CASE REPORTS

Left atrial myxoma in a child diagnosed by ultra-sound

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Summary

A case of left atrial myxoma in a child is presented. The diagnosis was made using ultra-sound and confirmed at operation.

The results of our investigations are reviewed in the light of other authors’ experience with this disease. The use of ultra-sound in recognizing this condition is discussed.

Left atrial myxoma is the commonest primary cardiac tumour and the clinical features are well documented (Yater, 1931; Goodwin, 1964).

The technique of mitral echocardiography using ultra-sound waves generated by a piezo-electric transducer directed at the anterior cusp of the mitral valve was first described in 1954 by Edler & Hertz. Effert & Domani (1959) showed that echoes arising from a left atrial myxoma were visible behind those of the anterior cusp of the mitral valve in diastole so that the normally echo-free space is filled by multiple dots. Reports of four atrial myxomas occurring in adults and demonstrated by this method (Schattenberg, 1968; Wolfe, Popp & Feigenbaum, 1969; Popp & Harrison, 1969) have appeared previously. This is the first record of a child with an atrial myxoma in whom the diagnosis has been made by ultra-sound.

Case report

A 7-year-old boy was referred by his general practitioner to the North Middlesex Hospital on the 21 February, 1970, with a 1-week history of pyrexia which had not responded to erythromycin. He had pain in the right lower leg and ankle and later similar

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pain on the left side severe enough to prevent him walking or standing. He had had intermittent ankle pain for 1 year. His general health was good and the family history was unremarkable.

On examination his temperature was 37-6°C and he had a malar flush. The pulse rate was 120/min. He was in sinus rhythm and had a blood pressure of 110/70 mmHg. There was no cardiomegaly. Auscultation revealed a variable presystolic murmur and loud first sound. No other abnormality could be detected on full systematic examination.

Investigations

The Hb was 11·3 g/100 ml; WBC 11,800 (52% neutrophils, 30% lymphocytes, 12% monocytes, some atypical, and 6% eosinophils); reticulocytes less than 2%; ESR 85 mm/hr (Westergren). Protein electrophoresis showed raised alpha1 and alpha2 globulins with a normal total protein and albumen content. The ASO titre was less than 50 units/ml and the SCAT was less than 1 in 8. The Paul Bunnel, ANF and Widal tests were negative. Blood and urine cultures were sterile. Urine microscopy showed ten red cells per high power field but no white cells or casts. Chest X-ray, IVP and ECG were normal.

He was treated as if he had rheumatic fever pending results of investigations and given a course of ampicillin and salicylates. However, he continued to have a sleeping tachycardia of 120–130 beats/min, a low grade pyrexia between 37.2°C and 38·7°C and intermittent proteinuria and haematuria.

On the 13 March he had a transient episode in which he became unconscious for 5 min and had a left facial weakness which resolved within a few minutes. On regaining consciousness he vomited and complained of a left frontal headache.

An ultra-sonic mitral echogram performed on the 25 March showed that the valve was not thickened.
and was normally mobile. During some beats the diastolic closure rate was normal at 85 mm/sec (Fig. 1a). At other times the diastolic closure rate was reduced to 30 mm/sec indicating that there was intermittent mitral obstruction (Fig. 1b). Throughout all cardiac cycles, multiple echoes were visible behind the anterior cusp of the mitral valve in diastole. This indicated the presence of a reflecting body behind the anterior valve cusp in diastole, where only blood should be. This is diagnostic of a left atrial myxoma.

On the 31 March he complained of chest pain, vomited and collapsed with pallor, bradycardia (30–40 apex beats/min), absent peripheral pulses and shallow respiration. ECG showed complete heart block with a ventricular rate of 40, ST depression and T wave inversion in lead III. The rate then increased and became nodal with ectopic beats. Sinus rhythm returned later, the ectopic beats disappeared and he recovered consciousness.

Following this episode the leg pulses below the femorals and the left arm pulses could not be felt. Fifteen hours later the SGOT was 97 units and the SLDH 680 units. ECG showed a posterior infarct—(Fig. 2). Cardiac catheterization was performed, and
following an injection of radio-opaque material into the main pulmonary artery, a large filling defect was demonstrated in the right side of the left atrium.

Operation

He was transferred to The Hospital for Sick Children, Great Ormond Street, and exploration of the left atrium was performed on the 4 April by Mr D. J. Waterston using cardio-pulmonary by-pass. A large myxoma measuring 5 cm \( \times \) 4 cm \( \times \) 4 cm arising from the posterior wall of the left atrium was found and excised. After the cardiac incision had been closed, the apex of the heart was elevated and a well demarcated infarct was found on the posterior surface. He had an uneventful convalescence and has now returned to school and normal activity.

The pedal pulses have returned but he still does not have any pulses in the left arm though the circulation appears to be adequate. The ECG has shown evolution of the posterior infarct, the T waves returning to their upright position; the ESR and SLDH have returned to normal levels and the post-operative echogram (Fig. 1c) has no echoes behind the anterior mitral cusp. The tachycardia prevents the diastolic closure rate being measured.

Comment

In spite of the absence of ECG evidence of carditis and lack of associated joint signs, rheumatic fever was initially diagnosed as the most likely cause of tachycardia, limb pains and a high ESR in a child. The ASO titre of less than 50 units was further evidence against rheumatic fever, an opinion recently reinforced by Davis’s view (1970) that a low ASO titre should stimulate the search for an alternative diagnosis.

Subacute bacterial endocarditis in the absence of a past history of rheumatic heart disease or cardiovascular defect is uncommon. It was the search for such a focus producing emboli, \( \text{viz.} \) an abnormal mitral valve, that led to the use of ultra-sound. This apparatus can record echoes from the anterior cusp of the mitral valve and detect abnormalities of anatomy and function. If the atrium contains a tumour protruding through the valve-ring in atrial systole, the characteristic picture seen in Fig. 1a and 1b will be produced. This picture is not seen when the tumour is wholly within the atrium.

Fever, tachycardia and raised ESR are non-specific and the mechanisms of their production in myxoma are obscure. Abnormal serum proteins have also been observed, the abnormality lying usually in an increased gamma globulin, but in this case it was alpha\(_1\) and alpha\(_2\) globulins. Goodwin \( \text{et al.} \) (1962) and Goodwin (1968) postulated a general poisoning of the body tissues by the myxoma, the mechanism being release of some substance (e.g. mucopolysaccharides) into the circulation by the tumour. Also suggested was hepatic anoxia and subsequent liver cell necrosis due to low cardiac output (Goodwin, 1964). Destruction of erythrocytes by the tumour or an auto-immune cause similar to the post-cardiomyotomy syndrome were suggested by Goodwin (1968) to account for the anaemia. A higher reticulocyte count than the 2% found in this child would have been expected.

Discussion

Syncopal attacks due to myxoma are commonly attributed to sudden decrease in cardiac output when the tumour obstructs the mitral orifice. In adults with an atheromatous coronary tree this decrease may be sufficient to cause anoxic myocardial damage and infarction without occlusion. Direct coronary occlusion is another alternative. The well demarcated infarct found at operation seems almost certainly the result of the syncopal attack and rhythm change 4 days earlier. The tumour was considered to be discharging emboli then and although rare, coronary emboli have been reported in this condition. Two cases of myxoma tumour emboli in coronary vessels were described by Harvey (1957) and one by Heath (1968). We feel that this represents a further case. However, Nadas & Curtis-Ellison (1968) described a 13-year-old child who died following sudden onset of heart block. Necropsy showed obstruction to the mitral valve by a myxoma but no coronary embolus.

Kroopf & Peterson (1957) described a patient with left atrial myxoma who developed acute anterior myocardial infarction. Necropsy failed to show coronary occlusion and the authors’ suggested compression of the left coronary artery by displacement by the large tumour in the left atrium.

Large vessel emboli have been reported in both adults and children. Embolectomy was rejected in this case because the circulation to the limb was good. This does raise the theoretical complication of aneurysm-formation at the site of the embolus; Burton & Johnson (1970) have described multiple cerebral aneurysms following embolization of myxoma tissue to the cerebral circulation with proliferation and fragmentation of the intima at the site of impaction.

Although left atrial myxoma is rare, this diagnosis must be considered in any patient who is embolizing without apparent cause, and the ultra sound technique is a simple and atraumatic means to investigate the left atrium.

Acknowledgments

We are grateful to Dr I. G. Wickes and Mr D. J. Waterston for permission to publish the details of this patient under their care, and to Dr B. G. Wells who performed the cardiac catheterization.
Carbenoxolone sodium and the parent substance, liquorice, have an aldosterone-like effect, and may give rise to oedema, hypertension and hypokalaemia (Baron & Nabarro, 1968; Hausmann & Tarnoky, 1968). The hypokalaemia may cause a myopathy, perhaps with myoglobinuria, or a nephropathy or death. Reports of these complications are listed in Table 1.

<table>
<thead>
<tr>
<th>Carbenoxolone sodium and liquorice</th>
<th>Muscle weakness</th>
<th>Nephropathy</th>
<th>Death</th>
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<tbody>
<tr>
<td>Liquorice preparations</td>
<td>Myopathy</td>
<td>With myoglobinuria</td>
<td>Chodkiewicz et al. (1), 1963</td>
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<td>Cayley (1), 1950; Heard, Campbell &amp; Hurley (6), 1950; Strong (2), 1951; Mollaret, Goulon &amp; Tournilhac (1), 1960; Garcia et al. (1), 1961; Giroire et al. (1), 1961; Jenny et al. (1), 1961; Salassa, Mattock &amp; Rosevear (1), 1962; Chodkiewicz, Clay &amp; Hecaen (2), 1963; Minvielle, Cristol &amp; Badach (2), 1963; Holmes et al. (1), 1970</td>
<td>Geerling (1), 1966; Gross, Dexter &amp; Roth (1), 1966</td>
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<td>Carbenoxolone sodium:</td>
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<tr>
<td>as Biogastrone</td>
<td>Myopathy</td>
<td>With myoglobinuria</td>
<td>Mohamed, Chapman &amp; Crooks (1), 1966</td>
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<td>Morgan, Donald &amp; McAndrew (1), 1966†</td>
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<td>Muir, Laithwaite &amp; Wood (1), 1969</td>
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* Serum potassium not estimated in two patients.
† Serum potassium not estimated.
‡ Myoglobinuria assumed because of positive urine occult blood test in the absence of haematuria or haemoglobinuria.
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