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Closed-cell stent for coil embolization of intracranial aneurysms: clinical and angiographic results

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Despite ongoing concerns about the durability of endovascular treatment of intracranial aneurysms, coiling has been increasingly accepted worldwide. The multicenter randomized ISAT, showing long-term improved safety and clinical outcome in patients treated with coil embolization compared with open clipping, has accelerated this trend, increasing the number of patients being referred for endovascular treatment and thus further emphasizing the need to enhance the ability to treat intracranial aneurysms effectively. However, large, giant, and wide-neck aneurysms can be difficult to treat because of the significant risk of coil herniation from the aneurysm into the parent artery. While the goal of endovascular treatment is complete exclusion of the aneurysm from the circulation while preserving the parent and side-branch artery lumen, this is often not possible. Even with adjunctive techniques such as balloon remodeling, large, giant, and wide-neck aneurysms remain challenging, with a significant number of subtotal occlusions. In addition, aneurysm recanalization is observed in 20%–40% of cases due to coil compaction, migration of coils into the aneurysm thrombus, or aneurysm growth.

Introduction of stents designed as an adjunct tool for coiling has been shown, in the short-term, to be of beneficial value. However, a variety of devices, different in their design and coating, was used with a high periprocedural mortality and morbidity. This study was designed to evaluate 1 specific stent available for aneurysm treatment, regarding the periprocedural and safety profile as well as efficacy. Our hypothesis was that stent-assisted coil embolization reduces recanalization and retreatment rates compared with historical controls without impacting procedural safety in a subgroup of wide-neck aneurysms or in those with a dome/neck ratio of <2.

Materials and Methods

Patients and Techniques

From June 2007 to June 2010, consecutive patients harboring intracranial aneurysms treated with Enterprise (Codman Neurovascular, Raynham, Massachusetts) SACE were enrolled in a prospective data base at 2 participating centers. All procedures were approved by the local institutional review board at each participating site. Inclusion criteria were the following: 1) wide-neck aneurysm: defined as one having a neck dimension that is >4 mm or a dome/neck ratio that is ≤2; 2) parent vessel diameter of ≤4 mm; 3) any use of a closed-cell stent-assisted coil embolization.
stent for coil embolization; and 4) patient or health care proxy informed consent. Patients younger than 18 years and pregnant women were excluded. Patients treated with open-cell stents were excluded. At the participating centers, all ruptured aneurysms are treated, and treatment of unruptured aneurysms is performed after careful assessment of perceived risk factors for rupture and consultation with our multidisciplinary cerebrovascular team and patients.

**Angiographic and Endovascular Procedures**

All endovascular procedures were performed by senior neurointerventionalists by using a biplane angiography unit with 3D rotational angiography capability (Allura Xper FD20/20; Philips Healthcare, Best, the Netherlands) with patients under general anesthesia. Patients were given ASA (80 mg) and clopidogrel (75 mg) for a minimum of 3 days before the procedure. Dual antiplatelet treatment was continued for a minimum of 6 months, followed by life-long continuation of ASA. In cases of acute aneurysm rupture, patients were administered 350 mg of ASA and 150 mg of clopidogrel through a nasogastric tube at a minimum of 1 hour before commencing the procedure. Most important, in ruptured cases, the neurosurgery and neurointensive care teams were consulted to assess the need for ventriculostomy before administering antiplatelet medication. Endovascular access was obtained by a standard transfemoral approach. Following access, patients were heparinized to maintain activated clotting time >250 seconds (for ruptured and unruptured aneurysms). The stent delivery microcatheter was first positioned across the neck of the aneurysm followed by placement of the coiling microcatheter within the aneurysm. All aneurysms were embolized by using the semi-jailing or jailing technique. Aneurysms were coiled as densely as possible with coils selected per the operator’s preference. Any technical complication, with or without clinical sequelae, was recorded.

Aneurysm occlusion was estimated by independent reviewers not involved in patient care, by using a 3-point RS (RS 1, complete obliteration of aneurysm and neck; RS 2, neck remnant without contrast filling the aneurysm sac; and RS 3, contrast filling the aneurysm sac) immediately after SACE and at follow-up. After the procedure, patients were transferred to the neurosurgical intensive care unit. Packing attenuation, the volume ratio of implanted coils to the volume of the aneurysm, was calculated as previously described. We calculated coil volume for each aneurysm, assuming that the coil was a solid cylinder by using the primary coil diameter given in the manufacturer’s specifications.

**Clinical Evaluation**

A complete neurologic examination was performed in all patients at baseline, immediately after the procedure, at discharge, and at follow-up by experienced physicians certified in stroke assessment. An mRS score was assessed at baseline for unruptured cases, at discharge, and at follow-up evaluations. For ruptured aneurysms, the Hunt and Hess grade was recorded at baseline. Primary adverse events included death and stroke. Secondary adverse events recorded were transient ischemic attack, the need for re-intervention, and the presence of hematomas. Residual aneurysm size was determined by angiography. Medical histories, procedural reports, and clinical outcomes were recorded in this prospective data base.

**Statistical Analysis**

All data are presented as the mean ± the standard error of the mean. Comparisons of categoric data were performed by using the Fisher exact test in GraphPad InStat (http://www.softpedia.com/get/Others/Finance-Business/GraphPad-InStat.shtml). Significant differences were established for \( P < .05 \).

**Results**

**Patient Information and Technical Results**

We enrolled 147 patients harboring 161 aneurysms that were embolized via SACE. Eighteen aneurysms (11%) were treated acutely following rupture. Patient demographic information and aneurysm characteristics/location are provided in Table 1. Most aneurysms were discovered incidentally (55%) and were of wide-neck saccular morphology (60%). The mean aneurysm and neck diameters were 6.5 ± 0.4 and 5.1 ± 0.3 mm, respectively; the dome/neck ratio was 1.3 ± 0.04. In 2 cases of a small aneurysm and a dissecting aneurysm, no coils were used in conjunction with the stent. In another 2 cases, the stent was reinserted following successful coil embolization by using the semi-jailing technique. In total, 167 stents were used (1.03 stents/aneurysm; 6 patients received 2 stents). The packing attenuation obtained was 43 ± 3\%. Of all stented vessels, 19 measured <2 mm (mean, 1.7 ± 0.04 mm).

During the procedure, clot formation on the surface of the...
Table 2: Neurologic complications and mortality (unruptured aneurysms)

<table>
<thead>
<tr>
<th>Change in mRS*</th>
<th>Frequency (No. of Patients)</th>
<th>Description (No.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>7</td>
<td>New or more severe headache (4), mild extremity weakness (3), visual field deficit (1)</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>Stroke after termination of antplatelet therapy 60 days posttreatment (1)</td>
</tr>
<tr>
<td>3</td>
<td>2</td>
<td>Anterior choroidal artery stroke (1); stroke after termination of antplatelet therapy 10 months post-treatment (1)</td>
</tr>
<tr>
<td>B</td>
<td>2</td>
<td>Sneddon syndrome, withdrawal of medical care after thromboembolism (1); stroke 6 months after treatment (1)</td>
</tr>
</tbody>
</table>

* Increase in mRS from baseline (mRS_{follow-up} − mRS_{baseline})

Table 3: Neurologic outcomes and mortality (ruptured aneurysms)

<table>
<thead>
<tr>
<th>mRS of Patients</th>
<th>Frequency (No. of Patients)</th>
<th>Description (No.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>8</td>
<td>Residual deficit after SAH (2); stroke due to stent occlusion after stopping antplatelet therapy for shunt placement 2 months after procedure (1)</td>
</tr>
<tr>
<td>2</td>
<td>3</td>
<td>Symptomatic intracranial hemorrhage following shunt revision while on antplatelet therapy (1)</td>
</tr>
<tr>
<td>3</td>
<td>1</td>
<td>Aneurysm rebleeding (2), complications of SAH/vasospasm (3)</td>
</tr>
</tbody>
</table>

* One patient lost to follow-up.

Aneurysm rebleeding (2), complications of SAH/vasospasm (3), p < 0.001)

Clinical Outcome

Seven patients (4.8%) died, including 2 with reruptures (Tables 2 and 3). There were 5 (3.4%) and 3 (2%) minor and major strokes, respectively, within 30 days of the procedure. During this period, 6 patients (4.1%) experienced a transient ischemic attack. The 30-day combined cardiac and pulmonary complication rate was 2.5%. Five patients (3.4%) had access site complications, including groin hematoma (4 patients) and suture-induced femoral artery constriction requiring surgical repair (1 patient). One patient developed hydrocephalus 30 days after SACE of a ruptured aneurysm.

Of the 140 surviving patients, 113 (80.7%) with 120 aneurysms were available for follow-up neurologic examination at a mean of 11.8 months (95% CI, 10.3–13.2 months). Of patients in the unruptured aneurysm cohort available for follow-up (n = 101), an increase in the mRS score from admission to follow-up by 1, 2, or 3 points was seen in 7 (6.9%), 1 (1%), and 2 (2%) patients, respectively (Fig 1). After adjudication of complications by the senior neurointerventionalists, procedure-related mortality and permanent neurologic deficits occurred in 2 (1.4%) and 5 patients (3.4%), respectively (Tables 2 and 3).

Angiographic Results

Follow-up angiography was performed in 120 aneurysms at a mean of 11.9 months (95% CI, 10.6–13.2 months) (illustrative case, Fig 2). Of the angiographic follow-ups, the number of aneurysms at each time point was the following: 3 at <6 months, 48 from 6 to 11 months, 62 from 12 to 24 months, and 7 at >24 months. Aneurysms were more likely to be completely occluded at follow-up, suggesting progressive thrombosis (P < .0001) (Fig 3). Recanalization, defined as any increase in the RS from the postprocedural angiogram, was seen in 12 aneurysms (10%). Of the recanalized aneurysms, 7 (5.8%) were subsequently retreated. No adverse events were seen as a result of the retreatment. Intimal hyperplasia producing a mild (<20%) or moderate (>20%; <70%) narrowing was seen in 6 (5%) and 1 (0.8%) of the stented segments, respectively. All patients with angiographic evidence of vessel narrowing remained asymptomatic. Of the stented segments in vessels that measured <2 mm, there were 2 (10.5%) mild and 1 (5.3%) moderate case of in-stent stenosis.

Discussion

Technologies such as balloon remodeling24,25 and stent-assisted coiling11,13,26,27 have enabled endovascular treatment of wide-neck complex aneurysms. A large multicenter prospective trial recently compared the safety profile of the balloon remodeling technique with that of coil embolization alone in unruptured aneurysms.28 The morbidity and mortality in the balloon remodeling group were 2.3% and 1.4%, respectively, and rates did not differ from those in the standard treatment group. A recent retrospective study of SACE in both ruptured and unruptured aneurysms reported procedure-related morbidity and mortality rates of 7.4% and 4.6%, respectively.15 This latter study included
all commercially available stent technologies, and the authors reported that procedure-related complications were dependent on the stent used. To remove this variable, a multicenter registry was performed to evaluate the acute clinical outcomes by using the Enterprise stent, and it found permanent morbidity and mortality rates of 2.8% and 2%, respectively.

We report the midterm clinical outcome for aneurysms treated by closed-cell SACE. In unruptured aneurysms, overall morbidity, defined as any increase in mRS at follow-up, and mortality were 10.3% and 1.6%, respectively. Permanent major morbidity (mRS score of ≥2 at follow-up) in the unruptured aneurysm cohort was seen in 4 patients (3.1%). If one uses similar criteria for comparison of SACE in this series with balloon-remodeling previously reported, mortality is similar and morbidity is slightly higher. Although it is convenient to compare these data with the balloon-remodeling experience, our centers frequently use the balloon-remodeling technique, and SACE is reserved for aneurysms with an exceedingly poor dome/neck ratio that approaches 1. Additionally, in this series, poor outcomes were mostly related to thromboembolic complications associated with premature termination of systemic dual-antiplatelet therapy rather than the SACE procedure.
We enrolled 18 patients in the acute period following subarachnoid hemorrhage, and the morbidity and mortality rates were 22% and 28%, respectively. These outcomes are similar to those reported for SACE in ruptured aneurysms. However, this complication rate does not compare favorably with that in the ISAT, in which 23.7% of patients were dependent or dead 1 year following endovascular coiling. One explanation for the lower rate of good outcomes in our study might be the inclusion of dissecting aneurysms (n = 6, 33%), which were excluded in the ISAT trial. Two significant complications occurred in our study when patients developed hydrocephalus 1–2 months following SACE of ruptured aneurysms. In 1 case, a patient underwent shunt revision 2 months following SACE of a ruptured aneurysm in the anterior circulation. The patient was treated with the patient on antiplatelet therapy, leading to symptomatic intracranial hemorrhage. Additionally, 2 deaths were associated with aneurysm rebleeding after SACE. One case of rebleeding was from a dissecting aneurysm, where the rebleed occurred 4 days after embolization. These data demonstrate the increased risk of using stents for aneurysm embolization versus coil alone or balloon-remodeling in ruptured aneurysms. Recently, a large meta-analysis reported aneurysm recurrence and retreatment following coil embolization in 20.8% and 10.3% of cases, respectively. Angiographic results reported herein demonstrate a high rate of progressive thrombosis of the aneurysm and complete aneurysm obliteration on follow-up. Aneurysm recurrence (10%) and retreatment (5.8%) were considerably lower with SACE despite the wide-neck morphology, which is associated with worse angiographic outcomes. Intimal hyperplasia induced by the presence of the stent within the parent artery leads to remodeling, thereby permanently excluding the aneurysm.

Limitations of this study are the lack of a direct control group and the relatively small number of ruptured aneurysms included in the study. Follow-up at a mean of 11.8 months was available in only 80% of patients. Patients were lost to follow-up either due to refusal to return for clinical examination or angiography or change of contact information with no forwarding instructions. Because our study enrolled prospectively all consecutive patients receiving SACE with a closed-cell device, our study included heterogeneous aneurysm morphologies including blister and dissecting aneurysms as well as both ruptured and unruptured aneurysms. We further included 2 cases in which the device was resheathed and not implanted, as well as 2 cases in which coiling was not possible.

Conclusions
SACE of wide-neck unruptured aneurysms with a closed-cell device is safe and produces durable and high occlusion rates; however, increased morbidity and mortality observed in ruptured aneurysms are associated with dual-antiplatelet treatment required for stent placement.

References
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