The haemodynamic response to myocardial ischaemia in ambulant patients with variant angina

RICHARD D LEVY, LEONARD M SHAPIRO, CHRISTINE WRIGHT, LORNA J MOCKUS, KIM M FOX
From the National Heart Hospital, London

Summary: The haemodynamic response to myocardial ischaemia in patients with variant angina during ambulatory activity is unknown. Ambulatory pulmonary artery pressure monitoring with a transducer tipped catheter and simultaneous frequency modulated electrocardiograms was used to assess changes in left ventricular function in five male patients (mean age 51.8 years) during variant angina; four patients had coronary artery stenosis and one had normal coronary arteries. Two hundred and seventy hours of ambulatory recordings were analysed. Twenty episodes (12 painful, 8 silent) of ST segment change > 1 mm occurred. Episodes tended to occur more frequently in the early morning hours. Six episodes of painful ST elevation were associated with a rise in pulmonary artery diastolic pressure. In the remaining episodes ST segment elevation was of shorter duration and there was no rise in pulmonary artery diastolic pressure. Pain was usually a late feature. Silent ST segment elevation occurred at rest and pulmonary artery diastolic pressure increased in all but one episode. Silent exertional ST segment depression was associated with a greater increase in pulmonary artery diastolic pressure than that seen during ST segment elevation. ST segment depression preceded or followed ST segment elevation in two episodes. The onset of ST segment elevation nearly always preceded the onset of a rise in pulmonary artery diastolic pressure. Ergometrine maleate provocation produced a rise in pulmonary artery diastolic pressure in three patients. In one there was no response to 1000 µg but spontaneous episodes of ST segment elevation were recorded during ambulatory monitoring. Treadmill exercise resulted in both ST segment elevation and depression with a similar haemodynamic response during both types of electrocardiographic change. When there is important coronary artery disease in two or more vessels ST segment changes may occur in different territories during treadmill exercise and during spontaneous episodes.

Ambulatory pulmonary artery diastolic pressure monitoring is a useful technique for the investigation of variant angina.

In 1959 Prinzmetal et al reported on a group of patients in whom angina occurred at rest and was associated with transient cyclic ST segment elevation. Termed variant angina, this syndrome was thought to be secondary to “temporary occlusion of a large diseased artery with a narrow lumen due to an increase in tonus of the vessel wall”. Later studies have shown that variant angina is part of a spectrum of “vasospastic myocardial ischaemia” and that variant angina may occur in normal coronary arteries or more commonly in those with atherosclerotic coronary artery disease. Myocardial ischaemia during variant angina is thought to result from a reduction in myocardial oxygen supply. ST segment depression during myocardial ischaemia in patients with coronary artery disease is associated with a rise in left ventricular end diastolic pressure.

Previous studies of the changes in left ventricular end diastolic pressure during variant angina have been confined to patients in the coronary care unit or catheter laboratory. In the present study we assessed the haemodynamic response during variant angina by means of ambulatory pulmonary artery pressure monitoring.
Patients and methods

Five male patients (mean age 51.8, range 40–62 years) were studied. Table 1 summarises the clinical, electrocardiographic, and angiocardiographic data. All patients had a relatively short history of anginal chest pain and four patients had rest pain which was mostly nocturnal. Variant angina was defined as ST segment elevation accompanied by chest pain on ambulatory monitoring or ergometrine maleate provocation. Patients 1, 2, 4, and 5 had confirmed coronary artery spasm at the time of ergometrine maleate provocation during coronary angiography. All antianginal medication other than glyceryl trinitrate was discontinued 48 hours before the study. The protocol was approved by the National Heart Hospital ethics committee and patients gave their informed and written consent.

PULMONARY ARTERY PRESSURE MONITORING
A polyurethane 6 French NIH type catheter with a miniature strain gauge transducer mounted on the tip was used. This was calibrated after immersion in saline for one hour before use. The transducer was driven and demodulated by an electrically isolated Gaeltec pre-amplifier. The catheter was introduced percutaneously via a subclavian vein to a proximal pulmonary artery under fluoroscopy in the cardiac catheter laboratory. The pulmonary artery pressure was recorded on an Oxford Medilog I miniature tape recorder that had been modified by the insertion of an AM4 pressure module.

ST SEGMENT MONITORING
ST segment was monitored on a frequency modulated recorder and leads CM2 and CM5 were recorded. The electrocardiogram was replayed on an Oxford MA20 scanner. Changes in the ST segment were measured 80 ms after the J point to an accuracy of 0.1 mm by means of a magnifying lens equipped with a graticule. A change of 1 mm in the ST segment that lasted >30 seconds was considered to be important. ST segment changes were analysed on a beat to beat basis for five minutes before each episode and again up to five minutes after the ST segment had returned to basal levels. The pulmonary artery recorder was linked to the frequency modulated electrocardiogram by an event button that marked both the frequency modulated electrocardiogram and the pulmonary artery trace.

CORONARY ARTERIOGRAPHY
Coronary arteriography was performed in all patients by the Judkins’ technique from the femoral artery with multiple angiographic projections.

ERGOMETRINE MALEATE PROVOCATION TEST
Patient 1 had a previously positive provocation test and this test was not repeated. Patients 2–5 underwent ergometrine maleate provocation in the catheter laboratory. A 12 lead surface electrocardiogram was recorded continuously (Siemens Mingograph). Incremental doses of ergometrine maleate were given intravenously every five minutes. The initial dose was 25 µg and this was increased to 50 µg and 100 µg and subsequently by 100 µg and 200 µg up to a maximum to 1000 µg. The test was terminated if there was chest pain, ST segment changes >2 mm, severe hypertension, headache, nausea, or vomiting. Angiographic assessment for the presence of coronary artery spasm was made at the end of the test and was followed immediately by 1–3 mg of intracoronary isosorbide dinitrate. Changes in blood pressure and pulmonary artery pressure were recorded continuously during provocation. Spasm at arteriography was defined as a transient total or subtotal coronary occlusion with delayed distal filling that was reversed by intracoronary nitrates.

AMBULATORY MONITORING
The patients returned to the ward with a pulmonary artery catheter in situ. Continuous monitoring of pulmonary artery pressure was performed with simultaneous recording of frequency modulated electrocardiograms from leads CM2 and CM5. Patients were instructed to keep diaries during the period of ambulatory monitoring and were asked to note the time of onset and severity of chest discomfort as well as their activity at the time of pain.
addition, they were asked to press the event marker so that the ST segment and pulmonary artery traces were marked.

**EXERCISE TEST**

Exercise testing was performed according to a modified Bruce protocol. Transducer tipped pulmonary artery pressure and frequency modulated electrocardiogram were recorded continuously. A 12 lead electrocardiogram was recorded on paper at one minute intervals. The exercise test was terminated by angina, dyspnœa, multiple ventricular extrasystoles, hypotension, ST segment depression or elevation >0.3 mV, or exhaustion.

At the end of the period of ambulatory monitoring the pulmonary artery catheter was removed and the pulmonary artery trace was replayed via a PB2 unit and PM3 amplifier. This was displayed on an SE laboratories 6008 oscillograph. The entire period of recording of the pulmonary artery diastolic pressure was displayed and measured on ultraviolet paper with a calibrated scale from a zero reference point. The pulmonary artery diastolic pressure was measured during both painful and painless episodes of ST segment depression on a beat to beat basis five minutes before the onset of the earliest change in pulmonary artery pressure and five minutes after the pulmonary artery diastolic pressure had returned to baseline. The pulmonary artery diastolic pressure was measured at the end of expiration.

**STATISTICAL ANALYSIS**

A change of at least three standard deviations from the mean level over a five minute period before any change in pulmonary artery pressure, heart rate, or ST segment was regarded as statistically significant. Wilcoxon rank sum testing was used for analysis.

**Results**

**ERGOMETRINE MALEATE PROVOCATION**

Four patients had angiographically confirmed coronary artery spasm after ergometrine maleate provocation. In patient 3 chest pain developed after a dose of 200 μg in the absence of ST segment changes and patient 4 was unaffected by a total dose of 1000 μg. Patients 3 and 4 had spontaneous episodes of ST segment elevation. The haemodynamic response to ergometrine maleate was variable (table 2). Pulmonary artery diastolic pressure tended to rise (range 1.5–12.4 mm Hg (18.8–117%)). Patient 3 had no ST segment change despite a rise in pulmonary artery diastolic pressure of 8 mm Hg (100%). The most pronounced haemodynamic response to ergometrine maleate occurred in patient 2 (fig 1). A significant rise in pulmonary artery diastolic pressure occurred 102 seconds after the dose of 100 μg; 42 seconds later the heart rate increased and 66 seconds later there was an abrupt rise in the ST segment. Pain occurred six seconds after the onset of ST segment elevation. The time to maximum pulmonary artery pressure was 220 seconds, maximum heart rate (145 beats/minute) 228 seconds, and maximum ST segment elevation 250 seconds. The episode of ST segment elevation was followed by ST segment depression and resolution of pain after isosorbide dinitrate.

**EXERCISE TESTING**

Patients 1–4 underwent treadmill testing (table 3). Three patients developed ST segment depression of >1 mm and one had ST elevation. In all cases there was a rise in pulmonary artery diastolic pressure. The greatest change in pulmonary artery diastolic pressure (12.4 mm Hg (775%)) occurred in the patient with ST segment elevation of 2.8 mm. This was the only patient in whom chest pain developed. There was a poor correlation between the magnitude and duration of ST segment change (r = 0.16, p >0.05) but there was a closer relation between the magnitude and duration of pulmonary artery diastolic pressure (r = 0.74, p < 0.05). Duration of exercise correlated with the duration of elevation in pulmonary artery diastolic pressure (r = 0.84, p < 0.05). In patient 3, who had ST segment elevation, the onset of the change in the ST segment and rise in pulmonary artery diastolic pressure were simultaneous. In two of the other three patients with ST
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Fig 1 Haemodynamic and electrocardiographic response to a total dose of 100 µg of ergometrine maleate in patient 2. A significant rise in pulmonary artery diastolic pressure occurred 102 seconds after the 100 µg dose of ergometrine. Forty two seconds later the heart rate increased and 66 seconds later there was an abrupt rise in the ST segment. Pain occurred six seconds after the onset of ST segment elevation. The time to maximum pulmonary artery pressure was 220 seconds and to maximum ST segment elevation 250 seconds, and a maximum heart rate of 145 beats per minute occurred at 228 seconds. The episode of ST segment elevation was followed by ST segment depression and resolution of pain after isosorbide (Iso).

Table 3 Response to treadmill exercise

<table>
<thead>
<tr>
<th>Patient</th>
<th>Magnitude of ST segment response (mm)</th>
<th>Rise in pulmonary artery diastolic pressure (mmHg) (%)</th>
<th>Pain (+/−)</th>
<th>Duration of exercise (min)</th>
<th>Duration of ST segment change (min)</th>
<th>Duration of raised pulmonary artery diastolic pressure (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>−2·0</td>
<td>10 (77)</td>
<td>−</td>
<td>12·3</td>
<td>36·5</td>
<td>20·1</td>
</tr>
<tr>
<td>2</td>
<td>−1·6</td>
<td>5·4 (207)</td>
<td>−</td>
<td>13</td>
<td>2·6</td>
<td>13·5</td>
</tr>
<tr>
<td>3</td>
<td>+2·8</td>
<td>12·4 (775)</td>
<td>+</td>
<td>5·2</td>
<td>14·2</td>
<td>5·3</td>
</tr>
<tr>
<td>4</td>
<td>−2·1</td>
<td>3·5 (12)</td>
<td>−</td>
<td>9·5</td>
<td>11</td>
<td>9·2</td>
</tr>
</tbody>
</table>

Twelve of these were painful and the remainder were not. Figure 2 summarises the diurnal distribution of episodes. Most episodes occurred between 4 am and 8 am. Table 4 shows the number of episodes and the duration of ambulatory monitoring for each patient.

AMBULATORY EPISODES

Two hundred and seventy hours of ambulatory recording of pulmonary artery pressure and frequency modulated electrocardiogram were analysed and 20 episodes of ST segment change were noted.

PAINFUL EPISODES

Eleven episodes were associated with ST segment elevation (median 2·15 mm, range 1–3·3) and one with 1 mm ST depression. Pain was a late feature in
nearly all cases, following the point of maximal ST segment elevation or depression. There was a rise in pulmonary artery diastolic pressure during six of the episodes of ST segment elevation. The onset of rise in pulmonary artery diastolic pressure was simultaneous with or followed the rise in the ST segment in all but one episode. Maximum pulmonary artery diastolic pressure and maximum ST segment elevation tended to coincide (p < 0.05). Ten of the eleven episodes of ST segment elevation occurred at rest. Glyceryl trinitrate was taken on one occasion and there was no change in pulmonary artery diastolic pressure. ST segment depression preceded or followed ST segment elevation in two episodes. Episodes of ST segment elevation not associated with a rise in pulmonary artery diastolic pressure were of shorter duration (median 1.15 minutes, range 0.6-2.6) than those episodes with a rise in pulmonary artery diastolic pressure (median 4.65 minutes, range 2–6). ST segment changes during different episodes usually occurred in the same territory in each patient.

PAINLESS EPISODES

Four episodes were associated with ST segment elevation (1.2–2.2 mm) and there was a rise in pulmonary artery diastolic pressure (66–344%) in three of these episodes that occurred simultaneously with or after the onset of ST segment elevation. There was no rise in pulmonary artery diastolic pressure in the remaining episode (patient 3) despite 27 minutes of ST segment elevation reaching a maximum of 2.2 mm. All episodes of ST segment elevation occurred at rest and one episode was followed by ST segment depression. ST segment elevation (median 1.25 minutes, range 0.8–27) lasted longer than the rise in pulmonary artery diastolic pressure (median 0.6 minutes, range 0.4–1) (p < 0.05). Four episodes of ST segment depression (1–1.2 mm) in the absence of ST segment elevation were recorded. All occurred on exertion and all were associated with a rise in pulmonary artery diastolic pressure (57–275%). The onset of rise in pulmonary artery diastolic pressure preceded the onset of ST segment depression in two episodes, followed it in one, and was simulta-

Table 4  Number of episodes of ST segment elevation during ambulatory monitoring

<table>
<thead>
<tr>
<th>Patient</th>
<th>Ambulatory monitoring (h)</th>
<th>Daytime painful episodes</th>
<th>Daytime silent episodes</th>
<th>Nocturnal painful episodes</th>
<th>Nocturnal painless episodes</th>
<th>Total</th>
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<td>0</td>
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<td>2</td>
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</table>
neous with the remaining episode. The duration of a raised pulmonary artery diastolic pressure on exertion (median 1·95 minutes, range 1·8–2) was longer than that during episodes at rest (median 0·6 minutes, range 0·4–1) (p < 0·05).

Discussion

We examined the haemodynamic response to myocardial ischaemia in patients with variant angina by means of simultaneous ambulatory pulmonary artery pressure and ST segment monitoring. Previous studies have been limited to patients confined to bed in the coronary care unit and have used fluid filled catheter systems.2 3 5 10 The transducer tipped catheter technique used in this study is both accurate and allows for full ambulation.8 We have shown that changes in pulmonary artery diastolic pressure do not precede ST segment elevation in spontaneous variant angina and this confirms the results of previous studies.2 11 12

Symptomatic and asymptomatic ST segment changes have been recorded by ambulatory monitoring13 and in the coronary care unit.2 In our study episodes of symptomatic ST segment elevation were not always associated with a rise in pulmonary artery diastolic pressure. The presence or absence of a rise in pulmonary artery diastolic pressure was related to the duration of ST segment elevation in some patients. Pain was a late feature in most cases.14–17 In asymptomatic episodes there was a different haemodynamic response in those with ST segment elevation at rest and in episodes with exertional ST segment depression. The magnitude of pulmonary artery diastolic pressure increase was greatest on exertion.

It has been suggested that episodes of ST segment depression and ST segment elevation occurring within a few minutes of each other show similar haemodynamic patterns.2 17 The interpretation of the haemodynamic changes may be difficult when ST segment depression and elevation occur sequentially.

Episodes of ST segment elevation tend to occur in the early morning.12 18–20 Eleven of the 20 episodes occurred between 4 am and 8 am. Coronary artery tone may be increased at night and is highest in the early morning21 22; however, it has also been shown that there is no increase in sympathetic activity to the heart before the onset of ST segment elevation.23 Nocturnal ST segment elevation has been associated with rapid eye movement sleep or changes in sleep pattern24 but others have refuted these findings.23 25 There is a diurnal variation in pulmonary artery diastolic pressure with a significant rise during the night with peak levels being reached in the early morning hours.26 This may be of importance because the raised left ventricular filling pressure in the early morning hours coincides with the largest number of episodes of ST segment elevation at rest.

Ergometrine maleate is an α adrenergic agonist that is used to assess the responsiveness of the coronary arteries to vasoconstrictor stimuli. This has been shown to be a highly specific test for coronary artery spasm in 90–100% of cases.12 Ergometrine maleate increases left ventricular dimension, with resultant poor contractility,27 and produces a dramatic reduction in coronary sinus flow consistent with an increase in coronary vascular resistance.28 29 We have shown that this test was positive in four of the five patients with variant angina. Patient 4 was totally unresponsive to a dose of 1000 μg of ergometrine maleate and yet had documented spontaneous episodes of ST segment elevation on ambulatory monitoring. Patient 3 developed chest pain after 200 μg of ergometrine maleate with no ST segment changes. One patient had a dramatic response to a total dose of 100 μg of ergometrine maleate with ST segment elevation of 6·5 mm. As a group the change in pulmonary artery diastolic pressure was variable but the response in patient 2 (fig 1) showed that the rise in pulmonary artery diastolic pressure occurred before the heart rate and ST segment changes.

Pharmacological interference with ergometrine maleate may produce coronary artery spasm but appears to have a different haemodynamic effect from that seen during spontaneous episodes. A study in which a transducer tipped catheter was used showed a rise in left ventricular end diastolic pressure which preceded pain and ST segment elevation in a single patient after 330 μg of ergometrine maleate.30 Patient 1 had only one episode and patient 5 no episodes of ST segment elevation during studies lasting 120 and 48 hours respectively. Both of these patients had positive ergometrine maleate provocation tests with coronary artery spasm at angiography and previously documented episodes of ST segment elevation. Thus disease activity can vary such that patients may have multiple daily attacks for some time followed by a pain free interval of several weeks.

Exercise testing in untreated patients with acute variant angina has been shown to produce ST segment elevation in approximately 30% of cases, ST segment depression in 30%, and no significant change in the remainder. The ST segment response is not predictive of the presence of or the severity of underlying fixed coronary artery disease.31 Four of the five patients in our study performed an exercise test. Three had ST segment depression >1·5 mm
with a rise in pulmonary artery diastolic pressure of 116–207% (3–5–10 mm Hg). This response to treadmill exercise was similar to that reported in a previous study of 15 patients with documented coronary artery disease and no documented spasm in whom there was a rise in pulmonary artery diastolic pressure (median 5 mm Hg, 69%; range 0–13–8 mm Hg, 0–262%). Only one patient in the latter group failed to have a rise in pulmonary artery diastolic pressure in the presence of important ST segment depression.7

ST segment depression during an exercise test has been shown to occur in different leads from those showing ST segment elevation at rest, suggesting that the same myocardial region may not be affected by ischaemia at rest and during exercise.18 32 In one patient (case 4) ST segment depression developed in lead CM5 and the anterior V leads during exercise whereas during several episodes at rest there was ST segment elevation in the inferior leads. Coronary arteriography in this patient showed irregularities in the left anterior descending and circumflex coronary arteries with a 50% stenosis in the proximal right coronary artery.

Various mechanisms for angina secondary to coronary spasm have been suggested since spasm was first suggested as a possible mechanism for angina pectoris.32 33 Angina has been described by Prinzmetal in patients with normal coronary arteries1 but is more commonly associated with fixed coronary artery obstruction.23 Only one patient in this study had normal coronary arteries.

 Coronary blood flow measured by thermodilution techniques has been shown to fall during spontaneous variant angina.29 A fall in coronary sinus oxygen saturation consistently precedes ST segment elevation, thus indicating that this factor has a causal role in the reduction in coronary blood flow.3 Myocardial scintigraphy with thallium–2014 34 and 1–131 labelled microaggregated human serum albumin13 has shown that transmural defects are present during ST segment elevation and absent in the resting state.

 Some workers have shown that coronary artery spasm is a frequent cause of angina18 36 but Quyyumi et al noted that of 100 consecutive patients referred with chest pain for coronary arteriography, only two had ST segment elevation on ambulatory monitoring.37 Our study was confined to five patients and has confirmed that ST segment elevation is infrequent and seldom, if ever, occurs other than in the resting state.

 The technique of ambulatory pulmonary artery pressure monitoring has provided a means of studying left ventricular function in ambulant patients with variant angina. We have confirmed that changes in pulmonary artery diastolic pressure, reflecting changes in left ventricular end diastolic pressure, do not precede the onset of ST segment elevation in most episodes of variant angina. The response is variable, however, when the changes are provoked by exercise testing or ergometrine maleate.

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