Immunological aspects of chronic venous disease pathogenesis

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Abstract

Chronic venous disease (CVD) is a very common health problem concerning up to 1/3 of the society. Although venous hypertension and valvular incompetence have been long known to be crucial for development of the illness, its exact aetiology remains unclear. Recent findings indicate that inflammatory processes may be crucial for development of incompetent valves and vein wall remodelling. One of the most interesting theories describes “leucocyte trapping” as the mechanism responsible for elevated vein wall permeability and oxidative stress in the veins. At the same time, the cytokine profile of the blood in incompetent veins has not been thoroughly examined. Popular anti-inflammatory drugs relieve some symptoms but do not have much proved effects in prevention and treatment.

We intend to summarize the existing knowledge of the immunological aspects of CVD in order to emphasize its importance for understanding the aetiology of this illness. We also wish to indicate some aspects that remain to be studied in more detail.

Key words: cytokines, chronic venous disease, varices, veins, leukocyte trapping.

Introduction

The lower extremity superficial veins (great saphenous vein – GSV and small saphenous vein – SSV) are responsible for draining the skin and subcutaneous tissue. They are connected with the deep vein system by perforators. All these veins are equipped with valves responsible for preventing the blood reflux. If any derangement of this system occurs, chronic vein incompetence sets in. Blood stasis causes symptoms like leg heaviness, oedema, pigmentation and ulceration and the most severe complication is thrombosis which may lead to pulmonary thromboembolism and death.

Chronic venous disease (CVD) is a very common problem, especially in Western countries, in industrial regions [1]. According to numerous papers, varices may concern as much as 1/5-1/3 of the population with a slight predominance of women [2-7]. The risk factors are family history [8], age, obesity [9], female gender and pregnancy [1, 4]. The hereditary factors and age seem to be the most important factors promoting the illness [3, 4, 10].

Aetiology of venous insufficiency remains obscure. Valvular insufficiency and reflux have been long considered crucial for formation of varices and indeed, valves of varicose veins are widened and hypotrophic and the vessel lumen is larger than in healthy veins [11]. Nevertheless, the primary reason for these changes remains unclear. Recent publications suggest that inflammation may be the fundamental element in the pathogenesis of venous insufficiency [12, 13], and the phenomenon of the “leucocyte trap” is of great interest [14-18].

The following review aims at summarizing the information on immunological processes which could be of importance for the venous disease pathogenesis. We included only a few of the publications on venous ulcers because immunological changes in this stage of venous insufficiency may be largely dependent on bacterial presence (including endogenous microflora and pathogenic microflora like Staphylococcus aureus and Pseudomonas aeruginosa) [19], therefore they may differ a lot from the immunological state underlying the disease.

Leukocytes involved in chronic venous disease

Multiple works point to the possibility of macrophage-released growth factors, proinflammatory cytokines, matrix metalloproteinases (MMP) and adhesion molecules being involved in the aetiology of varicose veins [15, 16]. Out of all white blood cells it is the macrophage/monocyte group which is represented in greatest numbers in incompetent veins of lower extremities. These cells are located differently than in healthy veins, i.e. in proximity of valves and vein walls [15, 16, 20]. They are present in adventitia and in vasa vasorum but not in the muscular layer [21].

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Macrophage infiltration is most intensive near the valves and through proteolytic enzymes it may contribute to their destruction and mechanical weakening [20]. Patients suffering from venous ulcers have a greater number of monocye-platelet complexes, both in lower and in upper extremities [22]. The number of these complexes does not diminish even after radical surgery of GSV, which indicates that they must be present before venous hypertension evolves and so they take part in early pathogenesis [23].

Most publications claim that the number of T lymphocytes in varices is the same as in healthy veins [15] and there are very few papers on their role in aetiology of venous insufficiency. Bujan et al. [24] observed an increase in CD4+ cells in the varicose vein tissue. The localization of these cells was different depending on age: in the older group (over 50 years of age) they infiltrated subendothelium and valves, while in the younger group they were found in the adventitial and medial layers. Lymphocytes B were also found in greater numbers in varices than in healthy specimens, however, this was true only in the older group of patients.

We would also like to mention here some studies of arterial diseases, which point to a significant role of T-cells in pathogenesis even though, just like in venous disease, their number is the same as in healthy patients. In arteriosclerosis the macrophage: lymphocyte T ratio is about 4:1. However, it is not the number, but the activity of T-cells that proves important. In atherogenesis mostly Th1 and interferon γ (IFN-γ), as well as interleukin (IL)-1, IL-12 and IL-18 promote the illness [25]. Th17 lymphocytes are still a novelty; their role in atherogenesis (and even more so in venous insufficiency) is unclear. In atherosclerotic plaques small amounts of IL-17 mRNA were found, but IL-17 protein was abundant [25, 26]. Ju et al. implicate that IL-17A has a large role in aortic aneurysm formation in mice [27]. These studies show that lack of significant differences in lymphocyte number in patients in comparison with healthy people does not necessarily mean that these cells have no impact on illness pathogenesis. Lymphocyte activity in venous illness may be an interesting target for future studies, just like it is in atherogenesis.

Neutrophil population in wall and in blood of varicose veins is very small [15, 28]. In one study flow cytometry showed lack of neutrophil activation in varices, regardless of ulceration [29]. There were no differences in neutrophil-platelet aggregation as well [22], although in the presence of elevated shear force adhesion between these cells was increased and followed by aggregation via P-selectin and its ligand, glycoprotein PSGL-1 [30].

It is postulated that neutrophils have an important role in venous insufficiency pathogenesis. Their absence in venous blood is explained by “trapping” in tiny vessels [17, 28]. The “leukocyte trap” mechanism is described as infiltration of activated neutrophils caused by hypertension, hypoxia and stasis [17, 28, 31, 32]. In normal environment neutrophils do not contact each other and their halftime is about 7 h [33]. Inflammation lengthens this time and causes neutrophil aggregation and their adhesion to endothelium [33, 34]. First they adhere to endothelium via P, E and L selectin and they roll along the vessel wall [35]. Next, the CD11/CD18 complex along with endothelial intercellular adhesion molecules ICAM-1 and ICAM-2 cause strong adhesion [36, 37]. As a consequence neutrophils produce free radicals and proteases [35]. The adhesion process may be initiated by IL-8, PAF (platelet activation factor), active complement complex, arachidonic acid metabolites and other cytokines like IFN-γ [34, 35, 38]. Tumor necrosis factor α (TNF-α) increases neutrophil activation (adhesion, degranulation and free radical creation) via p55 and p7 [39].

In accordance with the “leucocyte trap” theory it has been shown that indeed, more neutrophils adhere to endothelium in veins incubated in hypoxic conditions [32, 33]. What is more, neutrophils adhering to the endothelium are active, with elevated cytoplasm calcium levels and considerable NO and LTB4 production [34, 38, 40]. Apart from hypoxia, also venous hypertension in standing position is related to inflammatory reaction [41], elevated L-selectin plasma levels and its decrease in neutrophils and monocytes [17]. Adhesion of active neutrophils and monocytes decreases after resting [17, 31]. Venous stasis causes release of IL-1b, IL-6 and TNF-α as well [16], suggesting endothelial activation [16, 41]. It is suggested that in plasma of varicose veins an unknown neutrophil activating factor is present because neutrophils from healthy patients added to plasma of patients suffering from varicose veins become activated and create multiple pseudopodia [28].

Pathological factors of vein wall remodelling

The role of free radicals in venous disease is well described. Enhanced production of free radicals by neutrophils in varicose veins was noted in numerous publications [14, 18, 42-44]. It seems that this production is performed mostly by NADPH oxidase (NOX) and by nitric oxide synthase (NOS) [43, 45], and in thrombophlebitis the xanthine oxidase is also relevant [14]. At the same time antioxidative mechanisms are weakened – superoxide dismutase (SOD) has lowered activity and the total antioxidative potential measured by the FRAP test (Ferric Reducing Ability of Plasma test) is small [46]. Free radicals, especially reactive forms of oxygen and nitrogen produced by active neutrophils and macrophages, cause degradation of the main protein components of the venous wall – collagen and elastin. Elevated concentrations of lipid peroxidation products like malondialdehyde (MDA), myeloperoxidase (MPO) and reactive thiobarbituric acid derivatives (TBARs) which are found in varices indicate elevated lipid peroxidation which destroys cell membrane, setting
free additional proteases [14, 46]. Moreover, it has been shown that incompetent venous fragments contain more ferric ions in comparison to normal vein fragments, there is also a greater concentration of zinc and copper which are SOD cofactors [47]. In incompetent vein fragments more oxidative damage and TBARs were found. Authors of this study emphasize the role of iron in oxidative stress and in chronic inflammation [47]. Removal of the incompetent vein causes a prominent decrease in oxidative stress markers in venous blood [48].

Endothelial protein degradation is also caused by elevated activity and expression of MMPs connected to hypertension and stasis [13, 41, 49, 50]. These damages in turn cause leukocyte infiltration and activation, manifested by elastase and lactoferrin increase [51, 52]. Elastase content is elevated in plasma of varicose veins regardless of inflammatory complications (like lipodermatosclerosis or ulceration) [52]. For venous ulcers, MMP-2 and MMP-9 appear as crucial factors which impair healing and the MMP-9 levels are elevated in wound fluid as well as in plasma [50, 53]. As the chronic wounds heal, a decrease in MMP-2 and MMP-9 is noted [54].

Gomez et al. showed decreased levels of active MMP-1 and MMP-2 in varices with increase of tissue inhibitors of metalloproteinases (TIMP-1 and TIMP-2) [55]. Sansilvestri, however, noted increased production of MMP-1, -2 and -3 with no difference in MMP-7 and -9 or TIMP-1, -2 and -3 [56]. Decrease in mitochondrial prostaglandin E, and its receptor, EP4 accompanied by increase in 15-hydroxyprostaglandin dehydrogenase, which is its only degrading enzyme, were also found. The authors conclude that the observed changes take part in thickening of the venous wall and suggest that these changes may have a protective impact, strengthening the vein during blood stasis [55].

The deranged activity of leukocytes along with factors released from platelets and macrophages leads to proinflammatory activity of endothelium and it is considered a potential source of pathologic vein wall remodelling, vein lumen widening and valve incompetency with following varices formation [31, 57].

At the same time there is no correlation between the intensity of systemic inflammatory reaction and the symptoms noted by the patients [38].

A rat model of venous hypertension (arterio-venous fistula) showed that the isolated increase of pressure does not cause reflux: the venous diameter grows but the valves remain competent. Only after some time, with increasing migration of monocytes/macrophages and lymphocytes T to the valves (and elevated expression of selectin P and ICAM-1 on endothelial cells) venous insufficiency sets in. Authors of this study administered a flavonoid prior to creating the fistula which to some extent decreased expression of adhesins and lymphocyte T infiltration [59].

As a result of inflammatory changes, the whole internal layer of the vein is degenerated with deformed, fragmented structure [60]. Endothelium presents inflammatory profile and it is prone to adhesion and migration of blood components which is proved e.g. by elevated surface expression of CD146 [15, 60-62]. Endothelial cells release multiple inflammatory mediators and growth factors like von Willebrand factor (vWF) [12], osteoprotegerin [62], smooth muscle proliferation inducing factors (PGF2α – prostaglandin F2α), βFGF – fibroblast growth factor β) [63], adhesion molecules intercellular adhesion molecule 1 (ICAM-1), vascular cell adhesion molecule 1 (VCAM-1), platelet endothelial cell adhesion molecule (PECAM-1) [64]. PECAM-1 has an anti-inflammatory function but it also takes part in preservation of vessel wall continuity [62]. It has a substantial role in clot dissolving during acute episodes of deep vein thrombosis – its lowered expression correlates with longer existence of thrombus (less macrophage infiltration, decreased angiogenesis) and increased fibrosis and the soluble form (sPECAM-1) could be a useful prognostic marker of postthrombotic syndrome [65].

Elevated concentration of monocyte/macrophage attracting factors in blood of varicose veins is probably not associated with endothelial cells, because removal of the endothelium does not cause decrease in expression of mRNA of these chemokines (IL-8, MCP-1, IP-10, RANTES, MIP-1α and MIP-1β) [66]. Endothelium of incompetent veins is more resistant to starvation (culture without serum or cytokines/growth factors), with decreased migration of its cells [62].

Another important feature of incompetent veins is increased angiogenesis accompanied by increased vessel permeability. It is caused probably by leukocytes adhering to the capillary wall and obstructing the vessels, releasing proteolytic enzymes and toxic metabolites [67]. Loss of larger molecules like fibrinogen causes formation of so called “cuffs” around the vein. These “cuffs” are often present in varices but their postulated role in formation of ulcerations has not been proved [68].

Venous intima has been shown to include an important subendothelial layer of pericytes which, unlike endothelial cells, have a prothrombotic profile. These cells store the tissue factor and in contrast to endothelium they cannot activate protein C [69, 70]. Therefore, once endothelial barrier is damaged, its antiocoagulatory properties are lost and the procoagulatory pericyte layer is exposed.

**Cytokines in chronic venous disease**

Lately Tisato et al. published two studies on cytokine profile of incompetent veins. In a study of a small group of 11 patients, among 31 cytokines tested the following cytokines were found in larger amounts as compared to control: IL-8, TNF-α, GM-CSF, IFN-α2, MIP-1β, VEGF, EGF, Eotaxin, MCP-1, PDGF and RANTES [71]. In another experiment with 31 patients with incompetent veins, out of 27 cytokines and chemokines assessed in the super-
natant from endothelial cells culture only PDGF-BB was significantly correlated to the reflux time in the incompetent vein [72]. Administering pharmacological inhibitors of main PDGF-BB synthesis paths with TNF-α as a stimulator, it was shown that NF-κB signalling path was the main contributor to the PDGF-BB synthesis (TNF-α receptor activation starts multiple signalling pathways. As a result of one of them, NF-κB is activated. This pathway is known to take part in TNF-α inflammatory actions) [72].

Studies on correlation between plasma VEGF levels and severity of the illness according to CEAP scale show that despite VEGF increase with exacerbating symptoms, a statistical significance is noted only in patients with healed ulcers [73]. Nevertheless, an interesting paper on VEGF was published by Hollingsworth [74]. He assumed that macroscopic changes in incompetent veins are not continuous, so a molecular defect requiring additional factors must take part in the pathogenesis. He studied VEGF because of its crucial role for continuity and reactivity of vessels. Incompetent vein fragments were compared to competent fragments of the same varicose veins and with specimens from healthy people. Of all studied genes of vessels, in incompetent vein fragments the sapheno-femoral junction was incompetent. While transcription of s.flt-1 was not increased for which transcription was greater in incompetent vein fragments. After administration of the four mentioned genes in hypoxic conditions, escin could be useful rather at an early stage of the illness because in more advanced stages the vein wall becomes insensitive for contracting effect of the drug [80].

Flavonoids are considered effective in protection of cells from active oxygen forms. Their antioxidative function is based on scavenging of free radicals and other transient oxidative oxygen forms. They are also supposed to stop neutrophil infiltration and free radical creation through preventing the activating cascade induced by hypoxia [14]. Micronized purified flavonoid fraction (MPFF) significantly reduces venous reflux in an animal model of venous hypertension, protecting the valves from narrowing and decreasing the number of lymphocytes T on the valves [41, 80]. Flavonoids administered for 60 days seem to decrease the plasma concentrations of some endothelium activation markers (ICAM-and VCAM) [81].

Unfortunately, some studies did not confirm any important impact of flavonoids on inflammatory markers (CD11b, CD18, IL-6, IL-8, selectins) [82]. Thirty days of driosmin administration in patients with healed venous ulcers did not show any anti-inflammatory effect (monitored by levels of cytokines and soluble adhesive molecules), on the other hand, an increased expression of adhesive molecule CD 11b on circulating granulocytes was shown. This effect is attributed by the authors to changed interactions between leukocytes and endothelium or the direct impact of the flavonoid on granulocytes [82].

300 mg of acetylsalicylic acid administered daily ameliorates the healing of venous ulcers but its administration for 15 days prior to surgical removal of varicose veins did not decrease expression of monocyte attracting chemokines in vitro [83]. This fact along with the short time of drug administration may be responsible for the lack of effect [83].

Another substance examined for its anti-inflammatory effects and possibility of therapeutic use in chronic venous insufficiency is a glycosaminoglycan – sulodexide (SDX) [84]. It causes a significant decrease in the release of almost all cytokines, chemokines and adhesive molecules from macrophages stimulated by lipopolysaccharide. In
a rat model of venous hypertension (arterio-venous fistula) [85] sulodexide decreased the expression of angiopoetin-2, simultaneously promoting expression of angiopoetin-1. According to the authors this may indicate ability to reduce endothelial hypertrophy in veins with hypertension. Sulodexide has antithrombotic and profibrolitic properties and it can restore the endothelial glycocalyx [86]. Importantly, glycocalyx includes glycosaminoglycans which are larger than most adhesion molecules and as long as they are intact, they prevent contact between e.g. ICAM-1 and leukocytes, thus reducing the inflammatory processes [19, 86].

In the aforementioned study by Tisato et al. [71] in the next stages of the study two substances were examined: α-lipoic acid and Ginkgo biloba derivative (GinkgoSelect phytosome). They both decreased cytokine expression in proinflammatory environment created by TNF-α addition. α-lipoic acid which acts through NF-κB and p38/MAPK path was more effective in comparison with Ginkgo biloba (acting through Akt path).

In a study from Philadelphia [45], MJ33, an indirect inhibitor of NOX was placed in anti-PECAM-1 immunoliposomes (Ab-MJ33/IL). They bound specifically with endothelium and neutralized angiotensin induced ROS production in vitro and in vivo. Moreover, AB-MJ33/IL turned out to be inhibitors of inflammatory marker VCAM expression in endothelium subject to TNF; they also diminished the damage in endothelial barrier caused by exposure to VEGF.

Summarizing, it can be stated that the inflammatory process is a significant part of venous insufficiency pathogenesis. Despite numerous studies there is still no clarity concerning the role of immunological cells in the aetiology of this illness, there are also few works on cytokine profile in varices. Such studies could help in targeting pharmacotherapy and as a result, better effectiveness of treatment and prophylaxis could be achieved.

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