Chlamydia trachomatis Infection, Immunity, and Pregnancy Outcome

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ABSTRACT

Chlamydia trachomatis can ascend from the cervix to the fallopian tubes and survive for long periods of time without causing symptoms. The immune response to infection clears the extracellular organisms but leads to development of a persistent intracellular infection. Repeated cycles of productive infection and persistence eventually induce tubal occlusion and infertility. Persistently infected cells continue to synthesize the chlamydial 60 kD heat shock protein (hsp60). Immunity to conserved regions of hsp60 may result in autoimmunity to human hsp60. Expression of hsp60 by the embryo and decidua during early pregnancy may reactivate hsp60-sensitized lymphocytes, disturb pregnancy-induced immune regulatory mechanisms, and lead to immune rejection of the embryo. Due to this mechanism women with tubal infertility who are sensitized to the human hsp60 may have a decreased probability of successful outcome after undergoing in vitro fertilization and embryo transfer. Infect. Dis. Obstet. Gynecol. 5:128–132, 1997. © 1997 Wiley-Liss, Inc.

KEY WORDS

Chlamydia trachomatis infection; immunity; pregnancy outcome

Due to recent advances in assisted reproduction technology many conditions previously considered to eliminate any chance for fertility are now eminently treatable. In vitro fertilization (IVF) bypasses the requirement for patent fallopian tubes and has rapidly become the treatment of choice for women with occluded or damaged fallopian tubes. However, it is only beginning to be appreciated that the cause of the tubal blockage may also negatively influence postfertilization events. In this article, we focus on the major cause of tubal occlusion in the United States and western Europe, Chlamydia trachomatis infections. We delineate the unique aspects of intracellular chlamydial persistence, the immune-mediated mechanism responsible for C. trachomatis-induced tubal damage and, lastly, the negative consequences these immune events may have for subsequent fertility after IVF and embryo transfer.

C. TRACHOMATIS PERSISTENCE

C. trachomatis can exist in two distinct morphological types. The extracellular elementary body is the infectious morphotype, while the intracellular reticulate body is the replicating form of the organism. C. trachomatis can only divide and reproduce within mammalian cells and is therefore classified as an obligate intracellular microorganism. This ability to reside within mammalian cells allows C. trachomatis to evade detection by the host’s immune system. Furthermore, the organism has developed mechanisms to persist under adverse conditions. Untreated C. trachomatis infections have been shown to persist asymptomatically in the fe-

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male genital tract for years, and this organism has been isolated from fallopian tubes of infertile women with tubal obstruction who had no history of a symptomatic pelvic infection.

There is also substantial evidence that *C. trachomatis* can persist in vivo in an intracellular form that cannot be propagated by in vitro culture. *C. trachomatis* DNA and protein have been identified in culture-negative fallopian tubes from women with postinfectious tubal infertility. The mechanism underlying chlamydial persistence in an unculturable form has been elucidated in in vitro systems; evidence for a similar mechanism existing in vivo is still lacking. Exposure of chlamydial-infected cells to tumor necrosis factor alpha (TNF-α), interferon gamma (IFN-γ), or some antibiotics leads to development of large aberrant reticulate bodies which are incapable of replication. These forms remain viable, however, and when the cytokine or antibiotic is removed the aberrant forms convert back to typical reticulate bodies and the chlamydial life cycle resumes. With each repeated cycle of productive infection and persistence the tubal epithelium becomes increasingly damaged due to cell destruction and the elaboration of pro-inflammatory mediators. Eventually, scarring becomes sufficiently pronounced to occlude the tube. This scheme of tubal damage is outlined in Table 1.

### C. TRACHOMATIS 57 kD HEAT SHOCK PROTEIN AND IMMUNE PATHOGENESIS

In its persistent, unculturable form, *C. trachomatis* synthesizes very little, if any, of its major structural components. However, one chlamydial protein, of 57 kD, continues to be produced and at a rate even higher than under favorable growth conditions. This protein has been shown to be a member of the 60 kD heat shock protein (hsp60) family with extensive amino acid sequence homology to the hsp60s of other bacteria and man. Hsp60 is a highly conserved protein that functions in the intracellular transport and assembly of other proteins. Under conditions of stress or rapid growth its synthesis is greatly enhanced to prevent protein denaturation and incorrect folding and, thereby, maximize the cell’s ability to survive.

The chlamydial hsp60 is a potent inducer of inflammation. In guinea pigs and monkeys previously sensitized to *Chlamydia*, introduction of purified chlamydial hsp60 induced a delayed hypersensitivity response. This suggested that an immune response to hsp60 might be involved in chlamydial disease pathogenesis. In man, several investigations have established associations between immunity to *C. trachomatis* hsp60 and recurrent salpingitis, tubal occlusion, and ectopic pregnancy. Very recent studies have further demonstrated that the presence of antibody to the chlamydial hsp60 was a risk factor for subsequent development of fallopian tube inflammation.

The extensive amino acid sequence homology between the chlamydial and human hsp60 implies that under certain conditions a *C. trachomatis* infection might induce autoimmunity to one’s own hsp60. Cell-mediated immunity to synthetic peptides corresponding to regions of the chlamydial hsp60 that were also present in human hsp60 was not observed in women with a recent chlamydial cervicitis and only rarely in women with an initial episode of salpingitis. It was a frequent finding, however, in women with recurrent chlamydial salpingitis. This suggested that repeated or chronic exposure to the chlamydial hsp60 was required for development of specific immunity to hsp60 epitopes shared between *C. trachomatis* and man. Evidence of humoral immunity to conserved hsp60 epitopes has been obtained for women with *C. trachomatis*-associated ectopic pregnancy and infertile women undergoing IVF.

### TABLE 1. Proposed mechanism of *C. trachomatis*-induced fallopian tube occlusion

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
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<tbody>
<tr>
<td>1</td>
<td>Asymptomatic <em>C. trachomatis</em> infection ascends to the fallopian tubes.</td>
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<tr>
<td>2</td>
<td>Extracellular chlamydial elementary bodies activate the host’s immune response, and IFN-γ, TNF-α, and other pro-inflammatory cytokines are released.</td>
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<tr>
<td>3</td>
<td>The extracellular infection is cleared by the immune system.</td>
</tr>
<tr>
<td>4</td>
<td>IFN-γ and TNF-α suspend intracellular chlamydial reticulate body development but do not affect viability.</td>
</tr>
<tr>
<td>5</td>
<td>When the extracellular infection is cleared, IFN-γ and TNF-α production ceases and reticulate body development resumes.</td>
</tr>
<tr>
<td>6</td>
<td>The reticulate bodies differentiate into elementary bodies, the infected cell is lysed, and other epithelial cells are infected.</td>
</tr>
<tr>
<td>7</td>
<td>The elementary bodies again activate the immune response and the resulting cytokine production induces chlamydial persistence and further damage to tubal tissue.</td>
</tr>
<tr>
<td>8</td>
<td>Repeated cycles of productive infection and persistence eventually lead to fallopian tube occlusion.</td>
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Infectious Diseases in Obstetrics and Gynecology
IMMUNITY TO C. TRACHOMATIS, hsp60, AND PREGNANCY OUTCOME

The relationship between prior immunity to C. trachomatis and pregnancy outcome remains controversial. We, and others, have presented data consistent with increasing susceptibility to subsequent spontaneous abortion after a chlamydial infection. Circulating anti-chlamydial IgG has also been associated with IVF failures in some studies but not others. These studies were all performed prior to the knowledge that immunity to the chlamydial hsp60 induces pro-inflammatory responses. A recent review on the possible influence of immunity to hsp60 and infertility has been published.

Whether prior immunity to the chlamydial hsp60 and to hsp60 epitopes also expressed in human hsp60 affects pregnancy outcome has become a critically important question since many women with tubal factor infertility now seek to become pregnant via IVF. A negative influence of hsp60-mediated immunity on pregnancy is a realistic possibility since the embryo and epithelial cells in the maternal decidua both express hsp60 during the early stages of pregnancy. In addition, a murine hybridoma specific for hsp60 was shown to also react with the surface of human trophoblast, suggesting hsp60 expression by these cells.

In a recent study of 198 women undergoing IVF and embryo transfer, cervical IgA antibodies to the C. trachomatis hsp60 were identified in 41 (20.7%) patients. The occurrence of this antibody was strongly correlated with unsuccessful IVF outcome. Anti-hsp60 IgA was present in 26.3% of women who did not become pregnant after embryo transfer, 33.3% of women with only transient biochemical pregnancies, and only 7.3% of women with live births. There was no relation between hsp60 antibody status and numbers of oocytes retrieved or fertilized. In a subsequent analysis of these same patients utilizing a synthetic peptide to a region of the chlamydial hsp60 that is also expressed in man (amino acids 260–271), it was demonstrated that the presence of IgA antibodies to this peptide also correlated with IVF failure and was associated with only biochemical pregnancies after embryo transfer.

A proposed mechanism of hsp60 immunity-mediated postfertilization pregnancy loss is outlined in Table 2. Expression of hsp60 by the embryo and/or maternal cells could reactivate lymphocytes previously sensitized to conserved hsp60 epitopes as a result of an asymptomatic C. trachomatis infection. This would result in the elaboration of pro-inflammatory cytokines and induction of an inflammatory response that could interfere with implantation and early pregnancy development.
of a localized delayed hypersensitivity response. Interference with immunoregulatory mechanisms necessary for maintenance of the semi-allogeneic embryo would then lead to immune rejection of the embryo. Another possibility is that a persistent, inapparent, intracellular Chlamydia trachomatis infection is still present in some women and becomes reactivated as a result of pregnancy-induced immune alterations. The host’s response to infection would then facilitate immune rejection.

FUTURE PROSPECTS
The recently recognized negative effects of an asymptomatic female upper genital tract chlamydial infection on pregnancy, the capacity to develop a persistent infection, and the induction of immune sensitization to conserved regions of hsp60 highlight even more the necessity for comprehensive Chlamydia trachomatis screening of sexually active women. To preserve fertility, this infection must be detected and effectively treated prior to establishment of persistence in the fallopian tubes. Utilization of DNA amplification technology, such as polymerase chain reaction (PCR)-based assays, should also be mandatory since their sensitivity exceeds that of any other detection method. Recent advances, such as the demonstration that specimens obtained from the entrance to the vagina (introitus) were as sensitive as endocervical specimens for Chlamydia trachomatis detection if PCR was utilized, should greatly expand the numbers of women who can be tested for this organism.

Physicians and other health care professionals must become advocates for state-of-the-art preventative and diagnostic modalities for women and not capitulate to non-professionals interested more in finances than the quality of human lives.

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