Pathological view of sudden cardiac death

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SUMMARY The common cause of sudden cardiac death is ischaemic heart disease. Such patients may have an occlusive recent thrombosis in a major coronary artery but the largest group has no recent occlusion. Comparison of such patients without occlusion with non-cardiac death control hearts suggests that an area of stenosis of 85 per cent is the best discriminating level. Most subjects who die of ischaemic heart disease suddenly have this degree of stenosis in two or three major arteries. Non-ischaemic sudden cardiac death occurs in hypertrophic obstructive cardiomyopathy and in severe left ventricular hypertrophy particularly from aortic valve stenosis. When the heart is macroscopically normal, review of previous electrocardiograms is the most helpful guide and may disclose conditions such as a long QT interval or pre-excitation. When no such data are available examination of the conduction system histologically may be helpful but is often non-specific. Use of the term "cardiomyopathy" by pathologists to cover all non-ischaemic sudden cardiac death is clinically misleading.

The pathologist, when faced with investigating sudden cardiac death in an adult or a child over 1 year of age, can usually place the heart into one of three categories. In order of frequency these are ischaemic heart disease, conditions recognised macroscopically known to be associated with sudden death and, finally, hearts which are, at least to the naked eye, normal. There is an understandable temptation for the pathologist to transfer hearts in the third category to the first in adults since few coroners view "no clear cause of death" with enthusiasm. In children this way out is impossible and more exacting studies are more readily undertaken.

Ischaemic heart disease

Annually in the United Kingdom many thousands of patients dying suddenly from ischaemic heart disease come to a necropsy ordered by the coroner. It must be stressed that these forensic necropsies are to exclude unnatural death and not to provide accurate scientific data. In consequence, there is surprisingly little detailed knowledge of the pathology of ischaemic heart disease related to sudden death.

A proportion of these patients do have a recent occlusive thrombus in a major coronary artery. Death can be assumed to be from subsequent ventricular fibrillation. The proportion of cases with such a thrombus is reported to be from 4 to 64 per cent (Table 1). Such widely divergent figures are unlikely to be a true reflection of the pathology. The wide range reflects, in part, the degree of care taken or the beliefs of the individual pathologist. Even within a single pathological department the proportion of thrombi found by different individual consultant pathologists varies widely.16 Other factors which militate against comparability of series are different temporal definitions of the term "sudden", varying proportions of smokers to non-

<table>
<thead>
<tr>
<th>Author</th>
<th>Place</th>
<th>Definition of sudden</th>
<th>No. of Thrombosis cases (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Friedman et al.1</td>
<td>San Francisco</td>
<td>&lt; 1 min</td>
<td>27</td>
</tr>
<tr>
<td>Koller et al.1</td>
<td>Baltimore</td>
<td>&lt; 24 h</td>
<td>486 15</td>
</tr>
<tr>
<td>Lie and Titus4</td>
<td>Minnesota</td>
<td>&lt; 6 h</td>
<td>120 17</td>
</tr>
<tr>
<td>Spain and Bradess4</td>
<td>USA</td>
<td>&lt; 1 h</td>
<td>189 18</td>
</tr>
<tr>
<td>Titus et al.5</td>
<td>Minnesota</td>
<td>&lt; 1 h</td>
<td>286 19</td>
</tr>
<tr>
<td>Perpez et al.6</td>
<td>Pennsylvania</td>
<td>&lt; 24 h</td>
<td>171 22</td>
</tr>
<tr>
<td>Haerem1</td>
<td>Oslo</td>
<td>&lt; 10 min</td>
<td>47 30</td>
</tr>
<tr>
<td>Myers and Dewar4</td>
<td>Newcastle</td>
<td>&lt; 1 min</td>
<td>10 30</td>
</tr>
<tr>
<td>Mitchell and Schwartz7</td>
<td>Oxford</td>
<td>&lt; 6 h</td>
<td>31 66</td>
</tr>
<tr>
<td>Davies and Popple18</td>
<td>London</td>
<td>&lt; 6 h</td>
<td>120 33</td>
</tr>
<tr>
<td>Baba et al.11</td>
<td>Ohio</td>
<td>&lt; 24 h</td>
<td>121 38</td>
</tr>
<tr>
<td>Myers and Dewar4</td>
<td>Newcastle</td>
<td>&lt; 24 h</td>
<td>66 42</td>
</tr>
<tr>
<td>Rissnien et al.14</td>
<td>Finland</td>
<td>&lt; 24 h</td>
<td>141 44</td>
</tr>
<tr>
<td>Scott and Brigg13</td>
<td>New York</td>
<td>&lt; 1 h</td>
<td>183 46</td>
</tr>
<tr>
<td>Libetson et al.1</td>
<td>Miami</td>
<td>&lt; 15 min</td>
<td>220 58</td>
</tr>
<tr>
<td>Friedman et al.1</td>
<td>San Francisco</td>
<td>&lt; 24 h</td>
<td>37 59</td>
</tr>
<tr>
<td>Crawford et al.14</td>
<td>London</td>
<td>&lt; 1 h</td>
<td>75 64</td>
</tr>
</tbody>
</table>

Table 1: Frequency of recent occlusive coronary thrombosis in sudden death caused by ischaemic heart disease
smokers, those with and without hypertension, and different age and sex ranges. Even when the available published data are grouped, however, by a definition of deaths within a minute (instantaneous), within six hours, or within 24 hours no consistent figures emerge for the incidence of occlusive thrombi. Our own continuing study of 120 sudden deaths within six hours from ischaemic heart disease suggests the incidence to be exactly 33 per cent (Table 1).

It is hardly surprising that sudden ventricular fibrillation should follow a coronary artery occlusion. The situation can be mimicked experimentally by coronary ligation in the dog. It is observed in patients with acute myocardial infarction who reach hospital. There is some evidence suggesting that right coronary artery occlusions are more often associated with sudden death. The limited published data available suggest that the ratio of right to left anterior descending artery occlusions is lower in patients dying in hospital of infarction than sudden death patients not reaching hospital (Table 2). James et al. has reviewed the possible causes of this association which leads the right coronary to be regarded as the artery of sudden death. The major factor is the role of the right coronary artery in supplying both sinutrial and atrioventricular nodes.

![Graph showing comparison of worst area of coronary artery stenosis and age in male patients dying suddenly (<6 h) of ischaemic heart disease (open circles) and age matched non-cardiac deaths (closed circles). The majority of ischaemic deaths show stenosis at least at one point of over 85 per cent. Some control hearts, however, show an identical degree of stenosis.](http://heart.bmj.com/)

Table 2: Site of occlusive coronary thrombus in acute myocardial infarction and sudden ischaemic death

<table>
<thead>
<tr>
<th>No.</th>
<th>Main left</th>
<th>LAD</th>
<th>R</th>
<th>LC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute myocardial infarction (hospital deaths)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Plotz et al.</td>
<td>1495</td>
<td>71</td>
<td>834</td>
<td>379</td>
</tr>
<tr>
<td>Davies et al.</td>
<td>460</td>
<td>15</td>
<td>219</td>
<td>160</td>
</tr>
<tr>
<td></td>
<td>1955</td>
<td>86</td>
<td>1053 (54%)</td>
<td>539 (28%)</td>
</tr>
<tr>
<td>Sudden death</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Friedman et al.</td>
<td>21</td>
<td>1</td>
<td>11</td>
<td>7 (33%)</td>
</tr>
<tr>
<td>Davies and Popple</td>
<td>39</td>
<td>0</td>
<td>18</td>
<td>16 (41%)</td>
</tr>
<tr>
<td>Myers and Dewar</td>
<td>40</td>
<td>0</td>
<td>12</td>
<td>19 (48%)</td>
</tr>
<tr>
<td>Crawford et al.</td>
<td>48</td>
<td>0</td>
<td>27</td>
<td>15 (31%)</td>
</tr>
<tr>
<td>Bashe et al.</td>
<td>46</td>
<td>3</td>
<td>16</td>
<td>19 (41%)</td>
</tr>
<tr>
<td></td>
<td>194</td>
<td>4</td>
<td>84 (43%)</td>
<td>76 (39%)</td>
</tr>
</tbody>
</table>

All pathological studies agree in that there is a second large group of patients who do not have an occlusive thrombus or, indeed, any easily demonstrable recent morphological change in the coronary arteries. This second group of patients has coronary stenosis caused by atherosclerosis but for the pathologist the difficulty is in defining the minimal degree of disease which is sufficient to be causally related to death. Our own studies quantifying in detail the degrees of coronary stenosis suggest that for practical purposes 85 per cent stenosis at any single point is the lower limit (Fig.). Statistically, the best breakpoint between control and test groups is 75 per cent stenosis. The majority of the patients in the test group with over 75 per cent stenosis have double or triple vessel disease (Table 3), but a minority of 12 per cent has this degree of stenosis in only one major vessel. The problem with such figures, for the pathologist, is in that a group of age and sex matched control hearts from patients dying with intracerebral tumours or from trauma overlap the test group in the degree of coronary artery disease (Fig.). In such a control group 12 per cent

![Graph showing frequency of arteries with more than 75 per cent stenosis in any one 0-4 cm segment: age-sex matched controls](http://heart.bmj.com/)

Table 3: Frequency of arteries with more than 75 per cent stenosis in any one 0-4 cm segment: age-sex matched controls

<table>
<thead>
<tr>
<th>No. of arteries</th>
<th>Test (%)</th>
<th>Controls (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>+</td>
<td>52</td>
</tr>
<tr>
<td>1</td>
<td>12</td>
<td>21</td>
</tr>
<tr>
<td>2</td>
<td>18</td>
<td>19</td>
</tr>
<tr>
<td>3</td>
<td>49</td>
<td>8</td>
</tr>
</tbody>
</table>

* One death wrongly certified at necropsy.
of subjects had stenosis of over 85 per cent at some point in the coronary tree (Fig. 1). This overlap represents the high background level of atherosclerosis in our population. Comparing single, double, and triple vessel disease in the control and test groups, longer segments of diseased vessel are found in the latter but again there is overlap of the groups. The number of patients in the general population with over 85 per cent stenosis at some point is too high to be explicable by competing causes of death, that is if the patient had not developed a cerebral tumour he would have died the next year from ischaemic heart disease.

Clinical studies of resuscitated “sudden death” patients\(^{14} \)\(^{21} \)\(^{22}\) support the concept that there are two groups of patients, one which develops myocardial infarction and one which does not. The mechanism of death in this latter group once again seems to be the onset of ventricular fibrillation.\(^{21} \)\(^{26}\)

The myocardial resistance to ventricular fibrillation, that is its “electrical stability”, may be one factor separating test and control patients with identical degrees of coronary disease but one dying the other living.\(^{57}\) Epidemiological evidence suggests that the different mortality rates from ischaemic heart disease in hard and soft water areas\(^{28} \)\(^{29}\) are not the result of differences in the degree of stenosis and are, therefore, presumably mediated by the differences in electrical stability of the myocardium. This may reflect altered ion concentrations in the myocardium, particularly of calcium and magnesium.\(^{31}\) In patients whose electrical stability is lowered by coronary artery stenosis the precipitating factor to invoke ventricular fibrillation may be sympathetic stimulation or neural or psychological factors.\(^{57}\)

To demonstrate a morphological cause for myocardial instability is the wish of many pathologists. The reported proportion of patients with ischaemic heart disease dying suddenly who have a degree of myocardial necrosis (that is anything from a tiny focus of 0.1 mm to a large regional infarct) is very wide, from 12 to 85 per cent.\(^{3} \)\(^{32}\) The lower figures are from those who use naked eye examination only, the higher when refined “histochemical” stains are used. The difficulty with the latter technique is the occurrence of very high positive results in control patients, a failure to distinguish true from agonal myocardial changes.

The problem is also one of the timing of death. A proportion of those cases with an occlusive thrombus can be predicted to develop regional myocardial infarction but when death occurs within 12 hours it is difficult for the pathologist to demonstrate necrosis. Figures from Seattle\(^{26}\) indicate that only 19 per cent of patients resuscitated from “sudden death” do develop myocardial infarction demonstrable by the electrocardiogram. Since it is well known to pathologists\(^{33}\) that not all occlusive thrombi produce myocardial infarction the Seattle figures probably underestimate the incidence of occlusive thrombi. The occlusion when present does, however, act as a trigger to provoke ventricular fibrillation.

Platelet emboli into small intramyocardial arteries from more proximal, but non-occlusive, complex atheromatous plaques or thrombi are postulated to cause sudden death.\(^{7} \)\(^{34}\) Since there is extreme difficulty in demonstrating disintegrated platelets by conventional histological techniques it is impossible to confirm or refute the suggestion absolutely. It is certainly possible that a shower of disintegrating platelets may not only block small arteries but their “pharmacological” contents could provoke intense spasm more distally. Most pathologists have seen isolated cases where platelet emboli are found, but my experience suggests it to be rare, at least in any form demonstrable by conventional methods. The role of pure coronary artery spasm is clearly not amenable to investigation in the dead heart and must remain speculative.\(^{35}\) It is apparent that a great deal more work on sudden ischaemic death is needed. It is salutary to consider that despite many thousands of necropsies a year in the United Kingdom we still do not know if the incidence of thrombi differs in smokers and non-smokers and whether the high rate of sudden death in Scotland as compared with the rest of the United Kingdom\(^{36}\) is associated with differences in the coronary artery pathology.

Cardiac death not caused by coronary atheroma

The second major group of hearts are those with recognisable, if unrecognised, non-ischaemic cardiac disease. Any cause of severe left ventricular hypertrophy but particularly aortic valve stenosis may be associated with sudden death. In practice, hearts of a total weight over 550 g will have sufficient ventricular hypertrophy to be reasonably associated with sudden death. In cases of aortic stenosis sub-endocardial recent necrosis is usually demonstrable in the left ventricle. Gross right ventricular hypertrophy also carries a risk of sudden death usually associated with previously unrecognised pulmonary valve stenosis, obstructive cardiomyopathy, or primary pulmonary hypertension. All the examples seen personally have had isolated right ventricular weights of over 110 g.

Hypertrophic obstructive cardiomyopathy (HOCM) continues to be missed by pathologists at
necropsy largely because of a failure to appreciate
its wide morphological spectrum and by opening the
left ventricular outflow tract through the
anterior cusp of the mitral valve. This technique
successfully masks the encroachment of the septum
on the ventricular outflow tract. In my experience
sudden death is seen particularly in men (Table 4)
and has been reported to occur particularly in
certain families. I have found no morphological
features to distinguish those cases dying suddenly
from those not. I have seen examples from under
10 to over 80 years of age at the time of sudden
death. The pathologist should suspect hypertrophic
cardiomyopathy in any heart showing ventricular hypertrophy with a small left ventricular
cavity for which there is no obvious cause. Far too
many cases of HOCM are labelled as hypertensive
cardiomyegaly by pathologists. Inclusion as a
standard necropsy practice of measurement of the
septum and posterior wall with ratios over 1.6
confirms many of these cases to be hypertrophic
cardiomyopathy. I have not observed aortic valve
stenosis to produce this figure (Table 5). In
hypertrophic cardiomyopathy gross hypertrophy of
the free left ventricular wall, on occasions, tends to
mask the septal asymmetric hypertrophy leading to
an erroneous diagnosis of “hypertensive” cardio-
megaly. It is not widely appreciated that asymmetric
hypertrophy, in fact, may be symmetric at nec-
ropsy. When echocardiograms of such patients
are available the asymmetry is often seen to be
more apparent in diastole, and it is in the contracted
postmortem heart that problems in diagnosis arise.
On rare occasions the mass of abnormal muscle is
not septal. A reversal of the septal/posterior wall
ratio may also therefore indicate hypertrophic
cardiomyopathy provided that no old septal
infarction is present. Subaortic endocardial
thickening resulting from contact between the
anterior cusp of the mitral valve and the septum is
always a valuable confirmatory feature in cases of
HOCM with outflow obstruction. Cardiomyo-
pathy of the congestive form is not, in my experience
associated to any obvious degree with sudden death
without a prior long period of left ventricular
failure.

Deposition of amyloid can cause sudden death
particularly when advanced in degree and involving
the conduction system. Care must be taken by the
pathologist not to overinterpret the finding of
amyloid in the heart. A high proportion of elderly
patients at necropsy have nodules of a substance
staining as myloid in the left atrium. This appears
to have little functional effect other than a tendency
to be associated with atrial fibrillation.

Acute myocarditis of all forms is associated with
sudden death; commonly there is a history of some
days’ malaise, fever, and tachycardia or palpitation.
More rarely the patient drops dead as the presenting
feature. It is usually possible to suspect the diagnosis
macroscopically; the myocardium is mottled, the
left ventricle dilated but with no cardiomegaly.
Pericarditis is usually present in viral myocarditis.
Idiopathic giant cell myocarditis has serpiginous
areas of myocardial necrosis easily seen with the
naked eye. Myocardial sarcoidosis is associated
with obvious scarring, particularly in the septum,
and is easily confirmed histologically.

Once again caution must be used to avoid over-
diagnosis of myocarditis at necropsy. Isolated foci
of lymphocytes in the atrial myocardium, while
strictly termed myocarditis on morphological
grounds, are common in all elderly hearts and may
be erroneously related to death by pathologists
wishing to find a cardiac abnormality. In cases of
death actually caused by myocarditis virtually every
histological block from the ventricular muscle, and
often from the conduction system itself, is involved.

Sudden death may occur in patients with floppy
mitral valves, yet without severe mitral regurgita-
tion. The frequency of the valve abnormality in the
population is of the order of 5 per cent42 so the risk
of death to any individual patient with a floppy
mitral valve must be very small. It is my experience
as a pathologist that occasional patients with a mild
to moderate floppy valve are indeed found without
other ascertainable cause of death. On the very few
occasions that we have been able to obtain a
previous electrocardiogram it has been abnormal
with inversion of T waves in the inferolateral leads.
It seems likely that, as shown previously,43 this
pattern defines a subset of patients with floppy

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**Table 4 Sudden death**

<table>
<thead>
<tr>
<th></th>
<th>Undiagnosed HOCM</th>
<th>HOCM diagnosed symptomatic</th>
<th>HOCM coincidental finding</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>41 (85%)</td>
<td>10 (62.5%)</td>
<td>6 (50%)</td>
</tr>
<tr>
<td>Female</td>
<td>7</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>Heart weight</td>
<td>478 ± 136</td>
<td>526 ± 123</td>
<td>470 ± 95</td>
</tr>
<tr>
<td>S/P ratio</td>
<td>1.8 (1.4 - 2.6)</td>
<td>1.9 (1.6 - 2.3)</td>
<td>2.2 (1.7 - 2.3)</td>
</tr>
</tbody>
</table>

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**Table 5 Septal/posterior wall ratios—necropsy hearts**

<table>
<thead>
<tr>
<th></th>
<th>Non-cardiac deaths normal weight hearts (n = 108)</th>
<th>Cardiac deaths LV hypertrophy (n = 37)</th>
<th>HOCM (n = 73)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1.02 ± 0.24</td>
<td>1.10 ± 0.24</td>
<td>1.9 ± 0.86</td>
</tr>
</tbody>
</table>

Only 4/3 HOCM hearts had ratio < 1.5
valves who have a tendency to sudden death. Patients with a mild floppy valve and a normal electrocardiogram probably have no such risk.

The mechanism underlying these electrocardiographic abnormalities is debatable.\textsuperscript{44-46} It has been ascribed to be associated with a primary muscle abnormality, mechanical traction on papillary muscles, endocardial impact with the valve, anomalous coronary arteries, and interference with left circumflex flow. In my experience I have been unable to demonstrate any morphological abnormality in the left ventricle. If this is a subtle abnormality of myocardial repolarisation it is uncertain if the association with a floppy mitral valve is fortuitous or linked by a close chromosomal position of two genes.

Anomalous coronary artery anatomy\textsuperscript{47} may be perfectly benign or produce serious functional effects. Where the abnormality is a simple one of both coronary orifices arising from the same sinus or a single orifice there is no risk of sudden death. The commonest form is to find a single or two orifices in the right coronary sinus. The left anterior descending coronary artery crosses in front of the right ventricular outflow. A personal study of 2240 hearts disclosed five such cases, all asymptomatic and dying of non-cardiac disease.

Sudden death is a risk either when a segment of the coronary artery tree is aplastic or when there is a coronary shunt. Most frequently the former is a single right coronary orifice with the left anterior descending artery passing behind the pulmonary artery and being represented as a fibrous strand without a lumen. Coronary shunts occur with a fistula from an artery into the ventricles, atria, or coronary sinus. Aneurysmal dilatation of the coronary artery involved ensues and myocardial perfusion becomes abnormal. Anomalous origin of one coronary artery from the pulmonary artery also leads to an aortic-pulmonary shunt.

Coronary embolism is a cause of sudden death. Emboli occur most frequently from aortic valve thrombus as in bacterial endocarditis. The aortic Starr-type prosthetic valve with its peripheral flow into the sinuses seems almost to direct emboli into the coronary orifices. Syphilitic ostial stenosis as a postulated cause of sudden death\textsuperscript{48} has virtually vanished.

Small myxomatous polyps on the aortic valve may prolapse into a coronary orifice orifice but are extremely rare.\textsuperscript{49} Isolated dissection of coronary arteries occurs occasionally to produce sudden death in Marfan’s syndrome and also in pregnancy.\textsuperscript{50} Coronary arteritis occurs in polyarteritis nodosa\textsuperscript{52} and sudden death is well described.\textsuperscript{19} In Japan a striking syndrome of lymphadenopathy, skin rash, conjunctivitis, and fever in young children with a high risk of sudden death from coronary arteritis is relatively common but sporadic cases are now known to occur both in the United Kingdom and the United States.\textsuperscript{53}

When all other known causes of sudden death have been excluded macroscopical examination of the area of the conduction system is, on occasion, helpful. The small benign mesothelial tumour of the atrioventricular node\textsuperscript{44} \textsuperscript{48} is usually visible as a 1 to 2 cm cystic mass in the atrial septum anterior to the coronary sinus. Calcific masses originating from the anterior cusp of the mitral or the aortic valve may also transect the area of the main bifurcating atrioventricular bundle.

It is when the pathologist is faced with a heart apparently totally normal to external examination that practical problems arise.

The majority of pathologists have a very liberal interpretation of the term cardiomyopathy and, to the clinicians’ confusion, apply it indiscriminately for such cases. Fully developed congestive cardiomyopathy (CCM) does have recognisable morphological abnormalities. At macroscopical level these include raised ventricular muscle weights for both ventricles, dilatation of the ventricular cavities leading to a normal or even reduced wall thickness, mural thrombus in all chambers, and diffuse endocardial thickening in the left ventricle. Microscopical examination of the myocardium shows interstitial fibrosis, reduced muscle fibre diameters with a wide scatter in size, and vacuolated fibres with hyperchromatic large nuclei. It would clarify semantics if pathologists would confine the term congestive cardiomyopathy to patients known to have a degree of left ventricular failure and whose heart showed at least three of the morphological abnormalities listed. As stated above, my own experience is that such hearts are rarely causes of instantaneous sudden death.

A number of conditions deserving better recognition is also squeezed into a “cardiomyopathy” group by pathologists. Isolated increase in the heart weight is better termed idiopathic cardiomegaly. Most examples are probably an excessive hypertrophy response to unrecorded hypertension and are particularly seen in patients of West Indian origin.

Widespread interstitial or focal myocardial fibrosis without other morphological abnormality is best termed idiopathic myocardial scarring and may well be post-viral myocarditis. This group of patients, clinically, may be associated with arrhythmic problems entirely without evidence of abnormal myocardial contractile function.

Macroscopically, normal hearts are also en-
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countered in which a selective and progressive loss of conduction fibres occurs followed by replacement fibrosis. These hearts are again not associated with evidence of loss of contractile junction but develop arrhythmias and conduction defects. No name is available for this entity but “conduction myopathy” might be appropriate.

Those hearts which comprise the group where there is no macroscopical abnormality and no microscopical abnormality of the contractile myo-cardium do pose a major problem for the pathologist. In practice, many of these patients are simply transferred to the ischaemic or cardiomyopathy groups since a great deal of further effort and public cost are thus avoided. Hearts from patients under the age of 50, and particularly under 30, are those more extensively studied.

The initial step is to exclude once again unnatural death, in particular a concealed suicide. Blood should be screened for drugs by gas chromatography. After these steps have been carried out death can be presumed to be cardiac in origin. A detailed medical history from the family is sought and any electrocardiogram ever taken reviewed. The conduction system is examined histologically.

Personal experience of studying such patients, suggests that the majority of deaths are indeed cardiac. My experience is of necessity selected by pathologists who know of my interest. In 57 hearts referred for investigation of sudden death without macroscopical abnormality and a normal myo-cardium, 25 proved to have had previous electrocardiograms, all recorded in the investigation of palpitation or syncopal attacks of varying severity and frequency. The electrocardiographic abnormalities are shown in Table 6. Several of these, such as pre-excitation, have a well-defined risk of sudden death and with careful morphological study anomalous pathways, often multiple, can be found.56

The long QT interval syndrome has a similar well-recognised clinical association with sudden death but a histologically normal conduction system. James et al.58 however, record nerve fibre pathology in this group.

Patients with ectopic beats and supraventricular tachycardias universally proved to have a morphologically normal conduction system, and, clearly, the pathophysiological abnormality is beyond conventional histology. This group must, however, be pathologically distinguished from those patients with widespread myocardial scarring who have a similar clinical picture and can be postulated to represent a post-viral myocarditis aetiology.

Patients in whom the electrocardiogram showed conduction defects can be shown to have morphological abnormality. Destruction of the atrio-ventricular node is characteristic of the small mesothelioma which may be no more than 0.3 cm across and escape macroscopical examination.

The Lenegre and Lev forms of idiopathic bundle-branch fibrosis which characteristically lead to complete atrioventricular block in patients over 50 and often over 65 years of age are associated with steadily progressive electrocardiographic changes over many years.60-62 At the stage where right bundle-branch block and left axis deviation is present indicating bilateral bundle-branch damage sudden death occurs but the exact risk and detection of those at risk is debated.63-64

Congenital or familial conduction defects at bundle-branch level may be relatively stable and not progress. These appear to represent aplasia or extreme hypoplasia of one segment, usually the proximal right branch. The essential pathology is therefore akin to congenital atrioventricular block.65 Such conditions appear to have a relatively low risk of sudden death.66 In contrast, patients and families in whom the electrocardiographic abnormality is progressive are at risk of sudden death.67-70 Published reports and the four cases in Table 6 all show widespread loss of conduction fibres throughout the distal conduction system. It is to this group that the term “conduction myopathy” might be applied. It remains uncertain whether this abnormality in patients under 50 years of age is identical to the Lenegre form of bundle-branch fibrosis which produces complete atrioventricular block in older patients. It is probable that they are simply different ends of a clinical and age spectrum.

Consideration of the clinical history of those patients without a previous electrocardiogram suggests (Table 7) that many of these would have been abnormal. Retrospective questioning of relatives of 25 of 37 (68%) disclosed suspicious symptoms. In retrospect, one case was likely to be the clinical syndrome of undue catecholamine sensitivity.71

Table 6 Twenty-five patients without macroscopical abnormality of heart

<table>
<thead>
<tr>
<th>Electrocardiographic abnormality</th>
<th>No.</th>
<th>Conduction system morphology</th>
</tr>
</thead>
<tbody>
<tr>
<td>All &lt; 50 years of age obtained</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pre-excitation</td>
<td>7</td>
<td>Anomalous pathways</td>
</tr>
<tr>
<td>Long QT interval</td>
<td>1</td>
<td>Normal</td>
</tr>
<tr>
<td>Episodic supraventricular tachycardia</td>
<td>2</td>
<td>Normal</td>
</tr>
<tr>
<td>Multifocal ventricular/atrial ectopies</td>
<td>3</td>
<td>Normal</td>
</tr>
<tr>
<td>Some ventricular ectopic beats</td>
<td>7</td>
<td>Normal</td>
</tr>
<tr>
<td>AV block (partial)</td>
<td>1</td>
<td>Fibrosis and hypoplasia of bundle-branches</td>
</tr>
<tr>
<td>Bundle-branch block (partial or complete)</td>
<td>4</td>
<td></td>
</tr>
</tbody>
</table>
Table 7 Thirty-seven patients dying suddenly—no electrocardiographic data available. No macroscopical cause apparent—myocardium normal (ages 5 to 49)

<table>
<thead>
<tr>
<th>Clinical data</th>
<th>No.</th>
<th>Conduction system morphology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Palpitation alone</td>
<td>14</td>
<td>Mesothelioma AV node</td>
</tr>
<tr>
<td>Palpitation and syncopal or fainting attacks</td>
<td>4</td>
<td>Anomalous conduction path (Mahaim type)</td>
</tr>
<tr>
<td>Syncopal attacks alone</td>
<td>3</td>
<td>Absent right bundle-branch</td>
</tr>
<tr>
<td>Family history of sudden death</td>
<td>3</td>
<td>Hypoplasia main atrioventricular bundle or bundle-branches</td>
</tr>
<tr>
<td>No complaints known</td>
<td>12</td>
<td>Fibrosis bundle-branches and distal conduction fibres</td>
</tr>
<tr>
<td>Sibs with long QT</td>
<td>1</td>
<td>Normal</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

As a basic principle, examination of the conduction system where no electrocardiographic data are available is seldom scientifically profitable. The incidence of morphological abnormality in this group is, however, high, and closely mirrors that found in the patients with electrocardiographic data.

Mesotheliomata when present are clearly the cause of death. Where distal conduction fibre loss has occurred this again is likely to be significant. Mahaim tracts are of dubious significance since they are relatively common in children who die accidentally. Where the conduction system is morphologically normal no conclusions can even be attempted. It must be admitted that cases will occur where no firm cause of death can be ever ascertained.

There is strong pressure on the pathologist undertaking these studies to find a lesion and numerous individual cases of sudden death associated with minor morphological abnormalities become recorded.19 72 73 Before these can be scientifically established as cause and effect many more morphological studies must be made of patients in whom electrocardiographic and particularly 24-hour monitoring has been recorded. The forensic necropsy service in England and Wales, even so, is not set up to recover such hearts and, while its sole responsibility is to exclude unnatural death, will not do so.

I am grateful to numerous pathologists who send me material, without whom this work would be impossible. Dr A Leatham and Dr D Krikler have interpreted many of the electrocardiograms with great patience. Detailed coronary assessment was carried out by Dr A Popple as part of an MD thesis.

References

Pathological view of sudden cardiac death


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