
Nefize Turan, Emory University
Griffin R. Baum, Emory University
Christopher M. Holland, Emory University
Faiz Ahmad, Emory University
Oswaldo Henriquez Ajami, Emory University
Gustavo Pradilla, Emory University

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Upper Nasopharyngeal Corridor for Transnasal Endoscopic Drainage of Petroclival Cholesterol Granulomas: Alternative Access in Conchal Sphenoid Patients

Nefize Turan¹  Griffin R. Baum¹  Christopher M. Holland¹  Faiz U. Ahmad¹  Oswaldo A. Henriquez²  Gustavo Pradilla¹

¹ Emory University School of Medicine, Department of Neurosurgery, Atlanta, GA, United States
² Emory University School of Medicine, Department of Otolaryngology, Head and Neck Surgery, Atlanta, GA, United States

Address for correspondence Gustavo Pradilla, MD, Department of Neurosurgery, Emory University School of Medicine, 1365 Clifton Rd. NE, Suite 86166, Atlanta, GA, United States 30322 (e-mail: gpradil@emory.edu).

Abstract

Background  Cholesterol granulomas arising at the petrous apex can be treated via traditional open surgical, endoscopic, and endoscopic-assisted approaches. Endoscopic approaches require access to the sphenoid sinus, which is technically challenging in patients with conchal sphenoidal anatomy.

Clinical Presentation  A 55-year-old woman presented with intermittent headaches and tinnitus. Formal audiometry demonstrated moderately severe bilateral hearing loss. CT of the temporal bones and sella revealed a well-demarcated expansile lytic mass. MRI of the face, orbit, and neck showed a right petrous apex mass measuring 22 × 18 × 19 mm that was hyperintense on T1- and T2-weighted images without enhancement, consistent with a cholesterol granuloma. The patient had a conchal sphenoidal anatomy.

Operative Technique  Herein, we present an illustrative case of a low-lying petroclival cholesterol granuloma in a patient with conchal sphenoidal anatomy to describe an alternative high nasopharyngeal corridor for endoscopic transnasal transclival access.

Postoperative Course  Postoperatively, the patient’s symptoms recovered and no complications occurred. Follow-up imaging demonstrated a patent drainage tract without evidence of recurrence.

Conclusion  In patients with a conchal sphenoid sinus, endoscopic transnasal transclival access can be gained using a high nasopharyngeal approach. This corridor facilitates safe access to these lesions and others in this location.

Keywords
► cholesterol granuloma
► endoscopic
► nasopharyngeal
► petroclival
► petrous apex

Background

Cholesterol granulomas (CGs) are benign lesions of the temporal bone that develop most frequently at the petrous apex (PA).¹ These granulomatous lesions are cystic with a fibrous capsule and are often filled with a brown or yellow fluid. Extensive pneumatization of the PA has been identified as a major risk factor for CG formation and was observed in 10 to 30% of patients with CG. CGs are the most common primary lesions in this location² but can also be found in the mastoid bone, middle ear cavity, paranasal sinuses, orbitofrontal bone,
and in the petroclival region. They are distinct from cholesteatomas, which are also found in many of the same regions.

The pathogenesis of CGs has traditionally been explained by two theories: the obstructive-vacuum and the exposed marrow hypothesis. The former and more classic concept states that mucosal edema causes obstruction of air cell tracts and creates a vacuum effect resulting in extravasation of blood into these spaces and subsequent granulomatous reaction to cholesterol and blood products, forming the pathognomonic lesion, which grossly appears as an intraosseous cyst filled with dark, viscous, chocolate brown fluid and granulation tissue. Conversely, the exposed marrow hypothesis postulates that hyperpneumatization of the PA predisposes to bone marrow–air cell interface formation and bleeding from hypervascular bone marrow blebs.

Clinically these lesions are benign and often remain asymptomatic until significant mass effect occurs leading to cranial nerve dysfunction. However, most commonly, headache is the presenting symptom followed by hearing loss, vestibular dysfunction, tinnitus, diplopia, and ipsilateral retro-orbital pain. On radiologic imaging, CGs appear characteristically as expansile and erosive cystic lesions with well-defined margins on computed tomography (CT) and as high-signal intensity lesions on both T1- and T2-weighted images on magnetic resonance imaging (MRI) without contrast enhancement. These lesions have a similar appearance to cholesteatomas on CT; however, CGs are hyperintense on both T1- and T2-weighted imaging and cholesteatomas are T1 hypointense.

Traditionally CGs were treated surgically with open transcranial approaches, including infralabyrinthine, transcanal infracochlear, middle cranial fossa, transotic, translabyrinthine, and suboccipital retrosigmoid. The surgical exposure of the petroclival region necessary to access CGs, however, can be challenging due to the morphology of the petrous bone and the neurovascular contents of the peripetrous complex. Endoscopic and endoscopic-assisted endonasal approaches are increasingly used as less invasive alternatives that allow wide panoramic exposure and extended visualization.

Endonasal access is gained most commonly through a transsphenoidal corridor, which requires favorable sphenoidal anatomy. Four types of sphenoid sinus pneumatization have been described in the literature: conchal (nonpneumatized) (2%), presellar (21%), sellar (54.7%), and postellar (22.3%). In patients with a conchal sphenoid, a transsphe- noidal approach is technically dif- ficult. In this report, we present a case of a low-lying petroclival CG in a patient with conchal sphenoidal anatomy treated with an endoscopic transnasal transclival approach via a high nasopharyngeal corridor that illustrates the technique for nasopharyngeal access to these lesions.

**Clinical Presentation**

A 55-year-old woman presented with intermittent headaches and tinnitus without other neurological symptoms. Physical examination was unremarkable and formal audiometry demonstrated moderately severe bilateral hearing loss. CT of the temporal bones and sella revealed a well-demarcated expansile lytic mass with soft tissue density, centered in the right PA, with medial extension into the clivus and petroclival synchondrosis, inferior extension into the carotid space, and cephalic extension to the right petrous ridge. Medially, the mass scalloped the adjacent clivus but did not extend into the sphenoid sinus. Marked thinning of the medial wall of the vertical petrous segment and the posterior wall of the horizontal petrous ICA canals with large areas of complete bone density loss were concerning for extreme thinning versus dehiscence. The horizontal petrous ICA was also displaced slightly anteriorly by the soft tissue mass. Conchal anatomy of the sphenoid sinus was also noted. MRI of the face, orbit, and neck showed a right PA mass measuring 22 × 18 × 19 mm that was hyperintense on T1- and T2-weighted images without enhancement, consistent with CG. Given her symptomatic presentation treatment was recommended and in light of her sphenoidal anatomy, an upper nasopharyngeal corridor was selected to perform a transclival endoscopic approach to the lesion.

**Operative Technique**

Preoperative imaging studies including maxillofacial and sinus CT as well as a craniofacial MRI were acquired with a standard high-resolution protocol and loaded into a neuro-navigation system (Medtronic S7 StealthStation, Louisville, Colorado, United States). Three-dimensional images were reconstructed and the patient underwent co-registration of the stereotactic images with surface landmarks. Registration accuracy was verified using known anatomic landmarks and the scalp surface contour. A right maxillary antrostomy was performed, and a pedicled nasoseptal flap was elevated. Middle turbinatectomy was performed to maximize the exposure and the natural antrum was identified and opened. Following identification of the nasopharyngeal and eustachian tube landmarks, the mucosae of the nasopharyngeal area and adenoid bed were completely removed using coblation (ArthroCare ENT, Austin, Texas, United States) and drilling proceeded posteriorly and laterally until the cyst contents were reached. Inside the cystic cavity, membranes and brown/yellow material with an oil-like fluid

Fig. 1 A

Fig. 2 A

Fig. 1 B

Fig. 2 B

Fig. 1 C

Fig. 2 C

Fig. 1 D

Fig. 2 D

Fig. 1 E

Fig. 2 E

Fig. 1 F

Fig. 2 F
were encountered, and these were evacuated using a combination of 45- and 90-degree pituitary curettes as well as forceful irrigation with saline, and a combination of straight and angled suctions under direct visualization with the 0-degree endoscope. Once all the cyst contents were clearly evacuated, a 30-degree endoscope was advanced to inspect the cystic cavity and complete the decompression (►Fig. 2 D–J). Using the NICO Myriad aspirator (NICO Corporation, Indianapolis, Indiana, United States) additional areas of the cyst wall were resected and the diamond bur was used to extend drilling medially, superiorly, and inferiorly enlarging the caliber of the outlet tract to facilitate placement of the nasoseptal flap and tract mucosalization. Hemostasis was achieved and the nasoseptal flap was set in place along the decompression tract (►Fig. 2 K–O).

**Postoperative Course**

The patient’s postoperative course was unremarkable, and the postoperative MRI and CT demonstrated effective evacuation of the cyst contents and an open drainage tract from the resection cavity (►Fig. 3 A–D). She was discharged home in good condition on postoperative day 3. Tinnitus and headaches improved and outpatient follow-up evaluations were conducted at 1, 3, and 9 months postop. Endoscopic outpatient examination at 1 month revealed patency of the outflow tract. A subsequent endoscopic examination at 3 months showed partial stenosis of the outflow tract without recurrence of the lesion. Follow-up CT obtained 9 months after surgery showed a stable outflow tract without evidence of recurrence. Audiogram obtained at 9 months showed stable mild to moderate sensorineural hearing loss with 90 to 100% speech recognition preservation.

**Discussion**

The endoscopic transsphenoidal approach to petrous apex cholesterol granulomas (PACGs) was first described by Fucci et al in 1994. Subsequently, Mattox suggested using the endoscope as an adjunct during open surgical approaches and reported that the endoscopic tools were useful to mobilize and remove debris within long-standing CGs, remove septations, and drain multiloculated cysts. The addition of image guidance and improvements in endoscopic visualization and instrumentation enabled the widespread use of endoscopic transsphenoidal approaches, especially in patients with lesions neighboring the posterior wall of the sphenoid sinus.

Hearing preservation and permanent aeration of the cyst cavity constitute the main goals of treatment for CGs. Cyst drainage is therefore sufficient and complete removal of the cyst wall is not required. Scopel and colleagues have shown that endonasal approaches can provide drainage windows three times larger than those achieved with transcanal infracochlear approaches, and therefore may be associated with better outcomes and decreased recurrence. A recent
systematic review by Eytan et al included 53 patients with PACGs from 22 studies and showed a 20% restenosis rate and 7.5% overall recurrence after endoscopic treatment. Compared with 12.5% overall recurrence rate in open surgical approaches reported in the same study, the clinical outcomes of endoscopically managed CGs are thought to be favorable although prospective studies are needed to confirm these results. Although mini-flaps and stents have been used to decrease the recurrence of CGs, Eytan et al did not find significant decrease in recurrence rates with stenting. Surgical decision making and selection of the appropriate approach is dependent on hearing status of the patient at
presentation, pneumatization of the mastoid cavity and PA, location of the ICA in relation to the lesion, size and pneumatization of the sphenoid sinus, and ability to achieve permanent drainage. In particular, as the anatomic variations and the pattern of pneumatization in the sphenoid sinus can have a direct effect on surgical planning, careful preoperative evaluation of the anatomy of the sphenoid sinus by CT scan and MRI are of paramount importance. Endoscopic approaches such as the medial transsphenoidal approach with ICA lateralization and the transpterygoid infrapetrous approach have been described for the treatment of CGs. The medial transsphenoidal approach involves removal of the posterior septum and rostrum with wide sphenoidotomies to provide access to the lateral sphenoid. When the lesion is poorly visualized due to poor pneumatization of the sphenoid sinus, posterolateral localization, or lack of medial expansion of the lesion, a medial approach can be performed with additional bone removal and ICA lateralization. In that case, the ICA can be decompressed and lateralized using the vidian nerve and artery as landmarks to identify the paraclival ICA. In those cases, the clivus can be further drilled until the clival dura is exposed for augmentation of the surgical corridor. These approaches, however, rely on favorable sphenoidal anatomy for access.

When the sphenoid is not accessible due to poor pneumatization, a transpterygoid infrapetrous approach has also been described. This approach involves eustachian tube and foramen lacerum dissection, vidian artery and nerve exposure and mobilization, drilling of the pterygoid plates, and access through the eustachian tube. The PA is reached inferiorly by drilling the bone between the horizontal segment of the petrous ICA and the eustachian tube. Once the horizontal and vertical segments of the ICA are outlined, drilling is performed below the petrous ICA until the lesion is accessed. Compared with this approach, the operative technique in the present report describes an alternative transclival corridor using a high nasopharyngeal trajectory with intraoperative image and Doppler ultrasonography for ICA localization with the help of anatomic landmarks such as the vidian nerve and artery. This alternative approach can be implemented patients with conchal sphenoidal anatomy.

**Conclusion**

Conchal sphenoidal anatomy, although rare, occurs in up to 2% of patients. Access to the PA via endoscopic transsphenoidal approaches can be difficult in patients with a poorly pneumatized sphenoid as in the illustrative case reported. Therefore, use of a high nasopharyngeal corridor in conjunction with intraoperative navigation, Doppler ultrasonography, and external landmarks such as the vidian canal provides broad access to the PA for effective drainage of these lesions.
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