Evaluation of functional state of muscular system in children with diabetes mellitus

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A – research concept and design; B – collection and/or assembly of data; C – data analysis and interpretation; D – writing the article; E – critical revision of the article; F – final approval of the article

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Aim. To evaluate the functional condition of skeletal muscles in children with type 1 diabetes (T1D), according to the duration of disease based on studying static and dynamic physical endurance.

Materials and methods. 76 children with T1D from 11 to 17 years old were examined. The first group included 20 kids with the duration of diabetes less than 1 year. The second group – 27 patients with T1D from 1 to 5 years. The third group – 29 kids with T1D over 5 years. The group of control included 16 conventionally healthy children. Static muscular endurance with evaluation of total static muscular endurance (TSME) and dynamic muscular endurance with evaluation of wrist strength index (WSI) were evaluated. The index of muscle quality (IMQ) was evaluated too.

Results. The greater duration of T1D was accompanied by the redistribution of percentage ratio of body components. The TSME was decreasing along with increase of T1D duration. Reverse correlation between glycated hemoglobin and TSME (r = -0.50; P < 0.05) was shown. The dynamic endurance revealed the progressing decrease of WSI before and after the physical exertion in children with T1D in comparison with the control group. The revealed peculiarities of functional condition of muscles were connected with TSME (r = +0.43, P < 0.05) and IMQ (r = +0.52, P < 0.05).

Conclusions. Functional condition of skeletal muscles in children with T1D is characterized by decrease of static and dynamic muscular endurance. The first signs of changes are observed over the first year of disease and progress as the duration of a disease increases and leads to decrease of muscle mass and diabetic myopathy developing. The major factor contributing the worsening of functional condition of muscles is insufficient glycemic control.
It is well known that type 1 diabetes (T1D) is considered to be one of the specific risk factors of sarcopenia developing in adults. At the same time, there is lack of publications regarding to disturbances of musculoskeletal system in children with T1D. Although it is a junior age that is being a critical period in formation of skeletal system. According to the latest researches it is noticed that there is a reduction in level of physical training in children with T1D compared to healthy kids even in terms of the equal level of physical activity \[1\]. That may point to the diabetic myopathy progress – the condition characterized by both muscle mass and function reduction \[2\].

Considering the importance of skeletal muscles in metabolism of lipids and glucose from the blood, the progress of diabetic myopathy may lead to both insulin resistance and the further diminishing of KLF15 destruction, which depends on ubiquitin and thus, contributes to muscle atrophy \[4\].

The data of the effect of physical activity on glycemic control in T1D are controversial. But according to the data obtained by Chimen et al. \(2012\) physical exertion didn’t cause any positive effect on the level of glycated hemoglobin (HbAc1) in children with T1D. The other resources revealed the data about the credible reduction of level of HbAc1 in teenagers with T1D going through the dosed aerobic physical exertion \[5\].

The anxiety of hypoglycemia progressing after the physical training is considered to be one of the negative factors that restrict the physical activity in kids and teenagers with T1D. The other factor is lack of glycemic control. So that, tests applied for the early diagnosing of disturbances in skeletal muscles system, that always accompany the progress of diabetic myopathy, can be very useful.

**Aim**

The aim was to evaluate the functional condition of skeletal muscles in children with T1D, according to the duration of disease based on studying static and dynamic physical stamina.

| Material and methods |
|----------------------|
| 76 children with T1D from 11 to 17 years old were monitored. According to the duration of the disease all children were divided into three groups. The first group included 20 kids with the duration of diabetes less than 1 year. The second group includes 27 patients with the duration of diabetes from 1 to 5 years. The third group was formed of 29 kids with the duration of diabetes over 5 years. The group of control includes 16 conventionally healthy children. All groups were representative according to the age, gender and body mass index (Table 1).

| All the kids went through the measuring of body weight and length with the further evaluation of body mass index (BMI). The muscular mass in kids below 14 years old was estimated according to A.M. Peters’ formula \[6\]. The P. Boer’s formula was applied for kids above 15 years old and counted the gender \[7\]. In order to evaluate the condition of muscular system, the skeletal muscle index (SMI) was assessed according to the formula \[8\]: |
| \[
| SMI = \frac{(skeletal \ muscle \ mass/body \ mass) \times 100}{BFP} 
| \]
| \[
| \text{Where: BFP} = (1.51 \times \text{BMI}) - (0.70 \times \text{age}) - (3.6 \times S) + 1.4
| \]
| \[
| \text{Where: S = 1 for boys and 0 for girls; BMI – Body mass index; age – age in years.}
| \]
| In children over 15 years old the body fat percentage (BFP) was evaluated applying the following formula \[9\]: |
| \[
| BFP = (1.20 \times \text{BMI}) + (0.23 \times \text{age}) - (10.8 \times S) - 5.4
| \]
| \[
| \text{Where: S = 1 for boys and 0 for girls; BMI – Body mass index; age – age in years.}
| \]
| Body fat mass (BFM) was evaluated applying the following formula: |
| \[
| BFM = (BFP/100) \times \text{weight,}
| \]
| \[
| \text{Where: BFP – body fat percentage; weight – weight in kg.}
| \]
| The ratio between the fatty and muscle mass was evaluated in the following way: fatty mass in kg divided by muscle mass in kg and expressed in conventional units (CU). |
Table 1. Distribution of children by age and gender, observation groups

| Criteria, units   | Group 1, n = 20 | Group 2, n = 27 | Group 3, n = 29 | Group of control, n = 16 |
|-------------------|----------------|----------------|----------------|-------------------------|
| Average age (years), M±m | 13.51 ± 0.52 | 14.08 ± 0.41 | 14.17 ± 0.36 | 14.41 ± 0.47 |
| Number of boys, abs./ %  | 11/55.0  | 14/51.8 | 15/51.7 | 9/56.2 |
| Number of girls, abs./ % | 9/45.0 | 13/48.2 | 14/48.3 | 7/43.8 |

To define functional abilities of muscular system static and dynamic endureance of skeletal muscles was estimated while fixing the maximal time period of given testing position sustaining in seconds (sec).

At the time of the endurance functional tests of skeletal muscles of all children with T1D included in the research, feeling “before” and “after” testing were assessed as satisfactory. Tests were performed 1 hour after eating to determine the concentration of glucose before, immediately after and 1 hour after end samples (in accordance with the recommendations of the Protocol of rendering medical aid to children with diabetes mellitus approved by Order № 254 of the MH of Ukraine dated 04/27/2006 with changes made in accordance with the Ministry of Health Orders № 55 (v0055282-09) dated 02/03/2009; № 864 (v0864282-13) dated 7/10/2013). The criteria for inclusion of patients into the research were: the consent of the patient and his parents for participation in the research; lack of ketoacidosis (ketone bodies of the urine were determined before and after the samples); the maximal level of glycemia on an empty stomach did not exceed 10.5 mmol/L, and the minimal level of glycemia was 5.7 mmol/L. The average level of glycemia on an empty stomach in the first group was 7.34 ± 0.48 mmol/L, in the second group – 7.38 ± 0.52 mmol/L, in the third group – 7.84 ± 0.48 mmol/L. The research excluded patients with the lack of consent to participate in the research; children, with T1D with obesity and excess body weight; with the presence of acute inflammatory processes or congenital malformations in the stage of decompensation.

To define the endurance of neck flexor muscle groups all children were requested to hold the neck flexed when lying down on the back. The chest should be on the surface.

To define the static endurance of spinal extensors children were requested to lift the upper body and hold it above the surface while lying down on the abdomen with the head, the legs on the surface and hands crossed up behind.

To define the muscular endurance of abs children were supposed to lift the legs upwards to the angle of 45° and hold when lying on the back on the surface [10].

To define the endurance of the glutous medius children were supposed to lift the leg up to the side up to the possible height and hold without the rotation when standing [11].

Total static muscle endurance (TSME) was also evaluated. The range of muscle quality was evaluated as a ratio of TSME to skeletal muscle index.

Skeletal muscle strength was estimated using the wrist spring dynamometer, which was squeezed by the patients’ hand, in order to offset the age of a kid when estimating the muscle strength, one has applied the wrist strength index (WSI):

\[ WSI = \frac{\text{wrist strength}}{\text{body mass (kg)}} \times 100\% \] [2].

The test was carried out using the wrist rubber ring-like expander with the resistance equal to 20 kg, one had to squeeze the expander during 30 sec.

All the results were analyzed using the set of statistic programs Statistica 13.0 (StatSoft Inc., № JPZ2041382130ARCN10-J). Parametrical methods that helped to evaluate simple average, mean squared deviation and standard error were applied for normally distributed rates. The method of correlation analysis was used to calculate the Pearson correlation coefficient in the normal distribution of features and the Speaman’s rank correlation coefficient in their absence. The reliability of the differences in the results obtained for different groups in the normal distribution of characteristics was determined by the parametric (Student’s criterion) method. Differences were considered to be significant at P < 0.05 [12].

In planning this work the bioethical commission gave permission to conduct research. All patients signed informed consent to participate in the study. All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and national research committee and within the 1964 Helsinki declaration and its later amendments or comparable ethical standards. Informed consent was obtained from all individual participants included in the study. The full data set by children, their parents, and physician that support the findings of this study are not publicly available due to the ethics approval originally obtained.

Results

According to the results of research it was established that there was no significant difference in BMI in all examined groups and all the rates were considered to be in normal ranges (Table 2).

Instead, as the duration of T1D was increasing, a gradual decrease in skeletal muscle index in patients with diabetes mellitus was observed compared with controls, becoming statistically significant after 5 years of the disease (P < 0.05). Starting from the second year of the disease, there was a significant increase in the percentage of body fat (P < 0.05). Also, there is the increase in the dynamics of the disease ratio of “fat mass / muscle mass” (P < 0.05) compared with the indicator of both the control group and the 1st group of patients with diabetes (Table 2). That is, an increase in the “length of time” of diabetes was accompanied by a component redistribution of body composition.

Taking into the account that changes in structural component of body may affect the functional muscle condition the next step of our investigation included static and dynamic test to estimate the muscle endurance. According to the data obtained after carrying out static endurance tests for examined muscle groups it was established that there was a deterioration of ability to hold the test position.
in children with T1D regardless of duration of disease, which led to statistically significant decrease of total static muscular endurance compared to results obtained from testing the group of control (Table 3).

The worst rates of muscle endurance were observed in abdominal muscles. It was 2.5–2.9 times lower compared to rates of the control group. It should be noted that, as the duration of disease increased the rates of total static muscular endurance were getting worse through the neck muscles. If in the first year of the disease the endurance of this muscle group was reduced by 1.7 times compared with the indicators of the control group, then with a long course of diabetes (group 3) this indicator not only statistically differed from the values of the control group (P < 0.05), but was 1.5 times less than the same indicator in group 1 (P < 0.05).

It was noticed that there was also worsening of muscle quality rates coming with changes in static muscle endurance in patients with T1D. There was a progressive decreasing of given indicator from 2.00 ± 0.15 CU in the first year of disease down to 1.17 ± 0.04 CU in patients from the 3rd group, which was 2 times less than the same indicator obtained in the group of control.

The level of glycated hemoglobin (HbA1c) in the research group increased with increase of duration of T1D and amounted to first group 8.9 ± 0.6 %, in the second group – 9.12 ± 0.38 %, in the third group – 10.31 ± 0.28 %. It was established, that the reduced static muscular endurance in patients with T1D is related to the increased level of HbA1c (r = -0.35, P < 0.05). This correlation became the most significant during the first 5 years of disease (r = -0.50, P < 0.05). That means that the lack of diabetes control causes significant changes in glycemia and causes the development of chronic complications of diabetes mellitus, as well as impaired function of skeletal muscles.

When examining the dynamic endurance of wrist and arm muscles using the rates of BMI it was indicated that there was a decreased initial rate compared to the control group, as well as its even bigger decreasing after physical exertion (Table 4).

While WSI in the group of control decreased on average by 6.8 % after physical exertion, patients with T1D had 13 % less WSI index in the first 5 years of disease increasing up to 16 % in patients of the 3rd group (P < 0.05), which spoke about worsening of functional state of skeletal muscles and increased muscle fatigue, which was observed during the first years of disease and was in progress in terms of growing endurance. Additionally, the determined particularities of functional state of wrist and arm muscles were related to total muscular static endurance (r = +0.43 and r = +0.37, respectively for the left and right upper limbs, P < 0.05) and to the muscle quality rate (r = +0.52 and r = +0.47, respectively, P < 0.05).

Discussion

The results obtained show the children with T1D to have an obvious decrease of muscular endurance starting from the first year of disease. It is mostly affected by insufficient glycemic control which is proved by received correlation relationship between glycated hemoglobin

Table 2. The rates of BMI and fatty and muscle mass in children with T1D according the duration of disease, M ± m

| Criteria, units | Group 1, n = 20 | Group 2, n = 27 | Group 3, n = 29 | Group of control, n = 16 |
|-----------------|----------------|----------------|----------------|------------------------|
| Body mass index, kg/m² | 18.99 ± 0.7 | 20.78 ± 0.7 | 20.36 ± 0.57 | 20.51 ± 0.89 |
| Percent of fat in organism, % | 16.68 ± 0.91 | 19.09 ± 0.99 | 20.02 ± 0.86 | 16.74 ± 2.98 |
| Skeletal muscle index, % | 81.39 ± 1.08 | 78.56 ± 1.14 | 77.33 ± 1.10 | 81.1 ± 1.17 |
| Fatty mass/muscle mass, CU* | 0.21 ± 0.01 | 0.25 ± 0.01 | 0.26 ± 0.02 | 0.20 ± 0.02 |

Table 3. The rates of static muscular endurance in children with T1D counting the duration of disease, M ± m

| Group of muscles examined, units | Group 1, n = 20 | Group 2, n = 27 | Group 3, n = 29 | Group of control, n = 16 |
|---------------------------------|----------------|----------------|----------------|------------------------|
| Neck flexors, sec. | 53.57 ± 5.17 | 42.75 ± 2.91 | 35.84 ± 2.66 | 90.82 ± 9.47 |
| Muscles of abs, sec. | 18.67 ± 2.2 | 21.12 ± 1.87 | 18.22 ± 1.52 | 52.63 ± 6.06 |
| Spinal extensors, sec. | 21.18 ± 1.87 | 20.54 ± 1.76 | 18.33 ± 1.16 | 38.25 ± 3.78 |
| Left gluteus medius, sec. | 29.47 ± 2.99 | 29.46 ± 2.73 | 29.67 ± 2.59 | 40.46 ± 4.3 |
| Right gluteus medius, sec. | 30.59 ± 3.45 | 30.38 ± 2.21 | 31.46 ± 2.77 | 43.23 ± 4.7 |
| Total static muscle endurance, sec. | 157.21 ± 11.47 | 138.13 ± 6.52 | 127.12 ± 8.13 | 259.09 ± 18.18 |
| Rate of muscular quality, CU* | 2.00 ± 0.15 | 1.88 ± 0.10 | 1.59 ± 0.11 | 3.26 ± 0.28 |

Table 4. The rates of dynamic endurance of wrist and arm muscles in children with type 1 diabetes according to the duration of disease, M ± m

| Indicators, units | Group 1, n = 20 | Group 2, n = 27 | Group 3, n = 29 | Group of control, n = 16 |
|-------------------|----------------|----------------|----------------|------------------------|
| WSI left, CU | 38.11 ± 3.24 | 37.21 ± 1.28 | 36.51 ± 1.76 | 50.75 ± 2.12 |
| WSI1/WSI2, left, CU | 1.14 ± 0.02 | 1.16 ± 0.03 | 1.22 ± 0.03 | 1.06 ± 0.03 |
| WSI right, CU | 42.44 ± 3.22 | 39.73 ± 2.01 | 38.75 ± 2.02 | 55.15 ± 2.71 |
| WSI1/WSI2, right, CU | 36.79 ± 3.25 | 34.38 ± 1.75 | 32.63 ± 1.90 | 51.52 ± 3.05 |

§ P < 0.05 in the period with the highest indicator of the control group; * P < 0.05 in the period with the highest indicator of 1 group; CU*: conventional units.
and total static muscular endurance \((r = -0.35, P < 0.05)\). At the same time, statistically significant decreasing of muscle mass occurs only after the 5th year of the disease. Thus, disturbances in functional abilities of muscles on the background of chronic hyperglycemia already fixed during the first years of disease eventually lead to decrease of muscle mass, in other words to the development of diabetic myopathy. Moreover, in children with T1D an increase in the ratio between fat and muscle tissue in condition of normal body mass index was observed. One of the reasons causing increasing of fatty tissue percentage in patients with T1D is insufficient glycemic control ending up with chronic hyperglycemia. Wherein, excessed amount of glucose turns into a fat, because the fatty tissue is characterized by quite significant metabolic activity and is highly irresistible to insulin. The increasing percentage of fatty tissue leads to further metabolic response dysfunction to exogenous insulin, that is to say towards insulin resistance [13]. The receptors of insulin situated in muscles play the main role in glucose regulation, while muscle tissue itself is the main place of glucose utilization, because the glucose is considered to be the source energy material necessary for their functioning [14]. Monaco C. M. in his research supposed, that repeated insulin injections may cause periods of relapses intracellular hyperglycemia in myofibrils. Thus, patients with T1D have a greater dependence on glycolytic metabolism due to lack of mitochondrial function and accompanying changes in muscle structure [3]. At the same time, we taking into account, that insulin is considered to be a catabolic hormone promoting the synthesis of protein by facilitating the muscle proteins production. The defects of its signals transmitting leads to lack of protein production and excessed protein degradation, which eventually causes the muscle power and muscle mass deficiency [15].

Conclusions

1. Functional condition of skeletal muscles in children with type 1 diabetes is characterized by deficiency of static and dynamic muscular endurance.
2. The first signs of violation of functional condition of skeletal muscles are observed over the first year of disease and progress as the duration of a disease increases and eventually leads to decrease of muscular mass and diabetic myopathy developing.
3. The major factor of risk contributing the worsening of functional condition of muscles is insufficient glycemic control.

Prospects for further studies. We plan to determine the predictors formation of diabetic myopathy in children with T1D.

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