INTRASPHENOIDAL ENCEPHALOCELE ASSOCIATED WITH CEREBROSPINAL FLUID FISTULA AND SUBDURAL HEMATOMAS: TECHNICAL CASE REPORT

OBJECTIVE AND IMPORTANCE: Intrasphenoidal encephalocele is a rare clinical entity that is often complicated by rhinorrhea, recurrent meningitis, and headache, but in no case has the association of rhinorrhea with subdural hematomas been described. A surgical procedure to stop persistent cerebrospinal fluid leakage is reported.

CLINICAL PRESENTATION: A 59-year-old man sought care for intractable rhinoliquorrhea of 6 months’ duration. Cranial computed tomographic and magnetic resonance imaging scans revealed a basal posterior frontal bony defect and an evocative image suggesting intrasphenoidal encephalocele.

INTERVENTION: A transnasal transsphenoidal surgical procedure was performed; the encephalocele was removed, and the sphenoid sinus was filled with an inflatable pouch made of synthetic dura mater containing abdominal fat. Postoperative reduction of the rhinoliquorrhea, but not its total disappearance, was observed. Total disappearance was achieved only after endonasal, transmucosal inflation of the pouch with human fibrin glue. One of the subdural hematomas disappeared spontaneously, and the other was treated by a surgical procedure.

CONCLUSION: The possible role of the presented technique in the treatment of cerebrospinal fluid leakage is discussed.

KEY WORDS: Cerebrospinal fluid rhinorrhea, Intrasphenoidal encephalocele, Transsphenoidal approach

The patient, a 59-year-old male doctor, had experienced rhinorrhea and occasional retro-orbital pain and frontal headache for approximately 6 years. No previous trauma or other significant abnormalities were reported. The neurological examination revealed nothing abnormal. The leaking nasal fluid was confirmed by a glucose test to be cerebrospinal fluid (CSF) (47 mg/100 ml). The first contrast-enhanced encephalic magnetic resonance imaging (MRI) scan, obtained 20 days before admission to our department, failed to demonstrate the location of a fistula and did not detect a blood collection. However, it revealed a fusiform formation in the sphenoid sinus with signal characteristics similar to those of the cerebral parenchyma (Fig. 1). A high-resolution computed tomographic (CT) scan revealed a communication between the right posterolateral side of the sphenoid sinus and the sphenoid sinus (Fig. 2). Considering the length of the patient’s clinical history and the therapeutic procedures performed in the past, which included prolonged lumbar drainage performed at another institution 3 months earlier, surgery was proposed. Surprisingly, control MRI scans, which...
were taken before surgery to assess increased CSF leakage, revealed two large bihemispheric SDHs (Fig. 3).

**OPERATIVE TECHNIQUE**

The sphenoid sinus was exposed through a transnasal transsphenoidal microsurgical approach disclosing, in the right section, a fusiform mass (1.5 × 0.7 cm) with a fine translucent coating; the small mass protruded through a bony defect onto the inferoposterolateral ethmoidal cells. CSF leakage around the mass was also evident (Fig. 4). The lesion was removed, and duraplasty was performed with human fibrin glue and synthetic dura mater. Then a pouch was made by wrapping synthetic dura mater around abdominal fat; this was inserted into the sphenoid sinus as a “plug,” which resulted in reduction, but not disappearance, of the CSF leakage (Fig. 5). This was not evident during the operation, but only postoperatively, when the recurrence of rhinoliquorrhea, even though reduced, was observed. A histological examination revealed that the removed lesion consisted of central nervous system tissue.

The patient underwent an endonasal, transmucosal surgical procedure while under CT scan control, as previously described (12). A 12-gauge lumbar puncture needle was inserted through the sphenoid sinus into the pouch, and human fibrin glue was injected under pressure (Fig. 6). After this surgical procedure was performed, the fistula disappeared. During the next 2 weeks, CT and MRI control scans revealed progressive reduction until the SDH disappeared on the left side; the one on the right side persisted, so it was surgically drained. At a follow-up examination at 20 months, the patient showed no signs of rhinorrhea, and a CT scan demonstrated that SDHs were absent.

**DISCUSSION**

The association of rhinorrhea, SDHs, and intrasphenoidal encephalocele represents an event not described, to our knowledge, in the available literature. It raises a diagnostic problem regarding the location of the fistulous connection and therapeutic questions regarding the choice of surgical procedure. The bony defect was localized by high-resolution CT scan (4, 8, 23). The MRI scan did not visualize the fistulous connection, but it contributed to the identification of the intrasphenoidal encephalocele (10, 23).

The choice of the surgical approach was not simple, as we envisaged both a transcranial and a transsphenoidal approach. We were persuaded that the choice of the transsphenoidal approach was not contingent upon the particular expertise of one of us (BF) with this technique (11). In our opinion, it is advisable that a surgeon with an equivalent confidence in both the transsphenoidal and the transcranial approach be at first inclined toward a less invasive procedure. The additional risks of the intracranial procedure are not negligible, and they are both generic and related to the possible manipulation of vascular and nervous structures. Undoubtedly, the transcranial route should be the first choice when a wide cranial base bone defect requires an osteoplasty completed with a large pericranial vascularized flap; the transsphenoidal approach probably cannot ensure an adequate closure in these cases. Nevertheless, in intracranial procedures for the treatment of CSF fistula, recurrences are not rare (2, 4, 5, 25). In the present case, the bone defect was actually limited and easy to reach transsphe-
have progressively restored the craniocerebral proportions, possibly also reducing fibrinolytic activity. Finally, these events may have resulted in neomembrane atrophy and progressive hematoma fluid resorption (14, 15, 19, 21, 24). However, these hypotheses do not explain why the contralateral hematoma persisted in our patient. Finally, although the follow-up period in our case may be considered to be too short, the convincing rationale of the technique leads us to conclude that the result will be lasting.

REFERENCES


FIGURE 5. Drawing of the insufficient intrasphenoidal filling of autologous fat wrapped by synthetic dura mater. The CSF leakage is reduced but still exists.

FIGURE 6. Drawing of the CT-guided pressure injection of human fibrin glue into the customized dural pouch.
COMMENTS

This article describes treatment of an intrasphenoidal encephalocele with an original technique of using an inflatable pouch that is inserted into the sphenoid sinus. In our experience, using the transsphenoidal approach and filling the sphenoid sinus with fat and glue, while draining the cerebrospinal fluid (CSF) via a spinal catheter for several days, may also be successful in most cases. However, after unsuccessful lumbar drainage in intractable CSF fistula via the sphenoid sinus, the technique described in this article may be an interesting and novel solution.

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The authors have successfully applied their previously published transmucosal needle technique (1) to arrest a CSF rhinorrhea persisting after transsphenoidal treatment of an encephalocele associated with a CSF fistula. This report demonstrates that it is sometimes possible to find simple solutions to apparently complex problems. One should consider the authors’ conclusions regarding the superiority of the transsphenoidal approach over the transcranial approach for the treatment of intrasphenoidal encephalocele before attempting this technique.

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The report of Fraioli et al. is interesting because of the rarity of the anomaly, the concomitant subdural hematomas, and the method of surgical repair. An intrasphenoidal encephalocele presenting with rhinorrhea was treated by transnasal transsphenoidal resection of the encephalocele and obliteration of the sphenoid sinus with autologous fat contained within a pouch of synthetic dura. Persistent rhinorrhea associated with bilateral subdural hematomas was successfully treated by secondary expansion of the pouch with fibrin glue injected endonasally.

Basal encephaloceles, herniations of brain, and meninges through defects in the cranial base are rare congenital anomalies that arise from failure of midline fusion (1). In infancy, they present in association with other facial, cranial, and intracranial midline defects (e.g., cleft lip and palate, hypotelorism, abnormalities of optic disc, nerve, and chiasm, pituitary or hypothalamic insufficiency, and agenesis of the corpus callosum). In adults, CSF rhinorrhea (spontaneous or iatrogenic, from biopsy of a nasal polyp), visual field loss, endocrinopathy, and a soft tissue mass in the epipharynx can occur. Indications for treatment in the adult include persistent rhinorrhea, progressive neurological deficit, and respiratory obstruction from an epipharyngeal mass. The risk of meningitis mandates treatment when rhinorrhea occurs (2). Even prolonged drainage of lumbar CSF can fail. When the sphenoid sinus is the route of CSF egress, transnasal transsphenoidal repair is a preferred, less invasive alternative to craniotomy. The defect must be small enough to be fully accessed by the transsphenoidal route, and the surgeon must be careful to avoid injuring protruding and potentially critical neurovascular structures. The authors’ use of a dural pouch is a clever modification that permits expansion of an otherwise inadequate plug.

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Fraioli et al. present a case of intrasphenoidal encephalocele associated with CSF fistula and subdural hematoma that ultimately resolved by the injection of fibrin glue into a dural patch used for repair. This is an innovative use of fibrin glue to repair a CSF leak. The senior author (LNS) would have preferred an intracranial approach to repair the CSF fistula with fascia lata and a pericranial flap initially. If transsphenoidal surgery is elected, autologous material, such as fascia lata or abdominal fascia supplemented with fat, may be better for repair because it is more pliable than artificial dura; it is more adhesive and results in better healing.

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