Immunocytochemical features of obstructed saphenous vein coronary artery bypass grafts

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SUMMARY The peroxidase-immunoperoxidase immunocytochemical method was used on 27 saphenous vein coronary artery bypass grafts, which had been resected because of recurrent angina, to identify in situ cellular and humoral elements possibly associated with graft occlusion. Immunostaining was performed on paraffin wax embedded control saphenous vein and graft sections incubated directly with primary antibodies against von Willebrand antigen (vWFAg), fibrinectin, fibrinogen, leucocyte common antigen (LCA), lysozyme, vimentin, desmin, platelet factor 4, and thrombospondin. Antigens were visualised by a chromogen providing an orange-red immunoprecipitate at the site of epitope localisation.

The intraluminal, amorphous exudate present in most grafts was not composed simply of fibrin or fibrinogen, as previously thought, but was a multiprotein complex including vWFAg, fibrinectin, thrombospondin and platelet factor 4. Along with macrophages, these components probably enter the graft after haemodynamic, physical, and chemical injury to, and disruption of, the endothelial cell. Progressive myointimal proliferation and fibrosis of these grafts may be local repetitive responses to macrophages and platelets, cells previously known to participate in vascular disease.

The purpose of this investigation was to identify specific cellular and humoral constituents within surgically removed saphenous vein and coronary artery bypass grafts which may be associated with late graft obstruction.1 Prior studies of these vessels by routine morphological assessment and staining may not have determined all the elements involved in reducing graft patency.2 In this study, therefore, we used the peroxidase-immunoperoxidase immunocytochemical technique3 to delineate previously unrecognised graft components which might have been associated with varying degrees of vasocclusion.

Material and methods

The grafts for study were obtained fresh from 16 consecutive patients who had further surgery because of unstable angina with angiographically documented luminal changes considered to be of clinical importance. Twenty seven grafts were removed from 14 men and two women, ranging in age from 35 to 72 years, with a mean of 58 years. Time elapsing since first bypass surgery varied from four to 13 years, with a mean of 9·1 years. Saphenous veins for identical analysis, used as controls, came from a separate group of 13 consecutive patients (11 men and two women, with a mean age of 60 years) about to receive their first coronary artery bypass.

All graft specimens were fixed in buffered formalin, embedded in paraffin wax, and sectioned at 5 μ. Tissue blocks from patients undergoing graft revision contained transverse sections of all the biopsy specimens from the site or sites of graft obstruction. Unfortunately, the sections were not necessarily identified by the vessel of origin, if more than one graft was involved, nor was the location of the single transection from each graft noted. Control sections were made by cutting 5 mm slices from each end and the centre of an unused portion of each venous segment and having the tissues processed as outlined above.

All biopsy specimens and control sections were stained first with Masson's trichrome, primarily to view the overall appearance of these vessels. The features being assessed were: the thickness of the vessel wall; whether separate wall layers were identifiable;
the relative diameter of the lumen and its shape; and if
the lumen was clear or contained an exudate.

Immunostaining of entire tissue sections mounted
on slides was performed by the triple sandwich
peroxidase-immunoperoxidase method, using
aminoethyl carbazole as the chromogen which
produced an easily seen red end-product, precipitating
at the site of the antigen to be identified. The
polyclonal and monoclonal primary antibodies were
chosen particularly to identify graft constituents of
potential pathogenic importance. The following
primary polyclonal antibodies, directed against the
specified antigen were made in rabbits: von
Willebrand antigen (vWFAg), the immunoreactive
portion of the factor VIII molecule necessary for
platelet adhesion; fibronogen; lysozyme, a monocyte
marker (all obtained from Ortho Diagnostic Systems,
Raritan, New Jersey); thrombospondin, important as
an adhesive protein, graciously provided by Dr Adam
Asch, Cornell University Medical College, New York;
and platelet factor 4, a platelet marker released from z
granules during platelet activation (American Dia-
agnostic, Greenwich, Connecticut). Antibody to
fibronectin, a protein functional in cell to cell inter-
action and adhesion, was purchased from BioGenex
Laboratories, Dublin, California. The following were
murine primary monoclones: leucocyte common
antigen (LCA), which recognises all leucocytes;
vimentin, which shows the presence of epitopes on
endothelium, macrophages, and fibroblasts; and des-
min, which shows smooth muscle (all purchased from
BioGenex Laboratories, Dublin, California). All
antibodies were used in dilutions to provide optimal
staining without background non-specificity. Method
controls consisted of assays without the primary
antibody and a saline substitute, with completely
irrelevant primary antibodies, and with chromogen
alone.

Results

Fig 1 shows a control saphenous vein stained with
Masson's trichrome before implantation. The vein
had the normal structure of a comparatively thin
intima and a proportionally thicker media and an
outer adventitia. The endothelium was smooth, the
lumen ellipsoid, sometimes circular, and clear. The
three cuts taken from each venous segment were
structurally similar to one another. Some veins,
however, showed variable degrees of subendothelial
thickening. When the control vein was immunostained
the intima was found to be acellular, the media
contained mainly smooth muscle marked by anti-
desmin with interspersed fibrocytes stained with
antivimentin. Only vWFAg and fibronectin appeared
as thin immunoprecipitates, sometimes limited to the
inner circumference of the vessel, and at times also
directly visible within the flat, single-layered endo-

Fig 1 Preoperative control saphenous vein. Note relatively narrow intima (small arrows), thicker media (heavy arrow), and
the outer, fibrillar adventitia (arrow head). The lumen is relatively symmetrical and clear. The different coats may vary
somewhat in width in different sectors.
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...thelial cells themselves (fig 2). Staining did not penetrate into the subendothelium. Thrombospondin, fibrinogen, and platelet factor 4 were not detected in the control veins.

Several types of changes were observed in the occluded grafts. The most common (11 of 27) was intimal myoproliferation of the wall. The lumen was reduced in calibre with a luminal exudate composed of an amorphous matrix and a cellular infiltrate. Higher magnification and immunostaining identified the cellular constituents of the exudate as mainly monocytes and macrophages reacting with anti-LCA and antilysozyme (fig 3). A few scattered cells with round nuclei, and a high nuclear:cytoplasmic ratio, also reacting with anti-LCA, had the appearance of lymphocytes.

The acellular portion of this exudate comprised four major proteins deposited as an extensive network within the obstructing graft lumen. Figs 4a–c identify immunostained vWFAg, fibronectin, and thrombospondin, respectively, providing a scaffold on to which the mononuclear cells have clustered. The presence of platelets was confirmed by released platelet factor 4 (fig 4d). The variation in staining intensity may have reflected the different concentrations of these proteins in situ. Surprisingly, fibrinogen did not immunostain in any graft areas. The inability to locate this protein may have been related to circulatory wash-out, epitope degradation, and proteolysis, or to limited accessibility of the antibody to the antigen within the other proteins.

The predominant characteristics (fig 5) of the second most commonly observed graft abnormality (eight of 27) were an exudate-free, reduced calibre irregular lumen in which were seen large, undulating projections of a thickened graft wall. These may be larger, more extensive versions of the myointimal protrusions seen in the first graft type. Demarcations of the vessel wall layers were unclear. When stained with vimentin and desmin antibodies, this exuberant mural proliferation was shown to be composed of smooth muscle and fibroblasts.

The remaining grafts were multivariate in appearance. Some were a composite between the first two presentations—namely, increasing myointimal proliferation with a comparatively reduced intraluminal exudate. In other instances the vessel wall was devoid of smooth muscle and had become completely fibrotic. Finally, other grafts retained only a minuscule lumen with closure by mural fibrosis in which neovascularisation had occurred. Obvious red thrombi were not observed.

Different forms of graft obstruction were sometimes observed simultaneously, but separately, in the biopsy specimens taken from different grafts in the same patient. These changes did not seem to have occurred over related time. Had multiple, serial sections of individual grafts been available, however, they may have shown transitional gradations from one type of appearance to another.

**Discussion**

A major new observation of this study is that the intraluminal, acellular exudate previously thought to be simply fibrinogen or fibrin is actually a complex multiprotein network. The in situ components shown by immunocytochemistry probably enter the graft after haemodynamic, physical, and chemical injury to the endothelium. These repetitive adverse stimuli include hypertension, hyperlipidemia, excessive shear stress, trauma at the perianastomotic graft site or exposure of the venous segment to an anomalous environment with an unusually high and pulsatile blood flow.

The finding of von Willebrand antigen, fibronectin, and thrombospondin, all products of endothelial synthesis, in the same place, and the deposition of vWFAg and fibronectin as aggregates, rather than in...
Fig 3  A photomicrograph to show macrophages as the major cell embedded in the amorphous meshwork. The inset illustrates a fine peripheral immunoprecipitate (arrowheads) formed on the macrophage with anti-LCA.

Fig 4  Four photographs of a single area immunostained with (a), anti-vWFAg; (b), anti-fibronectin; (c), anti-thromboplastin; and (d), anti-platelet factor 4. 5d has been enlarged to emphasise that platelet factor 4 has been deposited in very close opposition to fibronectin. Note, too, the difference in appearance of vWFAg and fibronectin in this occlusive matrix from the fine immunoprecipitate in the control vein (fig 2) formed by the same antigens.
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**Fig 5** The second most common form of an occluded graft with a thickened intima-subendothelium (small arrows) into which extends a proliferative media (large arrow), ultimately leading to wall thickening, loss of layer demarcation, and reduced patency.

 linear form, attest to endothelial cell disruption.\textsuperscript{13-15} It must be borne in mind that platelets are an additional source of these three proteins. Furthermore, the codistribution of platelet factor 4, a platelet specific protein, within the intraluminal glycoprotein matrix confirms the presence of platelets and degranulation at the site of injury.\textsuperscript{16,17} Finally, the migration of macrophages into the extracellular adhesive network is evidence of a chemotactic response to endothelial damage.\textsuperscript{18,19}

The different morphological and immunocytochemical presentations of these grafts imply that reduction in luminal patency may not be based on a single source or pathogenic sequence. The relative frequency, however, with which the glycoadhesive proteins vWF\textsubscript{Ag}, fibronectin and thrombospondin were deposited within a large proportion of these grafts, coupled with their inherent physiological function, suggest that they may be of importance in the genesis of graft obstruction. They may do this not only by interfering with blood flow themselves but also by providing an extracellular matrix into which monocytes, macrophages, and platelets migrate. These cells are known to have a role in vascular disease\textsuperscript{12,20,21} and they may function similarly in this instance, possibly as a source of mitogenic and proliferation-inducing biokines. Such macromolecules may stimulate replication of smooth muscle and fibroblasts with migration of these cells into the intima, resulting in luminal narrowing and graft failure.

Able technical assistance was provided by Ms Elizabeth Ross and Ms Marion B Brody. The manuscript was typed and collated by Ms Marene Urbany.
References


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