Autoimmune and inflammatory diseases present with a wide variety of clinical manifestations. Often the cutaneous signs and symptoms do not accurately reflect the degree of immune activation and tissue damage. Adjuvant tests, such as those for serum C-reactive protein (CRP) level and erythrocyte sedimentation rate, can help clinicians assess the degree of inflammation and tailor management. During the active phases of certain inflammatory diseases, leukocyte-mediated damage of the blood vessels can occur, resulting in vasculitis. Damage to blood vessels can also lead to activation of the coagulation cascade, thrombus formation, and D-dimer release into the bloodstream. It follows that D-dimer levels might be a useful marker to track inflammation and blood vessel damage.

D-dimers are small protein fragments generated by fibrinolysis of a thrombus or blood clot. D-dimer assays are often used in the diagnosis of deep venous thrombosis or pulmonary embolism and have been used to predict the likelihood of recurrent venous thromboembolism. Previous reports have indicated that there is an increased risk of venous thromboembolic events in patients with vasculitis. Furthermore, D-dimer levels have been shown to be elevated in patients with vasculitis such as Henoch-Schönlein purpura, Kawasaki disease, and eosinophilic granulomatosis with polyangiitis (Churg-Strauss syndrome). The wider application of D-dimer levels as a marker of inflammation has been suggested by work in nonvasculitic conditions such as bullous pemphigoid and chronic urticaria.

Here for the first time, to our knowledge, we demonstrate disease activity correlated with D-dimer levels in a patient with cutaneous polyarteritis nodosa (PAN) and another with recurrent atypical urticaria. We suggest that measuring D-dimer levels in a variety of inflammatory conditions might be of clinical use in assaying disease activity, assessing response to treatment, and in potentially stratifying patient risk of venous thromboembolic events.

Report of Cases

We present 2 cases of cutaneous inflammation showing a correlation of disease activity with D-dimer levels.

Case 1

The first case is of a woman in her 50s who presented to dermatology clinic with a history of livedoid erythema (Figure 1A) and recurrent leg ulcers (Figure 1B). The patient first developed skin problems 6 years prior after hiking when she noticed blue skin discoloration on her legs. This discoloration extended to her buttocks, thighs, and arms during a period of...
Four years before presentation, she developed a foot drop and associated numbness. She next developed superficial ulcers of the legs that resolved with use of systemic and topical nifedipine and appropriate dressings. One year before presentation, the fourth and fifth toes of the right foot became ischemic and this resolved with dalteparin sodium anticoagulation therapy. The patient was observed by thrombosis clinic staff with a presumptive diagnosis of livedoid vasculopathy and D-dimer levels were checked from 2009 onward (Figure 2). When no active disease was present, her D-Dimer Levels and Cutaneous Disease Activity
baseline plasma D-dimer levels were in the range of 400 to 600 μg/mL (reference value, <500 μg/mL [to convert to micromoles per liter, multiply by 5.476]). At presentation, the patient had experienced a 2-month flare of her condition with the development of numerous ulcers of the legs and a marked rise in D-dimer levels to a maximum of 1839 μg/mL. A biopsy performed during this flare showed vasculitis involving deep dermal vessels. Other investigations including liver function, renal function, hepatitis serologic analysis, cryoglobulins, antineutrophil cytoplasmic antibodies, and antiphospholipid antibodies had negative results. Given the lack of systemic manifestations (kidney, heart, liver) usually seen in systemic PAN, a diagnosis of cutaneous PAN was made on the basis of the clinical presentation and biopsy result. The flare of the cutaneous PAN subsided with intravenous IgG and systemic steroid therapy and her D-dimer levels returned to her normal range. She was subsequently treated with azathioprine and has not had any further flares of her disease (Figure 1C and 1D).

Case 2
The second case involves a man in his 20s who presented to the emergency department with right foot erythema and swelling that was initially thought to be a deep venous thrombosis. His plasma D-dimer level was more than 4000 μg/mL, but there was no radiological evidence of thrombosis. On further review, a history of recurrent periodic episodes of swellings with associated erythema dating back 2 to 3 years was obtained. His lesions would affect various parts of the body including the torso, extremities, and genitals and lasted for hours to days (Figure 3). The patient was subsequently investigated by internal medicine staff for angioedema. Laboratory tests including those for levels of immune complexes, antinuclear antibodies, functional C1 inhibitor, C3, and C4 all had results within normal ranges. The D-dimer test was repeated a few weeks after the initial presentation when the erythema and swelling had resolved, and the level was found to be 501 μg/mL. Over the course of the next year, the patient had 2 more episodes of swelling with erythema, accompanied by elevations in D-dimer levels (with values of 2545 and 1771 μg/mL), and he had normal D-dimer levels during quiescent intervals (Figure 4). Interestingly, the serum CRP levels changed little during the flares of his disease. The patient is now being observed by the dermatology service with a working diagnosis of atypical recurrent urticaria based on clinical evidence of episodes of sustained swelling that are...
unresponsive to antihistamines. Skin biopsy revealed pathologically normal skin, without the neutrophil infiltrates noted in urticarial vasculitis or neutrophilic urticaria. Treatment with dapsone and colchicine has abolished further swelling episodes during 1 year of follow-up.

Discussion

C-reactive protein is a well-established biochemical marker of inflammation. Previous studies have suggested a link between inflammation, elevation of serum CRP levels, vascular damage, and elevation in plasma D-dimer levels. Herein we present 2 patients who had negative results on a CRP test and positive results on a D-dimer test during active disease, indicating that markers of acute inflammation may be less sensitive than markers of activation of coagulation. Medication use may play a role in these differences. Particularly, many of the patients included in the study of Nikpour et al were receiving anti-inflammatory medications that have been shown to suppress CRP levels. This would provide an additional reason to use D-dimer levels as a biochemical marker of disease activity because they are less likely to be directly affected by immunosuppressive medications and as such more directly reflect vascular inflammation-mediated activation of the coagulation cascade and fibrinolysis.

Although CRP is the more widely used biochemical marker of inflammation, there have been a few case reports and case series that correlate D-dimer levels with disease activity and risk of venous thromboembolic events. Perhaps the most studied is the association between systemic lupus erythematosus and the incidence of thrombosis. Arterial and venous thromboembolism has a prevalence of 10% in patients with systemic lupus erythematosus and is believed to be related to hypercoagulability, premature atherosclerosis, and increased incidence of vasculitis. Vasculitis seems to be a common pathophysiologic link between the other reports of inflammatory conditions and elevated D-dimer levels. These reports include Henoch-Schönlein purpura, Kawasaki disease, eosinophilic granulomatosis with polyangiitis (Churg-Strauss syndrome), and Behçet syndrome. In addition, several groups have shown that the coagulation cascade is activated in patients with chronic urticaria and atypical urticaria and that D-dimer levels were elevated in these patients and associated with disease severity. One study examined the activation of the coagulation cascade in patients with bullous pemphigoid and found a correlation between disease activity and increased D-dimer levels. When patients with bullous pemphigoid were treated and clinical manifestations of their disease subsided, their D-dimer levels decreased to normal. These findings from a wide variety of inflammatory conditions support the findings of this study and the use of D-dimer as a potential biochemical marker of disease activity.

Conclusions

We report, to our knowledge, the first documented case of cutaneous PAN that shows a direct correlation between disease activity and D-dimer levels. Furthermore, we confirm the previously reported association between recurrent atypical urticaria and elevated D-dimer levels. In both cases, CRP levels did not correlate with clinical disease activity. We propose that D-dimer measurements may play a role as a biochemical marker of inflammatory skin disease activity. Elevated D-dimer levels in patients with vasculocentric and/or vasculopathic inflammation suggest that vascular endothelial damage may be occurring and that these patients may be at higher risk of venous thromboembolic events. Whether elevated D-dimer levels in patients with inflammatory skin disease identify patients with increased risk of thromboembolism remains to be determined. Clinicians should be aware of the potential utility of D-dimer levels to evaluate disease severity, track patient response to treatment, and assess the need for anticoagulation therapy.
Solving the Mystery of Jimmy’s Red Sweat

Walter H. C. Burgdorf, MD; Leonard J. Hoenig, MD

In the 1950s, Harry Hurley and Walter Shelley, distinguished dermatologists at the University of Pennsylvania, devoted considerable attention to the apocrine gland. Their curiosity extended beyond human subjects, and their most fascinating subject was a rotund inhabitant of the Philadelphia Zoo, Jimmy the Hippopotamus.

Since ancient times, travelers to the Nile Valley in Africa had reported that hippopotamuses “sweat blood.” Hurley and Shelley noted that Jimmy, when annoyed, excreted a “bloody” red sweat, especially on his head and shoulders. Since he was not fond of his handlers, their mere appearance elicited this response. The dermatologists were not foolhardy enough to attempt a biopsy on Jimmy and were not allowed to administer drugs to stimulate or diminish sweating. However, the slimy, turbid nature of the red discharge, as well as its association with emotional stimuli, convinced them it was apocrine in nature.1 In his autobiography many years later, Shelley wrote, “Just contemplating any experiments on Jimmy gave our axillary a nice wash of apocrine and eccrine sweat.”2

Figure. A Poster Titled “Visit the Zoo—Philadelphia” Showing a Hippopotamus

Jimmy was born wild in Africa in 1934 and arrived at the Philadelphia Zoo, Philadelphia, Pennsylvania, in 1936. He was the subject of a famous Work Projects Administration (WPA) poster (Figure). His partner, Submarine, arrived in Philadelphia in 1950 from the Brookfield Zoo, the Chicago suburb of Brookfield, Illinois. The couple had 12 children before Jimmy died in 1977. In Submarine’s obituary in 1990, her long-term handler discussed the problems of keeping her weight below 4000 pounds.

Japanese investigators have studied the red apocrine sweat in hippos.3 They noted that the sweat was initially clear but turned red and then brown as the pigments polymerized. They collected the sweat by wiping a hippo’s face and back with gauze and extracted 2 aromatic acids derived from tyrosine precursors—red hippodisocadic acid and orange norhippodisocadic acid. These acids have antiseptic properties and also serve as sunscreens by scattering UV light. While there are still no biopsy studies clearly identifying the responsible glands as apocrine, this seems most likely.

Shelley’s insatiable curiosity led him to many interesting observations, including Jimmy’s red sweat. He was very pleased many years later when another, more courageous, group obtained sweat samples. Today, researchers hope that further study of the hippo’s red sweat may lead to development of more potent sunscreens. In the meantime, the hippopotamus remains one of our favorite animals, bringing smiles of delight to all who behold this behemoth, including the noted American poet Ogden Nash (1902-1971) who wrote:

The Hippopotamus
Behold the hippopotamus!
We laugh at how he looks to us,
And yet in moments dank and grim,
I wonder how we look to him.

Peace, peace, thou hippopotamus!
We really look all right to us,
As you no doubt delight the eye
Of other hippopotami.

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