Atherosclerotic coronary lesions with inadequate compensatory enlargement have smaller plaque and vessel volumes: observations with three dimensional intravascular ultrasound in vivo


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Atherosclerotic coronary lesions with inadequate compensatory enlargement have smaller plaque and vessel volumes: observations with three dimensional intravascular ultrasound in vivo


Abstract

Objective—To compare vessel, lumen, and plaque volumes in atherosclerotic coronary lesions with inadequate compensatory enlargement versus lesions with adequate compensatory enlargement.

Design—35 angiographically significant coronary lesions were examined by intravascular ultrasound (IVUS) during motorised transducer pullback. Segments 20 mm in length were analysed using a validated automated three dimensional analysis system. IVUS was used to classify lesions as having inadequate (group I) or adequate (group II) compensatory enlargement.

Results—There was no significant difference in quantitative angiographic measurements and the IVUS minimum lumen cross sectional area between groups I (n = 15) and II (n = 20). In group I, the vessel cross sectional area was 13.3 (3.0) mm² at the lesion site and 14.4 (3.6) mm² at the distal reference (p < 0.01), whereas in group II it was 17.5 (5.6) mm² at the lesion site and 14.0 (6.0) mm² at the distal reference (p < 0.001). Vessel and plaque cross sectional areas were significantly smaller in group I than in group II (13.3 (3.0) v 17.5 (5.6) mm², p < 0.01; and 10.9 (2.8) v 15.2 (4.9) mm², p < 0.005). Similarly, vessel and plaque volume were smaller in group I (291.0 (61.0) v 353.7 (110.0) mm³, and 177.5 (48.4) v 228.0 (92.8) mm³, p < 0.05 for both). Lumen areas and volumes were similar.

Conclusions—In lesions with inadequate compensatory enlargement, both vessel and plaque volume appear to be smaller than in lesions with adequate compensatory enlargement.

Intravascular ultrasound provides transmural images of coronary arteries in vivo. The coronary vascular wall, the cross sectional area of the atherosclerotic plaque, the consequences of plaque accumulation, and the mechanisms of lesion formation can be studied in humans in a manner previously not possible. 1–4

Methods

PATIENT POPULATION

The study population consisted of 35 patients with primary (not restenotic) atherosclerotic lesions examined using preintervention intravascular ultrasound. Inclusion criteria were: angiography documented non-curved lesion segments; limited plaque calcification throughout a lesion length of 20 mm; absence of a complete occlusion of the stenotic lumen during the ultrasound imaging run; and absence of major side branches. Thirty two men and three women (mean (SD) age 61 (9) years) were examined. Lesions were located in the left anterior descending coronary artery (n = 20), right coronary artery (n = 10), and left circumflex coronary artery (n = 5); 32 were proximal and three were in the mid-portion. The study was approved by the Local Council on Human Research. All patients signed a written informed consent form, approved by the local medical ethics committees.
The patients received 250 mg aspirin and 10 000 U heparin intravenously. If the duration of the entire catheterisation procedure exceeded one hour, the activated clotting time was measured, and intravenous heparin was given in order to maintain a clotting time of more than 300 s. Intravascular ultrasound imaging was performed after intracoronary injection of 0.2 mg glyceryl trinitrate, starting at least 10 mm distal to the lesion segment. A mechanical intravascular ultrasound system (ClearView, CardioVascular Imaging Systems, Sunnyvale, California, USA) and a sheath based imaging catheter were used. The catheter incorporated a 30 MHz bevelled, single element transducer rotating at 1800 rpm (MicroView, CardioVascular Imaging Systems). This catheter is equipped with a 2.9 F 15 cm long sonolucent distal sheath that has a common lumen that either houses the guide wire (during catheter introduction) or the transducer (during imaging after the guide wire has been pulled back), but not both. This design avoids direct contact of the imaging core with the vessel wall. The ultrasonic transducer was withdrawn through the stationary imaging sheath using a motorised pullback device at a constant speed of 0.5 mm/s. All intravascular ultrasound examinations were recorded on high resolution s-VHS videotape for later offline quantitative analysis. After the intravascular ultrasound examination, all patients were successfully treated by balloon angioplasty, coronary stent implantation, or directional coronary atherectomy; there were no procedural or postprocedural in hospital complications.

**INTRAVASCULAR ULTRASOUND IMAGE ANALYSIS**

Twenty millimeter long lesion segments (10 images/mm axial arterial length), centered on the target lesion site, were analysed off-line using a computerised intravascular ultrasound analysis system (fig 1). Reference images with the smallest plaque burden were acquired no more than 3 mm distal to the lesion segment. Cross sectional area measurements at the reference site were obtained with the computerised analysis system (single frame mode); care was taken to avoid any major side branch between the lesion segment and the reference site.

Cross sectional area measurements (mm$^2$) included the lumen and vessel cross sectional area. The vessel cross sectional area was measured by tracing the border between the hypoechoic media and the echoreflective adventitia. As in many previous studies using intravascular ultrasound, the cross sectional area (and thickness) of plaque plus media was used as a measure of atherosclerotic plaque area (and thickness) because ultrasound cannot measure media thickness accurately. Plaque cross sectional area was calculated as vessel cross sectional area minus lumen cross sectional area. The cross sectional area plaque burden was calculated as plaque cross sectional area divided by vessel cross sectional area.

Compensatory enlargement was considered inadequate (group I) if the vessel cross sectional area at the site of the minimum lumen cross sectional area was smaller than that at the distal reference site (fig 2). If the vessel cross sectional area at the site of the minimum lumen cross sectional area was larger than or equal to the distal reference site (fig 3), compensatory enlargement was considered adequate (group II).

Volume measurements (mm$^3$) of the lumen, vessel, and plaque (based on 10 intravascular ultrasound images/mm axial arterial length) were calculated according to Simpson’s rule as

$$\text{Volume} = \sum_{i=1}^{n} \text{cross sectional area}_i \times H$$

where $H =$ thickness of a coronary artery slice represented by a single tomographic intravascular ultrasound image, and $n =$ number of images in the three dimensional image set. The volume plaque burden (%) was calculated as plaque volume divided by vessel volume.

The overall plaque eccentricity index was calculated as mean value of the eccentricity indices of all individual image slices; these were derived as previously described (minimum plaque thickness divided by maximum plaque thickness). A higher value of that index indicated a more concentric plaque distribution (the maximum value 1.0 would indicate perfectly concentric plaque distribution along the entire lesion segment), whereas a lower
The overall lumen symmetry index was calculated as mean value of the lumen symmetry indices (minimum lumen diameter divided by maximum lumen diameter) of all individual image slices. A higher value of that index indicated a more symmetrical lumen shape (the maximum value 1.0 would indicate a perfectly circular lumen along the entire lesion segment).

**COMPUTERISED INTRAVASCULAR ULTRASOUND ANALYSIS METHOD**

The analysis was performed offline using a computerised intravascular ultrasound analysis system. The analysis system used the Windows™ (Microsoft, Redmond, Washington, USA) operating system on a personal computer. The computerised analysis required the digitisation of a stack of intravascular ultrasound images from videotape. Two longitudinal sections were automatically reconstructed (fig 1), and the contours corresponding to the lumen–tissue and media–adventitia interfaces were automatically identified. The longitudinal contours were visually checked and, if necessary, edited with computer assistance (see below). The longitudinal contours generated individual edge points on the planar images defining the centre and range of the automated boundary search on the planar images. Subsequently, contour detection of the planar images was performed. The axial location of an individual planar image (on the longitudinal contours) was indicated by a cursor; the cursor was used to scroll through the entire set of planar images while the planar contours were visually checked. Finally, the contour data of the planar images were used for the computation of the results.

Automated contour detection and computer assisted contour editing were based on the application of a minimum cost algorithm to detect the luminal and external vessel boundaries. Each digitised intravascular ultrasound image was resampled in a radial format (64 radii per image), and a cost matrix representing the edge strength was calculated from the image data. For the boundary between lumen and plaque, the cost value was defined by the spatial first derivative. For the external vessel boundary a cross correlation pattern matching process was used for the cost calculations. The path with the smallest accumulated value was determined by dynamic programming techniques.

The computer assisted editing differed considerably from conventional manual contour tracing. The computer mouse was used to indicate the correct boundary. This forced the contour through the manually entered point by assigning this point a very low value in the cost matrix. Editing the contour of a single slice caused the entire dataset to be updated (dynamic programming). Side branches with relatively small ostium and small calcified portions of the plaque were generally ignored by the algorithm as a result of its robustness, which means that the automated contour detection did not follow every abrupt change in the cost path.

This algorithm has been validated in tubular phantoms. A comparison between automated three dimensional intravascular ultrasound

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**Figure 2** Analysis of a 20 mm long mid-right coronary segment with inadequate compensatory vascular enlargement (group I). The vessel cross sectional area is smallest at the target lesion site. Markers indicate that site on the longitudinal sections (right upper panels) and the display of the cross sectional area measurements (right lower panel). Linear functions of the vessel and lumen cross sectional area form the upper and lower boundaries of the greyish area, which represents the plaque cross sectional area. Alternatively, the values of plaque cross sectional area can be derived directly from a linear function (single black line), which here partly overlaps the greyish area.

**Figure 3** Adequate compensatory vascular enlargement, as observed in one of the lesion of group II. The vessel cross sectional area is larger at the target lesion site (upper panel) than at the reference site (left mid panel). The lower panel illustrates and underlines the principle of compensatory vascular enlargement.
Table 1 Quantitative coronary angiographic measurements

<table>
<thead>
<tr>
<th>Variables</th>
<th>Group I (n = 15)</th>
<th>Group II (n = 20)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minimum lumen diameter (mm)</td>
<td>1.03 (0.33)</td>
<td>1.16 (0.33)</td>
<td></td>
</tr>
<tr>
<td>Interpolated reference diameter (mm)</td>
<td>3.04 (0.39)</td>
<td>3.31 (0.65)</td>
<td></td>
</tr>
<tr>
<td>Diameter stenosis (%)</td>
<td>66 (10)</td>
<td>63 (15)</td>
<td></td>
</tr>
</tbody>
</table>

Values are mean (SD); all non-significant.

Groups I and II, lesions with and without inadequate compensatory vascular enlargement, respectively.

Table 2 Cross sectional area (CSA) measurements with intravascular ultrasound

<table>
<thead>
<tr>
<th>Variables</th>
<th>Group I (n = 15)</th>
<th>Group II (n = 20)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minimum lumen site</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lumen CSA (mm²)</td>
<td>2.4 (1.0)</td>
<td>2.3 (1.5)</td>
<td>NS</td>
</tr>
<tr>
<td>Vessel CSA (mm²)</td>
<td>13.3 (5.0)</td>
<td>17.5 (5.6)</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>Plaque CSA (mm²)</td>
<td>10.9 (2.8)</td>
<td>15.2 (4.9)</td>
<td>&lt; 0.005</td>
</tr>
<tr>
<td>CSA plaque burden (%)</td>
<td>82 (7)</td>
<td>87 (7)</td>
<td>&lt; 0.05</td>
</tr>
<tr>
<td>Reference site</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lumen CSA (mm²)</td>
<td>7.2 (3.6)</td>
<td>7.4 (4.1)</td>
<td>NS</td>
</tr>
<tr>
<td>Vessel CSA (mm²)</td>
<td>14.4 (5.6)</td>
<td>14.0 (6.0)</td>
<td>NS</td>
</tr>
<tr>
<td>Plaque CSA (mm²)</td>
<td>7.3 (2.5)</td>
<td>6.5 (3.0)</td>
<td>NS</td>
</tr>
<tr>
<td>CSA plaque burden (%)</td>
<td>52 (15)</td>
<td>49 (16)</td>
<td>NS</td>
</tr>
</tbody>
</table>

Values are mean (SD).

Groups I and II, lesions with and without inadequate compensatory vascular enlargement, respectively.

Table 3 Volumetric measurements by intravascular ultrasound

<table>
<thead>
<tr>
<th>Variables</th>
<th>Group I (n = 15)</th>
<th>Group II (n = 20)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minimum lumen volume (mm³)</td>
<td>113.6 (44.0)</td>
<td>125.7 (49.9)</td>
<td>NS</td>
</tr>
<tr>
<td>Vessel volume (mm³)</td>
<td>291.0 (61.0)</td>
<td>353.7 (110.0)</td>
<td>&lt; 0.05</td>
</tr>
<tr>
<td>Plaque volume (mm³)</td>
<td>177.5 (48.4)</td>
<td>228.0 (92.8)</td>
<td>&lt; 0.05</td>
</tr>
<tr>
<td>Volumetric plaque burden (%)</td>
<td>61 (11)</td>
<td>64 (10)</td>
<td>NS</td>
</tr>
</tbody>
</table>

Values are mean (SD).

Groups I and II, lesions with and without inadequate compensatory vascular enlargement, respectively; segment length was 20 mm.
in group II (p < 0.05 for both) (table 3); there was no significant difference in lumen volume or in volumetric plaque burden.

The eccentricity index (minimum plaque thickness divided by maximum plaque thickness) was significantly higher in group I than in group II (0.28 (0.07) v 0.21 (0.11), p < 0.05), indicating a more concentric plaque distribution in group I. The lumen shape was significantly more symmetrical in group I; this was shown by the higher lumen symmetry index (minimum lumen diameter divided by maximum lumen diameter) in group I (0.86 (0.02) v 0.84 (0.03), p < 0.05).

Discussion
Three dimensional intravascular ultrasound was initially used for visual assessment of the spatial configuration of plaques and dissection membranes, whereas contemporary three dimensional intravascular ultrasound systems are equipped with algorithms for computer assisted analysis of the plaque or lumen volumes and dimensions. The three dimensional intravascular ultrasound analysis system used in the current study has been extensively validated; it permits the rapid automated analysis of lumen and plaque volumes and dimensions on a large number of planar image slices.

We used this three dimensional intravascular ultrasound analysis system to compare 15 lesions with inadequate compensatory enlargement to 20 lesions with adequate compensatory enlargement. Compared to lesions with adequate compensatory enlargement, lesions with inadequate compensatory enlargement had smaller plaque and vessel volumes, more concentric plaque distribution, more symmetrical lumen shapes, and similar lumen volumes and dimensions. Importantly, lumen volumes and dimensions were similar in both groups, whether assessed by intravascular ultrasound or by quantitative coronary angiography. This emphasises the significance of intravascular ultrasound in the assessment of human atherosclerosis in vivo.

Our observations corroborate recent studies, which showed that inadequate compensatory enlargement may contribute to the development of significant luminal narrowing. In these previous studies, despite intracoronary injection of nitrates before the ultrasound examination, local vasospastic activity could not be excluded; and a collapse of the coronary artery could have resulted from a decrease in coronary arterial pressure, attributable to subtotal occlusion of the residual lumen during the ultrasound imaging run. However, neither local vasospasm nor collapse of the coronary artery can explain the significantly smaller vessel and plaque volumes of lesions with inadequate compensatory enlargement, as observed in the current study.

LIMITATIONS AND POTENTIAL SOURCES OF ERROR
As all previous studies of the natural history of coronary atherosclerosis in human were performed at a single point in time, the time course and magnitude of vascular response to plaque growth remains unknown.

Imaging with intravascular ultrasound can be hampered by eccentric catheter position, non-uniform transducer rotation, and non-coaxial catheter position.

As the external vascular boundary cannot be seen in the acoustic shadow behind calcium, we did not include lesions with severe plaque calcification. As in all studies with intravascular ultrasound, intracoronary injections of nitrates were performed before the ultrasound examination to prevent vasospasm; no angiographic changes before and after the intravascular ultrasound imaging procedure were observed, but this does not exclude local vasospastic activity.

Linear three dimensional systems, as used in the current study, provide approximate volumetric indices because they do not account for the presence of vascular curvatures. In the current study, only relatively straight coronary segments on the angiogram were included to minimise the curve induced error in the volume calculation. Approaches that combine data obtained from angiography and intravascular ultrasound can provide information on the real spatial geometry of the vessel, but these sophisticated techniques are still subject to refinement and ongoing research.

CONCLUSIONS
Planar intravascular ultrasound analysis identified a population of coronary artery lesions with inadequate compensatory vascular enlargement. Volumetric intravascular ultrasound analysis showed that these lesions have less atherosclerotic plaque. Serial intravascular ultrasound studies will be required to determine whether the adaptive remodelling state of atherosclerotic lesions has any implication for the success of catheter based or pharmacological treatment strategies.

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References
von Birgelen, Mintz, de Vrey, et al


