The effect of the Ginger dry extract on the indicators of the carbohydrate metabolism under conditions of the experimental metabolic syndrome in Syrian golden hamsters

**Aim.** To study the effect of the ginger dry extract on the indicators of the carbohydrate metabolism in the experimental metabolic syndrome.

**Materials and methods.** The effect of the ginger dry extract on the carbohydrate metabolism was determined by the level of basal glycemia, basal insulinemia, HOMA-IR index, HbA1c level, the glycogen content in the liver and the body weight against the background of the metabolic syndrome induced by a high-calorie diet in Syrian golden hamsters.

**Results and discussion.** Consumption of high-calorie food for 6 weeks led to development of the metabolic syndrome, it was confirmed by an increase in the body weight, hyperglycemia, compensatory insulinemia, insulin resistance, increased glycolysis in the liver and glycosylation of proteins. The use of the ginger dry extract in the dose of 80 mg/kg over the period of 14 days reliably reduced blood glucose by 43.3 % and normalized insulinemia by 32.8 % affecting a decrease in the HOMA-IR index. The introduction of the ginger extract in the dose of 80 mg/kg was also accompanied by suppression of protein glycosylation by 29.6 % and restoration of glycogen-forming function of the liver. By its ability to restore the carbohydrate metabolism the ginger dry extract in the dose of 80 mg/kg did not differ from metformin and exceeded the effectiveness of the herbal drug “Arphasetin”. It is probably due to the powerful complex pharmacological action of phenolic compounds of ginger – gingerols and other components.

**Conclusions.** On the experimental model of the metabolic syndrome the use of the ginger dry extract normalized blood glucose, insulinemia, decreased insulin resistance and restored the glycogen content in the liver at the level of metformin. By the intensity of the pharmacological action the ginger extract exceeded the reference herbal drug “Arphasetin”. This fact is the basis for its further pharmacological study as a promising agent for the treatment of the metabolic syndrome and type 2 diabetes.

**Key words:** Ginger dry extract; metabolic syndrome; high-calorie diet; carbohydrate metabolism

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Вплив сухого екстракту імбиру на показники вуглеводного обміну за умови експериментального метаболічного синдрому у сирійських золотих хом’ячків

**Мета** – вивчення впливу сухого екстракту імбиру на показники вуглеводного обміну за умови експериментального метаболічного синдрому.

**Матеріали та методи.** Визначення впливу сухого екстракту імбиру на вуглеводний обмін проводили за рівнем базальної глюкемії, базальної інсулінемії, індексом HOMA-IR, рівнем HbA1c, вмістом глікогену у печінці та масою тіла на тлі метаболічного синдрому, індукованого висококалорійною дієтою у сирійських золотих хом’ячків.

**Результати та їх обговорення.** Споживання висококалорійної їжі впродовж 6 тижнів призвело до розвитку метаболічного синдрому, що підтверджувалося збільшенням маси тіла, гіперглікемією, компенсаторною інсулінемією, інсулінорезистентністю, посиленням процесів глікогенолізу у печінці та глікозилування білків. Застосування сухого екстракту імбиру дозою 80 мг/кг впродовж 14 днів достовірно на 43,3 % знижувало рівень глікози в крові та на 32,8 % нормалізувало інсулінемію, що відображалося зменшенням індексу HOMA-IR. Уведення екстракту імбиру дозою 80 мг/кг також супроводжувалося пригніченням процесу глікозилування білків на 29,6 % та відновленням глікогеноутворювальної функції печінки. За здатністю відновлювати показники вуглеводного обміну сухий екстракт імбиру дозою 80 мг/кг не відрізнявся від метформіну та перевищував ефективність збору «Арфазетин», що, можливо, обумовлено потужною комплексною фармакологічною дією фенольних сполук імбиру – гінгеролів та інших складових компонентів.

**Висновки.** На експериментальній моделі метаболічного синдрому застосування сухого екстракту імбиру нормалізувало глюкемію, інсулінемію, зменшувало інсулінорезистентність та відновлювало вміст глікогену у печінці на рівні метформіну. За виразністю фармакологічної дії екстракт імбиру переважав препарат порівняння збор «Арфазетин», що є обґрунтованням подальшого фармакологічного вивчення екстракту імбиру як перспективного засобу для лікування метаболічного синдрому та ЦД 2 типу.

**Ключові слова:** сухий екстракт імбиру; метаболічний синдром; висококалорійна дієта; вуглеводний обмін
According to the modern concepts the metabolic syndrome (MS) is a complex of metabolic, hormonal and clinical disorders that can be as factors of high risk for development of cardiovascular diseases. The main manifestations of MS are the abdominal type of obesity, insulin resistance and hyperinsulinemia, arterial hypertension, glucose tolerance/type 2 diabetes mellitus (DM), early atherosclerosis/ischemic heart disease, hemostasis disorder, hyperuricemia and gout, microalbuminuria, hyperandrogenemia [1].

Today MS attracts the attention of many experts since it precedes the emergence of type 2 diabetes mellitus (DM) and atherosclerosis – diseases that presently are major causes of mortality [2]. Increased interest is due to the fact that MS is a reversible condition – with the appropriate timely treatment the disappearance or a significant reduction of its main manifestations can be achieved.

The basic aim of the treatment of patients with MS is correction of the main components of of MS, such as insulin resistance and hyperinsulinemia, arterial hypertension, obesity and lipid metabolism disorders. Along with changes in the lifestyle and drug correction an important role in complex therapy of MS and type 2 DM is played by herbal products that due to a wide range of the pharmacological action can regulate the carbohydrate and the metabolism, blood pressure, blood rheology, normalize the body weight. The results of numerous experimental and clinical studies indicate the hypoglycemic, hypolipidemic, antioxidant properties that plants exhibit, including ginger, which is a famous spice, and the source of useful and medicinal substances [3-5]. Biologically active substances developed on the basis of ginger root are used for the prevention and treatment of different diseases as anti-inflammatory, antitumor, antioxidant, antiabetic drugs. It has been found that ginger decreases glucose levels in the blood, enhances glycogenogenesis, improves the synthesis of insulin and reparative properties of the pancreas, and normalizes the lipid metabolism [6, 7].

The aim of the work was to study the effect of the ginger dry extract on the indicators of the carbohydrate metabolism in the experimental metabolic syndrome induced in Syrian golden hamsters.

Materials and methods
A dry extract of common ginger (Zingiber officinale) produced by “Medagroprom”, Dnipro, was obtained from the ginger rhizomes by extraction with 50% alcohol and spray drying. The ginger dry extract (IMB) contains phenolic compounds – gingerol (not less than 5%) that provide its pharmacological activity, and a dry residue (at least 95%).

In the experiment Syrian golden male hamsters (20 weeks old at the beginning of the experiment) were used; they were kept under standard conditions in the vivarium of the Central Research Laboratory of the National University of Pharmacy at a temperature of 22 ± 1°C, humidity of 50-60%, in a room with changing the day or night light regimes.
The metabolic syndrome was modeled by keeping the hamsters on a high-calorie (hypercaloric) diet rich in energy (containing 29% of fats – mainly saturated lipids) and fructose (1 g a day per 100 g of the body weight) as an aqueous solution for 6 weeks [8].

Hamsters were divided into the following experimental groups: group 1 – animals of the intact control (IC); group 2 – animals of the control pathology (CP), they were kept on a high-calorie diet for 6 weeks; groups 3 and 4 – animals that starting from the 4th week of the experiment were injected intragastrically with an aqueous solution of the ginger extract in the doses of 50 and 80 mg/kg for 14 days (IMB50, IMB80) against the background of a high-calorie diet; groups 5 and 6 – animals that starting from the 4-th week of the experiment against the background of a high-calorie diet were injected intragastrically with metformin from the group of biguanides in the dose of 60 mg/kg for 14 days.

On the 1st day before the start and after 6 weeks of the experiment the body weight of animals was measured.

The state of the carbohydrate metabolism against the background of the model pathology was assessed by the level of basal glycemia and basal insulinemia [9].

The glucose concentration in the blood serum of animals was determined on an empty stomach by the glucose oxidase method using a “One touch ultra-easy” glucose meter (manufactured by LifeScan, Johnson & Johnson, USA).

The concentration of insulin in the blood serum was calculated based on the values of glucose and insulin levels is considered as a manifestation of insulin resistance. To confirm it the HOMA-IR index was calculated, it increased significantly by 2.33 times compared to IC (Table).

It is known that the simultaneous increase of glucose and insulin levels by 3.0 and 1.63 times, respectively, were developed compared to IC (Table). It is known that the simultaneous increase of glucose and insulin levels is considered as a manifestation of insulin resistance. To confirm it the HOMA-IR index was calculated, it increased significantly by 2.33 times compared to IC (Table). It indicated formation of insulin resistance as one of the main pathogenetic links of MS.

Results and discussion

It is known that consumption of high calorie food leads to obesity that is closely associated with the risk of developing MS and type 2 diabetes. Keeping animals on a hypercaloric high-fructose diet for 6 weeks resulted in obesity since the increase in the body weight in CP hamsters was significantly 3 times greater than the same indicator in IC group (Table).

At the same time, in CP animals the severe hyperglycemia and hyperinsulinemia with a significant increase of glucose and insulin levels by 3.0 and 1.63 times, respectively, were developed compared to IC (Table). At the same time, in CP animals the severe hyperglycemia and hyperinsulinemia with a significant increase of glucose and insulin levels by 3.0 and 1.63 times, respectively, were developed compared to IC (Table). At the same time, in CP animals the severe hyperglycemia and hyperinsulinemia with a significant increase of glucose and insulin levels by 3.0 and 1.63 times, respectively, were developed compared to IC (Table).

The statistical processing of the results was performed on a personal computer using Excel packages and Statistica 6.0 for Windows. The Newman-Keys and Mann-Whitney tests were used to determine the probable differences between the experimental groups. Differences were considered statistically significant at p < 0.05.

Table

| Indicator                      | IC      | CP      | IMB50   | IMB80   | “Arphasetin” | Metformin |
|-------------------------------|---------|---------|---------|---------|--------------|-----------|
| Glucose, mmol/l               | 5.18 ± 1| 13.57 ± 0.55* | 9.59 ± 0.57** | 6.70 ± 0.52** | 8.88 ± 0.52** | 6.40 ± 0.21** |
| Insulin pmol/l                | 82.55 ± 2.32 | 134.18 ± 2.22* | 113.17 ± 3.44* | 90.12 ± 1.25**/a | 95.93 ± 1.89**/a | 89.03 ± 1.25**/a |
| HOMA-IR                       | 1.55    | 3.61*   | 2.68*   | 1.93**   | 2.21**/a     | 1.81**    |
| HbA1c, %                      | 5.9 ± 0.22 | 9.8 ± 0.53* | 7.3 ± 0.43**/a | 6.9 ± 0.28** | 7.3 ± 0.52**/a | 7.2 ± 0.48**/a |
| Glycogen of the liver, mg/100 g of the tissue | 2326.5 ± 219.41 | 1098.3 ± 103.94* | 1873.5 ± 304.62** | 2146.2 ± 251.18** | 1439.5 ± 244.57**/a | 2165.0 ± 275.34**/a |
| Increase in the body weight within 6 weeks, g | 30.5 ± 3.3 | 92.3 ± 6.3* | 44.5 ± 4.2**/a | 34.5 ± 5.3**/a | 54.3 ± 7.3**/a | 33.5 ± 5.2**/a |

Notes: * – statistically significant difference in relation to the values of IC group (p ≤ 0.05); ** – statistically significant difference in relation to the values of CP group (p ≤ 0.05); a – statistically significant difference in relation to the values of the group taking “Arphasetin” (p ≤ 0.05).
Disorders of glucose utilization in CP hamsters were also characterized by a decrease in the glycogen stores in the liver by 2 times. It confirms development of liver resistance to the insulin action, which results in the processes of glycogenolysis and gluconeogenesis (Table).

On the background of chronic hyperglycemia the increase of HbA1c content by 66.1 % was observed compared to the IC indicators. It indicated a high level of glycosylation of proteins in this model of MS.

Therefore, changes in the indicators characterizing the state of the carbohydrate metabolism of golden hamsters kept on the high-calorie diet indicate simulation of the MS state characterized by hyperglycemia, compensatory hyperinsulinemia, insulin resistance, increased glycogenolysis in the liver and glycosylation of proteins.

According to the data presented in Table the body weight increased in all experimental hamsters, but dynamics of its increasing in animals receiving IMB in the dose of 80 mg/kg and metformin did not statistically differ from the group of IC. The ginger extract in the dose of 50 mg/kg moderately inhibited development of obesity in a similar way as the reference drug “Arphasetin”.

The use of IMB for 14 days simultaneously with the hypercaloric diet was accompanied with a possible decrease in the indicators reflecting the state of the glycemic profile. It should be noted that the maximum therapeutic effect was observed with the use of IMB in the dose of 80 mg/kg. Thus, the intragastric introduction of IMB in the dose of 80 mg/kg was accompanied by a significant reduction in the severity of hyperglycemia by 43.3 % compared to CP animals at the metformin level, while using it in the dose of 50 mg/kg resulted in decreasing of this indicator by 29.2 %. The level of glucose at the end of the experiment in the groups of IMB80 and metformin, unlike IMB50 and “Arphasetin”, did not significantly differ from the value of IC although it did not normalize completely. The data obtained reflect the expressive acceleration of glucose utilization under the effect of IMB80 in the conditions of the experimental MS.

There is clinical evidence that hyperinsulinemia is an independent risk factor for coronary heart disease [10]. Insulin has a direct atherogenic effect on the walls of vessels, adding a significant contribution to development and progression of atherosclerosis. Under the effect of IMB, probably due to decrease of hypoglycaemia, hyperinsulinemia decreased too, it was confirmed by decrease in the insulin level by 15.7 % compared to IC in case of using IMB in the dose of 50 mg/kg and by 32.8 % in case of introduction of IMB in the dose of 80 mg/kg. The process of normalizing insulin in the group of IMB80 was the same as in the metformin group and significantly exceeded the effect of “Arphasetin” and IMB50.

Confirmation of the corrective effect on insulin resistance manifestations was a significant decrease in the HOMA-IR index by 46.5 and 49.8 % in hamsters receiving IMB in the dose of 80 mg/kg and metformin in animals of the CP group. The percentage of HOMA-IR decreased under the effect of IMB in the dose of 50 mg/kg and “Arphasetin” compared to CP, and it was 25.7 and 38.8 %, respectively.

The determination of HbA1c is considered to be a criterion for normalization of the carbohydrate metabolism and the effectiveness of the metabolic control, which should be at the level of 6.5 %, and if the potential hypoglycaemic agent is used, it is the confirmation of its ability to reduce the risk of micro- and macrovascular complications of diabetes [11]. Under the conditions of our experiment the introduction of IMB in the dose of 50 mg/kg was accompanied with the decrease in the HbA1 level by 25.6 % compared to “Arphasetin”, while decrease of this indicator by 29.6 % when using IMB80 was consistent with the metformin effect.

Consequently, IMB did not differ from metformin and exceeded the effectiveness of the herbal drug “Arphasetin” by the effect on the level of glucose, the HOMA-IR index and HbA1c. It is probably due to the powerful complex pharmacological action of phenolic compounds of ginger – gingerols and other components having hypoglycemic, antioxidant, antidiabetic properties according to the literature data [3-5].

The ability to restore the glycogen-forming function of the liver IMB in the dose of 80 mg/kg was on the same level with metformin and reliably exceeded the effect of “Arphasetin”.

Thus, the results obtained indicate the ability of IMB to normalize the carbohydrate metabolism, reduce manifestations of insulin resistance under the conditions of the experimental MS and confirm the feasibility of its further study as an antidiabetic drug.

CONCLUSIONS

1. Keeping hamsters on a hypercaloric high-fructose high-calorie diet for 6 weeks led to manifestations of the metabolic syndrome, it was confirmed by development of hyperglycemia, hyperinsulinemia, the increase of the HOMA-IR index, the level of HbA1c and the decrease of the liver glycogen content.

2. The use of the ginger extract in the dose of 80 mg/kg normalized insulin glycaemia, decreased insulin resistance restored the glycogen content in the liver at the level of metformin. By the intensity of the pharmacological action the ginger extract exceeded the reference herbal drug “Arphasetin”.

3. The results obtained substantiate the further pharmacological study of the ginger extract as a promising agent for the treatment of the metabolic syndrome and type 2 diabetes.

Conflict of Interests: authors have no conflict of interests to declare.
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