

1 **Evaluation of the antioxidant/antimicrobial performance of *Posidonia oceanica* in comparison**
2 **with three commercial natural extracts and as a treatment on fresh-cut peaches (*Prunus***
3 ***persica* Batsch)**

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25 **ABSTRACT**

26 This research aimed at extending the choice of natural antimicrobials/antioxidants for food
27 applications. Four plant extracts, *Posidonia oceanica* (PO), Green Tea (GT), Grape seeds (GS) and
28 Grape skin (GK), were analyzed to determine their total phenolic content, antioxidant activity and
29 *in vitro* antimicrobial performance. PO extract showed the highest total phenolic content (711 mg
30 gallic acid/g extract) and antifungal activity against *Aspergillus niger* and *Penicillium chrysogenum*.
31 The highest antioxidant (3.81 mg/L EC₅₀) and antibacterial activities (bactericidal against Gram
32 positives and bacteriostatic against Gram negatives) were found for GT extract.
33 The best performing extracts (PO and GT) were applied by dipping on peach slices in storage trials.
34 Microbiological and pomological parameters were evaluated during 7 d storage. Total aerobic
35 count, *Pseudomonas* as well as yeasts and moulds populations, were reduced by about 0.5 log cfu/g,
36 mainly up to 5 d in all treated samples compared to the control. Total soluble solids, titratable
37 acidity and colour (L*a*b*) changes were also delayed in treated fruit.

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39

40 **KEYWORDS**

41 Ready-to-eat fruit, Green tea, *Posidonia oceanica*, dipping, antimicrobials, antioxidants

42 **1. Introduction**

43 One of the most important research areas as rated by a large majority of food companies is the
44 development of healthy foods, and the introduction of fresh cut produce onto the market, in order
45 facilitate fruit consumption, is rapidly growing (Jung and Zhao, 2016).

46 Nevertheless, the high perishability of minimally processed fruit may lead to an increase in food
47 waste and economic losses (Amani and Gadde, 2015). Throughout production process, cell
48 breakage takes place causing juice leakage and leading to microbial contamination and growth.
49 Moreover, the contact between enzymes and cell juice under oxygen exposure increases cell
50 respiration and activation of fruit senescence. Specifically, minimally processed fruit, and peaches
51 in particular, are very susceptible to flesh browning (Denoya et al., 2016). Therefore one of the
52 current challenges for the agro-food companies is to lengthen cut fruit shelf life, consequently
53 improving attractiveness to customers as well as food safety.

54 The food industry has been increasingly employing polyphenols to limit enzymatic oxidation which
55 affects the shelf life of ready-to-eat fruit (Gyawali and Ibrahim, 2014). The beneficial properties of
56 polyphenols on human health also have to be taken into account (Pandey and Rizvi, 2009).

57 As sources of polyphenols, several trials have been carried out using plant extracts from common or
58 endemic species (Perumalla and Hettiarachchy, 2011) or alternatively from by-products of the agro-
59 food industry (Balasundram et al., 2006). Nowadays, exploitation of by-products and/or residues
60 represents one of the environmental and economic priorities. Several substances discarded from
61 agro-food production can find alternative applications in different contexts. As examples, grape skin
62 (GK) and seeds (GS) are the main wastes from the wine industry, nevertheless they are appreciated
63 for their high phenolic content which includes flavonoids, phenolic acids and non-flavonoid
64 compounds (Poudel et al., 2008). The hydroxyl groups of gallic acid, present in grape by-products,
65 showed antimicrobial activity against *Bacillus cereus*, *B. subtilis*, *B. coagulans*, *Staphylococcus*
66 *aureus*, *Escherichia coli* and *Pseudomonas aeruginosa*; also all the substituents of the benzene

67 rings were found effective against *S. aureus* (evaluated by Minimal Inhibitory Concentration assay)
68 (Jayaprakasha et al., 2003).

69 The antioxidant capacity of polyphenols can also be used to prevent or slow down enzymatic
70 oxidation of vitamins and pigments contained in ready-to-eat fruit and vegetables (e.g. enzymatic
71 browning), thus preventing the loss of nutritional elements and increasing attractiveness to
72 consumers due to the maintenance of their sensorial characteristics (Rojas-Grau et al., 2009).
73 Moreover, they can be added as antimicrobials thus increasing product shelf life (Guillen et al.,
74 2013).

75 Flavonoids from plants have high antioxidant capacity and they are widely used substances,
76 including catechin, epicatechin, galocatechin, epigallocatechin, catechin gallate, epigallocatechin-
77 3-gallate (the most abundant and biologically active compound in green tea), galocatechin gallate
78 and epicatechin gallate (Sutherland et al., 2006). The hydroxyl groups in the ring structure of
79 catechin can be easily oxidized (Janeiro and Brett, 2004).

80 Green tea (GT) is one of the plant extracts with high antioxidant and antibacterial activities, and
81 with anti-tumor effects due to its catechin content. GT catechin showed antimicrobial activity
82 against Gram positive and Gram negative bacteria including certain pathogens of the
83 gastrointestinal tract such as *S. aureus*, *S. epidermis* and *Plesiomonas shigelloides*, but it was not
84 effective against *E. coli*, *Pseudomonas aeruginosa* and *Aeromonas hydrophila* (Kusmita et al.,
85 2014).

86 *Posidonia oceanica* (PO) is a marine endemic plant of the Mediterranean sea protected by the EU
87 (92/43 EEC Habitat Managerial and Community Board 97/62/EEC). It is an important species in
88 coastal waters defence, forming extensive marine grasslands (Foden et al., 2007). Twenty-three
89 phenolic compounds were identified in this species (Cuny et al., 1995; Agostini et al., 1997) and
90 several studies showed that PO extract is able to inhibit the growth of both Gram positive and Gram
91 negative bacteria, and it was particularly effective against *P. aeruginosa* and *S. aureus* (Berfad and
92 Alnour, 2014), as well as yeasts. PO extract was also assayed in the biomedical field, proving its

93 high anti-diabetic and anti-oxidant effects (Gokce and Haznedaroglu, 2008). However, some reports
94 found evidence for the transfer of toxins originating from toxic dinoflagellates which live as
95 epiphytes on PO leaves (Bellassoued et al., 2012).

96 The present research is aimed at extending the choice of natural antimicrobials/antioxidants for food
97 applications, derived from PO, GT, GS and GK. These extracts were analyzed to determine their
98 total phenolic content and antioxidant activity as well as *in vitro* antimicrobial performance. The
99 two best performing extracts were also used to set up fresh-cut storage trials on peach slices,
100 applying the dipping procedure. Peach (*Prunus persica* L. Batsch) is a climacteric fruit that contains
101 carbohydrates, organic acids, pigments, phenolics, vitamins, volatiles, antioxidants and trace
102 amounts of proteins and lipids, which make it very attractive to consumers (Kader and Mitchell,
103 1989). However, peaches are susceptible to physiological disorders (internal breakdown and
104 chilling injury), pathogen (moulds) and processing manipulation (browning of tissues) (Caceres et
105 al., 2016).

106

107 **2. Materials and methods**

108

109 **2.1 Chemicals**

110 2,2-Diphenyl-1-picrylhydrazyl (DPPH), (\pm)-6-Hydroxy-2,5,7,8-tetramethylchromane-2-carboxylic
111 acid (Trolox) 97%, Ethanol (\geq 99.8%), Ethyl acetate (anhydrous, 99.8%), Folin-Ciocalteu's phenol
112 reagent, hydrochloric acid, sodium hydroxide and sodium sulfate (\geq 99.0%, anhydrous), were
113 purchased from Sigma–Aldrich (Gallarate, MI, Italy).

114 Green tea, grape skin (*Vitis vinifera* L., Chardonnay variety) and seed extracts for oenological use
115 (antioxidants) were obtained from DAL CIN GILDO S.p.A. (Concorezzo, MB, Italy).

116

117 **2.2 *Posidonia oceanica* extract**

118 *Posidonia oceanica* (L.) Delile was collected by scuba diving from Palermo (Sicily, Italy),
119 Tyrrhenian Sea, in October 2014. Note that as this is a protected marine plant, for use in any
120 industrial application, it should be sourced from aquaculture systems under controlled growing
121 conditions. The epiphytes on the leaves were removed with paper towels without damaging the
122 organs, as reported by Gokce and Haznedaroglu (2008). Leaves were dried in the dark at $20 \pm 1^\circ\text{C}$
123 and then stored at $4 \pm 1^\circ\text{C}$ before use. The extract was obtained according to the method of Gokce
124 and Haznedaroglu (2008). Briefly, homogenized tissues were infused in 50% (v/v) ethanol-water
125 solution for 3 h in a water bath at 40°C with a reflux system in the dark. The homogenate was
126 filtered and acidified at pH 3 with hydrochloric acid 2 N. After evaporation of ethanol under
127 vacuum at 45°C , the aqueous residue was extracted with ethyl acetate. The organic phase was
128 filtered and evaporated under vacuum. The extract obtained, which was a green viscous material
129 (Figure 1), was freeze dried and finally stored at -20°C until use.

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132

133 **Figure 1.** Extract of *P. oceanica*

134

135 **2.3 Total phenolic content**

136 Total phenols (TP) levels of the PO, GT, GS and GK extracts were estimated colorimetrically by
137 the Folin-Ciocalteu method (Scalbert et al., 1989). Extracts (1 g/L) were dissolved in 50% (v/v)
138 methanol/water and appropriately diluted (1:2.5, 1:5 and 1:10 v/v) in the same solvent. The Folin-

139 Ciocalteau reagent was 10-fold diluted in water (v/v) and 2.5 mL were added to each 0.5 mL
140 sample. Two milliliters of 75 g/L sodium carbonate solution were added and tubes kept for 1 h at 20
141 $\pm 1^\circ\text{C}$ in the dark. In the meanwhile, the calibration curve for gallic acid (5-100 mg/L) dissolved in
142 50% (v/v) methanol/water was achieved. The absorbance at 765 nm was measured and results were
143 expressed as g gallic acid/100 g powder. Each formulation was analyzed in triplicate.

144

145 **2.4 Antioxidant assay**

146 Analysis of the antioxidant capacity of PO, GT, GS and GK extracts was carried out employing the
147 DPPH assay, following the method of Brand-Williams et al. (1995) with some modifications. The
148 DPPH solution was diluted in methanol to obtain 1.00 ± 0.03 absorbance units at 515 nm. The
149 extracts samples were dissolved (20 g/L) in 70 % methanol (v/v) and, after centrifugation, they
150 were serially diluted. The DPPH solution (2.94 mL) was placed in a cuvette where a 60 μL sample
151 was added. The absorbance readings were carried out after incubation for 50 min at $20 \pm 1^\circ\text{C}$. A
152 calibration curve was prepared by adding increasing concentrations of Trolox ranged from 50 to
153 1000 μM ; each concentration was assayed in triplicate. Results were expressed as mol Trolox per
154 100 g dry weight. Each formulation was analyzed in triplicate.

155

156 **2.5 Microorganisms and culture conditions**

157 Antimicrobial activity was analyzed by carrying out *in vitro* tests determining the Minimal
158 Inhibitory Concentration (MIC) against strains belonging to official collections, i.e. *Escherichia*
159 *coli* CECT 434 (Spanish Type Culture Collection), *Listeria innocua* DSM 20649 (Deutsche
160 Sammlung von Mikroorganismen und Zellkulturen), *Pseudomonas putida* ATCC 12633
161 (American Type Culture Collection), *Staphylococcus aureus* ATCC 29213, *Aspergillus niger*
162 NRRL 565 (Agricultural Research Service Culture Collection) and *Penicillium chrysogenum*
163 CECT 2802. These microorganisms were selected among the most common spoilage and/or
164 pathogen microorganisms that might be present in fresh food products (Mascheroni et al., 2014).

165 Bacterial strains were weekly maintained on TSB (Tryptic Soy Broth, Scharlau Chemie, Spain),
166 incubated at 30°C for 24 h and then stored at 4°C, while moulds were maintained on MEA solid
167 culture (MEB added with 15 g/L agar), incubated at 25°C for 5-7 d and then stored at 4°C until use.

168

169 **2.6 Determination of antimicrobial activity *in vitro***

170 Qualitative determination of antimicrobial activity was performed as follows: 30 mL of soft TSA or
171 MEA (TSB or MEB added with 8 g/L agar) were poured in a Petri Dish and inoculated with 300 µL
172 of a microbial suspension prepared in sterile distilled water (OD 600nm: 0.300 ± 0.050); moulds
173 were inoculated as spores suspension in sterile distilled water (OD 600 nm: 0.300 ± 0.050). Once
174 solidified, holes were made by using a sterile tip and 150 µL of extracts were poured inside.
175 Cultures were all incubated at each appropriate temperature for 24 h (up to 7 d for moulds). The
176 presence of a growth inhibition halo around holes indicates an antimicrobial activity.

177 Quantitative determination of antimicrobial activity was performed only with PO and tea extracts.
178 Ten mL of soft TSA or MEA were poured in a Petri Dish, to which aliquots of PO extract were
179 added in order to obtain a final concentration of 0.5-1.0-1.5-2.0-2.5 and 3.0 g/L. Tea
180 extract was tested at the following concentrations: 0.5-1.0-1.5-2.0 g/L. Once solidified,
181 plates were all surface inoculated with aliquots (3 µL) of appropriately diluted (OD 600nm:
182 0.300 ± 0.050) overnight microbial cultures on TSB (bacteria). Moulds were inoculated as spore
183 suspension in sterile distilled water (OD 600 nm: 0.300 ± 0.050). For each strain control trials were
184 also prepared without the addition of extracts. In order to highlight any possible inhibitory effect of
185 the solvent present in extracts, a set of solid cultures was also set up by adding ethanol (up to 3 g/L)
186 to the culture media.

187 Cultures were all incubated at each appropriate temperature for 24 h or up to 7 d for moulds. MIC
188 (Minimum Inhibitory Concentration) was determined as the lowest extract concentration (g/L) able
189 to inhibit microbial growth. Trials were repeated twice for each extract.

190

191 **2.7 Fresh-cut peach storage**

192 Peaches (*Prunus persica* L. Batsch cv. ‘Rich May’) were purchased at the wholesale market (3 kg),
193 24 h after harvesting at commercial maturity and stored for 1 d at 4°C until use. Peaches were
194 homogenous in weight (180 ± 5 g) and ripening (10.85 ± 0.07 °Brix). Fruits were pre-washed with
195 distilled water, sanitized for 2 min in chlorinated water (150 mg/L sodium hypochlorite), rinsed
196 with distilled water and gently dried by hand. Peaches with skin were cut into slices (8 slices per
197 fruit) of about 1.5 cm thickness (15 ± 2 g each slice), using a sterile stainless-steel knife, and dipped
198 for 3 min in the following solutions: PO extract (2% w/v), GT extract (1% w/v) and distilled water
199 for control samples (CTRL). Slices were then left in the air for 2 min in order to drain off the excess
200 solution. Three slices (45 ± 6 g) were placed into low-density polyethylene (LDPE) bags (22 x 15
201 cm, 25 µm thickness, bag volume 450 mL, ratio between fruit weight and container volume 100
202 g/L, surface film for each bag 660 cm², O₂ permeability 6200 cm³ m⁻² d⁻¹ bar⁻¹, CO₂ permeability
203 24000 cm³ m⁻² d⁻¹ bar⁻¹ at 10 °C). Bags were all stored at 4 ± 1 °C up to 7 d. A total number of 48
204 bags were prepared, 24 used for color evaluation, and the remaining ones for the other analyses.
205 Samples were then collected after 0, 3, 5 and 7 d. Each trial at each day was carried out in duplicate.

206

207 **2.8 Colour evaluation**

208 Flesh colour was evaluated using the CIE L*a*b* System by a Minolta CR-300 chromameter
209 (Konica Minolta Sensing, Inc., Japan). Three measurements were performed on each side of slices.
210 The instrument was calibrated using a standard white plate. The chroma (C) was calculated as
211 follows (1):

$$212 \quad C^* = \sqrt{a^{*2} + b^{*2}} \quad (1)$$

213

214 **2.9 Total soluble solids and titratable acidity**

215 Total soluble solids (TSS, %) and titratable acidity (TA, g/L) were measured on the juice obtained
216 from slices (30 g for each sample) by an electronic blender (Ariete, Italy). TSS were determined by

217 a digital refractometer (Atago Co., Ltd, Tokyo, Japan model PR-32), while TA was determined by
218 titrating 1:10 diluted juice using sodium hydroxide 0.1 M by an automatic titrator (Compact 44-00,
219 Crison Instruments, SA, Barcelona, Spain).

220

221 **2.10 Antimicrobial activity of green tea and *P. oceanica* on peach slices**

222 After 0, 3, 5 and 7 d, peach slices (15 g) were transferred aseptically into a Stomacher bag (400 mL
223 PE, Barloworld, France) containing 135 mL of sterile peptoned water (10 g/L bacteriological
224 peptone, Costantino, Italy) and blended in a Stomacher (Star Blender LB 400, Biosystem, Belgium)
225 at high speed for 3 min. Ten-fold dilution series of the obtained suspensions were made of the same
226 solution for plating. The following culture media were used: TSA (Merck, Germany) for
227 mesophiles, *Pseudomonas* Agar base (Himedia, India) for *Pseudomonas spp.*, VRBLA (Violet Red
228 Bile Agar, Merck, Germany) for *Enterobacteriaceae* and MEA for yeasts and fungi. Colonies were
229 counted after incubation at 30°C for 24 h for mesophiles, 30°C for 5 d for yeasts and fungi and
230 25°C for 24 h for *Pseudomonas*. Counts were performed in triplicate and reported as logarithms of
231 the number of colony forming units (log cfu/g peach), and means and standard deviations (SD) were
232 calculated.

233

234 **2.11 Statistical analysis**

235 Statistical analysis was carried out using the STATISTICA 7.1 software package (Statsoft Inc.,
236 Tulsa, OK, USA). One-way analysis of variance (ANOVA) was performed on mean values and
237 Tukey's test was carried out for the comparison of difference among treatments for each storage
238 time and for each treatment during storage. Differences were considered significant at $p \leq 0.05$.

239

240 **3. Results and discussion**

241

242 **3.1 *In vitro* evaluation of total phenolic index and antioxidant activity**

243 Total phenol index (TPI) and antioxidant capacity (AC) were evaluated for each investigated extract
244 (Table 1). PO showed the highest TPI (711 mg/g) followed by the GS extract (526 mg/g). The
245 highest AC was found for GT extract (3.8 ± 0.11 mg/L EC₅₀).

246 In case of PO, the highest polyphenols content corresponded to the lowest antioxidant capacity.
247 Such a low value could be attributable not only to the extraction procedure but can also be related to
248 the type of polyphenols present in the extracted matrix (Berfad and Alnour, 2014). The choice of
249 the solvent to use (50% v/v ethanol-water solution) was based on the best results obtained by Berfad
250 and Alnour (2014), who investigated the extraction performance of different solvent mixtures on *P.*
251 *oceanica*, as well as also taking into account its food-grade nature. As reported in the literature, the
252 principal polyphenols present in PO are acetosyringone, ferulic acid and acetovanillone, while those
253 in GT are gallic acid, catechin gallate and epicatechin (Agostini et al., 1998). These last compounds
254 are characterized by the presence of three proximal hydroxyl groups, able to efficiently delocalize
255 radicals present in the aromatic ring thus acting as radical scavengers. On the contrary, the
256 polyphenols present in PO hydroxyl groups are not in close proximity and a methyl moiety is often
257 present, thus reducing its antioxidant capacity (Agostini et al., 1998).

258 Nevertheless, the polyphenols values obtained were comparable with those reported in the literature
259 for other plants (Mensor et al., 2001). The highest antioxidant capacity of the GT extract is not
260 surprising since GT is particularly rich in phenols, as proanthocyanidins, with low redox potential
261 and it does not contain bi-flavanols (Lee et al., 2014).

262

263 **3.2 *In vitro* determination of antimicrobial activity**

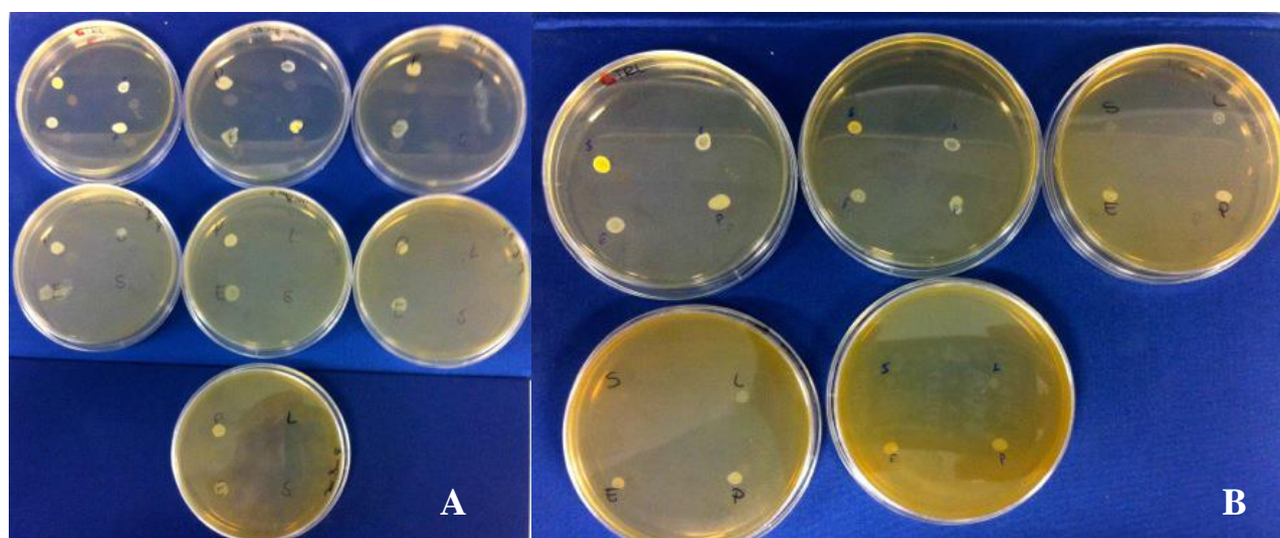
264 In qualitative trials performed employing 1 g/L extract solutions, all samples showed the highest
265 antimicrobial activity against *L. innocua*. GT proved to be the most powerful sample, while at the
266 tested concentration PO did not show any antibacterial activity but was the only extract possessing
267 an antifungal action against *Aspergillus niger* (Table 2).

268 Quantitative determination of antimicrobial performance found that MIC for GT and PO extracts
269 were 1 and 2 g/L respectively against the two Gram positive strains *L. innocua* and *S. aureus*. The
270 two Gram negative strains showed a reduction of microbial growth attributable to a bacteriostatic,
271 rather than a bactericidal, effect (Figure 2). *Aspergillus* and *Penicillium* showed a marked reduction
272 of hyphal growth (up to 30% for *A. niger* and to 70% for *P. chrysogenum* employing 3 g/L extract)
273 and sporulation when grown in presence of PO, while no effect was evident with GT (Figure 3).
274 Only in the control plate (no extract added) after 7 d incubation the two moulds showed an
275 antagonistic effect, while in all the other plates the hyphal growth of each strain was not influenced
276 by the presence of the other one. Note that ethanol, the solvent used to prepare extracts, was not
277 found to inhibit microbial growth at the tested concentrations, thus confirming literature results
278 (Dantigny et al., 2005).

279 Recently, Alkan and Yemenicioglu (2016) tested the *in vitro* antimicrobial activity of various plant
280 phenolics, finding that clove extract was the most potent antimicrobial, with MIC values of 10.24
281 g/L against the plant pathogens *Erwinia amylovora*, *E. carotovora*, *Pseudomonas syringae* and
282 *Xanthomonas vesicatoria*. The reported values are much higher than those found in the present
283 research (MIC of 1-2 g/L), at least for PO, highlighting that the tested extracts are of actual interest.

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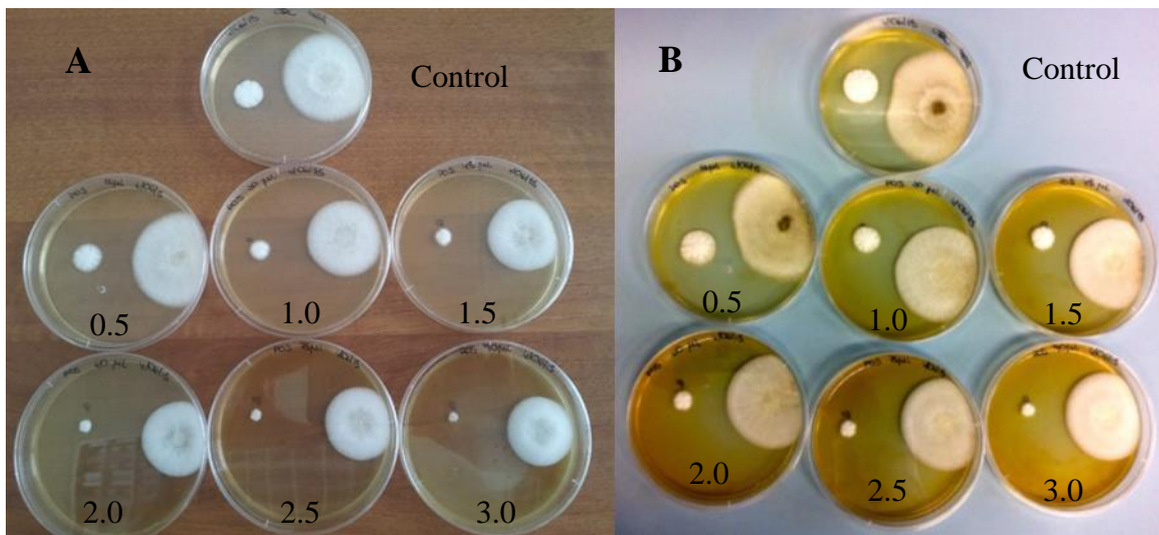
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287 **Figure 2.** Growth of bacterial strains in presence of *P. oceanica* (A) and Green Tea (B) extracts.
288 Concentrations used: 0-0.5-1-1.5-2-2.5-3 g/L PO; 0-0.5-1-1.5-2 g/L GT. Strains: *L-Listeria*
289 *innocua*, *S-Staphylococcus aureus*, *E-Escherichia coli*, *P-Pseudomonas putida*.

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295 **Figure 3.** Growth after 2 days (A) or 7 days (B) incubation of *Penicillium chrysogenum* (left in
296 each plate) and *Aspergillus niger* (right in each plate) in presence of different concentration of *P.*
297 *oceanica* extract (from 0.5 to 3.0 g/L). Control: medium without extract.

298
299

300 **3.2 Fresh-cut peach storage**

301 GT and PO extracts were chosen for the dipping of fresh cut peach slices: the first of these had the
302 most powerful antioxidant capacity, while the second was chosen for its high polyphenols content
303 and antifungal activity. Fruits are usually quite acid and hence quite resistant to invasion by
304 bacteria. Therefore spoilage of fruit and fruit products is often caused by fungi (Pitt and Hocking,

305 1999). However, recent studies have documented the exponential growth of bacteria on a variety of
306 fresh-cut fruit (Alegre et al., 2010; John et al., 2013).

307 In terms of pomological traits, after 7 d storage, TSS content was found almost constant in treated
308 samples of peach slices, while in CTRL samples this value sharply increased up to 11.8 %, mainly
309 from 5 to 7 d (Figure 4A).

310 Differences of TA content were significant between t 0 and 3 d, then reduced with storage. CTRL
311 samples evidenced a significant decrease from 3 to about 1 g/L. Peach slices dipped in the two
312 extracts showed a TA decrease of only 17 % with respect to the initial value, but with different time
313 courses: when PO was employed, a sharp decrease of TA in the last 3 d of shelf life was evident,
314 while in samples treated with GT the decrease was evident in the first 3 d.

315 Note that the sharp decrease of TA in CTRL samples occurred at the same time with the TSS
316 increase in the last 3 d of the shelf life. For the GT treatment, no significant changes were evident in
317 TSS between 3 and 7 d (Figure 4B).

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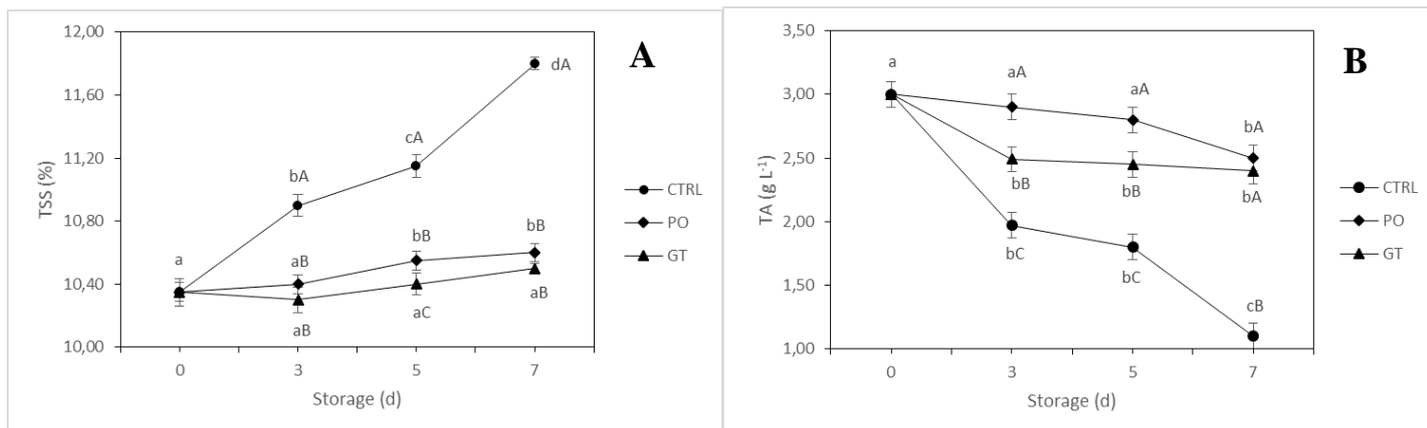
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327 **Figure 4.** Evolution of total soluble solids (%) (A) and titratable acidity (g/L) (B) in peach (*Prunus*
328 *persica*) cv. Rich May slices treated with *Posidonia oceanica* (PO), Green Tea (GT) and untreated
329 (CTRL). Data are means ± SE. Minor and capital letters show significant differences ($p \leq 0.05$) for
330 each treatment and among treatments for each storage time, respectively.

331

332

333 After three days, lightness (L^*) of peach slices decreased in all treatments (Table 3). When
334 prolonging the incubation time up to 7 d, lightness decreased much more in CTRL sample rather
335 than in peach slices treated by GT or PO. Caceres et al. (2016) developed a flesh browning
336 assessment methodology for fresh whole peaches stored for a long time, and reported that ΔL^*
337 values higher than 21 can be considered symptoms of extreme flesh browning. Even if our data
338 relate to peach slices, ΔL^* values calculated at 7 d storage were found higher than 21 only in the
339 control sample.

340 An indicator of chlorophyll degradation is the a^* value, which decreases when color changes from
341 green to red (Martín-Diana et al., 2008). An increase of a^* values was only found for PO treatment
342 (Table 3). This behavior can be attributed to the greenish colour imparted by the PO extract,
343 similarly to what was found by Martín-Diana et al. (2008), after a dipping treatment with natural
344 extracts, which they also correlated to lightness lowering.

345 The b^* parameter indicates the color changes from yellow to blue, and its values decreased in all
346 samples during storage (Table 3) due to phenolic degradation taking part on tissues (Fuentes-Perez,
347 2014). However, after 7 d storage, treated samples showed higher values than the control.

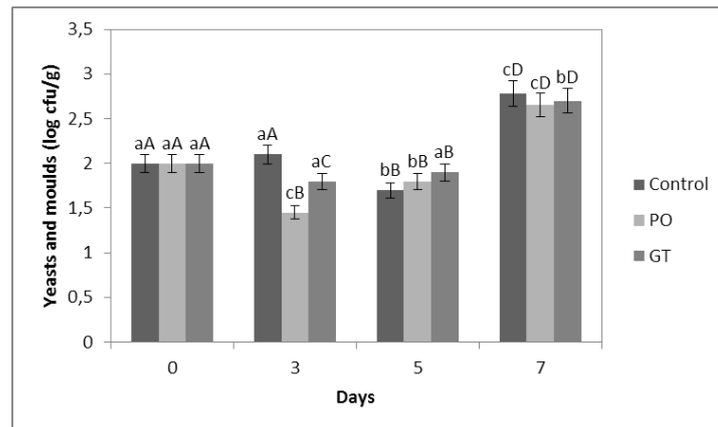
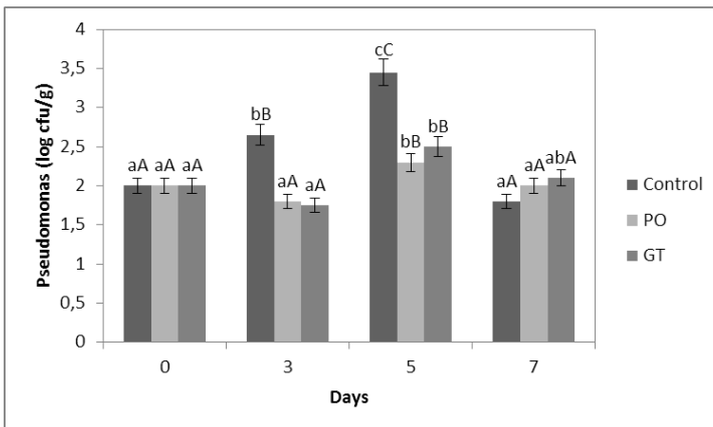
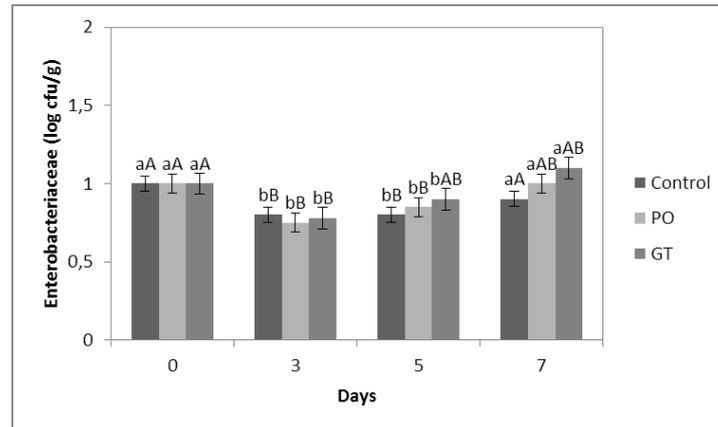
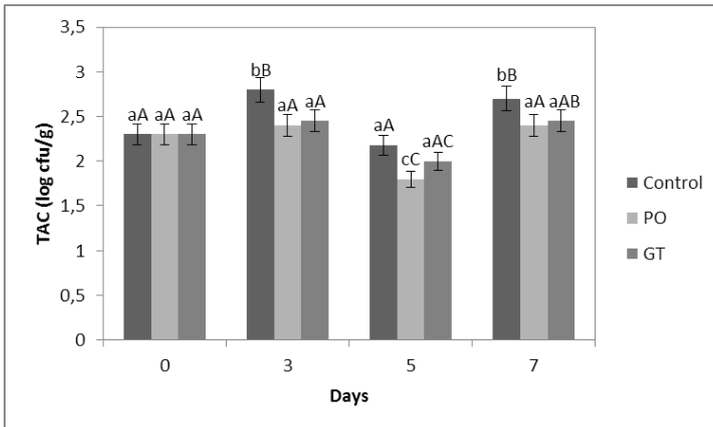
348 Finally, the Chroma decreased after 3 and 5 d storage more rapidly in PO samples compared to GT
349 ones (Table 3). After 7 d, GT showed a higher value compared with samples dipped in PO.

350 These results, in accordance with the findings reported by Oms-Oliu et al. (2010), highlighted that
351 dipping treatment after peeling and/or cutting can represent an effective way to control browning
352 phenomena in fresh-cut fruit, since it can affect enzyme activity or substrates availability for
353 enzymatic degradation. In particular, the high polyphenols content present in both extracts can
354 protect the cut surface of fruit products against oxidative rancidity, degradation and enzymatic
355 browning, thus slowing their senescence process (Rojas-Grau et al., 2009).

356

357 From the microbiological point of view, the applied dipping treatments were found effective in
358 lowering the total aerobic count (TAC) and *Pseudomonas* population present in peach slices of
359 about 0.5 log cfu/g mainly up to 5 d (Figure 5). This behavior may also be favored by the fruit
360 respiration process (Rojas-Grau et al., 2009). Yeasts and moulds were found significantly lower
361 than the CTRL only at 3 d, with the best performance being shown by peach slices dipped in PO
362 extract. No significant changes were found for the *Enterobacteriaceae* population. These results are
363 in accordance with those reported by Siroli et al. (2014) for minimally processed apples dipped in
364 different antimicrobials comparatively: shelf life of fresh cut fruit is limitedly affected by microbial
365 growth: independently from the addition of natural antimicrobials, the end of shelf life is mainly
366 determined by changes in colour.

367 Time course of CO₂ and O₂ during storage trials was not determined as it was assumed meaningless
368 because of the high gas permeability and the geometry of the packaging system used. Assuming an
369 oxygen respiration rate of the fruits of about 18 mg kg⁻¹ h⁻¹ at 10 °C and taking into account the
370 permeable surface of the package (0.066 m²), the headspace volume (450 cm³), the amount of the
371 product (0.045 kg) and the oxygen and carbon dioxide permeabilities (respectively 6200 and 24000
372 cm³ m⁻² d⁻¹ bar⁻¹, at 10 °C) using a common model for forecasting atmosphere changes in ready-to-
373 eat vegetables (Piergiovanni et al., 1999) we estimated after 7 d a maximum CO₂ concentration
374 equal to 0.8% and a minimum O₂ concentration of 18%.



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376

377 **Figure 5.** Time course of Total Aerobic Count (TAC), *Enterobacteriaceae*, *Pseudomonas* and
 378 yeasts and moulds presence (log cfu/g peach) in samples of peach slices subjected to dipping
 379 treatment with *P. oceanica* (PO) and Green Tea (GT) extracts and then stored at 4 °C. CTRL: peach
 380 slices dipped in sterile distilled water. Data are means ± SD. Minor and capital letters show

381 significant differences ($p \leq 0.05$) for each treatment and among treatments for each storage time,
382 respectively.

383

384 **4. Conclusions**

385 In this study *Posidonia oceanica* (PO) and green tea (GT) extracts were applied by dipping on
386 peach slices, once having shown their highest total phenolic content and antifungal activity, as well
387 as the highest antioxidant activity, respectively. Results showed that these natural extracts limited
388 microbial spoilage of fresh-cut peach, especially the *Pseudomonas* population, and maintained the
389 pomological parameters during storage at 4°C while not modifying their characteristic taste.

390 Overall, polyphenolic extracts derived from PO and GT could provide an additional post-harvest
391 benefit of fresh-cut produce. To the best of our knowledge, this paper represents the first report on
392 the application of *P. oceanica* extract on fresh-cut fruit, even if this marine protected plant never
393 should be collected from the sea for industrial application.

394 Although this work relates to the application of natural extracts directly on fresh-cut fruit in a
395 traditional package, future trials will be aimed at setting up innovative active packaging solutions
396 from which these extracts will be released. Further research will be also needed to complement
397 antioxidant activity of plant or other extracts with digestion simulation to assess parameters such as
398 bioaccessibility, bioavailability and *in vivo* antioxidant performance; sensorial analyses should also
399 be performed on treated fruit.

400 These results can pave the way to the use of innovative natural extracts to be applied on ready-to-
401 eat vegetables, thus improving the attractiveness for consumers of these healthy foods.

402

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405

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517

518 **Table 1:** Total Phenolic Index (TPI) and Antioxidant Capacity (AC) of the extracts investigated in
 519 this study. Data are presented as mean \pm SE.

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| Extract | TPI | AC |
|---------------------------|--------------------------|-----------------------|
| | mg gallic acid/g extract | mg/L EC ₅₀ |
| <i>Posidonia oceanica</i> | 710.6 \pm 20.1 | 72.42 \pm 22.95 |
| Green tea | 526.3 \pm 14.9 | 3.80 \pm 0.11 |
| Grape skin | 398.2 \pm 9.5 | 6.14 \pm 0.78 |
| Grape seeds | 596.4 \pm 15.6 | 4.10 \pm 0.14 |

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531 **Table 2.** Diameter (mm) of microbial growth inhibition halos around wells containing the analyzed
 532 extracts (1 g/L) after incubation.

| Microorganism | GT | PO | GK | GS |
|--------------------------------|------|-------|------|------|
| <i>Listeria innocua</i> | 12 | n.p.* | 4 | 6 |
| <i>Staphylococcus aureus</i> | 10 | n.p. | 2 | 4 |
| <i>Escherichia coli</i> | 6 | n.p. | 2 | 4 |
| <i>Pseudomonas putida</i> | 4 | n.p. | n.p. | 4 |
| <i>Aspergillus niger</i> | n.p. | 8 | n.p. | n.p. |
| <i>Penicillium chrysogenum</i> | n.p. | n.p. | n.p. | n.p. |

533 *n.p.: inhibition halo not present.

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538 **Table 3.** Color parameters changes during storage of fresh cut peach slices treated by *P. oceanica*539 (PO), Green Tea (GT) and untreated. Data are presented as mean \pm SE. Minor and capital letters540 show significant differences ($p \leq 0.05$) for each treatment and among treatments for each storage

541 time, respectively.

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| Treatment | Day | Lightness (L*) | a* | b* | C* |
|-----------|-----|---------------------------------|---------------------------------|--------------------------------|---------------------------------|
| | 0 | 64.07 \pm 1.30 ^a | -2.22 \pm 0.41 ^a | 41.56 \pm 0.40 ^a | 41.58 \pm 0.40 ^a |
| | 3 | 59.63 \pm 0.85 ^{bA} | -1.04 \pm 0.18 ^{bB} | 39.63 \pm 0.55 ^{aA} | 39.73 \pm 0.54 ^{aA} |
| CTRL | 5 | 50.01 \pm 1.14 ^{cA} | 0.40 \pm 0.22 ^{cB} | 29.60 \pm 1.63 ^{bA} | 29.61 \pm 1.63 ^{bA} |
| | 7 | 41.70 \pm 0.89 ^{dB} | 1.57 \pm 0.27 ^{dB} | 18.96 \pm 0.28 ^{cB} | 19.06 \pm 0.27 ^{cB} |
| | 3 | 57.11 \pm 0.19 ^{bB} | -3.15 \pm 0.19 ^{abA} | 33.88 \pm 0.72 ^{bB} | 34.04 \pm 0.71 ^{bB} |
| PO | 5 | 48.69 \pm 1.11 ^{cA} | -1.65 \pm 0.40 ^{bA} | 25.92 \pm 0.84 ^{cA} | 26.03 \pm 0.84 ^{cA} |
| | 7 | 42.99 \pm 0.62 ^{dAB} | -0.28 \pm 0.41 ^{cA} | 20.05 \pm 0.39 ^{dA} | 20.12 \pm 0.40 ^{dAB} |
| | 3 | 55.19 \pm 0.79 ^{bB} | -1.29 \pm 0.24 ^{bB} | 39.39 \pm 0.45 ^{aA} | 39.42 \pm 0.45 ^{aA} |
| GT | 5 | 48.56 \pm 0.58 ^{cA} | 0.43 \pm 0.41 ^{cB} | 29.02 \pm 1.59 ^{bA} | 29.07 \pm 1.59 ^{bA} |
| | 7 | 44.71 \pm 0.68 ^{dA} | 1.24 \pm 0.09 ^{cB} | 20.28 \pm 0.44 ^{cA} | 20.32 \pm 0.44 ^{cA} |

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