An Analysis of the proteomics approach to the glycated peptides of human milk

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Abstract: Many studies have shown that advanced glycation end-products (AGEs) and glycation adducts are significantly linked to aging and disease. Particularly, the level of glycation in human milk is important because the AGE intake is closely related to AGE levels in infants. In this study, we used human milk samples obtained from four primiparae and four multiparae. We isolated proteins using acetone and trichloroacetic acid (TCA) precipitation. A total of 67 glycated proteins and 122 glycated peptides was quantified; among them, 19 glycated peptides were differentially expressed. We confirmed that the degree of glycation differed according to fertility. The study provides a foundation for using proteomics to evaluate the mother’s milk quality and link between maternal health and human milk quality.

Key words: glycation, advanced glycation end-product, human milk, LC-MS/MS, fertility

1. Introduction

Milk is important for the health of infants because it contains nutrients, immune-related components, anti-infective factors, and metabolic enzymes, which greatly impact growth and development. The composition of human milk is dynamic and depends on various environmental factors such as breastfeeding and the mother’s diet.

The protein concentration in human milk is 8.5 g/L, which is lower than the 34.5 g/L found in bovine milk. The protein ratio of whey: casein found in bovine milk is between 50:50 and 80:20. Compared to bovine milk, the percentage of whey protein and ratio of carbohydrate in human milk are high. Thus, human milk is a better environment for glycation compared to bovine milk.

Glycation is a spontaneous post-translational modification that occurs at lysine, arginine, and the N-terminus of proteins. The active carbonyl group of the sugar and nucleophilic free amino group chemically react to form an unstable Schiff base. The Schiff base is rearranged to form Amadori compounds that form advanced glycation end-products (AGEs) through several reactions.

Levels of serum AGEs between the mother and
infant are strongly correlated. In addition, increasing the intake of dietary AGEs also increases the level of serum AGEs in infants. In many studies, it has been confirmed that toxic substances such as dichlorodi-phenyltrichloroethane, perfluorinated compounds, and polychlorinated biphenyls of human milk come from multiparae more than primiparae.

In a previous study, the automated boronate affinity enrichment method has been applied to bovine and human serum albumin using a liquid handling robotic system. In this study, we removed N-glycan that interfered with glycated peptide enrichment by performing PNGase F digestion during the filter-aided sample preparation (FASP) method. Glycated peptides from human milk were enriched and analyzed using liquid chromatography-tandem mass spectrometry (LC-MS/MS)-based label-free quantification. We analyzed glycated peptides in human milk and their relationship with fertility.

2. Experimental

2.1. Chemicals and reagents
Iodoacetamide, ammonium bicarbonate (ABC), MgCl₂, urea, Tris(hydroxymethyl)aminomethane (Tris), trichloroacetic acid (TCA), and ammonium acetate (NH₄OAc) were purchased from Sigma-Aldrich (St. Louis, MO, USA). Tris (2-carboxyethyl) phosphine (TCEP) was obtained from Thermo Fisher Scientific (Waltham, MA, USA). Sequencing-grade modified trypsin was acquired from Promega (Madison, WI, USA). Acetic acid (HOAc), methanol, acetonitrile, and high-performance liquid chromatography-grade water were purchased from J.T. Baker (Phillipsburg, NJ, USA). Formic acid was from Fluka (Charlotte, NC, USA). PNGase F was purchased from New England Biolabs (Ipswich, MA, USA).

2.2. Instrumental conditions
Experiments were conducted on a nano-LC-MS/MS system consisting of a Q Exactive™ Hybrid Quadrupole-Orbitrap MS (Thermo Fisher Scientific) coupled with a Dionex Ultimate 3000 HPLC (Sunnyvale, CA, USA). The solvent A as the mobile phase was composed of 0.1 % formic acid in water and solvent B as the mobile phase was composed of 0.1 % formic acid in 80 % acetonitrile. The glycated peptides were loaded on an Acclaim™ PepMap™ 100 C18 nano-trap column (75 μm × 2 cm, 3 μm particles, 100 °C pore size) by carrying out using a mobile phase at a flow rate of 3 μL/min for 10 min. The peptides were separated on a PepMap™ RSLC C18 nano-column (2 μm, 100 Å, 75 μm × 50 cm) at a flow rate of 300 nL/min. Data-dependent acquisition was performed, and the top ten precursor peaks were fragmented with higher energy collisional dissociation and normalized collisional energy was 27. Ions were scanned at 70,000 in MS1 (the first level of mass analysis) and 17,500 in MS2 (the second level of mass analysis) over an MS scan range of 400-2000 m/z for both the MS1 and MS2 levels. The injection quantity was 300 ng. A NanoDrop™ 2000 Spectrophotometer (Thermo Fisher Scientific) was used.

2.3. Whey protein extraction
Human milk samples and clinical information were obtained with informed consent, and procedures were performed with the approval of the Institutional Review Board (IRB 7001355-202108-E-153). Information on the clinical samples is summarized in Table 1. The overall protocol is shown in Fig. 1. Human milk samples (3 mL) were treated with 3 mL of acetone, sonicated for 30 min at room temperature, and centrifuged at 3,600 × g for 10 min. The protein pellet was separated from human milk and subjected to TCA precipitation. To remove casein protein, the protein pellet was treated with 60 mM CaCl₂, incubated for 1 h at 4 °C and centrifuged at 3,600 × g for 30 min. The organic layer was removed, and the sample was centrifuged again. The bottom layer of the sample was extracted and filtered through a 0.45-µm mixed cellulose ester syringe filter from Advantec (Tokyo, Japan). The filtered sample was treated with a three-fold volume of cold acetone/10 % TCA (v/w) and incubated for 2 h at -20 °C. The sample was washed twice with cold acetone and dried using a HyperVAC-MAX VC2200 centrifugal vacuum concentrator (Hanil Scientific, Inc., Gochon-eup, South Korea).
Human milk whey protein was digested using FASP on a Microcon YM-30 filter from Millipore (Billerica, MA, USA). Each 200 µg of human milk whey protein was adjusted to a volume of 100 µL lysis buffer (8 M urea, 0.1 M Tris and HCl buffer, pH 8.5). The protein was reduced by incubation with TCEP at 37 °C for 30 min and alkylated with iodoacetamide at 25 °C for 1 h in the dark. After washing with lysis buffer and 50 mM ABC sequentially, the protein was treated with PNGase F to cleave N-linked-glycan chains from the protein and incubated at 37 °C for 3 h. After washing with 50 mM ABC, the protein was digested with trypsin (enzyme to protein ratio of 1:50; w/w) at 37 °C for 18 h. The digested protein was collected into new tubes, and trypsin was inactivated by acidification with 20 µL formic acid. Collected peptides were dried using a centrifugal vacuum concentrator.

2.5. Boronate affinity enrichment
Glycated peptides were enriched using Cellufine PB (500 µL, JNC Corporation, Tokyo, Japan), a phenyl borate affinity ligand. Loading buffer (50 mM MgCl₂, 250 mM NH₄OAc, 0.1 M Tris, pH 8.1), washing buffer (50 mM ABC and 0.1 M Tris, pH 8.1), and elution buffer (0.1 M HOAc, pH 3) were prepared in high-performance liquid chromatography grade water. Cellufine PB was placed on the Microcon YM-30 filter and washed five times with 400 µL loading buffer for equilibration. The peptide sample was dissolved in 250 µL loading buffer and loaded into YM-30 filter with Cellufine PB. The filter was washed thrice with 400 µL washing buffer. After washing, the retained glycated peptide was eluted twice using a 200 µL elution buffer. Eluted glycated peptides were desalted with ultra-micro spin C18 columns (Harvard Apparatus, Holliston, MA, USA). Desalted glycated peptides were dried using a centrifugal vacuum concentrator.
2.6. Data analysis and statistical interpretation

Thermo MS/MS raw files of analysis were searched using Proteome Discoverer™ software (ver. 2.4), and the Human database was downloaded from Uniprot. The appropriate consensus workflow included a peptide-spectrum match validation step and SEQUEST HT process for detection as a database search algorithm. Search parameters were set up as follows: 10 ppm of tolerances of precursor ion masses, 0.02 Da fragment ion mass, and maximum of five missed cleavages with trypsin enzyme. The modifications on the peptide sequence were as follows: static carbamidomethylation of cysteine (+57.012 Da), dynamic modifications of methionine oxidation (+15.995 Da), carbamylation of protein in N-terminus (+43.006 Da), and acetylation of protein in N-terminus (+42.011 Da). Glycation of lysine (+162.053 Da) for analyzing glycated proteins and deamidation of asparagine and glutamine (+0.984 Da) for evaluating the effect of PNGase F were added as the dynamic modifications in the Proteome Discoverer™ software search. The results below 1% of the false discovery rate were selected and filtered for at least more 6 amino acids in a peptide length.

3. Results and Discussion

In this study, human milk samples obtained from eight mothers were collected and whey protein extraction and glycated peptide enrichment were performed. Whey protein was extracted using acetone precipitation and TCA precipitation and digested using FASP. PNGase F was used to remove N-glycan, and glycated peptides were enriched using boronate beads.

3.1. Qualitative analysis of human milk glycation enrichment

The numbers of glycated peptides and proteins identified in the human milk sample from primiparae were 102 and 52, respectively. The numbers of glycated peptides and proteins identified in the human milk sample from multiparae were 80 and 41, respectively. As shown in Fig. 2, average numbers of glycated peptides and proteins were greater in human milk samples from primiparae compared to those in multiparae.

3.2. Quantitative analysis of human milk peptides

We confirmed through quantitative analysis that the peptide level in human milk differs according to fertility. Fig. 3(a) shows the volcano plot obtained in the multiparae and primiparae groups, with 485 differentially expressed peptides (251 up and 234 down). Volcano plot was made of the ratio of normalized abundance between groups and P-value. The peptides identified and quantified according to fertility were visualized in a principal component analysis (PCA) plot in Fig. 3(b). PCA is based on signal intensity to compare samples. As shown in Fig. 3(c), a heatmap was generated for 3104 quantified peptides to compare...
the differences in the normalized abundance of peptides between groups.

3.3. Qualitative analysis of human milk glycation enrichment

As shown in Fig. 4, we quantitatively analyzed human milk according to fertility and found that 19 of 94 glycated peptides were differentially expressed. The 100% stacked bar chart shows the average values of the 19 differentially expressed peptides (DEPs) in the group divided by fertility. A list of the 19 glycated DEPs with primiparae and multiparae is shown in Table 2. The 14 glycated DEPs down-regulated and 5 glycated DEPs up-regulated in multiparae. The 19 glycated DEPs belong to 10 albumin proteins. glycated albumin was studied as potential factor for glycemic control.13 So, we confirmed that glycemic control was better in human milk of multiparae. The degree of glycation in patients with diabetes is higher than that in normal subjects, as reported in several previous studies.12,14 As the nutritional levels derived the human milk are related to the maternal health, the glycation degree is a potential factor that determining the quality of human milk.15 Studies have shown that the intake of AGEs increases the level of AGEs in the body.16 Based on these results, primiparae should consume food that includes less AGEs to positively impact the baby’s nutrition.
Fig. 4. The 100 % stacked bar chart of the 19 differently expressed glycated peptides was calculated as the average abundance in each group.

Table 2. List of 19 significantly different glycated peptides from human milk between multiparae and primiparae group

<table>
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<tr>
<th>Uniport accession ID</th>
<th>protein name</th>
<th>Annotated Sequence</th>
<th>Glycation sites</th>
<th># PSMs</th>
<th>Theo. MH+ [Da]</th>
<th># Missed Cleavages</th>
<th>Abundance Ratio Primiparae/Multiparae</th>
<th>P-Value: Multiparae / Primiparae</th>
<th>Peptide level change Multiparae/Primiparae</th>
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<td>E7EQB2</td>
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<td>[K]SPKFQLEFGPSGQK[D]</td>
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4. Conclusions

We isolated whey proteins from human milk using acetone and TCA precipitation. We quantified 67 glycated proteins and 122 glycated peptides using a modified FASP with PNGase F. Nineteen glycated peptides were differentially expressed in human milk samples by classifying according to fertility. We confirmed difference between the glycated peptide level in human milk and fertility. We hope to analyze more human milk samples to study of glycation for validation. Other omics studies should also be conducted to identify relationship between human milk-derived components associated with fertility.

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References


Authors’ Positions

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Jun Hwan Song : Professor
Nam Mi Kang : Professor