pathways are also affected by female sex hormones with the genes for TNF-α, IL-17C, and type I and II IFN receptors (among other cytokine genes) upregulated in progesterone-treated endometrial cells that were infected with *C. trachomatis*. The roles of sex hormones in modulating immune responses in the female genital tract and particularly in relation to chlamydial infections have recently been reviewed.

Ocular pathology is related to the collateral tissue damage that occurs following cytokines triggering the initial immune response to chlamydial infections, and these cytokines are prominent both in active and in scarring trachoma. In children with active trachoma disease, an increase in the neutrophil chemotactic factor CXCL5 has been reported and increased transcript levels of IL-1β have also been associated with chlamydial infection and active trachoma disease. Conjunctival scarring was also found to be strongly associated with a proinflammatory, innate immune response and with increased expression of (among other factors) IL-1β, in a study of 363 trachoma cases. There was also differential expression of MMP-7, -9, -10, and -12 in trachomatis scar tissue, although the expression of many of these genes was also significantly associated with the presence of non-chlamydial bacterial infection.

An increasing number of studies indicate that innate immune responses to *C. trachomatis* infections arising from the epithelium and innate immune cells, along with changes in MMP activity, are likely to also be important in the development of genital tissue damage and oviduct fibrosis in TFI, with MMP-9 expression enhanced in response to chlamydial infections in vitro in human fallopian tube organ cultures and from neutrophils in vivo in a murine chlamydial genital tract model. Fallopian tube scarring associated with *C. trachomatis* genital tract infections may also be the result of increased production of inducible nitric oxide synthase, and mediators such as activins, that are secreted by infected tubal epithelial cells. Host factors associated with scar formation and other fibrotic conditions were also recently reported as being released from epithelial cell monolayers as soon as 1 hour postinfection with *C. trachomatis* serovar E, and these findings have formed the basis of a hypothesis involving multiple feedback loops to explain *Chlamydia*-induced fibrotic scarring of fallopian tubes. These explanations for the pathology induced by chlamydial infections add to previous work showing an involvement of heat shock protein 60 and a proposed model of molecular mimicry which has been discussed extensively by Hafner.

### Adaptive immunity: protection and pathology

Cells of both the humoral (B-cells) and cell-mediated (T-cells) adaptive immune responses are also coordinated with cells of the innate immune response in attempts to protect against chlamydial infections.

Although chlamydiae are intracellular pathogens, and anti-chlamydial antibodies have a major role in protective immunity against chlamydial reinfections, there is increasing evidence that B-cells can protect against infectious agents using antibody-independent mechanisms.

In trachoma, the protective role of antibodies remains to be elucidated, although it is known that the presence or absence of clinical signs of ocular disease is associated with specific antibodies directed against certain chlamydial antigens.

In chlamydial infections of the murine genital tract, it has been reported that a deficiency in B-cells significantly reduced the endogenous *Chlamydia*-specific CD4+ T-cell responses within the local draining iliac lymph nodes and that infection was disseminated in these mice; thus, one possibility is that B-cells may also act as antigen-presenting cells during primary chlamydial infection and participate with the cell-mediated response in bacterial clearance to prevent chlamydial dissemination. Alternative possibilities for the potential roles of B-cells in the draining lymph nodes during a primary *Chlamydia* genital tract infection have also been presented and include an effector function of B-cells to produce cytokines in response to microbial stimulation that may contribute to local defense in the lymph node. Interestingly, data from studies of autoimmune disease and allergic responses also suggest that B-cells can cause pathology by antibody-independent mechanisms, and since 1) the ascension of chlamydiae from the lower to the upper genital tract and, eventually, infection of oviduct epithelium are prerequisites for the development of long-term sequelae and 2) B-cells are involved in bacterial dissemination and cytokine production, perhaps this is also the case for chlamydial-induced genital tract pathology.
Acquired immunity and pathology

Although antibodies are involved in protection against infections caused by *C. trachomatis*, the most potent host defense mechanism is T-cell immunity, particularly involving the release of IFN-γ from host CD4+ and CD8+ T-cells. It also known that IFN-γ-secreting CD4+ T-cells are required to home to the genital mucosa to drive T-cell-mediated protective immunity to chlamydial infections at this site, and that this is achieved by the adhesion receptor integrin α4β1 in the murine genital tract.

However, studies in humans have shown that the tissue damage and scarring sequelae of trachoma and TFI are the result of chronic, immunopathogenic responses to the chlamydial organism and are associated with T-helper (Th) cells such as Th1 and Th17.

In active trachoma, the expression of IL-17A, a proinflammatory cytokine (suggesting Th17 cell activity), is significantly increased, and IL-17A may contribute to fibrosis through epithelial–mesenchymal transition and increased collagen production. In scarring trachoma, there is also an increased expression of MMP-9 (gelatinase-B), a metalloproteinase that is induced not only by NK cells, but also induced and regulated by the Th1 cytokine, IFN-γ. Children with active trachoma also have increased amounts of conjunctival MMP-9.

In the cervical cells collected from *Chlamydia*-infected women, the levels of IL-17 and IL-22 were significantly higher than the levels of these cytokines observed in noninfected women. It has also been reported that the levels of IFN-γ were significantly higher in the cervical washes of women with recurrent chlamydial infection, as compared to the cytokine levels from women with primary chlamydial genital infections. In a recent study of 250 women, 59 of whom had TFI, *C. trachomatis*-infected cells produced significantly higher levels of the IL-12B subunit, p40 than uninfected cells, providing evidence that genetic variation in the IL-12 cytokine family affects *C. trachomatis*-specific immune responses.

Finally, as noted by Mabey et al., “an important difference between ocular and genital infection is that in the eye the damaging sequelae occur at the site of infection, the conjunctival epithelium” while “by contrast in the female genital tract, the major sequelae develop in the fallopian tubes and not at the cervix, which is the site of inoculation”. Nevertheless, the pathology of *C. trachomatis* infection is similar in the eye and genital tract and involves a complex interaction of innate and adaptive immune responses to the organism that influence both protection and pathology in ocular and sexually transmitted chlamydial infections. Vaccine efforts against this intracellular pathogen will need to ensure that any immune responses generated by the ocular and genital vaccines will include the appropriate populations of innate, B, and T-cells that will engender protection and minimize pathology in the host.

Current progress toward the development of a vaccine

Natural infections can induce partial protection, but may also induce immunopathology under some circumstances

Studies in mice and humans (that have been approved by the relevant animal and human ethics committees) have provided results that support the fact that immunity can develop following natural, live chlamydial infections. In the well-studied *Chlamydia muridarum*-mouse model, it is well established that following a live intravaginal infection, immunity does develop, with immunized animals very strongly protected against a subsequent live chlamydial challenge dose. Numerous experiments confirm the fact that major histocompatibility complex (MHC) class II-restricted CD4+ cells and IFN-γ are essential for this protection. Antibodies, particularly neutralizing antibodies, are suspected to play some role in the resolution of the primary infection but a much greater role in the resolution of secondary infections.

Animals that recover from an initial live chlamydial infection shed virtually no viable organisms upon challenge, a level of protection that, as yet, cannot be replicated using recombinant protein vaccines. However, along with protection against infection in these live infection studies comes severely increased immunopathology. A similar effect was observed in the early human trachoma vaccine trials in which inactivated whole chlamydial particles did confer some degree of protection, but unfortunately also led to a level of enhanced immunopathology in some vaccinated individuals. There is also good evidence in human genital tract infections that some women at least do develop a level of natural immunity, with this study showing that 18% of women attending sexually transmitted disease clinics for *C. trachomatis* infections naturally resolved their chlamydial infections prior to any antibiotic treatment. In a follow-up study, up to 22% of women spontaneously resolved their infections, and importantly, these women were significantly less likely to become reinfected, suggesting that their “natural immunity” not only helped them resolve their initial infection, but also protected them from subsequent infections.
If we can define the basis of this human protection, while avoiding any adverse immunopathology, then a vaccine against *C. trachomatis* genital tract infections in women should be possible.

The choice of immunogenic target antigens is rapidly expanding
While early vaccine trials used whole, inactivated chlamydial particles, these will not be acceptable for vaccines as they contain the deleterious antigens that can lead to adverse immunopathology. As a consequence, efforts are now directed primarily toward defined recombinant proteins. While the chlamydial major outer membrane protein (MOMP) dominated early vaccine efforts, the good news is that the era of genome sequencing and reverse genetics has enabled a host of new chlamydial vaccine targets to be investigated. Not surprisingly, surface-exposed membrane proteins such as the MOMP, Omp2, and several Pmps, including PmpD and PmpG, are a major focus.\(^{130}\) However, a wide range of intracellular proteins (NrdB, HtrA, GroEL, Nqr3, MAC–perforin, DnaK, IncA) as well as secreted proteins (Tarp, CPAF, CopB) are also showing promise.\(^{131-135}\) The strategies for identifying key antigens is also improving. The group of Brunham and colleagues used an immunoproteomic screening approach to identify antigens via their presented T-cell peptides in a *C. muridarum*-infected DC mouse model.\(^{136}\) Using this approach, they identified 13 *Chlamydia* peptides derived from eight novel epitopes presented by MHC class II molecules from bone marrow-derived DCs infected with *Chlamydia*. While some of the targets overlapped with previously identified targets (PmpG), others were novel (RplF, FabG, AasF, ClpP-1, Gap, PmpE). Other groups have used antigenic profiling of gene expression libraries to identify *C. trachomatis* proteins both with cell-mediated and with humoral immune responses in women with confirmed *C. trachomatis* genital tract infections, leading to several novel potential vaccine targets.\(^{137}\) When they evaluated selected antigens in the urogenital *C. trachomatis* mouse model, perhaps not surprisingly, the balance between cell-mediated and humoral responses differed markedly between the different antigens.\(^{138}\)

The rapidly expanding chlamydial genetic toolbox should expedite vaccine development
One aspect that has held the *Chlamydia* field back is the lack of a genetic transformation system and the ability to test gene mutants. This is one area that is rapidly changing and it should have a significant impact on vaccine development. The chlamydial genetic transformation conundrum was cracked in 2011 when the group of Clarke et al in Southampton, UK,\(^{139}\) showed that it was possible to develop a plasmid vector that contained both the chlamydial endogenous plasmid and an *Escherichia coli* origin of replication, so that it could shuttle between the two bacterial hosts. By paying careful attention to detail, the genetic transformation of several chlamydial species and strains has now become commonplace.\(^{140,141}\) An important addition around the same time was the use of chemical mutagens, such as ethyl methanesulfonate, to generate libraries of chlamydial mutants.\(^{142-144}\) Unfortunately, these mutants tend to have multiple mutations, but nevertheless, it is possible to screen such libraries to identify clones with your gene of interest disrupted, and subsequently to evaluate such mutants in vitro.\(^{144}\) By complementing the mutated gene with an intact gene on the plasmid shuttle vector, it is now possible to more definitively test the role of key virulence targets. Several of these newly characterized virulence targets are being evaluated as potential vaccine candidates.\(^{80}\) While the chlamydial genetic toolbox has expanded significantly over the past few years, it is likely to continue to become easier and easier to manipulate *Chlamydia*, with both forward and reverse genetics now being possible (eg, targetron approach), and more advanced techniques such as conditional gene silencing and transposon mutagenesis not far off.

Is a live attenuated vaccine the answer?
For the past 10 years or so, virtually all efforts toward vaccine development have focused on the use of recombinant proteins either singly, or occasionally, as mixtures. Despite all efforts, the levels of “protection” obtained have usually been relatively modest. The general model used to evaluate vaccines has been the *C. muridarum*-mouse model, although continued excellent work is progressing using the guinea pig model.\(^{145-147}\) Protection is usually assessed following live genital tract challenge and is measured by any improvements in infection load (either peak of infection, total infection burden, or days to clearance) or reduction in pathology (physical size of the hydrosalpinx). By all measures, the candidate vaccines have not been able to produce a protective response anywhere near close to elimination of the organism, as can be seen following live infection. The effect on pathology reduction, by comparison, has been more impressive. The other major challenge is how the mouse model data will eventually translate into human studies. Caldwell et al\(^{17}\) at the Rocky Mountain Laboratories have attempted to overcome these issues by producing a live, attenuated (plasmid-free) strain of *C. trachomatis* and have evaluated its effectiveness in the monkey model.
results have been by far the most promising for future vaccine development. Kari et al produced a plasmid-deficient strain of *C. trachomatis* and used it to infect the eyes of cynomolgus macaque monkeys. They showed that, compared to the virulent wild-type strain, the attenuated plasmid-free strain resulted in short-lived infections that resolved spontaneously and, importantly, did not produce ocular pathology. They then challenged the vaccinated monkeys with the virulent wild-type strain. All six unvaccinated animals developed severe pathology and extended infections. By comparison, three of the vaccinated animals were solidly protected, with virtually no infection and no adverse pathology. The other three vaccinated animals showed only partial protection, still having significant infections and pathology; thus, two major questions remain. First, will a similar protective effect be seen in humans and will it perhaps be HLA-restricted? Second, will it be possible to get approval from the authorities for a live, attenuated chlamydial vaccine?

**Should we target gastrointestinal infections in addition to genital tract and ocular infections?**

The final question relates to the site of infection. While there are similarities between ocular and urogenital sites of chlamydial infection, the strains of *C. trachomatis* that infect the sites are different (although there are only very minor genomic differences) and hence it is assumed that there will be two separate vaccines produced. One of the reasons is that the vaccines will most likely be used in very different settings, with the ocular chlamahoma vaccine being used in less-developed countries where cost and distribution will be major issues, whereas the genital STI vaccine will be used in more affluent regions. In 2013 though, Yeruya et al reported that animals, and presumably humans, are commonly infected in the gastrointestinal tract with *Chlamydia*. These infections are generally chronic and exist for long periods of time without evidence of clinical disease. They also proposed that females, in particular, may be at risk of autoinoculation. These findings suggest that for any chlamydial vaccine to be fully effective, it may make sense to target the chlamydial organisms present at all tissue sites, further challenging the type of immune response that is required.

**Future directions**

The pipeline for a protective and yet non-immunopathogenic vaccine against *C. trachomatis* ocular and genital tract infections is looking very promising. We now have a genetic toolbox to manipulate the obligate intracellular pathogen, and this augurs well for exciting rapid developments for a *C. trachomatis* vaccine. We continue to have an increased understanding of the potential vaccine targets that will best stimulate protective immunity against the pathogen while avoiding the deleterious adverse immunopathological reactions in the host. Microbiomes of the conjunctival and reproductive tracts are becoming increasingly well studied as are the effects of sex hormones estradiol and progesterone on genital and ocular microbiomes, chlamydial infections, and immune responses. The challenges that the organism presents to the host are becoming known, the host responses are being understood, and the progress toward a protective vaccine for *C. trachomatis* infections is occurring at an exponential rate.

Where to from here? The rapidly expanding genetic toolbox means that we should be able to understand and characterize more key chlamydial virulence factors. We should then be able to identify potential new vaccine targets that might enable improved levels of protection above those levels observed with the current vaccine targets. Live attenuated vaccines are certainly one promising direction. Whether it is the plasmid-free strains or specific gene knockouts (not yet possible but perhaps not far off), the level of protection should be significantly better than what can be achieved using recombinant proteins. However, the key question is, will authorities ever approve a live, attenuated chlamydial vaccine? Will partial protection from a vaccine be sufficient to prevent disease sequelae? In fact, should at least some vaccine efforts be directed at vaccines that only target disease, rather than infection? Will the vaccine be able to be delivered to the disease endemic areas of the world? Yes, there are many questions, but with the rapid developments in our knowledge and technological expertise, we are well on the way to deliver protection without the paradoxical pathology of infections caused by the “cloaked” pathogen, ie, *C. trachomatis*.

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**Disclosure**

The authors report no conflicts of interest in this work.

**References**


