Leukocytoclastic vasculitis (LCV), also termed hypersensitivity vasculitis, is a small-vessel vasculitis with a reported incidence rate of about 30 cases per million people per year and is thought to affect men and women in equal numbers. The skin is the organ most commonly involved in LCV. Typical presentation is a painful, burning rash predominantly in the lower extremities (Figure 1), with up to one-third of patients presenting with trunk and upper extremity involvement. The most common skin manifestation of LCV is palpable purpura. Other skin manifestations include maculopapular rash, bullae, papules, plaques, nodules, ulcers, and livedo reticularis. Patients with LCV may also present with arthralgias or arthritis involving the knees or ankles. The differential diagnosis for LCV includes drug reaction with eosinophilia and systemic symptoms (DRESS) syndrome, amyloidosis, antiphospholipid syndrome, atrial myxoma, Behcet disease, Churg-Strauss syndrome, granulomatosis with polyangiitis, Henoch-Schönlein Purpura, urticarial vasculitis, immune thrombocytopenic purpura, and meningococcemia. Multiple etiologic factors including drugs, infections, foods, autoimmune diseases, collagen vascular diseases, and malignancies have been associated with LCV. Although the exact pathogenic mechanism remains to be elucidated, circulating immune complexes are believed to be involved.

In the evaluation of patients with LCV, laboratory tests including a complete blood count, erythrocyte sedimentation rate, biochemistry profile with liver and renal function, and urinalysis are useful in excluding other vasculitides, determining the presence of systemic disease, and identifying an associated disorder, which can provide prognostic information. Patients with suspected LCV presenting to the Emergency Department may require parenteral analgesics for pain control, and those patients not requiring hospitalization should be referred to a dermatologist upon discharge. Diagnosis of LCV is confirmed on histologic examination of a biopsy from the affected area that demonstrates perivascular and vascular leukocytic infiltrates along with fibrinoid necrosis. Mild, skin-limited LCV does not require treatment apart from rest, elevation of the legs, ice packs to the affected area, and removal or treatment of the inciting cause. Presence of arthralgia or arthritis requires use of nonsteroidal antiinflammatory drugs, or a short course of oral steroids (eg, prednisone or methylprednisolone) at a dose of 1 mg/kg/day for 4 weeks followed by a steroid taper. Most patients respond to such treatment. A single-pulse dose of intravenous corticosteroids (eg, methylprednisolone, 15 mg/kg) may be required, followed by oral corticosteroids in more severe cases. Colchicine has also reportedly been useful in patients with skin and joint symptoms, though success in a small, randomized controlled trial was limited. Colchicine should be used with caution in persons with kidney disease and in pregnant women. Most patients with cutaneous leukocytoclastic vasculitis are treated in an outpatient setting. Inpatient care is needed in patients who have severe systemic vasculitic syndromes and severe organ dysfunction. In the absence of internal involvement, the majority of cases of LCV resolve within weeks to months, with approximately 10% of patients experiencing chronic or recurrent disease.
Dermatologic Diagnosis: Leukocytoclastic Vasculitis

References


Medicine

The critical sense and sceptical attitude of the Hippocratic school laid the foundations of modern medicine on broad lines, and we owe to it: first, the emancipation of medicine from the shackles of priestcraft and of caste; secondly, the conception of medicine as an art based on accurate observation, and as a science, an integral part of the science of man and of nature; thirdly, the high moral ideals, expressed in that most “memorable of human documents” (Gomperz), the Hippocratic oath; and fourthly, the conception and realization of medicine as the profession of a cultivated gentleman.

— Sir William Osler, MD, 1st Baronet, 1849-1919, Canadian physician and one of four founding professors of Johns Hopkins Hospital