THE CIRCULATORY EFFECTS PRODUCED IN A PATIENT WITH PNEUMOPERICARDIUM BY ARTIFICIALLY VARYING THE INTRAPERICARDIAL PRESSURE

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THE present understanding of the circulatory effects of pericardial effusion is based upon the work of Cohnheim. He noted a rise in venous pressure and a fall in the arterial blood pressure when he increased the intrapericardial pressure by introducing oil into a dog's pericardium. This has since been confirmed by others on many different laboratory animals. In man, however, studies of the effects of increased intrapericardial pressure on the dynamics of the circulation have been limited to a few isolated observations. On two occasions, in the same patient, Stewart, Crane, and Deitrich made simultaneous observations of the intrapericardial pressure and the venous pressure. They also demonstrated a reduction in the venous pressure and a proportional increase in the cardiac output following paracentesis of the pericardium. Caughey, and others, likewise demonstrated a reduction in the venous pressure after pericardial paracentesis in cases of pericardial effusion. Zuccola measured the reduction in intrapericardial pressure following pericardial paracentesis. We have had the opportunity of studying the effects of a sequence of changes in the intrapericardial pressure on the human circulation under conditions more readily controlled.

A twenty-year-old white woman was admitted to the University of Michigan Hospital in July, 1938, with tuberculous polyserositis. Therapeutic pneumopericardium and left-sided therapeutic pneumothorax were established. Following this she improved, and for three months prior to this study her condition was relatively stable. There was no dyspnea, cyanosis, or edema; the ascites, which was present on admission, disappeared after the establishment of the pneumopericardium and had not recurred. The liver, which had been considerably enlarged, became smaller, but could still be felt three fingerbreadths below the costal margin. The roentgenogram, which was taken twenty days before this study, showing the pneumopericardium and left-sided pneumothorax, is reproduced in Fig. 1. In the attempt to maintain a dry pneumopericardium, an opportunity was presented to establish the limits of pressure which would not unduly embarrass the circulation. Consequently, observations were made of the changes in pulse rate, respiratory rate, arterial pressure, venous pressure, and circulation time when the intrapericardial pressure was altered over a fairly large range by the injection of air.

METHODS

By fluoroscopic observation it was found that a needle could be introduced safely into the air-containing pericardial sac through the subxiphoid route with the patient in the supine position. With the needle in place, the intrapericardial pressure
could then be varied by the introduction or removal of air by means of a pneumothorax apparatus. A Y tube from the needle permitted measurements of the intrapericardial pressure on a water manometer and simultaneous tambour recording on a kymograph of the pressure changes produced by the respiratory and cardiac cycles. The Erlanger apparatus was used to make a kymographic record of the pulse in the leg. The principle employed in the Erlanger apparatus was also used to maintain a relatively constant sensitivity of the tambour connected with the intrapericardial space throughout the period of observation. To accomplish this, a small balloon in a glass chamber, which could be adjusted to atmospheric pressure, was placed in the circuit, and this permitted changes in the intrapericardial pressure without changes in the recording level of the tambour. The pulse rate was also counted by palpating the radial artery. Arterial pressure was measured with a mercury manometer by palpation of the radial pulse, since the auscultatory method was not satisfactory in this patient. The respiratory rate was counted for periods of one minute and also recorded on the kymograph. The venous pressure measurements were made by the direct method, as modified by Lyons, Kennedy, and Burwell, using the right antecubital vein. Circulation time was measured by the arm-to-tongue method, using decholin.

The intrapericardial pressure was increased in step-like fashion by the addition of air to the pericardial sac. After each change in pressure, time was allowed for apparent stabilization of the consequent circulatory changes.

It should be made clear that in this study the substance used to elevate the intrapericardial pressure was air. As air, in contradistinction to fluid, is easily compressible and has a lower viscosity, pneumopericardium and hydropericardium may well have different effects upon ventricular filling, even at the same pressure levels. The pressure relationships reported here, therefore, may not be directly transferable to pericardial effusion, but the direction of the changes will be the same.

At the time of these observations the patient suffered from some degree of impairment of cardiac function, as evidenced by the slightly elevated venous pressure, tachycardia, and hepatomegaly. The abnormally high intrapericardial pressure that was originally present had been somewhat reduced by the replacement of the effusion with air. Nevertheless, in order to maintain the pneumopericardium, the intrapericardial pressure had been kept above atmospheric pressure. Some impairment in cardiac function could therefore be expected, since Beck and Isaac have demonstrated that merely exposing the heart to atmospheric pressure produces a diminution in cardiac output in dogs.

**RESULTS AND DISCUSSION**

The variations of venous pressure, pulse rate, arterial pressure, and circulation time with changes in the intrapericardial pressure are illustrated in Graph I. There were no significant changes in the rate or character of respiration throughout the entire period of observation, hence it is not included in the graph.

*Intrapericardial Pressure.*—On entering the pericardium, the pressure was found to fluctuate between +50 mm. of water during inspiration and +100 mm. during expiration. This was in accord with many previous observations on this patient. Table I shows the step-like changes in intrapericardial pressure, the amount of air added or removed to produce each change, and the inspiratory, expiratory, and mean intrapericardial pressure at each level. For convenience in discussing the
Fig. I.—Posteroanterior and left lateral roentgenograms of the chest, taken seven months after admission to the hospital, four months after the establishment of pneumopericardium, and twenty days prior to the observations here reported.
various pressure levels hereafter, the mean intrapericardial pressure will be used.

Once a pressure level in the pericardium was established, it remained constant until altered by further addition or removal of air. There was no evidence of a change in pressure as a result of pericardial stretch, although this has been noted in experimental animals. The fluctuations in intrapericardial pressure caused by the heartbeat were too rapid to be measured on a water manometer, but they were well recorded on the kymograph (Fig. 2). The respiratory fluctuations in intrapericardial pressure, since they were slower, could be measured on a water manometer as well as recorded on the kymograph. Thus, a comparison of the recorded fluctuations caused by the heartbeat with those caused by respiration gave a rough index of the pressure changes within the pericardium produced by systole and diastole. As the intrapericardial pressure was increased, the changes in pressure caused by the heartbeat were less marked, suggesting that there was a decrease in diastolic filling.

Graph I.—Each measurement of intrapericardial pressure is represented by a rectangle which illustrates the range of pressure change caused by respiration. For the purpose of simplicity, the pulse rate and arterial pressure are placed on the same scale as intrapericardial pressure. This has the result of minimizing the graphic representation of the changes. The time, in minutes, is from the beginning of the observations.

As the intrapericardial pressure was increased, the fluctuations caused by respiration were decreased. Thus, with a mean intrapericardial pressure of 75 mm. of water the respiratory fluctuations were in the region of 50 mm., but at higher pressures these fluctuations were reduced to 25 mm. It seems possible in this case that the decrease of intra-
pericardial pressure accompanying inspiration facilitated diastolic filling of the heart. Thus, a reduction in the respiratory fluctuation of intrapericardial pressure may be a factor in determining the level of intrapericardial pressure at which severe tamponade will occur.

Venous Pressure.—It can be seen in Graph II that, at the beginning of the observations, the venous pressure was approximately 95 mm. of water above the mean intrapericardial pressure. Increasing intrapericardial pressure to 145 mm. did not affect the venous pressure appreciably. At this point, however, the difference between venous pressure and intrapericardial pressure was about 35 to 40 mm. of water. A subsequent increase in intrapericardial pressure produced a proportionate increase in the venous pressure. With the final elevation of the intrapericardial pressure to 265 mm., the increase in venous pressure was not proportionate, and the difference between them fell to 25 mm. After about six minutes under these conditions, unequivocal signs of severe tamponade became evident. The patient appeared rather pale, there was cyanosis of the mucous membranes, and she became quite faint. The pulse became rapid and thready, and the arterial pressure fell. It appears plausible that in this case a venous pressure of 35 to 40 mm. above the pressure in the pericardium was necessary for the maintenance of an adequate circulation.
When the pericardial pressure was reduced by the rapid removal of air, the venous pressure fell, but remained 40 to 50 mm. above the intrapericardial pressure, and the symptoms of severe tamponade were quickly relieved. Further reduction in the intrapericardial pressure, to approximately its original level, was accompanied by a fall in venous pressure, also to its original level.

**Circulation Time.**—As is shown in Graph III, the circulation time increased from nineteen seconds to a maximum of thirty-eight seconds. As was noted with venous pressure measurements, no significant changes were manifest until the intrapericardial pressure was elevated above 145 mm. of water. This is better illustrated in Graph IV, in which the venous pressure and circulation time are plotted against the mean intrapericardial pressure.

**Pulse Rate.**—With each increase in the intrapericardial pressure there was an increase in the pulse rate, but the rate tended to return toward its original level after a few moments (Graph V). Like the venous pressure and circulation time, the pulse rate was not markedly affected until a mean intrapericardial pressure of 145 mm. of water was attained. With further increases in pressure there was a decided increase in pulse rate, reaching a maximum of 148. With the sudden release of pressure in the pericardium, just after the onset of severe tamponade, there was

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<tr>
<th>VOLUME AIR ADDED OR REMOVED (C.C.)</th>
<th>TOTAL AIR ADDED OR REMOVED (C.C.)</th>
<th>INTRAPERICARDIAL PRESSURE MM. WATER</th>
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<td>+110</td>
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*Air released through stopcock—not measured.
Graph IV.—The venous pressure in mm. of saline and the circulation time in seconds (ordinate) are plotted against the mean intrapericardial pressure in mm. of water (abscissa). It can be seen that no significant changes occur in either venous pressure or circulation time until the mean intrapericardial pressure is elevated above 145 mm. of water.

Graph V.—See legend to Graph I.
a striking but momentary fall in the pulse rate from 148 to 116, which was slightly below the original rate. This accompanied a rather prompt fall in the venous pressure and a diminution in the visible and palpable distention of the neck veins. This temporary fall in the pulse rate must, therefore, have coincided with a marked increase in the diastolic filling of the heart.

Arterial Pressure.—The arterial pressure tended to fall with each increase in intrapericardial pressure, and to return to its previous level in the next few moments. When the mean pressure in the pericardium was 266 mm. of water, the arterial pressure fell slowly throughout the six minutes of observation. The low level of 74 mm. of mercury was recorded coincident with the release of air from the pericardium.

SUMMARY

1. Observations on changes in pulse rate, respiratory rate, arterial pressure, venous pressure, and circulation time were made with artificially produced changes in intrapericardial pressure in a patient with pneumopericardium. These relationships have been illustrated graphically.

2. The intrapericardial pressure fluctuated with respiration, and, as higher pressures were reached, these fluctuations decreased.

3. The intrapericardial pressure fluctuated with systole and diastole. Kymographic tracings were made of the fluctuations in intrapericardial pressure caused by respiration and the heartbeat. With the higher intrapericardial pressures, the changes in pressure produced by systole and diastole were less marked.

4. Significant changes in pulse rate, arterial pressure, venous pressure, and circulation time did not occur until the intrapericardial pressure was elevated to, or above, 145 mm. of water.

5. In order to maintain the circulation, it was necessary that venous pressure exceed intrapericardial pressure by at least 35 to 40 mm. of water.

REFERENCES


