Effect of long term treatment with metoprolol and sotalol on ventricular repolarisation measured by use of transoesophageal atrial pacing

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SUMMARY The effects of long term (4 weeks) treatment with oral metoprolol (100 mg twice daily) and sotalol (160 mg twice daily) on ventricular repolarisation time were compared in a double blind crossover study in 20 patients post-infarction. For QT interval studies transoesophageal atrial pacing was performed at a cycle length of 800 ms. Sotalol prolonged the QT interval by 5–7% compared with metoprolol. The prolongation reflects a change in the repolarisation time because there was no change in the QS interval. Measurements of heart rate at rest and during bicycle exercise indicated that metoprolol and sotalol in the doses selected were equipotent as beta blockers. Transoesophageal atrial pacing is a simple non-invasive method with few and mild side effects that is well suited to drug studies.

According to the Vaughan Williams classification, beta adrenergic receptor antagonists have class II antiarrhythmic properties that are characterised by a reduction of the spontaneous phase 4 depolarisation of pacemaker cells. Some beta blockers, for example alprenolol and propranolol, also have a direct membrane stabilising (class I) effect that leads to a reduction in the maximum rate of rise of cardiac action potentials. This antiarrhythmic effect has been seen when high concentrations of such drugs are used in vitro, but it is believed to be of no clinical importance.

Metoprolol is a cardioselective beta blocker which has no intrinsic sympathomimetic activity and membrane stabilising properties. Sotalol is a non-cardioselective beta blocker that has no important membrane stabilising or intrinsic stimulatory activity. In contrast to other currently available beta blockers, sotalol has class III antiarrhythmic activity in that it increases the duration of the action potential after short term administration. Some beta blockers including metoprolol have caused similar effects but only after long term treatment and to a lesser degree. Although recent studies in man have suggested that sotalol and other currently available beta blockers may have different effects on ventricular repolarisation after long term treatment as far as we know no double blind study has been performed to confirm this. Craemer et al performed a randomised crossover study but this was not double blind.

We have compared the effects of long term treatment with metoprolol (class II) and sotalol (classes II and III) on ventricular repolarisation in a double blind crossover study in patients who were already on chronic beta blocker treatment after a previous myocardial infarction. We used doses of metoprolol (100 mg twice daily) and sotalol (160 mg twice daily) with equipotent beta blocking effects.

Patients and methods

We studied 20 men aged 46–64 years (mean 60 years) who had had a myocardial infarction more than one year before. They were on long term metoprolol treatment (minimum daily dose of 150 mg). None was on any other antiarrhythmic agent. We excluded patients with clinical signs of heart failure or severe angina (New York Heart Association functional class III–IV) and those with either partial or complete bundle branch block. To ensure measurable QT intervals we did not study patients with biphasic negative T waves or U waves in the surface lead V2. The
study protocol was approved by the ethical committee of the hospital, and each patient gave informed oral consent.

METHODS
In this double blind crossover study the patients remained on their usual long term treatment with metoprolol during the run-in period. They were randomly allocated to receive oral sotalol 160 mg twice daily (10 patients) or metoprolol 100 mg twice daily (10 patients) for four weeks (period I). After this they were switched to treatment with the other tablet for another four weeks (period II). Investigations were performed during the run-in period and after both periods of treatment.

ERGOMETER TEST
The patients underwent a bicycle ergometer test starting at a workload of 50 W which was increased by 10 W per minute. Heart rate, blood pressure, and a 12 lead electrocardiogram were monitored each minute. When the heart rate reached 70% of the predicted maximum pulse rate the exercise test was stopped. The effect of the beta blockade was evaluated and compared by measurement of heart rates at rest and during exercise.

TRANSOESOPHAGEAL ATRIAL PACING
With the patient in a semirecumbent position, that is lying at approximately 40° to the horizontal, a bipolar electrode catheter was advanced into the distal oesophagus. We used flexible silicone rubber coated bipolar electrode catheters with an interelectrode distance of 18 or 28 mm (Medtronic 6992). The distal electrode was designated as the negative (cathode) electrode. Unipolar atrial electrograms were recorded from the catheter (Fig. 1a). The electrode position was adjusted to obtain a maximum positive unipolar atrial deflection from the distal electrode and a biphasic atrial electrogram from the proximal electrode. The minimum threshold current required to pace the atria reliably was determined by a technique described elsewhere. A burning sensation causing slight epigastric discomfort was common during stimulation, but no sedative or other premedication was required. Transoesophageal bipolar atrial stimulation was undertaken for approximately three minutes at a basic cycle length of 800 ms (Fig. 1b). Atrial stimulation was performed with a programmable stimulator made in our own laboratory that delivers a constant current square-wave pulse. The pulse width was fixed at 10 ms. A precordial electrocardiogram was displayed on an oscilloscope and recorded on a Mingograph 82 (Siemens-Elema) as well as on a tape recorder (TEAC) during atrial stimulation.

BLOOD SAMPLING
Blood samples were collected immediately after the investigations for later analyses of plasma concen-

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**Fig. 1**  (a) A representative example of unipolar atrial electrograms recorded from the oesophageal electrode catheter. Paper speed 50 mm/s. (b) Recording of transoesophageal bipolar atrial pacing from the same patient as in (a). Atrial stimulation was performed at a basic cycle length of 800 ms. The arrow indicates the stimulus spike. Paper speed 25 mm/s.
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**QTc calculation**

QTc is usually calculated by the Bazett formula from the QT interval measured during sinus rhythm. In the present study the QTc was calculated from the QT intervals measured during transoesophageal atrial pacing at a basic cycle length of 800 ms.

**Statistical methods**

We used the two-tailed Student's t test to compare paired differences. The level of statistical significance was set at a p < 0.05, but all p values are given.

**Results**

Fifteen out of 20 patients were included in the QT interval analysis. The reasons for exclusion were as follows: one patient was excluded during run-in because he developed hypotension, sweating, and dizziness and slight bradycardia when transoesophageal atrial pacing was started. The symptoms and physical findings were observed shortly after pacing was started, and stimulation was discontinued before reliable atrial pacing could be obtained. The patient reported no pain. Blood pressure returned to normal after the patient was tilted in the head down position. In another patient an intolerable headache developed during blind period 1 (sotalol) and the medication was discontinued after 11 days. Three patients had biphasic negative T waves or prominent U waves in one or more recordings that made subsequent comparisons of QT intervals impossible. They completed the study but were excluded from QT interval analyses.

Thus atrial pacing was performed without side effects in 18 of 19 patients on three consecutive occasions. The means of the thresholds for adequate pacing were: 19 mA (run-in), 21 mA (metoprolol), and 22 mA (sotalol) (range 7–36 mA).

**Heart rate**

The heart rates at rest and during exercise (measured at a workload of 100 W) were not significantly different on the two treatments (Table). This indicates that a similar degree of beta blockade was induced by the two agents.

**QS intervals**

There was no significant change in the QS interval, which was within the normal range (Table) on both agents.

**QTT and QTc intervals**

Treatment with sotalol was associated with a significantly longer QTT interval than treatment with metoprolol (mean (SD)) 319 (18) ms vs 301 ms.

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**COMPUTERISED ANALYSIS OF THE QRST INTERVALS**

We used a specially developed interactive computer program to analyse the following variables: QS, QT top, QT end, JT top, and JT end (as defined in Fig. 2). All measurements were made from lead V2 (except for one patient where V3 was used because of lack of signal). The intervals were all measured at a fixed basic atrial pacing cycle length of 800 ms. A representative sample of each recording saved on tape was displayed on the oscilloscope. For the first beat chosen, the operator defined the following seven points: start of the pacemaker stimulus, start of the Q wave, top of the R wave, negative top of the S wave, end of the S wave, top of the T wave, and end of the T wave. For the subsequent analysed beats these points were outlined by the program but could be corrected if necessary. Only signals that were preceded by constant and adequate pacing were analysed. All the results are the means of at least five representative complexes. The output consisted of a table of each individual variable for all accepted beats as well as the mean values and standard deviations.

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**Fig. 2. The following intervals were analysed:** QS (from the beginning of the Q wave to the end of the S wave), QTT (from the beginning of the Q wave to the top of the T wave), and QTc (from the beginning of the Q wave to the end of the T wave). The JT interval was calculated by subtracting the QS interval from the QT interval. All intervals were measured at a paper speed of 100 mm/s.
Table  Comparisons of heart rate and QRST intervals (ms) during run-in and after 4 weeks' treatment with metoprolol or sotalol

<table>
<thead>
<tr>
<th>Results</th>
<th>Run-in</th>
<th>M</th>
<th>S</th>
<th>M vs run-in (p)</th>
<th>S vs run-in (p)</th>
<th>S vs run-in (% change)</th>
<th>S vs M (p)</th>
<th>S vs M (% change)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart rate at rest (bpm)</td>
<td>58 (16)</td>
<td>55 (8)</td>
<td>57 (10)</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>Heart rate at 100 W (bpm)</td>
<td>96 (20)</td>
<td>92 (11)</td>
<td>95 (14)</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>QT top</td>
<td>300 (11)</td>
<td>301 (16)</td>
<td>319 (18)</td>
<td>NS</td>
<td>&lt;0.001</td>
<td>+6</td>
<td>&lt;0.001</td>
<td>+6</td>
</tr>
<tr>
<td>QT end</td>
<td>421 (17)</td>
<td>414 (25)</td>
<td>436 (25)</td>
<td>NS</td>
<td>&lt;0.001</td>
<td>+4</td>
<td>&lt;0.001</td>
<td>+5</td>
</tr>
<tr>
<td>JT top</td>
<td>202 (16)</td>
<td>202 (19)</td>
<td>219 (23)</td>
<td>NS</td>
<td>&lt;0.001</td>
<td>+8</td>
<td>&lt;0.001</td>
<td>+8</td>
</tr>
<tr>
<td>JT end</td>
<td>324 (24)</td>
<td>316 (26)</td>
<td>336 (28)</td>
<td>NS</td>
<td>&lt;0.001</td>
<td>+4</td>
<td>&lt;0.001</td>
<td>+6</td>
</tr>
<tr>
<td>QTc top</td>
<td>330 (10)</td>
<td>340 (20)</td>
<td>360 (20)</td>
<td>NS</td>
<td>&lt;0.001</td>
<td>+9</td>
<td>&lt;0.001</td>
<td>+6</td>
</tr>
<tr>
<td>QTc end</td>
<td>470 (20)</td>
<td>460 (30)</td>
<td>490 (30)</td>
<td>NS</td>
<td>&lt;0.001</td>
<td>+4</td>
<td>&lt;0.001</td>
<td>+7</td>
</tr>
</tbody>
</table>

Run-in, long term treatment with metoprolol at a minimum daily dose of 150 mg; M, after four weeks' treatment with metoprolol (100 mg x 2); S, after four weeks' treatment with sotalol (160 mg x 2).

(16) ms, +6%; p < 0.001. Sotalol also prolonged the QTc interval (436 (25) ms vs 414 (25) ms, +5%, p < 0.001) (Table and Fig. 3).

**PLASMA CONCENTRATIONS**

The plasma concentration of metoprolol in 15 samples about five hours after one oral dose was 41–700 nmol/l (mean 265 nmol/l). The plasma concentration of sotalol in 14 samples was 0.8–2.0 μg/ml (mean 1.4 μg/ml). One sample was lost.

Discussion

The antiarrhythmic efficacy of beta blockers may be the result of their ability to oppose the effects of

![Electrocardiograms](http://heart.bmj.com)
catecholamines on automaticity and conductivity. Animal studies have shown that several beta blocking agents with different pharmacological properties produce a moderate prolongation of the ventricular repolarisation time when they have been given for four weeks. This contrasts with that lack of effect after short term administration. In healthy volunteers metoprolol prolonged the right ventricular repolarisation time by about 13-17% after intravenous administration of 100 mg, and this effect was maintained after twelve weeks of oral treatment. This is consistent with other reports demonstrating a prolongation of QT and JT intervals after short term administration of sotalol. The mechanism by which beta adrenergic blockers such as metoprolol prolong the ventricular repolarisation time after chronic treatment is not yet known. It has been proposed that this effect represents a myocardial adaptation to prolonged beta blockade.

In order to clarify whether long term treatment with a beta blocker without class III activity would prolong the ventricular repolarisation time to the same extent as an antiarrhythmic agent with combined class II and III activity, it seemed logical to use sotalol as a standard for comparison. The ventricular repolarisation time may be assessed from the QT intervals, ventricular effective refractory periods, and ventricular monophasic action potentials. Recent studies with beta adrenergic blockers have shown consistent changes in these indices after long term treatment.

Difficulties with defining the end of the T wave make measurement of the QT interval uncertain. To ensure more easily defined intervals we also included the QT top interval. The QT interval is frequency dependent and the most widely applied rate correction factor has been Bazett's formula. The formula was intended to correct the values to a heart rate of 60 beats per minute, but will give an over-correction of the QT interval at high rates and an undercorrection at low heart rates. When Bazett's formula is used for correction during sinus rhythm, slight or even moderate drug induced changes might therefore be masked, as they were in other drug studies. If, however, the heart rate is kept constant, the measurements will contain a systematic error but will allow drug induced changes to be detected. To circumvent the use of a correction factor some previous studies have compared QT intervals at identical and constant paced heart rates by means of intracardiac atrial stimulation.

To ensure the detection of even minor drug induced changes and to avoid an invasive intervention, we have compared QT intervals at a fixed paced heart rate produced by transoesophageal atrial stimulation. We performed transoesophageal atrial pacing on 56 different occasions and on only one (1.8%) occasion was there a complication, judged to be a vasovagal reaction.

The electrode catheter insertion depth is carefully controlled to minimise the pacing threshold and the appropriate depth may be predicted by the site of the maximum atrial electrogram or the patient's height. In the present study, the threshold for adequate atrial pacing was found in most patients when the atrial electrogram from the proximal electrode showed a biphasic deflection and the distal electrode a highly positive atrial deflection, and this accords with other studies. The mean minimum pacing threshold was of about the same magnitude for all three pacing sequences in the same patient when pacing was performed from identical positions.

The minimum current required for atrial pacing varies among different investigators. In one study, atrial capture was achieved in 75% of the attempts when a current of 17.5 mA was used. Others have reported an average minimum pacing threshold of 10 mA (range 4.5-20 mA). The minimal thresholds obtained in the present study were somewhat higher than those reported by other investigators, but no patient complained of intense discomfort. The currents used in this study were considerably below those reported to produce hyperaemia or epithelial erosion in canine experiments. Epithelial erosion appeared after pacing at 60 mA at a pulse width of 2 ms for 4 hours, but no injury was noted when the pacing time was limited to < 30 minutes.

We found that four weeks' treatment with sotalol significantly prolonged the QT intervals by 5-7% more than metoprolol (Table). The prolongation was confined to the JT interval, as there was no significant difference in the QS interval, and this must therefore reflect a change in the ventricular repolarisation time. This is consistent with results from other studies, although the groups of patients studied were not comparable. The QT top and the QT end intervals showed consistent changes, which suggests that the former may be used with advantage, especially when the end of the T wave is obscured. The QTc showed similar and statistically significant changes when calculated from a fixed paced heart rate. The QTc was not calculated during sinus rhythm. The heart rates at rest and the heart
rates during exercise were not significantly different on the two treatments, hence the difference in the repolarisation time cannot be attributed to unequal beta blockade.

Even though long term administration of various beta blockers may result in a class III like mode of action, this study and those of others indicate a quantitative difference between these agents and sotalol. Unlike metoprolol, sotalol significantly and selectively prolonged ventricular refractoriness in infarcted zones compared with normal myocardium after short term administration in dogs.\(^2\) In the same study sotalol significantly prevented and slowed ventricular arrhythmias in ischaemic myocardium; metoprolol did not have this effect. Whether this effect is due to the class III property of sotalol or to other differences (beta\(_2\) blockade) is not known.

In conclusion, sotalol significantly prolonged the QT interval by 5–7\% compared with metoprolol after four weeks’ treatment. The prolongation reflects a change in the repolarisation time, since no significant change in the QS interval was noted. As the QT top and QT end intervals showed consistent changes, the former may be used when there are difficulties in defining the end of the T wave. No rate correction factor was required to obtain these QT interval changes because they were measured at a fixed heart rate. Transoesophageal atrial pacing was found to be a simple non-invasive method with few and relatively mild side effects. The method is well suited for drug studies requiring measurements at fixed heart rates.

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References