Hepatic and skeletal muscle glycogen content in rats treated with metformin and submitted to acute exercise by swimming

Abstract

The aim of this study was to evaluate the behavior of glycogen reserves over an acute exercise condition (50 minutes of swimming at low intensity), after treatment with metformin in rats. Forty Wistar rats (180-200g) adults were divided into four groups (treated or not for a fortnight) and represented as follows: Control; Acute exercise by swimming (perform a session of swimming, 50 min on light intensity); Treated with metformin (received the drug metformin at a dose 1.4 mg/ml during the experimental period); Treated with metformin and exercised by swimming (received the drug metformin at a dose 1.4 mg/ml and held a swimming, 50 min on light intensity). The acute exercise decreased the glycogen reserves, while animals treated with metformin showed an increase in their muscle and liver glycogen reserves (p>0.05). Additionally, the drug showed no increase glucose and corticosterone concentration compared to the control and treated with metformin groups (p>0.05). Treatment with metformin improved energy conditions and lowers the stress response, suggesting that an important pharmacological tool for the potentiation of performance.

Keywords: Glycogen; Performance; Biguanide; Rats.

Introduction

Recent studies have demonstrated the multifactorial action of the biguanide metformin, while pharmacological agent widely used in glycemic control in patients with diabetes mellitus\textsuperscript{1-2}. Within the broad spectrum of action of the substance, deserves spotlight the reduction in the production/release of liver reserves of glucose (gluconeogenesis, glycogenolysis), increased insulin sensitivity in (especially in muscles) peripheral tissues, increased peripheral glucose utilization (by capture glucose and insulin-stimulated glycogen synthesis) and increase the population of insulin receptors (reducing hyperinsulinemia) without weight gain\textsuperscript{3-4}.

Metformin formula was first published by Ungar et al.\textsuperscript{5} and used primarily by Mahler et al.\textsuperscript{6} in patients with type 2 diabetes mellitus, being introduced for use in insulin resistant states, even before the development of hyperglycemia\textsuperscript{7}. According to Diabetes Prevention Program Research Group\textsuperscript{8}, its effects include anti-hyperglycemic action without risk of severe hypoglycemia and a reversal of the insulin resistance process.

Metformin activates AMP-activated protein kinase (AMPK) that promotes glucose utilization\textsuperscript{9}, an increase in AMPK activity is associated with increased translocation of glucose transporter (GLUT)-4 to the plasma membrane, increase in hexokinase activity and glycogen content in cells muscle, in addition, there is increased oxidation of fatty acids\textsuperscript{10}.

Inside the metabolic profile of skeletal muscle, it is known that preferentially uses glucose as energy substrate, which is captured in basal conditions by the carrier glucose GLUT-1 whose action is not
dependent on insulin or the GLUT-4 that can be translocated from cytosolic reservoirs for membrane through the action of insulin or by the increase in contractile activity\(^1\). After uptake, glucose can be oxidized or reserved in the form of glycogen, while important determinant of the efficiency of reservoir contractile process, reaching minimum values when the muscle goes into fatigue\(^1\).

The antihyperglycaemic activity of metformin is related mainly to the suppression of gluconeogenesis, block in glycogenolysis, reduction of intestinal glucose absorption, increased kinase activity of the insulin receptor, stimulation of post-insulin receptor pathways, increased translocation of GLUT-4, glucose uptake consequent elevation of the enzyme glycogen synthase activity leading to induction of the formation of large reserves of glycogen\(^12-13\). Emerging hypothesis to evaluate whether the drug promotes increased metabolic resistance against physical exercises.

When considering the metabolic changes provided by the use of metformin, which promotes a rise in the glycogen content, the hypothesis arises that the previous use of metformin, at doses like those commonly prescribed for diabetics, may provide adjustments in energy metabolism that may be important in the practice of physical exercise. Thus the aim of this study was to evaluate, in rats eutrophic and previously treated with metformin, the behavior of glycogen reserves over a acute exercise during 50 minutes of swimming in low intensity. We emphasize that, in spite of that medication be prescribed for diabetics, recently, eutrophic individuals who begin to perform physical activity have indiscriminately used it. These individuals use the drug as a result of metabolic modulations associated with improved physical performance. So opens notorious perspective to research this use.

**Methods**

**Sample**

It was used forty *Wistar* rats (*Rattus novergicus var, albinus*, *Rodentia, Mamalia*) at 3 months of age, weighing 180-200 grams, from the Vivarium of the Methodist University of Piracicaba - UNIMEP and kept in the vivarium of the Faculty of Health Sciences (FACIS-UNIMEP) under ambient temperature of 23°C ± 2°C, submitted to light/dark cycle of 12 h light, water and food *ad libitum*.

The study lasted 15 days, the mice were randomly divided into four experimental groups (ten animals per group): 1) Control (C); 2) Stressed by swimming (E); 3) Treated with Metformin (M); 4) Treated with Metformin and stressed by swimming (ME). All procedures used in this study were in accordance with the principles of handling and care of laboratory animals recommended by COBEA (Brazilian College of Animal Experimentation) and approved by the Ethics Committee on Animal Use of UNIMEP protocol number 09/2013.

**Metformin treatment**

The experimental groups treated with metformin received the substance by the orogastric via at the concentration of 1.4 mg/ml during 15 days. This concentration was chosen following proposal that showed no toxicity at this concentration\(^14\). Furthermore, considering that the animal body weight changed by 180–200 grams, the correction of that delineated dosage for a 70 kg man would denote range of 490-544 mg, like the also delineated for insulin resistant individuals.

**Stress induced condition**

After fifteen days of metformin treatment in ME group and without metformin treatment in the E group, the animals underwent a single session acute exercise by swimming for 50 minutes in low intensity, performed in a tank containing water at 30 ± 2°C\(^15-16\). The training sessions were drawn without the use of additives, which corresponds to an intensity below the anaerobic threshold, or to a predominantly aerobic stress, according to prior publication\(^17\). Such volume and the intensity chosen are justified by the fact that the same are applicable to individuals, apparently healthy and beginners in the physical exercise.

**Biochemical reviews**

The animals were anesthetized with sodium pentobarbital (40mg/Kg, ip) immediately after the
exercise and the blood was collected by renal artery, this blood was centrifuged and the plasma was separated for evaluation of plasma glucose concentration by an enzymatic kit Labortest, corticosterone was performed by immunoassay kit from Assay Designs* (Ann Arbor, MI, USA). The following samples of liver, soleus and gastrocnemius white portion were carefully isolated and removed for determination of glycogen content by the phenol sulfuric acid method$^{18}$.

**Histological analysis**

For histological analysis of the liver, its ventral segment was fixed and then processed from which various cross-sections of non-serial 5μm thick, which were stained using hematoxylin-eosin (HE) were obtained.

**Statistical analysis**

The collected data were tabulated and analyzed using the statistical software “GraphPad Prism 6”. Statistical analysis was performed by applying the Shapiro-Wilk test to verify the normality of the data.

Thus, for comparisons between groups, the statistical test for analysis of variance (one-way ANOVA) followed by Tukey-Kramer multiple comparisons for parametric data. In all cases we adopted a value of p < 0.05 for statistical significance.

**Results**

Were observed that glycogen stores of soleus muscle were depleted by 60% due to the stress-inducing condition (E). On the other hand, in the group treated with metformin (M) the glycogen content presented was 19% higher compared to the control (C). In the group treated with metformin and subjected to acute exercise (ME) was observed that the glycogen content was 89% higher when compared to untreated and exercised group (E) (FIGURE 1).

Moreover, it is noted that behavior of muscle glycogen reserves of gastrocnemius white portion, on different experimental conditions, was 69% lower because of acute exercise (E). In turn, it is noteworthy that there was an increase of 21% on glycogen reserves in the group treated with metformin (M). In assessing, the group treated with metformin and subjected to acute exercise (ME) it was found that glycogen reserves showed up 150% larger compared to untreated and exercised group (E) (FIGURE 1).

After that, the hepatic glycogen content was evaluated, and found that the acute exercise decreases the reserves in 84% (E) if compared to the control group (C). In the group treated with metformin (M) the hepatic glycogen reserves was increased in 57%, on the other hand, in the metformin group subjected to acute exercise (ME) the reserves were 368% higher when compared to untreated and exercised group (E). In this condition, the values were 26% lower in ME compared to the content observed in the control group (C) (FIGURE 1).

In assessing of blood glucose was found that acute exercise (E) promoted an increase of 99.4% compared to the control group (C). In the group treated with metformin (M), the values showed no significant difference if compared to the control group (C). Already the treated and exercised group (ME) showed 38% lower blood glucose if compared to untreated and exercised group (E) (FIGURE 1).

Within the endocrine profile, plasma corticosterone was evaluated and observed that treatment with metformin did not change the plasma concentration. On the other hand, the acute exercise group showed (E) a significant increase of 185% in the plasma concentration if compared to control group (C), something observed in the treated and exercised group (ME), but ME had lower concentrations of corticosteroids if compared to untreated and exercised group (E) (FIGURE 1).

The histological analysis of hepatic glycogen content can show the glycogen granules, which are more abundant on the hepatocytes in treated group (M) and treated/exercised group (ME). In the liver of exercised animals (E), there was depletion of granules mainly in the perportal region. The same did not occur with the same intensity in the treated and exercised group (ME), indicating lower reserve mobilization in the presence of biguanide (FIGURE 2).
The results are expressed as the mean ± SD, n=10. C = Control, M = Treated with Metformin, E = Exercised by swimming, ME = Treated with Metformin and exercised by swimming. *p < 0.05 comparing with group C; #p < 0.05 compared with group M; ψp < 0.05 compared with group E. 227x293mm (300 x 300 DPI)

FIGURE 1 - Content of muscle glycogen, hepatic glycogen, plasma glucose and corticosterone concentration of experimental groups.

Discussion

The interrelationships between carbohydrate metabolism in the liver and muscles are studied by different scientists, since the energy reserves are determinants of resistance at physical exercise. However, it has been noted modifications at the mobilization of these reserves concomitant to elevation at physical activity, stress or in the presence of some drugs that modify metabolic homeostasis of the organism.

Initially we studied the effects of acute exercise induced by inescapable forced swim for 50 minutes in low
intensity, proposed by Sampaio-Barros et al.\textsuperscript{15}, which corresponds to an intensity below the anaerobic threshold, or to a predominantly aerobic stress, according to prior publication\textsuperscript{17}, on liver and muscle glycogen reserves. The data show that acute exercise condition can cause an intense depletion of hepatic glycogen stores (FIGURE 1), and can be explained by the increase in the neuroendocrine axis activity, represented by the increase in the secretion of epinephrine and norepinephrine, which activate adrenoceptor determining glycogenolysis in hepatocytes\textsuperscript{21}. Occurs sequentially increase in the secretion of growth hormone, glucagon, adrenocorticotropic hormone, prolactin and opioid peptides, which also manifest glycogenolytic action, characterizing a synergistic effect\textsuperscript{22}.

In order to avoid doubts and characterize the acute exercise, plasma corticosterone was evaluated and showed an increase in its concentration after physical training session (FIGURE 1), reflecting the activation of the neuroendocrine axis\textsuperscript{23}.

With respect to muscle glycogen reserves, it was observed that due to the increase in energy requirements generated by the inducing acute exercise condition the reserves were depleted (FIGURE 1). These results corroborate with recent studies, which suggest that the intensity and duration of the stressor may modulate the utilization of energy substrates, with a special energy source, the degradation of muscle glycogen\textsuperscript{15,24-25}.

Thus, metformin was used in this study to verify its efficiency in conditions of acute exercises, based on published reports that indicate that the biguanide promotes a significant improvement in muscle glucose metabolism\textsuperscript{26-28}.
It has been suggested that metformin does not alter the phosphorylation of the insulin receptor, and its centralized action in the post-receptor phenomena, where the biguanide promotes a rise in glucose uptake level, preserves the sensitivity of hepatocytes to insulin to regulate the activity of tyrosine kinase bound to the receptor, it enhances the dynamics of glucose uptake favoring the formation of glycogen reserves. In muscle, AMPK activation by metformin promotes the use of glucose, this increases the enzyme activity, causing the GLUT-4 translocation to the plasma membrane, promoting an increase in the activity of hexokinase and glycogen content in muscle cells. In addition, fatty acids undergo oxidation and reduction of the synthesis.

In the liver, AMPK activation inhibits transcription of enzymes phosphoenolpyruvate car-boxiquinase (PEPCK) and glucose-6-phosphatase (G6Pase), generating a reduction in gluconeogenesis. In vitro and in vivo studies demonstrated that phosphorylation of AMPK, caused by metformin, are modulated by the OCT1 activity, suggesting that this transporter has a key role in the cascade of events that result in decreased hepatic gluconeogenesis. In addition, AMPK also promotes improvement in lipid metabolism during treatment with metformin, so that activation of this enzyme inhibit acetyl-CoA carboxylase (ACC) and 3-hydroxy-3-methylglutaryl-coenzyme and reductase (HMG-CoA reductase), key enzymes in the synthesis of triglycerides and cholesterol, respectively. In addition, AMPK activation in the liver also suppresses the expression of SREBP-1 and SREBP-1c genes, which are involved in the pathogenesis of insulin resistance, dyslipidemia and diabetes mellitus.

It’s permissive to suggest that considering such molecular mechanisms associated with capitation of glucose by peripheral tissues resulting from metformin, this one has the potential to improve performance in physical exercise in predominantly aerobic conditions, due to increased energy substrate availability to meet demand from the effort. This increased availability of glucose to satisfy the demand of the aerobic effort, is crucial, since for metabolizing fats, requiring the continuous creation of oxaloacetate, recognized this metabolite is synthesized from own glucose. In his line of thought, studies with animal models has shown that skeletal muscle and liver glycogen reserves are related to higher physical performance, resulting in the increase in the threshold exhaustion, specifically against the effort in aerobic condition.

The assessment of hepatic glycogen content of rats treated with metformin (M) showed that stocks were high, the histological analysis showed it too, and is due to metformin suppress hepatic gluconeogenesis and activate the mechanisms of glycogen synthesis by activating enzymes flag system at the level of insulin post-receptor, especially the enzyme via phosphatidyl-inositol 3 kinase (Pi-3K) and glycogen synthase. Similar finding was observed in muscles treated with the biguanide, which also showed an increase in the concentration of glycogen synthesis whose actions should be elevated post-insulin receptor, similar to that described in hepatocytes. It is worth mentioning that mice treated with metformin (M) showed no change in the secretion of corticosterone (FIGURE 1), showing specificity in the action of the biguanide metabolic adjustment.

In the experimental result we started to evaluate the hepatic and muscle glycogen reserves of rats treated with metformin undergoing inducing acute exercise condition (ME) being observed that the reserves were significantly higher than those observed in the untreated and exercised group (E) (FIGURE 1). This result shows that the glycogenic activity of metformin gave better energy conditions both in liver as skeletal muscle, leading to a different reaction when animals to stressful conditions and yet, the plasma concentration of glucose and corticosterona were lower compared to untreated and exercised group (E) (FIGURE 1), culminating in increased fatigue threshold and, consequently, increasing the time to exhaustion, due to the delay in decrease of muscle and liver glycogen. This suggests that the treatment led to an increased resistance against acute exercises indicating that there may be a correlation between the energy status and response to physiological stress in the conditions of this study, with a session of swimming for 50 minutes in low intensity.

A limitation of the present study was the fact that the analysis of the evaluated parameters occurred under acute exercise by swimming condition. Thus, if the evaluations of these parameters had occurred under a chronic training condition, would surely bring greater subsidies for understanding the use of metformin.

Concluding, metformin treatment modified the metabolic responses classically observed against acute exercises conditions providing greater resistance exercise confirming the study hypothesis.
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initially proposed. It is noteworthy that the limiting factor of this study is the fact that the effects of the supplementation were evaluated solely on condition of acute stress. In this sense, future studies are which would be required, focusing on analysis of the chronic effects of such supplementation and during subsequent training sessions. Once the biguanide is not metabolized and is excreted in pharmacological form, this component still induces an increase of muscle and liver glycogen, enabling greater endurance. Furthermore, whereas a period of 15 days of supplementation with metformin, in eutrophic animals, has the potential to maximize the metabolic responses to acute stress by maintaining liver and musculoskeletal glycogen stores. Thus, it is necessary that the readily Word Anti-Doping Agency develop a technique for the diagnosis of use of metformin in order to identify a possible frame of doping. Further studies are required, and the physical therapists and fitness trainers shall see that the substance should not be used as an ergogenic agent.

Resumo

Reservas hepáticas e musculares de glicogênio de ratos tratados com metformina e submetidos ao exercício agudo por natação

O objetivo deste estudo foi avaliar o comportamento das reservas glicogênicas de ratos, submetidos a uma condição de exercício agudo (50 minutos de natação na intensidade leve), após o tratamento com metformina. Quarenta ratos Wistar (180-200g) adultos foram divididos em quatro grupos (tratados ou não por quinze dias) e assim representados: Controle; Exercício agudo por natação (realizaram uma sessão de natação, sendo 50 minutos na intensidade leve); Tratado com metformina (receberam o fármaco metformina na dosagem de 1,4 mg/ml, durante o período experimental; Tratados com metformina e submetidos a condição exercício agudo por natação (receberam o fármaço metformina na dosagem de 1,4 mg/ml e realizaram uma sessão de natação, sendo 50 minutos na intensidade leve). O exercício agudo diminuiu as reservas glicogênicas, já os animais tratados com metformina, apresentaram um aumento em suas reservas glicogênicas musculares e hepáticas em relação ao grupo que realizou o exercício sem suplementação (p<0,05). Adicionalmente, o fármaco demonstrou não aumentar a glicemia e a concentração de corticosterona em relação ao grupo controle e suplementado com metformina (p>0,05). O tratamento com metformina promoveu melhora nas condições energéticas e menor resposta ao estresse, sugerindo ser uma importante ferramenta farmacológica para a potencialização da performance.

Palavras-chave: Glicogênio; Desempenho; Biguanida; Ratos.

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