Discussion

Pauker et al. (1972) described 2 cases of BCG osteomyelitis and osteoarthritis of the knee joint with positive pathologico-anatomical findings but negative bacteriological findings. In the present case the infection seemed to be of low virulence, no regression was seen after several weeks' treatment with antibiotics effective against Gram-positive cocci, and fewer than 4 years had elapsed since the BCG vaccination. In accordance with the criteria of Foucard and Hjelmstedt (1971) we therefore concluded the lesion was probably BCG osteomyelitis. Since our experience and that of others (Paisseau et al., 1941; Backman and Wallgren, 1954; Felländer, 1963; Foucard and Hjelmstedt, 1971) was that even biopsy may not give the diagnosis with certainty we decided to try conservative treatment. The prompt healing with tuberculostatic treatment was in our opinion sufficient evidence that the diagnosis and management were correct.

On the basis of this case a 2-month-old baby girl with an osteolytic focus of the humerus near the site of the BCG inoculation was recently treated elsewhere in the same conservative way with good results.

References


M. Pauker,* M. Seelenfreund, and G. Morain Department of Orthopedic Surgery and Traumatology, Beilinson Medical Center, Sackler School of Medicine, Tel Aviv University, Petah Tikva, Israel.

*Correspondence to Dr. M. Pauker.

Summary

A 3-year-old boy presented with an osteolytic lesion and periosteal reaction in the distal metaphysis of the right femur which failed to respond to immobilization and intensive antibiotic treatment. Since the infection seemed to be of low virulence and fewer than 4 years had elapsed since BCG vaccination BCG osteomyelitis was suspected. Tuberculostatic treatment led to rapid recovery and surgical measures were unnecessary.

Symptomatic sinus arrest in a young girl

A recent report (Scott et al., 1976) on symptomatic sinoatrial node dysfunction noted its infrequency in children and no reported occurrence of the
syndrome in girls. The following case is therefore presented.

Case report

A 13-year-old girl was first seen at 9 years of age because of an asymptomatic persistent ductus arteriosus. At this time electrocardiograms (ECG) were normal, showing rates from 72–80/min, normal P waves, and PR interval 140 ms. Ligation of the ductus was accomplished uneventfully. Postoperatively there were no arrhythmias and an appropriate increase in heart rate to stress. One year postoperatively there was no clinical or x-ray evidence of heart disease and ECG was unchanged. She was therefore discharged.

She reappeared 3 years later because of recurrent syncopal attacks of 2 years' duration. Attacks were described as transient loss of consciousness, unrelated to activity and unaccompanied by other manifestations. These episodes had occurred two to three times a month during the first year and three to four times a month in the second year. There had been no known illness before the onset of the syncopal attacks. Drug use was denied. There was no family history of arrhythmias or syncope.

Examination at this time showed no clinical evidence of heart disease. Chest x-ray was normal. ECG showed a frontal plane axis of +70° and no evidence of chamber enlargement. The rate was 45–49/min, PR interval 200 ms, QRS interval 40 ms, and QT interval 400 ms (Fig. 1a). On exercise heart rate increased to 75/min and PR interval to 260 ms (Fig. 1b).

She was hospitalized for further diagnostic study and therapy. On Holter monitoring when the sinus rate was 71/min or less, there were frequent periods of sinus arrest, maximally of 1·6 s duration (Fig. 1c). The sinus arrest was unrelated to activity; syncope did not occur under observation. Additional labora-

Fig. 1  (a) Resting ECG, sinus bradycardia at a rate of 45–49/min with PR interval 200 ms; (b) after exercise, with an increase in heart rate to 75/min and PR interval 260 ms; (c) representative tracings over a 3-hour period of Holter monitoring showing frequent episodes of sinus arrest, maximum 1·6 s; (d) pacemaker tracing, showing periodic appearance of sinus beats. PR interval 260 ms.
Sinus node dysfunction with atrioventricular conduction disturbance has been reported in a number of adults (Eraut and Shaw, 1971) but only once in children (Nugent et al., 1974). Complete heart block with persistent ductus arteriosus, though uncommon, is second in frequency to its occurrence with L transposition in patients with congenital heart disease (Nakamura and Nadas, 1964; Michaëllson and Engle, 1972). Sinus node dysfunction in association with congenital heart disease (types unspecified) has been described in 4 children and attributed to a congenital defect of the sinus node or alterations on the sinus node due to right-sided volume or pressure overload (Nugent et al., 1974).

The aetiology of the sinoatrial dysfunction and disturbance in conduction in this patient is obscure. Such abnormalities have not been noted previously in patients with persistent ductus arteriosus. Haemodynamic abnormalities were not present. The time relationship of the two conditions reasonably negates a congenital origin. Evidence, however, for an acquired aetiology is also lacking.

Summary

Sinus node dysfunction, previously unreported in girls, occurred in a 13-year-old girl who required permanent pacemaker implantation because of recurrent syncopal attacks. In addition to periodic sinus arrest, the presence of significant atrioventricular conduction disturbance was also documented. Although she had had a persistent ductus arteriosus divided at an earlier age, the disturbance of rhythm and conduction cannot be ascribed to a congenital or haemodynamic abnormality. An acquired origin can only be postulated.

References


**DENNISON YOUNG* and ROBERT E. EISENBERG**
Department of Pediatrics, Montefiore Hospital and Medical Center, 111 East 210th Street, Bronx, New York 10467, USA.

*Correspondence to Dr D. Young*
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D Young and R E Eisenberg

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