Histopathology of 'fresh' human aortic valve allografts

J. B. Gavin, B. G. Barratt-Boyes, G. C. Hitchcock, and P. B. Herdson

Department of Pathology, University of Auckland, and Cardiothoracic Surgical Unit, Green Lane Hospital, Auckland, New Zealand

Gavin, J. B., Barratt-Boyes, B. G., Hitchcock, G. C., and Herdson, P. B. (1973). Thorax, 28, 482-487. Histopathology of 'fresh' human aortic valve allografts. Six aortic valve allografts were studied histologically after having functioned in patients as aortic valve replacements for 14 to 442 weeks. The grafts initially had been collected under sterile conditions from cadavers and stored in Hanks’s balanced salt solution for 2 to 24 days before use. All grafts showed a cellular reaction along the host graft interface characterized by macrophages, lymphocytes, and organizing granulation tissue, and there was a progressive replacement of the donor aortic sleeve by host collagenous tissue. Sheaths of cellular, avascular host tissue extended from the margins of all grafts over their intimal surfaces and, in those which had been in place more than 36 weeks, this tissue had resulted in thickening of the proximal parts of one or more cusps. In one graft this thickening extended almost to the free margins of the cusps. While the leaflets of the graft at 14 weeks were virtually acellular, older grafts contained cellular areas with active fibroblasts in the proximal regions of the cusps as well as acellular regions which generally were more distally placed. Macrophages were always present along the interface between cellular and acellular areas. These observations indicate that there is a gradual replacement fibrosis of the graft by the host which proceeds in different grafts, and even different cusps in the same graft, at different rates.

Allograft heart valves collected under sterile conditions from cadaver donors and surgically inserted with minimal preparatory procedures, such as brief storage in Hanks's balanced salt solution, have proved very successful. These so-called fresh or untreated allografts appear to be less commonly affected by cusp rupture and other undesirable sequelae of more rigorous preparatory procedures (Barratt-Boyes and Roche, 1969; Barratt-Boyes et al., 1969; Barratt-Boyes, 1971; Barratt-Boyes et al., 1972). Although allografts are immunologically foreign classic rejection does not occur, due, it has been suggested, to their lack of vascularity and anatomic site (Baue, Donawick, and Blakemore, 1968) or their low inherent antigenicity (Mohri et al., 1967b). In the case of fresh allografts it is thought that donor fibroblasts may continue to function in the recipient and that this may be an important factor in their prolonged survival (Mohri, Reichenbach, Barnes, and Merendino, 1968; Kosek, Iben, Shumway, and Angell, 1969; Buch, Kosek, and Angell, 1971).

In view of their good clinical behaviour, human fresh heart valve allografts are difficult to obtain for detailed study and there are few reports of their histopathology (Duran and Whitehead, 1966; Kosek et al., 1969). This, together with the current efforts to devise preparative procedures for allografts which will preserve donor fibroblast viability (Al-Janabi, Gonzalez-Lavin, Neirotti, and Ross, 1972), prompted this review of six 'fresh' human grafts and their comparison with chemically sterilized (Gavin, Herdson, and Barratt-Boyes, 1972a) and antibiotic treated (Gavin, Herdson, Monro, and Barratt-Boyes, 1972b) allografts from the same hospital.

MATERIALS AND METHODS

Between 1962 and 1964, 16 human aortic valve allografts, which had been removed under sterile conditions from cadavers, were stored in Hanks's balanced salt solution from 2 to 24 days before placement in patients in the aortic position in a freehand manner (Barratt-Boyes et al., 1969). Six of these 'fresh' grafts were recovered either at necropsy or at the time of surgical replacement at intervals from 14
to 442 weeks after insertion and form the basis of this report. They were examined macroscopically and then fixed in 4% phosphate-buffered formaldehyde or in 5% phosphate-buffered glutaraldehyde. Paraffin-embedded sections were prepared from each cusp and consecutive sections were stained with haematoxylin and eosin, or by the elastic van Gieson, von Kossa, Hale's colloidal iron, periodic acid-Schiff, Gram, or alcian blue techniques for light microscopic examination. Contiguous segments of one graft removed after 442 weeks were examined with a Philips EM300 electron microscope. Data relating to these grafts, including the reason for their removal, are summarized in the Table.

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Donor Age</th>
<th>Donor Sex</th>
<th>Recipient Age</th>
<th>Recipient Sex</th>
<th>Days in Hanks’s BSS</th>
<th>Weeks in BSS</th>
<th>Reason for Removal of Graft</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1 31</td>
<td>F</td>
<td>62 F</td>
<td></td>
<td>10</td>
<td>14</td>
<td>Death from myocardial fibrosis and pulmonary congestion; allograft competent</td>
</tr>
<tr>
<td>2</td>
<td>2 53</td>
<td>M</td>
<td>43 M</td>
<td></td>
<td>3</td>
<td>36</td>
<td>Incompetence due to a peripheral suture line leak</td>
</tr>
<tr>
<td>3</td>
<td>3 37</td>
<td>M</td>
<td>21 M</td>
<td></td>
<td>24</td>
<td>65</td>
<td>Death from heart failure; allograft competent</td>
</tr>
<tr>
<td>4</td>
<td>4 48</td>
<td>M</td>
<td>17 F</td>
<td></td>
<td>4</td>
<td>169</td>
<td>Incompetence due to severe shrinkage of cusp leaflets</td>
</tr>
<tr>
<td>5</td>
<td>5 16</td>
<td>F</td>
<td>56 M</td>
<td></td>
<td>5</td>
<td>262</td>
<td>Incompetence due to shrinkage of all cusps with prolapse of one cusp</td>
</tr>
<tr>
<td>6</td>
<td>6 56</td>
<td>M</td>
<td>48 M</td>
<td></td>
<td>3</td>
<td>442</td>
<td>Incompetence due to peripheral leak plus rupture of one prolapsed cusp</td>
</tr>
</tbody>
</table>

BSS = balanced salt solution

Allografts from patients dying at 14 and 65 weeks after operation were competent and unremarkable. The other four grafts required replacement because they were incompetent. Peripheral suture line leak was the sole reason in case 2 and a contributory cause in case 6. In case 4 the allograft was grossly incompetent due to marked shortening of all three leaflets. Shrinkage of leaflets also contributed to the prolapse of one leaflet of the allograft valve in case 5. The longest surviving allograft in this group (case 6) was from a patient with luetic calcification of the aortic root and was incompetent due to the prolapse and rupture of its non-coronary cusp (Fig. 1). On removal this allograft showed an oval hole measuring 0.75 × 0.5 cm in one cusp, several groups of calcific nodules in all cusps (Fig. 1) and extensive calcification of the donor aortic sleeve.

Macroscopically the allograft valve leaflets were pliable but there was considerable variation in cusp thickness. The basal parts of the cusps of all grafts removed after 14 weeks were opaque and thickened due the presence of intimal fibrous sheaths. In cases 2 and 5 this thickening extended almost to the periphery of the leaflets (Figs 2 and 3) but in cases 3 and 6 it was limited to the basal quarter or less of the cusps.

Microscopically all grafts showed a reaction at the host/graft interface, a proliferation of host tissues over the intimal surface of the graft to form intimal fibrous sheaths, and changes in the graft itself.

**HOST/GRRAFT INTERFACE** The region of junction between the donor aortic valve and the host aortic wall was characterized by the presence of an infiltrate of macrophages and lymphocytes with smaller numbers of plasma cells, fibroblasts, and endothelial cells forming small vessels. With increasing graft survival time this organizing granulation tissue extended further into the

**FIG. 1. Allograft (case 6) on removal from a luetic patient 442 weeks after insertion. Calcific nodules are present in two cusp leaflets (arrows) and the third contains an oval fenestration.**
acellar donor tissue. A similar collection of inflammatory cells usually was present around embedded suture material, and occasional foreign body giant cells were noted. Several small foci of calcification were present in the donor aortic wall in the graft removed after 65 weeks (case 3), and there was extensive calcification in the graft removed at 442 weeks from the patient with luetic calcification of the aorta (case 6).

**Intimal Fibrous Sheaths**

Sheaths of avascular, collagenous connective tissue extended from the graft margin over sutures and along the intimal surface of the graft, tapering toward or onto the bases of the cusps.

The intimal fibrous sheaths were variable in extent. With the graft removed at 14 weeks the sheaths just reached the bases of the three cusps. At 36 weeks the sheaths were in continuity with the thickened basal parts of the cusps which extended more than half-way out to the free margins of three cusps (Fig. 2). After 262 weeks all cusps showed similar, but more extensive thickenings (Fig. 3). However, in the graft examined after 169 weeks the intimal fibrous sheath extended well onto two cusps but only a short distance onto the third, and in that examined at 442 weeks the sheaths were poorly developed on all three cusps.

Often the line of demarcation between the intimal fibrous sheath and the underlying donor tissue was indistinct as fibroblasts appeared to extend from the sheath into the underlying graft tissue (Figs 4 and 5). The intimal fibrous sheaths were richer in acid mucopolysaccharides and they

**FIG. 2.** Section through one cusp of heart valve allograft removed after 36 weeks (case 2). An intimal fibrous sheath (arrows) thickens the proximal two-thirds of the cusp. *Haematoxylin and eosin × 9.*

**FIG. 3.** Sections through the cusps of an allograft removed after 262 weeks (case 5). The cusps are considerably thickened by intimal fibrous sheath except near their free margins (arrows) where they are of normal thickness. *Haematoxylin and eosin × 6.*

**FIG. 4.** Cusp of an allograft after 36 weeks (case 2) showing the junction between the thickened cellular proximal part (C) and the thinner acellular distal part (A). *Haematoxylin and eosin × 180.*
FIG. 5. Base of one cusp from an allograft after 36 weeks (case 2); fibroblasts from the more cellular intimal fibrous sheath (S) above extend into the less cellular graft (G) beneath. Haematoxylin and eosin ×165.

FIG. 6. Base of a cusp from an allograft after 14 weeks (case 1). Except for a few cells (arrow) in the vicinity of a mural deposit of fibrin, the graft is acellular. Haematoxylin and eosin ×82.

FIG. 7. Part of the cytoplasm of a fibroblast from a cellular region of cusp leaflet from an allograft removed 442 weeks after insertion (case 6). Rough surfaced endoplasmic reticulum (arrows) is prominent indicating active protein synthesis. Electron micrograph ×16,500.

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contained fewer collagen and elastic fibres than the underlying donor tissue.

Changes in graft tissue The 14-week-old graft was virtually acellular apart from fibroblasts and inflammatory cells along the host/graft interface and in the vicinity of occasional mural deposits of fibrin (Fig. 6). Longer surviving grafts all contained both cellular and acellular regions. The acellular regions were common in the donor aortic sleeve and in the distal parts of the cusp leaflets (Fig. 4). While the allograft removed at 262 weeks (case 5) was almost completely populated by cells and the shortest surviving graft (case 1) was virtually acellular, the degree of cellularity did not vary consistently with the duration of the graft. Thus the graft removed after 36 weeks (Figs 4 and 5) was cellular in the proximal two-thirds of all three leaflets, whereas all the cusps of that removed after 442 weeks were mostly acellular apart from a few focal infiltrations of inflammatory cells and fibrin. Variations in cellularity were also observed between the leaflets of individual grafts. The 169-week graft (case 4) had two cusps which contained many cells but the third cusp was virtually acellular in the plane sectioned.

Many cells within the cellular regions contained prominent amounts of rough surfaced endoplasmic reticulum in their cytoplasm, indicating active
showed minimal disruption of regions macrophages, lymphocytes, fibrous sheaths. The histological cells intimal cellular and inflammatory reactions elastic and polysaccharides are prominent along this interface. Haematoxylin and eosin ×650.

FIG. 8. The interface between intimal fibrous sheath (S) and the underlying much less cellular cusp (G) of a graft removed after 262 weeks (case 5). Macrophages (arrows) are prominent along this interface. Haematoxylin and eosin ×650.

protein synthesis (Fig. 7). These were surrounded by a matrix rich in acid mucopolysaccharides and a relatively sparse network of collagen fibres with macrophages, lymphocytes, and other inflammatory cells scattered among them. Macrophages were invariably present along the boundaries of cellular and acellular regions (Fig. 8) and often lay in lacunae within the cusp matrix, apparently resorbing it. Cellular areas were almost always continuous with more proximal cellular regions and ultimately with host/graf interface or with the intimal fibrous sheaths.

In contrast to the cellular areas the acellular regions of the cusps stained weakly for acid mucopolysaccharides and the dense meshwork of elastic and collagenous elements had normal staining reactions and a normal ultrastructure and showed minimal disruption and separation. Endothelial cells were absent from the acellular parts of the graft although many were present on the intimal fibrous sheaths.

DISCUSSION
The histological changes found in fresh allografts in the present study are essentially the same as those in chemically sterilized (Gavin et al., 1972) and antibiotic treated (Gavin et al., 1973) allo-

grafts, but there are striking differences in the degree to which they are expressed. In fresh allografts focal infiltration of fibrin and inflammatory cells into the cusps is comparatively uncommon and intimal fibrous sheaths are more extensive and merge almost imperceptibly with underlying donor tissue. These latter observations indicate that host fibroblasts can proliferate into and across allograft tissue which is untreated more easily than when it has been chemically sterilized or treated with antibiotics. It is likely that molecular cross-linking among protein and ground substance components by beta propiolactone or ethylene oxide and perhaps also by the beta lactam moiety of penicillin could be responsible for these differences (Seelye, 1972).

The development of extensive intimal fibrous sheaths and the replacement of the cusp leaflets of fresh allografts by ingrowing host tissue might appear to be advantageous. However, in the present study these changes were accompanied by considerable thickening and sometimes by shortening or shrinkage of the affected cusps. Leaflet shrinkage is probably due to contraction of the newly formed host collagen as it matures. While thickening may cause minor loss of leaflet pliability, leaflet shrinkage is a potentially more serious change. In case 4 severe cusp shortening was undoubtedly the cause of the serious incompetence, and in case 5 it was an important contributory factor.

How often it occurs is unknown but in our series of fresh allografts (Barratt-Boyes et al., 1969) it is now known to be the cause of late serious incompetence in 2 out of 13 patients followed for 44 months or longer. Its occurrence emphasizes the importance of ensuring adequate central cusp redundancy at the time of allograft insertion.

In previous reports from fresh allograft material leaflet calcification and rupture had not been noted (Barratt-Boyes et al., 1969). Their occurrence in case 6, where the graft was removed because of severe incompetence eight and a half years after insertion, indicates that fresh grafts are not immune to degenerative changes. It seems likely, however, that the severe syphilitic aortic root calcification present in this patient may have modified the host reaction.

Our morphological observations are similar to those of Kosek et al. (1969), who studied both human and canine fresh allografts, and of Mohri et al. (1967a, b), who reported on experimental allografts in dogs, but our interpretation is different. Mohri et al. (1967a) reported that 37%
of cells cultured from an aortic valve from a female dog had sex chromatin in their nuclei and that four and six months after transplantation of such a valve into a male dog sex chromatin was still present in 15% and 16% of the cells respectively. In a subsequent paper Mohri et al. (1968) reported that 20% of cells cultured from grafts after one year had donor sex chromatin. But what of the other 80% of cells? Kosek et al. (1969) identified such condensed chromatin in endothelial cells in male to male transplants in several instances, indicating their donor origin. There is no doubt that, while tissue culture methods provide direct evidence of cell viability, the results of associated sex chromatin analysis to determine host or donor origin of cultured cells must be interpreted with care. Basu (1966) has reviewed sex chromatin in transplanted tissues and states that a reduction in the expected percentage of sex chromatin-containing cells in a female transplant indicates replacement of the cellular elements of the graft by host cells. He also cites Klen (1964), who considered that 20% or more of corneal epithelial cells should contain condensed chromatin bodies before a female origin could be concluded. The development of a method, perhaps involving immunofluorescence techniques, which would specifically identify host and donor cells would be most useful in clarifying the role of the donor fibroblast in the heart valve allograft.

Other groups of workers also have described regions of acellularity in fresh allografts but considered them to be regions which were non-viable prior to transplantation (Kosek et al., 1969) or which were subject to mechanical injury at the time of insertion (Mohri et al., 1967a). In another paper the diffuse or segmental acellularity of viable grafts was considered to indicate a limited life-span of donor cells (Mohri et al., 1968). Thus all observations on untreated heart valve allografts are consistent with the loss of donor cells at some time after the insertion of the graft.

We believe that the essential pathological process affecting heart valve allografts is replacement fibrosis by the host. This conclusion is based on the acellularity of the fresh graft recovered after 14 weeks, the presence of acellular areas surrounded by macrophages in all grafts, and a generally progressive extension of cellular areas towards the periphery of the cusp leaflets with time. However, the reasons for the wide differences in the rate of replacement between similarly treated grafts remain obscure.

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REFERENCES
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