Lytic bacteriophages reduce *Escherichia coli* O157:H7 on fresh cut lettuce introduced through cross-contamination

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The role of lytic bacteriophages in preventing cross contamination of produce has not been evaluated. A cocktail of three lytic phages specific for *E. coli* O157:H7 (Ecoshield™) or a control (phosphate buffered saline, PBS) was applied to lettuce by either; (1) immersion of lettuce in 500 ml of Ecoshield™ 8.3 log pFU/ml or 9.8 log pFU/ml for up to 2 min before inoculation with *E. coli* O157:H7; (2) spray-application of Ecoshield™ (9.3 log pFU/ml) to lettuce after inoculation with *E. coli* O157:H7 (4.10 CFU/cm²) following exposure to 50 μg/ml chlorine for 30 sec. After immersion studies, lettuce was spot-inoculated with *E. coli* O157:H7 (2.38 CFU/cm²). Phage-treated, inoculated lettuce pieces were stored at 4°C for and analyzed for *E. coli* O157:H7 populations for up to 7 days. Immersion of lettuce in 9.8 log pFU/ml Ecoshield™ for 2 min significantly (p < 0.05) reduced *E. coli* O157:H7 populations after 24 h when stored at 4°C compared with controls. Immersion of lettuce in suspensions containing high concentrations of Ecoshield™ (9.8 log pFU/ml) resulted in the deposition of high concentrations (7.8 log pFU/cm²) of bacteriophages on the surface of fresh cut lettuce, potentially contributing to the efficacy of the lytic phages on lettuce. Spraying phages on to inoculated fresh cut lettuce after being washed in hypochlorite solution was significantly more effective in reducing *E. coli* O157:H7 populations (2.22 log CFU/cm²) on day 0 compared with control treatments (4.10 log CFU/cm²). Both immersion and spray treatments provided protection from *E. coli* O157:H7 contamination on lettuce, but spray application of lytic bacteriophages to lettuce was more effective in immediately reducing *E. coli* O157:H7 populations fresh cut lettuce.

**Introduction**

Two outbreaks of *Escherichia coli* O157:H7 infections in 2006 associated with the consumption of bagged baby spinach and bagged shredded lettuce were collectively responsible for sickening 254 people and causing 3 deaths.1,2 Romaine lettuce contaminated with *E. coli* O157:H7, grown in California, was responsible for causing a multi-state outbreak of 60 infections in 2011.3 A previous multi-state outbreak associated with *E. coli* O154 and contaminated Romaine lettuce grown in Arizona resulted in 34 cases of foodborne illness.4 Bagged spinach and shredded lettuce are especially vulnerable to bacterial contamination due to the lack of a kill-step during their harvesting, processing and packaging. Concentrations of chemical disinfectants like sodium hypochlorite used during the processing of fresh cut leafy green commodities may not be sufficient to kill foodborne pathogens attached to intact or cut surfaces of leafy greens, and are primarily used to prevent cross contamination of leafy greens through water.5,6

Outbreaks of foodborne illness associated with the consumption of leafy greens emphasize the need for an effective natural antimicrobial which can be directly applied to fresh cut leafy green to kill enteric pathogens. Bacteriophages are viruses that which can attack, infiltrate, lyse and kill bacterial cells.7 Phages are the most abundant organisms on earth, and they are highly specific for bacterial species with a lack of ability to cause deleterious effects on human health.8 Thus, they may provide a safe, natural, and effective alternative to chemical sanitizers. Lytic bacteriophages can also use a phenomenon termed “Lysis from without” (LO) to kill bacterial cells, where cell lysis is not mediated through the infective cycle but through a large number of phages adsorbing to the surface of a cell, resulting in cell wall damage, physiological stress and eventual cell lysis.7 The LO mechanism is the most likely route of biocontrol of enteric bacterial pathogens used on refrigerated produce commodities, where the storage temperature prohibits the completion of the full lytic cycle.9 A cocktail of phages specific for *Listeria monocytogenes* (Listshield™) has been approved by the Food and Drug Administration (FDA) for use on ready-to-eat foods Recently, a mixture of *E. coli* O157:H7-specific bacteriophages (Ecoshield™, Intralytix) has been approved through Food Contact notices by FDA and have been
granted a temporary exemption by the FSIS for use on meats and on food contact surfaces. Phages have been found to be naturally present in a variety of food commodities including chicken, pork sausage, deli meats, and also mushrooms, lettuce, refrigerated biscuit dough, and frozen chicken pot pies. Lytic phage activity has also been assessed on meats, and a cocktail of three O157-specific bacteriophages was effective in reducing populations by 2 log CFU/sample when stored at 4°C for 7 d. Different capital letters at the top of each bar indicate significant (p < 0.05) differences between control (PBS) and lytic bacteriophages treatment (EcoShield™) within each day.

Previous work evaluating the effectiveness of lytic bacteriophages against foodborne pathogens has shown that phages can be effective when applied before the introduction of the pathogen; however, the role of lytic bacteriophages in preventing cross-contamination—applying phages before the introduction of the pathogen to the produce surface—has not been evaluated. The effectiveness of using bacteriophages (EcoShield™) to prevent contamination of lettuce with E. coli O157:H7 introduced through cross-contamination was investigated.

**Results and Discussion**

Immersion of cut lettuce pieces in EcoShield™ suspension (9.8 log PFU/ml) for 2 min significantly (p < 0.05) reduced populations of E. coli O157:H7 when immediately applied to fresh cut lettuce stored at 4°C for 1, 3 and 7 d after inoculation, compared with lettuce treated with control (phosphate buffered saline, PBS) (Fig. 1). By days 6 and 7, E. coli O157:H7 populations were reduced to undetectable levels in EcoShield™-treated lettuce samples. However, immersion of lettuce pieces in EcoShield™ suspension for 2 min at a lower titer (8.3 log PFU/ml), and followed by inoculation of E. coli O157:H7, was not as effective as immersion in higher (9.8 log PFU/ml) EcoShield™ suspensions to reduce E. coli O157:H7 populations (Fig. 2). Although significant statistical differences in E. coli O157:H7 populations between control and bacteriophage-treated lettuce were not observed at the lower concentration (8.3 log PFU/ml), populations did decline after 48 h of storage at 4°C, and remained low until day 7. When lettuce pieces were immersed for shorter durations (30 sec) at 8.3 log PFU/ml (Fig. 3), no significant reductions were observed between bacteriophage treatments and control. It was also not as effective as the 2 min immersion. For example, immersion of lettuce in EcoShield™ suspensions at 8.3 log PFU/ml for 30 sec resulted in 4.5 log PFU/cm² on fresh cut lettuce pieces; but immersion of lettuce in EcoShield™ at higher concentrations (9.8 log PFU/ml) for 2 min resulted in 7.8 log PFU/cm² on the lettuce surface. MOI values at the lower concentration of EcoShield™ (8.3 log PFU/ml) were < 100; however, MOI at the higher concentration (9.8 log PFU/ml) were > 10000. The longer contact time between the lettuce and bacteriophages (2 min) allowed more bacteriophage particles to adsorb to the surface of the lettuce, leading to the rapid and more effective reduction of E. coli O157:H7.

Currently, fresh cut leafy greens are washed in chlorinated water before being de-watered and packaged into bags before shipment to retail outlets. These washes are not intended to kill bacteria on the surface of fresh cut leafy greens but rather to reduce the likelihood of cross-contamination through water on to fresh cut leafy greens. Lytic bacteriophages applied to lettuce either during a washing step or after dewatering step may provide a more direct mechanism to directly reduce E. coli O157:H7 on
Unlike our previous work, our current study introduces bacteriophages to fresh cut lettuce before the introduction of E. coli O157:H7 on to the surface of lettuce to determine if a pre-treatment of lytic bacteriophages can protect against cross-contamination.22-23 Our results show that application of phages by this method does not cause an immediate reduction of E. coli O157:H7 on the surface of fresh cut lettuce, but reductions do occur during several days of storage at 4°C (Figs. 1 and 2).

Previous work evaluating the effectiveness on leafy greens had utilized spraying of lytic bacteriophages on the commodity after the introduction of 4 log CFU E. coli O157:H7. Abuladze et al. (2008) sprayed ECP-100 (now EcoShield™), a mixture of three lytic bacteriophages specific for E. coli O157:H7, on to spinach, and reduced E. coli O157:H7 populations by 100% and 99% after 24 and 168 h, respectively.22 Similarly, a spray treatment of ECP-100 on fresh cut lettuce following the introduction of E. coli O157:H7, reduced bacterial populations by 1.82 log CFU/cm² on day 0.23 With both of these experiments, the effect of spray application was rapid and substantial in reducing E. coli O157:H7 populations on lettuce surfaces. The effectiveness of spray application was attributed to increased phage particle dispersion on lettuce leaves. Since the E. coli O157:H7 cells were pre-established on the leaf surface and the phage-treated lettuce was stored at 4°C, the phages are assumed to cause rapid E. coli O157:H7 reductions through the lysis from without (LO) mechanism, mediated by the attachment of multiple bacteriophages to a single cell, lysing the cell membrane without undergoing the entire infective cycle of the bacteriophage.

Other workers who used a spot inoculation technique, whereby bacteriophages specific for E. coli O157:H7 were applied to the same spot as E. coli O157:H7 was applied on a baby spinach leaf, also observed increased bacterial reductions after 24 h compared with 10 min or 1 h at all tested storage temperatures of 4, 8 and 23°C.24 These findings, coupled with the results presented here, support that immersion or spot application can cause reductions of E. coli O157:H7 populations after 24 or 48 h periods at 4°C, but not as rapidly as when bacteriophages are spray-applied to lettuce.

Unlike previous studies with a spray application after the introduction of E. coli O157:H7 to the lettuce surface, no immersion treatment in our current study resulted in the immediate reduction of E. coli O157:H7 on day 0 as was previously
observed. The immersion of pieces in the EcoShield™ solution may have impeded the mobility of the bacteriophage on the surface of the lettuce pieces compared with the spray application. The dispersion of bacteriophage particles on the surface of lettuce pieces achieved through spraying may compensate for the limited mobility of bacteriophages on cut lettuce, whereas immersion may not be as effective a method for optimal distribution of the phages on the lettuce surface. Recent work using model E. coli and coliphages with a bioluminescent reporting system to measure bacterial viability reported that lytic phages reduced the number of bacterial cells on the surface of the lettuce compared with controls (non-phage treated) lettuce within 5 h, as measured through reduced bioluminescence on the lettuce surface stored at room temperature after 24 h. In our immersion studies of lettuce using bacteriophage concentrations of 8.3 or 9.8 log PFU/ml for 2 min, the lack of an immediate bacteriophage-mediated decline in E. coli O157:H7 populations may be attributable to the more localized and limited dispersion of the phage particles on the lettuce surface. Spraying lettuce surface tissue with bacteriophages, as previous studies have shown, may have allowed the bacteriophages to be more active on the surface of the tissue and not penetrate deeply into the tissue, thus facilitating the rapid interaction between E. coli O157:H7 cells and the introduced bacteriophages. Furthermore, the storage of the lettuce at 4°C, with the refrigeration temperature limiting metabolic activity, may have also retarded the interactions between phage particles and bacterial cells. In an effort to simulate and evaluate the lytic bacteriophages in a fresh cut leafy green processing/washing regime, phages were sprayed on to lettuce after treatment with 50 μg/ml sodium hypochlorite (NaOCl) for 30 sec and then de-watered. On day 0 (immediately after treatment), lettuce treated with NaOCl only (3.00 log CFU/cm²), NaOCl and EcoShield™ (2.22 log CFU/cm²), or EcoShield™ only (3.28 log CFU/cm²) all had significantly lower E. coli O157:H7 populations than control treatment (4.10 log CFU/cm²) (Table 1). The control treatment was phosphate buffer, with no EcoShield™ or hypochlorite treatment. However, there were no significant statistical differences between E. coli O157:H7 populations on lettuce receiving hypochlorite or EcoShield™ treatments. After 24 h of storage at 4°C, the same trends were observed as on Day 1—lettuce which received no chemical or EcoShield™ treatments had significantly higher E. coli O157:H7 populations (4.86 log CFU/cm²) than those treated with NaOCl (3.48 log CFU/cm²), NaOCl and EcoShield™ (3.32 log CFU/cm²), or EcoShield™ alone (3.54 log CFU/cm²). On both day 0 and day 1, lettuce treated with sodium hypochlorite and EcoShield™ bacteriophages had lower E. coli O157:H7 populations than lettuce treated with either NaOCl or EcoShield™ alone (although differences between treatments other than control were not statistically significant). This finding suggests that lytic bacteriophages could be used synergistically with chemical sanitizers to reduce E. coli O157:H7 populations, and are more effective than employing solely a chemical or biological agent. Furthermore, bacteriophage titers on the surface of lettuce pieces sprayed with EcoShield™ were 7.2 and 6.4 log PFU/cm² on day 0 and 1, respectively. The effective MOI was > 1000. These titers are similar to those observed on lettuce pieces treated with 50 μg/ml hypochlorite for 30 sec followed by spraying with EcoShield™ (6.8 and 6.7 log PFU/cm²) on days 0 and 1, respectively. Apparently the residual chlorine on lettuce leaves did not inactivate the bacteriophage particle or reduce the lytic potential of bacteriophages applied against E. coli O157:H7 introduced to the lettuce surface. The MOI of the spray study was 10-fold lower (> 1000) than observed in the immersion study at 9.8 log PFU/ml (> 10,000), and therefore is unlikely responsible for the increased effectiveness of the spray treatment compared with the immersion treatment. These results are similar to those observed using lytic bacteriophages specific for Listeria monocytogenes in a suspension with a quaternary ammonium compound-based sanitizer, QUATAL. Those workers observed that a suspension containing three listeriapheres and QUATAL in combination was significantly more effective in reducing L. monocytogenes populations on stainless steel and polypropylene cylinders than separate treatments containing either listeriaphages or QUATAL. Listeriapher concentration were not reduced in the presence of QUATAL utilized at 50 μg/ml concentration.

The preliminary findings presented here indicate that an immersion treatment of lytic bacteriophages for fresh cut lettuce or a combination of a hypochlorite wash and E. coli O157:H7-specific bacteriophages have the potential to be applied to leafy greens in a processing facility and/or a field setting. The incorporation of lytic bacteriophages into washwater used to process fresh cut lettuce, or a subsequent treatment before dewatering, may not affect an immediate reduction in E. coli O157:H7 populations, but populations do decline after 24 h or 48 h at 4°C in our current study. Although the impact of the modified atmosphere packaging (MAP) used to preserve leafy green appearance and quality on the efficacy of lytic bacteriophages was not assessed in our work, other workers have shown that EcoShield™ was effective on fresh cut spinach stored under MAP conditions currently used for fresh cut leafy greens. Current harvesting and processing of iceberg lettuce intended for bagged salads or shredded lettuce packages begins with a head of iceberg lettuce being cut at the stalk in the field, trimmed of damaged leaves, and then cored before being placed on a conveyor belt and deposited into a bin to be transported to the processing facility. Lettuce, as a member of family Asteraceae is a latex producer, especially when its phloem cells are damaged, releasing lactucin-laden latex.
shown to predispose lettuce to opportunistic growth of *E. coli* O157:H7 and exacerbate contamination.30 Cored areas of lettuce heads are sprayed with 50–150 ppm chlorine (as hypochlorite) wash on the coveroy belt in the field to primarily remove latex, but such concentrations are insufficient to kill 100% of an enteric bacterial pathogen present.31 The demonstrated synergistic effect of the hypochlorite and lytic bacteriophages as studied under our experimental conditions provide an opportunity to test direct antimicrobial mitigation against potential *E. coli* O157:H7 contamination of fresh cut lettuce in a field processing setting.

In summary, the mode of application (immersion and spraying) affected the efficacy of lytic bacteriophage treatments to reduce *E. coli* O157:H7 on fresh cut lettuce. Although both methods provided a degree of protection from introduction of *E. coli* O157:H7 to fresh cut lettuce, spray application of lytic bacteriophages specific for *E. coli* O157:H7 to lettuce surfaces was more effective in immediately reducing *E. coli* O157:H7 populations on lettuce surfaces compared with immersion of lettuce in EcoShield™ solutions. The presence of *E. coli* O157:H7 on the surface of the leaf before spraying may have promoted more rapid and frequent interactions between phage particles and *E. coli* O157:H7 compared with the addition of *E. coli* O157:H7 to lettuce leaves prior to immersion in a lytic bacteriophage solution. A synergistic effect between lytic bacteriophage treatment and sodium hypochlorite wash was also observed, indicating the suitability of using lytic bacteriophages in the presence of such sanitizers. Immersion of fresh cut lettuce into a lytic bacteriophage solution reduced *E. coli* O157:H7 counts only after 24 to 48 h after of when stored at 4°C, indicating that bacteriophages required extended periods of time to reduce *E. coli* O157:H7 populations during refrigerated storage. The use of lytic bacteriophages can reduce *E. coli* O157:H7 on fresh cut lettuce. More research is needed to optimize the application of *E. coli* O157:H7-specific bacteriophages against *E. coli* O157:H7 on leafy greens under field- and facility-processing conditions.

**Materials and Methods**

**Strains and bacteriophage preparations used.** *Escherichia coli* O157:H7 gfp86, which expresses the green fluorescent protein, was obtained from the USDA-ARS Microbial Food Safety Laboratory (Fratamico et al., 1997).32 The isolate was streaked from frozen stock on to tryptic soy agar (TSA) (Becton Dickinson) supplemented with 100 μg/ml ampicillin (TSAA) (Sigma-Aldrich) and incubated at 37°C for 24 h. An isolated colony was then inoculated into tryptic soy broth (Becton Dickinson) supplemented with ampicillin (Sigma Aldrich) (TSBA) and incubated with agitation (175 rpm) for 24 h at 37°C. Cultures were then serially diluted in 0.1% peptone water for either dilution into sterile water or direct application to fresh cut lettuce. EcoShield™ consists of three *E. coli* O157:H7-specific lytic bacteriophages (ECML-4, ECML-117 and ECML-134) of the *Myoviridae* family isolated from fresh and salt water environments, and was obtained from Intralytix, Inc.31 EcoShield™ was diluted in phosphate buffered saline (PBS) (Sigma Aldrich) to various concentrations for immersion and spray applications. Treatment of fresh cut lettuce by immersion in lytic bacteriophage suspension. Iceberg lettuce (*Lactuca sativa*) was purchased at a grocery store in Beltsville, MD. The plastic packaging of heads of iceberg lettuce (*Lactuca sativa*) was rinsed with 70% ethanol before removal. Outer leaves of the lettuce were removed and inner leaves were cut with a sterile stainless steel scalpel. A sterile metal coupon was then used to cut 3 cm × 3 cm pieces of lettuce on a sterile metal tray. For immersion studies, 24 pieces of lettuce were immersed in 500 mL of sterile water containing EcoShield™ in a 1 L sterile glass beaker with a frequently spinning stir-bar to simulate flume water in fresh cut lettuce processing. Pieces of lettuce were immersed in 8.3 log PFU/ml for 30 sec or 2 min 9.8 PFU/ml EcoShield™ for 2 min. Twenty-four pieces of lettuce were also washed with phosphate buffer (PBS) as a control for the same periods of time. Excess EcoShield™ suspension or PBS was then removed from lettuce pieces by using a household salad spinner (Oxoid). Pieces of lettuce were allowed to dry for 10 min before each piece was spot inoculated with 50 μl of 5.1 log CFU/ml *E. coli* O157:H7. The inoculum was applied using a 100 μl pipet, dispensing 8–10 droplets on each piece of lettuce. Lettuce was allowed to dry for 30 min in a biosafety cabinet at 25°C. Pieces of lettuce were stored on a piece of wetted filter paper and sealed (with parafilm) in a plastic Petri dish for up to 7 d at 4°C.

Spray treatment of inoculated cut lettuce in combination with EcoShield™ and sodium hypochlorite solution. Pieces of lettuce (9 cm²) were spot inoculated with *E. coli* O157:H7 with 50 μl of 6.9 log CFU/ml and allowed to dry for 90 min. Pieces of lettuce were either placed in 500 ml of a 50 μg/ml sodium hypochlorite (NaOCl) solution (Sigma Aldrich), made up in 0.05 M potassium phosphate buffer, or 500 ml PBS. NaOCl solutions were prepared in sterile chlorine-saturated 1 L glass beaker. Free chlorine concentrations were measured using a DR 720 colorimeter (Hach Inc.) and appropriate chlorine testing methodology. Inoculated lettuce was washed in NaOCl solution for 30 sec. Pieces of lettuce were then either immediately placed in in 15 ml Dey Engley (Becton Dickson) to neutralize the NaOCl in a 50-ml disposable conical tube (Fisher Scientific), or in a salad spinner to removed excess moisture and ‘spin’ for 15 sec. Subsequently the pieces of lettuce were then sprayed immediately with EcoShield™ (9.3 log PFU/ml) or PBS. Pieces of lettuce were treated with either PBS or EcoShield™ administered using a hand-held air-brush (Model 200, Badger Air-Brush Co.) attached to an air compressor (Badger Air Brush Co.). The air brush was filled with 20 ml of EcoShield™ (9.3 PFU/ml), and calibrated to deliver 2.4 ml over a 500 cm² surface area,30 which resulted in 5.98 log PFU EcoShield™/cm² concentration in the manner described above. After lettuce pieces were sprayed with EcoShield™ suspension, individual pieces were either placed in 15 ml of PBS for microbial analysis or stored at 4°C for 24 h as described above.

Microbiological analysis of *E. coli* O157:H7 populations on cut lettuce. After appropriate duration, one piece of lettuce inoculated with *E. coli* O157:H7 treated with EcoShield™ or PBS and were added to 15 ml of PBS in a 50-ml conical tube and homogenized with a Polytron Pr 2100 laboratory blender.
(Kininematic AG) for 30 sec. Two pieces of lettuce from each treatment were analyzed on each day of analysis. The blender was rinsed with 70% ethanol and sterile deionized water to prevent cross-contamination between each sample. A volume of 1 ml of the lettuce homogenate was surface-plated (0.25 ml on four plates) on sorbitol MacConkey agar (SMAC) (Becton Dickinson) supplemented with ampicillin (SMACA), or homogenates were serially diluted in 0.1% peptone water and then enumerated (0.1 ml, in duplicate) on SMACA. Plates were incubated at 37°C for 24 h before enumeration, and fluorescent colonies of E. coli O157:H7 gfp86, from lettuce samples were counted under UV light (395 nm). Colony counts were converted to log CFU/cm² for statistical analysis. The limit of detection was 0.22 log CFU/cm². Previous work has indicated that separating bacteriophages by centrifugation does not affect the bacteriophage titers; therefore this was not performed in this experiment.23

Bacteriophage enumeration. Bacteriophage concentrations were enumerated from lettuce or solutions containing EcoShield™ by soft agar overlay methods. Lettuce pieces treated with bacteriophages by either immersion or spray but not inoculated with E. coli O157:H7 were homogenized as described above either on Day 0 or after appropriate storage duration at 4°C. Homogenates were then serially diluted and mixed with 0.1 ml of stationary phage cultures of E. coli (held at 50°C), and poured onto a TSAA plate. After soft agar had solidified, plates were incubated at 37°C for 24 h. Plaques resulting from bacteriophage lysis of the E. coli O157:H7 lawn were counted and titers were converted to log PFU/ml or log PFU/g.

Statistical analysis. All experiments were repeated at least three times. Populations of E. coli O157:H7 gfp 86 recovered from lettuce homogenates treated with either control or EcoShield™ were subjected to an analysis of variance and Duncan’s least-significant-difference test. The data were analyzed using Statistical Analysis Software (SAS 9.1.2).

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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