Case Report

Cavernous Carotid Aneurysms: To Do or Not To Do?

Sudha Menon, R. Girish Menon

Departments of Ophthalmology and Neurosurgery, Kasturba Medical College, Manipal, Karnataka, India

INTRODUCTION

Cavernous carotid aneurysms (CCAs) pose considerable dilemmas in management. It is still unclear as to whether an asymptomatic cavernous carotid aneurysm should be subjected to treatment. Similarly, the ideal management strategy for a symptomatic aneurysm is controversial. We present the case of a 60-year-old female with a giant CCA and discuss the management issues.

CASE REPORT

A 60-year-old hypertensive female presented with complaints of blurring of vision and left hemifacial and periorbital pain of 2 months duration. One month into her illness, she developed drooping of her left eyelid. On admission, the general systemic examination was normal. She had diminished visual acuity of her left eye and could only perceive hand movements. She had complete ophthalmoplegia of her left eye along with sensory blunting involving the first two divisions of the left trigeminal nerve. Rest of her nervous system examination was normal. Her magnetic resonance imaging [Figure 1] revealed a well-defined extra-axial altered signal intensity lesion measuring 4.0 cm × 4.1 cm × 3.4 cm in the left parasellar region which was iso to hyperintense on T1-weighted imaging, heterogeneous on T2-weighted/fluid-attenuated inversion recovery and did not show any diffusion restriction. On postcontrast studies, the central portion of the lesion showed homogeneous intense enhancement while rest of the lesion did not show enhancement. The left optic nerve appeared displaced and pushed medially and upward. The sphenoid bone was unremarkable. Digital subtraction angiography [Figure 2] revealed a giant cavernous aneurysm measuring nearly 3.5 cm × 2.4 cm with areas of thrombosis. An aneurysm was seen extending superiorly into the intradural subarachnoid compartment. Cross circulation studies following compression of the left carotid artery revealed inadequate filling of the ipsilateral middle cerebral vessels, but there was a venous filling delay of over 5 s in the ipsilateral side. She underwent a high flow extracranial-intracranial (EC-IC) (proximal internal carotid artery [ICA] to M2 segment of middle cerebral artery) bypass using a saphenous vein graft followed by ICA ligation. Postoperative computed tomography angiogram [Figure 3] revealed complete thrombosis of the aneurysm with no evidence of contrast enhancement. The bypass graft showed normal contrast opacification and there was no flow detected in the ICA distal to the occlusion.

DISCUSSION

Cavernous carotid aneurysms (CCAs) constitute 2%–9% of all IC aneurysms. These aneurysms can be idiopathic, traumatic, iatrogenic, or infectious in etiology. Traumatic, iatrogenic, and infectious aneurysms have an aggressive course and need urgent intervention. The natural history of idiopathic cavernous aneurysms is not well known. They often remain asymptomatic and are detected incidentally. They tend to become large to giant in size when they manifest with features of mass effect in the form of cranial nerve palsies of adjacent nerves. This could be in the form of diplopia, ptosis, ophthalmoplegia, or pain or paresthesia along the fifth nerve distribution. A Large transitional variant of these aneurysms which have an intradural component can press against the optic nerve and result in visual symptoms. These intradural variants also carry a risk of subarachnoid hemorrhage (0.2%–0.4%). Intracavernous rupture can result in direct carotico cavernous fistula although such instances are rare. Rarely, these aneurysms can erode into the sphenoid sinus and rupture resulting in fatal epistaxis. Spontaneous thrombosis of these aneurysms has also been reported as have been thromboembolic strokes originating from intra-aneurysmal thrombus.

The dilemmas in the management essentially pertain to (a). What are the chances that an asymptomatic CCA would turn symptomatic

Address for correspondence: Dr. R. Girish Menon, Department of Neurosurgery, Kasturba Medical College, Manipal - 576 104, Karnataka, India. E-mail: girish.menon@manipal.edu

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and is prophylactic treatment indicated in such aneurysms? (b) What are the treatment options for a symptomatic CCA?

**Unruptured asymptomatic aneurysms**

These aneurysms are known to have an overall low risk of rupture and life-altering complications. Being enclosed in a safe venous pouch, the cavernous sinus, these aneurysms tend to grow from large to giant size before they manifest clinically. In the ISUIA study, the 5-year cumulative rupture rate of unruptured CCAs was 0% for aneurysms with size ≤12 mm, 3% for aneurysms of size 13–24 mm and 6.4% for aneurysms >5 mm.[9] Similarly, risk prediction for bleed based on the PHASES score remains high for these aneurysms as the majority of these aneurysms tend to be giant.[9] However, for some reason, the common predictors of bleed for an incidental aneurysm such as hypertension, age, sex, and previous bleeds, do not apply strictly for CCAs There are several published series on successful conservative management of cavernous segment aneurysms the largest being by Stiebel-Kalish et al.[9] In their series of 132 aneurysms followed up for 4 years, 39 patients improved 21 remained unchanged, eight patients worsened and two died, suggesting a rather benign course compared to other aneurysms. The need for any prophylactic intervention purely based on the size of the aneurysm, thus, remains debatable.

Intervention for an asymptomatic aneurysm is justified in three situations. Aneurysms with an intradural component carry a risk of subarachnoid hemorrhage and need to be promptly excluded from the circulation. Such intradural components are often missed if not looked for specifically. Appropriate imaging of the dural rings is, therefore, mandatory for all CCAs. Similarly, an expanding aneurysm can result in sphenoid bone erosion, which can eventually result in fatal epistaxis. The presence of sphenoid bone erosion on bone window scans is the second indication for prophylactic intervention in an asymptomatic aneurysm. The third instance for preemptive treatment for a CCA would be an aneurysm seen to have increased in size on serial angiograms, especially in a young individual.

**Symptomatic aneurysms**

It is reasonable to assume that all symptomatic CCAs should be treated. Symptomatic aneurysms which have bled, which produce thromboembolism, which cause intolerable pain and deteriorating vision definitely merit treatment. Cranial nerve palsies other than vision, association with previously ruptured aneurysms, etc., are gray areas where treatment needs to be individualized. However, treatment is justified only if a successful cure can be offered without major complications. Both surgical and endovascular modalities of treatment essentially attempt to exclude the aneurysm from the circulation and initiate thrombosis. Except for direct clipping of the aneurysm none of the other modalities attempt to decompress the aneurysm and thereby reduce mass effect. Pain is one symptom which responds moderately well to treatment. Successful intervention certainly reduces the risk of further bleed and ischemic stroke. Intervention helps in arresting the progression of cranial nerve deficits and complete recovery seldom happens. The second cranial nerve is an exception and visual recovery following decompression has been reported in few cases.[12] Like other causes of cranial nerve deficits, the duration of the deficits before treatment can have an impact on the resolution of the deficit.

Unfortunately, all modalities carry a considerable risk of complications. Both endovascular and surgical options carry significant risks, and the overall mortality and morbidity varies from 3.2%–22.6% to 9.2%–14.8%, respectively.[11] The decision to intervene thus needs to be made judiciously and the type of treatment ought to be justified adequately.

**Treatment modalities**

Exclusion of the aneurysm from the circulation can be done either by surgery or by endovascular techniques. Treatment can be either occlusive or reconstructive. Occlusive strategies include parent artery ligation surgically or by endovascular techniques. Reconstructive strategies include
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The next however, in developing countries like [13-16] the risk of infarct with 15 cases of more than 0.5 s in venous filling in the vascular territory of the occluded vessel which indicates nontolerance. This simple test described by Müller-Forell and Valavanis in 1995 and later validated by van Rooij et al. is a simple safe and effective in predicting tolerance to carotid occlusion and has become the current standard of practice in many centers.[17,18] The risk of infarction comes down to 22% in patients who successfully complete all preoperative assessment tests.[19,20] The risk, however, does not completely disappear. With an additional bypass, this risk comes down to 14.6% (1.8–29.4). A bypass procedure before parent artery occlusion is, therefore, preferable to reduce the risks of postocclusion stroke even in patients who tolerate BTO successfully (universal bypass).

Bypass could be an augmentative Superficial temporal to Middle cerebral artery (low flow STMC) bypass for patients with moderate cerebrovascular reserve or a replacement bypass (high flow EC-IC) bypass for those with poor cerebrovascular reserve. The choice between an arterial radial artery graft and a venous saphenous graft depends on the surgeon’s preference. Radial artery graft lasts longer but has potential risk of spasm. Venous graft is easier to harvest but is prone to kinking and occlusion. Similarly, the choice between parent artery occlusion alone or trapping is difficult as it has been shown that there is no difference in complications and outcome between trapping and carotid occlusion.[13] Trapping supposedly prevents backflow into the aneurysm due to cross circulation and also prevents dislodged thrombi from reaching the IC circulation. Trapping is preferred to mere carotid occlusion for aneurysms with significant intradural extension and patients who demonstrate significant retrograde flow during the BTO.[21] The next controversy pertains to the choice of the proximal site for anastomosis. Common carotid artery, ICA and external carotid artery all have comparable results and the decision essentially depends on the surgeon’s preference. We prefer the ICA or the external carotid artery to avoid manipulation of the carotid bulb situated close to the bifurcation.

Our patient presented with poor vision and intolerable pain justifying treatment. We did not perform a BTO, SPECT, or PET study for our patient as the cross compression study revealed poor cross circulation. Saphenous vein graft and ICA for proximal anastomosis was chosen due to surgeon’s preference and past experience.

**Conclusion**

CCA are rare and pose considerable challenges in management. Their natural history is unclear, but they seldom cause life-threatening complications. The decision to treat needs to be made judiciously based on careful interpretation of the radiological images. Venous filling on cross compression is a good indicator for cross circulation and obviates the need for detailed cerebrovascular reserve studies in selected cases. Endovascular treatment strategies offer reasonable results but are expensive. Parent artery occlusion combined with a universal bypass is a cost-effective alternate option with comparable results.

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There are no conflicts of interest.
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