Pemphigoid gestationis (PG) is an uncommon autoimmune bullous disease that almost exclusively presents during pregnancy. Patients typically present with a diffuse blistering and intensely pruritic eruption that begins periumbilically and spreads to involve the rest of the body. Direct immunofluorescence demonstrating C3 in a linear pattern along the dermoepidermal junction confirms the diagnosis of PG. Corticosteroids remain the choice of therapy and early intervention is essential because of possible adverse effects of PG on the fetus. We report a case of PG and review the literature.

Case Report

A 27-year-old white woman (gravida 2, para 1) presented to our clinic at 32 weeks' gestation with a report of a severely pruritic eruption of 6 weeks' duration. She had been seen by her primary care physician a few weeks prior and was diagnosed with rhus dermatitis. She was treated with a 12-day prednisone taper of 30 mg for 3 days, 20 mg for 3 days, 10 mg for 3 days, and 5 mg for 3 days, which initially provided some relief. However, the eruption persisted and intense pruritus returned. Subsequently, she was referred to our clinic.

Physical examination revealed annular, urticarial, vesicular plaques covering her entire back, forearms, flank, chest, abdomen, and legs (Figure 1). Yellow crusting, bullae, and vesicles were present throughout the plaques (Figure 2). An initial clinical assessment of pemphigoid gestationis (PG) was made. Punch biopsy of the left forearm was taken for hematoxylin and eosin staining and a perilesional site was sampled for direct immunofluorescence. In the interim, we started the patient on triamcinolone acetonide cream 0.1%, a 30-day prednisone taper starting at 60 mg daily for 5 days and decreasing 10 mg daily every 5 days until a dosage of 10 mg daily was achieved, and cyproheptadine hydrochloride 4 mg 3 times daily. In addition, we gave her a 7-day course of erythromycin 500 mg twice daily and instructed her to apply mupirocin ointment to secondarily impetiginized areas.

Punch biopsy results showed subepidermal vesicles and papillary dermal edema (Figure 3). Scattered lymphocytes, histiocytes, and eosinophils were located in the dermis in a perivascular distribution (Figure 4). Direct immunofluorescence staining was negative for IgG, IgM, and IgA, but C3 was present in a linear fashion along the dermoepidermal junction, confirming the diagnosis of PG.

The patient responded well within 2 weeks of starting the prednisone taper and her lesions regressed. She delivered at 38 weeks' gestation and did not flare at the time of delivery. The infant was healthy and had no evidence of skin lesions or adrenal suppression at birth. However, at 1-week postpartum and at the onset of her first menses, the patient experienced a resurgence of her PG. Again, the patient was started on the same prednisone taper used during her pregnancy and her lesions completely resolved during both relapses. At the time of this publication, she had no further recurrence of PG.

Comment

Pemphigoid gestationis is a rare dermatosis that almost exclusively presents during pregnancy. It also has been associated with trophoblastic diseases such as hydatidiform mole or choriocarcinoma. Pemphigoid gestationis, known historically as herpes gestationis, has no known viral etiology or association with the
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herpes simplex virus. The term herpes gestationis was coined in recognition of the herpetiform lesions classically seen in PG. Interestingly, PG shares clinical, histological, and immunological characteristics with bullous pemphigoid, suggesting that the 2 diseases may be related.

Pemphigoid gestationis is a rare disease with a wide range of incidences reported. Several reports estimate the incidence of PG at 1 in 50,000 to 60,000 pregnancies. Higher incidences, up to 1 in 3000, also have been recorded, but these numbers have been criticized for accuracy secondary to referral bias and the inclusion of cases that fail to meet immunologic criteria for PG. In a study performed by Zurn et al, the incidence was estimated at 1 in 7000, which has led many investigators to speculate that cases of PG may be underreported. There is some predilection for race with considerably more cases documented in white individuals than other races.

Clinical Characteristics—The clinical presentation of PG is polymorphic and evolutionary in nature. Typically, patients initially present with erythematous papules, plaques, or targetoid lesions reminiscent of erythema multiforme. Over time, small vesicles and bullae appear on normal skin or on top of urticarial

Figure 1. Large urticarial plaques involving the abdomen and right flank, extending to the chest and back.

Figure 2. Urticarial plaques with multiple bullae and vesicle formation on the right anterior forearm.

Figure 3. Punch biopsy results showed subepidermal vesicles and papillary dermal edema (H&E, original magnification ×40).

Figure 4. Perivascular inflammatory infiltrate consisting of lymphocytes and eosinophils (H&E, original magnification ×400).
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plagues, reminiscent of urticarial bullous pemphigoid. There are some cases of PG in which blister formation never occurs.9,13-15 Pemphigoid gestationis lesions characteristically begin periumbilically and then spread outward to include the buttocks, trunk, and extremities, sparing the mucous membranes and face.14 Nonetheless, there have been case reports of PG with primary involvement of the oral mucosa.16

Intense pruritus is the consistent hallmark of PG, while the clinical course is variable. Pemphigoid gestationis typically presents during the second or third trimester with a mean onset of 21 weeks but may develop in the first trimester, the time of delivery (75% of cases),17,18 the first postpartum month,13-15 and with the onset of menses or the use of oral contraceptives in the postpartum period.13,14,19 In 20% of cases, the initial presentation occurs in the immediate postpartum period.5 Some studies have shown that breastfeeding may decrease the duration of the disease,9,19 with lesions for breastfeeding and bottle-feeding mothers lasting on average 5 weeks and 24 weeks, respectively.9

Most commonly, PG resolves spontaneously without scarring within weeks to months after delivery.20 Pemphigoid gestationis can arise in subsequent pregnancies and may have a tendency to occur earlier in gestation and be more severe at times. Mothers affected by PG do not always experience the disease in succeeding pregnancies. In fact, skip pregnancies have been noted in the literature, estimated at 5% to 8%.15,21 In multiparous patients with PG, the postpartum duration of PG may be extensive and, as shown in one unusual case, may last for several years after delivery.22

Pemphigoid gestationis also has been linked with an increased risk for autoimmune diseases, especially Graves disease.23 An estimated 10% of patients with PG also have a concurrent autoimmune disease.24 Cases of alopecia areata, vitiligo, and ulcerative colitis also have been reported in patients with PG.23

Differential diagnosis of PG includes pruritic urticarial papules and plaques of pregnancy (PUPPP) and bullous pemphigoid. Although pathologic analysis is the only way to clearly differentiate between these diagnoses, clinical features may lend support to one diagnosis over the other. Pruritic urticarial papules and plaques of pregnancy, the most common dermatosis of pregnancy, typically occurs in primigravid patients in the third trimester and lesions tend to lack bullae and appear more focally on the lower abdomen and thighs instead of periumbilically. In addition, the lesions of PUPPP characteristically appear in the striae. Bullous pemphigoid primarily affects elderly patients and both sexes equally. Other potential diagnoses such as erythema multiforme and dermatitis herpetiformis also resemble PG in presentation and should be included on the list of potential diagnoses.

Histopathology, Immunofluorescence, and Pathophysiology—The histologic findings in PG parallel bullous pemphigoid and consequently fail to distinguish it from this autoimmune subepidermal disease. In PG, there is a subepidermal bulla with an underlying inflammatory infiltrate consisting of histiocytes, lymphocytes, and eosinophils in the dermis. Blisters, if present, are found subepidermally. Direct immunofluorescence shows deposition of C3 in a linear pattern along the basement membrane and is considered pathognomonic for diagnosis of PG.17 In approximately 25% to 30% of cases, IgG deposition also can be found along the basement membrane.25

In addition to histologic analysis and direct immunofluorescence to aid in the diagnosis of PG, Powell et al26 demonstrated that a serologic test measuring the major immunoreactive portion of the NC16A domain of BP180 antigen can be used to verify the diagnosis of PG and differentiate it from PUPPP. Sitaru et al27 also supported the use of NC16A levels to measure the presence of autoantibodies and assist in the diagnosis of PG. Furthermore, because NC16A levels correlate with disease activity, they can be used to follow disease progression and consequently may impact the course of treatment.27

Pemphigoid gestationis is considered an autoimmune blistering disease in which autoantibodies develop against the basement membrane zone, the same mechanism believed to be responsible for bullous pemphigoid.28 Approximately 20% of patients with PG have a complement-fixing protein in their serum, known as the herpes gestationis factor, a revelation that was further defined by Jordon et al29 and Katz et al30 as complement-fixing anti-basement membrane zone IgG. Kelly et al31 identified IgG1 as the major subclass of PG anti-basement membrane zone autoantibodies. In contrast, Patton et al31 found that the predominant IgG subclass in PG is IgG4, the major subclass in bullous pemphigoid. Regardless, it is accepted that the PG autoantibodies recognize the BP180 antigen, also known as collagen type XVIII and BPAG2,32-38 and activate complement via the classical pathway.37 The bullous pemphigoid antigen is located in the basement membrane of normal skin and in the basement membrane zone of amniotic epithelium of both the placenta and umbilical cord.39,40 An allogenic autoimmune response, possibly facilitated by the abnormal expression of class II major histocompatibility complex antigens in the placenta, may explain the pathophysiologic phenomenon of PG.1

There appears to be a genetic basis for the development of PG, particularly with HLA-DR3 and DR4.
As many as 61% to 85% of patients with PG have HLA-DR3, 50% have HLA-DR4, and 43% to 45% have both HLA-DR3 and DR-5. In addition, 50% of husbands of PG patients have HLA-DR2. The paternal role in the pathogenesis of PG also has been studied. In a study by Holmes et al, a change in paternal role in the pathogenesis of PG also has been implied that the 2 events may be related and that the development of PG may depend on antigens provided by the father. In contrast, Jenkins et al asserted that their study showed no correlation between partner changes and the occurrence of PG; skip pregnancies could not be explained by partner changes either. The hypothesis that PG results from an allogeneic response to placental basement membrane zones raises the issue of the paternal role in the pathogenesis of PG because the placenta primarily is of paternal origin.

Fetal Risk—One of the most important considerations in approaching the patient with PG is the effect the disease may or may not have on the fetus. Patients with PG should be identified as a high-risk pregnancy and be followed accordingly. Most studies show that infants of PG mothers tend to be small for gestational age and have an increased likelihood to be born premature, findings consistent with low-grade placental insufficiency. Furthermore, a controversial study performed by Lawley et al claimed that PG was associated with an increase in fetal death. However, later studies have disputed this claim and did not find an increase in fetal morbidity or mortality in PG. Some infants of PG mothers, approximately 5% to 10%, have a transient subepidermal blistering eruption that resolves on its own with no known sequelae. This finding is most likely due to transplacental passage of antibodies from the mother to the baby. After delivery, infants of mothers treated with corticosteroids should be monitored for signs of adrenal insufficiency.

Treatment—Pemphigoid gestationis usually can be managed with the administration of topical or oral corticosteroids or a combination thereof. Mild lesions can be managed with topical steroids, while more severe cases require the use of systemic steroids. Most cases of PG respond well to low dosages (20–60 mg daily), but the need for higher dosages up to 180 mg daily have been reported. Multiple steroid tapers may be necessary for treatment because the eruption may flare right after delivery or with the onset of menses. It does not appear that treatment with systemic steroids adversely affects the fetus. Effective management is gauged by the disappearance of old lesions and the lack of new lesions erupting. Antipruritic drugs such as diphenhydramine, hydroxyzine, loratadine, or cetirizine are effective to help alleviate pruritus. Due to excoriations as well as bullae and vesicles breaking open, patients should be monitored for secondary infection and treated accordingly.

A mixture of other agents have been used to treat PG in the postpartum period. There has been variable success noted with the use of dapsone, sulfapyridine, gold, cyclophosphamide, pyridoxine, and methotrexate. Intravenous immunoglobulin in conjunction with cyclosporine also has been used in resistant cases. There also have been some promising results with tetracycline in postpartum PG. In addition, plasmapheresis has been applied in refractory cases with some clinical improvement.

Conclusion
In summary, PG is an autoimmune disease that occurs almost exclusively during pregnancy. Its clinical course is variable, but eruptions typically respond to steroid therapy. It is important to diagnose and treat PG early, not only to provide symptomatic relief to patients but to prepare for possible adverse outcomes on the fetus.

REFERENCES