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BACKGROUND AND PURPOSE: Dissection of the carotid artery can, in certain cases, lead to significant stenosis, occlusion, or pseudoaneurysm formation, with subsequent hemodynamic and embolic infarcts, despite anticoagulant therapy. We sought to determine the therapeutic value of stent-supported angioplasty retrospectively in this subset of patients who are poor candidates for medical therapy.

METHODS: Five men and five women (age range, 37–83 years; mean age, 51.2 years) with dissection of the internal (n=9) and common (n=1) carotid artery were successfully treated with percutaneous endovascular balloon angioplasty and stent placement. The etiology was spontaneous in five, iatrogenic in three, and traumatic in two. Seven of the treated lesions were left-sided and three were right-sided.

RESULTS: The treatment significantly improved dissection-related stenosis from 74±5.5% to 5.5±2.8%. Two occlusive dissections were successfully recanalized using microcatheter techniques during the acute phase. Multiple overlapping stents were needed in four patients to eliminate the inflow zone and false lumen and establish an angiographically smooth outline within the true lumen. There was one case of retroperitoneal hemorrhage, but there were no procedural transient ischemic attacks (TIAs), minor or major strokes, or deaths (0%). Clinical outcome at latest follow-up (16.5±1.9 months) showed significant improvements compared with pretreatment modified Rankin score (0.7±0.3 vs 1.8±0.44) and Barthel index (99.5±0.5 vs 80.5±8.9). One delayed stroke occurred in a treated patient with contralateral carotid occlusion following a hypotensive uterine hemorrhage at 8 months; the remaining nine patients have remained free of TIA or stroke.

CONCLUSION: In select cases of carotid dissection associated with critical hemodynamic insufficiency or thromboembolic events that occur despite medical therapy, endovascular stent placement appears to be a safe and effective method of restoring vessel lumen integrity, with good clinical outcome.

Dissection of the cervical carotid arteries occurs when the integrity of the intima and media layers of the vessel wall is compromised, resulting in hemorrhage in the vessel wall and consequent compromise of the native vessel lumen (1–4) (Fig 1). The formation of a false channel and associated anatomic disruption of the endothelial monolayer results in conditions favoring local thrombus formation, which may then embolize distally, and result in brain infarction (5). The accepted medical standard of therapy has consisted of systemic anticoagulation, and has been observed to lead to a significant rate of subsequent resolution and healing (6, 7). In a subset of patients, spontaneous healing, with restoration of native vessel anatomy, fails to occur (1, 8–10). Instead, the arterial dissection may result in total occlusion of the true vessel lumen by compression from the false lumen, created by entry of pressurized blood in the dissected medial defect (2, 9). In another subset of patients, the dissection site may develop into a focal or extensive stenosis, with formation of an associated pseudoaneurysm (1, 2, 8). In such cases, anticoagulation, although critical to prevent thromboembolic events in the initial stages of the disease, may be insufficient to overcome the flow-limiting nature of the dissecting lesion or may be contraindicated because of risk of pseudoaneurysm rupture (3). En-
Endovascular stents have been shown to be of benefit for the treatment of coronary atherosclerosis (11), and have been considered for the treatment of carotid atherosclerosis in certain high-risk patient subgroups (12). Stents also have been used to treat intimal dissection resulting from balloon angioplasty and trauma (13–16). By virtue of their design, stents provide the necessary centrifugal force to permit apposition of the dissected segment to the vessel wall to obliterate the false lumen and resolve the stenosis (13, 17). Stents can also provide mechanical support to serve as a scaffold for coil embolization of dissection-related wide-necked pseudoaneurysms (18, 19) (Fig 1). In order to determine the clinical and angiographic outcome of this form of therapy, we present our endovascular management of cervical carotid dissection, achieved using percutaneous stent angioplasty in a retrospective series of 10 patients.

Methods

Patient Selection Criteria

During the 18-month period between October 1997 and March 1999, 10 consecutive patients underwent endovascular treatment of carotid dissection performed using intraluminal stent placement. Informed consent was obtained from patients or medical guardians. Patients were considered eligible for stent therapy of carotid dissection only if they had either failed optimal medical therapy, were unable to undergo anticoagulation, or if they had no other therapeutic option of acceptably low risk. The specific indications (Table 1) included the presence of transient ischemic attacks (TIAs) despite anticoagulant or antilplatelet therapy (patients 4, 8, 10), contraindication to anticoagulation (patient 9), contralateral carotid occlusion or stenosis in a patient who was neurologically unstable or had clinical evidence of hemodynamic insufficiency (patients 3, 5, 5), documented poor collateral circulation (patient 1), need for elective occlusion of the contralateral internal carotid artery for aneurysm treatment (patient 2), and need to avoid flow increase through the anterior communicating artery because of the presence of an aneurysm (patient 7). Patients who underwent stent placement for angioplasty-induced intimal dissection during endovascular treatment of carotid atherosclerosis were excluded from the study.

Procedure

All patients underwent complete digital subtraction angiography of both anterior and posterior cerebral circulations, including the external carotid arteries. Patients were administered intravenous heparin to achieve an activated clotting time of greater than 250 seconds. A 7F or 9F guide catheter (Brite Tip; Cordis Endovascular, Miami Lakes, FL) was placed in the common carotid artery via a femoral vascular sheath (Avanti; Cordis). A 2.3F microcatheter (Transit; Cordis) was used coaxially over a 0.014- to 0.016-in microguidewire (Instinct 10; Cordis/Transend 14, Scimed, Maple Grove, MN) to enter the true arterial lumen under real-time high-resolution digital roadmap angiography (Toshiba, Tusinit, CA). A 300-cm-long exchange microguidewire (Stabilizer; Cordis) was then passed through the microcatheter and used to exchange the latter for the stent delivery catheter. Three types of stents were employed in this study: 1) a self-expanding stainless steel stent (Wallstent; Schneider, Plymouth, MN); 2) a premounted balloon-expandable stainless steel stent (GFX; Arterial Vascular Engineering, Santa Rosa, CA); and 3) a Nitinol shape-memory alloy self-expanding stent (S.M.A.R.T.; Cordis). After stent deployment, the patients were administered ticlopidine (250 mg po bid) or clopidogrel (75 mg po qd) for 6 weeks and aspirin (325 mg po qd) indefinitely. The vascular access sheath was removed on the day after the procedure, with the aid of an external femoral compression device (FemoStop; Radi, Uppsala, Sweden).

Outcome Analysis

Follow-up was performed using both neurologic examination and telephone interview. The modified Rankin score and Barthel index were used, as previously described (20–22). Evaluation of stent patency was obtained by sonography or repeat digital subtraction angiography.

Statistics

Severity of stenosis was computed using the NASCET method, with dimensions obtained either by using the relative fluorographic size of a reference object or the angiographic digital computer system (Toshiba). Specifically, the diameter of maximal stenosis (D_{Stenosis}) was measured along with the most proximal diameter of the distal normal vessel (D_{Normal}), and the degree of stenosis computed as Stenosis = \left\{1 - \left(\frac{D_{Stenosis}}{D_{Normal}}\right)\right\} \times 100. The paired Student’s t test and analysis of variance were used in the comparison of numerical variables. For categorical analysis, a Pearson’s chi-square test was used. Statistical significance was assumed for P<.05.

Results

Patient Population and Lesion Characteristics

There were five male and five female patients, ranging in age from 37 to 83 years (mean age, 51.2 ± 4.2 years) (Table 1). Seven (70%) of the 10 patients suffered from coexisting hypertension and seven patients (70%) had suffered a stroke prior to treatment as a result of the initial arterial dissection. The location of dissections treated was as follows: one in the left common carotid artery, six in the left internal carotid artery, and three in the right internal carotid artery. Two patients were treated within 12 hours of the initial angiographic or symptomatic dissection, three patients were treated between 3 and 10 days of symptomatic dissection, and the remaining five patients were treated for chronic dissection between 3 weeks and 10 years after initial detection of their dissection.

Therapeutic Interventions

The etiology of the carotid dissections encountered in this study was spontaneous, iatrogenic, or traumatic. Five patients (50%) had spontaneous dissections, one of which was chronic (discovered 10 years prior to admission), and three had iatrogenic dissections (30%) resulting during diagnostic angiography. Of these three patients, two underwent recanalization and stent angioplasty of a completely occluded internal carotid artery, with complete restoration of flow (patients 1, 7). The third (patient 2) underwent stent angioplasty of a narrowed and dissected left internal carotid artery in order to enable treatment of a contralateral giant cavernous carotid artery aneurysm that required...
balloon occlusion and carotid sacrifice. Two patients (20%) presented with traumatic dissection, one as a result of manual strangulation by her spouse (patient 6), and the other after vehicular whiplash (patient 10) (21) (Table 2). The angioplasty and stent treatment improved dissection-induced stenosis significantly from $74\pm5.5\%$ to $5.5\pm2.8\%$ ($P<.001$).

Two patients (patients 3 and 5) had bilateral involvement of the internal carotid artery. These patients suffered spontaneous dissection and had undergone diagnostic angiography at an outside hospital 2 to 3 days prior to admission, which showed unilateral flow-limiting dissection on one side and slight irregularity in the contralateral carotid artery. After transfer, repeat angiography revealed progression of the disease to unilateral complete occlusion of the previously dissected internal carotid artery and a severe flow-limiting lesion on the contralateral side. Eight lesions were left-sided and two were right-sided.

In four patients, multiple stents were deployed in an overlapping fashion in order to enable the elimination of the false lumen and to treat the residual stenosis. Two of these patients, with a severe form of bilateral dissection leading to unilateral occlusion at the time of treatment, required three tandem stents to reconstitute an angiographically smooth luminal outline (patients 3 and 5). In the other two patients, two tandem stents were used in both cases to allow spanning the entire length of the dissection, which included the carotid bulb with its greater diameter compared with the proximal common carotid (patient 4) or the cervical internal carotid artery (patient 9). In two patients (patients 8 and 10), a stent was deployed to serve as a scaffold through which a wide-necked pseudoaneurysm was subsequently treated using Guglielmi electrolytically detachable coils, in a manner similar to that reported on in previous reports (18, 19). These aneurysmal dilatations were located in the cervicopetrous junction (patient 8) and in the petrocavernous segment (patient 10) of the internal carotid artery.

Complications

There were no instances of postprocedural transient ischemic attacks (TIAs), new neurologic def-
<table>
<thead>
<tr>
<th>Patient (No.)</th>
<th>Age (yrs)/Sex</th>
<th>Presenting Symptoms</th>
<th>Duration between Symptoms or Dissection and Treatment</th>
<th>Infarct/Location</th>
<th>Etiology of Dissection</th>
<th>Indication for Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 45/F</td>
<td></td>
<td>L hemiparesis following diagnostic angiogram</td>
<td>12 hours</td>
<td>R posterior frontal infarct</td>
<td>Iatrogenic (angiography)</td>
<td>Restoration of flow, given poor collaterals</td>
</tr>
<tr>
<td>2 46/F</td>
<td></td>
<td>R ptosis, CN III palsy, diplopia from mass effect of cavernous ICA aneurysm</td>
<td>3 days</td>
<td>None</td>
<td>Iatrogenic (angiography)</td>
<td>Restoration of LICA flow, given need to sacrifice RICA</td>
</tr>
<tr>
<td>3 51/M</td>
<td></td>
<td>Sudden loss of vision in L eye 10 days prior to admission, headache, spontaneous bilateral ICA dissection</td>
<td>10 days</td>
<td>Bilateral internal capsule, L occipital lobe</td>
<td>Spontaneous</td>
<td>Restoration of LICA flow, given occluded RICA</td>
</tr>
<tr>
<td>4 83/M</td>
<td></td>
<td>L hemispheric TIAs, aphasia, R hemiparesis</td>
<td>3 weeks</td>
<td>None</td>
<td>Spontaneous</td>
<td>Obliteration of false channel to eliminate thrombus formation</td>
</tr>
<tr>
<td>5 51/M</td>
<td></td>
<td>Spontaneous dissection, aphasia, R hemiparesis</td>
<td>10 days</td>
<td>L perisylvian, insular and posterior frontal infarct</td>
<td>Spontaneous</td>
<td>Restoration of RICA flow to avoid occluded LICA</td>
</tr>
<tr>
<td>6 37/F</td>
<td></td>
<td>R hand and arm weakness, leg numbness, and dysphasia</td>
<td>3 months</td>
<td>Left posterior frontal infarct</td>
<td>Traumatic (strangulation)</td>
<td>Reduction of LICA stenosis given narrowed and hypoplastic RICA</td>
</tr>
<tr>
<td>7 40/F</td>
<td></td>
<td>Asymptomatic, immediate detection of dissection during follow-up angiogram for Acomm aneurysm coiling</td>
<td>30 minutes</td>
<td>Old lacune in posterior limb of R internal capsule</td>
<td>Iatrogenic (angiography)</td>
<td>Restoration of LICA flow to avoid high flow-mediated Acomm aneurysm expansion</td>
</tr>
<tr>
<td>8 64/M</td>
<td></td>
<td>R side hemiparesis and left ocular pain 6 months prior to admission. Diagnosed with LICA dissection but persistent TIAs despite oral anticoagulant therapy</td>
<td>6 months</td>
<td>Multiple infarcts in L capsule, basal ganglia, and frontal lobes</td>
<td>Spontaneous</td>
<td>Obliteration of pseudoaneurysm and reduction of LICA stenosis to decrease thrombus formation</td>
</tr>
<tr>
<td>9 51/M</td>
<td></td>
<td>Subarachnoid hemorrhage, intracranial vasospasm, headache</td>
<td>4 weeks</td>
<td>None</td>
<td>Spontaneous</td>
<td>Obliteration of false channel given contraindication for oral anticoagulant therapy because of planned abdominal surgery</td>
</tr>
<tr>
<td>10 44/F</td>
<td></td>
<td>Expressive dysphasia, R arm weakness and numbness</td>
<td>4 months</td>
<td>L MCA distribution</td>
<td>Traumatic (vehicular whiplash)</td>
<td>Restoration of LICA luminal integrity and caliber</td>
</tr>
</tbody>
</table>

Note.—L, left; R, right; ICA, internal carotid artery; LICA, left internal carotid artery; RICA, right internal carotid artery; Acomm, anterior communicating artery; MCA, middle cerebral artery.
TABLE 2: Summary of the angiographic characteristics of the lesion, including location, presence of collaterals, method of treatment, stenosis, and complication related to treatment

<table>
<thead>
<tr>
<th>Patient No.</th>
<th>Location of Lesion</th>
<th>Evidence of Angiographic Collaterals</th>
<th>Treatment</th>
<th>Pre/Post Luminal Compromise</th>
<th>Complications</th>
<th>Treatment Complication</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>RICA dissection and occlusion</td>
<td>Absent Acomm, perfusion from R Pcomm and R opthalmic artery</td>
<td>1—Recanalization of true RICA lumen 2—Placement of Single RICA WallStent 6 × 20 (mm)</td>
<td>RICA 100%&gt;0%</td>
<td>Transient vasospasm</td>
<td>Resolved spontaneously upon removal of guide catheter</td>
</tr>
<tr>
<td>2</td>
<td>LICA dissection and stenosis</td>
<td>Patent Acomm and bilateral Pcomm</td>
<td>1—Placement of single LICA WallStent 7 × 20 (mm) 2—Balloon occlusion of RICA 3 days later</td>
<td>LICA 80%&gt;20%</td>
<td>None</td>
<td>N/A</td>
</tr>
<tr>
<td>3</td>
<td>LICA dissection and occlusion</td>
<td>Occluded origin LICA</td>
<td>1—Placement of 3 RICA tandem stents: 1 WallStent 6 × 20 (mm) and 2 GFX 4 × 12 (mm) in RICA</td>
<td>RICA 60%&gt;15%</td>
<td>None</td>
<td>N/A</td>
</tr>
<tr>
<td>4</td>
<td>LCCA dissection</td>
<td>False lumen in CCA shows retrograde flow back down to aortic arch</td>
<td>1—Angioplasty of LCCA stenosis 2—Placement of 2 tandem LCCA WallStents 8 × 40 and 10 × 42 (mm)</td>
<td>LCCA 60%&gt;20%</td>
<td>Persistent minimal filling of false lumen</td>
<td>None</td>
</tr>
<tr>
<td>5</td>
<td>LICA dissection and occlusion</td>
<td>Patent Acomm and left Pcomm</td>
<td>1—Placement of 3 tandem RICA WallStents: 6 × 45, 7 × 20, and 8 × 20 (mm)</td>
<td>RICA 80%&gt;0%</td>
<td>None</td>
<td>N/A</td>
</tr>
<tr>
<td>6</td>
<td>LICA chronic dissection and stenosis</td>
<td>Patent bilateral Pcomm</td>
<td>1—Placement of single LICA WallStent 6 × 20 (mm)</td>
<td>LICA 85%&gt;0%</td>
<td>Retropertoneal hematoma</td>
<td>Transfusion and surgical repair of arteriotomy</td>
</tr>
<tr>
<td>7</td>
<td>LICA dissection and occlusion</td>
<td>Patent Acomm</td>
<td>1—Recanalization of LICA 2—Placement of 1 LICA WallStent 6 × 20 (mm)</td>
<td>LICA 100&gt;0%</td>
<td>Transient high cervical LICA vasospasm</td>
<td>Percutaneous balloon angioplasty</td>
</tr>
<tr>
<td>8</td>
<td>LICA dissection and stenosis with pseudoaneurysm formation</td>
<td>None</td>
<td>1—Stent placement of 1 LICA WallStent 6 × 20 (mm) 2—GDC coil-through-stent of LICA pseudoaneurysm</td>
<td>LICA 70%&gt;0%</td>
<td>None</td>
<td>N/A</td>
</tr>
<tr>
<td>9</td>
<td>LICA dissection at carotid bulb</td>
<td>None</td>
<td>1—Placement of 2 LICA tandem S.M.A.R.T. Stent 8 × 40, 10 × 20 (mm)</td>
<td>LICA 40%&gt;0%</td>
<td>None</td>
<td>N/A</td>
</tr>
<tr>
<td>10</td>
<td>LICA dissection at carotid bulb</td>
<td>None</td>
<td>1—Placement of 1 LICA tandem S.M.A.R.T. Stent 8 × 20 (mm) 2—GDC coil-through-stent of LICA pseudoaneurysm</td>
<td>LICA 75%&gt;0%</td>
<td>None</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Note.—RICA, right internal carotid artery; Acomm, anterior communicating artery; Pcomm, posterior communicating artery; CCA, common carotid artery.
icits, and no new minor or major strokes (0%) prior to patient discharge (Table 3). Two patients had intraprocedural vasospasm of the high cervical carotid artery; this vasospasm was severe enough to necessitate angioplasty in one case (patient 7) and resolved spontaneously after removal of the microguidewire in the other (patient 1). One patient suffered a retroperitoneal hemorrhage after removal of the vascular sheath and placement of an external compression device (patient 6) on postoperative day 1; this required blood transfusion (2 U) and was surgically repaired without complication. A single, major, delayed complication occurred in patient 2, who was referred with a left internal carotid dissection that occurred during angiographic work-up of a giant symptomatic cavernous aneurysm. The flow-limiting left internal carotid artery dissection was treated successfully using stent deployment. Three days later, permanent sacrifice of the right internal carotid artery was performed using detachable silicone balloons after the patient had tolerated successful balloon test occlusion (30 minutes with hypotensive challenge). The patient had an uncomplicated postprocedural course and was discharged home. Eight months later, the patient had a massive uterine hemorrhage with an associated prolonged hypotensive episode, which led to an ischemic stroke of the right hemisphere (contralateral to stent, ipsilateral to balloon occlusion), resulting in left-arm weakness. The deficit has mostly resolved at latest clinical follow-up (24 months).

Outcome
During the acute postprocedural period, no patients developed worsening of symptoms. Two patients treated in the acute phase of dissection showed significant and rapid improvement in symptoms (patients 1 and 5). Clinical follow-up was obtained from all patients at 16.5 ± 0.6 months (range, 8–27 months) and angiographic or sonographic follow-up was available for seven patients at 6.3 ± 0.9 months (range, 3–10 months). There was no evidence of stent occlusion or stenosis in any of the follow-up radiographic studies (0%). Clinical outcome was evaluated using the modified Rankin scale and was improved at latest follow-up (0.7 ± 0.26) compared with pretreatment values (1.8 ± 0.44) (P < .008). Similarly, the Barthel index showed improvement at latest follow-up (99.5 ± 0.5) compared with pretreatment values (80.5 ± 8.6). Except for patient 2 presented above, no other patient in this high-risk subset has suffered any delayed TIA or stroke since treatment at latest follow-up.

Illustrative Cases

Acute Symptomatic Occlusive Dissection Treated with Recanalization and Stent Placement

The patient (patient 1) is a 45-year-old woman with a history of hypertension, lupus erythematosus, and migraine headaches, who underwent diagnostic angiography at an outside institution for evaluation of recurrent headaches with associated left-arm weakness and numbness. During the procedure, the operator noted that an acute dissection had been induced by contrast injection in a spastic vessel. The procedure was halted and intravenous heparin was immediately administered, but the patient developed hemiparesis 1 hour later. She was transferred to our hospital, where emergent angiography revealed that the previously dissected left internal carotid artery now showed tapering to a complete occlusion (Fig 2). Collateral evaluation revealed no anterior communicating flow across the midline from the left internal carotid artery injection, weak flow from the posterior circulation via the posterior communicating artery, and retrograde flow through the right ophthalmic artery from the right external carotid artery. A Rapid Transit microcatheter was used in combination with an In- stencil-10 microguidewire to navigate through the true lumen carefully and to reestablish recanalization of the internal carotid artery, which was noted to harbor a tonsillar loop, a condition previously described to predispose to dissection (24). Evaluation of the major intracranial vessels by contrast injection through the microcatheter revealed no evidence of a large thrombus, and showed that most of the distal branches remained patent. A decision was thus made to treat the dissection flap by elimination of the inflow to the subintimal dissection. A single 8-mm × 2-cm Wallstent was deployed at the proximal inflow zone, after which control angiography revealed that blood flow had resumed to normal. The patient’s left hemiparesis improved rapidly after the procedure, and she was discharged 3 days later with a mild left pronator drift. The patient has had no further TIAs or stroke, and follow-up sonography at 6 months revealed the stent to be patent, with no abnormal flow characteristics. At latest neurologic follow-up (17 months), the patient still complained of poor fine-finger movement, numbness of the left face, and subjective left-leg hypesthesia, but is otherwise intact.

Bilateral, Subacute, Spontaneous Carotid Dissection Progressing to Unilateral Occlusion and Flow-limiting Contralateral Stenosis

This patient (patient 5) is a 51-year-old man, with no significant past medical history, who noted left arm and hand numbness while chopping wood. Four days later, he was found lying in bed, aphasic and disoriented, with right hemiparesis. He was admitted to an outside hospital, and imaging was performed. A head CT scan was unrevealing, but a brain MR scan showed a left posterior frontal infarct, and carotid sonography showed little to no flow in the left internal carotid artery (Fig 3). An angiogram obtained at an outside hospital showed a flow-limiting dissection of the left internal carotid artery and an irregular contour of a patent right
TABLE 3: Outcome of treated patients immediately after the procedure and at latest follow-up

<table>
<thead>
<tr>
<th>Patient (No.)</th>
<th>Immediate Outcome</th>
<th>Long-term Outcome</th>
<th>Anatomic Follow-up</th>
<th>Clinical Follow-up (months)</th>
<th>Pre/Post mRankin and Barthel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Rapid improvement of L hemiparesis, CT evidence of R posterior/superior frontal lobe infarct</td>
<td>1—Residual L hand weakness</td>
<td>6 months U/S patent stent</td>
<td>17</td>
<td>mRanking Pre/Post = 4/1 Barthel Pre/Post = 45/100</td>
</tr>
<tr>
<td>2</td>
<td>No new symptoms, tolerated RICA permanent balloon occlusion for R Cavernous aneurysm treatment</td>
<td>1—Improved R cavernous aneurysm mass effect symptoms 2—Stroke at 8 months following a massive uterine hemorrhage with associated prolonged hypotension</td>
<td>8 months MRA patent stent 10 months U/S patent stent</td>
<td>24</td>
<td>mRankin Pre/Post = 2/1 Barthel Pre/Post = 80/100</td>
</tr>
<tr>
<td>3</td>
<td>No new symptoms, no improvement in left ocular vision, persistent left frontal headache</td>
<td>1—Persistent headache 2—No further TIAs</td>
<td>3 months U/S patent stent</td>
<td>15</td>
<td>mRankin Pre/Post = 2/1 Barthel Pre/Post = 80/100</td>
</tr>
<tr>
<td>4</td>
<td>Resolution of TIAs</td>
<td>Asymptomatic</td>
<td>4 months U/S patent stent</td>
<td>16</td>
<td>mRankin Pre/Post = 2/0 Barthel Pre/Post = 90/100</td>
</tr>
<tr>
<td>5</td>
<td>Improvement of leg, arm, and face hemiparesis, minimal improvement in receptive aphasia</td>
<td>Partial expressive aphasia, receptive speech intact, normal motor and sensory exam</td>
<td>9 months U/S patent stent</td>
<td>20</td>
<td>mRankin Pre/Post = 4/2 Barthel Pre/Post = 20/100</td>
</tr>
<tr>
<td>6</td>
<td>No new symptoms</td>
<td>Asymptomatic</td>
<td>6 months A/G patent stent</td>
<td>27</td>
<td>mRankin Pre/Post = 1/0 Barthel Pre/Post = 100/100</td>
</tr>
<tr>
<td>7</td>
<td>No new symptoms</td>
<td>Asymptomatic</td>
<td>N/A</td>
<td>13</td>
<td>mRankin Pre/Post = 0/0 Barthel Pre/Post = 100/100</td>
</tr>
<tr>
<td>8</td>
<td>Resolution of TIAs</td>
<td>Asymptomatic</td>
<td>6 months A/G patent stent</td>
<td>17</td>
<td>mRankin Pre/Post = 1/0 Barthel Pre/Post = 95/100</td>
</tr>
<tr>
<td>9</td>
<td>No new symptoms</td>
<td>Asymptomatic</td>
<td>N/A</td>
<td>8</td>
<td>mRankin Pre/Post = 0/0 Barthel Pre/Post = 100/100</td>
</tr>
<tr>
<td>10</td>
<td>No new symptoms</td>
<td>1—Persistent mild expressive dysphasia 2—No further TIAs</td>
<td>N/A</td>
<td>8</td>
<td>mRankin Pre/Post = 2/2 Barthel Pre/Post = 95/95</td>
</tr>
</tbody>
</table>
intracranial carotid artery. Intravenous anticoagulation with heparin was begun, and the patient was transferred to our center. Examination disclosed that the patient was aphasic and had a right upper motor facial and right upper and lower extremity hemiparesis, with no response to visual threat from right-sided confrontation. A repeat diagnostic angiogram was obtained 2 days after the previous study. The new angiogram revealed progression of the previously noted left internal carotid artery dissection to complete occlusion. In addition, it showed the previous contour abnormality of the right internal carotid artery had progressed to a long-segment stenosis with an associated distal pseudoaneurysmal dilatation. Given the disease progression in both internal carotid arteries despite anticoagulation, a decision was made to proceed with securing a patent luminal conduit to prevent further progression of the right-sided dissection to complete occlusion. Recanalization of the left internal carotid artery was not attempted, given the duration and extent of the established left middle cerebral artery distribution infarct. Microcatheterization of the true lumen was performed with a Rapid-Transit microcatheter over an Instinct-10 microguidewire. A 6-mm × 4.5-cm Wallstent was deployed over the Stabilizer exchange microguidewire at the proximal inflow zone to tack down the dissection flap, followed by balloon angioplasty of the long-segment stenosis. An additional two Wallstents (7 mm × 2 cm and 8 mm × 2 cm) were needed to reconstruct the vascular channel to the same size as the native vessel. After the procedure, the patient had progressive improvement of his right-sided face and arm weakness and of his receptive aphasia. Neurologic follow-up at 20 months revealed persistent, moderate, expressive aphasia with naming difficulty, normal comprehension, symmetric face, absent pronator drift, and intact sensation with normal gait. Sonography confirmed persistent patency of the right internal carotid artery and stents, with no abnormal flow characteristics within the deployed stents.

**Treatment of a Symptomatic Carotid Dissection Related to an Aortic Dissection**

The patient (patient 4) is an 83-year-old right-handed man with previous type A aortic dissection (involving both the ascending and descending aorta) 10 years prior to admission, which was surgically repaired using a Dacron graft. Angiographic study of the arch after an episode of transient aphasia revealed extension of the repaired aortic dissection into the left common carotid artery. The patient was begun on aspirin and Coumadin and had been neurologically without symptoms until 3 weeks prior to admission. The patient began to have left hemispheric TIAs, resulting in episodes of right-hand weakness, with poor fine-finger movement, despite therapeutic oral anticoagulant therapy. The patient was begun on intravenous anticoagulation. Angiography of the aortic arch revealed an extensive left common carotid artery dissection involving the origin of the common carotid artery, the cervical carotid bifurcation, and the proximal internal carotid artery to C3 (Fig 4). Flow was antegrade in the true lumen of the left common carotid artery and retrograde in the false lumen, which exited at a poorly visualized area in the aortic arch. The retrograde flow back to the arch was the result of a pressure gradient from a Venturi effect at the false lumen exit into the aorta. A 5F Simmons-I catheter was used to catheterize the left common carotid artery selectively and was exchanged for a 9 F guide catheter. Two Wallstent devices were deployed in an overlapping fashion. The proximal stent measured 8 mm × 4 cm and was deployed in the common carotid artery across the inflow zone. The second stent measured 10 mm × 4.2 cm and was deployed distally in the carotid bulb and into the origin of the cervical internal carotid artery. Postdeployment angioplasty was performed for each stent by using an 8-mm then a 10-mm angioplasty balloon (Powerflex; Cordis) (Fig 4). Postprocedure angiography showed near-complete elimination of the false lumen and the retrograde flow component. The patient tolerated the procedure well, without further symptoms, and was maintained on Coumadin until 1.5 months after discharge, when he underwent open prostatectomy without complication. Carotid sonography performed at 6 months showed persistent patency of the stent, without evidence of retrograde flow in the previous false lumen. Neurologic follow-up at 16 months revealed no further TIAs.

**Discussion**

The annual incidence of spontaneous carotid dissection has been reported to be 2.6 per 100 000 (27), and has been estimated to account for up to 20% of strokes in the younger population (28). Most infarcts associated with dissection have been proposed to be embolic in nature, although a non-negligible proportion consists of hemodynamic flow-related infarcts (5, 6). Despite anticoagulation, a number of patients progress to have hemodynamically significant residual stenosis or develop pseudoaneurysms (2), leading to a risk of hemodynamically significant stenosis or to a danger of delayed distal embolization (10). Although the rate of repeat embolization in unilateral spontaneous carotid dissection lesions is low (6, 29), and, as such, does not justify placement of an intravascular stent as primary treatment, the patients presented in this series underwent stent placement only after they had fulfilled stringent criteria such as failure of medical therapy and absence of low-risk surgical options. Although surgery has been performed for direct repair of carotid dissection, it is difficult to identify the inflow zone or repair the entire extent of the dissection (1). Consequently, surgery for carotid dissection is associated with a significantly greater risk than is carotid endarterectomy for atherosclerosis (1).
Surgical options for the treatment of hemodynamic insufficiency as a result of carotid dissection have included an extracranial-to-intracranial bypass procedure for patients with hemodynamic insufficiency who have not responded to anticoagulant therapy (29). The surgical treatment of dissection-induced cervical carotid aneurysm has included bypass grafting (30) and ligation of the internal carotid artery for pseudoaneurysm repair (29).

One of the drawbacks of conventional and stent-supported balloon angioplasty of the carotid artery for atherosclerosis has been the risk of distal embolization of atheromatous debris during the balloon dilatation phase of the procedure. We did not encounter in this series any cases of distal embolization, perioperative stroke, or new deficits attributable to stent deployment despite performing angioplasty of the deployed stent in the majority of
cases. Unlike atherosclerotic stenoses, which require higher-pressure angioplasty and necessitate fracturing the plaque, the stent deployment for dissection is less traumatic and is performed at a lower pressure because the lesions are more compliant and are not usually atheromatous or calcified.

Additional advantages of endovascular therapy of carotid dissection are that it enables the identification of the true and false lumens by superselective catheterization and angiography, and further allows the recanalization of completely occluded vessels by use of microcatheter techniques, provided the thrombus burden is not prohibitive (Fig 2). The availability of flexible stents that conform to the contour of the artery enables the sequential reconstruction of an angiographically smooth luminal outline through the narrowed true lumen and obliteration of the false lumen (Fig 5). In addition, the endovascular approach circumvents the need for blood flow occlusion, which is required during di-
Fig 3. A 51-year-old man (patient 5) with a 10-day history of left-sided TIAs now presents with sudden onset of aphasia and right hemiparesis. Axial T1-weighted contrast-enhanced MR imaging shows a left frontal infarct (A). Digital subtraction angiography of the left common carotid artery showed progression of a left internal carotid dissection from a partial stenosis, 2 days prior, to a complete occlusion despite systemic anticoagulation (B). Digital subtraction angiography of the right common carotid artery revealed a dissection of the right internal carotid artery with an associated long-segment stenosis and an expansile pseudoaneurysm at the distal end (C). A Rapid Transit microcatheter was used to navigate the patent lumen of the right internal carotid artery and to deploy three Wallstents (6 mm × 45 mm, 7 mm × 20 mm, and 8 mm × 20 mm) in a tandem overlapping fashion with postdeployment balloon angioplasty (D). The procedure resulted in reconstitution of the normal lumen of the right internal carotid artery (E).

Fig 4. An 83-year-old man (patient 4) with a history of aortic dissection, which was surgically repaired 10 years previously, presents with left hemispheric TIAs and episodes of aphasia. Digital subtraction angiography of the left common carotid artery outlines a chronic extensive dissection, with antegrade flow through the true lumen and retrograde flow in the false lumen down to the aorta, by virtue of the Venturi effect at the aortic arch (A, early injection; B, late injection). Two stents were placed in tandem overlapping fashion, followed by postdeployment angioplasty, resulting in a near-complete elimination of the retrograde flow in the now-reduced false lumen (C, D).
A 51-year-old man (patient 3) developed sudden loss of vision in the left eye, a left Horner’s syndrome, and orthostatic light-headedness. Digital subtraction angiography with injection of the right common carotid artery shows intimal dissection of the right internal carotid artery in the high cervical region, with delineation of the true and false lumen (A, B). Injection of the left common carotid artery, which had evidence of dissection but was shown to be patent on an angiogram obtained 4 days earlier, now reveals complete left internal carotid artery occlusion (arrowhead) despite systemic anticoagulation (C). A Rapid Transit microcatheter was used to catheterize the true lumen, followed by deployment of a Wallstent (6 mm × 20 mm) in the proximal portion of the dissected segment (D). Persistent filling of the false lumen, however, required the tandem placement of two additional GFX stents (4 mm × 12 mm) in the petrous segment of the right internal carotid artery (E), with reconstitution of the normal luminal diameter (F).

rect surgical bypass procedures, a crucial factor that most patients in the current report would not have tolerated because of contralateral involvement or poor collateral flow. Finally, the endovascular approach enables the simultaneous treatment of any coexistent pseudoaneurysmal dilatations by embolization using Guglielmi electrolytically detachable coils, by the coil-through-stent technique, which was employed in two cases (patients 8 and 10) (Fig 1). Although the current series is the largest reported to date, the number of patients treated remains small. Nonetheless, the procedure has proved to be safe, because it was not associated with any periprocedural TIA or stroke. The success in reducing dissection-induced stenosis, the patency rate obtained at follow-up, and the lack of TIAs suggest that stent angioplasty offers a viable alternative to complex surgical bypass procedures. The long-term efficacy and durability of stent placement for carotid dissection remains to be determined. It is unknown whether intimal hyperplasia, which was not detected during the anatomic follow-up periods de-
scribed herein, will become of concern in the distant future.

Conclusion

The results of the intermediate-term follow-up provided in this report suggest that treatment of carotid dissection is a useful technique for select high-risk cases, such as when the lesion is hemodynamically significant from severe stenosis or contralateral occlusion, or when anticoagulation has failed to prevent TIAAs or is contraindicated. Additional, extended, long-term angiographic and clinical follow-up will help define the potential importance and relative value of this technique for the treatment of carotid dissection.

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References