Case report

Successful valve replacement for aortic incompetence in rheumatoid arthritis with vasculitis

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SUMMARY A patient with rheumatoid arthritis (RA), vasculitis, and aortic valve incompetence of the histologically nonspecific type is described. Valve replacement was undertaken, and an excellent haemodynamic result was achieved. Both arthritis and vasculitis subsequently remitted. Valve replacement can be successfully performed in RA despite active vasculitis.

Cardiac valvular disease in association with rheumatoid arthritis (RA) has been described for some time (Baggenstoss and Rosenberg, 1941, 1944). The aortic valve may be involved, either with granulomata (Iveson et al., 1975) or nonspecifically (Lebowitz, 1963). To date 10 cases of aortic valve replacement have been reported (Barker 1971; Iveson et al., 1975; Yates and Scott, 1975; Barracough et al., 1975; Iveson and Pomerance, 1977). Only 1 of these had concommitant vasculitis, and this patient died after surgery (Yates and Scott, 1975). The last authors rightly emphasised the danger of surgery when systemic features are present. We report a case of successful aortic valve replacement for incompetence in a patient with RA and active vasculitis.

Case report

A 54-year-old man was admitted because of progressive symptoms of the left ventricular failure due to aortic incompetence. He had a 10-year history of seropositive, nodular, erosive RA. The onset of arthritis was peripheral and symmetrical and gradually involved almost all joints. The course had been remitting but progressive and destructive. The only treatment was with nonsteroidal anti-inflammatory analgesics. Three years prior to admission he developed breathlessness to exertion and at rest, and these symptoms rapidly deteriorated. There was no history of rheumatic fever.

Physical examination revealed rheumatoid deformities with moderately active synovitis, nodules, and vasculitic lesions of hands and feet (Fig. 1). Blood pressure was 170/60 mmHg. Jugular venous pressure was raised 6 cm. He had a collapsing pulse. The heart was enlarged. An early aortic diastolic murmur and an Austin-Flint murmur were heard. There were basal crepitations. Severe aortic incompetence was confirmed by echocardiography and cardiac catheterisation. Other investigations showed Hb 13·9 g/dl, leucocytes 7·0 × 10⁹/l, ESR 52 mm in 1 hour, Rose-Waaler titre 1:256, antinuclear antibody positive, admission he developed breathlessness to exertion and at rest, and these symptoms rapidly deteriorated. There was no history of rheumatic fever.

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native DNA binding 2.8% by Farr assay. A chest roentgenogram showed congestive changes. An electrocardiogram showed left ventricular hypertrophy. Multiple blood cultures failed to isolate a pathogen. His cardiac failure was unsatisfactorily controlled with an anticoagulant regimen of digoxin 0.25 mg and frusemide 80 mg daily with potassium supplementation. Dyspnœa and physical signs of aortic regurgitation remained prominent. Aortic valve replacement was considered necessary and was performed 3 months after presentation.

At operation the valve leaflets were grossly thickened, shrunken, and retracted. There were no vegetations. Valve replacement by a Starr-Edwards prosthesis was successfully carried out. The valve histology showed fibrous thickening with foci of polymorphonuclear leucocytic, lymphocytic, and plasma cell infiltration (Fig. 2).

Postoperatively cardiac function has remained excellent. Sodium aurothiomalate (Myocrisin) was started just before surgery and in the subsequent 18 months follow-up period the patient has had a remission of his arthritis and the vasculitis has resolved.

Discussion

Valvular heart disease in RA is well recognised (Iveson et al., 1975). The aortic and mitral valves are most commonly effected (Roberts et al., 1968). The lesions appear clinically significant in about one-third of cases. The commonest symptom detected is congestive cardiac failure (Sokoloff, 1964). Of the 10 cases of aortic valve replacement reported granulomata were seen in 4 and a non-specific valvulitis in 6. Successful valve replacement was obtained in 9 of the 10 cases, and this made possible the continuation of physical therapy for arthritis.

The danger of surgery for valvular disease when systemic features accompany inflammatory disorders of connective tissue was stressed by Yates and Scott (1975). Of the 3 cases they reported 2 died after operation as a result of systemic disease. One had systemic lupus erythematosus and nephritis and the other RA and vasculitis.

Our patient had a disease which was seropositive, nodular, erosive, and active. There was obvious vasculitis at the time of surgery. There was no past history of rheumatic fever. The valvular changes were consistent with histologically nonspecific type. The result of aortic valve replacement was excellent, with return of normal exercise tolerance. Furthermore, with the subsequent continuation of gold therapy good remission was obtained, and vasculitis disappeared. Our case is the first illustration that if there is severe aortic incompetence requiring replacement, surgery is worthwhile and technically feasible despite accompanying rheumatoid vasculitis.

References


Fig. 2 Aortic valve leaflet showing mononuclear cell infiltration (×175)