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1 **Antifungal activity of GRAS salts against *Lasiodiplodia theobromae* in vitro and**
2 **as ingredients of hydroxypropyl methylcellulose-lipid composite edible coatings**
3 **to control Diplodia stem-end rot and maintain postharvest quality of citrus fruit**

4

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16 Highlights

- 17 • GRAS salts effectively inhibited mycelial growth of *Lasiodiplodia theobromae*.
- 18 • HPMC-BW coatings containing GRAS salts reduced severity of citrus Diplodia stem-end
19 rot.
- 20 • Quality of coated and cold-stored oranges was not adversely affected.

21

22 Abstract

23 A large amount of GRAS (generally recognized as safe) salts and concentrations were evaluated in
24 in vitro tests (inhibition of mycelial growth on PDA dishes) against *Lasiodiplodia theobromae*, the
25 causal agent of citrus Diplodia stem-end rot. Ammonium carbonate (AC, 0.2%), potassium sorbate
26 (PS, 2.0%), potassium carbonate (PC, 0.2%), sodium methylparaben (SMP, 0.1%), sodium
27 ethylparaben (SEP, 0.1%), sodium benzoate (SB, 2.0%), and potassium silicate (PSi, 2.0%) were
28 selected as the most effective. Disease control ability of edible composite coatings formulated with
29 hydroxypropyl methylcellulose (HPMC), beeswax (BW), and these selected antifungal GRAS salts
30 was assessed in in vivo experiments with ‘Ortanique’ mandarins and ‘Barnfield’ oranges artificially
31 inoculated with *L. theobromae*. Coatings containing 2% PS, 0.1% SEP, or 2% SB were the most
32 effective reducing disease severity (up to 50% reduction) and were also applied to non-inoculated
33 and cold-stored ‘Barnfield’ oranges to determine their effect on postharvest fruit quality. After
34 periods of 21 and 42 d at 5 °C followed by 7 d of shelf life at 20 °C, coatings containing SEP and
35 SB significantly reduced weight loss and did not adversely affect the physicochemical quality
36 attributes (firmness, soluble solid content, titratable acidity, and ethanol and acetaldehyde content)
37 and sensory flavor with respect to uncoated control fruit. Although the internal gas concentration
38 (CO₂ level) of coated fruit increased, the coatings did not induce off-flavors.

39

40 **Keywords:** orange, mandarin; Diplodia stem-end rot; nonpolluting postharvest decay control; food
41 additives; citrus coatings

42 1. Introduction

43 Citrus fruit are grown in more than 100 countries with tropical and subtropical climate. Spain,
44 producing more than 6 million tons in 2016, ranked sixth in the world, preceded by Brazil with a
45 production higher than 22 million tons, China, USA, India, and Mexico. While 80% or more of the
46 Brazilian production is devoted to the juice industry, in Spain this percentage is devoted to the fresh
47 fruit market. Spain, in fact, is the first worldwide exporter of citrus fruit for fresh consumption
48 (FAOSTAT, 2018).

49 One of the most important problems affecting both the fresh and juice citrus industries are
50 postharvest losses caused by fungal pathogens that infect the fruit before, during or after harvest,
51 but develop disease after harvest. Depending on many factors, these losses are estimated to reach
52 20-50% on developing countries and up to 25% in developed countries, even though the sector in
53 these countries has availability of major postharvest technologies for fruit preservation and quality
54 maintenance. Depending on the climatic areas where citrus are grown, the main pathogenic fungi
55 that cause the highest incidence of fruit postharvest disease differ. In high rainfall areas such as
56 Brazil and Florida the most relevant belong to genera that typically produce latent infection in the
57 orchard, e.g., *Lasiodiplodia*, *Phomopsis*, *Colletotrichum*, *Phytophthora*, *Alternaria*, and *Botrytis*,
58 among others. Nevertheless, in Mediterranean-type climate areas with lower summer rainfall such
59 as Spain, California, and South Africa, the most prevalent are wound pathogens such as
60 *Penicillium*, *Geotrichum*, and *Rhizopus* (Eckert and Eaks, 1989; Palou, 2014; Smilanick et al.,
61 2006).

62 *Lasiodiplodia theobromae* (Pat.) Griffon & Maubl. (Botryosphaeriaceae) [synonyms:
63 *Botryodiplodia theobromae* Pat., *Diplodia theobromae* (Patouillard) W. Nowell, *Diplodia*
64 *natalensis* Pole-Evans; teleomorph *Botryosphaeria rhodina* (Berk. & M.A. Curtis) Arx] is a
65 polyphagous and opportunistic fungus with reduced pathogenic specialization, which infects many
66 species of plants and can cause serious damage on fruit commodities both before and after harvest.
67 Important postharvest diseases caused by *L. theobromae* are stem-end rots of subtropical and

68 tropical fruits such citrus, avocado, mango, banana, papaya, pineapple, and persimmon, among
69 others (Palou et al., 2013; Ploetz et al., 1994). On citrus fruit, it causes the postharvest disease
70 commonly known as Diplodia stem-end rot (Zhang, 2014). Typically, the attack of the fungus
71 occurs at the stem end of the fruit before harvest, the infection remains latent in the calyx and disk
72 (button) and disease symptoms and fruit deterioration are only manifested when the fruit ripens
73 after harvest, the button senesces and abscises from the fruit, and the pathogen begins to actively
74 develop through both the central axis and the external rind surface of the fruit, where it causes
75 characteristic finger-like projections of brown tissue. The progress of the infection depends on the
76 growth and the enzymatic capability of the microorganism and the physiological and biochemical
77 status of the fruit (Brown and Wilson, 1968; Zhang, 2014). Disease incidence can be especially high
78 on early-season degreened citrus fruit because the presence of exogenous ethylene favors the
79 activity of the button abscission enzymes polygalacturonase (PG) and cellulase (CX) (Brown and
80 Burns, 1998). Decayed fruit tissue is initially firm, but later becomes wet and mushy (Brown and
81 Eckert, 2000). On artificial medium such as potato dextrose agar (PDA), the colonies of *L.*
82 *theobromae* are grayish to black, with abundant aerial cottony mycelia and globular pycnidia.
83 Conidia are ovoid to ellipsoidal, thick walled, initially hyaline, aseptate, but turn dark brown and
84 uniseptate as the colony ages (Luo et al., 2011, Pereira et al., 2006). Another fungal pathogen that
85 causes citrus postharvest stem-end rot, but not with the tear-staining symptoms of *L. theobroame*, is
86 *Phomopsis citri* H.S. Fawc. (teleomorph *Diaporthe citri* F.A. Wolf).

87 Chemical fungicides are commonly used as the main tool to control both preharvest and
88 postharvest diseases of citrus fruit. While postharvest fungicide application is mainly devoted to
89 reduce green mold, caused by *Penicillium digitatum* (Pers.) Sacc., and blue mold, caused by *P.*
90 *italicum* Wehmer, some common active ingredients such as thiabendazole, imazalil, fludioxonil, and
91 sodium o-phenylphenate have also shown effect against stem-end rots (Zhang, 2014). However,
92 their use is increasingly restricted due to public concerns about possible toxicological risks for
93 people and the environment associated with excessive chemical residues. In addition, continuous

94fungicide usage can lead to the proliferation of resistant pathogenic fungal strains that make the
95treatments ineffective. Therefore, alternatives to control citrus postharvest diseases including
96Diplodia stem-end rot are of great interest to ensure safe agricultural production and reduce
97environmental pollution (Palou et al., 2016; Wisniewski et al., 2016). One of such alternatives is the
98use of edible coatings formulated with food-grade antifungal compounds. Fruit can be directly
99coated with a thin layer of edible material in order to improve gas and moisture barriers, mechanical
100and sensory properties, convenience, and microbial protection and, as a consequence, prolong
101product shelf life (Janjarasskul and Krochta, 2010). Polysaccharides, proteins, and lipids are the
102main ingredients used to formulate edible coatings. In many cases, two or more of these ingredients
103can be mixed to produce composite edible coatings in order to reduce both water and gas exchange
104between the fruit and the surrounding environment (Hernández-Izquierdo and Krochta, 2008). In
105postharvest applications, different polysaccharides such as chitosan, methylcellulose,
106hydroxypropyl methylcellulose (HPMC), and carboxy methylcellulose (CMC) have been found to
107affect positively weight loss, firmness, brightness, taste, and other physicochemical and sensory
108quality attributes of citrus fruit (Arnon et al., 2015; Navarro-Tarazaga et al., 2007, 2008).

109 To wide and enhance the functionalities of these coatings and further extend the shelf life of
110coated fresh produce, additional antimicrobial ingredients can be added to the emulsions to provide
111activity against pathogens and contaminating microorganisms (Valencia-Chamorro et al., 2011a).
112Antimicrobial agents used for the formulation of edible coatings should be classified as food-grade
113additives or compounds generally recognized as safe (GRAS) by the competent authorities.
114International regulators are responsible for approving antimicrobials for use in food. In the
115European Union (EU), these compounds are regulated by the EU Framework Directive 89/107 (EU,
1161989), while in the United States (USA) by the title 21CFR172 promulgated by the US Food and
117Drug Administration (US FDA, 2009). According to Palou et al. (2016), the antimicrobial
118ingredients used for formulation of antifungal synthetic biopolymer-based coatings can belong to
119three different categories, depending on their nature: i) synthetic food preservatives or GRAS

120 compounds such as various inorganic and organic salts, ii) natural compounds such as essential oils
121 and other natural plant extracts, and iii) microbial antagonists as biocontrol agents (bacteria, yeast,
122 yeast-like fungi, and even some filamentous fungi). General advantages of GRAS salts are their
123 high solubility in water, availability, feasibility of use as postharvest treatments for fresh fruit, and
124 good price. Intensive previous work at the IVIA CTP has resulted in the development and
125 characterization of HPMC-lipid edible coatings containing GRAS salts with antifungal activity
126 against citrus postharvest green and blue molds (Palou et al., 2015; Valencia-Chamorro et al. 2008,
127 2009a). This type of coatings were also effective for the control of important postharvest diseases of
128 other fresh commodities such as plums (Karaca et al., 2014) and cherry tomatoes (Fagundes et al.,
129 2013, 2015). However, to our knowledge, no studies are available on the development of edible
130 coatings with antifungal activity against citrus postharvest stem-end rot caused by *L. theobromae*.

131 The aim of this research work was to evaluate the in vitro activity of GRAS salts against *L.*
132 *theobromae* and to develop novel stable HPMC-lipid coatings containing selected GRAS salts. The
133 ability of these coatings to effectively control *Diplodia* stem-end rot was assessed in in vivo
134 experiments with mandarins and oranges artificially inoculated with the pathogen. Furthermore, the
135 effects of selected antifungal edible coatings on fruit physico-chemical and sensory quality were
136 determined on oranges stored at 5 °C for up to 42 d.

137

1382. Materials and methods

1392.1. Pathogen and fungal inoculum

140 In this work, the strain *L. theobromae* NEU-1 from the IVIA CTP culture collection of
141 postharvest pathogens was used. It is an isolate obtained from a decayed orange found in a citrus
142 packinghouse in Valencia province (Spain). Before each experiment, the isolate was grown on PDA
143 medium (Sigma-Aldrich Chemie, Steinheim, Germany) in Petri dishes in the dark in an incubation
144 cabinet at 25 °C for 7-14 d. Five-mm diameter mycelial plugs were cut from these cultures with a
145 sterilized cork borer and used as described below for plate and fruit inoculations in in vitro and in

146vivo tests, respectively.

147

1482.2. GRAS salts

149 The compounds, acronyms, molecular formulas, and molecular weights of the antimicrobial
150agents used in this work are given in Table 1. They are inorganic and organic salts classified as
151GRAS or as food additives by the USA or EU competent legislation. Laboratory reagent grade
152preservatives (99% minimum purity) were purchased from Sigma-Aldrich Chemie, Fluka Chemie
153AG (Buchs, Switzerland), Panreac Química S.L.U. (Castellar del Vallés, Catalonia, Spain), and
154Merck KGaA (Darmstadt, Germany). Potassium silicate (PSi), as the commercial product Sil-Ma-
155trix[®] (29% PSi) was purchased from PQ Corporation (Valley Forge, PA, USA).

156

1572.3. In vitro antifungal activity of GRAS salts

158 The effect of GRAS salts on radial mycelial growth of *L. theobromae* was determined in Petri
159dishes with PDA medium as described by Karaca et al. (2014) for evaluation of the activity against
160the stone fruit pathogen *Monilinia fructicola* (G. Winter) Honey. Briefly, PDA was amended at 40-
16150 °C with sterile aqueous solutions of each salt at concentrations of 0.2, 1.0, and 2.0% (v/v) (0.01,
1620.05, and 0.1% in the case of paraben salts). PDA plates without salt were used as control dishes.
163One plug of a 7-14 d-old culture of *L. theobromae* was inoculated in the center of each PDA dish
164and incubated for up to 14 d at 25 °C. Radial fungal growth was measured every 2-3 d and results
165expressed as percentage of growth inhibition with respect to control dishes. For each salt and salt
166concentration, 4 replicates (4 PDA dishes) were used and each combination was tested twice.

167

1682.4. Preparation of antifungal coatings

169 The hydrocolloid of the composite coating matrixes, HPMC (Methocel E15), was purchased
170from Dow Chemical Co. (Midland, MI, USA) and the lipid of the matrixes, beeswax (BW) (grade
1711), was supplied by Fomesa Fruitech S.L. (Beniparrell, Valencia, Spain). Stearic acid and glycerol

172 were purchased from Panreac Química S.L.U. HPMC-BW composite edible emulsions were
173 prepared combining the hydrophilic phase (HPMC) with the hydrophobic phase (BW) suspended in
174 water. Glycerol and stearic acid were used as plasticizer and emulsifier, respectively. All the
175 emulsions contained 1.3% HPMC (w/w, wet basis, wb) and 3% BW (wb). HPMC-glycerol (2:1)
176 and BW-stearic acid (3:1) ratios were kept constant for all coatings. The concentrations of GRAS
177 salts in the formulations varied between 0.01 and 2.0% (wb) and were determined according to the
178 effective doses previously obtained in the in vitro tests. For emulsion preparation, an aqueous
179 solution of HPMC (5%, w/w) was prepared by dispersing the HPMC in hot water at 90 °C and later
180 hydration at 20 °C. Water, BW, glycerol, and stearic acid (Tween 80 in the case of emulsions with
181 sodium propionate) were added to the HPMC solution and heated at 98 °C to melt the lipids.
182 Samples were homogenized with a high-shear probe mixer (Ultra-Turrax model T25, IKA-Werke,
183 Steufen, Germany) for 1 min at 12,000 rpm and 3 min at 22,000 rpm. After adding the
184 corresponding salts at the indicated amounts, emulsions were cooled under agitation to a
185 temperature lower than 25 °C by placing them in an ice bath and agitation was continued for 25 min
186 to ensure complete hydration of the HPMC. Viscosity of the emulsions was determined with a
187 viscosimeter (Visco Star Plus R, Fungilab, S.A., Barcelona, Spain) and pH values using a pH-meter
188 (Consort C830 multi-parameter analyzer, Turnhout, Belgium). The formulations were tested for
189 stability according to the method described by Valencia-Chamorro et al. (2008). In brief, the
190 emulsions were placed in volumetric tubes and phase separation was assessed after 24 h at 25 °C.
191 Emulsions were kept overnight at 10 °C before use in the experiments.

192

193 2.5. Fruit

194 Hybrid mandarins [*C. reticulata* × (*C. sinensis* × *C. reticulata*) cv. ‘Ortanique’; synonym:
195 ‘Topaz’] and oranges (*Citrus sinensis* L. cv. ‘Barnfield’; Navel group) were used for in vivo
196 experiments and fruit quality assessments. Fruit were harvested in commercial orchards in the
197 Valencia area and no postharvest treatments were applied. Fruit with no wounds, bruises, or any

198external damage were selected, washed, disinfected by immersion in diluted commercial bleach
199(0.5% NaClO), rinsed thoroughly with tap water, allowed to dry at room temperature, and placed in
200plastic trays in carton boxes to be used in the experiments the next day.

201

2022.6. In vivo curative activity of antifungal coatings

203 For fruit inoculation, a circular wound (1-2 mm deep, 5 mm in diameter) was inflicted with a
204sterile cork borer in the stem-end of each fruit and a PDA plug of *L. theobromae* culture (5 mm in
205diameter) was deposited on the wound (the mycelium growing side in the inner part) and ensured
206with transparent tape to avoid desiccation. To favor infection, inoculated fruit were placed in humid
207chambers and incubated at 28 °C and high RH (>90%) for 48 h. After this period, the tape and the
208mycelium discs were removed using a small sterile spatula, and the fruit were individually coated.
209For coating application, 400 µL of the desired emulsion were pipetted onto each fruit and rubbed
210with gloved hands to simulate the industrial application of waxes and coatings in rotating brushes in
211the packinglines in citrus packinghouses (Bai et al., 2002). After draining and air-drying at room
212temperature, coated fruit were incubated for up to 15 d at 28 °C and 90% RH in a climatic walk-in
213room. Inoculated but uncoated mandarins or oranges served as controls. Each treatment was applied
214to 4 replicates of 10 fruit each and each trial was conducted twice.

215 The incidence (percentage of infected fruit) and severity (lesion diameter in mm) of Diplodia
216stem-end rot were assessed, depending on the experiment, after 3, 7, and 10 d of incubation at 28 °C
217in the case of mandarins and after 4, 8, 12, and 15 d of incubation in the case of oranges. Results
218were reported as disease reduction (%) with respect to the control treatments. The area under the
219disease progress stairs (AUDPS) was also calculated.

220

2212.7. Effect of coating application on fruit quality

222 Among the edible coatings tested for antifungal activity, the three most effective were selected
223to determine their effect on postharvest quality of non-inoculated and cold-stored oranges. HPMC-

224BW coatings containing the following GRAS salts and concentrations were selected: potassium
225sorbate (PS) at 2%, sodium ethylparaben (SEP) at 0.1%, and sodium benzoate (SB) at 2%. Selected
226healthy 'Barnfield' oranges were washed, coated, and stored at 5 °C and 90% RH for 21 or 42 d,
227followed by a shelf-life period of 7 d at 20 °C. Uncoated oranges served as controls. The following
228quality attributes were determined at harvest and after cold storage and shelf life.

229

2302.7.1. *Weight loss*

231Thirty oranges per treatment were individually marked and weighted. After storage and shelf life,
232they were weighted again and the percentage weight loss was expressed with respect to the initial
233weight.

2342.7.2. *Fruit firmness*

235Firmness of 20 oranges per treatment was determined as percentage of rind deformation with an In-
236stron Universal testing machine according to Valencia-Chamorro et al. (2009b).

2372.7.3. *Internal quality*

238Soluble solids concentration (SSC, %), titratable acidity (TA, %), and maturity index (MI, SSC/TA)
239of the juice from three previously weighed samples of 10 oranges each per treatment were deter-
240mined as described by Palou et al. (2007).

2412.7.4. *Internal CO₂ concentration*

242CO₂ concentration (%) in the internal cavity of 10 oranges per treatment was determined by gas
243chromatography using the methodology described by Valencia-Chamorro et al. (2009b).

2442.7.5. *Ethanol content (EC) and acetaldehyde content (AC)*

245The content of these volatile compounds (mg/L) in the headspace of juice from three replicates of
24610 oranges per treatment was analyzed by gas chromatography according to Valencia-Chamorro et
247al. (2011b).

2482.7.6. *Sensory evaluation*

249Taste (1-9 scale; 1 = very poor and 9 = optimal) and external appearance (1-3 scale; 1 = bad, 2 = ac-
250ceptable, and 3 = good) of four coated oranges per treatment were evaluated by 8-10 trained judges
251following the procedures described by Valencia-Chamorro et al. (2009b).

252

2532.8. Statistical analysis

254 Data from both in vitro and in vivo efficacy tests and fruit quality assessment were subjected
255to analyses of variance (ANOVAs; Statgraphics 5.1, Manugistics, Inc., Rockville, MD, USA). Since
256the experiment was not a significant factor, means of repeated experiments are presented. Data on
257percent inhibition of mycelial growth was subjected to one-way ANOVA with the concentration of
258the different GRAS salts as dependent variable. Disease reduction with respect to control fruit was
259calculated as percentage. When appropriate, means separation was performed by Fisher's protected
260least significant difference test (LSD, $P = 0.05$).

261

2623. Results and discussion

2633.1. In vitro antifungal activity of GRAS salts

264 The determination of antifungal activity in this study was based on the inhibition of the radial
265growth of fungal colonies of *L. theobromae* compared to growth on control plates (PDA with no
266salt addition) after 3, 5, and 7 d of incubation at 25 °C (Table 2). Further readings are not reported
267in this table because 7 d was the incubation period after which the pathogen entirely covered the
268control plates; however, after 14 d of incubation, it was observed that the fungal growth remained
269completely inhibited in those plates with 100% inhibition after 7 d.

270 Significant differences between treatments were found and the effect of each salt was clearly
271dependent on the concentration at which it was applied (Table 2). Ammonium bicarbonate (ABC),
272ammonium carbonate (AC), sodium carbonate (SC), and potassium carbonate (PC) were the most
273effective antifungal salts against the pathogen, with complete inhibition of fungal growth after 7 d
274of incubation at all the tested concentrations of 0.2, 1.0, and 2.0% and with no significant differ-

275ences among these concentrations (Table 2). In research work with banana postharvest pathogens,
276SC and sodium bicarbonate (SBC) also effectively reduced the in vitro growth of *L. theobromae*
277(Alvindhia, 2013). In a work similar to the present research, Karaca et al. (2014) also identified some
278of these salts, particularly AC and ABC, as the most effective at all tested concentrations (0.2, 1.0,
279and 2.0%) to inhibit the growth of *M. fructicola* on PDA dishes. These salts, but also SEP, sodium
280methylparaben (SMP), and potassium silicate (PSi) at similar concentrations greatly inhibited the
281growth of the fungi *Botrytis cinerea* Pers. and *Alternaria alternata* (Fr.) Keiss. in in vitro experi-
282ments (Fagundes et al., 2013). These pathogens have a very wide range of fruit hosts, including cit-
283rus fruit. Olivier et al. (1998) reported that ABC, SC, PC, and potassium bicarbonate (PBC) sub-
284stantially reduced the in vitro growth of *Helminthosporium solani* Durieu & Mont., the causal agent
285of potato silver scurf. In the present study, a second group of preservative salts including sodium bi-
286carbonate (SBC), PBC, SB, and PSi were also completely effective against *L. theobromae*, but only
287at the concentrations of 1.0 and 2.0%. SEP also completely inhibited the fungus at the two highest
288concentrations tested of 0.05 and 0.1%. Sodium propionate (SP), potassium sorbate (PS), and SMP
289were significantly more effective only at the highest concentration tested (2.0 or 0.1%). The least
290effective GRAS salt was ammonium phosphate (APh), with which growth inhibition after 7 d was
291lower than 50% at the highest concentration.

292 It is clear from this and previous research that some of these GRAS salts show a broad spec-
293trum antimicrobial activity since they are able to inhibit in vitro a large number of fungal pathogens.
294For instance, besides the examples just mentioned, different carbonate salts have been found effec-
295tive in other reports to inhibit the radial growth of strains of *B. cinerea* (Alaoui et al., 2017; Youssef
296and Roberto, 2014), *Geotrichum citri-aurantii* (Talibi et al., 2011), *Colletotrichum gloeosporioides*
297(Penz.) Penz. & Sacc. (Sivakumar et al., 2002), and *Penicillium expansum* L. (Lai et al., 2015).
298Nevertheless, the toxicity in vitro of a particular salt to different fungal pathogens greatly differs
299and it is influenced by different factors such as the pathogen species and strain, the salt composition
300(the nature of anions and cations) and concentration (typically the toxicity is concentration-depen-

301dent), the pH, the culture medium, and the incubation conditions. The addition of inorganic or or-
302ganic salts to the medium largely modifies its pH and, in general, the toxicity is also pH-dependent
303(Xu and Hang, 1989). However, the pH alone cannot account for the strong antifungal activity of
304these compounds since different salts with the same pH show different toxicity to the same fungal
305strain. Therefore, the salt cation also plays an important role and, in fact, sodium, potassium, and
306ammonium forms of the same salt can show large differences in toxicity to a particular fungal strain
307(Karaca et al., 2014; Palmer et al., 1997). General antifungal mechanisms of action of GRAS salts
308include the alteration of the integrity and permeability of the fungal cell membranes, disturbances in
309the transport of nutrients that eventually cause cell inactivation and death (Lucera et al., 2012), and
310reduction of cellular turgor pressure with collapse and shrinkage of conidia and/or hyphae (Talibi et
311al., 2011).

312

3133.2. In vivo curative activity of antifungal coatings

314 GRAS salts and concentrations to be used as ingredients of HPMC-BW edible coatings were
315selected according to the previous in vitro results and results from preliminary experiments con-
316ducted in the laboratory to evaluate their compatibility with the rest of coating ingredients. APh and
317SP were discarded due to their low toxicity to *L. theobromae* (Table 2). Effective salts such as
318ABC, SBC, SC, and PBC had to be discarded due to incompatibility with the coating matrix that led
319to the occurrence of phase separation, instability of the emulsions, or undesirable characteristics
320such as too high viscosity or presence of apparent salt residues on the surface of treated mandarins
321or oranges. Effective salts that formed stable emulsions with appropriate characteristics were used
322at their minimum effective concentration. Therefore, the selected salts and concentrations were the
323following: AC (0.2%), PS (2%), PC (0.2%), SMP (0.1%), SEP (0.1%), SB (2%), and PSi (2%).
324Composite coatings formulated with these GRAS salts all contained similar HPMC and BW
325amounts and a total solid content in the range of 6.1-8% depending on salt concentration. The most
326important properties of the emulsion formulations, pH and viscosity, are presented in Table 3.

327 In a first set of in vivo experiments with ‘Ortanique’ mandarins, fruit artificially inoculated
328 with *L. theobromae* were coated 24 h later with these HPMC-lipid composite coatings containing
329 the selected GRAS salts and incubated for 10 d at 28 °C and 90% RH. The effect of coating appli-
330 cation on the inhibition of *Diplodia* stem-end rot is shown in Fig.1. Due to the use of mycelium
331 plugs as artificial inoculation method, disease incidence (percentage of infected fruit) was 100% on
332 all inoculated fruit (data not shown) and the variables percent reduction of disease severity (reduc-
333 tion of lesion size with respect to control fruit) and AUDPS were used as means to assess the dis-
334 ease control ability of the coatings. Although all the tested antifungal coatings reduced somewhat
335 the severity of stem-end rot in comparison with uncoated controls, disease reduction was only im-
336 portant with coatings containing PS (about 50% reduction) and SB (about 40% reduction). Coatings
337 formulated with SEP and PSi reduced disease severity about 20-25% and the rest only 10% or less
338 (Fig. 1).

339 AUDPS on artificially inoculated and coated ‘Ortanique’ mandarins was determined with
340 readings of lesion diameter after 3, 7, and 10 d of incubation at 28 °C and 90% RH. Data showed
341 that the emulsion of HPMC-BW had no antifungal activity by itself since AUDPS was the same
342 (about 190) on fruit treated with this coating (without salt addition) than on uncoated control fruit.
343 In contrast, all the coatings formulated with GRAS salts significantly reduced AUDPS if compared
344 with the control ($P < 0.05$), although important reductions were only obtained with coatings contain-
345 ing PS at 2% (from 190 to 100) and SB at 2% (from 190 to 120). Coatings with SEP at 0.1% re-
346 duced AUDPS from 190 to about 150 (Fig. 1).

347 These three coatings at these concentrations were therefore selected to be tested again in a
348 confirmation experiment with ‘Barnfield’ oranges artificially inoculated with *L. theobromae*. In this
349 case, treated fruit were incubated for 15 d at 28 °C and 90% RH and readings of lesion size were
350 performed after 4, 8, 12, and 15 d of incubation. Similarly to the previous results with mandarins,
351 all coatings significantly reduced *Diplodia* stem-end rot severity on oranges, being the coatings con-
352 taining 2% PS or 2% SB significantly more effective (about 60% reduction) than the coating con-

353taining 0.1% SEP (about 30% reduction) ($P<0.05$) (Fig. 2). AUDPS results were in accordance with
354these observations, although in this case the AUDPS value on oranges treated with PS-coatings was
355significantly lower than on fruit treated with SB- or SEP-coatings.

356 To our knowledge, this is the first research work in which edible coatings with GRAS salts are
357applied to citrus fruit for the control of *Diplodia* stem-end rot. Likewise, previous references to the
358use for this purpose of coatings or waxes non-amended with conventional fungicides are really
359scarce. Early work by Waks et al. (1985) showed that while the application of citrus commercial
360waxes reduced the incidence of green and blue molds during cold storage, it increased that of stem-
361end rots caused by *L. theobromae* or *Alternaria citri* Ellis & N. Pierce. They discussed that the
362cause could be changes in the internal atmosphere of the fruit induced by waxing, particularly an in-
363crement of EC. A coating comprised of glycolchitosan and the biocontrol yeast *Candida saitoana*
364reduced the incidence of stem-end rot on semi-commercial trials with ‘Valencia’ oranges (El-
365Ghaouth et al., 2000). Recently, Xu et al. (2018) reported that chitosan and chitosan/montmoril-
366lonite coatings significantly reduced the accumulated decay rate of cold-stored tangerines, but they
367did not identify the actual pathogens causing decay. Nevertheless, some information is available on
368the postharvest use of coatings to control *Diplodia* stem-end rot on other fruit commodities. For in-
369stance, decay caused by *L. theobromae* on avocado was significantly reduced by a commercial coat-
370ing amended with essential oil from the plant *Lipia scaberrima* (Regnier et al., 2010) or by a pectin-
371based edible coating (Maftoonazad et al., 2007). The disease was also reduced on banana by coat-
372ings comprised of chitosan and cinnamon extract (Win et al., 2007), on mango by chitosan coatings
373formulated with the active antimicrobial lactoperoxidase system (Cissé et al., 2015), and on longan
374by chitosan coatings formulated with the GRAS salt PS (Apai et al., 2008). The noteworthy activity
375of PS is in agreement with the present results obtained with citrus fruit. The compound ethylparaben
376produced by the biocontrol agent *Brevibacillus brevis* was identified in recent work as effective
377against *L. theobromae* and mutants of the antagonistic bacteria producing more ethylparaben were
378promising for the biocontrol of *Diplodia* stem-end rot on apple fruit (Che et al., 2018).

379 The type of HPMC-BW antifungal edible coatings with GRAS salts tested here have been al-
380ready evaluated for the control of other important postharvest diseases of citrus fruit. Particularly,
381Valencia-Chamorro et al. (2008, 2009a) found that, among a large variety of GRAS salts tested,
382HPMC-BW coatings containing the salts PS, SB, SP, and their mixtures exhibited antifungal activ-
383ity in vitro against the citrus pathogens *P. digitatum* and *P. italicum* and were the most effective for
384green and blue mold reduction on ‘Valencia’ oranges and ‘Ortanique’ and ‘Clemenules’ mandarins
385artificially inoculated with these pathogens, coated 24 h later, and incubated at 20°C for 7 d to simu-
386late fruit shelf life. The curative activity of similar composite coatings has also been observed in
387other fresh fruit pathosystems. Fagundes et al. (2013, 2015) found significant reductions of black
388spot on cherry tomatoes artificially inoculated with *A. alternata* and coated 24 h later with HPMC-
389BW coatings formulated with SB, SEP, or SMP. Karaca et al. (2014) reported that HPMC-BW
390coatings containing PS, SEP, or SMP, among other salts, effectively reduced the incidence and
391severity of brown rot caused by *M. fructicola* on artificially inoculated plums. It seems clear from
392this research that the selection of the most appropriate antimicrobial agent in general and GRAS salt
393in particular to confer disease control ability to a coating is greatly dependent on each particular
394pathosystem (type and properties of the stored fruit and activity of the agent against the target
395pathogen), but also on issues such as the good interaction between the composite matrix and the an-
396tifungal salt, the matrix capacity to gradually release the salt during storage, the presence of other
397additives in the emulsion, and the environmental conditions during storage and shelf life (Valdés et
398al., 2017; Valencia-Chamorro et al., 2011). Emulsion attributes such as pH and viscosity (Table 3)
399clearly influence these properties and are dependent not only on the coating matrix (HPMC-BW)
400and the GRAS salt and their respective proportions, but also on the nature and amount of the other
401components of the coating (i.e., stearic acid and glycerol). The importance of the role of the coated
402fruit host (species and even cultivar) and the environmental conditions on the disease control ability
403of antifungal coatings may explain why results from in vivo experiments cannot often be anticipated
404by the in vitro antifungal activity of the GRAS salt (Fagundes et al., 2013; Karaka et al., 2014).

405 Tests in Petri dishes allow fully exposure of the fungal structures to the antifungal salt, while in in
406 vivo tests with coatings the contact can be limited depending on the emulsion properties, the charac-
407 teristics of the fruit peel, and the storage conditions. Therefore, it is important to tailor the formula-
408 tion of appropriate coatings for particular fruit species and cultivars and even specific postharvest
409 applications. On the other hand, the application of antifungal compounds such as GRAS salts as
410 coating ingredients can be advantageous if compared to their postharvest application in citrus pack-
411 inghouses as aqueous solutions in drench, dip or spray systems. Coatings could be designed to slow
412 down the diffusion of the active ingredient from the matrix to the commodity, which could contrib-
413 ute to maintain for longer periods of time the effective residue of the antifungal on the fruit surface
414 (Vargas et al., 2008). In addition, coating application could reduce in some cases phytotoxicity risks
415 or alleviate potential adverse effects on fruit quality associated with aqueous applications (Palou et
416 al., 2015).

417

418 3.3. Effect of coating application on fruit quality

419 HPMC-BW composite coatings containing PS at 2%, SB at 2%, and SEP at 0.1% were se-
420 lected for fruit quality evaluation because they showed the highest control ability in in vivo trials.
421 Weight loss of non-inoculated ‘Barnfield’ oranges uncoated (control) or coated and cold-stored at 5
422 °C and 90% RH followed by shelf life at 20 °C is shown in Fig. 3. As expected, weight loss was
423 higher after 42 d (about 3-3.5%) than after 21 d (2-2.5%) of cold storage followed by 7 d of shelf
424 life. After both storage periods, weight loss reduction on coated oranges was not high, but coatings
425 containing SEP significantly reduced weight loss with respect to uncoated control fruit ($P < 0.05$).
426 Coatings containing SB and PS slightly influenced weight loss (positively and negatively), but not
427 significantly. Working with ‘Valencia’ oranges, Valencia-Chamorro et al. (2009b) also found that
428 HPMC-BW-shellac coatings containing SB were superior to coatings containing PS for weight loss
429 reduction during cold storage at 5 °C. They discussed that PS-coatings exhibited higher water vapor
430 permeability than SB-coatings. Similarly, SB-coatings also reduced weight loss and maintained

431 firmness of 'Clemenules' clementine mandarins without adverse effects on the overall quality of
432 coated fruit (Valencia-Chamorro et al., 2011). Gunaydin et al. (2017) reported similar results work-
433 ing with plums, where HPMC-BW coatings, with or without antifungal agents, significantly re-
434 duced weight loss compared to uncoated control samples, and the coatings containing sodium
435 paraben salts such as SEP were more effective than those containing PS. In general, cellulose-lipid
436 composite coatings are reported to reduce fruit weight loss due to the moisture barrier exerted by
437 the lipid ingredients (BW, shellac, etc.) of the coating formulation (Contreras-Oliva et al., 2011).
438 Similarly to postharvest decay control, their effectiveness to reduce fruit weight loss depends not
439 only on the lipid composition and concentration, but also on the species and cultivars of the fruit.
440 Thus, similar HPMC-lipid coatings effectively reduced weight loss of 'Ortanique' and
441 'Chemenules' mandarins (Valencia-Chamorro et al., 2010, 2011b) but not of 'Valencia' oranges (Va-
442 lencia-Chamorro et al., 2009b).

443 Table 4 shows the external and internal physicochemical quality attributes of 'Barnfield' or-
444 anges at harvest and after treatment with HPMC-BW coatings containing the selected antifungal
445 GRAS salts. Firmness, SSC, TA, EC, and AC were evaluated in fruit stored at 5 °C for 21 or 42 d
446 followed by a shelf-life period of 7 d at 20 °C. After 21 d, rind deformation (expressing fruit firm-
447 ness) of all coated samples was in the range 1.0-1.5% and fruit coated with emulsions formulated
448 with 2% SB were less firm (higher percentage of deformation) than fruit treated with the other coat-
449 ings. After 42 d, oranges treated with SB- and PS-coatings were less firm than control fruit and fruit
450 treated with SEP-coatings ($P < 0.05$), although in all cases the percent deformation was low, in the
451 range of 2-3% (Table 4). According to previous results with HPMC-BW coatings, it seems that the
452 citrus cultivar plays an important role on the effect of coating on fruit firmness. For instance, while
453 HPMC-BW coatings amended with SB, PS, or mixtures did not affect the firmness of coated 'Va-
454 lencia' oranges (Valencia-Chamorro et al., 2009b), they significantly increased the firmness of
455 coated 'Clemenules' mandarins (Valencia-Chamorro et al., 2011), showing that the response of
456 coated fruit is not only dependent on the coating characteristics but also on the fruit inherent at-

457tributes. Polysaccharides such as pectin, starch, and hemicellulose, present in the cell wall, are im-
458portant for the maintenance of fruit firmness and the degradation of these compounds by hydrolyz-
459ing enzymes, such as pectin methylesterase and polygalacturonase, causes softening of the fruit dur-
460ing ripening and storage. Edible coatings can have the capacity to modify the internal gas composi-
461tion of the fruit in terms of O₂ and CO₂ concentrations, which might influence the activities of the
462cell wall degrading enzymes, reducing fruit softening (Gunaydin et al. 2017). Furthermore, there is
463usually a correlation between the effect of coatings on fruit firmness and weight loss, although this
464aspect is often dependent on the citrus cultivar. For instance, while, in accordance with the present
465results, no correlation was observed in the studies by Pérez-Gago et al. (2002) with ‘Fortune’ man-
466darins, a positive correlation was reported by Navarro-Tarazaga et al. (2008) for ‘Ortanique’ man-
467darins.

468 Regarding juice quality, coated oranges, irrespective of the coating used and the storage pe-
469riod, were less sweet than control fruit ($P<0.05$ for SSC parameter) and no significant differences in
470TA were observed between control and coated fruit (Table 4). The lower SCC of coated oranges
471could be related to changes in fruit respiration and gas exchange patterns induced by coating appli-
472cation. The concentration of internal CO₂ in ‘Barnfield’ oranges uncoated or coated with HPMC-
473BW coatings containing the selected salts is showed in Fig 4. All three coatings modified the inter-
474nal atmosphere of coated oranges, especially after 42 d at 5 °C, and internal CO₂ levels were signifi-
475cantly higher in coated fruit, which indicates that the coatings were effective as gas barriers
476($P<0.05$). After 21 d of cold storage, but not after 42 d, CO₂ levels were lower in oranges treated
477with the coating containing SB than in those coated with the coating formulated with PS ($P<0.05$).
478In this study, internal CO₂ values in coated oranges (4-6 kPa) were equivalent to those observed in
479coated ‘Valencia’ oranges (Valencia-Chamorro et al., 2009b) and ‘Clemenules’ clementines (Valen-
480cia-Chamorro et al., 2011b), but lower than those observed in ‘Ortanique’ mandarins coated with
481similar HPMC-lipid coatings containing GRAS salts (6-8 kPa) (Valencia-Chamorro et al., 2010),
482and in ‘Valencia’ oranges (Navarro-Tarazaga et al., 2007) and ‘Fortune’ mandarins (Pérez-Gago et

483al., 2002) coated with HPMC-lipid coatings without GRAS salts. Factors such as fruit peel mor-
484phology and physical properties of the coating, including the addition to the emulsion of GRAS
485salts or other food additives, should be considered as factors influencing fruit respiration when coat-
486ings are applied to citrus fruit. These factors could influence the coating flexibility or its capacity of
487adaptation to the fruit surface, affecting the gas barrier of the coating. The effect of edible coatings
488on delaying changes related to fruit ripening, such as softening, color change, decrease in acidity,
489and some physiological disorders has been associated with the gas barrier exerted on the fruit sur-
490face leading to reductions in respiration rate and/or weight loss (Fagundes et al., 2015; Valero et al.,
4912013). In the work conducted by Gunaydin et al. (2017), application of HPMC-BW matrixes con-
492taining paraben salts resulted in the lowest CO₂ production rates, showing the potential of these
493coatings as gas barriers on plums. In contrast, Valencia-Chamorro et al. (2008) reported an increase
494in O₂ permeability of HPMC-BW-shellac edible films amended with SEP and Fagundes et al.
495(2015), working with similar coatings amended with a variety of antifungal agents, observed the
496highest respiration rates in cherry tomatoes coated with emulsions containing SEP. This confirms
497that the capacity of an edible coating to create an effective gas barrier depends not only on the coat-
498ing composition and properties, but also on the commodity, cultivar, and storage conditions (Gunay-
499din et al., 2017).

500 In general, after both storage periods of 21 and 42 d, volatile content in ‘Barnfield’ oranges
501was not affected by coating application and ranged 200-600 mg/L of ethanol and 6-11 mg/L of ac-
502etaldehyde, in spite of the higher internal CO₂ concentration in coated than in uncoated fruit (Table
5034, Fig. 4). This was not an anticipated result since in previous studies fruit coating increased EC in
504the juice of coated oranges and mandarins cold-stored for long periods, indicating the creation of a
505modified atmosphere within the fruit (Pérez-Gago et al., 2002; Valencia-Chamorro et al., 2009b,
5062011). However, differences in the citrus cultivar, in the composition of the HPMC-BW coatings
507used in these studies (presence of shellac or different antifungal ingredients and concentrations),
508and in the total solid content and viscosity may explain this different behavior. In contrast to

509 ethanol, it is more frequent that the levels of acetaldehyde on coated and cold-stored citrus fruit do
510 not differ significantly from those in uncoated control fruit (Valencia-Chamorro et al., 2009b,
511 2010).

512 Results from the sensory evaluation of 'Barnfield' oranges treated with HPMC-BW edible
513 coatings containing antifungal GRAS salts and stored at 5 °C followed by 7 d of shelf life at 20 °C
514 are showed in Table 5. The evaluation was based on the assessment of fruit flavor and coating ap-
515 pearance by a trained panel of 8-10 members. Fruit flavor, evaluated in a 1-9 scale, was very
516 slightly modified by the application of HPMC-BW based coatings containing salts. After the first
517 storage period of 21 d at 5 °C, no significant differences were observed among control and coated
518 oranges, while after 42 d, the coating containing PS was rated as with the poorest flavor, although
519 with no significant differences with control fruit. It is well known that citrus off-flavor is due to the
520 accumulation of volatile components associated to anaerobic fermentation (Ke and Kader, 1990).
521 Among the different volatiles that may be present, ethanol has been found to be the component un-
522 dergoing the greatest change occurring in citrus during storage and the application of fruit coatings
523 can lead to enhanced anaerobic respiration as they restrict gas exchange through the rind surface
524 (Teitel et al., 2011). Prior research showed that off-flavor production from modified EC in citrus is
525 cultivar dependent. In general, mandarins are more sensitive to anaerobic conditions than other cit-
526 rus fruit, which has been attributed to differences in enzymatic activity and in peel permeability to
527 gases (Shi et al., 2007). Thus, for example, minimum EC associated with off-flavor has been re-
528 ported to be 2,000 mg/L in 'Valencia' oranges (Ke and Kader et al., 1990), whereas EC of 1,000
529 mg/L and 500-600 mg/L have been reported in 'Clemenules' (Navarro-Tarazaga and Pérez-Gago,
530 2006) and 'Murcott' mandarins (Shi et al., 2005), respectively. In this work, EC levels were lower
531 than those reported by other authors to induce off-flavor in oranges and in some mandarin cultivars
532 (Table 4), which can explain the sensory scores.

533 Coating appearance in a 1-3 scale was evaluated according to the presence or absence of
534 cracks, blemishes, stains, and homogeneity of the coating. In general, the appearance of coated or-

535anges was not optimal because scores after shelf life were 1-2, while they were 3 for uncoated con-
536trol oranges. Coatings containing SB at 2% were the worst qualified in terms of external appearance
537($P<0.05$). In general, HPMC-BW emulsion coatings are not characterized for providing significant
538gloss to coated fruit such as citrus and tomatoes, mainly due to the macro emulsion character of the
539coating formulation (Fagundes et al., 2015; Valencia-Chamorro et al., 2009, 2011). Furthermore,
540Valencia-Chamorro et al. (2010) also reported the presence of small white spots on the surface of
541coated mandarins that reduced the general good appearance of the samples when the HPMC-based
542coatings were amended with some GRAS salts.

543

544**Conclusion**

545 In this work, GRAS salts and concentrations were selected according to their in vitro antifun-
546gal activity against *L. theobromae* NEU-1, a strain of the pathogen isolated from citrus fruit in Va-
547lencia (Spain). The potential of these salts as ingredients of antifungal HPMC-BW edible coatings
548applied for stem-end rot control and quality preservation of citrus fruit was highlighted. Coatings
549containing PS, SEP, and SB were selected and significantly reduced the severity of Diplodia stem-
550end rot on artificially inoculated ‘Ortanique’ mandarins and ‘Barnfield’ oranges. Application of the
551coatings containing SEP and SB significantly reduced weight loss of coated ‘Barnfield’ oranges. All
552the coatings increased internal CO₂ concentration compared to uncoated oranges, but did not ad-
553versely affect EC and AC in the juice, the flavor, and the production of off-flavors. Further research
554should focus on the improvement of physical characteristics of the coatings to enhance water loss
555control and gloss of coated citrus fruit. Likewise, the combination of these antifungal coatings with
556other alternative nonpolluting control methods to synergistically improve the control of Diplodia
557stem-end rot in citrus packinghouses should also be explored.

558

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567

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744 **Table 1.** Characteristics of antifungal GRAS salts tested in vitro for inhibition of *Lasiodiplodia*745 *theobromae* and in vivo (as coating ingredients) for control of Diplodia stem-end rot.

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GRAS salt	Acronym	Molecular formula	E-code ^a	MW ^b
Ammonium phosphate	APh	NH ₄ H ₂ PO ₄	E-342 (i)	149.09
Ammonium bicarbonate	ABC	NH ₄ HCO ₃	E-503 (ii)	79.06
Ammonium carbonate	AC	(NH ₄) ₂ CO ₃	E-503 (i)	114.10
Potassium bicarbonate	PBC	KHCO ₃	E-501 (ii)	100.12
Potassium carbonate	PC	K ₂ CO ₃	E-501 (i)	138.21
Potassium silicate	PSi	K ₂ SiO ₃	E-560	154.26
Potassium sorbate	PS	C ₆ H ₇ O ₂ K	E-202	150.22
Sodium bicarbonate	SBC	NaHCO ₃	E-500 (ii)	84.01
Sodium benzoate	SB	C ₇ H ₅ O ₂ Na	E-211	144.11
Sodium carbonate	SC	Na ₂ CO ₃	E-500 (i)	105.99
Sodium ethylparaben	SEP	C ₉ H ₉ Na O ₃	E-215	188.16
Sodium methylparaben	SMP	C ₈ H ₇ Na O ₃	E-219	174.13
Sodium propionate	SP	CH ₃ CH ₂ COONa	E-281	96.06

747 ^a Code number for food additives approved by the European Union.748 ^b Molecular weight (g/mol).

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752 **Table 2.** Percentage inhibition of radial growth of *Lasiodiplodia theobromae* on PDA Petri dishes

753 amended with different concentrations of GRAS salts after 3, 5 and 7 d of incubation at 25 °C.

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GRAS salt	Inhibition of <i>L. theobromae</i> (%) ^a			
	Concentration (%)	Day 3	Day 5	Day 7
Ammonium phosphate	0.2	1.27h	0j	3.15j
	1	5.93h	0j	0j
	2	73.73de	56.08gh	48.06gh
Ammonium bicarbonate	0.2	100a	100a	100a
	1	100a	100a	100a
	2	100a	100a	100a
Ammonium carbonate	0.2	100a	100a	100a
	1	100a	100a	100a
	2	100a	100a	100a
Sodium bicarbonate	0.2	85.59bcd	79.47cde	72.6de
	1	100a	100a	100a
	2	100a	100a	100a
Sodium benzoate	0.2	73.73de	69.21ef	59.49f
	1	100a	100a	100a
	2	100a	100a	100a
Sodium carbonate	0.2	100a	100a	100a
	1	100a	100a	100a
	2	100a	100a	100a
Sodium propionate	0.2	52.54fg	54.89gh	47.22h
	1	77.11cd	74.22def	71.76e
	2	89.83ab	89.02abc	83.53bc
Sodium methylparaben	0.01	51.69fg	49.16hi	44.2h
	0.05	97.03ab	85.91bc	82.52bcd
	0.1	100a	100a	100a
Sodium ethylparaben	0.01	41.95g	38.9i	29.07i
	0.05	100a	100a	100a
	0.1	100a	100a	100a
Potassium sorbate	0.2	88.56abc	78.76cde	72.77de
	1	100a	94.03ab	90.75ab
	2	100a	100a	100a

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	0.2	100a	100a	100a
Potassium carbonate	1	100a	100a	100a
	2	100a	100a	100a
	0.2	61.44ef	62.77fg	57.48fg
Potassium silicate	1	100a	100a	100a
	2	100a	100a	100a
	0.2	86.44bc	81.62cd	75.63cde
Potassium bicarbonate	1	100a	100a	100a
	2	100a	100a	100a

756^a Colony diameter reduction with respect to control treatments (non-amended PDA plates).

757 Means in columns with different letters are significantly different by Fisher's protected

758 LSD test ($P < 0.05$) applied after an ANOVA.

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761 **Table 3.** pH and viscosity of edible coatings formulated with HPMC-BW and GRAS salts.

Coating with GRAS salt	pH	Viscosity (cp)
No salt (HPMC-BW)	5.76	45.2
Ammonium carbonate 0.2%	6.83	46.2
Potassium sorbate 2%	6.27	51.2
Potassium carbonate 0.2%	7.15	50.0
Sodium methylparaben 0.1%	7.15	46.7
Sodium ethylparaben 0.1%	7.03	45.9
Sodium benzoate 2%	6.07	46.7
Potassium silicate 2%	9.50	60.0

762 HPMC, hydroxypropyl methylcellulose; BW, beeswax

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766**Table 4.** Quality attributes of ‘Barnfield’ oranges coated with HPMC-BW composite edible coatings containing antifungal

767GRAS salts, stored at 5 °C followed by 7 d of shelf life at 20 °C.

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Coating with GRAS salt	21 d 5 °C + 7 d 20 °C					42 d 5 °C + 7 d 20 °C				
	Firmness (% deformation)	SSC (%)	TA (% citric acid)	EC (mg/L)	AC (mg/L)	Firmness (% deformation)	SSC (%)	TA (% citric acid)	EC (mg/L)	AC (mg/L)
At harvest	1.12	12.32	0.694	106.0	5.43					
Control	1.39ab	13.91a	0.636ab	434.2a	7.951ab	2.16c	13.9a	0.680ab	458.3a	8.852a
PS 2%	1.25b	10.38d	0.724a	382.5ab	8.838a	2.74ab	12.45b	0.706a	384.1a	10.813a
SEP 0.1%	1.22b	10.93c	0.721a	388.1ab	9.155a	2.33bc	12.35b	0.479b	448.2a	9.990a
SB 2%	1.49a	13.15b	0.574b	271.9b	6.013b	2.80a	11.91b	0.581ab	609.1a	10.532a

769 HPMC, hydroxypropyl methylcellulose; BW, beeswax; PS, potassium sorbate; SEP, sodium ethylparaben; SB, sodium benzoate.

770 SSC, soluble solids content; TA, titratable acidity; EC, ethanol content; AC, acetaldehyde content.

771 Columns with different letters are significantly different according to Fisher's protected LSD test ($P < 0.05$) applied after an ANOVA.

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 774 **Table 5.** Sensory evaluation of flavor and coating appearance of ‘Barnfield’ oranges coated with
 775 HPMC-BW edible composite coatings containing antifungal GRAS salts, stored at 5 °C followed by
 776 7 d of shelf life at 20 °C.

777

GRAS salts	21 d 5 °C + 7 d 20 °C		42 d 5 °C + 7 d 20 °C	
	Flavor (1-9 scale)	Coating appearance (1-3 scale)	Flavor (1-9 scale)	Coating appearance (1-3 scale)
Control	6.90a	3.00a	6.63ab	3.00a
PS 2%	7.00a	1.29c	5.90b	1.36b
SEP	6.72a	1.86b	6.09ab	1.63b
SB 2%	6.81a	1.00d	7.00a	1.00c

778 HPMC, hydroxypropyl methylcellulose; BW, beeswax.

779 PS, potassium sorbate; SEP, sodium ethylparaben; SB, sodium benzoate.

780 Columns with different letters are significantly different according to Fisher’s protected LSD test

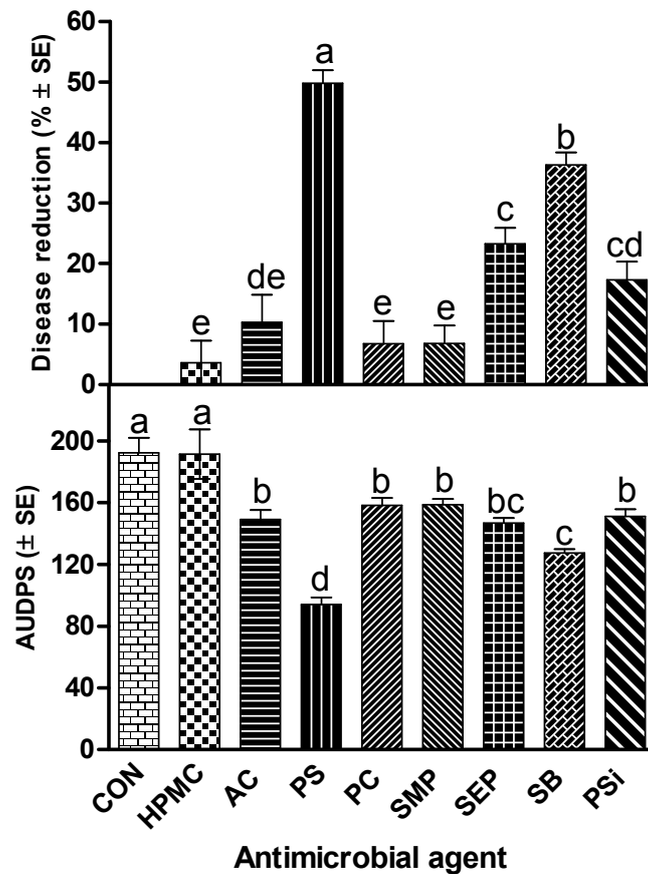
781 ($P < 0.05$) applied after an ANOVA.

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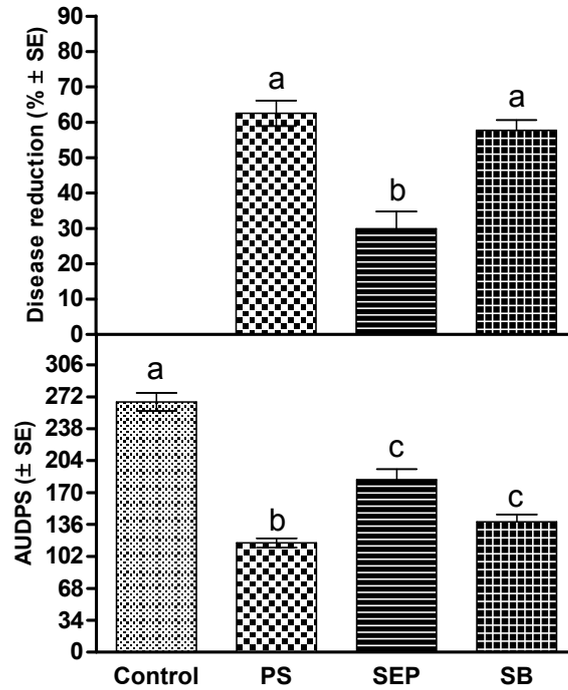


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790**Fig. 1.** Reduction of the severity with respect to control fruit (inoculated but uncoated) and area
791under the disease progress stairs (AUDPS) of stem-end rot on mandarins ‘Ortanique’ artificially
792inoculated with *Lasiodiplodia theobromae*, coated 24 h later with HPMC-BW composite edible
793coatings containing GRAS salts, and incubated for 10 d at 28 °C and 90% RH. GRAS salts and
794concentrations are those indicated in Table 3. AUDPS was determined with readings of lesion
795diameter after 3, 7, and 10 d of incubation. Columns with different letters are significantly different
796according to Fisher’s protected LSD test ($P<0.05$) applied after an ANOVA

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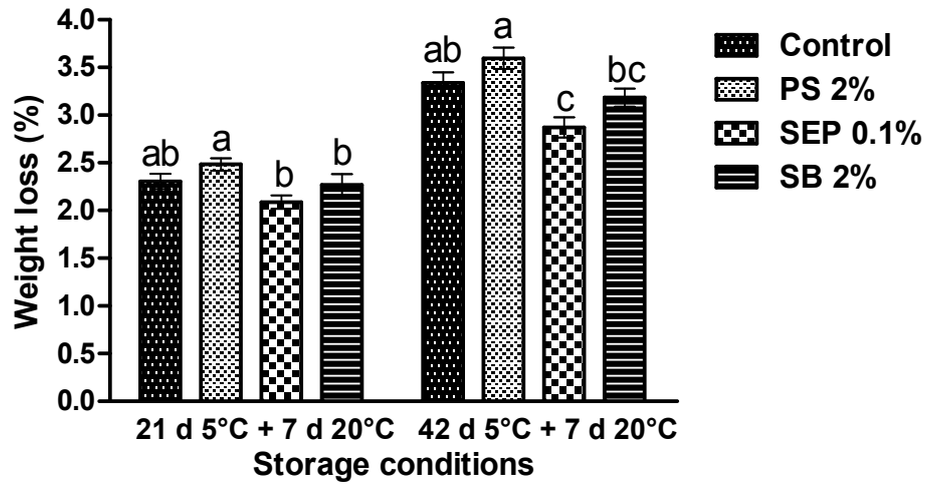


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802**Fig. 2.** Reduction of the severity with respect to control fruit (inoculated but uncoated) and area un-
803der the disease progress stairs (AUDPS) of stem-end rot on ‘Barnfield’ oranges artificially inocu-
804lated with *Lasiodiplodia theobromae*, coated 24 h later with HPMC-BW composite edible coatings
805containing GRAS salts, and incubated for 15 d at 28 °C and 90% RH. Coatings were formulated
806with 2% potassium sorbate (PS), 0.1% sodium ethylparaben (SEP), or 2% sodium benzoate (SB).
807AUDPS was determined with readings of lesion diameter after 4, 8, 12, and 15 d of incubation. Col-
808umns with different letters are significantly different according to Fisher’s protected LSD test
809($P < 0.05$) applied after an ANOVA.

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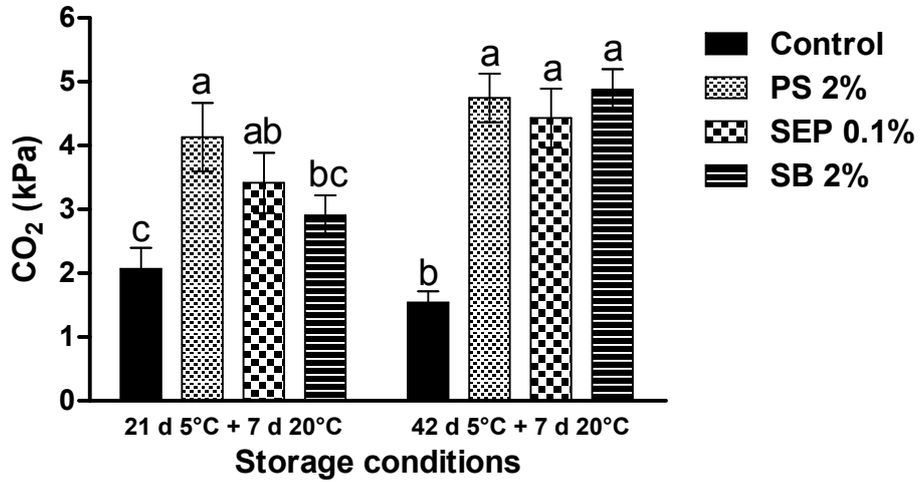
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815**Fig. 3.** Weight loss of 'Barnfield' oranges uncoated (control) or coated with HPMC-BW composite
816edible coatings containing GRAS salts at the indicated concentrations and stored for the indicated
817periods: Potassium sorbate (PS), sodium ethylparaben (SEP), sodium benzoate (SB). For each stor-
818age period, columns with different letters are significantly different according to Fisher's protected
819LSD test ($P < 0.05$) applied after an ANOVA.

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826**Fig. 4.** Concentration of internal CO₂ in 'Barnfield' oranges uncoated (control) or coated with
827HPMC-BW composite edible coatings containing GRAS salts at the indicated concentrations and
828stored for the indicated periods: Potassium sorbate (PS), sodium ethylparaben (SEP), sodium ben-
829zoate (SB). Columns with different letters are significantly different according to Fisher's protected
830LSD test ($P < 0.05$) applied after an ANOVA.

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