Longer 6-mm Diameter Stent Retrievers Are Effective for Achieving Higher First Pass Success with Fibrin-Rich Clots

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Fibrin clot · Solitaire\textsuperscript{TM} · First pass success · Stent retriever · Mechanical thrombectomy · Large vessel occlusion · Stroke

Abstract
First pass success (FPS) can be defined as in vitro retrieval of clot in a single pass during mechanical thrombectomy (MT) for acute large vessel occlusion (LVO). Despite advancements in MT technology, retrieval of fibrin-rich clots remains a challenge. Therefore, the effect of stent retriever length on FPS for fibrin-rich clots was investigated by using Solitaire\textsuperscript{TM} 6 × 40 versus 6 × 30 mm devices with a balloon guide catheter (BGC) or distal access catheter (DAC) and sheath, in an in vitro model of anterior circulation neurovascular anatomy. Additionally, vascular safety of the Solitaire\textsuperscript{TM} 6 × 40 versus 6 × 30 mm devices was evaluated in a porcine model for differences in: luminal thrombus, inflammation, endothelial coverage, fibrin deposits, smooth muscle cell loss, elastic lamina and adventitia disruption, intimal hyperplasia, and lumen reduction, at 0, 30, and 90 days post-treatment. In vitro overall FPS was measured as: Solitaire\textsuperscript{TM} 6 × 40 (95%) and Solitaire\textsuperscript{TM} 6 × 30 (67%). FPS for clot location in middle cerebral artery was: (a) BGC (6 × 40 mm: 100%; 6 × 30 mm: 100%; \(n = 8\)); (b) DAC with 088 sheath (6 × 40 mm: 83%; 6 × 30 mm: 33%; \(n = 12\)). FPS for clot location in internal carotid artery was: (a) BGC (6 × 40 mm: 100%; 6 × 30 mm: 80%; \(n = 11\)); (b) DAC with 088 sheath (6 × 40 mm: 100%; 6 × 30 mm: 67%; \(n = 10\)). Stent length had a significant effect (Fisher's exact test; \(p < 0.05\)) on FPS. In vivo evaluation in the porcine model showed no difference in vascular safety parameters between the Solitaire\textsuperscript{TM} 6 × 40 and 6 × 30 mm devices (\(p > 0.05\)) at all time points in the study. Longer stent retrievers may be safe and effective in improving FPS for fibrin-rich clots in in vitro and in vivo models of LVO.

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Introduction

Mechanical thrombectomy (MT) with stent retrievers has shown to be safe and effective for treatment of intracranial large vessel occlusion in acute ischemic stroke treatment [1–7]. First pass effect with stent retriever use has been suggested as a clinical metric of improved patient outcomes [8]. Higher fibrin content thrombi present in approximately 44% of clinical cases [9] are mechanically stiffer and typically more difficult to retrieve compared to softer red blood cell-rich clots. Recent studies have evaluated clot retrieval resistance as a function of clot composition and it has been found that stent retrievers do not engage well with high fibrin content clots [10, 11].

Newer-generation stent retriever devices (Solitaire™ Platinum 6 × 40 mm, Medtronic) are longer and larger in diameter. These modifications are hypothesized to allow the clot to engage more effectively with the stent with multiple planes of attachment at the lower range (2–3 mm) of diameters in the MCA (middle cerebral artery) and maintain this engagement at the higher range (3–6 mm) of diameters in the ICA (internal carotid artery) during clot retrieval. In vitro benchtop methods provide a controlled and reproducible method to assess device performance characteristics with a variety of clot types [12–18] and would therefore allow a comparison of clot retrieval efficacy between device lengths of the same diameter. Even though longer stent retrievers are hypothesized to exhibit a higher clot retrieval success rate, the vascular safety of these devices in an in vivo model also needs to be assessed. Previous in vivo studies in porcine models have shown that longer devices 4 mm in diameter (4 × 40 vs. 4 × 20 mm) are equivalent in terms of safety [19]; however, a similar comparison between the larger 6-mm diameter devices has not been conducted. Recent clinical data show higher first pass mTICI 2b/3 reperfusion with longer 4-mm diameter stent retrievers [20]. However, the mechanism of the improved efficacy with longer devices and correlation with fibrin-rich clots is difficult to decipher from clinical data alone.

The purpose of this study was therefore two-fold: (a) to evaluate in vitro any differences in clot retrieval efficacy as a function of stent retriever length for 6 × 40 mm versus 6 × 30 mm Solitaire™ devices for the fibrin-rich clot type, and (b) to evaluate in vivo the safety of the 6 × 40 mm Solitaire™ device compared with the 6 × 30 mm Solitaire™ device, in a porcine model.

Methods

First Pass Success: In vitro Study

First Pass Success

In the present study, we introduce first pass success (FPS) as a metric to measure in vitro clot retrieval efficacy for homogenous cylindrical fibrin-rich clots. FPS was measured as a binary outcome: 0 (fail) or 1 (pass) for each run completed. Therefore, the result is either a TICI3 recanalization (successful clot retrieval – pass) or no recanalization (no clot retrieval – fail). The result for each device is expressed as a percentage of successful clot retrievals (% FPS).

Test Model

A silicone elastomer-based tortuous anatomical model consisting of the right ICA extending from a type 3 aortic arch with clinically relevant vessel diameters ranging from 6.5 or 5.5 mm in the proximal ICA to 1.5 mm in the M2 segment of the MCA, was used (Fig. 1). A physiological solution of water and 0.2% v/v soap was circulated from the ICA and the posterior communicating artery (PCOM) to the anterior cerebral artery (ACA), MCA and ophthalmic arteries (outflow) at 37 ± 2 °C under clinically relevant pressures (150 ± 10 mm Hg [systolic] and 70 ± 10 mm Hg [diastolic]) and flow rate (300 ± 50 mL/min). Flow was driven with a Harvard Pulsatile Pump (Harvard Apparatus, MA, USA) operating at 60 bpm.
Clot Analog

One clot analog – fibrin rich – was utilized for this study based on properties known from clinically analyzed clots [21]. The fibrin-rich clot composition consisted of a combination of porcine blood (3 mL), barium sulfate (0.25 g), fibrinogen (4 mL of 10% w/v), calcium chloride (0.1 mL of 1 M stock), and thrombin (0.025 mL of 500 U/mL stock). The clot was evaluated for mechanical stiffness using a Dynamic Mechanical Analyzer (TA Instruments) per the methodology described previously to ensure that the range of values was consistent with thrombus seen clinically [21]. The modulus was found to be 0.138 ± 0.08 MPa, on average. The desired length of the thrombi was 20 and 25 mm for the MCA and ICA, respectively, for the 6 × 30 and 6 × 40 mm devices FPS comparison. These represent the upper limits of clot lengths commonly seen at these locations [22].

Clot Retrieval

After the clot was introduced, access to and through the clot to the distal MCA was achieved using a 0.014 in Avigo™ guidewire (Medtronic) and a Marksman™ 0.027 in microcatheter (Medtronic). The Solitaire™ device was delivered through the microcatheter until the distal tip of the Solitaire™ reached the end of the microcatheter and was deployed by pinning the delivery wire and withdrawing the microcatheter. The non-working lengths (proximal and distal ends of the devices) did not interact with the clot during device deployment. All experiments were conducted under full visibility of the model, clot, and device to the operators (no fluoroscopy), thereby allowing for precise deployments in all cases. After engagement with the clot, the Solitaire™ device was withdrawn into either (a) a distal access catheter (DAC; 060 Arc™, Medtronic) or (b) a balloon guide catheter (BGC; 9F Cello™, Medtronic). If a DAC was used, the microcatheter was removed prior to aspiration with a 60-ml syringe. If a BGC was used, the balloon was inflated prior to retrieval of the clot to arrest flow through the ICA.

Devices and Experiments

A total of 41 thrombectomy evaluations were conducted in vitro and were randomized between Solitaire™ 6 × 30 mm (n = 20) and 6 × 40 mm (n = 21) with both MCA and ICA as clot locations and both BGC or DAC with 088 sheath as access options.
Vascular Safety: In vivo Study

Test Model and Medication

Swine was used for the animal model as the arterial anatomy, hemodynamic and coagulation properties mimic those of a human [19, 23]. The in vivo study was conducted under an approved Institutional Review Board protocol. Eighteen Yorkshire cross swine were utilized in this study with 6 animals each at 0-, 30-, and 90-day time points. The Solitaire™ devices (6 × 30 and 6 × 40 mm) were deployed as per the instructions for use in the following arteries: vertebral, internal thoracic, and costocervical. The targeted vessels were <3.0 mm in diameter to ensure that the devices were deployed into worst-case vessel sizes at or below the minimum labeled vessel size. Dual antiplatelet therapy (aspirin: 81 mg and clopidogrel: 75 mg) was administered orally 1-day pre-procedure for all animals to reduce the risk of procedure-related thromboembolic complications. Nifedipine (30 mg) was administered as necessary to prevent vessel spasm intra-procedure only.

Test Devices and Deployments

For one complete test, a Solitaire™ device was deployed in the vessel seven times, resheathed into the microcatheter six times at the deployment location and retrieved through the target vessel into the guide catheter three times (i.e., multiple deployments of the devices were performed at the target location to mimic a clinical worse case). Angiographic assessments of the target vessel were performed prior to and following each pass, and presence of any emboli, vessel damage, thrombus formation, and/or vasospasm were noted. No clot analogs were utilized in the in vivo testing to reduce potential variation between tests and to provide the most direct evaluation of the stent retriever on the vessel wall; this aligns with previous preclinical work performed on Solitaire™ 4-mm devices [19].

Angiographic and Histopathology End Points

At 0, 30 ± 3, or 90 ± 3 days following the multiple deployment and recovery testing, the animals underwent a follow-up angiogram to evaluate target vessels for stenosis, thrombus formation, and vessel damage. Animals were euthanized following angiographic evaluation, and the treated vessels were evaluated histologically for the following attributes: luminal thrombus, inflammation, endothelial coverage, fibrin deposits, smooth muscle cell loss, elastic lamina and adventitia disruption, intimal hyperplasia, and lumen reduction, and graded on a scale of 0–4 at three locations along the treated zone (Table 1).

Statistical Analysis

Statistical comparisons for in vitro FPS for Solitaire™ 6 × 40 versus 6 × 30 mm were conducted with Fisher’s exact test. Statistical comparisons for histological parameters between Solitaire™ 6 × 40 and 6 × 30 mm were conducted with ANOVA at the three time points (0, 30, and 90 days). A significance level of 0.05 was used for all tests.

Results

FPS: In vitro Study

Solitaire™ 6 × 40 and 6 × 30 mm FPS Comparison

FPS with the 6 × 40 mm device (95%; n = 20) was significantly higher compared to the 6 × 30 mm device (67%; n = 21). The results are shown in Figure 2 for overall FPS and FPS by access type. FPS for clot location in MCA was: (a) BGC (6 × 40 mm: 100%; 6 × 30 mm: 100%; n = 8); (b) DAC with 088 sheath (6 × 40 mm: 83%; 6 × 30 mm: 33%; n = 12). FPS for clot location in ICA was: (a) BGC (6 × 40 mm: 100%; 6 × 30 mm: 80%; n = 11); (b) DAC with 088 sheath (6 × 40 mm: 100%; 6 × 30 mm: 67%; n = 10). In most cases, the clot was corked at the tip of the DAC or BGC prior to retrieval of the entire system.

Vascular Safety: In vivo Study

All animals were in good clinical health throughout the duration of the study. A gross necropsy evaluation found no thromboembolism in the heart or lungs of the animals.
Angiographic Evaluation

No thrombus formation, emboli, or vessel wall damage was seen angiographically during any of the procedures for all devices used.

Histopathologic Evaluation

There was no significant effect of stent length on any of the histopathology parameters of the treated vessels with the 6 × 40 and 6 × 30 mm Solitaire™ devices ($p > 0.05$). All groups

| Histopathologic attribute | Grading scale |
|---------------------------|---------------|
| Luminal thrombus          | 0: No luminal thrombus  
1: (Minimal): Occupies ~5–5% of the lumen area  
2: (Mild): Occupies ~5–35% of the lumen area  
3: (Moderate): Occupies ~35–70% of the lumen area  
4: (Severe): Occupies ~>70% of the lumen area |
| Endothelial cell coverage | 0: (Absent): ~<5% of the luminal surface  
1: (Minimal): ~5–25% of the luminal surface covered  
2: (Mild): ~25–50% of the luminal surface  
3: (Moderate): ~50–90% of the luminal surface  
4: (Complete): ~>90% of the luminal surface |
| Inflammation              | 0: No inflammatory response  
1: Minimal response  
2: Mild response  
3: Moderate response  
4: Severe response |
| Fibrin deposits           | 0: Not present  
1: Minimal  
2: Mild  
3: Moderate  
4: Severe |
| Medial smooth muscle cell loss | 0: No medial SMC loss  
1: Minimal ~<5% medial SMC loss  
2: Mild ~5–25% medial SMC loss  
3: Moderate ~25–50% medial SMC loss  
4: Severe ~>50% medial SMC loss |
| Disruption of the internal elastic lamina and external elastic lamina | 0: No disruption  
1: Minimal ~<5% disruption  
2: Mild ~5–25% disruption  
3: Moderate ~25–50% disruption  
4: Severe ~>50% disruption |
| Adventitia disruption     | 0: No changes  
1: Minimal ~<5% of the area with changes  
2: Mild ~5–25% of the area with changes  
3: Moderate ~25–50% of the area with changes  
4: Severe ~>50% of the area with changes |
| Neointimal hyperplasia    | 0: (Absent): ~<5% of the luminal surface  
1: (Minimal): ~5–25% of the luminal surface  
2: (Mild): ~25–50% of the luminal surface  
3: (Moderate): ~50–90% of the luminal surface  
4: (Severe): ~>90% of the luminal surface |
| Lumen area reduction due to hyperplasia | 0: (Absent): ~<5% of the luminal area  
1: (Minimal): ~5–25% of the luminal area  
2: (Mild): ~25–50% of the luminal area  
3: (Moderate): ~50–90% of the luminal area  
4: (Severe): ~>90% of the luminal area |

**Table 1.** Histopathologic evaluation attributes and grading scale
showed minimal or no response for the following at each time point investigated: luminal thrombus, inflammation, internal elastic lamina and external elastic lamina disruption, adventitia changes.

The effect of time point was significant for the following parameters: endothelial coverage, neointimal hyperplasia, lumen area reduction, smooth muscle cell loss, and presence of fibrin deposits \((p < 0.05)\). At the acute (0-day) time point, endothelial denudation and slight SMC loss was noted for all devices. By the 30-day time point, all vessels had complete or nearly complete endothelialization and no SMC loss. However, at this time point, there was minimal fibrin deposition and moderate to severe neointimal hyperplasia in all vessels with mild lumen area reduction. At the 90-day time point, both treatment groups saw a significant increase in the lumen area. These results are shown in Figure 3.

**Discussion**

The present data seem to indicate that increase in length of the 6-mm diameter Solitaire™ device may improve FPS for fibrin-rich clot retrieval without increasing the risk of vessel damage. Regarding safety, the 6 × 40 mm Solitaire™ device showed no significant effect of stent length on any in vivo vascular parameters evaluated in a porcine model compared with the 6 × 30 mm Solitaire™ device. No serious adverse events were noted angiographically during the procedures and during the 90-day survival phase of the study. Additionally, the histopathology results were similar to those seen in a previous study of 4-mm Solitaire™ devices and did not show any significant health risks \([19, 24]\). The effect of time post-treatment was significant for some vascular parameters – in particular the endothelial coverage – at the target location of the procedure. This has been reported previously with MT \([19]\) and is known not to have long-term clinically significant effects on the vessel \([23–25]\). Despite some
recent evidence collected in an animal model suggesting that alternative endovascular techniques may cause less vessel damage [26], a direct comparison in a clinical trial has shown no difference in adverse events when compared to MT with the use of stent retrievers [27].

Some limitations with the animal model are as follows: (a) the tortuosity of the target vessels was much lower than typically observed in the ICA and MCA; (b) the in vivo testing was also performed with no clot analog in order to most directly compare the stent effects on smaller-sized vessels alone (maximize the contact of the stent and the vessel, a worst-case scenario); (c) evaluation of the treated vessels was also limited to a 90-day window, which provides adequate information on the immediate effect of treatment and the subsequent near-term vascular healing process; and (d) the porcine arterial vessels in the in vivo study may have structural differences compared with human intracranial arteries.

Benchtop evaluation of FPS using a fibrin rich clot with the 6 × 40 versus 6 × 30 mm Solitaire™ devices demonstrated that there is a significant increase in FPS \( (p < 0.05) \) when using longer lengths of these devices. This is evident primarily with the fibrin-rich clot (clinical occurrence in approximately 44% of cases [9]) which tends not to engage as effectively with

**Fig. 3.** Summary of 0-, 30-, and 90-day vessel histology scoring for 6 × 40 and 6 × 30 mm Solitaire™ devices following multiple deployment and retrieval passes in a porcine model. No statistical difference was found due to stent length for any time point \( (p > 0.05) \). IEL, internal elastic lamina; EEL, external elastic lamina; SMC, smooth muscle cell.
the stent retrievers as red blood cell-rich or mixed clot compositions due to higher mechanical stiffness [21] and possibly differences in friction [10]. The longer-length devices provide better engagement for fibrin-rich clot after deployment of the stent retriever and enable corking at the tip of the DAC or BGC prior to retrieval of the entire system. Additionally, the FPS reported here with the BGC is higher than the DAC and sheath for all groups (Fig. 2) which also supports improved patient outcomes (lower mRS scores and higher first pass effect) reported clinically with BGC use [8].

The following limitations of the in vitro study should be noted. The FPS testing performed for this study was designed to simulate a single-sided ICA-MCA-ACA model with limited collateral flow. To enable effective 1:1 comparisons for FPS between stent retriever lengths, several assumptions were made in this study: (a) lack of full circle of Willis limits extensive flow collateralization; (b) viscosity of the fluid was similar to water (incorporation of soap is essential to mimic the friction between the stent and vessel wall and is not feasible when using glycerol); (c) homogeneous fully occlusive cylindrical shaped clots were used; (d) the DAC option was primarily used (since there is no flow arrest and potential for the clot to disengage from the stent retriever is higher); and (e) placement of the DAC at the ICA-T bifurcation (this is the clinical worst case if the DAC cannot be advanced to be en face with the clot in the MCA).

Although the clot tends to disengage with both devices during retrieval, the longer device helps maintain engagement until the clot is captured or corked at the tip of the DAC. The additional stent length is also relevant for retrieval of large clots at the ICA-T bifurcation, particularly in the case of BGC use, where the stent-clot engagement needs to be maintained until the system is retrieved up to the proximal ICA (location of the BGC). All these factors collectively result in demonstrating a higher FPS for the longer 6-mm diameter stent retrievers in this bench model.

Conclusion

The in vitro portion of the present study suggests a significantly higher FPS with the 6 × 40 mm Solitaire™ device compared to the 6 × 30 mm Solitaire™ device in fibrin-rich clot retrieval. The in vivo portion of this study seems to indicate that the 6 × 40 mm Solitaire™ device has a non-inferior vascular safety profile compared to the 6 × 30 mm Solitaire™ device.

Statement of Ethics

Animal experiments conform to internationally accepted standards and have been approved by the appropriate institutional review body.

Disclosure Statement

Dr. Amon Y. Liu and Dr. Italo Linfante are consultants for Medtronic.

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