The likeliest explanation for these fatty lesions is that they are produced by pancreatic enzymes which have entered the circulation either directly or via lymphatics. The latter is thought to be the more important route. Abdominal peritoneal lymphatic drainage and transdiaphragmatic lymphatic absorption of abdominal fluid are probably prime factors in the ingress of pancreatic enzymes to the thoracic duct and hence into general circulation (ScarPELLI, 1956; Perry, 1947). Circulatory trypsin may break down vessel wall integrity and allow lipase to hydrolyse neutral fat in fat cells to glycerol and fatty acid. It has also been suggested (HODSON-WALKER and Woods, 1970) that the lesions are secondary to vascular spasm or necrosis, which leads to local areas of fat necrosis. The primary insult on the vessel being from pancreatic enzymes or due to intravascular thrombosis sometimes seen in acute pancreatitis.

A further but unlikely factor is that the lesions are caused by emboli of necrotic pancreatic tissue (Blauvelt, 1946).

Peripheral fat necrosis has been described in acute pancreatitis and as a marker of pancreatic neoplasm. This case and the others described should serve as a reminder that pancreatic disease may present with bizarre and obscure symptoms and signs.

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Periodic migrainous neuralgia associated with an arteriovenous malformation

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Summary
A patient with periodic migrainous neuralgia is reported. An arteriovenous malformation was found on the affected side. This association has not previously been recorded.

Introduction
In 1926 Wilfred Harris wrote of a type of recurrent facial pain and headache which he called periodic migrainous neuralgia. The syndrome was described in greater detail by Symonds (1956). Paroxysms of headache occur in bouts lasting several weeks with long intervals of freedom. There is at least one paroxysm in each 24 hr period. The pain is agonizing and is felt mainly in the supra-orbital region, in and behind the eye, and it is strictly unilateral. The patient is woken from his sleep in the early hours of the morning and cannot lie still. It rarely lasts longer than 2 hr and there is complete relief from pain between the paroxysms. The interval between bouts is rarely less than 6 months and may be several years. Balla and Walton (1964) documented twenty-eight
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Cases and found it most common in males, their first symptoms appeared between the ages of 20 and 40 years. He listed many of the associated features of the syndrome including watering of the affected eye, blurred vision, occasional nausea, tender swollen face, and ptosis. No cause for the pain is discovered (Symonds, 1956).

A case is presented in which an arteriovenous malformation was found on the affected side. This association has not previously been described.

Case history

P.W., a 39-year-old male driving instructor, presented in 1970 with a 3-week episode of recurrent attacks of pain above and behind the right eye. These occurred each 24 hr at 3.00 a.m. and lasted up to 2 hr. They were severe enough to wake him and he paced the bedroom floor. He had not had migraine previously and there was no relevant family history. A diagnosis of periodic migrainous neuralgia was made and he was treated with ergotamine tartrate. His symptoms settled.

In June 1973 there was a recurrence of his symptoms, but now his headaches began in late evening and lasted all night. The pain was sharp and throbbing. He noticed that his right eye watered, his vision was blurred and that he had a feeling of swelling above his right ear.

General examination showed him to be in good health. There was slight right ptosis but no change in pupil size. A localized systolic bruit was heard above and behind the right ear, where there was a palpable thrill. An arteriovenous malformation was suspected and he was admitted for arteriography.

Right carotid angiogram (Fig. 1) showed considerable enlargement of the superficial temporal artery and there was an arteriovenous malformation in the soft tissue of the scalp above the ear. There was rapid venous drainage to the external jugular vein with no abnormality in the internal carotid system.

A neurosurgical opinion was taken and it was felt that operation should not be performed since he was then symptom-free and there was no danger to his life. He has remained free of pain.

Discussion

Since Harris's initial description, various authors have described the syndrome by a variety of terms—erythromelagia of the head (Horton, Maclean and Craig, 1939); histamine cephalgia (Horton, 1941); greater superficial petrosal neuralgia (Gardner, Stowell and Dutlinger, 1947); cluster headaches (Kunkle et al., 1952).

This has led to confusion in diagnosis but in recent times the syndrome has become better recognized (Balla and Walton, 1964). Apart from the
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unusual length of headache in his second bout, this case is typical. It demonstrates well the importance of listening for bruits in the head when examining a patient with headache.

The aetiology of migraine is uncertain and the literature on the subject is large and complex. It is thought that the headache is due mainly to stretching, by dilatation, of branches of the external carotid artery and possibly to a lesser extent of the internal carotid artery (Brain, 1969). The findings of vasodilator peptides as well as histamine and acetylcholine in painful regions during migraine headache points to these substances being involved (Wolff, 1972).

There is a division of opinion whether periodic migrainous neuralgia is a migraine variant or a separate entity (Bickerstaff, 1959; Balla and Walton, 1964). Broche et al. (1970) studied the cerebral and ocular circulation in several patients with the syndrome and their findings suggested that vasodilatation was present in the pulsatile intravascular bed during the pain attack being particularly pronounced on the headache side. It is probable that the two conditions share similar mechanisms of vasodilatation.

Carotid angiography was performed by Ekblom and Greitz (1970) in patients with periodic migrainous neuralgia. Four out of eighteen had generalized dilatation of the cerebral arteries and the remainder were normal.

The significance of the arteriovenous malformation in this case is a matter for conjecture. Even if the headache returned and the malformation was removed it would be difficult to judge whether operation had been successful since spontaneous remission can occur after several years (Bickerstaff, 1959).

It is of considerable interest that Heyck (1969) has tried to relate the signs and symptoms of migraine to the opening of arteriovenous shunts. He suggests the vascular changes caused in the regional circulation, together with tissue hypoxia and acidosis which lower the pain threshold, can be the cause of the migraine pain. No histological evidence of these arteriovenous vessels or their demonstration during an attack has, however, been accomplished.

The importance of examining for bruits is emphasized. This may lead to the finding of further vascular abnormalities in this form of neuralgia.

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