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ABSTRACT

BACKGROUND AND PURPOSE: Flow-diverter stents are emerging for the endovascular treatment of difficult-to-treat or otherwise untreatable cerebral aneurysms (wide-neck, fusiform, dissecting, blisterlike, or giant). We assessed the clinical safety and efficacy of the Flow-Redirection Endoluminal Device.

MATERIALS AND METHODS: This was an institutional review board–approved single-center observational clinical study in 29 patients with 34 aneurysms elected to be treated by endovascular intervention. After providing informed consent, patients were included according to the following criteria: aneurysm fundus-to-neck ratio \( \leq 2 \) or neck diameter \( \geq 4 \) mm, fusiform, dissecting, or giant aneurysms. The primary end point for clinical safety was the absence of death, absence of major or minor stroke, and absence of transient ischemic attack. The primary end point for treatment efficacy was complete angiographic occlusion according to the O’Kelly Marotta grading scale immediately after the procedure and at follow-up after 3 and 6 months (O’Kelly Marotta D: complete occlusion).

RESULTS: The Flow-Redirection Intraluminal Device deployment was technically successful in all cases. In 26/29 (89%) of patients, the primary end point of safety was reached; in the 3 remaining patients, 1 disabling ischemic stroke and 2 minor strokes with complete recovery at follow-up were observed. Angiographic (DSA and MRA) and clinical follow-up were available after 3 months in 29/29 (100%) and after 6 months in 25/29 (86%) patients (after 6 months, only MRA follow-up was performed according to our study protocol and institutional standard). At 3-month follow-up, complete occlusion was reached in 19/34 aneurysms (O’Kelly Marotta D: 19/34; 56%). At 6-month follow-up, aneurysm occlusion was complete in 22/30 aneurysms (O’Kelly Marotta D: 22/30; 73%).

CONCLUSIONS: Deployment of the Flow-Redirection Intraluminal Device flow-diverter stent is safe and effective in the treatment of difficult-to-treat or otherwise untreatable intracranial aneurysms.

ABBREVIATIONS: FD = flow-divider; FRED = Flow-Redirection Intraluminal Device; LTA = light transmission aggregometry; OKM = O’Kelly Marotta grading scale; PTA = percutaneous transluminal angioplasty.

Endovascular treatment of intracranial aneurysms by coiling has become an accepted alternative to surgical clipping, with increasing evidence for lower morbidity and mortality rates, especially in clinical equipoise. However, especially in wide-neck, fusiform, dissecting, and giant aneurysms, incomplete coiling and reperfusion are still a major limitation preventing stable long-term occlusion. Aneurysm recanalization and/or neck remnants may be observed despite further refinement in coil technology such as coated platinum coils and/or procedural modification such as the balloon-remodeling technique or stent-assisted coil embolization.

The development of flow-diverter (FD) stents has offered the potential of aneurysm occlusion through thrombosis triggered by the disruption of flow into the aneurysm sac. As a key element of construction, these stents have a braided mesh with a densely covered surface. Once the FD is expanded to cover the aneurysm neck, thrombosis is induced by stasis of flow within the aneurysmal sac. The porosity of the FD mesh and the pressure gradient between parent and smaller adjacent branch vessels preserve flow and patency of the latter even if covered. The Flow-Redirection Endoluminal Device (FRED; MicroVention, Tustin, California) is a new generation of FDs for reconstruction of the parent artery and aneurysm occlusion. Its unique dual-layer design composed of a low-porosity inner mesh and a high-porosity outer stent may provide potential advantages over other available FDs in safe deliverability and effective occlusion.
of the target lesion. We report our analysis of the clinical safety and efficacy of the FRED in 29 patients with 34 aneurysms.

MATERIALS AND METHODS

Patient Selection

Approval for prospective data collection of all interventional procedures reported in this study was given by the institutional review board of the medical faculty. Informed consent for study inclusion was obtained from the patients or their legal representatives. Patients were enrolled between February 2013 and July 2014.

Criteria of Inclusion/Exclusion

Patients with intracranial aneurysms were included in the study and treated with the FRED if the aneurysms satisfied the following morphologic and morphometric criteria of inclusion: 1) wide-neck saccular (fundus-to-neck ratio <2 or neck diameter >4 mm), 2) fusiform or circumferential, 3) dissecting, 4) blisters-like (broad-based at a typical location, eg, the suprachinoid side wall of the ICA and ≤2 mm), and 5) giant (defined as ≥25-mm maximum diameter) or those presenting with mass effect. We considered these aneurysms to have a high likelihood of technical failure and/or recurrence with conventional endovascular techniques. Five patients (5/29, 17%) with recanalization after previous coil ing were enrolled. Exclusion criteria were consensus about clipping as the appropriate treatment option in cases with unfavorable branch configuration with an acute angle in relation to the parent vessel or an unfavorably small diameter of the parent artery of ≤2.0 mm.

FRED

The FRED is an FD designed for the treatment of intracranial aneurysms. The device consists of a braided self-expandable closed-cell dual-layer stent (also referred to a “stent within a stent”) with a low-porosity inner mesh of higher pore attenuation (48 nitinol wires) and an outer stent with high porosity (16 nitinol wires). This dual-layer design is restricted to the midsection of the stent and encompasses 80% of its total length. An interwoven double helix of radiopaque tantalum strands attaches the inner mesh to the outer stent and improves visibility over its full length of dual-layer coverage. Each end of the stent is flared and marked by 4 radiopaque tips. Currently, the FRED is available in 5 different diameters (3.5, 4.0, 4.5, 5.0, and 5.5 mm) recommended for vessel diameters from 2.5 to 5.5 mm and at working lengths (dual-layer coverage) from 7 to 56 mm. The FRED is attached to a delivery microcatheter with a radiopaque distal tip and a proximal marker. If the device is not correctly positioned across the aneurysm neck, the pusher allows resheathing and repositioning of the stent as long as ≤80% of its length has been unsheathed/deployed. As soon as the 0.027-inch delivery-microcatheter (Headway 27; MicroVention) is fully withdrawn over the delivery microwire, the coupling wire releases and the stent is deployed.

Description of Technique

Every procedure was performed via a transfemoral approach with the patient under general anesthesia. A 6F–8F guiding catheter was introduced through a femoral sheath into the carotid or vertebral artery. The radiologic examination of the target vessel was performed by using a biplane angiographic system (Artis zee biplane; Siemens, Erlangen, Germany) and 3D rotational angiography. FD length was chosen according to the length of the aneurysm neck with the procedural goal to ensure arterial wall coverage with the inner mesh extending to at least 2 mm beyond both the distal and proximal limits of the neck. For FD delivery, a Headway 27 microcatheter was navigated past the aneurysm neck with a microguidewire, Traxcess (MicroVention) or Radiofocus Guide Wire GT 16 (Terumo, Tokyo, Japan). Under roadmap guidance, the FD was unsheathed by slowly withdrawing the delivery microcatheter for 2–3 minutes while the delivery wire was gently pushed on to facilitate complete opening and wall apposition of the FD. An in-stent percutaneous transluminal angioplasty (PTA) was performed with a balloon microcatheter (HyperGlide or HyperForm; Covidien, Irvine, California) if incomplete opening was observed on 2D angiography or 3D DynaCT (Siemens). In those cases in which additional aneurysm covering was performed, a microcatheter was initially positioned inside the aneurysmsacaclosedbyjailingofthemicrocatheter between the parent vessel wall and the FD at deployment.

Anticoagulation/Antiplatelet Regimen

Patients were prepared with aspirin, 100 mg, and clopidogrel, 75 mg (loading dose of 300 mg), 5 days before treatment. During the procedure, anticoagulation was initiated with a bolus of standard heparin (70–100 IU/kg) followed by intravenous administration to maintain an activated clotting time of ≥250 seconds. After the procedure, a daily dose of clopidogrel (75 mg) and aspirin (100 mg) was given for 3 and 6 months, respectively. Platelet inhibition was tested by using the light transmission aggregometry (LTA) method on the day before or immediately before starting the procedure. If according to this test, the patient was considered a nonresponder (LTA result of <6 µA), an LTA test with 2-MeS-AMP (selective P2Y12-adenosine diphosphate receptor antagonist) was performed. In case of an LTA result with 2-MeS-AMP ≤5 µA, a pharmacokinetic nonresponse was suspected and a reloading dose of clopidogrel (300 mg) was administered. In the case of repeated test failure even after reloading or an LTA result with 2-MeS-AMP >5 µA, pharmacodynamic resistance was assumed and prasugrel was substituted for clopidogrel. In patients with ruptured aneurysms, tirofiban was administrated intravenously during the endovascular procedure before FD deployment and aspirin and clopidogrel were started after the procedure.

Evaluation of End Points

Safety. Clinical evaluation by detailed neurologic examination was performed immediately after the procedure, at 2 hours after the procedure, and during the clinical visit on the following day and was finally scored by the mRS at discharge and follow-up by a board-certified neurologist not blinded to treatment. Technical safety, including navigation, visibility, deployment, opening, and vessel wall apposition of the FD, were rated by 2 senior neurointerventionalists (M.A.M., M.P.).

Efficacy. Each angiographic result was categorized according to the O’Kelly Marotta (OKM) grading scale for assessment of cere-
bral aneurysms treated by flow diversion. Each aneurysm was assigned an occlusion grade according to the initial degree of filling (A = total filling, B = subtotal filling, C = entry remnant, D = no filling) and the degree of stasis (prolongation of stasis into 1 = arterial, 2 = capillary, 3 = venous phase). Detailed follow-up examinations were performed with both DSA and MR imaging (3T, Magnetom Trio or Magnetom Verio; Siemens; TOF MRA with and without contrast agent) at 3 months. After 6 months, MRA was exclusively performed. To evaluate thrombosis of the aneurysm and shrinkage of the aneurysm sac, we measured the maximum diameters of all lesions on preprocedural MR images in the axial plane with 2 additional orthogonal planes and compared them with MR imaging at 3 and 6 months. Aneurysm morphometry on MR imaging was categorized into 3 groups: 1) complete or partial (≥20%) decrease, 2) stable, 3) progressive (≥10%).

RESULTS

Twenty-nine patients (14 female, 15 male; mean age, 58 years; age range, 32–80 years) with 34 aneurysms fulfilled the criteria of inclusion. Patient demographics and aneurysm features are shown in On-line Table 1, and clinical and angiographic details are listed in On-line Table 2.

After preparation with aspirin and clopidogrel, a pharmacokinetic resistance was suspected in 1 patient. Following a reloading dose of clopidogrel, responder status was confirmed. In another patient, pharmacodynamic resistance was assumed on the basis of LTA results and prasugrel was substituted for clopidogrel. Twenty-three aneurysms originated from the ICA/posterior communicating artery; 3, from the anterior cerebral artery; and 8 were located in the posterior circulation. We included 17 wide-neck saccular aneurysms, 3 fusiform/dissecting aneurysms, 4 blisterlike aneurysms, and 10 large/giant aneurysms (including 2 partially thrombosed aneurysms). Nine aneurysms were symptomatic, and 5 aneurysms were in acute/subacute stages of SAH. In all patients and for all aneurysms, the FRED could be navigated to the target area and was deployed successfully across the aneurysm neck with complete neck coverage.

Thirty-two of 34 (94%) aneurysms were treated by deployment of a single FD. Two patients had undergone previous endovascular treatment with stents for the same target lesion: One patient with reperfusion was previously treated by stent-assisted coiling (LEO+ stent; Balt Extrusion, Montmorency, France); the other patient with a 16.3-mm large-neck cavernous ICA aneurysm was pretreated with 2 overlapping LEO+ stents (3.5 × 50 mm). Median diameters of the stented parent vessel segments were proximally 3.8 mm (range, 2.0–5.3 mm) and distally 3.4 mm (range, 1.9–5.7 mm). In 10 cases, bioactive coils (Cerecyte; Codman Neurovascular, Raynham, Massachusetts) were used with a median packing attenuation of 20% (range, 9%–43%). A PTA before FD deployment was performed in 1 patient with a 50% stenosis directly proximal to a fusiform vertebral aneurysm. PTA after FD deployment was performed in 5 patients with paraophthalmic or supraclinoid ICA aneurysms. In all of these 5 patients, near-complete (≥90%) wall apposition of the FD had been achieved before PTA; however, PTA was elected to further improve wall apposition and opening of the vessel diameter. At termination of the procedure, an immediate change of angiographic filling (OKM grading scale) was achieved in 33/34 (97%) aneurysms with complete or near-complete early occlusion (OKM C or D) occurring in 9/34 (26%) within 30 minutes after FD deployment.

Clinical Safety and Procedure-Related Complications

The primary end point of safety (absence of any new transient or permanent neurologic deficit or death) was reached in 26/29 (90%) patients. A new permanent neurologic deficit by major stroke occurred in 1 of the treated patients after the procedure. Minor strokes were observed in 2 patients (1 immediately postprocedural and 1 after 2 months [see below]) who recovered completely and were each asymptomatic after 3 months of follow-up (NIHSS = 0, mRS = 0). Neither mortality nor transient ischemic symptoms occurred in any of the treated patients after the procedure and throughout follow-up.

The major stroke occurred in a patient with a ruptured aneurysm of a short paramedian circumferential perforator branch emerging from the basilar trunk. This very small aneurysm measured 1.6 mm and arose from the midsection of the basilar trunk just distal to the AICA offspring. It was unmasked only on the repeat angiogram 12 days after SAH and was not visible on the initial angiogram obtained on the day of SAH. Weight-adapted tirofibin was started intravenously with heparin after femoral access and was maintained for 12 hours. The shortest available FRED (3.5 × 7 mm) was successfully deployed to cover the midsection of the basilar trunk, and the control angiogram demonstrated immediate contrast stasis within the dome with preserved flow through the perforator. Nine hours after FRED deployment, the patient developed a right-sided high-grade hemiparesis and dysarthria. DWI MR imaging was performed immediately and confirmed a left paramedian pontine ischemic infarction at the level of the aneurysm. At discharge, the patient had a Glasgow Coma Scale score of 15 with substantially improved motor strength on her right side.

One minor stroke occurred in a patient with a superior cerebellar artery aneurysm 1 day after FRED deployment. DWI MR imaging confirmed a few microembolic infarcts in the ipsilateral superior cerebellar artery territory. Immediate control DSA showed intact duplicated superior cerebellar arteries and contrast stasis within the dome without any evidence of local thrombus formation inside the FD or thromboembolic branch occlusion distal to it. The patient was discharged home with a residual mild unsteadiness of gait, which had completely resolved at 3-month follow-up.

The second above-mentioned minor stroke occurred in a patient with a giant partially thrombosed ICA aneurysm 2 months after FD deployment. Because of infection of his ventriculoperitoneal shunt, which required conversion to a ventriculostial shunt, prasugrel (the patient was a nonresponder to aspirin and clopidogrel) was terminated. After 1 week of anticoagulation on heparin only, the patient developed sudden hemiparesis due to complete FD in-stent thrombosis with complete ICA occlusion confirmed by Doppler sonography and emergency DSA. Resolution of symptoms could be observed clinically during emergency DSA in the awake state after pharmacologic induction of controlled hypertension with DSA documentation of sufficient col-
lateral circulation through the circle of Willis and leptomeningeal anastomoses. MR imaging the next day showed development of only microembolic watershed infarcts and absence of any cortical or basal territorial infarction. Full resolution of symptoms was observed in this patient at discharge, and stable ICA occlusion without any evidence of residual antegrade flow was documented by Doppler sonography and MR imaging. None of the other patients developed any new transient or permanent neurologic deficits during follow-up.

An asymptomatic in-stent thrombosis in the acute stage after SAH Hunt and Hess 3° was observed in 1 other patient with a dissecting aneurysm of AICA origin. Appositional clot formation occurred within the FD at the level of the aneurysm neck 5 minutes after FD deployment. This patient had received prior weight-adapted tirofiban (30 minutes before deployment) and heparin, and the clot dissolved after pharmacologic induction of controlled hypertension. Subsequently, DWI MR imaging ruled out brain infarction, and the patient did not develop any new symptoms.

**DISCUSSION**

In this single-center prospective clinical study, the FRED was evaluated for the treatment of otherwise untreatable or difficult-to-treat intracranial aneurysms (ie, wide-neck, fusiform, dissecting, blisterlike, or giant). This new generation of intracranial FD appeared to be a promising safe and effective alternative treatment to other FDs for these complex aneurysms. The primary end point for clinical safety (ie, absence of death and major or minor stroke and absence of transient ischemic attack) was reached in 26/29 (90%) treated patients.

The observed overall permanent morbidity of 3.4% and mortality of 0% in our study was low. In other prospective clinical studies on different intracranial FDs (Pipeline Embolization Device; Covidien, Irvine, California; Silk+; Balt Extrusion; Surpass stent; Stryker Neurovascular, Fremont, California), similar end points occurred with 0%–15% permanent neurologic deficits and 0%–8% mortality.7–16

**Routine Follow-Up (Clinical, DSA, MRA)**

Complete angiographic (DSA and MRA) and clinical follow-up could be performed after 3 months in 29/29 (100%) and after 6 months in 25/29 (86%) patients (after 6 months only MRA was performed according to our study protocol and institutional standard). At 3-month follow-up, a complete occlusion (OKM D) was achieved in 19/34 (56%) aneurysms. A near-complete occlusion (OKM C) was reached in 12/34 (35%) aneurysms. Reduction of aneurysmal sac filling and prolongation of stasis were noted in 27/34 (79%) aneurysms from the initial postprocedural control DSA to the 3-month follow-up control DSA. At 6-month follow-up, 7/30 (23%) aneurysms showed a decrease of residual inflow, and none developed recurrence with enlargement of the aneurysm at the base. At 6-month follow-up, aneurysm occlusion was complete (OKM D) in 22/30 (73%) aneurysms. A near-complete occlusion (OKM C) was detected in another 8/30 (27%) aneurysms. On follow-up angiograms, in no case was any in-stent stenosis, in-stent thrombosis, or migration of the FRED implant observed.

MR imaging analysis at 3 months showed shrinkage or complete disappearance of the thrombosed aneurysm in 59% (20/34 aneurysms) and no change in size in 41% (14/34 aneurysms including 10 additional aneurysm coilings). On MR imaging at 6 months, shrinkage or complete disappearance was observed in 70% (21/30 aneurysms) with no change in 30% (9/30 of aneurysms, including 8 that were additionally coiled).

Two illustrative cases are demonstrated in Figs 1 and 2.
Altogether, we report in this study that the end point of clinical safety was not achieved in 2 patients with minor ischemic stroke, in whom complete resolution of symptoms was observed after 3-month follow-up and in 1 patient with major ischemic stroke. In the latter patient, the only one with a permanent neurologic deficit, an exceptionally rare and difficult-to-treat aneurysm was targeted. Only repeat DSA but not initial DSA revealed a ruptured aneurysm of a perforator branch of the trunk of the basilar artery. In the literature, only 13 cases were reported with this type of aneurysm. Because this aneurysm was not amenable to any coiling or neck-clipping technique, we decided in favor of treatment with flow diversion by using the FRED. Despite sufficiently effective antiplatelet therapy according to platelet function tests, however, the patient’s symptoms, consistent with a delayed perforator stroke 9 hours after FRED deployment, were confirmed by DWI. The exact location and anatomic extension of the ischemic pontine DWI lesion was consistent with delayed occlusion of the perforator branch harboring the aneurysm and was likely the inevitable effect of decreased flow in this vessel.

In one patient who had microembolic infarcts in the superior cerebellar artery territory, from which she recovered completely at 3-month follow-up, a similar flow-related pathogenesis seems most plausible. In a larger series, only observing 32 posterior circulation aneurysms treated with FDs, Phillips et al reported a permanent neurologic complication rate of 9.4%, with perforator territory infarctions occurring in 14% of the subgroup of patients with basilar artery aneurysms. These authors concluded that treatment with an FD is safe and effective but should be reserved for cases not easily or effectively treatable with conventional endovascular techniques, owing to the high rate of ischemic perforator infarcts following FD use in the posterior circulation and the basilar artery trunk in particular.

Two cases (2/29 patients, 6.9%) of in-stent thrombosis occurred in our study: one appositional clot formation in the acute stage of SAH, which was fully reversible on angiography and without clinical consequences under controlled hypertension together with weight-adapted tirofiban and heparin; and the other at 2 months after termination of prasugrel because of infection of his ventriculoperitoneal shunt requiring surgical intervention for conversion into a ventriculostrial shunt. No further delayed thromboembolic events after the procedure were detected in any other patient by occurrence of clinical symptoms or by follow-up MR imaging.

In the present trial, FD deployment was technically successful in all cases. Insufficient opening of the FD, which is reported for other available FDs to occur in up to 10% of cases, was not observed in any case for the FRED in our study. We elected to perform in-stent PTA of the FRED if the slightest suspicion of incomplete wall apposition arose on the immediate postdeployment control angiograms (5/29 patients, 17%). This operation standard was chosen because we consider the risk of thrombus formation and subsequent parent artery occlusion to be high if suboptimal wall apposition is tolerated. In those 5 cases in which we performed PTA within the FD stent (all were paraophthalmic or supraclinoid ICA aneurysms with sharp angulation of the carotid siphon), opening of the stent before PTA was still near-complete with at least 90% of the stent diameter and was further improved to 100% after PTA. To ensure proper opening and full deployment of the FRED, we ensured that unsheathing and release of the FRED occurred in a slow and controlled fashion during several minutes by gentle application of the push/pull technique with the delivery microcatheter tip properly held in a central position within the central vessel lumen. However, the operator should be aware that even if these precautions are taken, incomplete opening still may occur, which should prompt complete retrieval of the device by simply resheathing it into the delivery microcatheter.

On follow-up DSA at 3 months, no in-stent stenosis or changes in stent morphology such as “fish mouth” (ie, inward crimping of 1 or both ends of FD) or “foreshortening” phenomena were observed, which were described in an early series by Kocer et al in 5 cases with a FRED. In our series of 29 patients, we exclusively used the second-generation version of the FRED, which obtained approval for human clinical use within the European community by CE marking in December 2012. Evaluation of the efficacy of occlusion showed promising results for the FRED in otherwise untreatable or difficult-to-treat aneurysms during early follow-up until 3 and 6 months postoperatively. Diagnostic follow-up was achieved not only by invasive angiographic but also MR angiographic evaluation, including contrast-enhanced TOF-MR angiography considered to be sensitive for residual aneurysmal inflow, especially when nonenhanced TOF-MR angiography source images are used for subtraction. At 3-month follow-up, complete occlusion (OKM D) was achieved in 19/34 (56%) aneurysms. At 6-month follow-up, aneurysm occlusion was complete in 22/30 (73%) aneurysms defined as an OKM grade of “D” (complete occlusion), which, on the Raymond and Roy scale, would correspond to grade I.

Compared with other prospective clinical studies on different intracranial FDs (the Pipeline Embolization Device, SILK+; and Surpass), our efficacy rate of 86% in aneurysms of <10 mm was similar to the reported complete occlusion rate at 6 months of 99%–93%. However, in aneurysms of >10 mm, our occlusion rates were lower than the rates reported in the Pipeline Embolization Device for Uncoilable or Failed Aneurysms trial and in the recent meta-analysis by Brinjikji et al. One possible explanation is that the low number of aneurysms of >10 mm in our study might have led to deviations from the population mean, which, due to the larger group size, might have been estimated more accurately in a larger study such as the Pipeline Embolization Device for Uncoilable or Failed Aneurysms trial, for example. Another possible explanation is that we required the 6-month follow-up to include TOF-MRA and applied rigorous aneurysm evaluation of nonenhanced TOF, contrast-enhanced TOF and, importantly, also on subtracted TOF images on which we rated any suspicion of residual inflow or very slow inflow as noncomplete occlusion.

In experimental aneurysm models and also in human studies, it could be regularly observed that FD stents can induce complete and stable occlusion of the aneurysm sac by stasis of flow and subsequent thrombosis, even without additional introduction of coils. In 10 patients, aneurysm occlusion was observed after adjunctive use of coils, introduced by a jailed microcatheter during the same treatment session before FD deployment. In our current
practice, we use adjunctive coiling if aneurysms are large or giant, in the acute stage of SAH, or in cases of retreatment/recurrence of previously ruptured aneurysms pretreated by coiling. The primary rationale for adjunctive coiling in our view is to prevent aneurysm rupture and to enhance stable long-term occlusion. Additional coiling may not only prevent early or delayed rupture but also, especially for extremely wide-neck aneurysms, provide good mechanical support and stabilization for FD deployment and a stable long-term position, in that (partial) herniation or bulging of the FD into the aneurysm sac can be avoided.

The recently introduced FRED may offer several potential advantages over other currently available intracranial FDs: 1) Due to its dual-layer design and availability in long sizes of up to 56 mm, the scaffolding effect of the stent for obtaining outward stability toward the wall of the parent vessel seems to be enhanced and, most important, is achievable in a single treatment session, which makes this device especially useful in fusiform or giant aneurysms, whereas 2 sequential procedures might have been necessary before (scaffolding stent first followed by FD implantation). 2) The outer stent is woven with 16 wires leading to lower friction within the delivery microcatheter, thus facilitating deployment especially of longer stents—other available FDs, for example, are composed of 48 wires (Silk and Pipeline Embolization Device) or 96 wires (Surpass). 3) The additive radial force vectors of the inner and outer stent provide a high degree of reliability of stent opening, a feature especially critical when deployment includes ICA segments around the carotid siphon. 4) There is improved lamina
tion of blood flow through its dual-layer design and higher pore attenuation with 16 + 48 wires compared with 48 wires of the Pipeline Embolization Device and Silk. There is lower vessel wall coverage in the longitudinal direction because of the shorter inner stent (shortest available on the market), which is designed to limit the working layer mainly to the neck of the aneurysm and to spare adjacent branches or perforator arteries, to maintain the patency of these vessels. 6) There is enhanced visibility of the central stent segment and of its distal and proximal ends.

We acknowledge that the major limitations of this prospective study include the limited number of 29 patients, a short follow-up period, MRA only after 6 months, and lack of randomized comparisons with other potentially efficacious therapies. Furthermore, most aneurysms were located at the parapaphalamic segment of the ICA, the mean aneurysm size was relatively small, and not all cases were treated solely by FDs but additional coiling was performed in some cases. However, recruitment followed clearly defined criteria of inclusion/exclusion, and clinical/radiologic follow-up was rigorous, including not only invasive angiography but also MR imaging to measure accurately the shrinkage of the aneurysm sac and nonechanced and contrast-enhanced TOF-MR angiography to detect even small amounts of slow residual inflow that may be silent on invasive angiography. Furthermore, our cohort is the largest so far in which the safety and efficacy profile of this new generation of intracranial FDs has been investigated and may serve as a basis for subsequent larger and multicenter studies.

CONCLUSIONS

The FRED for treatment of difficult-to-treat or otherwise untreatable intracranial aneurysms was prospectively observed in this clinical study to provide a high degree of safety and efficacy, measured as complete aneurysm occlusion on DSA and MR imaging during 6-month follow-up. Long-term durability and safety still remain to be proved by larger series and after prolonged follow-up. We report that it is reasonable, safe, and effective to use intracranial FDs such as the FRED, especially for targeting otherwise untreatable or difficult-to-treat complex intracranial aneurysms.

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