Acute placentitis and spontaneous abortion caused by *Chlamydia psittaci* of sheep origin: a histological and ultrastructural study

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**SUMMARY** A sheep farmer's wife who had been assisting with lambing developed an influenza like illness in the 28th week of pregnancy. After five days of malaise she spontaneously delivered a stillborn infant; she became acutely ill during the immediate postpartum period with septicaemic shock, acute renal failure, and disseminated intravascular coagulation. The diagnosis was made by isolation and identification of *Chlamydia psittaci* from the placenta, fetal heart blood, and fetal lung, together with maternal serological evidence. The prominent histological and ultrastructural appearances of the chlamydial placentitis are described.

In the United Kingdom, ovine *Chlamydia psittaci* (group B chlamydiae) is a major cause of abortion in sheep.1 Anecdotal reports2–4 have suggested that the same group B chlamydiae may also be transmitted to pregnant women and cause abortion. Although chlamydial organisms were isolated from the products of human abortion by Schachter,5 no direct causal relation between the agent and the abortion was convincingly shown. The case reported here appears to be the first in which ovine *C psittaci* has been definitely shown to have caused human abortion. The clinical, microbiological, and serological findings have already been reported6 and the purpose of this communication is to record in detail the histological and ultrastructural appearances of human chlamydial placentitis.

**Case report**

A previously healthy 28 year old farmer's wife with a history of one previous normal pregnancy was admitted to the Aberdeen Maternity Hospital in the 28th week of pregnancy after spontaneous delivery of a stillborn baby boy at home, which was followed by postpartum haemorrhage. Some weeks before admission she had assisted with the lambing on her husband's farm, on which some ewes had aborted. For the five days before her own abortion she had had an influenza like illness with fever. Shortly after admission she became acutely shocked and rapidly developed acute renal failure and disseminated intravascular coagulation. She was transferred to the intensive care unit of Aberdeen Royal Infirmary. A chest radiograph showed pulmonary oedema and mild transient pleural effusions with no evidence of pneumonic consolidation. Ultrasonography revealed a bulky uterus, but no evidence of retained products, together with enlargement of the liver and spleen. The patient was treated empirically with intravenous broad spectrum antibiotics, comprising penicillin, cloxacillin, and cefoxitin; she gradually recovered during the next 10 days.

**MICROBIOLOGY**

Micro-organisms were not isolated from cultures of blood, urine, throat swabs, and vaginal swabs. Leptospirosis, listeriosis, brucellosis, toxoplasmosis, Q fever, and mycoplasmal infection were all specifically excluded. *C psittaci* was cultured from the fetal liver, lung, and heart blood and from the placenta. The isolates had the cultural and nutritional characteristics of an ovine rather than an avian strain of *C psittaci*. *C trachomatis* was not isolated from maternal cervical or high vaginal swabs or from fetal tissues and placenta.

Between the fifth and eighth days of the illness...
serological studies on maternal serum samples showed a rise in the titre of complement fixing antibody against chlamydial group antigen prepared from ovine strains of *C. psittaci* from 1/64 to 1/512. In addition, rising titres of neutralising antibody were shown against the chlamydiae which had been isolated from the placenta and fetal tissues. These rising titres were detected against an ovine, strain of *C. psittaci*. Furthermore, immunofluorescence tests showed rising titres of both IgG and IgM antibody to a strain of *C. psittaci* derived from a case of ovine abortion, but only a weak response to a genital (D-K) strain of *C. trachomatis*.

**PATHOLOGY**

Necropsy of the stillborn baby boy was performed about 48 h after death. Externally, the body (weighing 1340 g) was that of a premature, well nourished, non-macerated baby of 28 weeks' gestation, with no
external evidence of disease or malformation. Internally, occasional pinpoint petechial haemorrhages were noted in the thymus (6 g), on the subpleural surfaces of the lungs, and on the epicardium along the coronary sulcus; such haemorrhages are usually interpreted as being non-specific indicators of terminal anoxia. The lungs (right 16 g, left 14 g) were unexpanded. No macroscopic abnormalities were found in the remaining internal organs. The liver (85 g) and spleen (3.5 g) were not enlarged. No macroscopic abnormalities of the fetal, maternal, or cut surfaces. The umbilical cord contained three vessels. It was inserted centrally and also appeared normal.

Fig. 4 Direct immunoperoxidase technique using antibody prepared from serum of sheep experimentally infected with C. psittaci, which binds to the cytoplasmic inclusions (several arrowed) in this placenta tissue section. Direct immunoperoxidase. ×40.

Ovine C. psittaci was cultured from both fetal and placental tissue.

Histopathological examination showed that the fetal viscera were unremarkable apart from some cortical atrophy of the thymus gland, which suggested stress in utero for some time before death. The placenta, however, showed a severe placentitis (Fig. 1) with large numbers of acute inflammatory cells in the intervillous spaces together with inflammation of the decidual bed. Occasional fetal stem vessels showed early perivasculitis, but there was no amnionitis. The vessels of the umbilical cord were not affected. In view of the mother’s rising antibody titre to chlamydial antigen and the isolation of C. psittaci from fetal heart blood and tissues, a careful search was made for histological evidence of chlamydial involvement in the placentitis.

Fig. 5 High power field of the immunoperoxidase stained section clearly defines these chlamydial inclusions (arrowed). Direct immunoperoxidase. Original magnification ×640.

Fig. 6 Ultrastructural appearance of a large cytoplasmic chlamydial inclusion in syncytiotrophoblast. Uranyl acetate and lead citrate. ×2500.
Haematoxylin and eosin stained sections showed, lightly basophilic, granular inclusions with the morphology of Halberstaeder-Prowazek chlamydial inclusions (Fig. 2). In Giemsa stained sections these inclusions appeared dark blue with transmitted light and, when viewed by dark ground microscopy, brilliant bluish white (Fig. 3). Likewise, with methylene blue staining, the placental tissue sections showed these inclusions as sky blue on direct light microscopy and brilliant greenish blue using dark ground microscopy. In addition, the inclusions stained positively with a direct immunoperoxidase technique recently developed for detecting C. psittaci in tissue sections (Figs. 4 and 5). This method does not, however, differentiate between C. psittaci and C. trachomatis. Nevertheless, no glycogen matrix within the cytoplasmic inclusions could be identified in preparations stained with Lugol's iodine, periodic acid Schiff, or Best's carmine, suggesting that they were unlikely to be C. trachomatis organisms.

The ultrastructural appearance of the cytoplasmic inclusions was investigated by transmission electron microscopy (Figs. 6–8) and compared with previously published descriptions of chlamydiae in cells. The inclusions were usually surrounded by membrane and were densely packed with structures which had the appearance of chlamydial infectious elementary bodies, with diameters ranging from 300 to 500 nm. In addition, occasional structures characteristic of chlamydial reticulate and intermediate bodies were present. Each of these chlamydial bodies was enclosed by a rigid envelope representing the cell wall. No glycogen matrix could be found.

**Discussion**

The cultural and serological data, together with the histopathological, immunohistochemical, and ultrastructural findings, indicate that this placenta associated with infection by C. psittaci, which might
well have been acquired from the aborting ewes which the patient had handled during recent lambing. No attempt was made to ascertain the cause of lamb loss at the time, but subsequent investigation showed that C psittaci were cultured from the faeces of the affected ewes. It would therefore seem probable that infection with the organism was the cause of the initial influenza-like illness and the subsequent spontaneous abortion and disseminated intravascular coagulation. The histological pattern of the placenta suggests a haematogenous spread of the infection from the maternal circulation, rather than infection by an ascending cervical route. The primary site of maternal infection or carriage was not identified; there was, however, no clinical or radiological evidence of any pneumonic process.

While C trachomatis is recognised more frequently as a cause of a variety of diseases, especially genital infections, it is pertinent that C psittaci is widespread in this country and economically important to the farming community. Between 1980 and 1981 in more than 1000 farms in the UK lambing losses due to chlamydial infections were as great as 30%, a figure which appears to be increasing. Sporadic cases of human abortion in these areas will occur, and we hope that this report will alert the clinician and pathologist to the possibility that C psittaci may be a cause of human placenta and subsequent abortion. The diagnosis can be made even on formalin fixed material by the relatively simple methods outlined in this report.

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References


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