Changes in Cardiac Output and Stroke Volume During First Four Months After Cardiac Infarction

FRANK NAGER*, MICHAEL THOMAS, AND JOHN SHILLINGFORD

From the Medical Research Council’s Cardiovascular Research Unit, Royal Postgraduate Medical School, London W.12

The circulatory changes that occur within the first four months after cardiac infarction have not been studied to the same extent as those occurring during the acute phase of the illness. Measurements of cardiac output and stroke volume have been made in some patients before leaving hospital (Pritchard and Hellerstein, 1950; Gammill et al., 1955; Lee, 1957; Broch et al., 1959; Thomas, Malmcrona, and Shillingford, 1965b), but in comparatively few patients have measurements been made over a longer period of two months (Smith, Wikler, and Fox, 1954; Murphy et al., 1963). In a single report results of investigations made up to three to five months after the acute illness are given (Malmcrona and Varnauskas, 1964).

The purpose of this study was to define more completely the changes in cardiac output and stroke volume that may occur during the first four months after cardiac infarction. The conclusions are based on 126 hemodynamic investigations made in conjunction with clinical observations in 22 patients. At each investigation the circulatory response to a standard postural change (Thomas and Shillingford, 1965) was studied.

Patients and Methods

Studies were made of 22 patients, 18 men and 4 women, whose ages ranged from 32 to 80 years, mean 57 years. Five other patients in whom the follow-up studies were not completed owing to their death in hospital or during the early convalescent period have been excluded. All patients had a clinical history and electrocardiographic signs typical of myocardial infarction. The serum lactic dehydrogenase was raised in 17 patients. Four patients (8, 9, 11, 15) had had a myocardial infarction in the past. Four (5, 13, 16, 22) were known to have had systemic hypertension before myocardial infarction. Hypertension of varying degree became apparent during the follow-up period in six patients (9, 15, 17, 18, 19, 20). One (19) suffered from mild diabetes mellitus and three (1, 5, 11) had evidence of chronic bronchitis.

During the follow-up period three patients (7, 11, 13) were treated with digoxin and diuretics; one hypertensive patient (13) received bethanidine; another hypertensive patient (22) received chlorothiazide before the last investigation; and one (16) had a probable further myocardial infarction before the last investigation.

All patients were in sinus rhythm at the time of hemodynamic study. All 22 patients were studied serially in the early convalescent period and after discharge from hospital. In addition, 14 of the 22 patients were studied during the acute illness, as detailed. Measurements in the acute phase (first week following infarction) were made in a special intensive care unit with equipment permanently installed (Shillingford and Thomas, 1964). Subsequent measurements during the early convalescent period (second to fifth week) were conducted in a catheter room and were in general made during the second or third week, as detailed. After discharge from hospital, patients were studied during the periods 4–5 weeks, 6–8 weeks, 9–13 weeks, and 15–22 weeks, with a few exceptions as noted.

Each hemodynamic study consisted of measurements of heart rate, cardiac output, and arterial blood pressure. Cardiac output was measured by a dye dilution technique (Gabe and Shillingford, 1961; Thomas, Malmcrona, and Shillingford, 1965a), using the photoelectric earpiece (Cambridge Instrument Co.) and Coomassie Blue dye (I.C.I.). A known quantity of dye (approx. 50 mg.) was injected from a special syringe (Gabe and Shillingford, 1961) through a fine polyethylene catheter (PE 60 Intramedic U.S.A.) inserted percutaneously by Seldinger technique and advanced to the great veins. The first dye curve was calibrated by the tailpiece method. Dye was
extracted from the plasma of a blood sample taken three minutes after injection. Dye concentration was measured by spectrophotometry. Subsequent relative cardiac outputs were calculated from the reciprocal of the areas of the extrapolated dye curves.

Heart rate was derived from the dye curve by using slightly incomplete pulse rejection on the dye recorder. Arterial pressure was measured by sphygmomanometry.

In six patients (5, 6, 7, 8, 10, 11) the cardiac outputs during the acute phase were calculated by the direct Fick principle. Oxygen consumption was measured by collecting expired air in a Douglas bag. The O₂ and CO₂ tensions of the expired air were determined by use of a polarographic electrode (Instrument Laboratory IL 113) and the volume was measured by passing it through a dry gas meter. Arterial blood was obtained from the brachial artery and mixed venous blood by advancing the catheter into the pulmonary artery. The oxygen content of the arterial and mixed venous blood was calculated from the PO₂, pH, and Hb, using Dill’s oxygen dissociation tables.

Patients studied during the acute phase were lying flat. All other investigations in the convalescent period and after discharge from hospital took place with the patients lying on an electrically-driven tipping table. Initial measurements were made in a 20° feet down position. The procedure undertaken for follow-up studies was to insert the catheter with the patient lying flat. The patient’s position was then changed to the 20° feet down position, and after approximately 5 minutes the first two measurements of cardiac output, heart rate, and blood pressure were made. The patient’s position was then changed to the horizontal and the legs raised by means of a wooden support. After 3–4 minutes two further measurements of heart rate, cardiac output, and blood pressure were made.

RESULTS

Results of the haemodynamic measurements are summarized in the Table and Fig. 1–6.

Haemodynamic Changes

(1) Cardiac Output. Measurements of cardiac output in the acute phase (14 patients) ranged from 2·40 to 6·74 l./min.; at the beginning of the convalescent period (22 patients) from 2·97 to 6·8 l./min. and at the end of the period of follow-up (22 patients) from 3·10 to 6·86 l./min.

Acute phase—early convalescence. Of the 14 patients studied in the acute phase, 2 (20, 22) had a cardiac output above 5 l./min., 9 (5, 6, 10, 14, 15, 16, 17, 19, 21) had a cardiac output between 4·0 and 5·0 l./min., and 3 (7, 8, 11) had a cardiac output less than 4·1 l./min.

Within the period between the acute illness and the first study in the early convalescent period, of the two patients with initial cardiac output above 5 l./min., one (20) showed an increase and the other (22) a fall. Of the 9 patients with initial cardiac output between 4·0 and 5·0 l./min., 4 (6, 10, 14, 16) showed an increase: in patients 10 and 14 this was of the order of 2 l./min. Four (5, 15, 17, 19) of these 9 patients showed no change and one (21) showed a fall.

Early convalescence—end of follow-up. All 22 patients were studied serially in the period between early convalescence and the third or fourth month. At the time of the first study in the convalescent period, 12 patients (2, 3, 4, 6, 8, 9, 10, 13, 14, 16, 18, 20) had a cardiac output above 5 l./min. Seven patients (5, 7, 12, 15, 17, 19, 22) had a cardiac output between 4·0 and 5·0 l./min., and 3 (1, 11, 21) had a cardiac output of less than 4·0 l./min.

During the period of follow-up from the second week, of the 12 patients with a cardiac output above 5·0 l./min. in early convalescence, only one (9) showed a further rise. One (6) showed a transient increase and then a fall to the initial level. Four patients (2, 3, 13, 18) showed no change; six (4, 8, 10, 14, 16, 20) showed a fall in cardiac output, but in only 3 (14, 16, 20) did cardiac output fall below 5 l./min. In one (16) of these three patients a further myocardial infarction at the twelfth week may have contributed and in another (20) increase of blood pressure to 200/110 mm. Hg occurred during the follow-up period.

(2) Stroke Volume. Results of stroke volume measurements are given with reference to three ranges:—those patients with stroke volume above 60 ml. (Group 1), those patients with stroke volume between 40 and 60 ml. (Group 2), and those patients with stroke volume of less than 40 ml. (Group 3). Changes in stroke volume in patients who had values within these ranges at the first study in early convalescence are illustrated in Fig. 1, 2, and 3.

Measurements of stroke volume in the 14 patients studied in the acute phase ranged from 20 to 87 ml., in all 22 patients at the beginning of convalescence from 38 to 97 ml., and at the end of the period of follow-up from 35 to 105 ml.

Acute phase—early convalescence. Measurements of stroke volume in the acute phase in the 14 patients previously detailed, as compared with the first measurement in early convalescence, showed the following changes. Of 3 patients (14, 16, 20) with a lowest value of stroke volume in the acute illness of above 60 ml., one (14) remained of the same order and two (16, 20) subsequently showed an increase. Of 9 patients (5, 6, 8, 10, 15, 17, 19, 21, 22) with stroke volume between 40 and 60 ml. in the acute phase, 6 (5, 6, 8, 10, 15, 17) showed an increase
Cardiac Output and Stroke Volume after Cardiac Infarction

During the following week, one (21) showed a fall and in the remaining two there was no significant change. Two patients (7, 11) had a stroke volume between 20 and 40 ml. in the acute phase. In the following week both showed an increase, greater in patient 7 as compared with patient 11.

Early convalescence—end of follow-up. Thirteen patients (2, 3, 4, 5, 6, 8, 9, 10, 13, 14, 16, 18, 20) had a stroke volume greater than 60 ml. at the first investigation in early convalescence. Of these, 4 (5, 8, 9, 13) showed a further increase, 5 (2, 3, 6, 10, 14) showed no change, and 4 (4, 16, 18, 20)
### TABLE

**H/AEMODYNAMIC DATA**

<table>
<thead>
<tr>
<th>Patient No.</th>
<th>Days (D1-D7) or weeks (2-28) after infarction</th>
<th>Position 1</th>
<th></th>
<th>Position 2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Heart rate (beats/min.)</td>
<td>Cardiac output (l/min.)</td>
<td>Stroke volume (ml.)</td>
<td>Blood pressure (mm. Hg)</td>
</tr>
<tr>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>80</td>
<td>3:70</td>
<td>46</td>
</tr>
<tr>
<td>3</td>
<td>3</td>
<td>59</td>
<td>5:20</td>
<td>88</td>
</tr>
<tr>
<td>4</td>
<td>2</td>
<td>76</td>
<td>6:55</td>
<td>106</td>
</tr>
<tr>
<td>5</td>
<td>7</td>
<td>77</td>
<td>4:25</td>
<td>55</td>
</tr>
<tr>
<td>6</td>
<td>16</td>
<td>74</td>
<td>4:17</td>
<td>56</td>
</tr>
<tr>
<td>7</td>
<td>12</td>
<td>74</td>
<td>4:04</td>
<td>63</td>
</tr>
<tr>
<td>8</td>
<td>18</td>
<td>74</td>
<td>4:04</td>
<td>63</td>
</tr>
<tr>
<td>9</td>
<td>13</td>
<td>74</td>
<td>4:04</td>
<td>63</td>
</tr>
<tr>
<td>10</td>
<td>17</td>
<td>74</td>
<td>4:04</td>
<td>63</td>
</tr>
<tr>
<td>11</td>
<td>21</td>
<td>74</td>
<td>4:04</td>
<td>63</td>
</tr>
<tr>
<td>12</td>
<td>24</td>
<td>74</td>
<td>4:04</td>
<td>63</td>
</tr>
</tbody>
</table>

Table continued on next page.
showed a fall. In patients 4 and 18, the stroke volume remained above 60 ml. In patients 16 and 20 stroke volume was below 60 ml at the final follow-up study, possibly in relation to a further small myocardial infarction in the thirteenth week in patient 16, and hypertension rising to 200/110 mm Hg in patient 20. The mean stroke volume in the 13 patients at the first study in early convalescence was 80 ml and at the end of the follow-up period it was 78 ml.
Seven patients (1, 12, 15, 17, 19, 21, 22) had a stroke volume between 40 and 60 ml. at the first investigation of the convalescent period. Of these, 4 (1, 12, 17, 21) showed an increase during the follow-up period and 3 (15, 19, 22) showed no change. The average stroke volume in the 7 patients at the first study in early convalescence was 52 ml. and at the end of the follow-up period 58 ml.

Two patients (7, 11) had a stroke volume between 20 and 40 ml. (38 and 37 ml., respectively) when studied during the second week. In 7 a period of temporary clinical improvement following treatment with digitalis and diuretics was associated with an increase of stroke volume from 38 to 74 ml. Subsequent deterioration with a fall in stroke volume to 35 ml. occurred after discharge from hospital. In patient 11 the stroke volume at the end of the follow-up period was 62 ml.

(3) Heart Rate: Acute phase—early convalescence. Heart rate in the acute phase (14 patients) ranged from 55 to 120/min. Six patients (5, 7, 11, 17, 19, 22) had heart rates above 85/min., five (6, 15, 16, 20, 21) between 70 and 85/min., and three (8, 10, 14) between 55 and 70/min. Within the time between the acute illness and the first investigation of the convalescent period there was a general tendency for high and low heart rates to revert to the normal range. Patients with heart rates in the range 70–85/min. during the acute illness showed little change.

Heart rates at the time of the first study of the convalescent period were within the range 70–85/min. in all patients except six (2, 7, 8, 18, 19, 22). Of these six patients, three (2, 8, 18) had rates below 70/min. and three (7, 19, 22) above 85/min. Patient 7 had a stroke volume below 40 ml. and other evidence of circulatory failure. Patient 19 had a stroke volume of 50 ml. and subsequent to the follow-up period developed congestive cardiac failure. Patient 22 had a stroke volume of 54 ml.
Cardiac Output and Stroke Volume after Cardiac Infarction

**Fig. 5.**—Changes in stroke volume with time and the response to the standard postural change in those patients with stroke volume between 40 and 60 ml. at the first study in the early convalescent period.

**Fig. 6.** Changes in stroke volume with time and the response to the standard postural change in those patients with stroke volume of less than 40 ml. at the first study in the early convalescent period. Interrupted lines represent a fall in stroke volume. On the left, patient 7; on the right, patient 11.
Early convalescence—end of follow-up. During the period of follow-up from the second week the 16 patients with heart rate in the range 70-85/min. at the beginning of convalescence behaved as follows. Eight (1, 3, 4, 6, 9, 10, 16, 21) showed no significant change in heart rate; seven (5, 11, 12, 13, 14, 15, 17) showed a fall, most conspicuous in 11 and 12. One patient (20) showed a progressive increase.

Of the three patients (7, 19, 22) with heart rate above 85/min. in early convalescence, one (7) showed an initial fall to 66/min., subsequently rising again to 88/min. One (19) remained in the range 85-105/min., and one (22) showed a progressive fall to 80/min.

Of the three patients with heart rates below 70/min. at the beginning of convalescence, two (2, 18) subsequently increased to 70/min. or above and one (8) remained low (about 55/min.).

(4) Blood Pressure. In all except five patients (1, 2, 3, 4, 7) the blood pressure increased during the period of study. In some (9, 15, 17, 18, 19, 20) hypertension appeared to varying degrees. Four patients (5, 13, 16, 22) with known previous hypertension either maintained or ultimately reached high levels of arterial pressure. Patient 13 was treated with bethanidine throughout the study and patient 22 was given chlorothiazide for control of arterial pressure between the studies of the eleventh and sixteenth weeks.

Circulatory Response to a Standard Postural Change

The changes in cardiac output, heart rate, stroke volume, and blood pressure, which resulted from the standard postural test conducted at different stages in the follow-up period, are given in the Table. Patients were placed in three groups on a basis of the stroke volume found at the first investigation of the follow-up period (second week in most patients). Group I consists of patients with stroke volume greater than 60 ml. in the 20° feet down position; Group 2 consists of those with stroke volume between 40 and 60 ml., and Group 3 those with stroke volume less than 40 ml.

Group I. Patients 2, 3, 4, 5, 6, 8, 9, 10, 13, 14, 16, 18, and 20 were placed in this group. The average stroke volume of these patients at all investigations conducted between the second week and the end of the follow-up period was 80-8 ml. The average stroke volume in the second position of the posture test was 99.0 ml. The average percentage increase of stroke volume was 22.5 per cent. The Table shows that 8 of the 13 patients within this group responded with a large increase of stroke volume following the change in posture at all investigations during the period of follow-up. An example of a typical response is shown in Fig. 4. Two patients (3, 5) in whom stroke volume was progressively increasing during the period of follow-up showed a progressively greater stroke volume response to posture. Patient 9 showed only a small stroke volume response at the beginning of convalescence but a greater response at the final study. Patient 20 achieved a good stroke volume response between the fourth and eleventh weeks but this was less at the final study: at the 16th week arterial pressure had risen to 210/110 mm. Hg.

The two patients (16, 18) who clearly did not increase stroke volume following the change in posture were also exceptional in that patient 16 had very high levels of arterial pressure at all times and suffered a further small myocardial infarction before the final study; patient 18 had evidence of a large functional aneurysm of the left ventricle though maintaining a good resting stroke volume of more than 80 ml.

Group II. Patients 1, 12, 15, 17, 19, 21, and 22 were placed in this group. The average stroke volume of these patients at all investigations conducted between the second week and the end of the follow-up period was 54.4 ml. The average stroke volume in the second position of the posture test was 66.3 ml. The average percentage increase in stroke volume was 21.9. Fig. 5 illustrates the progress of each individual patient in terms of stroke volume and postural response. Three (1, 12, 17) showed increases in stroke volume during the period of study; 2 of these (1 and 12) showed improving stroke volume response to posture and a consistent moderately good response, respectively, but 17 showed a diminishing response. Patient 21 showed a moderate stroke volume response to posture except at the seventh week. Patients 15, 19, and 22 showed little change in stroke volume, but in patient 19 the stroke volume response to posture was greater as compared with patients 15 and 22 in whom the response was poor until 16 weeks after cardiac infarction.

Group III. Patients 7 and 11 were placed in this group (Fig. 6). The average stroke volume of these two at all investigations conducted between the second week and the end of the follow-up period was 51.8 ml. The average stroke volume in the second position of the posture test was 48.8 ml. The average percentage change of stroke volume was a fall of 6.
Cardiac Output and Stroke Volume after Cardiac Infarction

In patient 7 a transient clinical improvement was associated with increased stroke volume but thereafter the stroke volume remained low. Stroke volume either showed little change or decreased after change of posture. Patient 11 showed a progressive increase in stroke volume from 37 to 62 ml. but stroke volume fell on every occasion after posture change.

Correlation of Haemodynamic Measurements with Clinical Features

The cardiovascular physical signs and radiological features that were present at the time of haemodynamic investigations were routinely recorded. Particular note was made of auscultatory signs, the presence or absence of a raised jugular venous pressure, radiological evidence of cardiac enlargement, and signs of pulmonary venous hypertension and pulmonary oedema. Clinical features suggesting ventricular aneurysm were also noted. Results are presented with reference to the findings at the first study in the early convalescent period and patients are divided into three groups.

Group I: patients with stroke volume greater than 60 ml. at this time.

Group II: patients with stroke volume between 40 and 60 ml.

Group III: patients with stroke volume less than 40 ml.

Auscultatory Signs

Group I: Patients 2, 3, 4, 5, 6, 8, 9, 10, 13, 14, 16, 18, and 20 were placed in this group. Seven of them were studied in the acute phase at which time four (5, 6, 14, 16) had a fourth heart sound, but this was lost by the time of the first convalescent study.

Three (8, 13, 18) had a fourth heart sound at the first study in early convalescence. In patient 8 this was lost after five weeks, but in patients 13 and 18 it persisted throughout the whole period of study. Patient 13 had systemic hypertension and patient 18 had clinical evidence of a ventricular aneurysm. Patient 16 had a fourth sound in the acute illness, lost it until the seventh week, when it returned, ultimately to persist. This patient had systemic hypertension. Patient 20 developed a fourth sound at the end of the period of study; he also had systemic hypertension.

Group II: Patients 1, 12, 15, 17, 19, 21, and 22 were placed in this group. Five of them were studied during the acute phase. Four (15, 17, 19, 21) had a fourth sound at this time; patient 17 had a third sound also. Patient 22 had a third sound during the acute illness.

At the time of the first study in early convalescence, five (1, 12, 16, 17, 21) had a fourth sound which was lost permanently during the period of study in 3 (1, 15, 17) and temporarily in one (21). In 2 of these 3 patients (1, 17), stroke volume was increasing during the period in which the fourth heart sound was lost and in patient 21 stroke volume increased at the time the fourth sound reappeared. In patient 22, in whom the arterial pressure rose to hypertensive levels during the follow-up period, a fourth sound appeared at the sixteenth week.

Group III. Patients 7 and 11 were placed in this group. Both of them were studied in the acute phase at which time patient 7 had fourth and third heart sounds and 11 a third heart sound.

At the first study of the convalescent period patient 7 had a fourth heart sound which persisted until the 20th week; from the eleventh week a third heart sound was present. General clinical improvement with a greater stroke volume was present during the time in which the third heart sound was absent. Patient 11 had a third heart sound during the whole period of study, at the end of which some small clinical improvement in terms of exercise tolerance and greater improvement in stroke volume was evident.

Raised Jugular Venous Pressure

Five (6, 7, 11, 21, 22) of the 14 patients studied during the acute phase had a jugular venous pressure 5 cm. or more above the sternal angle at that time.

At the time of the first study in the convalescent period only 3 (7, 11, 18) had a raised jugular venous pressure. In patient 18, who had a functional cardiac aneurysm, the venous pressure was normal at subsequent studies. Patients 7 and 11, in whom the venous pressure remained raised to some extent for many weeks, had stroke volumes of less than 40 ml. at the beginning of the convalescent period and of 35 and 62 ml., respectively, at the end of the period of study (Fig. 3).

Heart Size

At the time of the first study in convalescence 7 patients (5, 7, 11, 12, 13, 15, 16) had radiological evidence of cardiac enlargement. This was particularly noticeable in 7 and 11 in whom the stroke volume was less than 40 ml. The stroke volume in the others ranged from 52 to 73 ml. Three (5, 13, 16) suffered from systemic hypertension and in 2 (12, 15) the blood pressure was significantly high at some stage in the follow-up period.
Radiological and Clinical Evidence of Pulmonary Venous Hypertension and Pulmonary Oedema

At the time of the first study in early convalescence only 2 patients (7, 11) had evidence of pulmonary venous hypertension which persisted during the whole of the period of study. Both had other clinical and haemodynamic evidence of poor cardiac function, with stroke volume of less than 40 ml. at the beginning of the follow-up period.

Ventricular Aneurysm

Four patients (2, 12, 13, 18) had evidence of a functional ventricular aneurysm. All had a pathological precordial pulsation and paradoxical ventricular movement as shown by screening. Three (12, 13, 18) had persistent S-T segment elevation in the electrocardiogram. The stroke volume at the beginning of the convalescent period was 88, 52, 62, and 87 ml., respectively. Three patients (2, 12, 13) responded to postural change by an increase in stroke volume but patient 18 showed only a very small stroke volume change. Patients 12, 13, and 18 had a persistent fourth heart sound but also had systemic hypertension.

Discussion

The long-term changes in the circulation which may follow myocardial infarction have not received the attention that has been directed towards the events of the acute illness. It may be that a knowledge of the circulatory changes occurring during the first months after myocardial infarction in relation to the physical signs shown by patients may assist in their clinical management and also aid in the assessment of prognosis with respect to physical activity.

Previous studies (Murphy et al., 1963) have pointed out that survivors from myocardial infarction usually have either a normal cardiac output during the acute illness or a rapid return of low cardiac output to normal levels during convalescence; whereas a persistently low cardiac output carries a poor prognosis for life. Malmcrona and Varnauskas (1964) studied patients during the acute phase of the illness, before they left hospital, and from 3 to 17 months after the infarction, and found no significant differences in cardiac output during these different stages of the disease. Stroke volume improved in the period between the acute phase and leaving hospital, but did not change thereafter.

While most haemodynamic studies in cardiac infarction have emphasized the importance of cardiac output, it is probable that from the point of view of the ability of the ventricles to contract, stroke volume is a more representative measurement. The results have, therefore, been presented with particular reference to stroke volume. It should be noted that measurements made in the acute phase were made in the horizontal position and subsequent values in the follow-up period were measured in a 20° feet down position. Corresponding levels for the horizontal position would be approximately 10 per cent greater in most patients with good cardiac function and less different in those with poor cardiac function.

A general appraisal of the values for stroke volume and the changes that took place introduces the difficulty of adopting criteria for normality. Since all strict conventions are open to some criticism, results have been given simply as absolute values, and division of patients into groups with stroke volume above 60 ml., between 40 and 60 ml., and below 40 ml., in the 20° feet down position, was made arbitrarily for the organization of results. The associated clinical features shown by patients were generally in support of these divisions being, respectively, representative of patients with good, moderately impaired, and poor cardiac function.

The patterns of change in stroke volume showed trends which may be distinct. Nearly all patients who achieved a stroke volume above 60 ml. during the follow-up period reached this level by the second week. Further change after the second week in these patients was variable, but in only two patients, in whom hypertension and further myocardial infarction may have been important, did stroke volume subsequently fall below 60 ml. This group of patients with the highest stroke volume showed the greatest and most rapid increase in stroke volume during the period between the acute illness and the second week. Their behaviour differed from that of the patients with a lower stroke volume at the second week who also had reduced stroke volume during the acute illness but in whom increase in stroke volume was much slower. In these latter patients, approximately half had stroke volumes less than 60 ml. at the end of the period of study. It appears, therefore, that assessment of patients at the second or third week gives a useful indication of prognosis with regard to stroke volume.

There was some relation between the severity of the acute illness in terms of degree of cardiovascular failure and the subsequent behaviour of stroke volume; in general an uncomplicated illness carried a good prognosis and overt cardiovascular failure a poor prognosis with respect to the stroke volume ultimately achieved. The essential reason for the difference in behaviour in stroke volume was not apparent from the study. A simple explanation in terms of bulk of myocardium affected by infarction...
Cardiac Output and Stroke Volume after Cardiac Infarction

was denied by the levels of serum lactic dehydrogenase and the extent of electrocardiographic changes. There was no apparent relationship to a history of previous myocardial infarction. It was notable, however, that the mean age of the patients with stroke volume above 60 ml. in early convalescence was about a decade lower than those in whom it was below 60 ml. (52 years; 63 years). Age may therefore be one factor tending to limit cardiac function after myocardial infarction.

The clinical features shown by the patients at the different times of study after the second week were in accord with the results of the haemodynamic measurements made, and in general reflected the level of stroke volume and degree of ventricular functional disease. Patients with normal stroke volume usually had normal cardiovascular physical signs though a fourth heart sound was present in those patients with systemic hypertension or ventricular aneurysm. In the absence of hypertension or signs of ventricular aneurysm, the presence of a fourth heart sound was a good indication of some reduction in stroke volume though not necessarily of a severe degree. A third heart sound persisted after the second week, however, which indicated a much reduced stroke volume and was associated with other clinical features suggesting very poor cardiac function.

Cardiac size was difficult to evaluate between the period of the acute illness and convalescence, since the initial X-ray films were taken with patients in the supine position using a ward unit, and, subsequent to the patient's mobilization, films were taken in the standing position. From the second week onward, when the technique was standard, it seemed apparent that cardiac size was normal or possibly only slightly enlarged in most non-hypertensive patients including those with moderately reduced stroke volume. Systemic hypertension was associated with cardiac enlargement in patients with normal stroke volume, but in people with normal blood pressure obvious cardiac enlargement was present only when stroke volume was very low.

The stroke volume response to a standard postural change was studied in all patients, with a view to exploring its use as a test of ventricular function. As had been observed in previous work (Thomas and Shillingford, 1965), there was a general relation between the ability to increase stroke volume and ventricular function, as assessed by the resting stroke volume and clinical criteria, but the test did not seem suitable for evaluating minor changes in cardiac function in individual patients. One patient with a ventricular aneurysm but a normal resting stroke volume failed to respond to the test, and in others the magnitude of the stroke volume response did not show a constant relation to the resting stroke volume. The irregularities may imply a multiplicity of factors involved in the test itself or possibly reflect a variety of types of ventricular dysfunction.

**Summary**

Serial haemodynamic measurements and clinical observations were made at regular intervals during the first four months after acute myocardial infarction in 22 patients. The natural course of changes in cardiac output and stroke volume was found to differ in patients who eventually had good cardiac function as compared with those in whom a cardiac functional defect remained. Increase in stroke volume was most rapid in those patients who returned to a normal stroke volume, and occurred much more slowly in those patients who failed to return to normal. Correlations between haemodynamic findings and clinical signs were found and are given in detail.

The use of a standard postural change as a test of cardiac function was explored and its place in objective assessment of the circulation was discussed.

The authors wish to thank Mr. Peter Burgess and Miss Sunja Marrington for technical assistance, and Miss Jean Powell who drew the diagrams.

**References**


Nager, Thomas, and Shillingford


Changes in cardiac output and stroke volume during first four months after cardiac infarction.
F Nager, M Thomas and J Shillingford

*Br Heart J* 1967 29: 859-870
doi: 10.1136/hrt.29.6.859

Updated information and services can be found at:
http://heart.bmj.com/content/29/6/859.citation

These include:

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/