PHYTOCHEMICAL AND PHARMACOLOGICAL STUDY OF POLYSACCHARIDE COMPLEXES OF PRUNUS DOMESTICA FRUIT

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1. Introduction

Nowadays, agricultural crops became the main focus of researchers due to fact they are a good sources of biologically active substances, yielding a sufficient raw material base. Most of them have been used in traditional medicinal practices since the prehistoric times. Phytochemical study of plants is used to find potentially promising plants for the development of new drugs. Plum (Prunus domestica, family Rosaceae) is widely cultivated in Ukraine as horticultural crops. Plum has long been used in traditional medicine and its usage is still actual [1].

2. Formulation of the problem in a general way, the relevance of the theme and its connection with important scientific and practical issues

During our preliminary study from dried plum fruits, polysaccharide complexes, water and alcohol extracts were obtained. Laxative activity of all obtained substances was studied by the Sticknay J. S. method [2]. Further investigation of polysaccharides complexes isolated from fresh plum fruits will help expand the raw material base and are necessary to determine constituents, safety and efficacy of the most effective extracts.

3. Analysis of recent studies and publications in which a solution of the problem and which draws on the author

The important role of fruits in human health and nutrition has been better understood with the recent studies on biochemical contents of fruits having antioxidant properties. In recent studies and publications total antioxidant capacity, phenolic compound, organic acid, and vitamin C contents of different plum species were determined and the correlation between the measured values was investigated [3, 4]. Analysis of phenolic compound indicated that chlorogenic acid was the predominant phenolic compound [3]. Laxative effect of different extracts from dried plum was confirmed [4]. For the standardization of plum fruits as herbal raw material, according to the methods of the State Pharmacopoeia of Ukraine the indexes of fresh plum were determined [5, 6].

Therefore, further study of influence of fibers and WSPC on histostucture of liver of rats in conditions of ethanol intoxication was actually, interesting and expedient.
4. Allocation of unsolved parts of the general problem, which is dedicated to the article

Plum is good sources of biologically active substances. At the base of review of modern research and our preliminary study it was confirmed, that plum – is rich on hydroxycinnamic acids, organic acids and polysaccharides. Moreover cathartic effect for row of isolated complexes were determined. Therefore, study of polysaccharide complexes in detail is prospective in development of new active pharmaceutical ingredients and creation of methods of quality control.

5. Formulation of goals (tasks) of article

The aim of our research was fractionation of polysaccharide complexes from fresh plum fruits, comparative study content of neutral sugars in it, element composition of plum fruit and investigation of influence of fibers and water soluble polysaccharide complex (WSPC) on histostructure of rats liver in conditions of ethanol intoxication.

6. Statement of the basic material of the study (methods and objects) with the justification of the results

In August 2017 the plant material (fresh plum fruits of varieties "Ugorka") were purchased on the market in Kharkiv, Ukraine. Plant raw material was pitted and grinded to the puree. Determination of the loss of drying of plum fruits were carried out according to the SPU requirement [6].

Three polysaccharide substances as fibers, WSPC and pectin were isolated from plum raw material sequentially.

For the isolation of fibers, hot water was added to the 100 g of pitted crushed plum fruit in the ratio 1:5, stirred for 1 hour at room temperature. Fibers were centrifuged (speed of rotation of 5000 rpm) during 10 minutes. Obtained fibers were dried in an oven to constant weight at the temperature 100–105 °C and yield was determined.

Water extract were decanted in a flask, evaporated to 50 ml and placed in a flask with capacity of 500 ml, after that 150 ml of 96 % alcohol was added and mixed. Precipitate (WSPC) was filtered through a paper filter. The filter with the WSPC was dried firstly in air and then to constant weight at the temperature 100–105 °C and yield was calculated.

The water extract remaining after receiving WSPC was evaporated to 50 ml and used to obtain the pectin fraction. Pectin was extracted with a mixture of 0.5 % solution of ammonium oxalate and 0.5 % solution of acid oxalate (1:1) at a ratio of 1:20. Extraction was performed twice at a temperature of 80–85 °C during 2 hours. The resulting extract was filtered, pooled and evaporated to 1/5 of the original volume. Pectin was precipitated by three volumes of 95 % ethanol, and settling for 12 hours in a cool place. The precipitate was filtered, washed with 95 % ethanol and dried in a drying oven to constant weight. Yield of pectin was determined.

For comparative study of obtained polysaccharide complexes 0.5 g of each obtained fractions were hydrolysed with a hydrochloric acid concentrated for 2.5 hours. The solutions were cooled and quantitatively transferred with water to volumetric flasks with a capacity of 25.0 ml, adjusted to the mark with the same solvent and mixed. Then were taken 5 ml from each obtained solution and neutralized by the solution of 30 % sodium hydroxide with universal indicator paper. The neutralized solution was filtered through a paper filter, transferred quantitatively to a 25.0 ml volumetric flask, diluted to volume with water, and stirred.

Then 2–5 ml of the solution were taken from the each volumetric flask into other 25.0 ml capacity volumetric flasks, 1.0 ml of 1 % picric acid, 3.0 ml of 20 % sodium carbonate were added in each flasks and heated at 100 °C for 20 minutes. After cooling, the volume was adjusted with water and stirred. In parallel, under the same conditions, 2.0 ml of a standard sample (SS) of glucose was prepared. The optical density of the test and glucose SS solutions were measured on a Hewlett Packard 8453 spectrophotometer at a wavelength of 463 nm in a cuvette with a layer thickness of 10 mm. A mixture consisting of 1.0 ml of 1 % picric acid, 3 ml of 20 % sodium carbonate and 1.0 ml of water was used as the reference solution [7].

The study of the content of macro- and microelements is important for further standardization and the development of quality control procedures. The elemental composition was determined with atomic emission spectrophotometer at State Scientific Institution “Institute for Single Crystals” of NAS of Ukraine. To obtain spectra and their registration plate spectrograph DFS-8 ISE 3500 (Thermo Scientific, USA) with a diffraction grating grating 600 line/mm and Trilinz Slot Lighting was used. The sample was evaporated from the craters of graphite electrodes in the discharge arc of an alternating current of 16 A with an exposure of 60 seconds. An excitation source of spectra, IWS-28 was used. Measuring intensities of lines in the spectra of analyzed samples and calibration samples was carried out with micro photometer MF-1 [8, 9].

The material for histological examination was the liver of rats after 7–10 days of intraperitoneal injection of 40 g of ethyl alcohol in a dose of 7 ml / kg – control pathology; liver of rats, which daily 1 hour after the introduction of ethyl alcohol for 7–10 days, received plum fibers or plum WSPC in a dose of 200 mg / kg, or Silibor (preparation of comparison) in a dose of 30 mg / kg; liver of intact rats - intact control. The withdrawal from the experiment of rats of all groups was carried out on the second day after the completion of the introduction of ethyl alcohol and investigated objects. The obtained liver samples from each rat were divided into several parts. Some samples, after fixation in a 10 % solution of formalin, were dehydrated in alcohols of increasing strength, poured into paraffin, transections were cut on a snuff-microtubule, stained with hematoxylin and eosin. Other samples, after fixation in formalin solution, were cut on a freezing microtome and sections were dyed with Sudan IV to detect neutral fat [10].

Illustrative material was prepared using light microscope Granum with digital camcorder Granum DCM 310. The photos were processed on a Pentium 2.4GHz computer using the Toup View program.
Loss of drying of fresh plum fruits were determined as 76.89±2.26 %.

The results of determination of the content of polysaccharide fractions obtained from the fresh plum fruits are given in Table 1.

### Table 1

<table>
<thead>
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<th>Polysaccharide fractions</th>
<th>Yield, %</th>
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<tr>
<td>Fiber</td>
<td>2.20±0.05</td>
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<tr>
<td>WSPC</td>
<td>1.13±0.05</td>
</tr>
<tr>
<td>Pectin</td>
<td>0.97±0.05</td>
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Obtained fiber was brown powder, WSPC – light brown and loose. The pectin was grayish brown and powdery texture.

The most content of neutral sugars were determined in the WSPS – 61.52±1.47 %. Their content in other complexes was: in fibers – 59.23±1.15 %, pectin – 22.85±0.55 %.

According to the analysis of the element composition, potassium has the most content in fruits – 2000 μg/100 g. The content of other elements was (μg/100 g): calcium – 52, magnesium – 70, sodium – 50, phosphorus – 30, silicon – 15, aluminum – 1.5, iron – 1.0, zinc – 0.2, strontium – 0.8, copper – 0.25, manganese – 0.5, molybdenum – 0.05, lead <0.03, nickel <0.03, Co <0.03, Cd <0.01, As <0.01, Hg <0.01.

Content of elements was calculated on dried raw material. Our research determined that the content of heavy metals in the raw materials does not exceed the norms established by the State Pharmacopoeia of Ukraine.

According to our result it was confirmed that plum is an important source of potassium, which is known as a systemic electrolyte, regulator of water-salt metabolism, acid-base state of the organism; it promotes removal of edema; activates enzymes; also it is necessary for normal muscle activity, in particular, for the proper functioning of the heart, improves the functioning of the myocardium in the case of metabolic disorders [1]. The presence and content other elements also play role for health. For example, copper maintains the normal blood composition, is contained in enzymes and participates in the delivery of oxygen to the cells. Zinc strengthens immunity, is important for growth, and supports the hormonal background. Magnesium has an antispasmodic and antiplatelet effect [1].

As microscopy showed, the liver of intact rats had a typical structure inherent in this species of animals. Connective tissue layers between the lobes were not expressed. The boundaries of the particles were determined by triads - portal paths (passageways of the branches of the hepatic artery, reverse vein and bile duct). The triad zones themselves are narrow. The hepatocytes (liver proteins) in the lobes had a clear radial orientation. Hepatocytes in different parts of the liver lobules were of a characteristic shape and size, the cytoplasm was uniformly stained, optically dense, and did not contain inclusions visible in light microscopy. The nuclei of hepatocytes were normochromic, centrally located, containing 1–2 nucleoli. The number of dual-core hepatocytes was sufficient. The intracellular hemocapillaries were moderately enlarged, containing the usual number of lymphoid cells. Kupffer cells without features (Fig. 1). Sudan coloring did not reveal the accumulation of fat in the cytoplasm of cells (Fig. 2).

Fig. 1. Liver of an intact rat. Normal state of the liver parenchyma. Hematoxylin-eosin. ×250

Fig. 2. Liver of intact rat. Absence of fat in the cytoplasm of hepatocytes frozen transection, Sudan IV. ×250

Sinusoidal hemocapillars are unevenly enlarged, cities are full-blooded. Hepatocytes and their nuclei vary somewhat in size. Cell nuclei were slightly hyperchromic, nucleoli differed not clearly. Kupffer cells were activated in cites. Vacuolation of cytoplasm of hepatocytes was diffusely or large focal part of the lobules. Inflammatory reactions, necrosis of hepatocytes are absent (Fig. 3). Sudan coloring revealed the fatty nature of vacuoles. Fat drops had a small drop nature, did not destroy the integrity of the cells, did not cause the dislocation of the nucleus (Fig. 4).

The microscopic picture above corresponds to parenchymal fatty degeneration of the hepatic parenchyma. Taking into account the previous “alcoholization” as an experimental condition, this picture may be a manifestation of the very initial stage of alcoholic hepatopathy [11].

Simultaneous administration ethanol and fibers of plum fruit positively influenced on the condition of the liver parenchyma. Hepatocysts contained juicy nuclei, nucleoli were clear, less pronounced anisonuclease. The vacuolation of cells was clearly reduced, with a small
focal nature. The parenchyma of the particles was predominantly normal in shape (Fig. 5).

Sudan coloring showed that fatty drops in the cytoplasm of hepatocytes were very small, often had a dusty character (Fig. 6).

Introduction to the background of alcohol of the WSPC of plum fruit visually had little effect on the expressiveness of the vacuolization of hepatocytes compared to the control pathology and the state of the hepatic parenchyma in general. The processes of microcirculation in the lobes are also comparable (Fig. 7). Fat drops did not break either the beam pattern or cell integrity (Fig. 8).

Comparison drug silibor also did not prevent the occurrence of fatty dystrophy of hepatocytes in rats with alcoholic loading. The expressiveness of changes was practically at the level of the last (Fig. 9, 10).

Thus, a light-optical study showed that after 7–10 days of administration of 40 ml of ethyl alcohol in rats at a dose of 7 ml / kg in the rats liver, morphological signs of parenchymal fatty degeneration appeared, which may be the very initial stage of alcoholic hepatopathy development.
Simultaneous administration of alcohol and fiber of fruit plums in a dose of 200 mg/kg prevents the development of fatty dystrophy of the liver parenchyma, in contrast to the similar scheme of the introduction of a WSPC of plum fruit in the same dose or drug comparison of silibor at a dose of 30 mg/kg.

7. Conclusions
1. Investigation of element composition of plum fruits was carried out.
2. Polysaccharide complexes of plum fruits were obtained and content of neutral sugars were determined there.
3. It was confirmed that simultaneous administration of alcohol with the fibers of plum fruits in a dose of 200 mg/kg prevents the development of fatty dystrophy of the liver parenchyma, in contrast to the similar scheme of the administration of the WSPC of plum fruit in the same dose or drug comparison silibor at a dose of 30 mg/kg.

Considering results of phytochemical and pharmacological research, we can assume that the plum fruit fiber is promising for further study in order to create a new effective and safe drugs for use in medical practice.

References
DYNAMICS OF EXCEPTION OF ORGANIC ACIDS FROM MIXTURE OF MEDICAL VEGETABLE RAW MATERIAL

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Introduction

Globally, there is a clear trend to higher level of psychopathological disorders, in particular a variety of psychogenic neurotic disorders. In Ukraine, the situation is complicated by various social-psychological problems, global informational oversaturation, chronic fatigue, environmental problems, the decline in the quality of life. All this leads to the onset of stress which manifests itself as fatigue, decreased performances, irritability, anxiety, worsened mood, disrupted sleep, etc. For the treatment of neurotic conditions, the most effective means are the phytopgenic sedatives of plant origin due to the similarity of many biochemical processes in the cells of plant and animal origin. This property becomes particularly important when it concerns treatment and prophylactic means. The particular interest of doctors and patients in these sedatives can be explained by the minimum of contraindications and side effects [1].