Involvement of *Acetobacter orientalis* in the production of lactobionic acid in Caucasian yogurt (“Caspian Sea yogurt”) in Japan

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**ABSTRACT**

Lactobionic acid was first found in a Caucasian fermented milk product popularly known as “Caspian Sea yogurt” in Japan. The presence of lactobionic acid in the fermented milk was indicated by the results of both high-performance anion-exchange chromatographic analysis with pulsed amperometric detection and mass spectrometric analysis. Thereafter, the acid was purified from the yogurt and analyzed by nuclear magnetic resonance. A substantial amount of lactobionic acid was found to be accumulated in the upper layer of the yogurt, especially within 10 mm from the surface. A total of 45 mg of lactobionic acid per 100 g of the upper yogurt layer was collected after 4 d of fermentation. The annual intake of lactobionic acid in individuals consuming 100 g of the yogurt every day would be 0.5 to 1.0 g. A lactose-oxidizing bacterium was isolated from the fermented milk and was identified as *Acetobacter orientalis*. Washed *A. orientalis* cells oxidized monosaccharides such as d-glucose at considerable rates, although their activities for substrates such as lactose, maltose, and cellobiose were much lower. When *A. orientalis* cells were cultivated in cow’s milk, they exhibited lactose-oxidizing activity, suggesting that this bacterium was the main organism involved in the production of lactobionic acid in the yogurt.

**Key words:** lactobionic acid, *Acetobacter orientalis*, Caspian Sea yogurt

**INTRODUCTION**

Lactobionic acid [β-D-galactopyranosyl-(1→4)-D-gluconic acid, *LacA*] is a component of the preservative solution used during organ transplantation (Wahlberg et al., 1987; Sumimoto et al., 1992) and is also used as an ingredient in the antibiotic erythromycin lactobionate (Neaverson, 1976). Lactobionic acid is thought to have several additional functions, such as in the formation of soluble salts with calcium to increase their intestinal adsorption (Brommage et al., 1993; Nakano et al., 2006). Thus, *LacA* might have potential for use in calcium supplements. Bacteria (Satory et al., 1997; Murakami et al., 2002, 2006) and fungi (Ludwig et al., 2004; Hua et al., 2007; Kiryu et al., 2008) with lactose-oxidizing activity have been studied as a means of producing LacA from lactose. The microorganisms used in these studies, however, are not suitable for food production. Therefore, it is necessary to find a bacterium used in food production, such as *Acetobacter* sp., for LacA production. *Acetobacter* sp. is one of the most important bacteria used in food production. It is used in the production of vinegar and the Caucasian yogurt studied here. The occurrence of LacA in foods and beverages also helps in the practical application of LacA, because it indicates that people have unconsciously consumed LacA for a long time. To our knowledge, the natural occurrence of LacA has not been demonstrated previously.

Caucasian yogurt, also known as “Caspian Sea yogurt” in Japan, was traditionally produced in the Caucasus region in the western part of Eurasia, and is believed to have been introduced into Japan from the village of Java, Georgia, in 1986 (Akatani and Ishida, 2002; Ishida et al., 2005). This yogurt has a lower acidity and greater viscosity than typical yogurts. In an early study, 3 bacteria, *Lactococcus lactis* ssp. cremoris, *Leuconostoc* sp., and *Gluconobacter* sp., were isolated from Caucasian yogurt (Akatani and Ishida, 2002). However, reexamination revealed that *L. lactis* ssp. *cremoris* and *Acetobacter orientalis* were the main bacteria involved in the production of the yogurt (Ishida et al., 2005). This yogurt is easily fermented after adding 1/10 to 1/20 vol of the seed yogurt to milk at room temperature, because the yogurt is fermented at a lower temperature (approximately 25 to 30°C) than typical yogurts. Because of this easy preparation, the yogurt has gained ground among Japanese people by word of mouth, and has become known as Caspian Sea yogurt. Fujicco (Hyogo, Japan), a major Japanese food company, has been selling seed powder of the yogurt (a mixture of dried cells of *L. lactis* ssp. *cremoris* and *A. orientalis*) since 2002.

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and the yogurt has become widely available in Japan. According to the Web site of the company (http://www.caspia.jp/), 1 million packages of powdered seed have been sold thus far. The yogurts derived from these powdered seed cells are fermented in Japanese homes.

**MATERIALS AND METHODS**

**Materials**

Commercial cow’s milk was used (butterfat content, 3.5%). A powdered culture of Caucasian yogurt was obtained from Fujicco. Authentic LacA was purchased from Wako Pure Chemical Industries (Osaka, Japan). Maltobionic acid [α-D-glucopyranosyl-(1→4)-D-gluconic acid] and cellobionic acid [β-D-glucopyranosyl-(1→4)-D-gluconic acid] were prepared by oxidizing maltose and cellobiose, respectively, by using a carbohydrate:acceptor oxidoreductase from *Paraconiothyrium* sp., as described previously (Kiryu et al., 2008).

**Preparation of the Caucasian Yogurt**

Powdered culture (3.0 g) was mixed with fresh cow’s milk (30 mL) and was incubated at 27°C for 24 h under static conditions to prepare a seed culture. The starter culture was prepared as follows: fresh cow’s milk was mixed with 1/10 vol of the seed culture and then incubated at 27°C for 24 h. This starter culture was stored at 4°C until seeding. The yogurt was fermented by adding the starter culture under the same conditions used to prepare the starter culture.

**High-Performance Anion-Exchange Chromatography with Pulsed Amperometric Detection Analysis**

After removal of the protein and other insoluble materials from the fermented milk samples, the samples were analyzed for the presence of LacA by high-performance anion-exchange chromatography with pulsed amperometric detection (HPAEC-PAD; Dionex Corporation, Sunnyvale, CA). To remove protein from the yogurt samples, water (same volume) and 1.0 M acetate (one-fifth the volume; special grade) were added to the yogurt sample and mixed vigorously. The precipitates thus formed were removed by centrifugation at 7,000 × g for 10 min at 18°C. The supernatant was then mixed with ethanol (one-half the volume) and immediately centrifuged again under the same conditions to remove the precipitate. The supernatant was subjected to HPAEC-PAD. Analysis was performed under the following conditions: system, Dionex DX-500; detector, Dionex Model PAD II pulsed amperometric detector; column, Dionex Carbopac PA-1 (40 × 250 mm); elution, linear gradient from 150 to 300 mM sodium acetate in 100 mM NaOH; flow rate, 1.0 mL/min; temperature, 35°C.

**Isolation of LacA from Caucasian Yogurt**

To prepare the yogurt, 50 g of the starter culture was added to cow’s milk (500 mL in a 3-L flask) and incubated for 5 d at 27°C. The fermented milk was centrifuged at 5,000 × g for 30 min at 10°C, and the precipitate was removed. Ethanol (one-half the volume; 250 mL) was added to the supernatant (500 mL), and the mixture was centrifuged again at 5,000 × g for 30 min at 10°C. The supernatant solution was then applied to an anion-exchange column (Diaion PA 406S column, diameter 20 × 30 mm; Mitsubishi Chemical Corporation, Tokyo, Japan) equilibrated with water. The fractions were eluted with a linear gradient of 0 to 0.01 N HCl at 20°C (flow rate, 1.0 mL/min) and the fractions were analyzed by HPAEC-PAD analysis. Excess methanol was added to the fractions containing LacA and evaporated to remove the HCl. Further purification was carried out by thin-layer chromatography (TLC) under the following conditions: TLC plates, Kaisel Gel 60 (Merck, Darmstadt, Germany); solvent system, ethyl acetate:acetic acid:water (3:1:1, by vol). The plate was heated to 150°C and sprayed with 50% (wt/wt) H2SO4 in methanol, revealing spots of sugars. The silica gel on the TLC plate (without spraying and heating), having approximately the same mobility as that of LacA, was recovered with a spatula and suspended in water (1.0 mL). After centrifugation, the supernatant containing the extracted LacA was collected and evaporated.

**Mass Spectrometry Analysis**

The LacA sample isolated from the Caucasian yogurt was diluted with 60% acetonitrile and analyzed with an API 2000 mass spectrometer (Applied Biosystems, Foster City, CA) as follows: ionization, electron-spray ionization; ion spray voltage, 5,000 V; detection, negative ion mode. The mass of LacA [(m/z)− = 357] was determined from the mass of pseudomolecular ions ([M − H]−). The signal derived from LacA [(m/z)− = 357] was fragmented by 25 V of collision energy.

**Nuclear Magnetic Resonance**

The 1H-nuclear magnetic resonance (NMR; 300 MHz) and 13C-NMR (75 MHz) spectra were recorded for a solution of 1% sample in deuterium oxide (D2O) by using a Jeol AL-FTNMR spectrometer (Jeol Ltd., Tokyo, Japan). 3-(Trimethylsilyl)-1-propanesulfonic acid sodium salt was used as the internal standard.
Analysis of LacA Concentrations in Various Layers of the Caucasian Yogurt

The starter culture (5 g) was used to inoculate 45 mL of fresh cow’s milk in a plastic tube (50 mL; diameter, 30 mm) with a silicone cap. After 5 d of static cultivation at 27°C under aerobic conditions, the fermented milk was frozen at −20°C and cut into disks of 10-mm layers. These layers of fermented milk were then defrosted, deproteinized, and subjected to HPAEC-PAD analysis.

Isolation of LacA-Producing Bacterium from the Caucasian Yogurt

Yogurt culture (10 mg) was used to inoculate a yeast extract-polypeptone-glucose-lactose (YPGL; 5.0 mL, pH 7.0) liquid medium consisting of 0.5% dried yeast extract (D-3, Nihon Pharmaceutical, Tokyo, Japan), 0.5% polypepton (Nihon Pharmaceutical), 0.5% lactose, 0.5% d-glucose, 0.1% MgSO4. The culture medium was shaken (240 oscillations/min) to selectively proliferate aerobic bacteria. After 24 h of culture at 27°C, 0.8 μL of the culture liquid was streaked onto YPGL plates (1.5% agar) and the plates were incubated for a further 24 h at 27°C. The isolates were then cultured in the YPGL liquid medium (5.0 mL) at 27°C for 48 h with shaking (240 oscillations/min). The cells were harvested by centrifugation (5,000 × g, 10 min, 18°C) and washed with 0.85% NaCl. The washed cells were incubated with 2% lactose solution (1.0 mL) containing CaCO3 (2.8 mg, 1/2 equimolar amount of lactose) at 27°C for 48 h with shaking (240 oscillations/min). After removal of the cells, the reaction mixture was analyzed by TLC to select the LacA-producing bacteria.

Conversion of Several Saccharides by Washed A. orientalis Cells

Acetobacter orientalis KYG22 was cultivated in YPGL liquid medium (5.0 mL) at 27°C with shaking (240 oscillations/min). The cells in 1.0 mL of culture medium were collected by centrifugation, washed with 0.85% NaCl, suspended in 100 μL of 0.85% NaCl, and incubated with 900 μL of the substrate solution, consisting of 10 mM of each saccharide (d-glucose, d-galactose, d-mannose, d-xylose, L-arabinose, maltose, lactose, cellobiose) and 100 mM of acetate buffer (pH 5.5) at 27°C with reciprocal shaking (240 oscillations/min). After 3 h, the reaction mixtures were boiled for 5 min and the oxidized products were analyzed by HPAEC-PAD. The oxidized disaccharides and monosaccharides were evaluated by using LacA and d-glucuronic acid as the standard, respectively.

Time Course of LacA Production by A. orientalis Cells

Acetobacter orientalis strain KYG22 was cultivated in 50 mL of YPGL liquid medium at 27°C for 48 h with shaking (120 oscillations/min) in a 500-mL shaking flask, and the cells were collected from 10 mL of culture broth by centrifugation at 7,000 × g for 30 min at 18°C. After washing with 0.85% NaCl, the cells were suspended in 1.0 mL of reaction mixture containing 2% lactose and 2.8 mg of CaCO3 (1/2 equimolar amount of lactose), then incubated at 35°C with shaking (240 oscillations/min) in test tubes. The cells were removed by centrifugation, and LacA production was measured by HPAEC-PAD.

Estimation of Daily and Annual Intakes of LacA

A starter culture for the Caucasian yogurt (100 g) was mixed with 600 mL of commercial fresh milk and fermented by using a household yogurt maker (Yogurtia Model YM-1200; Tanica Electric, Gifu, Japan) at 27°C for 12 h. The yogurt was stored at 4°C for 12 h, after which 100 g of the upper part of the sample was subjected to HPAEC-PAD analysis, as described above. In addition, 100 g of the yogurt was sampled every 24 h (36, 60, 84, 108, 132 h) for LacA measurement. Based on the assumption that 100 g of the yogurt would be eaten every day, the daily and annual intakes of LacA were estimated from the total amounts of LacA in the sampled yogurts (12 to 132 h), defined as 6 d of intake.

Lactose-Oxidizing Activity Assay Using a d-Glucuronic Acid Measurement Kit

The activity of the cells was calculated based on LacA production. Lactobionic acid was measured as d-glucuronic acid after acid hydrolysis as follows. A cell suspension or sonicated cell solution (250 μL) was mixed with a mixture (250 μL) containing 400 mM lactose and 200 mM acetate buffer (pH 5.5), then incubated at 40°C for 10 min. Fifty microliters of 1.0 N NaOH was added to the reaction mixture to stop the reaction. The mixture was neutralized by adding 50 μL of 1.0 N HCl. The reaction mixture was boiled with 500 μL of 2.0 N HCl for 40 min to hydrolyze LacA to D-glucuronic acid and D-galactose. After adding 2.0 N NaOH (500 μL), the amount of D-glucuronic acid was measured with a D-glucuronic acid measurement kit (F-kit, D-Glucuronic acid/D-Glucono-δ-lactone; Boehringer Mannheim, Mannheim, Germany). One unit of the cells was defined as the amount of cells that produced 1 μmol of LacA per minute.
Utilization of LacA as a Carbon Source by A. orientalis

Acetobacter orientalis strain KYG22 was cultured in YPGL liquid medium (5.0 mL) at 27°C for 48 h. Ten milliliters of the culture broth was diluted 1.0 × 10⁴-fold, and the diluted sample (200 μL) was plated on mineral medium plates (pH 5.5 to 6.0) consisting of NaCl (0.2%), NH₃NO₃ (0.2%), KH₂PO₄ (0.1%), K₂HPO₄ (0.1%), sugar (d-glucose or LacA, 1%), and agar (1.5%). The plates were incubated for 72 h at 27°C, and growth of the bacterium was observed.

Effects of pH and Temperature on A. orientalis Cells

Acetobacter orientalis strain KYG22 was cultured in YPGL liquid medium (1.0 mL) at 27°C with shaking (240 oscillation/min) for 3 d. The cells were collected by centrifugation (5,000 × g, 10 min at 18°C) and suspended in 250 μL of 0.85% NaCl. To determine the optimal pH, the activity of the cell suspension was measured with a d-gluconic acid measurement kit. The pH of the reaction mixtures was adjusted with 100 mM sodium acetate-HCl buffer (pH 2.0 to 4.0), acetate buffer (pH 4.0 to 5.5), phosphate buffer (pH 7.0 to 8.0), and glycine-HCl buffer (pH 9.0 to 11.0). To measure pH stability, the cell suspension (250 μL) was mixed with 50 μL of 250 mM sodium acetate-HCl buffer (pH 2.0 to 4.0), acetate buffer (pH 4.0 to 5.5), phosphate buffer (pH 7.0 to 8.0), and glycine-HCl buffer (pH 9.0 to 11.0), and was incubated at 40°C for 24 h. The cell suspension was mixed with the reaction mixture (300 mL) containing 400 mM lactose and 400 mM acetate buffer (pH 5.5), and the activity was measured. To determine the optimal temperature, activities of the cells were measured at 5 to 80°C. To measure thermal stability, cells in 100 mM acetate buffer (pH 5.5) were treated at 0 to 80°C for 30 min. The activity of the treated cells was measured by adding them to 1/5 vol of 1.0 M lactose.

Preparation of Soluble and Membrane-Bound Fractions of A. orientalis Cells

To investigate whether the lactose-oxidizing enzyme of A. orientalis was membrane bound or soluble, A. orientalis KYG22 was cultured for 2 d, collected by centrifugation (5,000 × g, 10 min at 5°C), washed, suspended in 100 mM acetate buffer (pH 5.5), and sonicated on ice by using an Ultrasonic Homogenizer 5202 with a CN 1005 horn tip (Otake Seisakusho, Tokyo, Japan). After 10 min of sonication, the homogenate sample was centrifuged (5,000 × g, 10 min at 5°C). The supernatant and the precipitates were used as the soluble and membrane fractions, respectively.

RESULTS

Detection and Identification of LacA in the Caucasian Yogurt

We suspected, based on the fact that 5.0 to 5.5% of milk is lactose, that LacA would be produced in fermented milk. We therefore analyzed 25 fermented milk
samples and eventually found a peak that appeared at the same retention time as that of the LacA standard in a Caucasian yogurt, popularly known as Caspian Sea yogurt in Japan (Figure 1A). When a LacA standard solution was added to the deproteinized yogurt sample, the peak was enlarged. The retention times of maltobionic acid and cellobionic acid were different from that of LacA. The peak was absent from the other yogurts tested. As shown in Figure 1B, mass spectrometry analysis also indicated the existence of LacA \([m/z]^{-} = 357\) in the yogurt. The compound corresponding to the peak (compound A) was purified from the yogurt by using an anion-exchanger and TLC for analysis with a mass spectrometer. The signal of purified compound A was the same as that of LacA \([m/z]^{-} = 357\). When the signal \([m/z]^{-} = 357\) was trapped and fragmented, the fragmentation pattern was the same as that of the LacA standard (Figure 2). Compound A was also isolated from the yogurt and further identified as LacA by \(^1\)H- and \(^{13}\)C-NMR. The carbon and proton signals of purified compound A coincided with those of the LacA standard (data not shown; Murakami et al., 2002).

**Production of LacA in Various Layers of the Caucasian Yogurt**

The oxidation of lactose is considered to proceed under aerobic conditions in fermented milk. Therefore, LacA production in various layers of the yogurt was measured (Figure 3). To measure the LacA concentration in each layer, the yogurt was prepared in plastic tubes and frozen. After the frozen yogurt was removed from the tube, it was cut into 4 layers as shown in Figure 3. Each layer was deproteinized, and the LacA concentration was measured by HPAEC-PAD. The first fraction (surface layer, 0 to 10 mm) contained the largest concentration of LacA (more than 10 mg/100 g of yogurt). The deeper layers contained noticeably less LacA.

**Time Course of Accumulation of LacA in Caucasian Yogurt and Estimation of Annual Intake of LacA**

The amount of LacA in the upper layer was greater than the amounts in the lower layers (Figure 3). Thus, to investigate LacA production in the upper layer of the yogurt, the yogurt was prepared from a thin (50-mm) layer of cow’s milk and the amount of LacA was measured after different culture periods (Figure 4). The starter culture (1.0 g) and fresh cow’s milk (10 mL) were mixed in a 100-mL flask (thickness, 50 mm) and incubated at 27°C. The pH of the yogurt was changed to 4.6 for 6 h and 4.2 for 10 h. The milk began to thicken after 8 h, and the surface of the yogurt turned a light reddish-brown color, probably because of the growth of the aerobic bacterium *A. orientalis* in the yogurt. During fermentation, the amount of LacA gradually increased, and 45 mg of LacA accumulated in 100 g of the upper layer of the yogurt after 4 d.

Next, the daily and annual intakes of LacA were estimated, assuming that the yogurt was eaten every day. The yogurt was prepared by using the Yogurtia Model YM-1200, a commercial fermented milk maker (n = 3). Starter culture (100 g) was inoculated into fresh milk (600 mL) and incubated at 27°C for 12 h, and the prepared yogurt was stored at 4°C. The upper part of the yogurt (100 g) was collected at various times, and the amount of LacA was measured by HPAEC-PAD (Figure 5). The isolated yogurt samples contained 1.4 ± 0.2 mg (12 h), 2.3 ± 0.4 mg (36 h), 2.8 ± 0.6 mg (60 h), 2.5 ± 0.7 mg (84 h), 1.7 ± 0.4 mg (108 h), 1.8 ± 0.5 mg (132 h), and 2.1 mg (average) of LacA in 100 g of yogurt. Assuming that 100 g of yogurt was consumed each day, the intake of LacA over a 6-d period (12 to 132 h) would be 12.6 mg, as shown in Figure 5. Thus, the annual intake of LacA was calculated to be approximately 760 mg.

**Identification of a Lactose-Oxidizing Bacterium, *A. orientalis*, and Its Sugar-Oxidizing Properties in Caucasian Yogurt**

The LacA-producing bacterial strain KYG22 was isolated from the fermented milk. Strain KYG22 is a...
gram-negative bacillus and was identified as *A. orientalis* based on its 16S ribosomal DNA and phenotypic properties as analyzed by TechnoSuruga Laboratory (Shizuoka, Japan; data not shown) NCIMB Japan. The properties of lactose oxidation by *A. orientalis* were then investigated. The bacterium was cultivated by using fresh cow's milk (2.0 mL) and YPGL liquid medium (2.0 mL) for 24 h at 27°C with and without shaking. The fermented milk did not thicken, probably because of insufficient lactic acid production. After 2 d of cultivation, the cells were collected and the growth of the cells (A₆₆₀ value) in YPGL medium with and without shaking was evaluated: these values were 1.2 and 0.2, respectively. The washed cells reacted with 1.0 mL of 2% lactose in 50 mM of acetate buffer (pH 5.5) for 7 h at 27°C with shaking (240 oscillation/min). Production of LacA was evaluated by TLC (Figure 6). The cells cultured in milk oxidized lactose at a rate very similar to that of cells in YPGL medium. The production of LacA with shaking, however, was much greater than that without shaking. The time course of lactose oxidation in washed *A. orientalis* cells is shown in Figure 7. When washed cells of the strain were incubated with 2% lactose with shaking, 0.72% of LacA was produced after 72 h of incubation.

**Characterization of the Sugar-Oxidizing Activities of A. orientalis Cells**

*Acetobacter orientalis* was cultivated on mineral medium plates containing D-glucose or LacA as the sole carbon source. The bacterium grew on plates containing D-glucose but produced no colonies on plates containing LacA. The effects of pH and temperature on the oxidizing activities of washed *A. orientalis* cells were investigated. The optimal pH for the oxidizing activity of the cells was 5.5, and the cells oxidized lactose most efficiently at 48°C. The pH stability range was
6.0 to 7.5, and thermostability was less than 55°C. The specificity of *A. orientalis* cells with respect to several sugars is shown in Figure 8. The bacterium efficiently oxidized D-glucose and several other monosaccharides; however, it exhibited low activities for lactose, maltose, and cellobiose (relative activities 0.04, 0.05, and 0.07%, respectively). The bacterium was sonicated, and the disrupted cells were separated into soluble and membrane fractions. Most of the lactose-oxidizing activity was detected in the membrane fraction, indicating that the enzyme involved in the oxidation of lactose was probably a membrane-bound enzyme.

**DISCUSSION**

Lactose oxidation by *A. orientalis* cells cultured in milk indicates an important role of *A. orientalis* in LacA production in Caucasian yogurt (Figure 6). However, in addition to *A. orientalis*, *L. lactis* ssp. *cremoris* is involved in the fermentation of yogurt prepared from the seed powder from Fujicco. The yogurts fermented by *L. lactis* ssp. *cremoris* alone are distributed as Caspian Sea yogurt in Japan, and we prepared fermented milks to use these yogurts as starter cultures. However, we failed to find LacA in these yogurts. On the other hand, LacA was produced by adding a small amount of *A. orientalis* cultured broth (10 μL) to these starter cultures (data not shown). Furthermore, contamination by unwanted bacteria did not significantly affect LacA production in the yogurt, because the pH of the yogurt became acidic for a few hours (Figure 4). In other experiments, the pH shifted rapidly to the acidic range. The conclusion that LacA in the Caucasian yogurt was produced by *A. orientalis* can be drawn from these results.

The LacA content of the yogurt suggested that the daily and annual intakes of LacA would be 1.0 to 3.0 mg and 0.5 to 1.0 g, respectively, if 100 g of yogurt prepared with a commercial yogurt maker were eaten every day (Figure 5). Because Caucasian yogurt is customarily eaten at every meal during the day in the Caucasus region, the people living in this region would likely have a much larger intake of LacA than Japanese people over a long period.

As shown in Figure 6, *A. orientalis* cells produced less LacA in static culture than in shaking culture because of the difference in oxygen supply. This result supports the results shown in Figure 3. Lactobionic acid was more abundant in the upper layer (aerobic conditions) of the yogurts.

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**Figure 5.** Estimation of daily intake of lactobionic acid (LacA). The Caucasian yogurt (approximately 700 mL) was prepared with a yogurt maker. After 12 h of fermentation at 27°C, the yogurt was stored at 4°C. One hundred grams of yogurt was sampled at 12 h and every 24 h thereafter (36, 60, 84, 108, 132 h) to measure the LacA concentration by high-performance anion-exchange chromatography with pulsed amperometric detection. Results are means ± standard deviation (n = 3).

**Figure 6.** Lactobionic acid (LacA) production by resting cells cultured in yeast extract-polypepton-glucose-lactose (YPGL) liquid medium and milk. *Acetobacter orientalis* was cultured in 2.0 mL of YPGL liquid medium (lane 3, 4) and milk (lane 5, 6) at 27°C for 2 d with (lane 4, 6) and without (lane 5, 7) shaking. After washing, the cells were reacted with 2% lactose and 0.026 g/mL of CaCO₃ at 27°C with shaking (240 oscillation/min). After 7 h of shaking, the reaction mixture was analyzed by thin-layer chromatography. Lanes 1 and 2 are the LacA and lactose standard, respectively.
yogurt than in the lower layers (anaerobic conditions), because the population of the aerobic bacteria would be larger at the surface. Furthermore, lactose oxidation is probably an aerobic process. The LacA content in the yogurt gradually increased during the fermentation period (Figure 4), because the pH of the yogurt and the optimal pH of the lactose-oxidizing activity were nearly the same (pH 4.0 to 4.5). Furthermore, _A. orientalis_ did not grow on a mineral salt medium containing LacA, but did grow on a medium containing d-glucose. These results indicate that LacA was not utilized as a carbon source by _A. orientalis_. Kontula et al. (1999) reported that _Lactococcus lactis_ ssp. _lactis_ does not utilize LacA. Neither _A. orientalis_ nor _L. lactis_ ssp. _cremoris_, another microbe in the Caucasian yogurt, appeared to utilize LacA. Lack of assimilation of LacA by _A. orientalis_ as well as by _L. lactis_ are also reasons for the accumulation of LacA in the fermented milk product.

_d_-Glucose was the preferred substrate for the membrane-bound enzyme of _A. orientalis_ (Figure 8). The specificity of this substrate was similar to that of a pyrroloquinoline quinone-dependent glucose dehydrogenase (GDH; EC 1.1.99.17; Cleton-Jansen et al., 1989a,b). Glucose dehydrogenase has been classified into 2 groups: soluble GDH (Dokter et al., 1986; Southall et al., 2006) and membrane-bound GDH (Matsushita et al., 1980, 1989; Ameyama et al., 1986). Soluble GDH has wide substrate specificity and oxidizes lactose at a considerable rate. In contrast, membrane-bound GDH specifically oxidizes monosaccharides and has no or extremely low activity against lactose. The oxidation activity of _A. orientalis_ on lactose was located in the membrane fraction, and it oxidized d-glucose and some monosaccharides (Figure 8). The activities on lactose, maltose, and cellobiose were extremely low. The substrate specificity and the location of the sugar-oxidizing activity were similar to those of membrane-bound GDH. In contrast, GDH from other acetic bacteria, _Acetobacter suboxydans_ (King and Cheldelin, 1956) and _Gluconobacter oxydans_ (Matsushita et al., 1981), have been reported to show no activity on lactose. It was therefore difficult to conclude whether the enzyme in this study was membrane-bound GDH or a different type of enzyme. Further investigation is necessary to characterize the lactose-oxidizing enzyme.
As shown in Figure 7, *A. orientalis* cells oxidized lactose to produce LacA. The conversion rate was slower during the first 24 h than that after 24 h. The optimal pH of the oxidation activity of the cells was 4.5, and the initial pH of the reaction mixture (approximately 7.6) was not suitable for the reaction. After 24 h, the production of LacA changed the pH to a more suitable pH (approximately 6.8). The reaction mixture contained 0.72% (720 mg/100 mL) of LacA after 72 h of incubation. This concentration was much greater than that shown in Figure 4 (45 mg/100 g of upper layer of yogurt) and Figure 5 (1.4 to 2.8 mg/100 g of yogurt). The primary cause of these differences is that the reaction mixture (1.0 mL) shown in Figure 7 contained a large amount of cells collected from 10 mL of the cultured broth. These differences also indicate the importance of oxygen supply both in the culture and in the reaction, because the experiment shown in Figure 7 involved shaking during both the culture and reaction, and the yogurt was prepared by static fermentation.

Several lactose-oxidizing enzymes from fungi, such as carbohydrate:acceptor oxidoreductase (Xu et al., 2001; Hua et al., 2007; Kiryu et al., 2008), glucooligosaccharide oxidase (Lin et al., 1991; Lee et al., 2005), and cellobiose dehydrogenase (EC 1.1.99.18; Ludwig et al., 2004), have been studied for their ability to produce LacA. Furthermore, bacteria (Nishizuka and Hayashi, 1962; Satory et al., 1997; Murakami et al., 2006), algae (Bean and Hassid, 1956; Sullivan and Ikawa, 1973), and citrus fruits (Bean et al., 1961) with lactose-oxidizing activities have been studied for LacA production. Most of the lactose-oxidizing organisms investigated, however, were not appropriate for the production of LacA for food. On the other hand, *Acetobacter* has been used in food production, such as in the production of vinegar, for a long time and safe production processes are in food production, such as in the production of vinegar. On the other hand, *Acetobacter* strains isolated from traditional Caucasian fermented milk. J. Nutr. Sci. Vitaminol. (Tokyo) 51:187–193.


