Endoscopic transnasal external fistulation in recurrent cystic subdiaphragmatic craniopharyngioma: a novel technique

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The authors present a technique for the persistent external drainage of intractable subdiaphragmatic cystic recurrences with the creation of a fistula between the cyst wall epithelium and epithelium on the nasal cavity, using a pedicled nasoseptal flap as a conduit. The long-term efficacy of endoscopic transnasal external fistulation (ETEF) in controlling cystic recurrences in this patient group is addressed through a retrospective observational review of 3 male patients aged 8, 22, and 45 years with the diagnosis of recurrent cystic subdiaphragmatic craniopharyngioma who underwent the ETEF procedure between 2006 and 2009. Clinical presentation, neuroimaging, surgical interventions, and follow-up were recorded. The main outcome measure was cyst reaccumulation on MRI.

Patients had a mean follow-up of 76 months (range 5–8 years) with no incidence of cystic recurrence. Follow-up imaging revealed sustained cyst involution contrary to the usual recurrent enlargement commonly seen in this patient group. Symptoms of headache and visual field defects improved post-ETEF. Long-term theoretical complications of a persistent fistula such as intracranial abscess, meningitis, or CSF leak were not observed. ETEF promotes nasalization of cystic recurrences in subdiaphragmatic craniopharyngioma. It is safe and effective, causing long-term involution of cysts and can be considered a definitive procedure.

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CRANIOPHARYNGIOMAS are tumors that arise due to embryogenic malformations in the sellar region and comprise 1%–4% of all brain tumors.21 Two main theories exist to explain these anomalies; the first suggests that craniopharyngioma originates from the embryonic remnants of Rathke’s pouch. The second proposes that such tumors arise from squamous cells of the par tuberalis of the pituitary gland. Indeed, the main pathologic subtypes of craniopharyngioma—adamantinomatous and squamous-papillary—may reflect the 2 respective hypotheses.21

The incidence of adamantinomatous craniopharyngioma has a bimodal distribution, with peaks being in childhood and also in adulthood, in contrast to the squamous-papillary subtype that is limited to adulthood.17,25 These lesions may be cystic, solid, or mixed in nature. Ninety percent of the adamantinomatous subtype and 50% of the squamous-papillary types are found to be cystic.1

The clinical significance of such midline tumors is that they exert mass effect, with patients presenting with visual and cognitive impairment, endocrinopathy, and symptoms of raised intracranial pressure.17

Historically, the management of craniopharyngioma has been challenging, as achieving successful, curative, gross-total resection of this lesion is rather difficult due to its proximity or adherence to surrounding critical neurovascular structures.2,4 Tumor and cystic recurrence is hence attributed to incomplete or subtotal resection, and thus rates of recurrence can be reduced with gross-total resection but at the cost of increased morbidity.2,20,26 Therefore, a widely accepted strategy in management of craniopharyngioma is a safe resection, which is as near total as possible without causing detriment by means of damage to critical neurovascular structures. Thereafter, adjuvant therapy such as radiotherapy may well be considered to augment the effects of a safe resection, which can be as effective as gross-total resection alone.9
Despite this, recurrence is common: as high as 50% in some series, and usually within 2–5 years of resection, but also reported after 30 years. There are a multitude of strategies directed at recurrence, the choice of which are influenced by cystic or solid predominance.

Strategies addressing recurrence include re-resection or cyst drainage via the transcranial or transsphenoidal routes; fractionated radiotherapy; Gamma Knife® and CyberKnife® stereotactic radiosurgery; and insertion of an intracystic Ommaya reservoir for repeated aspiration or administration of chemotherapy (bleomycin and interferon alpha). Although such strategies have been shown to be effective, they are not always definitive in addressing recurrence.

Our particular interest is that of recurrent cystic subdiaphragmatic craniopharyngioma. In light of the difficulty in achieving complete resection of the craniopharyngioma epithelium, our aims were to devise a persistent drainage pathway for cyst fluid in the context of recurrent cystic craniopharyngioma. This would provide a permanent drainage solution, thus preventing cystic reaccumulation and repeated surgery. Our technique would circumvent the need to completely excise the cyst epithelium, which can be risky, by providing a drainage pathway for the fluid continuously produced by the epithelium.

To create a permanent drainage pathway into which cystic fluid would be allowed to drain required the selection of an appropriate route. Given that one of the most common corridors of access for drainage of sellar cystic craniopharyngiomas is the endoscopic transnasal transsphenoidal route, drainage from the craniopharyngioma cyst into the sphenoid sinus and on to the postnasal space appeared to be a promising route. However, to maintain patency of this route, in contrast to transsphenoidal pituitary surgery, the sellar opening would not be closed or reconstructed after cyst drainage.

Within the setting of endoscopic transnasal transsphenoidal surgery of the skull base, the Hadad-Bassagasteguy pedicled nasoseptal flap has commonly been used to prevent or address CSF leaks within this region. Herein we propose a novel application for this pedicled mucosal flap, using it to line an area of craniopharyngioma cyst wall epithelium, with the aim of creating a fistula between the 2 epithelial surfaces.

The fistula created between the epithelium of the craniopharyngioma cyst cavity and the epithelium of the nasoseptal flap would allow persistent drainage of the cyst fluid out of the craniopharyngioma cyst, into the sphenoid sinus, and on to the postnasal space. However, the flap would need to be placed in such a manner as not to occlude the sphenoidotomy, sellar opening, and durotomy, and allow these apertures to remain open, thus providing a permanent drainage pathway and preventing cystic reaccumulation and its sequelae (Fig. 1).

Methods

Study Design and Patient Selection

We carried out a retrospective observational review of 3 cases involving male patients with recurrent cystic subdiaphragmatic craniopharyngioma, aged 8, 22, and 45 years at the time of undergoing endoscopic transnasal external fistulation (ETEF) between 1998 and 2014. All patients had previously undergone 1 or more established procedures (craniotomy and resection plus radiotherapy; microscopic transsphenoidal resection, endoscopic transsphenoidal drainage) to address predominant cystic craniopharyngioma of the sellar region at the Queen’s Medical Centre, Nottingham, UK (Table 1).

Preoperative Assessment

All patients received thorough endocrinological and ophthalmological review and optimization prior to surgery. Tumors were characterized by a neuroradiologist using contrast-enhanced MRI, and suitability for the ETEF procedure was determined by a neurosurgeon in conjunction with a rhinologist/endoscopic skull base surgeon.

Outcome Measures

Our main outcome measure was sustained cyst involution on follow-up MRI studies. Operative and postoperative complications together with neurological, endocrinological, and ophthalmological changes were sought from patient notes.

The ETEF Technique

The ETEF procedure is performed under a general anesthetic, and the nasal cavity is prepared with cocaine and adrenaline soaked cottonoids. The patient is positioned supine, with the head supported in a neutral position.

Using a 2-surgeon, 4-handed technique and the endoscopic transnasal transsphenoidal approach, the middle turbinates are lateralized to improve access and instrumentation. The sphenoid ostia are identified. A unilateral nasoseptal flap is raised upon the nasoseptal branch of the posterior septal artery, and placed aside for later use. Posterior septectomy is then performed, allowing for a wide sphenoidotomy. Mucosectomy is performed within the sphenoid sinus, followed by opening of the sella (Fig. 2).

A durotomy is performed and the cystic craniopharyn-
ETEF in recurrent cystic subdiaphragmatic craniopharyngioma

Results

Between June 2006 and May 2009, 3 male patients with a mean age of 25 years (8–45 years) underwent the ETEF procedure for recurrent cystic subdiaphragmatic craniopharyngioma. Each patient had 1 or more prior surgical attempts at addressing the lesion (Table 1). Two patients received radiotherapy after their initial surgery due to clinicoradiological disease progression. One patient did not receive radiotherapy, as the oncological perspective deemed the tumor to be predominantly cystic and symptomatic relief with stable disease was achieved after the first procedure. Stable disease was sustained for 3 years in this patient prior to performing ETEF. No radiotherapy was administered post-ETEF in this patient due to quiescent disease and no solid tumor expansion.

Preoperative MRI revealed predominantly cystic subdiaphragmatic craniopharyngioma. In all 3 cases, postoperative imaging revealed complete resolution of the cystic lesion, and this was sustained on subsequent imaging for a mean follow-up of 76 months, ranging from 60–96 months (Fig. 3). Follow-up nasoendoscopy at 4–6 weeks postoperatively revealed patent fenestrations for all 3 patients. Further nasoendoscopy at 3 months for Patient 3 and 2 years (longest endoscopic follow-up) for Patient 2 confirmed patent fenestrations. At 4-weeks postoperative-
ly for Patient 1, Fig. 4 demonstrates a patent fenestration, with the nasoseptal flap within the cyst cavity in continuity with the cyst epithelium and visible cyst effluent.

The incidence of CSF leak, meningitis, or cerebral abscess formation was 0%. Neurological deficits were not encountered by any patient. All 3 patients suffered from panhypopituitarism from the outset; they continued to receive hormone replacement therapy post-ETEF. Two patients had a continuation of DI that manifested after previous procedures.

Visual field defects were present in all patients upon initial diagnosis of craniopharyngioma. In 2 patients, this had improved after procedures prior to ETEF. A persistent small temporal scotoma in 1 of the patients did not change after ETEF. Symptoms of headache gradually improved in all patients postoperatively.

Discussion

The use of the endoscopic transnasal transsphenoidal technique is well established in the management of craniopharyngioma.\textsuperscript{3,5,8,26} Wide angled, superior views together with a binasal approach allowing a 4-handed technique permit resections equivalent to that of an open approach, without the increase in morbidity. Prior to the widespread use of endoscopy, the microscopic transsphenoidal route in the management of craniopharyngioma remained a longstanding successful alternative to traditional craniotomy in selected cases.\textsuperscript{13}

Despite the advantages of the endoscopic technique, achieving complete resection remains a challenge due to the intimate relationship of this tumor with critical structures such as the hypothalamus, pituitary, optic nerves, and cerebral vasculature. Recurrence of these lesions, therefore, remains a problem.

Although craniopharyngioma is considered a benign lesion, its behavior can appear malignant. The adverse, potentially reversible sequelae of this lesion arise due to the mass effect it exerts. Therefore, resection, whether complete or subtotal, together with cyst decompression can provide relief of symptoms, albeit transient in some cases.

Indeed, removing tumor tissue will reduce both mass effect and further production of cyst fluid and help to prevent cyst expansion. However, any residual tumor capsule also remains capable of secretion and subsequent cystic growth.

In the context of recurrent cystic subdiaphragmatic craniopharyngioma, our technique is focused less on achieving complete curative resection and more on striving for disease control. The ETEF procedure allows safe cyst decompression and also the creation of a persistent drainage pathway for cyst fluid, thus preventing reaccumulation. Such a technique circumvents the need to achieve total capsular clearance and, hence, avoids the inherent risks.

![Fig. 3. Pre-ETEF, post-ETEF, and most recent post-ETEF follow-up sagittal (left) and coronal (right) MR images obtained in all 3 patients. Preoperative (A) and postoperative (B) Gd-enhanced and 8 years postoperative (C) unenhanced T1-weighted MR images obtained in Case 1 (45-year-old male). Preoperative (D), postoperative (E), and 6 years postoperative (F) T1-weighted Gd-enhanced MR images obtained in Case 2 (22-year-old male). Preoperative (G) Gd-enhanced and postoperative (H) unenhanced T1-weighted MR images and 5 years postoperative (I) CISS sequence (left) and FLAIR image (right) in Case 3 (8-year-old male). Postcontrast sagittal and coronal images for C, H, and I were not available.](image-url)
A similar technique using the microscopic transsphenoidal route to fenestrate the cyst and maintain cyst patency with a silastic splint alone has, however, been previously described. What is novel about ETEF is the use of the pedicled nasoseptal flap to not only maintain patency of the fenestration but to create a true fistula. Such a communication between the nasal epithelium and cyst capsule epithelium using the nasoseptal flap results in a persistent epithelialized drainage route from cyst to the postnasal space.

Additionally, the use of the pedicled nasoseptal flap for the same purposes of providing a persistent drainage pathway has also been described in the context of endoscopic drainage of cholesterol granulomas of the petrous apex. A recent pediatric series involving the use of nasoseptal flaps revealed success for a variety of indications in patients ranging from age 5 to 17 years. The ETEF technique in our study has been performed in patients representative of both adult and pediatric groups (ages 8, 22, and 45 years), and it remains theoretically feasible in younger children, where endoscopic instrumentation is physically possible and a nasoseptal flap can be raised.

**Conclusions**

The use of established endoscopic transnasal transsphenoidal techniques, augmented by the expertise of both rhinologist and neurosurgeon, has resulted in the early success of ETEF. The creation of a fistula between the nasal epithelium and the craniopharyngioma cyst wall epithelium to create a persistent drainage pathway is a novel concept, resulting in nasalization of cystic recurrences. Thus far, ETEF appears to be a safe, effective, and definitive means of achieving disease control in patients with recurrent cystic subdiaphragmatic craniopharyngioma.

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**References**


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Supplemental Information
Previous Presentations
Interim follow-up results of this work have been presented in abstract form and published in the proceedings of the Society of British Neurological Surgeons Bi-Annual Conference (April 18, 2012, Aberdeen, Scotland, UK) and the World Federation of Skull Base Surgeons’ Conference (May 16–19, 2012, Brighton, England, UK).

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