RUPTURE OF THE INTERVENTRICULAR SEPTUM

BY
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Rupture of the interventricular septum is an uncommon complication of cardiac infarction. The seven cases reported here illustrate the clinical and pathological features of this condition.

In 1948, Fowler and Failey, in a survey of previously reported cases, found records of 60 patients who had myocardial infarction involving the interventricular septum with subsequent septal rupture; in 17 of these the condition was recognized clinically. Physicians are now becoming increasingly aware of this complication of cardiac infarction and in recent years a number of isolated case reports have appeared (Leonard and Daniels, 1938; Nareff et al., 1950; Black, 1952; Evans and Anderson, 1952; Hamilton et al., 1954; Arons and O'Rourke, 1954.)

The following seven instances of septal rupture occurred in patients admitted to the Royal Victoria Hospital, Belfast, and to the Belfast City Hospital. In the case of patients not seen personally by one of us, clinical and pathological records were made available by the physicians and pathologists concerned. The clinical and pathological features of all seven cases are summarized in tabular form, and those of one case are presented in some detail.

TABLE I

<table>
<thead>
<tr>
<th>Case</th>
<th>Age</th>
<th>Blood pressure (m.m/Hg)</th>
<th>Auscultation</th>
<th>Thrill</th>
<th>Cardiac rhythm</th>
<th>Pallor</th>
<th>Hb (g./%)</th>
<th>Cyanosis</th>
<th>Peripheral oedema</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>56</td>
<td>115/105</td>
<td>Loud systolic murmur maximal in fourth left intercostal space</td>
<td>Present</td>
<td>Regular</td>
<td>++</td>
<td>14·1</td>
<td>++</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>75</td>
<td>190/130</td>
<td>Systolic murmur heard over precordium</td>
<td>Regular</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Absent</td>
</tr>
<tr>
<td>3</td>
<td>74</td>
<td>90/?</td>
<td>Loud systolic murmur all over precordium</td>
<td>Fibrillation 150/min.</td>
<td>++</td>
<td>12·7</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>50</td>
<td>130/100</td>
<td>Harsh systolic and diastolic murmurs between apex beat and sternum</td>
<td>++</td>
<td>11·2</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>75</td>
<td>120/80</td>
<td>Heart sounds faint and almost inaudible</td>
<td>Gallop rhythm</td>
<td>++</td>
<td></td>
<td>++</td>
<td>Gross</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>87</td>
<td>130/80</td>
<td>Loud blowing systolic murmur maximal to left of sternum</td>
<td>Present</td>
<td>++</td>
<td>6·4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>74</td>
<td>135/90</td>
<td>Mitral systolic and diastolic murmurs</td>
<td>++</td>
<td></td>
<td></td>
<td></td>
<td>Gross</td>
<td></td>
</tr>
</tbody>
</table>

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## TABLE II

<table>
<thead>
<tr>
<th>Case</th>
<th>Heart weight (g.)</th>
<th>Occlusion</th>
<th>Area infarcted</th>
<th>Site of rupture</th>
<th>Edema lungs</th>
<th>Liver weight (g.)</th>
<th>Central zonal necrosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Case 1 (male)</td>
<td>550</td>
<td>Anterior descending branch left coronary artery</td>
<td>Antero-septal region</td>
<td>Below centre of anterior septal margin</td>
<td>Acute edema</td>
<td>1600</td>
<td>Present</td>
</tr>
<tr>
<td>Case 2 (male)</td>
<td>500</td>
<td>Anterior descending branch left coronary artery</td>
<td>Septum extending to posterior wall left ventricle</td>
<td>Posterior apical part of septum</td>
<td>Moderate oedema</td>
<td>1600</td>
<td>Present</td>
</tr>
<tr>
<td>Case 3 (female)</td>
<td>560</td>
<td>No satisfactory occlusion found. Gross atheroma present</td>
<td>Antero-septal region</td>
<td>Lower apical part of septum</td>
<td>Chronic venous congestion</td>
<td>1500</td>
<td>Present</td>
</tr>
<tr>
<td>Case 4 (male)</td>
<td>500</td>
<td>Anterior descending branch left coronary artery</td>
<td>Antero-septal region</td>
<td>Lower apical part of septum</td>
<td>Gross oedema</td>
<td>2400</td>
<td>Present</td>
</tr>
<tr>
<td>Case 5 (female)</td>
<td>480</td>
<td>Anterior descending branch left coronary artery</td>
<td>Antero-septal region</td>
<td>Lower apical part of septum</td>
<td>Acute oedema</td>
<td>1400</td>
<td>Present</td>
</tr>
<tr>
<td>Case 6 (male)</td>
<td>400</td>
<td>Atheroma and calcification left coronary. Right coronary abnormally small</td>
<td>Lower half of septum</td>
<td>One inch above apex in septum</td>
<td>Oedema with terminal pneumonia</td>
<td>900</td>
<td>Central zonal congestion</td>
</tr>
<tr>
<td>Case 7 (female)</td>
<td>350</td>
<td>Gross atheroma. Haemorrhage into plaque in left anterior descending coronary artery</td>
<td>Anterior wall left ventricle and apical portion of septum</td>
<td>Apical part of septum</td>
<td>Acute oedema</td>
<td>1600</td>
<td>Central zonal congestion</td>
</tr>
</tbody>
</table>

## CASE REPORT

G. McK., a 56-year-old man, was admitted to hospital with a diagnosis of internal haemorrhage. On examination he was seen to be shocked and cyanosed. His conjunctivae were blanched to such an extent as to suggest profound anaemia. The haemoglobin was 99 per cent (14.1 g./100 ml.). The cardiac rhythm was regular, the pulse volume poor, and there was gross jugular venous congestion. The blood pressure was not recordable. A loud systolic murmur was maximal in the fourth left intercostal space just to the left of the sternum. Pulmonary oedema was present and the liver was considerably enlarged. There were 15,000 leucocytes/cu. mm. A chest X-ray showed moderate cardiac enlargement and pulmonary congestion. An electrocardiogram showed a typical antero-septal left ventricular infarct (Fig. 1). The blood urea was 147 mg. per 100 ml. Four days later, the patient became disorientated and increasingly cyanosed, and icterus was noted. A further electrocardiogram revealed no extension of the infarction. He lapsed into coma and his condition deteriorated further. Three days later he died. The records of this patient revealed that he had been treated for hypertension several years previously at which time his blood pressure was 240/140 and an electrocardiogram showed only hypertensive changes. His heart sounds at that time were normal.

**Post-mortem Findings.** The body was icteric. There was a fibrinous pericarditis with an adhesion on the anterior surface of the left ventricle (Fig. 2). The heart weighed 550 g. There was ante-mortem
thrombus in the appendage of the dilated right atrium. The right ventricle had an opening in the antero-superior part of the septal wall, with surrounding discoloured and hemorrhagic myocardium. The left atrium was normal. The left ventricle was dilated, especially in the septal region where the dilatation was aneurysmal. It was here that perforation had occurred. The apical third of the septum, the anterior wall of the left ventricle and a small portion of the posterior wall at the apex were discoloured and thin. Apical endocardial thrombi were present. The anterior descending branch of the left coronary artery contained a thrombotic occlusion near its origin.

Histologically the infarcted areas showed extensive necrosis of the myocardium. In some parts the entire thickness of the ventricle was involved with the exception of a narrow surviving epicardial zone. The coronaries were markedly atheromatous and the presence of a recent thrombotic occlusion was confirmed. There was acute pulmonary oedema, macroscopically and histologically.

The liver weighed 1600 g. and had a "nutmeg" pattern. Histologically the liver showed intense central zonal congestion and necrosis (Fig. 3).

**Discussion**

Sager (1934) found that only 3 per cent of myocardial ruptures occurred through the interventricular septum. This infrequency of septal rupture has been ascribed by Gross (1921) to its rich anastomotic blood supply. Most authors agree that septal rupture is found in the presence of severe coronary arteriosclerosis. However, an assessment of this problem must include consideration of other factors. The splinting effect of the right ventricular pressure in reducing the pressure gradient across the septum tends to reduce the incidence of interventricular septal rupture.
On the other hand, if hypertension persists after the occurrence of septal or non-septal infarction, ventricular rupture is more likely to result (Edmondson and Hoxie, 1942).

The following factors appear to contribute to this complication of cardiac infarction.

1. Complete or almost complete infarction of the full thickness of the septal muscle.

2. The time factor. Most septal ruptures occur between the third and twelfth days after infarction. Edmondson and Hoxie (1942) found the average survival period to be 7·4 days.

3. Age. The average age of the patients in this series was 70 years.

4. Hypertension. This is difficult to assess clinically because many of these patients are admitted for the first time after the occurrence of infarction with a consequent fall in blood pressure. The heart weight provides an index of the degree of cardiac hypertrophy present at death. In this series the average heart weight was above 475 g.

The pathological effects of septal rupture are of considerable interest. Intense vascular congestion of the liver lobules occurs and in five of our cases it was associated with central zonal necrosis of the liver lobules (Fig. 3). In our first case the degree of liver damage was such as to result in obvious jaundice. The septal rupture is usually small. Sager (1934) commented on the fact that the precordial murmur is often loud, despite the patient's poor general condition and that it is usually maximal to the left of the lower end of the sternum.

Nareff et al. (1950) stated that it might be confused with that of aortic stenosis, mitral insufficiency, or rupture of chordae tendineae or papillary muscles. Fowler and Failey (1948), discussing
rupture of a papillary muscle, state that this complication is to be suspected when the condition of the patient suddenly becomes worse, the murmurs become louder and nearer the apex, and the ensuing heart failure is left-sided rather than right-sided. They considered that rupture of chordae tendineae was more likely in middle-aged or elderly patients who, suddenly, without dramatic incident, develop a loud systolic murmur, usually with a thrill, maximal at the cardiac apex and along the left sternal border. Congestive heart failure in these cases does not usually develop for months or years.

Rupture usually occurs in the apical end of the septum near one or other wall of the ventricle. This was held by Nareff et al. (1950) to account for the rarity of intraventricular conduction defects. They mention that rupture is often associated with a dramatic deterioration in the patient’s general condition with peripheral circulatory failure and the rather abrupt appearance of signs of right ventricular failure. In our experience, the shock and peripheral circulatory failure following septal rupture have been so marked as to lead to an erroneous clinical impression of anaemia on several occasions.

It would appear that septal rupture is more likely to occur in the elderly hypertensive patient with advanced coronary arteriosclerosis, who has suffered cardiac infarction during the previous fortnight. The prognosis seems to be uniformly poor. All the patients in this series died within a few weeks of septal rupture.

**SUMMARY**

The clinical and pathological findings in seven patients who had cardiac infarction involving the interventricular septum with consequent septal rupture are presented.

The causation and clinical features of this unusual complication of cardiac infarction are discussed. Suggestive clinical signs are the appearance of a loud praecordial systolic murmur, maximal to the left of the sternum in the fourth intercostal space, and often an associated systolic thrill in this area. At the time of septal rupture the patient’s condition deteriorates conspicuously; there is usually profound shock, with peripheral circulatory failure and right heart failure.

We are indebted to Professor Biggart and Professor Bull for their ready advice during the preparation of this paper. We wish to thank Professor Biggart, Dr. Robert Marshall, Dr. T. H. Crozier, Dr. T. A. Kean and Dr. George Adams for permission to publish details of these patients, and Mr. David Mehaffey, A.R.P.S., for the photography.

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