INTRACRANIAL MENINGIOMA IN A 13-YEAR-OLD MALE

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This case demonstrates the presence in a young child of a large intracranial meningioma showing minimal localizing signs clinically, but with the patient deteriorating progressively into a comatose state.

Case History

The patient, a male scholar, aged 13 years, was admitted to hospital complaining of right-sided headaches of 3 weeks' duration becoming progressively worse. There had been occasional fits.

Previous history. There was an episode when he had been butted on the forehead by a ram at the age of 6 years. This caused a large swelling which later subsided.

On examination, he was malnourished, apyrexial, had a large head, and neck rigidity was present. Bilateral papilloedema was noted, but his cranial nerves were intact. Motor power and tone of the limbs were normal and equal on both sides. All reflexes were present and equal. No abnormalities of the sensory system were found. Coordination was difficult to test.

The chest was clear clinically and on X-ray. The cardiovascular system showed a pulse of 64/min., regular and of good volume; the heart sounds were closed; BP was 100/160 mm.Hg. No masses were felt on abdominal examination. Liver, spleen and kidneys were not palpably enlarged.

An X-ray of the skull showed widening of the coronal sutures. No intracranial calcification or erosion was noted. A diagnosis of increased intracranial pressure, probably owing to hydrocephalus, was made.

Two days later he was seen by the neurological surgeon, who suggested ventriculography and bilateral carotid angiography.

The patient became increasingly drowsy, had attacks of vomiting and the neck rigidity became extreme, to the extent of opisthotonos. In view of the rapid deterioration, the absence of lateralizing signs and the possibility of tuberculous meningitis, a lumbar puncture was performed—proper precautions having been taken to deal with any complications arising out of this procedure. CSF was clear and under pressure. No pathological vessels or arterio-venous shunts were noted. The appearances were those of a large avascular space-occupying lesion in the right temporoparietal region. This confirmed the appearance of a provisional diagnosis of a large hydatid cyst or porencephalic cyst (Figs. 1-4). It was decided to proceed with a craniotomy.

Operation—Craniotomy

The lesion was approached through a liberal right temporoparietal osteo-plastic flap. The dura mater was found to be under extreme tension, and a brain cannula introduced into the lesion failed to get into the cyst. The lateral ventricle was opened and a large surfacing bluish tumour was encountered, the attachment to the anterior cerebral artery appeared stretched over a dilated lateral ventricle. The posterior temporal branch of the right middle cerebral artery was displaced laterally and the choroidal artery appeared to be displaced medially and slightly downwards. No pathological vessels or arterio-venous shunts were noted. The appearances were those of a large avascular space-occupying lesion in the right temporoparietal region. This confirmed the appearance of a provisional diagnosis of a large hydatid cyst or porencephalic cyst (Figs. 1-4).

Subsequent investigations. On the fourth day after admission urgent ventriculography was done under general anaesthesia. A brain biopsy done during this procedure showed a large hydatid cyst.

The ventriculogram showed ventricular dilatation and gross displacement of the lateral ventricular system from right to left. The third ventricle also appeared displaced and the findings suggested a large right-temporal or temporoparietal tumour.

The next day a right carotid angiogram was performed showing anterior and middle cerebral vessels displaced markedly upwards and to the left, there being some herniation of the terminal branches of the anterior cerebral artery underneath the free ends of the falx. The pericollosal branches of the anterior cerebral artery appeared stretched over a dilated lateral ventricle. The posterior temporal branch of the right middle cerebral artery was displaced laterally and the choroidal artery appeared to be displaced medially and slightly downwards. No pathological vessels or arterio-venous shunts were noted. The appearances were those of a large avascular space-occupying lesion in the right temporoparietal region. This confirmed the appearance of a provisional diagnosis of a large hydatid cyst or porencephalic cyst (Figs. 1-4).

The lesion was approached through a liberal right temporoparietal osteo-plastic flap. The dura mater was found to be under extreme tension, and a brain cannula introduced into the lesion failed to get into the cyst. The lateral ventricle was opened and a large surfacing bluish tumour was encountered, extending from the temporal to the parietal region, measuring 11 cm. x 16.5 cm. The tumour was easily freed from its bed and it was noted that there was only a small attachment to the extent of half an inch to the dura mater of the sphenoidal ridge. The tumour was removed in toto and the attachment to the dura mater was cauterized. The dura mater was then closed and the bone flap was replaced.

The postoperative course was uneventful and the patient was discharged from hospital after 6 weeks.

At the time of discharge he was symptom-free, intellectually normal and no abnormal physical signs were present. The papilloedema was disappearing. Subsequent follow-up for the

![Fig. 1. Plain lateral radiograph of skull demonstrates sutural diastasis mainly of the coronal sutures.](image1)
![Fig. 2. P-A view of ventriculogram demonstrating a gross displacement of the lateral ventricles and the third ventricle to the left.](image2)
![Fig. 3. Right cerebral angiogram, lateral position, demonstrating the upward displacement and compression and stretching of the middle cerebral group of vessels with some downward displacement of the posterior cerebral artery.](image3)
![Fig. 4. Right cerebral angiogram, A-P position, demonstrating gross displacement of middle cerebral group of vessels in a medial direction with moderate displacement of anterior cerebral artery across the mid-line to the left side and some downward displacement of the posterior cerebral artery.](image4)
duration of 1 year showed that he was symptom-free and had returned to school.

This patient presented with the classical triad of raised intracranial pressure—headache, vomiting and papilloedema. The minimal lateralizing signs presented in this case confirm the experience of Teng\(^3\) who, in reviewing the literature, commented upon several unusual features of intracranial meningioma in children, which were:

(a) Enlargement of head—which may be markedly enlarged with minimal neurological signs,
(b) The tendency to recurrence of growth,
(c) Higher incidence of sarcomatous change than in adults,
(d) Lack of dural attachment, and
(e) High operative mortality.

Cushing and Eisenhardt\(^4\) in a series of 284 intracranial meningiomas (1938) quote 6 occurrences in the pre-adolescent age group. Cushing contends that meningiomas are the most favourable of all intracranial tumours for surgery, but present a problem for successful removal because of the large size, the great vascularity and the tendency to recurrence.

There is usually a long history of symptoms before the initial examination, but Pineda and Coe\(^5\) described 4 cases with rapid progression to a comatose state, with absent lateralizing signs and stressed the need to bear the diagnosis in mind when an individual is admitted to hospital in a comatose state.

An interesting finding in this case was the normal cerebrospinal fluid protein, which contrasts strikingly with the high figure for this entity commonly found.

**SUMMARY**

1. This case illustrates a large intracranial meningioma in a 13-year-old male.
2. Minimal neurological signs existed in spite of the large tumour mass.
3. The patient deteriorated rapidly to a critical state.
4. The tumour was removed successfully.

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**REFERENCES**


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**THE BELGIAN MEDICAL DISPUTE IN RETROSPECT*\(^*\)**

**DR. J-R. GOSSET, Associate Editor, World Medical Journal**

A dispute between the Belgian medical profession and the Government broke out at the beginning of 1964. This dispute had considerable repercussions, and public opinion was so emotionally swayed that at times it was difficult during the conflict to ascertain the true facts. Now, at a distance, it is easier to see the events of spring 1964 in perspective.

Before the reforms to which the medical profession took exception, health insurance law was in the hands of politically or confessionally based organizations and was limited to cash payments in accordance with a tariff, the physicians' fees not being fixed. Sums reimbursed were small and various political committees had worked for two years on draft legislation for health insurance reform. On 8 May 1963 the draft was handed