Operator independent left ventricular function monitoring during pharmacological stress echo with the new peak transcutaneous acceleration signal

T Bombardini, E Marcelli, E Picano, B Borghi, P Fedriga, B Garberoglio, G Gaggini, G Plicchi

Abstract
Background—As the myocardium contracts isometrically, it generates vibrations that can be measured with an accelerometer. The vibration peak, peak endocardial acceleration (PEA), is an index of contractility.

Objective—To evaluate the feasibility of PEA measured by the cutaneous precordial application of the accelerometer sensor; and to assess the usefulness of PEA monitoring during pharmacological stress echocardiography.

Design—Feasibility study.

Setting—Stress echo laboratory.

Patients—34 consecutive patients underwent pharmacological stress (26 with dipyridamole; 8 with dobutamine) and PEA monitoring simultaneously.

Interventions—A microaccelerometer was positioned in the precordial region and PEA was recorded. Dipyridamole was infused up to 0.84 mg/kg in 10 minutes, and dobutamine up to 40 µg/kg/min in 15 minutes.

Results—A consistent PEA signal was obtained in all patients. Overall mean (SD) baseline PEA was 0.26 (0.13) g (g = 9.8 m/s²), increasing to 0.5 (0.36) g at peak stress (+0.24 g, 95% confidence interval (CI) 0.14 to 0.34; p < 0.01). PEA increased from 0.26 (0.16) to 0.37 (0.25) g in the dipyridamole group (+0.11 g, 95% CI 0.08 to 0.16; p < 0.01), and from 0.29 (0.1) to 0.93 (0.37) g in the dobutamine group (+0.64 g, 95% CI 0.37 to 0.91; p < 0.01).

Conclusions—Using precordial leads this method offers potential for diagnostic application in the short term monitoring of myocardial function. PEA monitoring is feasible during pharmacological stress and documents left ventricular inotropic response quantitatively in a non-invasive and operator independent fashion.

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Keywords: ventricular function; contractility; peak endocardial acceleration; stress echo

The contractile state of the heart can theoretically be defined by the maximum velocity of shortening of unloaded myocardial contractile elements. Changes in the maximum rate of rise of ventricular pressure are highly sensitive to acute changes in contractility. In an attempt to develop methods of monitoring long term changes in function of the human heart, implantable systems using pressure transducers—which detect changes in right ventricular dP/dt—have been developed. The long term use of pressure sensing devices has various technical limitations, and the long term stability of such sensors remains to be demonstrated. In an attempt to overcome these limitations, alternative approaches have been developed for direct measurements of myocardial vibrations during isovolumic systole. When the myocardium contracts isometrically, it generates vibrations which have audible components that are responsible for the first heart sound. Both the audible and the inaudible spectrum of these vibrations can be measured with an accelerometer. Recent studies have shown that the peak of these myocardial vibrations (PEA, peak endocardial acceleration)—occurring in the isovolumic contraction phase—is an index of myocardial contractility and that its directional changes mirror changes in left ventricular peak dP/dt very closely.

Our aims in this study were: first, to evaluate the feasibility of the PEA measurements by cutaneous precordial application of the accelerometer sensor; and second, to assess the usefulness of PEA monitoring during pharmacological stress echocardiography.

Methods
We used a micromass uniaxial acceleration sensor connected to a standard ECG monitoring electrode with foam tape and solid gel. The ceramic piezoelectric accelerometer has a frequency response up to 1 kHz and a sensitivity of 18 mV/g (g = 9.8 m/s²) (fig 1).

The monitoring electrode was temporarily positioned in the mid-sternal precordial region before starting the scheduled stress test. The accelerometer was connected to an external signal amplifier with a frequency range of 0.05–1000 Hz. An analogue peak to peak detector synchronised with the standard ECG scanned the first 100 ms following the R wave (fig 2). All the data were collected on a seven channel TEAC FM magnetic tape recorder with a frequency response of dc to 1 kHz; analogue data were then digitised using a 4 kHz...
sampling rate with 12 bit resolution and stored and processed by an Intel/80386 based computer (Santa Clara, California, USA).

All the patients who participated in the study gave informed consent.

Thirty four patients consecutively referred to our stress echo laboratory simultaneously underwent a pharmacological stress echocardiogram (with dipyridamole in 26 patients and dobutamine in eight) and PEA monitoring (with a cutaneous precordial lead). According to standard protocols dipyridamole was infused up to 0.84 mg/kg in 10 minutes, and dobutamine up to 40 µg/kg/min in 15 minutes.91 0

The PEA signal was continuously recorded on a magnetic tape and the peak of the acceleration signal (apex to nadir) occurring in the pre-ejection systolic period was automatically measured at baseline, during the test, and at peak stress.

The wall motion score index (WMSI) was calculated in each patient at baseline and peak stress, according to the recommendations of the American Society of Echocardiography, from 1 = normal to 4 = dyskinetic in a 16 segment model of the left ventricle.11

**STATISTICS**

Data are expressed as mean (SD). Intrigroup comparisons were performed using the paired Student t test. Intergroup comparisons were performed using the non-parametric Wilcoxon test for intragroup comparisons and the Mann–Whitney test for intergroup comparisons were performed to confirm significance. Relations between variables were assessed using linear regression analysis and Pearson’s correlation coefficient. A probability value of p < 0.05 was considered significant.

**Results**

**STRESS ECHOCARDIOGRAPHY DATA**

Of the 26 patients of the dipyridamole group, 15 had normal resting ventricular function (WMSI = 1) and 11 had regional left ventricular motion abnormality (WMSI = 1.5 (0.34)). Of the eight patients of the dobutamine group, three had normal resting ventricular function (WMSI = 1) and five had regional left ventricular motion abnormality (WMSI = 1.9 (0.33)). Heart rate increased from 67 (14) to 87 (13) beats/min after dipyridamole (+20 beats/min, 95% confidence interval (CI) 12 to 28 beats/min; p < 0.01), and from 73 (16) to 126 (30) beats/min after dobutamine (+53 beats/min, 95% CI 34 to 72 beats/min; p < 0.01).

**PEA DATA**

A consistent PEA signal was obtained in all patients (fig 2). In the patients as a whole, mean (SD) baseline PEA was 0.26 (0.15) g, increasing to 0.50 (0.36) g at peak stress (+0.24 g, 95% CI 0.14 to 0.34 g; p < 0.01). PEA increased from 0.26 (0.16) g to 0.37 (0.25) g in the dipyridamole group (+0.11 g, 95% CI 0.08 to 0.16 g; p < 0.01), and from 0.29 (0.1) to 0.93 (0.37) g in the dobutamine group.
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+0.64 g, 95% CI 0.37 to 0.91 g; p < 0.01). Mean PEA percentage increase was +46 (38)% in the 26 dipyridamole patients and +235 (107)% in the eight dobutamine patients (p < 0.01 between groups; 95% CI 100% to 279%) (fig 3).

In the dipyridamole group the PEA increase was similar in the 15 patients with normal resting left ventricular function and in the 11 patients with a regional left ventricular wall motion abnormality at rest (+45 (36)% v +48 (43%), NS; 95% CI −35% to +29%).

In the dobutamine group the PEA increase was greater in the three patients with resting normal left ventricular function than in the five patients with a regional left ventricular wall motion abnormality (+338 (73)% v +174 (68%), p < 0.05; 95% CI 39% to 288%).

In the group of patients as a whole, the PEA percentage increase from baseline to peak stress was significantly related to the percentage increase in heart rate (r = 0.7, p < 0.05).

A typical PEA trend during dipyridamole and dobutamine stress is shown in fig 4.


discussion

THE PEA SIGNAL
A stable, reproducible, and consistent PEA signal was obtained in all patients. The signal to noise ratio was high, and little beat to beat variation was observed. Myocardial vibrations show several peculiar time frequency components. An analogue peak to peak detector synchronised with the ECG R wave scanned an appropriate time interval containing the isovolumic contraction phase, and we labelled the maximum value detected during this window PEA. This high amplitude vibration is an expression of the tension wavefront produced during initial activation of the heart. It occurs at the onset of endocardial movement, an average of 20 ms before mitral valve closure, and was fairly consistent among patients.2–24

PROPOSITION OF THE SIGNAL
Baseline PEA value had an ample range (from 0.1 to 0.7 g), unrelated to age or left ventricular function. Cardiac vibrations propagate as mechanical shear waves, and the intervening viscoelastic thoracic tissue attenuates the higher frequencies and introduces a variable propagation delay. The absolute PEA value in the single patient can be related more to the transthoracic propagation of cardiac vibrations than to left ventricular function. In fact, when measured epicardially or endocardially, pre-ejection cardiac vibrations are up to 10 times more powerful than when measured on the chest.15–18 In a multicentre PEA feasibility study, mean (SD) basal endocardial PEA was 0.45 (0.25) g—more powerful than our precordial PEA values (0.27 (0.14) g).20 21

PEA SIGNAL AND INOTROPIC CHANGES
Pharmacological inotropic stimulation increased the strength of the vibration signal, in keeping with the experimental studies.9 It is known that dobutamine increases heart rate and ventricular contractility. Its major characteristic is that it exerts a potent inotropic effect. The inotropic response is co-mediated by β1 and α adrenoreceptors, the latter causing an inotropic component that is independent of any chronotropic effect.22 23

Dipyridamole has the well known coronary vasodilator effects mediated by the inhibition of adenosine cellular transport, eventually leading to extracellular adenosine accumulation and steal phenomena. There is a mild catecholamine release that is responsible for the inotropic effect of the drug.9 These data are consistent with our findings, where there was a more pronounced and prolonged increase in the PEA signal in the dobutamine group than in the dipyridamole group.

CHOICE OF PHARMACOLOGICAL ECHO STRESS TO TEST THE PEA SIGNAL
For testing the ability of transthoracic PEA to measure cardiac inotropic changes, a cutaneous sensor is the preferred type as it is very sensitive to vibrations from the myocardium. However, it is also sensitive to skeletal muscle vibrations, so during physiological upright bicycle or treadmill testing a considerable amount of noise occurs.

The relative advantages and disadvantages of transthoracic versus endocardial PEA are now clear. Transthoracic PEA is a non-invasive measurement for short term monitoring in resting patients; endocardial PEA is an invasive measurement for long term monitoring with an implantable device, to measure the signal in daily life, in moving or exercising patients, so as to follow the success or adverse effects of chronically administered cardioactive drugs.20 21
The clinical applicability of the method using precordial leads offers potential for diagnostic applications in the short term monitoring of myocardial function, to measure spontaneously occurring changes in heart function in patients in bed. It also represents a physiological, operator independent tool for assessing directional global contractility changes during pharmacological stress echo and for post-test inotropic recovery.

CONCLUSIONS

PEA monitoring is feasible during pharmacological stress and documents the left ventricular inotropic response quantitatively in a totally automatic, non-invasive, and operator independent fashion. The response is shown to be milder with dipyridamole and more substantial with dobutamine.

The data obtained in this single centre feasibility study suggest that transcutaneous PEA is a good signal for assessing intraindividual changes in myocardial contractility. To establish fully the advantages and limits of this new method, comparisons with left ventricular pressure-volume loops in humans and multicentre study data are needed.

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