A ngioplasty was first used in 1977 for dilating atherosclerotic plaques in coronary arteries. Although restenosis after angioplasty is a common occurrence in atherosclerosis cases, it has only been reported once after the use of angioplasty for cerebral vasospasm. Sedet et al. reported delayed stenosis of the middle and anterior cerebral arteries. In this paper, we report delayed stenosis of the ICAs. To our knowledge, this is the second reported case of delayed stenosis following angioplasty for treatment of cerebral vasospasm.

**Case Report**

This 53-year-old woman suffered severe SAH (Hunt and Hess Grade IV) due to a ruptured posterior communicating artery aneurysm. Initial CT angiography revealed widely patent ICAs and an associated right posterior communicating aneurysm. The patient underwent a frontotemporal craniectomy on an emergency basis and clip ligation of the aneurysm and evacuation of the intracerebral hematoma were performed. Seven days after admission, CT angiography revealed severe diffuse vasospasm. As a result, angioplasty of the left and right ICAs and middle cerebral arteries was conducted with a Hyperglide balloon (ev3, Inc.). After the angioplasty procedure, verapamil was infused intraarterially to dilate more distal branches. Angioplasty of the right M1 and M2 segments was then performed after exchanging the microcatheter for a Maverick balloon catheter (Boston Scientific Corp.). Bone plate replacement was performed 2 weeks later. The following week, the patient underwent placement of a ventriculoperitoneal shunt on her left side due to hydrocephalus resulting from SAH. At 1-month follow-up, the patient exhibited signs of fatigue, diminished vision in both eyes, and short-term memory loss.

Despite continual fatigue and short-term memory loss at 1 month follow-up, the patient improved neurologically and had no focal neurological symptoms at 3 months. A follow-up CT angiogram obtained 6 months postoperatively revealed bilateral ICA stenoses that were
especially prominent in the supraclinoid region. A CT perfusion study suggested diminished perfusion bilaterally. Repeat diagnostic cerebral angiography 9 months after angioplasty confirmed persistent stenosis in bilateral ICAs (Figs. 1 and 2).

Discussion

Vasospasm is common in patients with aneurysmal SAH. It is a leading cause of cerebral ischemia, and hence death and injury after an aneurysm rupture. Approximately 50% of SAH cases result in death while 15% result in disability. Research suggests that nitric oxide, endothelin-1 and bilirubin oxidation products, and inflammation are important contributors to cerebral vasospasm.

Angioplasty, although initially used in coronary atherosclerosis, was first shown to be an effective treatment for vasospastic arteries in 1984 by Zubkov et al. It works by dilating the constricted arteries via a balloon catheter and thus preventing ischemic conditions. Clinical studies by Newell et al. and Bejjani et al. have alluded to its efficacy and safety. However, the case report presented here, being only the second reported case of restenosis after cerebral vasospastic angioplasty, suggests that there can be delayed effects of angioplasty for the treatment of vasospasm. This finding is important as this effect may not be unique and continued surveillance imaging of patients who undergo angioplasty for vasospasm may be required. Indeed, we are considering bypass surgery or stent placement to restore normal cerebral blood flow in our patient, but given the unusual circumstances and the patient’s continued neurological recovery, we have not decided on therapy yet.

Pathophysiology

A study by Chavez et al. has shown that transluminal angioplasty leads to endothelial cell denudation, stretching and rupture of internal elastic lamina, and changes in smooth muscle cells of the medial layer, but these changes resolved within 7 days. Yamamoto et al., however, showed long-term connective tissue damage in the medial and adventitial layers from the disruption of the arrangement of collagen fibers due to its stretching and tearing. Since extracellular matrix is an important medium in the transmission of contractile forces, its disruption may explain the occurrence of delayed stenosis.

A study by Wakayama et al. investigated the pathophysiology of restenosis with the use of a transluminal vascular injury model in rodents. The idea behind the use of this model was to create a suture with a diameter larger than that of a rat’s ICA, such that the vessel would be dilated when the suture was advanced. Specifically, a 25-mm length of 3-0 surgical suture was modified with an epoxide resin–coated region at the tip, 0.6 mm in diameter and 5 mm in length. For 1 minute, the length of suture material was inserted into the external carotid artery and advanced 18–20 mm into the ICA, leading to rapid enlargement of the ICA (normal luminal diameter is 0.3 mm). Following injury using this intraluminal suture technique, there was loss of endothelial cells and the...
smooth muscle cells of the medial layer. However between 1 and 4 weeks afterward there was a reemergence of the neointimal layer composed of smooth muscle cells and the surrounding extracellular matrix. This new intimal constitution may explain the contractile properties of restenosis.

Rubin et al. explained that the occurrence of restenosis may be mitigated by inflammation. They showed that within 24 hours after injury macrophages begin migrating into the adventitial layer. Within 1 week, they spread to the medial and intimal layers but were more prominent in the latter. Neointimal hyperplasia was evident within 3 weeks, during which time the presence of macrophages could also be detected.

Finally, injury to blood vessels as a result of angioplasty can lead to activation of reactive oxygen species at the injury site from molecular oxygen. One such reactive oxygen species, the superoxide radical, exerts numerous effects such as inactivation of endothelium-derived relaxing factor, lipid hydroperoxide–mediated modulation of vascular tone, and platelet activation. Although generated as part of a repair reaction, these effects can lead to occurrence of restenosis.

Applicability of This Report

Angioplasty for vasospasm has been an important adjunct for the treatment of cerebral ischemia in the setting of SAH. Though the efficacy of the procedure has not been proven with Class I evidence, it is thought to be effective if administered within a short time after ischemic symptoms develop. This report simply reminds treating physicians that there can be delayed consequences to this therapy and follow-up surveillance imaging is important in patients who have undergone this treatment for vasospasm.

In addition, the use of the compliant softer balloons may play a role in the development of delayed stenosis. In both our case and the previously reported case, Hyperglide balloons were used. These soft balloons inflate to between 3 and 5 mm in diameter and so may stretch the vessel beyond its normal size. The advantage of this technique is that the vessel appears to be restored to normal size immediately and rarely requires retreatment for vasospasm at the same site, but the potential disadvantage is that stretching the vessel beyond its normal diameter may instigate the above-proposed mechanisms leading to the development of delayed stenosis. While noncompliant coronary balloons can also be used to treat vasospasm, the diameter of the balloon is selected to be smaller than the normal vessel (for example, for a 3-mm ICA, a 2.5-mm–diameter balloon may be selected). Hence, when the balloon is fully inflated the vessel is not stretched be-
yond its normal diameter. One potential risk associated with these types of balloons is the chance of vasospasm recurring in the vessel after treatment. If the chosen balloon diameter is too small (for example, if a 1.5-mm balloon were to be selected in the above example) then there remains a chance the vessel will not be stretched enough and repeated vasospasm may occur during the acute period. The noncompliant balloons are also stiffer, so they may be harder to track to the affected vessel. In addition, they tend to straighten the artery, so if the balloon is too long, it may kink or injure the treated vessel. In this reported case, we used both kinds of balloons, and the delayed stenosis only occurred in the area treated by means of the compliant balloons.

**Conclusions**

We report the unique finding of delayed stenosis following angioplasty of vasospastic ICAs using compliant balloons. The stenoses occurred bilaterally at the same site. Fortunately, there were no focal neurological symptoms attributable to these stenoses in our patient, but this case report should alert practitioners of this potential complication. Surveillance imaging is important after angioplasty for vasospasm and should be performed in a delayed fashion. Treatment of the delayed stenosis could include stent placement or intracranial bypass procedures.

**Disclaimer**

The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

**References**


Address correspondence to: Alan S. Boulos, M.D., Division of Neurosurgery, Albany Medical Center, 47 New Scotland Avenue, Albany, New York 12208. email: alan.boulos@gmail.com.