Effect of high-pressure processing on reduction of *Listeria monocytogenes* in packaged Queso Fresco


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**Received September 29, 2013.**
**Accepted November 28, 2013.**

1Mention of trade names or commercial products in this publication is solely for the purpose of providing specific information and does not imply recommendation or endorsement by the USDA.
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**ABSTRACT**

The effect of high-hydrostatic-pressure processing (HPP) on the survival of a 5-strain rifampicin-resistant cocktail of *Listeria monocytogenes* in Queso Fresco (QF) was evaluated as a postpackaging intervention. Queso Fresco was made using pasteurized, homogenized milk, and was starter-free and not pressed. In phase 1, QF slices (12.7 × 7.6 × 1 cm), weighing from 52 to 66 g, were surface inoculated with *L. monocytogenes* (ca. 5.0 log10 cfu/g) and individually double vacuum packaged. The slices were then warmed to either 20 or 40°C and HPP treated at 200, 400, and 600 MPa for hold times of 5, 10, 15, or 20 min. Treatment at 600 MPa was most effective in reducing *L. monocytogenes* to below the detection level of 0.91 log10 cfu/g at all hold times and temperatures. High-hydrostatic-pressure processing at 40°C, 400 MPa, and hold time ≥15 min was effective but resulted in wheying-off and textural changes. In phase 2, *L. monocytogenes* was inoculated either on the slices (ca. 5.0 log10 cfu/g; ON) or in the curds (ca. 7.0 log10 cfu/g; IN) before the cheese block was formed and sliced. The slices were then treated at 20°C and 600 MPa at hold times of 3, 10, and 20 min, and then stored at 4 and 10°C for 60 d. For both treatments, *L. monocytogenes* became less resistant to pressure as hold time increased, with greater percentages of injured cells at 3 and 10 min than at 20 min, at which the lethality of the process increased. For the IN treatment, with hold times of 3 and 10 min, growth of *L. monocytogenes* increased the first week of storage, but was delayed for 1 wk, with a hold time of 20 min. Longer lag times in growth of *L. monocytogenes* during storage at 4°C were observed for the ON treatment at hold times of 10 and 20 min, indicating that the IN treatment may have provided a more protective environment with less injury to the cells than the ON treatment. Similarly, HPP treatment for 10 min followed by storage at 4°C was the best method for suppressing the growth of the endogenous microflora with bacterial counts remaining below the level of detection for 2 out of the 3 QF samples for up to 84 d. Lag times in growth were not observed during storage of QF at 10°C. Although HPP reduced *L. monocytogenes* immediately after processing, a second preservation technique is necessary to control growth of *L. monocytogenes* during cold storage. However, the results also showed that HPP would be effective for slowing the growth of microorganisms that can shorten the shelf life of QF.

**Key words:** high-pressure processing, Queso Fresco, *Listeria monocytogenes*, microbial inactivation

**INTRODUCTION**

In the United States, fresh cheeses are manufactured from pasteurized milk to eliminate pathogens, if present, and to lower the levels of microflora that can reduce shelf life and affect the textural and sensory properties of the cheese. One of the most popular of the fresh cheeses is Queso Fresco (QF), which is a Hispanic-style cheese distinguished by its bright white texture, crumbliness, mild salty flavor and non-melting characteristics. However, its high pH and moisture content provide the ideal conditions for growth of bacteria and other microflora, which can limit its shelf life (Leggett et al., 2012). Despite the use of pasteurized milk, QF and other fresh cheeses made commercially have been subject to occasional recalls, most likely due to environmental contamination by *Listeria monocytogenes* (Lin et al., 2006; Soni et al., 2010).

Postpasteurization *L. monocytogenes* contamination likely occurs both at the surface and the interior of
the cheese due to processing steps that may involve handling of the curd and use of utensils (Sandra et al., 2004; Soni et al., 2010; Leggett et al., 2012) or during the milling step in manufacture, which is used to impart a crumbly texture to the curd (Van Hekken et al., 2012). Although larger, modern QF plants conduct manufacturing in closed vats so that the cheese has little contact with the environment, the hoops used for pressing and shaping the cheese may be packed by hand.

In a study to assess the viability of *L. monocytogenes* when introduced as an environmental pathogen, Leggett et al. (2012) demonstrated that QF inoculated with *L. monocytogenes* showed a maximum population density of 7.80 log<sub>10</sub> cfu/g of cheese after 20 d of storage with Leggett et al. (2012) demonstrated that QF inoculated with *L. monocytogenes* when introduced as an environmental pathogen, has little contact with the environment, the hoops used for pressing and shaping the cheese may be packed by hand.

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The application of pressure to a sample through a pressurized fluid via HPP is an effective method for inactivation of pathogens, such as *L. monocytogenes* and spoilage organisms under room temperature conditions, and spores at elevated temperatures, in a variety of food products such as guacamole, salsa, fruit juices, meats, and seafood (Rastogi et al., 2007; Zhang and Mittal, 2008; Simonin et al., 2012). High-hydrostatic-pressure processing is growing as a processing method or intervention technology of choice because of its demonstrated ability to economically extend shelf life and preserve the quality of food, as heat is not applied. Unlike interventions, such as antimicrobials or oils, which are usually applied to the surface of a food product, HPP operates according to the isostatic principle, in which the pressure applied to a sample through a pressurized medium such as water or oil is instantaneous and uniform throughout the sample, regardless of its volume or shape, thus inactivating microbes throughout a sample (Rastogi et al., 2007).

Although HPP is typically conducted under room temperature conditions, the temperature of the solid or liquid sample and the pressurizing fluid will increase when pressure is applied, assuming no loss of heat from the walls of the pressure chamber. This is due to the work of adiabatic compression, which interrupts the intermolecular forces of the pressurizing fluid and the sample, causing a temperature increase in both (Denys et al., 2000; Ardia et al., 2004; Knoerzer and Versteeg, 2009). The compression heating factor (∆T/∆P) is given by

\[ \Delta T/\Delta P = \beta T/\rho C_p, \]

where \( T \) is the temperature (absolute K), \( P \) is the pressure (Pa), \( \beta \) is the coefficient of thermal expansion (K<sup>−1</sup>), \( \rho \) is the density (kg/m<sup>3</sup>), and \( C_p \) is the heat capacity (J/kg·K).

In accordance with Le Chatelier’s principle, interruption of intermolecular forces by pressurization leads not only to an increase in sample temperature, which in the case of cheese, depends on composition (Hnosko et al., 2012; Van Hekken et al., 2013), but a decrease in the volume of water, which is a response by the sample to restore the various equilibria that were in operation before HPP (Huppertz et al., 2006). For cheese, adjustments in the intermolecular forces are reflected in changes in the microbiological, physicochemical, rheological, and sensory properties upon HPP treatment and after (Martínez-Rodríguez et al., 2012). High-hydrostatic-pressure processing treatment of cheese has been shown to lead to a reorganization of water molecules around the ions, changes in the amount of free and unbound water molecules (Martínez-Rodríguez et al., 2012), and more compact structures, which affect the mineral balance of cheese, enzyme interactions, and protein conformation (Knorr et al., 2006). The covalent bonds remain intact after HPP but the secondary structures are denatured and changes in the tertiary structures, maintained by the hydrophobic and ionic interactions, occur at applied pressures >200 MPa (Hendrickx et al., 1998; Chawla et al., 2011).

High-hydrostatic-pressure processing applied to QF does not significantly change its traditional properties. For QF in particular, which is known for its color and crumbliness, HPP conducted at 400 MPa resulted in QF that was more yellow compared with the control (Sandra et al., 2004). However, Van Hekken et al. (2013) found that QF made without starter culture and not pressed, treated by HPP at 200 or 400 MPa at an initial cheese temperature of 20°C, had the same color as untreated samples. Only QF treated at an initial temperature of 40°C by HPP at 200, 400, or 600 MPa was slightly more yellow than the control. Crumbliness of QF was not adversely affected with treatment at 400 MPa, a 20-min hold time, and 20°C compared with the control (Sandra et al., 2004). Van Hekken et al. (2013) also found that HPP at 200, 400, and 600 MPa and initial QF temperature of 20°C with up to a 20-min hold time resulted in a texture similar to the control. However, for QF manufactured with a starter culture and pH 5.0, HPP applied at 400 and 600 MPa for 1
to 25 min did not maintain the signature crumbliness (Hnosko et al., 2012).

High-hydrostatic-pressure processing has already been shown to be an effective option for reduction or inactivation of L. monocytogenes and spoilage microorganisms from cheese (Martínez-Rodríguez et al., 2012), but few studies have been conducted to examine the effectiveness of HPP for elimination of L. monocytogenes and other microflora on high-moisture cheeses such as QF. The extent of microbial inactivation using HPP is affected by applied pressure, initial substrate temperature, hold time, food substrate and composition, the presence of antimicrobial compounds, and the pressure resistance of the microorganisms (Smelt, 1998; Chen and Hoover, 2003; Patterson, 2005; Hayman et al., 2007). Although inactivation of microorganisms by HPP has been attributed to cell death because of loss of cytoplasmic membrane integrity, partial loss of membrane integrity does not necessarily lead to cell death but demonstrates sublethal injury to the cells (Ritz et al., 2001; López-Pedemonte et al., 2007) and accounts for the growth of cells after treatment. Hnosko et al. (2012) showed greater than 5-log reductions of Listeria innocua for HPP treatment pressures of 500, 550, and 600 MPa at hold times of 15, 3, and 1 min, respectively, but the inactivation was described as not permanent.

Thus, the objective of this study was to examine the effects of HPP temperature, pressure, and hold time on inactivation of L. monocytogenes inoculated both in and on packaged slices of QF and to follow growth of L. monocytogenes throughout refrigerated storage at 4 and 10°C. Total molds and yeasts and total aerobic mesophilic bacteria were also monitored throughout storage for select samples.

MATERIALS AND METHODS

QF Manufacture

One hundred and eighty kilograms of raw cow milk obtained from a local farm was stored overnight at 4°C and then standardized to 3.5% (wt/wt) fat content. The milk was then homogenized and pasteurized at 72°C for 15 s (Universal Pilot Plant; Waukesha Cherry-Burrell, Philadelphia, PA). Queso Fresco was manufactured in 5 separate trials according to the commercial procedure as previously described (Leggett et al., 2012; Van Hekken et al., 2013) in which starter cultures were not used and salt was added at 14.5 g of NaCl/L of milk. The salt was divided into 3 portions and added in 3 applications with 10 min of stirring between each to ensure that it was evenly distributed throughout the cheese. The cheese was not pressed. Curd yield ranged from 29 to 31 kg.

The curds were cut into approximately 5- to 10-cm-thick pieces, cooled to an internal temperature of 21°C, milled, and then packed into molds for overnight storage at 4°C (Leggett et al., 2012). The cheese was sliced into about 12.5 × 7.6 × 1.0-cm rectangular prisms, each weighing from 52 to 66 g.

High-Pressure Operations

High-hydrostatic-pressure processing experiments were conducted using an Avure model 2L-700 high-pressure food processor (Avure Technologies Inc., Columbus, OH). The pressure limit of the system is 690 MPa and the temperature operating range is from 10 to 90°C. The construction of the unit was described in Van Hekken et al. (2012). The basket for holding the cheese samples has dimensions of 24.1 cm high × 7.6 cm i.d. Two thermowells for type K thermocouples extend into the basket, with one extending approximately 5.1 cm to the top of the basket (measured as the length of the thermocouple extending from the upper closure of the HPP unit) and the other extending about 12.7 cm into the basket when the upper closure is in position. The pressure medium was filtered distilled water and was pumped through the lower closure during operation. A control system regulated heating of the pressure medium to the set-point temperature, pressurizing the vessel to the set-point pressure, setting the holding time, and controlling the vessel depressurization back to atmospheric pressure. The times to reach the pressures of 200, 400 or 600 MPa used in the experiments were 1.3, 1.9, and 2.5 min, respectively, with depressurization time of 0.5 s. The HPP unit was regulated by a control system that contains a recipe editor to define the set points of the system operation. System operation included steps for heating of the pressurizing fluid to the set-point temperature, pressurizing the vessel to the set-point pressure, setting the holding time for the chosen temperature and pressure, and vessel depressurization to atmospheric pressure. The real-time responses of pressure versus holding time and temperature versus holding time were displayed on a computer monitor.

Inoculation Procedures with Listeria monocytogenes Strains

As described previously (Leggett et al., 2012), approximately equal numbers of 5 rifampicin-resistant (Rif; 100 g/mL; Sigma Chemical Co., St. Louis, MO) strains of L. monocytogenes (MFS 53, MFS 1365, MFS 104, MFS 1363, and MFS 1394) were used in this study to inoculate both the cheese curds before molding and slicing (IN treatment) or the surface of cheese slices (ON treatment).
Preliminary experiments showed high levels of indigenous microflora in QF during an extended shelf life. To recover only the inoculated \textit{L. monocytogenes}, the indigenous microorganisms were suppressed by using selective agar supplemented with the antibiotic rifampicin. The 5-strain, rifampicin-resistant (10 mg/mL) \textit{L. monocytogenes} was obtained by streaking a portion (≤100 μL) of a semi-frozen suspension of each strain separately onto brain-heart infusion (BHI) agar and incubating overnight at 37°C. A single colony of each strain was then transferred individually into 10 mL of BHI broth and incubated at 37°C with shaking (100 rpm) for 24 h. One hundred microliters of the transfused was transferred into 10 mL of fresh BHI broth and incubated at 37°C for 18 h with shaking to attain a population of stationary-phase cells. To generate the rifampicin-resistant populations, 100-μL portions of each strain were spread-plated onto BHI agar containing 50 μg of rifampicin/mL and incubated at 37°C until colonies appeared on the plate. Then, a single colony was transferred and spread for isolation onto BHI agar containing 100 μg/mL levels of rifampicin using an inoculating loop. To prepare the cocktail, approximately equal volumes of cell suspensions of each rifampicin-resistant (10 mg/mL) strain of \textit{L. monocytogenes} were combined, and the final volume was adjusted to 200 mL with sterile 0.1% peptone water to yield ca. 5 log\textsubscript{10} cfu/mL.

**Preliminary HPP Trials**

In phase I of this study, preliminary experiments were conducted to identify the optimal temperature and pressure combinations for reducing the numbers of \textit{L. monocytogenes} inoculated on the cheese by HPP. Using the procedures described in Leggett et al. (2012), approximately 6 kg of the milled curds from each of 2 cheesemaking trials was removed from hoops, stored overnight at 4°C, and then cut into approximately 12.5 × 7.5 × 1.0-cm slices (ca. 52 to 66 g), with each slice weighing from about 52 to 66 g. Slices were individually surface inoculated with 50 μL of a semi-frozen suspension of each strain containing the 3 individually vacuum-packaged QF slices and incubated at 37°C until colonies appeared on the plate. After inoculation, individual slices of the cheeses were vacuum packaged (Ultravac; Koch Equipment LLC, Kansas City, MO) into sterile 10.1 × 20.3-cm 3-mil nylon-polyethylene bags (Prime Source Vacuum Products, San Jose, CA) and stored at 4°C until HPP treatment on the same day. Prior to each HPP treatment, 3 of the individually packaged cheese slices were placed into a second sterile nylon-polyethylene bag and then vacuum packaged.

The preliminary HPP trials were conducted with QF at 2 initial temperatures (20 and 40°C), 3 pressures (200, 400, and 600 MPa), and 4 holding times (5, 10, 15, and 20 min). Prior to being treated, the sample bag containing 3 individually vacuum-packaged QF slices was folded over twice and placed between the thermowells on the underside of the upper closure of the HPP apparatus. While holding the sample, the basket was screwed to the underside of the upper closure and then lowered into the pressure vessel. For experiments conducted at 20°C, the samples that were stored at 4°C were warmed in a water bath to 20°C before HPP. For experiments conducted at 40°C, the sample bag containing the 3 individually vacuum-packaged QF slices was warmed in a water bath for about 2 min until the center of the cheese reached approximately 40°C. The temperature of the pressurizing water was set to 40°C. After processing, the samples were stored at 4°C until assayed the following day.

**Storage of HPP-Treated QF**

The preliminary trials demonstrated that QF at an initial temperature of 20°C and treatment pressure of 600 MPa would decrease \textit{L. monocytogenes} below detection levels if on the surface of QF. In phase II of this study, in each of 3 cheesemaking trials, 8 kg of the QF milled curds were inoculated to a final concentration of approximately 5 to 7.0 log\textsubscript{10} cfu/g, mixed to distribute the inoculum throughout the curd, hand-packed into molds, and then stored overnight at 4°C as previously described (Leggett et al., 2012). On the following day, the cheese was removed from the molds and sliced as described above for the first phase of this study. These slices are referred to as the IN treatment slices. The remaining 8 kg of milled curds was prepared as described above in phase I for the ON treatment. Both the IN and ON slices were vacuum packaged as described above. Next, the IN and ON slices with an initial temperature of 20°C were treated at 600 MPa for holding times of 3, 10, and 20 min. After treatment, the slices were stored at 4 or 10°C and sampled on d 0, 7, 14, 28, 42, and 60.

**Enumeration of \textit{L. monocytogenes}**

The sampling and enumeration procedures for QF inoculated with \textit{L. monocytogenes} have been described previously (Leggett et al., 2012). Briefly, each slice was weighed after opening the ethanol-sterilized packages, combined in a filter bag (model XX-C003; Microbiology International, Frederick, MD) with 75 mL of sterile 0.1% peptone water (Becton, Dickinson and Co., Franklin Lakes, NJ) and then macerated for 60 s in a stomacher (model 400C; Seward, Cincinnati, OH). Ten milliliters of the homogenous mixture was transferred to a 15-mL conical tube, serially diluted (1:10) as needed in sterile
0.1% (wt/wt) peptone water, and then spread plated onto Modified Oxford (Becton, Dickinson and Co.) agar plates plus 100 μL/mL of rifampicin (MOX™). To recover sublethally pressure-injured cells of *L. monocytogenes*, the homogeneous mixture was serially diluted in sterile 0.1% peptone water, spread plated onto BHI agar (Becton, Dickinson and Co.), and then incubated at 37°C for 2 h. After incubation, ca. 10 mL of MOX™ was overlaid on the BHI agar, and then plates were incubated at 37°C for 48 h (Kang and Fung, 1999).

All plates were incubated at 37°C for 48 h and typical *L. monocytogenes* colonies were counted manually. Bacterial numbers were expressed as the logarithm of colony-forming units per gram. The detection limit was <0.91 log₁₀ cfu/g. The extent of sublethal injury was calculated as the percentage of the raw number of cells injured (Gurtler et al., 2010):

\[
\% \text{ injury} = \left(1 - \frac{\text{raw number of cells recovered on selective media}}{\text{raw number of cells recovered on non-selective media}}\right) \times 100\%.
\]

### Enumeration of Endogenous Microorganisms

Enumeration of the endogenous microflora in pathogen-free samples of QF was performed as previously described (Renye et al., 2008). High-hydrostatic-pressure-processing-treated QF samples were analyzed after storage at 4 or 10°C for 1, 7, 28, 56, and 84 d. A 10-g QF sample was homogenized in 90 mL of sodium citrate (2%) using a stomacher (230 rpm for 2 min; model 400C; Seward) and serial 10-fold dilutions of the homogenate were prepared in 0.1% peptone water. Plate count agar (Oxoid Ltd., Basingstoke, UK) and oxytetracycline glucose yeast extract (OGYE) agar base (Difco Laboratories Inc., Detroit, MI) containing 0.1% oxytetracycline (Calbiochem, San Diego, CA) were spread plated with 100 μL of each dilution. Plate count agar plates were incubated at 30°C for 48 h and 5°C for up to 14 d; and OGYE agar plates were incubated at 25°C for 72 h. Bacterial colonies were counted on duplicate plates for each dilution and the reported counts were the averages of 3 independent QF cheeses. The limit of detection was 2.0 log₁₀ cfu/g of QF. Two representatives for each colony morphology observed were grown in BHI broth (Difco Laboratories Inc.) and used as a template for PCR amplification of the 16S rRNA gene using the ebacterial oligonucleotide primers EubA and EubB (Cottrell and Kirchman, 2000). The 1.6-kb DNA fragment was sequenced with an ABI Prism 3730 DNA analyzer (PerkinElmer Inc., Wellesley, MA). The bacterial species were identified by comparing the 16S rRNA sequences with those available in GenBank using the National Center for Biotechnology Information BLAST search program (http://www.ncbi.nlm.nih.gov/BLAST/).

### Scanning Electron Microscopy

Samples of QF were prepared in thin sections approximately 5 × 5 × 10 mm for scanning electron microscopy analysis. Squares measuring 2 × 2 mm were marked on the surfaces of select samples with a pen. These squares were surface inoculated with *L. monocytogenes* (ON treatment) and then treated by HPP at 200, 400, and 600 MPa. The samples were then placed in a covered polystyrene nonpyrogenic Petri dish (no. 353004; BD Falcon, Franklin Lakes, NJ) and 10% glutaraldehyde (catalog no. 16210; Electron Microscopy Sciences, Fort Washington, PA) in a 0.1 M imidazole reagent (IMID) buffer (catalog no. 16905; Electron Microscopy Sciences) was applied to the squares and allowed to fix for 1 h. Then, the samples were thin sliced to reduce sample volume while retaining the surface of interest on the cheese. Three milliliters of 2.5% glutaraldehyde/0.1 M IMID buffer was additionally added to the plates and allowed to fix overnight.

Samples were then washed in 10 mL of 0.1 M IMID buffer twice for 1.5 h each the next day. Samples were then processed through an ethanol dehydration series of 50% (vol/vol) ethanol twice for 2 h (Warner-Graham Co, Cockeysville, MD) and 80% ethanol 3 times for 1 h each with an overnight hold at 80%. The final 100% dehydration step was done 3 times for 2 h each. The scanning electron microscopy images were obtained using a scanning electron microscope (Quanta 200 field emission scanning electron microscope; FEI Co., Hillsboro, OR).

### Statistical Analysis

In phase II of this study, a SAS ANOVA (version 9.12; SAS Institute Inc., Cary, NC) was used to analyze the growth data to determine the effects and interactions of type of treatments (IN vs. ON), temperature (4 or 10°C), time of treatments (3, 10, or 20 min), and days of storage (0, 7, 14, 28, 42, or 60 d) using PROC MIXED. Fisher’s least significant difference (LSD) was used to test for significant differences at the *P* = 0.05 significance level.

### RESULTS AND DISCUSSION

**Phase I**

**Thermal Inactivation of *L. monocytogenes* in QF.** Little information is available on thermal inacti-
vation of bacteria in soft cheeses because heat-induced quality changes tend to occur at temperatures ≥40 to 50°C (Capellas et al., 2000; López-Pedemonte et al., 2007), which are required for inactivation. Heat treatment at 63°C for 5 min of a brined white cheese with a production procedure similar to that of QF resulted in a ca. 1.5 log10 reduction of L. innocua (Al-Holy et al., 2012). Those authors did not indicate if heat treatment affected the quality attributes of the cheese. In our study, heat treatment of the packaged slices of QF at 40°C also resulted in a ca. 1.5 log10 reduction of L. monocytogenes. However, significant whey-off was observed and the QF was found to have different textural and rheological properties compared with a control at 20°C (Van Hekken et al., 2013).

**Temperature-Pressure Inactivation of L. monocytogenes in QF.** The phase-1 studies were conducted to determine the optimal pressure-temperature-hold time combinations for inactivating L. monocytogenes in QF. Each pressurization was accompanied by an increase in QF and pressurizing fluid temperatures compared with their initial temperatures of 20 or 40°C. The increase in temperature may be attributed to adiabatic compression generated during the come-up time (i.e., the time that the target pressure was reached). The come-up times at 200, 400, or 600 MPa were 1.3, 1.9, and 2.5 min, respectively, with corresponding maximum QF temperatures of 26.3, 35.7, and 43.2°C from an initial QF temperature of 20°C. The average increase in the ΔT/ΔP was about 3.6°C/100 MPa. The maximum temperatures of QF upon pressurization at 200, 400, or 600 MPa, from the initial temperature of 40°C, were 48.7, 59.4, and 67.7°C, respectively, corresponding to an increase in ΔT/ΔP of about 4.6°C/100 MPa.

The ΔT/ΔP is a function of temperature and the physical properties of QF, as shown in Equation 1. Van Hekken et al. (2013) reported that QF contains an average of 56.1% moisture, 15.4% protein, 22.3% fat, 2.9% lactose, and 3.3% ash, with about a 2.0% salt content. It could initially be assumed that because of its high moisture content of 56.1%, the ΔT/ΔP for QF would agree with those for water, with calculated values of 2.9°C/100 MPa at 20°C and 3.4°C/100 MPa at 40°C (Harvey et al., 2004). However, fats have higher ΔT/ΔP than those for water, with reported values of up to 8.7°C/100 MPa (Rasanayagam et al., 2003). Thus, the higher ΔT/ΔP for fat would raise the overall ΔT/ΔP for cheese, as the other components have ΔT/ΔP similar to that of water (Trujillo et al., 2002). There may also be a small contribution to the higher value of ΔT/ΔP (4.6°C/100 MPa) for QF with an initial temperature of 40°C (3.6°C/100 MPa) versus 20°C due to the loss of whey associated with preheating the cheese before HPP and a loss of 2% of whey during processing, which tended to increase with increasing pressure, and would shift the distribution of the components in the cheese (Van Hekken et al., 2013).

The levels of L. monocytogenes for pressures of 200, 400 and 600 MPa, with holding times of 5, 10, 15, and 20 min for initial cheese temperatures of 20 and 40°C are shown in Figures 1a and 1b, respectively. Examination of the curves for QF treated at 200 MPa and 20 and 40°C show what is often interpreted as shoulders at each of the hold times. This pressure, with adiabatic temperature increase of 26.3°C, had no effect (P > 0.05) on inactivation of L. monocytogenes relative to the control (Figure 1a). For treatment at 200 MPa for QF with an initial temperature of 40°C, the adiabatic temperature increase of 48.7°C (Figure 1b) did not contribute to an increase in reduction of L. monocytogenes levels relative to the 1.5-log reduction that occurred upon heating the inoculated QF from 20 to 40°C before pressurization, even at the highest hold time of 20 min.

At 400 MPa (Figure 1a), with QF adiabatic temperature increase of 35.7°C, a log-linear decrease in counts with holding time was noted, with an overall 1.78-log reduction in L. monocytogenes numbers at a hold time of 20 min (Figure 1a). The synergistic effects of pressure at 400 MP and adiabatic temperature increase of 59.4°C are shown in comparison in Figure 1b, with a log-linear decrease in L. monocytogenes counts of 2 log at a 5-min hold time. When the hold time was increased to 15 min, a tailing effect was observed, with only an additional 1.25-log reduction in pathogen numbers, reaching below the minimum level of detection of 0.91 log. At 600 MPa, at initial QF temperatures of 20 and 40°C (Figures 1a and 1b), L. monocytogenes numbers were reduced by 4.6 and 3.25, with final counts reaching the minimum level of detection at the lowest hold time of 5 min.

Although HPP processing conditions of 200 MPa were generally ineffective in reducing L. monocytogenes on QF, pressures of at least 300 MPa applied for 5 min with temperatures in the range from 5 to 25°C have been found to reduce yeast and molds, as these microorganisms are less resistant to HPP than gram-positive bacteria such as Listeria spp. (Martínez-Rodríguez et al., 2012). The susceptibility of L. monocytogenes to pressure, measured by the numbers of sublethally injured cells, was not determined for QF at an initial temperature of 20°C in this phase of the current study. For QF with an initial temperature of 40°C, sublethally injured L. monocytogenes were not detected when QF was treated at 400 MPa and a hold time ≥15 min, or at
600 MPa and all hold times. Thus, these temperature-pressure combinations were lethal to *L. monocytogenes* on QF.

For most bacteria, higher pressures and temperatures and longer hold times are necessary for achieving reductions greater than 5 log in cheese (Martínez-Rodríguez et al., 2012.) However, in the case of QF, temperatures ≥40°C and the pressure can negatively affect the sensory properties, causing whey losses (Capellas et al., 2000; Van Hekken et al., 2013), subtle textural changes (Sandra et al., 2004; Van Hekken et al., 2013), or severe changes in texture. These negative effects have been reported for a low-pH QF made using starter cultures, changing the signature properties of the cheese (Hnosko et al., 2012).

**Microstructure.** In addition to the use of nonspecific media, scanning electron microscopy is useful for indicating if microorganisms have been sublethally injured by physical treatments such as HPP (Ritz et al., 2001). Scanning electron micrographs at 5,000× magnification, showing *L. monocytogenes* on the surface of QF sections with an initial temperature of 20°C, after treatment at 200, 400, or 600 MPa, for a hold time of 5 min are shown in Figure 2a. Prior to treatment, the surfaces of *L. monocytogenes* cells were smooth. The cell surface remained smooth after treatment at 200 MPa, but showed surface damage after treatment at 400 MPa. After treatment at 600 MPa, the cell showed increased surface damage in the form of bud scars and appeared to shrink. At a hold time of 20 min (Figure 2b), cells of *L. monocytogenes* appeared to be intact after treatment at 200 MPa, but bud scars were evident on the cell surface, indicating that the membrane was disrupted or that a loss of membrane integrity occurred. Although the disruption was apparent, it was not lethal to the cells, as the decrease in pathogen levels at this pressure compared with the control (Figure 1a) was considered not significant (*P* > 0.05). According to Ritz et al. (2001), the number of buds is about proportional to pressure. At 400 and 600 MPa, bud scars were also evident on the intact cells and a few cells appeared to have lost their morphological characteristics. However, due to the qualitative nature of this approach it was difficult to discern if a difference existed in the number of cell disruptions at 400 versus 600 MPa. Figure 1a shows that pathogen levels were reduced to below the detection level after 5 min of HPP treatment at 600 MPa, whereas surviving cells were apparent after treatment at 400 MPa for 20 min. The differences in temperature due to adiabatic compression at 400 MPa (i.e., 35.7°C) and at 600 MPa (i.e., 43.2°C) were not likely to account for the differences in *L. monocytogenes* numbers after a hold time of 20 min.

The substrate, composition, and water activity may affect the response of microorganisms to HPP treatment (Considine et al., 2008; Martínez-Rodríguez et al., 2012). The changes in the QF substrate as the applied pressure was increased are apparent in Figures 2a and 2b. As described previously (Van Hekken et al., 2013), the increasing pressures from 200 to 600 MPa resulted in a progressive fusing of the protein matrix with individual micelles fusing into thicker strands. These changes were accompanied by increasing wheying-off observed on the surface of the cheese. It is doubtful that the increased microbial susceptibility to HPP at the higher pressures was directly affected by changes in the substrate relative to the high pressures applied because *L. monocytogenes* was on the surface and not the interior of QF. However, the increases in pressure altered the substrate, thereby forcing whey from the interior of the cheese to the surface. This alteration
Figure 2. Scanning electron micrographs of Queso Fresco (QF) surface inoculated with ca. 4.5 log_{10} cfu of *Listeria monocytogenes*/g of QF and then processed at 200, 400, and 600 MPa, respectively. Micrographs are at 5,000× magnification. The scale bars represent 5 μm. At 200, 400, and 600 MPa, QF temperatures during pressurization were 26.3, 35.7, and 43.2°C, respectively: (a) Hold time of 5 min; (b) hold time of 20 min.
may affect inactivation of *L. monocytogenes*, as low water activity environments support the survival of microorganisms due to reduced permeability of the cell membrane (Smelt, 1998; Martínez-Rodríguez et al., 2012).

Scanning electron microscopy was also conducted for QF at 20°C that was surface inoculated with *L. monocytogenes* and HPP treated at 600 MPa with hold times of 3, 5, 10, 15, and 20 min (not shown). The results showed damage to *L. monocytogenes* cells even at the lowest hold times, with no apparent differences in the extent of damage with increasing hold time. Scanning electron microscopy of QF at 40°C was attempted but *L. monocytogenes* cells were not found on the surface after treatment at any of the pressures.

Ritz et al. (2001) observed that pressurized cells did not behave as live or dead cells, even though they gave the appearance of total inactivation through cell count determination. They suggested that reversible damage was a possibility due to the variability in the extent of injury and cell heterogeneity. This observation suggests that assessment of HPP for food safety purposes should be followed by storage studies over the shelf life of a food product to ensure that populations of pathogens, such as *L. monocytogenes*, are inactivated and not injured.

**Phase II**

**Storage Study of *L. monocytogenes* in and on QF Slices.** The storage study was conducted using QF samples with an initial temperature of 20°C that were treated at 600 MPa and held for 3, 10, or 20 min at a temperature of 43.2°C following adiabatic compression. This pressure-temperature combination was found to be the most effective for inactivation of *L. monocytogenes* on the surface of QF while preserving the sensory and textural properties of QF (Van Hekken et al., 2013). The initial levels of *L. monocytogenes*-inoculated IN and ON slices of QF were 6.91 ± 0.15 and 4.85 ± 0.21 log_{10} cfu/g, respectively, to simulate environmental contamination after pasteurization (Leggett et al., 2012). The viability of *L. monocytogenes* following treatment at 600 MPa for 3, 10, or 20 min was followed by storage for 60 d at a refrigeration temperature of 4°C and abuse temperature of 10°C. The *L. monocytogenes* levels after HPP correspond to a time of 0 in Figures 3a to 3f. High-hydrostatic-pressure processing treatment at 600 MPa with a hold time of 3 min showed an average 3.31 ± 1.20 log reduction for the IN treatment QF and 4.12 ± 0.81 log reduction for the ON treatment.

For the IN treatment cheese, *L. monocytogenes* increased from 3.60 ± 1.05 log_{10} cfu/g on d 0 of storage at 4°C to 8.08 ± 0.20 log_{10} cfu/g at 60 d of storage (Figure 3a). In comparison, *L. monocytogenes* for the ON treatment increased from 0.73 ± 0.60 to 8.40 ± 0.19 log_{10} cfu/g (Figure 3b), showing no difference (*P > 0.05) in pathogen viability between treatments, even though the initial counts of *L. monocytogenes* were greater for the IN samples on d 0. Calculations of the percentage of injured cells for the IN and ON treatments for up to 2 wk of storage are shown in Table 1 for 1, 7, and 14 d of storage at 4°C or 1 d after 10°C. Regardless of treatment, at 4°C, most of the cells of *L. monocytogenes* were injured. A decrease in the percentage of injured cells was noted as storage time extended to 2 wk. By d 14 of storage at 10°C (Figures 3a and 3b), little difference (*P > 0.05) was observed in *L. monocytogenes* counts for QF or lag time (*P < 0.05) for either the IN and ON treatments, as the higher temperature facilitated growth of the pathogen. By 28 d, the value of the *L. monocytogenes* counts for the IN treatments at both storage temperatures began to converge but for the ON treatments, the data at 4 and 10°C did not converge until 42 d, in both cases indicating that HPP followed by storage at 4°C cannot control growth of the pathogen. Overall, although QF was treated at 600 MPa for a 3-min hold time, *L. monocytogenes* numbers were within 1.0 log after 42 d of the average of 8.24 ± 0.41 log_{10} cfu/g noted for QF at 60 d of storage.

Increasing the hold time to 10 min after pressurization at 600 MPa resulted in a decrease in *L. monocytogenes* to below detection levels for the IN and ON treatments at d 0 (Figures 3c and 3d, respectively). During storage at 4°C, growth of *L. monocytogenes* was observed by d 7 for the IN treatment but it appeared that there was a 7-d lag in growth for the ON treatment. Most of the cells for the IN (97.2%) and the ON (96%) treatments were injured 1 d following HPP treatment. After 2 wk, there was a decrease in the number of injured cells (Table 1) for the IN treatment and, as indicated in Figure 3c, a steady growth of *L. monocytogenes* was observed; but for the ON treatment, a high percentage of cells (97.7%) remained injured after 2 wk, with growth of *L. monocytogenes* delayed for 1 wk (Figure 3d). Little difference was observed in the trends observed for viability of *L. monocytogenes* for the IN or ON treatments when stored at 10°C.

Increasing the hold time to 20 min after pressurization to 600 MPa resulted in a decrease in *L. monocytogenes* levels to below the detection limit for both the IN and ON treatments (Figures 3e and 3f). An increase in the lethality of the process compared with the 3- and 10-min hold times was indicated by the 0% injured cells for the IN treatment and 29% for the ON treatment (Table 1). Regardless of the apparent lethality for the IN process on d 0, growth of *L. monocytogenes* occurred by 2 wk, after an apparent lag time of 1 wk (Figure 3e).
Figure 3. Growth of *Listeria monocytogenes* in storage at 4 or 10°C on Queso Fresco (QF) slices after high-hydrostatic-pressure processing (HPP) treatment at 600 MPa, initial cheese temperature of 20°C, and hold times of (a) 3, (c) 10, and (e) 20 min for the IN treatment (*L. monocytogenes* inoculated in QF curds, formed into a cheese block and sliced), and (b) 3, (d) 10, and (f) 20 min for the ON treatment (*L. monocytogenes* inoculated on QF slices). The error bars represent SD.
Injured cells were detected after 7 and 14 d. Although the ON process showed that 29% of cells were injured on d 0 and cells were injured on d 7 and 14, no growth of \textit{L. monocytogenes} occurred until d 28 (Figure 3f). For the IN treatment and storage at 10°C, the maximum growth of \textit{L. monocytogenes} was 7.51 ± 0.10 log\textsubscript{10} cfu/g and was achieved at 28 d. The ON treatment stored at 10°C showed maximum growth of \textit{L. monocytogenes} of 5.82 ± 2.62 log\textsubscript{10} after 14 d and was not significantly different than the ON treatment (\textit{P} < 0.05).

In general, as HPP hold time was increased to 20 min, a decrease in the numbers of injured cells occurred because of the increased lethality of the treatment. At 4°C, the growth rate for the IN treatment was approximately twice that of the ON treatment and a 7-d delay in growth was not observed for the IN treatment, except for the 20-min hold time. For the ON treatment, lag times in growth of \textit{L. monocytogenes} at 4°C were observed after the 10- and 20-min hold times, indicating that HPP treatment for \textit{L. monocytogenes} on the surface of QF appeared to be more effective than that for cells on the interior, showing the sensitivity of the pathogen to subtle changes in environment even within the same sample during processing. As indicated in Van Hekken et al. (2012), unpressed QF loses approximately 1.2 to 1.7% moisture because of syneresis (wheylogging) during storage. With HPP, the moisture content of the interior of QF decreases further, with increases in pressure and hold time resulting in visible wheylogging on the surface of packaged QF, due to structural changes shown in Figure 2. The increased moisture at the surface of the cheese enhances injury and inactivation (Smelt 1998; Martínez-Rodríguez et al., 2012). The pH and mineral content shifts throughout the cheese were not followed but may have also contributed to the differences in lag times between the IN and ON treatments, as described in Considine et al. (2008).

**Levels of Indigenous Bacteria.** As reported previously, the total aerobic mesophilic counts for pathogen-free QF cheeses were 3.5 ± 0.1 and 3.4 ± 0.3 log\textsubscript{10} cfu/g after 1 d, and 6.3 ± 0.8 and 7.7 ± 0.5 log\textsubscript{10} cfu/g after 84 d, when stored at 4 and 10°C, respectively (Leggett et al., 2012). High-hydrostatic-pressure processing treatment at 600 MPa for 3 min or 10 min decreased the mesophilic counts to below the detection level of 2 log\textsubscript{10} cfu/g (Figure 4A, d 1). These results are in agreement with previous reports that showed that HPP treatment at >500 MPa effectively reduced the number of microbial contaminants in Cheddar cheese (O’Reilly et al., 2000) and the number of starter lactococci in fresh curd cheese (Daryaei et al., 2006). Mesophilic counts remained at or below the level of detection for up 56 d in most HPP-treated QF samples stored at 4°C, with the exception of one 3-min HPP-treated cheese showing 3.2 log\textsubscript{10} (56 d) and one 10-min HPP-treated cheese showing 3.4 log\textsubscript{10} cfu/g (28 d). Following 84 d of storage at 4°C, the indigenous microflora was not detected in 2 of the 3 QF samples that were treated for 10 min, with the third cheese having 4.2 log\textsubscript{10} cfu/g. All QF samples that were HPP treated for 3 min had bacterial growth at 84 d, with an average of 4.1 ± 1.5 log\textsubscript{10} cfu/g, which was 2 log lower than the 6.3 ± 0.8 log\textsubscript{10} cfu/g observed in nontreated QF samples (Leggett et al., 2012). A delay in the recovery and growth of aerobic mesophiles was previously reported for fresh cheeses stored at 4°C after HPP treatment at 300 or 400 MPa for 5 min (Evert-Arriagada et al., 2012). However, HPP treatment at ≤400 MPa did not decrease the initial load of aerobic mesophiles in fresh cheese and the bacterial load was similar to the control cheese by d 21 of storage (Evert-Arriagada et al., 2012), suggesting that the higher pressure used in the current study offers a significant advantage for increasing the shelf life of cheeses stored at 4°C.
For QF samples stored at 10°C, the mesophilic counts remained at or below the level of detection on d 7 (Figure 4B). By d 28, bacterial growth was observed in 5 of the 6 QF samples, with an average of 4.4 ± 0.4 and 4.3 ± 0.7 log_{10} cfu/g for the 3- and 10-min HPP-treated cheeses, respectively. The bacterial load increased after 84 d of storage to 5.4 ± 1.7 and 6.0 ± 0.8 log_{10} cfu/g in the 3- and 10-min HPP-treated QF samples, but remained 2 log_{10} lower than the untreated sample (7.7 ± 0.5 log_{10} cfu/g stored at 10°C). These results emphasize the importance of the storage temperature following HPP treatment, as the bacterial load in both HPP-treated cheeses stored at 10°C for 28 d was similar to what was observed in the 3-min HPP-treated QF stored at 4°C for 84 d. Similar results were reported for HPP-treated fresh cheeses, where the bacterial load increased more rapidly when stored at 8 versus 4°C (Evert-Arriagada et al., 2012). Previously, we reported the identification of gram-negative contaminants, including *Escherichia hermannii*, *Pseudomonas aeruginosa*, and *Enterobacter asburiae*, in QF samples stored for 56 d at 4°C, or 84 d at 10°C (Leggett et al., 2012). However, these bacterial species were not isolated from any of the HPP-treated QF samples, demonstrating the effectiveness of this technique against these contaminants. The bacterial species isolated from the HPP-treated QF samples included *Enterococcus faecalis* (99% identity; GenBank accession no. HQ721277), *Enterococcus durans* (97%; GenBank accession no. HQ6778261), *Bacillus pumilus* (99%; GenBank accession no. HQ334985), *Bacillus simplex* (99%; GenBank accession no. JF496520), *Paenibacillus glucanolyticus* (99%; GenBank accession no. AB073189), and *Micrococcus luteus* (92%; GenBank accession no. EU379264), which have all been previously identified as contaminants within milk and dairy foods (Ogier et al., 2004; Foulquié Moreno et al., 2006; Renye et al., 2008; Ivy et al., 2012). The identification of gram-positive bacteria within the stored HPP-treated cheeses was expected, as they are generally considered more tolerant of high pressure than gram-negative bacteria, molds, and yeasts (Cheftel, 1995). Bacterial spores have been reported to withstand pressures of up to 1,000 MPa (Cheftel, 1995), which may explain why the *Bacillus* and *Paenibacillus* species persisted in HPP-treated QF samples. Due to the pressure tolerance of spores, other studies have shown that the use of additional hurdles, such as nisin (López-Pedemonte et al., 2003), with HPP treatment may improve the potential for inactivating spores within foods. The identification of *E. faecalis* and *E. durans* in the HPP-treated QF samples from the current study, along with previous reports that *Enterococcus hirae* was highly tolerant of pressure treatments up to 500 MPa (Szczawiński et al., 2003; Fonberg-Broczek et al., 2005) suggests that enterococci, in general, have a high pressure tolerance. *Micrococcus luteus* was also reported to have a high pressure tolerance with pure cultures having a similar tolerance level to spore formers *Bacillus cereus* and *Bacillus subtilis* (Arroyo et al., 1999). In a recent study, both enterococci and *Micrococcaceae* were shown to persist in an
HPP-treated (at 600 MPa) raw milk cheese stored at 4°C. Survival was dependent on the time of HPP treatment, as only Micrococccaeae were observed to survive for up to 240 d when pressure was applied after 5 wk of ripening (Calzada et al., 2013). In the current study, storage of QF at 4°C was required to extend the shelf life of 3-min HPP-treated cheese to 56 d, and 10-min HPP-treated cheese to 84 d.

High-hydrostatic-pressure processing treatment for either 3 or 10 min was also sufficient to prevent the growth of psychrotrophic bacteria, molds, and yeasts in QF samples stored at either 4 or 10°C for up to 84 d. In untreated QF samples, psychrotrophic bacteria, including Stenotrophomonas spp., Pseudomonas fluorescens, and Enterobacter aerogenes, were detected at 56 d (4.5 log_{10} cfu/g) when stored at 4°C, and 84 d (5.2 log_{10} cfu/g) when stored at 10°C; and yeast and molds (growth on OGYE agar) were detected by 28 d (2.7 log_{10} cfu/g) and 56 d (2.4 log_{10} cfu/g) when stored at 4 and 10°C, respectively (Leggett et al., 2012). The use of 600 MPa of pressure may be essential for inhibiting the growth of these microorganisms, as a previous study reported the recovery and growth of psychrotrophic bacteria, molds, and yeast in cheeses stored at 4 and 8°C after HPP treatment at 400 MPa (Evert-Arriagada et al., 2012).

Cost Analysis. In commercial operations, food processors prefer using the shortest hold times possible to maximize production and preserve food quality. In this study, HPP conducted at 600 MPa for QF at an initial temperature of 20°C, for hold times ranging from 3 to 20 min, was found to reduce L. monocytogenes by up to 7 log_{10} cfu/g, with repair and then growth of injured cells occurring from 7 to 28 d after treatment. The same treatment conditions (HPP with either a 3- or 10-min hold times) were effective in preventing growth of psychrotrophic bacteria, yeasts, and molds for QF stored at either 4 or 10°C for up to 84 d (Figure 4), making it an effective treatment for extending the shelf life and quality of this cheese.

To estimate the costs of HPP for commercial operations, it was assumed that for a plant processing approximately 18,180 kg/d of packaged QF, it could be processed at the plant or shipped to a contract facility using a system such as the Avure QFP 350L-600 (Avure Technologies Inc.), which has a high throughput rate. The costs of HPP depend on the pressure and hold time, with a lower hold time favoring increased production rate. For the commercial system above with an assumed 4.1-min loading and unloading time and a 3-min hold time, 7.71 cycles/h would be possible when processing 1,905 kg/h. With a 10-min hold time and the same time for loading and unloading, only 3.83 pressurizing cycles/h would be possible with a throughput of 959 kg/h.

The detailed costs for the commercial HPP system are shown in Table 2. Only the costs of the HPP treatment are given and do not include the costs per kilogram of making or packaging the cheese. Regardless of the cycle time, the capital costs for the HPP system are $2,200,000 and an additional $2,200,000 is assumed for installation, required infrastructure, the building, and auxiliary equipment, for a total capital cost of $4,400,000. Annual depreciation is $440,000. To calculate the operating costs, production of 5,500 h/yr are assumed, with annual production of 10,476,760 kg/yr for HPP applied for 3 min and 5,275,532 kg/yr for HPP applied for 10 min.

High-hydrostatic-pressure processing requires 2 operators/shift at $40.00/h, bringing annual labor charges to $440,000. Utility charges include costs of $4.52/h for chilled water at 16°C and a flow rate of 83.5 kg/min, with unit costs of $0.90/1,000 kg. Electric power was available at 50% of peak power (370 kVA peak), or an average 185 kW at a unit cost of $0.07/kWh, resulting in costs of $12.95/h. Annual utility costs are $96,066/yr. Maintenance and part replacement costs were assumed as 8% of the total capital costs. Administrative costs were assumed as 2% of capital costs at $88,000. The total annual costs for 3 or 10 min of pressurization at 600 MPa were $1,416,066. Unit costs per kilogram for pressurizing packages of QF at 600 MPa for 3 min are $0.13/kg and for 10 min are $0.26/kg. This is within the range reported in Rastogi et al. (2007) of $0.05 to $0.50/kg, with lower costs typically associated with thermal processing of foods.

CONCLUSIONS

High-hydrostatic-pressure processing of QF at 20°C, a maximum pressure of 600 MPa, and hold time of 20 min was effective for the immediate elimination of L. monocytogenes inoculated in the curds or on the surface of QF slices. However, growth of the pathogen resumed after a lag time, which differed depending on whether L. monocytogenes was inoculated in (7 d) or on the QF slices (28 d). High-hydrostatic-pressure processing in conjunction with antimicrobials may help limit the growth of L. monocytogenes during 4°C storage. Adjustments in pH, salt content, or water activity of QF may also help limit growth. However, applying the recommendations of Bolton and Frank (1999) for QF formulations that would have a <5% probability of L. monocytogenes growth would require a salt content ranging from 5 to 11%. High salt content would undoubtedly affect the properties and flavor of the cheese (Leggett
et al., 2012). Higher HPP pressures may be effective in eliminating *L. monocytogenes*, but would likely be cost prohibitive. Even though HPP did not eliminate *L. monocytogenes* under the conditions of this study, it was effective in eliminating the various microorganisms that can limit the shelf life of QF. This suggests that, at the conditions of this study (600 MPa, initial QF temperature of 20°C, and either a 3- or 10-min hold time), HPP is a promising postpackaging process that targets spoilage microorganisms throughout the cheese and not just at the surface. This would be an advantage over the use of antimicrobials, which would have to be applied throughout the cheese curd and possibly would alter the flavor of the cheese.

**ACKNOWLEDGMENTS**

The authors thank Douglas Soroka for assistance with the scanning electron microscopy studies, Sudarsan Mukhopadhyay for assistance, Andrew McAloon for cost studies, Winnie Yee (all from the US Department of Agriculture, Agricultural Research Service, Eastern Regional Research Center, Wyndmoor, PA) for assistance with the figures, and Nana Farkye (Dairy Products Technology Center, California Polytechnic State University, San Luis Obispo) for advice on the cheese-making procedure. This research was funded in part by Dairy Management Inc. and administered by the Dairy Research Institute (Rosemont, IL) through Agreement No. 58-1935-9-908.

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