COMPARATIVE STUDY OF TECHNOLOGIES FOR EXTRACTION OF BIOLOGICALLY ACTIVE SUBSTANCES FROM THE RAW MATERIAL OF ANIMAL ORIGIN

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Abstract
Technologies of isolation and concentration of biologically active substances, developed in the middle of the 20th century, need adjustment and adaptation to modern conditions both to increase the activity of substances and for greater economic efficiency. The aim of the research is the comparison of dynamics of biologically active compounds extraction from porcines pancreas in two methods: the saline method based on 0.9% sodium chloride solution, and the acidic method based on 2.4% trichloroacetic acid solution. Also the purpose of research is to assess the possibilities for further optimization of technologies. The total protein concentration based on the biuret reaction in the samples taken during the extraction, as well as the calculation and analysis of the point degrees of separation of substances, was determined. The predominant region of molecular weight was identified. The presence of some proteins absent in the final saline extract was noted. Also the proteomic profile of the extracts was studied. Wide range of compounds with molecular weight of less than 52 kDa was found in extracts based on physiological saline solution, and protein substances of whole presented range of molecular weights in trichloroacetic acid based extracts were considered. The predominance of low molecular weight protein fraction of interest was noted also in this method of extraction in comparison with the other methods of extraction. According to the UniProt database, we assume availability of probable compounds with a molecular weight of less than 30 kDa in the purified acidic extract. The presence of some proteins absent in the final saline extract was noted. The acidic erythrograms showed a weak degrading effect of both types of extracts on the membranes of rat erythrocytes, as well as the cytoprotective effect of acidic ultrafiltrates (less than 3 kDa). The obtained results prove a better efficiency of trichloroacetic acid extraction method used for obtaining a mixture of a wide range of compounds, including biologically active substances of low molecular weight.

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Introduction
The study and optimization of technologies for isolation of biologically active compounds from animal raw materials are today one of the most important aspects of modern biotechnology. Along with the fast development of the analytical chemistry and biochemistry spheres, the technological execution of methods, aimed to supervision and control of the observed processes, requires timely improvement to match the progress. In this regard, the permissible limits of sensitivity of the detection devices are improved, devices are designed with functional possibility to run combined experiments (for example, HPLC with a diode-matrix UV-visible spectrophotometric detector (DMD), fluorimetric MS-detector, capillary electrophoresis with DMD [1], low molecular weight electrophoresis combined with mass spectrometry of inductively coupled plasma [2]). The databases of discovered compounds are being replenished more and more (for example, refer to the UniProt protein database) [3]. The early studies, replicated on the currently obsolete equipment, were as accurate as it was possible for that time. Technologies based on such researches are still used today. By the way, these technologies often feature high cost, low efficiency and high labor intensity. In order to potentially reduce the economic costs of production, as well as to reduce the volume of waste, the dynamics of the process flow shall be studied in order to identify possible ways to optimize the methods.

In the 1920s insulin, pancreatin, glucagon, obtained from animal pancreas extracts, were patented. Later, the development of technologies for production of certain medical preparations of enzymes and hormones, extracted from porcine pancreas, went on. Pigs were used as a source of a whole range of important biologically active substances, and in the 30s of the last century, traditional approaches to the extraction of some specific compounds or their mixtures from the pancreas were developed. Those achievements serve as the basis of modern technologies [4,5]. Due to the rapid development of the technological execution of necessary compounds isolation, it can be assumed that the potential of this raw material is not fully achieved.
Objects and methods
The object of the study was porcine pancreas *Sus scrofa*, sampled at Limited Liability Company “Pushkinskiy Dvor” (Russia, Pushkino). The raw materials were subjected to extraction by two methods: aqueous saline (physiological saline) and acidic (trichloroacetic acid), followed by study of obtained extracts and their ultrafiltrates.

The raw materials were frozen and stored at a temperature of minus (40 ± 2) °C with further thawing immediately before the study at a temperature of (22 ± 2) °C. Before extraction, the pancreas was also subjected to fine grinding with the help of a meat grinder (Kenwood, England): mesh of 100 holes with a diameter of 3 mm. The extraction scheme is shown below in Figure 1.

Saline extraction [7] was conducted in 0.9% sodium chloride solution by way of mixing of the finely chopped raw materials and an extractant in hydromodule in ratio of 1:5. The process lasted for 330 minutes. After completion of the extraction, the obtained solution was centrifuged in a Centrifuge CM-6M (ELMI, Latvia) at 3500 rpm for 8 min. at a temperature of (22 ± 2) °C. Then, the supernatant liquid, separated from the sediment, was saved for further analyses. To study the extraction process, samples were taken throughout the entire process: at 0, 5, 10, 15 minutes; and further every 15 minutes till to 330th min. Each sample was centrifuged for 5 minutes immediately after its sampling at a temperature of 4 °C on a Centrifuge 5427R (Eppendorf AG, Germany) at a speed of 3500 rpm; the supernatant liquid was frozen.

In case of acid extraction [8,9], the finely chopped raw material was loaded into the processor bowl together with an extractant — 2.4% trichloroacetic acid solution with a hydromodule of 1.3:15. The acid extraction technology also included the stage of the final extract purification from ballast proteins: primary centrifugation was run in a Centrifuge 5427R (Eppendorf AG, Germany) at 6,861 rpm for 10 minutes with cooling the extract to 4 °C; the process also included the precipitation with sodium hydroxide solution at pH 6.2–7.2. The final centrifugation was run at 3,500 rpm for 10 minutes on a Centrifuge CM-6M (ELMI, Latvia) at a temperature of (22 ± 2) °C. Extraction was conducted in the same laboratory installation, that was used for saline extraction, for 2 hours 15 minutes with sampling at 0, 5, 15 minute and then every 15 minutes till to 135 min. Each sample was centrifuged at 6,861 rpm for 10 minutes. when cooled to 4 °C. The supernatant liquid was frozen at a temperature of minus (40 ± 2) °C.

In both cases, the extraction process was conducted by the laboratory unit in LDU (Labotex, Russia) along with cooling down to (4 ± 2) °C in order to avoid autolysis, at a stirring speed of 400 rpm. The supernatant liquids of the purified extracts obtained after centrifugation were frozen at a temperature of minus (40 ± 2) °C and stored for no more than a month.

To obtain a highly active low molecular weight fraction of the extracts, the extracts underwent ultrafiltration on filters with a permeability limit of 3 kDa (Amicon, Ireland) as the final stage of the technology process. The process was carried out on a Centrifuge 5427R (Eppendorf AG, Germany) at a speed of 11,481 rpm for 20 minutes at a temperature of 4 °C.
In the samples taken during the extraction, the total protein concentration, the total proteolytic activity of enzymes, and the dynamics of the yield of protein compounds were analyzed by the proteomic profile. The membranotropic effect of biologically active compounds in primary purified extracts, as well as in their ultrafiltrates were studied.

Sample of the raw material (pancreas native further PN) were prepared in a known way [7].

The total protein concentration and ultrafiltrates in the extracts were measured on the basis of the biuret reaction according to the Kingsley-Weixelbaum method [10]: 600 µl of the biuret reagent was incubated with the test sample, and after 10 minutes the optical density of the solution was measured at 540 nm by a photometer BioChem SA (HTI, USA).

On the basis of parameters of protein yield to the extractant the basic characteristics of the process like degree and rate of extraction were determined.

The degree of extraction was determined as the ratio of the extracted substance (protein) mass to the total mass of raw material and was calculated by the following formula:

$$D = \frac{M}{m} \times 100 \%$$  \hspace{1cm} (1)

where

$M$ — is the mass of the extracted substance, g;

$m$ — is the total mass of the initial raw material in the mixture, g.

**Figure 1.** Scheme of extractions based on saline and trichloroacetic acid

Legend: CP1 — control point for native pancreas sampling to obtain a comparative sample (CS); CP2 is the control point for sampling of extracts to study the dynamics of both extraction methods; CP3 — control point of the final extract sampling: after centrifugation of the total volume of the extract — for saline extraction; after purification and secondary centrifugation — for acid extraction.
The point rate of extraction was determined by change in protein concentration (extracted substance) along the vector of time. It was calculated at each moment of the extraction process. Calculations were conducted by the formula:

\[ v = \frac{\Delta c}{\Delta t} = \frac{dc}{dt} \]  

(2)

where

- \( c \) — is concentration of protein in extract, g/l;
- \( t \) — is corresponding time of the extraction, min.

The total proteolytic activity of the final extracts was determined by Leilian-Folgard method, which included heating in a test flask up to 37°C in a water bath of 20 ml of 5% alkaline casein solution with 10 ml of the test sample for 60 minutes. In a control flask, a solution of the same composition, immediately after mixing was precipitated first with a 0.2 N hydrochloric acid solution, and then with a 15% sodium sulfate solution. The precipitated fallout was filtered off. After one hour of sample exposure in a test flask, the same operations were carried out. Then 10 ml of the obtained filtrate was transferred into clean flasks and titrated with 0.1 N sodium hydroxide solution in presence of 1% cresol red solution till obtaining the saturated bright crimson color. The proteolytic capacity was calculated from the difference in the volumes of sodium hydroxide required for titration of the experimental sample and control sample.

Membranotropic activity of biologically active compounds in extracts and ultrafiltrates was determined by constructing acidic erythrograms of erythrocyte membrane degradation [11, 12]. 10 ml of the test sample with a protein concentration of 1000 ng/ml on the basis of 0.9% sodium chloride solution that was introduced into a test flask. The control flask contained pure physiological saline. In both cases, 20 μL of blood from rats’ tails vein was added to the flasks, the contents were mixed for uniform contact of active compounds with erythrocytes and left to incubate at room temperature for 1 hour. After expiration of this time, 2 ml of both samples were transferred into clean flasks and mixed with 2 ml of 0.004 N sodium chloride-based hydrochloric acid solution. The dynamics of changes in optical density was recorded by SF-2000 spectrophotometer (Spekttr, Russia) at 650 nm every 15 seconds from the moment of mixing until the end of hemolysis (when the optical density values reach a plateau), i. e. for at least 10 minutes.

The proteomic profile of the extracts was studied on the basis of electropherograms obtained by Laemmli method, i. e. by one-dimensional denaturing electrophoresis in 12.5% polyacrylamide gel in presence of SDS. The process was run in a VE-10 chamber (Helicon, USA) at a temperature of (22±2)°C at 60 V, until the samples reached the boundary of the separating gel, then at 130 V until the electrophoresis was completed. Standards (Fermentas, Lithuania) were used as comparison referential markers.

The analyzed samples were primary stained with Coomassie Brilliant Blue G-250 solution. Later the stain was removed by 10% acetic acid. To increase the sensitivity level of this method, silver staining was performed according to Bloom’s method by alternate incubation of the samples in solutions of sodium hyposulfite, of silver nitrate and sodium carbonate.

Density diagrams were obtained by the software ImageJ (National Institutes of Health, USA) [13] based on results of studying the proteomic profile of extracts selected during the extraction process at control points. The results were visually presented on the basis of automatic densitometric analysis of the scanned electrophorograms. In order to avoid errors in study of various extractions electrophrogram, the same parameters were set up. The image was converted to 8-bit type, then an area of 1280x1938 pixels was selected on the track. The molecular weight of the fractions was determined by correlating of the corresponding peaks on the density diagram of the test sample with the peaks of the standards [14,15].

The software STATISTICA 10.0 was used for statistical processing of the total protein content. The final results were calculated and presented as “mean ± standard error” (M ± SE). Significant differences were checked using one-way analysis of variance ANOVA followed by Tukey’s test. The deviations, when P values didn’t exceed 0.05, were considered as statistically significant [16].

Results and discussion

When measuring the total protein concentration (Figure 2) in the samples taken during the saline extraction, a sharp increase in the values from 0 to 5 minutes was observed. And the further decrease in the protein content in the samples was observed at 90th min. (21.2 ± 2.7 g/l), at 165th min. (19.6 ± 0.4 g/l), at 240th min. (19.7 ± 0.2 g/l), at 270th min. (19.5 ± 0.3 g/l). The observed maximum concentration values are explained by release of a new fraction of compounds into the extractant, for example, at 60th min. (23.1 ± 0.9 g/l), which together with the concentration value for 135th min. (23.3 ± 0.6 g/l) was the highest throughout the entire process of extraction. Minor increases in concentration were also observed at 195th and 255th min. In general, it is necessary to note the smooth way of change in the samples protein content, starting from 5th min. of the process.

TCA extraction featured a smooth increase in total protein content up to 75th minute, where it reached a maximum and amounted to 15.6 ± 0.4 g/l. A further slight decrease in concentration led to coming of protein amount, released to extractant, to a plateau. This plateau values were was observed until the very end of the extraction process.

According to the diagram of dependence of the total protein concentration on time of extraction, in case of saline extraction a higher yield of protein into the extractant is evident throughout the entire process. From the first moments of the process, a sharp increase in the amount of protein was observed in comparison with a smooth increase of protein concentration in case of acid extraction. Both methods of extractions feature an oscillatory process of the protein content changing in the extracts.
As result of processing the data on total protein concentration in the extracts, the degree (D) and point rate (v) of extraction were calculated for both methods. Diagram of the dependence of rate of proteins release into the extractant are shown below in Figure 3. It is necessary to note the high rate of saline extraction till the 5th minute, in contrast to TCA extraction, in which the rate gradually increased up to the 15th minute and then began to decrease uncritically. In both cases, the process showed an oscillatory change in the rate within the range of 0 g/(l*min), i.e. in general the release of protein into the extractant took approximately constant values with minor deviations. Thus, there was a slight decrease in rate of extraction at the 165th and 270th minutes in saline extraction, and at the 90th minute in case of TCA extraction.

The data on degree of extraction are shown above in the Table 1. It is necessary to note the correlation with the data on the total protein content in the extracts. The final degree of extraction of the protein component in the saline variant exceeded 2.25 times the value of D for acid extraction. It proves that the saline solution is more efficient as an extractant in terms of this regard. The highest degree of extraction (Table 1, light green line) was observed from the 30th till 150th minute, and at the 75th minute. (Table 1, red line) for extracts based on 0.9% sodium chloride solution and TCA, respectively.

The study of proteolytic activity of enzymes in the final extracts showed a higher value in the case of saline solution extraction and amounted to 94,276 U/ml in comparison with 33,748 U/ml for the TCA-based sample. The data indicate the best enzymatic activity in the saline extract, which is primarily associated with mild environmental conditions (neutral pH, the absence of denaturing agents that can negatively affect the active centers of enzymes, etc.). The low proteolytic capacity of the acid extract indicates an insufficient degree of purification of active enzymes from the inhibitory action of TCA, even after the completion of additional stages of purification. The reason may also be the removal of enzymes fraction together with the ballast proteins.
Table 1. Extraction rate of protein-peptide fraction in the process of saline and acid extraction

| D, %  | 1.30 | 11.17 | 10.63 | 10.40 | 11.11 | 11.03 | 11.33 | 10.61 | 10.34 | 10.55 | 10.56 | 11.25 | 10.73 |
|-------|------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
| Time, min | 0   | 5     | 10    | 15    | 30    | 45    | 60    | 75    | 90    | 105   | 120   | 135   | 150   |

| D, %  | 9.40 | 9.90 | 10.20 | 9.69 | 10.61 | 9.30 | 9.88 | 9.14 | 9.06 | 9.17 | 9.23 | 9.62 |
|-------|------|------|-------|------|-------|------|------|------|------|------|------|------|
| Time, min | 165  | 180  | 195   | 210  | 225   | 240  | 255  | 270  | 285  | 300  | 315  | 330  |

Table 2. Total protein concentration in extracts before and after ultrafiltration

| Extraction variation | Total protein concentration, g / l |
|----------------------|-----------------------------------|
|                      | Final extract | Ultrafiltrate |
| Saline               | 23.3 ± 0.6    | 6.03 ± 0.1    |
| Acidic               | 10.8 ± 0.1    | 8.04 ± 0.2    |

The results of the analysis of erythrograms obtained with acid inhibition of rat blood erythrocytes after its incubation with extracts (Figure 4) and their low-molecular-weight fractions (Figure 5) proved a short-term hemolysis and a tendency for a smooth decline in all samples.

precipitated at the stage of neutralization and centrifugation. It is necessary to keep in mind that a decrease in proteolytic activity may be a positive factor, indicating that many large and medium-sized proteins, that avoided the undesirable process of autolysis, are preserved in their native form.

To obtain information on the biological activity of the low molecular weight fraction, ultrafiltration was conducted. The data on the total protein concentration in the final investigated extracts and their ultrafiltrates are presented below in the Table 2.

![Figure 4](image1.png)

Legend: A-E — curve of incubation of erythrocytes with TCA extract; S-E — curve of incubation of erythrocytes with saline extract; C — curve of incubation of erythrocytes with saline solution without the sample

![Figure 5](image2.png)

Legend: LM A-E — incubation curve of erythrocytes with low molecular weight fraction of TCA extract; LM S-E — incubation curve of erythrocytes with low molecular weight fraction of saline extract; C — curve of incubation of erythrocytes with saline without a sample
Analysis of the diagram typical for incubation of erythrocytes with primary extracts (Figure 4) revealed particularly low optical densities for the TCA extract, which is explained by the presence in the sample of a large number of biologically active compounds that can penetrate through the membrane and accelerate the process of its degradation. In addition, this hemolysis curve is also spanned below the control sample curve, which confirms the overall negative effect of the extract on biological membranes. The results of determining the optical density of erythrocytes in the case of incubation with a saline extract were similar to the values in the control sample at the very beginning of the process and at the end of hemolysis. The decline in the diagram, that indicates the process of direct cells destruction, was found below the control curve and almost coincided with the values for the acid extract, which indicates a high degree of membrane degradation. Based on the results of the analysis, it can be concluded that the primary extracts are unable to provide cytoprotective effect on biological membranes, which is more profound in the TCA-based sample. In this regard, it is possible to assume the possible presence of highly active compounds in the samples, capable to transport substances through the membrane, which indirectly leads to its destruction.

A similar process was observed in the case of saline ultrafiltrate (Figure 5). The incubation diagram with erythrocytes also coincided with the control curve, which proves a weak membranotropic activity of the low molecular weight fraction in this sample. An obvious difference was observed when considering the acid ultrafiltrate curve. High values of the optical density of the sample based on TCA confirmed the cytoprotective effect of biologically active substances released into the extract, as well as an increase of erythrocyte membrane preservation level being exposed to degrading agent — hydrochloric acid.

Analysis of the proteomic profile of the final extracts (refer to Figure 6) showed the range of protein compounds on electropherogram at the final point of saline extraction (the 330th min. — Figure 6, No. 1) and track of the raw material (Figure 6, No. 2) corresponding to the pancreas sample, which was not subjected to extraction. Compounds within the range of 52 kDa and less were found in the extracts, which were discussed in detail in our previous research [7]. It should be noted that the fractional bands in this extract are more saturated than the primary electropherogram of acid extraction (Figure 6, No. 3, 4). In case of acid extraction only a few slightly expressed protein fractions were observed in the range of more than 250 kDa, 69–70 kDa, 52 kDa, 22–23 kDa and 15 kDa. The visual analysis of the native track of purified acid extract found no difference from the track of extraction 135 minutes long. The staining with silver (Figure 6, No. 5) revealed a wide profile of protein compounds in whole presented range of molecular weights. In the track of the purified extract (refer to Figure 6, No. 6), some medium-molecular fractions were absent: 70–95 kDa, 32–48 kDa; the fraction 15–16 kDa became less saturated, and the intensity of the low-molecular fraction of weight of less than 8 kDa increased slightly.

![Figure 6. One-dimensional electropherogram of porcine pancreas extracts](image)

Legend: ST — molecular weights standards: 250, 150, 100, 70 50, 40, 30, 20 15, 10 and 5 kDa; No. 1 — track of the endpoint of saline extraction; No. 2 — track of the sample not subjected to extraction (PN); No. 3 — track of the endpoint of acid extraction; No. 4 — track of purified acid extract; No. 5 — silver-stained acid extraction endpoint track; No. 6 — silver stained track of a purified acid extract.

The densitometric study of native electropherograms of acid extracts in the ImageJ software brought little information, since only 5 weak peaks were identified at the endpoint of extraction and in the purified extract too (Figure 7, No. 3, 4). After silver staining, a greater number of fractions were found on the corresponding tracks diagram within entire presented range of molecular weights. This range featured prevalence of fractions over 100 kDa, 83–90 kDa, 70–75 kDa, 43–65 kDa, 18–32 kDa, 14–16 kDa, less than 11 kDa. After purification of the extract some of the protein compounds were removed, and that provided better separation of the peaks in these specified ranges of masses. Thus, peaks at 137–140 kDa, 110 kDa, 100 kDa, 70 kDa, 52 kDa, 41–42 kDa, 27–31 kDa, and 21 kDa were clearly distinguishable. The absent peaks in the density diagram (49–50 kDa, 33–34 kDa, i6 and 11 kDa) most likely corresponded to the removed ballast proteins.

During saline extraction (Figure 7, No. 1) and analysis of the sample not subjected to extraction (Figure 7, No. 2), a large number of similar peaks were noted: a profound peak at 52 kDa, a row of clear minor peaks with varying degrees of manifestation within the range 32–47 kDa, at 30 kDa, the peaks at 24–25 and 29 kDa were expressed better in case of the saline extraction, as well as the low molecular weight fraction of 10–15 kDa which was better visible too.
Analysis of electropherogram and corresponding density diagram by ImageJ software showed a wide range of compounds in all types of extracts. In case of acid extraction the high molecular weight proteins prevailed, which was detected by silver staining. The saturation of the protein bands prevailed in the saline extract and the track of the initial PN sample, which protein bands correlated with the previously presented data on the total protein content. The analysis of dynamics of the biologically active protein substances extraction from porcine pancreas showed high values of the total protein concentration during saline extraction along with a wide range of compounds throughout the entire process. In case of acidic extracts the protein content values were lower; however, the proteomic profile of the samples taken during the process was characterized by a greater variety of fractions within the whole presented range of molecular weights. First of all, this may be explained by dependence of solubility [17, 18] and activity of many enzymes [19] secreted by the porcine pancreas on the acidity of the medium. Thus, as the pH approaches the isoelectric point of a protein, its solubility decreases when it reaches its minimum at pI value [20]. This indicator has long been established for many hydrolytic enzymes of the pancreas of cattle and pigs. It has been determined that most of the main enzymes are secreted by the organ under normal conditions in the form of pancreatic fluid. To the maximum extent, isoenzymes of α-amylase I and II, for example, precipitate at pH 6.5 and 6.1, trypsin at pH 10.2–10.8, chymotrypsin at pH 8.1, ribonuclease at pH 9.6 [6,21–23]. In this regard the acidic environment created by TCA provides better solubility and, accordingly, a greater variety of compounds in extracts. In saline samples, due to physiological solution, a neutral reaction of the medium is established, which leads, to varying degrees, to a decrease in the yield of some compounds into the extractant. The lower concentration of total protein in acid extracts samples is also associated with inhibition of many digestive enzymes, since the maximum activity of pancreatic lipase is observed at pH 8–9, trypsin 7–9, chymotrypsin 7.8–8.0, pancreatic elastase 8.5, carboxypeptidase A 8.0 [9,24–27]. Thus, a higher yield of pancreatic proteins, most of which are enzymes [28], is observed in case of neutral or slightly alkaline medium, since these pH values account for maximum activity of enzymes; this phenomena was also observed in case of saline extraction. On the other hand, high values of the protein concentration in these extracts can be caused by mild extraction conditions as those mild conditions provided extraction of active substances together with biologically active compounds of ballast proteins. The ballast proteins were removed from the extract in case of acid extraction by an additional purification stage.

Throughout the entire experiment, in both methods the oscillatory nature of the degree and rate of protein extraction from pancreas was observed. Decrease in these values at some moments of the process is explained by a decrease in amount of proteins released to the extractant or by their fermentative breakage as a result of autolysis caused by action of extracted proteolytic enzymes [29–32]. Decrease in proteases activity can be achieved by dropping down of the process temperature (which was done in this research), as well as by acidifying the extractants as indicated above. When using saline as an extractant, an instant increase in the extraction rate was observed due to absence of obvious inhibitory agents. For obvious reasons the extraction
of TCA was accompanied by smooth increase in rate of protein release into the extractant. Based on the obtained results, it is possible to note the greater dependence of the extraction method efficiency on availability of inhibiting factors, as well as on the type of the extractant.

The fraction of interest with molecular weight of less than 30 kDa in the final saline extract was researched in early works [7]. Using the UniProt database [3] the following low molecular weight compounds were presumably detected in the purified acid extract: chymotrypsin C (28.9 kDa), a member of family of chymotrypsin-like elastases 1 (28.8 kDa) and 2A (28.7 kDa), glutathione S — transferase omega-1 (27.4 kDa), kininogenase (27.2 kDa), proglucagon (21 kDa), secretin (14.6 kDa), trefoil factor 2 (13.8 kDa), somatostatin (12.7 kDa), colipase (12.1 kDa), inhibitor of serine protease of Kazal-type 4 (9.6 kDa) and 1 (6 kDa), precursor of pancreatic hormone (7.3 kDa) etc. Due to smaller amount of compounds in the saline extract, many of the above specified types of substances were not found there.

Conclusion

The study of dynamics of extraction based on 0.9% sodium chloride solution showed a higher concentration of total protein throughout the entire process with a wide range of compounds with molecular weight of less than 52 kDa. In TCA extraction, the protein content in the analyzed samples was lower than in the first extraction method; however, analysis of the proteomic profile of the samples showed a greater variety of compounds in the entire presented range of molecular weights. The low molecular weight fraction prevailed in acid extracts both before ultrafiltration (according to results of electropherograms study) and after ultrafiltration (based on analysis of protein content in the ultrafiltrates).

Using of the drawn acidic erythrograms enabled to detect the capability, peculiar for primary extracts, to destroy the biological membranes. The low molecular weight fraction of the acid extract possessed the highest cytoprotective effect, which proved the presence of highly active compounds in the extract.

When comparing the saline and acid extraction methods, the applicability of both modifications for obtaining a mixture of proteins with different molecular weights is noted. In this case, greater efficiency is observed in extraction with trichloroacetic acid. In result of this method it is possible to obtain a complex of low-molecular-weight highly active biological compounds.

It is necessary to note that the processes, considered in this work, require a comprehensive approach and are the subject of further extensive research. It is necessary not only to trace the dynamics of the process, but also to select target markers-compounds, which could be used to determine the efficiency of extraction, and also to study the potential for isolating some specific compounds while maintaining of their high activity. The study of other extractants, for example, alcohol or sulfuric acid, is also considered a promising direction of researches.

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