Clinical conditions responsible for hyperviscosity and skin ulcers complications

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Abstract. In this brief review, we have examined some clinical conditions that result to be associated to an altered hemorheological profile and at times accompanied by skin ulcers. This skin condition may be observed in patients with the following conditions, such as primary polycythemic hyperviscosity (polycythemia, thrombocytemia) treated with hydroxyurea, primary plasma hyperviscosity (multiple myeloma, cryoglobulinemia, cryofibrinogenemia, dysfibrinogenemia, and connective tissue diseases), primary sclerocythemic hyperviscosity (hereditary spherocytosis, thalassemia, and sickle cell disease). In addition, it may be present in patients with secondary hyperviscosity conditions such as diabetes mellitus, arterial hypertension, critical limb ischemia and chronic venous insufficiency.

Keywords: Hyperviscosity syndrome, blood viscosity, skin ulcers

1. Introduction

The blood flow differs from that running through microvessels and that through large vessels. These differences refer to the blood composition, haemodynamics, and specifically blood viscosity. Rheological alterations play a prominent role in microcirculation than in large vessels haemodynamics. When a potentially ischemic condition emerges, some changes develop in microcirculation in relation to the diameter and the wall permeability of microvessel, the cell metabolism and the haemorheological profile. Physiologically, the blood flow is influenced by blood velocity, vessel diameter, structure and blood viscosity. As for the blood viscosity, this is determined by the haematocrit, the plasma viscosity, and the red cell aggregation and deformability. Blood viscosity varies in relation with the shear rate. Results have clearly demonstrated that red cell deformability and plasma viscosity are very significant at high shear flow while red cell aggregation occurs at low shear flow.

2. Primary hyperviscosity condition

Primary hyperviscosity condition may be subdivided into polycythemic, plasma and sclerocythemic [8, 21, 26, 58, 59].

Skin ulcers are uncommon in polycythemic hyperviscosity caused generally by the bone marrow proliferative states (polycythemia, thrombocytemia, leukemia). Skin ulcers, nevertheless, may be caused

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by the treatment with hydroxyurea (DNA synthase inhibitor) in patients with polycythemia and thrombocytemia. Their pathogenesis is not clear, but there is a possible connection with a direct toxicity of this chemotherapeutic agent on the skin. According to other authors, skin ulcers are instead related to the prolonged use of this molecule [35, 41, 48, 82], even if previously had been demonstrated that this molecule acts on erythrocyte geometry and deformability [30].

Patients with plasma hyperviscosity may be affected by skin ulcers; this hemorheological disorder is most often associated with paraproteinemias, which are clinical condition characterized by the presence of an abnormal immunoglobulin secreted by malignant B-lymphoid cells of monoclonal origin. In some cases, more than one monoclonal paraprotein is present in the same patient and in some instances the paraprotein is the immunoglobulin light chain, while in fewer cases it is the heavy chain.

Multiple myeloma (MM) and Waldenstrom’s macroglobulinemia (WM) are blood diseases complicated with paraproteinemias; in MM patients there may be the presence of skin ulcers [25, 86]. One of our studies (unpublished data) carried out in a group of MM patients has highlighted not only an increase in plasma viscosity (especially at low shear rate) and a decrease in hematocrit, but also a reduction in erythrocyte deformability. This data may be explained by the alteration of the lipid composition demonstrated in the erythrocyte membrane and in the plasma of MM patients [46, 104]. In addition, an alternative hypothesis may be explained by the presence of a paroxysmal nocturnal hemoglobinuria-like defect in the erythrocyte membrane of MM subject group [19, 64, 94, 100].

Cryoglobulinemia is a clinical condition in which the presence of plasma hyperviscosity may be associated with skin ulcers and in particular with leg ulcers. Cryoglobulinemia is due to the presence in the plasma or in the serum of one or more immunoglobulins which precipitate at a temperature below 37°C and redissolve on rewarming. The composition of cryoglobulins is heterogeneous. Three basic types are recognized according to the clonality and the type of immunoglobulins. Type I consist of monoclonal immunoglobulins, generally either IgM or IgG. Type II is an association of monoclonal IgM and polyclonal IgG. Type III is a mixture of polyclonal IgM and polyclonal IgG. Type II and III are described as mixed cryoglobulinemia because they consist of polyclonal IgG and IgM [74].

A percentage of 2% to 50% of patients affected by circulating cryoglobulins develop clinical symptoms. At the onset of the disease the most frequent symptoms found on 80% of patients are purpura, arthralgia and weakness.

Ferri et al. demonstrated an alteration in plasma and serum viscosity and in cellular filtration index in a subject group with mixed cryoglobulinemia at 37°C and at 25°C [31]. In several papers regarding mixed cryoglobulinemia the presence of skin ulcers has been observed [5, 32, 33, 37, 78, 108]. The hemorheological alteration influences and worsens the skin ulcers through tissue ischemia even if the mixed cryoglobulinemia is a systemic vasculitis and the hemorheological impairment may favor the immunological damage in the vessel wall. The outcome of the clinical history of patients with mixed cryoglobulinemia depends on the presence of possible concomitant diseases and complications and on their response to treatment.

Cryofibrinogenemia, primary or secondary, is a rare disorder characterized by cryoprecipitation of the native fibrinogen in the plasma, which can cause thrombotic occlusions of the small to medium arteries [65, 83]. Clinically, it is possible to distinguish patients with isolated or primary cryofibrinogenemia from those with associated cryoglobulinemia. Patients with primary cryofibrinogenemia suffered more frequently from recurrent and necrotic skin lesions [76]. The mean concentration of plasma cryoprecipitate is generally higher in patients with primary cryofibrinogenemia. This clinical condition causes a variety of skin manifestation, including skin ulceration [3, 9–11, 51, 84, 85, 106].

Cryofibrinogenemia is a treatable and potentially reversible disease. The use of corticosteroids in association with low-dose aspirin is considered the specific treatment of moderate forms, even if stanozolol results an alternative therapy. Immunosoppressive therapy, plasmapheresis, and/or intra-venous fibrinolysis are useful for treating some severe forms of cryofibrinogenemia.
Dysfibrinogenemias are clinical disorders in which a condition of plasma hyperviscosity has been found [51, 67]. Dysfibrinogenemias are characterized by structural abnormalities in the fibrinogen molecule that alters its functional properties [2, 42, 43, 79]. As previously demonstrated, the diagnosis of dysfibrinogenemia is related to a fibrinogen with abnormal structure or function. Generally, a dysfibrinogenemia may be found by discovering an abnormal thrombin time, with or without an abnormal reptilase time. The presence of a dysfibrinogenemia is suggested by a normal or increased immunologic level of fibrinogen with a lower functional level. Congenital dysfibrinogenemia is caused by heterozygosity for a mutation within any of the three fibrinogen chain genes. Acquired dysfibrinogenemia is a rare abnormality that may be found in patients with conditions such as: liver disease, multiple myeloma, autoimmune disorders and cancer [99].

Clinically, subjects with dysfibrinogenemia (congenital or not) are frequently asymptomatic (55%) even if some subjects will exhibit bleeding (25%), thrombotic complications (20%), or both, and among the thrombotic complications, skin necrosis may be discovered [18, 89]. Skin ulcers may be present in patients with connective tissue diseases, not only in systemic sclerosis, but also in rheumatoid arthritis, systemic lupus erythematosus and in ankylosing spondylitis [20, 27, 39, 69, 87, 88, 92, 103]. In all these clinical conditions the rheological abnormality [57, 80], has been attributed to the presence of some protein of polyclonal origin. In previous studies regarding connective tissue diseases, data demonstrates an increase in plasma and serum viscosity, and in erythrocyte aggregation; in some cases also a decrease in whole-blood filtration has been detected. All these clinical conditions are characterized by an evident increase in plasma viscosity that may explain the alteration of the skin microvascular blood flow, even if its direct effect might be mitigated by the increase in endothelial nitric oxide synthesis, that reduces the vascular resistance [97]. Plasma viscosity, in fact, controls the blood flow resistance, which regulates the vascular tone and preserves the functional capillary density [13, 98]. The impairment of this microcirculatory aspect may contribute to the pathogenesis of skin ulcers.

In sclerocytic hyperviscosity, the presence of chronic and refractory skin ulcers can be often present in some diseases, such as hereditary spherocytosis, thalassemia and sickle cell disease.

Hereditary spherocytosis is a common inherited disorder characterized by anemia, jaundice and splenomegaly. The primary erythrocyte alteration of this type of anemia is the loss of membrane surface area, leading to a reduced deformability referable to the defects in the membrane proteins such as ankirin, band 3, β-spectrin, α-spectrin or protein 4.2 [7, 57, 61, 72].

An uncommon complication of spherocytosis is the refractory chronic leg ulcer (2%) that heals after splenectomy [1, 73]. From the study based on two patients (father and son) affected by spherocytosis who had already undergone a splenectomy, we found an increase in whole-blood viscosity at low shear rate, a decrease in whole-blood filtration and especially a reduction in elongation index evaluated with laser diffractometry.

Thalassemia is a congenital haemoglobinopathy caused by defective synthesis of the α or β hemoglobin chains. β-thalassemias are a group of hereditary blood disorders characterized by anomalies in the synthesis of the β-chain. β-thalassemias can be classified into major, intermedia, minor and carriers [17, 36]. In β-thalassemia major [60] there is an evident decrease in whole-blood viscosity at low shear rate, a reduction in hematocrit and an increase in erythrocyte rigidity without any variation of plasma viscosity. As for the β-thalassemia intermedia [34] there is a significant decrease in hematocrit and in whole-blood filtration. In relation to the β-thalassemia minor [71] there is a reduction in the relaxation time and in the erythrocyte deformability, without any significant variation of elastic modul and surface viscosity. The β-thalassemia carriers [101] present a decrease in the elongation index and in the erythrocyte aggregation; patients with β-thalassemia intermedia may have painful, indolent and refractory leg ulcers [12, 53, 68].
Sickle cell disease is a genetic disorder of the haemoglobin (homozigous HBB val6) responsible for acute deep tissue damage, such as vaso-occlusive crisis, acute chest syndrome and spleen infarction and for chronic involvement regarding bones, kidney and skin. [4, 6, 73, 102, 104].

Previous studies highlight the disabling complication and the severity indicator of sickle cell skin ulcers [45]. There are several reports that indicate how in sickle cell disease (SCA) there is an evident alteration of the hemorheological pattern [57, 68]. Hemorheologically patients with SCA present a decrease in the hematocrit, in the whole-blood viscosity at low and high shear rates, in the aggregation index and especially in the elongation index. In addition, some hemorheological parameters and some hemorheological indexes (except for the hematocrit), seem to discriminate SCA patients with frequent crisis from those with infrequent crisis.

Our study based on a small number of patients with microdrepanocytosis highlighted an evident decrease in erythrocyte deformability, examined with laser diffractometry. A programmed erythrocytapheresis treatment was conducted on SCA patients and it has demonstrated to control and improve the acute and chronic complications accompanying this clinical condition [29, 47].

All these clinical disorders (hereditary spherocytosis, β-thalassemia, sickle cell disease) are marked out by an evident decrease in erythrocyte deformability, which acts in the microcirculatory zone through an obstructive mechanism and also influences the oxygen delivery to tissue [8, 28, 70, 107]; both these components contribute to the pathogenesis of skin ulcers.

3. Secondary hyperviscosity condition

Diabetes mellitus, arterial hypertension, critical limb ischemia and chronic venous insufficiency are among the diseases that may cause a secondary hyperviscosity condition and skin ulcers.

Our studies related to these disorders have discovered the presence of an alteration of the hemorheological profile. This is characterized by an increase in whole-blood, plasma and serum viscosity, by an increase in red cell aggregation, and by a decrease in erythrocyte deformability. In addition, our investigation using the spectroscopic fluorescence and employing fluorescent probes, found evident alterations of the erythrocytes and polymorphonuclear membrane rheology [14–16, 45, 55, 56].

The clinical course and the treatment of diabetic foot syndrome (ischemia, ulcers, gangrene) obtained with fibrinogen adsorption [50] or with Heparin-induced extracorporal LDL Precipitation (HELP) improves the prognosis of the ulcers of diabetic patients [76, 77, 105]. Diabetic foot syndrome is a complication of long-standing diabetes. The combination of macrovascular and microvascular disease associated with neuropathy leads to the development of leg ulcers.

A microvascular disease worsens with the increase in plasma viscosity and the decrease in red cell deformability, as observed in diabetes mellitus conditions. The increase in plasma viscosity may be explained by the presence of hyperfibrinogenemia. Factors influencing erythrocyte deformability in diabetes mellitus are: decrease in the surface/volume ratio related to the sorbitol cytosolic accumulation and to the membrane lipid alterations, increase in the cytosolic viscosity related to the reduction in the organic phosphates, increase in the calcium and glycated hemoglobin, alteration of the membrane dynamic properties related to the qualitative and/or quantitative membrane alterations of lipids and proteins.

To be underlined is the association of the diabetic disease with thrombocytopenia related to the increased platelet adhesiveness and aggregability; these laboratory findings are present in this metabolic condition before the development of vascular lesions and depend on poor metabolic control. The increase in platelet aggregation found in diabetic subjects aggravates the microcirculatory blood flow and slows down the healing of skin necrosis.

All these hemorheological and coagulative alterations described in diabetes are decisive for the microcirculatory disorders. Moreover it must be underlined that the increase in glycated hemoglobin,
besides reducing the erythrocyte deformability, shifts the hemoglobin dissociation curve and diminishes the P50. This latter certainly acts negatively on the oxygen transport and contributes to the skin lesion.

Moreover, the arterial hypertension may be associated with skin ulcers (Martorell’s ulcers), frequently symmetric and located in the distal third and anterolateral surface of the lower limbs [38, 54, 102]. Martorell’s ulcers are noticeable for their painful red blisters, which soon become blue, purpuric and finally ulcerate. These ulcers may be preceded by “pigmented pretibial patches”. Pain, relented healing and poor clinical response to standard therapy are a distinguishing sign of Martorell’s ulcers. The study of microcirculation shows an increase in resistance of the arterioles associated to a limited compensatory mechanism. Other causal factors in the genesis of these ulcers may be the alterations in the sympathetic innervation, a persistent arteriolar hypertonia and an abnormal arteriolar vascular response to vasoactive substances. The impaired hemorheological profile has a possible role in the clinical course of these ulcers and may contribute to the organic complications of arterial hypertension, such as left ventricular hypertrophy and retinopathy. Another interesting point is related to the abnormalities in hemorheological parameters, which are observed mainly in the high-renin than in low-renin hypertensive subjects. In essential hypertension the impaired tissue oxygenation, that seems to accompany this clinical condition, may have a role in the pathogenesis of skin lesions.

In peripheral arterial disease and especially in subjects with non-diabetic and non-hypertensive critical limb ischemia, the hemorheological alteration is ascribable especially to the erythrocyte rigidity and after arterial reconstruction there seems to be no improvement in the hemorheological profile [44, 49, 90]. However, the importance of the hemorheological profile in critical leg ischaemia refers to the negative results that blood viscosity and fibrinogen levels have on the intermittent claudication, as well as the negative prognostic significance of hemoglobin levels in the healing of amputations due to critical leg ischaemia, including the negative prognostic significance of fibrinogen in critical leg ischaemia. There is a break between the microvascular flow and the microvascular defence systems in patients with critical leg ischaemia. A more rapid and evident rheological effect may be obtained by the prescription of normovolemic hemodilution, pharmacological defibrinogenation and plasma exchange. In critical leg ischemia the pathogenesis of skin ulcers results to be complex: endothelial injury, and neutrophil and platelet activation, that influence the hemorheological pattern, might be responsible for the damage in microcirculation [23, 24].

The progression of chronic venous insufficiency may develop venous leg ulcers [22]. Persistent venous stasis associated with increased venous pressure develops venous ulcers. The increase in capillary permeability leads to the extravasation of proteins and fibrinogen from the capillaries. High fibrinogen concentration causes a fibrin cuff composition blocking the diffusion of nutrients, an microcirculation impairment and subsequent skin necrosis. In the pathophysiology of these venous ulceration evident is the trapping and the activation of PMN cells; in patients with venous ulcers we observed a decreased PMN membrane fluidity, an increased PMN cytosolic Ca"++ content and an abnormal response to the PMN integrins (CD11b, CD11c, CD18), especially after in vitro activation with PMA and fMLP [55]. The clinical evolution of the chronic venous insufficiency with skin ulcers is referable not only to the marked microcirculatory disorder, but in particular to the functional alterations of the polymorphonuclear cells [63, 91].

4. Conclusions

Several clinical conditions responsible for primary or secondary hyperviscosity may be associated with skin ulcers. However, a clear impact of the hemorheological alteration on these ulcers cannot be demonstrated. An acceleration of the healing process of skin ulcers has been obtained through
pharmacological treatment, plasma exchange, erythroapheresis or fibrinogen adsorption (rheosorb). Consequently, resulting in an improvement of the hemorheological profile.

References


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