Mitral valve closure and left ventricular filling time in patients with VDD pacemakers

Assessment of the onset of left ventricular systole and the end of diastole

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SUMMARY  The effect of mitral valve closure on left ventricular filling time and its relation to the onset of systole were assessed from mitral valve echocardiograms and simultaneous apex cardiograms in 21 normal subjects, 11 patients with left bundle branch block, and 19 patients with VDD pacemakers programmed for atrioventricular intervals of 50, 150, and 250 ms. The interval between the electrocardiograph Q wave and the apex cardiogram upstroke was similar in normal subjects and patients with left bundle branch block, but was significantly longer in patients with VDD pacemakers at all atrioventricular intervals. Similarly there was little difference in the time interval between the Q wave and mitral valve closure in normal individuals and patients with left bundle branch block but this was considerably delayed in VDD pacemaker patients with the atrioventricular interval set at 50 ms. With increasing atrioventricular intervals the mitral valve closed significantly earlier, whereas the onset of left ventricular systole and the timing of mitral valve opening remained unchanged. Thus as a result of earlier mitral valve closure left ventricular filling time decreased progressively as the atrioventricular interval was increased. Since the onset of left ventricular systole, with respect to left ventricular stimulation, is considerably delayed in VDD pacemaker patients a short atrioventricular interval is required in these patients to maintain the normal time relations between atrial and ventricular contraction and hence maximise left ventricular filling.

Left ventricular filling provides the blood volume for subsequent left ventricular ejection. One of its important determinants is left ventricular filling time, which is determined by the mitral valve movement. This injection time depends not only on cardiac cycle length but also on the mechanisms which time mitral valve closure. The timing of mitral valve closure can be altered by electrical delay of ventricular depolarisation and in atrioventricular conduction disturbances. These two conditions coexist in patients with dual chamber pacemakers since ventricular depolarisation begins from an abnormal focus and is delayed and the PR interval can be altered by programming the pacemaker. Since the relative timing of atrial and ventricular contraction can be altered simply by reprogramming the pacemaker these patients offer a unique opportunity to study the effect of these events on mitral valve movements and left ventricular filling time.

This study was designed to examine non-invasively the relations between electrical and mechanical events in patients with VDD pacemakers over a wide range of atrioventricular intervals. The movements of the mitral valve and hence left ventricular filling time were detected by M mode echocardiography as was aortic valve opening which times the onset of left ventricular ejection. The onset of mechanical systole was detected by the upstroke

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of an apex cardiogram and electrocardiogram. A group of normal controls was studied. Finally patients with left bundle branch block were also studied to examine the possible effects of conduction disturbance of a type similar to that seen in right ventricular pacing.

This information is of theoretical interest because of the insight that it gives into the mechanisms underlying mitral valve motion and particularly mitral valve closure. It is also of considerable practical importance. Patients with myocardial impairment who receive dual chamber pacemakers supposedly benefit from the improvement in left ventricular diastolic function produced by coordinating atrial and ventricular contraction. This study attempts to establish the atrioventricular intervals that provide the longest left ventricular filling time and thus should maximise the clinical benefits of this type of pacing.

Patients and methods

We studied 51 patients who had been informed about the nature of the study and had consented to take part in it. These patients were divided into three groups:

Control group—There were 21 controls (12 men, 9 women; age range 24–57 years). Cardiac disease was excluded in these subjects by a normal history, physical examination, electrocardiogram, and chest x-ray.

Left bundle branch block—There were 11 patients (seven men, four women; age range 39–70 years). They had normal PR intervals but left bundle branch block in association with the following diseases: hypertension (2), hypertrophic cardiomyopathy (3), congestive cardiomyopathy (3), ischaemic heart disease (1) and aortic valve replacement (2). This group was included so that we could investigate the effects of abnormal ventricular depolarisation of a type similar to that occurring in patients with right ventricular pacing.

Pacemaker group—There were 19 patients (11 men and eight women, age range 21–75 years) in this group. All showed the electrocardiographic pattern of left bundle branch block due to right ventricular stimulation by a dual chamber pacemaker. The pacemakers, all programmed in VDD mode, had been implanted for complete heart block (17 patients) or second degree heart block (2 patients) 3–18 months before the study. Ten of these patients had presented with symptoms of left ventricular failure (New York Heart Association grade II–III); three had ischaemic heart disease, and two had undergone aortic valve replacement. In the other nine patients without symptoms of left ventricular failure the aetiology of the heart block was not clear apart from one patient with congenital heart block and another with a history of myocarditis. All patients were studied with their VDD pacemakers programmed for the following atrioventricular intervals: 50, 150, and 250 ms. The protocol began with an atrioventricular interval of 150 ms, and this was followed by the two other intervals in random sequence. All measurements were made on beats in which ventricular pacing was triggered by a P wave after the programmed delay.

Investigations

Echocardiograms to detect mitral and aortic valve movements were obtained from all patients by means of a Hewlett Packard 77020 A ultrasound system with a 3–5 MHz transducer. M mode echocardiograms were recorded both at the level of the aortic valve and the mitral valve at a paper speed of 100 mm/s. Electrocardiograms were recorded simultaneously in all patients. A simultaneous apex cardiogram was recorded in 10 randomly selected patients in each group. The apex cardiogram was recorded because the onset of its upstroke coincides with the onset of left ventricular pressure rise. Thus mitral valve movements and electrocardiographic events could be related to the onset of mechanical systole in these subgroups. Each observation was the mean value of measurements taken from 10 consecutive cardiac cycles.

The following time intervals were measured from the preceding electrocardiographic pacemaker spike or Q wave as appropriate: (a) the onset of the upstroke of the apex cardiogram; (b) mitral valve closure (MC); (c) aortic valve opening (AO); (d) mitral valve opening (MO).

![Fig. 1 Timing (mean (SD)) of upstroke of apex cardiogram, of mitral valve closure, and of aortic valve opening with reference to the Q wave or the pacemaker spike in normal individuals, patients with left bundle branch block, and in patients with VDD pacemakers set at an atrioventricular interval of 50 ms. *p<0.05, ***p<0.001.](http://heart.bmj.com/)
Mitral valve closure and left ventricular filling time in patients with VDD pacemakers

The onset of mitral A wave (A) was timed from the succeeding pacemaker spike or Q wave and the cycle length (R-R interval) was also recorded.

LV filling time (FT, ms) was calculated as follows:

\[ FT = RR - MO + MC \]

and normalised by expressing it as a percentage of cycle length (%FT).

\[ \%FT = \frac{FT}{RR} \times 100 \]

The statistical significance of differences between mean values in the three groups was analysed by the Kruskal Wallis test and by the Mann Witney test. Differences between mean values within the pacemaker group were analysed by the Friedman test and the Wilcoxon rank test. Correlation coefficients were calculated by the least squares method.

Results

Increasing delay of intraventricular depolarisation, its influence on the end of diastole and onset of systole

Figure 1 shows the intervals between the Q wave or pacing spike and the onset of the upstroke of the apex cardiogram, mitral valve closure, and aortic valve opening. In this section the results given were obtained with pacemakers programmed to an atrioventricular interval of 50 ms, from patients with left bundle branch block, and from normal controls. Mean cycle length was the same in all three groups of patients. The upstroke of the apex cardiogram occurred 44 (16) ms (mean (SD)) after the electrocardiographic Q wave in normal subjects and was similarly delayed (52 (18) ms) in patients with left bundle branch block. It was considerably delayed in pacemaker patients, occurring 119 (21) ms (p < 0.001) after the pacing spike. Mitral valve closure was 52 (11) ms after the Q wave in normal individuals and slightly later (65 (20) ms) in patients with bundle branch block (p < 0.05). It was markedly delayed in pacemaker patients being 127 (14) ms after the pacing spike (p < 0.001) compared with both other groups. Aortic valve opening demonstrated a similar pattern in the three groups. It occurred 111 (14) ms after the Q wave in normal subjects, 143 (19) ms after the Q wave in patients with left bundle branch block (p < 0.001), and 193 (20) ms after the pacing spike in pacemaker patients (p < 0.001).

A positive linear correlation was demonstrated between the onset of left ventricular systole (taken as the onset of the apex cardiographic upstroke) and the end of diastole (taken as mitral valve closure) (Fig. 2) in normal individuals, in patients with left bundle branch block, and in VDD pacemaker patients with an atrioventricular interval of 50 ms. The regression equation describing this relation for these patients with increasing delay of left ventricular depolarisation was \( y = 0.9565x + 16; r = 0.93 \).

Effects of increasing atrioventricular intervals on the end of diastole and onset of systole in paced patients

Mitral valve closure occurred significantly earlier as the programmed atrioventricular interval was...

Fig. 2  Correlation between onset of left ventricular systole (taken as timing of the apex cardiographic upstroke) and end of preceding diastole (taken as mitral valve closure) arranged in groups according to the increasing delay of intraventricular conduction—that is normal individuals, patients with left bundle branch block (LBB), and patients with VDD pacemakers set at an atrioventricular interval of 50 ms.

Fig. 3 Timing (mean (SD)) of upstroke of apex cardiogram, of mitral valve closure, and of aortic valve opening in VDD pacemaker patients at atrioventricular intervals of 50, 150, and 250 ms. ***p < 0.001.
Fig. 4  An M mode echocardiogram of the mitral valve and a simultaneous apex cardiogram during VDD stimulation with atrioventricular intervals of 50, 150, and 250 ms. Early mitral valve closure with respect to the pacemaker spike (marked by the vertical line) is present at an atrioventricular interval of 250 ms.

increased (Fig. 3). This is shown in an original recording from a patient with a low heart rate (Fig. 4) and at a faster heart rate even more pronounced changes were found (Fig. 5). As the atrioventricular interval was increased so the scatter of the individual values increased. This is shown by the rising standard deviation in Fig. 3.

In contrast, alterations in the atrioventricular interval had no effect on the timing of the upstroke of the apex cardiogram (mechanical systole), aortic valve opening (the onset of left ventricular ejection) (Fig. 3), or mitral valve opening (the onset of diastole) (Fig. 6).

Left ventricular filling time
Figure 6 shows the sequence of timing of mitral valve movements in all three groups with reference to the Q waves or pacing spikes and also shows cycle...
Mitral valve closure and left ventricular filling time in patients with VDD pacemakers

Fig. 6 Timing (mean (SD)) of mitral valve movements (MC, closure; MO, opening) throughout the cardiac cycle (with the Q wave or pacemaker spike (PS) as reference points) in the group of normal individuals (N), in patients with left bundle branch block (LBB), and in patients with VDD pacemakers set at atrioventricular intervals of 50, 150, and 250 ms.

Fig. 7 Correlation between left ventricular filling time and cycle length in the group of normal individuals, in patients with left bundle branch block (LBB), and in VDD pacemaker patients. The regression lines are given for each group.

Fig. 8 Correlation between left ventricular filling time and cycle length in 19 VDD pacemaker patients at atrioventricular intervals of 50, 150, and 250 ms.

length and the duration of left ventricular filling time. Mean cycle length was non-significantly different in controls, patients with left bundle branch block, and patients with pacemakers set with an atrioventricular interval of 50 ms. With increasing atrioventricular delay there was a small but significant decrease in cycle length (Fig. 6).

In the pacemaker patients left ventricular filling time significantly decreased with increasing atrioventricular intervals. It was 441 (94) ms at the atrioventricular interval of 50 ms, 372 (104) ms (p<0.03) when the interval was 150 ms, and 304 (103) ms (p<0.02 compared with a setting of 150 ms) when the interval was 250 ms. This reduction in left ventricular filling time was due to alterations in the timing of mitral valve closure since the timing of mitral valve opening was unchanged (Fig. 6).

Normalised left ventricular filling time was 53 (6)% in control subjects, somewhat less in patients with left bundle branch block (48 (7)%; p<0.025), and did not differ from normal in pacemaker patients with an atrioventricular interval of 50 ms (50 (6)%). With increasing atrioventricular intervals there was a significant reduction to 45 (9)% (p<0.007) when the atrioventricular delay was 150 ms, and to 38 (10)% (p<0.02 compared with an interval of 150 ms) when the atrioventricular delay was 250 ms; both these values were significantly less than normal (p<0.004 and p<0.001).
individuals was \( y = 0.857x - 292 \) \((r = 0.96)\). With longer atrioventricular intervals, however (Fig. 8), the line was shifted downwards. The equations were \( y = 0.909x - 334 \) for an atrioventricular delay of 50 ms, \( y = 0.892x - 350 \) at a delay of 150 ms, and \( y = 0.767x - 311 \) when the delay was 250 ms; the final value is significantly different from that seen with an interval of 50 ms \((p < 0.001)\).

**Mitral A wave**

The onset of the mitral A wave, signifying the mechanical effect of atrial contraction, occurred 81 (23) ms before the next Q wave in normal controls, 86 (28) ms before the next Q wave in patients with left bundle branch block, and only 5 (17) ms before the pacing spike in pacemaker patients in whom the atrioventricular interval was set at 50 ms \((p < 0.001)\) compared with both controls and left bundle branch block patients. As the atrioventricular interval was increased in the pacemaker patients the onset of the mitral A wave occurred, as expected, at increasingly longer intervals before the pacing spike. These intervals were 76 (24) ms \((p < 0.001)\) when the atrioventricular delay was 150 ms and 148 (38) ms \((p < 0.001)\) when the delay was 250 ms. As the mitral A wave occurred longer before the pacing spike it began to exert an effect on mitral closure not seen with shorter atrioventricular delays when mitral closure occurs as a result of ventricular systole. This resulted in the correlation between onset of left ventricular systole and mitral valve closure being lost when the atrioventricular delay was long.

**Discussion**

Because of the short time available in the cardiac cycle and the large number of events occurring during it, considerable synchronisation between myocardial and valve movement is necessary if the heart is to function efficiently. The study of time intervals in the cardiac cycle has a long history. \(^8\) It has mainly concentrated on systolic events, despite the fact that asynchrony during diastole is of great importance since any impairment in ventricular filling must result in an equivalent impairment or ejection.

In 1921 Wiggers defined the onset of systole as coinciding with the closure of the atrioventricular valves. \(^9\) Subsequent use of phono- and echocardiography has shown many exceptions to this definition. Considerable variation in the timing of mitral valve closure, which means that it does not necessarily coincide with the onset of mechanical systole, has been described in the presence of raised left atrial pressure, \(^10\) \(^11\) intraventricular conduction disturbances, \(^1\) \(^3\) atrioventricular block, \(^4\) \(^6\) and atrial fibrillation. \(^12\) Since mitral valve closure must be the end of left ventricular filling its relation to the onset of systole is of considerable importance.

The study of patients with dual chamber pacemakers is of great interest. Firstly, these patients represent a unique physiological model in which the timing of cardiac events may be studied. The interval between the normal P wave and the ventricular pacing stimulus which it triggers when the pacemaker is in the VDD mode can be altered by reprogramming the pacemaker in the absence of any other physiological change. Therefore the effect of atrial and ventricular contraction on mitral valve closure and its relation to the onset of mechanical systole can be studied over a wide range of atrioventricular intervals. Secondly, study of these patients provides practical information about the atrioventricular interval that allows the longest diastolic filling period and thus the most efficient cardiac function.

**Intraventricular conduction delay and the onset of systole**

The electromechanical delay, detected as the interval between the Q wave and the onset of the upstroke of the apex cardiogram, which coincides with the onset of left ventricular pressure rise, \(^7\) was about 50 ms. This accords with previous reports. \(^13\) In the paced patients we studied, this electromechanical delay lasted considerably longer (about 120 ms) and was the same for all three atrioventricular intervals. This delay may be partly because of slow initial spread of depolarisation from the right ventricular pacing site until it gains access to the faster specialised conducting system. If this is the case some increase in the electromechanical delay might be expected in the group of patients with left bundle branch block. This was not found by apex cardiography but other indicators suggest that a small delay does occur. Firstly, mitral valve closure was slightly delayed and is, as will be clear later, dependent on the onset of systole in these patients. Secondly, aortic valve opening, denoting the onset of left ventricular ejection was also delayed compared with values in normal individuals. Others have found a considerable delay of right ventricular contraction in right bundle branch block but they stated left ventricular contraction is not delayed in left bundle branch block because they claimed to have demonstrated normal timing of mitral valve closure. \(^2\) \(^3\)

Their point of reference, however, was closure of the tricuspid valve. When we reanalysed their results with reference to the electrocardiographic Q wave we found a delay in mitral valve closure that is similar to that observed in the present study.

Despite the differences in the intervals between
Mitral valve closure and left ventricular filling time in patients with VDD pacemakers

The phases of the cardiac cycle are often defined with reference to the electrocardiogram with end diastole being taken as coinciding with the Q wave or pacing spike. By this definition the period of up to 127 ms after the pacing spike would be assigned to systole, despite the fact that the mitral valve is still open. In theory this period is long enough to permit all of rapid left ventricular filling although in practice it is more commonly used for left ventricular filling due to atrial contraction. Similarly such traditional definitions of end diastole may cause considerable inaccuracy in the assessment of end diastolic left ventricular dimension by echocardiographic or scintigraphic techniques since some or all of the atrial contribution to ventricular filling will be neglected because it occurs after the Q wave. To avoid these problems we regard end diastole as occurring at the moment when the mitral valve closes, and the onset of systole as the moment when left ventricular pressure begins to rise.

Effect of various atrioventricular intervals on the end of diastole

Each increase in the atrioventricular interval by 100 ms in the VDD paced patients made the coaptation of the mitral leaflets in relation to the pacing spike occur 50 ms earlier. This cannot be due to the mitral valve being closed earlier by left ventricular contraction, since the timing of the onset of systole in relation to the pacing spike did not change. The most likely explanation is that this was the result of a change in the mechanism causing mitral valve closure as the atrioventricular delay was increased. When the delay is short, closure is effected by ventricular systole, and this mechanism accords with the strong linear relation between the timing of the two events and the delay of 50 ms. At longer atrioventricular intervals atrial contraction may become the most important determinant of mitral closure. This concept is supported by echocardiographic studies of mitral closure in patients with complete heart block in whom wide variations in the interval between atrial and ventricular contraction can be studied. It is suggested that atrial contraction raises left ventricular pressure so that as atrial systole ends there is a reverse pressure gradient which closes the valve unless ventricular systole intervenes to close it first. Therefore the mechanism of closure will vary depending on the atrioventricular interval.

In sinus rhythm a PR interval of more than 200 ms is required before mitral valve closure precedes ventricular contraction (that is as a result of atrial contraction). The situation is different in patients with ventricular pacing. Movement of the valve at various atrioventricular intervals was studied by cinefluorography in a patient with a Starr-Edwards mitral prosthesis, complete heart block, and a ventricular pacemaker. This showed that valve closure preceding ventricular systole occurred at much shorter atrioventricular intervals (> 100 ms). Our study also shows that this effect often occurs when the interval is set at 150 ms. This difference between the interval required for atrial contraction to produce mitral closure in paced patients and those required in sinus rhythm is due to the longer delay in the onset of systole with reference to the electrocardiogram (pacing spike and Q wave respectively) in paced patients. This additional delay, which is approximately 80 ms, means that the functional atrioventricular interval lasts 80 ms longer in paced patients than it does in patients in sinus rhythm with the same electrocardiographic PR interval. Therefore, theoretically, atrioventricular delay settings of 50–100 ms should be sufficient to maintain the normal intervals between atrial and ventricular contraction in paced patients.

These observations are of considerable practical importance. An atrioventricular interval of 150 ms is commonly used when programming VDD pacemakers because this is erroneously thought to correspond to the functional interval in sinus rhythm. Our study, however, shows that at this setting mitral valve closure frequently occurred before the onset of ventricular systole, and thus late diastole was not available for ventricular filling. It is also commonly thought that short atrioventricular intervals lead to the atrial A wave being cut off prematurely by ventricular systole. This study shows that this is not the case. At a setting of 50 ms the onset of the mitral A wave occurred 5 (17) ms before the pacing spike. The delay of ventricular activation until 127 ms after the pacing spike allowed sufficient time for a mitral A wave of normal duration (132 ms) to occur at this setting. This value for the duration of the A wave is almost identical with that obtained from simultaneous left ventricular and left atrial tracings in patients with both normal and abnormal left ventricular function. When the atrium and ventricle are...
paced sequentially, the delay between the atrial pacing spike and atrial activation, which may vary between 30 and 120 ms, should be assessed individually and added to the proposed range of 50–100 ms advocated here for the VDD mode.

The reversed pressure gradient by which atrial contraction closes the mitral valve is determined by the force of atrial contraction. It may be counteracted if atrial contraction is weak, atrial pressure is high, or if blood flow is increased. These additional influences may explain the wide individual scatter in the timing of mitral valve closure at long atrioventricular intervals. There were four patients in whom the timing of leaflet coaptation was unchanged despite increasing intervals. One had poor left ventricular function due to previous anterior myocardial infarction, and there was radiographic and clinical evidence of raised left atrial pressure. The other three were physically fit youths in whom the mitral valve reopened after the A wave. These four patients contribute to the exceptional individual values seen in Fig. 8.

Although in this study we examined only the sequence of haemodynamic events in the left side of the heart, there is no evidence that the sequence in the right heart is essentially different. It is well known that the right atrial P wave occurs about 30 ms before depolarisation of the left atrium. Phonocardiographic and invasive studies of patients with left bundle branch block or right ventricular pacemakers have shown that the onset of contraction of both ventricles is simultaneous. Thus the selection of the short atrioventricular interval, shown in this study to benefit left ventricular filling, is unlikely to affect right ventricular function in an adverse way.

Left ventricular filling time

Left ventricular filling time was significantly shortened in our patients with VDD pacemakers when the atrioventricular delay was set at 150 ms or longer. When the delay was 150 ms filling time was reduced by 60 ms and this reduction increased to 130 ms when the delay was set at 250 ms. Timing of mitral valve opening was the same at all atrioventricular intervals. With an atrioventricular delay of 150 ms left ventricular filling time was reduced by a combination of reduced cycle length and earlier mitral valve closure. With an increase of the programmed delay to 250 ms the further reduction in filling time was entirely due to even earlier mitral valve closure since the cycle length remained constant. The resulting interval between mitral valve closure and the onset of left ventricular systole of up to 100 ms serves neither left ventricular filling nor ejection.

This waste of time may be harmless at a normal heart rate and with a normal left ventricle because up to 80% of the left ventricular filling occurs during the initial rapid filling phase of diastole which lasts about 120 ms and the remaining 20% can easily be completed during diastasis and with atrial contraction. Elderly pacemaker patients, however, may have other cardiac abnormalities such as ischaemic heart disease or left ventricular hypertrophy that lead to disturbed left ventricular diastolic function. Seven of our 19 patients with VDD pacemakers had evidence of such abnormality with delayed mitral valve opening consistent with left ventricular relaxation abnormalities. In these circumstances the amount of left ventricular inflow during the early rapid filling period is reduced to 60% or less and thus the time available for left ventricular filling in late diastole becomes far more important and may determine the left ventricular end diastolic volume and stroke volume.

The wasted time in late diastole with long atrioventricular intervals also becomes important during exercise, particularly in the presence of abnormal left ventricular diastolic function when the heart rate increases and left ventricular filling time is reduced. Since left ventricular filling time is strongly dependent on cycle length, we analysed this relation in all groups (Fig. 7). The regression equations were identical for the normal controls, patients with left bundle branch block, and VDD pacemaker patients with an atrioventricular interval of 50 ms. These equations show that theoretically left ventricular filling time would be zero at mean cycle length of 320 ms. The relation between cycle length and filling time was shifted downwards as atrioventricular intervals increased. Thus with an atrioventricular interval of 250 ms the cycle length producing a theoretical filling time of zero increases to 405 ms. Therefore longer atrioventricular intervals have an increasingly greater negative influence on filling time as heart rate increases. We conclude that a short atrioventricular interval of 50 or 75 ms in patients with VDD pacemakers is likely to promote more physiological conditions by maximising left ventricular filling time both at rest and on exercise. This is likely to be of particular benefit in the groups of patients who most need this type of pacemaker; the young who require high cardiac outputs at high heart rates and patients with impaired left ventricular diastolic function.

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