Ambulatory scintigraphic assessment of transient changes in left ventricular function: a new method for detection of silent myocardial ischaemia

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Demonstration of ischaemic left ventricular dysfunction in the absence of chest pain should provide important confirmation of silent myocardial ischaemia in patients with asymptomatic ST segment changes. For this purpose, a new portable scintillation probe (VEST) similar to a miniaturized nuclear stethoscope combined with a Holter ECG was evaluated. After standard equilibrium radionuclide angiography with technetium-99m labelled red blood cells, the VEST was positioned under gamma-camera control and data were recorded from 1–12 h in 61 unselected patients. Ejection fraction (LVEF), relative changes in volumes, heart rate and ST segment changes were determined. Reproducibility of LVEF at rest (r = 0.91; variability 3.8 ± 3%, N = 19) and during exercise (r = 0.98; variability 3.2 ± 2%, N = 19) was good. In 15 asymptomatic exercise tests four different patterns of LVEF and ST segment responses were identified: (1) decrease in LVEF followed by significant ST depression (five times); (2) ST depression followed by decrease in LVEF (three times); (3) decrease in LVEF without significant ST changes (three times); and (4) ST depression without significant LVEF change (four times). In this small series, patterns (1) to (3) corresponded to patients with documented coronary artery disease, which was not the case for pattern (4). For detection of silent ischaemia at rest, a decrease in LVEF of > 5% lasting for > 1 min was defined as ischaemic LV dysfunction. Using this definition, four spontaneous episodes of silent LV dysfunction could be demonstrated in two of three CCU patients with unstable angina during 160–680 min of data recordings without simultaneous ST changes. Based on this initial experience, we conclude that VEST is a reproducible method to detect transient global LV dysfunction and will be useful to confirm silent ischaemia in otherwise uncertain ST segment changes.

Introduction

The diagnosis of silent ischaemia is limited by the lack of a gold standard for detection of myocardial ischaemia. Whereas there are invasive haemodynamic, metabolic and angiographic as well as non-invasive scintigraphic methods to confirm asymptomatic ST segment depression as a sign of myocardial ischaemia, these methods are restricted mostly to use in the laboratory. Tests to provoke (silent) myocardial ischaemia include mental stress, a cold pressure test and drug administration such as ergonovine maleate or dipyridamole besides physical exercise testing. However, spontaneous episodes of silent myocardial ischaemia rarely occur under laboratory conditions. Therefore, the only method for detecting such transient ischaemic episodes which has gained widespread application is ambulatory Holter ECG monitoring[11]. Critical analysis of ST segment changes during daily activities showed that despite high quality recorders and stringent criteria for diagnosis of silent ischaemia, there remains a fairly high percentage of false or uncertain positive (and negative) results[12]. These findings were confirmed by probability calculations indicating that even so-called significant ST segment depression in an asymptomatic patient with low probability of disease may not just be labelled 'silent ischaemia' but require confirmation by another method[3].

With the introduction of a new portable scintillation probe (VEST)[14,31], similar to a miniaturized nuclear stethoscope[6] combined with a Holter ECG recording system, there is the ability to detect transient left ventricular dysfunction during ischaemic episodes on an ambulatory basis. Demonstration of ischaemic left ventricular dysfunction in the absence of chest pain should provide important confirmation of silent myocardial ischaemia in patients with asymptomatic ST segment changes. The aim of this report is to describe and validate the 'VEST' method.
in view of this purpose and to present initial findings in patients with silent myocardial ischaemia.

Patients and methods

PATIENTS
To test and validate the VEST system, 61 unselected patients referred for diagnostic radionuclide angiocardiography were asked to participate in the protocol. The total patient population has been previously described in more detail\[7\]. Out of 98 exercise tests performed in these patients, 15 were totally asymptomatic without anti-ischaemic drugs but showed either significant ST depression and/or a significant drop in left ventricular ejection fraction (LVEF) during exercise. The presence or absence of coronary artery disease was confirmed by coronary angiography or a history/follow-up of myocardial infarction. Three additional patients with unstable angina were studied in the coronary care unit at rest over 160 to 680 min.

RADIONUCLIDE ANGIOCARDIOGRAPHY/VEST
Standard radionuclide angiocardiography was performed at rest and if required during exercise after in vivo labelling of red blood cells with 25 mCi of technetium-99m as previously reported and validated\[8\]. After this diagnostic test, the detector of the VEST system was positioned over the left ventricle under gamma camera control in a left anterior oblique projection such that the left atrium and the right ventricle were not covered by it. The VEST detector consisted of a 5-cm diameter sodium iodide crystal with a parallel hole collimator to maximize sensitivity and field uniformity. To hold the detector in place over the left ventricle and lung, it was fitted with a firm plastic vest-like garment. Counts detected by the crystal and signals from a chest lead of the ECG were recorded on a modified Holter ECG-recorder as previously described\[17\].

Data analysis consisted of reviewing radionuclide data for technical adequacy and to rule out major motion artefacts as indicated by a sudden shift in total counts. Thereafter, ECG and radionuclide data

Figure 1 Silent ischaemia during exercise (H.Z., 55 years, two-vessel CAD). Note decrease in LVEF during exercise before significant ST depression. Note also the overshoot increase in LVEF after exercise and the artefact after 57 min when the patient sat up: sudden drop in total end-diastolic volume. ex, Exercise.
were summed for 15-s intervals to determine ejection fraction, relative changes in end-diastolic and end-systolic volumes, heart rate, cardiac output and ST segment changes. On the basis of previous studies\(^{15-71}\), a fixed background correction of 75% of end-diastolic counts was used to calculate LVEF values. Data were displayed graphically and numerically for analysis. Based on results of variability measurements, a decrease of > 5% was defined as 'ischaemic' left ventricular dysfunction. For the ST segment, a depression of > 1 mm was assumed to be significant.

All exercise tests were performed on a supine bicycle ergometer up to symptom limitation (chest pain, dyspnea or fatigue). For determination of reproducibility of obtained results, some patients had two exercise tests according to previously described protocols\(^{71}\).

RESULTS

Reproducibility of LVEF determined by the VEST system at rest \((r = 0.93; n = 19)\) and during exercise \((r = 0.96; n = 19)\) was good with a low variability of repeated measurements (at rest: ±3%, during exercise: ±2%). Therefore, the exercise induced change in LVEF (delta EF% rest − exercise) was also good \((r = 0.88)\), indicating that transient changes in LVEF can be consistently determined by this method.

In the 15 asymptomatic exercise tests, four different patterns of LVEF and ST segment responses were identified: (1) decrease in LVEF followed by a significant ST depression (five times); (2) ST depression followed by a decrease in LVEF (three times); (3) decrease in LVEF without significant ST changes (three times); and (4) ST depression without significant LVEF change (four times). Examples of each of these are shown in Figs 1–4. In this still small series, patterns (1) to (3) corresponded to patients with documented coronary artery disease, which was not the case for pattern (4). For detection of silent ischaemia at rest, a decrease in LVEF of > 5% lasting for > 1 min was defined as ischaemic left ventricular dysfunction. Using this definition four spontaneous episodes of silent left ventricular dysfunction could be demonstrated at rest in two of three coronary care unit patients with unstable angina during 160–680 min of data recording without simultaneous ST changes.

\[ \begin{align*}
\text{Min} & \quad 0 & 10 & 20 & 30 & 40 & 50 & 60 & 70 & 80 & 90 \\
(\text{ST level (mV)}) & \quad 0 & -1 & -2 & -3 & -4 \\
(-\text{EDV(%)}} & \quad 40 & 80 & 120 & 160 \\
(\text{LVEF(%)}} & \quad 20 & 40 & 60 & 80 & 100 \\
(\cdots) \text{Heart rate (beats min}^{-1}\) & \quad 60 & 80 & 100 & 120 & 160 \\
(\cdots) \text{ST level (mV)} & \quad 0 & -1 & -2 & -3 & -4 \\
\end{align*}\]

\^{}Figure 2\^{} Silent ischaemia during exercise (T.S., 57 years, two-vessel CAD). Note decrease in LVEF only during second level of exercise after significant ST depression. Note again the overshoot increase in LVEF after exercise.
changes. An example is shown in Fig. 5. This patient had anginal chest pain while walking from the gamma camera to his bed, where pain even increased without significant ST depression at this point in time but with marked depression of LVEF. Furtheron, he had two episodes of prolonged (6 min and 20 min) depression of LVEF of > 5% without chest pain or ST depression.

Discussion

Demonstration of ischaemic left ventricular dysfunction in the absence of chest pain can provide important confirmation of silent myocardial ischaemia in patients with asymptomatic ST segment depression. This study demonstrates that using the new portable VEST system, transient changes in radio-nuclide LVEF can be determined reproducibly. Based on the initial experience of our laboratory, four different patterns of LVEF and ST segment changes could be identified and it was shown that even spontaneous episodes of silent ischaemic left ventricular dysfunction can be registered.

On the basis of experimental studies, characteristic changes of ischaemic events have been identified. If myocardial perfusion becomes insufficient, changes in metabolism are first detected, followed by diastolic and then systolic functional impairment; electrocardiographic changes occur only at a later phase, but still before appearance of chest pain. This chain of events may explain observed patterns (1) and (3) in this study where functional ischaemic changes appeared before or without significant ST segment shifts. For pattern (2), however, other factors have to be discussed: with a nuclear probe as used in the VEST system, only changes in global LVEF may be detected. In a previous study we could show that hypoperfusion even of a large area as the anterior wall does not have to induce a significant 5% fall in LVEF if the other parts of the myocardium are normally perfused and behaving. Since ischaemic heart disease is a regional
Figure 4  Silent ischaemia during exercise? (H.R., 52 years, normal coronaries). Note there is no significant change in LVEF during exercise despite a >1 mm ST depression. Note also again the marked overshoot increase in LVEF after exercise.

Figure 5  Silent ischaemia at rest (A.H., 70 years, unstable angina, three-vessel CAD). Note symptomatic and prolonged silent ischaemic episodes as detected by long-term LVEF monitoring (cf. text).
disease of the myocardium, there may be non-uniformity of contraction which may explain delayed impairment of global function as observed in pattern (2). Furthermore, this may or may not be detected by electrocardiographic changes depending on the leads registered. Finally, pattern (4), excludes ischaemia with some degree of certainty only if there is a normal increase in LVEF of >5% during exercise as in diagnostic radionuclide studies. These considerations lead to the conclusion that there are some limitations inherent to radionuclide probe measurements as presented here: (1) only global function and relative haemodynamic changes can be observed and (2) functional changes of the apical area of the left ventricle may be weighted more than basal changes due to the detector position. In addition, probe displacement during ambulatory recording remains an important problem.

On the other hand, Kayden and co-workers have recently demonstrated that transient ischaemic left ventricular dysfunction detected by the VEST system is clinically relevant\(^1\). In a prospective study, they reported that this finding was more sensitive than symptoms or electrocardiographic exercise testing to identify patients at risk for future ischaemic events after acute myocardial infarction and thrombolysis. On the basis of their and our initial experience, we conclude that VEST is a reproducible method of detecting transient global left ventricular dysfunction which will be useful to confirm silent ischaemia in otherwise uncertain ST segment changes and may have important prognostic power.

**References**


