Can the surface electrocardiogram be used to predict myocardial viability?

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Abstract

Objective—To investigate whether QRS morphology on the surface ECG can be used to predict myocardial viability.

Design—ECGs of 58 patients with left ventricular impairment undergoing positron emission tomography (PET) were studied. 13N-Ammonia (NH3) and 18F-fluorodeoxyglucose (FDG) were the perfusion and the metabolic markers, respectively. The myocardium is scarred when the uptake of both markers is reduced (matched defect). Reduced NH3 uptake with persistent FDG uptake (mismatched defect) represents hibernating myocardium. First, the relation between pathological Q waves and myocardial scarring was investigated. Second, the significance of QR and QS complexes in predicting hibernating myocardium was determined.

Results—As a marker of matched PET defects, Q waves were specific (79%) but not sensitive (41%), with a 77% positive predictive accuracy and a poor (43%) negative predictive accuracy. The mean size of the matched PET defect associated with Q waves was 20% of the left ventricle. This was not significantly different from the size of the matched PET defects associated with no Q waves (18%). Among the regions associated with Q waves on the ECG, there were 16 regions with QR pattern (group A) and 23 regions with QS pattern (group B). The incidence of mismatched PET defects was 19% of group A and 30% of group B (NS).

Conclusions—Q waves are specific but not sensitive markers of matched defects representing scarring myocardium. Q waves followed by R waves are not more likely to be associated with hibernating myocardium than QS complexes.

(J Heart 1999;82:663–667)

Keywords: electrocardiography; myocardial viability; positron emission tomography; myocardial scarring

The significance of pathological Q waves on the surface ECG is controversial. Some investigators believe that Q waves reflect scarred myocardium. Others state that the Q waves are dynamic rather than permanent changes after myocardial infarction. A third view suggests that evidence of viable myocardium can be found in many patients with pathological Q waves. Furthermore, it has been suggested that preservation of R waves after pathological Q waves is a marker for hibernating myocardium. This has not been tested against imaging techniques for myocardial viability.

In this study we investigated the use of the surface ECG in selecting patients for viability studies by evaluating the relation between QRS morphology and myocardial viability using positron emission tomography (PET).

Methods

Patients

The ECGs of 58 patients were studied (48 male, 10 female). These were an unselected group of patients with ischaemic heart disease and impaired left ventricular contraction who were undergoing PET. The index myocardial infarct occurred at a mean of 109 weeks (range six weeks to 13 years) before imaging. This was the most recent infarction in five patients (9%) who had multiple documented infarctions. Myocardial infarction was diagnosed retrospectively in 13 patients (22%), when they presented with angina or left ventricular impairment. The patients’ characteristics are summarised in table 1.

The Electrocardiograms

Surface ECGs were reviewed to assess the relation of pathological Q waves to myocardial scarring, and the predictability of myocardial hibernation by the maintenance of R waves after pathological Q waves. In addition, we assessed the impact of the size of the myocardial region with scarring or hibernating myocardium on the appearance of Q waves, or the preservation of R waves after Q waves, respectively.

Pathological Q waves were defined as wide (> 0.04 seconds) and deep (> 4 mm or > 25% of R wave).

Table 1 Demographic characteristics and risk factors

<table>
<thead>
<tr>
<th>Risk factor</th>
<th>Number of patients</th>
<th>%</th>
</tr>
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<tbody>
<tr>
<td>Hypertension</td>
<td>20/58 (34.5%)</td>
<td></td>
</tr>
<tr>
<td>Hypercholesterolaemia</td>
<td>50/58 (86.2%)</td>
<td></td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>16/58 (27.6%)</td>
<td></td>
</tr>
<tr>
<td>Family medical history</td>
<td>23/58 (39.7%)</td>
<td></td>
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<tr>
<td>History of smoking</td>
<td>43/58 (74.1%)</td>
<td></td>
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Accepted for publication

1 July 1999
of the height of the corresponding R wave, provided the R wave was > 5 mm), appearing in at least two contiguous leads.

The ECG leads were divided into two main regions:
1. The septal, anterior, and lateral walls region: leads I, aVL, and V1–6.
2. The inferior region: leads II, III, and aVF.

Lead aVR was excluded from analysis.

**Figure 1** Two polar maps representing the left ventricle. Each is divided into five regions. The numbers represent the following left ventricular regions: 1, the apex; 2, the septum; 3, the anterior wall; 4, the lateral wall; 5, the inferior wall. (A) The perfusion polar map. This shows an infarcted defect (arrowhead). (B) The metabolic polar map. This shows a matching metabolic defect in the inferolateral region (arrowhead). Therefore, the patient has a matched infarcted defect suggestive of myocardial scarring in that region. (The colour scale is representative of the radioactive activity in the region, with yellow being the highest count.)

**Figure 2** The importance of Q wave as a marker of myocardial scarring. The bars represent the patients with (group 1) and without (group 2) pathological Q waves on their surface ECG.

**POSTHON EMISION TOMOGRAPHY**

PET was carried out after obtaining informed written consent. The locally appointed ethics committee approved the protocol.

The methods of image acquisition and analysis have been described previously. In brief, using a Siemens Exact 31 scanner (Siemens, Knoxville, Tennessee, USA), 31 slices were produced through the left ventricle. The reconstructed image resolution was 10 mm. Attenuation correction was achieved by performing a transmission scan before the emission scan.

13N-Ammonia and 18F-fluorodeoxyglucose were used as the perfusion and metabolism markers, respectively. If-18F-Fluorodeoxyglucose was given after an oral glucose load, and insulin was given to achieve euglycaemia in diabetic patients.

Using the filtered back projection method, images were reconstructed to generate short axis sections of the myocardium. These sections were subjected to circumference profile analysis to produce polar maps representing the left ventricle (fig 1).

The level of highest 13N-ammonia uptake was considered to represent the reference perfusion level. The glucose uptake in the area with reference perfusion level was regarded as the reference 18F-fluorodeoxyglucose uptake for that patient. The perfusion and metabolism in the other areas were then compared with the perfusion and metabolism in the reference area.

Viable myocardium was defined by the presence of 18F-fluorodeoxyglucose uptake. The viable areas with reduced 13N-ammonia uptake and increased 18F-fluorodeoxyglucose uptake, compared with the reference region were labelled as mismatched defects. Mismatched defects are considered to represent hibernating myocardium (although the definitive diagnosis of hibernating myocardium relies on postoperative improvement in contraction in a previously impaired region).

Severe reduction (to a level less than 50% of the reference level) of both perfusion and metabolism activity defined a matched defect. Matched defects represent scarred myocardium. The size of a matched defect was estimated from the short axis sections of the left ventricular perfusion and metabolism images.

**STATISTICAL ANALYSIS**

Continuous variables are presented as mean (SD). Categorical data are presented as frequencies. The comparison between groups was performed using χ² tests for categorical data. Differences with a p value < 5% were considered to be statistically significant. Data were analysed using Microsoft Excel Release 4.0 and SPSS for Microsoft Windows Release 6.1 software packages.

**Results**

**Q WAVE AS A MARKER OF MYOCARDIAL SCARRING**

There were 116 electrocardiographic regions. These were divided into two groups according to the presence of pathological Q waves. Group 1 (n = 39) formed the regions with pathological Q waves, and group 2 (n = 77) the regions with no pathological Q waves. Matched PET defects, as evidence of scarring, were noted in 74 regions. Of these scarred regions, 30 were in group 1 and 44 in group 2 (fig 2). Therefore, the specificity of Q waves as markers of myocardial scarring was 79%. Their sensitivity, however, was low at 41%. In the cohort studied, the positive predictive value of Q waves for myocardial scarring was 77%, and the negative predictive value was 43%.

The effect of the extent of chest lead involvement by Q waves on the predictability of scarring was also assessed. The patients with pathological Q waves in these leads were divided into those with pathological Q waves in the leads V1–3 (septal), and those with pathological Q waves in the leads V1–4, 6 (extensive) (table 2). One patient with exclusively lateral lead involvement (V4–6) by pathological Q waves was excluded from this analysis. There was a greater chance that the patients with...
Table 2 The impact of the extent of lead involvement by Q waves on the predictability of myocardial scarring

<table>
<thead>
<tr>
<th>Septal Q waves</th>
<th>Extensive anterior Q waves</th>
</tr>
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<tbody>
<tr>
<td>Patient</td>
<td>Scar?</td>
</tr>
<tr>
<td>8</td>
<td>Yes</td>
</tr>
<tr>
<td>11</td>
<td>No</td>
</tr>
<tr>
<td>17</td>
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<td>27</td>
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<td>33</td>
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<td>30</td>
<td>Yes</td>
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<td>31</td>
<td>Yes</td>
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</table>

Patients with pathological Q waves in the chest leads, and the incidence of matched perfusion–metabolism defects (scars) in the anterior wall. The patients were divided into those with septal Q waves (V1–3) and those with extensive anterior Q waves (V1–4, 6).

extensive pathological Q waves had matched perfusion–metabolism defects than the patients with pathological Q waves in the septal leads alone (100% vs 67%). However, the difference did not reach statistical significance, the p value being just over 0.05.

To investigate the potential impact of posterior wall involvement on the absence of pathological Q waves, we studied the 31 patients with inferior scars. Among these, there were eight with posterior scars, four of whom had pathological Q waves in the inferior leads (table 3). When the cohort is considered, there were 44 scarred regions not associated with pathological Q waves. The absence of pathological Q waves in these regions with myocardial scarring cannot be ascribed to the posterior location of the scars except in four cases.

The mean area of the matched defects associated with pathological Q waves on the ECG was 20% of the left ventricular surface. This was not significantly different from the mean area of matched defects associated with electrocardiographic non-Q wave regions (18%).

R AFTER Q AS A MARKER FOR HIBERNATING MYOCARDIUM

To demonstrate any relationship between the presence of hibernating myocardium and the maintenance of R wave after pathological Q wave, we concentrated on the 39 regions with pathological Q waves. These were divided according to the preservation of R waves after Q waves, into two groups: group A (n = 16) comprised the regions with maintained R waves following the pathological Q waves; group B (n = 23) comprised the regions with QS complexes.

In group A, three regions (19%) showed metabolism–perfusion mismatch defects suggestive of hibernating myocardium. On the other hand, as many as seven regions (30%) of the group B regions were associated with evidence of hibernating myocardium on PET. The difference between the two groups was not statistically significant. Therefore, the presence of R waves following a pathological Q wave was not helpful for predicting the presence of hibernating myocardium (fig 3).

Discussion

We investigated the use of the QRS complex on the surface ECG in predicting myocardial viability. This was addressed through two clinical questions.

The first was whether pathological Q waves are useful markers of myocardial scarring. The second was whether preservation of R waves following pathological Q waves is a marker for hibernating myocardium.

We found that pathological Q waves have a high specificity and positive predictive accuracy for myocardial scarring. However, their absence does not exclude the presence of myocardial scarring, given their low sensitivity and low negative predictive accuracy. In addition, the appearance of pathological Q waves is not determined by the size of the matched defect.

In the second question, two groups of regions with Q waves were compared to test the role of preserved R waves as a marker of hibernating myocardium. We found that following pathological Q waves, R waves are not more likely to be associated with hibernating myocardium than QS complexes. Therefore hibernating myocardium is not more likely to be present if pathological Q waves are followed by R rather than S waves.

Detection of viable myocardium is important in the presence of impaired left ventricular contraction, where revascularisation can lead to functional improvement. This can be best predicted using PET.
However, viability studies are expensive, hence the need to find ways of selecting patients to undergo these studies, to limit expenditure. The selection process cannot rely on factors such as the frequency of angina, the presence of congestive heart failure, the extent of left ventricular impairment or the presence of serious arrhythmia. These variables do not reliably differentiate between patients with and without residual myocardial viability. 

It has been proposed that the preservation of R waves, following pathological Q waves, may be a marker for hibernating myocardium, but this has not been tested against imaging techniques for myocardial viability. As for pathological Q waves, their relation to myocardial viability has been controversial.

Those who believe that pathological Q waves represent myocardial scarring cite evidence of increased myocardial necrosis (increased creatine kinase levels), myocardial impairment, and a worse prognosis in association with Q wave myocardial infarction. 

Those who doubt the relation between Q waves and myocardial scarring, however, rely on the presence of myocardial viability in areas corresponding to the electrocardiographic regions with Q waves, and the presence of only mild to moderate degree of fibrosis in up to 20% of biopsies of the akinetic or dyskinetic anterior myocardial segments with Q waves on the ECG. 

One of the problems of investigating the role of the QRS complex in the prediction of myocardial viability is the unreliability of the ECG in localising pathological processes to specific myocardial regions. In our investigation, we avoided the detailed division of the rather complicated septal, anterior, and lateral regions of the left ventricular myocardium. The left ventricular myocardium was therefore divided into two major regions, anterior and inferior. However, we did look at whether extensive pathological Q waves in the chest leads (V1–4, 6) are more likely to predict scarring than the limited appearance of these waves in the septal leads (V1–3). There was no statistically significant difference in the incidence of scarring between the two subgroups. However, a potential confounding factor is the small number of patients in each subgroup (table 2).

In the setting of posterior myocardial infarction, overt Q waves may not develop on the classical surface ECG. The patient may have an increased R to S ratio anteriorly, and particularly when the infarct related artery is the left circumflex artery these patients may have diminution of the R wave amplitude in the lateral leads (I, aVL, and V5–6). However, this problem was encountered in four of the 44 regions with scars and no corresponding pathological Q waves.

A third potential confounding factor in the investigation of myocardial viability is the use of ECGs recorded in the acute or subacute phases of myocardial infarction. Q waves may be a temporary electrical expression of myocardial injury during these phases of myocardial infarction rather than a sign of permanent myocardial damage. Therefore, all ECGs studied were recorded at a minimum of six weeks after myocardial infarction, with the mean time from the index infarct to the ECG recording being 109 weeks (range six weeks to 13 years).

Some of the patients (22%) never had a clinically documented myocardial infarct. In these patients, the occurrence of infarction was evident from the demonstration of severe wall motion abnormality on non-invasive or invasive imaging, with or without an occluded infarct related artery on coronary angiography. In these patients, PET was carried out at least six weeks following their presentation, during which they neither experienced symptoms suggestive of myocardial infarction nor developed any new electrocardiographic evidence of infarction.

The present study extends the previous work of Brunken et al. In a small group of 20 patients, these investigators found that about 40% of regions with Q waves and severe hypokinesia (or worse function) on wall motion study will show metabolic activity on PET. In our study, a larger cohort of patients was studied, likely to be representative of a typical group of patients who will be considered for myocardial viability studies.

Ragosta et al used rest-redistribution planar thallium-201 to detect myocardial viability in 21 patients undergoing myocardial revascularization. There were 11 patients whose left ventricular contraction did not improve significantly after surgery (and therefore had scarred myocardium). These were significantly more likely to have Q waves on the ECG (91% of the patients in this group vs 40% in the remaining group, p = 0.02). Their finding suggests that Q waves are more likely to be associated with scarred than with hibernating myocardium. This is in keeping with our conclusion that Q waves have high specificity and positive predictive accuracy for myocardial scarring.

In conclusion, the presence of pathological Q waves does not exclude the presence of viable myocardium. However, it indicates a high probability of some scarring in the corresponding myocardium. The reverse is not true, as the negative predictive accuracy of Q waves as markers of scarring is poor. The presence of hibernating myocardium in areas with Q waves is not reflected by preservation of R waves. Further studies into simple clinical markers for myocardial viability are required.

IMAGES IN CARDIOLOGY

Left main coronary artery lesion detected by transoesophageal echocardiography before cardioversion

A 60 year old white man with history of hypertensive heart disease had complained of dyspnoea for a month. Newly developed atrial fibrillation caused symptomatic deterioration. Routine precardioversion transoesophageal echocardiography to rule out left atrial thrombus under oral anticoagulation had revealed a mosaic pattern in the left main coronary artery (LMCA) ostium. Colour Doppler imaging was performed to determine whether there was a significant left main coronary artery lesion. Coronary angiography showed ostial stenosis of the left main coronary artery without any other lesion in the left or right coronary artery system. We detected 35% diameter stenosis and 58% area stenosis in the LMCA ostium. Minimal lumen diameter and reference segment diameter were calculated to be 3.13 mm and 4.83 mm, respectively. Although the lumen diameter (> 3 mm) was sufficient for coronary artery perfusion, intravascular ultrasound imaging was performed, which showed a non-critical eccentric lesion with mixed plaque composition, causing 27% diameter stenosis and 46.5% area stenosis. The patient underwent cardioversion safely.

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