Acute pericarditis due to perforation of a benign mediastinal teratodermoid into the pericardial sac

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Rupture into the pericardium is a very rare complication of anterior mediastinal teratodermoid. There have been only four recorded cases, which either have been characterised by cardiac tamponade or have been an incidental finding at operation or necropsy. This report concerns a patient who presented with acute pericarditis and is considered to be unique.

Case report

A 32-year-old woman was admitted to hospital with a three-hour history of anterior chest pain. This was sudden in onset, radiated to the neck and arm, and was associated with nausea and vomiting. Before this episode, she had been entirely well.

On admission no abnormality was found during the physical examination. The white cell count was 15·0 × 10⁹/l with a normal differential count and the erythrocyte sedimentation rate was 22 mm in one hour (Westergren). The initial electrocardiogram and cardiac enzyme levels were normal but the following day there was widespread ST elevation and the aspartate aminotransferase activity was raised at 360 U/l. An anteroposterior chest radiograph taken shortly after admission showed segmental collapse at the left base but was otherwise considered normal. A presumptive diagnosis of acute pericarditis was made and an echocardiogram was performed. This showed 1·5 cm of pericardial fluid but no evidence of cardiac tamponade.

A posteroanterior chest radiograph (fig) taken two days after admission showed a small right pleural effusion; a left hilar mass, which on the lateral view was seen to occupy the anterior mediastinum; and a possible right hilar mass. Tomography of this area showed a lobulated mass affecting the left hilum and extending forwards to occupy most of the area between the mediastinum and the anterior chest wall. The appearances were thought to be those of enlarged lymph nodes and in view of the patient's age lymphoma was considered the most likely diagnosis.

Bronchoscopy and mediastinoscopy were performed in an attempt to obtain a tissue diagnosis but the results of both were normal. A left mediastinotomy was therefore carried out by excision of the second costal cartilage. This showed a large cystic mass in the anterior mediastinum, which was closely adherent to the pericardium and mediastinal pleura. When the cyst was opened pultaceous material containing hair was found; digital exploration of the cavity showed extension into the right side of the mediastinum. The skin incision was therefore extended across the midline and the sternum transected to allow removal. It was impossible to separate the cyst from the pericardium, so a large pericardial window was made and the cyst excised in toto.

When the pericardium was opened both serous surfaces were noted to be grossly inflamed with patches of adherent fibrin. The pericardial sac contained hair with pultaceous material similar to that found in the cyst cavity. Although it was obvious that the cyst had ruptured into the pericardial cavity before operation, the point of communication was not found. The histological features were those of a benign mediastinal teratodermoid and the cyst contents were sterile on culture. The patient has remained in good health since the operation.

Discussion

Perforation into an adjacent structure is an unusual but well recognised complication of anterior mediastinal teratoderoids. The bronchus is the most commonly affected site but rupture into the pleura, aorta, superior vena cava, and pericardium have also been described.† Marsten et al² estimated that rupture into adjacent cardiovascular structures by benign teratoderoids occurred in less than 1% of cases.
The first recorded case of rupture into the pericardium was described by Cordes in 1859.³ The patient had a large pericardial effusion with signs of tamponade and died seven weeks after admission when necropsy showed a mediastinal teratodermoid communicating with the pericardial cavity. More than 50 years later a second report concerned a patient with a large teratodermoid which had perforated into the pleural cavity, causing an empyema.⁴ The intrapericardial rupture was found as an associated incidental finding at necropsy.

The first case to be treated successfully by surgical removal of a perforated cyst was reported by Aktan in 1961.⁵ This patient presented with a five-month history of right-sided chest pain and productive cough and was found at operation to have a teratodermoid compressing the middle-lobe bronchus, causing lobar collapse. The cyst was found to have ruptured into the pericardium when this was opened to allow excision. Another case which presented with cardiac tamponade but which was treated successfully by removal of the cyst was described by Marsten et al in 1966.⁶

Thus in two of the previously recorded cases the predominant feature after rupture has been cardiac tamponade, whereas in the other two the primary problem has been related to the lung, the intrapericardial rupture being an incidental finding. This case is therefore unique since the patient presented with acute pericarditis but the volume of fluid in the pericardium was small and tamponade did not occur.

Although there can be little doubt that intrapericardial rupture has occurred in all the cases described, only Cordes was able to identify the point of communication.⁷ Presumably in other cases the site of rupture was small and sealed spontaneously.

Despite the rarity of the condition, the diagnosis of intrapericardial rupture of a teratodermoid should be considered if there are cardiovascular symptoms and signs with an anterior mediastinal mass on the chest radiograph. The accepted treatment, as in uncomplicated cases of teratodermoid, is surgical excision. This can be performed after rupture by removing a pericardial window with the teratodermoid. Such a procedure has led to a successful outcome in two previously recorded cases and in the present case.

References

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