A CASE OF SEVERE PEMPHİGOİD GESTATİONİS WITH CENTRAL NERVOUS SYSTEM PATHOLOGY İN THE NEWBORN AND REVIEW OF THE LITERATURE

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SUMMARY

A case of severe Pemphigoid Gestationis (PG), persisted postpartum despite high-dose corticosteroid therapy was reported. Small for gestational age, premature birth, and neonatal convulsions with confirmed central nervous system pathology occurred as an adverse fetal outcome. Is the disease as innocent as described earlier?

Key words: dermatose, herpes gestationis, pemphigoid gestationis

INTRODUCTION

Pemphigoid Gestationis (PG) is a rare autoimmune bullous disease that is associated with pregnancy and rarely with trophoblastic malignancy. On the grounds that this pregnancy-associated bullous dermatosis is not caused by any herpes virus and has much in common with the immunologically mediated disorder “Bullous Pemphigoid”; the new term ”Pemphigoid Gestationis(PG)” instead of the old term ”Herpes Gestationis” was proposed. The incidence of PG is estimated between 1 in 10000 and 1 in 50000 pregnancies. We report a case of severe PG, which required hospitalization during several months even after delivery and resulted significant decrease in patient’s quality of life due to high-dose and multiple medications. Besides a premature and intrauterine growth restricted fetus; temporary neonatal convulsions and pathological electroencephalography findings together with vesicular eruptions in early neonatal periods were observed. Present report indicates that most likely because of passive transfer of PG antibody, the disease may affect central nervous system of the newborn, in addition to the skin.
CASE

A 42-year-old woman (gravida six, parity three, abortion two) at 20 weeks’ gestation presented with extremely pruritic, blistering skin eruption localized around the umbilicus (Figure 1) and vulva. In her obstetrical history, there were two spontaneous abortions, two vaginal births and one preterm birth in 34th weeks in which similar but mild eruptions occurred at the beginning of the third trimester. There were no other risk factors to explain the preterm birth.

Lesions were polymorphic with urticaria-like erythematous and edematous papules, plaques, tense vesicles and bullae. Multiple skin biopsies were obtained and PG was confirmed as histopathological diagnosis after a sample processed for direct immunofluorescence. Subepidermal edema, vesicle, and perivascular infiltrate of lymphocytes, histiocytes and eosinophils were reported as histopathologic features (Figure 2).

Pruritus was quite severe. No relief obtained from topical corticosteroids and antihistamines. Despite administration of the oral prednisone, a generalized bullous reaction ensued that spares the face, mucous membranes, and palms and soles (Figure 3, 4).

Fluocortolone orally started and doses up to 60 mg daily administered. Doses regulated according to exacerbations and remissions throughout pregnancy. She delivered a viable infant at 35 weeks’ gestation following preterm premature rupture of the membranes. She continued to have recurrent severe symptoms during the peripartum and postpartum period. The patient required hospitalization in dermatology department and high-dose corticosteroid treatment applied, until remission observed in postpartum 12th weeks.

Antenatally, when presented, in 20th weeks’ gestation, there was no problem in both biometry and anatomic survey of the fetus. The woman despite advanced maternal age has not accepted amniocentesis. In the
ultrasound examination performed in 29th weeks, estimated fetal weight was in 10 percentiles. At 34 weeks’ gestation, estimated fetal weight was between 5 and 10 percentiles. Following preterm premature rupture of the membranes, she delivered a viable infant at 35 weeks with cesarean electively. The male neonate was 2120 g (10. percentile) with 1st minute and 5th minute apgar scores of 7 and 8, respectively. Clinical confirmation of prematurity and small for gestational age diagnoses were made.

At postpartum 2nd day papulo-vesicular eruptions similar with mothers lesions observed in the newborn on the forehead and facial region but they disappeared in a few days without any treatment. Because of lack of the parents consent, biopsy was not taken from eruptions. At the sixth day of life myoclonic jerks developed. Electroencephalography (EEG) examination revealed partial epileptic activity on both hemispheres, but no abnormal findings were observed in the cranial ultrasonography and magnetic resonance imaging. Following phenobarbital therapy, convulsions disappeared and control EEG was normal at postnatal 30th day when anticonvulsant therapy was discontinued.

DISCUSSION

PG is an autoimmune organ-specific blistering skin disease(1). It occurs only in the presence of placental tissue. The incidence was reported about 1 in 50,000 pregnancies(1). In our case, disease course was so severe that patient’s quality of life was severely impaired despite multiple and high dose medication. In addition to the estimated growth restriction and prematurity, convulsions of undetermined etiology were observed in the early neonatal period, which we couldn’t find such an adverse fetal outcome in the literature.

The disease manifests itself most commonly during the second or third trimester with an average onset at 21 weeks, although in 20% of patients initial onset occurs in the immediate postpartum period(2). Our patient presented at 20 weeks’ gestation. In her fourth pregnancy, same lesions attributable to the same disease did so later and less severe and skip pregnancy occurred for which there has been no satisfactory explanation could be done in the literature(3). Although our patient had advanced maternal age, no such association was described.

We could easily make a presumptive diagnosis of PG in the case with a typical distribution of vesicles and bulla. The typical blistering eruption had a herpetiform appearance, but the vesicles were not clustered and were located more peripheral than herpes. Skin biopsy, including a sample processed for direct immunofluorescence, confirmed the diagnosis. This confirmation distinguished PG from polymorphic eruption of pregnancy, which does not recur with subsequent pregnancies.

In the cases reported, the disease runs a variable clinical course. A flare at the time of delivery is a typical feature, seen in 75% of cases and in most patients; the disease spontaneously regresses in the postpartum period. Many patients with PG transiently improve during the last 6 to 8 weeks of pregnancy only to experience an uncomfortable flare of their disease within 24 to 48 hours after delivery(2). Nevertheless, in the present case although the flare had occurred peripartum, the disease persisted during the course of the pregnancy and the postpartum period. The bullous, extremely pruritic lesions persisted 12 weeks. We could not find association between severity of the disease and the perinatal outcomes in the previous reports. Moreover, the effects of breast-feeding and prolactin on prolonging the duration of PG deserve further investigation(5).
Although doses up to 180 mg of prednisone daily have been reported, most patients respond to lower doses (20-40 mg daily). Orally administered prednisone did not bring relief promptly and inhibit formation of new lesions in our case and with improvement of clinical symptoms following administration of fluocortolone and no new bulla formation, the dose then had tapered. Refractory cases during the postpartum period had been reported that respond to adjunctive cyclophosphamide, pyridoxine, gold or methotrexate\(^{13, 14, \text{and } 15}\). Intravenous immunglobuline combined with cyclosporine, plasmapheresis, chemical oophorectomy with goserelin, and ritodrine have been exceptionally used in chronic PG with some success. However, because the response to these agents has been variable and their safety questionable, we have not administered additional therapy to the high-dose corticosteroids. Because in the postpartum period patient had experienced a flare of pemphigoid gestationis requiring continued steroids, prednisone doses of 20 mg/day still was allowed safe breast-feeding.

Prior to the development of immunologic techniques that permitted an accurate diagnosis, PG was believed to have minimal adverse effects on the fetus\(^{16}\). In 1969, Kolodny reported no increase in preterm births, stillbirths, or abortions\(^{17}\). However, Lawley et al. subsequently reviewed 40 immunologically proven cases of PG and documented serious fetal morbidity and mortality: preterm births occurred at a rate of 22 percent and three women suffered stillbirths\(^{18}\). These authors suggested that high levels of circulating antibasement membrane antibody and peripheral eosinophilia caused the increased fetal risk. The pathophysiology of fetal complications was thought to be due to mild placental insufficiency, because placental antigens may be targeted by the immune responses that target the skin. Shornick and Black described 74 patients with PG confirmed by immunofluorescence studies\(^{5}\). They found an increased frequency of fetal growth restriction and prematurity, but not stillbirth or miscarriage\(^{5}\). No evidence currently exists to determine whether systemic therapy for PG reduces the risk of fetal prematurity. In the present report, all adverse perinatal outcomes reported that had association with the disease had been observed; premature birth, small for gestational age and neonatal transient eruptions. Nevertheless, it is the first case in the literature that central nervous system pathology, which has been confirmed with EEG findings, had reported. These convulsions may not have any association with the disease but, they were transient and no other metabolic or the other etiologies have been found so it can be suggested that this clinic finding may be result of PG antibody, which crosses placenta passively. In addition to the placenta and skin, same antibodies may have cross-reactions with neuronal tissue, too. Neonatal vesicles occur in 10% of cases, most likely because of passive transfer of PG antibody\(^{15}\). The eruption is usually mild and self-limited; like our case but the lesions may become super infected as the immune system of the neonate is not fully developed. Still once the diagnosis of PG is established, the pregnancy should be considered high risk. In addition, close surveillance is required for the possible adverse fetal effects. The patient had to be counseled about the recurrence risk of pregnancies in the future.

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Düzeltme

Uzmanlık Sonrası Eğitim ve Güncel Gelişmeler Dergisi 2005 yılı Cilt 2, Sayı 2, 103-106. sayfalarda yayınlanan makalede yazarlardan Dr. Aydan Biri’nin adı baskıda görülmemiştir. Düzeltilmiş yazar adı ve sırası aşağıdaki gibidir.

RALOXIFENE KULLANIMININ CİNSEL FONKSİYON ÜZERİNDE ETKİSİ.

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Uzmanlık Sonrası Eğitim ve Güncel Gelişmeler Dergisi 2005 yılı Cilt 2, Sayı 2, 103-106.