Case reports

well account for the phonocardiographic features of extreme variability of the A₂-OC interval and the intermittent absence of the OC. The systolic murmur was in keeping with mitral reflux with its rate related variability.

In a very ill patient with a malfunctioning mitral Starr-Edwards prosthesis we feel that the phonocardiographic findings can be a useful bedside adjunct to the physician and the cardiac surgeon in the assessment of prosthetic function when other methods may not be practicable.

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References


Peripheral gangrene in polycythaemia vera

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Summary
A case of peripheral vascular disease which became gangrenous and led to a diagnosis of polycythaemia vera is described. Chemotherapy has been successful and surgical intervention avoided. The authors stress the importance of full haematological examination of any patient presenting with peripheral digital gangrene.

Introduction
Vascular complications in patients with the myeloproliferative syndrome are common, but it is rare for digital ischaemia to be the presenting feature. It is the purpose of the authors to report such a case, to draw attention to the response to therapy, and to emphasize some of the surgical hazards in patients with this condition.

Case report
A 45-year-old Caucasian female first presented at the age of 36 with peripheral vascular disease. Her haemoglobin was 16 g/100 ml, WBC 15,100/mm³ and platelets 430,000/mm³. Bilateral lumbar sympathectomy provided symptomatic relief until the age of 43 when she presented with a similar 8-month history of pain and colour changes aggravated by the cold. Haemoglobin was 17.7 g/100 ml, PCV 53.8% and WBC 14,900/mm³. Response to Rheomacrodex was complete.

Two years later she was admitted with early gangrene of the right little toe, and both liver and spleen were palpable 4 cm below the costal margins. Haematology was essentially unchanged. Rheomacrodex led to improvement, but 2 months later she was re-admitted with extensive dry gangrene of the toe. Haemoglobin was 16.6 g/100 ml, PCV 48.4%, WBC 23,900/mm³, and platelets 625,000/mm³. Bone marrow aspiration and trephine biopsy confirmed the presence of myelofibrosis. She was started on chlorambucil 8 mg and aspirin 300 mg daily with venesection as required to maintain her PCV at 40–45%. On this regime her haemoglobin and platelets have fallen, and the white count decreased to just above the upper limit of normal. Surgical intervention has been avoided, and her toe has healed progressively.
Discussion

Untreated polycythaemia vera carries a high risk of intravascular thrombosis. In a recent review of 200 cases, Barabas, Offen and Meinhard (1973) noted that 49% presented with vascular complications. These were arterial in 34%, venous in 28% and both arterial and venous in 13%. Cerebrovascular accidents, preceded in 66% of the cases by transient ischaemic attacks (TIAs) and digital artery occlusions were the most frequent complications, although presentation with peripheral gangrene has been reported only rarely. In five of fifteen of their cases where only digital arteries were affected, femoral arteriograms showed localized narrowing in femoral or popliteal vessels. In all six patients reported by Gillespie (1973), co-incident atherosclerosis was present, and our patient had clinical evidence of vascular disease. It has been postulated that an underlying vascular abnormality may be a prerequisite for the development of peripheral vascular disease in patients with polycythaemia, and Barabas et al. (1973) feel that in some, if not all, embolism rather than local thrombosis is the mechanism of occlusion. These workers made the observation that in their series cerebrovascular accidents were four times more common than coronary artery disease while the reverse is generally true in arteriosclerosis, and drew a parallel with transient ischaemic cerebral attacks which have been shown to be due to microemboli.

The diagnosis of polycythaemia vera was not difficult in our patient, but where doubt exists, blood volume and red cell mass should be measured by standard radio-isotopic techniques.

The patient with a myeloproliferative syndrome offers particular problems at the time of surgery (Gilbert, 1973) and surgical intervention should be preceded by adequate control of the haematology as these patients are liable to bleeding or thrombotic manifestations. Failure of haemostasis may be seen in spite of a normal or even elevated platelet count, as myeloproliferative platelets are often qualitatively defective. It is recommended that no elective procedure be undertaken unless the patient has been haematologically stable for at least 4 months as they have been shown to have the most favourable post-operative course.

It should be emphasized that hypertonic solutions in the concentrations used for angiography may profoundly increase the viscosity to levels twice that suggested by the PCV, and patients undergoing this procedure should be venalected to reduce the PCV (La Celle and Weed, 1971).

Therapy of the underlying haematological condition is probably the most important therapeutic manoeuvre. It must be remembered that thrombocytopenia may be part of the syndrome and that reactive thrombocytosis is a normal response to bleeding. Fortunately, platelet function may be ablated by a number of agents, of which aspirin and dipyridamole are the two most commonly employed. Venasection should be employed as required to maintain the PCV in the range of 40-45%, and chlorambucil is extremely effective in controlling the elevated platelet count. In those patients reported by Gillespie (1973), four did extremely well after chemical sympathectomy using phenol, but the two patients in whom sympathectomy was not performed did as well on treatment of the haematological condition alone. Our patient responded excellently to the above therapy, and the gangrene has disappeared.

In the series of Barabas et al. (1973), ten patients had digital amputations before the diagnosis of polycythaemia vera was established. It is emphasized that in any patient presenting with peripheral digital gangrene, a full haematological work-up is mandatory, including marrow aspiration and marrow biopsy, and in the event of any suggestion of abnormality, radioisotopic investigations should be performed.

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References


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