Chemical treatment of water used to wash or crisp raw fruits and vegetables

Issue you would like the Conference to consider:
The consumption of fresh produce has increased in the US while at the same time outbreaks of public health significance related to fresh fruits and vegetables continue to occur. The safety of fresh produce remains a challenge for the food industry. As technologies to enhance the safety of fresh produce become more readily available, they should be utilized by all food establishments.

As specified in the 2013 Food Code in section 3-302.15 Washing Fruits and Vegetables, "...raw fruits and vegetables shall be thoroughly washed in water to remove soil and other contaminants before being cut, combined with other ingredients, cooked, served, or offered for human consumption in READY-TO-EAT form. [Emphasis added]"

Further, in the Food Code paragraph 3-302.15(B), it states, "Fruits and vegetables may be washed by using chemicals as specified under § 7-204.12." [Emphasis added]

Washing fresh produce, in this context, is required but using treated water is optional. It is well documented that raw agriculture commodities (RACs) may be contaminated with pathogens and, when soaked or submerged in water, there is a risk of cross-contamination. Various chemicals are available that can minimize and/or prevent cross-contamination and, to a lesser degree, reduce pathogen load on fresh produce.

Therefore, the Conference should consider that when produce is washed, crisped, re-hydrated or processed by soaking or submersion, the water used for these purposes shall be chemically treated to minimize the risk of cross-contamination.

Public Health Significance:
The use of chemicals for washing, treatment, storage and processing fruits and vegetables is specified in the Food Code as follows:

7-204.12 Chemicals for Washing, Treatment, Storage and Processing Fruits and Vegetables, Criteria.
(A) Chemicals, including those generated on-site, used to wash or peel raw, whole fruits and vegetables shall:

1. Be an approved food additive listed for this intended use in the Code of Federal Regulations, 21 CFR 173, or
2. Be generally recognized as safe (GRAS) for this intended use, or
3. Be the subject of an effective food contact notification for this intended use (only effective for the manufacturer or supplier identified in the notification), and
4. Meet the requirements in 40 CFR 156 Labeling Requirements for Pesticide and Devices.

The criteria for using chemicals for washing, treatment, storage and processing fruits and vegetables are designated in the Food Code as priority items. Sufficient controls are already prescribed to ensure the safe and effective use of these chemicals.

Washing raw fruits and vegetables can remove soil and other contaminants. Many food establishments use soaking or submersion as an approved, effective technique for washing produce. This method is often preferred for a variety of reasons, including:

- The contact time is better controlled
- All surfaces come in direct contact with the water
- It reduces the amount of waste water
- It allows for simultaneous washing and re-hydrating
- It helps minimize shrink and extends shelf life
- It improves the appearance of the product
- And, when chemicals are added, can provide an antimicrobial treatment for the reduction/prevention of cross-contamination.

It is well documented that pathogenic microorganisms may be present on the exterior surfaces of raw fruits and vegetables. The Food Code Annex 3, Chapter 3, Section 3-302.15 Washing Fruits and Vegetables states that "...more recent studies have demonstrated washing to fall short of their [pathogens] complete removal." There is currently no readily available treatment that can ensure removal or destruction of all pathogens on raw agriculture commodities (RACs) with the possible exception of irradiation.

Using chemically treated water to wash and/or process fresh produce can impact public health by minimizing the risk of cross-contamination and reducing pathogens if they are present. The Food Code Annex 3, Public Health Reasons, supports this position in Section 3-302.15 Washing Fruits and Vegetables as follows:

"All fresh produce, except commercially washed, pre-cut, and bagged produce, must be thoroughly washed under running, potable water or with chemicals as specified in Section 7-204.12, or both, before eating, cutting or cooking. Even if you plan to peel or otherwise alter the form of the produce, it is still important to remove soil and debris first" [Emphasis added] and "It is important to follow practices that minimize pathogens in the water or on the surface of produce." [Emphasis added]
The use of chemicals is equivalent, if not better, than rinsing under running water. Further, the use of chemicals will minimize pathogens in the water. It is estimated (unpublished data) that over three-quarters of grocery stores soak/submerge certain raw produce items to wash, crisp and/or re-hydrate them. Concerns about cross-contamination have led some experts to question the potential risk when soaking produce in untreated water. However, treated water has been shown to be very effective in minimizing/preventing cross-contamination.

In January 2014, the Food Marketing Institute published, in collaboration with the Produce Marketing Association and United Fresh Produce Association, "Produce Safety Best Practices Guide for Retailers" advising retailers to use sanitizers when soaking/submerging fresh produce. The following guidance was provided to retailers regarding crisping fresh produce:

- If a bath is used, follow sanitizer recommendations
- If using a bath, an appropriate sanitizer should be used in compliance with label directions. [Emphasis added]

(Treatment-wash-water-produce.pdf)

Treating produce wash water in the processing sector has been extensively studied. The FDA Guidance for Industry: Guide to Minimize Microbial Food Safety Hazards for Fresh Fruits and Vegetables (October, 1998) specifically addresses this issue. In Chapter 2, Section 2.2 it states, "...antimicrobial chemicals in processing water are useful in reducing microbial build-up in water and may reduce microbial load on the surface of produce. Thus, antimicrobial chemicals may provide some assurance in minimizing the potential for microbial contamination."

(www.fda.gov/food/guidanceregulation/guidancedocumentsregulatoryinformation/ucm064574.htm)

The failure to add antimicrobial chemicals in processing water has also been cited as a contributing factor in foodborne outbreaks attributed to fresh produce. For example, a U.S. House of Representatives, Committee on Energy and Commerce report on an investigation of an outbreak of Listeria monocytogenes in cantaloupe states that FDA officials found several deficiencies including not using an "antimicrobial solution such as chlorine in the water used to wash the cantaloupes."

Additional studies and research support the use of chemicals in water that comes in contact with RACs. For example, a study was conducted in November 2013, comparing 5 different sanitizer options and plain tap water. It was found that sanitizers can have a significant impact on food safety because they are effective in reducing pathogens in the wash water itself, which reduces opportunities for cross-contamination.[1] (attached)

At the 2013 Symposium of the Center for Produce Safety (CPS), a collaborative partnership of industry, government and academic communities, a Key Learning report on wash water concluded, in part:

*Many different products are washed, cooled or transported using water. Therefore it is important that the water is treated and maintained properly so that it does not become a source of cross contamination for human pathogens, should they be present. It is equally important to remember that simply washing products is not an effective mechanism for*
removing contamination, i.e. it cannot remove or kill pathogens that have had the opportunity to naturally seek out hidden surfaces on products and adhere to them. Therefore our focus is to manage contamination risks throughout production (e.g. GAPs, inspections, hygiene, equipment sanitation, training programs, etc.) and control wash, cooling and transport processes using water so that we do not create cross contamination scenarios. Improper control over wash, cooling or water-based transport systems can do harm, i.e. resulting in large-scale cross contaminations. Dr. Trevor Suslow vividly demonstrated this assertion using an inoculated cilantro load and washing it with uninoculated parsley on a commercial wash system. The improperly controlled wash system permitted cross contamination onto the parsley demonstrating the potential for cross contamination.


On January 22, 2013, the Center for Produce Safety (CPS) conducted a seminar on post-harvest water disinfection. Among the Key Learnings from this seminar was the following conclusion:

Disinfectants are used in water that contacts produce to prevent cross contamination and not necessarily to kill microorganisms that might be present on the surface of the fruit or vegetable.

If water is not properly treated with active disinfectant, after a period of time the water could become a source of contamination for any fruits or vegetables that are conveyed, cooled or washed in it. Therefore the primary reason for treating water with disinfectants is to keep the water clean of microbial build up. In most systems the level of microbial reduction on the surface of fruits or vegetables is generally thought to be 1-2 logs.

(http://www.pma.com/content/articles/2014/05/cps-wash-water-key-learnings)

The FDA Analysis and Evaluation of Preventive Control Measures for the Control and Reduction/Elimination of Microbial Hazards on Fresh and Fresh-Cut Produce, A Report of the Institute of Food Technologists for the Food and Drug Administration published September 30, 2001 provided this summary in Chapter V. Section 1:

It is well established that pathogenic microorganisms associated with whole or fresh-cut produce can cause disease outbreaks, thereby demonstrating the need for improved mitigation efforts to reduce risks associated with these products.

The best method to eliminate pathogens from produce is to prevent contamination in the first place. However, this is not always possible and the need to wash and sanitize many types of produce remains of paramount importance to prevent disease outbreaks. It should be noted that washing and sanitizing are unlikely to totally eliminate all pathogens after the produce is contaminated. Therefore, it is important to use washing and sanitizing protocols that are efficient.

(http://www.fda.gov/Food/FoodScienceResearch/SafePracticesforFoodProcesses/ucm091363.htm)

Finally, the technology and/or products used to treat water used to wash, crisp, re-hydrate or process fresh produce by soaking or submersion are not proprietary. Several antimicrobial compounds are readily available to the industry. No one product or supplier is advocated. Food establishments have the opportunity to select a water treatment that is
most appropriate to their circumstances. A comprehensive review of these chemicals can be found in the FDA Preventive Control Measures for Fresh & Fresh-Cut Produce, Chapter V., Methods to Reduce/Eliminate Pathogens from Produce and Fresh-Cut Produce. (www.fda.gov/Food/FoodScienceResearch/SafePracticesforFoodProcesses/ucm090977.htm)


**Recommended Solution: The Conference recommends...:**

that a letter be sent to the FDA requesting that section 3-302.15 of the 2013 Food Code be amended as follows (language to be added is underlined; language to be deleted is in strikethrough format):

3-302.15 Washing Fruits and Vegetables.

**(B) Fruits and vegetables may **shall be washed by using chemicals as specified under § 7-204.12 when soaked or submerged.**

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**Supporting Attachments:**

- "Efficacy of Commercial Produce Sanitizers against Nontoxigenic Escherichia"

*It is the policy of the Conference for Food Protection to not accept Issues that would endorse a brand name or a commercial proprietary process.*
Efficacy of Commercial Produce Sanitizers against Nontoxigenic Escherichia coli O157:H7 during Processing of Iceberg Lettuce in a Pilot-Scale Leafy Green Processing Line

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ABSTRACT

Chemical sanitizers are routinely used during commercial flume washing of fresh-cut leafy greens to minimize cross-contamination from the water. This study assessed the efficacy of five commercial sanitizer treatments against Escherichia coli O157:H7 on iceberg lettuce, in wash water, and on equipment during simulated commercial production in a pilot-scale processing line. Iceberg lettuce (5.4 kg) was inoculated to contain 10⁶ CFU/g of a four-strain cocktail of nontoxigenic, green fluorescent protein–labeled, ampicillin-resistant E. coli O157:H7 and processed after 1 h of draining at ~22 °C. Lettuce was shredded using a commercial slicer, step-conveyed to a flume tank, washed for 90 s using six different treatments (water alone, 50 ppm of peroxyacetic acid, 50 ppm of mixed peracid, or 50 ppm of available chlorine either alone or acidified to pH 6.5 with citric acid [CA] or T-128), and then dried using a shaker table and centrifugal dryer. Various product (25-g) and water (50-ml) samples collected during processing along with equipment surface samples (100 cm²) from the flume tank, shaker table, and centrifugal dryer were homogenized in neutralizing buffer and plated on tryptic soy agar. During and after iceberg lettuce processing, none of the sanitizers were significantly more effective (P ≤ 0.05) than water alone at reducing E. coli O157:H7 populations on lettuce, with reductions ranging from 0.75 to 1.4 log CFU/g. Regardless of the sanitizer treatment used, the centrifugal dryer surfaces yielded E. coli O157:H7 populations of 3.49 to 4.98 log CFU/100 cm². Chlorine, chlorine plus CA, and chlorine plus T-128 were generally more effective (P ≤ 0.05) than the other treatments, with reductions of 3.79, 5.47, and 5.37 log CFU/ml after 90 s of processing, respectively. This indicates that chlorine-based sanitizers will likely prevent wash water containing low organic loads from becoming a vehicle for cross-contamination.

In 2009, leafy greens were ranked as the riskiest food category regulated by the U.S. Food and Drug Administration, accounting for 363 outbreaks and 13,568 reported cases of illness (13). Between 1995 and 2006, leafy green–associated outbreaks increased by 38.6%, whereas consumption increased by only 9% (22). The nationwide outbreak of Escherichia coli O157:H7 that was traced to baby spinach in 2006 resulted in 205 confirmed infections, 103 hospitalizations, and three deaths (10, 17). Following two additional E. coli O157:H7 outbreaks in 2006 linked to shredded iceberg lettuce resulting in 150 illnesses (12), at least nine more outbreaks responsible for nearly 300 cases of E. coli O157:H7 infection have been documented in the United States through 2012 (14), heightening continued safety concerns surrounding fresh-cut leafy greens.

Bacterial pathogens can contaminate leafy greens at any point during the farm-to-fork continuum (31). Major on-farm areas of concern now recognized by the U.S. Food and Drug Administration include agricultural water, biological soil amendments (e.g., manure), domesticated and wild animals, field worker health and hygiene, and the cleanliness of harvesting equipment, tools, and buildings (47). However, leafy greens are also prone to contamination during commercial processing, packing (8), distribution, marketing (51), and in-home preparation (35). Regarding leafy greens, pathogens are most likely to attach to stomata, irregularities on intact surfaces, cut surfaces, or cracks on the external surfaces (20, 36, 38, 39, 42) and can be protected from sanitizers by biofilms (40). Because sanitizers in the wash water cannot be relied upon to inactivate attached or internalized pathogens during processing, it is imperative that growers and harvesters follow good agricultural practices and good handling practices to reduce the likelihood of contamination (19).

Washing of leafy greens remains important for removing soil and debris, decreasing the microbial load, improving quality and appearance, and enhancing product shelf life and safety (21). Numerous small-scale laboratory studies have shown that produce sanitizers reduce pathogen populations by 1 to 3 log CFU on lettuce (4, 18, 20, 36, 38), with water alone decreasing E. coli O157:H7 levels about 1 log CFU on lettuce during pilot-scale processing (6). Recirculation of this wash water during processing can further magnify the spread of contaminants at large, centralized processing facilities (21, 28). Hence, the

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addition of sanitizers to processing water is imperative to minimize cross-contamination during commercial production of fresh-cut leafy greens (2, 29, 38, 46).

Chlorine-based sanitizers are preferred for commercial flume washing systems because of their relatively low cost compared with other sanitizers and minimal negative impact on end-product quality (11, 21, 29, 33, 36). Since the active component of chlorine, hypochlorous acid (HClO), is most abundant at pH 6.5 to 7.0 (3), the pH of the wash water typically needs to be lowered by adding a weak acid, most commonly citric acid (21). A new, generally recognized as safe acidifying agent composed of phosphoric acid and propylene glycol, known as T-128 (SmartWash Solutions, Salinas, CA), has been developed to improve the stability of chlorine (25, 29, 33, 41). However, chlorine use has raised concerns regarding potentially hazardous by-products, worker safety, environmental damage, and most importantly, decreased efficacy in the presence of an increasing organic load in recirculating flume water, which has heightened interest in other alternatives such as peroxyacetic acid–based sanitizers (38, 43).

Numerous small-scale laboratory studies have assessed sanitizer efficacy against pathogens on leafy greens (1, 4, 23, 24, 27, 30, 34, 42, 52, 53). However, these findings are difficult to extrapolate to large-scale commercial production facilities. Previous work completed by our group was performed without chemical sanitizers to quantify E. coli O157:H7 transfer during pilot-plant production of fresh-cut leafy greens (6, 7). Since chemical sanitizers remain the sole intervention strategy to prevent cross-contamination during commercial production of fresh-cut leafy greens, it is imperative that these sanitizers be reevaluated under conditions that more closely resemble commercial operations. Consequently, the objective of this study was to assess the efficacy of five commercial sanitizer treatments against E. coli O157:H7 during processing of iceberg lettuce in a pilot-scale leafy green processing line.

MATERIALS AND METHODS

Experimental design. The efficacy of five different sanitizing treatments was assessed in triplicate against E. coli O157:H7 by processing a 5.4-kg batch of iceberg lettuce inoculated at 10⁶ CFU/g, with sanitizer-free water serving as the control. All lettuce was processed by shredding, conveying, fluming, shaker table dewatering, and/or centrifugal drying, during and/or after which various product, water, and equipment surface samples were collected and quantitatively examined for E. coli O157:H7.

Iceberg lettuce. Individually wrapped heads of iceberg lettuce (Lactuca sativa L.) (24 heads per case) were obtained from a local wholesaler (Stan Setas Produce Co., Lansing, MI), with the product originating from California or Arizona depending on the growing season. All lettuce was stored in a 4°C walk-in cooler and used within 5 days of delivery.

Bacterial strains. Four nontoxigenic (stx₁⁻ and stx₂⁻) strains of E. coli O157:H7 (ATCC 43888, CV287, 6980-2, and 6982-2) were obtained from Dr. Michael Doyle at the Center for Food Safety, University of Georgia, Griffin. These strains had been previously transformed with a pGFPuv plasmid containing a green fluorescent protein gene and ampicillin-resistance gene. All four strains were stored at −80°C in tryptic soy broth (Difco, BD, Sparks, MD) containing 0.6% (wt/vol) yeast extract (Difco, BD) (TSBYE) and 10% (vol/vol) glycerol (Sigma Chemical Co., St. Louis, MO) until needed. Working cultures were prepared by streaking each stock culture on tryptic soy agar plates (Difco, BD) containing 0.6% (wt/vol) yeast extract and 100 ppm of ampicillin (ampicillin sodium salt, Sigma Life Science, St. Louis, MO) (TSAYE plus amp). After 18 to 24 h of incubation at 37°C, a single colony was transferred to 9 ml of TSBYE containing 100 ppm of ampicillin (TSBYE plus amp) and similarly incubated.

Lettuce inoculation. A 0.2-ml aliquot of each nontoxigenic E. coli O157:H7 strain was transferred to 200 ml of TSBYE with amp and incubated for 18 to 20 h at 37°C. Based on similar growth rates as determined previously (6), the four strains were combined in equal volumes to obtain an 800-ml cocktail, which was added to 80 liters of municipal tap water (−15°C, <0.05 ppm of free chlorine) in a 121- liter plastic container (Rubbermaid, Wooster, OH) to achieve a level of ~10⁸ CFU/ml. Hand-cored heads of iceberg lettuce (~12 heads) were immersed in the E. coli suspension for 15 min and then drained or air dried for 1 h at 22°C before being spun in a dewatering centrifuge (described below) to remove residual inoculum from the interior of the heads. Duplicate 25-g samples were then aseptically collected to determine the initial inoculation level at the time of processing.

Lettuce processing line. The same small-scale commercial leafy green processing line consisting of a lettuce shredder, step conveyor, flume tank, shaker table, and dewatering centrifuge was used as previously described in detail by Buchholz et al. (6). For this work, a custom-made stainless steel gate with 1.25-cm-diameter holes spaced 0.65 cm apart (Heinzen Manufacturing, Inc., Gilroy, CA) was added at the end of the 3.3-m-long stainless steel flume tank to retain the product during 90 s of washing.

Wash water. Iceberg lettuce (0.5 kg) was homogenized in 500 ml of Michigan State University tap water using a mechanical blender (model BLC10650MB, Black & Decker, New Britain, CT) and then added to 890 liters of processing water at 12 to 15°C to achieve a low organic load. The following five commercial produce sanitizer treatments were assessed: 30 ppm of peroxyacetic acid (Tsunami 100, Ecolab, St. Paul, MN), 30 ppm of mixed peracid (Tsunami 200, Ecolab), 30 ppm of available chlorine (XY-12, Ecolab) at pH 7.85, 30 ppm of available chlorine (XY-12) acidified to pH 6.50 with citric acid (Sigma-Aldrich, St. Louis, MO), and 30 ppm of available chlorine (XY-12) acidified to pH 6.50 with T-128 (SmartWash Solutions) as measured with a pH probe (pHTest 30, Oakton, Vernon Hills, IL). Peroxyacetic acid test kit 311 (Ecolab) was used to confirm the peroxyacetic acid and mixed peracid sanitizer concentrations, and chlorine test kit 321 (Ecolab) was used to measure available chlorine. Sanitizer-free Michigan State University tap water (<0.05 ppm of free chlorine) served as the control.

Lettuce processing. Inoculated heads of cored iceberg lettuce (5.4 kg) were hand-fed into the shredder at a rate of about 0.5 kg per s, with the shredded product then step-conveyed at a rate of 2.85 m/s to the top of the conveyor. Processing was then halted for ~10 min to aseptically collect and bag five 25-g lettuce samples in red mesh produce bags (5 lb Header Bag, Pacon Inc., Baldwin Park, CA) for subsequent sampling. Thereafter, processing was resumed with the iceberg lettuce conveyed to the flume tank, washed in 890 liters of recirculating wash water with or without a
concentrated neutralizing buffer (BD) was filled with All lettuce samples (25 g) were E. coli O157:H7 counts. The three equipment surface % E. coli and were subjected to O157:H7. The one-ply composite tissue log CFU/g reduction seen for concentrated Difco neutralizing O157:H7 counts were converted # #. O157:H7. O157:H7 populations for 3). Means of the same wash water # #. Iceberg lettuce contained an average DAVIDSON ET AL. 0.05). concentration would neutralize # # #. E. coli ~ (wt/vol) phosphate buffer measuring 100 cm % E. coli value of (48) O157:H7 populations decreased 1.40 log CFU/g; E. coli # J. Food Prot., Vol. 76, No. 11 sanitizer for 90 s, partially dewatered on the shaker table, collected in a single centrifugation basket, and centrifugally dried.

**Sample collection.** During the 90 s of flume washing, three prebagged iceberg lettuce samples (25 g each) were retrieved at the flume gate at 30-s intervals and were immediately added to 100 ml of sterile Difco neutralizing buffer (BD, Franklin Lakes, NJ) in a Whirl-Pak filter bag (Nasco, Fort Atkinson, WI). In addition, nine 50-ml water samples were collected at 10-s intervals in 50-ml centrifuge tubes containing 38 × concentrated Difco neutralizing buffer (BD). After shaker table dewatering, product in the basket was washed in the preset 50-lb (110-kg) capacity Spin Dryer (model SD50-LT, Heinen Manufacturing). During centrifugal drying, four water samples (50 ml each) were similarly collected from the centrifuge drain at 10-s intervals for the first 40 s of the 80-s cycle. After centrifugation, two bagged lettuce samples (25 g each) were also retrieved from the centrifugation basket. Nine product contact areas on the equipment (three flume tank, three shaker table, and three dewatering centrifuge), previously described in detail by Buchholz et al. (6), measuring 100 cm² as previously identified using Glo Germ (Glo Germ Co., Moab, UT) were sampled immediately after processing as described by Vorst et al. (48) using one-ply composite tissues moistened with 1 ml of sterile Difco neutralizing buffer (BD).

**Microbiological analyses.** All lettuce samples (25 g) were homogenized in a stomacher (Stomacher 400 Circulator, Seward, Worthington, UK) for 1 min at 260 rpm and then either appropriately diluted in sterile 1% (wt/vol) phosphate buffer (8.5 g/liter NaCl, 1.44 g/liter Na₂HPO₄, and 0.24 g/liter KH₂PO₄; J.T. Baker, Mallinckrodt Baker Inc., Phillipsburg, NJ) and plated on TSAYE with amp (calculated minimum detection limit of 40 CFU/g) or processed using 0.45-µm-pore-size membrane filters (Millipore, Millipore Corporation, Billerica, MA) (calculated minimum detection limit of 0.04 CFU/g), which were placed on 60-mm-diameter petri plates containing TSAYE with amp to quantify E. coli O157:H7. The one-ply composite tissue samples were added to 15 ml of sterile Difco neutralizing buffer in a Whirl-Pak bag, homogenized for 1 min at 260 rpm, and then plated identically to the lettuce samples, giving a calculated lower detection limit of 1 CFU/100 cm². The 50-ml water samples were either appropriately diluted in sterile 1% phosphate buffer and plated on TSAYE with amp or processed by membrane filtration, which gave a calculated minimum detection limit of 0.02 CFU/ml. Following 20 to 24 h of incubation at 37°C, all green fluorescing colonies as seen under UV light (365 nm; Blak-Ray, Ultra-violet Product Inc., San Gabriel, CA) were counted as E. coli O157:H7.

**Sanitizer neutralization confirmation.** Triplicate 1-liter water samples containing 30 ppm of available chlorine (XY-12), 30 ppm of peroxyacetic acid (Tsunami 100), or 30 ppm of mixed peracid (Tsunami 200 ppm) were prepared and confirmed with chlorine test kit 321 or peroxyacetic acid test kit 311. Citric acid (Sigma-Aldrich) and T-128 were used to acidify the chlorine-based sanitizer solution to pH 6.5. A 50-ml centrifuge tube containing 3 ml of 38 × concentrated neutralizing buffer (BD) was filled with the sample containing sanitizer, agitated for 5 s, and then immediately assessed for neutralization of the sanitizer as previously described using the appropriate test kit. Preliminary experiments found that a 38 × concentration would neutralize various concentrations of the active component of each sanitizing agent used in this study without impacting E. coli O157:H7 counts.

**Statistical analysis.** E. coli O157:H7 counts were converted to log CFU per gram, milliliter, or 100 cm² and were subjected to analysis of variance using JMP 9.0 (SAS Institute Inc., Cary, NC). Values equaling half the limit of detection were used for samples without E. coli O157:H7 counts. The three equipment surface samples from each respective piece of equipment were averaged. A P value of ≤0.05 was considered significant for all tests. The Tukey-Kramer honestly significant difference test was used to identify significant differences in E. coli O157:H7 populations for individual lettuce, water, and equipment surface samples.

**RESULTS**

**Lettuce.** Iceberg lettuce contained an average E. coli O157:H7 inoculum of 5.93 log CFU/g at the time of processing (Fig. 1). After shredding, conveying, 90 s of washing, shaker table dewatering, and centrifugal drying, no significant difference (P > 0.05) was seen in populations of E. coli O157:H7 recovered from the finished product, regardless of sanitizer treatment. Using mixed peracid, E. coli O157:H7 populations decreased 1.40 log CFU/g; however, this decrease was not significantly different (P > 0.05) compared with the 0.75-log CFU/g reduction seen for water alone. Processing significantly reduced (P ≤ 0.05) E. coli O157:H7 populations on lettuce when mixed peracid, chlorine,
or chlorine plus CA were used, with reductions of 1.40, 0.77, and 0.89 log CFU/g, respectively. The reductions of 0.75, 0.93, and 0.97 log CFU/g seen for water alone, peroxyacetic acid, and chlorine plus T-128, respectively, were not significant ($P > 0.05$) (Fig. 1).

Flume water. Wash water containing chlorine, chlorine plus T-128, and chlorine plus CA had significantly lower ($P < 0.05$) $E. coli$ O157:H7 populations at all sampling times (maximum of 0.99 log CFU/ml) compared with 4.61 log CFU/ml in water alone. Using chlorine plus CA and chlorine plus T-128, $E. coli$ O157:H7 levels were below the limit of detection of 0.02 log CFU/ml by the end of processing. $E. coli$ O157:H7 populations were similar ($P > 0.05$) using water alone and peroxyacetic acid, with respective populations of 3.47 and 3.01 log CFU/ml recovered after 90 s of processing. Similar $E. coli$ O157:H7 populations were obtained using mixed peracid ($P > 0.05$) and peroxyacetic acid, with these populations rarely lower ($P < 0.05$) than those in water alone (Fig. 2).

Centrifugation water. Using peroxyacetic acid, mixed peracid, or chlorine, wash water exiting the centrifuge drain after spin drying yielded maximum $E. coli$ O157:H7 populations of 4.51, 4.36, and 5.48 log CFU/ml, respectively, which were not significantly different ($P > 0.05$) from those in water alone (maximum population of 5.58 log CFU/ml) during the 40-s sampling period. However, chlorine plus CA and chlorine plus T-128 resulted in $E. coli$ O157:H7 populations that were lower than those in water alone ($P < 0.05$) during the first 20 s of centrifugation. Water samples collected after 40 s of centrifugation yielded $E. coli$ O157:H7 populations that were not significantly different for any of the treatments (Fig. 3).
FIGURE 4. Mean (±SD) E. coli O157:H7 populations on equipment surfaces after processing iceberg lettuce inoculated at ~6 log CFU/g (n = 3). Half the limit of detection was used to calculate the mean log value when a sample did not yield any colonies by direct plating. Means of the same product type with different letters are significantly different (P ≤ 0.05).

Processing equipment surfaces. After processing iceberg lettuce, all five sanitizer treatments yielded significantly lower (P ≤ 0.05) E. coli O157:H7 populations remaining on the flume tank and shaker table as compared with the water control. Significantly lower (P ≤ 0.05) E. coli O157:H7 populations were recovered on the centrifugal dryer using peroxycetic acid (3.61 log CFU/100 cm²) and mixed peracide (3.49 log CFU/100 cm²) compared with the other treatments, with the highest level (4.98 log CFU/100 cm²) seen when water alone was used for washing (Fig. 4).

DISCUSSION

Due to the potential production of infectious aerosols during lettuce processing, the same four nontoxigenic strains of E. coli O157:H7 were used as in our earlier transfer studies (6, 7). The growth and adherence rates for these four nontoxigenic strains were previously shown to be similar to three strains from the 2006 leafy green outbreaks (6). As previously reported, green fluorescent protein labeling also allowed for easy differentiation of the inoculum from background bacteria (6, 7, 49).

Dip inoculation of the lettuce to contain 6 log CFU/g was crucial to ensure uniform distribution of E. coli O157:H7 throughout the heads as well as quantifiable results for subsequent mathematical modeling with this work to be reported elsewhere. Although this inoculation level clearly exceeds levels thought to occur on field-grown lettuce, feces from “super-shedding” cows can potentially contain E. coli O157:H7 at levels of 6 log CFU/g (15), with such fecal material potentially able to come in contact with lettuce through irrigation water. Preliminary experiments using a mixture of Glo Germ and water showed uniform fluorescence in dipped heads of iceberg lettuce. Additionally, Buchholz and others (6) found that E. coli O157:H7 populations were statistically similar in iceberg lettuce heads before and after shredding, indicating that the inoculation was homogenous. Dip inoculation of the cored lettuce heads may have allowed internalization of E. coli O157:H7 through the damaged tissues, with such cells protected from sanitizers (37). Since all lettuce samples were processed by stomaching, any internalized cells would have gone undetected with only the cells on the surface of the leaves recovered.

Commercial producers of fresh-cut leafy greens use different sanitizers, sanitizer concentrations, and contact times, depending on the design of the processing line. In this study, six different wash treatments were assessed during 90 s of flume washing. Processing inoculated iceberg lettuce resulted in E. coli O157:H7 reductions of 0.75 to 1.4 log CFU/g on the finished product. Both during and after processing, no significant differences in sanitizer efficacy (P > 0.05) were seen against E. coli O157:H7 on iceberg lettuce for any of the treatments, including water alone. However, three wash treatments—mixed peracide, chlorine, and chlorine plus CA—significantly reduced (P ≤ 0.05) E. coli O157:H7 populations after washing. Numerous small-scale laboratory studies have shown similar pathogen reductions (~1 log CFU/g) during washing of various fruits and vegetables with or without sanitizers (4, 5, 9, 50). Using a pilot-scale leafy green processing line, Luo et al. (29) also reported an E. coli O157:H7 reduction of <1 log after processing inoculated baby spinach (29). Consequently, produce sanitizers cannot be relied upon to ensure end product safety. Chemical sanitizers are routinely added to recirculating wash water to minimize the spread of microbial contaminants during flume washing (27). Regarding their use, peroxycetic acid–based sanitizers are limited to a maximum of 80 ppm of peroxycetic acid (16, 21), whereas free chlorine concentrations typically range from 10 to a maximum of 200 ppm (20, 36, 45). However, soil, debris, and vegetable latexes released during shredding of leafy greens will accumulate in the flume water over time (32), decreasing the efficacy of many sanitizers, most notably chlorine (2, 26, 38, 52). The wash water used in this study contained an organic load of ~0.0006% blended iceberg lettuce (wt/vol) to simulate wash water quality during the early stages of processing. Hence, higher E. coli O157:H7 populations would have been expected after 90 s of processing if the organic load in the wash water had been.
higher, especially for the chlorine-based sanitizer. *E. coli* O157:H7 populations recovered from the wash water were consistently lower (*P* ≤ 0.05) using chlorine, chlorine plus CA, and chlorine plus T-128 compared with water alone, peroxyacetic acid, and mixed peracid. Both chlorine plus CA and chlorine plus T-128 treatments yielded *E. coli* O157:H7 levels that were below the limit of detection, which is similar to the findings of López-Gálvez et al. (27) using 40 ppm of chlorine.

This study was designed to assess the efficacy of sanitizers during processing, not to assess long-term pathogen persistence in the wash water. Produce sanitizers are primarily used to minimize cross-contamination during flume washing, with their effectiveness dependent on the type of sanitizer, concentration, temperature, and organic load in the wash water. The pilot-scale processing line used in this study was not equipped with a chiller. Therefore, all processing needed to be conducted at our incoming tap water temperature of 12 to 15°C rather than at the targeted commercial temperature of 4°C. Since sanitizer efficacy against *E. coli* O157:H7 is enhanced at temperatures above 4°C (53), our *E. coli* O157:H7 reductions likely exceed those that would be expected in commercial operations.

Levels of *E. coli* O157:H7 recovered from spent centrifugation water containing sanitizers were rarely lower than those seen in sanitizer-free water. Similar *E. coli* O157:H7 populations were recovered from centrifugation water containing peroxyacetic acid, mixed peracid, chlorine, or no sanitizer at all four sampling times. The combination of chlorine and citric acid or T-128 was significantly more effective than the other sanitizers (*P* ≤ 0.05) against *E. coli* O157:H7 in centrifugation water collected during the first 20 s; however, after 40 s no significant difference was seen compared with the water control (*P* > 0.05). These results indicate that, whereas populations of *E. coli* O157:H7 may be close to or below the limit of detection in flume water, populations in the centrifugation water were not significantly different than the water control by the end of sample collection. Therefore, spent centrifugation water would be best suited for pathogen testing.

*E. coli* O157:H7 cells recovered from equipment surfaces after processing reflect those that were present in the film of water on the equipment surface. During processing, the flume tank was in continuous contact with the recirculating wash water, with water contact decreasing during shaker table dewatering and centrifugal drying. Numbers of *E. coli* O157:H7 recovered from surfaces in the centrifugal dryer were not significantly different from the water control when any of the three chlorine-based sanitizer treatments were used, indicating that those surfaces may also be well suited for pathogen testing, depending on the particular sanitizer used.

This study was done to assess the efficacy of commercial produce sanitizers against *E. coli* O157:H7 on lettuce, in wash water, and on equipment surfaces during small-scale processing of iceberg lettuce. Whereas none of the sanitizers were more effective than water alone against *E. coli* O157:H7 on iceberg lettuce at any point during or after processing, it is important to reiterate that sanitizers are designed to reduce the microbial load in wash water rather than on the product. Overall, the populations of *E. coli* O157:H7 recovered in wash water containing peroxyacetic acid or mixed peracid were rarely significantly different than those seen in water alone. However, the three chlorine-based treatments were significantly more effective than water alone at reducing *E. coli* O157:H7 populations in wash water during processing. The wash water used in this study replicated a “best-case” scenario for processors due to the extremely low organic load and freshly added sanitizers. Similar studies using higher organic loads will be needed to assess sanitizer efficacy against *E. coli* O157:H7 under conditions that more closely simulate commercial processing.

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