

1 **WASH WATER DISINFECTION OF A FULL-SCALE LEAFY VEGETABLES**
2 **WASHING PROCESS WITH HYDROGEN PEROXIDE AND THE USE OF A**
3 **COMMERCIAL METAL ION MIXTURE TO IMPROVE DISINFECTION**
4 **EFFICIENCY**

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21 **ABSTRACT**

22 Hydrogen peroxide (H₂O₂) was used to maintain the microbial wash water quality of a full-scale
23 leafy vegetables (radicchio, sugar loaf, curled endive, lollo, lollo rosso) wash water process.
24 Despite addition of 300 L/h of 1.8 % H₂O₂ to a 450 L washing bath (333 ± 50 kg/h fresh-cut
25 produce introduction speed), the H₂O₂ quickly decreased and a lower wash water contamination of
26 aerobic psychrotrophic plate count (APC) and enterococci than without addition of H₂O₂ could not
27 be maintained. There was no significant difference between the APC on fresh-cut leafy vegetables
28 washed with H₂O₂ and those washed with water.

29 In a second part, lab-scale experiments were performed to assess the impact of a commercial metal
30 ion formulation (Bacsan®, containing a. o. Cu²⁺, Zn²⁺, Ag⁺) on the stability of H₂O₂ in artificial
31 wash water, made from iceberg lettuce and tap water. Bacsan improved the stability of H₂O₂ in
32 artificial lettuce wash water and fresh-cut leafy vegetables wash water from a processing company
33 and synergistically increased the disinfection efficiency of APC and *Escherichia coli* (*E. coli*)
34 compared to H₂O₂ or Bacsan. Increasing chemical oxygen demand (COD) had detrimental effect
35 on the H₂O₂ stability and disinfection efficiency. Addition of Ag⁺ to Bacsan further synergistically
36 enhanced the H₂O₂ stability.

37 H₂O₂ is not suited as an *in situ* wash water disinfectant to avoid cross-contamination in fresh-cut
38 leafy vegetables washing processes due to the slow water disinfection kinetics and the rapid H₂O₂
39 consumption. However, H₂O₂/Bacsan shows potential for use in off-line processes.

40

41 **1. INTRODUCTION**

42 Among fresh produce, leafy vegetables are one of the commodities most frequently implicated with
43 food disease outbreaks, the culprit most often being *E. coli* O157: H7 or *Salmonella* spp. (Olaimat,
44 & Holley, 2012; Tomas-Callejas et al., 2012). Washing of fresh-cut lettuce is often the only
45 processing step able to reduce the microbial load (Artés et al., 2009). Current washing treatments
46 with the purpose of decontaminating fresh-cut produce for microbial safety or quality reasons, have
47 evolved from processes that were originally developed to remove soil from whole produce, to a
48 water disinfection process for removal of microbial targets from fresh-cut produce (Sapers, 2001).
49 The success of these washing processes to remove naturally present microorganisms from fresh-
50 cut produce is limited (1-3 log reduction), i.e. microbial reductions occur but total removal cannot
51 be achieved. The access of sanitizers to the target microorganisms is hindered by the presence of
52 microorganisms in biofilms, attachment near and within stomata, and internalization through cut
53 surfaces and other tissue wounds. Therefore it is preferable to avoid contamination wherever
54 possible by implementing good agricultural and manufacturing practices during the production and
55 processing of fresh produce (Holvoet et al., 2012; 2013; Keskinen, Burke, & Annous, 2009; Lopez-
56 Galvez et al., 2010; Sapers et al., 2001). The post-harvest washing water is a vehicle for microbial
57 cross-contamination and to counter this an *in situ* wash water disinfection can be performed. Water
58 disinfection can also be used to treat the wash water before reusing it (i.e. reconditioning) for a
59 similar or different purpose. The efficiency of wash water disinfection is not limited by the issues
60 that plague decontamination, but the effectiveness of chemical oxidants (a. o. chlorine, chlorine
61 dioxide, ozone, H₂O₂, peracetic acid) is hindered by the presence of organic matter in the wash
62 water, the degree depending on the properties of the chemical oxidant (Van Haute, Sampers,
63 Jacxsens, & Uyttendaele, 2013b).

64 H₂O₂ does not produce toxic fumes in the worker space and is an environmentally friendly
65 alternative to chlorine for decontamination of fresh produce, as it breaks down in water and oxygen
66 (Tofant, Vucemilo, Pavicic, & Milic, 2006), and does not form carcinogenic disinfection
67 byproducts (USEPA, 1997; Van Haute, Sampers, Jacxsens, & Uyttendaele, 2014). Considerable
68 research has been conducted on the use of H₂O₂ as produce decontamination agent against bacterial
69 and viral indicator organisms, pathogenic bacteria, or spoilage microflora on fresh (-cut) fruit and
70 vegetables (Parish et al., 2003; Ukuku, Bari, & Kawamoto, 2012), among which some experiments
71 have been performed on leafy vegetables (Allwood, Malik, Hedberg, & Goyal, 2004; Hadjok,
72 Mittal, & Warriner, 2008; Huang, & Chen, 2011; Li et al., 2011; Lin, Moon, Doyle, & McWatters,
73 2002). On the contrary, its use as a water disinfectant to control the wash water quality of fresh
74 produce washing processes is virtually unexplored. Earlier water disinfection studies that focused
75 on inactivating vegetative bacteria, bacterial spores, viruses, or protozoa have shown that H₂O₂ by
76 itself is a slow acting water disinfectant, requiring high dosages and contact times for microbial
77 inactivation (Barbee, Weber, Sobsey, & Rutala, 1999; Raffellini, Guerrero, & Alzamora, 2008;
78 Raffellini, Schenk, Guerrero, & Alzamora, 2011; Toledo, Escher, Ayres, 1973; Weir et al., 2002).
79 Combined with Ag⁺ and Cu^{1 or 2+}, performance of H₂O₂ can be enhanced (Batterman, Zhang, &
80 Wang, 2000; Orta De Velasquez, Yanez-Noguez, Jimenez-Cisneros, & Luna Pabello, 2008;
81 Pedahzur et al., 2000; Pedahzur, Shuval, & Ulitzur, 1997).

82 In this study, the use of H₂O₂ to maintain the microbial wash water quality in a full-scale industrial
83 fresh-cut leafy-vegetables washing process was assessed. To the knowledge of the authors, this is
84 the first published study that utilizes H₂O₂ as wash water sanitizer in a full-scale washing process
85 of fresh-cut leafy vegetables. Also, lab-scale experiments were performed to assess the use of
86 Bacsan (containing a. o. Cu²⁺, Ag⁺, and Zn²⁺) to improve the H₂O₂ disinfection efficiency in post-
87 harvest water disinfection processes.

88 2. MATERIALS AND METHODS

89 2.1. Water disinfection in a fresh-cut leafy vegetables processing company

90 2.1.1. Experimental setup

91 Experiments were executed in a Belgian fresh-cut leafy vegetables processing company. First, a
92 run was executed without addition of water disinfectant, i.e. the 'blank' run. A batch of 400 kg
93 mixed salad was processed, containing radicchio (33%), sugar loaf (*Chicorium intybus*) (33%) and
94 curled endive (33%). The leafy vegetables were cut (in pieces of 1 by 5 cm), and transported
95 through two subsequent immersion washing baths (washing bath 1: WB1 and washing bath 2:
96 WB2) with a volume of 450 L each, and a leafy vegetable residence time of 1 min in each washing
97 bath. The washing system consisted of bubble washers, i.e. production of agitation in the washing
98 baths by air bubble injection through underwater air nozzles. Subsequently they were transported
99 by a conveyer belt to a centrifuge for dewatering, followed by a weighing unit (computer controlled
100 weight proportioning scales). Both washing baths were filled with bore hole water, cooled on
101 beforehand to 2 °C. During the washing process, 300 L/h of bore hole water was added to each of
102 the washing baths. Wash water was recirculated within washing baths but not between washing
103 baths. The only water that was transferred from WB1 to WB2 was the water that was attached to
104 the transferred lettuce. Two wash water disinfection experiments were performed. In both
105 experiments, the same types of leafy vegetables were processed during the wash water disinfection
106 experiments of which the first batch (467 ± 55 kg) was the same leafy vegetables mix as in the
107 blank runs. In addition, a second batch (258 ± 31 kg) was processed, consisting of white lollo
108 (*Lactuca sativa* cv. Lollo Bianco) (50%) and lollo rosso (*Lactuca sativa* cv. Lollo Rosso) (50%).
109 For each type of leafy vegetable and experiment, the crops originated from the same farm, and the
110 crops were processed at the day of harvest. On average leafy vegetables were washed at 333 ± 50
111 kg/h. In the disinfection experiments, WB1 was operated identically to the blank runs. In the first

112 disinfection experiment, WB2 was filled with 1.8 % H₂O₂ (i.e. 4% EcoClearProx, ABT Belgium,
113 Belgium) and 300 L/h 1.8% H₂O₂ of bore hole water was added. In the second disinfection
114 experiment, WB2 was filled with 1.8% H₂O₂ and 300 L/h of bore hole water was added. During
115 processing, 300 L/h of wash water was tapped from the washing bath and 5.4 L/h H₂O₂ was dosed
116 (again to obtain addition of 1.8 % H₂O₂/L) and sent through a low pressure UV-C system (Aquadra
117 2, Wedeco, Belgium; 55 W) with fluence of 240 mJ/cm² at a flow of 300 L/h and 98% UV 254 nm
118 transmittance/cm, before recirculation to WB2.

119 *2.1.2. Sampling in the fresh-cut leafy vegetable processing company*

120 Samples of the fresh-cut leafy vegetables, water samples from WB1 and WB2, and samples from
121 the food contact surfaces of the conveyer belt and the weighing unit were taken five times
122 throughout the processing: at the start of batch 1, at the middle of batch 1, at the end of batch 1 =
123 start of batch 2, in the middle of batch 2, at the end of batch 2. About 250 g of fresh-cut leafy
124 vegetables was sampled and put directly into a sterile stomacher bag. For sampling the raw
125 material, each lettuce type was sampled separately per batch, and averaged as the microbial count
126 of the raw material. The water samples were collected into a sterile 1 L bottle according to ISO
127 19458:2006 (ISO, 2006). Excess H₂O₂ was quenched with sterile Na₂S₂O₃. The food contact
128 surfaces were sampled with sterile swabs. Aseptic templates covering 50 cm² were used and a
129 sterile swab moistened in 5 mL of buffered peptone water was used to swab a delimited area
130 vertically, horizontally, and diagonally. All the samples were stored and transported in the dark at
131 < 4°C to the lab for further handling and subsequent microbial analysis within 12 h. For each
132 measuring point two independent samples were taken. At each time point and operation unit, water
133 and food contact surfaces were sampled at two consistent points, and each of the two samples for
134 raw materials screening originated from two crops.

135 *2.1.3. Microbial analyses*

136 For the fresh-cut leafy vegetables samples and food contact surfaces, APC and *E. coli* were
137 enumerated, whereas in the water also enterococci were enumerated. For the fresh-cut leafy
138 vegetables samples, 10 g of fresh-cut leafy vegetables was weighed in a stomacher bag and
139 homogenized for 1 minute in 90 ml buffered peptone water. The enumeration of APC was done
140 with the reference method ISO 4833:2003 (ISO, 2003), with the exception that the plates were
141 incubated at 22°C for five days instead of at 30°C for 3 days. *E. coli* was enumerated with the pour
142 plate method on RAPID'*E.coli* 2 agar (BioRad, France), a selective chromogenic medium,
143 incubated for 24 h at 37°C. For the water and food contact surface samples, APC was measured
144 according to ISO 6222:1999 and incubated for 3 days at 22°C (ISO, 1999). The enumeration of *E.*
145 *coli* was done according to ISO 9308-1 (i.e. membrane filtration) with the exception that the tergitol
146 7 medium was replaced by RAPID'*E.coli* 2 agar (Biorad, France) (ISO, 2000a). The detection and
147 enumeration of enterococci was performed using the membrane filtration method ISO 7899-2 (ISO,
148 2000b).

149 *2.1.4. Physicochemical parameters*

150 Alkalinity was determined with acid titration, turbidity with a turbidimeter (HI98703, HANNA
151 Instruments, Belgium), COD according to the small-scale sealed-tube method (LCI 400, Hach
152 Lange, Belgium). H₂O₂ concentration, pH and T were determined at the fresh-cut leafy vegetables
153 processing company. H₂O₂ was determined with the spectrophotometric I₃⁻ method by Klassen et
154 al. (1994). H₂O₂ was determined immediately after sampling, to avoid further consumption due to
155 reaction with the water matrix components.

156 ***2.2. Water disinfection in standardized wash water***

157 *2.2.1. Standardized wash water*

158 The outer leaves of the iceberg lettuce (*Lactuca sativa L.*) were removed. The leaves were cut into
159 pieces of about 3 cm and 67 g of the cut salad was put in a stomacher bag to which 200 ml of tap
160 water was added. The mixture was homogenized for 2 min. The COD of this suspension was
161 determined, and subsequently, this mixture was diluted with tap water, to obtain standardized wash
162 water (SWW) with the desired COD.

163 2.2.2. *Industrial wash water*

164 Industrial wash water was collected at a Norwegian fresh-cut leafy vegetables processing
165 company. The water was collected immediately after a batch of mixed lettuce, i.e. iceberg lettuce,
166 rucola (*Eruca sativa*) and radicchio, had been washed with tap water.

167 2.2.3. *Bacterial inoculation*

168 *E. coli* ATCC 25922 was grown in nutrient broth (Oxoid, France) for 24 hours at 37° C. The *E.*
169 *coli* cells were washed in phosphate-buffered saline and subsequently added to the SWW to obtain
170 5-6 log CFU / mL.

171 2.2.4. *Physicochemical parameters*

172 COD and H₂O₂ were measured as described above. The Cu²⁺ ion concentration in SWW was
173 measured with a test kit (MD 200 2IN1 copper, Lovibond, Germany), based on the reduction of
174 Cu²⁺ to Cu⁺, the reaction of Cu⁺ with bicinchoninic acid, followed by spectrophotometric
175 measurement of the formed complex. Free chlorine was measured as described by Van Haute et al.
176 (2013a).

177 2.2.5. *Disinfection experiments*

178 H₂O₂ was diluted from a 30% stock-solution (Fluka Analytical, Germany). Bacsan (Labola,
179 Norway) is a patented, commercial formulation from Aqua Chemical Nutrients, marketed as water
180 disinfectant and containing a. o. Cu, Ag, and Zn. The content of the Bacsan solution was analyzed
181 with inductive coupled plasma emission spectrometry to determine the actual metal ion

182 concentrations and were found to be: 84.2 ± 1.1 g/L Cu, 7.3 ± 0.2 mg/L Ag, 23.7 ± 0.4 g/L Zn,
183 24.0 ± 0.4 mg/L Al, and 56.0 ± 0.2 g/L NO_3^- . 100 mL of continuously mixed, inoculated SWW at
184 4 ± 2 °C was exposed to 500 mg/L of H_2O_2 , with or without the addition of 2 or 10 mg/L Bacsan-
185 Cu (expressed as mg/L Cu^{2+} in Bacsan), or to 500 mg/L of H_2O_2 with the addition of 10 mg/L Cu^{2+}
186 from $\text{CuSO}_4 \cdot 5\text{H}_2\text{O}$ (Merck, Germany), or to 10 mg/L Bacsan-Cu without H_2O_2 . H_2O_2 residual
187 concentration was measured after 5, 30, and 120 min. Microbial samples were taken after 30 and
188 120 min and immediately quenched with $\text{Na}_2\text{S}_2\text{O}_3$. For each treatment and chosen COD level of
189 SWW, 3 independent experiments were executed. The industrial fresh-cut leafy vegetables wash
190 water was similarly treated as the SWW, except for the *E. coli* inoculation which was not executed.
191 Also, exposure of SWW to 500 mg/L of H_2O_2 with addition of 2 mg/L Bacsan-Cu and 0.1 mg/L
192 Ag^+ from AgNO_3 (Sigma-Aldrich, Germany) or to 500 mg/L of H_2O_2 with addition of 10 mg/L
193 Bacsan-Cu and 1 mg/L Ag^+ was assessed for H_2O_2 stability in SWW.

194 2.2.6. Microbial analyses

195 APC was enumerated with the pour plate method on Water plate count agar (Oxoid, England)
196 (incubated for 3 days at 22° C) and *E. coli* with the pour plate method, using RAPID'*E. coli* 2 agar
197 (Biorad, France) (incubated for 24 h at 37°C).

198 2.2.7. Assessment of the interaction of catalase and Bacsan

199 For investigating the effect of Bacsan and pH on the H_2O_2 consumption caused by the enzyme
200 catalase, SWW was rapidly heated to 80°C and maintained at 80°C for 10 min to inactivate catalase
201 (Hirvi, Griffiths, McKellar, Modler, 1996; Anderson, 2002). Thereafter, the SWW was rapidly
202 cooled to 4°C. The heated and unheated SWW were treated with 630 mg/L H_2O_2 (with or without
203 10 mg/L Bacsan-Cu) and also with 100 mg/L free chlorine for comparison with a disinfectant that
204 is no specific target of an enzyme. Free chlorine was diluted from a chlorine stock solution (28.4
205 g/L NaOCl, La Croix, Belgium).

206 Also, experiments using pure catalase from bovine liver (Sigma-Aldrich, Norway) were performed.
207 H₂O₂ was diluted in 0.05 mol/L phosphate buffer with pH 5.5, 6.0 or 7.2 to a final concentration
208 of 590 mg/L H₂O₂. Preliminary tests were done at pH 5.5 and pH 7.2 by adding 0, 2, 10 mg/L
209 Bacsan-Cu, or 10 mg/L Cu²⁺ (as CuSO₄.5H₂O) to the buffered H₂O₂ solutions, to which catalase
210 was added to a final concentration of 1.5 mg/L. The consumption of H₂O₂ caused by the added
211 catalase was assessed by measuring the H₂O₂ residual concentration after 5 and 30 min, at 4±2 °C,
212 and under continuous mixing. More detailed experimentation was done with 0 and 10 mg/L
213 Bacsan-Cu in the presence of 590 mg/L H₂O₂ in buffered solutions at pH 5.5, 6.0, and 7.2. The
214 H₂O₂ residual concentration was measured after 5 min and 30 min. The experiments were
215 performed at 4±2 °C and repeated 3 times.

216 2.2.8. *Oxidative browning*

217 Lettuce washing experiments were executed to determine the effect of the treatments (i.e. 500 mg/L
218 H₂O₂ with or without Bacsan) on oxidative browning of the lettuce and consisted of washing 30 g
219 of cut lettuce with mechanical agitation for 2 min at 4 ± 2 °C, followed by centrifugation and
220 storage in sterile plastic boxes at 4 ± 2° C. After 3, 4, and 5 days the lettuce samples were observed
221 for visible traces of enzymatic browning.

222 2.3. *Statistics*

223 Statistical analysis was performed with SPSS statistics 21 and Microsoft Excel. Comparison of
224 parameter levels was done with one-way ANOVA or Brown-Forsythe when equal variance could
225 not be assumed. Group comparison was done with post-hoc tests (Tukey or Games-Howell). For
226 comparing means of parameters, the Mann Whitney-U and Wilcoxon-signed rank tests were used
227 for unpaired and paired samples respectively. A level of significance of $p \leq 0.05$ was chosen for
228 all statistical analyses.

229 3. RESULTS

230 *3.1.H₂O₂ wash water disinfection in a fresh-cut leafy vegetables processing company*

231 Initially, the washing process turbidity and COD increased rapidly and subsequently, in general,
232 the increase diminished as a function of time (Figure 1). The turbidity in WB1 was significantly
233 higher than in WB2 during the trials. The COD in WB2 was not measured due to the interference
234 of H₂O₂. The H₂O₂ concentration rapidly decreased during the washing process (Figure 1), and a
235 significant negative correlation between turbidity and H₂O₂ concentration ($r^2 = 0.608$; $p < 0.0005$)
236 was observed. The temperature increased to higher values in WB2 compared to WB1. The pH of
237 the bore hole water was 7.4 ± 0.1 . The pH value rose to slightly higher values in WB2 compared
238 to WB1 towards the end of the water disinfection experiments, though not significantly (Figure 1).
239 The alkalinity did not change significantly as a function of time and was not significantly different
240 between treatments nor washing baths, with 6.36 ± 0.10 , 6.38 ± 0.14 and 6.51 ± 0.16 mmol/L
241 bicarbonate in the bore hole water, WB1 and WB2 respectively, indicating it originated
242 predominately from the bore hole water itself.

243 The *E. coli* contamination was below the limit of detection at all times and locations, i.e. < 1 log
244 CFU/g on the fresh-cut leafy vegetables, < 0.3 log CFU/100 mL in the water and < 0.7 log CFU/
245 50 cm² on the conveyer belt and the weighing unit. APC and enterococci contamination was
246 significantly higher in WB1 compared to WB2 in all experiments (Figure 2). To assess the impact
247 of the water disinfection treatments, the differences between the measured contamination in both
248 washing baths was calculated (WB1 – WB2) in order to be able to compare the wash water
249 disinfection efficiency of the treatments and to incorporate fluctuations in transfer of
250 microorganisms from fresh-cut leafy vegetables to the washing baths between the different water
251 disinfection trials. These differences were significantly higher for both disinfection treatments
252 compared to the blank during the first batch (first 60 to 80 min dependent on the batch size), i.e.
253 the wash water contamination was significantly reduced with the 1.8 % H₂O₂ (with or without UV)

254 treatment. For enterococci, no significant reductions in the wash water were found. The gradually
255 increasing APC concentration in the H₂O₂ treated wash water (Figure 2b) reflects the declining
256 H₂O₂ residual during the washing process (Figure 1), despite the continuous addition of 300 L/h of
257 1.8 % H₂O₂ in a 450 L washing bath, due to the build-up of organic matter in the washing bath,
258 indicated in WB2 as increasing turbidity (Figure 1). The APC and enterococci contamination after
259 the UV/H₂O₂ unit was reduced to below the detection limit (< 2 log CFU/100 mL and < 0.3 log
260 CFU/100 mL respectively) at all times. However, a decrease of the wash water contamination in
261 WB2 compared to the 1.8% H₂O₂ treatment was not observed.

262 For batch 1, the initial APC load of the fresh-cut leafy vegetables was 7.1 ± 0.4 , 6.8 ± 0.3 , and 6.8
263 ± 0.2 log CFU/g for the blank run, 1.8 % H₂O₂, and 1.8 % H₂O₂ + UV respectively, whereas for
264 batch 2, it was 7.4 ± 0.2 and 7.6 ± 0.2 log CFU/g for 1.8 % H₂O₂ and 1.8 % H₂O₂ + UV respectively.
265 The APC load on the fresh-cut leafy vegetables was reduced significantly with 1.8 % H₂O₂ (Figure
266 3), with or without UV, but also with a water wash. The introduction of organic matter lowered the
267 H₂O₂ concentration as the washing process advanced in time. However, processing time had no
268 influence on decontamination efficiency of any of the treatments, and 1.8 % H₂O₂ (with or without
269 UV) did not improve the decontamination efficiency (considering batch 1) compared to a water
270 wash (Figure 3). The APC contamination on the conveyer belt increased during processing from
271 3.1 to 4.8, 2.7 to 4.2 and 2.3 to 4.7 log CFU/ 50 cm² in the blank run, with 1.8 % H₂O₂, and with
272 1.8 % H₂O₂ + UV respectively, and on the weighing unit from 3.2 to 5, 2.3 to 4.7, and 3.5 to 5.5
273 log CFU/ 50 cm² respectively.

274 ***3.2.H₂O₂ stability in SWW***

275 For all treatments, both COD and contact time had a significant detrimental influence on the H₂O₂
276 concentration in the SWW (Table 1). The rate of H₂O₂ consumption was lowest with H₂O₂ + 10

277 mg/L Bacsan-Cu. The consumption rate was lower with H₂O₂ + 2 mg/L Bacsan-Cu than with H₂O₂
278 at COD 497 and 848 mg O₂/L, whereas at COD 1830 mg O₂/L no difference was observed (Table
279 1). The stability of H₂O₂ in SWW of COD 789 mg O₂/L was higher with H₂O₂ + 10 mg/L Bacsan-
280 Cu compared to H₂O₂ + 10 mg/L Cu²⁺ (from CuSO₄), which in turn was significantly higher after
281 30 min than in the absence of metal ions (Figure 4). The addition of 0.1 mg/L Ag⁺ to 2 mg/L
282 Bacsan-Cu and 1 mg/L Ag⁺ to 10 mg/L Bacsan-Cu in SWW of COD 753 mg O₂/L further enhanced
283 the stability of initially added 500 mg/L H₂O₂ in a synergistic fashion (Figure 5).
284 The Cu²⁺ concentration in solution decreased only moderately after 120 min treatment time, i.e.
285 9.5 ± 0.7 %, 3.6 ± 1.3 %, and 5.6 ± 3.1 % for H₂O₂ + 10 mg/L Bacsan-Cu, H₂O₂ + 2 mg/L Bacsan-
286 Cu and 10 mg/L Bacsan-Cu respectively. The pH of the SWW was 7.3 ± 0.2. Addition of 500 mg/L
287 H₂O₂ did not change the pH significantly, whereas addition of 10 mg/L Bacsan- Cu, H₂O₂ + 2 mg/L
288 Bacsan-Cu, and H₂O₂ + 10 mg/L Bacsan-Cu decreased the pH with 0.3 to 0.7, 0.3 to 0.9, and 0.5
289 to 1.0 respectively, the pH drop increasing with decreasing COD of the SWW, most likely due to
290 an increasing amount of pH buffering molecular species in SWW of higher COD. This pH drop
291 was in part due to the low pH of the Bacsan stock-solution i.e. below the detection limit of the pH
292 meter (pH 0).
293 When heating SWW of COD 819 mg O₂/L, the COD did not change significantly. Addition of 630
294 mg/L H₂O₂ to heated SWW led to an initial rapid decrease in the first 5 min, after which no
295 considerable further consumption occurred during the remaining 25 min (Figure 6a). Addition of
296 Bacsan had no influence on the H₂O₂ consumption. In the unheated water however, Bacsan
297 decreased the H₂O₂ consumption (as observed before). The H₂O₂ residual in heated SWW was
298 considerably larger than in the unheated SWW, whereas for free chlorine the difference in
299 consumption was much smaller (Figure 6b).

300 To explain the increased stability of H₂O₂ in the presence of Cu²⁺ and to larger extent in the
301 presence of Bacsan, the impact on the activity of bovine liver catalase activity was assessed. In the
302 absence of catalase the H₂O₂ concentration remained constant in all the phosphate buffered
303 solutions during the experimental period. The pH was measured and found to not be affected by
304 the addition of H₂O₂ and Bacsan. Preliminary experiments (without repeats) showed the following
305 order of H₂O₂ stability: H₂O₂ < H₂O₂ + 2mg/L Bacsan-Cu ~ H₂O₂ + 10 mg Cu²⁺ (from CuSO₄) <
306 H₂O₂ + 10 mg/L Bacsan-Cu (data not shown). More detailed experiments showed that at each pH
307 value (5.5, 6, or 7.2), the consumption of H₂O₂ was significantly lower in the presence of 10 mg/L
308 Bacsan-Cu than in absence of Bacsan (Figure 7). The pH affected the consumption of H₂O₂ by
309 catalase. The highest residual H₂O₂ concentration after 5 and 30 minutes was measured at pH 5.5
310 in the presence of 10 mg/L Bacsan-Cu (Figure 7).

311 ***3.3. Water disinfection in SWW***

312 At the start of the experiments in SWW, the APC was 6.0 ± 0.2 log CFU/ mL averaged among all
313 experiments, and the inoculated *E. coli* contamination was 5.4 ± 0.4 log CFU/ mL. *E. coli* was
314 more susceptible than APC to H₂O₂ combined with 2 mg/L or 10 mg/L Bacsan-Cu (Table 1). The
315 inactivation of APC was significantly higher with H₂O₂ + 10 mg/L Bacsan-Cu compared to the
316 other treatments, whereas for *E. coli*, the reduction was higher with H₂O₂ combined with 2 or 10
317 mg/L Bacsan-Cu compared to the other treatments. Exposure to increasing concentrations of COD
318 had a detrimental influence on the inactivation of *E. coli* and APC. A significantly higher
319 inactivation of APC and *E. coli* after 120 min compared to 30 min contact time was only observed
320 at COD 1830 mg O₂/L with H₂O₂ + 10 mg/L Bacsan-Cu and at COD 497 mg O₂/L with H₂O₂, and
321 of *E. coli* at COD 848 mg O₂/L with 10 mg/L Bacsan-Cu. The low improvement on disinfection
322 efficiency in the interval 30 to 120 min contact time is attributed to the low remaining H₂O₂ residual
323 (Table 1).

324 Synergy of H₂O₂ + 10 mg/L Bacsan-Cu was observed in SWW of COD 848 and 1830 mg O₂/L
325 (except for APC after 30 min in COD 1830 mg O₂/L) (Table 1). In SWW of COD 497 mg O₂/L,
326 observation of possible synergy was hindered by detection limit issues combined with overall
327 higher inactivation due to the lower physicochemical load of the SWW. Only the presence of
328 synergy in the case of APC after 30 min could easily be observed. The reduction of APC in SWW
329 of COD 789 mg O₂/L was higher with H₂O₂ + 10 mg/L Bacsan-Cu compared to H₂O₂ + 10 mg/L
330 Cu²⁺ (as CuSO₄) (Figure 4). Despite the H₂O₂ concentration being significantly higher after 30 min
331 when adding Cu²⁺ than in the absence of Cu²⁺ (as CuSO₄), the APC inactivation with H₂O₂ + 10
332 mg/L Cu²⁺ was not significantly different from that obtained with only H₂O₂ (Figure 4).

333 ***3.4.H₂O₂ consumption and water disinfection in industrial wash water from a processing*** 334 ***company***

335 3 CFU/ 100 mL *E. coli* were found in the industrial wash water and none were detected after the
336 disinfection treatments. The inactivation of APC was lower in industrial wash water of COD 509
337 mg O₂/L compared to in SWW of COD 497 mg O₂/L (Table 1) . Nonetheless, the H₂O₂ residual
338 was significantly higher in the industrial wash water than in SWW after 30 and 120 min contact
339 time with H₂O₂ + 2 and 10 mg/L Bacsan-Cu, but not with solely H₂O₂ compared to SWW. As in
340 SWW, both the highest H₂O₂ stability and a synergistic APC inactivation were observed with 500
341 mg/L H₂O₂ + 10 mg/L Bacsan-Cu.

342 ***3.5.Impact of washing treatments on browning of stored fresh-cut iceberg lettuce***

343 After washing the lettuce in the water disinfection solutions, some browning appeared after 3 days
344 of storage in the fresh-cut lettuce, and considerable more browning was observed when washing in
345 10 mg/L Bacsan-Cu and H₂O₂ + 10 mg/L Bacsan-Cu compared with washing in water, 2 mg/L
346 Bacsan-Cu, H₂O₂, and H₂O₂ + 2 mg/L Bacsan-Cu. When the lettuce was rinsed after treatment with

347 10 mg/L Bacsan-Cu (with or without H₂O₂), the amount of browning was similar to that of tap
348 water treatment (and the other treatments) for the 5 days storage duration.

349 **4. DISCUSSION**

350 The results in this study show that applying 1.8% H₂O₂ in the washing bath and dosing 300 L/h of
351 1.8% H₂O₂ in a 450 L washing bath is insufficient for maintaining the microbial wash water quality
352 in the washing process when washing 333 ± 50 kg/h fresh-cut leafy vegetables. This, due to the
353 build-up of organic matter in the washing bath which resulted in a rapid consumption of H₂O₂ due
354 to oxidation of this organic matter. The exothermic nature of these oxidation reactions (Klais, 1993)
355 explains the higher temperatures in the H₂O₂ treated washing bath compared to the untreated one.

356 Industrial washing of the leafy vegetables in ≤ 1.8 % H₂O₂ for 1 min did not improve the
357 decontamination efficiency compared to a water wash in this study. Other decontamination studies
358 of fresh-cut leafy vegetables have been conducted with H₂O₂ in the range of 1 – 3 %, with
359 concentrations ≥ 2% showing improved decontamination efficiency compared to washing in water.
360 However, these studies were performed at room or elevated temperature and with artificially
361 inoculated microorganisms (Allwood, Malik, Hedberg, & Goyal, 2004; Huang, & Cheng, 2011;
362 Lin, Moon, Doyle, & McWatters, 2002). Possible explanations for higher removals in those
363 experiments than in the present study are longer contact time, higher temperature, the use of
364 artificial inocula, the use of specific bacterial pathogens instead of general plate counts, and the
365 full-scale experiments in the present study *versus* lab-scale in the other studies. On the other hand,
366 Ramos et al. (2013) noted that at concentrations of 1-2%, H₂O₂ is not effective for produce
367 decontamination. The issue of artificial inocula was illustrated by Hadjok et al. (2008), who used
368 vacuum infiltration in order to achieve infiltration of inoculated *Salmonella* Montevideo in fresh-
369 cut iceberg lettuce, and observed a much lower inactivation with H₂O₂/UV of internalized

370 *Salmonella* than those bound to the surface. Improved decontamination efficiency of fresh-cut leafy
371 vegetables with H₂O₂ in current industrial processes seems unrealistic, due to the requirement of
372 long contact times, relatively high temperatures, high H₂O₂ wash water residual and the high
373 reactivity of H₂O₂ with wash water organics. The observed increase in APC contamination on food
374 contact surfaces illustrates that besides cross-contamination via the wash water, cross-
375 contamination via food contact surfaces is an issue in fresh-cut leafy vegetables processing
376 operations.

377 The executed case-study with UV/H₂O₂ in the fresh-cut leafy vegetables company shows the
378 inadequacy of using an off-line disinfection technique to attempt to control the microbial
379 contamination in a process with rapid and continuous influx of microbial contamination such as a
380 fresh-cut leafy vegetables washing process. The issue is that an off-line disinfection technique only
381 treats part of the water at any given time, whereas an *in situ* disinfection technique maintains a
382 residual, as such treating all the water at any given time. From the point of food safety, pathogenic
383 contamination is mostly not widely distributed among a batch of fresh-cut leafy vegetables,
384 although there are extreme cases, for example 12.1 % (4/33 positive samples) prevalence of
385 *Salmonella* spp. on cabbage sampled in a field in India irrigated with partially treated municipal
386 wastewater (Rai, & Tripathi, 2007), or 76.9 % (10/13) and 61.5 % (8/13) prevalence of respectively
387 *Cryptosporidium* spp. and *Giardia* spp. on lettuce sampled in a field in Spain irrigated with
388 contaminated water from an irrigation canal. Nonetheless, the prevalence of bacterial pathogens
389 (*Salmonella* spp., pathogenic *E. coli*, *Listeria monocytogenes*, *Campylobacter* spp.) is, as far as
390 microbial screening studies show, much lower (<1% of the crops) (Johannessen, Loncarevic, &
391 Kruse, 2002; University of Georgia, 2011). In those cases only a very minor part of the crops are
392 contaminated. However, the statement that *in situ* disinfection is needed to avoid cross-

393 contamination during point-contaminations holds for the same reasons as with a continuous
394 microbial build-up, i.e. i) the off-line disinfection occurs at another location than the actual
395 introduction of contamination in the wash water, and ii) the disinfection kinetics are too slow as
396 again only part of the water will be treated at a certain time and the microbial contamination will
397 rapidly disperse throughout the wash water.

398 The experiments in SWW showed that the stability of H₂O₂ was improved by the addition of
399 Bacsan and to lesser extent Cu²⁺. Elimination of heat labile molecules (80 °C, 10 min) greatly
400 increased the stability of H₂O₂ in SWW, which was much less the case for free chlorine. The most
401 obvious heat labile compound with a high and selective impact on H₂O₂ stability is catalase,
402 originating from the lettuce tissue and present in the SWW. This study showed the inhibiting
403 influence of Bacsan and to lesser extent Cu²⁺ on bovine liver catalase activity. The decrease in
404 catalase activity in the presence of certain metals may be related to direct binding of metal ions
405 (including Cu²⁺, Zn²⁺, Ag⁺) to –SH groups of the catalase enzyme, as such inhibiting the enzyme
406 (Atli, Alptekin, Tukel, & Canli, 2006; Atli, & Canli, 2007). Furthermore, peroxidase (from
407 vegetables) activity is reduced after heating (80°C, 10 min) (Morales-Blancas, Chandia, &
408 Cisneros-Zevallos, 2002), and inhibition of peroxidases by metal ions (Zn²⁺, Ag⁺) has been
409 reported (Splittgerber & Tappel, 1979). As such it could be that peroxidases were inhibited by
410 Bacsan. The less rapid H₂O₂ consumption by catalase at lower pH is attributed to catalase having
411 to operate at sub-optimal pH. Bovine liver catalase has optimal activity at about pH 7 and isoelectric
412 point (pI) of pH 5.4 (Maehly, & Chance, 1954; Samejima, Kamata, & Shibata, 1962; Shi et al.,
413 2008), while lettuce catalase has optimal activity in the pH range 7-8, and consists of two
414 izoenzymes (pI 5.8 and pI 6.2) (Bestwick, Adam, Puri, & Mansfield, 2001). Also, the H₂O₂
415 consumption rate is reduced at lower pH, especially below pH 3 (Ortiz, Angelica Rubio, & Lissi,

416 2000; Watts, Foget, Kong, & Teel, 1999) and the pH drops caused by the addition of Bacsan, and
417 amplified when combined with H₂O₂, potentially slightly contributed to the inhibition of lettuce
418 catalase in SWW.

419 Cu²⁺ in the presence of H₂O₂ generally does not induce free radical formation (through Fenton-like
420 reactions) in systems that contain biomolecules because of the tendency of Cu²⁺ to tightly bind
421 amino group containing compounds (Gutteridge, & Wilkins, 1983; Pham, Xing, Miller, & Waite,
422 2013). This is an additional explanation why Cu²⁺ did not accelerate the H₂O₂ consumption.
423 Therefore, the inactivation effect of H₂O₂/Cu²⁺ against bacteria is most likely due to the combined
424 attack of the two disinfectants, rather than the production of radical formation (Orta De Velasquez,
425 Yanez-Noguez, Jimenez-Cisneros, & Luna Pabello, 2008, Macomber, Rensing, & Imlay, 2007).

426 In accordance with the increased stability of H₂O₂ when combined with Bacsan, synergistic effects
427 were observed when these disinfectants were combined, both in SWW and in industrial leafy
428 vegetables wash water. The lower inactivation of the APC in the industrial wash water, despite the
429 higher H₂O₂ exposure, can be attributed to an overall more resistant microbiota than in the SWW,
430 which is plausible, as APC is a non-discriminative enumeration method. The lower H₂O₂
431 consumption might be due to the fact that COD is a general parameter that measures the amount
432 of oxygen that is necessary to oxidize the substances in the sample, and is used as an indicator for
433 the organic load of the water. As such, it does not directly inform about the reaction rate of specific
434 molecular species with H₂O₂ (as was observed to be significant in the heated SWW), nor the levels
435 of iron or phosphate, which can also influence the H₂O₂ consumption (Watts, Foget, Kong, & Teel,
436 1999). On the contrary, in a previous study, the COD was found to be a universal parameter that
437 effectively predicted the disinfection efficiency of free chlorine to inactivate inoculated *E. coli*
438 O157 in both SWW (made in the same fashion as in this study but with butterhead lettuce instead

439 of iceberg lettuce) and fresh-cut leafy vegetables wash water from 2 processing companies (Van
440 Haute, Sampers, Holvoet, & Uyttendaele, 2013). The chlorination trials in heated SWW in this
441 study show the relative independence of chlorine consumption on specific heat labile compounds
442 in the SWW.

443 The results in SWW confirm that H₂O₂ is a slow acting water disinfectant that quickly decomposes
444 in the presence of high COD of fresh-cut lettuce origin. The inactivation of *E. coli* in oxidant
445 demand free conditions requires a much higher exposure (concentration and contact time) to H₂O₂
446 (21) than to free chlorine (Rice, Clark, & Johnson, 1999; Van Haute, Sampers, Holvoet, &
447 Uyttendaele, 2013, Zhao et al., 2001) and ozone (Hunt, & Marinas, 1997). H₂O₂ is a non-radical
448 reactive oxygen species that is capable of penetrating most biological membranes yet directly
449 inactivate only few enzymes (Atli, Alptekin, Tukel, & Canli, 2006). It is the production of hydroxyl
450 radicals through the Fenton-reaction with free intra- and extracellular iron that enables damage to
451 membrane structures, DNA, and proteins (a. o. oxidation of Fe-S cluster proteins and more
452 generally of cysteine residues in proteins) (Brudzynski, Abubaker, St-Martin, Castle, 2011; Imlay,
453 2003; Raffellini, Schenk, Guerrero, &, Alzamora, 2011).

454 Bacsan itself also showed antimicrobial activity when dosed at 10 mg/L (as Cu). Ionic copper and
455 silver induce cell lysis and death by attaching to the negatively charged bacterial cell surface,
456 disrupting cell wall permeability, blocking cell respiration, and causing extra- and intracellular
457 protein denaturation (Agnihotri, Mukherji, & Mukherji, 2013; Feng et al., 2000; Huang et al., 2008;
458 Lin et al., 1998). Feng et al. (2000) observed that exposure of *E. coli* and *Staphylococcus aureus*
459 cells to Ag⁺ led to transformation of the DNA into a condensed form, leading to loss of replication
460 ability. Cu⁺, Ag⁺, and Zn²⁺ ions inactivate Fe-S cluster enzymes in *E. coli* and depletion of
461 antioxidant reserves, particularly glutathione, can occur due to metal ions including Ag⁺ and Zn²⁺

462 and exposure of *E. coli* to toxic doses of these metal species can lead to depletion of total cellular
463 thiols (Lemire, Harrison, & Turner, 2013; Macomber & Imlay, 2009). Also, combining sub-
464 inhibitory concentrations of Ag^+ with certain metal ions, including Cu^{2+} and Zn^{2+} , increased the
465 toxicity of those metal ions on *E. coli* by a factor of 10 (Pedahzur, Shuval, & Ulitzur, 1997). As
466 such, the combination of measured ions in Bacsan exhibit antimicrobial action.

467 The synergistic microbial inactivation due to H_2O_2 /Bacsan cannot be attributed to the presence of
468 Cu^{2+} alone, but is dependent upon the combination of metal ions added to the SWW, in the presence
469 of H_2O_2 . Contrary to this study, synergy of H_2O_2 (50 to 250 mg/L) with 1 mg/L Cu^{2+} was observed
470 for inactivation of vegetative bacteria in primary wastewater treatment effluent (Orta De
471 Velasquez, Yanez-Noguez, Jimenez-Cisneros, & Luna Pabello, 2008). $\text{Ag}^+/\text{H}_2\text{O}_2$ was more
472 effective than H_2O_2 alone for inactivation of bacteria and fungi (Tote et al., 2009) and
473 synergistically against *E. coli* (Batterman, Zhang, & Wang, 2000, Pedahzur et al., 2000, Pedahzur,
474 Shuval, & Ulitzur, 1997). To the knowledge of the authors, this is the first study that considers the
475 impact of water matrix catalase on H_2O_2 disinfection efficiency. Earlier research has shown that
476 H_2O_2 is less effective against microorganisms with high catalase activity level (Armon et al., 2000;
477 Lambert, Johnston, & Simons, 1999; Sacchetti, De Luca, & Zanetti, 2009; Watts et al., 2003).
478 Monofunctional catalases are not only produced by plants and animals, but also widespread among
479 bacteria (a. o. *E. coli*) and fungi, next to the catalase-peroxidases which are only distributed among
480 bacteria and fungi, and the non-heme catalases that have only been found in certain bacterial
481 species (Loewen, Klotz, & Hassett, 2000; Nadler, Goldberg, & Hochman, 1986). The synergy of
482 H_2O_2 and Bacsan can be explained by i) the higher stability of H_2O_2 in SWW in the presence of
483 Bacsan, allowing for higher exposure of the microorganisms to H_2O_2 and combined with the stress
484 from Bacsan itself, ii) the multiple damage mechanism, meaning both disinfectants attack different
485 targets in the microorganism, as such creating different stresses that could make it harder for the

486 microorganism to remain viable compared to separate addition of the damage from both
487 disinfectants (Koivunen, & Heinonen-Tanski, 2005), and iii) the metal ions in Bacsan might inhibit
488 the functioning of microbial catalases, as such rendering them more susceptible to H₂O₂ attack.
489 Research on inhibition of microbial catalases of target microorganisms to improve the disinfection
490 efficiency, as well as further research on inhibition of water matrix catalase might be interesting
491 towards enhancing the use of H₂O₂ as a disinfectant.

492 It could be that in the case of washing in 10 mg/L Bacsan-Cu, Cu²⁺ was transferred to the lettuce
493 in amounts that induced the activity of polyphenoloxidase, whereas a subsequent rinsing step
494 leached the Cu²⁺ back from the lettuce and as such avoided additional browning (Vandekinderen,
495 2009). 500 mg/L of H₂O₂ does not influence the rate of fresh-cut lettuce browning and is much
496 lower than the concentrations normally applied to decontaminate fruits and vegetables (mostly 1
497 to 5 %), and the concentrations that potentially cause sensorial quality issues towards the lettuce
498 (McWatters et al., 2002a, 2002b; Lopez-Galvez et al., 2013; Ukuku, Bari, & Kawamoto, 2012). In
499 case H₂O₂/Bacsan would be implemented as a reconditioning technique for fresh-cut leafy
500 vegetables wash water, most of the Cu²⁺ (from the Bacsan) would remain in the SWW after
501 reconditioning, and could come into contact with the fresh-cut leafy vegetables when the water is
502 reused. This is advantageous from a cost perspective, because it would enable the reuse of the metal
503 ions for water disinfection. Nonetheless, implementation of a rinsing step with tap water would be
504 required to avoid presence of Cu²⁺ on the packaged lettuce. Furthermore, regulations (for Cu) or
505 guidelines (for Ag and Cu) exist to govern their presence in drinking water (Council of the
506 European Union, 1998; USEPA, 2009, 2013; WHO, 2003, 2004) to avoid ingestion of aesthetically
507 altering (discoloration of the skin and white part of eye in the case of Ag intake) or toxic dosages
508 of these metals. From the perspective of the fresh-cut produce processor and the consumer safety,

509 the primary concern is the possible transfer of these metals to the fresh-cut leafy vegetables and
510 the influence of a rinsing step on the removal of these metals from the fresh-cut leafy vegetables;
511 more so than directly assessing the quality of the wash water itself as this is not consumed.

512 **5. CONCLUSIONS**

513 H₂O₂ is not suited as an *in situ* wash water disinfectant to avoid cross-contamination due to the
514 slow water disinfection kinetics and the rapid consumption in fresh-cut leafy vegetables water. As
515 such, excessive amounts of H₂O₂ would have to be dosed to maintain the microbial wash water
516 quality. The use of off-line disinfection systems (such as the UV/H₂O₂ system applied in this study)
517 are incapable of avoiding cross-contamination in a fresh-cut leafy vegetables washing process, due
518 to the fact that at any given time only part of the wash water is being ‘controlled’, i.e. disinfected.
519 When combined with Bacsan, H₂O₂ showed increased stability in SWW and industrial wash water
520 and synergistic inactivation of *E. coli* and APC in SWW and of APC in industrial wash water. As
521 such, H₂O₂/Bacsan shows potential for use in reconditioning processes, when the inactivation rate
522 is of less importance. The metal ions mixture (Cu²⁺, Zn²⁺, Ag⁺) in Bacsan stabilizes H₂O₂ through
523 a mechanism that most likely involves the inhibition of lettuce catalase present in the fresh-cut
524 leafy vegetables wash water, and potentially other heat labile compounds. When combined with
525 Ag⁺, the H₂O₂ stabilizing action of Bacsan is further enhanced in a synergistic fashion. Further
526 research towards the combination of H₂O₂ and metal ion mixtures such as Bacsan could greatly
527 improve the potential of H₂O₂ as a water disinfectant in general.

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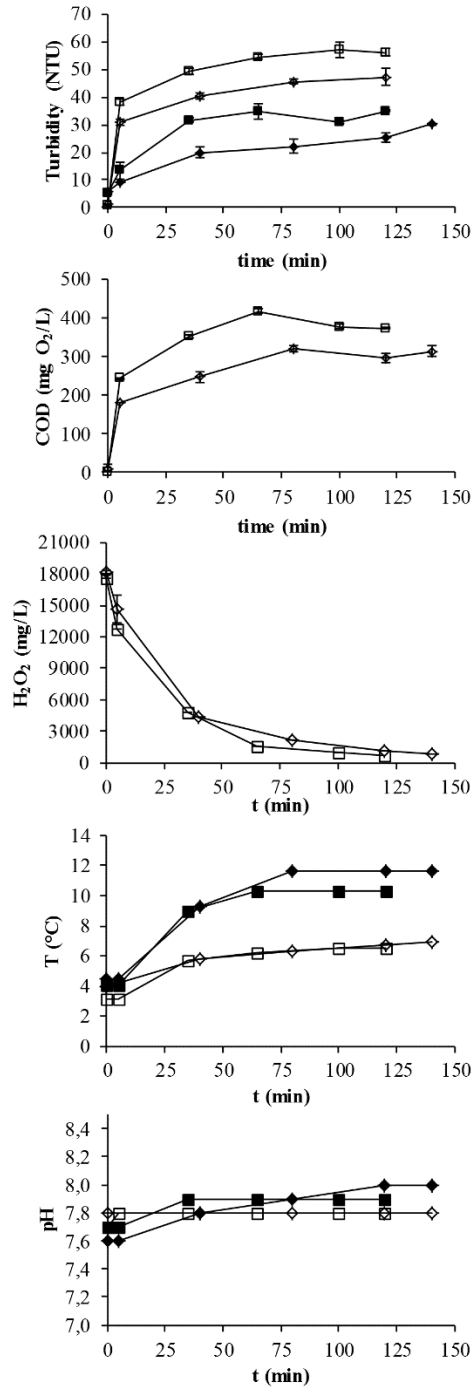
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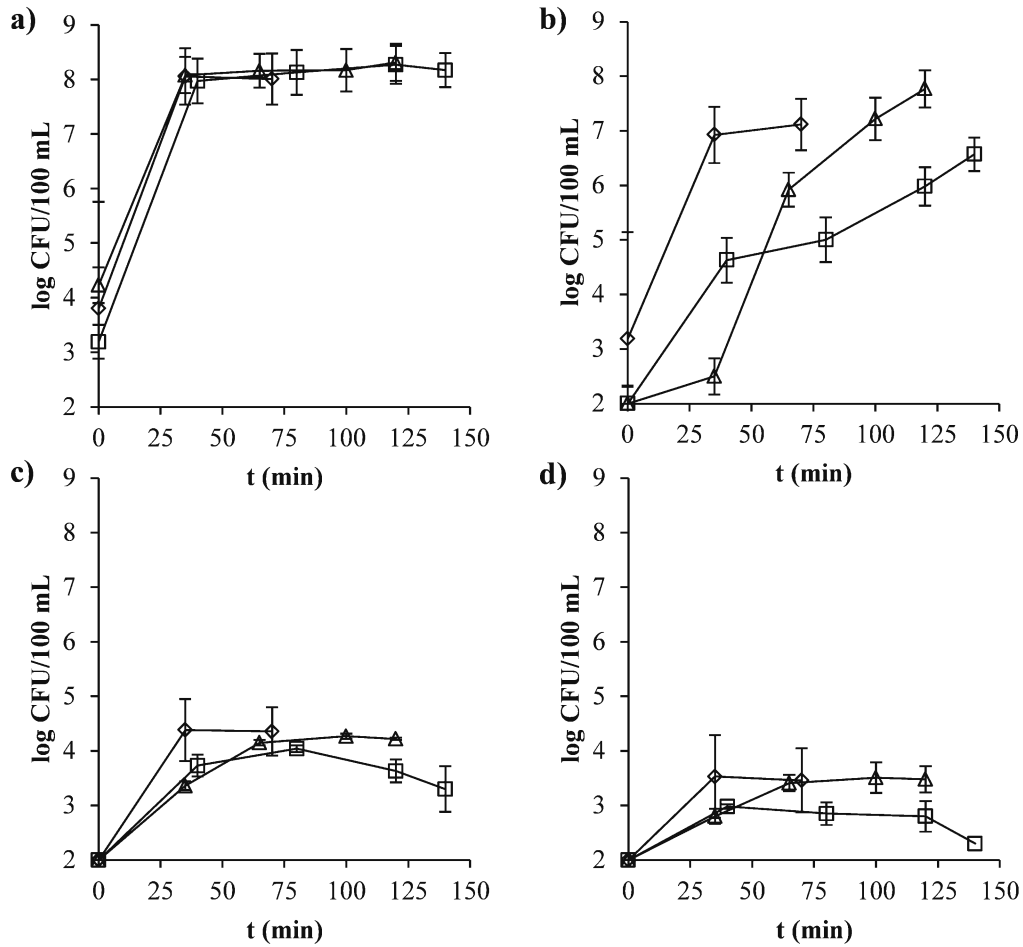
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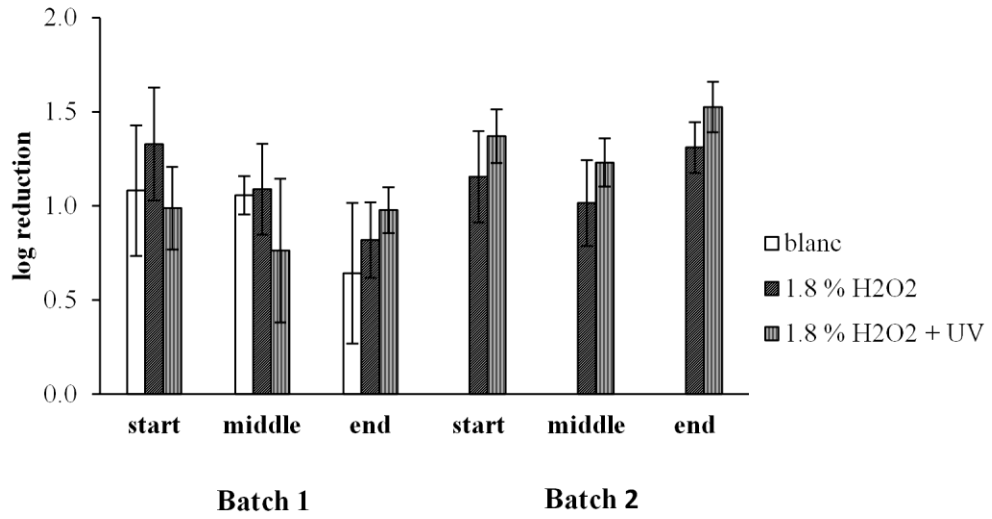
754 **FIGURES**



755
 756 Figure 1. Turbidity, COD, H₂O₂, T, and pH during the washing bath trials with 1.8% H₂O₂,
 757 measured in WB 1(◇) and WB 2 (◆), 1.8 % H₂O₂ + UV, measured in WB 1 (□) and WB 2 (■) (n =
 758 3).



759
 760 Figure 2. Washing bath contamination of APC during the washing trials in the screened company
 761 in a) WB1 and b) WB2 and enterococci in c) WB1 and d) WB2; during the blank run (◇), when
 762 WB2 was treated with 1.8 % H₂O₂ (□), when WB2 was treated with 1.8 % H₂O₂ + UV (△) (n = 2).
 763

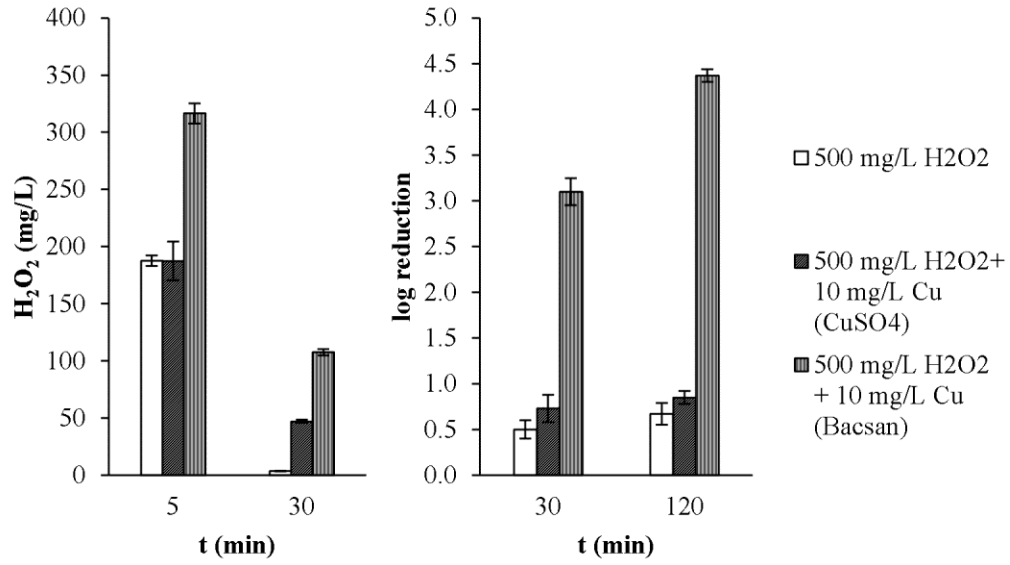


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765 Figure 3. APC reduction on the fresh-cut leafy vegetables due to the industrial washing processes

766 in the screened company (n = 2).

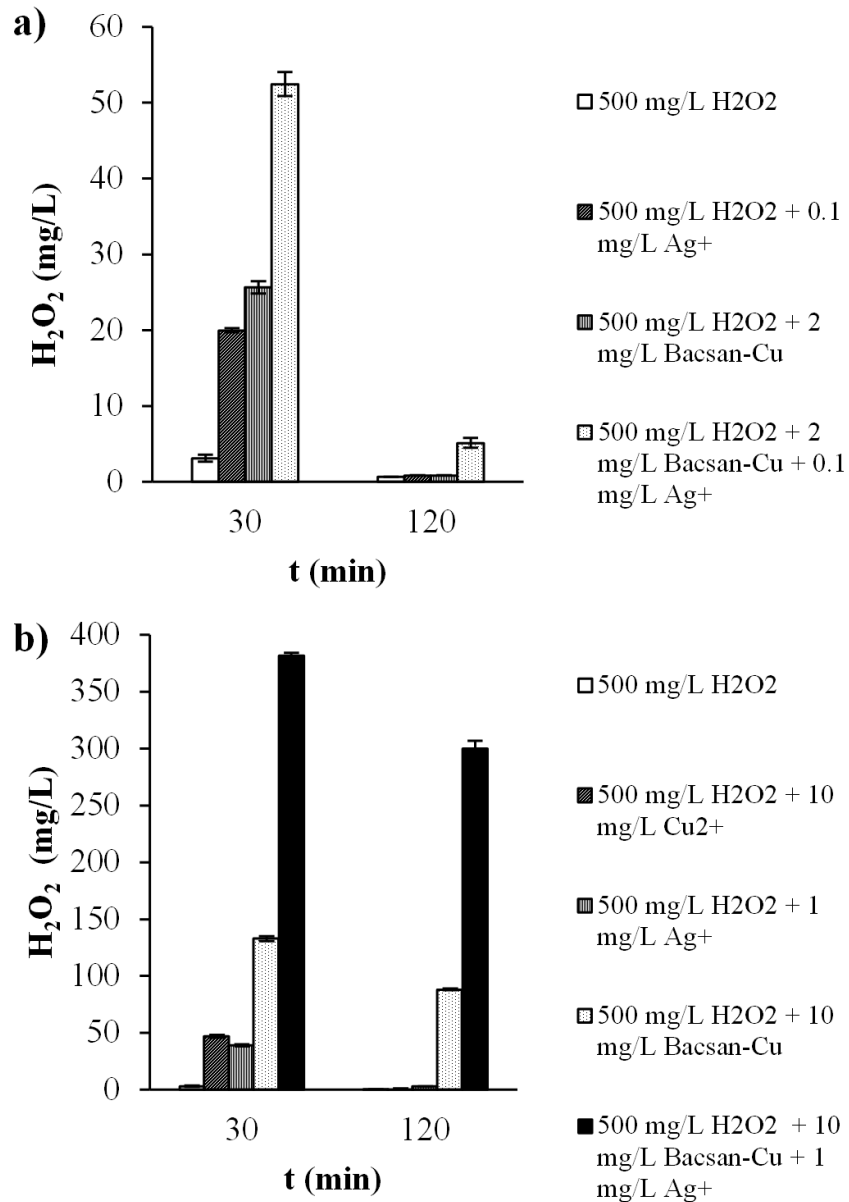
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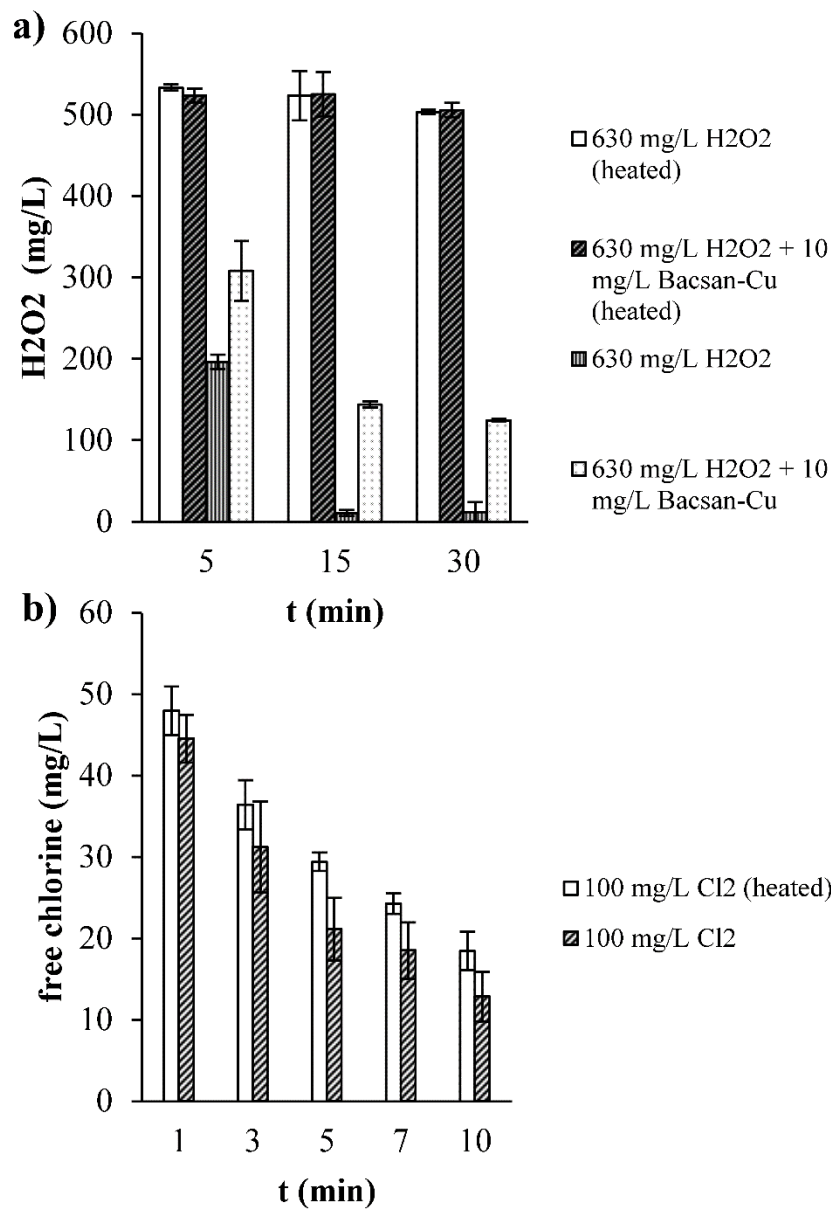
768

769 Figure 4. Residual H_2O_2 and APC reduction in SWW of $COD\ 789 \pm 7\ mg\ O_2/L$ when comparing
 770 treatment with 500 mg/L H_2O_2 , H_2O_2 + 10 mg/L Cu^{2+} (as $CuSO_4$) and H_2O_2 + 10 mg/L Bacsan-Cu
 771 (n = 3).

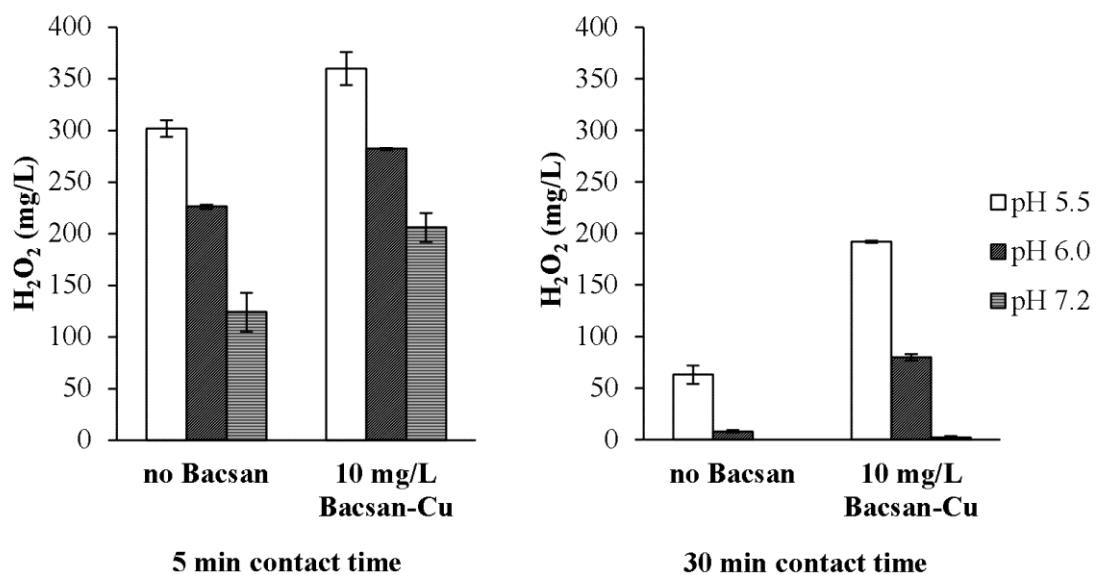
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773
 774 Figure 5. H₂O₂ residual in function of time, when adding Ag⁺ to Bacsan in SWW of COD 753 ± 5
 775 mg O₂/L, a) combinations with 2 mg/L Bacsan-Cu and 0.1 mg/L Ag⁺, b) combinations with 10
 776 mg/L Bacsan-Cu and 1 mg/L Ag⁺, (n = 3).



778
 779 Figure 6. a) Residual H₂O₂ concentration of initially added 630 mg/L H₂O₂ in SWW with COD
 780 819 ± 10 mg O₂/L, with or without heating at 80 °C for 10 min, and b) residual free chlorine
 781 concentration of initially added 100 mg/L free chlorine in SWW with COD 634 ± 2 mg O₂/L, with
 782 or without heating at 80 °C for 10 min (n = 3).
 783



784
 785 Figure 7. Residual H₂O₂ concentration of initially added 590 mg/L H₂O₂, measured after addition
 786 of 1.5 mg/L catalase to phosphate-buffered solutions at different pH and in presence and absence
 787 of 10 mg/L Bacsan-Cu (n = 3).

788

789 **TABLES**

790 Table 1. Microbial reduction and H₂O₂ concentration during water disinfection trials in SWW of varying COD and industrial wash
 791 water with 500 mg/L H₂O₂ and/or Bacsan dosage

COD (mg O ₂ /L)	SWW						Industrial wash water					
	497 ± 7			848 ± 6 mg			1830 ± 21			509 ± 3		
t (min)	5	30	120	5	30	120	5	30	120	5	30	120
APC (log reduction)												
H ₂ O ₂		0.7±0.1	2.2±0.5		0.5±0.2	1.3±0.3		0.0±0.1	0.0±0.2		0.5±0.2	1.0±0.1
H ₂ O ₂ +2 mg/L Bacsan-Cu		2.0±0.4	2.7±0.1		1.2±0.2	1.3±0.3		0.0±0.2	-0.1±0.3		0.4±0.1	0.8±0.1
H ₂ O ₂ +10 mg/L Bacsan-Cu		4.8±0.1	5.0±0.2		4.5±0.2	4.5±0.2		0.1±0.1	0.9±0.1		2.8±0.2	3.3±0.1
10 mg/L Bacsan-Cu		2.6 0.1	>2.7		0.8±0.3	1.1±0.2		0.1±0.2	0.1±0.1		0.3±0.1	0.8±0.1
<i>E. coli</i> (log reduction)												
H ₂ O ₂		0.7±1.1	3.0±0.1		0.6±0.2	0.7±0.2		0.0±0.1	0.1±0.1			
H ₂ O ₂ +2 mg/L Bacsan-Cu		3.8±0.2	4.3±0.5		3.0±0.3	3.0±0.3		0.2±0.1	0.4±0.1			
H ₂ O ₂ +10 mg/L Bacsan-Cu		>5	>5		>5	>5		1.2±0.1	1.8±0.2			
10 mg/L Bacsan-Cu		>2.5	>2.5		1.2±0.2	1.9±0.1		0.2±0.1	0.1±0.2			
H₂O₂ (mg/L)												
H ₂ O ₂	314±2	38±1	1±0.1	221±2	1±0.1	1±0.1	94±1	3±1	1±0.1	338±3	40±1	1±0.1
H ₂ O ₂ +2 mg/L Bacsan-Cu	358±1	57±3	16±2	241±2	27±2	1±0.1	92±2	6±1	1±0.1	352±6	88±1	35±1
H ₂ O ₂ +10 mg/L Bacsan-Cu	374±2	166±9	88±2	319±3	143±8	72±2	157±5	16±1	2±0.1	387±7	202±8	122±1

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