Case Report

Male patient, 73 years old, presented to the Emergency Department with coffee ground vomiting (5 times) associated with abdominal pain and melena. He was diabetic, hypertensive, and had chronic obstructive pulmonary disease and had past history of Coronary Artery Bypass Grafting (CABG). After three days, he developed septic shock which was associated with coma, purpuric rash and overt Disseminated Intravascular Coagulation (DIC), and renal impairment. He was managed by fluid resuscitation and intubation and was transmitted to Intensive Care Unit (ICU).

On Examination

Figure 1: Showed purpuric blackish edematous lesions over Right hand.
He was afebrile, his temperature was 36.8°C, his blood pressure was 104/68 mmHg with a heart rate of 71 beats/minute, respiratory rate of 17 breaths/minute, and oxygen saturation of 94% in room temperature air. He had generalized deep purpuric lesions on both hands, upper and lower extremities, some of lesions were blackish purpuric in color with tense blisters as in dorsum surfaces of both hands, as shown in Figure 1 & 2.

**Laboratory Investigations**

**As regard his complete blood count**

The clinically significant results are as follows:

- White blood cell count (WBC) of 6.97k/µL (N) (n4 – 10k/µL), with a neutrophilic predominance of 93.3% (n40-80%).
- Red blood cell count (RBC) of 4.02M/µL (L) (n4.5-505M/µL) with a hemoglobin of 9.9gm/dl (L), hematocrit of 32.90% (L) (n40-50%), Platelet count of 269 K/µL (N) (n150-410 K/µL).

**As regard his chemistry**

- His random glucose of 322mg/dL (high) (n70-140mg/dL), blood urea of 154.60mg/dL (high critical) (n 8.6-10.20mg/dL), creatinine of 2.99mg/dl (high) (n 8.6-10.20mg/dL), normal SGOT (AST) (22µ/L) (n 0-35µ/L), normal SGPT (ALT) (14µ/L) (n 0-4.1µ/L), normal bilirubin (total) (0.476mg/dL) (n 0-1.1mg/dL), low total protein (5.93g/dL) (n 6.6-8.7g/dL) and low albumin of 3.67g/dL (L) (n 3.97-4.94g/dL), low chloride (92.20mmol/L) (n 98-107mmol/L), low sodium (132mmol/L) (n 135-151mmol/L), and High critical potassium (8.02mmol/L) (n 3.4-5.1mmol/L).

**As regard coagulation profile**

- Partial thromboplastin time (PTT) was (84.40 sec.) (high) (n26-40 seconds), ThromboPlastin time (PT) of (16.60 sec.) (high) (n11-14.5 seconds), and INR of 1.30% (high) (n0.8-1.2%).

**Histological Findings**

Sections examined from skin biopsy showed dermis with small blood vessels surrounded by neutrophils, neutrophils dust, fibrinoid necrosis of the vessel walls, and extravasations of red blood cells. The overlying epidermis showed normal histology, as shown in Figure 3 & 4.
Discussion

The patient was admitted at the ICU, and his high blood sugar was controlled by insulin, his purpura was treated by the rheumatologist with methylprednisolone 750mg i.v. daily for three days, and was treated by heparin and broad-spectrum antibiotics, but unfortunately, he died due to severe sepsis after 8 days. Although, the histopathological examination suggested diagnosis of leuckcytoclastic vasculitis, the case was presented clinically as a purpura fulminans, and the methylprednisolone 750mg i.v. daily for three days which was prescribed by the rheumatologist was wrong because it flared the sepsis, and the best medication was fresh plasma and immunoglobulin G.

Differential diagnosis

Purpura fulminans

Purpuric Fulminans (PF) is a life-threatening disorder of acute onset characterized by cutaneous haemorrhage and necrosis caused by disseminated intravascular coagulation and dermal vascular thrombosis. Three distinct categories can be identified: inherited or acquired abnormalities of protein C or other coagulation systems, acute infectious PF, and idiopathic [1]. Its cardinal manifestations are presence of circumscribed ecchymosis of skin and symmetrical gangrene of the extremities with coagulation abnormalities suggestive of disseminated intravascular coagulation [2]. Late findings in necrotic areas are the formation of vesicles and bullae, which mark the development of haemorrhagic necrosis, and finally firm eschar, which ultimately sloughs. The distal extremities are often most severely involved, usually in a symmetric manner, probably due to fewer collateral channels for tissue perfusion, and the relatively greater impact of circulatory collapse on perfusion of distal vascular beds [3].

Development of systemic consumptive coagulopathy is a defining feature of PF, which distinguishes it from other forms of skin necrosis due to dermal vascular occlusion such as warfarin or heparin induced skin necrosis, thrombotic thrombocytopenic purpura, cryoglobulinaemia, antiphospholipid syndrome, or paroxysmal nocturnal haemoglobinuria. Fibrinogen, coagulation factors (factor V and VIII), and platelets are consumed in ongoing thrombosis and fibrinolysis. Prothrombin time, partial thromboplastin time are prolonged; fibrin degradation products are elevated, and protein C, protein S and antithrombin III are reduced. The histopathological hallmarks of PF are dermal vascular thrombosis and secondary haemorrhagic necrosis [4], which are identical to Schwartzman reaction [5]. Vasculitis, including a peri vascular neutrophilic infiltrate, is a characteristic feature of acute infectious PF which distinguishes it from other forms of PF.

Leukocytoclastic vasculitis

Leukocytoclastic vasculitis (LCV), also called hypersensitivity vasculitis, is a small-vessel vasculitis its incidence rate is about 30 cases per million people per year and affect men and women in equal numbers [6,7]. The skin is the organ most commonly involved by LCV and its presentation is a painful, burning rash predominantly in the lower extremities, with up to one-third of patients presenting with trunk and upper extremity involvement. [3] The most common skin manifestation of LCV is palpable purpura [8]. 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 [9].

Meningococcemia

The clinical pattern of meningococcemia varies. Persons with meningococcal disease may present with a nonspecific prodrome of cough, headache, and sore throat. After a few days of upper respiratory symptoms, the temperature rises abruptly, often after a chill, malaise, weakness, myalgia, headache, nausea, vomiting, and arthralgia are common presenting symptoms. A skin rash, which is essential for recognizing meningococcemia, is the characteristic manifestation. The skin rash may advance from a few ill-defined lesions to a widespread petechial eruption within a few hours. Meningococcemia’s potential rapidity of progression cannot be stressed enough. In fulminant meningococcemia, a hemorrhagic eruption, hypotension, and cardiac depression, as well as rapid
enlargement of petechiae and purpuric lesions, may be apparent within hours of the initial presentation [10].

**Warfarin-Induced Skin Necrosis**

Warfarin inhibits the activation of vitamin K–dependent clotting factors II, VII, IX, and X and the anticoagulant proteins C and S [11-13]. Warfarin-induced skin necrosis is a rare (0.01%-0.1%) but serious complication that usually occurs within the first several days following warfarin initiation, although case reports have described this phenomenon on later in therapy as well [12-15]. This adverse drug reaction appears to be associated with larger doses of warfarin [13]. The pathophysiology is not well understood; however, it is theorized that this complication is due to the initial procoagulant state that warfarin causes, leading to the formation of fibrin clots within the microvasculature. These blood clots cause interruption in blood supply to the skin, resulting in necrosis. The general progression of warfarin-induced skin necrosis is as follows: pain and erythema; petechial hemorrhages; red, purple, blue, or black discoloration; necrosis; and hemorrhagic blisters and/or bullae. Early recognition of symptoms and treatment are vital to preventing substantial morbidity associated with warfarin-induced skin necrosis; however, due to its rare occurrence, appropriate treatment has not been well established [12].

**Thrombotic thrombocytopenic purpura**

Thrombotic thrombocytopenic purpura (TTP) is a rare blood disorder characterized by clotting in small blood vessels (thromboses), resulting in a low platelet count [16]. In its full-blown form, the disease consists of microangiopathic hemolytic anemia, thrombocytopenic purpura, neurologic abnormalities, fever and Renal disease [17].

TTP can affect any organ system, but involvement of the peripheral blood, the central nervous system, and the kidneys causes the clinical manifestations. Patients with TTP typically report an acute or subacute onset of symptoms related to neurologic dysfunction, anemia, or thrombocytopenia. The neurologic manifestations include alteration in mental status, seizures, hemiplegia, paresthesias, visual disturbance, and aphasia. Fatigue may accompany the anemia. Severe bleeding from thrombocytopenia is unusual, although petechiae are common [18,19].

**Conclusion**

Purpura fulminans manifestations are confusing with many diseases, so we should differentiate it promptly from other diseases to avoid misdiagnosis, delay management, mistreatment, and sudden death as occurred in our presented case.

**References**


