The floppy mitral valve

Study of incidence, pathology, and complications in surgical, necropsy, and forensic material

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SUMMARY In the study of 1984 routine hospital necropsies the mitral valve was examined from the left atrium in the intact heart with a pressure head of water in the left ventricle. The valve was graded from grade O (normal) to grade 4: grade 1, expansion of a small part of one cusp only; grade 2, over one-third of the posterior cusp or one-half of the anterior cusp expanded, with intact chordae; grade 3, ruptured chordae; grade 4, chordal fusion to ventricular wall. The frequency of grade 2 to 4 floppy valves rose with age with an overall incidence of 3-9 per cent in men and 5-2 per cent in women. Grade 1 floppy valves have no clinical significance. Grade 2 floppy valves were found to be associated with auscultatory signs but often only contributed to cardiac failure or were coincidental findings. Grade 3 and 4 floppy valves were direct causes of death from bacterial endocarditis and/or severe mitral regurgitation.

The surgical series of floppy valves showed that chordal rupture was the event which most commonly made operation necessary in middle age: in a minority this was caused by bacterial endocarditis. Dilatation of the annulus was an important contributory factor but can produce significant mitral regurgitation without chordal rupture, particularly in inherited connective tissue disorders such as the Marfan syndrome.

Forensic necropsies confirm that sudden death occurs in patients with floppy valves. The majority have grade 3 or 4 floppy valves and presumably significant mitral regurgitation. A minority have minimal valve involvement and the mechanism of death is unexplained.

The exact magnitude of the risk for any patient with a floppy valve of developing bacterial endocarditis, or chordal rupture leading to significant mitral regurgitation, or of dying suddenly, is not known but must be very low considering the frequency of the valve lesion.

The essential pathology of the floppy mitral valve is weakening of the central fibrous core allowing cusp expansion and chordal elongation to occur. The weakness of the collagen is in part genetically determined, in part age related. Identical changes occur in the tricuspid valve, and in the aortic root, leading to aortic regurgitation. Similar pathological changes are well recognised in other mammals, particularly the aged dog.

The best name has not yet been decided for the entity known as the floppy valve (Read et al., 1965), redundant cusp syndrome (Hill et al., 1974), billowing sail deformity (Oka and Angrist, 1961; Bittar and Sosa, 1968), or balloononing mitral cusp (Behar et al., 1967). These names reflect the expansion of mitral cusp area and elongation of chordae which allow prolapse of the cusp into the atrium in ventricular systole, eventually resulting in mitral regurgitation. The terminology used by pathologists stresses the degenerative nature of the disease: the terms myxomatous (Read et al., 1965; Aslam et al., 1970), myxoid (Kern and Tucker, 1972), and mucinous degeneration (Frable, 1969) of the mitral valve are all in use. Further difficulty arises with the number of clinical terms in use for mitral valve prolapse (Abrams, 1976), including
The floppy mitral valve

mid-systolic click–late systolic murmur syndrome (Barlow et al., 1963) and prolapsed mitral cusp (Cirley et al., 1966). The floppy valve is probably the major cause of this syndrome, but in an unknown proportion of cases the syndrome is the result of ischaemic heart disease or cardiomyopathy (Aranda et al., 1975; Jeresaty, 1975; Nutter et al., 1975; Verani et al., 1976).

The diverse names and the preoccupation of groups of workers with selected aspects of their material have hindered the elucidation of an overall picture of the natural history of the disease. In the present study, we have attempted to present a composite view by examining material from medical, surgical, morbid anatomical, and forensic sources.

Material and methods

Material for study was available from three separate and unrelated sources. Forty-four floppy valves excised at operation for mitral regurgitation were obtained from two cardiothoracic units. A survey of 1984 consecutive routine necropsies carried out during the same period in four general hospitals in London provided 90 further examples of floppy mitral valve of moderate to severe degree. Over a period of 5 years, particular search for cases of sudden, unexpected death with a floppy valve was initiated by Professor R. D. Teare, who provided 13 examples from forensic necropsies performed for H.M. Coroner.

(a) CLINICAL FEATURES OF SURGICAL CASES

Of the 44 cases available for study, 31 were men and 13 women. The age range was wide, from 32 to 69 years (mean 51-4 years). Murmurs had been present for many years in some cases, the longest being 48 years, but 23 per cent had had a murmur for less than 1 year and 48 per cent for less than 5 years (Table 1). Symptoms had been present for days only or up to 18 years before operation, with a mean of 4-1 years. Four patients only were definitely known to have had a late systolic murmur or click; the remainder were recorded as having a pansystolic murmur when first seen by the clinician. Four patients had had proven bacterial endocarditis; 3 additional cases had had a presumptive diagnosis of bacterial infection of the valve. Three patients had skeletal stigmata of Marfan’s syndrome; 1 patient had osteogenesis imperfecta. Two patients were known to have had atrial septal defects of the secundum type.

(b) CLINICAL FEATURES OF HOSPITAL NECROPSY CASES

The incidence of all grades of the floppy valve is shown in relation to age and sex in the 1984 necropsies in Table 2. The relation between causes of death, age, sex, and clinical features of the 90 cases of significant degrees of the floppy valve is shown in Table 3. Floppy valves were considered to be the direct cause of death when present as the only discernible cause of left ventricular failure or when death resulted from bacterial endocarditis. Floppy valves were regarded as a possible contributory cause of death when associated with left ventricular failure, but with additional ischaemic heart disease, chronic bronchitis, or systemic hypertension. Deaths from, for example, carcinoma of the bronchus were recorded as having no relation to the floppy valve.

Table 1 Surgical series: length of known history of cardiac murmur

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<td>No.</td>
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Table 2 Routine necropsy series: sex and age incidence of all grades of floppy valve

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<tr>
<th>Men: Age at death (y)</th>
<th>&lt;40</th>
<th>40–49</th>
<th>50–59</th>
<th>60–69</th>
<th>70–79</th>
<th>80–89</th>
<th>90+</th>
<th>All ages</th>
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<tr>
<td>Total no. of necropsies</td>
<td>88</td>
<td>108</td>
<td>154</td>
<td>323</td>
<td>320</td>
<td>102</td>
<td>16</td>
<td>1111</td>
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<tr>
<td>Grade of valve abnormality</td>
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<td>0</td>
<td>8</td>
<td>8</td>
<td>18</td>
<td>20</td>
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<table>
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<tr>
<th>Women: Age at death (y)</th>
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<th>40–49</th>
<th>50–59</th>
<th>60–69</th>
<th>70–79</th>
<th>80–89</th>
<th>90+</th>
<th>All ages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total no. of necropsies</td>
<td>67</td>
<td>85</td>
<td>106</td>
<td>171</td>
<td>269</td>
<td>134</td>
<td>41</td>
<td>873</td>
</tr>
<tr>
<td>Grade of valve abnormality</td>
<td>1</td>
<td>2</td>
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<td>3</td>
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<td>0</td>
<td>0</td>
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<td>0</td>
</tr>
</tbody>
</table>
Clinical data extracted from the patients' notes included age, sex, blood pressure readings, presence and nature of any murmur, and the presence of cardiac failure. The causes of death as found at necropsy were recorded. Family history was not sought other than that recorded in the clinical notes: previous medical history was taken as recorded in the clinical notes.

(c) FORENSIC NECROPSY MATERIAL
The material has been exclusively drawn from necropsies carried out for H.M. Coroner and all were examples of sudden unexpected death, within minutes, of patients who had not seen their medical practitioner for some weeks. The search was only undertaken to determine if any appreciable number of patients with floppy valves died suddenly without other apparent cause. Cases of floppy valves with a clear and definite alternative cause of death were not included in this study. No analysis of the exact number of cases of sudden death in the population from which they are taken was possible, but only 13 examples of the floppy valve were encountered in 5 years, compared with an average of 250 cases of ischaemic heart disease with sudden death annually. All cases were screened for the presence of drugs. The histories of forensic necropsy cases were obtained from the Coroner's Officer and where possible the patients' general practitioners. Of the 13 cases, 5 had no history of any cardiac disease; 8 were known to have had cardiac murmurs for periods of 4 to 25 years, and 4 of these had in the past been treated for cardiac failure. The

Pathological methods

(a) SURGICAL MATERIAL
At the time of operation the surgeon specifically looked for and recorded the incidence of chordal rupture. The size of the mitral annulus was roughly assessed by inspection at operation and by the size of prosthetic valve (Starr or Björk-Shiley) inserted. The use of a number 4 Starr mitral valve was taken to indicate a dilated mitral annulus. After excision each valve was pinned flat and photographed (Fig. 1). Material for routine microscopy was

![Fig. 1 Surgically excised floppy valve pinned flat for examination. Anterior cusp shows chordal elongation with expansion of free margin of the cusp. Posterior cusp shows expansion and doming of whole cusp with ruptured chordae (arrow).](http://heart.bmj.com/br-heart-j-first-published-as-10.1136/hrt.40.5.468-on-1-may-1978. Downloaded from http://heart.bmj.com on September 17, 2023 by guest. Protected by copyright.)
fixed in 10 per cent formol saline, for estimation of acid mucopolysaccharides in acetone, and for electron microscopy in glutaraldehyde. Sections for light microscopy were stained by the elastic-Van Gieson technique to show elastic and collagen. Sections were examined for connective tissue mucopolysaccharides by staining with alcian blue/PAS at pH 3-0 (Mowry and Winkler, 1956). Material fixed for electron microscopy was post-fixed in osmium tetroxide for one hour before routine embedding and cutting. Thin sections were examined in the Zeiss EM9S electron microscope.

(b) NECROPSY MATERIAL
Before opening the ventricle, the mitral valve was inspected from the left atrium and ruptured chordae noted. The left ventricle was filled with water via a rubber tube introduced through the aorta, and the mitral valve closed either by squeezing the ventricle by hand or by tying off the aorta and applying high pressure from a mains water tap; inspection of the closed mitral valve from the left atrium then allows easy identification of cusp prolapse (Fig. 2, 3, 4). After opening the left ventricle, the mitral ring circumference was measured in centimetres. In all 1984 hearts the anterior and posterior cusps of the mitral valve were graded as follows: grade 0, within normal limits; grade 1, expansion of only a small portion of one cusp; grade 2, at least one-third of the posterior cusp or one-half of the anterior cusp involved, but with intact chordae (Fig. 5); grade 3, floppy valves complicated by chordal rupture (Fig. 6); grade 4, chordae fused to the ventricular wall (Fig. 7). Changes in the tricuspid valve were similarly noted. Subsequently the heart was dissected, isolated left and right ventricular weights were measured, and the presence of any additional pathology in the heart was noted. After routine fixation in formol saline, all valves were examined histologically in sections stained by the elastic-Van Gieson technique, alcian

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Fig. 2 Normal mitral valve fixed by pressure in closed position and viewed from left atrium. The three scallops of the posterior cusp are visible (L, lateral; C, central; M, medial). There is no cusp prolapse.

Fig. 3 Examination of mitral valve in unfixed intact heart. Valve viewed from left atrium with water at mains pressure in left ventricle. Posterior cusp prolapses into the atrium and water streams under its free edge. (Man aged 80. Discharged from Army in 1916 at age 20 with murmur which persisted. Death from carcinoma.)

Fig. 4 Same heart as Fig. 3, fixed with mitral valve in closed position to show prolapsed middle segment of posterior cusp with ruptured chordae.
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blue/PAS at pH 3.0, and haematoxylin and eosin.

Results

Pathological appearances

The macroscopical appearances of the floppy valve are characteristic and, with practice, easily recognised in both surgical and necropsy material. In the latter, recognition of a significant abnormality was best made from the left atrium before opening the ventricle. Indeed the best definition of the floppy valve is sufficient elongation of chordae and expansion of cusp area to allow prolapse into the atria on applying a pressure load to close the valve. Ruptured chordae can also be noted without risk of artefactual cutting.

The valve cusp was increased in area and in both longitudinal and transverse dimensions leading to

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Fig. 5  Grade 2 floppy valve at necropsy. One half only of anterior cusp is expanded and ballooned (arrows). Posterior cusp is unaffected.

Fig. 6  Grade 3 floppy valve at necropsy. Middle segment of posterior cusp is domed into left atrium with several chordae ruptured (arrow). An atrial jet lesion is present (J). Clinical diagnosis was significant mitral regurgitation.

Fig. 7  Grade 4 floppy valve at necropsy. Chordae of middle segment of posterior cusp fused into fibrous mass adherent to ventricular endocardium (arrows).
folding and convolution (Fig. 8) and upward doming of the cusp towards the atrium. Partial involvement of the posterior cusp involved the central (Fig. 4) or medial scallop, more rarely than the lateral scallop alone. Partial involvement of the anterior cusp was more common at the medial commissure (Fig. 5). More severe changes involved the whole anterior or posterior cusp or both. Chordal elongation, tortuosity, and thinning were usually present. Chordal rupture (grade 3) led to an accentuation of upward doming of the cusp. Ruptured chordae appeared filiform or, with time, became blunt-ended or folded back to adhere to the cusp surface. Chordae with areas of thinning, presumably preceding rupture, were seen (Fig. 9). Focal areas of thickening had developed on the mural ventricular endocardium in most cases of grade 2 or 3 floppy valves; these occasionally enlarge sufficiently to entrap and fix chordae (grade 4) (Fig. 7), as recognised by Salazar and Edwards (1970). Bacterial endocarditis was also a cause of chordal rupture and cusp destruction (Fig. 10, 11).

In some instances the valve appeared translucent and gelatinous, but in the majority very considerable secondary fibrosis had occurred on the atrial aspect of the cusp making the valve white and opaque; in the past this has caused misinterpretation and erroneous diagnosis of chronic rheumatic disease (Fig. 8). Elongated chordae also undergo secondary fibrosis with considerable thickening (Fig. 8), again simulating chronic rheumatic disease. The mitral ring circumference was normal, moderately,
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ischaemic damage (4.4%). Isolated left ventricular weights showed a considerable range in the 90 cases with floppy mitral valves. Forty-seven were below the upper limit of normal for our laboratory (185 g). Forty-five lay between 185 and 300 g, and 8 were in excess of 300 g. No histological evidence was found for coexistent cardiomyopathy though the 6 cases with dilated mitral annuli had isolated left ventricular weights over 300 g and a large left ventricular cavity.

In the surgical series mitral valves known to be derived from patients with other stigmata of Marfan's syndrome showed the same general range of macroscopical appearances as other floppy valves, with the exception that translucent rather fragile valves were more common in young subjects.

**MICROSCOPICAL FINDINGS**

The normal valve cusp has a dense collagenous central core (valve fibrosa) continuous with the valve ring and extending into the chordae. This is covered on both atrial and ventricular aspects by a thin layer of loose connective tissue and finally by endothelium. On the ventricular surface of the cusps at the sites of chordal insertion the superficial zone (ventricularis) of connective tissue is normally loose and myxomatous, i.e. contains stainable acid mucopolysaccharide (Gross and Kugel, 1931).

or grossly increased. The ring circumferences ranged from 6 cm to 13.6 cm, with a mean of 8.0 cm. The mitral ring circumference in hearts without any valve abnormality in the necropsy series was 5.6 to 10.3 cm (mean 7.8 cm). Of the 90 cases of floppy mitral valve (grade 2 and above) in the hospital necropsy series, 6 had a mitral ring circumference over 10.3 cm (6.7%). Valve calcification was virtually never present, only being seen if coexistent senile mitral ring calcification was present (3 cases, 3.3%). The papillary muscles were normal.

In the necropsy series overall 6.1 per cent of all hearts showed an area of ischaemic scarring in excess of 2 cm in size. Of the 90 hearts with significant floppy valves, 4 only had coexistent myocardial

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In contrast in the floppy valve the collagen of the central valve fibrosa is abnormal. Large areas show apparent loss of fibrous tissue with individual collagen bundles fragmented, coiled, and disrupted. Maximal destruction occurs around sites of chordal insertion but the changes may extend into the body of the cusp. Areas of destruction of collagen usually show easily demonstrable pools of acid mucopolysaccharide in which residual strands of collagen lie. These histological changes must not be confused with the rather loose myxomatous connective tissue normally found in the ventricularis zone near the chordae. Histological examination must be made of sections passing exactly through the long axis of the valve where the topography can be seen and the fibrosa identified. Over the atrial surface of the cusp a superficial zone of dense laminated collagen forms which often contrasts with the ‘moth eaten’ fibrosa. Within this zone of superficial dense fibrosis small areas of fibrin deposition occur on the surface, and the lesion as a whole results from friction of the excessively mobile cusps. Ultrastructural study does not do more than confirm the fragmentation of collagen and occurrence of pools of connective tissue mucin. We have not observed abnormal periodicity in collagen fibrils.

In the absence of a history of bacterial endocarditis the valve cusps are not vascularised nor do they contain chronic inflammatory cells. Both features enable histological distinction to be made from chronic rheumatic valve disease. A further point of distinction is the absence of calcification in the floppy mitral valve.

Small isolated areas of collagen destruction even within the fibrosa are very common in grade 1 floppy valves in otherwise normal hearts. The diagnosis of significant degree of floppy valve is made by naked eye observation of sufficient expansion of cusp or chordae to allow prolapse under pressure and microscopy to show major dissolution of the valve fibrosa in the prolapsed segment. Random histological sections without detailed examination of the intact fresh heart are useless.

**CLINICAL CORRELATION WITH MORBID ANATOMICAL FINDINGS**

(a) **Surgical material**

The onset of symptoms ranged from abrupt, requiring operation within a month, to gradual increase of disability over years. The mean period between onset of symptoms and operation was 4-1 years. The incidence of chordal rupture was high (77%) with the posterior cusp most commonly involved (Table 4). Of the 44 patients, 24 required a No. 4 valve prosthesis indicating a large mitral annulus. Only 1 patient had chordal fusion to the posterior cusp at operation.

(b) **Hospital necropsy material**

In the hospital necropsy series trivial degrees of the floppy mitral valve syndrome (grade 1) were very common. Review of the clinical records showed that only 15 per cent of these patients had had any

<table>
<thead>
<tr>
<th>Table 4 Surgical series: incidence of ruptured chordae</th>
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<tbody>
<tr>
<td>Anterior cusp</td>
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<td>---------------</td>
</tr>
<tr>
<td>No. of cases</td>
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Fig. 13 Histological section (×276) of the valve fibrosa stained by elastic Van Gieson technique (left, normal; right, floppy valve). The abnormal fibrosa has less densely packed collagen with areas of loose myxomatous tissue; collagen bundles appear broken and coiled.
murmur and the lesion was not clinically significant. Grade 2 to 4 floppy mitral valves are potentially more serious and were found in 3-9 per cent of all men and 5-2 per cent of all women (Table 2). In 80-1 per cent of these patients, murmurs had been recorded. The incidence of floppy valves rose with age to a maximum of 5-5 per cent in the eighth decade in men, after which it apparently fell. In women there was a steady rise in incidence to a maximum of 10-0 per cent over the age of 90.

The possible significance of the grade 2 floppy mitral valves in 67 patients without chordal rupture could be judged by the cause of death. Thirty-seven (55-2%) died of non-cardiac disease, 24 (35-8%) died of cardiac failure, but with additional causes including hypertension, chronic bronchitis, and ischaemic heart disease also present. Only 6 patients of the 67 (9-0%) died of cardiac failure only attributable to the floppy mitral valve. In these 6 patients the mitral ring circumference ranged from 10 cm to 13-5 cm, with a mean of 11-5 cm.

The significance of grade 3 and 4 floppy valves was more clear cut. In 22 of the 23 patients, loud pansystolic murmurs were recorded in the clinical notes; there was no record of heart sounds in the other case. Nine of the patients (39%) died directly from bacterial endocarditis or cardiac failure from mitral regurgitation because of the floppy valve; in 12 patients (52%) death was from cardiac failure but with coexistent hypertension, ischaemic heart disease, or chronic bronchitis. In only 2 patients (9%) was death the result of malignant disease and unrelated to the floppy valve.

The distribution of cusp expansion in grade 2 floppy valves and the anatomical sites of chordal rupture in grade 3 to 4 floppy valves are shown in Table 5. The posterior cusp is most commonly involved, with posterior chordae the most common site of rupture, though both cusps or, rarely, the anterior cusp alone can be affected.

Table 5 Necropsy series: pathological features

<table>
<thead>
<tr>
<th>Grade 2 (66 cases)</th>
<th>Grade 3 to 4 (24 cases)</th>
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<tr>
<td>Expansion of cusps</td>
<td>Ruptured chordae</td>
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<tr>
<td>Anterior</td>
<td>Posterior</td>
</tr>
<tr>
<td>Both</td>
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</tr>
<tr>
<td>3</td>
<td>44</td>
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<tr>
<td>19</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>5</td>
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</table>

*Associated with bacterial endocarditis in 5.

In many patients with grade 2 to 4 floppy mitral valves, the tricuspid valve also showed expansion of the cusp area with prolapse into the right atrium. The change maximally involved the anterior cusp. Of 67 patients with grade 2 mitral valves, 45 (67%) had an equivalent degree of abnormality in the tricuspid valve. The 23 patients with grade 3 to 4 mitral valves showed a higher incidence, with at least one cusp of the tricuspid involved in 18 (78%). One patient had ruptured chordae to the anterior cusp of the tricuspid valve. There was no clinical evidence to suggest that the tricuspid abnormalities were of any clinical significance.

(c) Forensic necropsy material

The 13 examples of unexpected death which could be ascribed to the floppy mitral valve had pathological findings listed in Table 6. Cases with ruptured chordae or bacterial endocarditis were likely to have had significant mitral regurgitation and their sudden death was, therefore, explicable. Four cases remained, 3 in young women, in which the mechanism and cause of sudden death was totally obscure. No histological evidence of ischaemic heart disease or cardiomyopathy was present.

Table 6 Forensic necropsy: pathological features

<table>
<thead>
<tr>
<th>Sex</th>
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<th>Ruptured chordae</th>
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<td>Old inferior infarct</td>
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<td>69</td>
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</tr>
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<td>M</td>
<td>79</td>
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<td>F</td>
<td>68</td>
<td>0</td>
<td>Bacterial endocarditis</td>
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Discussion

A generally acceptable name for the 'floppy valve' entity remains to be agreed (Abrams, 1976). The condition is certainly not new; a specimen was mounted in the Pathological Museum of St George's Hospital in 1936 under the name 'parachute deformity of the mitral valve'. A local tradition has existed for the use of this name, but it does lead to confusion with the congenital anomaly of chordal insertion into one papillary muscle (Shone et al., 1963) and has had to be suppressed. The term 'floppy valve' is gaining acceptance, though 'redundant cusp' is more accurate. The pathological names such as myxoid or myxomatous degeneration are less popular, stressing as they do only minor morphological features of the disease. The term 'blue valve syndrome' (McCarthy and Wolf, 1970), stressing the accumulation of acid mucopolysaccharides as shown by alcian blue stains, has also not proved popular. The term prolapse of
The floppy mitral valve

The mitral cusp is purely descriptive in relation to echocardiograms or angiograms and may result from the cusp and chordal abnormality of the floppy valve, but also occurs with ischaemic papillary muscle damage (Aranda et al., 1975; Nutter et al., 1975) or in cardiomyopathy (Liedtke et al., 1973). The frequency of the lesion shown in this necropsy series is, however, so close to that of mitral valve prolapse in echocardiographic studies (Procacci et al., 1976) that floppy valve is likely to be the commonest cause of prolapse.

The floppy valve is essentially the result of a weak cusp fibrosa allowing stretching and expansion to occur. The clinical course suggests that this process may begin very early in life and progress extremely slowly if at all, or begin late in life with more rapid progression over a few years; intermediate varieties are common. The pathogenesis of the weakening of the collagen in the cusp fibrosa remains unknown. In part it is genetically determined; familial cases are well recognised (Hunt and Sloman, 1969; Shell et al., 1969; Shappell et al., 1973), and the floppy valve is an integral part of the generalised connective tissue weakness of Marfan's syndrome (McKusick, 1955; Miller and Pearson, 1959; Shankar et al., 1967; Bowers, 1969; Simpson et al., 1969; Murdoch et al., 1972). Other diseases associated with generalised connective tissue defect in which the floppy mitral valve is known to occur include osteogenesis imperfecta (Crisciotti et al., 1965; Wood et al., 1973), the Ehlers-Danlos syndrome (McKusick, 1966; Brandt et al., 1975), and pseudoxanthoma elasticum (Huang et al., 1967). The occurrence of the floppy valve in Marfan's syndrome has led to the suggestion that all cases are hereditary and, in the absence of other stigmata, are 'formes frustes' of the fully developed disease (Read et al., 1965). No familial trend emerged in our series, with the exception of one patient in the forensic necropsy series with a cousin in the surgical series. The usual family history taken for clinical notes is, however, not exhaustive; moreover, the condition has not been adequately diagnosed until recently.

The incidence of a floppy mitral valve at necropsy tends to rise with age; there is work (Bashey et al., 1967) to suggest that the collagen content of the human mitral valve falls with age. The evolution of the floppy mitral valve may, therefore, in part be a process analogous to the change seen in the skin with increasing age. An identical process occurs in aged dogs leading to mitral regurgitation (Pomerance and Whitney, 1970) and is known to occur in other mammals in advanced age (Whitney, 1975). Pomerance (1969) has previously recorded the high incidence of floppy valves in human geriatric necropsies.

While the necropsy data tend to suggest the lesion increases in severity with age, individual clinical case histories illustrate how benign and chronic the condition may be. The reported increased incidence with age in necropsy series could partly result from the greater ease of recognition in long-standing cases when secondary cusp fibrosis has occurred.

No other pathological process has been implicated and the valves are not vascularised, inflamed, or replaced by dense fibrous tissue as in chronic rheumatic disease. It is not clear whether the weakness of the fibroa in the floppy valve is caused by weakness of the individual collagen fibres or by a decrease in total collagen. The morphological findings tend to support the former explanation and histological appearance of the floppy valve with its destruction of the fibroa is easily distinguishable from that of chronic rheumatic disease with its increased collagen. The superficial fibrosis over the atrial surface of the cusp and on the chordae as a result of friction does lead to a superficial resemblance to chronic rheumatic disease. Rheumatic disease, however, leads to cusp retraction not expansion.

If the total collagen content of the valve fibroa is diminished, it is not known if synthesis is diminished or lysis is increased. Morphological similarity between the focal areas of collagen loss in affected chordae and the action of elastase on collagen in vitro lends some support to a 'lytic' theory (Selzer et al., 1967; Caulfield et al., 1969; Sanders et al., 1971). It is probable that accumulation of connective tissue mucins is secondary to the abnormality of collagen and elastic synthesis or breakdown (Shappell et al., 1973) and no abnormal mucopolysaccharides is found (Sherman et al., 1970). This point is of some importance since accumulation of abnormal mucopolysaccharides within valve tissue in Hurler's syndrome (gargoylism) (Berenson and Geer, 1963) may lead to disruption of the fibroa, with a picture closely analogous to the floppy valve. Our own ultrastructural studies and those of others (Kern and Tucker, 1972) have confirmed the haphazard arrangement of collagen fibrils, with disruption or fragmentation, but the characteristic periodicity of the individual fibrils remains normal.

Our own work and that of others (Sanders et al., 1971; McKay and Yacoub, 1973) supports the view that degeneration of collagen within the central core of the chordae is responsible for chordal rupture, though it is claimed that increased chordal tension resulting from the enlarged area of the cusp may play a contributory role. In a small proportion of cases, chordal rupture results from superimposed bacterial endocarditis, which may occur in
the floppy valve syndrome (Lachman et al., 1975), and is well shown in the necropsy study by Pom-
erance (1969), in the present necropsy and surgical series, and in a prospective study of patients with
the late systolic murmur syndrome (Allen et al., 1974). Infection begins on the atrial surfaces of the
cusps in floppy valves where incorporation of fibrin is often seen as a secondary friction lesion.

Our hospital necropsy series shows that minor
degrees (grade 1) of floppy valve are very common
(Table 3) but are of no clinical significance. More
severe grades (2, 3) of floppy valve are potentially
more serious and were found in up to 7 per cent of
older individuals, though this figure is higher than
that recorded by Pomérance (1969) in a necropsy
series. Our necropsy series suggests that the in-
cidence rises with age and is a little higher in
women, though our surgical series is predominantly
male. Any hospital series is selected, contains a dis-
proportionate number of older patients, and does
not accurately reflect the incidence in the young or
very old. Our necropsy data are in keeping with a
number of recent echocardiographic studies (Proc-
cacci et al., 1976). In addition, the necropsy study
shows that the floppy valve has a spectrum of
severity from partial to whole cusp involvement.
This spectrum could well explain the very different
incidences reported in different echocardiographic
studies, in some of which partial involvement of the
posterior cusp may not have been detected. Our
necropsy study supports the view that floppy valves
of some degree are present in between 5 and 18 per
cent of the population (Brown et al., 1975;
Markiewicz et al., 1975; Procacci et al., 1976). The
necropsy data do not support the great preponder-
ance of women recorded in these echo studies,
though the incidence was higher in young and old
women than in men. Only one echo study (Higgins
et al., 1976) has so far described the common
occurrence of valve prolapse in middle-aged men,
but clearly echo studies are needed to confirm
or refute the necropsy evidence of a rise in incidence
with age.

The actual clinical significance of floppy valves
found at necropsy must be questioned. In the
hospital necropsy series, moderately severe floppy
valves, but without chordal rupture, were directly
responsible for death from cardiac failure as a result
of mitral regurgitation only in a small minority,
could have contributed to cardiac failure in about
one third of patients, and had no clinical significance in
over half of patients (Table 3). In contrast, more
severely affected floppy valves with chordal rupture
caus ed death directly in 39 per cent of patients and
contributed to cardiac failure in 52 per cent.

Dilatation of the mitral annulus in association with
the floppy valve will potentiate mitral regurgitation
(Bulkeley and Roberts, 1975) but was only found in a
small minority of cases in our necropsy series. Our
necropsy data do not show any difference in the
occurrence of ischaemic (coronary artery) disease
between those cases with and without floppy valves.
We have not confirmed the association of absence
of the left circumflex artery with floppy valve
(Gentzler et al., 1975).

The surgical series shows an age range somewhat
below that of the necropsy cases. Analysis of the
reasons for their clinical presentation with severe
mitral regurgitation at a younger age suggests that
chordal rupture is the common precipitating event,
but that bacterial endocarditis and annular dilata-
tion can also be responsible. The last is particularly
associated with generalised connective tissue dis-
orders such as Marfan's syndrome. Review of the
surgical cases shows the onset of symptoms to be
either gradual or sudden. Some patients were
known to have had systolic murmurs for up to 47
years, but the majority had significant mitral regur-
gitation when first seen by the surgeon. The
striking feature is that only 10 per cent were known
to have had late systolic murmurs or clicks before
their first presentation, though this may reflect
merely lack of recording or recognition at previous
routine medical examination. In a prospective
study of patients with late systolic murmurs
(Allen et al., 1974), only one has so far had to have
valve replacement for chordal rupture.

Sudden unexpected death has been increasingly
reported in the late systolic murmur syndrome
(Hancock and Cohn, 1966; Shell et al., 1969;
Jeresaty, 1976) and in some cases necropsy has
shown a floppy valve (Shappell et al., 1973). On
the other hand, a prospective study of patients with
isolated late systolic murmurs showed no increased
risk of sudden death (Allen et al., 1974). This
divergence of view may reflect the fact that a late
systolic murmur has a number of causes. In perhaps
a majority the cause is a floppy valve and some
evidence for this view comes with the association
of the late systolic murmur with minor skeletal
abnormalities (Rizzon et al., 1973; Salomon et al.,
1975). Other causes of a late systolic murmur
include ischaemic papillary muscle damage, primary
myocardial disease, and rheumatic mitral valve
disease (Epstein and Coulshed, 1973; Jeresaty,
1975), and it is possible that the recorded sudden
deaths came from patients with these conditions
rather than those with floppy valves. In the present
hospital necropsy series, half of the patients with
severe degrees of floppy valve died suddenly, but
only one of the moderately severe patients did.
This, however, merely reflects the fact that patients with severe mitral regurgitation may die suddenly.

In the United Kingdom sudden death is likely to result in a necropsy outside the hospital by forensic pathologists working for H.M. Coroner. These cases would not appear in our hospital necropsy series. Specific search has, therefore, been made over the same period for grade 2 to 4 floppy valves thought to be the direct and only cause of sudden death. In forensic practice only 13 cases emerged, the majority with significant mitral regurgitation as a result of chordal rupture. There were, however, 4 cases, 3 in young women, with floppy valves but no chordal rupture, where no other cause for death was found. Thus, it seems likely that sudden death can be a complication of the floppy mitral valve with only minimal regurgitation, but the risk to an individual patient is very low. The mechanism remains to be elucidated, but atrial and ventricular arrhythmias are well known in patients with mitral valve prolapse (Winkle et al., 1975; Ritchie et al., 1976).

A number of cardiac abnormalities are known to be linked with the floppy valve. The link with Marfan's syndrome is the best recognised, the two conditions overlapping in cases of floppy valves with minor skeletal abnormalities (Rizzon et al., 1973). Some authors have recorded an association of aortic regurgitation with floppy mitral valves and suggested the existence of 'floppy' aortic cusps (McKay and Yacoub, 1973). The mechanism of aortic closure and competence is, however, entirely different from that of the mitral valve. Simple expansion of the aortic cusp area will not lead to regurgitation and we have no morphological proof that a floppy aortic valve occurs. We have 3 cases of floppy mitral valve with aortic regurgitation in our necropsy series; all had increased aortic root diameters and the link between aortic regurgitation and floppy mitral valves lies in an associated weakness of the connective tissues of the aortic root and of the mitral valve fibrosa. Two patients in the surgical series had mild aortic regurgitation at the time of their mitral operation without overt stigmata of Marfan's syndrome.

Atrial septal defects of the secundum variety may be associated with late systolic murmurs and clicks (McDonald et al., 1971; Littler et al., 1973) or with prolapse of the mitral cusp (Betriu et al., 1975; Leachman et al., 1976), and occasionally mitral regurgitation develops. The mitral valve abnormality may be rheumatic or a secondary haemodynamic effect (Okada et al., 1969), but we have seen two typical floppy valves associated with atrial septal defect in the surgical series and the association is well recorded (Hynes et al., 1974; Betriu et al., 1975).

While the present study identifies the risks and complications of the floppy valve, it does not and cannot accurately assess the magnitude of each risk. If it is assumed that the floppy valve is the most common cause of an isolated late systolic murmur, the work of Allen et al. (1974) suggests that the risk of each complication is very small. The most contentious point is the risk of sudden death in patients with minimal mitral regurgitation from a floppy valve without coronary artery or myocardial disease. Enough cases have been seen in forensic practice for the diagnosis to be acceptable in the Coroner's court as a cause of sudden death, but only the cases followed in great clinical detail, as by Shappell and his colleagues (1973), will provide scientific proof of the association.

There is a real need for a large-scale, long-term study of patients shown by echocardiography to have prolapse of a mitral cusp. Unfortunately only necropsy may actually show whether the cause is a floppy mitral valve, ischaemic papillary muscle, or cardiomyopathy, and provide information about the risk of sudden death in each group. Thus, determination of the actual incidence of each complication in relation to each cause of mitral cusp prolapse might require life-long follow-up and necropsy study.

References


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