Introduction

Stress test has been used in Brazil since 1972 and its sensitivity and specificity for the diagnosis of chronic CAD are 50-72% and 69-74%, respectively.\(^1\) The QT interval dispersion (QTD) measurement is considered a promising instrument to improve the diagnostic accuracy of stress test. QTD was defined in the 1990s\(^5\) as the difference between maximal and minimal QT interval duration measured in 12 ECG leads. It has been proposed as a regional marker of ventricular repolarization dispersion (VRD) and correlates with the dispersion of action potentials (AP) in animals and humans.\(^6\) QT interval, measured from the beginning of the QRS complex to the end of the T wave, represents the time it takes for ventricular myocardial cells to depolarize and repolarize.\(^7\) However, U-wave should not be included in the measurement.\(^8\)

During exertion, patients with chronic CAD present increased ventricular repolarization heterogeneity,
which is reflected by increased QTD.²⁻¹¹ Koide et al.,¹² observed that even when a coronary patient, undergoing stress test, did not present ischemia criteria, the QTD was higher (62 ± 13 ms) compared to patients without coronary disease (40 ± 14 ms). Musha et al.,¹³ and col. showed an increase of QTD after exercise, which was not reduced by beta-blockers.¹³ Naka et al.,¹⁴ in a study with infarcted patients, found an increase in QTD due to residual ischemia; however, it did not increase in patients without residual ischemia.

There are several limitations to QT interval measurement technique. Among them, T-P fusion during higher heart-rates and changes in Qt interval rates in relation to men and women¹⁵,¹⁶ are worthy of note.

The aims of this study are to evaluate whether the QTD index is sensitive to action potential changes in the presence of stress induced myocardial ischemia, as well as to define a cutoff point for QTD that could become a diagnostic criterion for myocardial ischemia.

**Methods**

An observational analytical study, where 80 patients underwent exercise testing and coronary angiography (CAT), with a maximum interval of 6 months between the tests. The patients were aged 18 to 80 years. People who had any of the following conditions were excluded from the study: previous acute myocardial infarction (AMI), complete right (RBBB) or left (LBBB) bundle branch block, patients with long QT syndrome; patients with known ventricular dysfunction; unreadable ECG traces or ECG where less than eight electrocardiographic leads were available for QTI measurement.¹⁷ A total sample of 80 patients was defined at the convenience of the researcher. Treadmill test was performed with an analog-to digital converter of signals, Ergo PC 13 model in Micromed 2.3 version, with a simultaneous acquisition of twelve leads and record with speed of 25 mm/s and amplitude of 10 mm/mV. The protocols used for all patients were individualized aiming at reaching maximum heart rate. The test was considered suggestive of myocardial ischemia in case the patient presented at least one of the ischemia criteria defined by the III Guidelines on ergometric tests of the Brazilian Society of Cardiology,¹⁸ typical chest pain on exertion; ST-segment elevation or depression, equal to or greater than 1 mm, in relation to baseline ECG. The QT interval of each lead was calculated by the mean of the three beats with less artifact. By using a cursor, one point was marked at the beginning of the QRS complex and another at the end of the T-wave (the point where the T wave returned to the isoelectric line) for each of the three beats.¹¹ After the measurement, we calculated the mean of the three values found, which would be the value to be considered as the QTI of the mentioned lead. The same procedure was performed for the 12 leads. Thus, for each patient in the study, at least, 24 QT intervals were measured at rest (standing) and 24 on ECG obtained within the first minute of the recovery stage. We decided to “measure the peak stress” within the first minute of the recovery phase in order to minimize the technique’s artifacts. At the end of the measurement of the QT intervals of all leads, we marked the highest and the lowest measure found, in the two phases studied: rest and effort. From these values, we calculated the QTI dispersion of these two phases, and also a delta QT dispersion value by determining the difference of QTD between effort and rest.

In order to adjust the QT interval for the corresponding heart rate (HR), we used the Bazett’s formula. The adjustment enabled the calculation of the QTc (QT interval dispersion corrected for heart rate), and also the QTc “delta” – the difference between rest-stress QTc intervals. All the electrocardiographic measurements were done by a single observer. Figure 1 shows the sequence to measure a QT interval.

Interobserver variability was determined by measurements performed by a second researcher, who was blinded to the measures obtained by the first observer. The second ECG expert measured the QT interval in 12 patients randomly selected (patients were numbered from 1 to 63, and 12 numbers were raffled). The correlation between the measures was determined by Pearson’s correlation coefficient. The Bland-Altman test was also used to assess interobserver variability (Figure 2).

After catheterization, patients with a coronary stenosis of at least 70% of one or more arteries, or with ≥ 50% stenosis of the left coronary trunk (LCT), were classified as “people with obstructive coronaropathy”. In contrast, patients with with stenosis less than 70% in epicardial coronary arteries, or less than 50% in the left coronary trunk, were classified as “without obstructive coronary disease”.

The Medcalc software was used for the statistical analysis. The data were expressed in absolute numbers, percentages and standard deviation. The classificatory variables were presented as tables, and the propotions were compared using the chi-square
Figure 1 - Measurement sequencing of a QT interval.

Figure 2 - Reproducibility of QT dispersion measurements before (A) and during (B) effort, assessed using the Bland-Altman method.
The Kolmogorov-Smirnov was used to assess the normality of the continuous variables. To identify the best QT dispersion cutoff point for the diagnosis of obstructive coronaropathy, ROC curves were used both for QT dispersion values and for rest and stress QTc. We calculated the QTd, the QTc, and conventional stress test sensitivity and specificity for the diagnosis of chronic obstructive coronaropathy. We also found a QT delta – the difference between rest-stress QTc intervals, as well as the QT delta - the difference between rest-stress QTd intervals. Subsequently, patients were divided into three groups: true positive (TP) – patients with positive stress test for ischemia and coronary angiography showing stenosis ≥ 70% of at least one major epicardial artery, except for left coronary trunk lesions which were considered to be significant when the obstruction was > 50%. The false positive (FP) group was composed of patients with positive stress test and stenosis less than 70% in at least one major epicardial artery, except for left coronary trunk lesions which were considered to be significant when the obstruction was < 50%. Finally, the true negative (TN) group was composed of negative stress test patients and coronary angiography showing stenosis less than 70% in any epicardial coronary artery, except for left coronary trunk lesions which were considered to be significant when the obstruction was < 50%. p < 0.05 was considered to be statistically significant and one-way-variance analysis (ANOVA) was used to compare the three groups. A paired t-test was used to assess the QTc behavior during rest and effort.

All the patients signed the free and clarified term of consent and the research Project was approved by the Ethics Committee on Human Research of Federal University of Espirito Santo (UFES), by the protocol number: 06177412.1.0000.5071.

**Results**

The difference between the mean QTd values obtained by both observers, at rest, was only 0.8 ± 18.3 ms, which is quite satisfactory. However, we observed some data points of standard deviation away from the mean, which demonstrated the low reproducibility of the measures at rest. The difference between the means obtained by the two observers was 9.5 ± 12.5 ms in peak stress measurements. Additionally, we also found data points of standard deviation away from the mean, which confirmed the low reproducibility of QT dispersion for peak stress measures as well.

Table 1 - General characteristics of the groups

|                | TP (n = 26) | FP (n = 23) | TN (n = 14) | p-value |
|----------------|-------------|-------------|-------------|---------|
| Age (years)    | 58 ± 10     | 54 ± 12     | 56 ± 11     | 0.43    |
| Male sex (%)   | 81%         | 71%         | 71%         | 0.453   |
| Diabetes mellitus (%) | 27%         | 13%         | 7%          | 0.213   |
| Arterial hypertension (%) | 58%         | 58%         | 79%         | 0.372   |
| Beta-blockers (%) | 23%         | 33%         | 21%         | 0.633   |
| ACEI/ARB (%)   | 42%         | 21%         | 21%         | 0.188   |
| Statins (%)    | 12%         | 8%          | 21%         | 0.199   |
| Calcium antagonists (%) | 12%         | 0%          | 29%         |         |

Values expressed as mean ± SD or percentages; TP: true positive; FP: false positive; TN: true negative.
the stress QTd, the sensitivity was 44.4% and the specificity was 81.6% (AUC 0.585; CI 95% 0.465-0.699) with a cutoff of 46 ms. For stress QTc, the sensitivity was 58.3% and the specificity was 63.2% (AUC 0.593; CI 95% 0.472-0.706) with a cutoff of 57 ms. In relation to the sensitivity and specificity values of the traditional treadmill test, and considering the presence of ST segment depression or typical chest pain on exertion, we found a sensitivity of 72% and a specificity of 32% in our sample. Since the confidence intervals found for the QTd and QTc cutoff values included the 0.5 value, we decided not to aggregate the QT dispersion values into the traditional stress test, because any improvements in the sensitivity and specificity that we could possibly find would not have been reliable.

When we analyzed the three groups formed (TN, TP and FP), the following was found: the mean values of QT dispersion at rest did not show statistically significant difference between the three groups. Respectively, 58 ± 30 ms, 47 ± 22 ms and 43 ± 19 ms, for the TN, TP and FP groups (p = 0.172). In addition, we did not observe significant difference between the mean values of the QTc dispersion at rest: 67 ± 40 ms, 55 ± 26 ms and 49 ± 21 ms, respectively, for the TN, TP and FP groups (p = 0.163). Thus, we moved on to analyze the mean QT dispersion values found between the three groups during effort. Similarly to what happened in relation to QT dispersion at rest, we found close mean values with no statistical difference between the three groups: 32 ± 11 ms, 48 ± 28 ms and 42 ± 22 ms, respectively, for the TN, TP and FP groups (p = 0.124). However, when we analyzed the data of QTc dispersion of effort, we verified that the values of QTc between the three groups were different: TN (47 ± 17 ms), TP (72 ± 42 ms) and FP (61 ± 31 ms), with p = 0.003. When we compared TN and TP, we found p < 0.05; when comparing TN and FP, we also found p < 0.05; however, when VP and FP were compared, we found p > 0.05 (Table 2).

In order to better assess the changes in stress induced coronary depolarization, we created a delta QT dispersion value (ΔQTd) which was obtained by the following equation: ΔdQT = QTd stress – QTd rest. Likewise, we obtained a delta value of QTc dispersion by a similar equation: ΔdQTc = QTc stress – QTc rest. The ΔQT was ~25 ± 33 ms in the TN group, 1 ± 27 ms in the TP and ~2 ± 23 ms in the FP group, with statistical difference between the three groups, with p = 0.013. Comparing TN and TP, we found p < 0.05; the same was observed when TN was compared with FP (p < 0.05); in contrast, the comparison between TP and FP showed p > 0.05. The mean ΔQTc dispersion was ~20 ± 45 ms in the TN group, 17 ± 40 ms in the TP group and 11 ± 30 ms in the FP group. Again, the same "p" value of 0.013 was found between the three groups, as well as the same values of “p” for the other comparisons: TN vs VP (p < 0.05), VN vs FP (p < 0.05) and VP vs FP (p > 0.05). We did not find any statistical difference between TP vs FP. Due to the statistical difference found between the three groups, in relation to the mean values of ΔQTc, we decided to illustrate the behavior of QTc dispersion from rest to stress peak in the three groups. Figures 3, 4 and 5 show the behavior of the three groups.

**Discussion**

The aim of this study was to assess the relationship between QT interval dispersion and chronic CAD. The focus of our investigation was to evaluate the feasibility
of using the QTd (within the first minute of recovery and/or rest) for the diagnosis of significant coronary disease. Based on previous studies, we hypothesized that coronary obstruction would lead to prolonged action potential in the ischemic region, and that such increase could be identified by the QTd. Actually, it would reflect in theory the difference between repolarization in the ischemic region compared to the non-ischemic in the ventricular syncytium.

The first result that will be discussed concerns the reproducibility of QT dispersion measurements. One of the major problems concerning QT interval measurements is the difficulty in obtaining acceptable interobserver variability. In our study, interobserver
reproducibility was weak. Our impression is that, even with the measurements being performed with more modern software, the problem involving the reproducibility of QT intervals measurements remains a considerable one.

With the results of catheterization, QT dispersion, QTc and stress test, we built ROC curves in order to find the possible cutoff values of QTd and stress-rest QTc, so as to subsequently calculate the sensitivity and specificity of QTd for the diagnosis of chronic CAD. We did not achieve a minimaly satisfactory ROC curve for QTd and QTc at rest. For QT and QTc dispersion of effort, we obtained ROC curves somewhat better than those obtained at rest, and we found cutoff values for QTd (46 ms) and QTc (57 ms). As for the sensitivity and specificity of stress QTd and QTc it is possible to say that they were comparable to the traditional stress test ischemic criteria. However, what really called attention, was the low specificity of the classical diagnostic criteria of exercise induced ischemia in our sample (32%). On the other hand, the sensitivity was 72%. The high false-positive rate was quite high. We observed that most of the false-positive results were found in the presence of segment depression without a concomitant stenosis of at least 70%. The most plausible explanation found lies in the arbitrariness of considering as the possible cause of ischemia only stenosis with obstruction greater than 70%. Smaller plaques, but under the effect of vasoconstriction substances, can cause ischemia. Another factor to be considered is the possibility of microcirculation disease in patients without significant disease of large coronary arteries. Finally, we must consider that the degree quantification of coronary obstruction made by the doctor responsible for the catheterization is performed by visual method only and, hence, it is observer-dependent.

When we analyzed the data from the three groups formed: true positive (TP), false positive (FP) and true negative (TN), heart rate (HR) and systolic pressure (SP) at stress peak were not statistically different between the three groups (Table 2). The incidence of typical chest pain and ST depression at peak stress were higher among the false-positive and true-positive groups, and was not present in the TN group, as we already expected. In relation to the QT dispersion and QTc dispersion at rest, there was no significant statistical difference between the three groups. In contrast, stress QTc was significantly higher among the TP and FP groups, compared with the TN group. The TP and FP groups behaved so similarly that made us wonder about the real importance of considering “people with significant coronary artery disease” only those patients with stenosis of at least 70% in epicardial arteries, or 50% or more in the left coronary trunk. We can speculate that if myocardial perfusion scintigraphy had been performed, instead of cardiac catheterization, as gold standard for significant CAD, it is possible that our results would have been similar to those...
obtained by Stoletniy and colleagues. These authors showed that the QTc increased from rest to stress in patients with ischemia documented by myocardial scintigraphic imaging, and that it did not increase from rest to stress peak in patients without myocardial scintigraphic imaging using radioisotope techniques. We speculate that QTc is, ultimately, a marker of myocardial ischemia, with no strict connection with the degree of obstruction in large coronary arteries.

The QTc delta also showed significant statistical differences between the groups. However, in order to test the theory that supports the concept of QTd, we decided to sketch a line graph that represented each individual in the three groups. Our concern was that the QTc and QTc delta values would show only the statistical difference between the means of the three groups (extreme QTc values of few individuals of one group can affect the mean and not necessarily represent the behavior of the rest to stress condition variable). Corroborating the results of the QTc dispersion of effort, nineteen patients from the TP group presented increased QTc from rest to stress conditions (Figure 3), whereas seven individuals showed a reduction. In contrast, in the TN group, five people presented increased QTd and nine reductions (Figure 5). The FP group maintained similar behavior as the TP group. In FP, fourteen patients increased the QTc dispersion and nine reduced it (Figure 4). In order to statistically test the behavior of each group in relation to the QTc dispersion of effort and at rest, we used a paired T-test for each group, and obtained significant differences between the TP group means from rest to stress conditions, and a trend to increased QTc in the FP group. The TN negative group did not show significant changes between the QTc pre- and postexercise means. We conclude that QTc behaves, predominantly, with an increase on exertion in patients with myocardial ischemia; and tends not to change significantly in patients without ischemia.

Our study has some limitations that should be pointed out. First, our sample was small so we could not obtain a highly reliable cutoff point for QTc. Still, we must remember that our QTc cutoff point was quite similar to the ones found by other authors. Secondly, the methodology we used to measure the QTc – as far as we know - is unprecedented in the literature. Thus, our data must be confirmed by other similar studies. Our results can only be considered for a coronary population with no previous AMI or ventricular dysfunction. Finally, this is a database retrospective study with all the limitations inherent to this type of study.

Conclusions

Based on our results, we believe that QT dispersion is, despite being a “crude” marker of ventricular repolarization heterogeneity, is sensitive to stress-induced myocardial ischemia and can aid in the diagnosis of chronic CAD.

Author contributions

Conception and design of the research: Barcelos AM, Mill JG. Acquisition of data: Barcelos AM, Rodrigues SL, Mill JG. Analysis and interpretation of the data: Barcelos AM, Baldo MP, Rodrigues SL, Mill JG. Statistical analysis: Barcelos AM, Baldo MP, Rodrigues SL, Mill JG. Obtaining financing: Barcelos AM. Writing of the manuscript: Barcelos AM. Critical revision of the manuscript for intellectual content: Barcelos AM, Baldo MP, Rodrigues SL, Mill JG.

Potential Conflict of Interest

No potential conflict of interest relevant to this article was reported.

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Study Association

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Ethics approval and consent to participate

This study was approved by the Ethics Committee of the Hospital Universitário Cassiano Antônio de Moraes under the protocol number CAAE: 06177412.1.0000.7051. All the procedures in this study were in accordance with the 1975 Helsinki Declaration, updated in 2013. Informed consent was obtained from all participants included in the study.
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