Histopathologic Evaluation of Arterial Wall Response to 5 Neurovascular Mechanical Thrombectomy Devices in a Swine Model


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ABSTRACT

BACKGROUND AND PURPOSE: Five commercial devices are available for mechanical thrombectomy in acute ischemic stroke. This study evaluated and compared the resultant arterial damage from these devices.

MATERIALS AND METHODS: Wall damage after 4 wall-contact devices (the Merci retriever, Catch thromboembolectomy system, and Solitaire FR revascularization devices of 4 and 6 mm) and 1 aspiration device (the Penumbra System) was evaluated in the superficial femoral arteries of 20 male swine. Each device was tested with and without intraluminal clot. Twenty control vessels were not subjected to any intervention. Acute histopathologic changes were evaluated.

RESULTS: In the device samples, endothelial denudation (72.8 ± 29.4% versus 0.9 ± 1.9%, P < .0001), medial layer edema (52 ± 35.9% versus 18.1 ± 27.8%, P = .004), and mural thrombus (5.3 ± 14.2% versus 0%, P = .05) were found to a greater extent compared with the control samples. The aspiration device provoked more intimal layer ([100 ± 79.1% versus 58.8 ± 48.9%, P = .27] and medial layer ([75 ± 35.4% versus 46.3 ± 34.8%, P = .13]) edema than the wall-contact devices.

CONCLUSIONS: All devices caused vascular injuries extending into the medial layer. The aspiration device was associated with more intimal and medial layer edema, compared with the wall-contact devices except for the Catch thromboembolectomy system.

ABBREVIATIONS: EEL = external elastic lamina; IEL = internal elastic lamina; MET = mechanical endovascular thrombectomy; SFA = superficial femoral artery

Recanalization is a powerful predictor of stroke outcome in patients with arterial occlusion treated with either IV rtPA or an endovascular approach.1,3 IV thrombolysis has been limited by its low recanalization rate in the setting of large-artery occlusions,5 whereas mechanical endovascular thrombectomy devices can result in higher rates of recanalization in patients with stroke with similar proximal arterial occlusions.4 The recanalization rate with MET is approximately 80%, and approximately 40% of patients have a favorable clinical outcome.7

A number of new thrombectomy devices have recently been introduced, with varying theoretic mechanisms of action. They can be classified into 2 major groups, according to their mechanism of retrieving the clot. Aspiration devices apply force to the proximal base of the thrombus6,8; this group includes various aspiration catheters and systems. Wall-contact devices can be divided into distal devices (brushlike, basketlike, or coil-like devices)9 and stentlike devices (including various self-expandable stent retrievers).10 With both types, the force is applied to the distal base of the clot. So far, to our knowledge, no systematic study has been performed comparing various MET devices under standardized conditions in vivo, in terms of arterial wall response (MEDLINE data base research by using the terms “endovascular,” “mechanical thrombectomy,” “clot removal,” “arterial wall,” and “animal model”).

The purpose of this study was to evaluate and compare arterial wall responses to 5 neurovascular mechanical thrombectomy devices: 4 wall-contact devices (the Merci retriever; Concentric Medical, Mountain View, California; the Catch thromboembolectomy system; Balt, Montmorency, France; and the Solitaire FR revascularization devices, 4 and 6 mm; Covidien, Irvine, California) and 1 aspiration device (Penumbra System; Penumbra, Alameda, California).

MATERIALS AND METHODS

Animal Care

All procedures were conducted according to international guidelines and were approved by the responsible local authorities.
Twenty male swine ranging from 20 to 25 kg (Haute-Vienne Research and Analysis Department, Haute-Vienne, France) were used in this study. The procedures were conducted with the swine under general anesthesia. No further heparin bolus or other thrombolytic drugs were administered. Further details are described in previous studies.\(^9\) Vital parameters, such as arterial blood pressure, heart rate, and carbon dioxide levels, were continuously recorded. Expired carbon dioxide levels were maintained at 30–35 mm Hg. One hour after the end of the last retrieval, the animals were euthanized.

### Description of the Neurovascular Thrombectomy Devices

We tested 5 devices: 4 wall-contact devices and 1 aspiration device. Their properties are summarized in Table 1. The wall-contact devices were the following:

- **Solitaire FR**
  - **FR 4 mm**: Diameters of 4 mm (Solitaire FR 4 mm and Solitaire FR 6 mm). These are self-expanding, closed-cell nitinol stent devices that can be fully deployed and fully resheathed, and their design allows immediate blood-flow restoration and clot retrieval.
  - **FR 6 mm**: Diameters of 6 mm (Solitaire FR 4 mm and Solitaire FR 6 mm). These are self-expanding, closed-cell nitinol stent devices that can be fully deployed and fully resheathed, and their design allows immediate blood-flow restoration and clot retrieval.

- **Catch Thromboembolotmy System**
  - Consists of a self-expanding, basket-like device that has a maximum diameter of 4 mm and is fixed to a pusher wire. After the microcatheter, equipped with a microwire (SilverSpeed 14; ev3, Irvine, California), was placed in the common femoral artery, the microcatheter was navigated into the occluded vessel, past the thrombus. The basket was deployed distal to the thrombus. Then both the basket and the microcatheter were pulled back, simultaneously, into the guiding catheter.

- **Merci Retriever V 2.5 Firm**
  - The Merci retrieval system is a tapered wire with 5 helical loops of decreasing diameter at its distal end. In this study, the Merci retriever V 2.5 Firm was used, with a maximum loop diameter of 2.5 mm. The microcatheter was advanced beyond the thrombus. Both the microcatheter and the Merci retriever were pulled back to contact the thrombus, and the remaining loops were deployed within the thrombus. During retrieval, aspiration was applied through the guide catheter, by using a 50-mL syringe.
  - The aspiration device was the Penumbra System 0.32 (Penumbra). This is a platform that includes a reperfusion microcatheter connected to an aspiration pump via aspiration tubing, generating a suction force of 0.9 bar (700 mm Hg).
  - When the microcatheter tip was placed in contact with the thrombus, the aspiration was applied during the clot fragmentation by using a separator.

### Clot Preparation

Radiopaque clots were prepared in vitro, as recently described by Kan et al.\(^{11}\) In brief, 20 mL of whole blood was mixed with 2 g of barium sulfate powder to obtain radiopacity, and the mixture in the syringe was incubated at room temperature for 120 minutes. The solid component was separated from the serum component, and a piece of clot was resected. Each prepared thrombus, which measured 25 mm in length, was inserted into a Cole-Parmer silicone tube (Vernon Hills, Illinois) with saline and was prepared for injection.

### Study Protocol

The protocol is detailed in Fig 1. Twenty swine had their superficial femoral arteries dissected bilaterally. The 20 right SFA samples were not subjected to any kind of intervention (control samples). The 20 left SFA samples were randomly submitted to thrombectomy with 1 of the 5 devices (device samples) (Fig 2A).

Concerning the aspiration system, the clot retrieval was conducted according to 2 scenarios: 100 and 200 passages of the fragmentation separator under aspiration of 0.9 bar (similar to the clinical use).

### Angiography and Clot Placement

A short 6F catheter sheath (Terumo, Tokyo, Japan) was inserted into the right common femoral artery and continuously flushed with...
A 6F guiding catheter (Neuropath guiding catheter; DePuy, Raynham, Massachusetts) was introduced and continuously flushed with physiologic saline. Angiography of the left SFA was performed on a biplane system (Integris; Philips, Best, the Netherlands) (Fig 2B). Rotational angiography, followed by 3D reconstruction of the native projections (Fig 2C), was performed just before the thrombectomy procedure was begun. The diameters of the vessels were measured in 3D reconstruction images.

For selective thromboembolization, the preformed thrombus was injected into a 5F guide catheter (Chaperon; MicroVention, Aliso Viejo, California) positioned in the left SFA. After connecting the silicone tube containing a prepared clot into the guiding catheter, we injected 2 mL of saline into the tube to propel the clot into the catheter. The clot migrated to the SFA to achieve the occlusion. After an angiogram had been obtained to document occlusion, the guiding catheter was retrieved. The occlusion was maintained for 1 hour (± 30 minutes) before mechanical retrieval was attempted (Fig 2D).

All procedures were performed with the swine under general anesthesia through a 6F guiding catheter placed in the left SFA. The retrieval attempts were conducted according to the requirements of the manufacturers. During retrieval, only the microwire, microcatheter, and device entered the target vessel. For repeated attempts, the devices were cleaned, inspected, and, if still intact, reinserted.

**Study End Point**
The protocol was strictly designed to assess the degree of vascular damage. The recanalization rate, time to achieve recanalization, vasospasm, arterial perforation, and distal embolism were not evaluated.

**Histopathologic Evaluation**
Both of the SFAs were dissected away from the surrounding tissue, and each was cut transversely into 6 individual pieces, numbered proximally to distally, with regard to arterial blood flow. The most proximal and distal 2 cm of the SFAs were excluded to avoid ligature sites occurring during surgery. The vessel sections from each artery were processed and embedded in paraffin, according to standard laboratory operating procedures. They were subsequently sectioned at 5–7 μm and stained with hematoxylin-eosin and orcein (elastic fibers). The sections were examined with an optical microscope by using magnification ranging from 40 to 400 times by 2 independent board-certified pathologists who were blinded to the device used (A.G. and C.Y., who had 5 and 21 years of experience, respectively).

A grading system to evaluate the acute mural response to the MET devices was developed (Fig 3). We assessed the following: 1) amount of endothelial denudation (percentage of surface area), 2) presence of mural thrombus (vascular occlusion percentage), 3) intimal layer edema (inner intima circumference percentage), 4) medial layer edema (inner media circumference percentage), and 5) internal elastic lamina fracture (percentage).

**Statistical Analysis**
The data from the histologic grading were independently analyzed by using SAS, Version 9.1.3 (SAS Institute, Cary, North Carolina). The statistical differences in arterial damage grades among the 5 devices were analyzed by the Mann-Whitney test. A P value < .05 was considered significant.

**RESULTS**
Forty SFAs with diameters of 2.0–3.5 mm from 20 swine were studied. The model of arterial occlusion was effective in all target vessels. The five devices were applied to 20 vessels (4 vessels each)—2 with clots and 2 without—and they were successfully retrieved in every case.

**Histopathologic Results**
All of the MET devices caused lesions, which extended from the intimal layer to the medial layer. In the device samples versus the control samples, these were characterized by significant endothelial denudation and medial layer edema (Table 2). Intimal layer edema and fracture of the IEL were also found in all of the device samples but with insignificant differences for all device samples combined compared with the control samples. No other lesions were found deeper in the external elastic lamina or adventitia, nor were there perforations or dissections in any samples.
There were no statistically significant differences in the arterial damage among the 5 device groups. However, the aspiration device (Fig 4) created less endothelial denudation and mural thrombus than the wall-contact device samples (Fig 5), but more intimal and medial layer edema, though none of these differences were significant (Table 3).

**Mural Thrombus**

Overall, the MET devices caused more mural thrombus in the arterial lumen of the device samples compared with control samples (Table 2). Three MET devices (the Solitaire FR 4 mm, Merci retrieval system, and Catch thromboembolotomy system) caused mural thrombus after their use, but the Penumbra system and Solitaire FR 6 mm did not.

**Intraluminal Clot**

There was a predominance of intimal layer edema in arteries without intraluminal clots, compared with the clotted ones, but there were no other significant histopathologic differences (Table 4).

**Number of Attempts**

For each device, 2 and 3 retrieval attempts were performed in samples both with and without a clot (resulting in 4 procedures per device).
mean arterial damage for each retrieval attempt with and without clot is summarized in Table 4. The number of attempts with each device in both groups of arteries (with and without clot) did not influence the histopathologic results; there was no statistical difference shown, but variability was large and sample sizes were small.

**DISCUSSION**

The approved medical therapy in acute ischemic stroke is IV rtPA within 4.5 hours. Recently, mechanical thrombectomy has been demonstrated to successfully restore large-vessel patency and therefore provide an alternative and synergistic method for flow restoration. Thrombectomy devices offer many potential advantages over pharmacologic thrombolysis, including more rapid achievement of recanalization, enhanced efficacy in treating large-vessel occlusions, and a potentially lower risk for hemorrhagic events. However, mechanical manipulation of a thrombus in intracranial arteries might cause complications such as vasospasm, dissection, or perforation and may worsen clinical outcome at discharge.

Nonetheless, arterial wall changes after endovascular treatment with mechanical devices in acute stroke are poorly reported and compared. There are no standards to evaluate such devices. The first in vivo model dedicated to MET in ischemic stroke was reported by Gralla et al. It permits an angiographic evaluation and comparison of neurovascular mechanical thrombectomy devices. However, no study has evaluated the histopathologic responses induced by devices, to our knowledge. Recently, a model has been specifically developed in swine, by using the superficial cervical artery to evaluate arterial structural changes. Compared with this model, surgical access to swine SFA is not technically difficult, due to its superficial
position. The normal histologic characteristics of the porcine femoral artery are well known\textsuperscript{23} and raise the possibility of evaluating pathologic lesions. In a postmortem study of 5 patients who died acutely after MET procedures, vascular pathologic abnormalities affecting the proximal arteries were variable, but 1 patient (20%) showed a subintimal dissection with resultant occlusion of the middle cerebral artery.\textsuperscript{24} In the current study, 5 mechanical devices (4 wall-contact devices and 1 aspiration device) were evaluated and the grading scale analyzing the acute extent of the injury caused by the device (from the inner layer to the adventitia layer) and the presence of swelling or an intraluminal thrombus were reported.

It was observed that the devices caused significant endothelial and medial damage to the vessel walls. Intimal and subintimal damage seemed higher, but no intimal dissection was found histologically. Littie is known about the response of the arterial wall after mechanical thrombectomy, partly due to limited angiographic/histologic data from the treated arteries. Recently, an angiographic follow-up study in 261 patients with 265 embolic vessel occlusions revealed 26.0% vasospasm and 0.4% dissection in the acute phase.\textsuperscript{25} On follow-up, 0.9% target-vessel occlusion and 3.4% of de novo stenosis were reported. These data may suggest that arterial wall injuries are probably reversible in the long term.

The aspiration device was responsible for more intimal and medial damage to the wall-contact devices and the aspiration system. The authors found that the former appeared to reduce irritation in the vessel wall, thus lowering the risk and severity of vasospasm, because it does not require repositioning and passing procedures.\textsuperscript{26}

An important finding was the presence of less arterial wall damage in vessels with out clots (35 ± 33.4% versus 99.11 ± 57%, P = .009). Regarding these results, we deduced that the length of the device should be close to the length of the clot, to reduce the lesions on the wall. Also, wall damage was not significantly increased when the devices were passed 3 times rather than twice.

Another point was the presence of a mural thrombus in the vessel walls submitted to 3 of the wall-contact devices, specifically the Merci retrieval system, the Catch thromboembolotomy system, and the Solitaire FR 4 mm. However, the lesion was limited to the intimal layer attached to the endothelium, and it was not associated with significant reduction of the arterial lumen. The Solitaire FR 6 mm device caused less endothelial damage and no mural thrombus compared with the Solitaire FR 4 mm device. The Solitaire FR 6 mm exerts a greater radial force at all diameters compared with the Solitaire FR 4 mm. The increased intimal tearing could be attributed to the greater length of the Solitaire FR 4 mm (31.8 versus 26.2 mm).\textsuperscript{27} Having more surface area in contact with the vessel wall. Thus radial force could be considered a less important contributor to arterial wall injury.

Because this study was restricted to the acute phase, performing further studies at later stages—subacute and chronic—would be of great value to identify the progression of these lesions. Previous research has shown that the extent and depth of the vascular lesion may be contributing factors in promoting early atherosclerotic and accelerated hyperplastic intimal and medial changes.\textsuperscript{28} Large areas of endothelial denudation without substantial medial trauma caused only mild intimal thickening, whereas focal endothelial denudation with substantial medial trauma produces marked delayed intimal thickening.\textsuperscript{29} These results are substantiated by those of other studies that have shown that smooth muscle cells are normally quiescent with respect to proliferation but that deep injuries result in the expression of a phenotype of vascular smooth-muscle cells that exhibits a high proliferative response to other mitogens.\textsuperscript{28}

Given these results, the aspiration device resulted in a less traumatic acute injury in the vascular wall. However, this study did not analyze the vascular recanalization rate; therefore, deducing

<p>| Table 3: Comparison of arterial damage by the wall-contact devices and the aspiration device\textsuperscript{a} |
|---------------------------------|---------------------------------|----------------------|</p>
<table>
<thead>
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<th>Wall-Contact Devices (n = 16)</th>
<th>Aspiration System (n = 4)</th>
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<tbody>
<tr>
<td>Mural thrombus (%)</td>
<td>6.6 ± 15.7</td>
<td>0 ± 0</td>
</tr>
<tr>
<td>Endothelial denudation (%)</td>
<td>810.0 ± 16.8</td>
<td>401.0 ± 47.5</td>
</tr>
<tr>
<td>Intimal layer edema (%)</td>
<td>58.8 ± 48.9</td>
<td>100 ± 79.1</td>
</tr>
<tr>
<td>IEL fractured (%)</td>
<td>26.6 ± 51.2</td>
<td>12.5 ± 14.4</td>
</tr>
<tr>
<td>Medial layer edema (%)</td>
<td>46.3 ± 34.8</td>
<td>75 ± 35.4</td>
</tr>
</tbody>
</table>

\textsuperscript{a} Data are means ± standard deviations.  
\textsuperscript{b} Mann-Whitney test comparing wall-contact device samples combined with aspiration system (Penumbra) samples.

<p>| Table 4: Comparison of arterial damage by the device samples according to the presence or absence of a clot and the number of retrievals\textsuperscript{b} |
|---------------------------------|---------------------------------|----------------------|</p>
<table>
<thead>
<tr>
<th></th>
<th>With Clot (10 Samples)</th>
<th>Without Clot (10 Samples)</th>
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</thead>
<tbody>
<tr>
<td>Mural thrombus (%)</td>
<td>0.5 ± 1.6</td>
<td>10.1 ± 19.3</td>
</tr>
<tr>
<td>Endothelial denudation (%)</td>
<td>81.8 ± 18.9</td>
<td>63.8 ± 35.8</td>
</tr>
<tr>
<td>Intimal layer edema (%)</td>
<td>35 ± 33.4</td>
<td>99.1 ± 57.8</td>
</tr>
<tr>
<td>IEL fractured (%)</td>
<td>10 ± 17.5</td>
<td>37.5 ± 61.5</td>
</tr>
<tr>
<td>Medial layer edema (%)</td>
<td>45 ± 43.8</td>
<td>59.1 ± 26.5</td>
</tr>
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</table>

\textsuperscript{b} Data are means ± standard deviations.  
\textsuperscript{c} Mann-Whitney test for 2 versus 3 retrievals.
that the aspiration device would have better functional results is not appropriate. These findings warrant further study of these devices in a model with a longer follow-up.

We acknowledge several limitations of the current study and proposed animal model, as with any in vivo experimental model. First, the swine SFAs are less tortuous without atherosclerosis compared with intracranial arteries in humans. Second, the present study compared arterial wall responses in 5 mechanical thrombectomy devices uncorrelated with their angiographic efficacy. We can imagine that 1 passage may suffice with stent retrievalers, while many more passages may be necessary with aspiration devices. It is probably total injuries for effective recanilization that count. Another limitation is that endothelial denudation is difficult to quantify with routine pathologic stains. Further methodologic limitations of the study are the small number of experiments per device and the lack of follow-up angiographic and pathologic data to assess whether damage leads to stenosis. In addition, because of the differences in the arterial wall (lack of adventitia in intracranial arteries), the histologic results of this model may not be applicable to cerebral arteries in clinical practice. Our new model does not represent a counterpart to the entire phenomenon seen in patients with stroke. However, we believe that it may expand the analysis of the safety profile and offer the possibility of an experimental model for the preclinical development and evaluation of mechanical devices.

CONCLUSIONS
We have demonstrated in this swine model that both the aspiration device and the wall-contact devices caused vascular injuries extending into the medial layer. However, it would appear that the aspiration device was associated with more intimal and medial-layer edema, compared with the wall-contact devices except for the Catch thromboembolocytosystem.

ACKNOWLEDGMENTS
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