PLUM OIL CAKE PROTEIN ISOLATE: A POTENTIAL SOURCE OF BIOACTIVE PEPTIDES

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ABSTRACT: In this study some functional properties and the in vitro digestibility of protein isolates from plum oil cakes, obtained by supercritical fluid extraction and cold pressing, as a control, were described and compared. Amygdalin contents in the protein isolates were at amounts considerably lower than regulatory. Solubility profiles of both protein isolates were typical for this type of plant proteins. Proteins showed good digestibility, which was assessed by gastrointestinal proteases (pepsin and pancreatin). SDS-PAGE analysis was used for characterisation of protein digests which showed that the protein isolates were completely digested. In vitro antioxidant capacity by three complementary methods and enzyme inhibitory effects towards Angiotensin-I Converting Enzyme (ACE) related to the onset of hypertension were determined. All obtained protein hydrolysates acted as DPPH and ABTS scavengers, as reducing agents and also as ACE enzyme inhibitor. Hence, the protein isolates obtained from plum kernel cake showed to be a potential source of natural products for food applications, with good digestibility and beneficial bioactive properties.

Key words: plum kernel protein, supercritical CO2 extraction, in vitro digestion, ACE inhibition, antioxidant activity

INTRODUCTION

The plum (Prunus domestica) is a member of Prunus family, along with apricot, cherry and peach, and one of the most widespread stone fruits crops around the world. In Serbia, plums represent the most dominant fruit crops which is consumed fresh, dried or used for preparation of different value-added products. The processing of plums into the foodstuffs produces tonnes of by-products in the form of fruit pits which have great potential as an unconventional valuable source of oil, rich in bioactive compounds such as phyto-sterols, tocochromanols, carotenoids, and squalene (Rudzińska et al., 2016). The oil from the kernels could be extracted by conventional methods such as screw pressing, solvent extraction or pre-presing followed by solvent extraction. Developing the “green chemistry” concept, these conventional methods tend to be replaced with different novel extraction techniques such as microwave, ultrasonic, high pressure-assisted extraction, supercritical fluid extraction (SFE), etc. Supercritical fluid extraction is an extraction te-
The aim of this study was to evaluate the antithrombotic and ACE inhibitory activities such as antithrombotic (Sabbione et al., 2015) and ACE inhibitory (Kamel et al. 1992). The limitation of using POC for human consumption is the presence of amygdalin. Amygdalin, cyano-genic glycoside, is non-toxic but in the presence of enzymes, it is hydrolysed and produces a benzaldehyde (hydrogen cyanide) (Garcia et al. 2016).

Recently, plum protein isolate (PI) has been receiving more and more attention as an ingredient that could be incorporated into food, cosmetic and pharmaceutical products. This protein isolate, besides a emulsifying ability, has a potential for the production of bioactive peptides which can be released by in vitro enzymatic hydrolysis, gastrointestinal digestion by commercial enzymes (pepsin and pancreatin) and fermentation from the inactive parent proteins. Bioactive peptides from different plant sources can present diverse activities such as antithrombotic (Sabbione et al., 2015), cholesterol-lowering (Turpeinen et al., 2009), antimicrobial (Sedaghati et al., 2016), antioxidative (Sudhakar and Nazeer, 2015) and ACE inhibitory/antihypertensive (García-Tejedor et al., 2014) activities.

The aim of this study was to evaluate the potential of plum oil cakes, obtained after two different methods of oil extraction, to produce protein isolates as a source of bioactive peptides. The protein isolates have been compared and characterized by the solubility and electrophoresis profile. In vitro two steps hydrolysis process, with commercial digestive enzymes (pepsin and pancreatin), has been performed for determination of digestibility of plum protein isolate. The obtained digests were analysed for different biological activities, such as antioxidant and ACE inhibitory activity. Therefore, the present study would give new approach for valorisation by-product of oil industry as a nutraceutical protein source.

**MATERIALS AND METHODS**

**Materials**

The plum kernels were obtained from the Association of Fruit Brandy Producers (Kneževi vinogradi, Croatia). Oil cakes were obtained by cold pressed (CP) and supercritical CO2 oil extraction (SFE).

**Protein extraction**

Protein isolate (PI) from oil cakes obtained by cold pressing (PICP) and supercritical fluid extraction (PISFE) was extracted by alkali extraction with isoelectric precipitation. Plum cakes, defatted with n-hexane in ration 1:5, were suspended in water solution at alkali pH 10 in ratio 1:10 that was set with 1M NaOH. After 30 min of extraction the slurry was extracted and filtered to remove the insoluble material. The dissolved proteins were precipitated by setting pH to 5 with 1M HCl. After centrifugation (Sorvall® RC-5B Refrigerated Super-speed Centrifuge, Du Pont Instruments, Newtown, PA, USA) at 10,000 rpm and 4 °C for 20 min, the precipitate was dried at 30 °C for 24 hours. The dried precipitate was ground to obtain PI powder. The yield of PI was 1.7 g/10 g oil cake.

**Protein solubility**

The solubility of PI was determined at pH range 2 to 9 and ionic strength range 0.1-1M NaCl. Briefly, 10 mg of PI was weighted into Eppendorf tubes and added to 1 mL of a buffer solution. Samples were constantly stirred for 1 h at 25 °C, using a
Thermo Shaker TS- 100C (Bio-San, Latvia). After mixing the solutions were centrifuged at 14,500 rpm for 10 min (Eppendorf Mini-spin plus, Eppendorf AG., Hamburg, Germany). Proteins from supernatant were determined by the Lowry et al. (1951) method.

**Determination of amygdalin**

Method was made according to Bolarinwaa et al. (2014), with some corrections. Briefly, protein cakes and PI were measured into a round-bottom flask (100 mL), and then added ethanol and the mixture was boiled under reflux during 120 min. In the end of extraction, the extracts were filtered and evaporated under vacuum to remove ethanol. On this way prepared samples were analysed by HPLC (Agilent 1290 Infinity I HPLC system with an Agilent DAD detector). The column for separation was used a Supelco Analytical HS-C18 column (4.6 x 250 mm, 5 µm) Sigma-Aldrich. The chromatographic conditions were: flow rate 1 mL/min, temperature 20 °C, injection volume 20 µL and UV detection at 210 nm. Mobil phase consisted of distillation water: methanol (75:25 v/v).

**Protein digestion**

PIs were hydrolysed using gastrointestinal proteases in order to simulate the human gastrointestinal environment. Digestion was performed using a combination of two enzymes, pepsin (E/S 1/25) and pancreatin (E/S 1/50), at 37 °C during 4 h. Protein suspension in water (2.5 g/100 cm³) was pre-incubated at 37 °C, then adjusted to pH 2.5 and added pepsin. After 120 min of gastric digestion, pH of the solution was adjusted to pH 7 and added pancreatin and intestinal digestion was carried out next 120 min. The reaction mixture was heated at 100 °C for 5 min to stop reaction and centrifuged using Eppendorf Mini spin plus (Eppendorf AG., Hamburg, Germany) at 14,500 rpm for 10 min at room temperature. The collected supernatants were further analysed.

**Degree of hydrolysis**

The degree of the hydrolysis (DH) was determined by method described by Popović et al. (2013).

**SDS-PAGE electrophoresis**

Proteins and digests were separated using sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE), described by the method of Laemmli (1970).

**Determination of antioxidant activity**

Antioxidant activity of all samples were carried out by DPPH, ABTS radical scavenging and reducing power method, the methods was described by Čakarević et al. (2019).

**Assay of ACE-inhibitory activity**

The ACE- inhibitory activity of the hydrolysates was measured following the method described by Yoshi-Stark et al. (2004).

**Statistical analysis**

The data were in triplicate, and subjected to statistical analysis, using analysis of variance (ANOVA) to determine significant differences between the samples (p < 0.05). Differences between the treatment means were separated using Duncan's multiple range tests.

**RESULTS AND DISCUSSION**

The proximate compositions of the plum oil cakes and protein isolates obtained from them are shown in Table 1. Protein was the major component of both plum cakes, with 50.69% for CP and 48.94% for SFE. Carbohydrates and fibres were the followed components in both cakes. Both PIs obtained from plum oil cakes by isoelectric precipitation, have protein content 97.94% for PICP and 99.15% for PISFE. Generally, utilization of this by-product for protein and oil production could be limited by the presence of cyanogenic glycosides such as amygdalin therefore we determined its content. Toxicity of amygdalin comes from benzaldehyde, with a typical bitter taste, and cyanide, as a result of the hydrolysis of amygdalin. According to the European Food Safety Authority (EFSA) it was set a maximum level of cyanide of 50 mg/kg in nougat, marzipan or their substitutes or similar products, 5 mg/kg in canned stoned fruits, and 35 mg/kg in alcoholic beverages (EFSA, 2016).
Chemical composition of plum oil cakes (POC) and protein isolates (PI) obtained from oil extraction by cold pressing (CP) and supercritical fluid extraction (SFE)

<table>
<thead>
<tr>
<th>Composition</th>
<th>CP</th>
<th>SFE</th>
</tr>
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<tbody>
<tr>
<td>Moisture (%)</td>
<td>10.80±0.145&lt;sup&gt;b&lt;/sup&gt;</td>
<td>6.27±0.315&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Crude fibre (%)</td>
<td>5.86±0.315&lt;sup&gt;a&lt;/sup&gt;</td>
<td>11.57±0.345&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Crude lipids (%)</td>
<td>10.09±0.165&lt;sup&gt;a&lt;/sup&gt;</td>
<td>10.63±0.12&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Total carbohydrate (%)</td>
<td>12.77±0.015&lt;sup&gt;b&lt;/sup&gt;</td>
<td>12.59±0.05&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Reducing sugars (%)</td>
<td>10.40±0.025&lt;sup&gt;a&lt;/sup&gt;</td>
<td>10.90±0.025&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Crude protein (%)</td>
<td>50.69±0.53&lt;sup&gt;b&lt;/sup&gt;</td>
<td>48.93±0.625&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Amygdalin content (mg/g PI)</td>
<td>97.94±0.875&lt;sup&gt;a&lt;/sup&gt;</td>
<td>99.15±0.095&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Protein content (%)</td>
<td>0.027±0.001&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.020±0.001&lt;sup&gt;b&lt;/sup&gt;</td>
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Values represent average of triplicates ± SD. Different superscript letters (a and b) indicate significant difference (p < 0.05) within the same row.

Figure 1. Influence of pH and ionic strength (µ) on protein solubility of plum oil cake protein isolates obtained by a) cold pressing (PICP) and b) supercritical fluid extraction (PISFE)

Figure 2. (a) Degree of hydrolysis (DH) of plum oil cake protein isolates obtained by cold pressing (PICP) and supercritical fluid extraction (PISFE) during in vitro digestion (b) SDS-PAGE analysis of plum oil cake protein isolates and their digests: (1) molecular mass markers SDS 7, (2) PICP, (3) PISFE, (4) PICP pepsin, (5) PICP pancreatin, (6) PISFE pepsin, (7) PISFE pancreatin

Similarly, our previous reports (Čakarević et al. 2019) and literature data (Garcia et al. 2016) showed that protein isolates from apricot cakes either did not contain amygd-
Cold pressing (CP) and supercritical fluid extraction (SFE) of plum oil cakes (POC) and protein isolates (PI) obtained from oil extraction by CP (CP) and SFE (SFE) have been conducted. The obtained results showed that amygda林 content in PIs is in amount considered as safe (Table 1).

**Protein solubility**

The protein solubility patterns were similar for both samples. The lowest solubility for both PIs was observed at pH 4 (3.95 mg/mL for SFE and 4.85 mg/mL for CP), and the higher solubility was evident under extremely acidic (pH 2) and alkaline (pH 9) conditions. This result indicates the isoelectric point of the extracted PI was at pH 4, which is in correlation with general trend that most of food proteins are acidic, with isoelectric point at pH 4-5 (Roodsoaman and Sothornvit 2018; Chambal et al. 2013; Wu et al. 2009). Significant influence of ionic strength was observed mostly at the extreme acidic pH, manifested as the salting-out effect, with decreasing solubility of PI. Over pH range from pH 4 to pH 8 salting-in effect occurred with a minor increase in solubility. Similar trend was observed from our previous reports for PI from apricot kernel cake after CP and SFE oil extraction (Čakarević et al. 2019).

**Digestibility of protein isolate**

Digestibility of PIs was subjected to evaluate the capacity to release peptides which could have potential bioactive effects when PIs were directly consumed. In order to simulate digestion of human gastrointestinal tract, proteases, pepsin and pancreatin were used and process of digestion was monitored by measuring DH. The hydrolysis curves of PIs after digestion are shown in Figure 2a. During pepsin digestion the DH of PICP and PISFE reached 41.47±0.79% and 37.85±0.73%, respectively. Moreover, subsequent digestion by pancreatin led to an increase of DH in both PI digests, reaching similar DH values (53.99±0.60% for PICP and 55.04±1.00% for PISFE, respectively). The electrophoretic profiles of PIs and their products of *in vitro* digestion by pepsin and subsequently by pancreatin are shown in Fig. 2b. Electrophoresis of PIs exhibited the presence of two groups of intense bands, corresponding to polypeptide complexes with molecular weight between 36-45 kDa and 20-29 kDa (lines 2 and 3). Similar protein patterns were previously reported for other seeds of genus *Prunus* L. (Garcia et al., 2015). In the digests, both PIs were further degraded to apparent molecular weights lower than 20 kDa. After pepsin, digests of both PIs showed lower molecular weight fragments notably bellow 20 and 14 kDa (lines 4 and 6). During the second enzymatic step, after treatment by pancreatin, no visible bands were obtained in the gel (lines 5 and 7). These results showed that PIs are not resistant to digestive proteases. Similar results were reported for other digests of plant proteins obtained from almond (Sze-Tao and Sathe, 2000) and oil palm kernel (Tapal et al., 2016).

**Antioxidant capacity of digested PIs**

Antioxidant capacities of protein digests obtained from PIs after both in vitro digestion steps were studied by DPPH and ABTS scavenging assays as well as by reducing power assay (Figure 3a, b). In general, all digests showed radical scavenging capacity. Values of antioxidant activity depended on the chosen assay (Figure 3). The DPPH radical scavenging activity of PICP and PISFE digests after pepsin digestion was 12.33% and 24.43%, respectively. Pancreatin digestion of PICP and PISFE digests led to an increase of the scavenging activity to 30.33% and 23.38%, respectively. Similar trend was obtained with ABTS radical scavenging activity of PIs, with 24.96% to 27.82% after pepsin digestion and 31.81% to 36.67% following pancreatin digestion. Reducing ability of digests indicated their capacity to donate electrons and it increased with increasing DH values. The digests from both PIs after pepsin digestion showed significant increase in the reducing power, with continued increase during pancreatin digestion. Results proved that PIs were cleaved into small peptides and free amino acids by digestive proteases which are reported to exert greater antioxidant activity than their parent proteins or large polypeptides (Vaštag et al., 2013).
ACE inhibitory capacity of digests

Both PIs and associated digests were screened for their ACE inhibitory potential. Native PIs did not show ACE inhibitory activity (data not shown), but their digests exerted a significant inhibitory activity, indicating the release of ACE inhibitory peptides during in vitro digestion (treated by pepsin and pancreatin). Figure 4 shows the linear function of ACE inhibition activity and sample concentrations. When the obtained ACE inhibitory activities of digests were presented as IC_{50}, then values for PICP were at 0.125 mg/mL and at 0.127 mg/mL for PISFE. Moreover, the inhibitory activity increased following treatment with pancreatin, with IC_{50} values at 0.115 mg/mL for PICP and at 0.106 mg/mL for PISFE. Similar results have been reported for antihypertensive peptides obtained from protein isolates after simulated gastrointestinal digestion, from sunflower (Megias et al., 2004), apricot seed (Čakarević et al., 2019), and chia protein (Pablo Osorio et al., 2019).

CONCLUSIONS

Plum oil cake which remains after oil extraction process could be an alternative...
source of bioactive compounds which have different biological activities that could provide health benefits. This residue is an unused protein source with protein content of about 50% hence the protein exploitation from this source is a good way to increase the value of this agricultural by-product. Protein isolate extracted from defatted plum seed flour by alkali solution along with isoelectric precipitation contained about 90% of protein. Bioactive peptides, released from parent proteins during enzymatic hydrolysis, exhibited diverse bioactivities, such as antioxidant and blood pressure-lowering (ACE inhibitory) effect. Presented results indicated that plum kernel proteins could be used as a functional food additive, which could, following digestion, be the source of peptides that exert positive effects on human health.

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ПРОТЕИНСКИ ИЗОЛАТ УЉАНЕ ПОГАЧЕ СЕМЕНА ШЉИВЕ: ПОТЕНЦИЈАЛНИ ИЗВОР БИОАКТИВНИХ ПЕПТИДА

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Сажетак: У овој студији описане су и упоређене функционалне особине и сварљивост протеинских изолата добијених из уљаних погача шљиве поступцима суперкритичне екстракције у основи уљана шљиве, а ладним пресовањем. Садржај амидалних у протеинским изолатима се анализовали у количини мањој од прописане. Растворљивост оба протеинска изолата одговара профилима растворљивости биолошких протеина. Протеински изолати су показали добру сварљивост, што је доказано применом гастроинтестиналних протеаза (пепсин и панкреатин) у процесу in vitro дигестије. За карактеризацију сварљивости протеина коришћена је SDS-PAGE анализа, којом је доказана потпуна сварљивост протеинских изолата. Помоћу три комплементарне методе одређена је антиоксидативна активност добијених хидролизата, као и инхибиторно дејство на ангиотензин-І конвертирујући ензим (ACE) који директно утиче на појаву хипертензије. Сви добијени протеински хидролизати деловали су као хватачи слободних радикала и инхибитори ACE. Стога се показало да су протеински изолати добијени из уљаних погача шљиве потенцијални извор функционалних додатака с добром сварљивошћу и биоактивним својствима.

Кључне речи: протеини уљане погаче, суперкритична екстракција, ин витро дигестија, ACE инхибитори, антиоксидативна активност

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