The impact of type 2 diabetes on the cardiovascular system in male and female rats*

O. V. Ivanova

SI «V. Danilevsky Institute for Endocrine Pathology Problems of the NAMS of Ukraine», Kharkiv, Ukraine

ivolga3006@ukr.net

Type 2 diabetes mellitus (T2DM) is a non-communicable and the one of the fastest growing disease that affects millions of people worldwide. Compared to the general population, individuals with type 2 diabetes are at an increased risk of cardiovascular disease (CVD) [1]. While CVD is a leading cause of death in both women and men, accumulating evidence suggest that biological sex is a major determinant for the development and progression of CVD.

Women with T2DM have been reported to have significantly higher risks of both fatal and non-fatal coronary heart disease and stroke than men with diabetes [2–6]. The Framingham Heart Study first revealed that diabetic women have a 5.1-fold increase in heart failure, while diabetic men only have a 2.4-fold increase compared to non-diabetic women and men, respectively [7].

It is hypothesized that diabetes mellitus (DM) attenuates the general female biological advantage of protecting against cardiovascular complications. These mechanisms include the altered systemic glucose regulation and lower insulin sensitivity, more pronounced hypertension, elevated triglycerides, inflammation and higher prothrombotic profile in females compared to males [8]. Sex differences in body composition and fat distribution may be involved in prevalence of over-weight and obesity in women and therefore increase CVD risk [9].

Additionally, estrogen-related signalling within the female myocardium plays important role both in protection and in progression of DM and its complications. Many risk factors probably contribute discrepancies in cardiovascular complications between men and women [9].

Moreover, diabetic females feature greater susceptibility to diabetic cardiomyopathy then

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males due to more pronounced cardiac structural remodelling and development of diastolic dysfunction. Furthermore, female sex is the only independent predictor of left ventricular hypertrophy [10].

However, how T2DM modulates this effect in the heart tissues of women versus men is currently unclear. Future research leading to determination of these mechanisms may contribute to sex-specific treatment for diabetic macro-vascular disease.

The aim of this study was to determine the impact of type 2 diabetes on the functional state of the cardiovascular system in male and female rats.

**MATERIALS AND METHODS**

The present study was approved by the bioethics committee of the V. Danilevsky Institute of Endocrine Pathology Problems, National Academy of Medical Sciences of Ukraine (Kharkiv, Ukraine) and performed in accordance with the European Convention for the Protection of Vertebrate Animals Used for Experimental and Other Scientific Purposes (Strasbourg, 1986).

The experiments were performed on Wistar rats, which were housed in Plexiglas cages at a temperature of (22 ± 1)°C, in a constant 12-hour light/dark cycle.

The animal model of type 2 diabetes (T2D) induced by a high-caloric diet, combined with multiple low-dose streptozotocin (STZ) injections, was used. The control intact groups (n = 6) were fed a standard diet ad libitum during 14 weeks. The experimental groups (n = 6) were fed the high-caloric diet, containing 16 % fat, 28% carbohydrates, 6% proteins for 14 weeks. All groups had free access to water. In four weeks, rats in the experimental group were injected intraperitoneally with small doses of STZ (25 mg/kg body weight) twice per week [11].

At the end of the study electrocardiograms were recorded in leads II using the electrocardiography method. The mean R-R, Q-T, P-Q, QRS, T-P intervals and the amplitude and duration of T, P and R waves were calculated.

Data normality were rated using the Shapiro-Wilk test, and all normally distributed data are expressed as the means ± standard error of the mean (SEM). Group comparisons of quantitative variables were performed by one-factor analysis of variance (ANOVA). The Newman-Keuls test was used for multiple comparisons of the groups. Values were considered significant at p < 0.05.

**RESULTS AND DISCUSSION**

As shown in Table 1, the basal glucose level in diabetic rats was significantly higher in comparison with control rats. In addition, the level of basal hyperglycaemia in diabetic animals was independent from sex.

In our study T2D caused an increase in heart rate, as indicated by the shortening of R-R interval, in both female and male rats in comparison with intact control groups (see table 2). The results indicate the development of pathologically accelerated rhythm and sinus tachycardia in experimental animals of both sexes.

It has been established that T2D leads to the prolongation of electrical systole occurs as the result of an increase in action potential duration in both males and females, as evidenced by an increasing of the total length Q-T intervals, compared to control group’s data (see table 2).

<table>
<thead>
<tr>
<th>Index</th>
<th>Control group</th>
<th>Diabetic group</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>male</td>
<td>female</td>
</tr>
<tr>
<td>Basal glucose level, mmol/l</td>
<td>5.00 ± 0.16</td>
<td>4.27 ± 0.20</td>
</tr>
</tbody>
</table>

Data are shown as mean ± standard error of the mean (SEM). * significant versus match control group (p < 0.05).
It was found that the duration of P-Q interval reflected the signal conduction from the sinus to the contractile ventricular myocardium, was probably higher in both intact females and T2D females versus intact and diabetic males (see table 2). However, T2D did not change the duration of P-Q interval in animals of both sexes, indicating that there is no influence of this model of diabetes on the atrio-ventricular conduction.

Our results have shown that T2D promotes the development of diastolic dysfunction in females, in contrast to males. Our study clearly points out the signs of diastolic dysfunction such as prolongation of diastolic T-P interval and a decrease of amplitude and duration of the T wave, which reflects the processes of ventricular myocardial repolarization, in comparison with intact female rats (see table 2 and table 3).

The findings are referred to clinical data on the higher prevalence of diastolic dysfunction and/or left ventricular hypertrophy, as a structural and functional component of diabetic cardiomyopathy, in women than in men [12].

T2D was accompanied by multidirectional impact on the impairment of atrial activation in heart of both sexes rats. Thus, females were showed an increase of the P wave’ amplitude compare to intact control, which may be explained by the development of hypertrophic processes in the atria. Decreased the P wave’ amplitude was noted in males possibly due to the progress of ischemia in the myocardium (see table 3).

It has been established that duration of the R-wave doesn’t change in any of the experimental groups, but R-waves voltage is decreased only in male rats with T2D in comparison with

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**Table 2**

<table>
<thead>
<tr>
<th>Indexes</th>
<th>Control group</th>
<th>Diabetic group</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>male</td>
<td>female</td>
</tr>
<tr>
<td>R-R interval, msec</td>
<td>129.2 ± 1.7</td>
<td>131.1 ± 1.6</td>
</tr>
<tr>
<td>Heart rate, Hz</td>
<td>468.1 ± 5.6</td>
<td>459.0 ± 5.7</td>
</tr>
<tr>
<td>Q-T interval, msec</td>
<td>56.1 ± 4.1</td>
<td>58.1 ± 2.2</td>
</tr>
<tr>
<td>P-Q interval, msec</td>
<td>34.2 ± 1.1</td>
<td>40.3 ± 2.1 b</td>
</tr>
<tr>
<td>QRS interval, msec</td>
<td>39.4 ± 1.2</td>
<td>28.4 ± 2.3</td>
</tr>
<tr>
<td>T-P interval, msec</td>
<td>70.3 ± 3.2</td>
<td>72.4 ± 3.2</td>
</tr>
</tbody>
</table>

Data are shown as mean ± standard error of the mean (SEM).

* significant versus match control group (p < 0.05),

**Table 3**

<table>
<thead>
<tr>
<th>Indexes</th>
<th>Control group</th>
<th>Diabetic group</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>male</td>
<td>female</td>
</tr>
<tr>
<td>Amplitude of T wave, mV</td>
<td>0.097 ± 0.006</td>
<td>0.094 ± 0.004</td>
</tr>
<tr>
<td>Duration of T wave, msec</td>
<td>30.1 ± 0.4</td>
<td>32.2 ± 0.3</td>
</tr>
<tr>
<td>Amplitude of P wave, mV</td>
<td>0.046 ± 0.006</td>
<td>0.043 ± 0.004</td>
</tr>
<tr>
<td>Duration of P wave, msec</td>
<td>26.2 ± 0.1</td>
<td>27.3 ± 0.2</td>
</tr>
<tr>
<td>Amplitude of R wave, mV</td>
<td>0.166 ± 0.006</td>
<td>0.156 ± 0.008</td>
</tr>
<tr>
<td>Duration of R wave, msec</td>
<td>27.0 ± 0.2</td>
<td>23.3 ± 0.3</td>
</tr>
</tbody>
</table>

Data are shown as mean ± standard error of the mean (SEM).

* significant versus match control group (p < 0.05),

b significant versus male diabetic group (p < 0.05).
intact control group (see table 3). The results indicate the disturbances of the ventricular conduction were more strongly associated with diabetes in male than in female rats.

CONCLUSIONS

1. We conclude that Type 2 diabetes, independently of gender, caused cardiac arrhythmias, functional changes in atrial and ventricular conduction, but only in females, in contrast to males, was accompanied by the development of myocardial diastolic dysfunction.

2. This data justify the necessity of gender-specific therapy development for the prevention and management of diabetic cardiovascular complications.

REFERENCES


THE IMPACT OF TYPE 2 DIABETES ON THE CARDIOVASCULAR SYSTEM IN MALE AND FEMALE RATS

O. V. Ivanova
SI «V. Danilevsky Institute for Endocrine Pathology Problems of the NAMS of Ukraine», Kharkiv, Ukraine
ivolga3006@ukr.net

Introduction. Type 2 diabetes mellitus is one of the main factors of the cardiovascular risk, which leads to a disproportionate increase in cardiovascular events in women and men. While the greater excess risk of diabetic vascular complications in women compared with men has been described, mechanisms underpinning the sex difference have not been identified in full.

The aim of this study was to determine the impact of type 2 diabetes (T2D) on the functional state of the cardiovascular system in male and female rats.

Materials and Methods. T2D was induced in male and female Wistar rats by a high-caloric diet during 14 weeks combined with intraperitoneal injections of 25 mg/kg streptozotocin twice per week. At the end of the study electrocardiograms were recorded in leads II. A comparative analysis of changes in the functional state of the heart in male and female rats with experimental T2D was conducted.

Results. It was established that T2D, independently of gender, results in the formation of pathologically accelerated rhythm and sinus tachycardia in experimental animals. Experimental T2D led to the prolongation of the systole in rats of both sex and decreasing of R-wave voltage in males in comparison with control group. In addition, T2D was accompanied by multidirectional impact on the atrial function in the heart of both sexes: P wave’ amplitude was increased in females while it was decreased in males, what can indicate either right or left atrial enlargement. It was found that T2D promotes the development of myocardial diastolic dysfunction in females, in contrast to males, which was confirmed by prolongation of T-P interval and a decrease of amplitude and duration of the T wave in comparison with intact female rats.

Conclusions. T2D, independently of gender, caused cardiac arrhythmias, functional changes in atrial and ventricular conduction, but only in females, in contrast to males, was accompanied by the development of myocardial diastolic dysfunction. This data justify the necessity of personalized- and gender-specific therapy development for the prevention and management of diabetic cardiovascular complications.

Key words: type 2 diabetes, functional state of the cardiovascular system, sex differences, rats.
Вступление. Сахарный диабет 2 типа является одним из основных факторов риска сердечно-сосудистых заболеваний, приводящим к непропорциональному их увеличению у женщин и мужчин. Известно, что риск развития диабетических кардиоваскулярных осложнений у женщин повышен по сравнению с мужчинами, но механизмы, обуславливающие гендерные различия, установлены не в полной мере.

Целью данного исследования было определить влияние сахарного диабета 2 типа на функциональное состояние сердечно-сосудистой системы у самцов и самок крыс.

Материалы и методы. Диабет 2 типа индуцировали у самцов и самок крыс линии Вистар высококалорийной диетой в течение 14 недель в сочетании с двумя внутрибрюшинными инъекциями стрептозотоцина в дозе 25 мг/кг через неделю. В конце исследования функциональное состояние сердечно-сосудистой системы крыс было изучено методом электрокардиографии и проведен сравнительный анализ полученных показателей в зависимости от пола.

Результаты. Установлено, что сахарный диабет 2 типа независимо от пола сопровождается формированием патологически ускоренного ритма и синусовой тахикардии у экспериментальных животных. Диабет 2 типа приводил к удлинению систолического интервала у крыс обоих полов и снижению амплитуды зубца Р у самцов по сравнению с контрольной группой. Кроме того, диабет 2 типа сопровождался развитием аритмий, функциональными изменениями в предсердной и желудочковой проводимости, но только у самок. Амплитуда зубца R у самцов в сравнении с интактными контрольными группами, что может свидетельствовать об увеличении либо правого, либо левого предсердий. Установлено, что сахарный диабет 2 типа индуцирует развитие диастолической дисфункции миокарда у самок в отличие от самцов, что подтверждалось пролонгацией интервала T-P, а также уменьшением амплитуды и длительности зубца T по сравнению с интактными самцами.

Выводы. Установлено, что сахарный диабет 2 типа, независимо от пола, сопровождается сердечной аритмиеей, функциональными изменениями в предсердной и желудочковой проводимости, но только у самок, в отличие от самцов, вызывает развитие диастолической дисфункции миокарда у самок, в отличие от самцов, что подтверждалось пролонгацией интервала T-P, а также уменьшением амплитуды и длительности зубца T по сравнению с интактными самцами.

Ключевые слова: сахарный диабет 2 типа, функциональное состояние сердечно-сосудистой системы, половые различия, крысы.