

1	Effect of different chemical compounds as coadjutants of 4-hexylresorcinol on
2	appearance of deepwater pink shrimp (Parapenaeus longirostris) during chilled
3	storage
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9	ABBREVIATED RUNNING TITLE: Appearance of pink shrimp immersed in 4-
10	hexylresorcinol-based formulas
11	
12	ABSTRACT
13	Different chemical compounds (kojic acid, cumic acid phytic acid, sodium metabisulphite,
14	magnesium carbonate, sorbic acid, and different protease inhibitors) were used as
15	coadjutants in 4-hexylresorcinol-based melanosis-inhibiting formulas tested by immersion
16	on pink shrimp (Parapenaeus longirostris). Increasing concentrations of 4-hexylresorcinol
17	delayed the occurrence of melanosis during storage. However, they could not prevent the
18	appearance of yellow-greenish colouration in the cephalothorax, which diminished the
19	consumer acceptability of shrimps. The incorporation of protease inhibitors (EDTA,
20	disodium dihydrogen pyrophosphate, iodoacetic acid, egg white and PMSF) into the 4-
21	hexylresorcinol-based blends improved acceptability through storage, suggesting protease
22	activity post mortem contributes to the final acceptability of crustaceans.
23	Keywords: melanosis, pink shrimp, Parapenaeus longirostris, polyphenol oxidase, 4-
24	hexylresorcinol.

25 INTRODUCTION

26 Melanosis is a surface colouration that initiates as soon as crustaceans are removed from 27 water and come into contact with the oxygen in the atmosphere. The reaction involves the 28 oxidation of phenols to quinones by polyphenol oxidase (PPO), providing the precursors of 29 insoluble polymeric pigments (Otwell & Marshall, 1986). Currently, prevention of black 30 spots involves application of sulphite derivatives, although the authorised residual content 31 in flesh is not very effective to delay the appearance of melanosis in some species such as 32 pink shrimp (Parapenaeus longirostris). The use of high doses of sulphites is not a good 33 alternative given that it is related to allergic reactions, as well as other health risks 34 associated with its use (Taylor & Bush, 1987). It emphasizes the need for a safe effective 35 sulphite alternative. 4-hexylresorcinol (4-HR), is for example a competitive inhibitor of 36 PPO "generally recognized as safe" (GRAS) for use in the prevention of crustacean 37 melanosis (Frankos et al., 1991). The use of 4-HR is permitted in the US, Canada, 38 Australia, and some Latin American countries (Montero, Martínez-Alvarez & Gómez-39 Guillén, 2004). In 2003, the Scientific Committee on Food of the European Commission 40 considered 4-HR to be toxicologically acceptable for the prevention of melanosis in 41 shrimps under specific conditions of use (provided residues in crustacean meat do not 42 exceed 2 mg/kg). In 2004, a proposal of amending the EU Directive 95/2/EC on food 43 additives other than colorants and sweeteners was adopted by the European Commission 44 and presented to the European Parliament in order to authorize 4-HR as an alternative to sulphites for preventing the browning of crustaceans. Its effectiveness as melanosis-45 46 inhibiting compound has been demonstrated both in laboratory tests and on board 47 (McEvily, Iyengar & Otwell, 1991; Otwell, Iyengar & McEvily, 1992; Guandalini, Joppolo, 48 Mantovani, Stacchini & Giovannini, 1998; Montero, López-Caballero & Pérez-Mateos,

49 2001a; Montero et al., 2004). However, the effective doses are different depending on 50 several factors such as species, physiological states, and application method. In addition, 51 the experimental conditions cannot be held in the same account, as those used in 52 commercial boats, where fishermen manipulate a considerable amount of shrimp and 53 accurate weight measurement is difficult. That results in amount of melanosis-inhibiting 54 formulations used on board not corresponding to those used in the laboratory.

55 A commercial formulation composed of 4-HR and NaCl (Everfresh®, Opta Food 56 Ingredients) is used in different countries with positive results in delaying melanosis. 57 However, for deepwater pink shrimp (Parapenaeus longirostris) this product has proved to 58 be quite ineffective using the concentrations and application times recommended by the 59 manufacturer (unpublished data). That encourages the search for more potentially effective 60 4-HR-based formulations, mainly for species very susceptible to melanosis such as pink 61 shrimp. Montero et al. (2004) reported the application of citric acid, ascorbic acid and 62 acetic acid in association with 4-HR to enhance the appearance of shrimp, accentuating 63 their natural pink colouration. Despite the fact that this formulation inhibited melanosis 64 itself, the viscera turned yellowish during storage, which is attributed to primary post 65 mortem spoilage. The incorporation of coadjutants in 4-HR based formulations could result 66 in the prevention of the prevalence of yellow shades in the head, thereby improving 67 appearance. These chemicals could be different food-grade compounds for which anti-68 browning action has been demonstrated in vegetables, fruits and crustaceans. Phytic acid is 69 a natural antioxidant (Graf, Empson & Eaton, 1987) and chelating agent which 70 demonstrates melanosis-inhibiting activity, as been described in vegetables (Hicks et al., 71 2004), but rarely in crustaceans (Guang-Li, Yuan-hong, Shu-ging & Xue-lin, 1996). Kojic 72 acid is an antibiotic substance produced by several species of Aspergillus and Penicillium,

73 and is widely used in Japan as a food additive for preventing enzymatic browning. Kojic 74 acid acts by interfering with the uptake of O₂ required for the enzymatic reaction and/or 75 reduction of quinone compounds to diphenols to prevent melanin formation (Chen, Wei, & 76 Marshall, 1991a). Cumic acid, abundant in seeds of *Cuminum cyminum* (cumin), acts by 77 binding to the coupled binuclear copper active site with the carboxylic group and can be 78 classified as a non-competitive inhibitor (Kubo & Kinst-Hori, 1998). Another chemical, 79 magnesium carbonate, is decomposed into carbon dioxide, which interferes with the 80 utilization of oxygen by PPO (Gutierrez Alsina, 1976). Furthermore, several food 81 preservatives, such as potassium sorbate and sulphite derivatives, could also be used to 82 delay the appearance of melanosis mediated by some spoilage bacteria, according to 83 Chinivasagam, Bremner & Reeves, (1998). Other substances show a double role, such as 84 ethylenediaminetetraacetic acid (EDTA), and disodium dihydrogen pyrophosphate (PPi) 85 (Iyengar & McEvily, 1992; Montero et al., 2004). They can chelate the copper prosthetic 86 group at the PPO active site, slowing down the enzymatic reaction, and can also inhibit 87 digestive metalloproteases responsible for PPO activation. Moreover, the inhibition of 88 digestive proteases should diminish free tyrosine concentration in flesh, which is a natural 89 substrate for PPO. Egg white can also be used as protease inhibitor, as it contains serine and 90 cysteine proteases. Other specific non-food grade inhibitors such as phenylmethylsulphonyl 91 fluoride (PMSF) and Iodoacetic acid (IAA) could also be used to study the role of different 92 proteases in the decomposition of viscera during storage. PMSF is an inhibitor of serine 93 proteases, while Iodoacetic acid (IAA) is an inhibitor of cysteine proteases.

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95 The main objective of this work was to assess the effect of several chemicals used as 96 coadjutants in 4-HR-based formulations on the occurrence of black spots and yellow-green 97 shades in the head of the deepwater pink shrimp (*Parapenaeus longirostris*) during chilled 98 storage. All formulations were used by dipping on board, simulating conditions habitually 99 used by fishermen. Preservatives, natural melanosis-inhibiting chemicals and protease 100 inhibitors infrequently studied in crustaceans were studied. The effect of these formulations 101 on acceptability was also evaluated through storage.

102 MATERIALS AND METHODS.

103 Deepwater pink shrimp (Parapenaeus longirostris) were caught off the South coast of 104 Spain (Cádiz) by trawl in March and May. Mean shrimp weights were 8.03 ± 1.54 g. On 105 board they were separated from the by-catch, washed with seawater and separated into 106 groups. Different 4-HR based formulations (Table 1) were then applied by immersion for 1 107 hour (shrimp:seawater relation of 1:2), one group immersed in seawater being considered 108 as control. All melanosis-inhibiting blends (Table 1) included 0.1% or 0.25 % of 4-109 hexylresorcinol (Sigma-Chemical, St Louis, MO, USA), and in almost all blends it was 110 used in combination with 0.5 % L-ascorbic acid (Sigma Chemical, St Louis, USA), 0.5 % 111 citric acid (Panreac Chemical, Barcelona, Spain), and 0.3% acetic acid (Panreac Chemical, 112 Barcelona, Spain). For simplicity, these formulations were designated as ACRA 0. 1% 113 (with 0.1% 4-HR) and ACRA 0.25% (with 0.25% 4-HR). Resorcinol alone or these basic 114 mixtures were accompanied by different chemical compounds to determine their capacity 115 to improve the appearance of shrimps: magnesium carbonate, di sodium di hydrogen 116 pyrophosphate (PPi), and potassium sorbate (Panreac Chemical, Barcelona, Spain); 117 iodoacetic acid (IAA, Sigma Chemical, St Louis, MO, USA), phenylmethylsulphonyl 118 fluoride (PMSF, Sigma Chemical), kojic acid (5-hydroxy-2 hydroxy-methyl-y-pyrone, 119 Sigma Chemical), phytic acid, and ethylenediaminetetraacetic acid (EDTA) (Sigma 120 Chemical); cumic acid (4-isopropilbenzoic acid, Aldrich Chemical, Milwaukee, WI, USA),

121 sodium metabisulphite (Sigma Aldrich Chemie, Germany); and dried egg white powder 122 (Degussa, Barcelona, Spain). Each formulation was assayed in two lots (1.5 kg of shrimps 123 per lot). Once the treatment time was finished, the shrimps were removed, placed in 124 perforated polystyrene boxes with a capacity of 2 kg, and covered with flaked ice. On 125 arrival of the trawler at the port, the boxes were taken by refrigerated truck to the Instituto 126 del Frío in Madrid, where they were stored in ice at 2 °C.

127 Sensorial analyses

128 A taste panel composed of eleven trained panellists visually inspected 14 shrimp per 129 treatment group every two days during the time of chilled storage. This conventional 130 sensory assessment was used to identify those significant sensory appearance attributes 131 which best define the quality loss of pink shrimp during storage. Those included general outward appearance (acceptability), the condition of the cephalothorax-tail junction, the 132 133 presence of yellow-greenish colouration beneath the head cuticle and also that of black 134 spots on the shell. The outward appearance was assessed as total percentage of shrimps 135 considered as acceptable for being sold on the market. The presence of yellow-greenish 136 colourations was evaluated as percentage of individuals with visual presence of this shade 137 in the head. The condition of the cephalothorax-tail junction was evaluated as percentage of 138 individuals with cephalothorax and tail separated. Finally, the presence of black spots in the 139 shell was assessed according to a visual scale from 1 to 4 (Montero et al., 2004), in which 140 1 = complete absence of black spots; 2 = a few small spots on the carapace; 3 =141 considerable spotting on the carapace; 4 = substantial spotting over the entire shrimp. The 142 cephalothorax-tail junction was not affected by the different formulas tested, and for that 143 reason this parameter was not further discussed.

144 Statistical analyses

The melanosis scores were regressed on time in storage using the SPSS computer software program (SPSS Inc., Chicago, Ill., USA). Linear or polynomial regressions were plotted through data dispersion, showing in all cases regression coefficients $R^2 \ge 0.90$. The data from the different variables analysed were used as the data matrix in different principal component analyses (PCA).

150 **RESULTS AND DISCUSSION.**

151 Non treated samples (Control samples). A noticeable increase in all parameters was 152 observed during iced storage, but especially in the case of melanosis, which showed the 153 earliest and more pronounced changes. Firstly, melanosis was observed mainly in the 154 cephalothorax and pleopods. Later, melanosis was also detected in abdomen cuticle and 155 telson, covering the whole surface after seven days of chilled storage (Fig 1). Regarding 156 acceptability, black spots decreased the perceived quality of the shrimp by the test panel, 157 and only approximately 25 % of shrimps were considered as acceptable two days after 158 capture.

159 Inhibitory effect of 4-hexylresorcinol combined with acids. The use of the melanosis-160 inhibiting formula composed of 0.1% 4-HR, 0.5% L-ascorbic acid, 0.5 % citric acid, and 161 0.3% acetic acid led to an absence of black spots in shrimps for the first four days of chilled 162 storage (Fig. 1a). This formula with 0.25 % 4-HR proved to be more effective at inhibiting 163 melanosis (Fig 1b). The effectiveness of 4-HR as inhibitor of PPO activity from crustaceans 164 has been extensively reported (McEvily et al., 1991; Otwell et al., 1992; Guandalini et al., 165 1998; Montero et al., 2004), although the effect depends on the application method, the 166 quantity applied, and mainly on the crustacean specie. The presence of organic acids in the 167 4-HR-based formula produces an acidic pH, that should negatively affect the PPO activity 168 (Gökoglu, 2004). Furthermore, ascorbic acid causes the chemical reduction of the pigment precursors, and citric acid may have a dual inhibitory effect on PPO: lowering the pH and chelating the copper at the active site of the enzyme (Iyengar & McEvily, 1992). Both ascorbic and citric acid could synergistically interact in the inhibition process, according to Lee-Kim, Hwang & Kim (1997).

Despite the finding that ACRA formula inhibited the PPO activity, the heads (beneath cuticle) turned greenish-yellow during storage, affecting acceptability (Table 2). The factor(s) which promote(s) the appearance of this colouration is/are unknown. However, it coincides with the appearance of black spots on carapace, so it is possible that yellowgreenish colouration is in some way related to melanosis process. Autolysis of shrimp viscera, or spoilage bacteria capable of producing melanin (Chinivasagam et al., 1998) could also be involved in the appearance of yellow-greenish shades in heads.

180 Kojic acid was incorporated in 4-HR based formulas as it would complement the inhibitory 181 effect of 4-HR (Chen et al., 1991a). The activity of kojic acid in the prevention of 182 melanosis involves two mechanisms: direct inhibition of PPO, and the chemical reduction 183 of the pigment or pigment precursors to colourless compounds (Ividogan & Bayindirli, 184 2004). Direct inhibition of PPO is produced as kojic acid is a competitive, slow-binding 185 inhibitor, like 4-HR (Jiménez & García-Carmona, 1997). The melanosis-inhibiting effect of 186 kojic acid has been observed in several crustacean species, such as Penaeus monodon 187 (Chen, Wei, Rolle, Otwell, Balaban & Marshall, 1991b) Penaeus japonicus (Montero, 188 Avalos & Pérez-Mateos, 2001b), but the effect of this chemical together with 4-HR has not 189 been studied in crustaceans. In this context positive results of melanosis-inhibiting 190 formulations including 4-HR and kojic acid have been described on apples (Ividogan & 191 Bayindirli, 2004). The effectiveness of kojic acid against melanosis could be higher when 192 mixed with ascorbic acid and citric acid, and those chemicals constitute a Japanese product 193 for inhibiting PPO in foods (Chen et al., 1991b). A mixture of ascorbic acid and kojic acid 194 has also been patented for use as an anti-browning agent in foods (Fukusawa, Wakabayashi 195 & Natori, 1982). Nevertheless, the use by immersion of formulations with 4-HR, organic 196 acids and 0.1% kojic acid had no significant effect on appearance of shrimps from two days 197 of chilled storage (Fig 1 and Table 1). It is worth noting that the bleaching mechanism of 198 quinones by kojic acid seems to be related to a redox reaction leading to an oxidised vellow 199 derivative of the inhibitor (Kahn, 1995), and it could enhance the intensity of yellow colour 200 in head during storage. The author suggested that kojic acid may not be desirable in certain 201 products, but that would depend on the concentration of the inhibitor and the type of food 202 in which it is used.

203 The incorporation of 0.1% cumic and/or phytic acid in the 4-HR-based formulations was 204 also tested (Fig. 1a and 1b). Cumic acid is a non-competitive inhibitor of PPO (Kubo & 205 Kinst-Hori, 1998). Furthermore, phytic acid is an antioxidant and chelating agent, so it 206 would inhibit not only PPO, but also digestive metalloproteases. Intensity of melanosis was 207 attenuated only when those chemicals were included in 0.1% 4-HR-based formulations, and 208 for 9 days of chilled storage. When cumic and/or phytic acid were incorporated in the 209 0.25% 4-HR based formula, the intensity of yellow shade in the head was lower after 7 210 days of chilled storage. However, this was not enough to improve the general appearance of 211 shrimps, according to the test panel.

Inhibitory effect of 4-hexylrerorcinol combined with sulphites. Sodium metabisulphite was tested in association with 4-HR. This chemical is usually incorporated in commercial melanosis-inhibiting formulations. The use of sulphite derivatives is limited by the regulatory authorities of many countries, which have indicated a maximum concentration of sulphites and derivatives in different foods. However, the appearance of melanosis occurs

217 more rapidly in several shrimp species such as pink shrimp, and for that reason dosage of 218 sulphites exceeds the usually permitted concentrations (Gómez-Guillén, Martínez-Alvarez, 219 Llamas Marcos & Montero, 2005). The use of 4-HR together with sulphite-derivatives in 220 commercial formulations could therefore be of high interest to reduce sulphite 221 concentrations in the flesh of shrimp. Figure 2 depicts the effect of 0.1% 4-HR combined 222 with variable concentrations of sodium metabisulphite routinely used on board (0.62% and 223 1.25%). In both cases, their effect on melanosis and also on the evidence of yellow-224 greenish colouration beneath cuticle was similar to that achieved by dipping shrimps in a 225 solution of 4-HR alone (Table 3). Nonetheless, lower concentrations of 4-HR were more 226 effective to prevent melanosis than that used for sodium metabisulphite.

227 Inhibitory effect of 4-HR combined with acids and magnesium carbonate. Magnesium 228 carbonate exerts a positive effect on melanosis prevention when it is used together with 229 citric, ascorbic, and sodium metabisulphite (Gutierrez Alsina, 1976). Magnesium carbonate 230 is decomposed into carbon dioxide, which interferes in the utilization of oxygen by PPO, 231 preventing decomposition of the viscera. Therefore, the effect of both magnesium 232 carbonate and 4-HR on melanosis could be complementary. The incorporation of 0.5% 233 magnesium carbonate in ACRA-0.1% formulations slightly prevented melanosis, but only 234 after 9 and 11 days of storage (Fig 3). This effect was not noticeable when magnesium 235 carbonate was used together with 0.25 % 4-HR. Moreover, the intensity of yellow-greenish 236 colouration was not reduced (Table 3), registering figures of 100% after 4 days of storage. 237 After 7 days, this colouration was not observed by the panellists due to the intensity of 238 melanosis in the head. Higher concentrations of magnesium carbonate (7 %) decreased 239 final quality after 7 days of storage and did not have any effect on the other parameters 240 judged.

241 Inhibitory effect of 4-hexylrerorcinol combined with acids and protease inhibitors. 242 The use of protease-inhibitors (EDTA and PPi) together with 4-HR and organic acids 243 increased acceptability, as shrimps were preserved well during nine days of chilled storage. 244 It should also be noted that, although exhibiting a decrease in sensorial appraisal during 245 storage, melanosis did not reach the threshold rejection value (≤ 2) after 9 days of iced 246 storage, except when egg white was used (Fig. 4). Despite the presence of vellow shades 247 beneath cuticle registering figures higher than 56% after 4 days of storage, almost 100% of 248 shrimps were surprisingly considered acceptable by the panellists during the whole period. 249 That is because the use of protease inhibitors decreased the intensity of the yellow shade in 250 head (Table 4), maintaining the fresh pink appearance. This result is ascribed to the natures 251 of EDTA and PPi, which are metalloprotease inhibitors and consequently may, at least 252 partially, inhibit the formation of free tyrosine and phenylalanine, substrates for the action 253 of PPO. In addition, they can inhibit the PPO activity by chelating the copper prosthetic 254 group at the PPO-active site or reducing the level of copper available for incorporation into 255 the enzyme (McEvily, Iyengar & Otwell, 1992).

On the other hand, egg white was added as it contains several inhibitor compounds, including ovomucoid and ovoinhibitor, which are serine-protease inhibitors, and cystatin, which is a cysteine-protease inhibitor (Stevens, 1991). Nonetheless, egg white exerts a detrimental effect on appearance (Table 5). It could be due to the fact that egg white could not be completely dissolved into the dipping solution. Egg white also contains proteins which can be hydrolyzed by endogenous and microbial proteases, producing free aminoacids which could be good substrates for PPO.

263 PMSF and IAA were also incorporated in the 4-HR-based formulas. Both were used only to264 gain scientific knowledge about their possible mechanism of action in the context of

265 enzymatic browning of crustacean, since they are non food grade chemicals. PMSF is a 266 characteristic inhibitor of serine proteases, while IAA is a thiol-blocking reagent used to 267 inhibit cysteine proteases in many studies (Hameed & Haard, 1985; Yamashita & 268 Konagaya, 1990; Gómez-Guillén, Hurtado & Montero, 2002). Shrimps treated with this 269 formulation showed the lowest melanosis index during storage, probably because the use of 270 protease inhibitors could at least partially inhibit the formation of free tyrosine and 271 phenylalanine, both substrates for the action of PPO. On the other hand, overall quality was 272 improved, but only after 9 days of chilled storage (Table 5), probably because the formula 273 with PMSF and IAA attenuated the yellow shade of head. The slight improvement in the 274 visual aspect could indicate that the appearance of greenish colouration and decomposition 275 of viscera are connected, since serine and cysteine proteases are the main enzymes implied in this process. The use of cystein and serine protease inhibitors should also be important as 276 277 these proteases are implied in the mechanism of activation of pro-PPO post mortem 278 (Söderhall & Cerenius, 1992; Zotos & Taylor 1997).

279 Effect of preservatives in 4-HR-based formulations. Spoilage bacteria could cause 280 melanosis in stored prawns, according to Chinivasagam et al. (1998). To detect the effect of 281 microorganisms on appearance of melanosis or greenish colouration, the preservative 282 potassium sorbate was incorporated in formulations containing 4-HR plus acids. Melanosis 283 in samples treated with and without preservatives evolved similarly during storage (Fig 5), 284 which evidenced the negligible effect of microorganisms on appearance of melanosis. 285 Similar results were observed regarding greenish colouration (Table 5). Furthermore, 286 almost 100 % of shrimps were considered acceptable by the panellists after 7 days of 287 storage.

Principal Component Analyses. Different principal component analyses (PCA) were performed in addition to sensorial analyses. The aim here was to identify significant differences that might have been overlooked previously in simple sensorial analyses. These further analyses were performed separately for samples treated with formulations including the same chemical compound. The corresponding principal component analyses (PCA) data matrix are shown in Table 7. In all cases, the global data matrix was reduced to 2 principal components (PC), which together account for 84-92 % of the explained variance.

295 The analyses of PC1 showed in all cases that yellow-greenish colouration correlated 296 positively with the time of storage and negatively with the acceptability. That means none 297 of the chemicals used prevented the appearance of yellow shades beneath the head cuticle, 298 which led, together with the appearance of melanosis, to the progressive loss of quality. 299 Even though 4-HR could not stop the appearance of the yellow-greenish colouration, the 300 acceptability and melanosis inhibition were increased with higher concentrations of this 301 chemical, as it is shown by the second component (PC 2, 36 % variance, Table 7a). The 302 incorporation of magnesium carbonate, phytic acid, kojic acid, cumic acid or sulphites in 303 the 4-HR-based formulas did not improve the general acceptability of shrimps (Table 7b-f). 304 On the other hand, the presence of protease inhibitors in the melanosis-inhibiting formulas 305 was related to better general acceptability (Table 7g, PC 2, 26% variance). However, PC 1 306 confirm that parameter acceptability is greatly influenced by both dark and yellow spots. 307 As yellow-greenish colouration was judged only as presence or absence, it is possible that 308 the protease inhibitors improved general appearance by reducing the intensity of this tone 309 beneath the head cuticle.

310 CONCLUSIONS

311 The use of increasing concentrations of 4-hexylresorcinol is related to better consumer 312 acceptability of pink shrimp (Parapenaeus longirostris) during chilled storage, also 313 inhibiting the evidence of melanosis. Although 4-hexylresorcinol can lessen the appearance 314 of black spots in the carapace, it can not prevent the previous manifestation of yellow-315 greenish colourations underneath head cuticle. The incorporation of protease inhibitors as 316 coadjutants in 4-hexylresorcinol-based formulations improved general appearance, but this 317 effect was not statistically significant. This suggests that the appearance of greenish 318 colourations could be related to decomposition of the viscera mediated by proteases, which 319 could facilitate liberation of substrates for PPO such as tyrosine. On the other hand, the 320 incorporation of another natural chemicals as kojic, cumic and phytic acid in 4-321 hexylresorcinol based formulations did not improve the overall acceptability of shrimps 322 during storage. Finally, the incorporation of preservatives did not have any effect on 323 appearance, suggesting micro-organisms do not participate in the appearance of anomalous 324 colourations in the cephalothorax.

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414 FIGURE CAPTIONS

Figure 1: Melanosis during the chilled storage of shrimp treated with 4-HR based
formulas including kojic acid (0.1%), cumic acid (0.1%) or phytic acid (0.1%). a) Formulas
with 0.1% 4-hexylresorcinol plus organic acids; b) Formulas with 0.25% 4-HR plus organic
acids. ACRA: 4-HR+ascorbic acid (0.5%)+acetic acid (0.05 N)+citric acid (0.5%); Koj:
kojic acid; Cum; cumic acid; Phy: phytic acid.
Figure 2: Intensity of melanosis during the chilled storage of shrimp treated with 4-

421 hexylresorcinol (0.25 % or 0.1%) combined with citric acid (0.5%), ascorbic acid (0.5%),

422 acetic acid (0.05 N), and sodium metabisulphite (1% or 0.625%). ACRA: 4-

- 423 HR+ascorbic+acetic+citric acid; Sul: sodium metabisulphite.
- 424 Figure 3: Appearance of melanosis during the chilled storage of shrimp treated with 4-
- 425 hexylresorcinol (0.25 % or 0.1%) combined with citric acid (0.5%), ascorbic acid (0.5%),

426 acetic acid (0.05 N), and magnesium carbonate (7% or 0.5%). ACRA: 4-427 HR+ascorbic+acetic+citric acid; Mg Car: magnesium carbonate.

Figure 4: Intensity of melanosis during the chilled storage of shrimps treated with 4hexylresorcinol (0.25 % or 0.1%) combined with organic acids, and the protease inhibitors
EDTA (225 ppm), PPi (1%), egg white (1% and 2%), Iodoacetic acid (210 ppm), and PMSF
(1 %)). ACRA 0.25%: 4-HR (0.25%)+ascorbic acid (0.5%)+acetic acid (0.05 N)+citric acid
(0.5%); EDTA: Ethylenediaminetetraacetic acid; PPi: Sodium pirophosphate; Ew: Egg

433 white; IAA: Iodoacetic acid; PMSF: Phenylmethylsulphonyl fluoride.

Figure 5: Melanosis during the chilled storage of shrimp treated with 4-hexylresorcinol
(0.25 % or 0.1%) combined with organic acids, the protease inhibitors EDTA (225 ppm)
and PPi (1%) and potassium sorbate. ACRA 0.25%: 4-HR (0.25%)+ascorbic acid
(0.5%)+acetic acid (0.05 N)+citric acid (0.5%). Sor: Potassium sorbate (2%).

439 **TABLES**

440 Table 1. Melanosis-inhibiting blends used by immersion after capture. ACRA: 0.5%

- 441 ascorbic acid+0.5% citric acid+0.3% acetic acid+0.1% 4-HR (ACRA 0.1%) or 0.25 % 4-
- 442 HR (ACRA 0.25%). R: 4-HR; Cit: Citric acid; Koj: Kojic acid; Cum: Cumic acid; Phy:
- 443 Phytic acid; Mg Car: Magnesium carbonate; Sul: sodium metabisulphite; EDTA:
- 444 Etilenediaminetetraacetic acid; Sor: Potassium sorbate; PPi: Sodium pirophosphate; Ew:
- 445 Egg white; IAA: Iodoacetic acid; PMSF: Phenylmethylsulphonyl fluoride.
- 446

MELANOSIS-INHIBITING	Koj	Cum	Phy	Mg	Sul	Sor	EDTA	PPi	EW	IAA	PMSF
BLEND	(%)	(%)	(%)	Car	(%)	(%)	(ppm)	(%)	(%)	(%)	(%)
				(%)							
CONTROL											
ACRA 0.1%											
ACRA 0.1%+Koj	0.1										
ACRA 0.1%+Cum		0.1									
ACRA 0.1%+Phy			0.1								
ACRA 0.1%+Cum+Phy		0.1	0.1								
ACRA 0.1%+Mg Car				0.5							
ACRA 0.25%											
ACRA 0.25%+Koj	0.1										
ACRA 0.25%+Cum		0.1									
ACRA 0.25%+Phy			0.1								
ACRA 0.25%+Cum+Phy		0.1	0.1								
ACRA 0.25%+EDTA+PPi							225	1			
ACRA 0.25%+EDTA+PPi+Ew 1%							225	1	1		
ACRA 0.25%+EDTA+PPi+Ew 2%							225	1	2		
ACRA							225	1		1	1
0.25%+EDTA+PPi+IAA+PMSF											
ACRA 0.25%+Sor						2 2					
ACRA 0.25%+EDTA+PPi+Sor						2	225	1			
ACRA 0.25%+Mg Car				0.5							
ACRA 0.25%+Mg Car 7%				7							
R 0.1%											
R 0.1%+Sul 1.25%					1.25						
R 0.1%+Sul 0.625%					0.625						
Sul 1.25					1.25						

- Table 2. Prevalence of yellow-greenish colourations and acceptability (%) of shrimp treated
 with 4-HR based formulas including kojic acid (0.1%), cumic acid (0.1%) or phytic acid
 (0.1%). a) Formulas with 0.1% 4-hexylresorcinol plus organic acids; b) Formulas with
 0.25% 4-HR plus organic acids. ACRA: 4-HR+ascorbic acid (0.5%)+acetic acid (0.05
 N)+citric acid (0.5%); Koj: kojic acid; Cum; cumic acid; Phy: phytic acid.
- 452 a)

		DA	YS OF	STOR	AGE	
Yellow-greenish shades (%)	0	2	4	7	9	11
Control	0	12.50	12.50	33.33		
ACRA 0.1%	0	16.67	42.86	70	75	100
ACRA 0.1%+Koj	0	16.67	100	100	100	100
ACRA 0.1%+Cum	0	41.67	50	80	75	100
ACRA 0.1%+Phy	0	33.33	57.14	80	100	100
ACRA 0.1%+Cum+Phy	0	33.33	57.14	80	100	100
Acceptability (%)	0	2	4	7	9	11
Control	100	27.58	0	0	0	0
ACRA 0.1%	100	100	100	80	0	0
ACRA 0.1%+Koj	100	100	100	0	0	0
ACRA 0.1%+Cum	100	83.3	100	50	8.33	0
ACRA 0.1%+Phy	100	83.3	100	50	0	0
ACRA 0.1%+Cum+Phy	100	100	100	50	0	0
b) Yellow-greenish shades (%)	0	2	4	7	9	11
Control	0	12.5	12.5	33.33		
ACRA 0.25%	0	8.33	28.57	80	100	100
ACRA 0.25%+Koj	0	33.33	14.29	80	91.67	100
ACRA 0.25%+Cum	0	33.33	21.43	50	90	81.25
ACRA 0.25%+Phy	0	16.67	14.29	30	80	87.5
ACRA 0.25%+Cum+Phy	0	33.33	21.43	50	80	87.5
Acceptability (%)	0	2	4	7	9	11
Control	100	27.58	0	0	0	0
ACRA 0.25%	100	100	100	90	8.33	0
ACRA 0.25%+Koj	100	100	100	100	8.33	6.25
ACRA 0.25%+Cum	100	100	100	100	16.67	0
ACRA 0.25%+Phy	100	100	100	100	8.33	6.25
ACRA 0.25%+Cum+Phy	100	100	100	100	0	0

454

455 Table 3. Presence of yellow-greenish colouration beneath cuticle and acceptability (%) of

456 shrimps dipped in melanosis-inhibiting solutions including 4-HR and/or sodium

457 metabisulphite. R: 4-HR; Sul: sodium metabisulphite.

458

DAYS OF STORAGE

Yellow-greenish shades (%)	0	2	4	7	9
Control	0	12.50	12.50	33.33	
R 0.1%	0	14.29	59.38	98.61	88.89
Sul 1.25%	0	24.40	25	33.33	
R 0.1%-Sul 1.25%	0	9.52	75	77.78	100
R 0.1%-Sul 0.625%	0	15.48	62.50	84.26	100
Acceptability (%)	0	2	4	7	9
Acceptability (%) Control	0 100	2 27.58	4 0	7 0	9 0
I V V	0	_		7 0 0	7
Control	100	27.58	0	7 0 0 0	0
Control R 0.1%	100 100	27.58 80.95	0 69.64	0	0 0

- 460 Table 4. Presence of yellow-greenish colouration beneath head cuticle (%) and
 461 acceptability (%) of shrimps treated with different melanosis-inhibiting blends including
 462 magnesium carbonate. ACRA: 4-HR (0.1 or 0.25%)+ascorbic acid (0.5%)+acetic acid (0.05
- 463 N)+citric acid (0.5%); Mg Car: Magnesium carbonate.
- 464

			DAY	YS		
Yellow-greenish shades (%)	0	2	4	7	9	11
Control	100	27.58	0	0	0	0
ACRA 0.1%	100	100	100	80	0	0
ACRA 0.25%	100	100	100	90	8.33	0
ACRA 0.1%+Mg Car 0.5%	100	100	100	20	0	0
ACRA 0.25%+Mg Car 0.5%	100	100	100	60	0	6.25
ACRA 0.25%+Mg Car 7%	100	100	100	0	0	0
Acceptability (%)	0	2	4	7	9	11
Control	100	27.58	0	0	0	0
ACRA 0.1%	100	100	100	80	0	0
ACRA 0.25%	100	100	100	90	8.33	0
ACRA 0.1%+Mg Car 0.5%	100	100	100	20	0	0
ACRA 0.25%+Mg Car 0.5%	100	100	100	60	0	6.25
ACRA 0.25%+Mg Car 7%	100	100	100	0	0	0

Table 5: Presence of yellow-greenish shades beneath cuticle (%) and acceptability (%) of
shrimps treated with different melanosis-inhibiting blends. ACRA 0.25%: 4-HR
(0.25%)+ascorbic acid (0.5%)+acetic acid (0.05 N)+citric acid (0.5%); EDTA:
Etilenediaminetetraacetic acid; PPi: Sodium pirophosphate; Ew: Egg white; IAA:
Iodoacetic acid; PMF: Phenylmethilsulphonil fluoride.

]	DAYS		
Yellow-greenish shades (%)	0	2	4	7	9	11
ACRA 0.25%	0	8.33	28.6	80	100	100
ACRA 0.25%+EDTA+PPi+Ew 1%	0	58.3	57.1	70		
ACRA 0.25%+EDTA+PPi+Ew 2%	0	41.7	71.4	40	40	
ACRA 0.25%+EDTA+PPi+IAA+PMSF	0	0	56.3	56.3	60	80
Acceptability (%)	0	2	4	7	9	11
ACRA 0.25%	100	100	100	90	8.33	0
ACRA 0.25%+EDTA+PPi+Ew 1%	100	100	100	0	0	0
ACRA 0.25%+EDTA+PPi+Ew 2%	100	100	83.3	0	0	0
ACRA 0.25%+EDTA+PPi+IAA+PMSF	100	100	100	83.3	83.33	67

- 473 Table 6: Development of yellow-greenish colouration in the cephalothorax and
 474 acceptability of shrimps dipped in different melanosis-inhibiting solutions. EDTA:
 475 Etilenediaminetetraacetic acid; PPi: Sodium pirophosphate; Sor: Potassium sorbate.

		DA	YS OF	STOF	RAGE	
Yellow-greenish shades (%)	0	2	4	7	9	11
ACRA 0.25%+Sor	0	41.7	42.9	75	100	100
ACRA 0.25%+EDTA+PPi+Sor	0	41.7	57.1	80	80	100
ACRA 0.25%	0	8.33	28.6	80	100	100
Acceptability (%)	0	2	4	7	9	11
ACRA 0.25%+Sor	100	100	85.7	100	0	0
ACRA 0.25%+EDTA+PPi+Sor	100	100	85.7	100	16.67	0
ACRA 0.25%	100	100	100	90	8.33	0

Table 7: Principal Component Analyses (data matrix) on the basis of the increasing
concentrations of 4-HR in the melanosis-inhibiting formulas (a), or else of the presence of
kojic acid (b), cumic acid (c), phytic acid (d), sulphites (e), magnesium carbonate (f), and
protease inhibitors (EDTA, PPi, Ew, IAA, PMSF) (g) in the 4-HR-inhibiting formulas.

a)	PC 1	PC 2
, ,	(49% variance)	(36 % variance)
4-hexylresorcinol content	0.316	0.864
Storage period	0.971	-0.199
Yellow-greenish colouration	0.966	0.056
Acceptability	-0.593	0.689
Melanosis score	0.349	-0.745
•	DC 1	DCA
b)	PC 1	PC 2
	(62 % variance)	(27 % variance
Kojic acid	0.008	0.985
Storage period	0.960	-0.047
Yellow-greenish colouration	0.908	0.109
Acceptability	-0.833	0.339
Melanosis score	0.960	0.095
c)	PC 1	PC 2
,	(68 % variance)	(22 % variance)
Cumic acid	0.035	0.990
Storage period	0.972	0.043
Yellow-greenish colouration	0.938	0.104
Acceptability	-0.886	0.251
Melanosis score	0.893	0.159
d)	PC 1	PC 2
,	(68 % variance)	(24 % variance
Phytic acid	0.017	0.987
Storage period	0.967	-0.003
Yellow-greenish colouration	0.952	-0.069
Acceptability	-0.832	0.458
Melanosis score	0.927	0.181

e)	PC 1	PC 2
-	(67 % variance)	(22 % variance)
Sulphites	-0.002	0.968
Storage period	0.978	-0.070
Yellow-greenish colouration	0.886	-0.211
Acceptability	-0.972	-0.089
Melanosis score	0.824	0.322
f)	PC 1	PC 2
,	(67 % variance)	(23 % variance
Magnesium carbonate	0.004	0.989
Storage period	0.965	-0.012
Yellow-greenish colouration	0.923	0.002
Acceptability	-0.888	0.341
Melanosis score	0.873	0.276
g)	PC 1	PC 2
8/	(58 % variance)	(26 % variance
	0.033	0.979
Protease inhibitors		
	0.944	-0.054
Protease inhibitors Storage period Yellow-greenish colouration	0.944 0.856	-0.054 -0.038
Storage period		