Spontaneous CSF Rhinorrhea as Clinical Symptom of an Idiopathic Temporal Encephalocele in a 64 Year Old Man

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Abstract

A rare case of idiopathic temporomesial encephalocele in a 64 years old female is presented. The clinical symptoms of this adult patient were spontaneous rhinorrhea and headaches. A temporomesial encephalocele could be diagnosed and the cele was removed surgically in two operative procedures. In case of clinical symptoms like rhinorrhea a surgical removal of the encephalocele is recommended to prevent infections like meningitis.

Keywords: Encephalocele; Temporal lobe; Elderly; CSF rhinorrhea

Introduction

Idiopathic temporal encephaloceles are rare conditions with a reported incidence of approximately 1 in 35,000 people [1]. Encephaloceles were etiopathologically described since 1888 as a prolonged persistence of the so-called Sternberg canal [2].

Newer anatomical and physiological findings combine arachnoid pits, pneumatisation of the lateral sphenoid sinus and an (partially) empty sella [3].

Therefore a classification of spontaneous lateral sphenoid cephaloceles into 2 subtypes have been proposed by their anatomical location: Type 1 with Herniation of the cele into a pneumatized sphenoid sinus often present with a CSF leakage and Type 2 which herniates into or through the greater wing of the sphenoid via e.g. a bony defect. In this group patients often present with seizures or headache and not as in our case with a CSF Leakage [4].

Case presentation

We present a rare case of an idiopathic temporomesial encephalocele in a 64 years old patient who complained about 3 weeks of headaches and rhinorrhea without any trauma in his history (Figure 1A and B).

The Ear Nose Throat (ENT) colleagues could not find a source of the rhinorrhea, but the patient has been treated with oral antibiotics because of a sinusitis.

After 1 week of persisting “water-like” secretion a computer tomography (CT) scan was performed with no pathological finding according to the outpatient colleagues. Indeed there has been a bony defect and finally after 3 weeks, a cranial magnetic resonance imaging (MRI) has been performed (Figure 1) and the patient was admitted to our department.

In the clinical neurological examination and lab analysis there were no signs of meningitis or other neurological deficits. The rhinorrhea could be provoked by leaning the patient’s head forwardly. The CSF rhinorrhea was confirmed with a β2-transferrin test.

We decided to surgically treat the encephalocele. In the surgical procedure we used a right pterional approach, removed the encephalocele and sealed the defect with a temporal muscle and fascia graft.

The CSF rhinorrhea did not occur again after surgery and the patient could be discharged without any complains on the 7th postoperative day. The postoperative cranial CT scan showed no remnants of the cele and no CSF-leakage. Six months later the patient presented again in our outpatient clinics with a recurrence of CSF-rhinorrhea. We performed a cranial MRI as well as a high resolution skull base CT-scan and diagnosed the leakage of the used autologous dural graft (Figure 2A and B).

We decided to re-operate and used a fibrin coated collagen sponge as well as small bone fragment, which we fixed on the temporal bone as an abutment to stabilize the sealing (Figure 3A and B).

Afterwards the patient recovered well and could be dismissed 6 days after the second surgical procedure without any complains. In the period of the last 2 years she was not re-admitted.
Figure 1: A) Preoperative MRI scan (coronal section T2-weighted) B) T1 weighted axial section

Figure 2: A) MRI. T2-weighted scan before 6 months after surgery showing a new leakage on the operated region B) CSF leakage scan before 2nd procedure
Discussion

There are a few cases of temporal encephaloceles described in literature [4-6].

The encephalocele of the present case is located anteromedial of the temporal lobe. A pneumatisation of the lateral sphenoid sinus is retrospectively visible on an MRI scan before the surgical procedure, as well as a partial empty sella in the coronal MRI scan. The difficulty in this case has been to localize the point of the CSF leakage, which finally has been solved via a high resolution skull base CT scan. Additionally the postulated intracranial hypertension or different CSF dynamics might be the reason of recurrence of the CSF-fistula.

CSF-rhinorrhea put the patient on a significantly higher risk for meningitis [7].

In cases of symptoms like CSF-rhinorrhea with an increased risk for meningitis, surgery should be performed after detailed high resolution CT scan and multiplanar MRI scan to avoid a second surgical procedure. Whether it should be performed endoscopically or microscopically depends on the expertise of the surgeon and on the anatomical localisation.

References

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