Carotid-cavernous fistulas

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Carotid-cavernous fistulas (CCFs) are vascular shunts allowing blood to flow from the carotid artery into the cavernous sinus. The characteristic clinical features seen in patients with CCFs are the sequelae of hemodynamic dysfunction within the cavernous sinus. Once routinely treated with open surgical procedures, including carotid ligation or trapping and cavernous sinus exploration, endovascular therapy is now the treatment modality of choice in many cases. The authors provide a review of CCFs, detailing the current classification and clinical management of these lesions. Therapeutic options including conservative management, open surgery, endovascular intervention, and radiosurgical therapy are presented. The complications and treatment results as reported in the contemporary literature are also reviewed.

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**Key Words**
- angiography
- carotid artery
- carotid-cavernous fistula
- cavernous sinus
- embolization
- shunt

Located just lateral to the sella turcica, the cavernous sinus is a trabeculated venous cavity invested by the dura mater and contains several neural and vascular structures. Named by Jacobus Winslow in 1734, the term “cavernous sinus” is a misnomer, as it is neither cavernous nor is it truly a sinus. Dwight Parkinson is credited for his extensive efforts in the 1960s and 1970s to further characterize the anatomy of the cavernous sinus and to define CCFs. Parkinson, along with Hashimoto and colleagues, emphasized the inappropriateness of the term “cavernous sinus,” preferring the more anatomically correct term “lateral sellar compartment.” In any event, the term “cavernous sinus” remains common in medical parlance today.

On average, the cavernous sinus measures 2 cm anteroposteriorly, 1 cm in lateral width, and 1.3 cm in vertical dimension. The major neural and vascular structures traversing the cavernous sinus include the ICA and CNs III, IV, V1, V2, and VI (Fig. 1). The oculomotor nerve (CN III), trochlear nerve (CN IV), and the first and second divisions of the trigeminal nerve (V1 and V2) course between the 2 dural layers of the lateral wall of the cavernous sinus. The abducens nerve (CN VI) runs inside the cavernous sinus itself, lateral to the ICA.

The cavernous sinus has been described as both a trabeculated venous channel and as a venous plexus. Harris and Rhoton divided the cavernous sinus into the following 4 compartments: medial, lateral, anteroinferior, and posteroinferior with respect to the ICA. Normally, the cavernous sinus receives drainage from the sphenoparietal sinus, the superior ophthalmic vein, and the superficial sylvian vein. The middle meningeal vein empties into the cavernous sinus laterally, while the superior and inferior petrosal sinuses drain the cavernous sinus posteriorly. Various venous anastomoses connect the 2 cavernous sinuses, including an intercavernous sinus and a basilar plexus of veins. These anastomoses are clinically relevant, both in terms of the spread of extracranial infections and neoplasms and as alternative routes for venous drainage in patients with obstruction of the cavernous sinus.

**Fistula Classification**

Carotid-cavernous fistulas are abnormal vascular shunts, allowing blood to flow either directly or indirectly from the carotid artery into the cavernous sinus. Carotid-cavernous fistulas have been classified according to the hemodynamic properties, etiology, or anatomy of the fistula (Table 1). Hemodynamic classification separates CCFs into high-flow and low-flow fistulas. Etiological classification distinguishes spontaneous lesions from those occurring due to trauma. Anatomical classification defines direct CCFs as those arising directly from the carotid artery, while indirect CCFs are those originating from carotid artery branch vessels.
Based on the nomenclature of Peeters and Kröger, Barrow et al. defined 4 types (Types A–D) of CCFs (Fig. 2). Type A CCFs are direct, high-flow lesions connecting the ICA directly to the cavernous sinus. Type A CCFs often result from a single tear in the carotid artery wall, caused either by trauma or aneurysm rupture. These are by far the most common type of CCFs, accounting for approximately 75%–80% of CCFs overall. Type A CCFs are now less commonly seen in part due to increased motor vehicle safety measures such as the standardization of airbags and widespread usage of seat belts. Type B, C, and D CCFs are all indirect, low-flow lesions that arise from meningeal branches of either the ICA or ECA. Type B CCFs arise from meningeal branches of the ICA, Type C CCFs arise from meningeal branches of the ECA, and Type D CCFs arise from meningeal branches of the ICA and ECA.

In a study of 132 patients presenting with CCFs, 100 CCFs (75.8%) were classified as Type A, none were classified as Type B, 4 (3%) were classified as Type C, and 28 (21.6%) were classified as Type D. Meyers et al. characterized the arterial supply of 135 dural (Types B–D) CCFs and showed that the internal maxillary, middle meningeal, and meningo-hypophyseal trunks and capsular arteries each supplied more than 50% of the fistulas.

**Epidemiology and Etiology**

**Traumatic CCFs**

Traumatic CCFs are the most common type, accounting for up to 75% of all CCFs. They have been reported to occur in 0.2% of patients with craniocerebral trauma and in up to 4% of patients who sustain a basilar skull fracture. Consistent with the demographics associated with traumatic injuries, traumatic CCFs are most commonly seen in young male patients. They typically occur as a result of a closed head injury–associated basilar skull fracture. There are various theories regarding the mechanism of CCF formation after head trauma. One theory holds that the carotid artery is directly torn either by a bony fracture or by shear forces during the traumatic incident. Finding no skull fractures in a series of 42 patients with posttraumatic direct CCFs, Helmke et al. proposed the following alternative theory: that there is a sudden increase in intraluminal pressure of the ICA with concurrent distal artery compression, which forces rupture of the vessel wall and results in a CCF. While the vast majority of CCFs caused by trauma are direct CCFs, posttraumatic indirect CCFs have been reported as well.

Traumatic CCFs may also result from projectile or slash injuries that result in laceration of the cavernous carotid artery, allowing high-flow communication with the cavernous sinus. Although rare, there have also been reports of CCFs caused by iatrogenic injury during craniotomy, carotid endarterectomy, transsphenoidal exploration, endovascular procedures, and sinus surgery. Bilateral CCFs are seen in 1%–2% of patients with posttraumatic CCFs.

**Spontaneous CCFs**

Spontaneous CCFs, which account for approximately 30% of all CCFs, are typically found in older, female patients. Ruptured cavernous ICA aneurysms are an often-cited cause of spontaneous, direct CCFs. Cavernous-carotid fistulas are reported in anywhere from 3% to 24% of patients with cavernous carotid aneurysms. In addition to cavernous aneurysms, genetic conditions such as fibromuscular dysplasia, Ehlers-Danlos syndrome, and pseudoxanthoma elasticum are known to predispose patients to spontaneous CCFs. It is thought that arterial wall defects in these patients predispose them to CCF formation after minor stress, such as coughing or Valsalva maneuver. In individuals without cavernous aneurysms or predisposition syndromes, it has been theorized that microscopic venous thrombosis or increases in venous sinus pressure may facilitate fistula formation by causing microscopic breaks in dural vessels to the cavernous sinus. Factors thought to contribute to the breaks in these vessels include arterial hypertension, atherosclerotic vascular disease, pregnancy, minor trauma, straining, diabetic vascular disease, and collagen vascular disease.

**Clinical Presentation**

**Direct CCFs**

Consistent with their etiology either from traumatic

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**TABLE 1: Classification of CCFs**

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<th>Hemodynamic</th>
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<td>Etiological</td>
<td>Spontaneous vs traumatic</td>
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<td>Anatomical</td>
<td>Direct vs indirect</td>
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**Fig. 1.** Schematic organization of the cavernous sinus in coronal section. The major neurovascular structures including the ICA and CNs III–VI within the cavernous sinus are shown. Printed with permission from Jason A. Ellis.
Carotid-cavernous fistulas

or aneurysmal rupture, high-flow, direct CCFs often present acutely. They typically progress rapidly, necessitating urgent treatment. The most common presenting signs and symptoms include proptosis in 72%–98%, chemo-
sis in 55%–100%, orbital bruits in 71%–80%, and headache in 25%–84% of patients. Additionally, the majority of patients complain of visual disturbances, including diplopia reported in 88% of patients, blurry vi-
sion, and orbital pain. These visual complaints may be due to retinal ischemia and may indicate the need for urgent intervention. Ophthalmoplegia has been report-
ed in 23%–63% of patients, while other cranial nerve deficits have been reported in 17%–44% of patients. Less common presentations include intracerebral or sub-
arachnoid hemorrhage in 5% of patients.

Indirect CCFs

In contrast to high-flow CCFs, so-called low-flow, indirect CCFs tend to be more insidious in onset. Conjunc-
tival injection is often the most prominent feature, and patients are commonly treated for other conditions, such as conjunctivitis, before a correct diagnosis is made. The disease course may be chronic or relapsing and remitting in nature, leading to a delay diagnosis and treatment of a year or more in more than 50% of patients. In a series of 135 patients with indirect CCFs, common signs and symptoms included arteriovenous shunting of conjunctival veins in 93%, chemosis in 87%, proptosis in 81%, diplopia with ophthalmoplegia in 68%, cranial bruit in 49%, retroor-
bitai headache in 34%, elevated intraocular pressure in 34%, and decreased visual acuity in 31%.

The precise pattern of symptoms is dependent on the rate of flow, the location of venous drainage of the CCF, inflammation, and pressure within the venous sinus. Ad-
ditionally, the pattern of venous drainage may change with the development and resolution of a thrombosis, resulting in an inconsistent pattern of symptoms. Anterior draining CCFs, the most common type of indirect CCF, usually present with ocular and orbital symptoms such as chemos-
is, conjunctival injection, and proptosis. Diplopia may be caused by CN III, IV, or VI palsy, and loss of vision may be due to increased intraocular pressure due to orbital ve-
nous congestion with resultant glaucoma, venous retinop-
thly, and ischemic optic neuropathy. Cavernous-carotid fistulas associated with retrograde cortical venous flow often present with neurological symptoms referable to the area of venous congestion or infarction. This may occur in the supratentorial or infratentorial compartments.
Radiographic Diagnosis

While cerebral angiography is the gold-standard imaging modality in the diagnosis of CCFs, patients typically undergo noninvasive cerebral imaging with CT scanning, MRI, or CT/MR angiography first. Evidence of cavernous sinus enlargement, proptosis, extraocular muscle enlargement, superior ophthalmic vein dilation, or dilation of cortical or leptomeningeal vessels, as well as associated skull fractures, may be seen on CT or MRI and are suggestive of CCF.1,10 However, the absence of abnormalities on noninvasive imaging studies does not exclude the diagnosis of CCF.

If there is a high degree of clinical suspicion and/or imaging studies are consistent with the presence of CCF, the patient should be referred for diagnostic catheter cerebral angiography. Typically, this will be performed using transfemoral arterial catheterization with imaging of the bilateral CCAs, ICAs, and ECAs, as well as the vertebral arteries.11,48 Ipsilateral carotid artery compression during vertebral artery injection, known as the Huber maneuver, may be helpful to demonstrate shunting to the distal aspect of the fistula from the posterior circulation vessels. Similarly, ipsilateral common carotid artery compression during ICA injection, known as the Mehringer maneuver, may help visualize very–high flow CCFs by limiting the concurrent amount of unopacified blood shunting through the fistula.

Treatment

The goal of CCF treatment is to completely occlude the fistula while preserving the normal flow of blood through the ICA. Historically, ligation of the CCA was the surgical intervention of choice for the treatment of patients with CCFs. Benjamin Travers28 is credited with performing the first successful surgical CCA ligation in 1809 for a patient with pulsatile exophthalmos, attributed to an “aneurism by anastomosis.” A century later, in 1908, DeSchweinitz and Holloway13 reported the results of 114 patients treated with CCA ligation for pulsatile exophthalmos and found a 56% success rate and an 11.7% mortality rate. In 1934, Dorrance and Loudenslager14 reported the results of 151 patients treated with either CCA ligation (82 patients) or cervical ICA ligation (69 patients) and found that 52% of the patients had a successful outcome, 12.6% of the patients developed hemiplegia, and 5.3% died. The high morbidity and mortality rates seen with these procedures clearly necessitated further treatment refinements.

Hamby21 and Hamby and Gardner22 were the first to propose the technique of intracranial ICA ligation, followed by insertion of a muscle embolus in the cervical ICA as a way to close the fistula. Brooks is credited as being the first to use intravascular flow-directed embolization alone, without carotid artery ligation, to treat a CCF in 1931.52 Although Brooks reported a good result, the patient lost vision in the ipsilateral eye. In 1937, Browder7 was perhaps the first to perform direct surgery on the cavernous sinus, packing it with muscle to treat a cavernous carotid aneurysm. In the 1970s, Fedor Serbinenko paved the way for the modern era of CCF treatment with the development of balloon catheters that could be navigated endovascularly to the fistula site.25

Conservative Management

Conservative management, consisting of external manual compression of the ipsilateral cervical carotid artery several times a day for 4–6 weeks, may be effective in the treatment of indirect, low-flow CCFs.27,33 However, this is ineffective in the treatment of direct, high-flow fistulas. Higashida et al.27 reported complete CCF occlusion without clinical or angiographic evidence of recurrence at the 1-year follow-up in 30% of patients with indirect CCFs but in only 17% of patients with direct CCFs. Similar results for manual compression in the treatment of indirect CCFs have been reported by other groups.20,34 Interestingly, it has been reported that 20%–60% of patients with indirect CCFs have spontaneous fistula closure.10,17,18,62

When conservative management is used, it is important to have close ophthalmological follow-up, with serial vision tests, intraocular pressure measurements, and funduscopic examinations.49 Progressive visual decline, papilledema, and refractory elevation of intraocular pressure are all indications for emergency endovascular intervention. Additionally, cortical venous drainage on diagnostic angiography, neurological symptoms, or intractable headache or eye pain are indications of increased risk of intracranial hemorrhage, and should prompt urgent endovascular intervention. Patients presenting with a failure of CCF closure after compression therapy should be considered for alternative treatment strategies.12,30 It is worth noting that in our experience, conservative management of CCFs has generally been ineffective.

Endovascular Intervention

Transarterial or transvenous embolization is the first-line treatment modality for the treatment of most CCFs (Table 2).19,61,71,83 Metallic coils and/or liquid embolic agents are now most commonly used for this purpose after the withdrawal of detachable balloons from the US market in 2003.56,83 Transarterial access is often used when the CCF originates from branches of the ECA, as well as in select cases of direct fistulas (Fig. 3). When the CCF originates from branches of the ICA, transarterial embolization is significantly more difficult and carries an increased risk of stroke due to embolic reflux into the ICA. In these cases, a transvenous approach is used, and the fistula is occluded using either a coil or liquid embolization of the cavernous sinus.36,37,48,49

Transvenous access to the cavernous sinus can be achieved using standard transfemoral vein cannulation techniques with navigation through the inferior petrosal sinus or the facial and superior ophthalmic vein in most cases. Even an occluded inferior petrosal sinus can often be catheterized with careful microcatheter and microguide manipulation.38 The superior petrosal sinus, the basilic plexus, and the pterygoid plexus are alternative transvenous access routes if the inferior petrosal sinus or superior ophthalmic vein cannot be accessed. In rare circumstances, direct cannulation of the superior ophthalm-
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<td>Wang et al., 2011</td>
<td>54 posttraumatic direct CCFs in 51 pts</td>
<td>pulsating exophthalmos (82%), chemosis (78%), retro-orbital bruit (71%), ophthalmoplegia (63%), decreased visual acuity (57%), ICH (20%), incidental SAH (10%), spinal fracture (6%)</td>
<td>all pts treated endovascularly via transarterial approach: 40 w/ DBs alone, 8 w/ DBs &amp; stents, 2 w/ stents alone, 2 w/ DBs &amp; coils, 1 w/ coils alone, 1 w/ DBs, stents, &amp; coils</td>
<td>overall, 98% cure w/ ICA preservation, 85% cure w/ DBs alone</td>
<td>22% required 2nd &amp; 3rd endovascular interventions; 18% of pts treated w/ DBs alone developed pseudoaneurysms, 2 of which were symptomatic &amp; 3 progressively enlarging (these 5 were subsequently treated w/ stents); 5 pts had persistent moderate decreased vision or CN deficits; no procedural complications or delayed neurological/vascular complications</td>
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<td>Yoshida et al., 2010</td>
<td>44 indirect CCFs</td>
<td>chemosis (100%), ophthalmoplegia (36%), proptosis (14%), decreased visual acuity (11%)</td>
<td>initial Tx w/ transvenous endovascular embol using detachable coils, 4 pts required add’l transarterial embol, 2 pts required add’l radiosurgery</td>
<td>82% cure, 14% minor residual shunt, 4% significant residual shunt</td>
<td>9% recurrence, 7% permanent morbidity (2 brainstem infarcts, 1 ICH), 14% transient CN palsy</td>
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<td>Gupta et al., 2006</td>
<td>91 direct CCFs in 89 pts (85 traumatic, 6 ruptured aneurysms)</td>
<td>chemosis (100%), proptosis (98%), HA (84%), tinnitus (72%), neurological deficits including CN deficit (44%), visual deficit (26%), seizure (6%), abnormalities of mentation (6%), IC bleed (5%)</td>
<td>79 treated endovascularly w/ transarterial DBs, 12 treated w/ transarterial coil occlusion, as a balloon could not be negotiated through the fistula</td>
<td>of the 79 CCFs treated w/ DBs, 91% cure; of the 12 treated w/ coils, 75% cure; overall at 1 mo, 89% cure, 10% significant improvement, 1% remained static</td>
<td>minimal complications in 3.4%, w/ 1 pt developing multiple CN palsies, w/ partial recovery w/ steroid Tx; 1 pt had asymptomatic ICA occlusion; of pts who did not achieve complete CCF occlusion, 9 had persistent minor HA &amp; diplopia on extreme lateral gaze, &amp; 2 had persistent mild proptosis &amp; diplopia; of the 23 pts w/ visual deficits at presentation, 4 had fixed deficits</td>
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<td>Kirsch et al., 2006</td>
<td>141 indirect CCFs</td>
<td>chemosis (94%), increased IOP (60%), exophthalmos (87%), CN palsy (54%), diplopia (51%), decreased vision (28%), bruit (19%), ptosis (13%)</td>
<td>initial Tx w/ transvenous coil placement, 23% required add’l partial arterial embol</td>
<td>81% cure, 13% minor residual shunt, 4% significant residual shunt, 2% Tx failed</td>
<td>14% required &gt;1 transvenous procedure to achieve cure; 11% of pts had residual CN palsy or diplopia</td>
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<td>Luo et al., 2006</td>
<td>176 traumatic direct CCFs</td>
<td>chemosis (89%), bruit (89%), decreased visual acuity (78%), proptosis (67%), impairment of CN function (33%), epistaxis (6%)†</td>
<td>initial Tx w/ endovascular transarterial embol w/ a DB or a coil, 22 required add’l transarterial balloon-assisted NBCA embol</td>
<td>80% cure w/ ICA preservation following initial procedure, 82% of pts who underwent transarterial balloon-assisted NBCA embol achieved CCF occlusion w/ ICA preservation on immediate postembol angiography</td>
<td>of the 18 pts w/ successful occlusion following transarterial balloon-assisted NBCA embol, 1 died of SAH on Day 2 postembol; 1 had a recurrent fistula on Day 2 postembol &amp; presented w/ massive epistaxis requiring balloon occlusion of ICA &amp; fistula; 4 had asymptomatic migration of the NBCA mixture distally to the superior ophthalmic vein (n=3) or the inferior petrous sinus (n=1); 5 had asymptomatic small false sac or pseudoaneurysms in the cavernous ICA</td>
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<td>Meyers et al., 2002</td>
<td>135 indirect CCFs (93%), chemosis (87%), proptosis (81%), diplopia w/ ophthalmoplegia (68%), cranial bruit (49%), retroorbital HA (34%), elevated IOP (34%), diminished visual acuity (31%)</td>
<td>arterialization of conjunctival veins; embol achieved using a combination of metallic coils (87%), silk suture fragments (13%), or liquid adhesive (3%)</td>
<td>133 treated endovascularly, 76% via a transvenous approach; 90% cure, 97% good clinical recovery, 1% moderate disability, 2% severe disability</td>
<td>29% of pts required ≥2 endovascular procedures to achieve cure; 2.3% had procedure-related permanent morbidity; 6% of pts had symptomatic complications (1 cerebral infarction, 2 pts w/ decreased visual acuity, 1 pt w/ DI, 1 w/ orbital ecchymosis, 1 w/ retroperitoneal hematoma, 2 w/ deep femoral vein thrombosis w/o PE)</td>
<td>9% of pts required ≥2 endovascular procedures to achieve cure; 2.3% had procedure-related permanent morbidity; 6% of pts had symptomatic complications (1 cerebral infarction, 2 pts w/ decreased visual acuity, 1 pt w/ DI, 1 w/ orbital ecchymosis, 1 w/ retroperitoneal hematoma, 2 w/ deep femoral vein thrombosis w/o PE)</td>
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<td>Tu et al., 1997</td>
<td>78 CCFs (66 direct, 12 indirect)</td>
<td>pulsating exophthalmos, chemosis, ophthalmoplegia, decreased visual activity</td>
<td>initial Tx w/ endovascular transarterial DBs; 5 pts required subsequent transvenous intervention; 18 pts in whom endovascular Tx failed &amp; 1 who presented w/ acute bleeding underwent direct surgical repair (9 treated w/ sinus packing, 2 treated by seal w/ fascia &amp; glue, 2 w/ IC-IC bypass, 1 w/ clipping, w/ suture, 4 w/ multiple techniques)</td>
<td>of the 77 pts who had endovascular Tx, 77% cure; of the 19 pts who underwent direct surgical repair, 100% cure, w/ 73% ICA patency; overall, 100% cure w/ 94% ICA patency</td>
<td>of the 19 pts who underwent direct surgical repair, 42% experienced transient CN III palsy &amp; 1 experienced postop CN VI palsy w/ only partial recovery; 1 pt had a wound infection requiring prolonged antibiotic administration</td>
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<td>Lewis et al., 1995</td>
<td>100 direct CCFs in 98 pts (76 traumatic, 22 ruptured aneurysms, 2 iatrogenic)</td>
<td>orbital bruit (80%), proptosis (72%), chemosis (55%), CN VI palsy (49%), conjunctival injection (44%)</td>
<td>initial Tx w/ transarterial embol w/ DBs</td>
<td>88% cure, w/ 75% ICA preservation; 4 pts had eventual spontaneous closure following unsuccessful balloon occlusion; 5 pts required direct op to achieve CCF occlusion; 2 pts had CCF closure w/ non-DBs</td>
<td>1 pt died of trauma-related injuries; 4% of pts experienced permanent neurological complications (1 pt w/ cerebral infarction, 1 w/ ICH, 1 w/ vision loss); 1 death related to balloon shift causing cerebral infarction; 3 pts experienced transient ischemia</td>
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<td>Higashida et al., 1989</td>
<td>213 posttraumatic CCFs (206 direct, 7 indirect)</td>
<td>pulsating exophthalmos, chemosis, retroorbital bruit, ophthalmoplegia, decreased visual activity</td>
<td>Direct CCFs initially treated endovascularly w/ transarterial DBs; addl endovascular interventions included use of liquid tissue adhesives, microcoils, &amp; silk sutures; 3% required transvenous approaches. Indirect CCFs treated w/ embol using healon particulate emboll &amp;/or liquid tissue adhesives; half required addl transvenous intervention</td>
<td>of the 206 direct CCFs, 88% cure w/ ICA preservation; of the 7 indirect CCFs, 100% cure</td>
<td>6 pts experienced transient cerebral ischemia, 5 had pseudoaneurysm formation, 5 experienced strokes, 1 had a peripheral nerve injury after Tx</td>
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mic vein after surgical exposure may be necessary to gain access to the cavernous sinus.86

Endovascular placement of covered or flow-diverting stents is being investigated as an alternative to embolization for the treatment of CCFs. Flow-diverting stents are deployed within the cavernous ICA, over the site of the fistula, redirecting blood flow away from the cavernous sinus.76,84 While these devices demonstrate dramatic results initially, delayed thrombosis remains a problem with the use of currently available covered stents in the cerebral circulation. The use of flow diverters alone or in combination with other embolic materials has been used anecdotally.

Reported complications of endovascular treatment include cerebral infarction, decreased visual acuity, diabetes insipidus, retroperitoneal hematoma, femoral vein thrombosis, and ophthalmoplegia in 2%–5% of patients.16,48 Other complications include subarachnoid or intracerebral hemorrhage, sinus rupture, extradural extravasation of contrast, and CN palsies. It is notable that up to 42% of patients will experience a transient worsening of symptoms after cavernous sinus embolization that generally resolves over time.68 Resolution of preexisting symptoms is related to their duration and severity.

More than 80% of patients who undergo endovascular treatment for direct and indirect CCFs will experience a complete cure.3,17,42,52,88 This is demonstrated both clinically by reversal of the signs and symptoms of the CCF and angiographically by fistula obliteration with reversal of retrograde cortical venous flow and vascular steal. In a
series of more than 200 patients presenting with posttraumatic direct CCFs, Higashida et al.\textsuperscript{26} reported complete fistula occlusion with preservation of the ICA in 88% of patients. Similarly, in a series of 135 patients presenting with indirect CCFs, Meyers et al.\textsuperscript{48} reported complete fistula occlusion in 90% of patients.

**Surgical Intervention**

In cases in which endovascular treatment is not possible or is unsuccessful, open surgical intervention may be warranted (Table 2). Surgical intervention may involve suturing, clipping, or trapping the fistula, packing the cavernous sinus to occlude the fistula, sealing the fistula with fascia and glue, ligating the ICA, or a combination of these procedures. Isamat and colleagues\textsuperscript{31–33} are proponents of direct intracavernous obliteration by muscle packing combined with transmural injection of fibrin glue, citing good results with less complexity, less time, and a higher probability of preserving the patency of the ICA compared with other surgical procedures.\textsuperscript{81}

Overall success rates using surgical intervention in the treatment of CCFs have been reported at between 31% and 79%. Some practitioners have reported 100% obliteration rates in small series.\textsuperscript{9,17,66,79} Tu et al.\textsuperscript{79} reported a series of 19 patients in whom direct surgical intervention was used after endovascular treatment failed. Closure of the CCF was achieved in all 19 patients, with a 73% ICA patency rate. Day and Fukushima\textsuperscript{9} reported similar success using direct surgical intervention after endovascular failure in the treatment of 9 patients with Type D CCFs. In this series, there was a 100% cure rate with no deaths. Transient diplopia and trigeminal hypesthesia were noted in all 9 patients, 1 patient suffered from temporary hemiparesis, and another patient had a permanent hemiparesis.

**Radiosurgical Intervention**

First performed by Barcia-Salorio and colleagues\textsuperscript{2} in 1977, radiosurgical intervention may play a role in the treatment of patients with indirect, low-flow CCFs (Table 3). Barcia-Salorio et al.\textsuperscript{1} reported complete or near-complete reversal of the neuroophthalmological symptoms associated with indirect CCFs in 91.6% of patients with spontaneous, low-flow lesions treated with radiosurgery. However, in patients presenting with previously treated direct CCFs, radiosurgery was effective in only 2 of 5 patients. Other small series of patients with indirect CCFs have shown long-term obliteration of the fistula after treatment with radiosurgery in 75%–100% of patients.\textsuperscript{3,17,26,53,64} Radiosurgery should not be used in emergency cases, as there is a latency of several months to years before complete obliteration of the CCF is accomplished.\textsuperscript{3}

**Prognosis**

After successful intervention with complete closure of a CCF, symptoms such as chemosis and proptosis generally resolve within hours to days.\textsuperscript{36,48} Cranial nerve palsy typically resolve over the course of several weeks.\textsuperscript{47,83} The degree of vision recovery, if vision loss was experienced prior to intervention, is largely dependent on the

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<td>Pollock et al., 1999</td>
<td>20 indirect CCFs</td>
<td>chemosis/proptosis (90%), decreased vision (40%), HA/eye pain (30%)</td>
<td>7 pts treated w/ radiosurgery alone; 13 pts treated w/ radiosurgery &amp; transarterial embol</td>
<td>95% symptomatic improvement; of 15 pts who underwent FU angiography, 13 showed complete obliteration of the fistula, 1 pt showed near-total obliteration of the fistula</td>
<td>2 pts developed new neuroophthalmological symptoms related to radiosurgery</td>
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<td>Hirai et al., 1998</td>
<td>26 indirect CCFs</td>
<td>diplopia (65%), decreased vision (40%), HA/eye pain (30%)</td>
<td>12 pts treated w/ radiation alone; 14 pts treated initially w/ endovascular embolization, w/ subsequent radiation in 6 w/ endovascular failure</td>
<td>of the 12 pts treated w/ radiation alone, 75% complete cure &amp; 8% w/ improvement, w/ a mean FU of 62 mos; of the 6 pts treated w/ radiation after endovascular embolization, 67% (4/6) complete cure, w/ an additional 1 pt having improvement w/ a mean FU of 24 mos</td>
<td>of the 12 pts treated w/ radiation alone, 75% complete cure &amp; 8% w/ improvement, w/ a mean FU of 62 mos; of the 6 pts treated w/ radiation after endovascular embolization, 67% (4/6) complete cure, w/ an additional 1 pt having improvement w/ a mean FU of 24 mos</td>
</tr>
<tr>
<td>Barcia-Salorio et al., 1994</td>
<td>25 CCFs (22 indirect, 3 previously treated direct)</td>
<td></td>
<td></td>
<td>20.22 (91.4%) low-flow CCFs completely obliterated, 1/3 (33.3%) previously treated direct CCFs aquired</td>
<td></td>
</tr>
</tbody>
</table>

* FU = follow-up.
pathogenesis, severity, and duration of the preintervention deficit. Recurrence of CCFs due to recanalization postembolization is uncommon but can typically be treated by repeat embolization.

**Conclusions**

Catheter cerebral angiography is the gold standard imaging modality used in the diagnosis and classification of CCFs. Although historically difficult to treat, these lesions are now routinely managed with low rates of morbidity and mortality. Endovascular intervention with a goal of complete fistula occlusion while preserving normal blood flow through the internal carotid artery has emerged as the treatment of choice. In select cases, open surgery, radiosurgery, or conservative management are also treatment options. Symptom resolution with low rates of recurrence can be expected in most cases after appropriate therapy.

**Disclosure**

The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

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