Abnormal subendocardial function in restrictive left ventricular disease

Michael Y Henein, Derek G Gibson

Abstract

Objective—To study possible disturbances in left ventricular long axis function in patients with a restrictive filling pattern.

Design—Prospective examination of the left ventricular transverse and longitudinal axes, transmitral flow, and the apexcardiogram.

Setting—A tertiary referral centre for cardiac diseases.

Subjects—21 normal subjects, age (SD) 51(11); 30 patients of similar age with a restrictive left ventricular filling pattern, defined as short early diastolic deceleration time less than the lower 95% confidence limit of the normal value (120 ms). 20 patients had a normal and 10 had an increased left ventricular end diastolic cavity size.

Results—Mitrail Doppler echocardiography: E wave velocity was high only in patients with a normal cavity size. A wave velocity was greatly reduced in the two groups (P < 0.001) so that the E/A ratio was abnormally high. The relative A wave amplitude on the apexcardiogram was greatly increased in the two groups: 46(15)% (mean (SD)) and 54(4)% v 15(5)%. Minor axis: Fractional shortening was reduced from 30(10)% to 17(7)% in patients with normal cavity size and to 13(4)% in those with a dilated cavity (P < 0.001), as was the posterior wall thickening fraction from 100(30)% to 42(20)% and 50(25)% respectively (P < 0.001). Total systolic epicardial motion was normal and isovolumic relaxation time was short in the two groups. Long axis: Left ventricular abnormalities included reduced total amplitude of motion and its component during atrial systole (P < 0.001 for the two groups at both sites). Peak long axis shortening and lengthening were decreased at both left ventricular sites (P < 0.001). The time intervals from q wave of the electrocardiogram and A2 (aortic valve closure) to the onset of shortening and lengthening respectively were increased (both P < 0.001). Right ventricular long axis function was similarly affected but to a lesser extent.

Conclusion—Left ventricular long axis function is consistently abnormal in patients with restrictive disease whether or not cavity size is increased. Not only are the extent and peak velocity of shortening reduced, but during diastole the peak early diastolic lengthening rate and amplitude during atrial systole are impaired. Early diastolic long axis motion is asynchronous with respect to transmitial flow and left ventricular minor axis. These effects will impair the overall left ventricular systolic and diastolic function independently of any decrease in passive cavity compliance. Unlike fibrosis, these long axis abnormalities are potentially amenable to treatment.

Ventricular restriction is usually considered to be a disorder of the endocardium or myocardium which spares systolic function, but which reduces passive ventricular compliance enough to compromise filling. One of its most prominent features is endomyocardial fibrosis, which is specifically recognised as occurring in restrictive cardiomyopathy in the World Health Organization and International Society and Federation of Cardiology definition and classification of the cardiomyopathies. The aim of this study was to investigate the possibility that this endomyocardial fibrosis might impair left ventricular subendocardial function in such patients. This would have consequences in both systole and diastole, which should be separable from a simple reduction in passive compliance, and which might contribute significantly to the overall pathophysiology of the disorder.

Patients and methods

Patients

We studied 30 patients with a restrictive left ventricular filling pattern, in whom a clinical diagnosis of heart failure had been made. Their mean (SD) age was 55(13) years, and five were women. A restrictive filling pattern was defined as an early diastolic mitral flow deceleration time shorter than the lower 95% confidence limit of the normal value, 120 ms, as measured by pulsed Doppler echocardiography. Patients with echocardiographic features of amyloid were deliberately excluded. No patient had Loeffler's disease, Churg-Strauss syndrome, hypereosinophilia, endomyocardial fibrosis, or haemochromatosis. Patients were divided into two groups according to their left ventricular end diastolic dimension obtained from the M mode record-

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ing of the left ventricular minor axis. Twenty patients had a left ventricular cavity size less than the upper 95% confidence limit of normal, 5.8 cm; two had had aortic valve homograft replacement; one had aortic valve stenosis; coronary artery disease was present in two; hypertension in two; and the other patients (13) had a diagnosis of congestive heart failure of unknown cause. The remaining 10 patients had a large ventricular cavity, with an end diastolic dimension of 6 cm or more; six had a documented history of ischaemic heart disease and the rest had unexplained congestive cardiac failure. Two patients had a family history of sudden cardiac death. None of the patients had evidence of pericardial disease. Eight patients were in atrial fibrillation and the rest in sinus rhythm. The results from these patients were compared with those from 21 normal controls of mean (SD) age 51(11) years, none of whom had any history of cardiac disease, hypertension, or diabetes.

METHODS

Mitral flow velocities were recorded from the apical four chamber view with pulsed Doppler echocardiography with either a Hewlett-Packard or a Doptek system with a 2 MHz transducer, and were recorded with frequency shift calibration. M mode and cross sectional echocardiograms were recorded using a Hewlett-Packard model 77020 A Sonos 1000 echograph with a 2.5 MHz transducer for the minor and long axes with the patient lying in the left lateral position. M mode echocardiograms of the left ventricular minor axis were recorded from the long axis parasternal view with the cursor at the tips of mitral valve leaflets. Long axis measurements were taken from the apical four chamber view with the cursor at the left, septal, and right sites of the atrioventricular rings.

Electrocardiograms were recorded using a Hewlett-Packard 12 lead electrocardiograph with a filter of 0.05-100 Hz. The PR interval and QRS duration were computed using built in software.

Apexcardiograms were obtained with a Cambridge pulse transducer (time constant four seconds). This was possible in 17 patients from the position of a well defined palpbale apex. All recordings were made separately on a Hewlett-Packard or a Honeywell (Echoline 22) strip chart recorder at a paper speed of 100 mm/s with a simultaneous electrocardiogram and phonocardiogram. In patients with atrial fibrillation measurements were taken from beats with a cycle length equal within 50 ms. Minor and long axis echocardiograms were digitised and analysed.5

MEASUREMENTS

All values are based on the mean (SD) of three beats.

DOPPLER

From the mitral pulsed Doppler trace we measured early and late diastolic peak flow velocities and derived the E/A ratio in patients with well identified E and A waves. The early diastolic accelerations time was taken from the onset of the E wave to its peak, and the deceleration time from the peak to its end. We measured time intervals from A1 of the phonocardiogram to the onset and to the peak of the E wave.

M MODE

From the left ventricular minor axis traces we measured end diastolic and end systolic dimensions at the onset of the q wave and A1 (the first high frequency component of the second heart sound) respectively, and thus derived fractional shortening. The amplitude of epicardial motion as well as the change in posterior wall thickness between end diastole and end systole were measured, and the thickening fraction was derived as the increase in thickness divided by end diastolic thickness, expressed as a percentage. The isovolumic relaxation time was taken as the interval between A1 and the onset of mitral cusp separation in early diastole. The peak rate of minor axis dimension change was derived by digitisation.3 The left atrial diameter was measured from the M mode trace of the aortic root.

From the original long axis traces we measured the following at the left, septal, and right sites: the time intervals from the q wave of the electrocardiogram to the onset of long axis shortening and from A1 of the phonocardiogram to the onset of long axis lengthening, as well as the overall amplitude of excursion and its component in late diastole due to atrial contraction.6 Peak rates of shortening and lengthening were derived by digitisation.3

On the apexcardiogram, the A wave excursion was measured as a percentage of the total amplitude of pulse wave excursion.

From the electrocardiogram recorded by a Hewlett-Packard machine the PR interval and QRS duration were measured from the built in software.

STATISTICAL ANALYSIS

The tables give individual mean (SD) values. We used an unpaired t test to compare mean values from normal subjects with those from patients in the two groups and also to compare the two patient groups.

Results

Tables 1-4 show the results obtained.

DOPPLER ECHOCARDIOGRAM

The mitral E wave deceleration time was less than 120 ms, the lower 95% confidence limit of normal, in all 30 patients, by definition, whether or not cavity size was increased (table 1). Its mean (SD) value was 95(15) ms and 95(10) ms in the two groups respectively compared with the normal value of 160(20) ms. The peak E wave velocity was normal in patients with a dilated left ventricular cavity and increased in those with normal end diastolic cavity size. Peak A wave velocity was
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Table 1  Doppler and apexcardiogram characteristics. Values are mean (SD)

<table>
<thead>
<tr>
<th>Variables</th>
<th>Normal (n = 20)</th>
<th>Dilated (n = 10)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peak E wave (m/s)</td>
<td>0.7 (0.1)</td>
<td>0.9 (0.16)*</td>
</tr>
<tr>
<td>Acceleration time (ms)</td>
<td>75 (10)</td>
<td>50 (30)*</td>
</tr>
<tr>
<td>Deceleration time (ms)</td>
<td>160 (20)</td>
<td>95 (15)*</td>
</tr>
<tr>
<td>E/A ratio</td>
<td>0.5 (0.1)</td>
<td>0.14 (0.1)*</td>
</tr>
<tr>
<td>A2 to onset of E wave (ms)</td>
<td>85 (15)</td>
<td>70 (20)†</td>
</tr>
<tr>
<td>Apexcardiogram</td>
<td>165 (30)</td>
<td>130 (30)*</td>
</tr>
<tr>
<td>Percentage A wave to total</td>
<td>15 (5)</td>
<td>54 (4)*</td>
</tr>
</tbody>
</table>

*P < 0.001 v controls.
†P < 0.01 v controls.
§Normal size restricted v dilated restricted.
¶P < 0.05.

Fig. 1  (A) Apexcardiogram from a patient with restrictive left ventricular filling. (B) Mitral flow pattern from the same patient. Note the large A wave pressure increase in the apexcardiogram corresponding to the small Doppler A wave. ECG, Electrocardiogram; and PCG, phonocardiogram.

The PR interval and QRS duration on the electrocardiogram were normal in the two groups of patients.

APEXCARDIOGRAM
In patients with sinus rhythm, the relative A wave excursion was considerably higher than normal in the two groups (table 1), particularly in patients with a normal end diastolic cavity dimension (fig 1).

M. MODE
Total epicardial motion of the minor axis (table 2) was normal in the two groups but fractional shortening was equally reduced whether or not the end diastolic dimension was increased; the same applied for posterior wall thickening. The isovolumic relaxation time was shorter than normal in the two groups. Peak shortening and lengthening rates were significantly reduced only in patients with normal end diastolic cavity size. The left atrium was dilated in the two groups.

Long axis function (tables 3 and 4) was similarly affected in the two groups. The total amplitude of motion and its late diastolic atrial component were significantly reduced at all sites with the exception of the A wave on the right side, which remained normal in patients with a normal end diastolic dimension. Peak long axis shortening and lengthening rates were also decreased at all sites, again with the exception of the right side, which was normal in patients with dilated ventricles. The time intervals from the q wave of the electrocardiogram to the onset of long axis shortening and from A2 of the phonocardiogram to the onset of lengthening were abnormally increased in the two groups and at all sites including the right side.

In normal subjects, the minor axis starts to lengthen before blood flow can be detected across the mitral valve. The long axis behaves similarly, starting to lengthen 27(7) ms and 25(5) ms at the left and septal sites respectively before the earliest detectable mitral flow. In the two patient groups, however, the onset of long axis lengthening followed that of early diastolic flow by 15–35 ms and 5–15 ms (P < 0.001) respectively at the two sites (fig 2).

Discussion
Although the diagnosis of restrictive left ventricular disease has traditionally been made invasively,2 mitral deceleration time is consistently reduced in this disorder to less than the lower 95% confidence limit of normal.7 We used this criterion to identify 20 such patients who originally presented with clinical evidence of heart failure. We excluded those with amyloid disease, as these have been previously studied in detail.7 As reported elsewhere,4 the most common clinical association was with...
coronary artery disease and mild ventricular hypertrophy, though in a substantial proportion, of patients no obvious underlying cause was found, so these patients could be described as having restrictive cardiomyopathy. We compared these patients with uncomplicated restriction with a second group of 10 patients with a restrictive filling pattern, but in whom the left ventricle was enlarged at end diastole. These patients would not therefore have fitted the orthodox invasive criteria of restriction, though the Doppler approach, being independent of cavity size, allowed them to be recognised.

Left ventricular systolic function was consistently abnormal in uncomplicated restriction even when the end diastolic cavity size was normal. In particular, the decrease in the minor axis during ejection was reduced, so that the shortening fraction was consistently low. Minor axis decrease during ejection because of the combined effects of inward motion of the epicardium and an increase in myocardial thickness, particularly that of the posterior wall. In our patients, the epicardium moved normally during systole, but the extent of posterior wall thickening was consistently reduced.

If the decrease in the minor axis was due entirely to the shortening of circumferentially arranged myocardial fibres, systolic wall thickening would be exactly determined by the inward motion of epicardium, as the myocardium is incompressible. The extent of thickening in normal subjects is too great to be explained simply by circumferential fibre shortening; it is necessary to invoke the additional contribution of obliquely or longitudinally directed fibres. The combination of normal epicardial motion and reduced thickening thus strongly suggests that such longitudinal shortening, and hence fattening, is selectively lost in patients with restriction. We confirmed this by showing directly that the extent and peak velocity of longitudinal shortening were reduced. Further, the onset and the termination of long axis shortening were delayed with respect to that of the minor axis. Motion in the two axes could not, therefore, have simply been due to transverse and longitudinal components of homogenous obliquely arranged myocardium, but must have been caused by anatomically separate fibres. As most longitudinally directed fibres are subendocardial, we suggest that the characteristic systolic abnormality in uncomplicated restriction is the direct effect of selectively impaired subendocardial function.

Table 2 Minor axis dimensions and velocities. Values are mean (SD)

<table>
<thead>
<tr>
<th>Variables</th>
<th>Normal</th>
<th>Restricted left ventricular filling</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Normal size (n = 20)</td>
<td>Dilated (n = 10)</td>
</tr>
<tr>
<td>Minor axis</td>
<td>End systolic dimension (cm)</td>
<td>3.3 (0.5)</td>
</tr>
<tr>
<td></td>
<td>Fractional shortening (%)</td>
<td>30 (10)</td>
</tr>
<tr>
<td></td>
<td>PW thickening fraction (%)</td>
<td>100 (30)</td>
</tr>
<tr>
<td></td>
<td>Systolic epicardial motion (cm)</td>
<td>0.4 (0.1)</td>
</tr>
<tr>
<td></td>
<td>IVR time (ms)</td>
<td>55 (10)</td>
</tr>
<tr>
<td></td>
<td>IVR dimension change (cm)</td>
<td>0.13 (0.1)</td>
</tr>
</tbody>
</table>

PW, Posterior wall; IVR, isovolumic relaxation.
*P < 0.001 vs normal controls.
†P < 0.01 vs normal controls.

Table 3 Long axis excursions and velocities. Values are mean (SD)

<table>
<thead>
<tr>
<th>Variables: long axis</th>
<th>Restricted left ventricular filling</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Normal size (n = 20)</td>
</tr>
<tr>
<td>Total excursion (cm)</td>
<td>Left</td>
</tr>
<tr>
<td></td>
<td>Septal</td>
</tr>
<tr>
<td></td>
<td>Right</td>
</tr>
<tr>
<td>Atrial excursion (cm)</td>
<td>Left</td>
</tr>
<tr>
<td></td>
<td>Septal</td>
</tr>
<tr>
<td></td>
<td>Right</td>
</tr>
<tr>
<td>Peak shortening rate (cm/s)</td>
<td>Left</td>
</tr>
<tr>
<td></td>
<td>Septal</td>
</tr>
<tr>
<td></td>
<td>Right</td>
</tr>
<tr>
<td>Peak shortening rate (cm/s)</td>
<td>Left</td>
</tr>
<tr>
<td></td>
<td>Septal</td>
</tr>
<tr>
<td></td>
<td>Right</td>
</tr>
</tbody>
</table>

*P < 0.001 vs normal controls.
†P < 0.05 vs normal controls.

Table 4 Long axis timing. Values are mean (SD)

<table>
<thead>
<tr>
<th>Variables</th>
<th>Restricted left ventricular filling</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Normal size (n = 20)</td>
</tr>
<tr>
<td>Q to onset of shortening (ms)</td>
<td>Left</td>
</tr>
<tr>
<td></td>
<td>Septal</td>
</tr>
<tr>
<td></td>
<td>Right</td>
</tr>
<tr>
<td>A2 to onset of shortening (ms)</td>
<td>Left</td>
</tr>
<tr>
<td></td>
<td>Septal</td>
</tr>
<tr>
<td></td>
<td>Right</td>
</tr>
</tbody>
</table>

*P < 0.001 vs normal controls.
†P < 0.01 vs normal controls.
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Figure 2  (A) Mitral Doppler recording from a normal subject (left) and a patient with restrictive filling (right). Vertical scale: one division represents 20 cm/s.  
(B) Minor axis recording.  
(C) Long axis recording from the left and septal sites. Vertical scale in centimetres. Note that lengthening normally starts before the onset of transmisral flow (left), but in the patient its onset delayed, particularly at the septal site. Vertical line corresponds to A2; broken line to onset of early diastolic flow. ECG, Electrocardiogram; PCG, phonocardiogram; and A, aortic valve closure.

axis (fig 1) shows increased passive stiffness of the left ventricular myocardium in the longitudinal direction.

The disturbances of long axis function in the patients in whom cavity size was increased at end diastole were similar to those in whom it was normal. The overall amplitude of long axis shortening was reduced, and time relations were disturbed in exactly the same way as in patients with uncomplicated restriction. The amplitude of epicardial motion was normal, and so were the peak rates of shortening and lengthening of the ventricular minor axis. During atrial systole, a reduced A wave amplitude on the transmisral Doppler and long axes was again combined with an increase in the pressure A wave. In spite of obvious differences in cavity architecture, therefore, similar long axis abnormalities were associated with a similar disturbance to filling in early and late diastole in the two patient groups. The right ventricle also played a part, though to a lesser extent than the left. The proportional impairment in the overall amplitude of motion and, in particular, of shortening and lengthening velocities, were not as obvious, whereas values for atrial systole were often close to or within the normal range.

We thus found direct and indirect evidence for depressed longitudinal function during systole and diastole, particularly on the left side of the heart in patients with a restrictive filling pattern. This was not the effect of abnormal ventricular activation as the QRS duration was normal. The disturbances in the timing of movement were similar to those in patients with coronary artery disease, suggesting an ischaemic basis. Some of our patients did have large vessel coronary artery disease, but even in its absence, increased ventricular diastolic pressures predispose to subendocardial ischaemia. This ischaemia
may also underlie the fibrosis which is such a characteristic feature of restrictive ventricular disease.

We believe that these long axis abnormalities may play a part in restrictive disease. Impaired longitudinal shortening is largely responsible for the reduced shortening fraction that was consistently seen in these patients. A major feature of normal early diastolic filling is the change in cavity shape towards a more spherical configuration as the volume increases. As a sphere has the maximum volume per unit surface area, such a shape change reduces the need for myocardial distension. The left ventricle becomes less spherical at end systole because of preferential shortening of the minor axis. As myocardial thickening was reduced by impaired long axis function, the reduced long axis shortening will have reduced the shape change, thus increasing the proportion of stroke volume to be accommodated by stretching of the abnormal ventricular wall during the succeeding diastole. Delayed onset of long axis relaxation implies myocardial tension persisting into the period of early diastolic filling, thus increasing myocardial stiffness by a mechanism independent of any change in its passive properties. Reduced changes in cavity shape will add to the effects of the abnormal myocardium in reducing cavity compliance, and may explain it completely in those patients in whom myocardial histology is normal.

Disturbances of early diastolic filling are usually accompanied by an increase in the proportion of the stroke volume entering the ventricle with atrial systole. The reduced extent of longitudinal lengthening, however, the dominant mechanism by which ventricular cavity size increases during atrial systole, impairs this compensating mechanism. Loss of the atrial contribution means that filling must take place almost exclusively in early diastole in spite of major relaxation and geometrical abnormalities. This can be achieved only at the expense of a high left atrial pressure, reflected in our patients by a short isovolumic relaxation time. Further evidence that impaired long axis function may directly affect early diastole is provided by the effects of intravenous milrinone in patients with cavity dilatation and a restrictive filling pattern. Its administration causes an increase in the amplitude of mitral ring motion, which is accompanied by increased wall thickening, along with prolongation of isovolumic relaxation and return of the filling pattern towards normal.

We conclude, therefore, that far from being an incidental finding in restriction, abnormal long axis function is common and contributes significantly to the physiological disturbances so characteristic of the disorder. Although established myocardial fibrosis is probably inaccessible to treatment, long axis abnormalities may be at least potentially reversible, as shown by the effect of milrinone, thus opening an avenue of treatment in these difficult cases.

5 Upton MT, Gibson DG. The study of left ventricular function from digitized echocardiograms. Prog Cardiovasc Dis 1978;20:359-44.
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