Pericapillary fibrin in the ulcer-bearing skin of the leg: the cause of lipodermatosclerosis and venous ulceration

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Abstract

Forty-one biopsy specimens, taken from the ulcer-bearing skin of 41 legs of 21 patients attending the varicose vein clinic, were selectively stained for fibrin with phosphotungstic acid haematoxylin before being blindly assessed. Layers of fibrin were found surrounding the dermal capillaries in all 26 legs with lipodermatosclerosis. None of the specimens from the 15 legs with clinically normal skin contained fibrin. There was also an increased number of dermal capillaries cut in cross section per high powered field in 24 of the 26 legs with lipodermatosclerosis compared with two of the 15 legs with normal skin (p < 0.001). The mean reduction in foot vein pressure during exercise was significantly less in the 26 limbs with pericapillary fibrin than in the other 15 limbs (p < 10^-4). Lipodermatosclerosis is synonymous with pericapillary fibrin deposition and is associated with, and probably secondary to, both a persistently raised venous pressure and an increase in the size of the dermal capillary bed. This extravascular deposition of fibrin probably stimulates tissue fibrosis and blocks the diffusion of oxygen to the overlying epidermis, producing cellular death and venous ulceration.

Introduction

The mechanism by which venous disorders of the leg produce ulceration around the ankle is unknown. Ulceration is the final event in a well-recognized series of changes in the skin and subcutaneous tissues. The first changes are cutaneous pigmentation, mild ankle oedema, and the appearance of dilated subdermal venules. Later the skin and subcutaneous fat becomes thickened and hard. Since the tissues are often red and tender they are mistakenly thought to be infected or to be the site of superficial thrombophlebitis. At this stage minor trauma will cause an ulcer. If ulceration does not occur the skin and fat contract and the patient develops a tight narrow gaiter of hard skin. We prefer to call this whole process “lipodermatosclerosis” rather than give each clinical phase a separate name.

In a series of experiments designed to elucidate the underlying cause of these changes we have shown that legs with poor calf pump function, as shown by the failure of exercise to reduce foot vein pressure, have an increased number of capillary loops within the ulcer-bearing skin. Confirmation that this was a causal relation was obtained from an animal study in which an increase in the venous pressure produced a similar increase in the size of the capillary bed of the calf skin. Increasing the capillary pressure opens the intercellular endothelial pores and increases capillary permeability. Studies in animals showed that the high intraluminal venous pressure and increased endothelial surface area available for exchange did not change the rate at which albumin and sodium escaped from the vascular compartment into the subcutaneous tissue fluid, but fibrinogen, a much larger molecule, accumulated significantly faster in the tissue fluid around the enlarged capillary bed. The object of this study was to determine whether chronic venous hypertension and lipodermatosclerosis were associated with the accumulation of fibrinogen-fibrin in the perivascular interstitial spaces of human limbs.

Patients and methods

We studied 42 legs of 21 patients (10 men and 11 women) without venous ulceration who were attending our varicose vein clinic. The limbs were examined for evidence of superficial, communicating, and deep-vein incompetence. The presence of lipoderosclerosis was noted. Foot vein pressure during rest and exercise was measured successfully in 41 of the 42 legs. The foot veins of one limb could not be cannulated so this limb was excluded from the study. The method of pressure measurement has been described in detail elsewhere. The results for each leg were obtained by expressing the fall in pressure (mm Hg) as a percentage of the resting pressure. Bipedal phlebograms were obtained from all patients using the method described by Lea Thomas. The results of the clinical examination and phlebography were combined to classify all limbs into one of four categories, each

References


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reflecting an increasing degree of chronic venous hypertension: (a) normal limbs (not excluding those with occasional minor varicosities); (b) limbs without liposclerosis but with saphenous vein (long or short) incompetence and normal deep veins; (c) limbs with liposclerosis, saphenous vein (long and short) incompetence, and normal deep veins; and (d) limbs with liposclerosis and damaged deep veins.

**Assessment of extravascular fibrin**: Small biopsy specimens of skin (5 mm x 5 mm square) were taken from the medial side of the foot vein (long and short) of all 41 legs 7.5 cm vertically above the medial malleolus, a point which invariably coincided with the most severe liposclerosis when this was present. All the biopsies were stained with phosphotungstic acid haematoxylin (PTAH), which colours fibrin and related substances blue. The sections were then examined by one observer (HWA) who was unaware of the other findings. Each specimen was reported as showing fibrin present or absent on this examination. The number of capillary vessels cut in cross section in each high-powered field was also assessed in arbitrary units, as previously described.1 Biopsy specimens from two limbs with liposclerosis were examined for fibrin by an immunofluorescent technique using a rabbit-raised antifibrin and also studied with transmission electron-microscopy.8 The study was approved by the hospital ethical committee and all patients gave their informed written consent to the procedure.

**Results**

**Clinical phlebographic categories and venous hypertension**—Ten legs were normal; six legs had no liposclerosis but had saphenous vein incompetence; six had deep veins; five legs had liposclerosis, saphenous vein incompetence, and normal deep veins; and 20 legs had liposclerosis and phlebographic evidence of damaged deep veins. Thus 26 of the 41 limbs had liposclerosis. The mean percentage reductions in foot vein pressure during exercise in these four groups were: 58%, 55%, 59%, and 12%, respectively. These results confirmed the relationship between the physical signs and phlebographic assessment on the one hand and calf pump inefficiency on the other.

**Pericapillary fibrin**—A layer of fibrin (blue fibrillar material on PTHA staining) was seen surrounding the dermal capillaries in 26 of the biopsy specimens while no fibrin was seen in the other 15. Two specimens that showed the blue-staining pericapillary material on the PTAH sections were tested with an immunofluorescent antibody to fibrin. Both were found to have a halo of fluorescence around individual capillaries corresponding to the area that stained blue with PTAH, confirming the presence of fibrin. Electronmicroscopic examination of specimens from the same patients showed a material with the fibrillar structure of fibrin around the capillaries. The biopsy specimens of all 26 legs with liposclerosis contained pericapillary fibrin, while those from the 15 legs without liposclerosis did not contain pericapillary fibrin (p = 1.58 x 10^-13, Fisher's exact test). Twenty-four of the 26 specimens that contained pericapillary fibrin showed an increased number of dermal capillaries in cross section, while 13 of the 15 without fibrin had a normal number of capillaries (χ^2 = 25.57; p < 0.001).

**Fibrin and calf-pump efficiency**—The mean reduction of foot vein pressure during exercise in the 26 legs with pericapillary fibrin was 17.2 ± 18% (SD), which was significantly less than the mean reduction of 54.6 ± 20 in the 15 limbs without liposclerosis (t = 6.006; p < 10^-4).

**Discussion**

These studies show that fibrin is present in the skin of the ulcer-bearing area of the legs of all patients with lipodermatosclerosis. The fibrin is laid down as a cuff around the enlarged dermal capillary bed. We have previously presented experimental evidence that chronic venous hypertension increases the leakage of large molecules into the tissue spaces,9 and so we postulate that this accumulation of fibrin within the tissues is the result of greater quantities of fibrinogen escaping through capillary pores which are enlarged by the raised venous pressure.4 Since the fibrinolytic activity of the blood and the tissues is deficient in patients with lipodermatosclerosis,4 any fibrinogen which is converted to fibrin in the interstitial spaces is less likely to be broken down or reabsorbed.

The high venous oxygen tensions found in the venous blood of ulcerated limbs10-13 could be explained if it were shown that fibrin prevents the passage of oxygen into the tissues, so turning the capillaries into physiological shunts. The Appendix contains the details of a simple experiment which shows that a 1-mm layer of fibrin reduces the diffusion of oxygen across a layer of water by a factor of 25. We therefore suggest that the pericapillary layer of fibrin acts as a diffusion barrier, leading to local tissue ischaemia, which is manifested as lipodermatosclerosis or ulceration.

If fibrin deposition can be reduced at an early stage, before it starts to kill tissue or stimulates irreversible fibrosis, venous ulceration may be prevented. This may be accomplished by surgical restoration of calf-pump efficiency, by the use of elastic compression, or by the use of drugs which enhance tissue fibrinolysis.14

**Appendix**

The permeability of a 1-mm thick commercial fibrin membrane (Ethicon Ltd) was tested for air, oxygen, and carbon dioxide in a specially designed diffusion chamber. A fugacity gradient was produced for each gas to maximise diffusion velocity through the aqueous media, and gas movement was measured by a displacement velocity of a meniscus formed within a volumetric capillary tube connected to the gas reservoir. Each gas was tested five times after initial measurements conducted with air diffusion and no membrane. The diffusion capacity of the fibrin was obtained by dividing the rate of gas diffusion in millilitres standard temperature and pressure in dry air per minute by the pressure difference across the membrane in mm Hg.

The diffusion capacity of both oxygen and air with no membrane in place was 5.04 x 10^-4 cm^3/min/mm Hg, which fell to 0.228 x 10^-4 cm^3/min/mm Hg when the membrane was inserted. The diffusion of carbon dioxide (0.699 x 10^-4 cm^3/min/mm Hg) was not affected by the membrane and was very similar to the diffusion capacity of carbon dioxide through the alveolar membrane (0.598 x 10^-4 cm^3/min/ mm Hg).

A sheet of fibrin appears to allow carbon dioxide to pass through it relatively freely but is relatively impermeable to the passage of oxygen.

**References**


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