Case reports

Ankylosing spondylitis and an aortic arch syndrome

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SUMMARY A 63 year old woman with a 16 year history of ankylosing spondylitis and peripheral joint involvement later presented with a large vessel arteritis affecting the branches of the arch of the aorta.

Cardiovascular complications of ankylosing spondylitis are well recognised. Most often they appear clinically as aortic incompetence and more rarely as cardiac conduction defects.1 2 Arteritis of large vessels such as the branches of the arch of the aorta is rare. We report the case of a patient with ankylosing spondylitis with peripheral joint involvement in whom an arteritis later developed affecting the subclavian and vertebral arteries. Although the aortic wall appeared slightly thickened on an echocardiogram, the aortic valve was normal.

Case report

A 63 year old woman first presented in October 1967 at the age of 47 years with left sacroiliac pain and local tenderness. Pelvic x ray findings and an erythrocyte sedimentation rate were normal. In December 1975 she developed low backache radiating to the buttocks. The erythrocyte sedimentation rate was 120 mm/first hour with a normal blood count. The Rose-Waaler and rheumatoid arthritis latex test results were negative, and x ray films of the spine and pelvis showed no abnormality.

In February 1976 she developed a generalised illness with myalgia, malaise, stiffness of her shoulders and hips, and back pain radiating to the thighs. She had a systolic murmur over the aortic area radiating to the neck. Apart from a proximal weakness there were no other neurological signs. Her haemoglobin concentration was 10·6 g/dl; white blood cell count was 13·9×10⁹/l and the film showed rouleaux with a normochromic normocytic anaemia. The erythrocyte sedimentation rate was 108 mm/first hour. Serum protein concentrations and protein electrophoresis were normal. She remained seronegative; Paul Bunnell, brucella, and viral studies were negative. Antinuclear antibodies were absent. X ray films of the pelvis showed calcification at the insertion of the hip adductors.

Treatment with prednisolone was started, but two years later she developed pain, stiffness, and swelling of the finger joints of both hands, and effusions of both knees, suggesting that the episode of 1976 was a myalgic onset of a seronegative arthritis. In October 1978 she was admitted because of pain in her left hip, knees, and hands. There was reduced internal rotation of both hips and swelling of the proximal interphalangeal joints of her hands. The blood pressure in the right arm was 120/80 mm Hg. Her erythrocyte sedimentation rate remained raised, and her serological tests remained negative. X ray films showed joint space narrowing in the metacarpophalangeal joints and erosions of both ulnar styloids. There was also joint space narrowing in the left hip and sclerosis of the sacroiliac joints with possible erosions. There was osteophytic lipping between the third and fourth lumbar vertebrae. She was treated with indomethacin.

In 1979 she had an episode of vaginal bleeding and was subsequently found to have an adenocarcinoma of the uterus. This was treated by hysterectomy and postoperative radiotherapy. There has been no clinical evidence of recurrence. Over the next two years she had an episode of tenosynovitis of the flexor tendons of the right hand and also pain in her left subtalar joint. Her left hip continued to be painful and further x ray films showed sclerosis and joint space narrowing with lesser changes on the right. The sacroiliac joints showed periarticular sclerosis with erosive changes, and her lumbar spine showed syndesmophytes consistent with the diagnosis of ankylosing spondylitis (Fig. 1).

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In October 1981 at the age of 61 years she developed pain in her arms while wringing out some washing. The pain in her right arm continued, and her hands felt cold. No subclavian, brachial, or wrist pulses could be felt. All other pulses were present. Her arms were warm and appeared adequately perfused. By January 1982 she had developed bilateral claudication of the forearms with delayed capillary filling. The erythrocyte sedimentation rate was 150 mm/first hour. An arch aortogram showed occlusion of both subclavian arteries beyond the origin of the vertebral arteries. The vertebral arteries themselves showed considerable narrowing at their origins (Fig. 2). The carotid arteries were normal. There were poor collateral channels with no evidence of vessel filling distal to the block. These changes were consistent with an aortic arch syndrome. She was treated medically with aspirin.

She was again admitted in September 1982 with increasing pain in her left hip. The aortic murmur was unchanged but she had developed bruits in both carotid arteries. An echocardiogram showed some thickening of the aortic wall, but the aortic valve was normal and there was no evidence of stenosis. There was reduced movement of the distal and proximal interphalangeal joints. Hip movements were reduced. The haemoglobin concentration was 10.6 g/dl; white blood cell count $10.5 \times 10^{9}$/l and platelet count $851 \times 10^{9}$/l. The erythrocyte sedimentation rate was 115 mm/first hour, and she remained seronegative. In November 1982 she had a left total hip replacement. Blood pressure during the operation was monitored with a thigh cuff and a Doppler method. Histology of the joint capsule showed extensive synovial damage with a fibrinous inflammatory exudate overlying granulation tissue. She was typed for the HLA antigens and showed A28, Aw19, B5, and B40.

Discussion

Cardiac manifestations of ankylosing spondylitis in the form of aortic and mitral valvular disease and heart block have been well documented.\(^1,2\) Involvement distal to the aortic root has, however, been reported only rarely.\(^3-6\) Ankylosing spondylitis in association with an aortic arch syndrome has been reported by Paloheimo et al\(^7\) in four patients and Ghozlan et al\(^8\) in one (Table).

Although these patients were described as having Takayasu's arteritis, a disease generally of young women,\(^9\) they include a man of 46 years. Our patient was 61 years old at onset, whereas the other women were aged less than 25 years. The older age of our patient and the history of a myalgic episode at the onset of her arthritis raises the possibility of a giant cell arteritis. The two diseases may be difficult to differentiate histologically,\(^10\) but the subsequent joint involvement does not favour giant cell arteritis. The delay between the onset of the arteritis and the

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**Fig. 1** Radiograph of pelvis showing sacroilitis and lumbar syndesmophytes.

**Fig. 2** Arch aortogram showing absence of filling of the subclavian arteries and narrowing of the origins of the vertebral arteries.
Ankylosing spondylitis and an aortic arch syndrome

Table  Reported cases of ankylosing spondylitis and an aortic arch syndrome

<table>
<thead>
<tr>
<th>Author (Case No)</th>
<th>Sex</th>
<th>Age (yr)</th>
<th>Arterial stenosis and murmurs</th>
<th>Changes on radiology</th>
<th>ESR (mm/h)</th>
<th>Delay in aortitis after arthritis (yr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paloheimo et al</td>
<td>M</td>
<td>46</td>
<td>Both subclavian, left carotid, both vertebral, both renal</td>
<td>Sacroiliitis, syndesmophytes, lumbar spine, peripheral joints</td>
<td>67</td>
<td>4</td>
</tr>
<tr>
<td>2</td>
<td>F</td>
<td>24</td>
<td>Right subclavian, systolic murmur</td>
<td>Sacroiliitis</td>
<td>112</td>
<td>3</td>
</tr>
<tr>
<td>3</td>
<td>F</td>
<td>25</td>
<td>Both subclavian, both carotid, systolic murmur</td>
<td>Sacroiliitis, lumbar spine</td>
<td>20-120</td>
<td>4</td>
</tr>
<tr>
<td>4</td>
<td>F</td>
<td>24</td>
<td>Both subclavian, left vertebral, both carotid</td>
<td>Sacroiliitis, lumbar spine</td>
<td>74</td>
<td>No arthritis</td>
</tr>
<tr>
<td>Ghozlan et al</td>
<td>F</td>
<td>22</td>
<td>Left subclavian, left vertebral, left carotid, left renal</td>
<td>Sacroiliitis, knees</td>
<td>30</td>
<td>3</td>
</tr>
<tr>
<td>Present case</td>
<td>F</td>
<td>61</td>
<td>Both subclavian, both vertebral, systolic murmur</td>
<td>Sacroiliitis, lumbar spine, peripheral joints</td>
<td>115</td>
<td>3</td>
</tr>
</tbody>
</table>

development of joint symptoms was 3-4 years in all but one case. All cases except one had a raised erythrocyte sedimentation rate during the acute phase of the arteritis. Most had characteristic systemic symptoms. Three patients developed sacroiliitis concurrently with the arteritis and, in two, these features were asymptomatic. It is apparent that some patients develop sacroiliitis concurrently with the systemic phase of the arteritis, whereas others may manifest an arteritis several years after the onset of their ankylosing spondylitis. Our patient was negative for the HLA B27 antigen, and it is of interest that the only other case tissue typed was also negative for this antigen.8

A patient has recently been reported with a seronegative spondyloarthropathy presenting with Reiter's syndrome and sacroiliitis, who subsequently developed a histologically confirmed aortitis proximal to the aortic bifurcation.11 Eighteen months later after surgical excision and grafting the patient developed a large joint arthritis and sacroiliitis. Aortic arch syndromes have also been associated with rheumatoid arthritis in the past.1213 Our patient showed evidence of an arteritis of the major arch vessels, lumbar syndesmophytes, and sacroiliitis. She also had a symmetrical peripheral arthritis with erosion of the ulnar styloid. Although this would suggest rheumatoid arthritis, erosions have been reported in seronegative spondyloarthropathies including ankylosing spondylitis14 and do not preclude the diagnosis.

Joint pains and synovitis may occur with the onset of the acute phase of the arteritic disease and is estimated by Serre et al to occur in 25% of cases.15 This present case represents the sixth case of ankylosing spondylitis in association with an aortic arch syndrome presumed to be inflammatory in origin.

References

4 Duvernoy WFC, Schatz J. Rheumatoid spondylitis associated with aneurysmal dilatation of the entire thoracic aorta. Henry Ford Hospital Medical Bulletin 1966; 14; 309–12.
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