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ABSTRACT

BACKGROUND AND PURPOSE: Although flow diverters have been reported with favorable clinical and angiographic outcomes in various literatures, randomized trials determining their true effectiveness and safety are still in lack. The Parent Artery Reconstruction for Large or Giant Cerebral Aneurysms Using the Tubridge Flow Diverter (PARAT) trial was designed to evaluate the safety and efficacy of the Tubridge flow diverter in the treatment of large or giant aneurysms in comparison with Enterprise stent-assisted coiling.

MATERIALS AND METHODS: This prospective, multicenter, randomized trial was conducted at 12 hospitals throughout China. Enrolled adults with unruptured large/giant intracranial aneurysms were randomly assigned (1:1) to receive either Enterprise stent-assisted coiling or Tubridge flow diverter implantation. The primary end point was complete occlusion at 6-month follow-up, while secondary end points included technical success, mortality, target vessel–related stroke, aneurysm bleeding, in-stent stenosis, parent artery occlusion, and the frequency of all adverse events.

RESULTS: Among 185 enrolled subjects, 41 withdrew before procedure initiation. Overall, 82 subjects underwent Tubridge implantation, and 62 subjects were primarily treated with stent-assisted coiling. The results of 6-month follow-up imaging included complete occlusion rates of 75.34% versus 24.53% for the Tubridge and stent-assisted coiling groups, respectively, with a calculated common odds ratio of 9.4 (95% confidence interval, 4.14–21.38; P < .001). There was a higher, nonsignificant frequency of complications for Tubridge subjects. Multivariate analysis showed a decreased stroke rate at the primary investigational site, with a marginal P value (P = .051).

CONCLUSIONS: This trial showed an obviously higher rate of large and giant aneurysm obliteration with the Tubridge FD over Enterprise stent-assisted coiling. However, this higher obliteration rate came at the cost of a nonsignificantly higher rate of complications. Investigational site comparisons suggested that a learning curve for flow-diverter implantation should be recognized and factored into trial designs.

ABBREVIATIONS: FD = flow diverter; LTF = lost-to-follow-up; PARAT = Parent Artery Reconstruction for Large or Giant Cerebral Aneurysms Using the Tubridge Flow Diverter; SAC = stent-assisted coiling

arge (10−25 mm) or giant (≥25 mm) aneurysms are a very challenging subtype among intracranial aneurysms, with

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much higher risks of rupture and poorer outcomes compared with small aneurysms.^{1,2} Despite recent technical advances, the treatment of large or giant aneurysms remains technically disappointing, with high complication and/or recanalization rates.³⁻⁶

Jin-min Liu and Yu Zhou contributed equally to this work.

The PARAT investigators are listed in the On-line Appendix.

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In recent years, various flow diverter (FD) devices, such as the Pipeline Embolization Device (Covidien, Irvine, California), the Silk flow diverter (Balt Extrusion, Montmorency, France), the Flow-Redirection Endoluminal Device (FRED; MicroVention, Tustin, California), and the Surpass stent (Stryker Neurovascular, Kalamazoo, Michigan), have been increasingly used in >50 countries. These devices were believed to improve long-term effectiveness, due to their capability to alter intrasaccular hemodynamics.⁷ Unfortunately, reported clinical results have varied significantly, with aneurysm occlusion rates and periprocedural complication rates ranging from 49% to 93.4% and 2.8% to 11%, respectively. On the other hand, there is only 1 randomized controlled trial reported until now (Flow Diversion in Intracranial Aneurysm Treatment [FIAT]), which showed very high complication rates and below-expectation effectiveness.8 These results confused neurointerventionalists about the true effectiveness and safety of FDs. The recent publication of Raymond et al⁸ echoed these sentiments by suggesting that more randomized trials are needed to determine the role of flow diversion in the management of aneurysms.

In 2012, a lack of prospective data and reports of significant adverse outcomes associated with FD use in treating intracranial aneurysms fueled our interest in leading a multicenter, prospective, randomized, controlled trial (Parent Artery Reconstruction for Large or Giant Cerebral Aneurysms Using the Tubridge Flow Diverter [PARAT]) assessing FD treatment of large and giant intracranial aneurysms. In contrast to the FIAT trial, the PARAT trial focused on unruptured large or giant internal carotid artery or vertebral artery aneurysms, the traditional indication for FD implantation. By comparing it with a well-established treatment, stent-assisted coiling (SAC), we attempted to characterize the safety and effectiveness of the Tubridge FD (MicroPort Neuro-Tech, Shanghai, China) in this specific subset of intracranial aneurysms.

MATERIALS AND METHODS

Study Design and Participants

The PARAT trial used a prospective, multicenter, parallel-group design, with balanced randomization (1:1). The primary trial purpose was to compare outcomes of subjects with unruptured large/giant intracranial aneurysms who were treated with either Enterprise SAC (Codman & Shurtleff, Raynham, Massachusetts) or Tubridge FD implantation. The study was conducted in 12 centers throughout China, each of which was required to have performed >30 stent-assisted coiling procedures per year during the previous 5 years. The PARAT trial protocol was approved by all relevant local ethics boards.⁹ Site investigators generated trial data, with monitoring and data base maintenance completed by a commercial clinical research organization. The corresponding author had full access to all trial data and had final authority for key decisions relevant to this publication.

At each trial center, prospective subjects were screened for trial eligibility, based on having an unruptured ICA or vertebral artery saccular aneurysm (including recanalized aneurysms) measuring \geq 10 mm in maximum diameter and \geq 4 mm across the aneurysm neck. Investigators recruited only those who met all inclusion criteria and none with the exclusion criteria. Patients with rup-

tured aneurysms or other intracranial diseases were excluded. Detailed inclusion and exclusion criteria are listed in On-line Table 1. Investigational sites did not record or retain a log of patients who were screened for eligibility. Written informed consent was obtained from each participant before enrollment.

Randomization and Masking

After we obtained each prospective subject's informed consent, treatment allocation was initiated by a researcher contacting a clinical research associate who was independent of the patient-recruitment procedure. Subject randomization was accomplished through an interactive Web response system, which was developed by information technology managers from an independent clinical research organization under the instruction of a clinical trial statistician. The randomization scheme included built-in stratification by the participating center, aneurysm size (≤ 15 mm versus >15 mm), and aneurysm location (anterior circulation versus posterior circulation). The on-line central randomization ensured that the allocation sequence was concealed from investigators who were recruiting patients before the decision to randomize. Treatment groups were balanced using minimization criteria.

Masking of the randomization results to local investigators or trial participants was impossible to achieve. However, members of an independent Core Laboratory and those who followed up with trial subjects via telephone or face-to-face on-site interviews were blinded as to treatment allocation.

Interventions, Treatment, and Evaluation Procedures

For all recruited patients, dual-antiplatelet drugs (300 mg/day of aspirin plus 75 mg/day of clopidogrel) were prescribed for at least 3 days before the pivotal procedure. All FD placement procedures were performed with the patient under general anesthesia and via a transfemoral approach. After sheath placement, heparin (100 u/kg) was administered to maintain an activated clotting time of 250–350 seconds throughout the procedure. Next, a suitable guiding catheter (7F for subjects to be treated with Tubridge and coils) was placed in the distal internal carotid or vertebral artery. Subsequently, treatment group subjects underwent Tubridge FD implantation alone or in combination with bare coils, while control group subjects were treated by Enterprise stent-assisted coiling with bare coils.

The Tubridge Flow Diverter, developed by MicroPort Neuro-Tech, Shanghai, China, is designed to encourage the formation of intra-aneurysmal clot, while concurrently repairing and reconstructing the parent artery. Previously, we have characterized the Tubridge FD and its structural differences from the Pipeline and Silk flow diverters, such as the platinum-iridium material used for the radiopaque microfilaments, more braided microfilaments for the large-sized FD, and a decreased incidence of shortening (Table 1).⁹ Deployment of the Tubridge FD requires a combination "push" and "pull" technique to ensure full opening of the device and to increase the metal coverage across the aneurysm neck, as previously described.⁹

The approach and timing of coil placement and the number of Tubridge devices implanted were left to the discretion of each operator. For control subject aneurysms, Enterprise stents were

Table 1: Structural comparison between different FDs

		Braided Microfilaments ^a		Radiopaque	Flared	Metal	
FD Type	Size	No.	Material	Microfilaments	End	Coverage	Retrievable
Pipeline	3–5.5 mm	48	75% cobalt chromium and 25% platinum	NA	No	30%–35%	Yes ^b
Silk	2.5–5 mm	48	Nickel-titanium alloy	4 Platinum wires	Yes	35%–55%	Yes
Surpass	2.5–5 mm	2.5 mm (36) 3 and 4 mm (60) 5 mm (84)	Cobalt-chromium	12 Platinum wires	No	30%	NA
FRED	2.5–5.5 mm	Inner layer (48) Outer layer (16)	Nickel-titanium alloy	2 Tantalum wires	Yes	NA	Yes
Tubridge	2.5–6.5 mm	<3.5 mm (46) ≥3.5 mm (62)	Nickel-titanium alloy	2 Platinum-iridium wires	Yes	30%–35%	Yes

Note:—NA indicates not applicable.

^a Braided microfilaments in this table mean those main wires excluding microfilaments especially for radiopaque usage.

^b Pipeline Flex embolization devices are retrievable.

implanted according to the instructions for use of the product, and all aneurysms were embolized (coiled) to achieve maximum packing density. All subject treatments were well-documented, and details included aneurysm shape and dimensions, width of the aneurysmal neck, diameter of the parent artery, and all relevant treatment outcomes. The postoperative antiplatelet regimen was as follows: <6 weeks: 300 mg of aspirin +75 mg of clopidogrel; 6 weeks to 3 months: 100 mg of aspirin +75 mg of clopidogrel; and, \geq 3 months: 100 mg of aspirin indefinitely.

Digital copies of angiograms, including 3D rotational angiography as well as working projection images and other anteroposterior/lateral angiographic images, were collected by the clinical research organization-assigned clinical research associate of each site. These were sent to an independent Core Laboratory for analyses by up to 3 experienced neurointerventionalists ("analysts"). On the first pass, 2 analysts separately reviewed each subject's imaging. If the initial 2 interpretations conflicted, a third analyst provided a tie-breaking assessment. Follow-up angiographic results were classified into 4 categories, according to the immediate degree of embolization: 1) occluded, defined as no contrast filling into the aneurysm sac; 2) improved, defined as decreased contrast filling into the aneurysm sac; 3) stable, defined as unchanged contrast filling into the aneurysm sac; and, 4) recanalized, defined as increased contrast filling into the aneurysm sac. Analysts also documented instances of parent artery occlusion or in-stent stenosis.

At 1, 3, and 6 months postimplantation, experienced site investigators followed up with each subject by telephone or by inperson clinical interviews. To provide some level of objectivity, these interviewers were blinded to the treatment allocation. If any adverse events were identified, investigators documented, in full detail, the following information: symptoms, event duration and severity, possible causes and associations, actions taken, and event resolution/final outcomes. A fully independent Clinical Events Committee evaluated all investigator-documented adverse events and categorized them according to the most likely causal relationship: procedure-related, device-related, disease-related, or unrelated to either disease, device, or procedure.

Study Outcomes

The primary end point was complete aneurysm occlusion at the 6-month follow-up. During reviews of 3D rotational angiography and other angiographic images, members of the Core Laboratory consistently followed precise analytic criteria for establishing the final effectiveness end point. Only those aneurysms exhibiting no contrast filling in the aneurysm were judged as meeting the complete occlusion designation. End point analysis was expressed as percentage occurrence within the test and control groups.

Secondary end points included the following:

 Immediate technical success rate, which included successful device delivery, exact stent positioning, and full expansion of the devices.

2) Death or stroke related to target vessel (assessment time points: 30 days, 90 days, and 1 year postoperation). Stroke was defined as sudden symptoms and signs of a focal cerebral function deficit associated with cerebral circulation disorders and including hemorrhagic or ischemic stroke. Hemorrhagic stroke was defined as acute extravasation of blood into the brain parenchyma or subarachnoid space with associated neurologic symptoms, whereas ischemic stroke was defined as rapid onset of a new focal neurologic deficit or rapid worsening of an existing focal neurologic deficit with clinical evidence of infarction not attributable to a nonischemic etiology (not associated with brain infection, trauma, tumor, seizure, severe metabolic disease, or degenerative neurologic disease).

3) Aneurysm bleeding rate (assessment time points: 30 days, 90 days, and 1 year postoperation), including intraoperative rupture and delayed aneurysm rupture confirmed by CT.

4) The rate of in-stent stenosis (assessment time point: 6 months postoperation).

5) The rate of parent artery occlusion (assessment time point: 6 months postoperation).

6) General adverse events (assessment time points: 30 days, 90 days, and 1 year postoperation).

Statistical Analysis

On the basis of an assumption of a 20% lost-to-follow-up (LTF) rate, a 124-subject sample size was planned (62 per group) to obtain a valid result (2-tailed test, significance level of $\alpha = .05$, power of $1-\beta = 0.80$). However, because postrandomization subject withdrawals were higher than anticipated, the independent Data Monitoring Committee of the trial suggested a larger sample size to reach the required minimum number for each group (62 subjects). Differences between the PARAT trial protocol and a final report were included in the statistical analysis plan.

Data analyses included a modified intention-to-treat approach based on a population of subjects who were recruited and treated (full analysis set). Those who left the trial immediately after randomization and who did not undergo the pivotal procedure were removed from this analysis. Those who completed treatment and the entire follow-up protocol constituted the perprotocol set, while those who were treated and had at least 1 safety evaluation were included in the safety set. To verify intergroup balance, the statisticians compared the baseline characteristics of subjects between the 2 study arms. Proportions were used for categoric variables, and medians with interquartile ranges were used for continuous variables. The categoric variables were compared between study arms using the χ^2 or Fisher exact test. For continuous variables, the *t* test or Mann-Whitney rank test, as applicable, was used according to the distribution of the data.

Analysis of the primary effectiveness end point involved a comparison of complete occlusion rates at the 6-month follow-up between the treatment and control groups in the full analysis set using the χ^2 method in combination with multivariable logistic regression adjusted for subject age (60 years or younger versus older than 60 years) and aneurysm size (\leq 15 versus >15 mm). A sensitivity analysis was performed to assess the effect of missing data (ie, those who were partially or wholly LTF after undergoing the pivotal procedure). A per-protocol set analysis was also performed excluding the following: subjects who were enrolled in the trial but who did not undergo the index procedure, those in whom the procedure failed, and those who did not reach the primary end point because they were deemed LTF. Secondary outcomes were compared between groups in the safety set using the χ^2 or Fisher exact test, as applicable. An additional multivariable logistic regression was performed to explore factors potentially affecting subject outcomes.

Statistical analyses were conducted by statisticians at the Institute of Clinical Evaluation, affiliated with Beijing University, and the data were analyzed using Statistical Analysis System software, Version 9.2 (SAS Institute, Cary, North Carolina). All tests were 2-sided, and a P value < .05 was considered statistically significant.

An independent Data and Safety Monitoring Committee had unrestricted access to trial data, to allow periodic monitoring of trial progress. This trial was registered on the Chinese Clinical Trial Registry: ChiCTR-TRC-13003127.

RESULTS

Subject Characteristics and Disposition

Between December 18, 2012, and May 9, 2014, one hundred eighty-five subjects were enrolled and randomized. The distribution of subjects among the investigational sites is shown in On-line Table 2. Of considerable surprise and prompting a midstudy revision of enrollment strategy was the high proportion of subjects (n = 41) exiting the study after randomization. Details of these LTF subjects are found in the final part of this article in a discussion of trial limitations as well as in the Figure.

Pivotal treatments were initiated in the remaining 144 subjects who formed the full analysis set based on a principle of modified intention-to-treat. Due to a tortuous parent artery or a wide aneurysm neck, investigators failed to catheterize the parent artery or deliver devices across the aneurysm neck in 6 instances (3 per trial group). Of 138 treated subjects, 5 died; 1 withdrew after serious procedure-related complications; and 6 were defined as LTF without reason. The remaining 126 subjects were included in the per-protocol analysis. There were no subject crossovers among trial groups. Treatment and follow-up details are shown in On-line Table 3.

A comparison of baseline characteristics between the 2 groups in the full analysis set showed similar distributions except for subject age (P = .036, Table 2), which was not considered a prognostic factor for the primary outcome. Because a high proportion of participants quit after randomization, we also compared baseline characteristics between the 41 excluded subjects and the 144 subjects in the full analysis set and determined that the distributions were comparable (On-line Table 4).

Primary Outcome

As shown in Table 3, in the 6-month angiographic follow-up, aneurysms treated with Tubridge FDs were associated with a favorable shift toward a complete occlusion rate. In the per-protocol analysis of 126 cases for the primary effectiveness end point, the complete occlusion rate was 75.34% (55/73) for the Tubridge group and 24.53% (13/53) for the Enterprise control group, with an adjusted common odds ratio of 9.31 (95% confidence interval, 4.00–21.66; *P* < .001). Unadjusted results and sensitivity analyses also showed superior results for the Tubridge group. Table 4 lists 6-month 100% aneurysm occlusion rates in the context of aneurysm location and size.

Secondary Outcomes

Immediate technical success rates did not differ significantly between the 2 trial groups. Three procedures in each group failed due to difficulties in advancing a microcatheter into the distal arteries. The calculated technical success rates were 96.34% (79/82) and 95.16% (59/62) in the Tubridge FD group and Enterprise control group, respectively, with a common odds ratio of 1.34 (95% CI, 0.26–6.87; P = .726).

Trial statisticians calculated a trend toward an increased but nonsignificant risk of complications for subjects in the Tubridge group. Hemorrhagic stroke occurred in 7 subjects, including 5/82 (6.1%) in the Tubridge group and 2/62 (3.23%) in the control group. Ischemic stroke, related to target vessels, occurred in 8/82 (9.76%) of the Tubridge group and 6/62 (9.68%) of the control group. These complications resulted in 4 deaths: Three were Tubridge subjects and 1 was an Enterprise-treated subject. In addition, 1 subject in each of the 2 trial groups died during follow-up for reasons unrelated to target vessels. Overall, the rates of death or stroke related to target vessels in 1-year follow up were 14.52% and 17.07% in the control and Tubridge groups, respectively. Statistically, there was no significant difference between the groups (P = .678).

The aneurysm bleeding rate was comparable between groups, with 3.66% versus 1.61% for the Tubridge group and Enterprise control group, respectively (P = .634). Overall, general adverse events occurred in 46/82 (56.10%) of the Tubridge

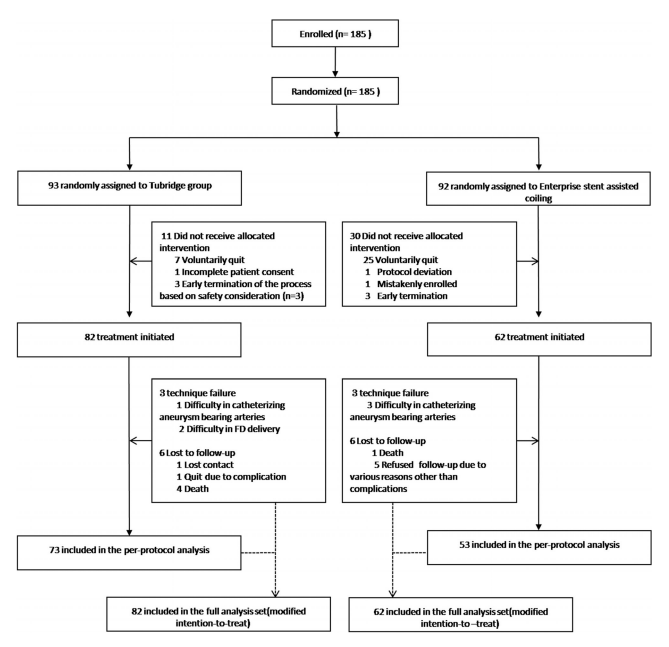


FIGURE. Flow diagram of the PARAT trial.

group and 33/62 (53.23%) of the Enterprise control group; among these, most were symptoms unrelated to the device, index procedure, or aneurysmal disease, such as headache, vomiting, or fever and there was no significant difference between the groups at the 30-day, 90-day, or 1-year follow-up. There was a trend toward a higher rate of in-stent stenosis or parent artery occlusion in the Tubridge group, but the difference was not statistically significant (Table 5).

Additional Complication Details

As shown in On-line Table 5, target vessel–related stroke, both ischemic and hemorrhagic, occurred in 21 subjects. Eight occurred during the procedure; 9, within the first 30 days after the index procedure (procedure-related); and the remaining 4 cases, at 1, 2, 3, and 7 months posttreatment.

Intraoperative bleeding occurred in 3 subjects. One Tubridge

subject had a microwire injury to a distal artery and died, while the other 2 subjects with intraoperative bleeding (1 per group/mRS of 1 at follow-up) recovered well after prompt coil deployment immediately after the onset of rupture. The Clinical Events Committee concluded that these complications, though unrelated to the Tubridge and Enterprise devices per se, were classified as procedure-related complications.

Periprocedural bleeding occurred in 4 cases (3 Tubridge/1 Enterprise). Delayed aneurysm rupture accounted for 2 complications (both large or giant aneurysms treated with FD alone). Of the remaining 2 cases, 1 Enterprise subject presented with cerebral hematoma and 1 Tubridge subject presented with mild subarachnoid hemorrhage in the Sylvian fissure. Both were thought to be unrelated to aneurysm rupture. The patient with a cerebral hematoma died due to prehospital delay, while the other one recovered well.

Overall, there were 14 cases of target vessel-related ischemic

stroke (8 Tubridge/6 Enterprise). Five subjects experienced intraoperative ischemia; another 5, within the first 30 days; and the remaining 4, at 1-7 months posttreatment.

Among those with intraoperative ischemic events, 3 were Tubridge subjects. One had a stroke after anterior choroidal artery occlusion due to coil introduction; another had a frontoparietal acute infarction due to parent artery occlusion caused by intrastent thrombosis; and, the other one was thought to be the result of a thromboembolic event. All of them resulted in contralateral limb weakness, and symptoms resolved in 1 case. In the 2 control subjects, patients had ischemic symptoms after uneventful procedures. Thromboembolic events were considered after further CT and angiographies, and both recovered well after medical therapy.

Among 5 subjects with periprocedural thromboembolic events (2 Tubridge/3 Enterprise), ischemic symptoms were all identified within 1 week after the procedure. Symptoms resulting from these periprocedural ischemia events were characterized as

Table 2: Baseline characteristics of the full analysis set^a

Characteristics

Large (10-15 mm)

Very large or giant (>15 mm)

period.

Overall, among all these complications, there were 4 deaths and 2 disabilities identified during the last follow-up. The procedure-related mortality rate was 3.66% (3/82) in the Tubridge group versus 1.61% (1/62) in the Enterprise group, while the procedure-related morbidity rate was 2.4% (2/82) in the Tubridge group versus zero in the control group in 1-year follow-up.

Multivariate Analysis

To further explore the factors affecting angiographic outcomes and complication occurrences, trial statisticians perned multivariate analysis, including iect age, aneurysm size, aneurysm loon (anterior circulation versus posor circulation), and treatment techue (FD alone; FD + coils; and erprise stent-assisted coiling). The nary investigational site (Shanghai nghai hospital) showed a decreased oke rate compared with other trial ters, with a marginal P value (OR = ; 95% CI, 0.991-23.335; P = .051).

DISCUSSION

Since the introduction of flow diverters to the global neurovascular community,

^a Data are No. (%), median (interquartile range), or mean (SD)

^b Of the 6 patients with failed procedures, locations of 2 aneurysms were absent.

Table 3: Sensitivity analyses on the composite primary outcome

	6-Mo Occlusion Rate		U	Unadjusted Result		Adjusted Result ^a		
	Tubridge Control			Р				Р
Factoring in Subjects Who Missed F/U	Group	Group	OR	95% CI	Value	OR	95% CI	Value
Missing F/U excluded (PPS)	55/73, 75.34%	13/53, 24.53%	9.40	4.14–21.38	<.001	9.31	4.00–21.66	<.001
Missing F/U counted as complete occlusion	64/82, 78.05%	22/62, 35.48%	6.47	3.09–13.52	<.001	6.14	2.92–12.91	<.001
Missing F/U counted as not completely occluded	55/82, 67.07%	13/62, 20.97%	7.68	3.57–16.51	<.001	7.34	3.36–16.05	<.001
Missing F/U counted as complete occlusion for	55/82, 67.07%	22/62, 35.48%	3.70	1.85–7.42	<.001	3.49	1.73–7.03	<.001
control and incomplete occlusion for Tubridge								

Note:-F/U indicates follow-up; PPS, per-protocol set.

Adjusted for each subject's age and aneurysm size (\leq 15 versus >15 mm).

Table 4: Data correlating aneurysm location and size with aneurysm complete occlusion rate

			Proportion of	100% Occlusion Rate at 6-Mo F/U	
Aneurysm Location	No.	Aneurysm Size (mm)	Giant Aneurysms	Tubridge	Controls
ICA communicating	19	18.0 ± 7.6 (10.0–33.9)	4/19	76.9%, 10/13	40%, 2/5
ICA ophthalmic	39	15.8 ± 5.6 (10.0–30.0)	4/39	83.3%, 15/18	33.3%, 6/18
ICA paraclinoid or cavernous	73	21.8 ± 7.5 (10.0–45)	25/73	75.7%, 28/37	13.8%, 4/29
ICA petrous	3	21.3 ± 9.3 (15.0–32.0)	1/3	50%, 1/2	0
Vertebral artery	4	14.3 \pm 2.2 (12.0–16.3)	0/4	33.3%, 1/3	100%, 1/1

Note:-F/U indicates follow-up.

Mean age (yr)	52.11 (10.31)	55.66 (9.53)
Sex ratio (male/female)	21:61	13:49
Medical history		
Cerebrovascular stroke	9 (10.98%)	8 (12.90%)
Coronary artery disease	1 (1.22%)	4 (6.45%)
Hypertension	41 (50.0%)	25 (40.32%)
Hypercholesterolemia	4 (4.88%)	3 (4.84%)
Diabetes	5 (6.10%)	2 (3.23%)
Current/previous smoking	12 (14.63%)	7 (11.29%)
Systolic blood pressure (median) (mm Hg)	129.50 (120.0–143.0)	129.50 (120.0–138.0)
Aneurysm location ^b		
Anterior circulation	77 (96.25%)	59 (98.33%)
Posterior circulation	3 (3.75%)	1 (1.67%)
Aneurysm size (median)	18.00 (13.14–26.0)	17.14 (12.44–24.74)
Aneurysm size classification		

30 (36.59%)

52 (63.41%)

Tubridge Group

n = 82

Enterprise Group

n = 62

24 (38.71%)

38 (61.29%)

mild and transient, with all events resolving well after aggressive antiplatelet therapy.

There were no hemorrhagic events after the periprocedural period. However, delayed ischemic stroke occurred in 4, including 3 Tubridge subjects and 1 Enterprise subject. For the 3 Tubridge subjects, 2 strokes were identified with parent artery occlusion after further examination and the patients were treated conservatively; and the other 2 patients (1 per group) had ischemic symptoms 1 and 2 months postprocedure, with patent parent arteries. All these symptoms improved during the follow-up

Table 5: Secondary outcome evaluation

	Enterprise	Tubridge		
Secondary Outcome	n = 62	n = 82	Odds Ratio (95% CI)	P Value
Immediate technique success rate	59/62 (95.16%)	79/82 (96.34%)	1.34 (0.26–6.87)	.726
Death or target vessel–related stroke				
30 days	7/62 (11.29%)	11/82 (13.41%)	0.821 (0.299–2.258)	.703
90 days	8/62 (12.90%)	12/82 (14.63%)	0.864 (0.330-2.263)	.766
1 yr	9/62 (14.52%)	14/82 (17.07%)	0.825 (0.332–2.051)	.678
Mortality rate				
30 days	1/62 (1.61%)	3/82 (3.66%)	0.432 (0.044-4.253)	.634
90 days	1/62 (1.61%)	4/82 (4.88%)	0.320 (0.035–2.934)	.391
1 yr ^a	2/62 (3.23%)	4/82 (4.88%)	0.650 (0.115–3.668)	.699
Hemorrhagic stroke related to target vessel				
30 days	2/62 (3.23%)	5/82 (6.1%)	0.531 (0.096–2.738)	.699
90 days	2/62 (3.23%)	5/82 (6.1%)	0.531 (0.096–2.738)	.699
1 yr	2/62 (3.23%)	5/82 (6.1%)	0.531 (0.096–2.738)	.699
Ischemic stroke related to target vessel	,		, , , , , , , , , , , , , , , , , , ,	
30 days	4/62 (6.54%)	6/82 (7.32%)	0.874 (0.236–3.239)	.874
90 days	5/62 (8.06%)	6/82 (7.32%)	1.111 (0.323–3.822)	.867
1 yr	6/62 (9.68%)	8/82 (9.76%)	0.991 (0.325–3.109)	.987
Aneurysm rupture	,		, , , , , , , , , , , , , , , , , , ,	
30 days	1/62 (1.61%)	3/82 (3.66%)	0.357 (0.072–1.783)	.634
90 days	1/62 (1.61%)	3/82 (3.66%)	0.357 (0.072–1.783)	.634
1 yr	1/62 (1.61%)	3/82 (3.66%)	0.357 (0.072–1.783)	.634
General adverse events				
30 days	26/62 (41.94%)	42/82 (51.22%)	0.808 (0.293-2.229)	.269
90 days	26/62 (41.94%)	42/82 (51.22%)	0.808 (0.293–2.229)	.269
l year	33/62 (53.23%)	46/82 (56.10%)	0.938 (0.644–1.365)	.88
Rate of intrastent stenosis	2/53 (3.77%)	4/73 (5.48%)	0.676 (0.119–3.837)	1
Rate of intrastent thrombosis	1/59 (1.69%)	6/79 (7.59%)	0.215 (0.025–1.839)	.273

^a Two subjects died during follow-up for reasons unrelated to the target vessels (1 each in Tubridge and control groups).

many clinical studies were conducted to evaluate the safety and effectiveness of FDs, such as the Pipeline for Uncoilable or Failed Aneurysms (PUFS), Aneurysm Study of Pipeline in an Observational Registry (ASPIRe), and PITA trials.8,10-18 Additional effort to evaluate the effectiveness and safety of FDs versus conventional treatments included the Multicenter Randomized Trial on Selective Endovascular Aneurysm Occlusion with Coils versus Parent Vessel Reconstruction using the SILK Flow Diverter (MARCO POLO), FIAT, Flow Diverter Stent for Endovascular Treatment of Unruptured Saccular Wide-necked Intracranial Aneurysms (EVIDENCE), and Complete Occlusion of Coilable Aneurysms (COCOA) trials.¹⁹⁻²² However, the number of studies that focused on FD treatment of large or giant intracranial aneurysms is low. The purpose of the PARAT trial was to compare the safety and effectiveness outcomes in the treatment of ICA or vertebral artery large or giant aneurysms with the Tubridge FD (with and without coiling) versus a more conventional approach using Enterprise SAC. The PARAT trial results suggested that in subjects with ICA or vertebral artery large or giant aneurysms, Tubridge FD implantation had a significantly higher 6-month complete occlusion rate compared with conventional stent-assisted coiling. However, there was a trend toward increased risk of stroke with FD implantation.

The treatment of large or giant intracranial aneurysms has evolved significantly during the past few years. Before the emergence of FDs, parent artery occlusion, coiling alone, and stentassisted coiling were the major treatment modalities for intracranial large or giant aneurysms. According to a meta-analysis by Turfe et al,²³ parent artery occlusion can result in a complete occlusion rate of 93.0% (95% CI, 86.0%–97.0%). However, parent artery occlusion may be a viable treatment option only when there is sufficient compensating blood flow. Even when a balloon occlusion test finding is negative, a 4%-15% complication rate is possible.²³ In addition, there have been concerns about de novo aneurysm occurrence after carotid occlusion.²⁴ In a report by Arambepola et al,²⁵ 4.3% of patients developed de novo aneurysms within a mean of 9 years. Bypass surgery may reduce the incidence of ischemic events, but the procedure may be too complicated, leading to morbidity and mortality rates as high as 7% and 13%, respectively.²⁶⁻²⁸ In this trial, parent artery occlusion was not selected as a feasible control treatment because parent artery sacrifice is considered, throughout most hospitals in China, a last and somewhat futile option for treating intracranial aneurysms. Thus, only those cases posing considerable difficulty or possible failure in parent artery reconstruction would be treated with this method as a salvaging effort. Other than with simple and uncomplicated cases, coiling of large or giant aneurysms without a stent is undertaken far less frequently because of anticipated high recanalization rates.²³

In many prospective multicenter studies, complete occlusion rates at final follow-up varied from 49% to 93.4%, and 6-month complete occlusion rates ranged from 55.7% to 93.3% (On-line Table 6). In the treatment of large or giant aneurysms with FDs, Becske et al¹⁸ reported a complete occlusion rate of 76.4% at 180 days. Chalouhi et al²⁹ reported a complete occlusion rate of 86% at a median angiographic follow-up of 7 months. On the basis of a meta-analysis of 29 studies, including 1451 patients with 1654 aneurysms, Brinjikji et al³⁰ reported that the complete occlusion rate after FD implantation was 74% (95% CI, 63%–83%) for large aneurysms and 76% (95% CI, 53%–90.0%) for giant aneurysms.

Generally, the complete occlusion rate of 75.3% in the Tubridge group at 6 months seems comparable with rates reported in previous publications. The complete occlusion rate of 24.5% for the control group is lower than that reported in previously published articles. In 2 relatively large studies involving large or giant aneurysms, complete occlusion rates of 31% and 41% were reported by Sluzewski et al⁵ and Chalouhi et al,²⁹ respectively. The discrepancies between the latter reports and other published results, as noted above, may be related to variations in patient-selection criteria, study methods, trial design, aneurysm characteristics, and evaluation specifications and procedures. Nevertheless, the effectiveness of Tubridge FD treatment appears to be superior to that of conventional stent-assisted coiling and comparable with reported effectiveness outcomes for other FDs.

Other concerns after FD implantation include technical success rates and overall safety, as well as how FD treatment compares with conventional treatment options. The Enterprise stent is thought to be a safe device for aneurysm treatment, and hemorrhagic complications are assumed to be uncommon.³¹ Although exact causes are not well-understood, delayed aneurysm rupture or intraparenchymal hemorrhage after FD implantation has been recognized.

As shown in On-line Table 6, the hemorrhagic stroke and aneurysm rupture rates were as high as 6.9% and 5.2%, respectively, as reported for multicenter prospective studies. In the International retrospective study of the Pipeline embolization device (IntrePED) study, capturing data from 793 patients among 17 centers, the intraparenchymal hemorrhage rate was 2.4%, while the aneurysm rupture rate was 0.6%.³² In a recent meta-analysis of 3125 treated subjects, the calculated intraparenchymal hemorrhage and aneurysm rupture rates were 2.9% and 1.8%, respectively. These complications are customarily thought to be higher in large or giant aneurysms.33 Calculated intraparenchymal hemorrhage and aneurysm rupture rates were 5.4% and 7.5%, respectively, in giant aneurysms, and 2.1% and 1.3%, respectively, in small and large aneurysms. The IntrePED study generated similar results, with intraparenchymal hemorrhage and aneurysm rupture rates of 5.8% and 5.8%, respectively, in giant ICA aneurysms; 2.6% and 0.5%, respectively, in large ICA aneurysms; and, 1.9% and 0%, respectively, in small ICA aneurysms. We observed an overall hemorrhagic rate of 6.1% (5/82) in the Tubridge group, of which 2 hemorrhagic occurrences were thought to be the result of procedural injury. Excluding these 2 cases, 2/82 (2.4%) manifested as aneurysm rupture. We did not encounter intraparenchymal hemorrhage, but we observed 1 (1.2%) lateral Sylvian fissure SAH, unrelated to aneurysm rupture. The results of our study are very comparable with those of the above-described studies.

The prospect of ischemic stroke presents additional concerns. A complication rate of 9.68% in the Enterprise control group was consistent with that reported previously by Chalouhi et al,²⁹ as well as our own single-center experience, in the range of 7.58%–11.4%.³⁴ Overall, ischemic rates after FD implantation vary among multiple publications, with a rate of 0%–10.3%. Two recent meta-analyses indicated ischemic rates ranging from 5.5% to 7.5%.^{33,35} However, the occurrence was increased in large or giant aneurysms due to intra-aneurysmal thrombosis or a prolonged procedure time, with a rate ranging from 5.2% to 13.5%.³² Ye et

al³³ indicated an ischemic rate of 9.5% for giant aneurysms. Among Tubridge subjects of the PARAT trial, ischemic complications occurred in 8 (9.76%) subjects. Although the PARAT trial rates are comparable with reported thromboembolic rates in published studies, a 9%-10% incidence of ischemic complications should be anticipated when using FDs for large or giant intracranial aneurysms. In the FIAT study, 12/75 subjects (16.0%; 95% CI, 8.9%–26.7%) treated by flow diversion were dead (n = 8) or dependent (n = 4) at ≥ 3 months postimplantation.⁸ These results differ from those in our previous experience with FDs. We believe this discrepancy may reflect "real world" early experience with flow diverters for large or giant aneurysms in multiple centers with adjudication by an external imaging Core Laboratory. Similar reports of low procedure-related risks are seen commonly with single-center or retrospective studies, as best exemplified by the recent the Stenting vs. Aggressive Medical Management for Preventing Recurrent Stroke in Intracranial Stenosis (SAMMPRIS) trial.³⁶

To better understand why these complications occurred, we undertook a detailed review of the PARAT trial complications. One possible factor is the role of the learning curve with flow diversion, and physicians' experience may still play a role in determining the relative safety of FD therapy. The prospect of a learning curve not only encourages a higher skill level in device handling and stent deployment but also leads to more appropriate subject selection. After gaining more experience with a new flow diverter in the treatment of large or giant aneurysms, physicians can avoid many technical failures, make more appropriate FDsize selections, better understand the necessity of appropriate postdeployment balloon dilation to avoid poor apposition or overdilation, and better identify the need for simultaneous coil insertion in the treatment of large or giant aneurysms. As an example, the PARAT trial experienced high rates of parent artery occlusions, and these contributed to 3 ischemic events in the Tubridge FD group. The possibility of malapposition may exist when the devices are underdeployed. In such instances, balloon dilation after initial FD deployment may protect against in-stent thrombosis and parent artery occlusion.

The concept of a PARAT trial learning curve was also supported by our multivariate analysis, which showed significantly lower complication rates in the primary investigational site, where the leading physicians already had gained considerable experience with the Tubridge FD in a previous single-center study. We believe that intensive training of inexperienced physicians on all flow diverters should be advocated before launching a study or introducing use in general neurointerventional practice. We believe that the role that learning curves can play in trials should be increasingly emphasized.

Several recent studies have suggested that antiplatelet regimens may play a role in the occurrence of hemorrhagic or thrombotic complications.^{37,38} In the PARAT trial, investigator management of antiplatelet therapy and subject compliance with prescribed study medications were not investigated.

We have identified some key limitations of the PARAT trial. The most obvious one is the high number of postconsent subject withdrawals (n = 41) in both arms, highlighting the delicate balance between physicians and subjects as to what treatment might be better suited for optimal outcomes in the treatment of highly challenging aneurysms. Seven prospective Tubridge subjects voiced their concerns about the potential adverse effects of flow-diverter use, while 15 subjects did not want to undergo stent-assisted coiling because they feared that the long-term durability was inferior to that anticipated with flow diverters. These subjects preferred to withdraw from the trial and wait to be treated with a commercially available flow diverter. The concept of new or apparently complicated technology can be intimidating for some patients.

Among the remaining subjects who departed from this trial prematurely, 1 dropped out after an incomplete subject consent, 6 had their index procedure terminated early after angiography examination indicated safety concerns, 1 was enrolled mistakenly, 1 violated the trial protocol and did not qualify for the index procedure, and 10 were terminated or withdrew from the trial for unknown reasons. Incomplete trial participation by the abovementioned subjects led to a need to expand the sample size and prevent unbalanced subject numbers and age. We compared the baseline data of subjects who were LTF with those from the full analysis set and determined that their removal had not affected the overall distribution of characteristics. Moreover, although subject age was not thought to affect outcomes, we used multivariable logistic regression, adjusted for subject age, to eliminate the effect of this imbalance. Second, the results of this study only represent the treatment of a specific subtype of aneurysms and a preliminary application of a novel FD. As discussed above, increasing familiarity with the Tubridge device is expected to lower the rate of clinical complications. Finally, although we included treatment of vertebral artery aneurysms in this trial, only 4 subjects with these aneurysms were enrolled. Thus, generalizations about the use of flow diverters in vertebral arteries should be made with caution. Clearly, additional vertebral artery aneurysm studies are warranted.

CONCLUSIONS

This trial showed that there was a significantly higher obliteration rate of aneurysms treated with the Tubridge FD versus conventional Enterprise SAC treatment in a selected group of patients with large or giant intracranial aneurysms. However, this higher obliteration rate comes at a cost of nonsignificant higher complication rates. For inexperienced operators who are either entering flow-diverter clinical trials or preparing for real-world clinical use, the importance of a learning curve should be emphasized.

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