Prediction of No-Reflow Phenomenon After Successful Percutaneous Coronary Intervention in Patients With Acute Myocardial Infarction

Intravascular Ultrasound Findings

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Plaque characterization by intravascular ultrasound (IVUS) before percutaneous coronary intervention (PCI) was evaluated in 81 consecutive patients with acute myocardial infarction (AMI) to establish if IVUS can predict the occurrence of the ‘no-reflow’ phenomenon. Angiographic no-reflow was defined as TIMI flow grade 1 or 2 without any mechanical obstruction in the epicardial artery. Patients were divided into 2 groups according to the post-PCI angiograms: normal flow (group R, n=60) and no-reflow (group NR, n=21). Although the incidence of either soft or noncalcified plaque was not statistically different between the groups, positive vessel remodeling was more frequent in group NR than in group R (57.1% vs 31.6%, p<0.05). Lipid core was also more frequently found in group NR than in group R (61.9% vs 25.0%, p<0.01). Positively remodeled vessels with lipid-rich plaques as characterized by IVUS before PCI predicted the occurrence of angiographic no reflow with a sensitivity of 43% and a specificity of 60%.

Key Words: Acute myocardial infarction; Intravascular ultrasound; Lipid core; No-reflow; Remodeling

Percutaneous coronary intervention (PCI), if performed adequately, limits infarct size and improve prognosis in patients with acute myocardial infarction (AMI).1–2 Patency of the infarct-related artery, however, does not always guarantee restoration of normal coronary blood flow. This ‘no-reflow’ phenomenon is related to a poor clinical outcome3 and although a considerable number of patients with AMI who undergo successful PCI are at risk of no-reflow, the mechanisms remain unclear and reliable prediction of the occurrence of no-reflow before PCI is performed has not been established4

No-reflow is frequently encountered in the setting of elective PCI for degenerated, plaque-rich saphenous vein graft stenoses.5,6 Hence, we postulated that the plaque characteristics of the infarct-related coronary artery lesion may be closely related to the occurrence of no-reflow.7–12 However, intracoronary plaque characterization is not possible using coronary angiography alone. Recently, intravascular ultrasound (IVUS) has emerged as a powerful tool for the evaluation of coronary artery lesion morphology13 so in the present study we decided to perform IVUS prior to PCI in patients with AMI to characterize the infarct-related coronary artery lesion morphology and investigate its relationship to the angiographic no-reflow phenomenon.

Methods

Study Population

The study prospectively enrolled 81 consecutive patients with AMI (58 men, 23 women) who underwent emergency PCI from May 2000 through October 2001. The Ethics Committee of Kansai Rosai Hospital approved the study protocol and all patients gave written informed consent. Clinical data including age, gender, risk factors for coronary artery disease (eg, diabetes mellitus, hypertension, total cholesterol, triglyceride, blood pressure, smoking history, previous MI) were collected. Patients satisfying all of the following criteria were given a diagnosis of AMI: (1) chest pain ≥30 min duration; (2) ST elevation ≥0.1 mV in at least 2 adjacent leads on ECG; (3) increase in creatine kinase (CK) concentration to double the normal value. The examined vessels were 44 left anterior descending arteries, 28 right coronary arteries and 9 left circumflex arteries.

Angiographic Definition of No-Reflow

Coronary flow was angiographically assessed by TIMI flow grade, as defined in the Thrombolysis in Myocardial Infarction (TIMI) trial.14 Flow reduction includes both slow flow and no-reflow. No-reflow was defined in this study as transient or permanent flow reduction as compared with the flow obtained immediately after aspiration with a Rescue™ catheter, equal to or more than TIMI grade 1 that was neither attributable to abrupt closure, severe dissection of the original target lesion, nor epicardial spasm. Slow flow or TIMI grade 2 even after successful PCI was also included as no-reflow. The average TIMI flow grade among all patients was 0.56 initially, 0.97 after wire crossing, 2.26 after...
Rescue, 2.38 after PCI, and was 2.69 by final angiography (Fig 1). Although the overall TIMI flow grade increased after the PCI procedure, TIMI flow grade worsened in some patients after PCI despite additional balloon dilations. Angiographic no-reflow was found in 21 cases (flow reduction: 11 cases; final no-reflow: 10 cases). Patients were accordingly divided into 2 groups based on the post-PCI angiograms: angiograms with normal flow (group R, n=60) and those with no-reflow (group NR, n=21).

**IVUS and PCI Procedures**

For diagnostic coronary angiography, intravenous heparin (3,000 units) and intracoronary isosorbide dinitrate (0.4–0.6 mg) were administered, and the right and left coronary arteries were imaged by the femoral artery approach using a 5Fr catheter to identify the infarct-related coronary artery. After the routine diagnostic catheterization was completed, an 8Fr guide catheter was inserted into the responsible coronary artery followed by an additional 7,000 units of heparin. Next, a 0.014-inch guidewire was passed through the lesion. According to the hospital’s strategy for the treatment of AMI, a Rescue™ catheter was used to aspirate as many intracoronary thrombi as possible to prevent distal embolization. On completion of the aspiration, in all patients underwent IVUS before the PCI procedure. An ultrasound transducer (30 MHz, 3.2F, Ultracross, Boston Scientific Corporation, Natick, MA, USA) was advanced as distally as possible over the guidewire and withdrawn by both slow manual and auto pullback (1 mm/s) methods. While pulling back the catheter, we manually infused a contrast medium to exclude the residual thrombus by improving the lumen/vessel contrast. Ultrasound images were recorded on videotapes.

Cineangiographic documentation of the lesion location was used for site identification.

**IVUS Image Analysis**

After the procedure, the ultrasound images were analyzed using computer software (3D Netra IVUS, Scimage Inc, CA, USA). The culprit lesion site and a proximal reference site were selected for measurement. The culprit lesion was defined as the site with the smallest lumen diameter on the IVUS images. Lumen area was determined by tracing the boundary between the lumen and the leading edge of the intima, excluding residual thrombus. Total vessel area (TVA) was determined by tracing the leading edge of the media–adventitia boundary. Plaque area was calculated as: total vessel area – lumen area. Percent plaque area was computed using the formula: \( \frac{(\text{total vessel area} - \text{lumen area})}{\text{total vessel area}} \times 100 \). Plaque morphology was classified into 3 types: soft, hard and calcified. A soft plaque was defined as plaque that was less echogenic than the reference adventitia; hard plaque produced a similar level of echo intensity as the adventitia; and calcified lesions were defined as echogenic plaque with acoustic shadowing occupying >90° of the vessel wall circumference. The remodeling index (RI) was defined as the ratio of the TVA at the lesion site to the TVA at the proximal reference site: \( \text{RI} >1.05 \) was regarded as positive remodeling.

### Table 1 Patient Characteristics

<table>
<thead>
<tr>
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<th>No-reflow (–)</th>
<th>No-reflow (+)</th>
<th>p value</th>
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<tbody>
<tr>
<td>Age (years)</td>
<td>64.7</td>
<td>65.2</td>
<td>NS</td>
</tr>
<tr>
<td>M/F (%)</td>
<td>42/18 (30%)</td>
<td>16/5 (23.8%)</td>
<td>NS</td>
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<tr>
<td>Previous MI</td>
<td>5 (8.3%)</td>
<td>3 (14.2%)</td>
<td>NS</td>
</tr>
<tr>
<td>Previous CABG</td>
<td>5 (8.3%)</td>
<td>3 (14.2%)</td>
<td>NS</td>
</tr>
<tr>
<td>Cardiogenic shock</td>
<td>3 (5.0%)</td>
<td>3 (14.2%)</td>
<td>NS</td>
</tr>
<tr>
<td>Peak CK</td>
<td>3378±2761</td>
<td>3776±3107</td>
<td>NS</td>
</tr>
<tr>
<td>Peak CK-MB</td>
<td>29±222</td>
<td>336±266</td>
<td>NS</td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>15±20 (25%)</td>
<td>6±21 (28.5%)</td>
<td>NS</td>
</tr>
<tr>
<td>Hypertension</td>
<td>26±60 (41.7%)</td>
<td>13±21 (61.9%)</td>
<td>NS</td>
</tr>
<tr>
<td>Hyperlipidemia</td>
<td>20±60 (33.3%)</td>
<td>9±21 (42.9%)</td>
<td>NS</td>
</tr>
<tr>
<td>Smoking</td>
<td>34±60 (56.7%)</td>
<td>12±21 (57.1%)</td>
<td>NS</td>
</tr>
<tr>
<td>Final stent use</td>
<td>17±60 (28.3%)</td>
<td>13±21 (61.9%)</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

CABG, coronary artery bypass grafting; CK, creatine kinase; MI, myocardial infarction.
Statistical Analysis
Numerical data were expressed as mean ± SD. Chi-square analysis was used to assess the difference among categorical variables. Student’s t-test was used to assess differences in numerical data. In this study p<0.05 was considered to be significant.

Results
Patient Characteristics
Background characteristics including age, gender, responsible vessel, and coronary risk factors (eg, diabetes mellitus, hypertension, hyperlipidemia, and smoking) were not different between group R and group NR, except for the stent use. Lesions could be crossed with the Rescue™ catheter in 78 patients (96.3%), except for 3 patients, thereby eliminating the residual thrombus at the culprit lesion in the majority of patients. Stents were more frequently used in group NR than in group R (61.9% vs 28.3%, p<0.01, Table 1). Multiple complex coronary stenoses were found 18 (22.2%) of the culprit lesions.

Quantitative IVUS Measurements
A calcified lesion was found in 15 cases (18%), soft plaque in 47 cases (58%) and lipid core in 29 cases (35%). Positive vessel remodeling, as represented in Fig 3, was more frequent in group NR than in group R (57.1% vs 31.6%, p<0.05). A lipid core was also more frequently found in group NR than in group R (61.9% vs 25.0%, p<0.01). Calcification tended to be less frequent in group NR, although it did not reach statistical significance. The presence of both vessel remodeling and a lipid core on IVUS predicted angiographic no-reflow with a sensitivity of 43% and specificity of 60%.

Discussion
We demonstrated in this study that the occurrence of angiographic no-reflow can be predicted by IVUS performed before emergency PCI in patients with AMI. The presence of both positive vessel remodeling and a lipid core was the major predictor of angiographic no-reflow. The no-reflow phenomenon has been attributed to multiple factors including damage to the endothelial cells, tissue
edema, neutrophil plugging of the microvessels, microvascular spasm, oxygen-free radicals, embolization caused by the thrombus burden at culprit lesion, and distal embolization by plaque from the culprit lesion. Considering the experience reported from elective PCI for degenerated saphenous vein grafts, which are often associated with large plaques, distal embolization by materials from the lesion site may play an important role in the occurrence of no-reflow in the acute setting. Thus, it can be anticipated that there is a close relationship between lesion morphology and the incidence of angiographic no-reflow. Accordingly, we used IVUS to characterize lesion morphology before the PCI procedure. Although the lesion vessel area was similar between group NR and group R (15.3 ± 5.7 mm² vs 14.4 ± 4.7 mm², NS), vessel remodeling was more significant in group NR than in group R (57.1% vs 31.6%, p < 0.05). The present study thus demonstrated that both lesions that are lipid rich and remodeled large plaques are associated with the development of angiographic no-reflow.

We performed thrombus aspiration prior to IVUS to exclude as much as possible the occurrence of distal embolization by intracoronary thrombi. Nevertheless, no-reflow still occurred in 21 cases (25.9%), which strongly suggests that the plaque contents, which will be destroyed during the PCI procedure, are also important in the occurrence of no-reflow, in addition to thrombus.

The characterization of plaque was done by IVUS in this study, but although this methodology is well established, it cannot always distinguish thrombi from underlying plaque, and its accuracy depends on the instrument settings and the expertise of the interpreter. Because of this limitation, we used a Rescue™ thrombectomy catheter before IVUS and PCI to eliminate thrombi as much as possible.

Tanaka et al have independently reported that large vessels associated with a lipid pool-like IVUS image indicate a high risk for no-reflow after PCI for AMI, also suggesting that plaque content may play a role in the occurrence of no-reflow. In their study, however, distal embolization by the thrombus burden at the lesion site was not sufficiently ruled out.

Study Limitations

The no-reflow phenomenon was angiographically defined in this study; however, the microcirculation was not directly evaluated, for example, by contrast echocardiography. Hence, the mechanism of the phenomenon remains speculative. Classification of plaque morphology is based on a computer-assisted visual interpretation, which depends on the instrument settings and the expertise of the interpreter. Although aspiration with a Rescue™ catheter had been extensively performed prior to IVUS and PCI, intracoronary thrombus may not have been completely eliminated.

Clinical Implications

New PCI strategies (eg, distal protection devices) could be used to prevent the no-reflow phenomenon if its occurrence can be predicted prior to PCI in patients with AMI.

In conclusion, the presence of both lipid core and vessel remodeling without calcification on IVUS are possible predictors of the occurrence of angiographic no-reflow after PCI in patients with AMI. Destruction of the plaque and subsequent distal embolization may be a cause of no-reflow.

References


