Fistula between three main cerebral arteries and a large occipital vein

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The literature contains few references to direct communications between cerebral arteries and veins. Dandy (1945) was the first to describe the passage of a branch from the middle cerebral artery into a vein leading into the Sylvian group. Jaeger and Forbes (1946) reported direct communications between the posterior cerebral and posterior choroid arteries and the basilar vein, together with multiple anastomoses with the great vein of Galen. Haberland (1950) published an account of the accidental finding of a communication between the branches of the posterior cerebral artery and the dilated vein of Rozenthal and proved it microscopically. Bergstrand, Olivecrona, and Tönnis (1936), Röttgen (1937), Sorgo (1938), and Verbiest (1951) reported observations of intradural direct arteriovenous communications. Some cases of vascular malformations of the great vein of Galen may also belong to this category, although they are generally regarded as a separate entity.

The following report describes an abnormality which is unique in our experience, and we have not encountered a similar one in our survey of the literature.

CASE REPORT

A 26-year-old man was first seen in September 1961. From childhood he had suffered from right-sided headaches, mainly in the temporal region and below the eyebrow, the pain being treated as migraine. As time went on, the headache ceased to trouble him, but he then began to complain of attacks of vertigo, associated with nausea or vomiting, usually lasting about one hour and being so severe that they forced him to go to bed for fear of falling. It was for these attacks that he sought advice.

The neurological examination and fundoscopy at that time were normal. Skull films and tomograms revealed a hyperostosis in the left fronto-parietal region, growing from the outer table, with a circular defect, limited sharply by sclerotic bone. In the tangential view there was thickening of the inner table in the base of the hyperostosis. These appearances were suggestive of a menigioma, and angiography was undertaken with this possibility in mind.

LEFT CAROTID ANGIOMGRAM (Fig. 1) This showed a dilatation of the internal carotid artery at its siphon and of the anterior cerebral artery in its first part. The majority of the contrast medium was drawn through the anterior communicating artery to the right anterior cerebral artery, which was dilated to a width of 5 mm in the early part of its course and 7 mm further posteriorly. It terminated directly into an enormous occipital vein running to the sagittal sinus. There was poor visualization of the territory supplied by the left anterior and middle cerebral arteries, indicating that most of the blood was going to the fistula.

RIGHT CAROTID ANGIOMGRAM (Fig. 2) This also showed a dilatation of the internal carotid artery. The beginning of the anterior cerebral artery was not seen. The middle cerebral artery was greatly enlarged, elongated, meandering, and terminated in the same anomalous occipital vein as had been seen in the previous angiogram. The vein wound in large loops to the sagittal sinus. The right sagittal and both transverse sinuses were also dilated. As on the left side, the blood supply of the right hemisphere appeared to be deficient, because most of the circulation was going to the fistula.

ANGIOMGRAM OF THE LEFT VERTEBRAL ARTERY (Fig. 3a, b) This showed that the vertebral and basilar arteries were enlarged, elongated, and winding. Almost all the contrast material passed into the right posterior cerebral artery, which was dilated and ran in big curves into the anomalous occipital vein. The cerebellar vessels were poorly visualized.

Total serial angiography confirmed these circulatory anomalies. The patient refused surgical treatment, as he did not consider that his complaints were troublesome enough at this time. He was readmitted on 2 February 1964 because the headaches had recurred, and because he had had several attacks of transient weakness of the right hand, and on two occasions loss of consciousness. The neurological state remained normal.

At operation on 25 February 1964 a right occipitoparietal craniotomy was performed. On the surface of the brain an anomalous vein, as large as a thumb, ran from the occipital to the parietal lobe, forming a large varix just before its termination in the sagittal sinus. The arachnoid was thickened over the whole of this region. From all directions large arteries entered the vein so that a thrill could be felt on palpation. When all these arteries had been ligated, the tension in the vein...
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FIG. 1. Before operation.

FIG. 2. Before operation.
greatly decreased and the thrill disappeared. Then the
gigantic vein with a part of the brain was removed.
The procedure was extremely difficult, as the walls of the
vessels were very fragile and, in spite of careful dissection,
often tore with violent bleeding, serious blood loss, and
drop of the arterial pressure. The patient received a
blood transfusion of 4,100 ml. The procedure took four
hours.

For some days after the operation the patient was
somnolent and confused and had a left-sided hemiparesis.
This complication cleared up rapidly, but a superficial
wound infection required antibiotic therapy for several
days. He was discharged in good mental and physical
condition. Control angiography was carried out on
1 April 1964 (Fig. 4a, b, c, d). The cerebral circulation in
both hemispheres now appeared to be virtually normal. All
of the main cerebral arteries on the right side had
diminished in size and the arteriovenous communications
were no longer visible. Both of the carotid arteries were
smaller and irregularly narrowed, but there was an
aneurysmal dilatation of the origin of the posterior
communicating artery on the left side.

**COMMENT** The vascular anomaly described above would
explain the chronic headache of which the patient had
complained since his childhood. All three main cerebral
arteries on the right side terminated in an occipito-
parietal vein which emptied into the sagittal sinus. All
these vessels were enormously dilated, and the fistulous
communication between them led to serious deprivation
of the blood supply to the whole brain. The radical
removal of the malformation led to a complete relief of
symptoms.

**DISCUSSION**

This vascular anomaly is unique in its form, and in
the literature no similar observation could be found. It is the nearest to the one described by
Dandy (1945), but in comparison is more com-
plicated and larger. In Haberland’s (1950) case the
arteriovenous communication was microscopic. In
other cases described in the literature the lesion has
been situated in the dura mater and the arterial
communications were mostly from the meningeal
arteries. The malformation which we have described
is more primitive than in the typical arteriovenous
malformations. Between the feeding arteries and
draining vein there was no system of transit vascular
channels, and so the arterial blood stream ran
directly into the enormously dilated cerebral vein.

This arteriovenous fistula is the result of a
developmental defect of the capillary system in the
early stage of embryonic development of the cerebral
vascular system, as in arteriovenous malformations.
It is possible that embryonic vessels, destined to be
arteries and veins, may cross in such a way that they
are separated only by two layers of endothelial cells
and that this barrier gradually disappears under the
influence of the arterial pulse.

It is probable that the abnormality, present at
birth and enlarging steadily thereafter, was respon-
sible for the long-standing headaches. Progressive
enlargement of the contributing vessels increased
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the power of the shunt, and led to a deficit in the cerebral and cerebellar blood supply, witnessed by vertigo, transient hemianopia, and attacks of loss of consciousness. Haberland's findings suggest that arteriovenous communications, at first only microscopic, open successively during life. In our case, the normal development of the brain, as indicated by intact mentality and the absence of neurological signs, suggests that the shunt, as seen at the time of angiography and operation, may have been present at birth, but enlarged progressively throughout life.

The indication for surgery was clear, not only because of severe headache but also because of the prospect of the cerebral circulation becoming more embarrassed, leading to the grave mental changes which are known to occur in large arteriovenous malformations.

Fig. 4 After operation.
SUMMARY
A case of unique multiple arteriovenous fistulae between all three main cerebral arteries—anterior, middle, and posterior arteries—and a cortical occipito-parietal vein, is reported. The malformation disturbed the cerebral circulation and caused severe headaches, vertigo, transient hemianopia, and attacks of loss of consciousness. Surgical ablation of the fistula relieved these symptoms and restored the circulation to normal. A possible explanation of the abnormality is offered.

REFERENCES


