Aged American Cheddar Cheese as Source of Protein Derived Compounds that Modulate Obese Mice Fecal Bacteria and Colon Inflammation In Vitro

Luis Condezo-Hoyos1,3, Giuliana D Noratto1,2*

1 School of Food Science, Washington State University
2 Current address: Department of Nutrition and Food Science, Texas A&M University
3 Current address: Faculty of Medicine, Universidad Autónoma de Madrid, Spain

Abstract

Intestinal bacteria are key component in obesity and obesity-induced inflammation that leads to chronic diseases. Lactobacillus helveticus, a highly autolytic and proteolytic strain, has been associated with dairy products with health benefits.

We investigated protein changes in American cheddar cheese manufactured with Lactobacillus helveticus (Cougar Gold brand, CG) during ripening, and their potential as modulators of fecal bacteria and colon Reactive Oxygen Species (ROS) in vitro.

Results confirmed that casein-derived products and proteins/peptides released after In Vitro Digestion (IVD) shifted DNA abundances of Enterobacteriaceae, Enterococcus, and Bifidobacterium sp. in obese mice feces to resemble lean mice. CG fecal fermentation metabolites protected human colon fibroblast CCD-18Co cells against production of ROS.

Overall, proteins in aged cheese might help to improve obese fecal bacteria balance and protect against obesity-induced colon inflammation. Further in vivo studies are needed to validate these findings and to confirm the intestinal beneficial value of ripened cheese.

Keywords: Cheese Proteins; Cheese Ripening, Obese Fecal Bacteria, Colon Inflammation

Abbreviations

CG: Cougar Gold brand; ROS: Reactive oxygen species; IVD: In vitro digestion; SLAB: Starter lactic acid bacteria; NSLAB: Non-starter lactic acid bacteria; MRPs: Maillard reaction products; AGEs: Advanced glycation end-products; PCR: Polymerase chain reaction; SCFAs: Short-chain fatty acids; RFU: Relative fluorescence units; PCA: Principal component analysis;

Introduction

Proteolysis of cheese caseins can be attributed to enzymes from milk and coagulant, as well as proteolytic activity of Starter Lactic Acid Bacteria (SLAB) and Non-Starter Lactic Acid Bacteria (NSLAB). In addition, cheese produced with Lactobacillus helveticus as adjunct culture has been reported to produce novel flavors, to improve acceptability of reduced-fat cheddar cheese, and to increase concentrations of low molecular weight peptides due to its high proteolytic activity and ability to autolyze [1]. The use of the L. helveticus strain as an adjunct to traditional starter cultures has been studied in cheese ripening. Results have shown that it produces a strongly flavored cheese within a relatively short period (2 mo). The main feature of the adjunct culture L helveticus is its ability to autolyze during cheese ripening which contributes to improved flavor [1]. This is evidenced by the release of intracellular enzymes such as d-lactate dehydrogenase and prolinase [2].

In addition, peptides produced by L. helveticus have been demonstrated to possess biological activity as angiotensin converting enzyme inhibitors in fermented dairy products. L. helveticus has genes responsible for key metabolic functions such as proteolysis, lipolysis, and cell lysis.

Thus, some health benefits of milk fermented with L. helveticus include the blood pressure lowering effect in a human clinical trial associated with the peptides-induced inhibition of Angiotensin Converting Enzyme (ACE), a molecule that promotes vasoconstriction. A study examined the effects of 8 mo ripened Gouda-type cheese produced using L. helveticus in addition to the starter culture bacteria. Results showed the suppression of abdominal adipose tissue accumulation in rats fed freeze dried cheese on a 20% fat diet compared with rats given butter oil and casein in isocaloric diet [3]. The cheese diet also led to changes in metabolic syndrome markers and inflammation such as reduction of serum cholesterol, triglycerides, very low density lipoprotein, low-density lipoprotein, and reduction on production of...
adiponectin from abdominal adipose tissue. However, the effects exerted by cheese consumption in preventing the development of obesity-related metabolic disorders might be attributed to the combination of SLAB and the peptides and fat components formed during the cheese ripening process.

Cheese ripening results in production of nucleophilic amino groups that react with carbonyl groups of reducing sugars and their degradation products through a series of non-enzymatic browning reactions called Maillard reactions. The series of reaction pathways that occur during cheese ripening are accompanied by decrease in reducing sugars, non-enzymatic browning, and the formation of early and advanced Maillard Reaction Products (MRPs) as described in detail [4].

The controversies around human health effects of MRPs have been extensively reviewed [5]. Harmful effects of dietary MRPs are related to their ability to promote glycation in vivo and to contribute to the progression of diabetes, cardiovascular complications and Alzheimer’s disease. The pathophysiological state in diabetes increase Advanced Glycation End-products (AGEs) load due to the impaired renal function and limited urine excretion. This indicates that, diabetic subjects are more sensitive to dietary MRPs, especially if they don’t keep blood glucose levels under control. This became evident when intake of severely heat-treated foods by healthy subjects did not increase endogenous levels of AGEs nor impaired renal function compared to the intake of a diet with fruit and vegetables [6]. On the other hand, the health benefits of MRPs are linked to their antioxidant, chemo preventive, and antiutagenic activities as reviewed in vitro and in vivo [5, 7]. Even though the effects of MRPs reported in the literature may not translate to humans, the antioxidant and chemo preventive activities of MRPs are associated to the induction of phase II detoxifying enzymes due to their ability to bind to an antioxidant responsive element. MRPs’ antiutagenic activity is related to their ability to bind to heterocyclic aromatic amines and other mutagens, thus preventing their digestion, absorption, and harmful physiological effects. In addition, glycated proteins are more resistant to gastrointestinal enzymes and can reach the lower gastrointestinal tract where they have the potential to modulate the physiology of the intestinal ecosystem [8, 9]. Less complex MRPs structures, which are preferentially produced in the presence of water and during shorter heating times, have shown to promote Bifidobacteria growth and the production of short chain fatty acids in a fermenter containing human fecal bacteria [5].

Our aim was to investigate how the ripening process influences the content of protein derived bioactive compounds in cheddar cheese produced with L. helveticus as adjunct culture and their potential to modulate gut bacteria communities and colon oxidative stress levels in vitro.

**Materials and Methods**

**Chemicals**

Acetone, glycerol, hydrochloric acid, methanol, sodium potassium tartrate and Tris base were purchased from Fisher (Pittsburgh, PA). Acetic acid, propanol and sodium hydroxide were acquired from JT Baker (Center Valley, PA). Bacterial growth media and supplements were obtained from BD (Franklin Lakes, NJ). The remaining chemicals and reagents were obtained from Sigma–Aldrich Co. Ltd. (Saint Louis, MO). All reagents were ACS grade, ≥ 99% purity. Primers for PCR were acquired from Integrated DNA Technology (San Diego, CA).

**Cheese manufacture**

The American type cheddar cheese Cougar Gold brand (CG) was manufactured in production scale at the Washington State University (WSU) creamery, according to the standard cheddar cheese making procedures as previously reported [10]. Briefly, the bacterial starter cultures *Lactococcus lactis ssp. lactis*, *Lactococcus lactis ssp. cremoris* were provided frozen by Chr. Hansen’s Laboratory Inc. (Milwaukee, WI). The Lactococcus cultures were grown in Delvolac LS-6 internally buffered starter medium (DSM Food Specialists B.V., Netherlands), and were inoculated at a concentration range of 0.31% to 0.47% (10⁹ colony forming units (cfu mL⁻¹) in milk. The Lactobacillus helveticus WSU19 adjunct culture maintained in liquid nitrogen at the WSU Creamery, was activated in sterile homogenized milk up to 10⁸ cfu mL⁻¹, and inoculated in a ratio of 45 mL/24.5 Kg of milk. Curd was pressed overnight (38 x 10⁴ Pa pressure on dial and 4.6 x 10⁴ Pa pressure on cheese), vacuum sealed in cans and maintained at 7°C for ripening for up to 24 mo. The samples were taken from batches produced under same conditions (starter culture, milk, and processing), and maintained under inventory by the WSU creamery, to keep ripening time as the only variable among samples. CG cheese samples on day 0 of ripening had 38.05% moisture, 1.55% salt, 35.15% fat, and pH 4.98 analyzed following official standards methods.

**L. helveticus counting**

For selective growth of mesophilic lactobacilli (*L. helveticus*) and inhibition of *Lactococcus lactis ssp. cremoris, and Lactococcus lactis ssp. Lactis* used as SLAB, cheese samples (10 g) were homogenized with 90 mL of peptone saline solution in stomacher Lab-Blender 400 Circulator (Seward Medical, London, UK) for approximately 3 min and analyzed by plating appropriate ten-fold dilutions onto MRS agar adjusted to pH 5.4 and incubated for 48 h at 37°C [11]. Results were expressed as *L. helveticus* cfu g⁻¹ of cheese.

**Reducing sugars**

Reducing sugars were extracted from cheese by homogenizing 5 g of cheese samples with 10 mL of phosphate buffer (10 mmol/L, pH 7.0) and centrifugation (5000 g for 15 minutes at 4°C) to remove fat. The supernatant was extracted with 4 volumes of chloroform:methanol (2:1 vol:vol) and centrifuged at 15000 g for 15 min (4°C) [12]. Reducing sugars were quantified in supernatant by the 3,5-dinitrosalicylic acid (DNS) assay [13]. Briefly, 30 µL volume of the glucose standards (0.5, 1.0, 1.5 and 2.0 mg/mL) or samples diluted in 0.1 mol/L sodium acetate buffer pH 5.5 were added to 60 µL 1% DNS reagent, heated at 95°C for 5 min, and cooled to 20 °C in a T100 Thermal cycler (Bio-Rad,
CA, USA). The reaction volume was transferred to 96-well plate and absorbance was measured at 540 nm in a plate reader (Biotek Instrument, Rochester, VT, USA). Reducing sugars levels were expressed as fold of 0 mo CG cheese.

**Extraction of caseins**

Caseins from CG cheese at different ripening stages were extracted as previously reported [4]. Briefly, 2.5 g of cheese was homogenized with 30 mL of acidified water with acetic acid (pH 5-6) followed by precipitation at pH 4.6 with 25% v/v acetic acid solution and centrifugation (4000 rpm, 10 min). The supernatant was discarded and the residue was homogenized with 20 mL of acidified water with acetic acid (pH 5-6) and 10 mL of methylene chloride, followed by centrifugation. Caseins located in the boundary phase were recovered and the process was repeated twice, washed with acetone, and were freeze dried. Extracted proteins were dissolved in Tris/2-Mercaptoethanol/Urea buffer pH 8.6 (2 mg/mL). Protein content was assessed by Bradford assay (Bio-Rad, CA, USA) using casein from bovine milk as standard (0.1-0.5 mg/mL). Electrophoresis and image analysis was performed with 20 µg protein diluted with Laemmli’s loading buffer (Bio-Rad, Hercules, CA), boiled, loaded on a 15% sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE) at 100 V for 10 min and at 200 V for 50 min in a Bio-RAD Mini-PROTEAN Tetra Cell (Bio-Rad). After separation, gels were soaked in prefixed solution containing methanol/acetic acid/water 50:10:40 v/v/v for 30 min and stained with 0.25% (w/v) Coomassie Brilliant Blue R250 at 55°C for 5 minutes. The gel was distilled at the same temperature with methanol/distilled water/acetic acid 25:65:10 (v/v/v) for 45 min and washed three times with de ionized water [14]. Scanned images were analyzed using Image J software to calculate band intensity. A linear equation of log molecular weight versus relative migration distance for protein standards (Precision plus Protein Western Standard, Bio-Rad) was used to estimate the molecular weight of proteins in samples.

**Protein carbonyls and AGEs-fluorophores in CG**

Extracted caseins were assayed for protein-bound carbonyls content by the 2,4-Dinitrophenylhydrazine (DNPH)- assay [15]. Briefly, freeze dried extracted caseins were dissolved in Tris/2-Mercaptoethanol/Urea buffer pH 8.6 (100 mmolL−1 Tris, 2 mmol L−1 2-mercaptoethanol, 8 mmol L−1 urea) to 10 mg mL−1 and reaction was carried out by mixing 250 µL of this solution with 250 µL of 10 mmol L−1 DNPH solution in 2.5 mol L−1 HCl for 15 min at room temperature in dark with constant vortexing. Protein carbonyls were precipitated with 10% v/v TCA on ice-bath for 30 min followed by centrifugation at 9000 g for 2 min (4°C). Pellets were washed three times with cold ethanol/ethyl acetate solution (1:1 v/v) to eliminate the excess of DNPH and dissolved in 6 mol L−1 guanidine-HCl. Absorbance was measured at 370 nm against a blank without DNPH. Protein-bound carbonyls were determined using extinction coefficient of DNPH (ε = 22000 mol L−1 cm−1) and expressed as nmol mg−1 protein quantified by Bradford assay (Bio-Rad, Hercules, CA).

AGE-fluorophores were assayed by measuring Relative Fluorescence Units (RFU) at 360 nm, excitation 460 nm emission of freeze dried extracted caseins dissolved in Tris/2-mercaptoethanol/urea buffer pH 8.6 (100 mg mL−1). A pQuant microplate reader (Biotek Instrument, Rochester, VT) was used. Results were calculated as RFU % of the 0 mo control [16].

**IVD of CG cheese**

CG cheese samples were successively incubated with digestive enzymes to simulate salivary, gastric, and duodenal digestion at 37 °C and agitation (200 rpm) as previously reported [17]. Briefly, cheese samples (10 g/200 mL) were homogenized with sodium maleate buffer (0.05 mol L−1, pH 6.0); an aliquot of this homogenate was subjected to salivary digestion simulation with 3.9 U of amylase mL−1 for 5 min. The gastric digestion was mimicked adjusting the pH to 1.5 with 6 mol L−1 HCl and incubating with 71.2 U of pepsin mL−1 for 90 min. Finally, the pH was adjusted to 7.5 using 6 mol L−1 NaOH and sample was incubated with pancreatin (9.2 mg mL−1) and bile extract (55.2 mg mL−1) for 150 min to simulate the duodenal digestion. IVD cheese samples were dialyzed overnight in a cellulose membrane with a cut off of 12.2 KDa (Sagmo-Aldrich, St. Louis, MO) against nanopure water at 4 °C, freeze dried, stored at -20°C for further use.

**In vitro fecal fermentation of CG cheese after IVD**

Fecal samples were collected from three obese one year old diet-induced obese mice (Body Mass Index (BMI) = 5.7 ± 0.7) and from three lean mice (BMI = 3.6 ± 0.4). Obese mice were under high fat (45% kcal from fat), diet (#D12451), containing 24% protein, 41% carbohydrate, 24% fat, plus minerals and amino acids (Research diets, Inc., New Brunswick, NJ). Lean mice were under 2018 standard diet (18% kcal from fat), containing 18% protein, 44.2% carbohydrate, 6.2% fat, plus minerals and amino acids (Teklad Diets, Madison WI). For the fecal culture experiments, a pre-culture was prepared with fresh feces diluted with anaerobic phosphate buffer 1:10 (w/v) (1 mol L−1, pH 7.2), homogenized to produce fecal slurries from each animal donor (no pooled fecal samples), and used for inoculation of pre-culture (10% v/v) in a pre-reduced sterile medium at pH 7.0 (peptone (2 g L−1), yeast extract (2 g L−1), NaCl (0.1 g L−1), K2HPO4 (0.04 g L−1), KH2PO4 (0.04 g L−1), NaHCO3 (2 g L−1), MgSO4.7H2O (0.01 g L−1), CaCl2.6H2O (0.01 g L−1), Tween 80 (2 mL L−1), hemin (50 mg L−1), vitamin K (10 µL L−1), L-cysteine (0.5 g L−1), bile salts (0.5 g L−1), resazurin (1 mg L−1), and distilled water) [18]. After incubation for 12 h at 37 °C under agitation (200 rpm) and anaerobiosis, culture was prepared with 2.5 % (v/v) of pre-culture in culture medium containing the freeze dried CG cheese (0, 6, or 12 mo ripening or casein, 0.2% w/v) after IVD in replacement of the yeast extract (obese control group). Feces from lean mice were cultured with casein after IVD (lean control group). After 24 h incubation under anaerobiosis generated by GasPak EZ® Gas Systems (BD Diagnostic Laboratory, NJ, USA) and constant agitation, fecal bacteria and supernatants were recovered by centrifugation and stored at -80 °C for further analysis.

**Analysis of relative abundances of DNA fecal bacterial**

DNA relative abundances of fecal bacteria were analyzed by PCR. Briefly, DNA was extracted from pellets recovered after
fecal fermentation using QIAamp DNA Stool Mini Kit (QIAGEN Inc, CA, USA) according to the manufacturer's protocol. The concentrations of DNA and absorbance ratio 260/280 nm as measurement of purity were assessed using a NanoDrop 2000 spectrophotometer (Thermo Scientific, NC, USA). PCR reaction mixtures (total of 10 μl) contained 5 μl of Sso Advanced™ SYBR® Green Supermix (Bio-Rad, CA, USA), 0.4 μl of primer mix (forward and reverse, final concentration 200 nM), 2 μl of adjusted (2 ng μl⁻¹) template DNA or non-template control, and 2.6 μl nuclease-free water (Qiagen GmbH, Hilden, Germany). PCR conditions were as follows: 95 °C for 30 s and 40 cycles at 95 °C for 5 s and 55 °C for 30 s. A melt curve analysis was performed to verify the specificity of the primers using the following conditions: 65 to 95 °C at 0.5 °C increments. Relative bacterial population was quantified by the Livak Method (2⁻ΔΔCt) using as reference the universal primer CT values. Data was normalized to cultures fermented with casein as substrate. Primer sequences obtained from [19] are presented in Table 1.

### Short-chain fatty acids (SCFAs)

Propionic, butyric, and acetic acids were quantified in supernatants from fecal fermentation as previously reported [20]. Briefly, supernatants filtered through a 0.22 μm PVDF filter (Thermo Scientific, TN, USA) were analyzed by High Performance Liquid Chromatography (HPLC) using the Aminex HPX-87H strong cation-exchange resin column (300 x 7.8 mm i.d.) and ion exclusion microguard refill cartridge (Bio-Rad, CA, USA) in a Waters 600S Separation Module equipped with an auto injector 717 Plus, a photodiode array detector (999 PAD) (Waters, MA, USA) and the Millennium software was used for HPLC analysis. A sample of 20 μl was eluted with H₂SO₄ (0.016 mol L⁻¹) at 0.6 mL min⁻¹ and 35 °C. SCFAs were identified and quantified by comparing retention times, UV-visible spectral data, and peak areas to known standards.

### Cell culture

The human colon fibroblast CCD-18Co cells were purchased from ATCC (Manassas, VA). Cells were cultured using high glucose Dulbecco’s Modified Eagle Medium, supplemented with 1% penicillin/streptomycin solution, 1% non-essential amino acids (10mM), 1% sodium pyruvate (100 mM) and 20% of FBS (Invitrogen, Carlsbad, CA). Cells were maintained at 37 °C with a humidified 5% CO₂ atmosphere.

### Cell Proliferation

Cells were seeded (2 x 10⁴ onto a 24-well plate) and incubated overnight to allow cell attachment. Cell number was determined using an electronic cell counter (Z1™ Series, Beckman Coulter, Inc) after 48 h incubation in culture media supplemented with sterile filtered supernatant (2.5-10%, v/v) from fecal fermentation. Results were expressed as % of controls incubated with culture media supplemented with sterile culture medium used for fecal fermentation.

### Generation of Reactive Oxygen Species (ROS)

Metabolites produced during fecal fermentation of CG cheese (0, 6, or 12 mo ripening) after IVD as detailed in section 2.7 were assessed for their effects in protecting human fibroblast CCD-18Co colon cells from production of ROS when challenged with E. coli lipopolysaccharides (LPS) mimicking obesity-induced inflammation. ROS levels were quantified using the dichlorofluorescein diacetate (DCFH-DA) fluorescent probe as previously reported [21]. Briefly, cells seeded (15 x 10⁴ cells/well) in a 96-well plate were incubated overnight to allow cell attachment followed by incubation with culture media supplemented with sterile filtered fecal culture supernatants (2.5% v/v) for 1 h. ROS generation was induced with LPS (2 μg/mL) for 2 hours. ROS levels are proportional to Relative Fluorescent Units (RFU) detected with DCFH-DA (10μM) after 30 min reaction and monitored at 520 nm emission and 480 nm excitation with a µQuant micro plate reader (Biotek Instrument, Rochester, VT, USA). Fecal culture supernatants of feces fermented with casein after IVD were used as control. Values were normalized to RFU of controls treated with casein fecal culture supernatants no challenged with LPS.

### Statistical analysis

Quantitative data represent mean values with the respective standard deviation (SD) or standard error of the mean (SE) of three or more replicates. One-way analysis of variance (ANOVA)

---

### Table 1: Sequences of primers used for PCR analysis of DNA

<table>
<thead>
<tr>
<th>Phylum</th>
<th>Class/Family/Genus</th>
<th>Species/Group</th>
<th>Primer sequence (5’-3’)</th>
<th>Approximately amplicon size (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Universal</td>
<td>All</td>
<td>All</td>
<td>ACTCTACGGAGGGACGAGGT</td>
<td>174-179</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>GTATTACCGCCGGTCGCTGCGAC</td>
<td></td>
</tr>
<tr>
<td>Bacteroidetes</td>
<td>All</td>
<td>All</td>
<td>GGAACATGGTGTTATATCGGATGAT</td>
<td>126</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>AGCTGACGAGAACCCATGAGCAG</td>
<td></td>
</tr>
<tr>
<td>Actinobacteria</td>
<td>Bifidobacterium</td>
<td>spp.</td>
<td>GCCCTGCTAAACACATCGAAGTC</td>
<td>126</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>CACCGGTTCACGAGGAGTATT</td>
<td></td>
</tr>
<tr>
<td>Firmicutes</td>
<td>All</td>
<td>All</td>
<td>TGAACACTAAGAATTTGAGGACC</td>
<td>157</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>ACCATCGACCACCTGTC</td>
<td></td>
</tr>
<tr>
<td>Proteobacteria</td>
<td>Enterobacteriaceae</td>
<td>group</td>
<td>CATTACGTGATACCCGAGAAGAAC</td>
<td>195</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>CTTTACGGACTCAAGACTTTGC</td>
<td></td>
</tr>
<tr>
<td>Firmicutes</td>
<td>Enterococcus</td>
<td>spp.</td>
<td>CCTTTATTGTATTGGAACATCTT</td>
<td>144</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>ACTCGTTGACTTTGCCATTGT</td>
<td></td>
</tr>
</tbody>
</table>
followed by a Bonferroni post-hoc test were performed with Graph Pad Prism (San Diego, CA). A value of $P < 0.05$ was considered statistically significant. Principal Component Analysis (PCA) was performed using SPSS version 15.0 (SPSS Inc., Chicago, IL). Figures were created with Graph Pad Prism and SigmaPlot™ (San Jose, CA).

### Results

#### Viability of L. helveticus and reducing sugars during ripening

Results showed the viability of L. helveticus decreased to levels < 30 cfu g\(^{-1}\) numbers within 12 mo of ripening (Fig 1a). The content of reducing sugars increased during the first 6 mo to ~2-fold of 0 mo and decreased within 6 to 24 mo down to levels found in the 0 mo control (Fig 1b).

#### Effects of ripening on CG caseins

CG caseins analyzed by electrophoresis showed that bands of higher molecular size tend to decrease throughout the ripening as illustrated in Fig 2a. The band intensities of P1 and P2, 25.0-29.4 kDa, decreased significantly at 6 and 12 mo ripening compared to 0 mo (Fig 2b). The bands of intermediate molecular size P3 and P4 (22-23.1kDa) increased at 6 mo ripening as a result of proteolysis of larger molecular size proteins and decreased to 0 mo levels at 12 mo ripening and beyond. The proteins of lower molecular size (17.9-19.4 kDa) did not change throughout ripening up to 18 mo, but decreased significantly at 24 mo ripening.

#### Effects of ripening on protein carbonyls and AGEs contents in CG cheese

Results showed that content of protein carbonyl products increased significantly with ripening (up to 2.7-fold of 0 mo control at 12 mo, Fig 2c); this was accompanied by an increase in the production of AGEs-fluorophores that reached maximum concentrations at 12 mo ripening (1.7-fold of 0 mo control, Fig. 2d).

### Modulation of fecal bacteria and production of SCFAs by CG cheese after IVD

Results showed that CG cheese after IVD did not change relative proportions of Firmicutes in feces from obese mice (ranged from 0.96-fold to 1.1-fold of obese control) (data not shown); however, the Bacteroidetes/Firmicutes ratio tended to increase in feces when fermented with 0 and 6 mo CG cheese after IVD (Fig 3a).

Enterococcus, a genus of lactic acid bacteria and common commensal organisms in the intestines, were significantly less abundant in feces from lean controls compared to the obese controls. The levels of Enterococcus in obese feces fermented with 0 mo CG cheese were similar to obese control fermented with casein after IVD. However Enterococcus DNA levels were significantly lower after fermentation with 6 and 12 mo ripened CG cheese after IVD and similar to the levels found in lean controls (Fig 3b). In addition, the DNA levels of Enterobacteriaceae were lower in feces from lean controls and the fecal fermentation with cheese (6-12 mo ripening) also decreased their relative abundance to levels similar to lean controls (Fig 3c). The lack of statistical significance among experimental groups on DNA relative abundances of the bacteria assessed by PCR was partially due to the high variability of fecal samples obtained from three different mice donors.

However, the PCA analysis showed that obese and 0 mo are located close while the 6 and 12 mo shift closer to the lean group indicating that bacteria abundances are similar to the lean group (Fig 3e).

The production of SCFAs has been associated with the growth of probiotic bacteria and seems to play an important role for epithelial proliferation in the gut [22]. Results showed that SCFAs concentrations in supernatants from fecal fermentation of CG after IVD were similar among ripening periods (0-12 mo) and ranged from 313 ± 17 to 324 ± 21 µmol L\(^{-1}\) for butyric acid, and 70 ± 5 to 75 ± 6 µmol L\(^{-1}\) for acetic acid with non-detectable levels of propionic acid.

### Metabolites produced by fecal fermentation of CG after IVD protected colon cells against oxidative stress

Protection of colon fibroblast CCD-18Co cells against oxidative stress was assessed with fecal culture supernatant at doses that cell proliferation was not inhibited 2.5% (v/v) (data not shown).

ROS are key signaling molecules which play an essential role in the progression of inflammatory disorders. Results showed that LPS challenge increased ROS levels in CCD-18Co cells up to 1.4-fold of cells pre-treated with supernatants from obese fecal fermentation of casein after IVD and not challenged with LPS. Pretreatment with 6 and 12 mo CG obese fecal culture supernatants prevented the ROS production and maintained levels down to 1.1- and 0.9-fold of control cells pre-treated with casein obese fecal supernatants and not challenged with LPS (Fig 4a).

---

Figure 2: Cheese ripening involves casein proteolysis and the generation of protein carbonyls and AGES. (a) SDS-PAGE and (b) Band intensities of CG cheese caseins. Proteins were extracted from a composite of three cheeses. To illustrate differences in protein concentrations, the relative intensity of bands P1 to P6 were determined using the Image J software as outlined in materials and methods. (c) Protein carbonyls and (d) AGE-fluorophores. Protein carbonyls were analyzed by the 2,4-dinitrophenylhydrazine (DNPH)-assay. AGE-fluorophores levels were determined based on the relative fluorescence units (RFU) at 360 nm excitation and 460 nm emission, as outlined in materials and methods. Values are average ± SD, (n = 3), different letters indicate statistical difference at p < 0.05.

Figure 3: CG cheese proteins after IVD change relative abundances of bacteria in feces from obese mice. (a) Bacteriodetes/Firmicutes DNA abundance ratio, (b) Enterococcus, (c) Enterobacteriaceae, (d) Bifidobacterium sp, and (e) PCA of relative DNA levels of the analyzed bacteria. Circles in PCA represent experimental units that are alike in DNA levels of bacteria. Obese control and 0 mo are enclosed by continuous line circle, lean, 6 and 12 mo are enclosed by dashed line circle. In vitro fecal fermentation was prepared using fecal slurries from obese mice (n = 3, no pooled fecal samples) in prerduced modified medium containing CG cheese (0, 6, and 12 mo) or casein (obese and lean controls) after IVD for 24 h under anaerobiosis and constant agitation as outlined in materials and methods. DNA was extracted from fecal material and analyzed for relative abundances of bacterial populations by PCR as detailed in materials and methods. Values are average ± SE, ***, p < 0.01 compared to the obese control.
Discussion

Counts of L. helveticus were reduced within 6 mo ripening. These results are in agreement with bacteria counts in Swiss-type cheeses produced with L. helveticus during ripening processes [23]. Literature reports that survival of SLAB is reduced during cheese ripening due to the decrease of lactose availability and autolytic activities [24]. The substrate changes that cause autolysis and release of bacterial enzymes during cheese ripening are associated with the cheese making process involving heat, acidic, osmotic, oxidative stresses on microorganisms and changes in the pH, water activity (aw), and redox potential gradients in the matrix [24, 25]. However, NSLAB might be viable and become the dominant microflora in the ripened cheese due to resistance to environmental stresses [24, 25].

The increase of reducing sugars during the first 6 mo was likely related to the conversion of lactose to glucose and galactose or galactose-6 phosphate by starter and NSLAB, which can occur at a higher rate than their use as source of energy, and as a result of the activity of enzymes, released by autolysis of bacteria cultures. Reducing sugars decreased after 6 mo ripening, which suggests their reaction with proteins to produce the protein carbonyls and other precursors of MRPs as demonstrated in “Formation of early and advanced Maillard reaction products correlates to the ripening of cheese” [4]. In addition, reducing sugars can be used as a source of energy by NSLAB during ripening, that were shown to survive the environmental changes occurring during cheese ripening [25].

Changes in CG caseins of higher molecular size (P1-P2) are associated with the composition of starter cultures and proteolysis occurring during the ripening period [26]. In addition, the residual activities of coagulants (chymosin, pepsin, or fungal acid proteinase) or at less extent, milk enzymes (plasmin and perhaps cathepsin D), produce large and intermediate-sized peptides. Specially, enzymes released due to autolysis of starter cultures and the adjunct L. helveticus are highly proteolytic and have shown to play a critical role in texture and flavor formation in cheese as well as the synthesis of bioactive peptides in situ [1, 23]. During ripening, CG caseins of lower molecular size (P3-P6) indicates that cheese ripening is a continuous dynamic process in which proteolysis of larger molecular size proteins counterbalances the continuous degradation of lower molecular size proteins. These changes in cheese caseins are associated with proteolytic activity of SLAB and NSLAB that contribute to the breakdown of proteins to produce low molecular size caseins, small peptides and free amino acids [23]. In general, significant changes in the protein band intensities occurred within 6-12 mo ripening, implying that an extended ripening beyond 12 mo might not be necessary to develop flavor and texture; when a highly proteolytic strain such as L. helveticus is used as adjunct culture.

Protein carbonyls are formed only in the presence of reducing sugars and may act as early precursors for the later production of AGEs- fluorophores [16]. These compounds have been used as a parameter of ripening in cheeses [4, 27, 28] and their formation was accompanied by the loss of galactose in Manchego cheese [29].

Also, results showed that CG after 24 h cheese production and day 0 of ripening had a pH = 4.9 and remained almost constant throughout ripening (0 to 24 mo) (pH = 5.08 ± 0.06) (data not shown). Lactose conversion to lactic acid occurs during the first 24 h of cheese production, coupled with growth of SLAB, lactic acid fermentation, and decrease in pH to approximately 5.1 to 5.25 [24].

Furthermore, glycated proteins were shown to be less susceptible to pepsin hydrolysis when compared to non-glycated ones and to potentially exert a beneficial effect in the lower gastrointestinal tract by modulating gut microbiota [8, 9]. Likewise, sodium caseinate glycated via Maillard reaction with galactose and lactose and corresponding peptides (which might resemble reactions occurring in cheese) were quickly fermented after simulated gastrointestinal digestion by strains of Lactobacillus, Streptococcus and Bifdobacterium. The bacteria growth was promoted at a greater extent than β-lactoglobulin complexes or even glucose [30]. This study suggested that conjugation of milk proteins with galactose and lactose via the Maillard reaction could be an efficient method to obtain novel food ingredients with a potential prebiotic character. These reactions occur during cheese ripening [4] and contribute to the bioactive potential of ripened cheese. This has been supported by an in vitro study demonstrating that MRPs from sodium caseinate lactose and glucose, exerted higher antioxidant activity than the intact milk protein and protected against...
oxidative damage [31]. Furthermore, the same research group demonstrated later in vivo that MRPs from sodium caseinate and their correspondent products after fermentation by Lactobacillus enhanced cardiovascular health in mouse and rat models [32]. The antioxidant activity and cardiovascular protective potential of MRPs derived from milk proteins increased after enzymatic hydrolysis [33] and after fermentation with Lactobacillus strains [34]. However, the bifidogenic activity of MRPs has been questioned since caseinomacropetide nonenzymatically glycosylated with prebiotic carbohydrates exerted a similar bifidogenic activity to that of the corresponding prebiotic carbohydrates in cultured human feces after being subjected to in vitro gastrointestinal digestion [35].

Gut microbiota composition can be altered by diet and influence the host health. Alterations in the gut microbiota of obese mice and humans have been associated with increase of Firmicutes and decrease of Bacteroidetes; the two dominant bacteria phylum in the human gut. A previous study reported that Bacteroidetes/Firmicutes ratio is low in obese state and this ratio increased with weight loss [36]. These alterations contribute to the increased efficiency of energy harvest in obesity [37]. Even though only an increasing trend in relative DNA abundances of Bacteroidetes to Firmicutes was found; the changes in Enterococcus DNA relative abundances were consistent with a previous study reporting that Enterococcus population is reduced in feces from lean rats compared to feces from obese rats [38]. Enterococcus abundance was positively correlated with genes and metabolites promoted by high protein diets and generally observed to be involved in colonic disease [39]. However, some Enterococcus species such as Enterococcus faecalis FK-23 exhibit anti-obesity effects in mice fed high fat diet [40]. Further in vivo research is needed to elucidate the specific Enterococcus species that seem to be inhibited by consumption of ripened cheese.

Likewise, a promising trend indicating decreasing levels of Enterobacteriaceae in obese feces fermented with cheese after IVD are supported by a previous study reporting that significantly higher levels of Enterobacteriaceae are present in feces of obese/overweight children [41]. Similarly, a promising trend to increase the levels of Bifidobacterium will need to be further investigated in vivo due to its known role as probiotic beneficial bacteria and its negative association with BMI in a study of feces from obese and lean subjects [42].

In general, only trends were observed among experimental groups regarding relative abundances of these bacteria assessed by PCR (p > 0.05) due to the high variability of fecal samples obtained from three different mice donors. Similar results have been previously reported [35]. Similarly, concentrations of SCFAs in fecal culture supernatant were not statistically different from the obese controls, though they have the tendency to increase in cultures with 6 and 12 mo ripening. The main effects of glycosylated caseino macro peptides seems to be their bifidogenic activity, no the production of SCFAs as is reported for dietary fiber [43].

The microbial metabolites in supernatants from obese feces cultured with 6 and 12 mo CG after IVD seemed to protect the fibroblast CCD-18Co cells from LPS-induced ROS at similar extent and maintained ROS levels down to the levels of controls not challenged with LPS. These results are supported by a previous study demonstrating that longer ripened dairy products impact positively levels of oxidative stress [44]. Furthermore, recent evidence indicates benefits of intestinal bacteria and metabolic activities, partly due to controlled release of ROS. Thus, gastrointestinal bacteria has been considered as an active adjuvant treatment for mood disorders [45].

In conclusion, our results demonstrated that proteins and peptides produced during the ripening process in CG cheddar cheese and those released after in vitro gastric digestion, can act as modulators of bacterial growth in feces. There was a promising trend to shift relative abundances of specific genera and phyla of obese mice towards a profile that resemble the lean mice fecal bacteria proportions. In addition, metabolites produced by fecal bacteria fermentation of CG cheese after IVD might protect colon fibroblast cells from chronic inflammation induced by ROS. However, further in vivo studies are needed to validate the potential of ripened cheese consumption for manipulation of gut microbial communities in obesity.

Acknowledgement

The authors are grateful for the financial support of the Build Dairy Program, managed by the Western Dairy Center at Utah State University. The authors would to thank Paulina Pilla, student from Pan-American School of Agriculture, Zamorano, Honduras for her technical assistance and the WSU creamery personnel for kindly providing the CG cheese samples kept in inventory up to 24 mo for our study.

References

8. Swiatecka D, Narbad A, Ridgway KP, Kostyra H. The study on the impact...


Page 9 of 10

Copyright: © 2016 Noratto, et al.
Aged American Cheddar Cheese as Source of Protein Derived Compounds that Modulate Obese Mice Fecal Bacteria and Colon Inflammation In Vitro


44. Geurts L, Everard A, le Ruyet P, Delzenne NM, Cani PD. Ripened dairy products differentially affect hepatic lipid content and adipose tissue oxidative stress markers in obese and type 2 diabetic mice. Journal