In Vitro Studies of the Neuroform Microstent Using Transparent Human Intracranial Arteries

For better understanding of relevant morphology and mechanics, direct visualization of a Neuroform microstent (NFM) within an actual human intracranial artery is essential. Twelve NFM were deployed into 8 various segments of formaldehyde-fixed cadaver intracranial arteries. The arteries were then dehydrated and cleared in methyl salicylate to create transparency. The morphology of NFM was studied by digital macro-photography with a back illumination system. The possible limitations and important findings of the NFM were discussed.

The Neuroform microstent (NFM; Boston Scientific/Target, Fremont, Calif) is the first microcatheter-delivered stent used for treatment of intracranial wide-neck aneurysms. Despite our substantial clinical experiences with the NFM since its availability, a variety of technical difficulties have been encountered that are difficult to resolve, at least in part because of the poor visibility of most of the NFM under fluoroscope and angiography. The purpose of this study is to better assess relevant morphology and mechanics of the NFM stent by direct visualization within actual human intracranial arteries, by using a novel adaptation of a method for creating transparent vessels.

Description of Technique

Intracranial arteries were harvested by dissection from formaldehyde-fixed cadaver brains and skull bases available from donors to the Deeded Body Program at The University of Iowa, College of Medicine. Multiple arterial segments, including the cavernous portion of the internal carotid artery (ICA), the distal-ICA-to-middle-cerebral-artery (MCA) segment, the distal-ICA-to-anterior-cerebral-artery (ACA) segment, the vertebrobasilar artery (VBA), the basilar artery (BA), and the BA-to-posterior-cerebral-artery (PCA) segment were obtained. The vessels were then cleaned by removal of blood clots and any attached arachnoid tissue, followed by storage in a 1% formaldehyde solution.

Understanding the structure of the NFM is essential for assessment of its use (Fig 1). The NFM is made of a nickel-titanium alloy known as nitinol, which possesses a high degree of elasticity and deformability because of its self-expandable properties. Most the stent is radiolucent except the bilateral platinum end markers. The stent is constructed by segmental connections of a variable number of sinusoidal crown segments, the so-called open cell design. Each crown segment length is approximately 2.5 mm. Therefore, a 20-mm length stent used for treatment of intracranial wide-neck aneurysms.

NFM has 8 crown segments. Adjoining crown segments have 2 connection points (struts) that are approximately 180° in orientation and approximately 90° difference in between adjacent struts. Neuroform2 microstent (NF2) is the second generation of NFM with a minor modification of the end platinum markers and improvement of the microdelivery system. The structure of the stent body and the characteristics of NF2 are the same as NFM.

Using the push-pull technique similar to clinical application, NFM (including NF2) were deployed into specific locations of arteries immersed in a Petri dish filled with warm saline. The temperature was maintained at 38°C for obtaining maximum expansion of the NFM. There were 2 purposes for the selected locations of NFM deployment; first, the location of common aneurysms seen clinically, and second, those locations that would mimic the neck of an aneurysm. The junction at larger caliber side branches, such as the conjunction point of a vertebrobasilar artery or a major arterial bifurcation, could be considered the neck of an aneurysm. Stent deployment across major side branches of a parental artery was used to simulate an artery with an aneurysm. To simulate the locations of NFM deployment commonly seen in the clinical setting as well as the mimicked neck of an aneurysm, the intrapetrous portion of the ICA, the cavernous portion of the ICA, the distal ICA to MCA, the ACA from A1 to A2, and the distal BA to PCA were selected as NFM deployment locations. Stent diameter was selected to be either nearly exact or oversized. No actual intracranial arterial aneurysms were available for our study.

Arteries with NFM in position underwent a standard dehydration process beginning with 70% alcohol solution and followed by 95% and 100% alcohol solutions at room temperature. The artery was kept in each solution for at least 24 hours. The dehydrated arteries were then immersed in 100% methyl salicylate (Fisher Scientific, Fair Lawn, NJ) and became transparent within 20 minutes (Fig 2). The transparent arteries with
NFM in place were kept in a Petri dish filled with methyl salicylate and put on a homogenous ground-glass, back-illumination system at room temperature. Macro-digital photography was taken from multiple angles to view the intravascular NFM morphology. The relationship between the stent and vascular wall as well as the stent and the mimicked aneurysm neck (the side branch that the NFM crosses) were studied. Different locations of the stents with their characteristic morphologic changes in specific arterial locations were also investigated.

Twelve NFMs were deployed into 8 various intracranial arterial segments (Figs 3–8). Good conformity as well as a smooth profile was generally observed within most of the arterial segments. However, increasing vessel curvature caused focal intercrown separation (diminished scaffolding) along the greater curvature of an artery. Conversely, along the lesser curvature of an arterial segment, there was a tendency for crowns to imbricate (dot-line arrows). Part of the crown of the stent protrudes into the major side branch, which is mimicking the neck of an aneurysm (arrowhead).

Owing to the “open cell design,” markedly asymmetric enlargement of a cell or multiple cells can occur, particularly when the stent was deployed into the segments with angulation. This was commonly seen in stents deployed across the distal BA to the PCA continuum (Figs 4 and 5) or within the anterior genu of the carotid siphon as well as petrous or/and cavernous ICA (Fig 6). In addition to larger gaps along the greater curvature of an artery, the crown segments consistently did not attach well to the vessel wall along the lesser curvature of cavernous and petrous ICA. In segments where there was an abrupt change in diameter within the curvature (eg, BA-PCA continuum), the connecting struts could rotate to within less than the ideal 180° orientation, which resulted in

Fig 2. Comparison of a vertebrobasilar artery with its major branches and proximal posterior cerebral arteries before (A) and after (B) the artery was made transparent. Two Neuroform microstents are in position (arrowheads). One stent is across the left vertebral and basilar artery and another is located across the distal basilar artery and posterior cerebral artery. The stents are clearly visualized through the artery after arteries are made transparent (B). Using this method, assessment of the mechanical behaviors of Neuroform microstent in actual human intracranial arteries is possible. Extensive atherosclerotic plaques (arrow) are noted in this specimen that involve the right vertebrobasilar junction, the distal basilar artery, and the bilateral posterior cerebral arteries.

Fig 3. A 4.5 × 15-mm Neuroform microstent in a transparent basilar trunk (A) and another 4.5 × 15-mm Neuroform microstent across another transparent vertebrobasilar junction (B). Overall, the Neuroform microstents show very good conformity to the vessel wall. Increasing curvature may cause focal intercrown separation (diminished scaffolding) along the greater curvature of an artery (arrow). Conversely, along the lesser curvature of an arterial segment, there is a tendency for crowns to imbricate (dot-line arrows). Part of the crown of the stent protrudes into the major side branch, which is mimicking the neck of an aneurysm (arrowhead).
a large and asymmetric intersegmental gap at the opposite end of the stent (Fig 4). Good end-crown attachment to the vessel wall was not always achievable, especially when the stent met a curve (Fig 7). Atherosclerotic plaques were observed in several specimens, which caused focal areas of poor stent attachment to the vessel wall.

We observed a connecting strut positioned at the origin of a major side branch and considered that this could happen at the neck of a true aneurysm (Fig 8). The continuity of the stent profile was smooth without protruding part of the crown into the side branch. The NFM is self-expandable and maintains a straight, tubular configuration once expanded. Although flexible and able to adjust to bends or curves present in the vasculature, the stent will automatically straighten out provided an opportunity, and so a small longitudinal pressure is exerted along the vessel wall at any points of contact that inhibit the stent from straightening out to its original configuration. This longitudinal rebounding force straightened a small arterial stump in one of the models (ACA complex, A1 to A2 in Fig 8).

To minimize this, a thin sheath was fixed between the trunk of the BA and the PCA before stent deployment to maintain the original angulation (Fig 5).

**Discussion**

Wide-neck aneurysms have been a challenging scenario for endovascular treatment. Several techniques and endovascular devices have been developed for the treatment of wide-necked aneurysms, including balloon-assisted neck remodeling, 3D coils, and stent assisted coiling. The NFM is the most recently developed endovascular microstent with self-expandable and micro-delivery properties that are specially designed for the treatment of intracranial wide-neck aneurysms by using stent-assisted coiling. However, the NFM is radiolucent under fluoroscope except the bilateral platinum end markers. Simulations of arteries with transparent characteristics such as silicon models have been attempted. This model may provide similar vascular morphology, elasticity, and the anatomic location of aneurysms, but the biologic characteristics of the arterial wall are completely different. The deployment and positioning of the NFM in this model is not adequate to simulate actual human intracranial arteries. Glass tube models have perfect transparency but do not simulate the vascular morphology, anatomic location, or vessel wall characteristics of actual intracranial arteries.

Methods for producing transparent arteries were developed several decades ago. This is the first study to use such methods for investigation of the NFM in human cadaver intracranial arteries. Some of the physical properties of “transparentized” arteries are different from in vivo human arteries, which may be a limitation of this study. First, the calibers of the arteries become smaller (approximately 10%–20%) because of fixation and dehydration processes. Second, the arteries become...
much less elastic and more rigid compared with the in vivo arteries, again because of the fixation and dehydration. Another greater limitation is that the transparent arterial model cannot simulate the dynamic status of in vivo arteries because their physical properties changed. The simulation of pulsatile blood flow was not re-established in our transparent arteries when we deployed the NFM and observed the morphologic changes of the NFM. Although this was the case, however, their anatomic-geometric characteristics are largely well preserved compared with their original configuration. In addition, the NFM was deployed before the dehydration process, when the artery was much less rigid and possessed its nearly normal caliber. Thus, the current study of NFM is capable of providing important insights for the device’s postdeployment steady behavior in actual human intracranial arteries, which cannot be achieved in the clinical setting.

The NFM is made of nitinol, a shape memory alloy. The austenite temperature of the NFM is body temperature, at which the stent will transform from martensite to austenite, the original cylindrical configuration of the stent. The deployment of the NFM occurred at body temperature for obtaining maximum expansion of the stent. Once a stent was deployed,
the crown segments of the stent attached to the vessel wall would not easily move or change shape because the dehydration and clearing as well as the observation process in our study is a steady state. The configuration and morphology of NFM in transparent arteries at room temperature were believed to be nearly the same as those initially deployed at body temperature.

Although no actual aneurysms were present in the specimens used, NFM deployed across large-caliber side branches or bifurcation points can be viewed as an anatomic equivalent to an aneurysm neck. As such, this study clearly showed that stent elements can either protrude significantly into the aneurysm or be poorly oriented to provide inadequate scaffolding. The ability of the NFM to properly bridge the neck of an aneurysm seems to be affected by at least a few factors, including the degree of curvature of the artery, stent sizing, location of the connecting strut, and vessel diameter transitions. Poor scaffolding of the NFM was frequently observed within curved segments of the vessels, particularly along the region of greater curvature or at points of angulation. The connecting strut of the stent could randomly abut the origin of a side branch; although in most cases this would probably not have adverse consequences, very small branches could suffer direct or indirect flow obstruction or disturbances. Conversely, the strut may be positioned at the neck of an aneurysm and thereby provide an ideal scaffolding support for Guglielmi detachable coiling (Fig 8B).

A NFM deployed across an artery with angulation exerts a certain longitudinal rebounding force and could straighten the angle of the parent artery (Fig 8). This phenomenon could have adverse consequences, such as proximal or distal arterial kinking, dissection, asymmetric mechanical strain on the aneurysm, or increased compliance mismatch at the stent ends, which might cause marginal or in-stent stenosis. The manufacturer recommended that the NFM needs 4 mm of length in both proximal and distal ends of the parental artery for securing the stent in position. We believe that in arteries with angulations, the NFM may need more “secure length” on both ends because of the longitudinal rebounding force. Therefore, if the secure length is not long enough, the force may cause the distal end of the stent to dislodge and jump into the aneurysm. We have experienced one such case but fortunately the aneurysm was safely coiled after a second NFM was deployed by using a stent-through-stent method similar to that described by Broadbent et al.

Although atherosclerosis and arterial aneurysm are 2 different disease entities, the intracranial artery harbored with an aneurysm may rarely be associated with atherosclerosis of the parental artery. In atherosclerotic arteries, the lumen is narrow and irregular, and we observed that the attachment of NFM end-markers to the vessel wall was not very good and that the NFM might experience less expansion. In addition, in arteries with angulations, the NFM has larger gaps in its body than expected. This seems to be the case in Fig 3, of the BA-PCA continuum, which also reveals extensive atherosclerotic plaques. In this case, the gaps of the NFM are probably larger than what one would observe in cases involving nonatherosclerotic associated aneurysms at this particular location.

The strut number for each segment of the NFM is an important factor that may cause such larger gaps within the stent along arteries with angulations. As described previously, there are only 2 struts for connecting each segment of the NFM. To improve this adverse effect, the third generation Neuroform3 (NF3; Boston Scientific/Target) was recently made available. The NF3 has 3 struts for connecting each segment over the middle crowns of the NFM. Ideally, increasing one strut will improve the stent profile and perhaps produce better scaffolding for containing coils in aneurysms that are located at curves or within angulated arteries.

Conclusions

Transparent human arteries are excellent models for evaluating certain important physical properties and the morphology of radiolucent stents such as the NFM. Although this model is not a complete simulation of in vivo human arteries because of fixation and dehydration processes producing smaller, more rigid arteries, the ability to directly visualize the stent and its relationship with vascular walls provides very important information that may not be obtainable in clinical settings. Our preliminary studies revealed several significant findings regarding the NFM. Proper sizing seems to be important for ensuring good stent conformity within the arteries, particularly when significant curvature is encountered. NFM deployment in arterial segments or branches with angulation could be problematic, potentially resulting in excessive gaps in between stent crown segments that probably will not provide adequate scaffolding for containing coils within an aneurysm. The operators should be aware that longitudinal rebounding force exists when the NFM is deployed at curved or angulated positions. Different complications may arise from this longitudinal rebounding force.

References