The effect of glucose variability on QTc duration and dispersion in patients with Type 2 Diabetes Mellitus

Yasar Sertbas¹, Ali Ozdemir², Meltem Sertbas³, Akin Dayan⁴, Seda Sancak⁵, Cihangir Uyan⁶

ABSTRACT

Objective: Glycemic variability (GV) is a new term with the episodes of hyper and hypoglycemia in diabetic patients. Both prolonged QT interval and QTd are potential risk factors for malignant ventricular arrhythmias affecting the mortality of different groups of patients including diabetes mellitus. In this study, we aimed to evaluate if the glucose variability increasing the QTc interval and QTc dispersion in type 2 diabetes mellitus.

Methods: We included 275 consecutive patients with type 2 diabetes. We quantified the GV with standard deviation (SD) and coefficient of variation (CV) from 7 point glucose measures. We investigated the relationship of GV parameters with QT parameters.

Results: The prevalence of prolonged QTc duration was 21%, no patients have prolonged QTc dispersion (> 80 ms). SD of the patients with prolonged QTc duration was significantly higher than the others (45.14 ±24.45 vs. 37.78 ±9.03 p<0.05). There was also a significant relationship between SD and QT dispersion (r: 0.164; p: 0.007). There were no relationship between the QT parameters and microvascular diabetic complications. SD and HbA1c levels were significantly higher on the patients having peripheral neuropathy (p<0.005).

Conclusion: The result of this study demonstrates that increased glycemic variability is associated with prolonged QTc duration and QTc dispersion. It is important to focus on targeting optimal glycemic control with GV as an additional goal point along with the traditional following parameters such as fasting-postprandial blood glucose and HbA1c.

KEY WORDS: Glucose variability, QTc dispersion, QT duration.

doi: https://doi.org/10.12669/pjms.331.11440

How to cite this: Sertbas Y, Ozdemir A, Sertbas M, Dayan A, Sancak S, Uyan C. The effect of glucose variability on QTc duration and dispersion in patients with Type 2 Diabetes Mellitus. Pak J Med Sci. 2017;33(1):22-26. doi: https://doi.org/10.12669/pjms.331.11440

This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/3.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

INTRODUCTION

Glycemic control is essential for the management of type 2 diabetes mellitus. Glycemic variability (GV) is a term in which the diabetic patients have similar mean glucose levels, although they exhibit differences in terms of both the number and degree of glucose excursions.¹ The broad definition of GV takes into account the intraday glycemic excursions including episodes of hyper and hypoglycemia.² Recently, many clinical studies have shown that glycemic variability might be related with cardiovascular diseases and mortality.³⁻⁶

The QT interval on the electrocardiogram (ECG) reflects the total duration of ventricular depolarization and repolarization. The interlead variation in duration of the QT interval on the surface ECG has been referred to as QT dispersion (QTd). Both prolonged QTc interval and QTc dispersion are potential risk factors for malignant ventricular arrhythmias affecting the mortality...
of different groups of patients including diabetes mellitus.7-9 There are also many studies about the influence of changes in glycemia on the length of QT parameters. Either hypoglycemia or hyperglycemia both can increase the QT duration and QTd.10,11

In this study, we aimed to evaluate the effect of glucose variability on QT interval and dispersion in type 2 diabetic patients as a predictive markers of arrhythmias and mortality.

**METHODS**

The present study included 275 consecutive patients with type 2 diabetes treated in the internal medicine department of Fatih Sultan Mehmet Education and Research Hospital, Turkey between January and March 2016. The study was approved by the institutional Ethics Committee and all the participants gave written informed consent (FSM EAH-KAEK: 2015/21).

Medical documentation of all patients were systematically recorded with respect to age, sex, body weight, diabetes duration, medication and presence of diabetic microvascular complications (retinopathy, nephropathy and peripheral neuropathy).

Blood pressure was measured two consecutive times in the sitting position by an appropriate cuff size. Hypertension was diagnosed as systolic blood pressure (SBP) ≥140mmHg or diastolic blood pressure (DBP) ≥90 mmHg or the current use of antihypertensive drugs.

Diabetic retinopathy was evaluated by an ophthalmologist; diabetic nephropathy was diagnosed if patient had creatinin above 1.5mg/dl or urine protein excretion was > 300 mg/day. Peripheral neuropathy was diagnosed on the basis of neuropathy symptoms and signs of objectively abnormal results, including intensitivity to a 10g monofilament and abnormal electromyography (EMG) findings.

We quantified the glycemic variability with standard deviation (SD) and coefficient of variation (CV) from the multiple self monitoring of blood glucose (SMBG) readings taken over the course of the day with seven point glucose measures (before breakfast, two hour after breakfast, before lunch, two hour after lunch, before dinner, two hour after dinner and before sleep) CV was defined as the ratio of SD to mean glucose values expressed as a percentage.12 Blood glucose concentration was measured in capillary blood obtained by finger stick with a point of glucometer.

A single investigator blind to the clinical data interpreted QT intervals using the program Adobe Photoshop CS64. The QT interval was measured from the beginning of the QRS complex to the end of the downslope of the T wave (crossing the isoelectric line); when U wave present, QT interval was measured to the nadir of the curve between the T and U waves. The QT interval corrected for the previous cardiac cycle length (QTc) was calculated according to Bazett’s Formula. QTc = QT/(RR)1/2. QTc>440 ms was considered prolonged. QTc dispersion was calculated using the difference between the maximum and the minimum QTc in any thoracic lead. QTc dispersion >80ms was considered prolonged.13

Blood samples were obtained after 12 hours fasting period. Glucose was carried out by the glucose oxidase method in the auto analyzer (Architect, 16000, Abbot Diagnostic) and that of HbA1c was measured with high performance liquid chromatography (HPLC) (Trinity Biotech Premier Hb9210).

Analyses were performed using statistical package for social sciences (SPSS) version 22.0 for Windows. Data are expressed as mean (standard deviation. One-sample Kolmogorov-Smirnov test was performed to assess the distribution of data. Numerical variables in different subjects were compared by t-test or Mann-Whitney U test. Bivariate correlation analyses were made by Pearson correlation test. Probability values were two tailed, and a p-value of less than 0.05 was considered significant.

**RESULTS**

There were 275 participants which included 172 female and 103 male diabetics with a mean age of 56.45±9.95. Patients’ demographic, clinical and metabolic data are presented in Table-I.

| Table-I: Patients demographic, clinical and metabolic characteristics. |
|---------------------------------------------------------------|
| Number of patients:                                           |
| Gender (male /female)                                         |
| Age (years)                                                  |
| Diabetes duration (years)                                    |
| Body mass index (kg/m²)                                      |
| Hypertension (%)                                             |
| Retinopathy (%)                                              |
| Nephropathy (%)                                              |
| Peripheral neuropathy (%)                                    |
| Metformin usage (%)                                          |
| Sulphonylurea usage (%)                                      |
| Insulin usage (%)                                            |
| Fasting glucose (mg/dl)                                      |
| HbA1c (%)                                                    |
| SD                                                          |
| CV                                                          |
| QTc                                                         |
| QTc dispersion                                               |
| 275                                                         |
| 103/172                                                     |
| 56.45±9.95                                                  |
| 9.76±7.65                                                   |
| 30.80±5.37                                                  |
| 38.9                                                       |
| 11.6                                                       |
| 17                                                         |
| 28.7                                                       |
| 92.7                                                       |
| 20                                                         |
| 52.3                                                       |
| 165.13±55.78                                               |
| 8±1.68                                                     |
| 39.3±18.75                                                 |
| 24.03±9.38                                                 |
| 419.07±28.54                                              |
| 26.43±10.1                                                 |
There was a significant positive correlation between SD and QTc dispersion ($r$: 0.164; $p$: 0.007). In linear regression analysis, the relation of the SD and QTc dispersion was as $\beta$ = 0.088, $p$<0.001. (Table-II).

While the prevalence of prolonged QTc duration was 21% (QTc>440 ms), no patients have prolonged QTc dispersion (> 80 ms). We compared the clinical and metabolic characteristics of patients with prolonged QTc duration ad normals, it is shown in Table III that SD was the only parameter significantly higher on patients with prolonged QTc duration than normals ($p$<0.05). No other metabolic parameters were affected from the prolongation of QTc duration (Table-III).

There were no relationship between the QT parameters and microvascular complications. GV or QT did not have any effect on nephropathy. Retinopathy was more prominent on patients with higher. HbA1c levels. SD and HbA1c levels were significantly higher in among patients having peripheral neuropathy ($p$<0.005) (Table-IV).

**DISCUSSION**

GV with the components of the hypo and hyperglycemia attacks increase the cardiac mortality in type 2 diabetic patients. By looking the relationship of GV and the increase of QT duration and QTc dispersion, we can get information about the effect of GV on malignant ventricular arrhythmias. This study demonstrates that ventricular depolarization and repolarization are affected by glycemic variability in type 2 diabetic patients.

Table-II: The association between glucose variability and QT parameters.

|       | QTc duration | QTc dispersion |
|-------|--------------|---------------|
| SD    | 0.116        | 0.164**       |
| P     | 0.055        | 0.007         |
| CV    | 0.045        | 0.093         |
| P     | 0.457        | 0.124         |

**correlation is significant at the 0.01 level.

Table-III: The effect of QTc prolongation on the glyemic parameters and complications.

|       | QTc ≤ 440 ms   | QTc> 440 ms   | P    |
|-------|----------------|--------------|------|
| SD    | 37.78 ±16.63   | 45.14±24.45  | 0.008|
| CV    | 23.48 ±9.03    | 26.09±10.39  | 0.6  |
| Fasting glucose (mg/dl) | 165.22 ±7.49 | 164.79±49.31 | 0.958 |
| HbA1c (%) | 7.97 ±1.71 | 8.23 ±1.63 | 0.366 |
| Diabetes duration | 9.67 ±7.01 | 10.09±9.73 | 0.714 |
| Neuropathy | 22.5% | 6.2% | 0.507 |
| Retinopathy | 9% | 2.6% | 0.602 |
| Nephropathy | 11.2% | 2.5% | 0.306 |

Robinson et al. and Marques et al. showed that experimental hypoglycemia causes an acquired long QT syndrome. Robinson et al. and Marques et al. showed that experimental hypoglycemia causes an acquired long QT syndrome. Later on it was also shown that QTc interval lengthens significantly during spontaneous clinical episodes of hypoglycemia. The cause of prolonged QTc intervals and QTc dispersion during hypoglycemia is complex and multifactorial. Increased sympathoadrenal activity, high levels of catecholamines and hypokalemia induced by hyperinsulinemia are claimed as factors to be involved.

Hyperglycemia is another component of GV, has also been shown to be related with prolonged QT interval and QTc dispersion. Marfella et al. and Gordin et al. showed that acute hyperglycemia produces significant increments of QTc and QTc dispersion both in the diabetic patients and healthy individuals. Reduction of nitric oxide bioavailability and stimulation of protein kinase C were suggested as the mechanisms that cause QT prolongation in hyperglycemia.

In our study the prevalence of prolonged QTc (QTc >440 ms) was 21%. On the other hand, no patients have prolonged QTc dispersion (> 80 ms). These findings were in accordance with the previous studies as the prevalence of high QTc duration and QTc dispersion ranged from 15.4 to 67% for

Table-IV: Relationship of GV and QT parameters with microvascular complications.

|       | SD        | CV         | HbA1c  | QTc      | QTc dispersion |
|-------|-----------|------------|--------|----------|----------------|
| Retinopathy (11.6%) | 38.82±19.19 | 24.17±9.75 | 7.93±1.69 | 417.4±27.63 | 26.22±10.66 |
| (+)   | 42.90±19.82 | 23.21±8.96 | 8.8±1.90 | 425.9±22.89 | 27.56±10.13 |
| P     | 0.266     | 0.603      | 0.009  | 0.100    | 0.508          |
| Nephropathy (13.7%) | 39.10±19.29 | 24.01±5.3 | 7.96±1.71 | 419.70±27.99 | 26.21±10.06 |
| (+)   | 40.88±17.51 | 23.91±9.09 | 8.21±1.62 | 415.87±23.93 | 28.31±10.09 |
| P     | 0.592     | 0.954      | 0.417  | 0.422    | 0.293          |
| Peripheral Neuropathy (28.7%) | 36.77±17.88 | 23.68±9.46 | 7.72±1.44 | 418.96±27.32 | 26.26±10.98 |
| (+)   | 44.01±21.13 | 24.54±9.96 | 8.66±2.09 | 418.34±27.04 | 26.78±9.70 |
| P     | 0.006     | 0.523      | 0.000  | 0.868    | 0.724          |
QTc and from none to 33% for QTc dispersion.\textsuperscript{25-26} Ninkovic et al. mentioned about the GV as a risk factor of prolonged QT and QT dispersion.\textsuperscript{27} They used the mean amplitude of glycemic excursion (MAGE) to represent the GV. MAGE was originally developed using hourly glucose samples and it has emerged as the preferred method for assessing continuous glucose monitoring (CGM) data. We used the SD and CV as the simplest method of assessing intra-day variability of serum glucose from the multiple (7 point) SMBG readings taken over the course of the day. When we compared the glycemic parameters of the patients who have QTc parameters longer than 440 ms and below, we saw that SD was the only parameter significantly higher in the group having a longer QTc duration. Since no patient had QTc dispersion above 80 ms we looked into the correlation between the QTc dispersion and glycemic variability. SD was significantly correlated with QTc dispersion (p: 0.007; r: 0.164). Although there was no correlation between QTc duration and SD (p: 0.055), because of its proximity to the level that can be considered significant (p<0.05), we thought that the reason might be the low number of study population.

It has been suggested as the glucose variability may have been responsible for microvascular complications of diabetes. Braged et al. showed that GV was an independent predictor of peripheral neuropathy, however, no significant relationship was found between GV and the other microvascular complications (retinopathy, nephropathy).\textsuperscript{28} In another study painful neuropathy was found to be related to increased glucose flux.\textsuperscript{29} Retinopathy was also found to be related to M-FBG and HbA1c.\textsuperscript{30} Diabetic complications were also suggested to be higher among the patients with long QTc duration. Although some studies have suggested a higher relationship between diabetic cardiovascular autonomic neuropathy (CAN) and QT prolongation, later studies have not confirmed an association of prolonged QT and CAN.\textsuperscript{31,32} Li et al. claimed that microalbuminuria was the only diabetic microvascular complication related to QTc prolongation other than retinopathy and nephropathy.\textsuperscript{33} On the other hand Ninkovic et al. showed a significantly higher prevalence of retinopathy and polyneuropathy among patients with high QTc duration.\textsuperscript{25} In our study, we couldn’t find any relation between microvascular complications and QT parameters. While retinopathy was only found to be related with HbA1c, nephropathy hasn’t any relationship with GV parameters. SD and HbA1c were found to be significantly higher among the patients with diabetic neuropathy.

**Limitations of the study:** It includes the manual assessment of the QTc interval and dispersion. Another limitation is the continuous glucose monitoring could be done instead of multiple self monitoring blood glucose. It should also be kept in mind that the superiority of available autonomic methods has not been proven for measurement of QT parameters and multiple SMBG is more feasible than CGM in daily life.\textsuperscript{33}

In conclusion, the result of this present study suggests that increased glycemic variability associated with prolonged QTc duration and QTc dispersion, which means ventricular depolarization and repolarization are affected by glycemic variability in type 2 diabetic patients. It is important to focus on targeting optimal glycemic control with GV as an additional goal point along with the traditional following parameters such as fasting-postprandial blood glucose and HbA1c.

**REFERENCES**

1. Jung HS. Clinical Implications of Glucose Variability: Chronic Complications of Diabetes Endocrinol Metab (Seoul). 2015;30:167-174.
2. Satya Krishna SV, Kota SK, Modi KD. Glycemic variability: Clinical implications. Indian J Endocrinol Metab. 2013;17:611-619.
3. Gordin D, Rönnback M, Forsblom C, Mäkinen V, Saraheimo M, Groop PH. Glycemic variability, blood pressure and arterial stiffness in type 1 diabetes. Diabetes Res Clin Pract. 2008;80:4-e7.
4. Mugggeo M, Verlato G, Bonora E, Zoppini G, Corbellini M, de Marco R. Long-term instability of fasting plasma glucose, a novel predictor of cardiovascular mortality in elderly patients with non-insulin-dependent diabetes mellitus: The Verona diabetes study. Circulation. 1997;96:1750-1754.
5. Kota SK, Mahapatra GB, Kota SK, Naveed S, Tripathy PR, Jammula S, et al. Carotid intima media thickness in type-2 diabetes Mellitus with ischemic stroke. Indian J Endocrinol Metab. 2013;17:716-722.
6. Bag Bagshaw SM, Bellomo R, Jacka MJ, Egi M, Hart GK, George C, et al. The impact of early hypoglycaemia and blood glucose variability on outcome in critical illness. Crit Care. 2009;13:91.
7. Rossing P, Breum L, Major-Pedersen A, Sato A, Winding H, Pietersen A, et al. Prolonged QTc interval predicts mortality in patients with type 1 diabetes mellitus. Diabet Med. 2001;18:199-205.
8. VandeLoo A, Arendts W, Hohnloser SH. Variability of QT dispersion measurements in the surface electrocardiogram in patients with acute myocardial infarction and in normal subjects. Am J Cardiol. 1994;74:1113-1118.
9. Buja G, Miorelli M, Turrini P, Melacini P, Nava A. Comparison of QT dispersion in hypertrophic cardiomyopathy between patients with and without ventricular arrhythmias and sudden death. Am J Cardiol. 1993;72:973-976.
Yasar Sertbas et al.

10. Robinson RT, Harris ND, Ireland RH, Macdonald IA, Heller SR. Changes in cardiac repolarization during clinical episodes of nocturnal hypoglycaemia in adults with type 1 diabetes. Diabetologia. 2004;47:312–315.

11. Marfella R, Nappo F, De Angelis L, Siniscalchi M, Rossi F, Giugliano D. The effect acute hyperglycaemia on QTc duration in healthy man. Diabetologia. 2000;43:571–575.

12. Jung HS. Clinical Implications of Glycemic Variability: Chronic Complexes of Diabetes. Endocrinol Metab (Seoul). 2015;30:167-174.

13. Veglio M, Giunti S, Stevens LK, Fuller JH, Perin PC, EURODIAB IDDM. Complications Study Group. Prevalence of Q-T interval dispersion in type 1 diabetes and its relation with cardiac ischemia: The EURODIAB IDDM Complications Study Group. Diabetes Care. 2002;25:702-707.

14. Robinson RT, Harris ND, Ireland RH, Lee S, Newman C, Heller SR. Mechanism of abnormal cardiac repolarization during insulin induced hypoglycemia. Diabetes. 2003;52:1469-1474.

15. Marques JL, George L, Peacey SR, Harris ND, Maconald IA, Cochrane T, et al. Altered ventricular repolarization during hypoglycaemia in patients with diabetes. Diabet Med. 1997;14:648–654.

16. Christensen TF, Tarnow L, Randle J, Kristensen LE, Struik Jj, Eldrup E, et al. QT interval prolongation during spontaneous episodes of hypoglycaemia in type 1 diabetes: the impact of heart rate correction. Diabetologia. 2010;53:2036-2041.

17. Zhang Y, Han H, Wang J, Wang H, Yang B, Wang Z. Impairment of human ether-à-go-go-related gene (HERG) K+ channel function by hypoglycemia and hyperglycemia. Similar phenotypes but different mechanisms. J Biol Chem. 2003;278:10417–10426.

18. Nordin C. The case for hypoglycaemia as a proarrhythmic event: basic and clinical evidence. Diabetologia. 2010;53:1552–1561.

19. Gordin D, Forsblom C, Rönnback M, Groop PH. Acute hyperglycaemia disturbs cardiac repolarization in Type 1 diabetes Diabet Med. 2008;25:101-105.

20. Arslan A, Uzun M Does the lower nitric oxide level cause cardiovascular changes in major depressed women? Eur Rev Med Pharmacol Sci. 2008;12:309-313.

21. Tesfamariam B, Brown ML, Cohen RA. Elevated glucose impairs endothelium-dependent relaxation by activating protein kinase C. J Clin Invest. 1991;87:1643-1648.

22. Davis FB, Davis PJ, Nat G, Blas SD, MacGillivray M, Gutman S, et al. The effect of in vivo glucose administration on human erythrocyte Ca2+-ATPase activity and on enzyme responsiveness in vitro to thyroid hormone and calmodulin. Diabetes. 1985;34:639-646.

23. Christensen PK, Gall MA, Major-Pedersen A, Sato A, Rossing P, et al. QTc interval length & QT dispersion as predictors of mortality in patients with non-insulin-dependent diabetes. Scand J Clin Lab Invest. 2000;60:323-332.

24. Veglio M, Bruno G, Borra M, Macchia G, Bargero G, D’Errico N, et al. Prevalence of increased QT interval duration and dispersion in type 2 diabetic patients and its relationship with coronary heart disease: a population-based cohort. Intern Med. 2002;41:317-324.

25. Kumar R, Fisher M, Whitaker R, Macfarlane PW. Effect of controlling hyperglycaemia with diet on QT abnormalities in newly diagnosed patients with type 2 diabetes. Diabetes Care. 2004;27:2767-2768.

26. Li X, Ren H, Zhang-Rong X, Liu YJ, Yang XP, Liu JQ. Prevalence and risk factors of prolonged QTc interval among Chinese patients with type 2 diabetes. Exp Diabet Res. 2012;2012:234084. doi:10.1155/2012/234084