Multi-lead electrocardiogram in relation to serum enzymes in acute myocardial infarction

Andrzej Krotkiewski, Janina Gajewska-Lipka, Jacek Szelementko, and Jaroslaw Ruszkowski
From Department 1 of Medicine, Central Clinical Hospital, Warsaw, Poland

This study was made in an attempt to analyse the ST segment deflection on the map of a multi-lead chest electrocardiogram, and the changes in that area in the course of time. The deflection was regarded as an indication to the extent of the infarction. The sum of ST elevation (Σ ST) was considered as a measure of the extent of tissue damage. A relation was sought between these parameters and the activity of certain serum enzymes.

The prognosis of myocardial infarction depends to some extent on its site and size, both of which, however, are sometimes difficult to assess.

Maroko et al. (1972) determined the extent of the ischaemic lesion caused by experimental ligation of the coronary artery in dogs by recording the electrocardiogram from several points of epicardium. These recordings were subsequently compared to the multi-lead electrocardiograms taken from the surface of the chest and a reasonable similarity was found as regards the size of the lesion.

Kjekshus, Maroko, and Sobel (1972) performed similar investigations in order to elucidate the relation between changes in the ST segment in epicardial recordings and the cell injury as reflected by creatinine phosphokinase (CPK) depletion in different layers of the muscle of the left ventricle. Regions of acute ischaemia were delineated with the use of epicardial electrocardiograms from multiple sites, 15 minutes after coronary occlusion. The authors found a relation between ST elevation and in CPK depletion.

The same authors also performed human studies assessing the size of the ischaemic area of the heart muscle by multi-lead electrocardiograms. It has been proved that the area on the chest surface where raised ST segments are recorded is in some way a projection of the ischaemic lesion, thus reflecting its size: provided that the chest electrocardiogram is a multi-lead one. Body surface isopotential maps of the QRS have been reported previously by Spach et al. (1966) and Tazawa and Yoshimoto (1969) in children. The same was true for the R-ST segment (Rakita et al., 1954; Sayen et al., 1958; Braunwald et al., 1969). The surface mapping of the RS-T segment was performed by Reid, Pelides, and Shillingford (1971), and Taccardi (1963) in normal adults.

Apart from these experimental investigations we have been unable to find any human studies on the precise comparison of the size of the area depicted by ST elevation (or the sum of ST elevations: Σ ST) with the serum enzyme changes in patients with acute myocardial infarction.

The present study is an attempt to analyse such a relation and its behaviour in time.

Subjects and methods

Twenty men aged 41–72 years admitted to the Intensive Care Unit with a clinical diagnosis of acute myocardial infarction were studied. The patients studied were those who were admitted to our hospital within the first 72 hours after the beginning of anginal pain, usually within 10 to 15 hours.

Electrocardiograms were recorded in the supine position with Mingograph 81 apparatus at 50 mm/sec recording speed. The electrocardiographic examination consisted of a routine 12-lead electrocardiogram followed by multi-lead recordings from the chest. These recordings were taken on admission and repeated every day for a week and then once a week for 5 or 6 weeks.

Multi-lead recordings were taken from 72 electrodes (Wilson type) arranged in 8 transverse rows, 9 electrodes in each row, from the 2nd to the 6th intercostal. The transverse rows were designated as A, B, C...H, A from above down. The longitudinal columns were arranged with the figures 1 to 9, starting from the right mid-clavicular line and proceeding to the left mid-axillary line (Reid et al., 1971). The position of each electrode was permanently marked on the skin to make sure...
that the arrangement of electrodes on repeated recordings would be exactly the same. The distances between individual electrodes in every particular patient were the same in both directions, but depended on the size of the chest.

In the multi-lead electrocardiogram the elevation of the ST segment was analysed. The deflection of the ST segment from the isoelectric line was measured exactly 0.06 sec after the S wave. Segments TP or PQ were considered as the isoelectric reference. The values so obtained were plotted on a blueprint of the electrocardiogram. Elevation of ST was measured in millimetres (1 mm = 0.1 mV), and the sum of ST elevations (Σ ST) was considered as a measure of the extent of tissue damage. The number of sites where the ST elevation was more than 0.5 mm represented the size of the ischaemic area. The points at which the ST deflection was the same were connected by a contour line. Such maps depicting the areas of similar deflection of the ST segment have been designated isodeflectograms.

In all patients determinations were made of GOT, GPT, CPK, and LDH, parallel to the electrocardiographic examinations.

Blood was sampled for initial enzyme determination, 2 to 16 hours after admission and subsequently at 24-hour intervals, on the 2nd, 3rd, 4th, 5th, and 6th day. Then the enzyme determinations were repeated at weekly intervals for 5 to 6 weeks. Enzyme determinations were made in parallel with electrocardiogram multi-lead recordings. SGOT, SOPT, and LDH were estimated using the Behringer-Mannheim reagent kits (Reitman and Frankel, 1957; Wróblewski and LaDue, 1955), and the results were expressed in international units (mU/ml).

CPK was assessed according to the method of Hacker, Krüger and Augustin (1967). The same scheme of blood sampling intervals and of electrocardiographs was strictly followed in all patients, the only exception being the initial examinations which differed to some extent regarding the time elapsing from the beginning of the attack.

**Results**

In 20 patients 193 multi-lead electrocardiograms were recorded and evaluated by means of isodeflectogram curves.

The anterior wall of the heart muscle was involved in 10 patients, posterolateral in 5, posterior in 4, and 1 patient had multifocal infarction.

In all 10 patients with infarction of the anterior wall a similar electrocardiographic pattern was seen. The ST elevation ranged from 0.5 to 10 mm. The sum of ST (Σ ST) elevations in 6 patients conspicuously diminished as soon as the day after admission, e.g. from 239 to 123 mm. In 4 patients in whom imminent infarction was diagnosed the sum of ST elevations was initially little increased with infarction. In all 10 patients the sum of ST elevations gradually decreased in the 5th or 6th week: ST elevation, less than 1 mm, was seen only in few recordings.

Drawings were made representing, graphically, the results of multi-lead electrocardiograms. In infarction of the anterior wall the contour line connecting the points with the highest elevation of the ST segment was surrounded by contours representing ST elevations of lower and lower grade. On repeated examinations following the clinical course, the original picture changed usually by stepwise diminution and then disappearance of the central area of highest ST elevation. A good correlation with decreasing activities of serum enzymes was found.

It has to be emphasized that the enlarging of the infarcted area was accompanied by spreading of isodeflectographic contours and by increasing the sum of ST elevation (Σ ST). There was a very good correlation between the results of multi-lead electrocardiography and the clinical picture and serum enzyme activity, whereas the routine 12-lead electrocardiogram revealed only inconspicuous changes or none.

Fig. 1–4 illustrate such changes. For reasons of clarity only two enzymes (LDH and GOT) were plotted.

**Patient S.K., aged 42 (Fig. 1a).** Infarction of the anterior wall. Ventricular fibrillation, clinical death, effectively resuscitated. Fig. 1a presents the isodeflectograms made on the 1st, 28th, and 125th day of the infarction. Differently marked curves represent the isodeflectographic contours. On the first day the area of ST elevation was the largest as compared to the following examinations, as was the area of maximal ST elevation (10 mm) covering a great central part of the picture. The isodeflectographic contours shrank on the 28th and 125th day. On the 125th day no elevations higher than 3 mm were found.

Fig. 1b refers to the same patient. The sum of ST elevations is compared with the activity of serum GOT and LDH. The sum of ST elevations is drawn as a continuous line, SGOT as a dotted line, and LDH as a dashed line. Normal values of SGOT are 12 IU/ml, and for LDH 195 IU/ml.

On the first day after infarction the sum of ST elevations was 239 mm, SGOT 50 IU/ml, and LDH 100 IU/ml. The highest SGOT activity occurred on the second day, 93 IU, the highest LDH activity on the fourth day, 299 IU/ml. At the same time the sum of ST elevations showed a decrease and then all three values decreased gradually.

**Patient K.W., aged 56 (Fig. 2a).** Infarction of the anterior wall. Isodeflectograms made on the 1st, 4th, and 30th day of the attack show an enlargement of the particular isodeflectographic contours and the values of ST elevation. This correlates well with the clinical course: recurrence of severe retrosternal pain and a rise of serum enzymes activity. On the 30th day there was a significant diminution of the abnormalities.
**Multi-lead electrocardiogram in relation to serum enzymes**

**Fig. I (a)** Anterior myocardial infarction (S.K. aged 42). Isodeflectograms on the 1st, 28th, and 125th day of the disease. Persistent ST elevation. (b) The same patient. The sum of ST elevations (continuous line) compared with the SGOT (dotted line) and SLDH (dashed line) activity.

Fig. 2b shows the curve of the ST elevation sum together with the curves of SGOT and SLDH. Here again the rise of three parameters is seen on the fourth day with subsequent gradual lowering.

Patient K.R., aged 44, anterior myocardial infarction. Fig. 3 shows isodeflectograms made on the 1st, 7th, and 29th day. A conspicuous decrease is seen in the area and absolute values of ST elevation. The last picture shows only a minute central spot of ST elevation not exceeding 1 mm.

Patient S.A., aged 48 (Fig. 4a). Posterolateral infarction. The dominant change in this patient with posterior myocardial infarction was ST segment depression. The enzyme changes however were evident and parallel to the depression of the ST segment, as they were parallel to the ST elevation in the other cases. The isodeflectogram presented in Fig. 4a shows the area of ST depression in the left upper part of the map. The same examination on the 7th day shows spreading of the isodeflectographic contours and an increase of the values for ST depression. In the clinical course there was a recurrence of pain, and an increase of serum enzyme activity. On the 35th day all the parameters mentioned diminished conspicuously.

Fig. 4b refers to the same patient. Comparison of the sum of the curves of ST depression (at the bottom), of SGOT and SLDH shows a distinct second rise of parameters followed by a gradual lowering.

In the remaining patients with other than anterior localization of the infarction, the changes on the isodeflectogram were less clear. Depression as well as elevation of ST was found at various points studied. The deflection of the ST segment from the isoelectric line was slight. This was especially so in relation to infarction of the posterior wall of the heart.

**Discussion**

ST elevation in the routine 12-lead electrocardiograms strongly suggests myocardial infarction. The subsequent evolution confirms the diagnosis. According to Maroko et al. (1971), the degree of ST elevation may reflect the size of damage due to myocardial ischaemia. We recorded our multi-lead electrocardiogram at 72 points on the chest surface (Reid et al., 1971) and analysed the changes in the level of ST. By this method we were able to present schematically the areas of ischaemia of various degree reflected by ST deviation.

The parallelity of the enzymatic changes with the sum of ST elevation suggests that the latter reflects the size and severity of ischaemic damage to the heart muscle in myocardial infarction. We hope to
have proved it by comparing the clinical and biochemical data with the sum of ST elevations as shown in Fig. 1-4. The earlier experiments of Kjekshus et al. (1972) seem to justify our assumption.

We believe that multi-lead electrocardiography analysis is a valuable, noninvasive method for determining and evaluating ischaemic heart damage. This method may perhaps enable the study of the effect of drugs on the size and severity of the infarction (Maroko et al., 1971). Pelides et al. (1972) used the surface mapping of the ST segment to evaluate

![Diagram](Fig. 2) (a) Anterior myocardial infarction (K.W. aged 56). Isodeflectograms on the 1st, 4th, and 30th day. On the 4th day enlargement of the ST elevation area and significant diminution on the 30th day. (b) The same patient. Comparison of the $\sum$ ST with the SGOT and SLDH activity (Legend as Fig. 1b). On the 4th day the rise of three parameters with subsequent gradual lowering.

![Diagram](Fig. 3) Anterior myocardial infarction (K.R. aged 44). Isodeflectograms on the 1st, 7th, and 29th day. A conspicuous decrease of the area and absolute value of ST elevation. On the 29th day only a very small area of ST elevation.
the effect of practolol on the size of the ischaemic area in myocardial infarction; observing after practolol administration the decrease in the extent and height of the ST segment elevation they suggested that it could be used as an index of the extent and degree of ischaemia. The standard electrocardiogram may be normal in acute myocardial infarction (Johnson et al., 1959) and there may be a delay in appearance of the electrocardiographic features (Short 1968, 1970). Probably the multi-lead electrocardiogram will be of value in diagnosing cases of myocardial infarction with a normal standard electrocardiogram. It has to be stressed that there are certain limitations in the method at present, because satisfactory results are obtained only when analysing anterior myocardial infarction. The problem of posterior localization needs further study.

It is hoped that further investigation of multi-lead electrocardiography with isodeflectographic analysis will develop better and simpler techniques of examination and be of assistance to the physician in evaluating the course of myocardial infarction.

Conclusions

1) Multi-lead electrocardiograms recorded from electrodes arranged on the anterior surface of the chest provide a graphic image of the extent and severity of ischaemic damage to the heart muscle.

2) The results of our studies, especially those relating to anterior myocardial infarction, are in good agreement with the clinical picture and biochemical parameters, and indicate greater sensitivity of the multi-lead electrocardiogram as compared with the routine 12-lead one.

3) The sum of ST elevations in particular leads measured in mm may also be a reasonable index of the size and severity of infarction.

4) The present method may be applied in clinical studies on the effect of drugs on the size and severity of ischaemic damage.

5) The results obtained in cases of posterior myo-
cardiac infarction cannot reliably be interpreted with the present arrangement of electrodes.

References

Requests for reprints to Dr. Andrzej Krotkiewski, Department I of Medicine, Central Clinical Hospital, 02-507 Warsaw, Komarowa 137, Poland.
Multi-lead electrocardiogram in relation to serum enzymes in acute myocardial infarction.

A Krotkiewski, J Gajewska-Lipka, J Szelemeiko and J Ruszkowski

Br Heart J 1973 35: 991-996
doi: 10.1136/hrt.35.10.991

Updated information and services can be found at:
http://heart.bmj.com/content/35/10/991.citation

These include:

Email alerting service

Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/