Aneurysms of the petrous internal carotid artery: anatomy, origins, and treatment

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Aneurysms arising in the petrous segment of the internal carotid artery (ICA) are rare. Although the causes of petrous ICA aneurysms remain unclear, traumatic, infectious, and congenital origins have been implicated in their development. These lesions can be detected incidentally on routine neuroimaging. Patients can also present with a wide spectrum of signs and symptoms, including cranial nerve palsies, Horner syndrome, pulsatile tinnitus, epistaxis, and otorrhea. The treatment of petrous ICA aneurysms remains challenging. Treatment options include close observation, endovascular therapies, and surgical trapping with or without revascularization. Management dilemmas exist, particularly for incidental lesions found in asymptomatic patients. The authors review the literature and discuss the anatomy of the petrous ICA as well as the pathophysiological features of aneurysms arising in this region, and they propose a management paradigm with current treatment options.

KEY WORDS • petrous internal carotid artery aneurysm • cerebrovascular bypass • balloon occlusion • endovascular therapy • stent

ANATOMY

Aneurysms of the petrous segment of the ICA are rare, and their true incidence is unknown. Most are considered congenital and their morphology is fusiform. Many are discovered incidentally in patients who require CT scans for other reasons. Typically, these lesions are large at the time of diagnosis, often with intraluminal thrombus. Petrous ICA aneurysms are thought to arise because of developmental weaknesses of the arterial wall at the sites of origin of regressed embryonic arteries including the caroticotympanic, pterygoid, vidian, stapedia, or the hyoid vessel. Histological examination demonstrates degeneration of the internal elastic lamina and medial aplasia as seen in intracranial berry aneurysms. The management of these lesions remains challenging, particularly in those that present incidentally in asymptomatic patients. Current treatment options include close observation, proximal occlusion or trapping with endovascular balloon occlusion, coil embolization, stent placement with or without coil embolization, and surgical trapping followed by revascularization through a high-flow bypass. In this review we discuss the anatomy, pathophysiology, presentation, treatment options, and management for petrous ICA aneurysms.

Abbreviations used in this paper: CT = computerized tomography; ECA = external carotid artery; ICA = internal carotid artery; OphA = ophthalmic artery.
foramen lacerum and anastomoses with branches from the ECA. The caroticotympanic artery, a vestige of the embryonic hyoid artery, is a small branch that arises near the genu of the petrous ICA and passes superiorly through the stapes to supply the middle ear cavity.

Knowledge of the anatomical relationship of the petrous ICA and its neighboring neuro-otological structures is very important. The greater and lesser superficial petrosal nerves lie lateral and superior to the horizontal segment of the petrous ICA. Deeper within the temporal bone are the tensor tympani muscle and eustachian tube, which are also lateral to the horizontal petrous ICA. The cochlea is situated posterior to the genu of the petrous ICA. As this vessel angles anteromedially it is limited superiorly by the cochlea, the tympanic cavity, and the geniculate ganglion, and laterally by the greater superficial petrosal nerve. The inferior petrosal sinus lies medially and slightly posteriorly to the petrous ICA. The ICA then exits the carotid canal above the foramen lacerum at the petrous apex (Fig. 2).

**PATHOPHYSIOLOGY**

The origin of petrous ICA aneurysms remains unclear, although several sources have been proposed, including traumatic, mycotic, and congenital. Atherosclerosis, which plays an important role in the rise of dolicho-ectatic and fusiform aneurysms, tends to spare the petrous ICA. Lesions in the petrous ICA can be further classified as true aneurysms or pseudoaneurysms. True aneurysms are characterized by walls that are continuous with the unaffected portion of the parent vessel and can develop from a traumatically weakened arterial wall. Pseudoaneurysms lack a true wall and develop when a thrombus and fibrous tissue capsule forms in response to injury to all layers of an arterial wall. Such injury may result from several mechanisms, including blunt or penetrating trauma, infections, inflammation, or radiation. Pseudoaneurysms form as cavities within adjacent blood clots and communicate with the lumen of the parent vessel. The lack of structural integrity of the wall of a pseudoaneurysm may result in rapid expansion if continued exposure to turbulent flow in the parent artery occurs.

Trauma is a significant cause of petrous ICA aneurysms and has been well documented. The anatomical arrangement of an immobilized petrous ICA segment distal to a mobile cervical ICA predisposes the petrous ICA to stretch forces and traumatic injuries that may induce formation of aneurysms. Patients may also present with a petrous aneurysm as a result of closed head trauma with or without associated fractures into the carotid canal; these are generally pseudoaneurysms resulting from dissection of the ICA. Iatrogenic injury to the petrous ICA during myringotomy and temporal bone surgeries has been reported.

Petrous ICA aneurysms of mycotic origin probably arise secondary to adventitial infection of the artery because of its close proximity to the middle ear and eustachian tube, rather than from hematogenous seeding. Infectious processes, including chronic otomastoiditis, tonsillar and pharyngeal infections, and cholesteatomas have been implicated in the development of these aneurysms. Chronic infection and the natural progressive enlargement of the aneurysm itself result in osseous erosion, rendering the ICA more susceptible to trauma or aneurysm dilation.

In the majority of cases, there is often no obvious cause and such lesions may be considered congenital in origin. In such cases, it has been postulated that aneurysms of the petrous ICA may arise directly from the main artery or from petrous ICA branches such as the caroticotympanic or vidian artery, the periosteal artery, and the persistent stapedial arteries. These aneurysms are thought to form as a result of developmental defects in the muscular, middle layer of the vessel. There are some arguments for a congenital origin because petrous ICA aneurysms have been diagnosed in children. There is one report of a petrous ICA aneurysm associated with von Recklinghausen neurofibromatosis.

**PRESENTATION**

Petrous ICA aneurysms are usually asymptomatic. Nevertheless, they can produce a wide range of clinical signs and symptoms, depending on the aneurysm’s size, direction of growth, and location within the carotid canal. Typically, patients present with symptoms in the third de-
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Fig. 2. Diagram illustrating the petrous (C2) ICA as well as its branches and adjacent structures. As it enters the carotid canal, the ICA lies anterior to the internal jugular vein. The eustachian tube (ET) is lateral and anterior. Note that the petrous ICA has an ascending vertical segment, a posterolateral genu, and a longer horizontal segment. The posterior genu in the intrapetrous ICA lies just in front of and below the cochlea. The two significant petrous ICA branches are the carotidocochlear canal (CTA) and the vidian artery (VA). The lacerum (C3) ICA segment is a short segment that extends from the endocranial opening of the carotid canal to the petrolingual ligament. It typically lies just above the foramen lacerum (FL, indicated by stippling), which in living patients is filled with fibrocartilage. Cranial nerves are indicated by Roman numerals, and carotid segments are indicated by Arabic numbers. CT = chorda tympani; GSPN = greater superficial petrosal nerve; IMA = internal maxillary artery (ECA branch); OC = optical canal; PTPF = pterygopalatine fossa; SOF = superorbital fissure; SPG = sphenopalatine ganglion. (Reprinted with permission from Osborn AG: Diagnostic Cerebral Angiography, ed 2, Lippincott Williams & Wilkins, Philadelphia, 1999, p 72.)

cade of life; these commonly include headaches, Horner syndrome, facial numbness, facial palsy, pulsatile tinnitus, dizziness, or hearing loss. Symptoms arise as a result of hemorrhage into or mass effect on neighboring structures of the carotid canal. Various combinations of single or multiple cranial nerve deficits associated with epistaxis, otorrhagia, vertigo, and dizziness have been reported. Because the ICA is surrounded by the sympathetic plexus, aneurysms in this location can present with Horner syndrome. Neurological deficits may occur due to compressive cranial nerve palsy or thromboembolic ischemic events.

Lateral extension of the lesion into the middle ear cavity may lead to pulsatile tinnitus, progressive hearing loss, and vertigo. Patients who present with aural symptoms frequently present initially to an otolaryngologist, and a vascular retrotympanic mass can be detected in most cases. These often can be mistaken for a glomus tympanicum tumor, and biopsy sampling of these lesions can produce a massive hemorrhage. A report exists of one rare case of a patient who had a giant petrous ICA aneurysm that presented as a cerebellopontine angle mass as a result of medial erosion of the petrous bone. When these aneurysms spontaneously rupture, dramatic hemorrhage into the eustachian tube or middle ear can present as massive epistaxis and/or otorrhagia. This presentation is thought to occur in 25% of patients with petrous aneurysms, but the true incidence is not known. Such bleeding episodes can be self limited or dramatic, sometimes requiring blood transfusions to reverse hypotension and anemia. Overall, these hemorrhages are always extracranial, and there have been no reported cases of intracranial hemorrhage from a ruptured petrous ICA aneurysm. Unlike most other intracranial aneurysms, petrous ICA lesions likely do not present with subarachnoid hemorrhage because of their extradural location.

TREATMENT

Once the diagnosis of a petrous segment ICA aneurysm is confirmed using cerebral angiography, it is important to decide if there is an indication for intervention or treatment. Patients with ruptured petrous ICA aneurysms have
traditionally undergone treatment to prevent future bleeding or to stop an active hemorrhage. In particular, the treatment for patients with pseudoaneurysms tends to be more aggressive because of the higher risk of hemorrhage and thromboembolism.

Typically, patients with unruptured petrous ICA aneurysms who experience unremitting symptoms including cranial nerve palsies have undergone procedures to attain symptomatic relief. In contrast, there have been cases in which patients had only mild symptoms that resolved without intervention.\(^{23,25}\) It is difficult to determine the best management of these lesions in an asymptomatic patient or in one with only mild symptoms. In these cases, it is relevant to compare the risk associated with a proposed treatment with that related to the natural history of the disease. It is necessary to take into consideration the age of the patient, preexisting medical comorbidities, risk of hemorrhage, likelihood of morbidity and death posthemorrhage, risk of a thromboembolic event, and the risk of aneurysm growth and the resulting neurological compromise from mass effect on neighboring vascular and neural anatomy.\(^{25,29}\)

Currently, treatment options for symptomatic and asymptomatic petrous ICA aneurysms include conservative management with serial imaging, endovascular ICA balloon occlusion, endovascular coil placement or stent-assisted coil insertion, placement of a flexible covered stent, or surgical trapping and revascularization with a high-flow bypass. The treatment should be tailored to the individual, depending on the condition of the patient and the nature of the lesion. The fusiform shape of these aneurysms has made it difficult or in many cases impossible to isolate them from the carotid circulation. Therefore, early treatments were aimed at occlusion of the parent vessel, which was achieved using a number of techniques.\(^{10}\) Although open surgical ligation of the common carotid artery or ICA is a well-described treatment for these aneurysms,\(^{12,24,52,55}\) a similar result can be achieved by a less invasive endovascular occlusion.\(^{10,14,21,25,53}\)

### Balloon Occlusion Test and Endovascular Therapies

Aneurysms in the petrous segment have been treated with parent artery occlusion by using endovascular balloon embolization since it was first described in 1978,\(^{10,14,21,25,53}\) and more recently with coil occlusion. This method has been successful in relieving the symptoms from petrous ICA aneurysms.\(^{10,21,25,29}\) In this procedure, a balloon is directed into the ICA immediately proximal to the aneurysm through a guide catheter from the common femoral artery.\(^{10,25,53}\) First, a balloon occlusion test is performed before parent artery occlusion of the ICA to evaluate for adequate collateral flow.\(^{10,25,53}\) While the balloon remains inflated for 30 minutes during the test occlusion, the patient is awake and is monitored with serial neurological examinations to evaluate for new deficits and with angiography to assess the collateral circulation, including the vertebral arteries and contralateral ICA.\(^{10,25,53}\) (Other monitoring options include electroencephalography and transcranial Doppler ultrasonography.\(^{10,53}\) If there are no adverse effects with the occlusion and if contralateral circulation is satisfactory, a detachable balloon is directed to the same site as in the occlusion study, inflated under direct fluoroscopic visualization, and detached. Then a second balloon is released proximally as a safeguard against deflation of the first balloon.\(^{10,25,53}\) A final injection of contrast material through the guide catheter is used to confirm complete occlusion of the aneurysm and patency of the collateral circulation.\(^{9,10,25}\)

Others have used embolization coils instead of the second balloon.\(^{12}\) Also, it is possible to perform an ICA occlusion exclusively with coil embolization.\(^{9}\) Nevertheless, there is a risk of coil dehiscence out of the vessel into the ear canal, which can cause life-threatening hemorrhages, especially in patients with ruptured aneurysms.\(^{9}\) Acute iatrogenic pseudoaneurysms located at the mesostympoma typically have no bone to encase the artery or pseudoaneurysm, thus precluding the option of coil embolization.\(^{1}\) Alternatively, balloon trapping procedures (balloons deposited on either side of the aneurysm) can be performed and may result in a decreased incidence of thromboembolic complications compared with proximal balloon occlusion.\(^{4,29}\) Nevertheless, the risk of dislodging a clot while traversing a partially thrombosed aneurysm with a detachable balloon must be strongly considered.

The major risk associated with ICA occlusion is ipsilateral cerebral infarction as a result of hypoperfusion, even in patients who have tolerated the test occlusion.\(^{10}\) There have been attempts to identify patients with borderline collateral flow by inducing hypotension during the test occlusion, by measuring ICA stump pressures, or by using various measures of regional cerebral perfusion.\(^{10}\) The risk of immediate complications from a balloon occlusion test ranges from 2 to 7%.\(^{9,49,59}\) Interestingly, the incidence of ischemic complications or cerebral infarctions after ICA occlusion, even with the use of a balloon occlusion test, ranges from 2 to 22%.\(^{16,17,30,38,49}\) Therefore, the risk of immediate or late ischemic complications and death in patients who tolerate the balloon occlusion test can be considerable, and there is no perfect way to predict accurately the safety of an ICA occlusion.\(^{39}\) Some patients may also suffer strokes years after undergoing this treatment.\(^{39,54}\)

An ICA occlusion is not a perfect treatment and there is a low incidence (3%) of continued filling of the aneurysm after proximal balloon occlusion.\(^{38}\) The aneurysm may also remain patent from filling through the distal ICA or via caroticotympanic, vidian, or periosteal branches.\(^{10}\) Another concern with ICA occlusion is the risk of cerebral aneurysm formation or growth after the procedure.\(^{3,34,35,62}\) These aneurysms may appear either in the arterial territory of the occluded vessel or remote from it, and this phenomenon is estimated to occur in 0 to 10% of cases.\(^{28,39,48}\) Overall, endovascular vessel sacrifice may be a less invasive procedure for petrous ICA aneurysms that are unsuitable for clip occlusion, but it is not without risks, even after a successful balloon occlusion test; therefore, other, newer treatment options should be considered.

In select cases it is possible to perform a balloon or coil embolization of a petrous ICA aneurysm and to preserve the parent vessel, but because most are giant fusiform aneurysms with no definable neck, this approach is not always feasible.\(^{29,63,66}\) There are increasing reports of petrous ICA aneurysms being successfully treated with Guglielmi detachable coils with or without the use of a stent to preserve the parent artery, to reconstruct the incompetent arterial wall, and to prevent future complications.\(^{19,37,40,45}\)
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New, flexible intracranial stents can be used to treat ICA aneurysms in difficult-to-access areas such as the petrous segment. The stent disrupts the aneurysm inflow tract, thereby inducing stasis and facilitating intraneurysmal thrombosis.35 Also, the stent acts as a scaffold to prevent coil herniation into the parent artery, which allows tight packing of wide-necked and irregularly shaped aneurysms.39 The stent also serves as a matrix for endothelial growth.

Among several limitations of stent-assisted coil insertion is stent-induced intimal hyperplasia, which can result in hemodynamically significant stenosis.65 The high porosity of intravascular stents is another limitation for aneurysm thrombosis after stent placement.37 The novel use of covered stents allows for immediate reconstruction of the arterial wall while preserving the flow of the parent artery. This technology may be applied to cases of artery laceration or pseudoaneurysms of the petrous ICA. Although endovascular stent treatment with or without coil embolization of petrous ICA aneurysms appears promising, close follow up will be necessary to assess long-term patency rates and to detect delayed complications from thromboembolic events, stent-induced stenosis, or thrombosis. Overall, the use of stents is a work in progress, and with new technology its application in the treatment of petrous ICA aneurysms has increased.

Cerebral Revascularization With High-Flow Interpositional Bypass

In patients with inadequate collateral flow to allow an ICA occlusion, a cerebrovascular high-flow bypass procedure can be performed, followed by endovascular aneurysm embolization, occlusion of the parent vessel, surgical trapping of the aneurysm, or resection of the lesion.29,44,50 Direct surgical exposure of petrous ICA aneurysms is a complex procedure because of the difficulties involved in accessing the lesions deep within the petrous bone.21 Early attempts at surgery were associated with high rates of mortality.27

Cerebral revascularization with a high-flow interpositional bypass is a feasible strategy in patients with complex aneurysms that cannot be directly clipped and who do not tolerate ICA occlusion, as demonstrated with a balloon occlusion test.30,32,53 As mentioned earlier, the balloon occlusion test produces false-negative results in predicting which patients will experience complications related to a surgical ICA occlusion.29 Some surgeons select revascularization procedures, even if the patient tolerates a balloon occlusion test without neurological deficits, if the individual exhibits a marked asymmetrical decrease in hemispheric cerebral blood flow (<30 ml/100 g/min) during the study.37 For younger patients with a longer life expectancy, the goal of preserving patency of the ICA should be strongly considered.

Revascularization procedures in the hands of skilled microsurgeons yield a low morbidity and mortality rate and reduce the risk of stroke that arises with isolated ICA occlusion procedures. In a series of 61 patients requiring cerebral revascularization who were described by Lawton and coworkers,39,57 patients had good outcomes and there was only one death. In five of these patients an aneurysm was identified along the petrous segment of the ICA. Overall, there was a 95% graft patency rate immediately postoperatively and a 92% late graft patency rate at a mean of 5.1 months after surgery. In this series, aneurysms located at the junction of the cervical and petrous ICA were resected and the parent vessel was reconstructed with a saphenous vein interposition graft. Aneurysms located in the petrous ICA were trapped proximally in the neck and distally along the petrous ICA. An entirely extradural cervical–petrous ICA bypass was performed. In another report, one patient with a petrous ICA aneurysm was successfully treated with a superficial temporal artery–middle cerebral artery bypass followed by balloon entrapment of the aneurysm without complication.64 In yet another series, ischemic symptoms were eliminated in four patients after bypass procedures for aneurysms of the petrous ICA, but these were all traumatic pseudoaneurysms.64

Several other approaches for the direct repair of petrous ICA aneurysms have been described. The first successful petrous ICA resection and primary anastomosis was reported by Glasscock and colleagues in 1983.34 It was performed via a modified infratemporal fossa approach, based on the Fisch approach for surgically removing extensive glomus jugulare tumors.19,20 After proximal and distal occlusion of the aneurysm, the intervening affected portion of the artery was removed and a primary end-to-end anastomosis was performed.24 This approach is quite invasive because it involves extensive petrous bone drilling, facial nerve transposition, mandibular condylectomy, and transection and closure of the external ear canal. Alternatively, Sekhar et al.56 advocated a combined subtemporal and preauricular infratemporal fossa approach, which affords excellent access to this region while preserving hearing and facial nerve function. Contraindications to direct repair of a petrous ICA aneurysm include poorly controlled active bleeding that requires immediate ICA occlusion or aneurysmal extension past the carotid canal, which precludes adequate exposure.24

Submandibular Cervical–Supraclavicular ICA Saphenous Vein Bypass

At our institution, we prefer the submandibular cervical–supraclavicular ICA interpositional saphenous vein graft bypass (Fig. 3). The details of this method have been previously described by the senior author (W.T.C.).13,61 This approach is less invasive than the subtemporal and infratemporal fossa approaches and does not require direct exposure of the aneurysm, which involves extensive petrous temporal bone drilling. We also prefer to tunnel the graft via the submandibular route. One advantage of this method is that it permits a more direct routing of the bypass graft to the recipient ophthalmic segment of the ICA. When the graft is tunneled superiorly and then cut to the proper length, its overall length is shortened, thereby promoting graft patency. In addition, the submandibular placement of the graft provides physical protection by the mandible, temporal muscle, and zygoma, in contrast to pre- or postauricular tunneling of the graft.

For this approach, a standard pterional craniotomy is performed on the side of the lesion. The lateral aspect of the sphenoid wing and the anterior clinoid process are removed extradurally by using a high-speed drill. This al-
lows adequate exposure to the ophthalmic segment of the ICA, which serves as the recipient vessel for the saphenous vein bypass. A neck incision is made along the anterior border of the sternocleidomastoid muscle and the cervical ICA is isolated. A segment of the greater saphenous vein measuring approximately 20 cm is harvested and perfused with heparinized saline.

Prior to making the submandibular pass, the zygoma is removed and reflected inferiorly with the masseter and temporalis muscles. A bone trough is also made at the middle fossa skull base to provide room for the graft. This step is important in avoiding graft compromise by mandibular movement. After induction of burst suppression with intravenous propofol and administration of 5000 U intravenous heparin, the cervical ICA is ligated as high in the neck as possible (this will ultimately shorten the graft length). An end-to-end proximal anastomosis is performed using the saphenous vein graft. The ICA may be temporarily mobilized inferiorly, which enables an easier technical proximal anastomosis. By using a short ventriculoperitoneal shunt passer or large cannula to accept the graft, the distal end is advanced beneath the mandible and through the skull base atraumatically. The graft is sized appropriately and anastomosed to the ophthalmic segment of the ICA between the OphA and posterior communicating artery branches. This is usually performed end-to-side to preserve the ophthalmic branch, or, if the OphA is not preserved, the anastomosis is performed end-to-end. A permanent vascular clip is placed, or suture ligation is performed just proximal to the OphA branch. This particular bypass technique has proved to be useful, with excellent long-term patency rates, and is an effective alternative route for interpositional vein graft placement in cases in which high-flow revascularization is desired (a case is illustrated in Figs. 4–7).

**MANAGEMENT PARADIGM**

Guidelines for the management of aneurysms of the petrous ICA are not well defined. There are three different situations that warrant discussion, as follows: 1) the asymptomatic patient or one with nondebilitating or mild symptoms; 2) the patient with significant symptoms including cranial nerve palsies or other neurological deficits related to mass effect from the aneurysm; and 3) the patient with a ruptured aneurysm.

**Asymptomatic Patients**

When deciding how to treat an asymptomatic patient, it is important to consider what is known about the natural history of petrous ICA aneurysms. Although up to 25% of patients may present with a hemorrhage, typically these are not subarachnoid in nature, patients come to medical attention in a timely manner, and most do not suffer life-threatening bleeding episodes. Although large fusiform aneurysms sometimes have associated thrombi, there have been no reports of thromboembolic strokes as a result of these lesions. It is reasonable to recommend conservative management with serial imaging for these incidental aneurysms. If, however, the aneurysm enlarges on follow-up imaging (with or without increased symptoms from mass effect), treatment should be considered because of radiographic progression.

**Symptomatic Patients**

For a patient who is symptomatic from cranial neuropathies and mass effect, we recommend treatment. The options include endovascular therapy or a high-flow bypass, as discussed earlier, given that the patient is medically sta-

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Fig. 3. Drawing showing a cervical–supraclinoid submandibular saphenous vein bypass. GA = giant petrous aneurysm (excluded); OA = OphA; SVG = saphenous vein graft. (Reprinted with permission from Couldwell WT, et al: J Neurosurg 94: 806–810, 2001.)

Fig. 4. Left: Preoperative CT scan with bone windows obtained in a 47-year-old man who presented with a left sixth cranial nerve palsy. The CT scan demonstrates erosion of the petrous apex and adjacent clivus on the left side. Right: Preoperative magnetic resonance image (fast low-angle inversion-recovery study) demonstrating high signal intensity within the giant petrous ICA aneurysm, with likely impingement of the abducent nerve near the region of the Dorello canal. (Reprinted with permission from Couldwell WT, et al: J Neurosurg 94:806–810, 2001.)
ble and able to tolerate these treatments. Endovascular therapies with balloon occlusion should be considered first if the patient is able to tolerate balloon occlusion testing. The patient should be informed of the risks related to ICA occlusion, however, especially a young patient with a long life expectancy. Endovascular stent placement with or without coil occlusion is an evolving therapy and may provide immediate reconstruction of the ICA wall while preserving the parent artery. A high-flow bypass should be considered if endovascular therapies are not feasible.

Patients With Ruptured Aneurysms

Finally, we recommend urgent treatment for a patient with a ruptured petrous ICA aneurysm. In the acute setting of an active hemorrhage, immediate endovascular balloon occlusion may be used to stabilize the patient. The use of coil embolization is not recommended by some authors because of the risk of coil extrusion into the ear canal in aneurysms presenting at this location. A high-flow bypass may be performed if necessary.

CONCLUSIONS

The treatment of petrous ICA aneurysms remains a challenge. Management dilemmas exist when incidental lesions are found in asymptomatic patients. Treatment should be tailored to the individual and patient selection is critical. Endovascular strategies appear to be the first-line therapy if balloon occlusion testing is tolerated by the patient. Novel endovascular stent technologies, including stent-assisted coil placement and covered stents, may eliminate the aneurysm while preserving flow in the parent artery. Surgical revascularization with an interpositional high-flow bypass graft is an important strategy for lesions that are not amenable to endovascular therapy.

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