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1 *Impact of Pseudomonas graminis* strain CPA-7 on respiration and
2 ethylene production in fresh-cut ‘Golden delicious’ apple according to
3 the maturity stage and the preservation strategy

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11 **Abstract**

12 The effect of the biocontrol agent (BCA) *Pseudomonas graminis* CPA-7 on the
13 accumulation of CO₂ and ethylene (C₂H₄) in fresh-cut apples at two maturity stages
14 was evaluated in refrigerated conditions. The influence of factors involved in the
15 preservation strategy upon commercial conditions such as the antioxidant (AOX)
16 treatment and the storage system was included in the analysis. Regardless of the
17 maturity stage, the BCA reduced C₂H₄ levels within the MAP atmosphere in AOX-
18 untreated apples wedges, by 29 and 43 % in immature and mature apples,
19 respectively. Nevertheless, the addition of ascorbate as antioxidant counteracted this
20 effect. *In vitro* tests suggested that the reduction of C₂H₄ levels was not associated to
21 the uptake of this molecule by CPA-7. Interestingly, in non-inoculated samples AOX
22 treatment showed contradictory effects on C₂H₄ production in MAP conditions by
23 significantly reducing C₂H₄ levels in immature apples (by 23 %) while increasing it in
24 mature ones (by 40 %). Similarly, CPA-7 had opposite effects on the CO₂ accumulation
25 pattern depending on the storage system or the fruit maturity stage. In this sense, CPA-
26 7 was associated to higher fruit respiratory activity at advanced maturity stages yet

27 without inducing the fruit fermentative metabolism or altering fruit quality during a week
28 of refrigerated storage. Overall, these results show that CPA-7 may contribute to the
29 overall maintenance of the microbiological and physicochemical quality of fresh-cut
30 apple by modulating the fruit ethylene production and/or respiration.

31 **Key words:**

32 **Antagonist, fresh-cut fruit physiology, passive modified atmosphere packaging, MAP**

33 **1. Introduction**

34 The effect of the application of antagonists on fresh-cut produce in commercial
35 conditions is influenced by internal factors like the type and the maturity stage of the
36 commodity and external factors such as temperature, preservative treatments as well
37 as oxygen and carbon dioxide concentrations within packages. From the physiological
38 stand, processed products essentially behave as wounded tissues where the disruption
39 of cell compartmentalization lead to the mixture of cellular components with an
40 increase of enzymatic and respiratory activities as well as an elevated production of
41 ethylene (C₂H₄) (Hodges and Toivonen, 2008; Mahajan et al., 2014). C₂H₄ is also
42 indeed, the main hormone controlling ripening in climacteric fruit (Reid et al., 1973) and
43 its biosynthesis involves the transformation S-adenosylmethionine into the precursor 1-
44 aminocyclopropane (ACC) mediated by the enzyme ACC synthase (ACS). ACC is later
45 converted to ethylene by ACC oxidase (ACO) (Yang and Hoffman, 1984). In apple, as
46 a fruit with a climacteric behavior, the regulation of these two steps is auto-inhibitory
47 during fruit development prior to ripening and auto-stimulatory at the onset of ripening
48 (Tatsuki, Endo, & Ohkawa, 2007; Wang et al., 2009; Lelièvre, Latché, Jones,
49 Bouzayen, & Pech, 1997). Processing implies the mechanical injury of fruit tissues
50 which induces the activity and synthesis of ACS leading to the formation of “wound
51 ethylene” whose accumulation may be enough to activate the climacteric phase
52 depending of the size and permeability of packages (Lamikanra, Imam, & Ukuku, 2005;
53 Yu & Yang, 1980). However, the low availability of ACO in pre-climacteric apples is a

54 limiting factor regulating ethylene production upon cutting due to the reduced capability
55 for the conversion of ACC into ethylene (Lara and Vendrell, 2000). In the post-
56 climateric stage the capacity for ethylene production is also reduced and response to
57 wounding is more limited than in the climateric stage (Abeles et al., 1993). Therefore,
58 the physiological stage of the commodity constitutes a key factor that should be taken
59 into account in the production of fresh-cut fruit products (Toivonen and Dell, 2002).

60 Respiration also shows a biphasic rise during the development of climateric
61 commodities, the first one early in development and the second one during ripening or
62 senescence. The second peak usually precedes the autocatalytic ethylene synthesis
63 stage (Fonseca et al., 2002). Moreover, respiration is also induced by cutting due to the
64 loss of compartmentalization of the enzymes involved in the respiration pathways and
65 its substrates, and the activation of key regulatory steps of glycolysis and the
66 tricarboxylic acid cycle (Rolle and Chism, 1987). The mechanical injury of cell
67 membranes also activates the enzymatic degradation of its lipid components, with the
68 formation of long-chain fatty acids whose α -oxidation also causes a rise in respiration
69 (Rolle and Chism, 1987). It is also well established that 'wound ethylene' induces fruit
70 respiration (Yu and Yang, 1980). Furthermore, an increase in CO₂ production also
71 occurs in fresh-cut tissues due to the activation of cell repair processes, not only for
72 obtaining energy but for the synthesis of replacement structural compounds (Gomez-
73 Lopez, 2012). The accelerated oxidative breakage of organic substrates and the loss of
74 structure of membranes entailed by the above mentioned processes are detrimental to
75 the nutritional properties and the overall quality of fresh-cut fruit (Soliva-Fortuny &
76 Martín-Belloso, 2003).

77 To reduce both respiration and ethylene production several methods comprising
78 chilling conditions and modified atmosphere packaging (MAP) are amongst the most
79 currently used in the fresh-cut produce industry (Rupasinghe and Yu, 2013). The
80 addition of biocontrol agents (BCA) such as *Pseudomonas* spp. is another method that

81 could contribute to modulate ethylene levels thereby extending the shelf-life of fresh
82 ready-to-eat products. Mechanisms for the modulation of plant ethylene metabolism by
83 *Pseudomonas* spp. have been already documented and may imply both its
84 exacerbation or its reduction (Fatima & Anjum, 2017; Glick, 2014; Hase et al., 2003).
85 To accomplish the first mentioned effect, Pseudomonads enhance the plant capacity to
86 transform the precursor ACC into ethylene, inducing the expression of C₂H₄-responsive
87 genes (Hase et al., 2003). Consequently, systemic induced resistance (ISR) is
88 triggered or primed allowing plants to respond better to a subsequent infection by a
89 broad spectrum of pathogens (Van Wees et al., 1997). The C₂H₄-reducing effect has
90 been observed in plants upon treatment with pseudomonads with ACC deaminase
91 (ACD) activity (Hernández-León et al., 2015; Singh et al., 2015). ACD cleaves ACC
92 into ammonia and α-ketobutyrate (Honma and Shimomura, 1978) lowering the amount
93 of available ACC and therefore limiting ethylene synthesis (Glick, 2014). As a
94 consequence of this process pseudomonads can delay ripening and senescence,
95 promote growth, prime resistance mechanisms and alleviate deleterious ethylene-
96 mediated plant stresses (Eckert et al., 2014; Glick, 2005; Wang, Knill, Glick, & Défago,
97 2000). Belonging to this bacterial group is *Pseudomonas graminis* CPA-7, an apple
98 epiphyte biopreservative strain which controls foodborne pathogens such as *Listeria*
99 *monocytogenes*, *Escherichia coli* and *Salmonella enterica* on fresh-cut fruit (Alegre et
100 al., 2013a, 2013b; Abadias et al., 2014; Collazo et al., 2017) and modulate oxidative
101 metabolism in fresh-cut apple (Collazo et al., 2018). In an attempt to clarify its mode of
102 action we also investigated the possibility for CPA-7 to modulate the ethylene
103 metabolism in fresh-cut apple thereby influencing the defense response and/or the
104 senescence of this fruit. With this in mind, we monitored the effect of the antagonist in
105 fresh-cut apples as affected by several factors involved in production and commercial
106 conditions (antioxidant treatment, packaging headspace gas composition, and the
107 maturity stage of the commodity). In addition, to assess the effect of CPA-7 metabolic

108 activity in supplying exogenous ethylene or metabolizing the produced by the fruit, the
109 ability of the antagonist to either produce or consume C₂H₄ was tested *in vitro*.

110 **2. Materials and methods**

111 **2.1 Antagonist inoculum preparation**

112 *Pseudomonas graminis* strain CPA-7 (Alegre et al., 2013b) inoculum was obtained by
113 centrifugation of 50 mL of an overnight culture in tryptone soy broth (TSB, Biokar,
114 Beauveais, France) at 25 °C in agitation (Collazo et al., 2017). The concentration of
115 the solution was checked by viable plate count of appropriate ten-fold dilutions in saline
116 peptone (8.5 g L⁻¹ NaCl, 1 g L⁻¹ peptone) onto TSA plates after incubation at 30 °C for
117 48 h.

118 **2.2 Fruit processing**

119 Apples (*Malus domestica* Borkh. cv. 'Golden delicious') used in this study were grown
120 in local farms (Lleida, Catalonia, Spain) and collected in august, 2017 at two maturity
121 stages (with a week of difference between harvests). Prior to experimental assays,
122 apples were washed with running tap water, surface disinfected with 700 mL L⁻¹
123 ethanol and either stored as such or processed (peeled with an electric fruit peeler and
124 cut into eight wedges with a handheld corer/slicer). Wedges were kept in chilled (5 °C)
125 chlorinated tap water (pH 6) until treatment and/or packaging.

126 **2.3 *In vitro* analysis of ethylene production or consumption by CPA-7**

127 **2.3.1 Preparation and inoculation of liquid culture media**

128 *In vitro* assays were performed in order to evaluate the putative ethylene production or
129 consumption by CPA-7 in a culture medium with a similar composition to the fruit but
130 discarding the changes due to the apple's native microbiota. For that, analysis glass
131 tubes containing 10 mL of apple juice were inoculated with CPA-7 to a concentration of
132 10⁵ CFU mL⁻¹. Aliquots of TSB were prepared, inoculated and analyzed in the same
133 way, as a control treatment. For juices preparation, apple wedges were previously
134 dipped in 6% NatureSeal[®] AS1 solution (AS1, AgriCoat Ltd., Great Shefford, UK), a

135 calcium ascorbate-based product or in cold deionized water for 2 min in agitation (15.7
136 rad s^{-1}) in a tabletop orbital shaker (Unimax 1010, Heidolph, Germany). Then, juices
137 were obtained in a commercial blender, subsequently filtered through cloth gauzes and
138 either adjusted to pH 6.5 with 1 mmol L^{-1} NaOH or sterilized as such at $215 \text{ }^\circ\text{C}$ for 5
139 min and stored at $5 \text{ }^\circ\text{C}$ until use. Non-inoculated aliquots of each culture medium were
140 also prepared and used as controls. Inoculated and non-inoculated tubes were stored
141 in aerobic conditions at $5 \text{ }^\circ\text{C}$ or $25 \text{ }^\circ\text{C}$ in agitation for 7 d. *In vitro* assays were repeated
142 twice and included three replicates per treatment.

143 **2.3.2 *In vitro* microbial dynamics**

144 CPA-7 population dynamics in each culture medium was tracked by viable plate count
145 on TSA at 0, 1, 3, 6, and 7 d post-inoculation, as described in section 2.1.

146 **2.3.3 *In vitro* CO₂ accumulation pattern**

147 The headspace gas composition (percentages of O₂ and CO₂) of each culture tube was
148 measured at 0, 1, 3, 6, and 7 d post-inoculation using a handheld gas analyzer
149 (CheckPoint O₂/CO₂, PBI Dansensor, Denmark). Before each measurement tubes
150 were hermetically closed for 12 h. CO₂ accumulation was expressed in mg mL^{-1} liquid
151 culture medium.

152 **2.3.4 *In vitro* ethylene accumulation**

153 The ethylene accumulation patterns of cultures tubes previously sealed for 12 h were
154 determined at 0, 1 and 3, 6 and 7 d post-inoculation. For that, 1 mL of gas sample was
155 withdrawn daily from each jar or tray with a syringe and injected into a gas
156 chromatograph (Agilent Technologies 6890, Wilmington, Germany) fitted with a FID
157 detector and an alumina column F1 80/100 ($2 \text{ m} \times 1/8 \times 2.1$, Tecknokroma, Barcelona,
158 Spain). The injector and detector were kept at $180 \text{ }^\circ\text{C}$ and $280 \text{ }^\circ\text{C}$, respectively.
159 Quantification was carried out by comparing the gas chromatography signal of the
160 samples to that of a $21 \text{ } \mu\text{L L}^{-1}$ C₂H₄ standard (Carbueros metálicos SL, Aragón, Spain).
161 Ethylene accumulation within the storage atmosphere of the tubes was expressed as
162 $\mu\text{L mL}^{-1}$ culture medium. Putative ethylene degradation by CPA-7 was assessed by

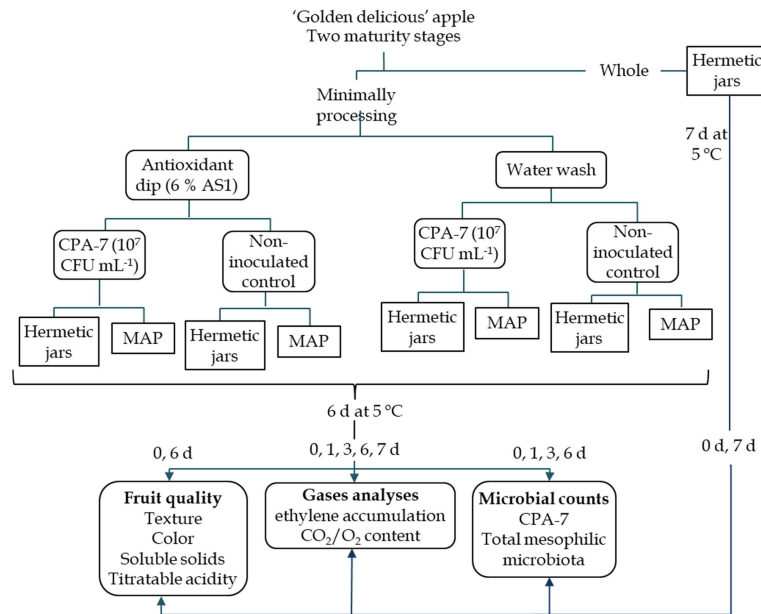
163 tracking the reduction of 1 mL of 21 $\mu\text{L L}^{-1}$ C_2H_4 standard injected with a syringe in
164 hermetically sealed tubes (containing 10 mL of either apple juice or TSB medium and
165 12 mL of headspace).

166 **2.4 *In vivo* analysis of ethylene and CO_2 accumulation patterns in CPA-7-** 167 **inoculated fresh-cut apples**

168 **2.4.1 Fruit treatment and packaging**

169 The experimental setup of *in vivo* tests is shown in Figure 1. For dip inoculation of
170 apple wedges, CPA-7 suspensions at a concentration of 10^7 CFU mL^{-1} were prepared
171 in cold deionized water (4 °C) or in cold 60 g L^{-1} AS1 antioxidant aqueous solution.
172 Fruit wedges were dipped in the bacterial suspensions or in non-inoculated water or
173 antioxidant solution as controls, at a ratio of 1:2 (weight of fruit: volume of solution) as
174 described in section 2.3.1.

175 After the drainage of the excess of water, the treated apple wedges were packaged in
176 two storage systems; hermetic jars (static system) and commercial trays (MAP). For
177 the static system, 1 kg of intact apples was stored in 3.4 L jars and 500 g of apples
178 wedges was stored in 1.7 L jars. Jars were equipped with a silicon septum for sampling
179 the gas of the headspace. For MAP, 120 g of processed fruit were placed in 400 mL
180 polyethylene terephthalate ShelfMaster™ Pronto™ trays (PlusPack, Denmark) and
181 thermosealed with 181.7 cm^2 of a 3-holed (60 - 80 μm diameter, 75 mm spacing)
182 multilayered microperforated film (polyester anti-fog film, OALF (14 μm of thickness) +
183 oriented polypropylene, OPP (20 μm of thickness) (PDS Group, Murcia, Spain) to
184 achieve passive modified atmosphere. Trays and jars were stored statically at 5 °C in
185 darkness. Each tray/jar was considered as a replicate and three replicates per
186 treatment and sampling time were included.



187

188 Figure 1. Experimental workflow of *in vivo* trials

189 **2.4.2 *In vivo* microbial dynamics**

190 CPA-7 as well as total mesophilic microorganisms' population dynamics on apple
 191 wedges stored in trays were analyzed at 0, 1, 3 and 6 d post-inoculation. For this, 10 g
 192 of apple from each tray was homogenized in 90 mL buffered peptone water (BPW,
 193 Biokar, Beauvais, France) and analyzed by viable cells count on TSA plates for CPA-7
 194 and on plate count agar plates (PCA, Biokar, Beauvais, France) for total mesophilic
 195 microbiota. Plates were incubated at 30 °C for 48 h. In the same way, microbial
 196 populations in jars were determined at 0 and 7 d post-inoculation. Microbiological data
 197 were expressed as colony forming units per gram of fresh weight of fruit (CFU g⁻¹ FW)
 198 and transformed to log₁₀ for subsequent statistical analyses.

199 **2.4.3 *In vivo* CO₂ accumulation pattern**

200 The headspace gas composition (percentages of O₂ and CO₂) of each replicate, stored
 201 in the static system (jars) or in MAP (trays), was measured at 7 h post-inoculation and
 202 then at 1, 2, 3, 6, and 7 d as described in section 2.3.3. CO₂ accumulation was
 203 expressed relative to the fresh weight of fruit (mg kg⁻¹).

204 **2.4.4 *In vivo* ethylene accumulation pattern**

205 To determine *in vivo* ethylene accumulation samples were taken from trays and jars at
206 0, 1, 3, 6, and 7 d post-treatment. Ethylene accumulation within the storage
207 atmosphere was expressed as $\mu\text{L kg}^{-1}$ fresh weight fruit.

208 **2.2.5 Fruit quality parameters**

209 Texture of whole and processed apples as well as color, pH, soluble solids and
210 titratable acidity of apple wedges were determined as described elsewhere (Alegre et
211 al., 2013a). Quality parameters were measured initially and at the end of storage in the
212 case of whole apples and wedges stored in jars while in processed apples stored in
213 MAP they were measured at 0, 1, 3 and 6 d post-treatment. Texture and pH were
214 measured in five wedges per replicate per treatment at each sampling time. Two
215 measures of color, one per side, were performed on five wedges per replicate per
216 sampling time. Soluble solids and titratable acidity of each replicate were measured
217 initially and at the end of storage regardless of the storage system. Low values of CIE
218 color parameter L^* and high values of a^* were considered as indicators of surface
219 browning intensity (Sapers and Douglas, 1987). The concentration of soluble solids at
220 20 °C was expressed as mass fraction (%). Titratable acidity was measured in 10 mL
221 of pulp and was expressed as malic acid content (g L^{-1} juice).

222 **2.5 Statistical analysis**

223 Data were analyzed using the general linear model procedure to determine the
224 treatment and interaction effects, with the statistical software JMP (version 11 SAS
225 Institute Inc., NC, USA). All data were verified for normal distribution and
226 homoscedasticity of residues. Results were schematically represented as means \pm
227 standard deviation. Means were compared by analysis of variance (ANOVA) and
228 separated by Tukey's test ($P < 0.05$).

229 **3 Results and discussion**

230 **3.1 In vitro analysis of microbial dynamics and ethylene production or**
231 **consumption by CPA-7**

232 To test the ability of CPA-7 to growth and produce or consume ethylene in sterile apple
233 juice, microbial populations and gas headspace composition were measured during 7
234 days. The same analyses were performed in TSB, a synthetic media usually used for
235 CPA-7 culture in the laboratory, as a control treatment (Abadias et al., 2014; Alegre et
236 al., 2013a). Results showed that CPA-7 initial populations (10^5 CFU mL⁻¹) increased by
237 1 log₁₀ in TSB after 24 h at 5 °C and afterwards they remained stable up to day 6 (Fig.
238 2).

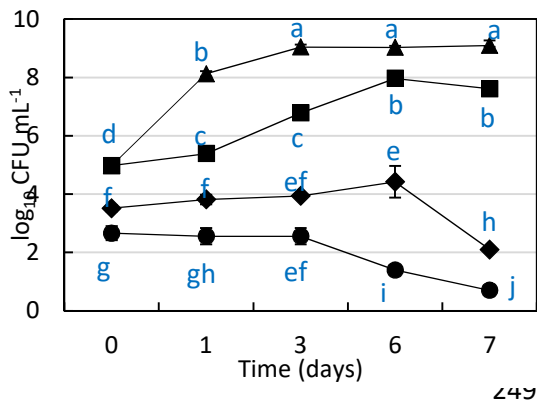


Figure 2. CPA-7 in vitro population dynamics in different growth conditions: apple juice pH 6.5, at 25 °C (▲), apple juice pH 6.5, at 5 °C (■), TSB medium at 5 °C (◆), and apple juice pH 4.5, at 5 °C (●). Symbols represent means and error bars represent standard deviation of the mean (n = 6). Different letters represent significant differences among treatments according to analysis of variances (ANOVA) and Tukey's test with a 95 % confidence (p < 0.0001).

250 No growth was observed in AOX-untreated apple juice pH 4.5 but instead, the
251 population gradually decreased to levels close to detection limit (5 CFU mL⁻¹), thus the
252 gas analysis of these samples was stopped after day 3. In accordance with this result,
253 previous studies shown that CPA-7 failed to growth in synthetic liquid media adjusted
254 with different organic acids to pH ranging from 4.5 to 5 (Iglesias, 2017). CPA-7 also
255 showed limited growth in melon juices compared to fruit pieces, which is a fruit with pH
256 close to neutrality but rich in citric acid (Collazo et al., 2017) . Therefore, the
257 subsequent analyses were performed in apple juices with pH adjusted to 6.5. Results
258 showed that CPA-7 populations quickly increased (by 3 log₁₀ in the first 24 h) in apple
259 juice pH 6.5 incubated at 25 °C reaching the stationary phase (9 ± 0.1 log₁₀ CFU mL⁻¹)
260 before day 3 as expected for the optimal growth temperature (Alegre et al., 2013b).

261 When incubated at 5 °C CPA-7 populations in apple juice pH 6.5 increased slower than
262 at 25 °C, reaching the stationary phase on day 6 and attaining levels 2 log₁₀ higher than
263 when grown in TSB (pH 7.3). In general results showed that apple juice pH 6.5 a
264 suitable culture medium for CPA-7, showing similar population dynamics to that
265 observed in fresh-cut melon (pH 6.4) incubated at a temperature within the optimal
266 range (20 °C) (Abadias et al., 2014).

267 CO₂ production was close to zero during the whole evaluated period in the samples
268 grown in TSB as well as in apple juice pH 4.5 (data not shown), which was in
269 accordance with the lower growth observed in those culture media compared to apple
270 juice pH 6.5. Likewise, greater CO₂ levels correlated with the populations dynamics
271 observed in AOX-treated apple juices (pH 6.5) stored at 5 °C or 25 °C, where the CO₂
272 production remained stable around 0.04 ± 0.04 mg L⁻¹ h⁻¹ or increased from 1.5 ± 0.4 to
273 2.9 ± 0.9 mg L⁻¹ h⁻¹, respectively.

274 Regardless of the temperature of storage no ethylene production was observed for
275 CPA-7 whether it was grown in TSB or in apple juices (pH 6.5 or pH 4.5). In the same
276 way, no differences in ethylene accumulation or in the antagonist population levels
277 were detected in the inoculated samples after supplementation with exogenous C₂H₄
278 compared to the non-supplemented control at any of the analyzed sampling points in
279 any of the culture media or incubation temperatures assayed (data not shown). Overall,
280 results obtained from *in vitro* assays showed that although CPA-7 is able to use the
281 nutrients present in the apple-based food matrix tested to grow, it is unable to produce
282 or consume ethylene in the conditions tested.

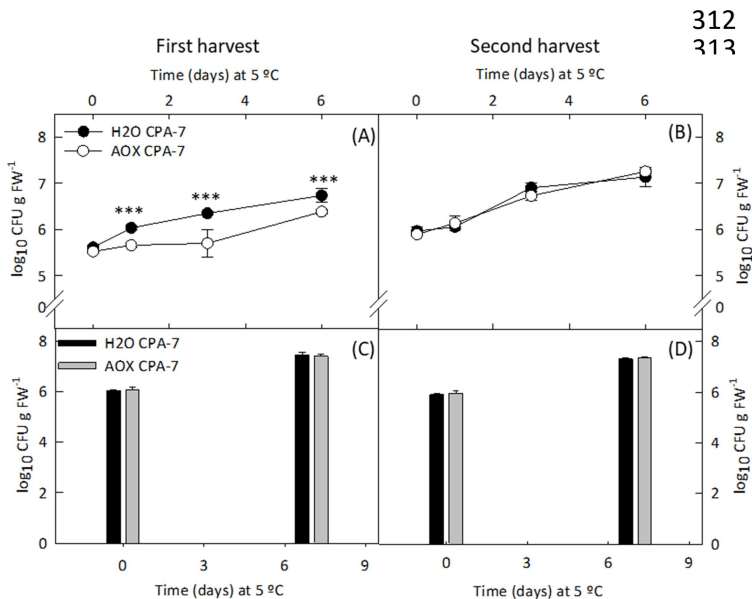
283 **3.2 *In vivo* analysis of ethylene and CO₂ accumulation patterns in CPA-7-** 284 **inoculated fresh-cut apples**

285 **3.2.1 *In vivo* microbial population dynamics**

286 In MAP-stored apples CPA-7 populations were initially 5.53 ± 0.04 and 5.90 ± 0.04
287 log₁₀ CFU g⁻¹ FW in samples from the first (H1) and the second harvest (H2),

288 respectively. Afterwards, in samples upon the antioxidant effect, CPA-7 populations
289 showed a slower growth and reached lower levels than in AOX-untreated samples (Fig.
290 3 A). These results agreed with the those observed for this bacteria when grown on
291 fresh-cut pear upon another antioxidant treatment but in similar storage conditions
292 (Iglesias et al., 2018). Conversely, in apples from the second harvest no differences
293 were observed in AOX-treated and untreated samples during the whole storage period
294 (Fig. 3 B). Likewise, Alegre et al. (2013a) selected the AS1 antioxidant treatment for
295 commercial assays with CPA-7 in 'Golden Delicious' fresh-cut apple wedges as they
296 observed no differences in growth between AS1-treated or untreated samples
297 incubated at 10 °C during 2 d. Furthermore, they observed increases by 2 log₁₀ AS1-
298 treated samples after 7d of storage at 10 °C. Discrepancies in the effect of antioxidant
299 treatments on CPA-7 growth between that study and ours might be related to the
300 maturity stage of the commodity used for the different experiments. However, CPA-7
301 grew more in H2 (by 1.2 ± 0.2 log₁₀) than in H1 apples (by 0.8 ± 0.1 log₁₀) regardless of
302 the antioxidant application. On the other hand, when stored in jars, CPA-7 populations
303 on apple wedges showed similar dynamics regardless of the harvest date or the
304 antioxidant treatment (Fig. 3 C and D).

305 Mesophilic bacteria populations were initially at the same levels in samples from both
306 harvest dates (2.3 ± 0.2 log₁₀ CFU g⁻¹ FW). As observed for CPA-7, mesophilic bacteria
307 grew more in the AOX-untreated H1 apples (by 1.9 ± 0.4 log₁₀) than in the rest of the
308 samples (by 0.7 ± 0.1 log₁₀) (data not shown). Those results confirmed the inhibitory
309 effect that antioxidant agents have on microbial growth as previously observed in
310 several studies performed with fresh-cut fruit treated with different antioxidant
311 compounds and mixtures (Iglesias et al., 2018).



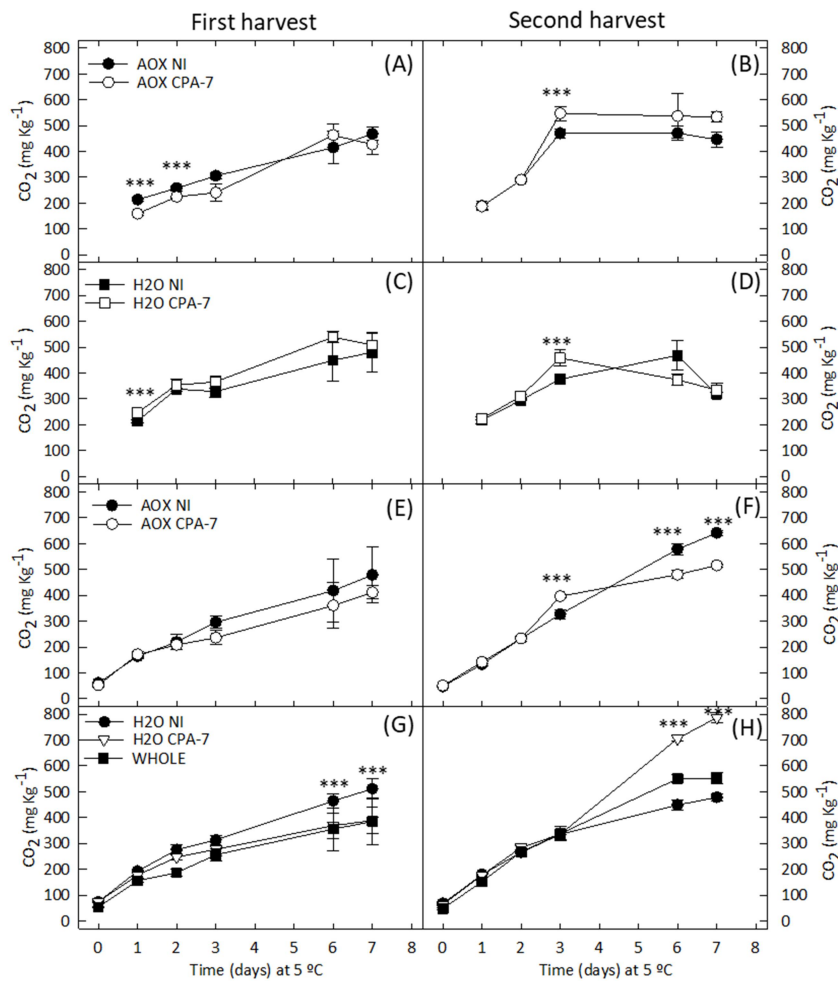
312 Figure 3. CPA-7 population
 313 dynamics on antioxidant-treated (AOX) or untreated (H₂O) fresh-cut 'Golden Delicious' apple wedges from the first (left column) and the second harvest (right column) dates during storage in trays (MAP conditions) (A-B) or in hermetically sealed jars (C-D) at 5 °C. Symbols represent means and error bars represent standard deviation of the mean (n = 6). Asterisks represent significant differences among treatments according to analysis of variances (ANOVA) and Tukey's test with a 95 % confidence (***) for p < 0.0001).

330 3.2.2 *In vivo* CO₂ accumulation pattern

331 The headspace gas composition of packages was highly influenced by the permeability
 332 of the storage system and the physiological stage of the fruit (Supplementary Table 1).
 333 As expected, apples from the second harvest (H2) showed higher respiration rates
 334 than those from the first one (H1) thereby depleting faster O₂ levels. This led to anoxic
 335 conditions (0.4 ± 0.3 %) by day 7 of storage in the hermetic system in H2 inoculated
 336 samples while in H1 apples O₂ levels only decreased to 12 ± 1 % in the same period. In
 337 the permeable system a stable O₂ concentration ranging from 15 to 17 % was
 338 maintained throughout storage.

339 Like ethylene production, respiration is activated both during ripening and upon
 340 wounding-stress thus, during the processing of fresh-cut produce (Saltveit, 2016).
 341 Accordingly, our results showed an initial rise in CO₂ production by 40 % in processed
 342 apples compared to intact apples in both maturity stages (Fig. 4 G-H). The magnitude
 343 of the wound-response in the respiration rate was significantly higher in mature than in
 344 immature apples exceeding by 2.5-fold and 1.4-fold, respectively, the respiration rate of
 345 whole apples. The exacerbation of respiration rate in fresh-cut apple slices by 2 - 3
 346 times compared to whole fruit had previously been reported (Lakakul, Beaudry, &
 347 Hernandez, 1999). The increased respiration in minimally processed fruit is mostly due

348 to a physiological response to wounding which is tightly linked to the maturity stage,
 349 since the removal of the peel barely reduce the resistance to O₂ diffusion in apples
 350 (Fonseca et al., 2002). In general, we observed that respiration rate reached the
 351 highest values immediately after processing and subsequently declined throughout
 352 storage regardless of the combination of factors analyzed.



353

354 Figure 4. CO₂ accumulation patterns in CPA-7-inoculated and non-inoculated (NI) fresh-cut 'Golden delicious' apple
 355 wedges from the first (left column) and the second harvest (right column) dates when stored in MAP (A-D) and in
 356 hermetic jars (E-H). Graphs A-B and E-F: apples treated with the antioxidant (AOX); graphs C-D and G-H: AOX-
 357 untreated samples. Symbols represent means of three biological replicates and error bars represent standard
 358 deviations. Asterisks represent significant differences according to an analysis of variances ANOVA and Tukey's test
 359 with a 95 % confidence (* for p < 0.05, ** for p < 0.01, *** for p < 0.001). Asterisks below the lines represent
 360 differences only between whole and processed apples, regardless of the inoculation with CPA-7.

361 In MAP storage, CO₂ accumulation in H1 AOX-untreated samples showed an increase
 362 by 16 % in the presence of CPA-7 compared to non-inoculated control, 24 h post-

363 processing (Fig. 4 C). The addition of the antioxidant reversed this behavior, being CO₂
364 production in inoculated samples up to 25 % lower than the control during the first two
365 days in MAP conditions (Fig. 4 A). In H2 samples differences in CO₂ accumulation
366 pattern were only observed on the 3rd day of MAP storage being remarkably higher in
367 response to the antagonist than in the control whether they were treated (by 16 %) or
368 not (by 22 %) with the antioxidant. This could be associated to the climacteric peak of
369 the product as it was correlated with a rise in ethylene production (Lelièvre et al.,
370 1997). According to this result, CPA-7 may enhance the climacteric peak in processed
371 apples at more advanced maturity stages in agreement with previous findings showing
372 that once the autocatalytic phase of ripening has begun the effectiveness of
373 preservative methods is considerably reduced and shelf-life is shortened (Rojas-Graü
374 et al., 2007; Soliva-Fortuny, Oms-Oliu, & Martin-Belloso, 2002). Previous experiments
375 performed with fresh-cut 'Fuji' apple slices showed that according to storage
376 conditions, reducing agents such as ascorbic acid influence other physiological
377 processes in addition of preventing oxidation (Gil et al., 1998). In that study, ascorbic
378 acid dips reduced respiration rate as well as ethylene production in 'Fuji' apples stored
379 in air (21% O₂, 0% CO₂) while increased respiration in MAP- stored (0% O₂, 0% CO₂,
380 100% N₂) apple slices (Gil et al., 1998).

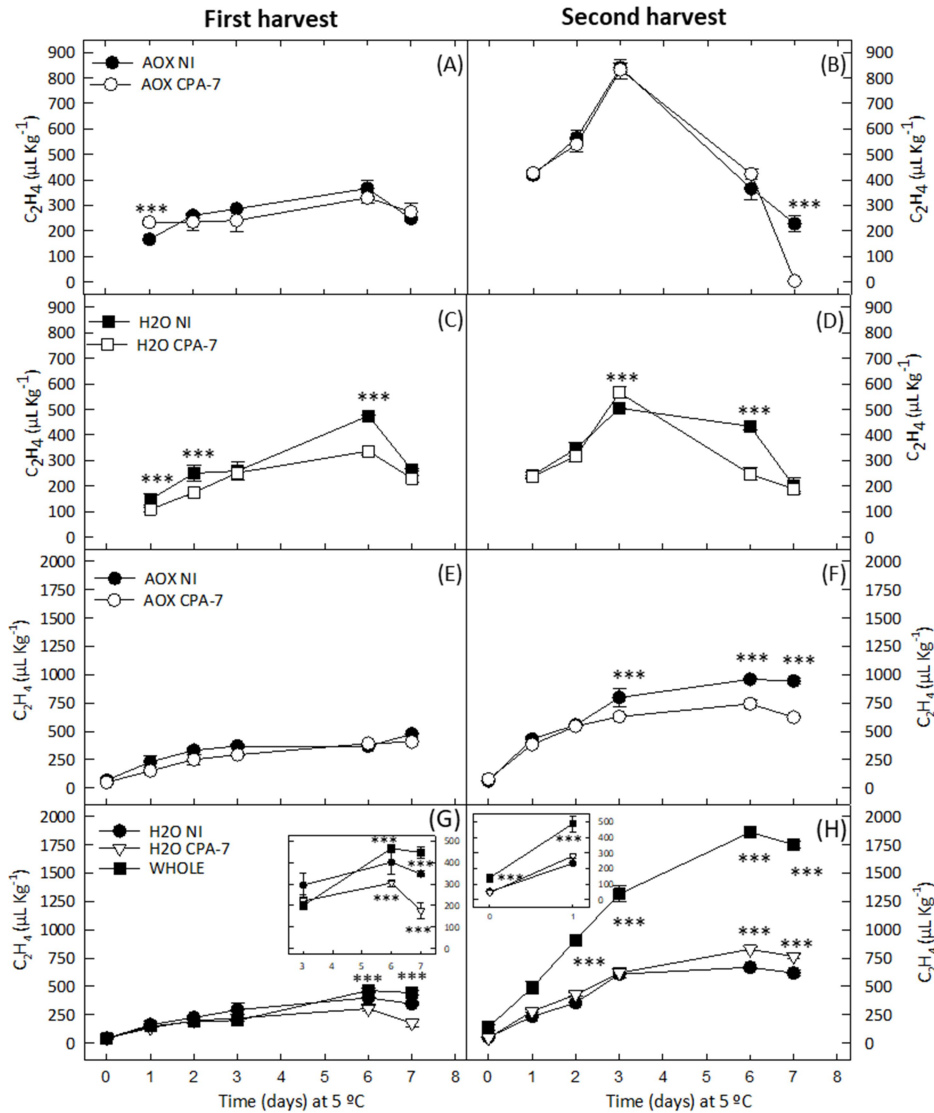
381 Contrastingly in the hermetic system (jars), no differences were observed between
382 inoculated and non-inoculated apples during the first two days of storage regardless of
383 the addition of the antioxidant or the harvest date. However, contradictory effects of
384 CPA-7 on CO₂ accumulation according to the maturity stage and the treatment with the
385 antioxidant were observed in the hermetic system from the 3rd day of storage even
386 when population levels were not influenced by the mentioned factors. For instance a
387 decrease (by 22 %; Fig. 4 G) or increase (by 67 %; Fig. 4 H) in CO₂ accumulation was
388 observed in CPA-7-inoculated AOX-untreated samples from H1 and H2, respectively,
389 compared to non-inoculated controls. The impact of CPA-7 on CO₂ accumulation in the

390 hermetic system was either eliminated (no differences in H1 samples) or inverted
391 (higher by 22 % than the H2 control) upon the antioxidant treatment in both maturity
392 stages (Fig. 4E-F).

393 The influence of external factors such as temperature and the gas headspace
394 composition (O₂ and CO₂) of the packages on the respiration of fresh-cut commodities
395 is generally recognized and thus it has been included in several models for the design
396 of MAP technologies (Fagundes, Carciofi, & Monteiro, 2013; Lakakul et al., 1999).
397 However the combination of these factors and biological or chemical preservatives has
398 not been so well studied. Investigation on this matter is needed for developing models
399 to predict the shelf-life of fresh-cut produce in commercial conditions.

400 *3.2.3 In vivo ethylene accumulation pattern*

401 The initial ethylene production of intact apples from the second harvest date (H2) was
402 3.7-fold higher than those from the first one (H1) evidencing their more advanced
403 physiological stage (Lelièvre et al., 1997), thus they will be henceforth referred as
404 mature and immature apples, respectively. In the same way, C₂H₄ production in intact
405 H2 apples remained 3-fold higher than processed ones during the whole storage while
406 it showed no differences in H1 fruit (Fig 5G-H) suggesting the pre-climateric stage of
407 the fruit from the earliest harvest (Chaves and de Mello-Farias, 2006; Oetiker and
408 Yang, 1995). The immediate effect of processing in C₂H₄ production was highly
409 influenced by the maturity stage, showing a reduction in mature apples wedges
410 compared to intact fruit and not differences in immature fruit. Contradictory effects of
411 cutting on ethylene production, as previously observed in several climateric
412 commodities, have been explained by differences in the physiological stage of the fruit
413 and linked to differential expression patterns of the genes involved in the ethylene
414 metabolism, before, during and after, the climateric phase (Bapat et al., 2010;
415 Toivonen & Dell, 2002; Vilanova et al., 2017).



416

417 Figure 5. Ethylene accumulation in CPA-7-inoculated and non-inoculated (NI) fresh-cut 'Golden delicious' apples
 418 from the first (left column) and the second harvest (right column) dates when stored in MAP (A-D) or in hermetic
 419 jars (E-H). Graphs A-B and E-F: apples treated with the antioxidant (AOX); graphs C-D and G-H: AOX-untreated
 420 samples. Inserts in graphs G and H are plots of the same data in a smaller scale. Symbols represent means of three
 421 biological replicates and error bars represent standard deviations. Asterisks represent significant differences
 422 according to an analysis of variances ANOVA and a Tukey's test with a 95 % confidence (* for $p < 0.05$, ** for $p <$
 423 0.01 , *** for $p < 0.001$).

424 The impact of the application of CPA-7 on fresh-cut fruit ethylene metabolism cannot
 425 be separated from that of other factors such as the gas internal composition of
 426 packages, the maturity stage and the preservative chemical treatment which has
 427 previously shown to have significant influence on the physiology of this kind of products
 428 (Rojas-Graü et al., 2007). In MAP storage, the ethylene accumulation pattern in
 429 processed AOX-untreated apples was different according to the maturity stage (Fig. 5,

430 C-D) showing a peak on day 6 in H1 apples and on day 3 in H2 ones. Similarly, during
431 the first days of storage, differential effects of the application of CPA-7 on the ethylene
432 accumulation within the MAP atmosphere were observed in processed AOX-untreated
433 apples, causing reduced C₂H₄ levels (by 30 %) in immature apples on day 2 while it
434 enhanced them by 11 % in mature apples on day 3. However, as time passed, CPA-7
435 was associated to a reduction in C₂H₄ accumulation by about 29 % and 43 % in H1 and
436 H2 apples, respectively. This may have in turn contributed to a reduction of the
437 ethylene-mediated fruit senescence (Czarny et al., 2006) and hence, partially explain
438 the improved quality of CPA-7 inoculated apples.

439 The antioxidant treatment effectively reduced C₂H₄ levels in H1 apples upon MAP
440 conditions during the whole storage while markedly enhanced it in H2 apples (Fig. 5A-
441 B). In general, CPA-7 effect on the C₂H₄ levels within MAP packages was suppressed
442 by the antioxidant, except for the increase by 29 % during the first 24 h post-processing
443 in H1 apples. Likewise, no differences were observed regarding C₂H₄ accumulation
444 within the MAP atmosphere in mature apples until day 6 when it dropped by 98 % in
445 the presence of CPA-7.

446 In our trials, the reduction of C₂H₄ levels cannot be explained by the consumption of
447 ethylene by the antagonist since the levels of this molecule remained invariable after its
448 supplementation *in vitro*, both in TSB medium and in apples juices. Alternatively, the
449 reducing effect of this bacterium on the fruit ethylene production could be associated to
450 1-aminocyclopropane-1-carboxylic acid (ACC)-deaminase activity. This hypothesis is
451 suggested by the presence in the bacterial genome of a gene fragment encoding a
452 protein (Collazo et al., unpublished results) that shares 98 % homology with the ACC-
453 deaminase (ACD) enzyme (GenBank Acc. No. WP_065986140.1) belonging to *P.*
454 *graminis* strain P 294/08 (Behrendt et al., 1999). However, further analysis should be
455 performed in order to demonstrate the functionality of this gene in CPA-7. Following
456 this thought, no remarkable reduction of C₂H₄ can be attributed to ACD activity soon

457 after processing since the initial response to wounding has been suggested to deplete
458 the existing pool of ACC available within plant cell through the rapid action of ACC
459 oxidase (ACO), which has greater affinity for ACC than ACD (Glick, 2014). However,
460 ACD activity could explain the reduction of the second ethylene peak, probably
461 associated to *de novo* production of ACC, that was observed in CPA-7-inoculated
462 samples after 6 d of storage. The decrease in C₂H₄ levels at that sampling point was
463 observed in both storage systems (MAP and hermetic) when O₂ concentration was
464 similar (14 – 16 %).

465 In the hermetic system, changes in the accumulation of ethylene in CPA-7-inoculated
466 AOX-untreated apple wedges were only observed at the end of storage and they
467 showed to be highly influenced by the maturity stage (Fig 5, G-H). In H1 apples CPA-7
468 reduced the C₂H₄ accumulation pattern by 24 and 49 % compared to the non-
469 inoculated control after 6 and 7 days of storage, respectively (Fig 5G). In H2 apples
470 inoculation with CPA-7 triggered higher C₂H₄ levels (by up to 19 %) after 6 d of storage
471 (Fig. 5H). The CPA-7 effect was either annulled (no differences in H1 apples) or
472 reversed (decreased by 34 % at day 7 in H2 apples) upon antioxidant treatment (Fig. 5
473 E-F). Similarly, a reduction of C₂H₄ levels in wounded climacteric melon inoculated with
474 the biological control agent *Bacillus subtilis* EXWB1 was also observed by Wang et al.
475 (2010) after 4 d of storage in hermetically sealed containers at 24 °C, which correlated
476 with a subsequent increase in the fruit defense response to fungal decay. In addition to
477 the maturity stage, the differences in the C₂H₄ accumulation patterns observed at the
478 end of storage in the hermetic system could also be influenced by O₂ availability, which
479 varied from 12 – 13 % for immature apples to 0.4 – 1.2 % for mature ones. However,
480 reduced C₂H₄ accumulation in respect of the control was observed in mature apples
481 stored in the hermetic system (2.9 – 1.7 % of O₂, 14 % CO₂) when treated with the
482 antioxidant. O₂ is a key factor in the regulation of *AcdS* gene, which encodes ACD
483 (Singh et al., 2015) but limited information is available regarding the influence of

484 antioxidants in this process. Results showed that antioxidant (AOX) dipping of mature
485 apples enhanced the accumulation of C₂H₄ both in the hermetic and the MAP storage
486 system, regardless of the inoculation with the antagonist. Ascorbic acid-based
487 antioxidant dipping has previously shown to increase ethylene production in fresh-cut
488 Fuji apples upon refrigerated conditions at a different timing and extent according to
489 oxygen availability and ripening stage, showing higher ethylene accumulation in MAP
490 than air conditions and in ripe than in unripe fruit (Rojas-Graü et al., 2007).

491 **3.2.4 Fruit quality parameters**

492 The initial firmness of whole apples was similar regardless of the moment of harvest
493 (67 ± 3 N) but it declined throughout storage in more extent in apples from the second
494 harvest (H2) than in those from the first one (H1): by 45 and 26 %, respectively. In
495 general, no significant differences in firmness were observed among treatments or
496 throughout storage in MAP or in jars, with values ranging from 13 ± 2 and 10 ± 1 N
497 (data not shown). No significant reduction of texture was either observed upon
498 inoculation with CPA-7 and MAP storage for 5 to 10 d at 5 or 10 °C in previous
499 experiments performed in fresh-cut apples, melons and pears (Abadias et al., 2014;
500 Alegre et al., 2013a; Iglesias et al., 2018). Those results suggest that CPA-7 does not
501 show an enhanced pectinolytic activity affecting fruit quality.

502 Soluble solids content (SSC) in unprocessed apples was initially higher in H2 apples
503 than in H1 ones (Table 1). It subsequently remained invariable throughout time in
504 apples from both harvests. Interestingly, samples treated with the antioxidant showed
505 an initial increase in the SSC in contrast with untreated samples regardless of the
506 maturity stage. Higher contents in soluble solids were also observed for AOX- treated
507 samples at the end of MAP storage in apples from the second harvest date. In general,
508 inoculation with CPA-7 did not alter SSC in any of the evaluated conditions. Similarly,
509 apple pH was about the same (4.0 ± 0.2) and remained stable throughout the studied
510 period for all the conditions tested. SSC was not significant altered by CPA-7 when

511 inoculated on fresh-cut melon, apples or pears as observed in previous experiments
 512 (Abadias et al., 2014; Alegre et al., 2013a; Iglesias et al., 2018).

513 Titratable acidity was initially lower in CPA-7 - inoculated H2 samples compared to the
 514 control while no differences were observed between inoculated and non-inoculated H1
 515 apples (Table 1). The opposite effect was observed at the end of storage in the
 516 hermetic system. In MAP conditions, this parameter did not show differences in the
 517 presence of CPA-7 at the end of storage. For the AOX-treated samples no differences
 518 were observed between inoculated and non-inoculated samples regardless of the
 519 maturity stage.

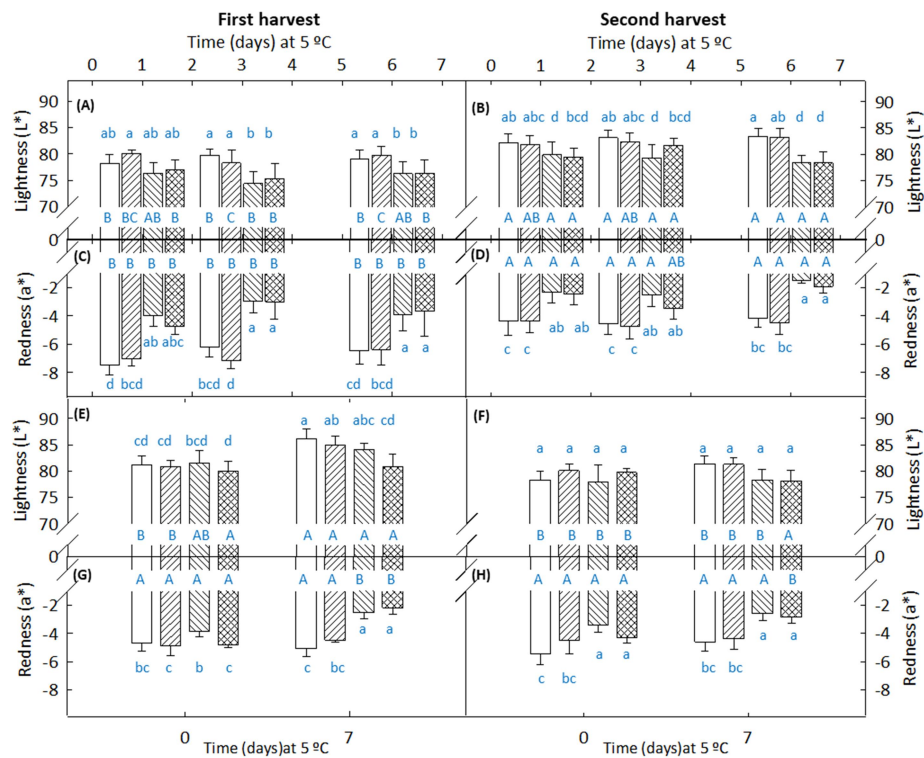
520 Table 1. Chemical quality parameters of intact (Whole) and fresh-cut ‘golden delicious’ apples from the 1st and the
 521 2nd harvest dates, inoculated with CPA-7 (CPA) or non-inoculated (NI), upon antioxidant treatment (AOX) or not
 522 (H₂O), when stored in trays (MAP) or in hermetic jars.

Parameter	Treatment	Jars and trays		Trays		Jars	
		day 0		day 6		day 7	
		1 st harvest	2 nd harvest	1 st harvest	2 nd harvest	1 st harvest	2 nd harvest
soluble solids (%)	Whole	12.5 ± 0.1 A	13.3±0.01aA			14±1 aB	13±0.9 aA
	AOX NI	12.8±0.01aA	13.6±0.2 aA	12.7±0.04aA	13.3 ±0.03aA	12.3±0.4aA	12.5±0.2 aB
	AOX CPA	12.9 ± 0.1aA	12.8±0.1 bA	12.6±0.1 aA	12.6 ±0.01aA	12.8±0.8aA	13.1±0.2 aA
	H2O NI	12.4 ± 0.1bA	12.4±0.04cA	12.5±0.03aA	11.6 ±0.04bB	12.1±0.2aA	12.4±0.4 aA
	H2O CPA	12.4 ±0.01bA	12.3±0.02cA	12.3±0.2 aA	11.7 ±0.01bB	11.2±0.4aA	12.6±0.1 aA
Titratable acidity malic acid (g L ⁻¹)	Whole	3.2 ± 0.1 aA	4.4±0.2 aA			2.9±0.4cA	4.6±0.4 aA
	AOX NI	3.3 ± 0.2 aA	3.3±0.1 aA	3.4 ±0.2 aA	3.8 ±0.3 aA	3.8±0.1aB	3.2±0.1 bA
	AOX CPA	3.22 ±0.02 aA	3.37±0.03aA	3.2 ±0.4 aA	3.6 ±0.5 abA	3.8±0.2aB	3.5±0.1 bA
	H2O NI	3.2 ± 0.1aA	2.8±0.5 abA	3.4 ±0.1 aA	3.0 ±0.2 bcA	4.1±0.1aB	3.2±0.1 bB
	H2O CPA	3.4 ±0.1aA	3.1±0.04bA	3.4 ±0.3 aA	3.1 ±0.3 bcA	3.3±0.1bA	3.1±0.04 bA

Results represent mean ± standard deviation. Different lowercase letters represent significant differences among treatments at each sampling time and uppercase letters represent significant differences throughout time for each treatment according to analysis of variances (ANOVA) and Tukey’s test (p < 0.05)

523 Inoculation with CPA-7 had no influence on a* and L* color parameters compared to
 524 non-inoculated control in any of the conditions tested (harvest date, storage system,
 525 antioxidant treatment and/or storage period) (Fig. 6). Similar results were obtained after
 526 14 d of storage at 5 °C in a previous study testing the effect of the inoculation with
 527 CPA-7 on the quality of fresh-cut apples in semi-commercial conditions resembling the
 528 ones assayed in the present work (Alegre et al., 2013a). In general, antioxidant

529 treatment resulted in a reduction of a^* after 1 d of MAP storage in inoculated and non-
 530 inoculated apples from both harvests (Fig. 6, C-D). However, this effect was more
 531 marked in H1 apple wedges than H2 ones. As observed for a^* , antioxidant dipping
 532 significantly improved and preserved lightness (L^*) throughout storage in both storage
 533 systems and harvest dates regardless of the inoculation with the antagonist (Fig. 6, A-
 534 B, E-F). The effect of AS1 on color maintenance of apple wedges was not that marked
 535 in the above mentioned study performed by Alegre et al., (2013a) and could be related
 536 to the influence of other parameters related with a different film used for MAP.



537

538 Figure 6. CIE color parameters: L^* (lightness) and a^* (redness) of fresh-cut apple wedges from the 1st (left column)
 539 and the 2nd (right column) harvest dates when stored in trays (MAP): A-D, or in hermetic jars: E-F. Columns
 540 represent means of 10 measures per each of three replicate treatments of: non-inoculated antioxidant-treated
 541 apples (□), CPA-7-inoculated antioxidant-treated apples (▨), non-inoculated water-washed apples (▩), CPA-7-
 542 inoculated water-washed (■). Error bars represent standard deviations. Different capital letters represent
 543 significant differences among sampling times for each treatment. Different lowercase letters represent significant
 544 differences among treatments for each sampling time, according to analysis of variances (ANOVA) and Tukey's test
 545 ($p < 0.05$).

546 In general, CPA-7 showed no negative effect on fruit quality parameters when
 547 combined with the antioxidant in the tested storage systems. AOX - dipping contributed

548 to maintain physical-chemical fruit quality in the conditions tested in agreement with
549 previous experiments showing that a calcium and ascorbate-containing antioxidant
550 treatment preserved fruit titratable acidity and soluble solids content in fresh-cut 'golden
551 delicious' apples (Soliva-Fortuny, Ricart-Coll, & Martín-Belloso, 2005). However, in
552 agreement with the results obtained by Soliva-Fortuny et al. (2002), we observed that
553 antioxidant dipping was more effective at preserving color parameters in apple wedges
554 from the first harvest than in those from the second one.

555 **Conclusions**

556 CPA-7 reduced the ethylene production in fresh-cut 'golden delicious' apples after 6 d
557 in MAP which could potentially delay fruit senescence increasing the shelf-life of these
558 kind of products in commercial conditions. However, the inhibition of this effect by the
559 ascorbate-based antioxidant treatment (AS1) limits the possibilities of the use of CPA-
560 7-AS1 combination as hurdle technology. Further studies on the effect of combined
561 strategies including this biocontrol agent and different types of natural chemical
562 preservatives and/or physical methods on the ethylene metabolism of fresh-cut apple,
563 could lead to the advent of better ecofriendly alternatives to improve the feasibility of
564 minimally processed products at a commercial level.

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Supplementary table 1. Headspace composition of packages of intact (Whole) and fresh-cut 'golden delicious' apples from the 1st and the 2nd harvest dates, inoculated with CPA-7 (CPA) or non-inoculated (NI), upon antioxidant treatment (AOX) or not (H2O), and when stored in trays (MAP) or in hermetic jars.

Treatment	O ₂ (%)							CO ₂ (%)						
	0.3 d	1 d	2 d	3 d	6 d	7 d	0.3 d	1 d	2 d	3 d	6 d	7 d		
Jars (hermetic)	1 st harvest	Whole	20.1 ± 0.1	17.8 ± 0.0	17.2 ± 0.4	16.5 ± 0.4	15.8 ± 0.1	15.6 ± 0.1	0.8 ± 0.1	2.5 ± 0.1	3.4 ± 0.3	4.9 ± 0.2	6.4 ± 0.2	6.5 ± 0.4
		NI AOX	18.3 ± 0.1	14.9 ± 0.1	13.6 ± 0.8	13.4 ± 0.7	12.3 ± 2	11.2 ± 0.5	1.3 ± 0.0	3.4 ± 0.0	5.8 ± 0.1	6.2 ± 0.1	8.2 ± 1.2	8.6 ± 0.4
		CPA AOX	18.4 ± 0.3	19.6 ± 0.2	15 ± 2	15 ± 2	15.6 ± 2	14.3 ± 3	1.1 ± 0.2	0.4 ± 0.0	4.0 ± 1	4.9 ± 2	5.4 ± 2	5.8 ± 3
	2 nd harvest	NI H2O	19.4 ± 0.2	15.9 ± 0.3	13.9 ± 0.3	13.0 ± 0.6	9.6 ± 1	8.9 ± 1	1.5 ± 0.2	4.1 ± 0.1	5.9 ± 0.5	7.2 ± 0.5	10.4 ± 0.7	10.3 ± 0.6
		CPA H2O	19.0 ± 0.0	16.1 ± 0.4	15.2 ± 0.6	14.2 ± 0.8	13.0 ± 1	11.9 ± 0.9	1.6 ± 0.1	3.8 ± 0.2	5.3 ± 0.3	6.2 ± 0.6	8.0 ± 1	8.2 ± 0.7
		Whole	18.7 ± 0.1	15.9 ± 0.4	13.4 ± 0.8	11.1 ± 0.4	7.6 ± 0.9	6.6 ± 1	1.5 ± 0.1	3.9 ± 0.2	5.7	0.1	7.7 ± 0.3	11.0 ± 1
Trays (MAP)	1 st harvest	NI AOX		19.2 ± 0.0	16.9 ± 0.2	16.0 ± 0.5	16.3 ± 0.6	15.6 ± 0.1		2.0 ± 0.1	3.1 ± 0.1	4.1 ± 0.2	4.9 ± 0.8	5.9 ± 0.0
		CPA AOX		18.6 ± 0.3	17.7 ± 0.3	17.5 ± 0.5	15.3 ± 0.8	17.2 ± 0.3		1.5 ± 0.1	2.7 ± 0.1	3.2 ± 0.4	5.5 ± 0.5	5.1 ± 0.5
		NI H2O		19.2 ± 0.0	17.3 ± 0.2	16.5 ± 0.3	15.9 ± 0.6	15.0 ± 0.6		2.0 ± 0.1	3.8 ± 0.3	4.4 ± 0.3	5.3 ± 1.0	5.7 ± 0.3
	2 nd harvest	CPA H2O		18.9 ± 0.1	16.3 ± 0.3	16.0 ± 0.4	14.8 ± 0.5	14.8 ± 0.8		2.3 ± 0.1	4.2 ± 0.3	4.9 ± 0.3	6.4 ± 0.3	6.0 ± 0.6
		NI AOX		17.4 ± 0.6	16.3 ± 0.7	13.6 ± 0.5	14 ± 1	15.4 ± 0.8		2.4 ± 0.3	3.5 ± 0.5	6.1 ± 0.3	6.2 ± 0.9	5.6 ± 0.6
		CPA AOX		17.3 ± 0.2	16.0 ± 0.2	12.5 ± 0.8	13.4 ± 0.1	14.5 ± 0.1		2.4 ± 0.2	3.7 ± 0.1	7.1 ± 0.6	7.5 ± 0.1	6.8 ± 0.1
2 nd harvest	NI H2O		17.9 ± 0.4	17.1 ± 1	15.4 ± 0.6	14.7 ± 0.8	16 ± 2		2.9 ± 0.4	3.6 ± 0.8	5.1 ± 0.4	5.1 ± 0.8	5 ± 2	
	CPA H2O		17.8 ± 0.1	16.7 ± 0.0	14.2 ± 0.8	15.8 ± 0.6	16.1 ± 0.4		2.9 ± 0.1	4.0 ± 0.1	5.9 ± 0.7	5.4 ± 0.7	5.0 ± 0.6	

Initial concentrations were 21 % O₂ and 0 % CO₂. NI: non-inoculated, CPA: CPA-7-inoculated, AOX: AOX-treated, H2O: AOX-untreated. Results are expressed as mean ± standard deviation.

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