Complex stenosis morphology and vasomotor responses to inhibition of nitric oxide synthesis

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Abstract

Objective—To assess the relation between coronary vasomotor effects of N⁶-monomethyl-L-arginine (LNMMA) administration and coronary stenosis morphology, length, and severity in patients with stable angina.

Design—In 28 patients (24 male, four female) with coronary artery disease and chronic stable angina, intracoronary normal saline and 4 μmol/min LNMMA were infused for four minutes each, followed by an intracoronary bolus of 250 μg glyceryl trinitrate. Coronary stenoses were classified as concentric (smooth), eccentric (smooth), or complicated (irregular). The diameters of these stenoses and their adjacent reference proximal segments were measured by quantitative angiography.

Results—During LNMMA infusion a significantly larger proportion of complicated stenoses than concentric and eccentric stenoses constricted by ≥5% (p < 0.01) and the magnitude of vasoconstriction was greater in complicated than in concentric and eccentric stenoses (p < 0.05). For complicated stenoses the magnitude of constriction (in mm) with reference to normal saline was greater than that of the concentric and eccentric stenoses (p < 0.05), whereas concentric and eccentric stenoses constricted similarly. Irrespective of the type of morphology, there was a correlation (p < 0.05) between both the severity and the length of stenoses and the magnitude of vasoconstriction to LNMMA. A similar proportion of concentric, eccentric, and complicated stenoses showed ≥5% increase in diameter with glyceryl trinitrate, and the magnitude of the response was similar in the three groups.

Conclusions—In patients with coronary artery disease, the response to LNMMA is greater when stenosis morphology is complex, indicating greater nitric oxide activity. This provides further evidence that plaques with complex morphology are in an active state.

Keywords: endothelium; nitric oxide; coronary artery disease; stenosis morphology

Nitric oxide is an endogenous vasodilator produced by vascular endothelial cells. Its synthesis can be specifically and competitively antagonised by arginine analogues, such as N⁶-monomethyl-L-arginine (LNMMA), which inhibit the enzymatic pathway in a process that can be reversed by increased availability of L-arginine. Inhibition of nitric oxide synthesis has been shown to decrease the basal diameter of angiographically normal and diseased epicardial coronary arteries and stenoses.

Evidence suggests that the obstruction associated with a coronary artery stenosis is often dynamic rather than fixed. A 10% circumferential shortening of the normal wall may lead to a greater change in luminal calibre at the site of an eccentric plaque than at the site of a concentric plaque. Furthermore, coronary stenosis morphology may be an important factor determining the degree of vasomotor response to stimulation. Previous studies in animals have shown that the intracoronary activation of platelets and their interaction with the vascular endothelium may contribute to or precipitate myocardial ischaemia, both by causing mechanical obstruction of the arterial lumen and by releasing potent vasoconstrictor substances such as endothelin, thromboxane, or serotonin. Further studies provide evidence of a role for these mechanisms in man. Experimental studies show that the effects of serotonin or acetylcholine differ depending on whether the stenosis is compliant or fixed. A clinical study has shown that the response of stenoses to serotonin is related to the morphology of the lesion. The relation between nitric oxide activity and coronary stenosis morphology has not been studied.

To study the vasomotor effects of LNMMA administration and their relations to coronary stenosis morphology, we examined the effect of intracoronary LNMMA infusion on coronary luminal diameter, using quantitative angiography, at the site of concentric, eccentric, and complicated lesions in patients with chronic stable angina.

Methods

PATIENTS

We studied 28 patients (24 male, four female, mean (SD) age 58 (8) years) with chronic stable angina, coronary artery disease, and a positive treadmill exercise test result (≥0.1 mV ST segment depression) at between 5 and 7 METS using the modified Bruce protocol. All patients were subjected to coronary angiography to assess their suitability for percutaneous transluminal coronary angioplasty (PTCA) or coronary surgery, because their angina limited their physical activity despite drug treatment. Patients were excluded from the study if they had diabetes mellitus, recent myocardial infarction (within six months), left ventricular hypertrophy (on
echocardiography), left ventricular dysfunction (left ventricular ejection fraction < 50%) or valvar heart disease, were current smokers, or had uncontrolled hypertension. Antihypertensive drug treatment was stopped 24 hours before the study. The patients were allowed to use sublingual glyceryl trinitrate as necessary, but no study was performed within three hours of its administration. The protocol was approved by the research ethics committee and each patient gave written informed consent.

PROTOCOL

Following the diagnostic coronary angiogram an optimal radiographic projection was selected and kept constant for subsequent angiograms (study operators CT, GJD). Thirty eight stenoses and their reference proximal segments were examined. Two ECG leads were monitored continuously throughout the study. Patients received a single four minute infusion of 0.9% saline (2 ml/min) followed by a four minute intracoronary infusion of LNMM (4 µmol/min) in saline, using a syringe pump, followed after five minutes by an intracoronary bolus dose of glyceryl trinitrate (250 µg in 2 ml of saline). Femoral arterial pressure and heart rate were recorded during the last 30 seconds of each infusion period. Angiography was performed with a hand injection of 6–8 ml non-ionic contrast medium at baseline, immediately after each infusion, and two to three minutes after glyceryl trinitrate. Before each angiogram, the catheter was emptied to avoid bolus administration of the infusate.

QUANTITATIVE CORONARY ANGIOGRAPHY

End diastolic frames from each arteriogram were selected for analysis. Each frame was analysed, in random order, using quantitative computerised analysis with an automated edge contour detection analysis system (Computerised Angiographic Analysis System (CAAS), version 2V2; Pie Data Medical, Maastricht, the Netherlands). Thirty five per cent of frames were analysed and kept constant for subsequent analysis. This classification of stenosis morphology was compared with that obtained by computed symmetry analysis (CAAS symmetry index) of the same coronary lesions. Concentric stenoses were defined as those producing symmetrical narrowing, with smooth borders or only very mild irregularity (symmetry index > 0.5–1), that looked similar in orthogonal projections. Eccentric stenoses were defined as asymmetrical narrowing with irregular borders and/or overhanging edges (type II of Ambrose and colleagues) or with an “abrupt proximal face” or a “rough” or “saw-tooth” component. Illustrations of these three morphological types of stenosis have been published previously. Symmetry index was calculated from two parameters, dl and dr, the smaller divided by the larger, where: dl = the distance between the detected and estimated left hand contour of the vessel; and dr = the distance between the detected and estimated right hand contours. The symmetry measure is computed as the minimum of dl and dr divided by the maximum of dl and dr. Accordingly a symmetry index of 1 denotes a concentric stenosis and the value decreases with increasing eccentricity.

Quantitative analysis of coronary arteriograms was carried out by two independent observers (DT, GG), who blindly reanalysed the films at a remote time for reproducibility of the method. No significant intraobserver variability (analysis of variance F = 0.4, p = 0.7) or interobserver variability (analysis of variance F = 0.35, p = 0.82) variability was found. The findings of computed and qualitative assessments of symmetry were concordant in 35 stenoses (92%) and discordant in three, which were classified by consensus. In a previous study, we analysed the intraobserver and interobserver variability of the measurement of coronary lumen diameter using the CAAS system in our institution. Twenty four arterial segments were analysed by two independent observers and reanalysed at a remote time. The mean intraobserver variation, expressed as the standard error of the estimate (SEE), was 0.12 mm; the interobserver variation (SEE) was 0.10 mm.

STATISTICAL ANALYSIS

Data are expressed as mean (SEM). Analysis of variance (ANOVA) and the Scheffe F test for repeated measures were used to compare serial changes in heart rate and blood pressure and in the diameter of coronary stenoses. To test for differences in percentage response to LNMM and nitrates of eccentric, concentric, and complicated stenoses, a two way ANOVA for repeated measures was applied. Associations between responses to LNMM and stenosis length, severity, and eccentricity ratio were assessed by performing linear regression analysis and calculating a correlation coefficient. Multivariate analysis was performed between LNMM response and stenosis length, severity, and eccentricity ratio. Student’s t test was used to compare paired and unpaired data.
between groups, and the responses to glyceryl trinitrate and to LNMMA. Differences between proportions were analysed by a Yates corrected $\chi^2$ test. A probability value of $p < 0.05$ (two tailed) was significant.

**Results**

Mean (SEM) systolic aortic pressure increased from 134.0 (3.8) to 143.6 (4.4) mm Hg ($p < 0.01$) during intracoronary administration of LNMMA, but heart rate remained unchanged, at 70.0 (1.3) to 70.8 (1.4) beats/min at baseline. ST segment depression occurred in two patients and was promptly relieved by intracoronary administration of isosorbide dinitrate. Intracoronary glyceryl trinitrate had no significant effect on systolic aortic pressure (134.9 (4.0) to 132.0 (3.6) mm Hg at baseline) or heart rate (71.0 (1.2) to 72.1 (1.1) beats/min at baseline).

**STENOSIS MORPHOLOGY**

Thirty eight of the 41 coronary stenoses (14 concentric, 12 eccentric, and 12 complicated) observed in these 28 patients were suitable for quantitative analysis, and the results below refer to these stenoses. Two patients had two stenoses within the same vessel (left anterior descending coronary artery). The severity of coronary stenoses for the whole group ranged from 20.2% to 91.4% (mean 46 (2.5)%) lumen diameter reduction. There were 13 stenoses of $> 50%$: four concentric, three eccentric, and six complicated. There was no significant difference in mean per cent minimum luminal diameter at baseline (44 (3.7)%), 43 (5.2)%, and 50.3 (4.3)% respectively) or in stenosis length (3.50 (0.30), 3.04 (0.31), and 3.72 (0.38 mm, respectively) between concentric, eccentric, and complicated stenoses.

### Table 1 Proportion of stenoses and their reference segments with reactivity (≥ 5% change from baseline) to intracoronary N$^\text{\textregistered}$-monomethyl-L-arginine (LNMMA) and nitrates

<table>
<thead>
<tr>
<th>Stenoses</th>
<th>LNMMA</th>
<th>Nitrates</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline</td>
<td>Ref segment</td>
<td>Stenoses</td>
</tr>
<tr>
<td>Concentric (n=14)</td>
<td>3/14 (21%)</td>
<td>2/14 (14%)</td>
</tr>
<tr>
<td>Eccentric (n=12)</td>
<td>3/12 (25%)</td>
<td>1/12 (8%)</td>
</tr>
<tr>
<td>Complicated (n=12)</td>
<td>8/12 (67%)*</td>
<td>1/12 (8%)</td>
</tr>
</tbody>
</table>

Figures given as $x/y (z)$, where $x = \text{number of reactive stenoses or reference segments}$, $y = \text{total number of stenoses or reference segments}$, and $z = \text{percentage of stenoses or reference segments which are reactive}$.

Values are mean (SD). Values in italics are percentage change from baseline.

### Table 2 Reactivity of coronary stenoses and their reference segments to intracoronary administration of LNMMA and nitrates

<table>
<thead>
<tr>
<th>Morphology</th>
<th>Minimum lumen diameter (mm)</th>
<th>Stenoses</th>
<th>Reference segments</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Baseline</td>
<td>LNMMMA</td>
<td>Nitrates</td>
</tr>
<tr>
<td>Concentric (n=14)</td>
<td>1.38 (0.11)</td>
<td>1.36 (0.11)</td>
<td>1.57 (0.14)</td>
</tr>
<tr>
<td></td>
<td>-1.9 (1.2)**</td>
<td>+13.3 (2.5)</td>
<td>+8.7 (1.7)</td>
</tr>
<tr>
<td>Eccentric (n=12)</td>
<td>1.64 (0.15)</td>
<td>1.62 (0.16)</td>
<td>1.80 (0.18)</td>
</tr>
<tr>
<td></td>
<td>-1.8 (1.1)**</td>
<td>+10.1 (2.6)</td>
<td>+8.6 (2.2)</td>
</tr>
<tr>
<td>Complicated (n=12)</td>
<td>1.33 (0.12)</td>
<td>1.24 (0.11)</td>
<td>1.55 (0.13)</td>
</tr>
<tr>
<td></td>
<td>-7.0 (1.6)</td>
<td>+16.5 (3.7)</td>
<td>+10.8 (2.1)</td>
</tr>
</tbody>
</table>

Values are mean (SD). Values in italics are percentage change from baseline.

$p < 0.05$; **$p < 0.01$ for complicated stenoses.
between stenosis severity and LNMMA responses as well (p = 0.05).

STENOSIS MORPHOLOGY AND RESPONSE TO GLYCERYL TRINITRATE
There was no difference in the effect of glyceryl trinitrate on proximal reference segments of concentric, eccentric, and complicated stenoses (table 2). A similar increase in lumen diameter of concentric, eccentric, and complicated stenoses (table 2, fig 1) occurred, and a similar proportion showed > 5% increase from baseline (table 1).

Discussion
The results of this study show that complicated (irregular) coronary stenoses are more likely to constrict significantly and constrict to a greater degree in response to LNMMA than eccentric (smooth) and concentric (smooth) stenoses in patients with chronic stable angina. Furthermore, the enhanced constriction is focal at the site of the stenosis, as indicated by the results showing a larger change in lumen diameter at the site of the stenosis than at the adjacent reference arterial segment. They provide evidence to support the hypothesis that complex morphology is characteristic of an active lesion producing nitric oxide. There was also a correlation between the response to LNMMA and stenosis severity and length, but no correlation with eccentricity ratio.

ATHEROMATOUS STENOSES AND NITRIC OXIDE ACTIVITY
Nitric oxide reduces vascular smooth muscle tone by stimulation of soluble guanylate cyclase and has an antioxidant effect. However, when it combines with equimolar amounts of superoxide, peroxynitrite is formed, which is a strong oxidant. Inducible nitric oxide synthase can produce large amounts of nitric oxide and is present in human atherosclerotic lesions. It has been suggested that diseased arteries may be relatively deficient in the substrate L-arginine. Apart from limiting nitric oxide production, substrate deficiency could lead to the generation of superoxide by both inducible and endothelial nitric oxide synthase.

The absence of significant response to LNMMA of smooth (concentric and eccentric) stenoses despite a dilator response to nitrates suggests that very little basal production of nitric oxide occurs at this site. This would be consistent with the destruction or malfunction of endothelium which is known to occur at the site of atheromatous plaques. There does appear, however, to be significant variation between patients, as 20–25% of these stenoses constricted by more than 5%. The correlation of stenosis severity with the response to LNMMA could indicate enhanced nitric oxide production related to greater shear stress at the site of severe stenoses. However, increased severity of stenosis may be related to greater bulk and complexity of the plaque and it is therefore difficult to identify any contribution from stenosis severity as an isolated entity. The response of the six most severe complicated stenoses was not significantly different from the six least severe stenoses.

Figure 2 Individual lumen diameter per cent change from baseline in concentric, eccentric, and complicated stenoses after LNMMA infusion. Horizontal bars indicate the mean values.

Figure 3 The magnitude of stenosis and of reference (ref) segment constriction are expressed as diameter after saline minus diameter after LNMMA (mm) for concentric, eccentric, and complicated stenoses. Complicated stenoses showed significantly greater constriction (p < 0.05) than concentric or eccentric stenoses.

Figure 4 Correlations between stenosis length (right panel), stenosis severity (middle panel), stenosis eccentricity ratio (left panel), and the magnitude of LNMMA response, expressed as per cent change in coronary artery diameter from baseline. Linear regression analysis showed a significant correlation between stenosis severity and LNMMA response, and between stenosis length and LNMMA response.
Stenosis morphology and nitric oxide synthesis

from that of the six least severe complicated stenoses.

**COMPLEX PLAQUES AND NITRIC OXIDE ACTIVITY**

Clinical studies have suggested that stenoses with a complex morphology rapidly progress to total or subtotal occlusion and are often the substrate for acute coronary syndromes. Our data indicate the presence of enhanced nitric oxide synthase activity possibly the inducible form at the site of complex plaques. A possible explanation for this is that nitric oxide production at the site of stenosis occurs in the endothelium of the new microvessels in the wall of the artery around the atheromatous plaque or in other cell types in the vascular wall such as macrophages and smooth muscle cells. Such neovascularisation of atheromatous plaques is well documented and a recent study showed the presence of inducible nitric oxide synthase in the neomicrovasculature. Nitric oxide production from endothelial cells lining the epicardial segment lumen could be increased because of greater shear stress at the site of complex stenoses. Furthermore, platelet derived nitric oxide activity, as has been shown recently, could be increased by greater platelet adhesion and aggregation at the site of complex stenoses.

There may be a link between enhanced inducible nitric oxide activity (produced by macrophages) and plaque instability. An abundance of macrophages is often found in such plaques. The macrophages and the T lymphocytes may induce nitric oxide synthase mediated by cytokines. Furthermore, metalloproteinases produced in macrophages in vulnerable regions of human atherosclerotic plaques may weaken the fibrous cap of the plaque. There is also growing evidence that macrophages may be involved in smooth muscle cell death by apoptosis occurring in plaques, perhaps reflecting a decline in growth factors or the activity of macrophages. It is well known that macrophages have the ability to initiate or enhance the degradation of collagen and to promote the propensity of those plaques to rupture and trigger thrombosis. It might appear that our results indicating enhanced basal nitric oxide activity within complex irregular plaques are in conflict with our previous observation of an enhanced constrictor response of complex stenoses to intracoronary serotonin or acetylcholine. However, a constrictor response to serotonin and acetylcholine does not necessarily indicate a reduction in endothelial dilator function as these two agents have a direct constrictor effect on vascular smooth muscle. The constrictor response could reflect smooth muscle cell hyperreactivity, and the coexistence of such hyperreactivity with increased nitric oxide synthase activity cannot be excluded. The results of this study provide further evidence that complex plaques are active structures. Irrespective of increased nitric oxide activity and vascular smooth muscle reactivity, the resting tone appears to be similar for both smooth and complicated stenoses, as evidenced by the similar magnitude of dilatation in response to nitrate administration.

**CLINICAL IMPLICATIONS**

As the incidence of complex stenosis morphology is about 60% in patients with unstable angina before myocardial infarction and approximately with about 20–30% in patients with chronic stable angina, greater emphasis should be placed on the detection of complex stenosis. This will require the development of new and preferably non-invasive methods of detecting active plaques. An improved understanding of the cellular and molecular mechanism of plaque instability is required.

**CONCLUSIONS**

Basal nitric oxide activity in atheromatous plaques is closely related to the morphology of stenosis. It is greater at the site of complicated (irregular) stenoses than at the site of smooth stenoses. This may be related to the cellular composition of complex lesions.

A 65 year old man with a five month history of low grade fever, fatigue, and asthenia was admitted to our hospital. Physical examination findings were normal except for a grade III/VI systolic murmur at the apex. There was anaemia (6.3 mmol/l) with an elevated erythrocyte sedimentation rate (54 mm/hour). Blood cultures grew *Enterococcus faecalis*, and the patient was treated with penicillin and gentamicin for six weeks. Transthoracic and transoesophageal echocardiography demonstrated a vegetative mass in the aortic valve, aortic regurgitation across the mitral valve (LA, left atrium; LV, left ventricle; RA, right atrium; RV, right ventricle). Doppler examination revealed a moderate to severe mitral regurgitation across the mitral valve aneurysm. Because of mitral and aortic regurgitation and haemodynamic deterioration, the patient underwent aortic and mitral valve replacement. The aneurysm of the anterior mitral leaflet was confirmed during surgery. Histologic examination of the resected segments showed focal fibrin deposits and acute inflammation foci, consistent with the diagnosis of infective endocarditis. The postoperative course was uneventful and subsequent blood cultures were negative.

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