Protein concentrates obtained from pig by-products using pH-shifting technique: a preliminary study

Claudine Garcia Chaves da Costa¹* Marielle Maria de Oliveira Paula¹
Armando Abel Massingue² Robledo de Almeida Torres Filho³
Eduardo Mendes Ramos¹ João de Deus de Souza Carneiro¹

¹Departamento de Ciência dos Alimentos, Universidade Federal de Lavras (UFLA), 37200-000, Lavras, MG, Brasil. E-mail: claudine.gchaves@gmail.com.
²Corresponding author.
³Escola Superior de Desenvolvimento Rural, Universidade Eduardo Mondlane (UEM), Vilankulo, Mozambique.

ABSTRACT: This study aimed to investigate the extraction and characterization of protein concentrates from pig by-products (heart, liver and kidneys) using the pH-shifting technique. From the solubility profiles (pH 2 to 12), the protein extraction was performed at alkaline pH (10.0 to 11.5), obtaining two fractions: soluble (SC) and insoluble (IC). Higher protein content (71 to 77%) and extractability (214 to 459 mg/g) were observed in heart and liver concentrates; whereas, for water holding capacity (WHC) the highest values (4.20 to 4.54 g water/g protein) were for the heart (SC) and kidney (SC and IC) concentrates. All concentrates had high emulsion stability and higher WHC than commercial protein extenders (whey and soybean). The concentrates obtained from the soluble fraction were redder (higher a* values and lower h values) and darker (lower L* values) than insoluble fraction, especially heart and liver concentrates. Use of concentrates in sausage production slightly altered the color chroma (C*) of samples. It was concluded that the pig by-products protein concentrates had great potential of use as extenders in sausage production.

Key words: heart, kidney, liver, protein extensor, pH-shifting.

INTRODUCTION

In recent years, consumption of pork and further processed pork products has greatly increased in Brazil. Brazil’s pork production from 2010 to 2017 increased from 2.3 to 3.8 million tons, being 81.5% of this production sold on the domestic market (ABPA, 2018). Considering this production, large quantities of by-products are generated at the slaughter process. Although, the pork consumption (14.7 kg/per capita in 2017) is high in Brazil, offal’s are little used for direct consumption, with most of the production destined for export (10.8% of pork products exported in 2017) or for use in processed meat products.

Despite their low acceptance, offal’s have relatively low cost, with high nutritional value, rich in proteins, and contribute to improve sensorially, technologically and increase the stability of processed products (TOLDRÁ et al., 2012). However, in
Brazilians also tend to use large offal’s content, being their addition in traditional meat products, as in emulsified sausages, limited to 10% of the product. This leads to a surplus of these by-products, especially in the small-scale abattoir operating in regional and local markets, which are considered the main suppliers of pork.

Certainly, there is an underutilization of pork offal’s such as liver, heart and kidney from which a more rational use is possible, especially due to their high protein content. One of the alternatives to increase the value of slaughterhouse by-products and make it more suitable for further processing would be to extract their proteins for use as functional ingredients in meat products, for instance as emulsifying or gelling agent, using similar technologies (SELMANE et al., 2008; HRYNETS et al., 2011). There are differences in functionality of proteins, which are closely related to their water holding capacity, emulsify capacity and gelling agent, being influenced by several factors, including protein source and process utilized. Therefore, a method that maintains protein functionality is necessary for the successful production of protein concentrates that have this desirable property (DEWITT et al., 2002).

Protein recovery using a pH-shifting process is a promising technique to recover meat proteins economically, at a relative high recovery rate and with improved functionality (HRYNETS et al., 2011). The process also known as isoelectric solubilization/precipitation (ISP) utilizes the principle that the solubility of a comminute protein-containing material homogenized in water is affected by the pH of the mixture. During ISP processing, meat proteins are first solubilized at low or high pH (acid or alkali), followed by precipitation at the isoelectric point to separate neutral and membrane lipids and further pH adjustment to the original meat pH (NOLSOE et al., 2009; TAHERGORABI et al., 2011). Despite fish material, the ISP processing has been applied to beef (heart and lungs), pork (lungs) and poultry (dark chicken-meat and mechanically deboned chicken and turkey meat) processing by-products (DEWITT et al., 2002; SELMANE et al., 2008; MOAYEDI et al., 2010; HRYNETS et al., 2011; TAHERGORABI et al., 2011).

Therefore, the overall objective of this study was to obtain and characterize protein concentrates from three pig by-products, namely liver, heart and kidney, by ISP process to initial screening for subsequent application as food ingredients.

MATERIALS AND METHODS

Fresh pig by-products (kidney, heart and liver), were purchased vacuum-packaged from a local abattoir and frozen (-18 °C) until used. Commercial ingredients, whey protein concentrate (WPC, 54.82% protein, 10.41% moisture, 3.47% fat, 5.18% ash and 26.13% carbohydrate; Tate & Lyle Gemachom Tech, Juiz de Fora, MG, Brazil) and soy protein isolate (SPI; 84.88% protein, 9.47% moisture, 3.54% fat and 3.09% ash; New Nutrition, São Paulo. SP, Brazil), were used as reference materials.

In order to find the effect of different pH on the solubility of proteins in raw material, a solubility curve was created by each by-product as described by HRYNETS et al. (2011) with minor modifications. Six grams of raw material were homogenized (TE-102; Tecnal Equipamentos Científicos, Brasil) with 300 mL deionized water (at 2 °C) for approximately 2 min. Aliquots of 30 mL of the homogenate were adjusted from pH 1.5 to 12.0 in 0.5 intervals, using HCl or NaOH solutions (0.2 M or 1.0 M), with a pH meter (DM 20; Digimed, Brazil). The homogenate was centrifuged (Hettich EBA 21 centrifuge; Hettich GmbH & Co.KG, Germany) at 5000×g for 15 min. Protein concentration of each supernatant was determined using the Biuret colorimetric method, using bovine serum albumin (BSA, Sigma Chemical Co.) as standard. Protein solubility was expressed as milligram per milliliter (mg/mL). Two replications were carried out for each measurement. The pH in which the highest protein solubility was obtained was selected to obtain the proteins concentrates.

Protein extractions were conducted using a pH-shifting with modification of the process described by HRYNETS et al. (2011). The alkaline-aided protein recovery was chosen after protein solubility profile analysis. For each by-product, 200 g of raw material were homogenized with cold (2 °C) distilled water (1:5 meat: water ratio, w/v) in an industrial blender for 15 min. The pH of each homogenate was adjusted by drop-wise addition of NaOH 0.2 or 1.0 M to reach the maximum solubility points with pH values of 10.0 (liver) and 11.5 (heart and kidney), as determined from the solubility profile. The homogenate was centrifuged (Heraeus™ Megafuge™; Thermo Fisher Scientific, USA) at a rotational speed of 19,000×g for 20 min at 4 °C. Upper layer (neutral lipids) was manually removed with the help of a stainless-steel spatula and two fractions were obtained: the supernatant fraction (soluble proteins); and the pellet (insoluble proteins). It was decided not to carry out the isoelectric precipitation in the supernatant, which was
conducted directly for lyophilization. Both fractions were frozen (at -20 °C) and subjected immediately to a lyophilization procedure in bench-top freeze drier (4.5L Labconco, USA). Dry concentrates were subjected to the ball milling process, for particle reduction and homogenization, vacuum packed and stored (at room temperature) until analysis.

Six protein concentrate were obtained: heart supernatant concentrate (HSC); heart pellet concentrate (HPC); liver supernatant concentrate (LSC); liver pellet concentrate (LPC); kidney supernatants concentrate (KSC); and kidney pellet concentrate (KPC). The by-products raw material and the concentrates were characterized according to its proximate composition (moisture, protein, fat and ash), by the AOAC procedures (AOAC, 2012).

The protein extractability was evaluated according to modifications of the procedures described by NOLLET e TOLDRÁ (2011). About 0.2 g of each protein concentrate was diluted in 10 mL of 0.05 M phosphate buffer solution (pH 7.4), vortexed and centrifuged (EBA 21; Hettich GmbH & Co.KG, Germany) at 5,000×g for 15 min. Protein concentration of the supernatant was determined by Biuret method and water-soluble proteins (WSP) expressed as mg protein/g sample. The pellet was suspended in 10 mL of 0.05 M phosphate buffer (pH 7.4) with 1.1 M NaCl, vortexed and centrifuged at 5,000×g for 15 min. The protein concentration of the new supernatant was determined by Biuret method and the salt-soluble proteins (SSP) expressed as mg protein/g sample.

The water holding capacity (WHC) was evaluated by modification of the centrifuge method described by NOLLET e TOLDRÁ (2011). About 0.2 g of each protein concentrate was diluted in 10 mL of 0.05 M phosphate buffer solution (pH 7.4), vortexed and centrifuged (EBA 21; Hettich GmbH & Co.KG, Germany) at 5,000×g for 15 min. Protein concentration of the supernatant was determined by Biuret method and water-soluble proteins (WSP) expressed as mg protein/g sample. The pellet was suspended in 10 mL of 0.05 M phosphate buffer (pH 7.4) with 1.1 M NaCl, vortexed and centrifuged at 5,000×g for 15 min. The protein concentration of the new supernatant was determined by Biuret method and the salt-soluble proteins (SSP) expressed as mg protein/g sample.

The WHC was calculated as the difference between hydrated weight and original weight of sample, being expressed as g water/g and as g water/g protein.

The CIE lightness (L'), redness (a') and yellowness (b') were obtained from the specular light excluded (SCE) mode readings using a spectrophotometer (CM-700d; Konica Minolta, USA), with 8 mm aperture port, illuminant D65 and 10° stander observer. The pig by-products protein concentrates and the commercial proteins ingredients (WPC and SPI) were placed in Petri glass dishes and the analysis was performed at six different points on the surface. The color coordinates were also expressed on the CIELCH system, with chroma (C*) and hue angle (h) being calculated as (RAMOS e GOMIDE, 2017): $C^* = (a'^2 + b'^2)^{0.5}$ and $h = \tan^{-1}(b'/a')$.

To evaluate the practical application of the pig by-product concentrates in meat products, an emulsion-type sausage model was elaborated according to MASSINGUE et al. (2018). Sausages (62% mechanically deboned poultry meat, 13% pork meat, 20.4% water, 2% salt, 0.5% sodium tripolyphosphate, 0.02% sodium nitrite and 0.1% sodium erythorbate) were prepared with 2% of each pig-by-product protein concentrate, WPC or SPI. The raw batter mixture was used to evaluate emulsion stability and the cooked sausage obtained to the instrumental color evaluations.

Stability of the emulsion was evaluated by HUGHES et al. (1997), with minor modifications. About 25 g of the raw emulsion batter was weight into 50 mL polyethylene centrifuge tubes, centrifuged (EBA 21; Hettich GmbH & Co.KG, Germany) at 3,000×g for 1 min to compact the batter. The tubes were heated in a water bath (at 70 °C) for 30 min, cooled in tap water and again centrifuged at 3,000×g for 3 min. All tubes were then dropped for 30 min in pre-weighted porcelain crucibles, to collect all released supernatant (fluid), which were dried overnight at 105 °C. The volumes of total expression fluid (TEF) and the percentage fat (% Fat) were calculated as follows: $\text{TEF} = (\text{Weight of crucible + fluid}) – (\text{Weight of empty crucible})$; $\%\text{TEF} = 100 \times \frac{\text{TEF}}{\text{sample weight}}$; and $\%\text{Fat} = 100 \times \left(\frac{[\text{Weight of crucible + dried fluid}]}{\text{TEF}}\right)$ (Weight of empty crucible)/TEF.

The CIE color was performed on cooked samples obtained after the emulsion stability analysis. Samples were cut crosswise at different points, obtaining three cylinders of 2.0 cm height, and the color indices obtained by three readings at different points of the internal surface of each cylinder as described for the protein concentrates.

The experiment was carried out in a completely randomized design with one factor, containing six (pig by-products concentrates) or eight (pig by-products concentrates plus WPC and SPI) levels, and three replicates. Main effects were evaluated by F-test (ANOVA) and means were separated using Tukey's test ($P<0.05$). Statistical analyzes were performed in the SAS software (Statistical Analysis System, Institute Inc., Cary, NC, USA).

RESULTS AND DISCUSSION

The protein solubility profile for pig liver, heart and kidney with pH ranging from 1.5 to 12.0 are described in the figure 1. The lowest solubility (or
highest precipitation) was observed in the pH range of 4.5 to 5.5, with lower values recorded by heart at pH 5.5 (0.38 mg/mL), followed by kidney at pH 4.5-5.0 (mean of 0.83 mg/mL) and by liver at pH 5.0 (mean of 1.18 mg/mL). At the isoelectric point, protein-water interactions are minimal, whereas aggregates of weak protein-protein hydrophobic interactions are present, causing precipitation of proteins (FENNEMA, 1996). The pH observed for the lower solubility of the by-products is close to the isoelectric points (pH 5.5) observed for the majority of muscle proteins (XIONG, 2018).

Notably, an increase in solubility was observed during acidification or alkalization of the extraction media. The protein solubility was higher at higher pH values for liver (5.23 mg/mL between pH 9.0 and 12.0) and kidney (3.81 mg/mL between 9.0 and 10.0) than at lower pH values (2.89 mg/mL liver and 3.53 mg/mL kidney, both at pH 2.5). For heart, however, the highest protein solubility (mean of 3.42 mg/mL) was observed in low (2.0) and high (12.0) pH values. These differences in the protein solubility profile might be attributed to differences in the ionizable groups of the polypeptide chains of proteins (FENNEMA, 1996) of each by-product. Based on the solubility study, alkaline-aided protein recovery (pH between 10.0 and 12) were selected for the protein extraction.

Table 1 summarize results from physicochemical and technological characterization of the pig by-products concentrates. Concentrates with higher protein content were obtained from sources that contained lower moisture (75.72% heart and 73.90% liver) and highest protein content (20.76% heart and 22.57% liver) and, except for the kidney concentrates, higher protein was retained in the soluble fraction. Higher amounts of insoluble stromal proteins (collagen content) in kidney (OCKERMAN and HANSEN, 2000) may have contributed to the absence of different between fractions of protein concentrates for this raw material. Moreover, lower values of protein content for kidney concentrates seems to be due to the lower values of protein in raw material (16.23%) and greater retention of water (due to the higher moisture content – 81.08%) after drying. In the raw material, higher fat contents were observed in the liver (3.81%), followed by kidney (3.27%) and heart (2.57%).

In relation to the ash content, higher values were observed in the heart (1.22%) than in the kidney and liver raw material samples (mean of 0.82%). However, a greater concentration in the soluble fraction of the concentrates of heart and kidney was observed. CHEN et al. (2007) demonstrated that basic treatments during solubilization of trout filleting processing by-products (contained bones with left-over fish meat, skin, scales, and fins) yielded protein concentrate with less minerals (Mg, Ca, and P) than the insoluble fractions.
in the present research, it should be considered that the raw material used to obtain the concentrates is richer in heme iron, which are associated with water-soluble heme pigments (hemoglobin and myoglobin) (RAMOS e GOMIDE, 2017). Therefore, this would explain the higher values in the soluble fraction. From the nutritional point of view, it is of interest to assess how these minerals are separated from the recovered protein during the pH-shifting processing.

The total extractable proteins and their fractions (water- and salt-soluble proteins) were affected (P<0.05) by the treatments. Extractability is an important property since the amount of protein available in the solution affects the functional properties expected from proteins (HRYNETS et al., 2011). Overall, the total extractable proteins were higher in the soluble concentrates from by-products with highest protein content (liver and heart). Also, higher water-soluble proteins (WSP) were observed in all concentrates than salt-soluble proteins (SSP), which confirmed that a sizeable amount of WSP is recovered, during pH-shifting processes (HRYNETS et al., 2011; MATAK et al., 2015). Liver concentrate from soluble fraction (LSC) had higher values of WSP than salt-soluble proteins (SSP), for kidney concentrates, which may also be due to the greater presence of insoluble stromal proteins in this by-product. Moreover, there was no difference in the WSP extractability between concentrate fractions and.

Table 1 - Physicochemical and technological characteristics of pig by-products protein concentrates and its effects in sausage formulation.

<table>
<thead>
<tr>
<th></th>
<th>Heart (HSC)</th>
<th>Kidney (KSC)</th>
<th>Liver (LSC)</th>
<th>Heart (HPC)</th>
<th>Kidney (KPC)</th>
<th>Liver (LPC)</th>
<th>WPC</th>
<th>SPI</th>
<th>SEM</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Composition</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Moisture (%)</td>
<td>9.56°</td>
<td>14.78°</td>
<td>8.90°</td>
<td>8.20°</td>
<td>11.50°</td>
<td>8.19°</td>
<td></td>
<td></td>
<td>0.58</td>
</tr>
<tr>
<td>Protein (%)</td>
<td>77.30°</td>
<td>67.45°</td>
<td>73.48°</td>
<td>73.51°</td>
<td>66.22°</td>
<td>70.32°</td>
<td></td>
<td></td>
<td>0.94</td>
</tr>
<tr>
<td>Fat (%)</td>
<td>5.51°</td>
<td>10.46°</td>
<td>14.81°</td>
<td>14.80°</td>
<td>14.18°</td>
<td>14.87°</td>
<td></td>
<td></td>
<td>0.88</td>
</tr>
<tr>
<td>Ash (%)</td>
<td>8.98°</td>
<td>8.19°</td>
<td>4.85°</td>
<td>3.92°</td>
<td>6.94°</td>
<td>4.84°</td>
<td></td>
<td></td>
<td>0.46</td>
</tr>
<tr>
<td><strong>Extractable proteins</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total (mg/g)</td>
<td>339°</td>
<td>177°</td>
<td>460°</td>
<td>249°</td>
<td>114°</td>
<td>214°</td>
<td></td>
<td></td>
<td>28</td>
</tr>
<tr>
<td>WSP (mg/g)</td>
<td>180°</td>
<td>123°</td>
<td>280°</td>
<td>193°</td>
<td>76°</td>
<td>150°</td>
<td></td>
<td></td>
<td>16</td>
</tr>
<tr>
<td>SSP (mg/g)</td>
<td>159°</td>
<td>54°</td>
<td>179°</td>
<td>56°</td>
<td>38°</td>
<td>64°</td>
<td></td>
<td></td>
<td>14</td>
</tr>
<tr>
<td>WHC</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(g water/g sample)</td>
<td>3.30°</td>
<td>2.83°</td>
<td>1.67°</td>
<td>2.63°</td>
<td>2.92°</td>
<td>2.28°</td>
<td></td>
<td></td>
<td>0.15</td>
</tr>
<tr>
<td>(g water/g protein)</td>
<td>4.27°</td>
<td>4.20°</td>
<td>2.27°</td>
<td>3.58°</td>
<td>4.54°</td>
<td>3.24°</td>
<td></td>
<td></td>
<td>0.20</td>
</tr>
<tr>
<td><strong>Emulsion stability</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>%TEF</td>
<td>0.26°</td>
<td>0.52°</td>
<td>0.45°</td>
<td>0.30°</td>
<td>0.66°</td>
<td>0.54°</td>
<td></td>
<td></td>
<td>0.32</td>
</tr>
<tr>
<td>%FAT</td>
<td>8.94°</td>
<td>5.88°</td>
<td>8.76°</td>
<td>8.62°</td>
<td>6.76°</td>
<td>6.88°</td>
<td></td>
<td></td>
<td>0.34</td>
</tr>
<tr>
<td><strong>CIE color: Concentrates</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Luminosity (L*)</td>
<td>60.21</td>
<td>70.88</td>
<td>64.07°</td>
<td>76.34°</td>
<td>78.35°</td>
<td>73.19°</td>
<td></td>
<td></td>
<td>2.58</td>
</tr>
<tr>
<td>Redness (a*)</td>
<td>15.42°</td>
<td>10.08°</td>
<td>13.41°</td>
<td>9.11°</td>
<td>8.90°</td>
<td>9.51°</td>
<td></td>
<td></td>
<td>0.79</td>
</tr>
<tr>
<td>Yellowness (b*)</td>
<td>14.14°</td>
<td>18.19°</td>
<td>19.33°</td>
<td>16.08°</td>
<td>18.32°</td>
<td>18.97°</td>
<td></td>
<td></td>
<td>0.38</td>
</tr>
<tr>
<td>Chrome (C*)</td>
<td>20.92°</td>
<td>20.80°</td>
<td>23.52°</td>
<td>18.48°</td>
<td>20.37°</td>
<td>21.22°</td>
<td></td>
<td></td>
<td>0.53</td>
</tr>
<tr>
<td>Hue angle (h°)</td>
<td>42.53°</td>
<td>61.00°</td>
<td>55.26°</td>
<td>60.47°</td>
<td>64.09°</td>
<td>63.83°</td>
<td></td>
<td></td>
<td>2.14</td>
</tr>
<tr>
<td><strong>CIE color: Sausages</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Luminosity (L*)</td>
<td>61.95</td>
<td>62.76</td>
<td>61.66</td>
<td>63.44</td>
<td>64.27°</td>
<td>61.69°</td>
<td></td>
<td></td>
<td>0.32</td>
</tr>
<tr>
<td>Redness (a*)</td>
<td>6.08°</td>
<td>5.87°</td>
<td>7.04°</td>
<td>5.78°</td>
<td>5.31°</td>
<td>5.93°</td>
<td></td>
<td></td>
<td>0.17</td>
</tr>
<tr>
<td>Yellowness (b*)</td>
<td>18.08</td>
<td>16.94</td>
<td>18.21</td>
<td>17.27°</td>
<td>17.87°</td>
<td>17.90°</td>
<td></td>
<td></td>
<td>0.19</td>
</tr>
<tr>
<td>Chrome (C*)</td>
<td>19.08°</td>
<td>17.98°</td>
<td>19.55°</td>
<td>18.22°</td>
<td>18.64°</td>
<td>18.86°</td>
<td></td>
<td></td>
<td>0.19</td>
</tr>
<tr>
<td>Hue angle (h°)</td>
<td>71.40</td>
<td>70.73</td>
<td>68.75</td>
<td>71.53°</td>
<td>73.44°</td>
<td>71.63</td>
<td></td>
<td></td>
<td>0.54</td>
</tr>
</tbody>
</table>

WPC = whey protein concentrate; SPI = soy protein isolate; S.M.E = standard error of means; WSP = water-soluble proteins; SSP = salt-soluble proteins; WHC = water holding capacity; %TEF = total expressive fluid; and %FAT = fat content in expressible fluid.

*Means followed by different letters in the same row differ (P<0.05) by Tukey’s test.
for each type of by-products, but higher SSP extract ability was observed in the soluble fraction of heart and liver concentrates; with the same behavior as observed for the protein content.

Contrary to that observed for total protein extractability, higher protein content did not guarantee a higher WHC; although, the highest WHC were observed in the HSC when expressed as g water/g sample. Taking into account the protein content of the concentrate, higher values of WHC (g water/ g protein) were observed in the soluble fraction of heart and liver concentrates; with the same behavior as observed for the protein content.

However, it is interesting to note that all pig by-products concentrates had WHC higher (P<0.05) than the whey concentrate (WPC) and, except for the LSC, than the soy protein isolate (SPI). Moreover, the emulsion stability (total expressive fluid; % TEF) of the elaborated sausages was not influenced (P>0.05) by the type of raw material or extraction fraction, being not different from sausages elaborated with WPC, but higher (lower % TEF values) than SPI. Emulsion stability differences between the by-products concentrate were observed only in relation to the amount of fat (% FAT) present in the exudate. These results suggested that the by-products concentrates had higher potential of use as extensors in food products, with good emulsify capacity.

Since in the pH-shifting process is expected extraction of a large amounts of sarcoplasmic proteins, among then the heme pigments (myoglobin and hemoglobin), the color of the concentrates becomes very important when applied in food. As expected, all color indexes (P<0.05) were affected by treatments. Overall, concentrates obtained from the soluble fraction are redder (higher a* values and lower h values) and darker (lower L* values) than the insoluble ones, which can be justified by the fact that the sarcoplasmic proteins are readily soluble in water and low ionic strength solutions (XIONG, 2018). In this sense, it is also justifiable that the reddest and darkest concentrates are those obtained from sources with higher contents of heme pigments, namely heart and liver (OCKERMAN e HANSEN, 2000).

All pig by-products concentrates had redder and darker color than the commercial extensors WHC and SIP commonly used in food products and these differences are illustrated in figure 2a. Nerveless, despite these differences in the concentrate’s colors, color of sausages on which they were used was very little affected by the type of concentrate used. Only redness (a’) and chroma (C*) were affected (P<0,05) by the treatments, but these differences are probably imperceptible (as can be observed in figure 2b).
CONCLUSION

Protein concentrate extracts processed from pig offal’s (liver, heart and kidney) is an enriched protein content above 60%, with water holding capacity and emulsion stability equal to or better than the commercial protein extensor used as ingredient in meat processing, namely soybean protein isolate and whey protein concentrate. The color of the by-products protein concentrates seems not affected the color of the meat product in which it is used.

Thus, it can be concluded that the production of protein concentrates from pig by-product is an effective alternative for its harnessing and added commercial value, but their development should be improved and other characteristics (nutritional, sensory and even technological) evaluated.

ACKNOWLEDGEMENTS

This study was financed in part by the and Coordenação de Aperfeiçoamento de Pessoal de Nível Superior (CAPES) for the scholarship grants to the first (Master Degree) author. The authors would like to thank also the Fundação de Amparo a Pesquisa do Estado de Minas Gerais (FAPEMIG) and the Conselho Nacional de Desenvolvimento Científico e Tecnológico (CNPq) for the financial support.

DECLARATION OF CONFLICT OF INTERESTS

The authors declare no conflict of interest. The founding sponsors had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, and in the decision to publish the results.

AUTHORS’ CONTRIBUTIONS

All authors contributed equally for the conception and writing of the manuscript. All authors critically revised the manuscript and approved of the final version.

REFERENCES


