2DGE-coomassie brilliant blue staining used to differentiate pasteurized milk from reconstituted milk

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ABSTRACT

Differentiating pasteurized milk and reconstituted milk by scientific approach was necessary to defend consumer from economic fraud of wrong labeling. In this paper 2DGE (2 Dimensional Gel Electrophoresis)-coomassie brilliant blue staining method was employed and significant color intensity changing was observed among raw milk, pasteurized milk, UHT milk and reconstituted milk. For example, the intensity of 10 protein spots including casein and lactoglobulin reduced more than two folds from pasteurized milk to reconstituted milk. However, DIGE (Differential Gel Electrophoresis) assay showed that the majority protein remained similar level from pasteurized milk to reconstituted milk. Therefore the color fading of coomassie brilliant blue stained 2D gels may be due to other biochemical reaction, such as Maillard reaction, instead of protein degradation. Stability of 2DGE pattern was confirmed by running six gels of the same sample in parallel and software analysis showed that all proteins were at similar level. Two commercialized pasteurized milk samples and one reconstituted milk sample were tested by 2DGE-coomassie blue staining method and reconstituted milk could be easily identified.

Keywords: 2DGE; Coomassie Brilliant Blue; Pasteurized Milk; Reconstituted Milk

1. INTRODUCTION

China has recently become one of the ten biggest milk producers in the world [1]. Fluid milk products include pasteurized milk and reconstituted milk. Chinese product standard of pasteurized milk (GB5408.1) commanded that only fresh raw milk could be used as raw material for pasteurized milk, while reconstituted milk made by resolving milk powder in water and sterilization was categorized as sterilized milk (GB5408.2). Compared to pasteurized milk, reconstituted milk underwent more complicated thermal process including spray drying, pasteurization, UHT-treatment or in-bottle sterilization [2]. Because of higher cost at factory location, seasonal variation and transportation, pasteurized milk claims higher price than reconstituted milk. It is reported that the price for 8 tons of raw milk in China is about 20,000 YUAN, while the price for 1 ton of imported milk powder is 14,000-15,000 YUAN which could be made into 8 tons of reconstituted milk [3]. However, as a lot of literature pointed out, intensive thermal treatment would compromise milk nutrition and flavor [4], thus consumers prefer pasteurized milk to reconstituted milk and were concerned at possible economic fraud by labeling reconstituted milk as pasteurized milk.

A few analysis techniques such as CE (Capillary Electrophoresis), HPLC (High performance liquid chromatography), ELSD (E evaporative Light-scattering Detector) have been applied in differentiating pasteurized milk and reconstituted milk [2,5]. In these methods, individual protein or sugar ingredient, for example furosine, lactoglobulin, HMF (hydroxymethylfurfural) is quantified, which demands complicated pre- procession of milk sample. Results of above-mentioned studies revealed significant change of protein component during the procession of milk. In this paper, we reported the application of 2DGE (2 Dimension Gel Electrophoresis) technique in an overall analysis of protein profile change related to milk thermal procession, revealing a significant alteration of protein component between pasteurized milk and reconstituted milk. Compared to other methods, 2DGE is characterized by simplicity in sample preparation, ability of parallel treatment of several samples and being information-rich. In recent years, 2DGE have been widely applied for food analysis [6-8]. A number of research work have been done in milk proteome such as Equidae milk [9], marsupial Trichosurus vulpecula milk [10], early lactation milk of the tammar wallaby [11], κ-casein micro-heterogeneity in bovine milk [12] and whey protein [13]. In our proteomic
study of milk product, it showed that 2D patterns after coomassie brilliant blue staining could differentiate pasteurized milk and reconstituted milk according to the change of color intensity of some protein spots.

2. MATERIAL AND METHODS

2.1. Material

Pooled raw milk sample was collected from Sanyuan Dairy Company and immediately sent to milk processing laboratory in Food Institute of China Agricultural University for heat processing. Dry milk powder was also collected from the company and reconstituted in accordance with the original milk/water ratio, then pasteurized. After preparation, total protein concentration of each sample was determined. Raw milk was centrifuged at 1100g, 20mins and fat cream was removed. Three commercialized milk samples were bought from local supermarket including two pasteurized milk samples from different supplier and one reconstituted milk sample. All samples were stored at 4°C for immediate use or at -80°C.

2.2. Total Protein Concentration Determination

Total protein concentration was determined using Protein Assay Kit (NoVagen, Merk, Darmstadt, Germany) following the instruction. The optical absorbance value was recorded on ELISA reader (Thermo Fisher Scientific, MA, USA). Protein concentration was calculated on the basis of Absorbance-Concentration curve of reference BSA standards.

2.3. 2DGE

In preparation for IEF running, 10μL milk was mixed with 440μL of solubilization buffer consisting of 8 M urea, 400mg/L CHAPS, 40 mM Tris, 50mg/L pH 4.7–5.9 carrier ampholytes (Bio-rad, Hercules, California, USA) and 100 mM DTT. The sample was used to hydrate a 17cm pH4.7–5.9 IPG strip for 12 h at room temperature. Hydrated IPG strips were focused in a PROTEAN IEF System (Bio-rad, Hercules, California, USA) at 100 V for 1 h followed by 500 V for 1 h and 1 kV for 1 h before the voltage was increased to 8 kV for a total of 100 kVh. In the second dimensional SDS-PAGE assay, focused strips were first balanced in equilibrium buffer I and buffer II, then embedded with 0.5% agarose on top of 14% polyacrylamide gels (18×18 cm). Electrophoresis was performed in PROTEAN II XL Cell (Bio-rad, Hercules, California, USA) at 100 mA/gel for 2 h followed by 20 mA/gel for 16 h. Gels were stained with Coomassie Brilliant Blue G-250 and destained in 1% acetic acid. Images were captured on Versadoc Imager (Bio-rad, Hercules, California, USA) in transmission mode.

2.4. DIGE

Milk samples were labeled with Cy dye (CyDye DIGE Fluors, GE Healthcare, Buckinghamshire, UK) according to the instruction. Sample pooling strategy was modified as 5μL milk labeled with 1μL cy working solution (400pmol/μL). All of the 10μL labeled sample comprised with 5μL pasteurized milk and 5μL reconstituted milk was pooled together. 2DGE was run following above-mentioned procedure.

2.5. Data Analysis

2DGE profiles caught by Versadoc imager were analyzed with PDQuest software 7.4.0 (Bio-rad, Hercules, California, USA). After automatic spot detection, spot view was performed to display quantity of protein spots. For DIGE imaging, specific cy channel was selected.

2.6. Spot Digestion

In-gel digest was conducted following procedure of literature (Holland et al. 2004). Digestion product was purified using Ziptip C18 pipette tip (Millipore, Danvers, MA, USA) following instruction. In the final elution step, peptide was dissolved in 10mg/mL a-cyano-4-hydroxycinnamic acid in 0.1%TFA/50%ACN and directly applied to MALDI-TOF analysis.

2.7. MALDI-TOF and Database Search

One microliter purified peptide solution was spotted onto a stainless steel MALDI target. Spectra were acquired using a 4700 MALDI-TOF mass spectrometer (Applied Biosystems, Foster City, CA, USA) in delayed extraction mode. Tryptic digests were analyzed in positive ion reflectron mode with an accelerating voltage of 20 kV, grid voltage at 64% and a delay time of 165 ns. One hundred laser shots were accumulated for each spectrum. Peptide mass fingerprint (PMF) of cut protein spot was analyzed by MS-fit program of the ProteinProspector software (University of California, USA). SwissProt.20071010 database was searched and searching parameter was set as follow: Bos Taurus species, Tol 1 Da, Min matched peptide set as 6.

3. RESULTS AND DISCUSSION

Raw milk sample was taken from pooled milk container to minimize heterogeneity of protein composition. Milk was processed in lab to ensure the authenticity of processing condition. Different PI ranges were tried to determine the best 2DGE condition (2DGE of methods section). As shown in Figure 1, pH4.9-5.7 IPG strip produced the most satisfied 2D pattern in terms of protein spot quantity and separating size among these spots.

Total protein concentration of raw milk, pasteurized milk, UHT milk and reconstituted milk was calculated by Biuret method as shown in Table 1. After comparison of the 2D-coomassie brilliant blue staining patterns of the four milk samples shown in Figure 2, we found that for the majority of protein spots, color intensity decreased significantly from raw milk, pasteurized milk, UHT milk to reconstituted milk.
SDS-PAGE was run in 7cm×7cm gels.

Figure 1. 2DGE profiles at different IPG range.

Ten microliter milk of each sample the protein concentration of which has been modulated at the same level was loaded.

Figure 2. 2DGE profiles of raw milk, pasteurized milk, UHT milk and reconstituted milk.

Those spots were begot by PDQuest software analysis. The analysis set manager was defined as pasteurized milk two times higher in quantity than reconstituted milk.

Figure 3. Quantitative comparison of ten protein spots among raw milk, pasteurized milk, UHT milk and reconstituted milk.

reconstituted milk. Quantitative analysis by PDQuest confirmed the trend. As shown in Figure 3, after the analysis set template was defined as pasteurized milk being two fold above reconstituted milk, ten spots were detected. The quantity of the protein spots as represented by the Y axis decreased when milk was heat treated and the difference between pasteurized milk and reconstituted milk was remarkable.

In order to prove that under standard operation procedure 2DGE pattern was characterized by good repeatability, six 2D gels of pasteurized milk were run in parallel. Raw maps of coomassie blue staining were shown in Figure 4. All gels presented similar pattern and PDQuest analysis showed similar quantity level for all protein spots.

Protein spots were extracted, digested and identified by MALDI-TOF. As shown in Figure 5, 16 spots were successfully identified as casein and its isomers, lactoglobulin and lactate dehydrogenase-like protein.

The above experiments demonstrated that after coomassie blue staining, color intensity of most proteins decreased when milk sample was processed under thermo condition and the difference between pasteurized milk and reconstituted milk was significant enough to be used in product identification. As for the reason behind the changing trend, Maillard reaction should be considered other than proteins degradation since Maillard was the most significant biochemical process during heat treatment of milk [4,14]. In Maillard reaction, the reducing sugar covalently binds to the epsilon amide residue of lysine. We inferred that the intensity decrease of protein spots was related to coomassie brilliant blue staining. DIGE (2D Difference Gel Electrophoresis) was then conducted to understand the true situation of protein

<table>
<thead>
<tr>
<th>Assay</th>
<th>Raw milk</th>
<th>Pasteurized milk</th>
<th>UHT milk</th>
<th>Reconstituted milk</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>0.227</td>
<td>0.237</td>
<td>0.233</td>
<td>0.228</td>
</tr>
<tr>
<td>A2</td>
<td>0.224</td>
<td>0.235</td>
<td>0.227</td>
<td>0.231</td>
</tr>
<tr>
<td>A3</td>
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<td>0.224</td>
<td>0.24</td>
<td>0.225</td>
</tr>
<tr>
<td>A</td>
<td>0.227</td>
<td>0.232</td>
<td>0.233</td>
<td>0.228</td>
</tr>
<tr>
<td>C(mg/ml)</td>
<td>23.5</td>
<td>23.8</td>
<td>23.8</td>
<td>23.6</td>
</tr>
</tbody>
</table>
2DGE profile was shown as raw map. Spot quantity was shown in the small box below the map.

**Figure 4.** Six 2DGE profiles of pasteurized milk run in parallel to confirm the stability of the method.

<table>
<thead>
<tr>
<th>SSP ID</th>
<th>Protein function</th>
</tr>
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<tbody>
<tr>
<td>3701</td>
<td>Beta-casein</td>
</tr>
<tr>
<td>5901</td>
<td>Alpha-S2-casein</td>
</tr>
<tr>
<td>4901</td>
<td>Alpha-S2-casein</td>
</tr>
<tr>
<td>7901</td>
<td>Alpha-S2-casein</td>
</tr>
<tr>
<td>8901</td>
<td>Alpha-S2-casein</td>
</tr>
<tr>
<td>5701</td>
<td>L-lactate dehydrogenase A-like 6B</td>
</tr>
<tr>
<td>9401</td>
<td>Kappa-casein</td>
</tr>
<tr>
<td>3302</td>
<td>Beta-lactoglobulin</td>
</tr>
<tr>
<td>4101</td>
<td>Beta-lactoglobulin</td>
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<tr>
<td>1701</td>
<td>L-lactate dehydrogenase A-like 6B</td>
</tr>
<tr>
<td>4701</td>
<td>L-lactate dehydrogenase A-like 6B</td>
</tr>
<tr>
<td>6901</td>
<td>Alpha-S2-casein</td>
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</tr>
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<td>7601</td>
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</tr>
<tr>
<td>3903</td>
<td>Alpha-S2-casein</td>
</tr>
</tbody>
</table>

The left figure was peptide mass fingerprint of one protein after in-gel digestion. The right table showed identification results of several proteins after database search.

**Figure 5.** Identification of milk proteins by MALDI-TOF.

quantity changing between pasteurized milk and reconstituted milk. As shown in **Figure 6**, after pre-stained with cy3 and cy5 respectively, pasteurized milk and reconstituted milk were mixed together and run in the same 2D gel. Images of cy3 and cy5 showed that the majority of protein spots had similar level of fluorescence intensity.
Profiles were shown as Gaussian map. Spot quantity was shown in the small box below the map. In each box, left column corresponded to reconstituted milk and right column to pasteurized milk.

**Figure 6.** DIGE profiles of pasteurized milk and reconstituted milk.

These three samples were bought from local supermarket.

**Figure 7.** 2DGE-coomassie brilliant blue staining profiles of three commercialized milk samples.
PDQuest analysis indicated that the quantity difference between pasteurized milk and reconstituted milk was less than two folds. Thus we suppose the remarkable diverge between cydye labeling and coomassie blue staining was due to different staining mechanism. Cydye DIGE fluor were designed to covalently attach to the epsilon amino group of lysine of proteins in a “minimal labeling” way which means the dyes labeled only on a single lysine per protein molecule. Therefore Maillard reaction would not affect staining efficiency. As for coomassie blue, the mechanism of staining is still not well understood since it was used in protein staining 45 years ago. However some literatures proved that coomassie blue varied widely in its ability to bind proteins due to its affinity for protein rich in basic amino acids such as lysine, arginine and histidine and its poor ability to bind with glycoprotein [15]. Moreover, since coomassie blue staining was not a “minimal labeling” technique, its labeling efficiency would be affected significantly by Maillard reaction. In fact the content of available lysine has often been used as marker of heat damage affecting dairy protein [16].

In order to verify the practicality of the method, three commercialized milk samples plus the reference pasteurized milk sample were tested in parallel. As shown in Figure 7, two pasteurized milk samples resulted in similar 2D pattern as reference sample. The reconstituted milk sample showed remarkably different 2D pattern. The seriously reduced level of protein staining indicated that the reconstituted milk sample might undergo stronger thermo treatment than pasteurization.

Therefore, 2DGECoomassie brilliant blue staining technique was proved to be very useful in differentiation between pasteurized milk and reconstituted milk in that color intensity of most proteins decreased significantly after heat treatment. DIGE experiment showed that true quantity of these proteins did not change much. Change of color intensity of different samples may be due to the decrease of labeling efficiency of coomassie blue caused by Maillard reaction after heat treatment. Therefore we could apply 2DGECoomassie brilliant blue staining method for identification of pasteurized milk and reconstituted milk. Moreover 2DGEC could be used to detect adulterated milk product according to different 2D pattern of protein from different organism or tissue.

4. ACKNOWLEDGEMENT

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