Left ventricular function during balloon dilatation of the aortic valve in elderly patients: a blind study of echocardiograms

Henrik Egeblad, Alf Wennevold

Abstract
Subcostal echocardiography of the left ventricle was continuously recorded during balloon dilatation of the aortic valve in ten consecutive elderly patients. Left ventricular wall motion decreased gradually during a minute of maintained complete balloon inflation. Blind analysis of M mode echocardiograms showed a significant reduction in fractional shortening. Deflation of the balloon resulted in global left ventricular hypokinesia. There was a transient overshoot in fractional shortening, wall thickness, and blood pressure approximately 40 s after deflation of the balloon. The myocardial reaction was similar to that known to occur regionally with brief coronary artery obstruction, whereas the timing of the events seemed to be dissociated from the timing of changes in left ventricular load. Thus it is likely that the left ventricular response during valve dilatation reflects not only changes in load but also an ischaemia-reperfusion effect on the myocardium. The systolic thickening of the hypertrophied interventricular septum was slightly reduced after balloon dilatation. This finding might be a result of repeated episodes of ischaemia and reperfusion. However, the potential for myocardial injury seemed to be unimportant clinically.

Balloon dilatation is increasingly used to treat elderly patients with aortic valve stenosis.1-3 The cardiovascular response to the repeated inflations and deflations of the balloon is complex. Transient circulatory obstructions cause considerable changes in left ventricular load. In addition, ischaemia and reperfusion may have a direct and potentially injurious effect on the myocardium. Brief obstruction of a coronary artery is known to result in characteristic abnormalities in regional wall motion.4,5 The aim of this study was to examine whether there are similar global myocardial reactions in the left ventricle during balloon dilatation of the aortic valve.

Patients and methods
We studied 10 consecutive patients (four men, six women; median age 76 years (range 72-87 years)) who underwent balloon dilatation of the aortic valve. All had clinical, electrocardiographic, radiological, and echocardiographic findings indicating severe stenosis of the aortic valve. Three of the patients had coexistent aortic valve incompetence that was considered to be haemodynamically insignificant on the basis of physical examination. None of the patients had primary mitral valve disease. Open heart operation for aortic valve stenosis was considered inappropriate because of the patient's advanced age or coexistent disease.

Invasive procedure
A 7 F pigtail catheter was introduced percutaneously into the femoral artery and advanced to the ascending aorta and on to the left ventricle with the aid of a guide wire. The peak gradient over the aortic valve was calculated as the difference between peak left ventricular systolic pressure and peak systolic pressure in the ascending aorta. The guide wire was used to replace the pigtail catheter by a Meditech 15 mm (3-4 cm) balloon catheter. The aortic valve was dilated with the centre of the balloon at the level of the aortic valve, which was apparent from the calcifications seen by fluoroscopy. The balloon was inflated with a mixture of saline and Urografin 76%. Inflation was maintained at constant maximum manual pressure for 60 s unless the patient showed signs of insufficient cerebral perfusion. On fewer than 10 occasions was it necessary to deflate the balloon after 40 or 50 s because of vertigo, obvious dyspnoea, or when the patients speech became slowed. None of the patients lost consciousness or developed chest pain during the procedure.

Repeated dilations were performed every 1-2 min, or more if the balloon catheter was replaced by a balloon of a larger size. Where possible dilations were routinely carried out with balloons of increasing sizes from 15 to 18 or 20 mm in diameter. The procedure was stopped after 3-5 dilations with the largest balloon. A median of eight (range 3-11) inflations of a balloon in the aortic ostium were carried out in the 10 patients.

Blood pressure was continuously recorded through an arterial sheath. Cardiac output was not measured. Because these were elderly patients we did not perform ventriculography, aortography, or coronary arteriography.

Non-invasive procedure
Cross sectional and M mode echocardiograms from the parasternal, apical, and subcostal positions were recorded on video tape before and after balloon dilatation. During dilatation
simultaneous cross sectional and M mode echocardiography were performed from the subcostal position. An M mode diameter through the left ventricle at the level of the chordae of the mitral valve was selected on the cross sectional four chamber view. Echocardiograms were continuously recorded on video tape before each inflation of the balloon, during maintained complete balloon inflation, and during and after deflation of the balloon (25 mm/s or 50 mm/s). Polaroid photographs were taken from the video tapes including recordings obtained before and after dilation and immediately before and at the end of each balloon inflation (fig 1). In addition, Polaroid recordings were obtained from the video tape after each deflation of the balloon when left ventricular wall motion was re-established and seemed from the cross sectional image to be at its maximum (fig 1). Exaggerated respiratory movements occasionally distorted the echocardiogram. In these patients photographs were taken of the video tapes as close to the scheduled time as possible.

The identity of the patient and the timing of the recording were marked on the reverse of each photo. They were later shuffled and analysed in random order without the knowledge of at what stage of the procedure they were obtained. We measured left ventricular diameter at end diastole (Dd) and end systole (Ds) and septal (ST) and lateral wall (LW) thickness. Left ventricular fractional shortening (FS = (Dd-Ds)/Dd × 100%) and thickening of the septum and lateral wall from diastole to systole were calculated. All measurements were obtained as mean values of 3–5 cycles.

The heart rate and number of runs of three or more ventricular extrasystoles in relation to inflation of the balloon were calculated from the continuously recorded electrocardiogram.

We used the Wilcoxon test for statistical analysis and a p value below 5% was regarded as significant.

**Results**

In the subcostal four chamber view the left ventricle appeared gradually to develop diffuse hypokinesia or akinesia over the 60 s period of maintained balloon inflation, with subsequent gradual development of global hyperkinesia after deflation of the balloon. These changes accorded with the measurements taken from the M mode recordings (fig 1 and 2). The median values and ranges shown in fig 2 include all values obtained immediately before inflation of the balloon, 60 (15–60) s after complete inflation, and 40 (10–60) s after the deflations. The most pronounced echocardiographic changes were associated with fractional shortening. There was a significant reduction from 30 (12–72)% before obstruction to 26 (9–56)% during obstruction and a significant increase to 43 (15–79)% after deflation. A reduction in systolic thickening of the lateral wall from 6 (1–13) mm before obstruction to 4 (0–10) mm during obstruction was statistically significant; systolic thickening of the septum was reduced from 4 (0–18) mm to 3 (0–14) mm (p > 0.05). After deflation systolic thickening of both the lateral wall and septum increased to 7 (0–15) mm and 7 (0–18) mm respectively (p < 0.05 for both).

The diastolic thickening of the septum and lateral wall did not change significantly during obstruction (before obstruction 16 (8–30) mm
for lateral wall thickness and 18 (10–35) mm for septal thickness compared with 15 (9–24) mm for lateral wall and 18 (10–38) mm for septal thickness during obstruction). Lateral wall thickness increased to 16 (8–25) mm and septal wall thickness to 20 (13–38) mm after deflation of the balloon (p < 0·05 for both). Left ventricular diameter at end diastole did not change significantly, either from before obstruction (52 (32–60) mm) to the value measured after 60 s of obstruction (52 (32–60) mm) or that measured 40 s after deflation of the balloon (51 (34–64) mm).

Heart rate increased from 92 (64–135) beats/min before to 100 (65–122) beats/min during obstruction and slowed to 95 (64–135) beats/min 40 (10–60) s after deflation, but these variations were not statistically significant. Blood pressure fell from 105 (68–175)/50 (30–70) mm Hg to 68 (24–95)/38 (17–55) mm Hg from baseline to balloon obstruction (p < 0·05 for systolic and diastolic blood pressure) and increased to 135 (65–195)/50 (30–100) mm Hg 40 (30–90) s after deflation of the balloon (p < 0·05 for both). Only one episode of three or more runs of ventricular extrasystoles was recorded before balloon inflation whereas runs of ventricular extrasystoles were seen in 43% of the periods after inflation of the balloon (5 (0–12) s after inflation) and in 30% of the periods after deflation (45 (18–80) s after deflation).

Values recorded for the first cycle of balloon inflation and deflation and for the last cycle were not significantly different (table 1).

Data recorded before the procedure were compared with those obtained after the treatment (table 2). Aortic valve opening was significantly increased and the valve gradient and septal thickening decreased significantly.

### Discussion

Previous studies showed that echocardiography may be a useful aid to guiding the balloon catheter and for evaluation of the result of balloon dilatation of the aortic valve.1,9 Transoesophageal echocardiography in particular may facilitate stable and clear monitoring of the valve and left ventricle.6 However, this technique requires general anaesthesia, which is best avoided in elderly patients and impedes the evaluation of the tolerance of decreased cerebral perfusion during balloon inflation. Parasternal M mode echocardiography of the left ventricle may be difficult to perform unless the patient is in the left lateral position, and interference between the ultrasound transducer, the hand of the examiner, and the fluoroscope, which is used to control the position of the balloon, is unavoidable.10 Therefore, in this study we monitored left ventricular wall motion from the subcostal view. Continuous cross-sectional echocardiography was helpful for maintenance of the direction of the M mode echocardiogram although optimum definition of the cardiac walls could not be obtained in all patients (fig 1). However, the reliability of the variations actually detected was endorsed by blind analysis of the echocardiograms and by previously reported experimental and clinical data.

Left ventricular hypokinesia during balloon obstruction—this study recorded by echocardiography (figs 1 and 2)—has recently been confirmed by angiography.11 Hypokinesia may occur because of the considerable increase in left ventricular afterload, but may also be a result of ischaemia. Reduction of cardiac output and aortic pressure owing to the balloon inflation together with possible obstruction of the coronary artery ostia may diminish myocardial oxygen supply. The ensuing reduction of the pump function might complete a vicious circle. The hypertrophy of the left ventricular myocardium (table 2) and possible coronary artery disease in some of the patients could further increase the sensitivity to reduced blood pressure, with the subendocardium being particularly jeopardised.12 Coronary arteriography was not performed in our

![Figure 2 Variation in left ventricular variables measured by subcostal echocardiography during balloon dilatation of the aortic valve. Median values and range are shown in 10 patients who had 8 (3–11) balloon inflations performed. Values measured after 60 (15–60) s obstruction of the aortic ostium are compared with preobstructive values obtained immediately before balloon inflation and with values measured 40 (10–60) s after deflation of the balloon. See footnotes to tables for abbreviations. *p < 0·05 compared with preceding value (rank sum test for paired data).](image)
| Table 2 | Data (median range) measured before and after balloon dilatation of the aortic valve |
|-----------------|---------------------------------|-----------------|
| **Before**      | **After**                       |                 |
| Time of examination | in relation to procedure (days) |                 |
| 3 (1-7)          | 4 (1-11)                        |                 |
| Aortic cusp separation | 4 (3-6)                     | 6 (3-9)*        |
| Dd (mm)          | 52 (39-62)                      | 58 (37-65)      |
| Ds (mm)          | 38 (24-52)                      | 43 (15-51)      |
| FS (%)           | 32 (17-40)                      | 29 (14-59)      |
| ST (mm)          | 15 (12-18)                      | 15 (12-20)      |
| PW (mm)          | 12 (8-17)                       | 13 (10-15)      |
| **Subcostal echocardiography** |                 |                 |
| Dd (mm)          | 53 (37-63)                      | 55 (37-59)      |
| Ds (mm)          | 38 (18-52)                      | 39 (18-48)      |
| FS (%)           | 28 (14-51)                      | 24 (16-53)      |
| ST (mm)          | 17 (13-34)                      | 17 (14-37)      |
| LW (mm)          | 14 (10-21)                      | 13 (12-21)      |
| ΔST (mm)         | 5 (0-9)                         | 3 (0-7)*        |
| ΔLW (mm)         | 5 (2-8)                         | 5 (2-12)        |
| HR (beats/min)   | 85 (65-107)                     | 90 (72-107)     |
| BP (mm Hg)       | 130 (100-155)                   | 130 (100-165)   |
| Aortic valve gradient | 80 (30-125)                   | 30 (0-55)*      |

Left ventricular characteristics, heart rate (HR), and blood pressure (BP) before and after balloon dilatation of the aortic valve. Dd, left ventricular diastolic diameter; Ds, left ventricular systolic diameter; FS, left ventricular fractional shortening; LW, lateral wall thickness in diastole; ΔLW, systolic thickening of lateral wall; PW, posterior wall thickness in diastole; ST, thickness of the interventricular septum in diastole; ΔST, systolic thickening of the interventricular septum. *p < 0.05.

patients, none of whom developed angina pectoris during balloon dilatation. However, transient ST-T depression was often seen on the monitor electrocardiogram. The ST-T depressions were not measured because electrocardiographic mapping would have interfered with the transcoronary anastomosis. Furthermore, a bundle branch block and treatment with digoxin may interfere with the measurement and interpretation of ST-T changes in many elderly patients with aortic stenosis.

Complete coronary artery obstruction by ligation leads to regional dyskinesia within a minute. Dyskinesia is often noticed in anterior wall infarction with Q waves but rarely in non-Q wave infarction and in patho-anatomical subendocardial infarction. Most of our patients showed hypokinesia during balloon inflation but a few showed akiniesia (fig 1). Dyskinesia did not occur. Left ventricular wall motion gradually became hypokinetic or akinetic within 60 s of complete inflation of the balloon. The left ventricular diameter at end diastole did not change significantly (fig 2). Thus the increase in left ventricular load remained constant while the balloon was kept inflated. Increased afterload doubtlessly contributed to the left ventricular response when the balloon was kept inflated. However, the timing and character of the wall motion abnormality were compatible with the concept of subendocardial ischaemia as a determinant of the development of hypokinesia and akiniesia during the balloon obstruction. This hypothesis is supported by the recent demonstration by catheterisation of the coronary sinus of transient low flow ischaemia during balloon inflation. The response of the left ventricle to deflation of the balloon included transient hyperkinesia of the left ventricular walls that reached a maximum 40 s after the start of deflation. There was no significant change in the left ventricular diameter at end diastole. Deflation of the balloon led to a reduction of the left ventricular afterload. A decrease of peripheral vascular resistance is likely to occur at the same time. Cardiac output was not monitored in this study but a reduction in peripheral vascular resistance was recognised in studies of declamping after aortic surgery. The reduction in afterload resulting from the balloon deflation and the potential reduction of peripheral vascular resistance accord with the suggestion that there is left ventricular hyperkinesia after obstruction by the balloon. However, reperfusion of the myocardium may contribute significantly to the increased wall motion. A hyperkinetic wall motion response, which was attributed to sudden cellular calcium influx, was seen in animal experiments ligation of the coronary artery for a minute resulted in maximum hyperkinesia after 40 s of reperfusion. A similar phenomenon of hypokinesia and hyperkinesia was seen in variant angina. Furthermore, the significant increase in diastolic wall thickness after deflation of the balloon resembles the findings at coronary artery reperfusion in the experimental setting. The increase in wall thickness has been attributed to engorgement of the myocardial vascular bed during reactive hyperaemia. The gradual increase in left ventricular wall motion and systolic blood pressure during reperfusion, with an overshoot in both after 40 s and a simultaneous diastolic blood pressure value at the pre-obstructive level are likely to reflect myocardial reperfusion rather than a decrease of afterload. The reduction in afterload may be greatest immediately after the balloon deflation. Finally, the occurrence of arrhythmias after deflation of the balloon indicates a reperfusion effect on the myocardium. The variation in fractional shortening was further analysed to obtain a clearer idea of the time course of the myocardial response to inflation and deflation of the balloon (fig 3). To limit the number of measurements, the analysis was confined to two patients. Their M mode video recordings of the left ventricle were stable and during all aortic valve dilatations the balloon was completely inflated for 60 s. The number of balloon inflations in both these patients resembled the median number of dilatations (eight) in the study population. The two patients are not those shown in fig 1. An additional 161 photographs were obtained from the two patients' video tapes immediately before the procedure and every 15 s during and after 60 s of maintained complete inflation of the balloon. Fractional shortening was measured without knowledge of the timing of the recordings. There was no significant reduction of fractional shortening until late during the 60 s period of maintained inflation (fig 3). Other data from our laboratory showed that the time needed for complete filling and emptying of the balloon was approximately 20 s. Figure 3 indicates that there was no reduction of fractional shortening during filling of the balloon.
left ventricular response to balloon dilatation of the aortic valve we found significant changes in left ventricular wall motion and wall thickness in relation to inflation and deflation of the balloon. The changes in left ventricular load undoubtedly contributed to the myocardial reactions but their character and time course resembled those seen in experimental and clinical ischaemia and reperfusion. Thus myocardial ischaemia and reperfusion may be important determinants of the left ventricular response to balloon dilatation of the aortic valve, and the potential impact on the myocardium of repeated ischaemia and reperfusion should be considered when percutaneous balloon dilatation of the aortic valve is performed.