Case report

Chylopericardium: a rare complication of a Waterston shunt

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SUMMARY A case of chylopericardium occurring after a Waterston shunt was successfully managed by pericardial drainage of chyle and substitution of dietary fats with medium chain triglycerides.

Chylous pleural effusion is a well recognised complication after thoracic surgery. The accumulation of chyle in the pericardium sufficient to cause tamponade, however, is a rare complication of cardiac surgery and was first described in 1971 by Thomas and McGoon.1 The aetiology of chylopericardium is uncertain and because the condition is so rare the correct treatment has yet to be established. We report a case of this unusual complication in a 5 month old infant after a Waterston shunt and discuss the management of this condition.

Case report

A male infant weighing 1.8 kg was born at 33 weeks' gestation after the spontaneous onset of labour. He was small for dates, but there was nothing unusual in the antenatal history. Cyanosis was not clinically apparent, an electrocardiogram showed sinus rhythm with left ventricular dominance, and the chest x ray film showed an increased cardiothoracic ratio. A systolic murmur was heard and he was referred for assessment. Cardiac catheterisation showed pulmonary atresia with a hypoplastic right ventricle, widely patent ductus arteriosus, and a large atrial septal defect. As his condition was stable, surgical intervention was not warranted and he was discharged. He was readmitted at the age of five months because of failure to gain weight and mild cyanosis, and it was felt that a palliative shunt was required.

A standard Waterston shunt by the retrocaval approach was carried out through a right lateral thoracotomy. The surgical procedure was uneventful and a good shunt murmur was audible after operation. The pleural drain was removed the following day. Two days later, however, an increase in heart size was noted on the chest x ray film; one week postoperatively the presence of a large pericardial effusion was confirmed by echocardiography but there were no specific features to suggest chylopericardium. Despite increased diuretic therapy the size of the effusion as assessed by echocardiography continued to increase.

Three weeks postoperatively surgical drainage was performed through a small sub-xiphoid incision; 200 ml of milky fluid was removed from the pericardium and a pericardial drain was left in situ. Laboratory analysis of the aspirated fluid showed the presence of chylomicrons and this confirmed the clinical diagnosis of chylopericardium. The patient was started on a diet in which dietary fats were replaced by medium chain triglyceride supplements. About 20 ml of chyle drained away each day for the next five days; drainage ceased after one week and the drain was removed. An echocardiogram two weeks later confirmed that no fluid had reaccumulated in the pericardium. Follow up three months later showed a heart of normal size on chest x ray film and normal echocardiographic appearance.

Discussion

The first case of chylopericardium after cardiac surgery was reported in 1971 by Thomas and McGoon.1 Feteih et al reviewed nine further cases and added their own.2 Since this review, further
cases have been added by Rose et al and Pugliese et al. These 13 cases span the full range of ages from infants to adults and cover the whole spectrum of cardiac operations from palliative shunts to definitive procedures.

Our case, like those reported by Hawker et al in 1972 and Jacob and colleagues in 1974, occurred in a patient who had a Waterston shunt. The aetiology of chylopericardium after cardiac surgery is not clear but it seems that there is a definite risk of this complication after a Waterston shunt. Transection of an important lymphatic channel is obviously a factor, but such an event has only been described occasionally, despite the high frequency of anatomical variations in the course of the thoracic duct. In the case described by Thomas and McGoon, thrombosis of the left subclavian vein secondary to insertion of an external jugular vein cannula was described and they believed that this thrombosis had obstructed the end of the thoracic duct and produced high lymphatic pressures. A similar aetiology was possible in the case described by Kansu et al and Rose et al; however, in our patient venous access was maintained by a right internal jugular cannula.

The interval from operation to appearance of the chylopericardium ranges from four days, as in our case, to five weeks. This suggests that the aetiologica development of the condition depends on whether damage to the major pericardial lymphatic occurs at operation or is a result of later fibrosis. Those who have reported cases of chylopericardium have not agreed on the management of such patients. Providing that no major lymphatic vessel had been damaged, as shown by excessive pericardial drainage or rapid reaccumulation of chyle after aspiration, most cases settled with conservative management. Adequate drainage of the pericardium is essential and this is most effectively established by the insertion of a pericardial drain via the sub-xiphoid approach. Chyle formation can be reduced by controlling the dietary intake of fats. Fats are hydrolysed in the gut to glycerol and fatty acids and these long chain fatty acids are re-esterified to triglycerides and passed into the lymphatics by chylomicra. Medium chain triglycerides, however, are not re-esterified but pass directly into the portal venous blood. Thus the substitution of dietary fats with medium chain triglycerides is an effective method of reducing chyle formation, and this approach was successful in our case and that of others.

References