Distal Shower Embolization During Directional Coronary Atherectomy and Stenting for Diffuse Stenosis of Right Coronary Artery

— Current Limitations of Intravascular Ultrasonography for Evaluating Fragile Plaque —

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A 74 year-old male with old anterior and inferior myocardial infarctions was treated with staged percutaneous coronary intervention. A chronic total occlusion of the middle segment of the left anterior descending branch was successfully stented during the first stage, and during the second stage, preprocedural intravascular ultrasoundography (IVUS) revealed that the proximal segment of the right coronary artery was diffusely stenosed by mixed plaque. Directional coronary atherectomy under IVUS guidance was performed, but coronary slow flow appeared during the procedure. After successfully bailing out with intracoronary nicorandil, percutaneous thrombectomy and manual blood pumping, 2 coronary stents were implanted to fully cover the lesion. Quite contrary to expectation, the no-reflow phenomenon appeared just after post-dilatation and a repeat of the same maneuver could not completely recover coronary flow. Plaque fragility could not be predicted from the IVUS examination. (Circ J 2004; 68: 257–262)

Key Words: Coronary intervention; Intracoronary imaging; Intravascular ultrasonography; No reflow phenomenon; Plaque

Intravascular ultrasonography (IVUS) has been accepted as a useful tool for assessing target lesion plaque, selecting the most suitable strategy and deciding on the endpoint of a procedure. We report a patient with the complication of distal shower embolization both during directional coronary atherectomy (DCA) and just after stent implantation despite preprocedural IVUS images suggesting that the target plaque was solid enough to be treated with those devices. This is a rare case in which the current limitation of IVUS examination was manifested.

Case Report

A 74 year-old male with diabetes and a past history of smoking was admitted with symptoms of chest discomfort on effort. Seventeen years earlier, he suffered from an acute anterior myocardial infarction and ventricular septal perforation (VSP), which were successfully treated with left ventricular aneurysmectomy and VSP closure after conservative therapy. Despite careful follow-up, he again suffered an acute inferior myocardial infarction 5 years before the current admission and was treated medically.

Coronary angiography (CAG) revealed that the middle segment of the left anterior descending branch (LAD) was totally occluded and the proximal segment of the right coronary artery (RCA) was diffusely stenosed with poor collateral circulation to the LAD (Fig 1). We planned a

![Fig 1. Diagnostic coronary angiography. (a) Left coronary artery from the antero-posterior cranial view: the middle segment of the LAD is totally occluded. (b) RCA from the LAO cranial view showing diffuse stenosis of the proximal segment. (c) RCA has poor collateral circulation to the LAD.](image-url)
staged percutaneous coronary intervention (PCI) for jeopardized multivessel disease in order to achieve complete revascularization.

A week later, the first stage of the PCI successfully dilated the chronic total occlusion of the mid LAD and a 3.0 mm Multilink-plus stent (Guidant) was implanted.

Another week later, he underwent the second stage. After insertion of a 10 Fr catheter introducer, the patient was heparinized and the activated clotting time (ACT) was maintained above 300 s throughout the procedure. The RCA was engaged with a 6 Fr JR3.5 Zuma-II guiding catheter (Medtronic USCI) via the right femoral artery. The target lesion was easily crossed with a 0.014-inch Hi-torque Balance Middle Weight guidewire (Guidant) and was examined with a 2.5 Fr Atlantis SR plus imaging catheter (SciMed, Boston Scientific). On that basis, we planned DCA and provisional spot stenting because plaque quality was assumed to be mixed and less calcified (Fig 2c). A short lesion around the bifurcation of the right ventricular branch was superficially calcified, and simple stent placement was thought adequate for dilating this stenosis rather than DCA–stenting (Fig 2c). After the removal of all devices, the RCA was re-engaged with a 10 Fr JCR3.5 Tourguide guiding catheter (Guidant), and the lesion was easily crossed again with a 0.014-inch Neo’s Gland-Slam guidewire of 300 cm length (Getz Bros). After making a small channel with a 2.0 mm Aerocross Fighter balloon catheter (Kaneka Corp), a 7 Fr-GTO Simpson Atherocath...
(Guidant) was advanced. Because the IVUS images suggested that the target plaque was eccentrically distributed toward the major curvature (Fig 2c), the atherocatheter was manipulated to turn its window upward, projecting from left anterior oblique view. The first 3 cuts with the directional balloon inflated at 10 psi were tentatively performed, rotating the atherocatheter clockwise by 45 degrees on the second cut and by the same degree counterclockwise for the third cut. The atherocatheter was removed leaving the long Neo’s guidewire crossing the lesion because there was little resistance between the guidewire and the wire lumen of the atherocatheter. We found only a small amount of plaque debris in the nose cone and when debulked it looked white, suggesting that its quality was fibrous as indicated by the IVUS images. The atherocatheter was again advanced to the target lesion and manipulated in the same manner except for balloon pressure of 20 psi not 10 psi (Fig 3). The atherocatheter was pulled back into the guiding catheter, and the subsequent angiography showed slow flow phenomenon (Fig 4a). After removal of the atherocatheter, 2 mg of nicorandil were transluminally injected and additional percutaneous thrombectomy with a 4.3 Fr Rescue-PT catheter (SciMed, Boston Scientific) and manual blood pumping for a few minutes recovered the coronary flow. Although a small dissection was detected at the debulked lesion on both angiography and IVUS (Fig 4b), antegrade flow in the RCA began to be disturbed at the acute margin, not at the target lesion. The possibility that debulked plaque debris had dislodged from the gap in the guidewire lumen at the nose cone was weak because the atherocatheter was carefully manipulated and the resistance between the atherocatheter and guidewire had been low after the 20-psi debulking. Collapse of fragile plaque that appeared solid on preprocedural IVUS images (Fig 2c) was considered to be the cause this complication, and dilatation of the
residual stenosis by stent placement was thought more appropriate treatment, on the expectation that stenting would improve forward flow and salvage coronary microcirculation. A 3.5 mm Multilink-plus stent (Guidant) and a 4.0×24 mm S670 stent (Medtronic AVE) were deployed at the distal and proximal segments of the lesion, respectively, but the no-reflow phenomenon appeared quite unexpectedly just after post dilatation with the system balloon of the previous S670 stent (Fig 5). Distal shower embolization by a massive amount of plaque debris squeezed out between the stent struts was considered the cause of this serious complication. Transluminal injection of nicorandil, percutaneous thrombectomy and manual blood pumping were again performed, but could only restore the coronary flow to a mild slow-flow level (Fig 6a). Postprocedural IVUS images revealed that the plaque burden had significantly reduced compared with the preprocedural images (Fig 6b). Insufficient expansion of the S670 stent was also revealed, but we did not carry out supplementary high-pressure ballooning because of the risk of recurrence of impaired coronary flow. The risk of in-stent thrombosis was considered low because the struts were adhered closely to the vessel wall and the minimum stent diameter was over 3.0 mm. A 9.5 Fr intraaortic counterpulsation catheter was inserted via his left femoral artery and he was transferred to the coronary care unit.

Discussion

The target lesion of the second stage of the present PCI was diffuse and the indications for and strategy of PCI for long or diffuse lesions are controversial. Although simple stenting without debulking has been reported as efficacious against long or diffuse stenosis, even in the setting of multiple short-stent implantations or single long-stent implantation, the fact that long or diffuse lesions belong to a sub-
Distal Shower Embolization During DCA Stenting

Several kinds of covered stent are commercially available, such as saphenous vein grafts (SVG), with the no-reflow phenomenon having been reported in up to 7.9% of cases, even without thrombus. Several kinds of covered stent are commercially available and one recently proved to be effective in reducing SVG flow impairment. Although the target lesion in the present case was located in a native coronary artery, simple implantation of a coronary stent graft might have been a therapeutic option. In cases of acute coronary syndrome (ACS), this phenomenon also occurs with considerable frequency even in native coronary arteries. However, the frequency of embolized debris in routine PCI of native vessel has not been evaluated. The first clinical results of the effectiveness and safety of the AngioGuard Emboli Capture Guidewire (Cordis, Johnson and Johnson) was recently reported in a study by Eberhard et al in which only 11 SVG lesions, but also 15 native vessel lesions were enrolled and all cases were electively treated. Surprisingly, there were particles with a mean size of 0.10±0.5 mm² and a range of 0.015–20 mm² in the filter basket and capture sheath of all cases. Therefore, there is a risk of distal embolization even with PCI in a native vessel, although the relationship between cardiovascular imaging and plaque fragility that will probably lead to distal embolization has yet to be elucidated fully.

In the setting of PCI for ACS, unstable or vulnerable plaques with sonolucent echogenicity are often found in an IVUS study. If conventional balloon angioplasty is performed for treatment of such lesions, distal embolization by atheromatous debris or intracoronary thrombus is often complicated and may lead to ischemic complications. Mixed fibrocalcific plaques such as in the present case do not appear to be destroyed by balloon compression, but in our case distal shower embolization was a quite unexpected complication. We used a 40 MHz transducer for the IVUS assessment because, compared with the conventional 20–30 MHz transducers, the high frequency IVUS transducers have been reported by Prati et al to be more effective in identifying lipid/necrotic pools in atherosclerotic plaques. In that study, lipid pools were observed by histology in 30 sections, but IVUS with a 40 MHz transducer revealed the presence of lipid pools in only 19 of them, so it must be emphasized that not all unstable plaques are detected during IVUS examination.

The present case may be in a minority subgroup in which IVUS cannot provide precise information. Optical coherence tomography (OCT) is a new modality for high-resolution, cross-sectional, intravascular imaging. IVUS is currently of little use in determining the stability of subendothelial plaque components because of its limited resolution (>100 µm even for 40 MHz systems) and the confounding influence of surrounding tissues. If IVUS systems become available with a 50 MHz transducer, the maximum resolution will still be no greater than 70 µm. In contrast, the maximum resolution of OCT is limited physically only by the diffraction limit of light at 1,300 nm, which is <100 nm, and systems are already available with axial resolutions less than 4 µm. Such systems may ultimately be of use in cardiology for the identification of subcellular structures and in fact, postmortem examination of human coronary arteries has shown that catheter-based OCT systems give superior delineation of vessel layers and lack of the ring-down artifact with higher resolution than a 30 MHz IVUS system. The greatest limitation of OCT is its reduced effectiveness when imaging through blood, which also applies to Raman spectroscopy and near-infrared spectroscopy.

The mechanism of the no-reflow phenomenon is probably multifactorial and involves small-vessel vasospasm and potentially platelet-mediated loss of capillary autoregulation. In the setting of PCI for acute myocardial infarction, endothelial damage, including endothelial swelling and myocyte edema, leads to initial no reflow zones, and additional edema, myocyte contraction, platelets, fibrin and leukocyte plugging result in expansion of the no reflow regions. However, in the setting PCI of routine native vessel such as in the present case, initial embolization of the atheromatous debris, as well as blood clots and platelet plugs, seems to have been the trigger for the cascade of events that lead to microvascular vasoconstriction. In particular, several investigators have suggested that platelet activation plays a pivotal role in this cascade and so the use of glycoprotein IIb/IIIa inhibitors might improve distal runoff.

A variety of new, innovative devices for distal protection are now in clinical studies and the research phase, and have proven to be effective in preventing impaired flow during catheter therapies. The introduction of these devices, as well as more potent antiplatelet agents, into real clinical settings will help reduce the incidence of impaired coronary flow during catheter interventions. However, precise procedural assessment with intracoronary imaging of plaque fragility that could lead to distal embolization systems is most essential.

**Conclusion**

IVUS assessment is essential for safe PCI. In most cases the images are consistent with histopathological findings,
but it should not be ignored that there are a small number of cases in which that principle cannot be applied. New intravascular imaging systems with higher resolution and less artifacts will improve both detection and evaluation of atherosclerotic plaques before interventional procedures, and new antiplatelet agents and distal protection devices are expected to reduce the no reflow phenomenon.

References