Editorial

Silent ischaemia: clinical implications in 1988

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Within the past few years the concept of silent ischaemia has become widely known. But silent ischaemia is not a new finding. Paul Wood and his colleagues described the development of electrocardiographic changes during exercise testing in the absence of chest pain and suggested that this could represent latent angina.1 In 1974 Stern and Tzivoni described asymptomatic alteration in the ST segment recorded during ambulatory monitoring in patients with coronary artery disease.2 Subsequent studies showed that such ST segment changes were common.34 Initially, concern that these ST segment changes represented a “false positive” response was fuelled by the debate over the use and value of exercise testing as a diagnostic tool.56

Later, invasive monitoring in the catheter laboratory, positron tomography, and ambulatory pulmonary artery monitoring showed that most of these episodes of ST segment depression reflect myocardial ischaemia,78 and it has been proposed that routine clinical management of patients with angina pectoris should include the identification and treatment of silent ischaemia. In the United Kingdom, however, routine ambulatory monitoring to detect and treat silent ischaemia is seldom performed.

It has been suggested that about 60–80% of patients with chronic stable angina have silent ischaemia.910 Unfortunately, most of these data were obtained from highly selected patient populations. Recently we studied 150 unselected patients with angina pectoris on no medical treatment.11 Silent episodes of ST segment depression were about three times as frequent as angina pectoris: one third of the patients had both silent episodes and painful episodes of ST segment depression and 20% had silent episodes without angina pectoris. Also, about a third of the patients had 90% of the silent episodes of ST segment depression. So silent ischaemia is most likely to be important in about one third of patients with chronic stable angina; other workers also have come to similar conclusions.12

This does not imply that silent ischaemia is unimportant. In the United Kingdom about a million patients have chronic stable angina and so about 300 000 would have frequent episodes of silent ischaemia. But these data came from patients with chronic stable angina who were not on medical treatment and who had cardiac catheterisation. Similar studies have not been performed in medically treated patients with stable angina; such patients are likely to have fewer episodes of silent ischaemia.

Workers who compared changes in heart rate with changes in the ST segment during ambulatory monitoring and exercise testing concluded that silent ischaemia is predominantly related to an alteration in coronary blood flow.1311 But analysis of the heart rate during silent episodes and painful episodes of ST segment depression during ambulatory monitoring does not support this conclusion.14 The 24 hour diurnal distribution of silent and painful ischaemia is similar; in patients with chronic stable angina there are more episodes between 9 am and 10 pm than during the rest of the day and night,15 while in patients with variant angina, silent episodes and painful episodes are most common at night.16 Analysis of heart rate changes has shown that both silent episodes and painful episodes of ST segment depression can on occasion be preceded by increase in the heart rate while other episodes within the same patient are not. Mental stress can provoke myocardial ischaemia; often this is silent.17 The pathophysiological mechanisms for this picture are unclear but an increase in myocardial oxygen demand and coronary vasoconstriction are likely to be important. Thus in chronic stable angina, silent ischaemia could be caused by various mechanisms as is angina pectoris. These mechanisms will vary from patient to

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patient and even from hour to hour within the same patient. The prediction that if the pathophysiological mechanisms of silent ischaemia and angina are similar both conditions will respond to the same treatment is borne out by reports that drugs that relieve angina can also suppress silent ischaemia. 18-20

Although silent ischaemia can be treated with antianginal agents it is still far from certain that such treatment is necessary or justifies the extra cost and staff it would require. Silent ischaemia is best identified by 24 hour ambulatory monitoring of the electrocardiogram. Amplitude modulated systems give a more stable baseline and are more widely available but frequency modulated systems give a more appropriate frequency response. 21 The new generation of solid state systems that are now available gives computerised reports and these need little attention from a technician. 22 Although solid state systems are less labour intensive they require validation and they are expensive.

Must ambulatory monitoring be used to identify silent ischaemia? Perhaps the investigations could be targeted more appropriately? Selwyn and Ganz suggested that angina pectoris does not reflect the true severity of myocardial ischaemia and that ambulatory monitoring may be needed to identify those patients with the worst prognosis. 23 But in the same issue of the Journal of Cardiovascular Nursing, Epstein et al claimed that electrocardiographic monitoring added little to the prognostic information provided by exercise testing. 24 Certainly, silent ischaemia is rare in those patients with a negative exercise test and in those whose exercise test is positive only at high workload. 25,26 Also there seems to be a close linear correlation between the time to 1 mm ST segment depression recorded during exercise testing and the frequency of silent ischaemia; perhaps therefore ambulatory monitoring could be restricted to those patients with ST segment depression at low workloads on the exercise test.

Although silent ischaemia can be treated and the use of exercise testing with ambulatory monitoring might not overstretch resources, we still do not know whether silent ischaemia in patients with chronic stable angina should be identified and treated. Prognostic data are woefully lacking. In the 1970s Eriksson et al investigated 2014 symptom free male factory workers by exercise testing and coronary angiography. 27 Fifty men had a positive exercise test and confirmed coronary artery disease without chest pain. In the next eight years 10 (20%) developed an acute myocardial infarction or died. But it is still not certain that these subsequent cardiovascular events had anything to do with silent ischaemia; they may have merely reflected the underlying coronary artery disease. More importantly, would treatment of silent ischaemia have prevented these events? Acute myocardial infarction is usually caused by rupture of an atherosclerotic plaque and the same mechanism almost certainly causes sudden death. 29 What effect will antianginal agents have? At the present time there is no evidence to suggest that preventing ischaemic episodes either by reduction of myocardial oxygen demand or by increasing coronary blood flow will have any effect on plaque rupture and thrombosis. Perhaps the frequency of silent ischaemia together with a diminution of exercise tolerance are markers that a stable plaque has become unstable; in such circumstances it might be better to stabilise the plaque and thrombotic process with drugs that prevent surges in blood pressure, smooth muscle relaxants, antithrombotic and anti-platelet agents, or to operate or perform angioplasty.

Another suggestion is that repeated episodes of myocardial ischaemia lead to small areas of myocardial cell death and subsequent reduction of left ventricular function. 29 If this is true, drugs that reduce the frequency and duration of silent ischaemic insults may be beneficial. Preliminary data support this theory, 30 but further studies are needed to resolve the question.

In contrast, in unstable angina evidence suggests that episodes of painful and silent ischaemia do have important prognostic implications. Gottlieb and his co-workers showed that silent ischaemia in patients with unstable angina was associated with an increased number of cardiovascular events, particularly myocardial infarction and death. 31 The most likely mechanism for the angina that precedes infarction is an unstable plaque with superadded platelet aggregation and possibly thrombosis. 28 Appropriate treatment of such ischaemia may prove rewarding. Again, however, although drugs that reduce myocardial oxygen demand and smooth muscle relaxants permit the myocardium to withstand greater ischaemic insults, they are unlikely to prevent myocardial infarction. In the recent Holland Interuniversity Nifedipine/Metoprolol trial the risk ratio of preventing acute myocardial infarction with β blocker alone was 1:07 and that for calcium antagonist was 1:51. 32 The use of drugs or interventions that directly influence the pathogenesis may, therefore, have to be considered—such as antiplatelet agents, coronary angioplasty, and bypass surgery. 33 Most clinicians would probably consider exercise testing unwise in patients with unstable angina, and repeated ambulatory monitoring to assess the frequency, duration, and severity of silent and painful ischaemia may well prove to be the most suitable investigation.

Silent ischaemia may be an important feature for identifying patients who will require further investigation and careful consideration of treatment. But
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much of this is speculation and until firm data have been collected routine ambulatory monitoring of the ST segment cannot be advised. In the United States there are devices (Q Med) to monitor the ST segment; ST segment depression sets off an alarm that prompts the patient to take a tablet of glyceryl trinitrate. The barometer of silent ischaemia has yet to settle; I hope that it will not be at this extreme.

References


K M Fox

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