Mechanism of Electrocardiographic T-Wave Flattening in Diabetes Mellitus: Experimental and Simulation Study

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Summary

In the present study we investigated the contribution of ventricular repolarization time (RT) dispersion (the maximal difference in RT) and RT gradients (the differences in RT in apicobasal, anteroposterior and interventricular directions) to T-wave flattening in a setting of experimental diabetes mellitus. In 9 healthy and 11 diabetic (alloxan model) open-chest rabbits, we measured RT in ventricular epicardial electrograms. To specify the contributions of apicobasal, interventricular and anteroposterior RT gradients and RT dispersion to the body surface potentials we determined T-wave voltage differences between modified upper- and lower-chest precordial leads (T-wave amplitude dispersions, TWAD). Expression of RT gradients and RT dispersion in the correspondent TWAD parameters was studied by computer simulations. Diabetic rabbits demonstrated flattened T-waves in precordial leads associated with increased anteroposterior and decreased apicobasal RT gradients (P<0.05) due to RT prolongation at the apex. For diabetics, simulations predicted the preserved T-vector length and altered sagittal and longitudinal TWAD proven by experimental measurements. T-wave flattening in the diabetic rabbits was not due to changes in RT dispersion, but reflected the redistributed ventricular repolarization pattern with prolonged apical repolarization resulting in increased anteroposterior and decreased apicobasal RT gradients.

Key words

Diabetes • Dispersion of repolarization • Repolarization gradients • T-vector • T-wave

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Introduction

Diabetes mellitus (DM) is a recognized public health burden, characterized, among other complications, by increased risk of life-threatening ventricular arrhythmias and sudden cardiac death (Spooner 2008). An underlying functional arrhythmic substrate requires the presence of several conditions, including an increased dispersion of repolarization (DOR) defined as a maximal difference between repolarization times (RT) irrespective of lead sites within the ventricular myocardium. Relationship between RT differences and T-wave generation on the body surface ECG provides a rationale for using estimate electrocardiographic T-wave indices to heterogeneities of repolarization as well as arrhythmia susceptibility.

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Diabetic patients demonstrate changes in both QT interval duration and T-wave amplitude (Kittnar 2016). However, the electrophysiological basis for these alterations is not fully established. The mechanism of prolongation of rate-corrected QT intervals is relatively clear and is attributed to lengthening of action potential durations (Magyar et al. 1992). However, the mechanism of T-wave voltage changes in DM has not been directly studied. As T-wave genesis is usually attributed to ventricular repolarization gradients (RT differences on definite axes, such as apicobasal, interventricular, anteroposterior and transmural) (Meijborg et al. 2014), it is reasonable to suggest that flattening of the T-wave in diabetics results from RT redistribution within heart ventricles. However, such repolarization pattern could correspond to two conditions: either 1) DOR decreases but the proportion between RT gradients is the same; or 2) a predominant repolarization gradient changes but the DOR magnitude preserves. Distinguishing between these two alternatives is important, as the former is expected to affect the reentrant arrhythmogenesis, while the latter could reflect involvement of specific myocardial regions. Therefore, elaborating on how the different ventricular repolarization gradients and DOR contribute to the DM-related T-wave changes is needed for a comprehensive ECG interpretation.

T-wave amplitude parameters have been studied less extensively than temporal markers; however, evaluating T-wave amplitude could be a promising approach for assessing repolarization heterogeneity. Our previous study (Sedova *et al.* 2015) demonstrated that DOR changes were associated with T-wave amplitude dispersion (TWAD), defined as the difference in the T-wave peak voltages between upper- and lower-chest precordial leads. However, a concept of T-wave generation in a given condition is required for clearly interpretating its amplitude changes.

An objective of the present study was to determine the contribution of apicobasal, interventricular and anteroposterior repolarization gradients and DOR to body surface T-wave parameters in an experimental rabbit DM model. Previous investigations (Arteyeva *et al.* 2013, Meijborg *et al.* 2014, Arteyeva *et al.* 2015) suggested that transmural gradient contribution to generating T-waves was much less pronounced, if at all present, as compared to the contribution of other gradients. Therefore, here we did not address the transmural gradient, instead focusing studying the epicardial repolarization pattern.

Overview

We experimentally recorded (i) cardiac potentials on body and epicardial surfaces in control and diabetic rabbits; (ii) described differences between the groups in apicobasal, interventricular two and anteroposterior RT gradients; (iii) constructed x-, y-, and z- components of T-vector for the diabetics and controls on the basis of experimentally-measured RT gradients; (iv) simulated body surface potential distributions using these constructed T-vector components; (v) compared computed and measured body surface potentials in both groups; (vi) compared T-vector components derived from measured body surface potentials between control and diabetic groups; and (vii) tested if a relationship between these measured T-vector components is similar to the relationship between experimentally-determined RT gradients.

Experimental study

The experiments were performed on Chinchilla breed rabbits of either sex, age of 6-8 months and body mass of 2.5-4.0 kg. The procedures conformed to the *Guide for the Care and Use of Laboratory Animals, 8th Edition* published by the National Academies Press (US) 2011 and to the *Declaration of Helsinki* and were approved by the ethical committee of the Institute of Physiology of the Komi Science Centre, Ural Branch of the Russian Academy of Sciences. Type 1 DM was induced in 11 animals (7 males) by a single dose of alloxan (120 mg/kg body weight, i.v.) and 9 (6 males) animals served as controls.

After one month, DM and control rabbits were anesthetized with Zoletil (15 mg/kg body weight, i.m.) and put on mechanical ventilation. ECGs were recorded with subcutaneous needle electrodes from conventional limb leads and six modified precordial leads described elsewhere (Sedova et al. 2015). In short, there were three upper-chest leads (J1-J3) shifted to the jugular notch level, and three lower-chest leads (J4-J6) positioned at the inferior costal margin level. J1 and J6 leads and J3 and J4 leads were placed in the right and left anterior axillary lines, respectively; while J2 and J5 leads were in the midline. After ECG recording, the heart was exposed via midline incision, and heart temperature was maintained at 37-38 °C by irrigation with warm saline and warmed indoor air. At spontaneous sinus rhythm, unipolar ventricular electrograms were registered from a 64-lead

epicardial sock array (3-5 mm interelectrode distance) in reference to Wilson's central terminal with a customdesigned mapping system (16 bits; bandwidth 0.05 to 1,000 Hz; sampling rate 4,000 Hz).

In each epicardial unipolar electrogram, activation time (AT) and repolarization time (RT) were measured as the instants of the minimum of first time derivative of potential (dV/dt min) during QRS complex and the maximum of the first time derivative of potential (dV/dt max) during T-wave, respectively, in relation to QRS complex onset in the II limb lead (Coronel et al. 2006). The activation-recovery interval (ARI) serving as a measure of local repolarization duration was taken as the difference between RT and AT. Apicobasal, interventricular and anteroposterior RT gradients were calculated using average RT values in the appropriate epicardial leads (for example, apicobasal RT gradient equaled to the difference between the average basal RT and the average apical RT, etc, is in Fig. 1). Total DOR was determined as the difference between maximal and minimal RT values on the ventricular epicardium (Fig. 1). T-wave amplitude dispersion (TWAD) parameters were measured as the difference between T-wave voltages in body surface ECG leads (J1-J6) using formulas suggested for calculating T-vector components (see below).



Fig. 1. Schematic presentation of determining epicardial repolarization parameters. Each map displays anterior (left part) and posterior (right part) aspects of the heart. The margins between left and right ventricles (anterior and posterior interventricular grooves) are depicted by thin solid lines. (**A**) Lead distribution on epicardial surface. Dispersion of repolarization is calculated as the difference between the earliest and the latest repolarization times on the overall surface. Apicobasal (**B**), interventricular (**C**) and anteroposterior (**D**) repolarization gradients are determined as differences between the average RT values obtained from areas designated by small and large lead points.

Statistical analysis was performed using SPSS 11.5. Data are given as medians and interquartile intervals. The Mann-Whitney U-test was used to compare control and diabetic groups of animals. Student's paired and unpaired t-tests were used to assess intra- and interobserver variabilities, and agreement between measurements was evaluated with the Bland-Altman plot. Differences were considered significant at P<0.05.

T-vector components

Direct determination. In order to relate ventricular repolarization gradients to body surface ECG, we simulated the electrical vector of ventricular repolarization (T-vector) as a single dipole located in the center of the heart ventricles. T-vector direction was based on experimentally measured ventricular repolarization gradients: the component Tx was proportional to the interventricular gradient, the component Ty was proportional to the apicobasal gradient, and the component Tz was proportional to the anteroposterior gradient (in vectorcardiographic coordinate system X, Y, Z). A potential distribution produced by this T-vector was calculated for an elliptical cylinder imitating the torso of a rabbit, taking into account the heart's realistic position within the rabbit torso, using the following formula:

$$\varphi = (R \cdot T) / R^3 \tag{1},$$

where φ is a potential value in an observation point, *T* is T-vector, *R* is a vector directed from the observation point to the T-vector origin, R is the length of *R*. The computed potentials were compared with measured body surface ECGs.

Inverse determination. T-vector components were then also inversely calculated from body surface potentials. Since there is no generally accepted method of calculating VCG from ECG for the rabbit, we used the following empirical formulas based on differences in potential magnitude in modified precordial leads J1-J6 taking into account their mutual positions on the torso.

Longitudinal:
$$Ty = ((J6-J1) + (J4-J3)) / 2$$
 (2),
Left-to-Right: $Tx = ((J3-J1) + (J4-J6)) / 2$ (3),
Sagittal: $Tz = 10 \times ((J2-J1) + (J2-J3) + (J5-J4) + 2 \times (J5-J6)) / 4$ (4).

Results

Fasting plasma glucose concentration was significantly higher in the diabetic animals than in

controls [26.6 (17.8; 27.5) mmol/l and 5.7 (4.9; 6.0) mmol/l, P<0.05, respectively], although both groups were of the same sex, age, body mass and heart mass.

Body surface ECGs

The diabetic rabbits demonstrated flattened T-waves in modified upper-chest and lower-chest precordial leads (Fig. 2). T-wave amplitudes in individual precordial leads were lower in the DM group than in the control group (Table 1). Longitudinal, left-to-right and sagittal TWAD indices calculated using formulas (2-4) (suggested for the corresponding T-vector components) differed between the control group and the DM group. The diabetic rabbits had greater TWAD_{sagittal} and lower TWAD_{longitudinal} than controls (Table 1).



Control

Diabetes mellitus

Fig. 2. Representative ECG tracings in modified precordial leads J1-J6 (solid black points) in control and diabetic rabbits. The heart position is indicated by thick arrows depicting predominant apicobasal and anteroposterior RT gradients for non-diabetic and diabetic animals, respectively.

Table 1. Cardiac and body surface ECG indices of ventricular repolarization in healthy and diabetic rabbits [Median and interquartile intervals (25 %, 75 %)].

Parameter	Control, n=9	DM, n=11	Р
Apicobasal RT gradient (ms)	17 (10; 18)	2 (-7; 4)	0.002
Interventricular RT gradient (ms)	-6 (-12; 0.2)	7 (-0.3; 10)	0.087
Anteroposterior RT gradient (ms)	-7 (-12; 9)	11 (6; 17)	0.016
DOR (ms)	51 (43; 57)	56 (41; 75)	0.351
Maximal lower-chest T-wave voltage (μV)	162 (144; 226)	119 (0; 143)	0.020
Minimal upper-chest T-wave voltage (μV)	-242 (-266; -212)	-170 (-218; -121)	0.052
$TWAD_{Longitudinal} (\mu V)$	197 (120; 236)	105 (88; 158)	0.023
$TWAD_{Left-to-Right} (\mu V)$	64 (53; 84)	45 (18; 66)	0.119
$TWAD_{Sagittal} (\mu V)$	58 (-144; 355)	-210 (-303; -95)	0.045
$TWAD_{Sum}(\mu V)$	338 (234; 471)	282 (178; 428)	0.340

DM – diabetes mellitus; RT – repolarization time; DOR – dispersion of repolarization; TWAD – T-wave amplitude dispersion.

Epicardial potential mapping

AT measurements were done automatically, while determining RTs included inspection hv experienced observers (AOO, MAV). Therefore, the data were tested for inter- and intraobserver reproducibility. Sample measurements (n=83) done by the same observer on different days demonstrated no significant differences in RT interval (values are given for the combined group of healthy and diabetic animals): 186.6±32.1 vs. 185.42±32.9 ms, p=0.91 and S.D. of the difference was 18.6 ms or 9.9 % of the mean. The difference between the first and the second observer's data was also insignificant: 186.1±38.8 vs. 187.0±40.6 ms, p=0.44 and S.D. of the difference 10.8 ms or 5.8 % of the mean.

At one-month follow-up, rabbits with DM and controls had similar sequences and durations of ventricular epicardial activation, but the epicardial repolarization pattern differed between the two groups (Fig. 3). In control animals, repolarization durations (measured as ARIs) were shorter at the apex and longer at the base manifesting in a dominant apicobasal RT gradient in respect to interventricular and anteroposterior gradients (Table 1). In diabetic animals, repolarization durations were redistributed with an area of prolonged ARIs development mainly on the anterior apical portion (mainly, the heart apex and the adjacent right ventricular area). This effect resulted in a pattern with decreased apicobasal and increased anteroposterior RT gradients in the diabetic hearts (Table 1). Despite the differences in ventricular repolarization gradients, there was no difference in total DOR between the control and DM groups.

Computer simulations

We simulated body surface potential distributions for both non-diabetic and diabetic rabbits (Fig. 4) according to experimentally-measured epicardial RT gradients. For the control rabbit, the T-vector was oriented forward, downward and to the left, with a dominant longitudinal component. For the diabetic rabbit, T-vector was oriented backward, downward and to the left, the longitudinal component was decreased, and the anteroposterior component was dominant. T-vector length was the same because total RT dispersion in DM was found to be close to that of non-diabetic rabbits.

The different T-vector directions in non-diabetic and diabetic animal models resulted in different simulated body surface potential distributions (Fig. 4). In the points corresponding to leads J1-J6, potential magnitudes decreased, but their polarity remained unchanged. These changes were similar to the changes observed in leads J1-J6 in diabetic rabbits (Fig. 2). T-vector directions calculated from the differences in measured potential magnitude in leads J1-J6 were similar to T-vector directions set in the model on the basis of measured RT gradients (Fig. 4).



Fig. 3. Representative maps of epicardial ventricular distribution of ARIs in control and diabetic (DM) rabbits. See the prolonged repolarization area on the anterior apical portion in the diabetic rabbit heart.

Discussion

In order to explain the T-wave changes in DM, we first studied the spatial patterns of ventricular repolarization, and then reconstructed the expression of these cardiac repolarization patterns in body surface potentials using computer simulations. Finally, we tested the predictions of simulations by comparing measured body surface potential distributions in non-diabetic and diabetic animals.

Fig. 4. Simulated body surface potential maps during T-wave peak for the control and diabetic rabbits. Simulated voltages in J1-J6 leads are shown. The transversal and frontal projections of the T-vectors set directly (bold gray) and calculated from the TWAD (black) are displayed to the right of each map.

Our primary objective was to find out the DM-related T-wave flattening mechanisms as they relate to changing ventricular gradients of repolarization. We found that the decrease in T-wave amplitudes in modified precordial leads was associated with altering two repolarization gradients, namely: 1) increase in the anteroposterior gradient with early posterior and late anterior RTs, and 2) decrease in the apicobasal gradient. These two differences between non-diabetic and diabetic rabbits were accounted for by repolarization duration prolongation at the RV apex, located mostly anteriorly. specific involvement of RV in diabetic The cardiomyopathy was reported earlier (Karamitsos et al. 2007, van den Brom et al. 2010, Vaykshnorayte et al. 2012, Olsen et al. 2013, Axelsen et al. 2015). Our study suggests that T-wave flattening was the electrocardiographic expression of the RV electrophysiological alteration, at least in the present DM model. The repolarization prolongation in DM is usually ascribed to the down-regulation of potassium currents (Zhang et al. 2007, Lengyel et al. 2008, Gallego et al. 2009); however, the exact mechanism of such spatially specific electrophysiological effect is largely unknown. Previous studies (Palova et al. 2010, Ovechkin et al. 2014) showed that an abnormal sympathetic tone possibly plays a role.

Thus, a redistributed spatiotemporal ventricular repolarization pattern in diabetic rabbits was associated with flattened body surface T-waves. Theoretically, the observed changes in body surface T-wave voltages associated with DM could be ascribed to three different myocardial alterations, namely: 1) decrease of the apicobasal RT gradient; 2) increase of the anteroposterior RT gradient; and 3) replacement of the apicobasal RT gradient by the anteroposterior RT gradient as dominant RT gradient. In the а context of arrhythmogenesis, it is important to note that these scenarios implied decreased, increased or unchanged DOR, respectively. Thus, further specification of the contribution of the different repolarization gradients to body surface potentials is required. As the next step of the present study, we performed computer simulations in order to test if T-wave voltage parameters could distinguish between the different variants of ventricular repolarization changes in DM.

Despite the *de facto* disappearance of vectorcardiography from routine clinical practice, vectorial characteristics of the repolarization cardiac electric field derived from ECG records are considered useful for diagnosis and prognosis (Man *et al.* 2015). Such indices could be obtained by reconstructing a vectorcardiogram from ECGs (Schreurs *et al.* 2010, Engels *et al.* 2015). The resultant T-vector is a characteristic related to the vectorcardiographic loop, and is considered to be a simple and useful representation of the ventricular repolarization process (Waks *et al.* 2015, Cortez *et al.* 2016), where its direction reflects the



gross repolarization sequence, its length (magnitude) is proportional to the resultant intensity of electric generators in the heart ventricles (indirectly total DOR), and its apicobasal, interventricular and anteroposterior components are related to the apicobasal, interventricular and anteroposterior RT gradients, respectively. The modified precordial leads J1-J6 were positioned on the body surface so that the heart was the center of symmetry, which allowed us to easily obtain the relationship between the apicobasal and interventricular T-vector components and the correspondent longitudinal and left-to-right TWAD indices. Since the heart is located within the thorax anteriorly, and the distance between the medial and lateral leads on the sagittal projection axis is small, scaling coefficients were required for estimating the sagittal T-vector component.

As the diabetic rabbits had the redistributed ventricular repolarization pattern with the increased anteroposterior and decreased apicobasal RT gradients, and, consequently altered T-vector components, we were able to test our approach to estimating the T-vector components. We found that the longitudinal TWAD was lower and the sagittal TWAD was higher in the DM group than in the control group, suggesting that TWADs reflected the RT gradients in the correspondent directions, and could therefore be used for assessing the electrophysiological properties in specific myocardial regions. Our data suggest that T-wave changes in DM rabbits reflected RV involvement. However, our epicardial measurements showed similar total DOR in non-diabetic and diabetic animals; accordingly, the TWAD_{sum}, which was suggested as an estimate for T-vector length, and indirectly for total DOR, was no different for the DM and control groups.

Our findings demonstrated that precordial T-wave voltage decrease could be associated with unchanged DOR and oppositely changed different RT gradients. Measuring temporal ECG indices of ventricular repolarization, such as QT and T_{peak}-T_{end} intervals or their dispersions for assessing DOR and predicting arrhythmic events often give unsatisfactory results (Zabel et al. 1998, Porthan et al. 2013), due at least in part to technical problems with determining T-wave end. In our study, we could not measure QT and T_{peak}-T_{end} intervals in a number of animals because of overlapped and T-waves. However, T-vector components Р specifically express the corresponding repolarization gradients, contain useful information on DOR, and may thus be tested as predictors of ventricular arrhythmias.

Limitations

We intended to establish the relationship between diabetic electrical myocardial remodeling and T-wave changes in body surface ECGs quantified as magnitudes of T-vector components to be further used as a noninvasive predictor of functional disturbances. From this point of view, several issues should be taken into account. Where there is attenuation of T-wave signals due to subcutaneous fat, a correction procedure may be required. It is also expected that TWAD_{sum} (T-vector magnitude) may differ somewhat, with DOR remaining unchanged, in settings where action potential durations uniformly prolong or shorten, such as in electrolyte disturbances. We demonstrated that TWAD reflected the magnitude of the RT gradients being the differences in RTs between the definite ventricular regions, such as the apex and the base, etc. However, we did not test local electrophysiological heterogeneities, which may affect the DOR, but perhaps not affect the RT gradients.

Conclusions

Our study suggests that T-wave flattening in modified precordial leads in diabetic rabbits was associated with repolarization prolongation in the apical portion of the RV myocardium, resulting in decreased apicobasal gradient and increased anteroposterior gradient of RTs. These findings provide the basis for assessing RV functional changes in DM. The contribution of apicobasal, interventricular and anteroposterior repolarization gradients could be estimated by calculating the correspondent TWAD parameters. In diabetic rabbits, we found two associations: 1) the augmented anteroposterior RT gradient and the increased sagittal TWAD; 2) the diminished apicobasal RT gradient and the decreased longitudinal TWAD. The obtained results show that T-wave flattening does not necessarily imply changes in DOR, which in turn could be assessed by T-vector length and TWAD_{sum}.

Conflict of Interest

There is no conflict of interest.

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