Balloon-mounted stenting for ICAS in a multicenter registry study in China: a comparison with the WEAVE/WOVEN trial

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ABSTRACT

Background The outcome of deploying balloon-mounted stents for symptomatic intracranial atherosclerotic stenosis (ICAS) has not been fully investigated. In this study we evaluate the safety and long-term outcome of using balloon-mounted stents to treat symptomatic ICAS in comparison with the WEAVE/WOVEN study.

Methods In a multicenter registry study of stenting for symptomatic intracranial artery stenosis in China, 159 patients treated with an intracranial balloon-mounted stent approved by the China Food and Drug Administration were evaluated. The morphological features of the lesions were categorized by Mori classification. The endpoints, including periprocedural and long-term clinical and radiological outcomes, were the same as those in the WEAVE/WOVEN study.

Results In the present study the mean percent stenosis before and after stenting was 84.0% and 6.1%, respectively. The proportions of Mori A, Mori B, and Mori C lesions were 33.3%, 52.2%, and 14.5%, respectively. The 72-hour rates of stroke and mortality after the procedure were 0%. The 1-year rates of any stroke, ischemic stroke, hemorrhagic stroke, and death were 6.3% (10/159), 5.7% (9/159), 0.6% (1/159), and 0.6% (1/159), respectively. The 1-year rate of in-stent restenosis (ISR) was 23.4% (15/64). The rate of ISR in Mori C lesions (53.8%, 7/13) was significantly higher than that in Mori A (15.8%, 3/19) or Mori B lesions (15.6%, 5/32) (p = 0.024).

Conclusions The short-term and long-term outcomes of using a balloon-mounted stent for symptomatic ICAS with focal and non-angular lesions (Mori A and B type) and smooth arterial access were comparable to the results of the WEAVE/WOVEN trial.

INTRODUCTION

Stroke is a common and disabling disease worldwide, and intracranial atherosclerotic stenosis (ICAS) is one of the leading causes of ischemic stroke in Asian populations.1–3 Intracranial stenting has been evaluated for the treatment of patients with symptomatic ICAS unresponsive to aggressive medical management, to improve the cerebral perfusion irrigated by the culprit arteries and thereby reducing the risk of recurrent stroke and improving neurological outcome.4 Even though registry studies demonstrated low rates of perioperative complications (4.5–6.9%) for ICAS stenting using the self-expanding Wingspan stent system,5,6 high perioperative complication rates in the stenting group were reported by the SAMMPRIS (Stenting and Aggressive Medical Management for Preventing Recurrent Stroke in Intracranial Stenosis) study (14.7%) and by the VISSIT (Vitesse Intracranial Stent Study for Ischemic Stroke Therapy) study (24.1%).7,11 The WEAVE (Wingspan Stent System Post Market Surveillance) and WOVEN (Wingspan One-year Vascular Events and Neurologic outcomes) study also demonstrated low rates of perioperative complications (2.6%) and long-term recurrent stroke (8.5%) for ICAS stenting using the self-expanding Wingspan stent system with experienced interventionalists and proper patient selection.12,13

Compared with the self-expanding Wingspan stent, balloon-mounted stents are placed at the same time as angioplasty, avoiding the time and risk of over-the-wire exchange. In addition, balloon-mounted stents have shorter cone tips and are easier to navigate through small and tortuous vessels distal to the target lesions, reducing the risks of vascular injury. Since the failed VISSIT study, studies on the feasibility and long-term efficacy of balloon-mounted stenting for symptomatic ICAS have been scarce.11 The Apollo stent (MicroPort Neuro Tech, Shanghai, China) is a balloon-mounted stent with a metallic surface area of about 14.0%, approved by the Chinese Food and Drug Administration for use in patients with symptomatic ICAS. Although several case series have supported its safety and efficacy since 2003,14–18 a prospective multicenter registry study has not been carried out.

We have previously reported the short-term and long-term outcomes of a prospective multicenter registry on intracranial angioplasty and stenting for symptomatic stenosis, which included both self-expanding and balloon-mounted stents.
Many balloon-mounted stents were deployed for focal and non-angulor lesions (Mori A and B type) with smooth arterial access. We therefore performed this study to re-evaluate the safety and long-term outcome of balloon-mounted stenting for symptomatic ICAS by using the same endpoints as the WEAVE/WOVEN study.

**MATERIALS AND METHODS**

**Study design and population**

This study is a retrospective review of a subgroup of patients from a prospective single-arm registry study with 20 participating sites. Details of the registry have been published elsewhere. Approval by each site’s institutional review board or ethics committee was obtained. Written informed consent was obtained from the patients or their legally authorized representatives. All reported endpoints were evaluated and confirmed by a central adjudication committee consisting of neurologists, neurosurgeons, and radiologists who were blinded to the treatment choices. An independent data and safety monitoring board oversaw the conduct, safety, and efficacy of the study.

In this subgroup analysis, we focused on the clinical and radiological outcomes of the balloon-mounted stenting group in comparison with the WEAVE/WOVEN study. We used the same primary endpoints as in the WEAVE/WOVEN study, including any stroke, ischemic stroke, hemorrhagic stroke, and death within 72 hours and at 1 year after the procedure and restenosis at 1 year.

**Enrollment of patients**

Inclusion and exclusion criteria were established by the executive committee. Patients were aged 18–85 years and had a symptomatic ICAS of 70–99% with a lesion length of ≤1.5 mm and target vessel diameter of ≥2.0 mm in the intracranial internal carotid artery (ICA), middle cerebral artery (MCA), intracranial vertebral artery (VA), or basilar artery (BA) (table 1). The measurements were made on digital subtraction angiography (DSA) using the WASID (warfarin–aspirin symptomatic intracranial disease) method with normal distal vessels as the reference. The symptoms included transient ischemic attacks (TIA) or ischemic stroke within the past 90 days that were attributable to hypoperfusion in the territory of the target lesion. Ischemic stroke referred to transient ischemic attacks (TIA) or ischemic stroke or hemorrhagic stroke, and death within 72 hours and at 1 year after the procedure and restenosis at 1 year.

<table>
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<td>Symptomatic ICAS of 70–99%</td>
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<td>Age 18–85 years</td>
<td>Age 22–80 years</td>
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<td>Baseline mRS score ≤3</td>
<td>Baseline mRS score ≤3</td>
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<td>Stenting of the lesion ≥3 weeks after the last stroke</td>
<td>Stenting of the lesion ≥8 days after the last stroke</td>
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<td>Ischemic symptoms within the past 90 days attributable to hypoperfusion in the territory of the target lesion</td>
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American Society of Interventional and Therapeutic Neuroradiology/Society of Interventional Radiology Collateral Flow Grading System score of ≤3 on DSA; systolic blood pressure of ≥200 mmHg and ≤100 mmHg; and (4) hemodynamic ischemic lesion by MRI defined as small ischemic infarcts in a watershed distribution in the culprit vessel territory. The ischemic mechanisms that could be entirely explained as an embolic phenomenon or perforator occlusion were excluded from this registry. The images were centrally reviewed by at least two physicians, who were allowed to resolve the disagreement through discussion. The patients were excluded from the study if the raters could not agree on the classification.

Patients were excluded if they had acute infarcts within 3 weeks, severe vessel tortuosity precluding the employment of endovascular devices as determined by the executive committee, a non-atherosclerotic lesion confirmed by high-resolution MRI (HR-MRI), embolic or perforator stroke based on MRI or CT, or baseline modified Rankin Scale (mRS) score of >3. Only patients without risk factors for intracranial atherosclerosis or those with a lesion suspected to be non-atherosclerotic by regular CT, MRI, or DSA were subjected to HR-MRI. All clinical and imaging data were reviewed centrally by the executive committee to decide whether the patient was eligible for enrollment.

**Endovascular treatment strategy and device selection**

The procedures were performed under local or general anesthesia by experienced neurointerventionists (performing more than 100 intracranial endovascular procedures per year). Intravenous heparin was administered after the placement of vascular access using a bolus of 75 U/kg followed by half of the dose after an hour. Two endovascular strategies were used when treating ICAS, and the operators were instructed to choose the devices based on the following guideline, taking into consideration their experience and preference as well as selecting the device which they thought was best suited to the patient. The ICAS lesions were categorized into three types according to the Mori classification: Mori A, short (≤5 mm in length) concentric or moderately eccentric lesions with ready accessibility; Mori B, tubular (5–10 mm in length) extremely eccentric lesions with moderately angulated segment (≥45°, <90°) and moderate tortuosity of the proximal segment; and Mori C, diffuse lesions (>10 mm in length), extremely angulated (≥90°) lesions with excessive tortuosity of the proximal segment (figure 1). For patients with smooth arterial access and a Mori A lesion, the Apollo balloon-mounted stent was preferred. For patients with tortuous arterial access or a Mori C lesion or a lesion with a significant mismatch in the diameter between the proximal and distal segments, balloon dilatation plus a self-expanding stent (Gateway balloon plus Wingspan stent system; Stryker, Maple Grove, Minnesota, USA) was preferred, but the balloon-mounted stent could also be used based on the operators’ experience. Mori C lesions tend to...
occlusion was treated with a 2.5 mm × 13 mm Apollo stent. Five-year DSA suggested no significant in-stent restenosis. (C) A patient presented with weakness of the left limbs for 48 days. A basilar arterial stenosis classified as Mori C type was treated with a 2.5 mm × 13 mm Apollo stent. Five-year CTA suggested no significant in-stent restenosis.

Figure 2 (A) A patient presented with weakness of the right limbs for 70 days. A right middle cerebral arterial stenosis classified as Mori A type, which was treated with a 2.5 mm × 8 mm Apollo stent. One-year DSA suggested no significant in-stent restenosis. (B) A patient presented with dizziness for 48 days. A basilar arterial stenosis classified as Mori B type was treated with a 2.5 mm × 8 mm Apollo stent. One-year computed tomography angiography (CTA) suggested no significant in-stent restenosis. (C) A patient presented with weakness of the left limbs for 36 days. A right middle cerebral arterial stenosis classified as Mori C type was treated with a 2.5 mm × 8 mm Apollo stent. One-year CTA suggested no significant in-stent restenosis.

Periprocedural management
Perioperative systolic blood pressure was kept between 100 and 120 mmHg during the procedure and for 3 days after the procedure. Non-contrast head CT was obtained to exclude hemorrhage after the procedure. All patients were given a weight-based dose of 0.4–0.6 mL nadroparin (Fraxiparine; Sanofi Winthrop Industry) every 12 hours subcutaneously for 3 days and monitored closely until discharge.

Medical treatment
All patients received aspirin (100 mg/day) and clopidogrel (75 mg/day) for more than 5 days before the procedure. Resistance testing for antiplatelet drugs was not performed. Other aggressive medical therapy was implemented to achieve the following goals: systolic blood pressure <140 mmHg (or <130 mm Hg in patients with diabetes mellitus), low-density lipoprotein <70 mg/dL (1.81 mmol/L) or a decrease by 50%, smoking cessation, lifestyle modification for obesity and sedentary state.

Clinical and radiological outcomes
Follow-up information on clinical outcomes was collected and reviewed by trained personnel who were blinded to treatment assignment at study entry, the day of discharge, 30-day follow-up, and face-to-face interview every 3 months. All follow-up visits were in person unless the patient could not return for a visit, in which case telephone follow-up was completed. If necessary, brain imaging studies including MR angiography (MRA) and CT angiography (CTA) were obtained in patients who developed neurological symptoms. DSA was recommended to patients at 12 months follow-up.

In this subgroup analysis we focused on the short-term outcomes including any stroke, ischemic stroke, hemorrhagic stroke, and death within 72 hours after stenting, and long-term outcomes including any stroke, ischemic stroke, hemorrhagic stroke, and death within 1 year. The definitions of ischemic stroke, hemorrhagic stroke, and TIA were the same as in the previous studies. For patients evaluated with DSA, in-stent restenosis (ISR) was defined as >70% stenosis within or immediately adjacent (within 5 mm) to the implanted stent (figure 2). For patients evaluated with CTA, the stents were considered as ISR if the stented segment or the proximal and distal parent vessel could not be well visualized or showed an apparent filling defect on CTA (figure 1).

Statistical analysis
Continuous variables were presented as mean±SD (normal distribution data) or median with interquartile range (skewed distribution data), as appropriate. Categorical variables were presented as number and percentage. The χ² test was used to analyze differences between the present study and the WEAVE/WOVEN study and differences in ISR rates among patients with different Mori types of lesion and between patients with anterior and posterior circulation lesions. The statistical analysis was performed using a commercial statistical software package (SPSS for Windows, Version 25.0, IBM-SPSS, Chicago, Illinois, USA).

RESULTS
Patient baseline characteristics
From September 2013 to January 2015 a total of 300 patients were recruited in 15 sites in China, including 159 patients treated with balloon-mounted stenting and 141 patients treated with balloon dilatation plus self-expanding stenting. In this study the 159 patients (aged 59.4±9.5 years) treated with balloon-mounted stenting (recruited in 14 sites) were analyzed, including 120 (75.5%) men and 39 (24.5%) women. Of the 159 patients, 110 (69.2%) had hypertension, 53 (33.3%) had diabetes...
mellitus, 60 (37.7%) had hyperlipidemia, and 45 (28.3%) were current smokers. The treated lesions were located in the ICA in 25 cases (15.7%), MCA in 38 cases (23.9%), VA in 48 cases (30.2%), and BA in 48 cases (30.2%). The main qualifying event included stroke in 52.8% and TIA in 47.2% of the patients. The median time to stenting from the qualifying event was 20 days (12 days for TIA and 26 days for stroke). The baseline characteristics are shown in table 2.

### Periprocedural characteristics

Of the 159 patients, 88 (55.3%) were operated under general anesthesia and 71 (44.7%) under local anesthesia. The proportions of Mori A, Mori B, and Mori C lesions were 33.3% (53/159), 52.2% (83/159), and 14.5% (23/159), respectively. The mean percent stenosis before and after stenting was 84.0% and 6.1%, respectively, and the median percent stenosis before and after stenting was 85.0% and 5.0%, respectively. The mean stent diameter and stent length were 2.8±0.4 mm and 10.2±2.9 mm, respectively, and the median stent diameter and stent length were 2.5 mm and 8.0 mm, respectively. The technical procedural characteristics are shown in table 3.

### Clinical and radiological outcomes

In the present study the stroke rates at 72 hours, 30 days, and 1 year after the procedure were 0% (0/159), 2.5% (4/159), and 6.3% (10/159), respectively; the rates of ischemic stroke at 72 hours, 30 days, and 1 year after the procedure were 0% (0/159), 2.5% (4/159), and 6.3% (10/159), respectively; and the mortality rates at 72 hours, 30 days, and 1 year after the procedure were 0% (0/159), 2.5% (4/159), and 5.7% (9/159), respectively; the rates of hemorrhagic stroke at 72 hours, 30 days, and 1 year after the procedure were 0% (0/159), 2.5% (4/159), and 6.3% (10/159), respectively; the rates of ischemic stroke in lesions of the anterior and posterior circulation were 26.7% (4/15) and 22.4% (11/49) respectively.

### DISCUSSION

The VISSIT study reported high rates of perioperative complications (24.1%) and 1-year recurrent stroke or hard TIA (36.2%) for ICAS stenting using a balloon-mounted stent. We found that the short-term and long-term outcomes of balloon-mounted stenting for symptomatic ICAS with focal and non-angular lesions (Mori A and B type) and smooth arterial access were comparable to the results of the WEAVE/VOVEN trial.
Balloon-mounted stents have been used in symptomatic ICAS treatment since 2001. Initially, off-label coronary balloon-mounted stents (including bare stents and drug-eluting stents) were used, but the rigidity of the delivery system impeded their clinical application. The Apollo balloon-mounted stent was designed for treating ICAS lesions and has been approved for use in China since 2004. The Apollo stent system offers 40 different stents of various combinations of six different lengths (8, 10, 13, 15, 18, and 23 mm) and seven different diameters (2.5, 2.75, 3.0, 3.5, 4.0, 4.5, and 5.0 mm). Other parameters of the Apollo stent system are as follows: the metallic surface area is about 14.0%; the nominal expansion pressure is 6 atm; the length of the delivery system is 1450 mm; the maximum guide wire diameter is 0.014 inch; and the minimum guide catheter is 6 French. The delivery system has a longer distance from the tip to the rapid-exchange outlet (300 mm) than that of the coronary stent systems (usually about 250 mm). Moreover, the Apollo balloon material (Pebax 70D) is softer than that of typical coronary balloons (usually Pebax 72D or 74 D). These two features make the Apollo stent more flexible than typical coronary stent delivery systems. Several single-center studies have shown its safety and efficacy for patients with ICAS. These improved features of the Apollo stents could have contributed to the low complication rate compared with the Vitesse stents.

In this study the inclusion criteria in the degree of symptomatic stenosis (ranging from 70% to 99%), patient age (18–85 years vs 22–80 years), baseline mRS score (≤3), and time to stenting from last stroke (≥3 weeks vs ≥8 days) were similar to the WEAVE/WOVEN study. The median time to stenting from the qualifying event was comparable in the present study and the WEAVE/WOVEN study (20 days vs 22 days). However, there were several differences between the present study and the WEAVE/WOVEN study. First, in the WEAVE/WOVEN study all of the qualifying ischemic events were ischemic stroke, while in this study the qualifying ischemic events also included TIA (47.2%). The present study enrolled the patients with ischemic symptoms within the past 90 days due to hypoperfusion in the territory of the target lesion, while the WEAVE/WOVEN study enrolled the patients with a recurrent stroke in the territory of the same lesion after receiving medical therapy. Second, regarding atherosclerotic risk factors of the enrolled patients, in the WEAVE/WOVEN study there was a higher prevalence of hypertension, hyperlipidemia, and diabetes mellitus (p<0.001) but a lower proportion of smokers (p=0.004). Third, compared with the WEAVE/WOVEN study, the proportion of anterior circulation stenting (39.6% vs 55.0%, p<0.001) and baseline mRS ≥2 (9.4% vs 62.5%, p<0.001) were lower.

In this study, technical success was achieved in 100% (159/159) of the patients. The preprocedural mean percent stenosis was comparable to that in the WEAVE/WOVEN study (84.0% vs 83.2%) but the postprocedural mean percent stenosis was relatively lower than that in the WEAVE/WOVEN study (6.1% vs 28.3%), which indicated that the technical success achieved using the balloon-mounted stent was not inferior to the self-expanding Wingspan stent system. Jiang et al performed a multicenter analysis of stenting in symptomatic ICAS which suggested that the 30-day periprocedural stroke rate was 6.0% in patients treated with balloon-mounted stents. The proportion of Mori A type was 32.0% (147/454) in the balloon-mounted group, which is similar to that in the present study (33.3%, 53/159). It should be noted that their study also included patients in the acute phase of stroke, which may increase the risk of perioperative complications. In the present study, both the stroke rate and mortality rate within 72 hours after the procedure were 0% while, in the WEAVE study, both the stroke rate and mortality rate within 72 hours after the procedure were 1.3%. Compared with the results of the WEAVE study, the safety of balloon-mounted stenting for symptomatic ICAS was acceptable.

It should be noted that, in this study, the proportion of Mori A or Mori B lesions was relatively high (85.5%) whereas Mori C lesions accounted for only 14.5%. However, in the WEAVE/WOVEN study, the morphological features of the ICAS lesions were not described. Balloon-mounted stenting requires the stent and the balloon to be in place at the same time, so it is more suitable for lesions that are relatively simple in shape, short in length, and with approximate diameters in the distal and proximal vessels, such as Mori A or Mori B lesions. This study suggests that endovascular treatment for patients with symptomatic ICAS (especially Mori A or Mori B lesions with smooth arterial access) using a balloon-mounted stent was feasible and safe.

In the VISION study the 1-year primary outcome of stroke or TIA (or hard TIA) occurred in 36.2% (21/58) of the patients in the stent group. However, few studies have investigated the long-term outcomes after balloon-mounted stenting for symptomatic ICAS with experienced interventionalists and proper patient selection. The 1-year rates of any stroke, ischemic stroke, hemorrhagic stroke, and death were 8.5%, 8.5%, 0%, and 1.6% in the WOVEN study, and comparable values were found using balloon-mounted stents in the present study.

In the present study the 1-year rate of ISR was 23.4% (15/64) with a follow-up rate of 40%, which was higher than that in the VISION study (2.9%, 1/34) with a follow-up rate of 58.6% and in the WOVEN study (17.6%, 18/102) with a follow-up rate of 79.1%. However, the 1-year rate of ISR in this study was similar to the 6-month rate of ISR in patients treated with balloon-mounted stents in the study by Jiang et al (20.3%, 60/295). The rate of ISR in Mori C lesions (53.8%, 7/13) was significantly higher than that in Mori A (15.8%, 3/19) or Mori C lesions in the WOVEN study, the morphological features of the ICAS lesions were not described. Balloon-mounted stenting requires the stent and the balloon to be in place at the same time, so it is more suitable for lesions that are relatively simple in shape, short in length, and with approximate diameters in the distal and proximal vessels, such as Mori A or Mori B lesions. This study suggests that endovascular treatment for patients with symptomatic ICAS (especially Mori A or Mori B lesions with smooth arterial access) using a balloon-mounted stent was feasible and safe. In the VISION study the 1-year primary outcome of stroke or hard TIA occurred in 36.2% (21/58) of the patients in the stent group. However, few studies have investigated the long-term outcomes after balloon-mounted stenting for symptomatic ICAS with experienced interventionalists and proper patient selection. The 1-year rates of any stroke, ischemic stroke, hemorrhagic stroke, and death were 8.5%, 8.5%, 0%, and 1.6% in the WOVEN study, and comparable values were found using balloon-mounted stents in the present study.
B lesions (15.6%, 5/32), which has not been described in the VISST or WOVEN studies. The 1-year rate of symptomatic ISR was 3.1% (2/64) in this study, which might have been underestimated because only near half of the patients with recurrent stroke had available follow-up images (44.4%, 4/9). This study indicated that the long-term efficacy of endovascular treatment using balloon-mounted stents for symptomatic ICAS with focal and non-angular lesions (Mori A and B type) and smooth arterial access were comparable to the results of the WEAVE/WOVEN trial. Previous studies have indicated that the long length of ICAS lesions or stents was associated with an increased risk of ISR. In this study, the higher rate of ISR in Mori C lesions may also be related to the longer length of the ICAS lesions and the placed stents.

Recently, balloon-mounted stents (including the Apollo stent) and self-expanding stents have been reported to be used for acute intracranial large vessel occlusion due to presumed atherosclerotic disease. These studies suggest that intracranial stent implantation may be one of the rescue treatments for acute ischemic stroke due to ICAS, but its safety and efficacy require confirmation in future studies.

This study has several potential limitations. First, the defining criteria for hemodynamic impairment were heterogeneous. Four different imaging modalities were allowed to evaluate patients, which made a homogenous evaluation of subgroups of patients difficult. Second, as a 1-year angiogram was performed in only 40% of the recruited patients on a voluntary basis, potential selection bias may be inevitable, especially for evaluation of ISR. Third, the patients were all Chinese, so the results cannot be generalized to other ethnicities. Whether the results of this registry would stand examination by a randomized trial remains to be seen.

CONCLUSIONS
The short and long-term outcomes of balloon-mounted stenting for symptomatic ICAS with focal and non-angular lesions (Mori A and B type) and smooth arterial access were comparable to the results of the WEAVE/WOVEN trial.

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