Ventricular Electrical Heterogeneity in Experimental Diabetes Mellitus: Effect of Myocardial Ischemia

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Summary
Aims of the study were to compare the development of electrocardiographic responses of the ischemia-induced heterogeneities of activation and repolarization in the ventricular myocardium of normal and diabetic animals. Body surface ECGs and unipolar electrograms in 64 epicardial leads were recorded before and during 20 min after the ligation of the left anterior descending artery in diabetic (alloxan model, 4 weeks, n=8) and control (n=8) rabbits. Activation times (ATs), end of repolarization times (RTs) and repolarization durations (activation-recovery intervals, ARIs) were determined in ischemic and periischemic zones. In contrast to the controls, the diabetic rabbits demonstrated the significant prolongation of ATs and shortening of ARIs (P<0.05) during ischemia in the affected region resulting in the development and progressive increase of the ARI and RT gradients across the ischemic zone boundary. The alterations of global and local dispersions of the RTs in diabetics correlated with the T peak-Tend interval changes in the limb leads ECGs. In the ischemic conditions, the diabetic animals differed from the controls by the activation delay, significant repolarization duration shortening, and the increase of local repolarization dispersion; the latter could be assessed by the Tpeak-Tend interval measurements in the body surface ECGs.

Key words
Cardiac electrophysiology • Diabetes mellitus • Regional ischemia • Myocardium • Electrocardiography

Introduction
The excessive myocardial electrical heterogeneity dictates the level of electrical instability of the heart, whereas the myocardial ischemia is a major event exacerbating electrical inhomogeneities and leading to malignant ventricular arrhythmias. Clinical studies have found that diabetes mellitus (DM) enhances the risk of ventricular tachyarrhythmias and mortality in acute myocardial infarction even more (Cho et al. 2002, Dziewierz et al. 2010, Sanjuan et al. 2011). On the other hand, experimental studies of diabetic hearts demonstrated either the increased (Hekimian et al. 1985, Bakth et al. 1986, Wang et al. 2012) or decreased (Kusama et al. 1992, Ravingerová et al. 2000, Galagudza et al. 2007, Matejiková et al. 2008) susceptibility to ventricular arrhythmias in ischemic conditions. Spooner (2008) pointed out that DM while worsening the long-term prognosis does not affect the susceptibility to ventricular arrhythmias in acute settings in patients with cardiovascular diseases. Some clinical observations suggested that the susceptibility to ventricular arrhythmias could be related to concomitant factors such
as hypoglycemic episodes or medication (Chow et al. 2014, Curione et al. 2014, Pistrosch et al. 2015). These conflicting data on the electrical stability of the heart in diabetes mellitus under ischemia suggest that the diabetic myocardium may have a specific electrophysiological response to the ischemic insult.

The diabetic cardiomyopathy is associated with the electrophysiological alterations in the myocardium. The DM-related changes in the cardiac electrical properties, specifically the prolongation of action potential durations, have been well documented at the cellular level (Magyar et al. 1992, Zhang et al. 2007, Lengyel et al. 2008, Gallego et al. 2009). Equally important would be largely lacking thus far data on the distribution of the electrophysiological properties throughout the myocardium and its dynamical changes in ischemia. This spatiotemporal electrical pattern is quantitated as a dispersion of repolarization that is, in turn, attempted to be assessed by ECG. For this purpose, several ECG indices were considered, including Tpeak-Tend interval, a promising index for the estimation of the dispersion of repolarization. Clinical investigations either supported (Panikkath et al. 2011, Hetland et al. 2014, Mozos 2015) or opposed (Smetana et al. 2011, Porthan et al. 2013) to its prognostic utility suggesting that electrophysiological information content of Tpeak-Tend interval could vary in different conditions. Tpeak-Tend interval has been also tested in diabetic patients (Clemente et al. 2012, Miki et al. 2014). However, the behavior of the Tpeak-Tend and its relation to the dispersion of repolarization in an arrhythmogenic stress conditions, such as myocardial ischemia, is largely unknown in the setting of DM.

The objective of the present study was to compare the ischemia-induced changes of ventricular epicardial activation and repolarization patterns and their expressions in the parameters of the body surface ECGs in normal and diabetic rabbit hearts.

Methods

The experiments were carried out in adult Chinchilla rabbits (age from 7 to 9 months, body mass from 2.9 to 3.3 kg). The investigation conformed to the Guide for the Care and Use of Laboratory Animals (2011). Experimental type 1 DM was induced in 8 animals (5 females) by a single intravenous alloxan injection (120 mg/kg body mass, 4 weeks follow-up), and 8 healthy animals (4 females) served as controls. DM was confirmed by at least double determinations of fasting venous blood glucose level more than 7 mmol/l with OneTouch glucometer (LifeScan Inc, USA). The open-heart experiments were consistently done during the daytime hours (from 11-00 to 13-00) on the animals anesthetized with zoletil (15 mg/kg body mass, intramuscular injection), intubated and mechanically ventilated. The heart was exposed by a midsternal incision. The temperature of the heart was maintained at 37-38 °C by the irrigation with warm saline and warming the room air. An electrode sock with 64 leads (3-5 mm interelectrode distance) was placed on the ventricular surface, and the unipolar electrograms were simultaneously recorded in reference to Wilson’s terminal at spontaneous sinus rhythm. The data acquisition was done using a custom-designed mapping system (16 bits; bandwidth 0.05 to 1000 Hz; sampling rate 4000 Hz).

Electrograms were recorded in the baseline and in the course of 20-min ischemia, which was produced by the ligation of the left anterior descending coronary artery (LAD). Evans blue dye (Sigma-Aldrich GmbH, Germany, 0.5 %) was injected postmortemly into the aorta. The ischemic zone was determined by the absence of Evans blue perfusion, and its size was estimated as the area of the figure circumscribed by the leads on the border of the ischemic zone.

Limb lead ECGs were monitored in the course of the experiment, and the QRS and QT durations were measured in the limb lead II. The corrected QT interval was calculated by the equation QTc=QT-0.175×(RR-300) (Carlsson et al. 1993). The total duration of the Tpeak-Tend interval was measured as a period between the earliest Tpeak and the latest Tend in the limb leads. In each epicardial lead, the activation time (AT) and the end of repolarization time (RT) were determined as dV/dtmin during the QRS complex and dV/dtmax during the T wave, respectively (Coronel et al. 2006). The activation-recovery interval (ARI) serving as a measure of local repolarization duration was measured as the difference between the RT and AT. The averaged values of these variables were calculated for different myocardial regions and specifically for the ischemic zone and the adjacent 3 to 5 mm width band of perfused myocardium referred to as a periischemic zone. The differences in the ATs, ARIs, and RTs between the ventricular regions were referred to as gradients (e.g. boundary gradient) and the difference between the earliest and the latest RT values throughout the ventricular epicardium was referred to as...
the dispersion of repolarization.

Statistical analysis was performed with the SPSS 11.5 software packages. The data are given as medians and interquartile intervals. Wilcoxon test and Fridman test with Dunnett post-hoc procedure were utilized for the single and multiple comparisons within the groups, respectively. Mann-Whitney U-test was used to compare different groups of animals. The correlation between epicardial mapping indices and ECG parameters was evaluated with Spearman rank correlation test as the evaluated parameters were not normally distributed. The differences were considered significant at P<0.05.

**Results**

The control and DM groups matched with each other for sex, age, body mass and heart to body mass ratio. As expected, the glycemia level was significantly higher in diabetic animals as compared to controls [15.3 (8.8; 20.0) mmol/l vs. 5.9 (5.7; 6.3) mmol/l, respectively, P<0.01]. RR, QT, QTc and Tpeak-Tend intervals were longer in animals with DM, whereas the QRS duration was similar in both groups (Table 1). The epicardial AT sequences in normals and diabetics were similar and directed from the left ventricular (LV) apex to the LV base and from the right ventricular (RV) free wall to the LV free wall. The spatial distribution of RTs was relatively uniform in healthy animals and heterogeneous in diabetics. In the latter group, the RV RTs were longer than the LV RTs [DM: 134 (119; 142) ms vs. 106 (104; 114) ms, P=0.012; Control: 115 (106; 127) ms vs. 114 (99; 119) ms, P>0.05, respectively].

<table>
<thead>
<tr>
<th>RR, ms</th>
<th>QRS, ms</th>
<th>QT, ms</th>
<th>QTc, ms</th>
<th>Tpeak-Tend, ms</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Baseline</strong> control</td>
<td>230 (220; 247)</td>
<td>33 (31; 36)</td>
<td>159 (147; 160)</td>
<td>163 (159; 171)</td>
</tr>
<tr>
<td>DM</td>
<td>269 (249; 282)</td>
<td>32 (30; 35)</td>
<td>182 (171; 186)</td>
<td>188 (180; 190)</td>
</tr>
<tr>
<td>P</td>
<td>0.03</td>
<td>ns</td>
<td>0.003</td>
<td>0.009</td>
</tr>
<tr>
<td><strong>10' ischemia</strong> control</td>
<td>229 (209; 247)</td>
<td>34 (30; 38)</td>
<td>147 (139; 171)</td>
<td>160 (153; 180)</td>
</tr>
<tr>
<td>DM</td>
<td>252 (239; 264)</td>
<td>32 (31; 36)</td>
<td>160* (151; 169)</td>
<td>170* (160; 175)</td>
</tr>
<tr>
<td>P</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
</tr>
<tr>
<td><strong>20' ischemia</strong> control</td>
<td>225 (205; 240)</td>
<td>33 (31; 39)</td>
<td>147 (134; 166)</td>
<td>160 (151; 177)</td>
</tr>
<tr>
<td>DM</td>
<td>252 (243; 267)</td>
<td>33 (30; 38)</td>
<td>155* (148; 168)</td>
<td>164* (157; 173)</td>
</tr>
<tr>
<td>P</td>
<td>ns</td>
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* P<0.05 vs. baseline; DM, diabetes mellitus; ns, nonsignificant.

The size of the ischemic area was similar in the diabetic and control animals [75.6 (56.5; 86.0) mm² vs. 55.3 (42.4; 63.5) mm², for control and DM animals, respectively, P>0.05]. Isolated premature ventricular beats were sporadically observed in both groups. During ischemia, the QT and QTc intervals shortened in the DM group (Table 1). Coronary occlusion induced changes in electrograms recorded from the ischemic zone (Fig. 1), but the alterations found in this area differed in the control and diabetic animals (Fig. 2). The statistically significant activation delay in the ischemic zone was found only in diabetics (P<0.05). At 20-min of coronary occlusion, there were no statistically significant effects on ARIs in the control group. At least in part, this could be ascribed to variable individual profiles of ARI changes in the control group, which demonstrated either ARI shortening or prolongation, while the consistent shortening was found in the DM group (Fig. 3).
Ischemia induced the development of specific epicardial activation and repolarization patterns (Fig. 4) characterized by the abrupt spatial differences in ATs and RTs corresponding to the area of ischemia (Fig. 4). Such differences were more pronounced in the DM group as compared to controls. The substantial ARI shortening in diabetics resulted in the development of a boundary ARI and RT gradients between the ischemic and adjacent periischemic zones, which progressively increased during the ischemic episode (Table 2). A significant correlation was found between the $T_{peak}$-$T_{end}$ interval duration and the global RT dispersion in the diabetic rabbits under 20-min ischemia ($r=0.86$, $P=0.007$). In turn, the global RT dispersion was associated with the magnitude of the boundary RT gradient ($r=0.857$, $P=0.007$) which accordingly explains the correlation between the $T_{peak}$-$T_{end}$ interval duration and the boundary RT gradient found only in the DM group ($r=0.714$, $P=0.047$).

**Table 2.** The changes of activation time gradients, activation-recovery interval gradients and end of repolarization time gradients between ischemic and periischemic (i/p) myocardium in control and DM groups [Median and interquartile intervals (25 %, 75 %)].

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>DM</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>AT i/p gradients</td>
<td>ARI i/p gradients</td>
</tr>
<tr>
<td><strong>Baseline</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>future</td>
<td>0.24(-2.1;0.8)</td>
<td>5.1(0.1;13.2)</td>
</tr>
<tr>
<td>ischemic zone</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1′ ischemia</td>
<td>-0.27(-1.1;1.8)</td>
<td>2.8(-2.2;7.6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10′ ischemia</td>
<td>0.4(-1.0;2.5)</td>
<td>0.6(-12.5;6.0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>20′ ischemia</td>
<td>0.4(-0.7;2.6)</td>
<td>-1.2(-12.2;4.3)</td>
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* $p<0.05$ for Dunnett post-hoc test (vs. baseline), † $p<0.05$ for Mann-Whitney (vs. control).
Fig. 3. Individual ARI changes (%) in the periischemic and ischemic zones during the ischemic episode in the control and diabetic animals. The baseline value is 100%. Dashed lines identify for the means.

Fig. 4. Representative isochronal maps of activation and end of repolarization times in the control and diabetic animals at 20 min of coronary occlusion that produced the regional ischemia (arrows). Numbers on the scales indicate the time elapsed from the QRS onset. The left, and right sides of each map correspond to the anterior and posterior surfaces of the ventricles, respectively. The greater contrast and “denser” isochrones at the margins of ischemic area demonstrates the greater effects ischemia produced in the diabetic animals. DM, diabetes mellitus.
Discussion

The prolonged QTc found in the diabetic rabbits is consistent with our previous observations (Vaykshnorayte et al. 2012, Ovechkin et al. 2015) and data obtained in other animal species (Howarth et al. 2011) or humans (Zákovičová et al. 2014). This suggested that the average ventricular repolarization duration prolonged, presumably due to the downregulation of potassium repolarizing currents (Magyar et al. 1992, Zhang et al. 2007, Lengyel et al. 2008, Gallego et al. 2009). The observed interventricular RT difference implied that the action potential duration lengthening should be heterogeneous in ventricular myocardium and be predominantly expressed in the RV at least at 1-month follow-up from the DM induction. Likewise, the shortening of QT and QTc during the ischemic exposure was consistent with the repolarization duration shortening in the DM group.

During ischemia, ATs and ARIs did not change significantly in the control group, and the marked interindividual variations in the ischemia-induced ARI response were observed. It is noteworthy that several control animals demonstrated the prolongation of ARIs. Verkerk et al. (1996) showed that more than a half of rabbit cardiomyocytes demonstrated phasic prolongation of action potential duration in ischemic conditions due to Ito inhibition. This initial prolongation was followed by the shortening of repolarization presumably due to the increase of I_{KATP}. The similar reaction was observed in human cells, which makes these observations clinically relevant. The above considerations could explain the varied ARI response to ischemia observed in the control group. As a result, the slight changes in activation and the variable alterations of repolarization caused the RT dispersion in the controls to be statistically unchanged. Therefore, the difference between diabetics and nondiabetics in the repolarization response to ischemia was possibly caused by the different relationships between I_{Na}-dependent prolongation and I_{KATP}-dependent shortening. This concept remains to be further evaluated.

On the other hand, the diabetics demonstrated a significant though small AT delay in the ischemic zone. This effect was not expressed in the QRS prolongation but could reflect the local changes of the depolarization process, for example, the reduced I_{Na} in the diabetic rabbit myocardium (Stables et al. 2014). In contrast to the control group, repolarization durations significantly shortened in the ischemic zone that was associated with the shortening of QT and QTc intervals on the body surface ECG. The difference between the controls and diabetics in the reaction of the repolarization duration to ischemia could be due to the down-regulation of I_{Na} current in the diabetic animals (Gallego et al. 2009). The inhibition of this current was shown to be responsible for the transient action potential duration prolongation in ischemia (Verkerk et al. 1996). Accordingly, such an effect was not observed in the DM group. The effects of ATs and ARIs oppositely influenced the local RTs thus reducing their change. As the magnitude in milliseconds of the ARI decrease was greater than the AT increase, the boundary RT heterogeneities developed in the diabetic myocardium at ischemia.

Since the diabetic rabbits differed from the normals in their electrophysiological response to ischemia, the abovementioned alterations of ventricular repolarization were likely to be differently expressed in the ECG indices in the control and DM groups. Theoretically, a correlation between the global RT dispersion and the T_{peak}-T_{end} interval duration could be expected (Arteyeva et al. 2013) and such a correlation was indeed found, but only in diabetics during the ischemic exposure. This observation at least in part could be explained by the fact that the global RT dispersion in the ischemic diabetic myocardium was significantly correlated with the newly developed boundary RT gradient absent in the control group and the DM group before exposure. Our further observation of the correlation between the T_{peak}-T_{end} interval duration and the boundary RT gradient in diabetics under ischemia supports the above explanation. Furthermore, it suggests that the measurements of the T_{peak}-T_{end} interval could be more effective in the assessment of the local ischemic electrical heterogeneities in a subset of patients with DM as compared to the nondiabetics.

Limitations

The limitations of the present study concerned at least the short follow-up of DM and the short ischemia exposure. It is not excluded that the longer ones could lead to the more pronounced electrophysiological effects and possibly the spontaneous arrhythmia incidence. Repolarization parameters are often reported to be subject to sex differences. In the present study, we could not specify the effects in males and females. However, the diabetic and control groups did not differ significantly from each other in the male-to-female ratio, and therefore we believe that the observed results could not be
attributed to gender effects. Anesthesia could have affected the myocardial electrophysiological properties; however, we believe that these effects should be similar in controls and diabetics and should not significantly modify the major findings of the study.

**Conclusion**

Thus, the more pronounced ischemia-related prolongation of activation and shortening of repolarization in diabetic animals presumably due to the $I_{Na}$ and $I_{to}$-down-regulation in DM led to the increase of local electrical inhomogeneities that in turn could be assessed by the $T_{peak}$-$T_{end}$ interval. The findings of the present study suggest that the $T_{peak}$-$T_{end}$ interval can be of different diagnostic utility in different pathological conditions.

**Conflict of Interest**

There is no conflict of interest.

**Acknowledgements**

The study was supported by the Ural Branch of the Russian Academy of Sciences (Project 13-4-032-KSC) and the Russian Foundation for Basic Research (Grant 14-04-31070, young_a).

**References**


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