Migrating emboli of retinal arteries in thrombocythaemia

GUSTAV SINGER
Eye Department, Hospital ÚVN, Prague-Stresovice, Czechoslovakia

Case report
A 37-year-old man, in the early months of 1965 complained of repeated attacks of temporary blindness in the right eye, occurring on an average of three or four times a week, and lasting for 2 to 3 minutes.

History
The patient had had two cerebral concussions in 1952 and in 1955. In 1958 he had a slight attack and in April, 1964, a more serious attack of aphasia. The neurological diagnosis was syndrome of the temporal lobe with aphasia, probably of vascular aetiology.

The visual acuity at this time was normal. On fundus examination a small white stripe was seen along the upper temporal branch of the right central retinal artery some 2 disc diameters from the edge of the papilla. This phenomenon was considered to be an arteriolar spasm. The surrounding retina showed an area of slight oedema 1 disc diameter in size. The patient had no symptoms. The oedema of the retina disappeared after 2 minutes as did the white stripe and the fundus became normal in appearance. Other clinical examinations were negative except for the neurological findings which consisted of the residue of expressive aphasia.

Examination
After discharge from hospital episodes of misty vision in the right eye occurred with greater frequency and the patient was re-admitted in December, 1965, when a photograph was taken of a spasm, this time progressing from the papilla to where the blood vessel divided into three branches above the papilla. This spasm remained visible for about half a minute (Fig. 1) and at the same time

FIG. 1 Two emboli are seen, one at the bifurcation of the retinal artery, the other more distally

Received for publication August 19, 1968
Address for reprints: Dr. Gustav Singer, Eye Department, Hospital ÚVN, Prague-Stresovice, Czechoslovakia
another vascular spasm was visible at the bifurcation at 12 o'clock on the edge of the papilla. Immediately after this a white stripe about one-third disc diameter long was observed in the upper temporal branch of the central retinal artery, and this stripe moved slowly from the bifurcation at the upper edge of the papilla towards the periphery (Figs 2 to 4). After about 2 minutes the fundus became normal in appearance.

**FIG. 2** The same emboli as in Fig. 1. One is migrating to the temporal periphery

**FIG. 3** Further migration of the temporal embolus seen in Fig. 2. The other has disappeared peripherally

*Progress*

In spite of the use of vasodilators the patient's condition remained unchanged until January, 1966, when he was admitted to the hospital's surgical department with suspected appendicitis. Several days after the removal of a chronically-inflamed appendix, a right hemiplegia developed and on the following day the patient's left leg was found to be cold and without a pulse. A thrombus, 12 cm. in length, was removed from the iliac artery, but as gangrene of the left leg developed a below-knee amputation was performed. The patient recovered very slowly during the following months.
Diagnosis
In May, 1966, a routine blood count showed 1,144,000 platelets per cu. cm. and a sternal puncture revealed large numbers of megakaryocytes in all stages of development confirming a diagnosis of essential thrombocythaemia.

Treatment
In view of these findings, treatment was commenced with the cytotoxic drug Mylecytan. After several weeks the number of platelets decreased to half a million. No visual disturbances have occurred since then.

Comment
It was concluded that this patient did not suffer from simple spasms of the retinal arteries but that the vascular occlusions were the result of so-called platelet-thrombi, which were producing the symptoms. Around these emboli the arterioles showed a spastic contraction.

That these emboli did not produce a permanent occlusion of the retinal arterioles may be due to the fact that, consisting of platelets, they are relatively soft and easily broken up.

Essential thrombocythaemia is a very rare disease. Lutz (1966) found only eighty cases of genuine essential thrombocythaemia. Most frequently repeated haemorrhage was cited as one of the symptoms, but one-third of all cases showed no haemorrhage at all, and the various symptoms of the patients were produced by some form of thromboembolic occlusions. It is assumed that the patient reported above belongs to this group.

Similar migrating emboli in the retinal arteries were described by Donaldson (1967) and Zweifach (1967), but not in patients with thrombocythaemia.

Summary
Migration of emboli in the retinal arteries is described and documented by photographs in a patient suffering from temporary blindness. The cause proved to be thrombocythaemia, and treatment with Mylecytan was followed by disappearance of the symptoms.

References
ZWEIFACH, P. H. (1967) Arch. Ophthal. (Chicago), 78, 705
Migrating emboli of retinal arteries in thrombocythaemia.

G Singer

doi: 10.1136/bjo.53.4.279