ADAPTATION OF TWO-DIMENSIONAL ELECTROPHORESIS FOR MUSCLE TISSUE ANALYSIS

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ABSTRACT

It is important to understand the molecular mechanisms that take place in muscle tissues and to predict meat quality characteristics. One of the most popular methods is two-dimensional electrophoresis, which allows us to visualize, share and identify different molecules, including meat proteins. However, the standard conditions of this method are not universal for all types of raw material, so the authors suggest a new variation of two-dimensional electrophoresis for muscle tissue analysis. Samples were tested by the classical version of isoelectric focusing (cathode buffer in the top and anode buffer in the bottom chamber of the electrophoresis cell) and its variation (anode buffer in the top and cathode buffer in the bottom chamber of the electrophoresis cell). Next, extruded gels were incubated in two different buffer systems: the first was equilibration buffer I (6 M urea, 20% w/v glycerol, 2% w/v SDS and 1% w/v Dithiothreitol in 375 mM Tris-HCl buffer, pH 8.8) followed by equilibration buffer II (6 M urea, 20% w/v glycerol, 2% w/v SDS and 4% w/v iodoacetamide in 375 mM Tris-HCl buffer pH 8.8 and the second, buffer A, consisting of 5 M urea, 2% w/v SDS, 5% w/v mercaptoethanol, 62.5 mM Tris-HCl buffer, pH 6.8 and 0.01% w/v bromophenol blue. Electrophoretic studies of muscle tissue revealed the best protein separation after changing the direction of the current (authors’ variation), while no differences were detected after changing incubation buffers.

Keywords: two-dimensional electrophoresis; muscle protein; isoelectric focusing; meat

INTRODUCTION

Meat and meat products will always play an important role in the human diet as a source of high-grade protein. There is a wide variety of meat processing products, the range of which is regularly updated. Depending on the processing technology, meat proteins undergo extensive modifications that affect their quality, shelf life, nutritional properties, and health effects. At the same time, tenderness and juiciness are the criteria most judged by consumers (Cao et al., 2020; Hollung et al., 2007).

These criteria are influenced by the following factors: genetics, environment, animal welfare, and further processing. The molecular mechanisms underlying such processes are still of interest. However, genetic information remains static throughout the life of the body, while the protein composition is dynamic and changes depending on factors affecting protein synthesis or degradation (Peng and Gygi, 2001). Thus, proteomics analysis makes it possible to better understand the molecular mechanisms that occur in tissues and to predict the most important characteristics, particularly in the case of meat (Bendixen, 2005; Bendixen et al., 2005; Zamaratskaia and Li, 2017).

Proteomic methods over the past decade have found increasing application in various fields of science and agriculture. To date, proteomic tools have improved significantly, and mass spectrometric methods are being developed very rapidly. Most proteomic methods are based on the separation of proteins in at least two directions, using chromatographic or electrophoretic methods, with further identification using mass spectrometry methods (Soares et al., 2012; Suman et al., 2014). In this paper, attention is focused on the method of two-dimensional gel electrophoresis (2-DE), because it is still an excellent way to visualize protein components and widely used in meat science, with an emphasis on sample preparation and aspects of isoelectric focusing (IEF).

Traditionally, 2-DE is carried out by fractionation of proteins according to two different physicochemical parameters. In the first direction, proteins are separated by charge according to their isoelectric point with IEF, and then in the second direction by their molecular weight with polyacrylamide gel electrophoresis (PAGE). Some variations of conducting 2-DE are discussed below to identify the most optimal variant for analysis of the meat proteome.
**Scientific hypothesis**

The classical variation of two-dimensional electrophoresis does not fully reveal muscle tissue proteins. It is necessary to select optimal conditions for the analysis. Changing the IEF parameters and additional incubation in lysis buffers will increase the resolution of the 2-DE method for muscle proteins.

**MATERIAL AND METHODOLOGY**

**Materials**

Chemical reagent: Urea, Thiourea, Dithiothreitol, Sodium hydroxide (NaOH), Glycerol, Sodium dodecyl sulfate (SDS), Tris, Acrylamide, Ammonium persulfate (APS), 2-Propanol, Acetic acid (PanReac, Spain); Bis-acrylamide, Tetramethylethylenediamine (TEMED), Mercaptoethanol, Bromophenol blue, Glycine, Coomassie Brilliant Blue G-250, Triton X-100 (Helicon Russia); Ampholyte (Serva, Germany) and Phosphoric (V) acid (H3PO4) (Component-reactiv, Russia).

The object of the study was the Vietnamese Pot-bellied from healthy females of 60 – 65 days old pig L. dorsi muscle. Samples were taken within 20 minutes after slaughter and placed in dry ice. Frozen muscle tissues (50 mg) were homogenized in 1 mL 7 M Urea, 2 M Thiourea, 1% Dithiothreitol, 0.4% Triton X-100, 2% pH 3-10 Ampholyte. Homogenates were centrifugated with an acceleration of 20 000 g for 20 minutes. Three samples, obtained from different animals, were studied by two-dimensional electrophoresis in two variations of isoelectric focusing (IEF) and two methods of gel incubation after IEF. Figure 1 shows the experimental design.

Two-dimensional gel electrophoresis (2-DE)

IEF in the first dimension was performed at 3650 V.h⁻¹. The anodic and cathodic electrode solutions used for IEF were 0.01 M Phosphoric (V) acid and 0.02 M Sodium hydroxide, respectively, in 2.4 mm × 160 mm tube gels. In the first classical version of IEF, the cathode buffer (0.02 M sodium hydroxide) was in the upper chamber of the electrophoresis cell, and the anode buffer (0.01 M orthophosphoric acid) was in the lower one. In the second variation of IEF, the electric current direction was changed: the anode buffer was in the upper chamber, and the cathode buffer was in the lower one.

After the IEF, gels were incubated in two different ways: extruded tube gels were incubated for 10 min, in 2.5 mL of equilibration buffer I (6 M urea, 20% w/v glycerol, 2% w/v SDS and 1% w/v Dithiothreitol in 375 mM Tris-HCl buffer, pH 8.8), followed by equilibration buffer II (6 M urea, 20% w/v glycerol, 2% w/v SDS and 4% w/v iodoacetamide in 375 mM Tris-HCl buffer, pH 8.8); extruded tube gels were incubated for 10 – 15 min, in 2.5 mL buffer A (5 M urea, 2% w/v SDS, 5% w/v mercaptoethanol, 62.5 mM Tris-HCl buffer, pH 6.8 and 0.01% w/v Bromophenol blue). For SDS-PAGE (12% T, 2.6% C) equilibrated tube gels were transferred to a 12.5% polyacrylamide gel (170 mm × 180 mm × 1.5 mm). Electrophoresis was carried out with a gel running buffer containing 25 mM Tris-HCl, 192 mM glycine and 0.1% w/v SDS at 30 mA per gel until the bromophenol blue front had reached the lower edge of the gel. Experimental molecular weights were detected relatively to marker proteins – 250; 150; 100; 70; 50; 40; 30; 20; 15; 10; 5 kDa (Thermo Scientific, Lithuania).

**Protein visualization and image analysis**

Protein spots were visualized staining by solution Coomassie (0.05% w/v Coomassie Brilliant Blue G-250, 10% v/v Acetic acid and 25% v/v 2-Propanol) for 1 hour. The destaining procedure was carried out by incubating in 10% v/v Acetic acid for 15 minutes several times until the background of the gel became transparent.

For computerized densitometry, two-dimensional electropherograms were used in a wet state. Their full digital images and/or images of individual fragments were obtained using a Bio-5000 plus scanner (Serva, Germany). Scanned images were analyzed with ImageMaster™ 2D Platinum software powered by Melanie 8.0 (GE Healthcare and Genebio, Switzerland). Spots were detected and quantified automatically with minimum thresholds: saliency – 11, min area – 5 and smooth – 3. The relative optical density (OD) and relative volume were computed to correct for differences in gel staining. These measures take into account variations due to protein loading and staining, by considering the total OD or volume over all the spots in the gel. The digitized 2DE images of the cortex were then compared by the matching method (Grove et al., 2006).

Protein spots on muscle tissue two-dimensional electropherograms were interpreted following the Swiss-Prot database (O'Donovan et al., 2002) and the Muscle organ proteomics database (Kovaleva et al., 2013).

**Statistical analysis**

The experimental data were analyzed using ordinary one-way ANOVA (between gels, obtained with different variation IEF) by ImageMaster™ 2D Platinum software powered by Melanie 8.0 (GE Healthcare and Genebio, Switzerland).

A p value <0.05 was considered to indicate a significant difference. All results are presented as mean ± SD from at least three independent experiments.

**RESULTS AND DISCUSSION**

In most scientific works, IEF is performed using a thin layer of gel deposited on a plastic substrate (IPG Dry Strip) to speed up and simplify the process. (Naveena et al., 2017; Di Luca et al., 2016; Lee, Saraygord-Afshari and Low, 2020). However, the glass-tube IEF, despite process complexity, has an advantage in the resolution and protein loading volume in the gel (Matsumoto et al., 2019). Based on the foregoing, for this experiment, we selected glass-tube IEF.

In accordance with the experimental design, depicted in Figure 1, four 2-DE variations were performed; the obtained electropherograms are presented in Figure 2. A different distribution of pig muscle protein was noted for molecular weights (MW) and isoelectric points (pI).
Figure 1 Experimental design of pig *L. dorsi* 2-DE.

Table 1 Results of the densitometry analysis.

| № Spot | Name of gel | A (vol ± SD) | B (vol ± SD) | C (vol ± SD) | D (vol ± SD) |
|--------|-------------|--------------|--------------|--------------|--------------|
| 1      |             | 8.19 ±1.1622 × 10^7 | 5.75 ±0.90 × 10^7 | 7.47 ±0.06 × 10^7 | 7.34 ±0.10 × 10^7 |
| 2      |             | 10.18 ±0.2622 × 10^7 | 10.70 ±0.14 × 10^7 | 11.48 ±0.26 × 10^7 | 12.00 ±0.34 × 10^7 |
| 3      |             | 9.37 ±0.2022 × 10^7 | 8.97 ±0.44 × 10^7 | 8.58 ±0.53 × 10^7 | 9.64 ±0.63 × 10^7 |
| 4      |             | 4.39 ±0.2522 × 10^7 | 9.47 ±1.05 × 10^7 | 7.51 ±0.15 × 10^7 | 7.20 ±0.23 × 10^7 |
| 5      |             | 1.92 ±0.39 × 10^7 | 2.71 ±0.72 × 10^7 | 2.95 ±0.54 × 10^7 | 4.03 ±0.66 × 10^7 |
| 6      |             | 10.65 ±0.62 × 10^7 | 9.42 ±0.84 × 10^7 | 14.42 ±0.60 × 10^7 | 13.23 ±0.98 × 10^7 |
| 7      |             | 8.19 ±0.41 × 10^7 | 5.75 ±1.22 × 10^7 | 7.47 ±0.07 × 10^7 | 7.34 ±0.64 × 10^7 |
| 8*     |             | 0.64 ±0.16 × 10^7 | 1.04 ±0.88 × 10^7 | 13.30 ±0.50 × 10^7 | 11.82 ±0.97 × 10^7 |
| 9*     |             | 0.34 ±0.05 × 10^7 | 0.42 ±0.14 × 10^7 | 6.35 ±0.60 × 10^7 | 6.23 ±0.74 × 10^7 |
| 10*    |             | 2.42 ±0.61 × 10^7 | 1.26 ±0.58 × 10^7 | 5.84 ±0.64 × 10^7 | 4.63 ±0.43 × 10^7 |
| 11*    |             | 1.24 ±0.02 × 10^7 | 1.57 ±0.27 × 10^7 | 10.68 ±0.74 × 10^7 | 9.68 ±0.85 × 10^7 |
| 12*    |             | 0.40 ±0.25 × 10^7 | 0.89 ±0.31 × 10^7 | 13.37 ±1.20 × 10^7 | 10.78 ±0.92 × 10^7 |
| 13*    |             | 0.29 ±0.08 × 10^7 | 0.45 ±0.11 × 10^7 | 4.91 ±0.27 × 10^7 | 5.45 ±0.59 × 10^7 |
| 14*    |             | 0.24 ±0.04 × 10^7 | 0.13 ±0.02 × 10^7 | 3.16 ±0.54 × 10^7 | 3.19 ±0.71 × 10^7 |
| 15*    |             | 4.23 ±0.25 × 10^7 | 0.24 ±0.08 × 10^7 | 5.61 ±0.44 × 10^7 | 2.47 ±0.37 × 10^7 |

Note: Spot Vol** were normalized by total valid spot volume and mean of value from duplicate analytical gels from three replicates. Data represented are means ±SD of three independent experiments. *Significant differences were found between IEF variations in the distribution of values between gels within the same sample with *p* <0.05. **Vol: The volume of a spot is the sum of the background-subtracted gray values of all pixels delimited by the spot border. By default, the background is defined as the minimum gray value on the spot border.
In the classical version of IEF (O’Farrell, 1975; Hirano, 1982; Kimura et al., 2003), when 0.02 M NaOH is in the upper chamber of the electrophoretic cell (Figure 2A), good protein separation is observed within a pI range of 5 to 6.5, such as tropomyosin beta chain (Mw 33.5 kDa, pI 4.80) (D’Alessandro et al., 2011; Peng et al., 2013), tropomyosin alpha-3 chain (Mw 33.5 kDa, pI 4.71) (Davoli et al., 2000), myosin light chain 3-like (Mw 22.0 kDa, pI 5.24) and myosin light chain 1/31 (Mw 21.0 kDa, pI 5.80) (Montowska and Pospiech, 2012; Kovaleva et al., 2013).

Moreover, on two-dimensional electropherograms with incubation in buffer A (Figure 2B), these proteins are most clearly identified (Ros et al., 2002; Montowska and Pospiech, 2007; Paredi, Mori and Mozzarelli, 2018). At the same time, in the alkaline zone of the gel, a blurred image of protein fractions is observed. A completely different gel was obtained when the current direction changed during the IEF, when 0.01 M orthophosphoric acid was in the upper chamber of the electrophoretic cell. Clear and well-defined protein spots were obtained with uniform distribution over the entire gel area (Chernukha et al., 2017). Presumably, this was because proteins with acidic and neutral pI are more highly represented in muscle tissue, resulting in a higher proportion of positively charged proteins. Since the current always goes in the direction from “plus” to “minus”, when changing the direction of the current flow, it is easier for positively charged proteins, which are always layered on top to move to the lower chamber of the electrophoretic cell, where a negative charge (cathode) is created (Colangeli et al., 2018).

Figure 2 2-DE of pig L. dorsi. Note: A – 0.02 М NaOH were from above, 0.01 М H₃PO₄ from below, tube gels were incubated in equilibration buffer I and II; B – 0.02 М NaOH were from above, 0.01 М H₃PO₄ from below, tube gels were incubated in buffer A; C – 0.01 М H₃PO₄ were from above, 0.02 М NaOH were from below, tube gels were incubated in equilibration buffer I and II; D – 0.01 М H₃PO₄ were from above, 0.02 М NaOH were from below, tube gels were incubated in buffer A.
Interestingly, during incubation in equilibration buffer I and II (Figure 2C), protein spots with a molecular weight greater than 50 kDa were better visualized (Vasilevskaia and Akhremko, 2019). In contrast, when incubated in buffer A (Figure 2D), protein spots with a molecular mass of less than 40 kDa were better visualized. The latter variation was used in a study by Kovalev (Kovalyov et al., 2006; Zvereva et al., 2015), and analysis of the results found that this method, as in the case of incubation in equilibration buffers, allows the best detection of muscle tissue proteins.

Major structural proteins of pig muscle tissue (Figure 3) were found (Montowska and Pospiech, 2012) and subjected to densitometric analysis (Table 1). The Fold change index (Persike et al., 2018) was calculated. Fold change index is the ratio between the volume of protein spots with the highest average value and the lowest average value. Eight fractions were detected (≥2-fold change, p < 0.05) with increased spot volume by at least 2 times compared with the classical version of the IEF (marked * in Table 1).

Thus, the variation of the IEF, when sodium hydroxide is in the upper chamber of the electrophoresis cell, has a lower resolution and does not properly detect protein fractions in the alkaline zone. So, in Figures 2A and 2B, proteins in the right half of the gel are noticeably out of focus. It is necessary to increase the number of volt-hours by at least 40% (up to 7 – 8 hours) for better protein separation in this area.

After changing the direction of electric current during the IEF, the process takes 4 hours. As a result, at least two times more protein spots are detected on 2-DE, including major structural muscle proteins, such as glyceralddehyde-3-phosphate dehydrogenase (Han et al., 2019), troponins group (Mora et al., 2016; Drousiotis et al., 2020), 3-hydroxyacyl-Coenzyme A dehydrogenase, beta-enolase and others (Nolan et al., 2019).

CONCLUSION

The results of electrophoretic studies showed that the best option for the separation of meat proteins is to change the direction of the current. Thus, when the anode buffer (0.01 M orthophosphoric acid) is in the upper chamber of the electrophoretic cell, and the cathode buffer (0.02 M sodium hydroxide) is in the lower one, the most informative picture is obtained. Incubation in both buffer A and equilibration buffers can also be used.

Densitometric analysis showed that the use of new parameters allows us to identify a larger number of proteins (almost 2 times). An increase in the color intensity of certain fractions is also noted. Thus, the proposed variation of the IEF can be used as the main one for muscle proteins electrophoretic analysis, since it requires less time and has a higher resolution.

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Acknowledgments:
This work was supported by a grant from the Russian Science Foundation No. 19-76-10034.

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