

This document is a postprint version of an article published in Postharvest Biology and Technology © Elsevier after peer review. To access the final edited and published work see <u>https://doi.org/10.1016/j.postharvbio.2017.12.013</u> Pseudomonas graminis strain CPA-7 differentially modulates the
 oxidative response of fresh-cut 'Golden delicious' apple
 depending on the storage conditions.

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13 Abstract

Oxidative response of fresh-cut antioxidant-treated 'Golden delicious' apples during 14 15 chilling storage was differentially modified by the biopreservative bacterium CPA-7 depending on the conditions; passive modified atmosphere packaging (MAP) or 16 aerobic, the latter being more favorable for the antagonist growth. Inoculation with 17 CPA-7 had no influence on fruit quality parameters in any of the conditions tested. 18 19 During the first 24 h both in air or in MAP, ascorbate peroxidase (APX) activity triplicated the initial level in response to CPA-7 inoculation, reaching up to 4-fold the 20 activity of non-inoculated fruit (control). From 24 h of storage in MAP, polyphenol 21 22 oxidase (PPO) activity was sharply enhanced in response to CPA-7 attaining values up 23 to 6-fold higher than that of the control at the end of storage. Contrastingly, from 1 d to 3 d of storage in MAP CPA-7 suppressed catalase (CAT) activity by 1.5-fold. 24 Subsequently, after 3 d in such conditions, superoxide dismutase (SOD) and PPO 25 activities were almost duplicated in the presence of CPA-7 if compared to the control. 26 27 On the other hand, when stored in air, polyphenol oxidase (POX) showed a biphasic 28 induction in response to CPA-7 after 3 d and 6 d of incubation. At day 6, the activity of 29 this enzyme reached values 8-fold higher in response to CPA-7 than the observed for 30 the control regardless of storage conditions, yet it was not paired to an increase of browning. Inoculation with CPA-7 led to the slowdown of the decline of antioxidant 31 capacity in air, which contrasted with the response upon MAP conditions. These results 32 suggest that CPA-7 may trigger the activation of the fruit defense-response thereby 33 34 mitigating its oxidative damage. Such activation may play a role as a putative biocontrol 35 mechanism against foodborne pathogen infections.

36 **1. Introduction**

Control of foodborne diseases through bio-friendly strategies circumventing the usage 37 38 of chemicals that produce potentially harmful residues in fresh-cut fruit is an important area of research. The use of antagonistic epiphytic microorganisms is a method that 39 fulfills this characteristic which has been widely explored over the past (Belak and 40 41 Maraz, 2015; Leverentz et al., 2006). In addition to the maintenance of microbiological 42 quality, the growing demand of fresh processed fruit and vegetables has challenged 43 food industry to also guarantee the physicochemical acceptance of these products in spite of the metabolic changes inherent to this type of commodities, mainly associated 44 45 to abiotic stress caused by both refrigerated storage and processing. In this sense, cold storage in regular controlled atmosphere conditions is known to be stressful to 46 harvested fruit inducing the accumulation of ROS (Chiriboga et al., 2013). Thus said, 47 scarce information is available about the oxidative changes of fresh-cut fruit at the 48 enzymatic or metabolic level (Larrigaudiere et al., 2008). 49

50 Besides, cutting of fruit flesh compromise compartmentalization in nearby cells allowing 51 phenolic compounds located in vacuoles to get in contact with polyphenol oxidase 52 (PPO), located in plastids, triggering the reaction known as enzymatic browning 53 (Holderbaum et al., 2010). Other antioxidant enzymes such as peroxidase (POX) has 54 also been linked to polyphenol-associated browning through a coordinated mechanism

55 involving the accumulation of its substrate, hydrogen peroxide (H₂O₂), stimulated by the PPO-mediated generation of quinones (Jiang and Miles, 1993). Concomitantly, H₂O₂ is 56 57 highly oxidant and may damage membrane and other cellular components through several mechanisms. Thus, in both fresh-cut and intact fruit, its removal is rapidly 58 activated in order to protect plant cells from these damages. H₂O₂- scavenging 59 mechanisms include enzymatic reactions which involve superoxide dismutase (SOD), 60 catalase (CAT) and ascorbate peroxidase (APX) found in the cytosol chloroplasts and 61 62 mitochondria of higher plants (Hung et al., 2005) as well as non-enzymatic antioxidants including phenolic compounds (vitamin E, flavonoids, phenolic acids and other 63 phenols); nitrogen compounds (alkaloids, amino acids and amines), carotenoids and 64 chlorophyll derivatives (Kumar, 2012). 65

The control of enzymatic browning in the fresh- cut produce industry is currently mainly based on the modulation of PPO activity through antioxidant formulations that commonly contain reducing compounds such as ascorbic acid and its derivatives, cysteine and glutathione (Rupasinghe et al., 2005). These agents are hypothesized to control PPO activity either by reducing quinones to the native diphenols or by reacting irreversibly with o- quinones to form stable colourless products (Nicolas et al., 1994; He and Luo, 2007).

73 In addition, plant oxidative metabolism is also activated in response to biotic stress as a 74 part of induced defense mechanisms (Pieterse et al., 2000, 2014). Under the light of 75 the increasing knowledge about plant-microorganism interactions, induced resistance 76 has arisen as a method for controlling postharvest diseases. The investigation of these processes has revealed that biocontrol agents (BCA) activate defense-related enzymes 77 such as phenylalanine ammonia lyase (PAL), POX, PPO and chalcone synthase 78 79 among others, therefore contributing to better antioxidant defenses and tolerance to pathogens attack (Chen et al., 2000; Jain et al., 2012; Alkan and Fortes, 2015). 80

Pseudomonas graminis strain CPA-7 is an aerobic epiphyte bacillus which has been 81 used as BCA against foodborne pathogens such as Escherichia coli O157:H7, 82 83 Salmonella enterica and Listeria monocytogenes on fresh cut apples, melons and peaches (Abadias et al., 2014; Alegre et al., 2013a, 2013b; Collazo et al., 2017). 84 Although attempts to elucidate the mechanisms underlying its antagonistic activity have 85 been carried out, they still remain poorly understood (Collazo et al., 2017). Thus, we 86 87 hypothesized that a putative mode of action for CPA-7 could be the activation of defense mechanisms in fresh-cut fruit leading to a better control of foodborne 88 pathogens populations. Accordingly, the oxidative metabolism of inoculated fresh-cut 89 apples treated with an antibrowning agent both upon aerobic conditions, suitable for 90 91 CPA-7 growth, and in semi-commercial conditions (modified atmosphere packaging, MAP) were investigated. 92

93 2. Materials and methods

94 2.1 Fruit processing

Apples (*Malus domestica* L. cv. Golden Delicious) were obtained from local packinghouses (Lleida, Catalonia, Spain). Prior to experimental assays, the fruit were washed with running tap water, surface disinfected with 700 mL L⁻¹ ethanol, peeled with an electric fruit peeler and cut into eight wedges with a handheld corer/slicer. Wedges were kept in chilled (5°C) chlorinated tap water (pH 6) until they were subsequently treated or stored.

101 2.2 Antagonist culture conditions

For the antagonist inoculum preparation, a single colony of *Pseudomonas graminis* strain CPA-7 grown in tryptone soy agar plates (TSA, Biokar, Beauveais, France) during 48 h at 30 °C, was inoculated into 50 mL of tryptone soy broth (TSB, Biokar, Beauveais, France) and incubated in aerobic conditions for 20-24 h in agitation (15.71 rad s⁻¹) at 25 °C. Bacterial cells were harvested by centrifugation at 9800 × g for 10 min at 10 °C and re-suspended in sterile distilled water. The concentration of the suspension was checked by plate count of appropriate 10-fold dilutions in saline peptone (8.5 g L⁻¹ NaCl, 1 g L⁻¹ peptone) onto TSA plates after incubation at 30 °C for 48 h.

111 2.3 Inoculation, sampling and microbiological analysis

Suspensions of CPA-7 at 10⁷ CFU mL⁻¹ were prepared in a 60 g L⁻¹ NatureSeal® AS1 112 113 AgriCoat Ltd., Great Shefford, UK) antioxidant aqueous cold solution (4 °C), as 114 recommended by the manufacturer. Non-inoculated antioxidant solution was included as control. Fruit wedges were dipped-inoculated at a ratio 1:2 (weight of product: 115 116 volume of suspension) for 2 m in agitation (15.71 rad s⁻¹) in tabletop orbital shaker 117 Unimax 1010 (Heidolph, Germany). After drainage of the excess of water, 10 g of apple wedges were used to determine initial microbiological quality in triplicate and three 118 119 samples of about 70 g were frozen with liquid nitrogen, grinded in a commercial grinder (Minimoka 6R-020, Coffeemotion, Lleida, Spain) and stored at -80 °C until biochemical 120 121 analysis. The rest of the treated fresh-cut fruit was packaged (120 g per replicate) in 400 mL polyethylene terephthalate ShelfMaster™ Pronto™ trays (PlusPack, Denmark). 122 Each tray was considered as a replicate and three replicates per treatment and 123 sampling time were used. Trays were sealed with a peelable plastic film (polyester anti-124 125 fog film (OALF) of 14 µm of thickness + oriented polypropylene (OPP) of 20 µm of thickness) with a line of holes of 60 - 80 µm each and 75 mm separation spacing 126 127 between them, to achieve passive modified atmosphere. In a parallel set of trays the 128 film was macro perforated (nine extra holes per tray) in order to attain aerobic 129 conditions. Trays were stored at 5 °C in darkness and biochemical and microbiological analyses were performed at 0, 1, 3 and 6 d post-inoculation. Population dynamics of 130 CPA-7 as well as total mesofils of inoculated and non-inoculated samples were 131 analyzed at each sampling time. For this, 10 g of apple from each tray was 132 133 homogenized in 90 mL of buffered peptone water (APT, Biokar, Beauvais, France) and tested by viable cells count onto TSA and plate count agar plates (PCA, Biokar, 134

Beauveais, France) incubated at 30 °C (for CPA-7) and at 25 °C (for mesofils) for 48 h
and 72 h, respectively.

137 2.4 Fruit physicochemical quality parameters

Firmness, color and pH of fresh-cut wedges were determined initially and at each 138 sampling time as described elsewhere (Alegre et al., 2013a). Low values of CIE 139 140 coordinate L* and high values of a* were considered as indicators of surface browning intensity (Sapers and Douglas, 1987). Soluble solids and titratable acidity were 141 measured in two samples per tray initially and at the end of storage. Soluble solids 142 concentration at 20 °C was expressed as mass fraction of sugars relative to the fruit 143 144 (%). Acidity was measured in 10 mL of pulp and was expressed as malic acid content $(g L^{-1} juice).$ 145

146 2.5 Biochemical analyses

147 2.5.1 Total antioxidant capacity (TAC) and total phenolic content (TPC)

Extracts for total phenolic content (TPC) and total antioxidant capacity (TAC) determination were prepared by mixing 3 g of frozen fresh pulp with 10 mL of a solution containing 19.7 mol L⁻¹ methanol, 0.05 mol L⁻¹ HCl. The mixture was held in agitation (20.94 rad s⁻¹) for 2 h and centrifuged at 24 000 x g at 4 °C. TPC was quantified by measuring the optical density (OD) at 765 nm in a spectrophotometer (EONC, Biotek Instruments, Highland Park, VT, USA) after the reaction of 0.05 mL of each extract with 0.25 mL of Folin-Ciocalteu reagent and 0.5 mL of 1.9 mol L⁻¹ Na₂CO₄.

TAC was determined by measuring OD at 593 nm of the above mentioned extracts
following the Ferric Reducing Antioxidant Power (FRAP) protocol (Giné-Bordonaba and
Terry, 2016).

158 2.5.2 Hydrogen peroxide production and malondialdehyde (MDA) content

Malondialdehyde (MDA) was quantified in 0.5 g of frozen fresh pulp as a marker of lipid
 peroxidation using the thiobarbituric acid reactive substrates (TBARS) assay described

by Martínez-Solano et al. (2005). The OD of the supernatants after reaction was
 measured at 532 nm and substracted to the inespecific absorption read at 600 nm.
 Quantification of TBARS was calculated by its extinction coefficient of 155 mmol L⁻¹ cm
 ⁻¹.

To determine hydrogen peroxide levels, 5 g of frozen fresh pulp was homogenized in 165 7.5 mL of 0.5 mol L⁻¹ trichloroacetic acid, filtered through two layers of Miracloth (Textil 166 Planas Oliverassa, Manresa, Spain) and centrifuged at 20 000 x g for 15 min at 4 °C. 167 H₂O₂ content was determined using the aqueous peroxide colorimetric assay, 168 169 PeroxiDetect[™] Kit (Sigma-Aldrich, St Louis, MO, USA) following the manufacturer's instructions. This procedure is based on the measurement of the color change at 560 170 nm of adduct formed by Fe^{3+} ion and xylenol orange (XO, PubChem CID 73041), 171 during the oxidation by peroxide of Fe²⁺ to Fe³⁺ ions at acidic pH. The molar extinction 172 coefficient of the XO-Fe³⁺ colored adduct in aqueous solution at 560 nm is 15 000. 173

174 2.5.3 Activities of enzymes involved in antioxidant metabolism

175 2.5.2.1 Peroxidase (POX) y Polyphenol oxidase (PPO)

Total peroxidase (POX, EC 1.11.1.7) and polyphenol oxidase (PPO, EC 1.14.18.1) 176 were extracted from 10 g of frozen fresh pulp as described by Giné-Bordonaba et al. 177 178 (2017). POX activity was measured as the optical density at 470 nm following the reaction of the extract with 10 mmol L^{-1} guaiacol and 10 mmol L^{-1} H₂O₂, according to 179 the method described by Lurie et al. (1997). PPO activity was determined at 400 nm in 180 a reaction mixture containing 0.1 mol L⁻¹ potassium phosphate buffer (pH 6) and 65 181 mmol L⁻¹ pyrocatechol together with the correspondent enzyme extract as described by 182 183 Vilaplana et al. (2006).

184 2.5.2.2 Superoxide dismutase (SOD) and catalase (CAT)

For the extraction of superoxide dismutase (SOD, EC 1.15.1.1) and catalase (CAT, EC 1.11.1.6), 5 g of frozen fresh pulp were homogenized in 15 mL 0.1 mol L^{-1} potassium phosphate buffer (pH 7.8), 2 mmol L^{-1} dithiothreitol (DTT), 50 g L^{-1}

polyvinylpolypyrrolidone (PVPP), 0.1 mmol L⁻¹ ethylene diamino tetracetic acid (EDTA) and 1.25 mmol L⁻¹ polyethylene glycol. The homogenized was filtered, centrifuged and purified as described by Giné-Bordonaba et al. (2017). In turn, SOD activity was determined by measuring its ability to inhibit the photochemical reduction of nitrobluetetrazolium (NBT) at 560 nm following the method of Gianopolitis and Ries (1977).

194 2.5.2.3 Ascorbate peroxidase (APX)

For ascorbate peroxidase (APX; EC 1.11.1.11) extraction, 10 g of frozen fresh pulp 195 was homogenized with 30 mL of 0.1 mol L⁻¹ base phosphate buffer (pH 7.5) containing 196 0.8 mmol L⁻¹ HCI, 1 mmol L⁻¹ EDTA, filtered through two layers of Miracloth and 197 centrifuged at 10 000 x g for 15 min at 4 °C. APX activity was determined at 290 nm 198 199 during 10 min by monitoring the H₂O₂-dependent decomposition of ascorbate in a mixture containing twenty microliters of the recovered supernatant and 280 µL of a 200 reaction solution (0.22 mmol L⁻¹ ascorbic acid, 1 mmol L⁻¹ EDTA, 1 mmol L⁻¹ H₂O₂) 201 202 (Nakano and Asada, 1981).

203 2.5.2.4 Protein content

Protein content of all extracts was determined by the Bradford method at 595 nm using a protein assay kit (Bio-Rad, München, Germany). Protein reagent was mixed with the correspondent phosphate buffer (pH 6, 7 or 7.8) used for each enzyme extraction in a ratio 1:3.6. Bovine serum albumin (BSA, Sigma-Aldrich, St. Louis, MO, USA) was used as standard.

209 2.6 Expression of results and statistical analysis

Enzyme activity was expressed in units of enzyme activity per milligram of protein (U mg⁻¹ protein). For SOD, 1 U represents the amount of enzyme required to inhibit NBT reduction by 50 % of initial amount. For the rest of the analyzed enzymes, 1 U represents the quantity of enzyme responsible for a change in 1 absorbance unit per minute. Microbiological data was estimated as microbial counts per gram of fruit (CFU

g⁻¹ fruit). Non-enzymatic antioxidant activities were expressed as g of the measured 215 analyte (i.e. Gallic acid (GAE) or Fe³⁺) per kilogram of fruit. Oxidative stress markers 216 (H₂O₂ and MDA) were expressed as µmol kg⁻¹ fruit. The data were analyzed using the 217 general linear model procedure to determine differences among treatments and 218 219 interaction effects using the statistical software JMP (version 11 SAS Institute Inc., NC, USA). All data were verified for normal distribution and homoscedasticity of residues 220 221 and accordingly, enzymatic activities and microbiological data were transformed to log₁₀ of U mg⁻¹ protein or CFU g⁻¹, respectively. Transformed results were schematically 222 223 represented as back-transformed means ± standard error of the mean. Transformed and non-transformed means were compared by analysis of variance (ANOVA) and 224 separated by Tukey's test (P < 0.05). 225

226 **3 Results**

227 3.1 Microbial dynamics

Initial populations of CPA-7 on the fruit were 5.53 \pm 0.03 log₁₀ (Fig. 1A). Low oxygen 228 229 concentration associated to modified atmosphere packaging (MAP) slowed down CPA-7 growth if compared to the samples stored in air, delaying the start of the exponential 230 231 phase from 1 d to 3 d post-inoculation. On day 6, samples stored in modified atmosphere packaging showed a 0.5 log₁₀ reduction of CPA-7 growth in respect of the 232 samples stored in aerobic conditions. Mesofils populations were initially the same (1.7 233 \pm 0.2 log₁₀) in all samples (Fig. 1B). However differences in growth (1 log₁₀ CFU g⁻¹ 234 fruit) among aerobic and modified atmosphere-stored samples became evident on day 235 236 6.

237 3.2 Fruit physicochemical quality parameters

No significant differences were observed among values of L* (lightness) and a* (redness) in the presence of CPA-7 (Table 1). Likewise, no differences were observed in the content of soluble solids or pH in response to the antagonist under any of the tested conditions throughout storage. Firmness was better maintained throughout the studied period in the presence of CPA-7 if compared to non-inoculated fruit, in which itwas reduced at the end of storage.

244 3.3 Biochemical analyses

245 3.1.1 Total antioxidant capacity (TAC) and total phenolic content (TPC)

At the beginning of the experiment total antioxidant capacity (TAC), measured by the content of Fe³⁺, was 13.6 \pm 0.2 g kg⁻¹ fruit in all samples and gradually decreased during storage (Fig. 2A). However, the drop of TAC was significantly slower in presence of CPA-7 in aerobic conditions where it remained at 12.3 \pm 0.4 g kg⁻¹ fruit until day 6. No differences were observed among MAP-stored samples in the first three days of the experiment. However, on day 6 in MAP conditions, it was recorded a reduction by 0.8-fold in response to CPA-7 if compared to the control.

Total phenolic content (TPC) was initially 5.1 ± 0.1 g kg⁻¹ fruit in all samples regardless of the presence of the antagonist (Fig. 2B). On day 3 and for both storage conditions, the inoculation with CPA-7 was associated to a significantly higher amount of TPC (4.5 ± 0.1 g kg⁻¹ fruit), than in the control samples (4.2 ± 0.04 g kg⁻¹ fruit in air and 3.7 ± 0.04 g kg⁻¹ fruit in MAP). In agreement to TAC, on day 6, TPC was reduced in MAP-stored samples in response to CPA-7 (4.1 ± 0.1 g kg⁻¹ fruit) if compared to the control (4.9 ± 0.1 g kg⁻¹ fruit).

260 3.1.2 Hydrogen peroxide production and malondialdehyde content (MDA)

No sign of peroxidation of the membranes could be estimated from the measurement of the content of the marker malondialdehyde ($0.019 \pm 0.04 \mu$ mol kg⁻¹ fruit) under any of the conditions tested during the analyzed period (data not shown). When combined to the antioxidant, CPA-7 was able to erase the oxidative burst or at least that referred to H₂O₂ accumulation in the fruit stored in air (Fig. 3). Contrastingly, in the samples stored in MAP H₂O₂ drastically increased in the first 24 h reaching up to 1.4-fold higher in response to CPA-7 than in the non-inoculated control. In general, metabolization of this molecule was significantly slower in the MAP-stored samples in the presence of CPA-7 than in the rest of the samples. In this sense, values remained 13-fold higher if compared to the control at the end of storage. Contrastingly, in inoculated samples stored in air, the levels of H_2O_2 remained close to zero throughout the whole experiment.

273 3.1.3 Activities of enzymes involved in antioxidant metabolism

Changes in the enzymatic antioxidant potential of 'Golden delicious' fresh-cut apple 274 differed throughout storage according to the inoculation with the antagonist and/or to 275 the storage conditions. SOD activity decreased from $5554 \pm 42 \text{ U mg}^{-1}$ protein at initial 276 277 time to 4863 ± 111 U mg⁻¹ protein after 24 h in all samples, except for the CPA-7inoculated MAP-stored sample, in which it remained stable up to day 3. This contrasted 278 279 with the drop (0.7-fold lower than the CPA-7-inoculated sample) observed for the 280 correspondent control (Fig 4A). In contrast, on day 6, SOD activity was oppositely modulated by CPA-7 depending on the storage conditions; it was 0.7-fold lower than 281 282 the control when stored in MAP and 0.6-fold higher than the control when stored in air.

Catalase activity immediately after processing was similar in all samples ($237 \pm 17 \text{ U}$ mg⁻¹ protein) regardless of any of the factors assayed (Fig. 4B). However, it differentially changed in response to the antagonist in different storage conditions. It increased by 0.7-fold compared to the initial value when stored in air, reaching the same level than the control after 24 h. In contrast, CAT activity in inoculated samples was inhibited in response to the antagonist when stored in MAP (by more than 0.6-fold if compared to the control) during the first 3 d of storage.

Initially, PPO activity was $243 \pm 7 \text{ U mg}^{-1}$ protein which was reduced by 0.5-fold in the first 24 h (Fig. 4C). Thereafter, it drastically increased in response to CPA-7 in MAP conditions reaching almost 8-fold the activity showed by the correspondent noninoculated fruit. However, when stored in air, CPA-7 had no significant effect in PPO activity throughout the analyzed period. Activity of APX was initially around $348 \pm 43 \text{ U mg}^{-1}$ protein in all samples (Fig 4D). During the first 24 h post-inoculation, APX activity was induced up to 3-fold, if compared to the initial values, in response to the antagonist when stored in air. At this sampling time, peaks of APX activity in response to CPA-7 triplicated the activities recorded for the controls, regardless of the storage conditions. An enhanced APX activity associated to the antagonist was maintained until day 3 (1.8 fold higher than the control) in air-stored samples.

302 Immediately after inoculation, POX activity was inhibited in response to CPA-7 (by 0.6-303 fold if compared with the control) (Fig. 4E). Thereafter, this enzyme showed a biphasic activation in response to CPA-7 in aerobic conditions reaching values 1.5-fold and 1.7-304 305 fold higher than those observed for the control after 1 and 6 d post-inoculation, 306 respectively. In contrast, POX activity was reduced in all MAP-stored samples during 307 the first 24 h post-inoculation, but it attained significantly lower values (98 ± 10 U mg⁻¹ protein) in the presence of the antagonist than in the control ($84 \pm 5 \text{ U mg}^{-1}$ protein). As 308 for air-stored samples on day six, POX activity was enhanced in the CPA-7-inoculated 309 MAP-stored samples, reaching values eight fold higher than the control. 310

311 **4 Discussion**

Processing of fresh produce implies the removal or damage of most of the constitutive 312 313 barriers of plants and generally activates inducible defense mechanisms to protect 314 themselves against pathogens (Reviewed by Hodges and Toivonen, 2008). 315 Pseudomonads that have been used as biocontrol agents (BCA) have shown to 316 enhance plant response including cell-wall straightening, oxidative burst and 317 expression of defense related genes that are usually activated by pathogens (Van der 318 Ent et al., 2009). This phenomenon 'primes' plant metabolism allowing it to react faster 319 and stronger upon a subsequent pathogen attack (Pieterse et al., 2001). One of the 320 enzymes associated to resistance responses is peroxidase (POX) which mediates the 321 last steps of the biosynthesis of lignin and hydrogen peroxide (H₂O₂) (Hiraga, 2001).

322 POX also catalyzes single-electron oxidation of phenolic compounds in the presence of hydrogen peroxide. This latter molecule is one of the first molecules generated in plant 323 324 reaction to wounding and pathogen infection and has been postulated to play a dual 325 role as signaling factor to induce defense genes related to wound-stress and 326 hypersensitive response and it is also involved in fine tune regulation of oxidative stress caused by reactive oxygen species (ROS) both locally and systemically (Hung et 327 328 al., 2005; Sewelam et al., 2014; Baxter et al., 2014). However, little is known about the oxidative response in fresh-cut fruit associated to BCA. 329

330 The results obtained in the present study showed that oxidative response of 'Golden 331 delicious' fresh-cut apples was differentially modified in the presence of CPA-7 332 depending on the storage conditions. For instance, immediately after processing and 333 treatment with the antioxidant, H₂O₂ quickly began to accumulate in all samples to up to 334 the first 24 h in the MAP-stored samples and up to 3 d in the air-stored control. Similar 335 H₂O₂ accumulation pattern has been previously observed in non-inoculated fresh-cut 'Fuji' apples treated with ascorbic acid and stored in aerobic conditions (Larrigaudiere 336 et al., 2008). However, in the presence of CPA-7, the oxidative burst was practically 337 erased in the air-stored samples, and H₂O₂ levels were kept close to zero during the 338 339 whole experiment. Activation of ROS-scavenging enzymes correlated with the initial oxidative burst associated with processing, thus the levels of SOD activity were initially 340 elevated in all samples. Subsequently, SOD activity decreased in all samples and APX 341 was sharply and transiently induced by CPA-7 as a first line of defense against 342 elevated H₂O₂ production in both MAP and air-stored samples after the first 24 h. On 343 the other hand, when stored in air CPA-7 was also associated to an early increase of 344 CAT and POX activities which could contribute to quickly eliminate H₂O₂ as soon as it 345 346 was produced. The decline of SOD activity 24 h after processing was also observed by 347 Larrigaudiere et al. (2008) in antioxidant-treated Fuji apples.

348 In the conditions assayed herein, as time passed, stress associated to mechanical 349 damage was added to that caused by low-oxygen atmosphere as in MAP-stored 350 samples, hence leading to changes in the modulation of some scavenging enzymes aiming to maintain redox homeostasis. In this way, from day 1 to 3, SOD activity 351 peaked in the CPA-7-inoculated MAP-stored samples and subsequently, PPO and 352 POX were sharply induced until the end of storage counteracting the inhibition of CAT. 353 354 In contrast, in the air-stored samples the activation of SOD due to CPA-7 lasted until day 6 and was paired to the induction of POX. Thus, stressful conditions were 355 associated not only to increased ROS-scavenging enzymes activities but to changes in 356 their balance. It has been previously observed that reduced CAT activity is sometimes 357 358 compensated by induction of APX and glutathione peroxidase (GPX) (Apel and Hirt, 2004). Similarly, Wang et al. (2014) observed that the treatment of loquat fruit with the 359 antagonist Bacillus cereus AR-156 induced SOD activity while reduced the activities of 360 CAT and APX enhancing the plant response to infection by *Colletotricum acutatum*. 361

362 The modulation of oxidative metabolism in fruit and vegetable in response to 363 Pseudomonas spp. as observed in the present study, has been previously reported and correlated to the induction of plant resistance to several plant pathogens 364 365 (Sangeetha et al., 2010). For instance, combinations of a non-fluorescent Pseudomonas spp. strain NFP6, P. fluorescens and B. subtilis induced up to 4-fold 366 PAL, POX and PPO activities and increased by 3.6-fold the phenolic content in banana 367 fruits challenged with crown rot pathogens, 5 d after treatment with the antagonists 368 (Sangeetha et al., 2010). Likewise, Pseudomonas corrugate strain 13 and 369 Pseudomonas aerofasciens strain 63-28 significantly induced PAL, POX and PPO in 370 cucumber 2-5 d following treatment and elicited the defense response to subsequent 371 372 infection with Phytium aphanidermatum (Chen et al., 2000). Similarly, mixtures of fluorescent rhizosphere and endophytic Pseudomonads induced these enzymes when 373 374 sprayed on banana plants which subsequently showed enhanced systemic resistance

against bunchy top virus (Harish et al., 2009). Similar response regarding POX, PPO and SOD activities was previously observed for other biocontrol agents such as *Bacillus amyloliquefaciens* LJ02 after being sprayed on three leave-stage seedlings (Li et al., 2015). In general accordance with our study, they observed a 2-fold increase of POX activity on day 6 as well as an enhanced SOD activity in aerobic conditions until day 5 priming plants for resistance to powdery mildew caused by the fungus *Spaheroteca fugilinea.*

382 Changes in PPO activity in response to CPA-7 has been previously investigated in 383 fresh-cut 'Piel de sapo' melon stored in air and in MAP conditions during 8 d of storage 384 at 5 °C (Plaza et al., 2016). In contrast to our results, no variation was recorded in 385 response to CPA-7 for MAP-stored samples. Furthermore, although an induction of this 386 enzyme associated to the antagonist was observed by Plaza et al. (2016) after 8 d of 387 storage in aerobic conditions compared to the initial activity, this increase did not differ 388 from the control. Differences in sampling days between that study and ours could suggest that PPO could peak after 6 d of MAP-storage and drop before day 8 and in 389 390 the same way, the increase of PPO activity in aerobic samples could be produced after 6 d of storage. Alternatively, discrepancies between both studies could be associated 391 392 to the differential behavior of the bacterium in relation to different fruit matrices.

The question still remains on whether the increase of ROS-scavenging enzymes 393 394 activities such as superoxide dismutase, catalase and peroxidase is only a plant response or is also a strategy deployed by the antagonist. This would confer this 395 396 bacterium with the ability to tolerate highly oxidant conditions generated after 397 processing of fresh produce and could act as a pivotal mechanism of action to outcompete foodborne pathogens or spoilage microorganisms. Differences in the 398 399 modulation pattern of the oxidative metabolism throughout storage observed in the present study did not correspond to great changes in CPA-7 populations as a function 400 of storage conditions. The tracking of population dynamics showed that CPA-7 was 401

able to tolerate the changes produced in the fruit oxidative metabolism during the 402 403 assayed period. In this sense, CPA-7 showed similar growth upon refrigerated 404 conditions both in air and in MAP with a slight inhibition corresponding to lower oxygen 405 availability after 6 d of incubation. Previous semi-commercial trials performed with 406 fresh-cut apples in conditions resembling those assayed herein showed similar CPA-7 407 population dynamics and revealed an inhibitory or bacteriostatic effect of this 408 antagonist on *L. monocytogenes* depending on the temperature of storage rather than on MAP or air storage conditions (Alegre et al., 2013a). From this perspective, high 409 resistance to ROS of the antagonistics yeasts (Cryptococcus laurentii LS-28 and 410 Rhodotorula glutinis LS-11) inoculated in wounded apple and peach fruits has been 411 412 associated with enhanced antagonistic activity against Botrytis cinerea and Penicilium 413 expansum (Castoria et al., 2003; Zhang et al., 2017).

414 On the other hand, concomitant with enzymatic changes, increased non-enzymatic 415 antioxidant activity provides host tissue with additional defense tools to counteract ROS 416 production induced by both biotic and abiotic stress (Hung et al., 2005). The initial 417 accumulation of phenolic compounds, as observed in our trials regardless of the presence of the antagonist, has been studied for fresh-cut produce (Saxena et al., 418 419 2009). It could be related to the high production of H₂O₂ upon mechanical injury through an enhanced respiration or due to the activation of several metabolic pathways 420 421 such as the hexose -monophosphate shunt pathway, the acetate pathway and/or the release of bound phenolic compounds by hydrolytic enzymes (Harish et al., 2009). 422 However, the subsequent reduction of total phenolic content was slowed down in the 423 presence of CPA-7 when stored in aerobic conditions at least until day 3 which could 424 also account for the maintenance of beneficial nutritional properties of the fruit. 425 426 Contrastingly, in MAP conditions, CPA-7 was associated to a peak in the production of phenolic compounds on day 3 with a concomitant increase of PPO activity. 427 428 Interestingly, under such storage conditions, the drastic accumulation of this enzyme at the end of the storage period was correlated to a significant drop of its substrates, yet itdid not correlate with a significant increase of browning.

Regarding the effect of the antagonist on fruit quality, CPA-7 had a positive effect on 431 432 the maintenance of physical parameters such as firmness throughout storage which 433 contrasted with control samples. This result agreed with the maintenance of the cellular membranes integrity in the presence of CPA-7 as estimated chemically from the 434 measure of MDA. No significant effect on pH or soluble solids was observed due to the 435 436 antagonist during the studied period. The combination of the antagonist with the 437 antioxidant agent resulted in the maintenance of luminosity and redness but similar to the results observed for the control samples at all times of analysis. Similarly, in CPA-438 7-inoculated fresh-cut 'Golden delicious' apples treated with 60 g L⁻¹ Natureseal® 439 (AS1), as in the present study, L* and a* Hunter values were maintained during 14 d of 440 441 storage at 5 °C (Alegre et al., 2013a). The same antioxidant treatment has been 442 previously shown to efficiently maintain color properties (with a* values from -2.55 to -3.14) and to increase the firmness of fresh-cut of apple wedges from ten cultivars after 443 storage for 5 d at 2 °C – 4 °C (Roble et al., 2009). In contrast, when using a different 444 antioxidant treatment (5 g L⁻¹ calcium chloride), a decrease of luminosity of CPA-7-445 446 treated fresh-cut melon after 8 d of storage at 5 °C was observed by Plaza et al. (2016). 447

448 **5** Conclusions

The results showed herein indicated that CPA-7 was associated to a differential modulation of the oxidative response of antioxidant-treated fresh-cut apple during chilling storage in MAP or aerobic conditions. CPA-7-inoculated fruit showed a higher antioxidant capacity, a higher accumulation of phenolic compounds and an enhanced activity of key antioxidant enzymes (POX, SOD and APX) which were related to the plant response to either mechanical injury or microbial invasion, at certain moments of storage. The antagonist ability to tolerate or to mediate these changes and to grow 456 upon such conditions could imply high resistance to oxidative stress which would, in turn, provide CPA-7 with an adaptive advantage to outcompete putative pathogen 457 458 infection. The differential temporal enhancement of the analyzed enzymes depending on the storage conditions suggests a synergistic activation of different pathways in 459 response to the several factors associated to production and commercialization of 460 fresh-cut fruit such as mechanical injury, low-temperature, antioxidant-treatment and 461 462 low-oxygen conditions. Further studies including transductional analysis of the 463 accumulation patterns of different antioxidant isoenzymes, which may be activated by the plant and/or the antagonist upon challenge with foodborne pathogens would 464 improve the understanding of such mechanisms. 465

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 stress on biocontrol activity of *Cryptococcus laurentii* against blue mold on peach
 fruit. Front. Microbiol. 8, 1–10. doi:https://doi.org/10.3389/fmicb.2017.00151

- 620 Table 1. Quality parameters of fresh-cut antioxidant-treated 'Golden delicious' apple
- 621 wedges inoculated with *P. graminis* CPA-7 (CPA) or non-inoculated (NI) during storage
- 622 in aerobic conditions (AIR) or in modified atmosphere packaging (MAP).

Parameter	Treatment -	Time (days) at 5 ºC										
		0		1		3			6			
Lightness (L*)	AIR NI	78.1±0.5	a B	80.9 ± 0.2	a A	79.7 ±	0.4	а	AB	78.5±0.6	abc	В
	AIR CPA	78.5 ± 0.9	a B	77.5 ± 0.4	bc B	78.8 ±	0.6	ab	В	78.5±0.7	abc	В
	MAP NI			78.2 ± 0.6	bc A	79.8 ±	0.5	а	Α	79.1±0.5	ab	А
	MAP CPA			80.1 ± 0.3	ab A	79.0 ±	0.5	ab	Α	79.7±0.4	а	Α
Redness (a*)	AIR NI	-6.6±0.4	b B	-6.4 ± 0.4	a A	-6.4 ±	0.3	cd	А	-5.8±0.5	bc	Α
	AIR CPA	-6.1 ± 0.6	ab B	-6.0 ± 0.3	bc B	-6.5 ±	0.2	ab	В	-6.2±0.4	bc	В
	MAP NI			-6.4 ± 0.5	bc AB	-7.7 ±	0.1	d	В	-6.4±0.1	bc	А
	MAP CPA			-7.2 ± 0.5	ab A	-6.0 ±	0.3	cd	Α	-6.5±0.3	С	Α
Firmness (N)	AIR NI	13.1 ± 1.0	аA	12.1 ± 0.8	a A	12.0 ± 0).9	а	А	11.5±0.6	а	В
	AIR CPA	13.0 ± 0.8	аA	12.8 ± 0.8	a A	12.5 ± 0).6	а	Α	13.0 ± 0.6	а	А
	MAP NI			14.3 ± 0.2	a A	12.2 ± 0).6	а	AB	11.7±0.3	а	В
	MAP CPA			11.3 ± 1.5	a A	11.7 ± 0).5	а	Α	13.1±0.9	а	Α
рН	AIR NI	3.8 ± 0.1	аA	4.0 ± 0.1	a A	3.9 ± 0).02	а	А	3.7±0.1	а	Α
	AIR CPA	4.1 ± 0.1	аA	3.9 ± 0.1	a A	4.1 ± 0).1	а	Α	3.9±0.2	а	А
	MAP NI			3.85 ± 0.03	a A	3.9 ± 0).1	а	А	3.9±0.1	а	А
	MAP CPA			3.9 ± 0.1	a A	3.9 ± 0).1	а	Α	4.15 ± 0.04	а	Α
Titratable	AIR NI	3.3 ± 0.2	аA							3.8±0.3	а	Α
acidity as malic acid content (g L ⁻¹ juice)	AIR CPA	3.22 ± 0.02	аA							3.6±0.5	а	А
	MAP NI									3.5 ± 0.3	а	А
	MAP CPA									3.6±0.2	а	Α
Soluble solids (%)	AIR NI	12.93 ± 0.04	аA							13.3±0.9	а	А
	AIR CPA	12.80 ± 0.06	аA							12.6 ± 0.1	а	А
	MAP NI									13.3 ± 0.1	а	А
	MAP CPA									12.6±0.3	а	А

Values are means \pm standard error of the mean. Different lowercase letters represent significant differences among treatments at each sampling time. Different capital letters represent differences for the same treatment throughout time according to analysis of variances (ANOVA) and Tukey's test (P < 0.05).

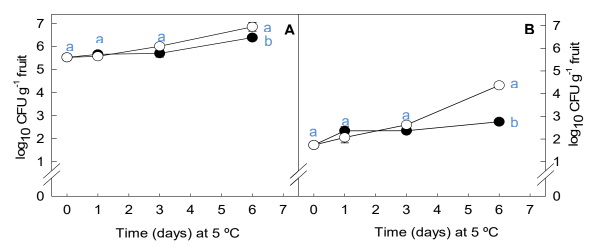


Figure 1. Population dynamics of (A) *P. graminis* CPA-7 and (B) total mesofils on 630 631 'Golden delicious' fresh-cut apple wedges treated with antioxidant, during six days of 632 storage at 5 °C in modified atmosphere packaging (MAP) (●) or in air (○). Each point represents the mean and error bars represent the standard error of the mean (n = 6). 633 634 Different letters represent significant differences among treatments at each sampling point according to analysis of variances (ANOVA) and Tukey's test (P < 0.05). 635 Underlined letters represent equal means that correspond with overlapped symbols in 636 637 the graph.

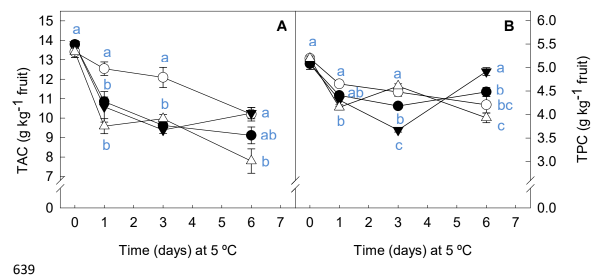


Figure 2. Changes in the levels of (A) Total Antioxidant Capacity, TAC, estimated from 641 the amount of Fe³⁺ and (B) Total Phenolic Content, TPC, estimated from the amount of 642 643 Gallic acid, in fresh-cut 'Golden delicious' apple wedges treated with antioxidant after 644 inoculation with CPA-7, during storage in air (O) or in MAP (\triangle) and in non-inoculated 645 control fruit during storage in air (\bullet) or in MAP (\mathbf{v}) . Each point represents the mean 646 and error bars represent the standard error of the mean (n = 9). Different letters represent significant differences among treatments at each sampling point according to 647 analysis of variances (ANOVA) and Tukey's test (P < 0.05). Underlined letters 648 represent equal means that correspond with overlapped symbols in the graph. 649

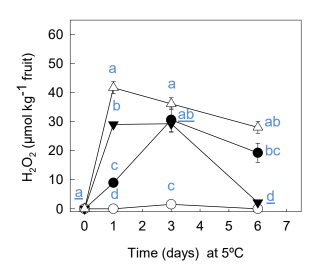
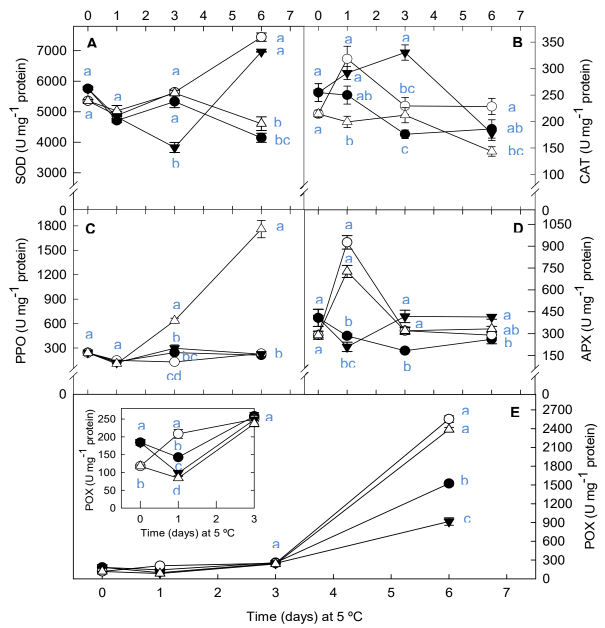


Figure 3. Changes in the levels of hydrogen peroxide (H₂O₂) in 'Golden delicious' freshcut apple wedges treated with antioxidant after inoculation with CPA-7, during storage in air (O) or in MAP (\triangle) and in non-inoculated control fruit during storage in air (\bullet) or in MAP (▼). Each point represents the mean and error bars represent the standard error of the mean (n = 9). Different letters represent significant differences among treatments at each sampling point according to analysis of variances (ANOVA) and Tukey's test (P < 0.05). Underlined letters represent equal means that correspond with overlapped symbols in the graph.



665 Figure 4. Changes in the activity of the enzymes A) superoxide dismutase (SOD), B) 666 catalase (CAT), C) polyphenol oxidase (PPO), D) ascorbate peroxidase (APX) and E) 667 peroxidase (POX) in 'Golden delicious' fresh-cut antioxidant-treated apple wedges, after inoculation with CPA-7, during storage in air (\bigcirc) or in MAP (\triangle) and in non-668 inoculated control fruit during storage in air (\bullet) or in MAP (\mathbf{v}). The internal plot in 669 graph E represents the first three sampling times at a smaller scale. Each point 670 671 represents the mean and error bars represent the standard error of the mean (n = 9). 672 Different letters represent significant differences among treatments at each sampling

point according to analysis of variances (ANOVA) and Tukey's test (P < 0.05).
Underlined letters represent equal means that correspond with overlapped symbols in
the graph.