Effects of natural antimicrobials with modified atmosphere packaging on the growth kinetics of *Listeria monocytogenes* in ravioli at various temperatures

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Abstract

The objective of this study was to investigate the antimicrobial effects of cultured sugar/vinegar (CSV) blend and nisin to control the risk of *Listeria monocytogenes* in ready to cook (RTC) ravioli. Ravioli dough was prepared with 0.1, 0.3, 0.5, 1% CSV blend and 0.1, 0.2, and 0.3% nisin. Inoculated spinach or artichoke raviolis with 2.0 $\pm$ 0.5 log cfu/g of *L. monocytogenes* were packed aerobically or using modified atmosphere packaging (MAP), and then stored at 4, 10, 17, and 24°C for 60 days. Growth kinetic parameters of the observed data fit well to the Baranyi equation. Ravioli with spinach filling materials yielded a higher risk than that with artichoke. *L. monocytogenes* was able to survive in ravioli with artichoke, but did not grow. The addition of 1% CSV blend or 0.3% nisin in spinach ravioli with the combination of MAP effectively controlled the growth of *L. monocytogenes* at the temperature below 10°C. The organoleptic quality of spinach ravioli was not also affected by the application of 1% CSV blend. Therefore, the CSV blend can be recommended to improve the microbial safety and quality of natural RTC ravioli at retail market.

Practical applications

The risk of ravioli was affected by the filling materials of ravioli at retail market. Addition of 1% cultured sugar/vinegar blend in dough substantially contributes to the extension of shelf-life of MAP spinach ravioli. Classification and regression tree analysis results indicate that refrigeration temperature is the main control factor to affect lag time and growth rate, while packaging method is critical for maximum population density.

1 | INTRODUCTION

With the increase of consumer demand for foods with more convenient, affordable and easy-to-cook portions, a wide range of new products, such as chilled/filled pasta, including ravioli, have been introduced to the market. According to the Euromonitor International (2012), the market for chilled/filled pasta, such as ravioli, grows faster than other product markets with the total sales worth U.S. $348.8 million by 2016. As consumption of these foods increases, safety of refrigerated/filled ravioli at retail market should be ensured. Manufacturing environment has been linked to many cases of foodborne listeriosis and frozen ravioli with spinach was recalled for potential *Listeria* contamination (Food Safety News, 2015). Although the chilled products are cool-processed, *L. monocytogenes* is frequently able to survive and grow even under refrigeration temperature (Larson, Johnson, & Nelson, 1999). *L. monocytogenes* is widely distributed in nature and causes serious concerns to food industries and regulatory agencies due to its high hospitalization rate (94%) and fatality rate (Warriner & Namvar, 2009).

Several methods are being employed to control *Listeria* in food industries, mostly by adapting antimicrobials. In recent years, consumers are seeking natural or organic foods (Organic Trade Association, 2015). Around 73% of all U.S. households occasionally purchase organic...
foods (Organic Trade Association, 2009). Thus, to emphasize the safety of used ingredients, producers are seeking clean-label alternative ingredients with less "chemical sounding" names. Cultured sugar and vinegar blend (CSV blend, Purac verdad NV55), a natural alternative to lactic acid salt compound, is generally regarded as safe and can be added to food products during processing. A natural blend of CSV consists of fermentation with specially curated cultures, such as sugars, organic acids, peptides, and aromas. The reagent is designed to have a high level of antimicrobial effect of 3% CSV blend on behavior of Campylobacter jejuni and Salmonella Typhimurium in chicken breast was also reported (Park, Hong, & Yoon, 2014).

In the present study, we investigated the antimicrobial effect of CSV blend on the growth control of L. monocytogenes in raviolis filled with spinach or artichoke and compared to that of nisin. Combined effects of modified atmosphere packaging (MAP) and storage temperature as hurdle techniques at retail market were also evaluated.

## 2 | MATERIALS AND METHODS

### 2.1 | Bacterial culture for inoculation study

L. monocytogenes strains (ATCC 19111, ATCC 19115, and ATCC 15313) were purchased from the Korean Culture Center of Microorganisms (KCCM, Seoul, Korea) and maintained at −80°C with tryptic soy broth (TSB, Difco, Sparks, MD) with 0.6% yeast extract (TSBYE) (Oxoid, Basingstoke, United Kingdom) in beads. For each experiment, each stock culture was thawed and inoculated in 10 ml of TSBYE, followed by incubation at 37°C for 24 hr. Viable cell counts of L. monocytogenes ranged between 8.5 and 9.0 log cfu/ml after incubation. An aliquot of 1 ml of the initial culture in stationary phase was transferred into 9 ml of 0.85% NaCl, which was serially diluted. A mixture of three L. monocytogenes strains was prepared before inoculation into the ravioli samples.

### 2.2 | Preparation and inoculation of ravioli

As antimicrobial agents for ravioli dough, a natural blend of CSV (Purac verdad NV55) and nisin were purchased from Purac (Lincolnshire, IL) and Sigma-Aldrich (St. Louis, MO), respectively. Imported durum semolina flour (Miller milling company, MN) was purchased from samjinFS (Kimpo-si, Gyeonggi-do, Korea), and then ravioli dough was prepared with flour, water, and egg (7:2:1 ratio) by an electronic mixer (K45SS, Kitchen aid, MI).

Spinach or artichoke base materials were provided from Giloy ravioli manufacturer (Pulmuone Co., CA). The CSV blend was uniformly added to the spinach or artichoke ravioli dough at the concentration of 0.1, 0.3, 0.5, or 1%, while nisin was added to only spinach ravioli dough at the concentrations of 0.1, 0.2, or 0.3%. Control was prepared without antimicrobial agents. The dough samples were formed into regular ravioli shape (8.5 ± 0.5 g) and then filled with 18 g of spinach or artichoke base materials. The prepared ravioli were then heated at 121°C for 10 min to remove any background microflora. After cooling, the ravioli were transferred into petri dishes (9 × 2 cm). The surface of each ravioli was uniformly inoculated with 0.1 ml of the diluted cultures of L. monocytogenes mixture using a sterile repeater pipette to reach the target inoculation levels (2.0 ± 0.5 log cfu/g). The inoculated spinach ravioli were packaged aerobically or in modified atmosphere with gas mixture (0% O2, 0.3% CO2, and 99.7%) (Composite Deoxidation Desiccant, Lipmen, Incheonm, Korea). The inoculated artichoke ravioli were also packed under MAP. Gas-tight wrapper type bags filled with a deoxidizer (PPEC Eumsung Fresh Noodle Co., Ltd., Eumsung, Korea) were used for MAP. The packaged samples were separately stored at 4, 10, 17, and 24°C for 60 days.

### 2.3 | Enumeration of L. monocytogenes

At the specific time intervals during storage, each sample was homogenized (Bag-Mixer 400, Interscience, Paris, France) in 90 ml of 0.85% NaCl (vol/vol) 2 min. One milliliter of the homogenized sample was diluted with 9 ml of 0.85% NaCl (vol/vol) and 0.1 ml aliquots of two dilutions of each sample was spiral plated (Automatic spiral plater, Interscience, Saint Nom, France) in duplicate on PALCAM agar (Oxoid, Basingstoke, United Kingdom) for L. monocytogenes and then incubated aerobically at 36°C for 48 hr. The colonies on the duplicate plates of each sample were counted by an automated colony counter (Scan 300, Interscience, Saint Nom, France). The results were expressed as log cfu/g and each experiment was repeated twice.

### 2.4 | Growth kinetics of primary modeling for L. monocytogenes in ravioli

Growth representing viable cell counts (log cfu/g) of L. monocytogenes as a function of time was iteratively fit to the Baranyi equation using the DM Fit 3.5 curve-fitting program (Institute of Food Research, Norwich, United Kingdom). The equation used was as follows (Baranyi & Roberts, 1994):

\[
y = y_0 + \frac{\mu_{\text{max}}}{\ln(10)} A \ln \left( \frac{1}{1 + \frac{q_0}{\ln(1 + \frac{1}{q_0})}} \right)
\]

\[
A = t + \frac{1}{\mu_{\text{max}}} \ln \left( \frac{\ln 1 + \frac{1}{q_0}}{1 + \frac{1}{q_0}} \right)
\]

where \(y\) is the logarithm of the cell numbers (log cfu/g), \(y_0\) is the initial cell number, \(\mu_{\text{max}}\) is the final cell number, \(A\) is the time variable, \(\mu_{\text{max}}\) is the specific growth rate (SGR; log per day), \(q_0\) is the physiological state of the inoculum; \(t_{\text{lag}}\) is the lag time (LT); and \(t\) is the sampling time. The goodness of fit of the data was evaluated based on the coefficient of determination (\(R^2\)). Three parameters, namely, LT, SGR, and maximum population density (MPD) were calculated from the equations described by Baranyi and Roberts (1994) and used for evaluation criteria.

### 2.5 | Secondary modeling for the effect of temperature on the growth kinetics of L. monocytogenes

Response surface equations as a function of temperature and the concentration of the CSV blend were developed for LT, SGR, and MPD of
L. monocytogenes in spinach ravioli with MAP by multiple regressions using the SAS (V 9.3) General Linear Models Procedure:

\[ \ln y = a_0 + a_1 A + a_2 B + a_3 A \times B + a_4 A \times X + a_5 B \times X + e \]  

where \( \ln y \) is the natural logarithm of the modeled growth parameters (LT, SGR, and MPD), \( A \) is the temperature, \( B \) is the concentration of the CSV blend, \( a_0 \) to \( a_5 \) are regression coefficients, and \( e \) is the random error.

### 2.6 Sensory evaluation

A triangle test was performed to determine whether a significant flavor change occurred in spinach ravioli with 1% CSV blend. The minimum number of panelist needed for the test was determined from the table of significant test for triangle test (Kim, Kim, Sung, & Lee, 1993). To be >95% certain (\( \beta = 0.05 \) and \( \alpha = 0.05 \)) that no more than 50% (\( \mu_d = 50\% \)) of consumers would be able to detect a difference if CSV the blend was applied to spinach ravioli, the required number of panelists is \( n = 19 \). In the present study, 40 trained panelists from the department of research and development at Pulmuone Co., Ltd. participated in sensory evaluation. Spinach ravioli with 1% CSV blend were compared with two controls without antimicrobial treatment. Three samples were randomly numbered with 3-digit codes and placed in a random order. Two of three samples were identical and the other was different. The panelists were asked to state which product they believed was the odd one, which was different based on appearance, smell, taste, etc. Testing was conducted in individual booths in the Pulmuone R&D center. The panelists were equally spaced throughout the room and instructed not to speak to one another during the test.

### 2.7 Statistical analysis

Three growth kinetics parameters for L. monocytogenes—namely, SGR, LT, and MPD were used to analyze the data. ANOVA model was applied to analyze the effect of multiple treatments for each of the factors: temperature (4, 10, 17, 24°C) and concentration of antimicrobials (CSV blend: 0.1, 0.3, 0.5, or 1%, nisin: 0.1, 0.2, or 0.3%). For each ANOVA, complete randomized block design was used by setting the other factor combinations as a block using statistical analysis system SAS V 9.3 (SAS Institute Inc., Cary, NC). R package (version 3.1.2) was also used for the classification and regression tree (CART) (Breiman, Friedman, Stone, & Olshen, 1984) analysis. The CART analysis was applied to visualize the suggested hierarchy of variables with respect to the three measured parameters (SGR, LT, and MPD). The CART analysis generates a tree-like structure by the set of decision points yielding partitions of the original group. The more important variables to affect the response, the higher the position on the tree they have as decision points.

### 3 RESULTS AND DISCUSSION

#### 3.1 Comparison of the growth kinetics of L. monocytogenes by the kind of ravioli, antimicrobial agent, and packaging

The growth curves of L. monocytogenes in both spinach and artichoke ravioli for 60 days well fitted to the Baranyi model. The mean values of the growth kinetics of L. monocytogenes were compared according to the kind of ravioli (spinach versus artichoke), antimicrobial agents (CSV blend versus nisin), and packaging (modified atmosphere versus aerobic) (Table 1). There was a higher growth potential of L. monocytogenes in spinach ravioli as compared to artichoke ravioli. Although no significant differences were observed in the SGR and LT values, the MPD values were differed significantly between spinach (5.420 log cfu/g) and artichoke ravioli (4.569 log cfu/g). These results indicate that the different filling in ravioli affects the level of stationary phase of L. monocytogenes and ravioli with spinach filling has a higher risk than that with artichoke filling. L. monocytogenes in artichoke was able to survive, but did not grow, with remaining counts at the initial concentration (around 4.5 log cfu/g) during the storage period (Sanz, Giménez, & Olarte, 2003). In previous research, extracts of artichoke exhibited antimicrobial activity against bacteria species, yeasts, and molds (Emanuel, Adrian, Sultana, & Sventtiana, 2011; Ionescu et al., 2013; Zhu, Zhang, & Lo, 2004). 7.7% contamination rate of L. monocytogenes in spinach was reported, whereas the pathogen was not isolated from artichokes (Cordano & Jacquet, 2009). Other reports also emphasized the hazard of spinach as a food ingredient for humans. Specifically, Pingulkar, Kamat, and Bongirwar (2001) and Yolanda Moreno et al. (2012) reported that the higher percentage of L. monocytogenes positive was detected in spinach samples. L. monocytogenes prevalence under such environment was attributed to the cross-contamination and ability of growth even at refrigeration, as well as to ambient temperature of processed foods (Beuchat, 1996; Wilks, Michels, & Keevil, 2006). The population of L. monocytogenes increased from 2.4 to 8.8 log cfu/ml in autoclaved spinach powder cultures at 30°C, indicating that spinach products provide a good environment for the growth of L. monocytogenes (Babic, Watada, & Buta, 1997).

In addition, antimicrobial effects of 0.1% CSV blend and 0.3% nisin on the control of L. monocytogenes growth in spinach ravioli were compared. No significant differences in the LT, SGR, and MPD values were observed between 0.1% CSV blend and 0.3% nisin. Despite this comparable activity, the CSV blend has its advantage of allowing the hazard of humans. Specifically, Pingulkar, Kamat, and Bongirwar (2001) and Yolanda Moreno et al. (2012) reported that the higher percentage of L. monocytogenes positive was detected in spinach samples. L. monocytogenes prevalence under such environment was attributed to the cross-contamination and ability of growth even at refrigeration, as well as to ambient temperature of processed foods (Beuchat, 1996; Wilks, Michels, & Keevil, 2006). The population of L. monocytogenes increased from 2.4 to 8.8 log cfu/ml in autoclaved spinach powder cultures at 30°C, indicating that spinach products provide a good environment for the growth of L. monocytogenes (Babic, Watada, & Buta, 1997).
would result in cell damage, death, or exhaustion due to expenditure of energy and resources to exclude the acid.

Response of a microbial cell depends on the nature and amount of direct inhibitors and the influence of the environment (Caillet, Millette, Salmieri, & Lacroix, 2006; Kostaki, Giatrakou, Savvidis, & Kontominas, 2009). One of the factors contributing to the microbialic effectiveness of antimicrobials is the nature of the atmosphere in contact with the target cells. It is well known that the composition (e.g., O2, N2, and CO2) of modified atmosphere systems can be an effective means to restrict or inhibit the growth of aerobic spoilage organisms of perishable foods, as well as to maintain the visual quality of products (Narasimha Rao & Sachindra, 2002; Stanbridge & Davies, 1998). In this study, we compared the growth kinetics of L. monocytogenes in spinach ravioli under two packaging conditions (MA and aerobic). Under the same concentrations of the CSV blend, inhibitory power was improved when extra hurdle of MAP was applied. SGR of L. monocytogenes under MAP was approximately a half (0.582 log/day) of that under aerobic condition (1.005 log/day, Table 1). Moreover, the LT in spinach ravioli with MAP was significantly extended up to 15.5 days, as compared to 2.5 days in aerobic packaged spinach ravioli. MPD was also lower in spinach ravioli with MAP (5.68 log cfu/g) than with aerobic packaging (7.51 log cfu/g). The combined treatment of low oxygen and high concentration of CO2 or N2, can provide adequate suppression of the growth of L. monocytogenes (Kostaki et al., 2009). According to Whitley, Muir, and Waites (2000), N2 MAP in the absence of O2 increases the LT of L. monocytogenes in cheese up to 3 weeks and retards the growth of L. monocytogenes.

3.2 | Modeling of L. monocytogenes growth in spinach ravioli as a function of the CSV blend and storage temperature

To predict the growth restriction of L. monocytogenes by adding the CSV blend in spinach ravioli with MAP, the primary growth models were generated at various temperatures. The impacts of storage temperature and the level of the CSV blend on SGR, LT, and MPD are summarized in Table 2. Adding the CSV blend at 0.3, 0.5, or 1% resulted in significant differences (p < .05) in the SGR values of L. monocytogenes in spinach ravioli, as compared with control (0%) or 0.1% (p < .05). Adding more than 0.3% CSV blend significantly decreased SGR from 0.15 to 0.05 log cfu/hr at 4°C and from 0.23 to 0.06 log cfu/hr at 10°C (p < .05). These results indicate that 0.3% CSV blend significantly affects SGR of L. monocytogenes in spinach ravioli at temperatures up to 10°C; however, it was not sufficient to decrease SGR at ambient temperature. At 17 and 24°C, addition of 0.5% CSV blend reduced SGR by 68 and 61%, respectively, as compared to the control.

In the case of LT, the addition of 1% of CSV blend in spinach ravioli completely inhibited the growth of L. monocytogenes at 4°C for 60 days storage. At 4°C, LT of L. monocytogenes in spinach ravioli were extended from 25.89 (control, 0% CVS blend) to 35.75 with 0.3% CVS blend and 56.30 days with 0.5% CVS blend. At 10°C, a significant extension of LT was still observed with the addition of 0.3 (25.86 days), 0.5 (36.42 days), and 1% (50.21 days) CSV blend compared to control (17.70 days). The LT values were significantly shortened with the increase of storage temperature. Abou-Zeid et al. (2007) also reported that lactate-diacetate mixture could lengthen the LT of L. monocytogenes only under refrigerated temperatures with pH below 6.5. In general, lag periods increase as the environmental conditions become less favorable for the growth of the pathogen. Robinson, Ocio, Kaloti, and Mackey (1998) hypothesized that lag could be determined by two parameters—namely, the amount of work to be done to adapt to a new environment and the rate at which that work can be done. Maintaining homeostasis and repairing cellular damage resulting from the antimicrobials and low temperature often require the activation of several mechanisms, which is metabolically demanding for injured cells and, therefore, leads to an extended lag phase.

The values of MPD of L. monocytogenes in spinach ravioli decreased significantly with the decrease of storage temperatures and the addition of the CSV blend (p < .05). The addition of more than 0.3% of the CSV blend in dough for spinach ravioli significantly decreased MPD at 4°C (2.47 log cfu/g), 10°C (4.55 log cfu/g), and 17°C (4.86 log cfu/g), as compared to the control (p < .05). Thus, to achieve effective inhibition of Listeria growth in spinach ravioli, it is recommended to add at least 0.3% of the CVS blend in dough. In our study, when 1% CSV blend was added into spinach ravioli, MPD of L. monocytogenes did not exceed 3 log cfu/g for 60 days under refrigerated temperatures. At 17 and 24°C, the behavior of L. monocytogenes demonstrated a significant change with 0.5 and 1% CSV blend. Other studies also reported the antimicrobial effect of the CSV blend as the

**TABLE 1** Comparison of the growth kinetics of L. monocytogenes according to the kind of ravioli, antimicrobial agent, and packaging

| Raviolia | Antimicrobialb | Packagingc |
|----------|---------------|------------|
| Spinach  | CSV | MAP  | Aerobic |
| Artichoke| Nisin |        |         |
| SGR      | 0.568 ± 0.650| 0.949 ± 0.827| 0.582 ± 0.627*| 1.005 ± 0.845 |
| LT       | 17.636 ± 18.459| 7.765 ± 11.084| 15.545 ± 15.766*| 2.510 ± 2.356 |
| MPD      | 5.420 ± 1.893*| 7.434 ± 0.1603| 5.682 ± 1.832*| 7.507 ± 1.454 |

LT = lag time (day); SGR = specific growth rate (log/day); MPD = maximum population density (log cfu/g).

aModified packaged ravioli with CSV blend at 0, 0.1, 0.3, 0.5, and 1% was compared.
bCSV blend and nisin at 0.1 and 0.3% was compared in spinach ravioli.
cOnly spinach ravioli treated CSV blend was compared.

*Significant differences between the mean values of growth kinetics were determined by t test (p < .05).
function of amount. For example, Glass and Sindelar (2010), Sullivan (2012), and Schrader (2010) also reported that the 3% CSV blend reduced L. monocytogenes growth on meat products. In addition, the survival of other pathogens, such as C. jejuni and S. Typhimurium, on precooked chicken breasts was also controlled with 3% CSV blend (Park et al., 2014). The observed concentration-discrepancy regarding the effectiveness of the antimicrobial could be explained by the difference in food products and food formulations during the course of experiment. The CSV blend used in the present study (up to 1%) in dough was lower than the recommended levels (3–5%) by the PURAC and other studies. There were significant interactions between the concentrations of the CSV blend and storage temperatures in spinach ravioli with MAP in the present study. The combined treatment of the CSV blend in dough with MAP packaging was effective at pathogen reduction, without significantly affecting the quality, and demonstrated its potential as a novel method to increase the microbial safety in ravioli, possibly other products, such as pasta and noodle, etc. Of all available options for ensuring safety, maintenance of low temperature throughout this food chain is likely to be the most effective.

The response surface models were also developed to describe the effects of the CSV blend concentration and storage temperature on SGR, LT, and MPD (Table 3). The growth kinetic values in the primary model of the present study were subjected to the response surface analysis using the SAS general linear model procedures. The developed secondary model was sufficient to predict the potential effect of storage temperature and CSV blend on L. monocytogenes growth in spinach ravioli. The $R^2$ values of SGR, LT, and MPD were .98, .94, and .93, respectively, indicating that the models effectively predicted the growth of L. monocytogenes in spinach ravioli with MAP.

In the present study, the growth of L. monocytogenes was not observed in MAP packed spinach ravioli with 1% CSV blend at 4°C (Table 2). Therefore, these conditions were excluded from the development of the secondary growth model in the present study. Increased amounts of CSV blend resulted in a significant growth control of L. monocytogenes in spinach ravioli with MAP.

### Table 2: Comparison of the growth kinetics of L. monocytogenes in CSV blend treated spinach ravioli with MAP

| Concentration of CSV (%) | Temperature (°C) |
|--------------------------|------------------|
|                          | 4                | 10               | 17               | 24               |
| SGR                      |                  |                  |                  |                  |
| 0                        | $^{b}0.15 \pm 0.06^a$ | $^{b}0.23 \pm 0.08^a$ | $^{b}1.08 \pm 0.37^a$ | $^{b}1.85 \pm 0.19^a$ |
| 0.1                      | $^{b}0.09 \pm 0.02^a$ | $^{b}0.14 \pm 0.05^a$ | $^{b}0.94 \pm 0.09^a$ | $^{b}1.96 \pm 0.06^a$ |
| 0.3                      | $^{b}0.05 \pm 0.00^a$ | $^{b}0.06 \pm 0.00^a$ | $^{b}0.65 \pm 0.05^a$ | $^{b}1.64 \pm 0.06^a$ |
| 0.5                      | $^{b}0.02 \pm 0.01^a$ | $^{b}0.06 \pm 0.00^a$ | $^{b}0.35 \pm 0.01^a$ | $^{b}0.73 \pm 0.00^a$ |
| 1                        | $^{b}0.01 \pm 0.00^a$ | $^{b}0.05 \pm 0.00^a$ | $^{b}0.27 \pm 0.00^a$ | $^{b}0.80 \pm 0.01^a$ |
| LT                       |                  |                  |                  |                  |
| 0                        | $^{b}25.89 \pm 2.05^b$ | $^{b}17.70 \pm 0.00^d$ | $^{b}1.23 \pm 0.03^c$ | $^{b}1.04 \pm 0.69^c$ |
| 0.1                      | $^{b}26.30 \pm 0.26^b$ | $^{b}25.83 \pm 0.02^d$ | $^{b}1.75 \pm 0.11^c$ | $^{b}1.26 \pm 0.01^c$ |
| 0.3                      | $^{b}35.75 \pm 0.02^b$ | $^{b}36.42 \pm 0.07^d$ | $^{b}5.16 \pm 0.07^b$ | $^{b}2.78 \pm 0.01^b$ |
| 0.5                      | $^{b}56.30 \pm 7.99^a$ | $^{b}50.21 \pm 2.4^a$ | $^{b}7.18 \pm 0.00^a$ | $^{b}6.28 \pm 0.03^a$ |
| 1                        | NA               |                  |                  |                  |
| MPD                      |                  |                  |                  |                  |
| 0                        | $^{b}4.73 \pm 0.00^a$ | $^{b}7.02 \pm 0.05^a$ | $^{b}7.85 \pm 0.04^a$ | $^{b}8.09 \pm 0.00^a$ |
| 0.1                      | $^{b}5.14 \pm 0.71^a$ | $^{b}6.82 \pm 0.00^a$ | $^{b}8.72 \pm 0.08^a$ | $^{b}8.24 \pm 0.21^a$ |
| 0.3                      | $^{b}3.35 \pm 0.00^a$ | $^{b}4.55 \pm 0.00^a$ | $^{b}8.46 \pm 0.00^a$ | $^{b}7.71 \pm 0.07^a$ |
| 0.5                      | $^{b}2.76 \pm 0.04^a$ | $^{b}3.70 \pm 0.00^a$ | $^{b}8.16 \pm 0.00^a$ | $^{b}8.75 \pm 0.00^a$ |
| 1                        | $^{b}2.47 \pm 0.41^a$ | $^{b}2.97 \pm 0.00^a$ | $^{b}8.18 \pm 0.00^a$ | $^{b}5.72 \pm 0.00^a$ |

**LT =** lag time (day); **SGR =** specific growth rate (log/day); **MPD =** maximum population density (log cfu/g).

* Means (n = 4) ± SD in the same column with different superscripts are significantly different by Duncan’s multiple range test at p < .05.

*–* Means (n = 4) ± SD in the same row with different superscripts are significantly different by Duncan’s multiple range test at p < .05.

### Table 3: Response surface models for effect of temperature and CSV blend concentration (0–1%) on lag time (LT), specific growth rate (SGR), and maximum population density (MPD) of L. monocytogenes in spinach ravioli

| Parameter | SGR | LT | MPD |
|-----------|-----|----|-----|
|           | Estimate | SE | Pr > | | Estimate | SE | Pr > | | Estimate | SE | Pr > |
|           |        |    | $|$ |    |        |    | $|$ |    |        |    | $|$ |
| Intercept | 0.0479 | 0.1189 | .6932 | .98 | 41.3454 | 5.3445 | <.0001 | .94 | 4.7068 | 0.6134 | <.0001 | .93 |
| temp      | 0.0172 | 0.0174 | .3388 | .66 | −3.7115 | 0.8064 | .0005 | .02066 | 0.0897 | .0372 |
| anti      | −0.7709 | 0.2969 | .0211 | .04 | 64.6007 | 13.6507 | .0004 | −8.5007 | 1.5319 | <.0001 |
| temp × temp | 0.0027 | 0.0006 | .0004 | .08 | 0.0852 | 0.0283 | .1000 | −0.0012 | 0.0031 | .6960 |
| anti × anti | 0.9708 | 0.2497 | .016 | .23 | −2.0734 | 11.8314 | .8484 | 5.4556 | 1.2884 | .0008 |
| temp × anti | −0.0652 | 0.0098 | <.0001 | .26 | 2.6137 | 0.5730 | .0005 | −0.0237 | 0.0504 | .6455 |
refrigeration temperature. In summary, the models in the matrix of conditions described in the present study can be used as a tool to estimate the impact of CSV blend and temperature on the growth of *L. monocytogenes* in retail ravioli (Table 3).

### 3.3 Determination of important control factor for the growth of *L. monocytogenes* by the CART analysis

We used the CART analysis to discriminate important variables (concentration of antimicrobials, packaging method, and storage temperature) in determining the growth kinetics of *L. monocytogenes* in spinach ravioli.

Figure 1 shows the results of the CART analyses using the mean absolute deviation criterion for the SGR (Figure 1b), LT (Figure 1b), and MPD (Figure 1c) values of *L. monocytogenes* in spinach ravioli. From the three variables (concentration of antimicrobials, packaging method, and storage temperature), storage temperature was the most important variable to affect SGR, followed by packaging method. Storage temperature of 4 and 10°C discriminated SGR in spinach ravioli at a split value of SGR, 0.223 log/day, whereas SGR was sixfold greater (SGR; 1.341 log/day) at 17 and 24°C. In the case of storage temperatures below 10°C, packaging method discriminated SGR 0.086 and 0.360 log/day under MAP and aerobic packaging, respectively, indicating that packaging method is an important control factor for *listeria* growth at the refrigeration temperature, while packaging method was not an important control factor any more at ambient temperature (Figure 1a).

Also, storage temperature was the most important variables in discriminating LT. The results of the CART analysis identified that storage temperature is the only important variable to affect the LT value. At temperature less than 10°C, LT of *L. monocytogenes* in spinach ravioli was estimated to 17.938 day, while LT at 17°C above was approximately one-eight shorter (2.218 day) than that at the refrigeration temperature. The results also indicated that, although packaging method and the CSV blend concentration were significant factors to affect LT, their effects were very low.

At the first split of MPD, the MPD value of *L. monocytogenes* in spinach ravioli with MAP was 5.42 log/g, while that with aerobic packaging was 7.64 log/g. The results show that packaging method is an important control factor for *listeria* growth at the refrigeration temperature, while packaging method was not an important control factor any more at ambient temperature (Figure 1a).

![FIGURE 1](image) Cart analysis result of (a) SGR, (b) LT, and (c) MPD. The number in the top node indicates parameter value with the corresponding sample number in parenthesis. By following the tree structure, partitions are generated with sequential criteria of the selected factors. The root nodes are shaded in grey with final predicted SGR values for the portioned samples.
packaging was 7.64 log/g, which is 1.4 times higher than MAP. This result indicates that packaging method is the main control factor to affect MPD. In L. monocytogenes in spinach raviolis with MAP, the samples treated with above 0.1% CSV blend showed the smallest MPD value (4.34 log/g), which was 38% lower than that of raviolis without CSV blend (7.05 log/g). In the case of spinach raviolis with aerobic packaging, 0.5% concentration of CSV blend was the critical level. The MPD value of L. monocytogenes in spinach raviolis with less than 0.5% CSV blend with aerobic packaging was 1.3 times (8.50 log/g) higher than that of the CSV blend concentration equal to or above 0.5% (6.36 log/g). It should be noted that the threshold concentration for the MAP partition was between 0.1 and 0.3%, while that for the aerobic packaging ranged between 0.3 and 0.5%. These results indicate that low concentration of antimicrobial agent can be effective to control L. monocytogenes in spinach raviolis with MAP. Although the degree of importance varied, temperature was the most important factor, followed by packaging method and concentration of the CSV blend to control the growth of L. monocytogenes in spinach ravioli.

Consistently with the results of previous studies, our CART analysis results further emphasized that the storage at the refrigeration temperature is a critical factor in controlling L. monocytogenes growth in food products. The combined treatment of the CSV blend with MAP was effective in increasing L. monocytogenes reduction without significantly affecting the quality, and demonstrated its potential as a novel method to increase the microbial safety in ready-to-cook refrigerated ravioli.

3.4 | Sensory evaluation

In the triangle test, 16 of 40 (45%) panelists correctly chose spinach sample treated with the 1% CSV blend that was different from the other two samples. Thus, it can be concluded with 95% confidence that not more than 50% of the population would be able to detect a difference in color, smell, or taste of spinach ravioli with 1% CSV blend. Our results also indicated that the product was well accepted by the panelists. Other studies reported that many conventional antibacterial interventions (e.g., irradiation) can result in undesirable alterations to the appearance, taste and smell of food (Gecgel, 2013) and make those foods less desirable to the consumer. However, our results showed that the organoleptic quality of spinach ravioli was not affected by application of 1% CSV blend and that no differences in taste, color, or appearance were detectable.

4 | CONCLUSIONS

The results of this study confirm the combined efficacy of the CSV blend as a clean label/natural antimicrobial and other hurdle factors, such as low temperature and MAP, on decreasing the risk of L. monocytogenes without significantly affecting the quality of RTC ravioli at retail market. In addition, the developed growth model for L. monocytogenes can predict the growth parameters of L. monocytogenes as a function of the CSV blend at varying concentration (0–1%) and storage temperature (4–24°C) to determine the optimum conditions for controlling the growth of L. monocytogenes in spinach ravioli at retail stores, thus replace the challenge study. According to the results of our CART analysis, although the microorganism is psychrotrophic, refrigeration is the primary factor to control the rate of L. monocytogenes growth. In addition, MAP was an effective hurdle technique to control L. monocytogenes in ravioli products. According to the results of sensory evaluation, the CSV blend substantially contributes to the extension of shelf-life of MAP spinach raviolis, delaying their spoilage while imparting a pleasant flavor.

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