Introduction

Scientists and researchers have constantly been trying to find alternative protein foods to combat malnutrition and diseases due to protein inadequacy in exponentially increasing population. Shortage of protein foods is a serious threat to mankind especially in under-developed and developing nations. Hence it has become utmost important to find non-conventional seeds as new protein source and nutritional supplements. As a consequence, during the last few decades the food industries have paid special attention on utilizing seed protein from both conventional and non-conventional sources [1]. For plant proteins to be acceptable and successful in food application, it must behave ideally in some physico-chemical properties, referred to as ‘functional properties’, such as solubility, viscosity, emulsifying property, foaming property, gelation capacity etc. [2].

Our intention was to find new proteins for human and animal consumption from non-conventional seeds. One of such non-traditional seeds is Crotalaria pallida, (also known as ‘smooth rattle pod’) which belongs to the family ‘Fabaceae’. This plant is widely distributed in the tropical and subtropical region across the world, though, very little effort has been made to isolate and characterize protein from the seeds of this plant. This plant is a good cover crop in tea, coconut and rubber plantations [3].

Flowers of this plant are used as vegetables. Seeds of this plant are roasted and grounded to prepare a sort of coffee beverage. The seeds of this plant are rich in nitrogen and protein content. The main objective of our group is to examine various functional properties of its seed protein isolate as a key step to check its usability as a non-traditional protein source. This present study also deals with the first time report of functional properties of Crotalaria pallida seed protein.

Materials and Methods

Materials: Fresh and mature fruits of C. pallida were collected from the Golapbag campus of the University of Burdwan and identified by A. Mukherjee, Department of Botany, The University of Burdwan, Burdwan, West Bengal, India. Voucher specimen (labeled as ‘BURD Sushobhan 203’) has been preserved at the herbarium of Botany Department. Collected seeds were air-dried and then finely powdered and kept in a refrigerator at 4°C for future uses. All the chemicals used in our present study were of analytical standard. Corn oil was obtained from Sigma-Aldrich, USA.

Methods

Preparation of Total Protein Isolates (TPI): Powdered seed material was defatted using pet ether (60°C-80°C) in a Soxlet apparatus for 72 hours, followed by washing with the help

Abstract

Solubility and functional properties of Crotalaria pallida seed protein isolate have been evaluated. The effects of pH on some of these properties were also investigated. Solubility of this protein was minimum at pH 5 (which is supposed to be close to its iso electric point) and maximum at pH 8. The protein content of the total seed protein isolate was 62.43%. Oil holding capacity and water holding capacity were estimated to be 4.62 g/g and 3.21 g/g respectively. The emulsifying activity and emulsion stability as well as foaming capacity and foam stability were largely affected by pH levels. Viscosity of the seed protein isolate was also found to be pH and concentration dependent. Gelation properties of the protein isolate have been investigated and least gelation concentration was found to be 10% in distilled water and 8% in 0.5 (M) NaCl solutions respectively.

Keywords: Crotalaria pallida; Seed Protein Isolate; Nitrogen Solubility; Functional Properties
of Chloroform: Methanol (1:1) solvent mixture. As the nitrogen solubility is maximum at pH 8.0, powdered seed was extracted using distilled water at the same pH. The defatted flour was stirred with distilled water (10 volumes than the weight of seed flour taken) for 30 minutes at constant pH (pH 8.0) and filtered through Whatman-41 filter paper and finally centrifuged at 5000g for 20 minutes. The supernatant was collected and precipitate was obtained from this solution by lowering the pH to a 4.0 using Trichloroacetic acid(TCA). The precipitate thus obtained was recovered by centrifuge at 5000g for 20 min. The precipitate was re-dissolved in deionizer water at pH 7.0 and then freeze-dried. The freeze-dried protein (total protein isolate) was preserved in a refrigerator at 4°C until further use.

Proximate analysis: The ash and moisture contents of the Total protein isolate (TPI) were estimated according to the method of AOAC [4]. Micro-Kjeldahl methodology was followed to determine its nitrogen content (AOAC, 1990)[4].The percentage of nitrogen was converted to the percentage of crude protein by multiplying by a factor of 6.25. All the results were placed in (Table 1).

Water-holding capacity (WHC) and Oil-holding capacity (OHC): Water- and Oil-holding capacity was measured according to the method of Carcea Bencini with slight modification [5]. One gram of protein sample in each case was stirred in 10 ml distilled water at pH 7.0 or 10 ml corn oil in a centrifuge tube. Then the samples were allowed to stand at room temperature for 30 min and centrifuged at 5000g for another 15 min. The volume of the supernatant was measured in a graduated cylinder. The number of grams of water or the number of grams of oil held by 1g of protein represents WHC and OHC respectively. Density of the corn oil was found to be 0.9 g/ml. The triplicate determinations of the capacities (WHC & OHC) were carried out and the results were placed in (Table 2).

Nitrogen solubility: The method of Were et al. was used to evaluate nitrogen solubility of the TPI at different pHs (2-12) [6] (Figure 1). Sample (125 mg) in each case was dispersed in 25 ml distilled water and the solution was adjusted to 2, 3, 5, 5, 6, 8, 9, 10, 11 and 12 using either 0.5 (M) HCl or 0.5 (M) NaOH. The dispersion was stirred for 30 min with the aid of a magnetic stirrer and then centrifuged at 10,000g for 15 min. The solubility was determined by nitrogen estimation with the aid of Kjeldahl method of AOAC (1990) [4]. Results are represented as mean ± SD, n=3. Solubility profile was obtained plotting protein solubility (%) against pH. The percentage of soluble protein was calculated as follows:

\[
\text{Solubility} = \frac{\text{amount of nitrogen in the supernatant}}{\text{amount of nitrogen in the sample}} \times 100
\]

Emulsifying activity (EA) and Emulsion stability (ES): Emulsions were prepared according to the method described by Sathe and co-workers with a little modification [7]. 5 ml of protein suspensions at different pHs (2, 4, 6, 8 and 10) were homogenized at 10,000 g for 1 minute and then with corn oil for another 1 minute. The emulsions obtained were centrifuged at 3000 g for 5 minutes and the volumes of the emulsified layers were noted.

Emulsion activity (EA) can be calculated as follows:

\[
E.A. \% = \frac{\text{Volume of emulsified layer}}{\text{Volume of the whole layer in the centrifuge tube}} \times 100
\]

Determination of Emulsion stability involves heating of the above emulsion at 80°C for 30 min, followed by cooling at room temperature and measuring the volume of remaining layer after rotating at 3000 g for 5 min in a centrifuge.

\[
E.S. \% = \frac{\text{Volume of remaining emulsified layer}}{\text{Original emulsion volume}} \times 100
\]

Foaming properties: The method described by Sze-Tao and Sathe[8] was used with slight modification for the determination of Foaming capacity and Foam stability. 1.5% aqueous solution (w/v) of TPI was prepared using ultra-sonic vibrator (Model No. Transsonic T660/H, Elma, Germany) and pH of the suspensions was adjusted to 2, 4, 6, 8 and 10 with 0.5 (M) Hcl or 0.5 (M) NaOH solutions. Then it was whipped with an electrical stirrer at medium speed for 5 minutes and the foam volume was recorded after 30 s. Foam stability was determined by measuring the decrease in foam volume as function of time up to a period of 120 minutes.

\[
\text{Foam capacity} (%) = \frac{\text{Volume after whipping (ml)} - \text{Volume before whipping (ml)}}{\text{Volume before whipping (ml)}} \times 100
\]

\[
\text{Foam stability} (%) = \frac{\text{Volume after standing (ml)} - \text{Volume before whipping (ml)}}{\text{Volume before whipping (ml)}} \times 100
\]

Viscosity: An appropriate sample was dispersed in distilled with the help of the ultra-sonic vibrator (aforementioned model). Each sample was prepared at concentrations of 1, 1.5, 2, 2.5, 3, 4, and 5% (w/v) and relative viscosity of each sample was measured in an Ostwald type viscometer. In another set of experiment, relative viscosity of 1.5% protein solution was
recorded at various pHs [4,6,8,10]. All these experiments are repeated thrice and means are presented in Table 3.

**Least Gelation concentration:** This was measured with slight modification of the method of Abbey and Ibeh [9], where TPI was suspended in 5 ml distilled water to obtain 4%, 6%, 8%, 10%, 12%, 14%, 16%, 18% and 20% (w/v) concentrations with the use of ultrasonic vibrator(same model used earlier). The test tubes containing the solutions/suspensions were heated in water bath for an hour, followed by rapid cooling in running tap-water. These test tubes were kept in a refrigerator for another 2 hours at 4°C. The least gelation capacity was regarded as the concentration at which the sample from the inverted tubes did not fall down or slip.

**Results and Discussion**

**Proximate analysis:** The moisture, ash and protein content of *Crotalaria pallida* seed isolates are given in (Table 1). The protein content of *Crotalaria pallida* seed protein isolate is sufficiently high, which makes it a good source of vegetable protein.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Values in % (±S.D)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture</td>
<td>10.61 ± 0.17</td>
</tr>
<tr>
<td>Ash</td>
<td>2.14 ± 0.16</td>
</tr>
<tr>
<td>Protein (N × 6.25)</td>
<td>62.43 ±0.12</td>
</tr>
</tbody>
</table>

*Values are means of triplicate samples.

**Table 1:** Proximate chemical analysis of seed protein isolate of *Crotalaria pallida*.

**Table 2:** Water- and oil- holding capacities of *Crotalaria pallida* seed protein isolate.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Values in g/g (±S.D)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water holding capacity</td>
<td>3.21 ±0.09</td>
</tr>
<tr>
<td>Oil holding capacity</td>
<td>4.62 ±0.08</td>
</tr>
</tbody>
</table>

*Values are mean of triplicate determinations.

**Water- and Oil-holding capacity:** Water- and oil holding capacity of TPI were found to be 3.21 g/g and 4.62 g/g respectively (Table 2). WHC for this protein is higher than Lupin seed protein concentrates (1.37 g/g) and Cashew nut protein concentrates (1.74 g/g) [10-11]. This value resembles with Soybean protein concentrates (3.06 g/g) and three Chinese indigenous legume seeds as reported by Chau et al. [12]. Such high value of WHC makes it a suitable ingredient for meat, bread and cakes industries. Oil holding capacity of *Crotalaria pallida* seed protein isolates is found to be higher than that of Soybean protein concentrate (1.37 g/g) [12]. High value of OHC for *Crotalaria pallida* seed protein concentrates is probably attributed to a higher level of non-polar side chains in the molecule. This is in agreement of the report of Campell, Shih and Marshal that the value of OHC increases with increasing protein concentration in sunflower and soya products [13]. Such high OHC makes the protein potentially useful in structural interactions in food, especially in flavor retention, improvement of palatability and extension of shelf life in meat products through lowering of fat and moisture loss. The ability of protein to bridge fat and water molecules simultaneously is an important criterion for cold-meat industry especially for sausages.

**Nitrogen solubility:** The nitrogen solubility profile of *Crotalaria pallida* seed protein isolate indicates a dip at pH 5.0 which is close to its isoelectric point. On either side of pH 5.0, there is a sharp increase in the nitrogen solubility of protein isolate having maximum solubility at pH 8. At pH above 8.0, solubility does not change significantly with increase in pH value. The percentage of soluble protein is 58.12 (± 0.22) at pH 5.0 and 96.28 (± 0.06) at pH 8.0. These are comparable to the solubility of bitter lupin concentrate (98.43%) and sweet lupin concentrate (98.79%) as reported by El-Adawy et al. [14]. Also the data indicates that solubility is slightly greater in alkaline medium than that of acidic medium *Crotalaria pallida* seed protein exhibits good solubility in both acidic and alkaline pH ranges, an important characteristic for food formulation. Moreover, the study of nitrogen solubility over a wide range of pH values is an important guide to protein functionality, since it is directly related to many important properties such as emulsification, foaming capacity, gelation etc [15].

**Emulsifying properties:** The effects of pH on emulsifying
activity (EA) and emulsifying stability (ES) are shown in Figure 2 and Figure 3 respectively. Minimum emulsifying activity is observed at pH 4.0 with coincidental decrease in solubility, and is slightly increased to attain maxima at pH 10.0. Emulsifying stability is also found to be pH dependant, where ES is 87.36% at pH 2.0, 84.54% at pH 4.0 and 97.80% at pH 10.0 resulting in a V-shaped pattern (Figure 3). A good number of studies have shown that the pH-emulsifying properties profile for different proteins including soya protein is quite similar to their corresponding pH-solubility profile [12]. This means that solubility and/or electrostatic charge largely affects emulsifying properties. Most food proteins are generally poor emulsifier near their is electric pH and they cannot move rapidly to the interface due to lack of electrostatic repulsive force at this pH. These proteins may, however, be effective emulsifier when moved away from their isoelectric pH [16-17].

Viscosity: Viscosity measurement over a wide range of pH and concentration is very important in food formulation. Concentration dependency of viscosity was also reported for sunflower protein isolate and soybean protein isolate [20]. The result of viscosity of Crotalaria pallida has been presented in (Table 3). Generally it has been observed that viscosity increases with the increase in concentration at room temperature (26°C). It is also observed that the viscosity of Crotalaria pallida seed protein isolate in slightly greater in alkaline range than that of acidic range. As the conformation of protein chain changes with pH, viscosity becomes pH dependent also.

**Table 3:** Effect of protein concentration and pH on viscosities of C. pallida seed protein isolate at room temperature (26°C).

<table>
<thead>
<tr>
<th>Protein Concentration (g/100 ml)</th>
<th>Viscosity (cp)</th>
<th>pH</th>
<th>Viscosity (cp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.94 ± 0.11</td>
<td>4</td>
<td>0.91 ± 0.15</td>
</tr>
<tr>
<td>2</td>
<td>0.99 ± 0.03</td>
<td>6</td>
<td>0.90 ± 0.20</td>
</tr>
<tr>
<td>1.5</td>
<td>0.99 ± 0.03</td>
<td>6</td>
<td>0.90 ± 0.20</td>
</tr>
<tr>
<td>2</td>
<td>1.04 ± 0.01</td>
<td>8</td>
<td>1.00 ± 0.03</td>
</tr>
<tr>
<td>2.5</td>
<td>1.12 ± 0.03</td>
<td>10</td>
<td>1.09 ± 0.02</td>
</tr>
<tr>
<td>3</td>
<td>1.23 ± 0.01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>1.57 ± 0.01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>1.96 ± 0.02</td>
<td></td>
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</tr>
</tbody>
</table>

*Values are mean of triplicate determinations (± S.D).

**Foaming properties:** The results of foaming capacity (FC) indicate that pH of the protein solution is an important factor governing the extent of foam formation as well as its stability in the medium. Foam formation is dependent on three crucial factors: transportation, penetration and reorganization at the air-water interface. Therefore, high foaming capacity is normally being observed apart from its is electric pH, where net charge increases on the protein resulting in unfolding of peptide chains. This allows protein to diffuse rapidly into the air-water interface to encapsulate air particles and then enhance foam formation [18]. The lowest FC (122%) is observed at pH 4.0 and the highest FC (178%) at pH 10.0. Maximum foam stability is observed at pH 4.0 and it gradually decreases with increase in pH value (alkaline medium). Maximum foam stability near the isoelectric pH is is mainly attributed to the formation of stable molecular layers in air-water interface where the net charge on peptide is minimal. Such pH dependency is also observed for Soybean and Sunflower protein [19]. The results are presented at (Figures 4 & 5).

**Gelation capacity:** The least gelation capacity (LGC) for Crotalaria pallida seed protein isolate is found to be 10% in distilled water medium and 8% in 0.5 (M) NaCl solution as a medium. These values are in agreement with the values reported by Okezie and Bello for soybeans powder (14%) [21], Lqari et al. reported that LGC of lupin protein concentrate is 12% [22], while Schmidt reported 7.5% for wheat protein isolate [23]. According to Schmidt, for a given type of protein, a critical concentration is required for the formation of a gel and the type of gel varies with the protein concentration. Considerably, higher protein concentration is usual required for the gelation of globular proteins. Thus, C. pallida seed protein may be globular in nature.

**Conclusion**

Functional properties of Crotalaria pallida seed protein isolate have been evaluated to screen its usability in food systems. Due to its high solubility in wide range of pH values, it can be a good source of protein. The values are quite comparable with commercially available protein foods such as soybean, sunflower, cashew nuts, pea nuts etc. Experimental results...
reveal that it can be useful in food systems such as whipped
toppings, chiffon deserts, angel and sponge cakes etc. (owing
to its high FC and FS), meat processing industry and sausages
(require high EA and ES), doughnuts (requiring high OHC)
etc. For these mentioned properties, the seed protein isolate of
*Crotalaria pallida* is very attractive as functional ingredients in
food systems but sensory and texture analysis of the product
would be necessary.

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**Statement**

This article has not been submitted anywhere else for
publication. Neither it has been published before in any
other journal nor has it been presented in any conference or
symposium.

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