CASE REPORT

Intracranial thrombectomy using the Solitaire stent: a historical vignette

Marta Aguilar Pérez, Elina Miloslavski, Sebastian Fischer, Hansjörg Bätzner, Hans Henkes

ABSTRACT

Introduction The endovascular treatment of acute ischemic stroke has been revolutionized in the past years by the introduction of new devices for mechanical thrombectomy. Several tools were already available in 2008. The majority allowed the recanalization of acutely occluded intracranial arteries with acceptable levels of safety and efficacy, and with occasional failures.

Case presentation On 3 March 2008, a 67-year-old woman was treated 3.5 h after the clinical onset of a right hemispheric stroke due to an embolic middle cerebral artery (MCA) M1 occlusion. The National Institutes of Health Stroke Scale (NIHSS) score prior to treatment was 10. Mechanical thrombectomy with a microbrush yielded a significant amount of thrombotic material without recanalization. Given the urgency of the situation, the uncertain outcome in the case of a persistent occlusion of the right M1 segment and the fact that no other device was available, a Solitaire stent was deployed within the occluded right M1 segment. After several minutes of incubation, the expanded stent was slowly withdrawn under continuous aspiration with instantaneous removal of the entire thrombus and complete recanalization of the right MCA with reperfusion of the whole MCA supply territory. Digital subtraction angiography showed neither peripheral emboli nor vasospasm. The patient made a complete clinical recovery with an NIHSS score of 0 at the 30 day follow-up.

Conclusion The Solitaire stent was initially developed for the endovascular treatment of wide necked intracranial aneurysms but has been demonstrated to be safe and efficacious for intracranial thrombectomy. This was the first successful human clinical use of a Solitaire stent for this purpose and the ignition spark for the development of a whole generation of new devices, now called stent retrievers.

INTRODUCTION

Early recanalization of occluded vessels in acute ischemic stroke by intravenous thrombolysis or endovascular recanalization is associated with improved clinical outcome and reduced mortality. The concept of endovascular treatment of acute ischemic stroke was initially based on the local injection of fibrinolytic agents. Clinical benefits of local intra-arterial fibrinolysis (LIF) were reported in case series and from controlled trials. The method was, however, not further developed when a series of randomized controlled trials of intravenous fibrinolysis (IVF) versus placebo led to the approval of the recombinant tissue plasminogen activator alteplase for use in ischemic stroke within 3 h of symptom onset. The difference between the two arms in these studies was quite sobering and <15% of all stroke patients were eligible for IVF.

The general acceptance of IVF, however, was pivotal in anchoring acute ischemic stroke in neurology and was instrumental for the installation of dedicated stroke units. Apart from these merits, the practice of IVF undermined the momentum of further development of endovascular treatment concepts. The first dedicated tool approved by the Food and Drug Administration (FDA) in 2004 for endovascular intracranial thrombectomy was the Merci retriever (Concentric Medical, Mountain View, California, USA), a nitinol spiral. In 2005, Catch (Balt Extrusion, Montmorency, France), a braided nitinol basket proximally attached to a wire and distally occluded, was CE marked (CE=Conformité Européenne, the CE logo indicates free marketability of an industrial product within the European Economic Area) in Europe for the same indication. Both devices were limited in their efficacy and had some safety issues. The Merci retriever was evaluated in large US trials and achieved a certain acceptance both in the USA and in Europe. Competing developments were brushes and braided cages for intracranial thrombectomy.

In 2006, Penumbra (Alameda, USA) came up with an aspiration catheter together with a dedicated aspiration pump and a modified wire (‘separator’). The Penumbra system was shown to be both efficacious and safe for endovascular intracranial thrombectomy in European and US trials and received FDA approval in 2008.

In 2008, completely independent of the efforts on engineering stroke devices, a new stent for the endovascular treatment of intracranial aneurysms was developed by Dendron (Bochum, Germany). This stent received the CE mark in 2004. Dendron was acquired by MTI (Irvine, California, USA) in 2002. The Solitaire stent, now manufactured and distributed by ev3 (Irvine, California, USA), became available for the treatment of wide necked aneurysms in Europe in December 2007.

This was the situation when we treated a patient with an acute embolic M1 occlusion in March 2008 and by chance made the observation described below. Starting from this point and with immediate enthusiasm of other colleagues, it became apparent that a new era of the endovascular treatment of acute ischemic stroke had begun. Here we describe the first successful usage of a Solitaire stent.
as a device for clot retrieval (‘mechanical thrombectomy’) from an acutely occluded intracranial human artery.

CASE PRESENTATION

This 67-year-old woman presented on 3 March 2008 at 16:00 to the neurological emergency room. Earlier that day (at 14:50), her relatives had witnessed the acute onset of a fluctuating paresis of her left arm and leg. Diffusion weighted MR images, carried out at 16:30, revealed an area of restricted diffusion in the right basal ganglia and the deep periventricular white matter and an MR angiogram demonstrated complete occlusion of the right middle cerebral artery (MCA) in the M1 segment. The emergency stroke team decided to initiate intravenous thrombolysis, which started within a 2 h window (16:45). As the patient did not show any clinical improvement within the next 30 min, the attending stroke physician decided to proceed with a bridging concept and endovascular treatment was initiated, after the patient was informed of the proposed procedure. This information included a description of the procedure, the potential risks and benefits and an explanation that extensions of the procedure (eg, surgical craniectomy) or the off-label use of medical devices (eg, coronary stents, etc) might become necessary. The patient gave informed consent to the operator. The neurological examination before digital subtraction angiography (DSA) showed a severe dysarthria and a left-sided hemiplegia with facial nerve palsy (National Institutes of Health Stroke Scale (NIHSS) score of 10). The emergent angiography began 3 h and 20 min after the onset of symptoms and the first DSA run was recorded at 18:12.

Using a 4 F diagnostic catheter, selective injection of both vertebral arteries and internal carotid arteries (ICAs) was carried out to show the extent of leptomeningeal collaterals and to rule out distal emboli, eventually not visible on MR angiography. Complete embolic occlusion of the right MCA at the M1 segment with moderate leptomeningeal collaterals between the right anterior cerebral artery and MCA was confirmed (figure 1A,B).

Systemic anticoagulation with 5000 U of heparin was initiated. Via a 6 F Envoy guiding catheter (Cordis, Miami, USA) in the right ICA, a straight Rebar27 was gently advanced to the thrombus under the guidance of a SilverSpeed16 microguidewire (both ev3). Under road map, a thrombectomy brush (pCR, phenox; Bochum, Germany) was deployed in the right angular artery and subsequently slowly withdrawn under continuous aspiration. The brush was loaded with clot material but no recanalization of the target vessel was observed (TICI 1). During the catheterization, the thrombus appeared organized and firm. The chances of success of either LIF or repeated pCR passages were considered low as previous experience has shown very low recanalization rates with pCR and LIF in firm organized emboli. It appeared evident to the operator (HH) that in this situation, more interaction between clot and retrieval device was required. While contemplating the use of either a Catch or a Merci retriever (at this time the only other approved devices for mechanical thrombectomy), we recognized that neither of these

Figure 1  Embolic occlusion of the right middle cerebral artery (MCA), treated by mechanical thrombectomy using a Solitaire stent. (A) Digital subtraction angiography (DSA) with injection of the right internal carotid artery (ICA) shows embolic occlusion of the right M1 segment. (B) In the late phase of this DSA run with injection of the right ICA, the leptomeningeal collaterals between the right anterior and MCA were visualized. (C) Solitaire stent loaded with thrombus, which was removed from the right M1 segment. (D, E) DSA with injection of the right ICA after Solitaire thrombectomy confirms the recanalization of the short M1 and both M2 segments (D, magnified view) without vasospasm or distal emboli (E, whole head run). There might be a minor remnant of the previously occluding thrombus in the inferior trunk to the MCA/M2 (D). (F) CT 22 months after the Solitaire thrombectomy of the right MCA shows a small post-ischemic scar of the right basal ganglia.
tools was in stock. After balancing the potential risks and benefits to this patient and given the uncertain outcome if there was no recanalization, we decided to use a Solitaire stent in the way we would have otherwise used a Catch or a Merci device. The right angular artery was again gently catheterized with a Rebar18 catheter. After removal of the microguidewire, a self-expanding Solitaire stent (4 mm diameter, 20 mm length) was inserted under continuous slow flush with saline solution to the distal end of the microcather and was deployed by slowly pulling back the microcather. Correct deployment of the Solitaire stent over the whole length of the thrombus was observed under continuous fluoroscopy. In order to avoid mechanically induced vasospasm, 2 mg of glycerin trinitrate were injected intra-arterially. After several minutes of incubation and with the rotating hemostatic valve tightly closed, a 50 ml Luer lock syringe was used for high volume aspiration through the 6 F guiding catheter during withdrawal of the open Solitaire stent and the microcather into the guide catheter. Withdrawal was possible with minor effort and was observed under continuous fluoroscopy. As soon as the device became visible, the rotating hemostatic valve was quickly removed from the guiding catheter and another 50 ml of blood were aspirated. The Solitaire stent was loaded with a thrombus (figure 1C). Contrast injection of the right ICA after removal of the Solitaire confirmed total recanalization of the right MCA (figure 1D,E; TICI 3). Neither peripheral emboli nor vasospasm at the previously occluded vessel site was encountered.

The entire procedure (from the first to the final DSA run) lasted 59 min. The patient awoke from general anesthesia with only mild neurologic deficit (NIHSS 2). The patient was discharged to a rehabilitation center. At the 30 day follow-up, NIHSS was 0.

The patient was heparinized and received 100 mg of acetylsalicylic acid daily as secondary prophylaxis. ECG monitoring revealed atrial bradillation as a potential source of cardiac ignition spark for these long awaited developments.

In the meantime, the safety and efficacy of the Solitaire FR and the Merci retrieval device were compared in a randomized trial which started in the USA in March 2010 (Solitaire FR With The Intention For Thrombectomy, SWIFT; http://clinicaltrials.gov identifier: NCT01054560). In March 2011, enrollment for this trial was stopped as the sponsor (Covidien/ev3) expects 510 (k) approval by the FDA based on the trial results.

Competing interests The senior author is the co-inventor of the Solitaire stent but has no further interest in this product. ev3 has paid the open access charges for this article.

Patient consent Obtained.

Contributors MAP, EM and SF wrote the first draft of the manuscript, which was revised by HB and HH. HH performed the endovascular treatment. HB reviewed the neurological aspects.

Provenance and peer review Not commissioned; externally peer reviewed.

REFERENCES