Characterization of Hempseed Protein in Cheungsam from Korea

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Abstract The nutricheinal analysis of hempseed from Cheungsam was performed in order to apply hempseed to food industry. Cheungsam hempseed was mainly composed of crude fat (30.52%) and protein (30.02%). The hempseed fat contained about 88% of polyunsaturated fatty acids, and the omega-6 to omega-3 ratio was determined to be approximately 3:1. Hempseed from Cheungsam was rich in essential amino acids. Moreover, hempseed protein from Cheungsam, extracted by acid-base precipitation method, showed free radical scavenging activity. The results suggest that Cheungsam hempseed protein could be utilized as a superior source of protein nutrition.

Key Words : Hempseed, Cheungsam, Characterization, Protein, Nutrient

1. Introduction

Hemp (Cannabis sativa L.) is a widely cultivated plant with industrial importance for thousands of years by prehistoric humans, especially as an valuable source of food, fiber and medicine[1]. In addition to the medical potential of cannabinoids in the leaves and flowers of the plant itself, hempseed has been utilized in the human diet for thousand of years. However, the effects of dietary hempseed intake have yet to be fully understood, owing largely to the politics prohibiting its cultivation, although hempseed has been recognized for thousands of years as an excellent sources of nutrition[2]. Recently hempseed has been legally used as food for humans in both Canada and the United States[3]. Hempseed has also been utilized as a valuable food source for farm animal, fish and cattle[4].

Hempseed, produced as a byproduct during commercial utilization of hemp fibre, is known to be rich source of high quality oil and protein[3]. The hempseed oil was known to contain over 80% of polyunsaturated fatty acids. A variety of studies demonstrated that hempseed is perfectly balanced in regards to the ratio of the two essential fatty acids, named linoleic (omega-6) and α-linolenic acid (omega-3) acids for human nutrition[3,5]. Besides the nutritional value, the hempseed oil also has some potential health benefits, such as lowering of cholesterol level, reducing high blood pressure, and treating human atopic dermatitis[6].

Hempseed protein was known to mainly consist of
high-quality storage protein, named edestin (globulin) and albumin. The edestin and albumin in the hempseed had superior essential amino acid composition and was easily digested[1,3,7].

In Korea, a new variety of non-drug fiber hemp, named “Cheungsam”, was developed by crossing IH3 Netherland variety and Korea local variety. The new variety of “Cheungsam” is regarded as non-drug type hemp, with a low level of δ-9 tetrahydrocannabinol (THC).

There are a few reports about hempseed protein to utilize them as a source of protein nutrition. The information on the hempseed protein in Cheungsam from Korea is not available. The objective of this work was to characterize the hempseed protein in Cheungsam from Korea. The neutrochemical and antioxidant properties of hempseed protein from Cheungsam were investigated.

2. Materials and Methods

2.1 Hempseed preparation

Dehulled hempseeds from non-drug fiber hemp, named “Cheungsam”, were ground in liquid nitrogen and then dispersed in deionized water at low temperature of less than 35°C.

2.2 Amino acid analysis

The amino acid composition of the hempseed protein samples was determined by an automatic amino acid analyzer (Waters M510, USA), using PICO, TAG column[1]. The determination was carried out at 37°C, with the detection wavelength of 254 nm and flow rate of 1.0 ml per min. The samples were hydrolyzed with 6 N HCl for 24 h at 110°C in sealed tube. The amino acid composition was reported as mg/100 g hempseed.

2.3 Preparation of hempseed protein

The hempseed protein was extracted as follows. 1 g of hempseed flour was mixed with 100 ml deionized water, and the mixture was adjusted to pH 10.0 with 1.0 N NaOH. After continuously stirring for 1 h, the suspensions were centrifuged at 8,000 g for 30 min and the precipitates were discarded. The pH of the supernatant was adjusted to pH 5.0 at 4°C with 1 N HCl, and the precipitates were collected by centrifugation. The precipitates were washed with pre-cooled deionized water, and dispersed in the deionized water. The dispersions were adjusted to pH 7.0 with 1 N HCl, and then dialyzed at 4°C.

2.4 Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE)

SDS-PAGE was conducted according to the discontinuous buffer system of Laemmli[8] at 5% stacking gel and 12.5% separating gel using gel electrophoresis apparatus. The protein samples were directly dissolved in the sample buffer, namely 0.125 M Tris-HCl buffer (pH 8.0) containing 0.1% (w/v) SDS, 0.05% (w/v) bromophenol blue, 30% (v/v) glycerol and 5% (v/v) β-mercaptoethanol (2-ME). The electrophoresis was run at 20 mM in stacking gel and at 40 mM in separating gel until the tracking dye reached the bottom of the gel. Before electrophoresis, the samples were heated at 100°C for 5 min. The SDS-PAGE under nonreducing condition was carried out as mentioned above, just using the samples dissolved in the sample buffer without the addition of 2-ME[7,9].

2.5 Periodic acid–Schiff (PAS) staining for carbohydrate

After finishing SDS-PAGE, gels were fixed overnight in 25% isopropanol, 10% acetic acid and 65% water. Gels were washed for 1 h in running water, soaked in 1% NaIO₄, 3% acetic acid for 1 h and then washed for 2 h in running water and twice for 0.5 h with distilled water. The gels were then stained for 2 h with Schiff’s reagent prepared from Basic Fuchsin[10]. Excess Schiff’s reagent was removed by soaking in 0.5% sodium metabisulphite. The gels were then soaked for two times 3 h in 7% acetic acid.

2.6 DPPH radical scavenging activity

The DPPH (1,1-diphenyl-2-picrylhydrazyl) radical scavenging activity was determined by the method described earlier[11]. 2 ml of the sample solution was mixed with freshly prepared 2 ml of 0.2 mM DPPH ethanolic solution and vortexed for about 10 sec. The
resulting solution was then left to stand for 30 min, prior to reading the absorbance at 517 nm. A low absorbance at 517 nm indicates a high DPPH scavenging activity. Ethanol was used as a blank.

2.7 Protein solubility

Protein solubility was determined according to the method of Tomotake et al.[12], with a few modifications. Protein dispersions (1 mg/ml, w/v) were prepared in 0.01 M phosphate buffer adjusted to pH 2-10. To achieve desirable pH above 10, the protein dispersion was adjusted directly using 1 N NaOH. For total soluble protein contents (control), the samples were solubilized in 0.1 N NaOH. The solubilized proteins were stirred at room temperatures for 1 h, centrifuged at 8,000 g for 30 min, and filtered through filter paper. Protein contents of the filtrate were determined according to the Bradford method using bovine serum albumin as a standard [13]. Each measurement was performed in triplicate.

3. Results and Discussion

3.1 Nutritional chemical analysis of hempseed from Cheungsam

The contents of crude protein, crude fat, fatty acids (saturated and unsaturated) and two essential fatty acids (linoleic acid and linolenic acid) of hempseed from Cheungsam were indicated in Table 1. The hempseed was mainly composed of crude protein (30.02%) and crude fat (30.52%). In addition, the hempseed fat contained about 88% of polyunsaturated fatty acids. Moreover, it was an exceptionally rich source of the two essential fatty acids such as linoleic acid (18:2 omega-6) and linolenic acid (18:3 omega-3).

Other hempseeds were also reported to contain over 30% fat and about 25% protein, with considerable amounts of dietary fiber, vitamins and minerals[3]. As shown in Table 1, the omega-6 to omega-3 ratio of hempseed from Cheungsam was determined to be approximately 3:1, indicating an excellent ratio for human health.

3.2 Amino acid composition of hempseed from Cheungsam

The amino acid compositions of hempseed from Cheungsam were determined as shown in Table 2. The hempseed had higher levels of glutamic acid, aspartic acid and arginine, which are consistent with the previous reports of other hempseeds [1,3,7,14]. Hempseed protein has been reported to have good amounts of the sulfur-containing amino acids methionine and cysteine[15]. However, the sulfur-containing amino acids methionine and cysteine might be in some content destroyed due to the HCl-hydrolysis method used in this experiment.

<table>
<thead>
<tr>
<th>Amino acids</th>
<th>mg/100 g hempseed</th>
</tr>
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<tbody>
<tr>
<td>Isoleucine*</td>
<td>1168.83</td>
</tr>
<tr>
<td>Leucine*</td>
<td>2029.29</td>
</tr>
<tr>
<td>Lysine*</td>
<td>1053.81</td>
</tr>
<tr>
<td>Methionine*</td>
<td>475.87</td>
</tr>
<tr>
<td>Cysteine</td>
<td>95.85</td>
</tr>
<tr>
<td>Tyrosine</td>
<td>691.13</td>
</tr>
<tr>
<td>Phenylalanine*</td>
<td>1369.67</td>
</tr>
<tr>
<td>Threonine*</td>
<td>1113.71</td>
</tr>
<tr>
<td>Valine*</td>
<td>1450.23</td>
</tr>
<tr>
<td>Histidine*</td>
<td>887.01</td>
</tr>
<tr>
<td>Tryptophan*</td>
<td>318.17</td>
</tr>
<tr>
<td>Aspartic acid</td>
<td>3271.25</td>
</tr>
<tr>
<td>Glutamic acid</td>
<td>6273.65</td>
</tr>
<tr>
<td>Serine</td>
<td>1602.9</td>
</tr>
<tr>
<td>Glycine</td>
<td>1402.41</td>
</tr>
<tr>
<td>Arginine*</td>
<td>3470.47</td>
</tr>
<tr>
<td>Alanine</td>
<td>1375.59</td>
</tr>
<tr>
<td>Proline</td>
<td>1229</td>
</tr>
<tr>
<td>Asparagine, Glutamine : N.D</td>
<td></td>
</tr>
</tbody>
</table>

Essential amino acids are indicated by asterisk*
Notably, hempseed from Cheungsam were rich in essential amino acids such as arginine, isoleucine, leucine, lysine, threonine, valine, phenylalanine, histidine and methionine. Hempseed protein from Cheungsam had more essential amino acids than that from Finola hemp [3].

Moreover, it had high quality proteins in common with soy bean and rape through a direct comparison of amino acid compositions from egg white, soy bean, rape seed, potato and rice [3]. The results suggest that the hempseed from Cheungsam has superior essential amino acid composition and thus it might be used as a good source of protein nutrition. The results also indicated that hempseed has good profile of essential amino acids required for infants [16,17].

3.3 SDS–PAGE profiles of hempseed protein

The hempseed protein was prepared at room temperature by using acid-base precipitation method [1]. SDS-PAGE profiles of protein constituents in the presence and absence of 2-mercaptoethanol were presented in Fig. 1. Under reduced condition, there were three major kinds of protein constituents, corresponding to 36.4 kDa, 20.4 kDa and 18.3 kDa Fig. 1A. In the absence of 2-ME, the major protein band of about 52.5 kDa was found, probably suggesting that the disulfide bonds between proteins detected on the denatured gel will not be disrupted, and as the result, the protein bands would be in the form of different size. Hempseed proteins were reported to mainly consist of edestin and albumin, and edestin is composed of six identical subunits, and each subunit consists of acidic subunit and basic subunit linked by one disulfide bond. In the present study, the band about 36 kDa was homogeneous, while the other heterogeneous bands mainly consisted of two bands of about 20 and 18 kDa in hempseed protein from Cheungsam. From the molecular size and relative content, the major protein is assumed to be edestin [18,19].

Fig. 1B shows the glycoprotein staining for hempseed protein isolate with periodic acid–Schiff (PAS) staining. As shown in Fig. 1B, only the bands assumed as edestin were detected by PAS staining in reduced and unreduced conditions. These results suggest that the suspected edestin in hempseed protein from Cheungsam might contain carbohydrate moiety in the structure.

3.4 DPPH free radical scavenging assay

The DPPH free radical scavenging ability of hempseed protein was presented in Fig. 2. The stable DPPH radical in ethanol has been widely used to test the ability of some compounds to acts as free radical scavengers or hydrogen donors [10,20]. DPPH assay evaluates the ability of hempseed protein to scavenge free radicals of DPPH. The present DPPH assay showed that hempseed protein from Cheungsam possess free radical scavenging activity. The free radical scavenging ability of hempseed protein gradually increased with increasing protein concentrations Fig. 2. The IC$_{50}$ value of the hempseed protein, meaning the concentration that causes a decrease in initial DPPH concentration by 50%, was determined to be approximately 1.5 mg/ml as shown in Fig. 2. The IC$_{50}$ value of wheat germ protein hydrolysate was 1.3 mg/ml and hempseed protein hydrolysate after hydrolysis using neutrase was reported to be 2.3-3.5 mg/ml[4]. The antioxidant properties of various food proteins, named bovine caseins [21], whey proteins [21], soy proteins [22-24], fish proteins [25,26] and wheat proteins [27] have been reported. The antioxidative ability of the food proteins could enlarge the application and utilization of food proteins for human nutrition.
3.5 Solubility of hempseed protein

Fig 3 shows the solubility of hempseed protein between pH 2 and pH 10. The protein solubility was minimum at pH in the range of 4.0-8.0 and increased gradually below pH 4.0 and above pH 8.0 Fig. 3. At pH 7.0, only 10% of protein was solubilized in 0.01M phosphate buffer. However, at above pH 8.0, the solubility increased up to more than 90%. The data suggests that hempseed protein is a kind of typical alkali soluble protein. The underlying mechanism of solubilization at alkaline pH was reported to be related to the dissociation of edestin[28], while the low solubility at pH less than 7.0 might be due to the aggregation of edestin[29].

![Fig. 3] Protein solubility profile of hempseed protein from Cheungsam at different pH values.

The data obtained in this investigation implicate hempseed protein from Cheungsam as a prominent source of nutrients, which had good profiles of essential amino acids. Moreover, the omega-6 to omega-3 ratio of HPI was determined to be approximately 3:1, indicating an excellent ratio for human nutrition. Hempseed protein isolate was composed of a major component, probably edestin on the SDS-polyacrylamide gel, and it has a DPPH free radical scavenging ability. Taken together, hempseed protein from Cheungsam might be utilized as a superior source of protein nutrition, and it can be applied to food industry as a good source of nutrition.

4. Acknowledgement

This research was supported by Technology Development Program for Agriculture and Forestry, Ministry for Food, Agriculture, Forestry and Fisheries, Republic of Korea.

References


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