Measuring QT dispersion: man versus machine

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Abstract

Objective—To compare manual and computer automated techniques for measuring QT dispersion.

Design—Assessment of the ability of manual and automatic measurements of QT dispersion to discriminate between a normal group and two cardiac groups.

Subjects—12 simultaneous electrocardiogram leads were recorded from 25 healthy volunteers, 25 subjects after myocardial infarction, and 25 with cardiac arrhythmias.

Main outcome measures—For each subject, QT dispersion was measured as the difference between the maximum and minimum QT from all 12 leads and separately for only those leads with T amplitudes of >100 μV and for those >250 μV.

Results—Manual QT dispersion (T > 100 μV) was greater (P < 0.02) in the arrhythmia patients (mean ± SD), 45 (21) ms, but not the infarction patients (54 (36) ms), than in the normal subjects (39 (13) ms). There were no significant differences when all T waves were included. QT dispersion was significantly reduced by an average of 30% when T waves < 100 μV were excluded, and by 51% when those < 250 μV were excluded. Automatic techniques gave different measurements for dispersion in comparison with manual measurements. Three of the four automatic techniques detected significant differences between normal and both patient groups when no leads were excluded (P < 0.01) as well as when T waves < 100 μV were excluded (with increased significance, P < 0.002).

Conclusions—Measurements of QT dispersion from small T waves increases measurement variability and reduces the potential for detecting clinical differences. Automatic measurement of QT dispersion gives different results from manual measurement, but can satisfactorily discriminate between normal and abnormal groups with good quality electrocardiograms.

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Dispersion of the QT interval across the 12 standard electrocardiogram leads can provide valuable clinical information. Greater than normal levels of dispersion are associated with death, especially sudden death in heart failure,¹ and with a poor prognosis in hypertrophic cardiomyopathy.² In addition, increased dispersion has been associated with a risk of ventricular fibrillation in acute myocardial infarction³ and with drug arrhythmogenesis.⁴

Traditionally, manual QT dispersion measurement has involved assessment and measurement of 12 leads of the electrocardiogram. This process is tedious and, as the end of T is often ill defined, may be subject to human error. Reliable automatic measurement of QT dispersion is desirable. Although modern electrocardiograph recorders can measure the QT interval automatically, this is not always undertaken in all 12 leads and as yet there is no standardised algorithm for this measurement. Various different algorithms have been developed⁵ for use in these recorders or for research.⁶ Earlier work from our group⁷ has shown that, because of the difficulty in identifying the end of the T wave, different automatic techniques can result in widely different QT measurements from the same ECG complexes.

This study was designed to assess the effectiveness of automatic techniques for the measurement of QT dispersion, and to determine the effect of including or excluding measurements from small amplitude T waves.

Methods

Our QT measurement methods have been described before,⁸ and are therefore presented in outline only.

DATA COLLECTION

All 12 leads of the electrocardiograms were sampled simultaneously by a computer with a resolution of 2:44 μV at a sampling rate of 500 Hz for each lead channel. Electrocardiograms were obtained from 25 subjects in each of the following three groups: healthy volunteers with no history of heart disease, patients with myocardial infarction, and patients with known ventricular or supraventricular arrhythmias but excluding atrial fibrillation. Electrocardiograms were collected from as diverse a range of subjects as possible to ensure that a wide range of T wave shapes was included.

MANUAL ANALYSIS

The electrocardiograms were plotted to paper using a laser printer with a resolution of 118
Figure 1  Manual measurement of QT dispersion for the three clinical groups, for all T waves, and separately for T waves > 100 μV. Values are means, error bars = SD. *P < 0.02.

Figure 2  Manual measurement of QT dispersion for the three clinical groups for all T waves, T waves > 100 μV, and T waves > 250 μV, separately for all 12 leads, at least eight leads, and at least four leads. Values are means, error bars = SD. *P < 0.05; †P < 0.005.

dots per cm. The plots had a vertical (voltage) scale equivalent to 10 mm/mV and a horizontal (time) scale equivalent to 50 mm/s. Using a digitising tablet and a previously validated technique,10 manual QT measurements were performed by an experienced researcher.

QT DISPERSION MEASUREMENT
QT was measured from two consecutive complexes in each electrocardiogram lead and the mean taken. Dispersion was defined as the difference between the maximum QT and the minimum QT across the 12 leads. Dispersion was then recalculated after excluding from analysis leads with T wave amplitudes less than 100 μV, and then again with the exclusion of all leads with T waves less than 250 μV. When T waves were excluded, dispersion measurements were calculated when there were at least eight leads remaining, and also when there were at least four leads remaining.

DATA ANALYSIS
The non-parametric Mann-Whitney test was used when making comparisons between groups. Whenever data could be paired, the paired differences were assessed using the signed Wilcoxon rank sum test. Data presented in the figures are shown as mean (SD).

Results
MANUAL MEASUREMENT
The results for the manual measurement of QT dispersion for all three clinical groups are shown in figure 1. Measurements are shown for all T waves and separately for measurements from T waves which were more than 100 μV in amplitude. When manual measurements were made in all 12 leads there were no significant differences between the patient groups. However, small T waves are known to introduce measurement variability. When T waves of 100 μV or less were excluded, significant differences between dispersion measures for the normal group (mean (SD), 39 (13) ms) and the arrhythmia group (45 (21) ms) were observed (P < 0.02). There was no significant difference between the normal and post-myocardial infarction groups (54 (36) ms, P = 0.1), because of the very large range of measurements in the infarction group. The numbers of subjects in each subgroup are given in table 1.

The effects of introducing different exclusion criteria for T waves and requiring the availability of different minimum numbers of
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leads are shown in figure 2. When small T waves were excluded there were significant reductions in QT dispersion. Across all subject groups there was an average fall of 30% when T waves with an amplitude of < 100 µV were excluded, and by 51% when those < 250 µV were excluded. There were no significant differences between measurements which included at least eight leads and those which included at least four leads.

AUTOMATIC MEASUREMENT

Figure 3 shows the automatic measurements of QT dispersion for the three clinical groups for T waves of > 100 µV when at least eight leads were available. All automatic techniques except the threshold technique showed very significant differences between the normal subjects and the two patient groups (P < 0.002).

COMPARISON BETWEEN MANUAL AND AUTOMATIC MEASUREMENT

Figure 4 plots the differences between the manual and automatic dispersion measurements for the same data as in fig 3 (T waves of > 100 µV and at least eight leads). The mean (SD) of these paired differences are tabulated, along with the other intercomparisons, in table 2.

Discussion

When the three groups were compared using manual measurements from all 12 leads no significant differences were detected. However, small T waves are known to introduce greater variability in measurement. When these were excluded, QT dispersion was significantly greater in the arrhythmia patients than in the normal subjects. Although the mean dispersion in the infarct group tended to be greater (not significantly), there were no significant differences between the normal and infarct group because of the large range of dispersion values. These reflect the range of dispersion, but may also be influenced by measurement difficulties with more diverse T wave shapes.

We have previously suggested that accurate QT measurement demands a T wave amplitude of at least 250 µV. However, excluding small T waves may jeopardise the reliability of QT dispersion analysis. The selection of such a threshold level is therefore a compromise between excluding too many small T waves and maximising the number of leads available for analysis. From the results obtained,
100 μV is a reasonable compromise for dispersion analysis. The number of leads available for analysis is important. For the main analysis of this study we required at least eight measurable leads. Nevertheless, including results from four or more leads had a smaller effect on QT dispersion than in changing the amplitude threshold for T wave inclusion. This was particularly evident in the post-myocardial infarction group. Including all T waves of > 100 μV, there were 23 patients with at least four leads available for analysis and 14 with at least eight leads, but there was no significant difference in QT dispersion between the four and eight lead data. Currently there are different views on the validity of correcting or adjusting QT dispersion for missing leads. Hnatkova et al15 accepted that correction was difficult, but concluded that the formula used in their study could be used to adjust QT in healthy subjects. Our study was not designed to determine whether a correction should be applied to compensate for missing leads, but it highlights the considerable problems that a correction factor would introduce. This is supported by Glancy et al14 who concluded that lead adjustment formulas for QT dispersion are not appropriate in patients with myocardial infarction. In clinical QT dispersion studies T waves are often excluded because they were small or difficult to measure. As they were not randomly selected, the statistical rules for correcting random sample data cannot be applied.

Our data provide strong support for the view that QT dispersion can be measured automatically, albeit with a different range of values. QT measurement based on intersection of the T wave with a threshold level above the isoelectric level, however, cannot be recommended. This accords with our previous work showing that variability using this technique was unacceptably high.16 The three other techniques gave discriminating results, and in that respect were superior to manual measurement. As in our study all electrocardiograms were of good quality, it is possible that as noise increases some or all of these techniques would suffer in comparison with manual measurement.

A normal range of QT dispersion is as yet to be defined, but it will be very dependent on the assessment technique employed. We have shown that, with certain limits, automatic QT dispersion analysis can be powerful, but whatever technique is finally adopted, it must be described accurately. Different techniques will produce different dispersion results, and new reference ranges will need to be adopted.

Our findings that the QT measurement approach based on the slope intercept technique can give useful results indicates that information on dispersion is contained not just in the final components at the end of the T wave, but also on other aspects of the T wave. This raises the prospects of capturing even more information from the surface electrocardiogram T wave. This observation is important for improving our understanding of the electrophysiological factors influencing QT dispersion.

CONCLUSION

QT dispersion can be assessed automatically in electrocardiograms of good quality. Small amplitude T waves pose a particular problem and we would recommend that those smaller than 100 μV are not used. Our results suggest that the T wave may contain more information than is currently addressed even by the relatively new approach of QT dispersion. Recovering QT and T wave information is very dependent on ECG quality. Studies and clinical assessments based on these features must employ meticulous methods to acquire the electrocardiogram and hence maximise the opportunities for analysis. In addition, we have shown that global features of T waves provide valuable information on dispersion, and this should help to improve our understanding of the clinical course of QT dispersion.

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12 McLaughlin NB, Campbell RWF, Murray A. Accuracy of four automatic QT measurement techniques in cardiac patients and healthy subjects. Heart 1996;76:422-6.
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