Variations of Remodeling in Response to Left Main Atherosclerosis Assessed With Intravascular Ultrasound In Vivo

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Histopathologic studies have demonstrated that vessels enlarge to compensate for an increase in plaque burden; this has been confirmed in vivo using intravascular ultrasound (IVUS). The initial studies suggested a biphasic course of lesion formation with (1) preservation of lumen dimensions up to a plaque burden of approximately 40%, and (2) luminal narrowing as plaque burden further increases. In this study, we used IVUS and angiography to assess the extent of left main (LM) atherosclerosis in 107 patients undergoing catheter-based procedures of the left anterior descending or left circumflex coronary arteries. Using IVUS, atherosclerotic plaques were found in all LM arteries, but only 26 (24%) had varying degrees of luminal narrowing on the angiogram. Nevertheless, there was an inverse relation (r = −0.62, p < 0.0001) between the minimal lumen area and the plaque burden (i.e., plaque + media divided by total vessel area) that was not restricted to plaque burden values >40% (or >30%), but persisted at plaque burden values of 20% to 40%. In addition, LM arteries with a plaque burden <40% had a similar total vessel area as did LM arteries with a plaque burden ≥40% (22.9 ± 6.1 vs 21.8 ± 4.8 mm², p = 0.30). These data suggest that lumen dimensions may not be preserved even if plaque occupies no more than 20% to 40% of the total vessel area. Thus, there is more variation in remodeling response during earlier stages of plaque accumulation within the LM artery than is commonly suggested. ©1997 by Excerpta Medica, Inc.

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Coronary angiography is used to assess lesion formation; however, early atherosclerosis is angiographically silent.2–5 Histopathologic studies of diseased arterial segments have demonstrated that compensatory enlargement of the vascular wall occurs to compensate for the accumulation of atherosclerotic plaque.5,6 Intravascular ultrasound (IVUS) provides transmural images of coronary vessels in vivo including the coronary vascular wall, the area of atherosclerotic plaque, accurate lumen dimensions, and the serial changes that occur with the atherosclerotic disease process.7–10 Intravascular and epicardial ultrasound studies of atherosclerotic coronary3,5,11–13 and peripheral arteries14–17 have provided insights into the consequences of plaque accumulation and mechanisms of lesion formation. Earlier studies have confirmed the initial histopathologic findings of a biphasic course of lesion formation: (1) early preservation of lumen dimensions until a plaque burden of approximately 40% is reached, and (2) luminal narrowing as plaque burden further increases.3,4,6 More recent studies have shown evidence that inadequate or absent compensatory vascular enlargement may also be important in the development of arterial stenoses.12,13,15–18 The left main (LM) coronary artery is one of the most important targets of atherosclerotic plaque accumulation.5,6,10–25 In this study we used IVUS and quantitative coronary angiography to systematically assess the extent and characteristics of LM atherosclerosis in 107 patients undergoing catheter-based interventions of significant lesions of the left anterior descending or left circumflex coronary arteries.

METHODS

Patient population: Between August 1, 1995, and July 31, 1996, a prospective IVUS examination of the LM artery was performed in 107 patients. There were 84 men (78%) and 23 women (22%) who ranged in age from 32 to 80 years (mean 58 ± 11); all of them were symptomatic (87 had chronic stable angina and 20 unstable angina). This patient population represented a consecutive series of patients with IVUS-guided coronary interventions of significant narrowings in the left anterior descending or circumflex coronary arteries. None were considered to have significant disease of the LM artery. This study was approved by the local council on human research. All patients signed a written informed consent form approved by the medical ethical committee of the University Hospital Rotterdam-Dijkzigt.

Intervention procedures and coronary angiography: All patients received 250 mg of aspirin and 10,000 U of heparin intravenously. If the duration of the entire catheterization procedure exceeded 1 hour, the acti-
dural in-hospital complications in these patients.

Using IVUS. There were no procedural or postprocedural complications.

Orthogonal views without overlapping side branches or foreshortening was then performed off-line. End-diastolic frames acquired after intracoronary application of nitrates and having homogeneous opacification of the coronary lumen were selected for analysis. The computer-based Coronary Angiography Analysis System (CAAS II, Pie Medical, Maastricht, The Netherlands) was used for scanning of the entire LM artery. All studies were performed using a motorized transducer pullback at 0.5 mm/s. All IVUS studies were recorded on 0.5-inch high-resolution sVHS tape for off-line analysis.

Intravascular ultrasound imaging: After intracoronary injection of 2 mg of isosorbide dinitrate, the LM artery was examined using a mechanical IVUS system (CardioVascular Imaging Systems Inc, Sunnyvale, California) and a sheath-based IVUS catheter. The IVUS catheter incorporates a 30-MHz beveled, single-element transducer rotating at 1,800 rpm within a 2.9Fr long monorail imaging sheath. The distal 15-cm-long sonoluent segment of the imaging sheath has a common lumen that alternatively houses the guidewire (during catheter introduction) or the transducer (during imaging after the guidewire has been pulled back), but not both. This design avoids direct contact of the IVUS imaging core with the vessel wall. Before the IVUS imaging run, the guiding catheter was withdrawn from the ostium of the LM artery to assure complete clotting time was measured; intravenous heparin was administered in order to maintain an activated clotting time of $\geq 300$ seconds. In all patients, successful catheter-based interventions of lesions in the left anterior descending ($n = 91$) or left circumflex ($n = 16$) coronary arteries were performed (63 with stent placement, 25 with directional atherectomy, and 19 with balloon angioplasty). At the end of the interventional procedure, intracoronary injection of 2 mg of isosorbide dinitrate was administered, $\geq 2$ angiograms of the LM were recorded from opposite (ideally orthogonal) views without overlapping side branches or foreshortening, and the LM artery was interrogated using IVUS. There were no procedural or postprocedural complications.

Angiographic analysis: The angiograms were first interpreted by 3 experienced coronary angiographers who were blinded to IVUS data. All angiographic projections were reviewed for the presence of atherosclerosis of the LM artery. If during the visual assessment of the LM artery, an indentation of the opacified luminal silhouette or a lumen narrowing of any degree was noticed, that LM artery was classified as “angiographically abnormal.” Accordingly, only LM arteries without such findings were classified as “angiographically normal.” Decisions for classification required at least 2 votes.

Quantitative coronary angiography of 2 angiographic views (ideally orthogonal views without overlapping side branches or foreshortening) was then performed off-line. End-diastolic frames acquired after intracoronary application of nitrates and having homogeneous opacification of the coronary lumen were selected for analysis. The computer-based Coronary Angiography Analysis System (CAAS II, Pie Medical, Maastricht, The Netherlands) was used for measurement by IVUS has been reported previously. The site of the minimum lumen cross-sectional area was identified by carefully scrolling the tape back and forth; if there were multiple image slices with the same minimum lumen cross-sectional area, then the image slice with the largest plaque burden was selected. This was the image slice on which the IVUS measurements were performed (Figure 1). The cross-sectional area measurements included the lumen and total vessel cross-sectional area (interobserver differences: $0.1 \pm 0.8$ mm$^2$ [lumen], $0.0 \pm 1.1$ mm$^2$ [total vessel]). Plaque + media cross-sectional area was calculated as total vessel minus lumen cross-sectional area, and the plaque burden was calculated as plaque + media divided by total vessel cross-sectional area. The total vessel diameter was calculated as: square root of $[(4 \cdot \text{total vessel cross-sectional area})/\pi]$. The minimum and maximum lumen diameters (interobserver differences: $0.0 \pm 0.3$ mm [minimum lumen], $0.0 \pm 0.3$ mm [maximum lumen]), and the minimum and maximum plaque + media thickness (interobserver differences: $0.01 \pm 0.02$ mm [minimum plaque + media], 0.01 $\pm$ 0.09 mm [maximum plaque + media]) were measured. The total vessel cross-sectional area represents the area within the border between the hypoechoic media and the echoreflective adventitia. As in many previous studies using
IVUS, plaque + media cross-sectional area was used as a measure of atherosclerotic plaque, because ultrasound cannot measure media thickness accurately.\textsuperscript{27} The circumferential arc of the LM artery containing plaque was measured (in degrees) using a protractor centered on the lumen. The plaque burden has been termed cross-sectional area obstruction, cross-sectional narrowing, or percent plaque area by other investigators.

**QUALITATIVE ANALYSIS:** Plaque composition was assessed visually. The presence of significant amounts of calcium, dense fibrous tissue, or soft plaque was tabulated. Calcium produced bright echoes (brighter than the reference adventitia) with acoustic shadowing (attenuation) of deeper arterial structures;\textsuperscript{c} the largest arc of calcium within the LM artery was identified and measured (in degrees) using a protractor centered on the lumen. Although initially an arc of calcium >120° was predetermined to be an exclusion criterion (to assure reliable measurement of the total vessel and plaque + media cross-sectional area), no patient had this degree of LM calcification. Plaque tissue producing echoes that were as bright as or brighter than the reference adventitia, but without acoustic shadowing was classified as “fibrous.” Tissue being less dense than the reference adventitia was classified as “soft.” Plaques containing more than 1 type of tissue were classified as “mixed.”

**Statistical analysis:** Categorical variables were presented as frequencies. Continuous variables were presented as mean ± 1 SD. Categorical variables were compared using chi-square analysis. Continuous variables were compared using the 2-tailed Student’s t test and linear and nonlinear regression analyses. A p value <0.05 was considered statistically significant.

**RESULTS**

**Overall angiographic and intravascular ultrasound data:** With use of coronary angiography, only 26 LM arteries (24\%) were classified as abnormal; however, IVUS revealed atherosclerotic plaques in all 107 LM arteries (p <0.0001 vs angiography). Twenty-five plaques were classified as soft, 22 were fibrous, and 50 were mixed. Plaque calcification was found in 30 LM lesions with an arc of calcium ranging from 20° to 120° in circumference. Data provided by both quantitative coronary angiography and IVUS are listed in Table I.

**Data of angiographically normal versus abnormal main stems:** The IVUS and quantitative coronary angiographic findings in patients with angiographically abnormal (n = 26) and normal LM arteries (n = 81) were then compared. The 2 groups were similar with regard to age (59 ± 10 vs 57 ± 11 years) and gender (77% vs 79% men). Patients with an angiographically abnormal LM artery had (1) a smaller quantitative angiographic minimal lumen diameter (3.30 ± 0.68 vs 4.02 ± 0.73 mm, p <0.0001), (2) a smaller quantitative angiographic reference diameter (4.01 ± 0.74 vs 4.23 ± 0.75 mm, p <0.0001), (3) a higher quantitative angiographic diameter stenosis (17.60 ± 6.85% vs 8.48 ± 3.15%, p <0.0001), and (4) a smaller IVUS minimal lumen cross-sectional area (13.0 ± 4.9 vs 15.7 ± 4.7 mm², p <0.05). In patients with an angiographically abnormal LM artery, the IVUS plaque burden tended to be higher (37.2 ± 14.2% vs 32.1 ± 14.2%, p = 0.10); however, the total vessel cross-sectional area tended to be even smaller (20.9 ± 6.4 vs 23.1 ± 5.4 mm², p = 0.10). There was no significant difference in plaque composition.

**Assessment of vascular remodeling:** The total vessel cross-sectional area by IVUS correlated directly with the plaque + media cross-sectional area (r = 0.54, p <0.0001, Figure 2).

In addition, the IVUS minimal lumen cross-sectional area correlated inversely with the plaque burden (r = −0.62, p <0.0001, Figure 3); this relation was very similar for the LM artery with a plaque burden ≥40% (r = −0.42, y = −0.017x + 19.0, p <0.01) and for the LM artery with a plaque burden <40% (r = −0.41, y = −0.21x + 22.3, p <0.001). Similar correlations were found for the LM artery with a plaque burden ≥30% and <30% (r = −0.48 and −0.42, respectively, both p <0.01). Various nonlinear regression approaches were tested and did not reveal relations with higher or equal significance.

**TABLE I Quantitative Coronary Angiography and Intravascular Ultrasound Data**

<table>
<thead>
<tr>
<th></th>
<th>All Patients [n = 107]</th>
<th>Plaque Burden &lt;40% [n = 71]</th>
<th>Plaque Burden ≥40% [n = 36]</th>
<th>p Value*</th>
</tr>
</thead>
<tbody>
<tr>
<td>IVUS</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lumen CSA (mm²)</td>
<td>15.1 ± 4.8</td>
<td>16.8 ± 4.7</td>
<td>11.6 ± 2.8</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>P + M CSA (mm²)</td>
<td>7.5 ± 3.6</td>
<td>6.1 ± 2.9</td>
<td>10.3 ± 3.2</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Total vessel CSA (mm²)</td>
<td>22.6 ± 5.7</td>
<td>22.9 ± 6.1</td>
<td>21.8 ± 4.8</td>
<td>0.30</td>
</tr>
<tr>
<td>Plaque burden (%)</td>
<td>33.3 ± 12.9</td>
<td>26.6 ± 9.4</td>
<td>46.6 ± 7.3</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Minimum lumen diameter (mm)</td>
<td>3.91 ± 0.75</td>
<td>4.18 ± 0.71</td>
<td>3.38 ± 0.54</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Maximum lumen diameter (mm)</td>
<td>4.65 ± 0.75</td>
<td>4.92 ± 0.70</td>
<td>4.12 ± 0.54</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Minimum P + M diameter (mm)</td>
<td>0.16 ± 0.22</td>
<td>0.08 ± 0.15</td>
<td>0.31 ± 0.25</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Maximum P + M diameter (mm)</td>
<td>0.99 ± 0.42</td>
<td>0.85 ± 0.37</td>
<td>1.26 ± 0.40</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Total vessel diameter (mm)</td>
<td>5.32 ± 0.65</td>
<td>5.36 ± 0.68</td>
<td>5.24 ± 0.57</td>
<td>0.34</td>
</tr>
<tr>
<td>Arc of plaque (%)</td>
<td>296 ± 83</td>
<td>270 ± 90</td>
<td>348 ± 22</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Quantitative coronary angiography</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Minimum lumen diameter (mm)</td>
<td>3.84 ± 0.78</td>
<td>3.98 ± 0.70</td>
<td>3.59 ± 0.86</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>Reference diameter (mm)</td>
<td>4.17 ± 0.75</td>
<td>4.30 ± 0.68</td>
<td>3.93 ± 0.84</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>Diameter stenosis (%)</td>
<td>8.0 ± 7.0</td>
<td>7.6 ± 6.2</td>
<td>8.8 ± 8.3</td>
<td>0.43</td>
</tr>
</tbody>
</table>

*Plaque burden <40% versus plaque burden ≥40%. Data are expressed as mean ± 1 SD. CSA = cross-sectional area; IVUS = intravascular ultrasound; P + M = plaque + media. 

\textsuperscript{c} Calcium produced bright echoes (brighter than the lumen). Although initially an arc of calcium measured (in degrees) using a protractor centered on the lumen. 

The circumferential arc of the LM artery containing plaque was measured (in degrees) using a protractor centered on the lumen. The plaque burden has been termed cross-sectional area obstruction, cross-sectional narrowing, or percent plaque area by other investigators.
Thirty-six of the 107 LM arteries had an IVUS plaque burden $\geq 40\%$; 71 had a plaque burden $<40\%$. Both groups were similar with respect to age (58 ± 11 vs 57 ± 11 years) and gender (79% vs 78% men). Results of quantitative coronary angiography and IVUS analysis of these 2 subsets are listed in Table I. In the group with plaque burden $\geq 40\%$, IVUS plaque composition was less frequently soft (8% vs 31%, $p < 0.01$); the frequency of plaque calcification was higher (47% vs 18%, $p < 0.005$), and the arc of diseased vessel wall was significantly larger (348 ± 22º vs 270 ± 90º, $p < 0.0001$). In addition, in vessels with plaque burden $\geq 40\%$, the lumen cross-sectional area and diameters were significantly smaller; the plaque + media cross-sectional area and diameters were larger, but the total vessel cross-sectional area was similar (22.9 ± 6.1 vs 21.8 ± 4.8 mm², $p = 0.30$).

**DISCUSSION**

In the present study, careful IVUS examination during catheter-based interventions of major epicardial left coronary arteries in a consecutive series of 107 patients revealed the presence of LM plaques in all and demonstrated the superiority of IVUS in detecting early atherosclerotic changes in vivo.1,2,8,19,20 These data are in agreement with previous IVUS findings in a smaller study group.19

**Adaptive vascular remodeling:** The absence of angiographic lumen narrowing despite the presence of ultrasound-confirmed plaque formation is generally thought to result from compensatory vascular enlargement. In addition, there were other signs of adaptive remodeling. First, the total vessel cross-sectional area measured 22.6 mm²; this is comparable to previous measurements in diseased LM arteries (22.0 to 23.3 mm²),3,20 but larger than measurements in nondiseased LM arteries (19.0 mm²).20 Second, despite an average plaque burden of 33%, the average lumen cross-sectional area was reduced by 20% compared with that in nondiseased LM arteries.20

As in previous histopathologic and IVUS studies,2–6 we found a significant relation between plaque + media and total vessel cross-sectional area. This relation, which has previously been understood as an arithmetic expression of the adaptive remodeling process, appears to be less strong in the LM artery ($r = 0.46$ to $0.56$)3,5 than has been reported in other major epicardial coronary branches or in peripheral vessels ($r = 0.63$ to 0.85).2,3,14–16 In addition, the value of the relation between plaque + media and total vessel area appears to be limited because (1) correlating plaque + media with total vessel cross-sectional area should have a significant positive relation because this is a correlation between $a$ and $a + \beta$, and (2) the regression equation will always show steeper slopes for mildly diseased vascular segments, falsely suggesting overcompensation of total vessel area in relation to plaque accumulation.28

**Variations of remodeling response:** However, the results of the current study also suggest that even during earlier stages of atherosclerotic plaque accu-
mulation (i.e., plaque burden of 20% to 40%), compensatory vascular enlargement may be inadequate to preserve lumen dimensions in a considerable number of cases. For instance, there was a negative correlation between lumen cross-sectional area and plaque burden ($r = 0.62$), a relation that was not restricted to patients with a plaque burden $> 0.30$ or 40%, as previously described. In the present study, there was also no significant difference in total vessel cross-sectional area between (1) LM artery with a plaque burden $< 40\%$ versus LM artery with a plaque burden $\geq 40\%$, or (2) angiographically normal versus angiographically abnormal LM arteries. The total vessel cross-sectional area was even slightly higher in the group with angiographically normal LM arteries. We used IVUS normal values from published data, obtained from 61 LM arteries with IVUS-documented absence of plaque formation ($19.0 \pm 6.5 \text{ mm}^2$), to evaluate the preservation of the lumen cross-sectional area. Figure 4 shows data of the current study compared with the mean $\pm 1$ SD of the normal size. In patients with a LM plaque burden $< 20\%$, the lumen size was evenly distributed on both sides of the mean normal value; in all patients the lumen was larger than the $-1$ SD margin. This may, to some extent, result from adaptive remodeling. Conversely, in several main stems with a plaque burden between 20% and 40%, the lumen was smaller than the $-1$ SD margin, which may reflect inadequacy or early cessation of the adaptive remodeling process.

Variability of the adaptive remodeling process has previously been suggested by others. Nevertheless, because these studies in humans were observations at one point in time, the time course and magnitude of vascular response to plaque growth remains unknown.

**Limitations and potential sources of error:**

(1) All patients underwent catheter-based coronary interventions. Therefore, we were limited to assessment of LM arteries in patients without angiographically significant LM lesions, but with significant lesions of the left anterior descending or circumflex coronary arteries.

(2) As in previous histopathologic or IVUS studies of vascular remodeling, this study was observational and provided only a “snapshot-like view” on coronary artery disease.

Clinical implications: Although IVUS inspection reveals the presence of plaque formation within the LM artery in patients undergoing procedures of major left coronary branches, the extent of plaque accumulation is variable and should not be equated with hemodynamically significant disease. Nevertheless, our findings underline the standard recommendation to perform any cannulation of the LM artery with care. The current study demonstrates that compensatory vascular enlargement is a variable phenomenon, explaining a smaller lumen cross-sectional area in some cases even before the plaque occupies 40% of the total vessel cross-sectional area. Serial IVUS studies will be necessary to gain insight into (1) the time course of vascular remodeling, and (2) whether plaques with different patterns of remodeling require specific or different therapy with respect to the acute procedural success and long-term outcome.


2. Stiel GM, Stiel LSG, Schofer J, Donath K, Mathey DG. Impact of compen-