Organic acid based sanitizers and free chlorine to improve the microbial quality and shelf-life of sugar snaps

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ARTICLE INFO
Article history:
Received 7 June 2013
Received in revised form 29 August 2013
Accepted 14 September 2013
Available online 21 September 2013

Keywords:
Sugar snap
Microbial quality
Organic acid
Chlorine
Decontamination
Water disinfection

ABSTRACT
A screening in a sugar snap packaging company showed a converged build-up of aerobic psychrotrophic plate count (APC) (ca. 6.5 log CFU/100 mL), yeasts and molds (Y&M), and lactic acid bacteria (LAB) (both ca. 4.5 log CFU/100 mL) in the wash water in the absence of water sanitizer, and a low build-up of chemical oxygen demand (30 ± 5 mg O2/L) and turbidity (5.2 ± 1.1 NTU). Decontamination experiments were performed in the lab with Purac FCC 80® (80% L(+)-lactic acid), two other commercial water sanitizers based on organic acids (NATRApHASE-ABAV®, and NATRApHASE-FVS®) and chlorine to evaluate their performance in reduction of the sugar snap microbial load as well as their functionality as disinfectant of the wash water to avoid cross-contamination. An additional 1 log reduction of APC on the sugar snaps was achieved with lactic acid in the range 0.8 to 1.6%, ABAV 0.5 to 1.5% and free chlorine 200 mg/L when compared to a water wash, while no significant difference in the numbers of Y&M was obtained when washing in sanitizer compared to water. There was no significant influence of the studied concentration and contact time on decontamination efficiency. Treatment with lactic acid 0.8% resulted in a lower APC contamination on the sugar snaps than on the untreated and water washed samples for 10 days. Chlorine 200 mg/L was the only treatment able to maintain the Y&M load lower than the untreated samples throughout the entire storage duration. The use of water sanitizers could not extend the sensorial shelf-life. Microbial loads were not indicative/predictive of visual microbial spoilage (shelf-life limiting factor), whereas maturity and amount of damage at the calyx end of the pods were.

The APC wash water contamination (5.2 log CFU/100 mL) was reduced significantly by chlorine 20 to 200 mg/L (to 1.4 log CFU/100 mL), ABAV 0.5 to 1.5% (to 2.7 log CFU/100 mL), FVS 0.5% (to 2.7 log CFU/100 mL) and lactic acid 0.8 to 1.6% (to 3.4 log CFU/100 mL). Only the use of chlorine enabled the reduction of the Y&M wash water contamination significantly (from 3.4 to 1.4 log CFU/100 mL). The low physicochemical build-up of the sugar snap wash water during the industrial washing process makes free chlorine attractive as a water disinfectant to prevent bacterial and fungal cross-contamination, whereas the sanitizers based on organic acids are not, due to their weak water disinfection efficiency.

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1. Introduction

Most grown sugar snaps (Pisum sativum var. macrocarpon) in the world are produced for local markets. In the last decades however, there has been a rise in the production of non-traditional export crops, including sugar snaps. Industrialized countries import large quantities of sugar snaps from tropical developing countries (such as Kenya and Guatemala), in order to have a year round supply and because of the high labor costs involved with picking (Humphrey et al., 2004; Messiaen et al., 2004). The main spoilage microorganisms on beans and peas are Pythium butleri, the fungal plant pathogens Rhizoctonia solani, Sclerotinia spp., and Botrytis cinerea and the pectinolytic bacterium Erwinia carotovora that break down the pectic substances of the middle lamella, with consequential loss of mechanical protection and rigidity (Brumwell, 2006; Tournas, 2005; Walker et al., 1998). The production of acid or antimicrobial compounds by native microbial flora may interfere with the colonization, survival and proliferation of foodborne pathogens (a.o. Salmonella spp., pathogenic Escherichia coli, Listeria monocytogenes) (Johnston et al., 2009; Liao and Fett, 2001; Shi et al., 2009; Teplitski et al., 2011). On the other hand, the chances of bacterial pathogen proliferation on and internalization in fresh produce are improved by the disruptive actions of certain fungal and bacterial spoilage microorganisms on the plant tissues (Brandl and Sundin, 2013; Critzer and Doyle, 2010; Ryser et al., 2009). Sugar snaps from Guatemala, which can be consumed either raw or cooked, have been the suspected vector of a Shigella dysenteriae outbreak in Sweden (May–June 2009) (Lofdahl et al., 2009) and a second Cyclospora cayetanensis outbreak

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http://dx.doi.org/10.1016/j.ijfoodmicro.2013.09.007
(June 2009) was also reported in Sweden, with sugar snaps from Kenya as the suspected source (Insulander et al., 2010).

Organic acids applied at relatively low concentrations exhibit inhibitory effects on microbial growth and are used to preserve acid foods and beverages. At higher concentrations organic acids can be used as decontaminants of food products such as fruits and vegetables and meat carcasses to improve food safety and quality (Virto et al., 2006). Organic acids are weak acids, and therefore they exist in a pH dependent equilibrium between the dissociated and undissociated states. The uncharged, undissociated acid can diffuse across the plasma membrane of microorganisms. Inside the cell the organic acid deprotonates, causing a pH drop and accumulation of toxic anions. As such, membranes can be disrupted, the proton motive force dissipated, essential metabolic reactions inhibited, and the intracellular pH homeostasis stressed (Brul and Coote, 1999; Capozzi et al., 2009).

Chlorine is the most used water disinfectant in fresh produce washing processes because of the low cost, the proven ability to rapidly inactivate suspended bacteria, and the minimal impact on the nutritional and sensorial fresh produce quality. Drawbacks of chlorination are the possibility of chlorine gas generation in the working environment when incorrectly applied (i.e. below pH 5 and excessive dosing), the rapid decomposition in the presence of organic matter, and most notoriously the possibility of creating harmful disinfection by-products in the wash water. However, studies on uncut carrots and fresh-cut lettuce have shown that only negligible or undetectable amounts of disinfection by-products were detected on the final product when a final rinse with tap water is applied (Klaiber et al., 2005; Van Haute et al., 2013).

After transport by airplane or container ship, sugar snaps are washed for rehydration and removal of materials from the pod surface. The objective of this study was to evaluate the use of water sanitizers for the reduction of the sugar snap microbial load and extension of shelf-life as well as their functionality as disinfectant of the wash water to prevent cross-contamination. To the knowledge of the authors, hitherto, no studies have been published regarding the use of water sanitizers to improve the microbial quality of sugar snaps, extend the shelf-life and maintain the wash water quality of sugar snap wash water. As decontamination efficiency depends in part on the produce surface and the way microorganisms attach to it, i.e. presence of stomata, surface roughness and hydrophobicity of the produce (Gomez-Lopez et al., 2008), the lack of knowledge on decontamination of pod vegetables (especially of snow peas and sugar snaps which can be consumed raw) makes sugar snap decontamination a topic of interest. In this study, an on-site screening of an industrial sugar snap washing process in the absence of water sanitizers was performed in a packaging company to observe the evolution of microbial and physicochemical parameters in function of processing time. In a second step, lab-scale experiments were performed with commercial formulations based on organic acids as produce and wash water sanitizers, and with chlorine as reference method.

2. Materials and methods

2.1. The sugar snaps

The sugar snaps used in the experiments originated from Peru and Guatemala. They were transported to the packaging company by container ship during 19 to 22 days at 3 ± 1 °C in modified atmospheric packaging (O2 < 10%, CO2 > 0.5%, Xtend®, StePac, Israel). Experiments were performed on 5 different batches (1 from Peru for the evaluation of the washing process in the packaging company and 4 for the decontamination experiments, of which 3 from Peru and 1 from Guatemala) that were sampled at different dates in the period October–December 2012. The acquired sugar snaps for the decontamination trials were as they were delivered to the packaging company, i.e. unwashed in crates of 4.5 kg.

The state of the pod at the calyx end, the amount of mechanical damage on the sugar snaps and the size of the seeds in the pod were compared among batches at reception, in order to be able to observe the impact of these characteristics on microbial number and growth and the onset of visual microbial spoilage. Seed size is an indicator of maturity status. In the immature state, seeds do not fill the hull, in the mature state they fill the hull without deforming it, and in the overmature state they deform the hull (Basterrechea and Hicks, 1991).

2.2. Evaluation of the washing process in the packaging company

The packaging company applied a bubble washer of 750 L volume with a replenishing rate of 400 L/h. 1000 kg of sugar snaps was washed in 188 min, air dried, screened with machine vision to remove pods showing excessive browning, and packaged in 300 g consumer units. At several time points during the washing process, samples of both sugar snaps before and after washing (after 0, 18, 54, 96, and 188 min) and of the wash water (after 0, 10, 18, 30, 54, 96, 120, 188 min) were taken. Also, samples of the tap water were taken at the point of entrance in the washing bath. Temperature, pH, and conductivity (all with HQ40d meter, HACH LANGE, Belgium) of the wash water were measured at the packaging company. The residence time of the sugar snaps in the washing bath was measured (n = 12) by labeling individual sugar snaps with fluorescent tape and timing the period from entrance till exit from the washing bath. The samples were transported under refrigerated conditions to the lab for further analysis. Alkalinity, turbidity, and chemical oxygen demand (COD) of the wash water were determined. Water samples were also analyzed for aerobic psychrotrophic plate count (APC), yeasts and molds (Y&M), and lactic acid bacteria (LAB). Sugar snaps, collected before and after washing, were analyzed for moisture content and water activity (aW). Samples of the sugar snaps (250 g) were stored in plastic bags for 22 days at 5 ± 1 °C under normal atmospheric conditions and periodically sampled (after 0, 6, 10, 15, and 22 days) for APC, Y&M, and LAB and judged for onset of visual microbial spoilage (i.e. fungal rot, bacterial slime formation).

2.3. Evaluation of water sanitizers to improve shelf-life of sugar snaps and maintain wash water quality

Sodium hypochlorite (28.4 g/L NaOCl, La Croix, Belgium), acetic acid (Sigma-Aldrich, Belgium), Purac FCC 80® (80% L(+) lactic acid, Purac, The Netherlands), NATRAPHASe-ABAv® (fine powder containing natural acids, Natural Biotechnology, Belgium) and NATRAPHASe-FVS® (blend of EU and FDA food approved organic acids and vitamins, Natural Biotechnology, Belgium) were used as water disinfectants. The experimental disinfectant concentration–contact time settings are shown in Table 1. For Purac FCC 80, the added concentration is expressed as active compound, i.e. L(+) lactic acid. For chlorine, the pH was adjusted to 6.5 using HCI (1 M). Each experiment (i.e. disinfectant; concentration; contact time) was executed on three different batches in order to incorporate possible influence of variation in microbiology and physical and physiological differences among sugar snaps in different batches.

Portions of 250 g of sugar snaps were washed by mechanical agitation in 4 L of tap water (5 ± 1 °C) with added water disinfectant. After washing, water samples were immediately quenched with Na2S2O3 (0.1 M) for quenching sodium hypochlorite, or phosphate buffer (pH 7.5) for Purac FCC 80, acetic acid, ABAv and FVS. Microbial analyses (APC, Y&M, LAB) were performed on the water samples. The sugar snaps were rinsed (0.1 L/kg·s for 10 s) with tap water. At the highest exposure conditions in the experiment (i.e. highest contact time and disinfectant concentration), samples of washed sugar snaps were either rinsed or not rinsed to observe the effect of residual disinfectant on discoloration, off-odors, damage and texture loss. ABAv samples were never rinsed and FVS samples were always rinsed because these patented formulations were recommended to be used respectively with or without a final rinsing step by the manufacturers. All samples were dried with sterile absorbent paper, and subsequently, samples were screened for discoloration, and sugar snaps showing browning were discarded. The samples were stored
in plastic bags at 5 ± 1 °C for 22 days under normal atmospheric conditions and periodically sampled (after 0, 6, 10, 15, and 22 days) for microbial analyses (APC, Y&M, LAB), and at the same time monitored for the presence of visual microbial decay, discoloration, off-odors, damage and texture loss due to the decontamination treatments.

### 2.4. Physicochemical parameters

Alkalinity was determined with acid titration, turbidity with a turbidimeter (HI 98703; HANNA Instruments; Belgium), chemical oxygen demand (COD) according to the small-scale sealed-tube method (LCI 400; HACH LANGE; Belgium), and absorbance at UV 254 nm with a UV-spectrophotometer (UV-1601, Shimadzu, Belgium) and quartz cuvettes with a 1-cm path length (Helima, Belgium) after filtration through a 0.45 μm polytetrafluoroethylene filter (Macherey-Nagel, Belgium). aw of the sugar snaps was measured with a dew point water activity meter (AquaLab Series 4:4TE, Decagon Devices, The Netherlands). Moisture content of the sugar snaps was determined through homogenization of 5 g of sample (T18 Basic ULTRA-TURRAX, IKA, Germany) and drying in an air circulation oven of 105 °C for 3 h. Free chlorine was measured with the N,N-diethyl-p-phenylenediamine (DPD) colorimetric method (Eaton et al., 2005).

### 2.5. Microbial analyses

The sugar snap samples were prepared by weighing 25 g of sugar snaps in a sterile stomacher bag with full-surface filter (0.5 mm pore size) (VWR, Belgium) which was homogenized in 225 mL peptone water (Oxoid, Belgium) for 1 min. Sugar snaps were analyzed for APC, Y&M, and LAB. APC was enumerated with plate count agar (Oxoid, Belgium) using the pour-plate method (incubation at 22 °C, 5 days). Y&M were enumerated with Rose Bengal Chloramphenicol agar (Oxoid, Belgium) containing 150 mg/L chloramphenicol and using the spreading plate method (incubation at 22 °C, 5 days). Membrane filtration was used to lower the detection limit of Y&M to 1 log CFU/g. LAB were enumerated with MRS (De Man, Rogosa, Sharpe) agar (Oxoid, Belgium), containing 1.4 g/L sorbic acid and with a final pH of 5.7, adjusted with NaOH (1 mol/L), using the pouring plate method with an additional cover layer of agar (incubation at 22 °C, 5 days). The water samples were analyzed for the same microorganisms, using the same enumeration methods. In addition, membrane filtration of 10 or 100 mL water was used to lower the detection limit for microbial enumeration to respectively 1 or 0 log CFU/100 mL.

### 2.6. Statistics

Data analysis was performed with SPSS Statistics 21. Influence of disinfectant type, concentration and contact time was assessed with one-way ANOVA or Brown–Forsythe when equal variance could not be assumed. Group comparison was done with post-hoc tests (Tukey or Games–Howell) when all relations among groups were of interest. However, when only certain relations were of interest, i.e. a significant reduction of the wash water contamination, or a significantly lower contamination on the sugar snaps compared to water washed or untreated sugar snaps, simple contrast analysis was performed. A level of significance p ≤ 0.05 was chosen for all statistical analyses.

### 3. Results

#### 3.1. Evaluation of the washing process in the packaging company

The average residence time of the sugar snaps in the washing bath was 25 ± 15 s. The microbial contamination in the wash water increased till about 25 min of exploitation after which the wash water contamination remained relatively stable (Fig. 1). For APC and Y&M, the microbial contamination of the municipal water used in the washing bath immediately at the tap was significantly lower than the water samples taken in the washing bath immediately before the start of the washing process, indicating the presence of some microbial contamination on the washing equipment prior to the start of operation. The microbial load on the sugar snaps before washing (3.0 ± 0.8, 2.7 ± 0.4, 2.3 ± 0.5 log CFU/g for APC, Y&M, and LAB respectively) was not significantly different from that on the washed sugar snaps (3.5 ± 0.4, 2.6 ± 0.4, 2.2 ± 0.8 log CFU/g for APC, Y&M, and LAB respectively). The microbial contamination on the washed sugar snaps did not change significantly in function of...
exploitation time. The COD and the turbidity of the wash water were significantly correlated (Spearman’s rho = 0.668; p = 0.005). Both became relatively stable from about an hour of exploitation until the end of operation: COD of 30 ± 5 mg O₂/L and turbidity of 5.2 ± 1.1 NTU (Fig. 2). The temperature was 7.7 ± 0.7 °C, the pH 8.0 ± 0.1, the conductivity 425 ± 4 μS/cm, and the alkalinity 3.05 ± 0.02 mmol/L bicarbonate. All these parameters did not significantly change in function of processing time. The washing process increased the water content of the sugar snaps (from 81.6 ± 3.5% to 86.0 ± 1.8%, p = 0.304), though not significantly. Also, the aw increased significantly (from 0.986 ± 0.001 to 0.990 ± 0.001, p = 0.004).

3.2. Microbial and visual quality of untreated sugar snaps at reception and during storage

The variation of APC and LAB among different batches of sugar snaps at reception and during storage was more pronounced than for the Y&M contamination (Fig. 3). The onset of visual microbial decay was not directly related to the overall microbial contamination degree of the sugar snaps. As the LAB were below or close to the detection limit (1 log CFU/g) in some batches, it was hard to make statistical claims concerning disinfection of LAB and therefore no such conclusions were made. The initial seed size, and to a lesser degree the integrity of the calyx end, seemed to have an impact on the onset of visual microbial decay (Table 2). Violation of calyx end integrity became most apparent through brown discoloration and loss of firmness. The visual microbial decay manifested itself in the pod tissue towards the calyx end and on major mechanical wounds. Except for batch 1 where all samples and most individual sugar snaps within samples showed signs of microbial decay, the onset of microbial decay in the other batches was mostly only visual on 1 sugar snap within a decaying sample (comprising ca. 20–50 remaining sugar snaps dependent on the storage time), which at a later date could become visible on one or more other sugar snaps.

3.3. Evaluation of water sanitizers to improve shelf-life of sugar snaps

Certain of the tested disinfectant settings caused damage to the sugar snaps, i.e. brown discoloration and formation of irregularities on the pod surface, more specifically, pit formation. In addition, acetic acid caused off-odors (Table 1). The data of the settings that caused damage to the product were not further incorporated in the shelf-life analyses. A water wash did significantly lower the concentration of Y&M (0.6 ± 0.4 log reduction; p = 0.003) on the sugar snaps, yet not of APC (0.5 ± 0.7 log reduction; p = 0.059) (Table 3). However, it is important to not blindly accept the statistical analysis. Reductions of 0.5 log compared to the untreated samples are very low in microbiological terms, both from the point of food spoilage (no considerable impact on the regrowth) and of food safety (the human dose–response and the associated increase in risk) (FDA, 2001). Duration of washing had no influence

<table>
<thead>
<tr>
<th>Batch</th>
<th>Maturity (seed size)ᵃ</th>
<th>Damage to calyx endᵇ</th>
<th>Other mechanical damageᶜ</th>
<th>Visual microbial decay (days storage)ᵈ</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 I-M</td>
<td>+</td>
<td>+</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>1 OM</td>
<td>+++</td>
<td>+</td>
<td>6 (3/3)ᵇ</td>
<td>/</td>
</tr>
<tr>
<td>2 M</td>
<td>+</td>
<td>++</td>
<td>10 (1/3)</td>
<td>/</td>
</tr>
<tr>
<td>3 M</td>
<td>++</td>
<td>++</td>
<td>15 (1/3), 22 (2/3)</td>
<td>/</td>
</tr>
<tr>
<td>4 I</td>
<td>+</td>
<td>+</td>
<td>/</td>
<td>/</td>
</tr>
</tbody>
</table>

ᵃ I: immature, M: mature, OM: overmature.
ᵇ Number of ’+’ expresses relative severity of characteristic.
ᶜ No visual microbial decay observed in 22 days of storage.
ᵈ Fraction of samples that showed decay.

Fig. 2. COD (♦) and turbidity (■) in the washing bath during sugar snap washing. The data points at t = −25 min show the COD/turbidity of the used tap water. Error bars denote standard deviation (n = 3).

Fig. 3. Microbial load of the untreated sugar snaps in function of experimental batch and storage time. Numbers in the graphs indicate the batch on which visual microbial decay was observed at that storage time. Error bars denote standard deviation (n = 3).
on reduction efficiency of any of the washing treatments, including a water wash, and these values were pooled to increase sample sizes for statistical analysis. Lactic acid in the range 0.8 to 1.6% (1.4 ± 0.5 log reduction), ABAV 0.5% (1.6 ± 0.2 log reduction), and free chlorine 200 mg/L (1.4 ± 0.5 log reduction) caused a significantly higher reduction of APC than a water wash (Table 3). In the studied concentration ranges, there was no relation between concentration and decontamination efficiency of APC for chlorine (p = 0.648) and FVS (p = 0.759) and some, yet no significant relation for ABV (p = 0.069) and lactic acid (p = 0.057). None of the decontamination treatments removed Y&M significantly more effective from the sugar snaps than a water wash. Free chlorine (range 20 to 200 mg/L) had the highest reduction of Y&M (on average 1.0 ± 0.7 log reduction).

None of the treatments maintained the APC contamination lower than the untreated and water washed samples for the whole storage duration (Table 3). Treatment with lactic acid 0.8% resulted in a lower APC contamination on the sugar snaps than on the untreated or water washed samples for 10 days. Chlorine 200 mg/L was the only treatment able to maintain the Y&M load lower than the untreated samples throughout the entire storage duration. For the other treatments, any significance in microbial reduction on the sugar snaps was lost in less than 10 days of storage.

Visual microbial decay occurred more or equally rapid on untreated than treated (including water washed) sugar snap samples (Table 4). Disinfection concentration had no effect on delaying the occurrence of visual microbial decay. In batches 1 and 2, the samples which showed microbial decay had a more rapid growth of APC than the other samples, and overall high counts were reached during storage (Fig. 4). Batch 3 showed similar visual microbial decay as batches 1 and 2, but a relatively high proportion of batch 3 was Y&M, and no differences in counts between decayed and other samples were observed for Y&M or APC. Batch 4 also had a high relative abundance of Y&M, yet none of the samples of batch 4 showed any visual microbial decay during storage. The visual decay manifested itself in the same way as with the untreated samples. There was no significant difference in disinfection efficiency of the disinfectants between the different treated batches, despite the differences in initial microbial load as well as microbial growth during storage (Fig. 4).

3.4. Evaluation of water sanitizers to maintain the wash water quality

Washing sugar snaps for up to 3 min had only minimal influence on the physicochemical water quality: turbidity increased from 0.41 ± 0.03 to 1.16 ± 0.71 NTU and absorbance at UV 254 nm (0.45 μm filtered) from 0.020 ± 0.003 to 0.047 ± 0.015. The pH value of the washing solutions did not change significantly after 3 min washing, the free chlorine concentration diminished 1.46 ± 0.08 mg/L when adding 20 mg/L free chlorine and no significant changes were observed when washing with 200 mg/L free chlorine for 3 min. The initial microbial load of the used tap water was 3.6 ± 1.0 log CFU/100 mL APC and 0.5 ± 0.6 log CFU/100 mL Y&M. The degree of microbial contamination transferred from the sugar snaps to the water during washing in water was independent of both washing time and experimental batch.

The washing time had no significant influence on the water disinfectant efficiency to lower the wash water contamination in any of the washing setups, so these values were pooled to increase sample sizes for statistical analysis. On the other hand, the disinfectant concentration had a significant influence on the water disinfectant efficiency of APC and Y&M for ABV and FVS, although for chlorine (20 to 200 mg/L) and lactic acid (0.32–1.6%) this was not the case (Fig. 5). The APC wash concentration diminished 0.18 ± 0.06 log CFU/100 mL when adding 20 mg/L free chlorine and no significant changes were observed when washing with 200 mg/L free chlorine for 3 min. The initial microbial load of the used tap water was 3.6 ± 1.0 log CFU/100 mL APC and 0.5 ± 0.6 log CFU/100 mL Y&M. The degree of microbial contamination transferred from the sugar snaps to the water during washing in water was independent of both washing time and experimental batch.

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0.8 log CFU/100 mL), whereas the Y&M wash water contamination (3.4 ± 0.6 log CFU/100 mL) was only reduced significantly by chlorine 20 to 200 mg/L (to 1.4 ± 0.5 log CFU/100 mL). Y&M were more resistant to water disinfection with organic acids than APC. For chlorine, both Y&M and APC were reduced to similar numbers and one could argue that the remaining microorganisms were mostly Y&M.

Lactic acid in the range 0.32 to 1.6% reduced APC in the water and on the sugar snaps with on average 1.3 ± 0.6 and 1.3 ± 0.5 log respectively and Y&M with on average 0.1 ± 0.4 and 0.6 ± 0.4 respectively. For the commercial sanitizers based on organic acids, ABAV and FVS, the same pattern was observed, i.e. no significant difference between disinfection efficiency of sugar snaps and water disinfection efficiency, which contrasts the much better performance of free chlorine to inactivate microorganisms in the suspended state (Wilcoxon signed rank test; p > 0.05 for lactic acid, FVS, and ABAV, p < 10E − 5 for chlorine).

4. Discussion

Visual microbial spoilage was the limiting factor of shelf-life. Nonetheless, APC and Y&M numbers were ineffective to indicate this visual microbial spoilage, as a large variation in microbial counts of the sugar snaps was found. The heterogeneity among microbial contamination of individual sugar snaps and the scarcity of sugar snaps that actually show signs of microbial spoilage make it unlikely to pinpoint excessive

<table>
<thead>
<tr>
<th></th>
<th>Batch 1</th>
<th>Batch 2</th>
<th>Batch 3</th>
<th>Batch 4</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Day</td>
<td>Day</td>
<td>Day</td>
<td>Day</td>
</tr>
<tr>
<td>Untreated</td>
<td>3/3a</td>
<td>1/3</td>
<td>1/3</td>
<td>0/3</td>
</tr>
<tr>
<td>Water</td>
<td>3/3</td>
<td>0/3</td>
<td>0/3</td>
<td>0/3</td>
</tr>
<tr>
<td>Chlorine</td>
<td>12/12</td>
<td>1/12</td>
<td>0/4</td>
<td>0/8</td>
</tr>
<tr>
<td>Lactic acid</td>
<td>12/12</td>
<td>0/12</td>
<td>2/8</td>
<td>0/4</td>
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<td>ABAV</td>
<td>/b</td>
<td>0/9</td>
<td>3/9</td>
<td>7/9</td>
</tr>
<tr>
<td>FVS</td>
<td>/</td>
<td>0/9</td>
<td>2/9</td>
<td>6/9</td>
</tr>
</tbody>
</table>

a Fraction of samples that showed visual microbial decay.
b Was not executed in that batch.
microbial growth through sampling. Also, APC and Y&M generally do not provide information about individual species and the growth of specific spoilage microorganisms might be masked by these broad-spectrum microbial analyses (Gram et al., 2002). Nonetheless, identifying and measuring the specific spoilage microorganisms would not solve the problem of variation in microbial counts within a batch as observed in this study. Although some studies observed that the level of total microbial counts or specific spoilage microorganisms in fresh produce were not related to the product quality and shelf-life (Allende et al., 2008; Bennik et al., 1998; Gimenez et al., 2003; Ragaert et al., 2007; Zagory, 1999), others have shown a good correlation between sensory shelf-life and microbial numbers, such as Chen et al. (2010) (correlation with aerobic mesophilic and psychrotrophic bacteria and yeasts and molds on fresh-cut asparagus lettuce) and Jacxsens et al. (2003) (correlation with yeasts and lactic acid bacteria on mixed bell peppers and grated celeriac). The type of microbial spoilage and sensory quality deterioration depends on the type of fresh produce (Jacxsens et al., 2003). It is plausible that increased understanding of the specific spoilage microorganisms, spoilage mechanisms, and produced metabolites in a certain type of fresh produce, will enable a better prediction of shelf-life through microbial measurements, not considering the microbial variability issues observed in this study.

Characteristics explaining physical damage and the physiological status of the sugar snaps, i.e. the maturity of the sugar snaps and the integrity of the pod at the calyx end, were more predictive towards the visual shelf-life of the sugar snaps. This illustrates that harvesting at the right stage of maturity, avoiding damage during harvesting and more thorough visual selection before processing could lead to an end product with longer shelf-life. Fungal spoilage usually originates from latent infections established in the field or wound infections during harvesting and handling (Terry and Joyce, 2004). *Pseudomonas* spp. and *Erwinia* spp. that colonize plant surfaces adhere preferentially in the natural depression of stomata or in the intercellular junction, or cracks or crevices formed through damage, after which diverse biofilms can arise, composed of gram-negative and gram-positive bacteria, yeasts and filamentous fungi (Carmichael et al., 1999). The weakening natural defense mechanisms of overmature or mechanically damaged sugar snaps and the loss of structural integrity at the calyx ends, potentially leading to increased solute leakage, improve the growth conditions of phytopathogens (Elghaouth et al., 1992; Nunes et al., 2010). Regardless of the observed microbial related issues of overmature sugar snaps, sugar snaps should always be harvested before physiological maturity, becoming tougher and fibrous (Basterrechea and Hicks, 1991; Sams, 1999).

Chlorine was confirmed to be an efficient, fast acting water disinfectant against vegetative bacteria as observed in previous studies (Lee et al., 2010; Luo et al., 2011; Van Haute et al., 2013). Chlorine also effectively removed yeasts and molds from the wash water, but did not significantly enhance the Y&M reduction on the sugar snaps compared to a water wash. Pereira et al. (2013) reported fungi to be more resistant to chlorination in drinking water than bacteria and viruses, but less resistant than Cryptosporidium oocysts. Beuchat et al. (1998) suggested a large abundance of chlorine resistant cell types among fungi. Contrary to its efficiency to remove suspended, vegetative microorganisms, chlorine is much less efficient as fresh produce decontaminant, a behavior shared among the chemical water disinfectants chlorine, chlorine dioxide, and ozone and observed in a myriad of studies. Decontamination processes are compromised by the presence of microorganisms in biofilm, attachment to and internalization through surface wounds and stomata, internalization through the plant roots and subsequent migration throughout the plant, and increasing surface roughness of the produce at microscopic level (Gomez-Lopez et al., 2008; Huang et al., 2006; Jahid and Ha, 2012; Luo et al., 2011; Takeuchi and Frank, 2001; Wang et al., 2009). Although gaseous chlorine, chlorine dioxide, hydrogen peroxide and ozone have a higher diffusion capacity than when dissolved in water and have a higher potential for decontaminating injured and other hard to reach produce surfaces, gaseous disinfectants do not solve the problems of microbial internalization in fresh produce (Gomez-Lopez et al., 2008). Han et al. (2001) observed an increased reduction of spot inoculated *L. monocytogenes* of 3 log reduction when applying chlorine dioxide as gas treatment (3 mg/L, 10 min, 20 °C) compared to an aqueous chlorine dioxide treatment (3 mg/L, 10 min, 20 °C) of both uninjured and injured green peppers. However, Hadjok et al. (2008), who used vacuum infiltration in order to achieve internalization of inoculated *Salmonella* Montevideo in fresh-cut iceberg lettuce, observed that gas exposure of the produce to 1.5% H₂O₂ at 50 °C resulted in 2 log reduction on the lettuce, whereas only 0.5 log of the internalized *Salmonella* Montevideo was inactivated.

The behavior of weak organic acids is fundamentally different from chemical oxidants such as chlorine as weak organic acids are not compromised as severely when inactivation of microorganisms is needed in the presence of organics in the water or food matrix, or exopolymeric substances in biofilms. Where chlorine is decomposed through reaction with organic matter, the loss of the ‘active substance’ of weak organic acids is synonymous with deprotonation, and buffer capacity in the vicinity of the produce surface, as well as alkalinity of the water (both inorganic and organic such as from anions of organic acids with pKa ≥ 4) could theoretically pose a disinfection barrier (Hemond, 1990). However, the alkalinity did not change significantly during the 3 hour washing trials.

![Fig. 5. Microbial wash water contamination during sugar snap decontamination experiments. Error bars denote standard deviation (n = 9).](image-url)
in the packaging company. Also, during washing of fresh-cut radicchio, sugar loaf, and endive, a process which generates a higher converged COD ($295 \pm 8 \text{ mg O}_2/\text{L}$), the alkalinity was stable throughout the 135 min washing process at on average 6.38 $\pm$ 0.12 mmol/L (own data not published). Unless considerable amounts of buffering substances are introduced during washing, efficiency of organic acids will not be severely influenced by the water matrix during produce washing operations. However, weak organic acids in general are inefficient water disinfectants, and the results in this study show that, given the contact times used, the efficiency to inactivate microorganisms in suspended state is not better than the reduction of microorganisms on sugar snap surfaces. Virto et al. (2006) modeled the inactivation of *L. monocytogenes* and *E. coli* in function of concentration of citric or lactic acid, temperature and contact time. To achieve a 3 log reduction in sterile distilled water at 5 °C and with 1.6% lactic acid (the most severe lactic acid settings applied in this study) would take 25 min and 35 min of contact time for *E. coli* and *L. monocytogenes* respectively. In this study, the short contact times applied (range 30 to 180 s), in combination with the experimental variability, masked the water disinfection kinetics of lactic acid. For comparison of lactic acid and free chlorine as water disinfectants, according to Chick–Watson kinetics, it would take 10 s for 1 mg/L free chlorine at pH 6.5 in oxidant demand free buffer to reduce *E. coli* O157 by 3 log (Van Haute et al., 2013). Lactic acid, PBS, and ABAV failed at effectively reducing Y&M, both on the produce and in the wash water. The resistance of spoilage fungi to organic acids is related to the membrane ATPase activity and pH homeostatic mechanisms such as acid anion efflux pumps (Brul and Coote, 1999; Smits and Brul, 2005).

Within the studied parameter ranges, contact time had little influence on the decontamination effectiveness of the sugar snap decontamination treatments. The same was observed for concentration, except for ABAV and lactic acid which showed increased (yet not significant) reduction of APC with increased concentration. The decontamination behavior in function of time and concentration and observed in numerous studies can be described as the following: the microbial load can initially be reduced quite effectively with limited exposure (concentration $\times$ contact time), after which increased exposure is less successful in achieving further microbial reduction (Akbas and Olmez, 2007b; Ayhan et al., 1998; Beuchat et al., 1998; Chen and Zhu, 2011; Mahmoud et al., 2008; Olmez and Akbas, 2009). This again can be explained by the state/location of the microorganisms on fresh produce, comprising of microorganisms that are easily, hard, or virtually impossible to inactivate with water disinfectants. Easily removable microorganisms that are vulnerable against the respective disinfectant require relatively little exposure, whereas those which reside in thick biofilms and stomata, require a much higher exposure. The severity of the exposure might be limited by produce damage or engineering issues such as long duration of produce washing steps, or might be virtually futile in the case of internalized microorganisms. The lack of influence of concentration on decontamination efficiency observed in this study can be explained by i) working in a concentration range in which all concentrations inactivated the easily reachable microorganisms, ii) the lack of disinfection efficiency of a certain disinfectant to remove hard to reach microorganisms, iii) the masking of the possible influence of concentration on decontamination efficiency of difficult to remove microorganisms due to the high variability in microbial counts, and iv) the presence of recalcitrant internalized microorganisms. The lack of influence of contact time on decontamination efficiency could be explained by i) working in a too small range of contact times to observe differences, ii) decontamination kinetics of disinfectants (suppose that contact times $>30$ s would result in no further significant inactivation), or iii) interference of high variability in microbial counts. Some studies (Akbas and Olmez, 2007b; Ayhan et al., 1998; Beuchat et al., 1998; Olmez and Akbas, 2009) show a more severe limitation of further fresh produce decontamination (and as such less influence of concentration and contact time beyond the initial effective decontamination stage) than others (Chen and Zhu, 2011; Mahmoud et al., 2008) in which further increase in exposure resulted in a more successful further microbial reduction. Different inactivation behaviors can be due to several causes: i) fresh produce type and whole vs fresh-cut produce, ii) inoculation method or naturally present microflora, iii) the microorganism type, iv) (related to i, ii, and iii) the relative abundance of easily reachable, hard to reach, and infiltrated microorganisms, v) characteristics of the disinfectant (inherent disinfection potential and disinfection kinetics, liquid or gas form), and vi) the applied experimental conditions and execution such as the created turbulence during the washing process.

Except for treatment with lactic acid 0.8% or chlorine 200 mg/L, gained reductions of the other treatments compared to the untreated sugar snaps were lost in less than 10 days of storage. Microbial regrowth can potentially occur quickly after decontamination due to reduction of competition (Delaquis et al., 1999; Gomez-Lopez et al., 2008; Ragaert et al., 2007). In this study, free chlorine, lactic acid, and ABAV were more effective than a water wash for reduction of APC but not of Y&M on the sugar snaps. Comparison of organic acids and chlorine as fresh produce decontaminants to reduce spoilage microorganisms has also been studied on rocket leaves (Martinez-Sanchez et al., 2006), fresh-cut iceberg lettuce (Akbas and Olmez, 2007a; Allende et al., 2008), fresh-cut escarole (Allende et al., 2008) and fresh-cut cilantro (Allende et al., 2009). Based on those studies, there is no clear, discernable pattern as to whether Y&M are less efficiently removed from fresh produce than mesophilic or psychrotrophic counts, whether lactic acid or citric acid is more/less efficient than chlorine to remove fungal or bacterial microorganisms from fresh produce, and whether these disinfectants improve the shelf-life of the produce.

The consequences of slow water disinfection kinetics by organic acids, as confirmed in this study, are that organic acids cannot be used to control cross-contamination, which Lopez-Galvez et al. (2009) demonstrated for *E. coli* transfer from inoculated to non-inoculated fresh-cut iceberg lettuce during washing with 2% Purac or 0.5% Citrox®. Therefore, it seems that organic acids are not suitable for washing applications of fresh produce, although there might be potential for their use in decontamination applications through spraying or electrostatic spraying on fresh produce (Ganesh et al., 2010, 2012), as such bypassing the low water disinfection efficiency by using a method without water immersion. This especially has potential when applied as a warm/hot spray, as research on *E. coli* and *L. monocytogenes* suspended in water (4 °C vs 20 °C vs 40 °C) as well as *E. coli* O157:H7 inoculated on baby spinach (22 °C vs 40 °C) has shown that the disinfection efficiency of lactic acid is significantly enhanced by increased temperature (Huang and Chen, 2011; Virto et al., 2006).

Washing of whole produce such as sugar snaps, introduces exudates in the wash water (most probably from wounded surfaces) to a much lesser extent than washing of fresh-cut produce. As such, the transfer of organic materials depends in greater part on foreign organics and particles present on the sugar snaps. The converged COD values (30 ± 5 mg O$_2$/L) in this study were low compared to the converged COD measured in two fresh-cut leafy vegetable companies by Van Haute et al. (2013), COD 465 ± 2 and 1405 ± 57 mg O$_2$/L. Therefore, considering the high microbial build-up during washing, the inability of the tested water sanitizers to prolong the shelf-life, the absence of detrimental effects of chlorine on the sensorial quality of the sugar snaps, the high performance of free chlorine as a water disinfectant, and the low physicochemical load of the sugar snap wash water which would minimize the disinfection by-product generation, maintaining a free chlorine residual seems to be a suitable strategy to avoid cross-contamination of vegetative bacteria and fungi in the washing process of sugar snaps.

**Acknowledgments**

The research leading to these results has been facilitated by the European Community’s Seventh Framework Program (FP7) under grant agreement no 244994 (project VEGI-TRADE). The authors would like to thank the Howeest-ALGent Master Students Ferielle Leveque and Rens Piccavet for the practical assistance.
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