Late complications of hemispherectomy:
report of a case relieved by surgery

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SUMMARY A case of Sturge-Weber disease treated with left hemispherectomy presented, 11 years later, with complications related to delayed intracranial haemorrhage. A loculation syndrome of the right lateral ventricle was detected and it was corrected by a ventriculoatrial shunt operation. The side of the hemispherectomy was evacuated of all the chronic products of haemorrhage, including the subdural membrane. The patient was relieved of her symptoms. It is considered that complications related to delayed haemorrhage after hemispherectomy are remediable.

Immediate and delayed complications occur after hemispherectomy. Early complications include obstructive hydrocephalus and herniations of the remaining hemisphere (Cabieses, Jeri, and Landa, 1957; Laine, Pruvot, and Osson, 1964). A syndrome of delayed intracranial haemorrhage was reported by Oppenheimer and Griffith (1966). The essential features of the syndrome are (1) an infantile hemiplegia treated by hemispherectomy; (2) a trouble-free period lasting for some years; (3) a period of deterioration, extending over several years and ending in death; during this period, evidence of bleeding into cerebrospinal fluid (CSF) pathways and later of obstructive hydrocephalus is found; (4) post-mortem findings of superficial haemosiderosis of the central nervous system; chronic granular ependymitis, leading to obstruction of cerebrospinal fluid pathways; and evidence of multiple bleeding points in the membrane which had replaced the missing hemisphere and in the extension of this membrane on to the lining of the ventricular system.

Falconer and Wilson (1969) reviewed the literature and reported four cases of delayed complications after hemispherectomy, of which three were relieved by surgery. They described a hitherto undescribed complication of hemispherectomy—namely, 'loculation' of the residual temporal horn.

This paper reports a case with complications related to delayed haemorrhage after hemispherectomy which was successfully relieved by ventriculoatrial shunt and reoperation.

CASE REPORT

A.L., a 17-year-old girl, was admitted to the Neurology service of Christian Medical College Hospital, Vellore, on 10 July 1969, with persistent headache, vomiting, and increasing drowsiness of three weeks' duration. She was born with a Sturge-Weber syndrome and had had a left hemispherectomy performed in another country 11 years before. She was free from seizures and major behavioural problems and was attending a school for backward children till November 1968, when she developed severe constant headache, vomiting, and drowsiness. She was admitted elsewhere in early December 1968, where brownish yellow fluid with a protein content of 1,150 mg/100 ml was tapped from the left subdural space. Fluid from the right lateral ventricle had a protein content of 1,100 mg/100 ml. Lumbar puncture revealed xanthochromic CSF, with a protein content of 352 mg/100 ml. In six weeks, after repeated ventricular and lumbar punctures, her symptoms subsided. The lumbar CSF protein came down to 160 mg/100 ml. She remained well from the middle of January 1969 to the end of June 1969. There was then a recurrence of symptoms, and a lumbar puncture on 28 June 1969 revealed clear fluid with a protein content of 73 mg/100 ml. Left subdural tap on 4 July 1969 revealed yellow blood-stained fluid with 3,000 mg/100 ml. protein. She was referred to our hospital for further management.

On admission she was bedridden, irritable, and extremely drowsy. She was disoriented in time and place and resented examination. Ocular fundi were pale. Left 6th nerve palsy was noticed. There was hemiatrophy of the right side with flaccid hemiparesis. A left facial naevus of Sturge-Weber syndrome was present on the left cheek. Vision was 20/200 in the left eye and 20/15 in the right. There was an asterixis of the left hand. The blood pressure was 110/60 mm Hg. The heart sounds were normal, and the liver was palpable 2 cm below the right costal margin. The right lower limb was flaccid and showed Babinski's sign. The left lower limb was normal. The left upper limb was flexed and the right arm was normal. There was no dysmetria. The gait was ataxic.

Lumbar puncture was performed on 11 July 1969. The opening pressure was 30 cm H2O. The fluid was straw coloured and the clear underlaying film was absent. The glucose level was 69 mg/100 ml, and the protein content was 5 mg/100 ml. No organisms were identified by smear examination or by culture. The fluid was sterile on culture.

On 7 August 1969, a left ventriculoatrial shunt was performed. The fluid was clear; the glucose level was 60 mg/100 ml, and the protein content was 5 mg/100 ml. Post-operatively, the patient was alert with a clear fundus and no evidence of papilloedema. She was referred to the Department of Ophthalmology.

On 28 June 1969, she was readmitted to the hospital with recurrent headache and vomiting. On admission, she was bedridden, irritable, and extremely drowsy. She was disoriented in time and place and resented examination. Ocular fundi were pale. Left 6th nerve palsy was noticed. There was hemiatrophy of the right side with flaccid hemiparesis. A left facial naevus of Sturge-Weber syndrome was present on the left cheek. Vision was 20/200 in the left eye and 20/15 in the right. There was an asterixis of the left hand. The blood pressure was 110/60 mm Hg. The heart sounds were normal, and the liver was palpable 2 cm below the right costal margin. The right lower limb was flaccid and showed Babinski's sign. The left lower limb was normal. The left upper limb was flexed and the right arm was normal. There was no dysmetria. The gait was ataxic.

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The patient was discharged on 31 October 1969.
Radiographs of the skull revealed a left hemiatrophy with silver clips in various locations. A right carotid angiogram showed considerable shift of the anterior cerebral artery across the midline to the left side, a hydrocephalic pattern, and elevation of the middle cerebral artery (Fig. 1). A left carotid angiogram showed an ipsilateral shift of the anterior cerebral artery. Air injected into the right lateral ventricle filled it, but could not be induced to enter the left side. All parts of the right lateral ventricle were markedly dilated and showed a contralateral shift. Air introduced into the left subdural cavity showed multiple loculated spaces (Figs. 2, 3, and 4). The right and the left sides were not communicating. The 3rd ventricle, aqueduct, and 4th ventricle were not visualized. A diagnosis of delayed haemorrhage after hemispherectomy and resultant obstructive hydrocephalus involving the right lateral ventricle was made.

**OPERATIVE PROCEDURES** A ventriculooatrial shunt was performed on the right side on 18 July 1969 using a Pudenz-Heyer valve. The patient showed remarkable improvement after the shunt operation. Headache gradually disappeared and she became more alert.

A left parietal craniotomy was done on 23 July 1969. A very thick subdural membrane was found which in places was calcified. Multiple loculated areas containing xanthochromic fluid were seen to fill most of the cavity. Blood clots of varying ages were also found. The membrane resembled that seen in chronic subdural haematoma (Fig. 5). Most of the membrane was excised except the medial part which was covering the basal ganglia. The whole cavity was evacuated and irrigated with Ringer solution. The dura was closed and the bone flap replaced.

Post-operatively, there was remarkable improvement. In 10 days after the operation she was talking and walking without support. There was no headache or vomiting. She showed interest in reading while in the hospital. She was discharged on 7 August 1969 and was reported to be doing well up to the preparation of this paper.

**DISCUSSION**

Oppenheimer and Griffith (1966) who first described the syndrome of delayed intracranial haemorrhage after hemispherectomy cited Falconer as saying that the condition can be treated surgically. Falconer and Wilson (1969) attributed the successful management of their cases primarily to removal of the products of haemorrhage rather than to attempts to create a normal CSF circulation by a shunt operation. They emphasized the point that these
patients may be regarded as having a chronic encapsulated or membranous subdural haematoma. They have tried many surgical procedures including simple evacuation of the fluid compartment, operative removal of solid clot, and reduction of the area of potential bleeding and exudative surface by excision of as much membrane as is safely feasible.

In our case, the lumbar CSF and the CSF of the right ventricle were clear and were of a relatively

FIG. 4. Ventriculogram. Brow-up lateral view: dilated anterior and temporal horns of right lateral ventricle and loculated air spaces on the left side.

FIG. 3. Ventriculogram. Posteroanterior view: dilated right lateral ventricle, especially the occipital horn.

FIG. 5. Left parietal craniotomy. Thick subdural membrane is indicated by the straight arrow and the shiny loculated area of xanthochromic fluid collection by the curved arrow.
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normal composition, whereas the fluid in the left subdural cavity was brownish, blood stained, and had a high protein content. Ventriculography confirmed that the left cavity was not communicating with the right ventricle. The right ventricle was abnormally dilated and CSF was under increased tension, indicating that this patient had a 'loculation syndrome' of the right ventricle. 'Loculation syndrome' of the residual temporal horn requiring ventriculoatrial (VA) shunt was described by Falconer and Wilson (1969). In our case the loculation was not confined to the temporal horn, but involved the whole of the right lateral ventricle. Though the patient had considerable relief of symptoms after VA shunt, reoperation and evacuation of the left cranial cavity gave total relief of her symptoms.

We feel that in a case presenting with delayed complications after hemispherectomy, evidence of 'loculation syndrome' should be looked for and an appropriate shunt procedure performed. The cavity with haemorrhage and membrane should be dealt with separately, the best procedure being removal of as much membrane as possible and evacuation and lavage of the cavity. We agree with Falconer and Wilson (1969) who emphasized that the syndrome of delayed complications related to haemorrhage years after hemispherectomy is a remediable condition, provided a correct diagnosis is made early.

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