1 The effect of cinnamon, oregano and thyme essential oils in marinade on 2 the microbial shelf life of fish and meat products

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15 Abstract

16 Fresh and minimally processed fish and meat are easy targets for microbial spoilage. The 17 demand for natural alternatives to synthetic additives increases. In this study essential oil (EOs) 18 in marinades were used on fish and meat and the effect on the microbial growth during storage 19 was assessed. EOs from Oreganum compactum (oregano), Cinnamomum zeylanicum 20 (cinnamon), and *Thymus zygis* ct. Thymol (thyme) were chosen. The marinade was composed 21 of water, Na-lactate/lactic acid buffer (2 w/w %), NaCl (10 w/w %), and EO emulsified with 22 Tween 80 and with a pH of 4.5. The necessary Tween 80 to emulsify the EOs in the marinade depended on the EO type and was increased more than tenfold by the NaCl and lactate buffer. 23 24 The treatment consisted of immersion of meat (pork filet, pork bacon, chicken filets, chicken 25 skin), salmon or scampi for 2 min in marinade solution. The samples were stored at 4°C in air. 26 Samples were analyzed for microbial counts (dependent on matrix: total coliforms, *Escherichia* 27 *coli*, lactic acid bacteria, yeasts and molds, total aerobic psychrotrophs). Growth inhibition was 28 achieved with some EO + marinade treatments but marinade itself did not slow down the 29 microbial growth. Most notably, the growth of yeasts and molds was inhibited by immersion 30 of all food matrices in 1 w/w % cinnamon EO. Use of (1 w/w % for all EO) cinnamon EO (+ 31 marinade) led to microbial shelf life increase of all matrices (except the chicken matrices as the 32 end of the shelf life was not reached during the experimental duration), oregano EO to shelf life increase of pork filet and salmon, and thyme EO of pork filet and scampi. Sensorial analysis 33 34 on pork filet and salmon showed that immersion in 3 % EO (resulting in 0.09 g EO / 100 g 35 pork filet and 0.05 g EO / 100 g salmon) resulted in an acceptable odor after 24 h of storage. 36 The results in this study show that the sensorial properties of the meat/fish are inevitably 37 affected when the necessary EO concentrations to extend the microbial shelf life are applied.

38 Keywords

39 Essential oil, microbial shelf life, odor, fish, meat, marinade

40 **1. Introduction**

41 Due to the high water content and availability of important nutrients on the product surface, 42 fresh and minimally processed fish and meat are vulnerable to microbial spoilage (Iturriaga et 43 al., 2012; Casaburi et al., 2014). The dominating microbiota on cooled fish products consists 44 of psychrotolerant Gram-negative bacteria (Pseudomonas spp., Shewanella spp.). When 45 additional stress is created by additional antimicrobial practices (e.g. adding acid, salt, 46 antimicrobial food additives), the harsher environment can lead to a shift in spoilage 47 microorganisms to lactic acid bacteria, yeasts and molds (Gram & Dalgaard, 2002). In meat 48 products, the situation is basically the same although the species of spoilage microorganisms 49 that grow to the highest numbers and dictate the shelf life will differ because the microbial 50 growth rate depends on the nutrient constitution of the food product (Gram et al., 2002).

Marinating is defined as the preincubation of raw meat/fish products with a fluid (Quelhas et al., 2010), aiming to create an additional sensorial value (flavor, tenderness, moistness of the cooked product) and to extend the shelf life (Pathania et al., 2010). Marinades are water-based solutions that can contain sugar, salt, oil, organic acids, herbs and food additives such as aroma enhancers, antioxidants and antimicrobials (Bjorkroth, 2005). The antimicrobial properties of marinades are due to lowering of the pH, lowering of the water activity and addition of certain herbs and antimicrobial food additives (Pathania et al., 2010).

The demand for natural alternatives to synthetic additives increases and the replacement, in foodstuffs, of synthetic antimicrobials such as sorbate and benzoate by essential oils (EOs) is getting considerable attention (Salvia-Trujillo et al., 2014). The active compounds in EOs with antimicrobial properties can be divided as: terpenes, terpenoids, phenylpropenes and others (Hyldgaard et al., 2012). Depending on the active compound in the EO, different microbial targets or processes, especially cellular membranes and cellular energy production, but also 64 less known actions such as inhibition of cell division have been observed or proposed 65 (Hyldgaard et al., 2012). There are indications that the microbial shelf life of certain meat and fish products can be increased by treatment of the foodstuff with certain EOs, and often EO 66 67 from Origanum vulgare or Thymus vulgaris has been studied in that context because they contain the antimicrobial compounds thymol and carvacrol (Burt, 2004; Mexis et al., 2009; 68 69 Radha Krishnan et al., 2014; Tao et al., 2014). There are precedents that show the potential of 70 EOs for use in marinades. Due to addition of EOs to marinades, both the possibility of reducing 71 pathogens, such as Salmonella Enteritidis and Campylobacter coli on broiler breast fillet and 72 whole wings (Thanissery & Smith, 2014b), and of inhibiting growth of spoilage 73 microorganisms, such as total mesophilic counts (Thanissery & Smith, 2014a) or Pseudomonas 74 spp. and yeasts (Carlos & Harrison, 1999) on broiler breast fillet, have been observed.

Three EOs (from *Origanum compactum*, *Thymus zygis* ct. thymol and *Cinnamomum zeylanicum*) were selected for use in marinades. The effect of the marinades on the spoilage microflora of marinated meat, salmon and scampi was assessed during storage in normal atmospheric conditions at 4°C.

79 **2. Materials and methods**

80 **2.1. Raw materials**

Chicken skin, chicken breast fillet, pork (*Longissimus thoracis et lumborum* (LTL)), pork backfat, salmon (*Salmo salar*) and scampi (*Penaeus monodon*) were acquired from producers and transported (4°C) to the lab. The used EOs in this study were *Cinnamomum zeylanicum* (cinnamon EO) from the bark (Biover, Belgium), *Origanum compactum* (oregano EO) from the flowering top (Pranarôm, Belgium) and *Thymus zygis* ct. thymol (thyme EO) from the flowering plant (Biover, Belgium).

87 **2.2. Marinade solutions**

The marinade consisted of 10 w/w % NaCl and 2 w/w % Na-lactate/lactic acid buffer in deionized water with pH 4.5. Tween 80 was added to emulsify the EO (i.e. EO + marinade) in the marinade solution and the appropriate amount of Tween 80 (added as w/w %) was based on the outcome of the stability tests as described in 2.3. Mixing was done at 12500 rpm for 2 min (T18 digital ultra turrax, IKA, Belgium).

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2.3. Stability of essential oil in marinade emulsions

94 Amounts of Tween 80, EO, NaCl and Na-lactate/lactic acid were varied and the influence on 95 emulsion stability during 24 h of storage at 22°C was observed. Sunflower oil was added at a 96 concentration of 0 to 15 w/w %. All emulsions that contained lactic acid were kept at pH 4.5. 97 Ten mL of the emulsions were poured in glass tubes (internal diameter 9 mm) and stored at 98 22°C. The stability of emulsions of EO in marinade was assessed by visual observation, i.e. 99 whether a visual (0.5-1 mm layer) creaming layer occurred during the 24 h of storage. At that 100 moment the emulsion was considered unstable. For sensorial and microbial experiments, the 101 optimal settings from the stability experiments (i.e. lowest amount of Tween 80 to emulsify the 102 applied EO concentration and reach a stable emulsion) were applied. The particle size 103 distribution of the emulsions was determined by laser light diffraction (Mastersizer 2000, 104 Malvern, Belgium), with the laser emitting at 633 nm. The Sauter mean diameter for a 105 distribution of discrete entities (d_{32}) was used as this links the area of the dispersed phase to its 106 volume and as such to the mass transfer of the antimicrobial compound (Pacek et al., 1998):

107
$$d_{32} = \frac{\sum_{i=1}^{k} n_i d_i^3}{\sum_{i=1}^{k} n_i d_i^2}$$
(1)

108 in which:

109 n_i is the number of particles with diameter d_i .

110 The particle size distribution can be represented by its span:

111
$$span = \frac{d90 - d10}{d50}$$
 (2)

112 in which:

dx0 is the diameter corresponding to x0 volume % on a relative cumulative particle sizedistribution curve.

115 **2.4. Sample preparation and marinating process**

For salmon, pork LTL, chicken skin, chicken breast fillet, 10 g of sample was used with a fairly 116 117 constant surface to volume ratio among samples. The sample was completely immersed in 30 118 mL of (1 w/w % EO +) marinade for 2 min. The sample was removed from the marinade and 119 left to leak for 5 s. The sample was stored in a sterile stomacher bag (VWR, Belgium) at 4°C 120 with a small opening to allow gas exchange, i.e. stored in normal atmosphere. For pork back-121 fat the same was done but with 25 g of sample in 75 mL of (EO+) marinade. The larger sample 122 size was used to assure that the different layers of the pork back-fat (fat layers and meat layers) 123 were represented in each sample.

124 **2.5. Measuring pick-up**

125 The pick-up, i.e. the mass of marinade solution that remains on the sample after marinating,126 was measured by weighing the sample before and after the immersion and the leaking:

127
$$pick up = \frac{mass_{after} - mass_{before}}{mass_{before}} \times 100\%$$
 (3)

128 in which:

129 pick up is expressed in g/ 100 g,

130 mass_{after} = mass of the sample after immersion in marinade (+EO) solution,

131 mass_{before} = mass of the sample before immersion in marinade (+EO) solution.

132 **2.6. Microbial analyses**

133 Ten g of sample was put in a sterile stomacher bag (filter 0.5 mm pore size) (VWR, Belgium)

134 and homogenized during 1 min in 100 mL buffered peptone water (Oxoid, Belgium). Total

135 coliforms and Escherichia coli (E.coli) were enumerated with Chromocult Coliform-agar 136 (Merck, Germany) using the spreading plate method (incubation at 37 °C, 24 h). Yeasts and 137 molds (Y&M) were enumerated with Rose Bengal Chloramphenicol agar (Oxoid, Belgium) 138 containing 100 mg/L chloramphenicol (incubation at 22 °C, 5 days). Lactic acid bacteria (LAB) were enumerated with MRS (De Man, Rogosa, Sharpe) agar (Oxoid, Belgium), containing 1.4 139 140 g/L sorbic acid and with a final pH of 5.7, adjusted with NaOH (1 mol/L), using the pouring plate method with an additional cover layer of agar (incubation at 22°C, 5 days). Total aerobic 141 142 psychrotrophs (TAP) were enumerated with plate count agar (Oxoid, Belgium) using the 143 pouring plate method (incubation at 22°C, 5 days).

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2.7. Sensorial analyses to assess odor acceptability

Sensorial analysis was used to assess whether human subjects could distinguish, based on odor, 145 146 between samples that were treated with different concentrations of the same EO + marinade (0) 147 to 5 w/w %). For sensorial analyses, triangle tests (ISO 4120:2004) were used in an adjusted 148 form. The subject was asked not only to select the sample that differed from the other two, but 149 also to place the samples on a continuous hedonic scale (0 = very bad, 10 = very good) to assess 150 for the acceptability of the odor of the samples. This value was called the "hedonic value". The 151 samples were prepared as described in section 2.4 and stored for 24 hours in the fridge. After 152 that, samples were assessed by the subjects (raw samples) or baked (baked samples). Baked 153 samples were baked for 1 min at both sides in 1 g butter/ 10 g of meat/fish and subsequently, 154 during baking, turned on the other side every 30 s until the core of the sample reached 72 °C. 155 After baking, these samples were left to cool for 30 min and assessed by the subjects. The 156 control sample consisted of a sample treated with 1 w/w % sunflower oil + marinade and 157 emulsified with 0.1 w/w % Tween 80. The sunflower oil was added in order to avoid visual 158 differentiation by the sensory panel between samples treated with EO + marinade and samples 159 treated with marinade.

2.8. Gas chromatography/mass spectrometry (GC/MS) analysis

GC/MS analysis of the EOs was executed on a 6890 series GC-system (Agilent, Belgium) 161 162 equipped with a 7683 series injector (Hewlett Packard) and coupled to a 5973 Mass Selective 163 detector (Hewlett Packard, Belgium) in the electron impact ionization mode (70 eV) in the m/z164 range 40 to 550. The analysis was carried out using a HP-5ms column (methylpolysiloxane, 30 165 m x 0.25 mm inner diameter, 0.25 µm film thickness, Agilent, Belgium). A time-temperature profile, as described by Espina et al. (2011) was used. The flow of helium, the carrier gas, was 166 167 kept at 1 mL/min. The EOs were diluted 100 times in n-hexane, and 1 µL was injected in the 168 split mode (ratio 1:100). The analysis was executed three times for each EO. Data acquisition 169 was carried out with GC/MSD ChemStation software (Agilent, United States). Identification 170 was done by matching recorded mass spectra with reference spectra in the computer library 171 (NIST 98 Mass Spectral Library). Carvacrol (Sigma-Aldrich, Belgium), and (E)-172 cinnamaldehyde (Sigma-Aldrich, Belgium) were also dissolved in n-hexane and injected as 173 described for the EOs in order to use the observed retention times to distinguish between 174 carvacrol and thymol, and between (Z)- and (E)-cinnamaldehyde respectively. For 175 quantification the signal area percentage contribution of each identified compound to the total 176 signal area was used.

177 **2.9. Statistics**

To statistically assess the possible presence of growth inhibition due to the treatment solutionsthe log reduction was used as dependent variable:

$$log reduction = log(blank as cfu/g) - log(treatment as cfu/g)$$
(3)

181 in which:

182 blank = stored sample that was not treated (at day x)

183 treatment = stored sample that was treated with marinade (+EO) (at day x)

assessed with contrast analysis using SPSS Statistics 22 (IBM, United States). As in most cases 185 186 less importance was given to comparing e.g. 1% cinnamon + marinade with 1% oregano + 187 marinade, contrast analysis was chosen instead of ANOVA or the non-parametric alternatives. 188 Statistics concerning pick-up and sensorial analyses (hedonic values) were executed with 189 ANOVA, Welch or Kruskal-Wallis (dependent on the presence of normal distributions and/or 190 equal variances between groups) and, if relevant, the respective post-hoc analyses (i.e. Tukey, 191 Games-Howell and Dunn's multiple comparison test). To assess for equal variance among 192 groups Levene's test was used, and for normality Shapiro-Wilk. The probability of a false 193 positive result in the triangle tests was determined via the binomial distribution. The standard 194 deviation was used throughout the manuscript to represent data variation unless otherwise 195 stated.

Significant growth inhibition compared to the blank or marinade (without EO) samples was

196 The microbial shelf life was determined based on microbial shelf life criteria by Uyttendaele 197 et al. (2010). A conservative approach was taken. If, for any measured microbial parameter, 198 the mean log CFU/g food sample, increased with the standard deviation (of the three 199 independent repeats), exceeded the microbial limit for that microbial parameter, the shelf life 200 duration was over. If a treated sample resulted in microbial counts that remained below the 201 microbial limit for a longer duration than the untreated sample, the treatment increased the 202 shelf life. For the meat matrices the following limits were used: 7 log CFU LAB / g, 5 log CFU 203 Y&M /g (and no visible mold growth), 3 log CFU E. coli / g. For salmon and scampi the same 204 limits for LAB and Y&M were used, and in addition 7 log CFU TAP /g (Uyttendaele et al., 205 2010).

3. Results

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3.1. Composition of the essential oils

The composition of the *Cinnamomum zeylanicum*, *Origanum compactum* and *Thymus zygis* ct. Thymol used in this research is given in Table 1. Major components (> 5 % abundance) for cinnamon EO were (E)-cinnamaldehyde (66.28 %) and cinnamyl acetate (10.54 %), for oregano EO these were carvacrol (47.80%), thymol (21.41 %), γ -terpinene (13.44%) and pcymene (8.53 %), and for thyme EO these were thymol (55.91 %), p-cymene (20.61 %), and γ -terpinene (5.59 %).

214 **3.2. Emulsion stability**

215 Cinnamon EO was effectively emulsified in distilled water with a Tween 80:EO ratio of 1:100, 216 whereas a ratio of 1:10 was necessary for oregano and thyme EOs, and for oregano and thyme 217 EO a bimodal particle size distribution was observed at these settings (Table 2), indicating that 218 a small part of the particles had a significantly larger size, and as such indicating a less stable 219 crude emulsion compared to the cinnamon EO-in-water emulsion. More than 10 times the 220 concentration of Tween 80 was required to produce stable EO emulsions in the presence of 10 221 % NaCl or marinade than in demiwater. The addition of sunflower oil to the EO-in-water 222 emulsions lowered the necessary concentration of Tween 80 for cinnamon and thyme EO but 223 not for oregano EO. The Tween 80:EO ratio and mean particle size of the EO + marinade 224 emulsions that were selected for use in the sensorial and antimicrobial tests are shown in boldface in Table 2, and for each EO the ratio was chosen as the lowest Tween 80:EO ratio 225 226 that resulted in stable crude emulsions.

3.3. Pick-up

There was a large variability of the pick-up values among food matrices (Table 3), with an order of magnitude difference between the highest (on chicken skin) and lowest pick-up (on scampi). The concentration and type of EO did not influence the pick-up. The pick-up correlated weakly positive with fat (r= 0.453, p < 5. 10^{-4}), and weakly negative with both protein (r= -0.440; p < 5. 10^{-4}) and water (r= -0.438; p < 5. 10^{-4}).

3.4. Influence of essential oils + marinade on the microbial shelf life of fresh meat and fish

Marinade without EO did not reduce the microbial parameters during storage of any researched food matrix except for the reduction of total coliforms on pork back-fat for at least 1 day of storage.

238 On both chicken matrices, immersion in 1% cinnamon + marinade reduced the counts of some 239 microbial parameters after 6 days of storage (Table 4), i.e. Y&M and LAB in the case of 240 chicken breast fillet and total coliforms, Y&M and LAB in the case of chicken skin. Immersion 241 in 1% oregano + marinade and 1% thyme + marinade were only moderately effective in one case, i.e. a small reduction of Y&M on chicken breast filet was achieved after 6 days. As the 242 243 microbial shelf life of the chicken matrices was not reached within the duration of the 244 experiment, a potential shelf life increase due to the treatments could not be observed (Table 245 4).

On pork back-fat, total coliforms were reduced for at least 16 days with 1% cinnamon + marinade and at least 6 days with 1% oregano + marinade and 1% thyme + marinade (Table 5), whereas total coliforms did not grow on pork LTL. *E. coli* did not grow on both the pork matrices. Y&M were reduced during at least 16 days by 1% cinnamon + marinade on both pork matrices and for at least 10 days on pork LTL by 1% oregano + marinade and 1% thyme + marinade. LAB on pork LTL were only reduced after 10 days when treated with 1% oregano + marinade and at least 1 day on pork back-fat with 1% of all three EO + marinade. The 253 microbial shelf life of pork LTL was increased with all three EO + marinade, and that of pork
254 back fat with cinnamon EO + marinade (Table 5).

255 On salmon, Y&M were reduced for 6 days with 1% cinnamon +marinade, LAB were not 256 reduced, and TAP were reduced for at least 3 days with 1% cinnamon + marinade and 1% 257 oregano + marinade (Table 6). On scampi, there was no growth of Y&M and as such the 258 possible influence of 1% EO + marinade could not be established (Table 6). LAB were reduced 259 for at least 6 days on scampi with 1% oregano + marinade and 1% thyme + marinade and TAP 260 for at least three days for all EO + marinade and at least 6 days for 1% thyme + marinade. The microbial shelf life of salmon was increased with cinnamon and oregano EO, and that of scampi 261 262 with cinnamon and thyme EO and the marinade treatment (Table 6).

3.5. Sensorial analysis

264 There is a strong indication that for both the raw and baked pork LTL muscle and salmon a difference in odor was observed between samples treated with 1% sunflower oil + marinade 265 and 1% EO + marinade and between 1% EO + marinade and 5% EO + marinade but not 266 267 between 1% EO + marinade and 3% EO + marinade (Table 7). For the raw matrices, the 268 samples that were treated with 1 to 5% EO + marinade had a significantly lower hedonic value 269 than those treated with sunflower oil + marinade, except for one instance in the case of salmon 270 (Table 8). For raw salmon, 1% EO + marinade scored higher than 5% EO + marinade. Baking 271 of samples that were treated with EO + marinade increased the acceptability (i.e. hedonic value) 272 of the odor. For the baked matrices the differences in hedonic values between samples treated 273 with EO + marinade and sunflower oil + marinade were mostly insignificant, except for baked 274 pork LTL where oregano EO + marinade scored lower than sunflower oil + marinade. For 275 baked salmon the odor of samples treated with 1% sunflower oil + marinade scored higher than the odor of the samples treated with 5 % EO + marinade. When considering individual 276

treatments (e.g. 1% oregano EO + marinade), some treatments scored lower than 1% sunflower
oil + marinade for the raw matrices, but no significant differences were observed for the baked
matrices.

4. Discussion

281 The goal of the EO emulsion stability trials was to create crude EO-in-water emulsions that 282 remained stable during the marinating process, and not to study in detail the influence of the 283 marinade components on the EO emulsion stability. As such, this was not studied nor discussed 284 in depth. However, the detrimental influence of ionic strength on the formation of EO-in-water 285 emulsions is remarkable and an issue that could be relevant for practical application of EOs in 286 certain (food) emulsion systems. The reported used ratios of Tween 80:EO to emulsify EOs 287 are in general between 1:10 to 2:1 (Donsi et al., 2011, 2012; Chang et al., 2012; Terjung et al., 288 2012; Salvia-Trujillo et al., 2013, 2014; Loeffler et al., 2014; Sugumar et al., 2014; Hashtjin & 289 Abbasi, 2015). Concerning the influence of ionic strength and pH on the stability of EO-in-290 water however, next to nothing has been published. For non-ionic surfactants such as Tweens, 291 the presence of cations (especially monovalent cations) can be detrimental to the formation of 292 oil-in-water microemulsions due to dehydration of the polar groups which leads to separation 293 of the surfactant from the solution along with the oil (Binks & Dong, 1998; Warisnoicharoen 294 et al. 2000; Hsu & Nacu, 2003). However, in this study the stability of sunflower oil-in-water 295 emulsions was not significantly compromised by the presence of 10 % NaCl. EOs have a 296 relatively low interfacial tension and relatively high polarity. This makes EOs susceptible to 297 Ostwald ripening (i.e. growth of larger droplets at the expense of smaller ones due to diffusion 298 of oil through the aqueous phase) and more susceptible to coalescence (McClements & Rao, 299 2011). Use of a carrier oil to increase the hydrophobicity of the dispersed phase is a possible 300 strategy for increasing the emulsion stability. Unfortunately, some studies show that, when 301 keeping the absolute concentration of antimicrobial EO (component) constant, a relative increase of carrier oil can decrease the antimicrobial performance of the EO/carrier oil in water
emulsion (Chang et al., 2012; Suriyarak & Weiss, 2014). Another strategy would be to apply
another surfactant type to prevent coalescence (McClements & Rao, 2011).

305 The GC/MS results are in line with previous observations that cinnamaldehyde, carvacrol and 306 thymol are the most prevalent compounds in cinnamon EO (Yang et al., 2005; Unlu et al., 2010), 307 oregano EO (Lamiri et al., 2001; Bouchra et al., 2003; Mezzoug et al., 2007), and thyme EO of 308 the thymol type (Bagamboula et al., 2004; Burt, 2004) respectively. Also, p-cymene and γ -309 terpinene are major compounds of oregano and thyme EOs (Burt et al., 2005), which was also 310 the case in the present study. Most consistent in this study, is the antifungal efficiency of 311 cinnamon EO on all food matrices. In addition to its major abundance in cinnamon EO (> 66312 % in this study), cinnamaldehyde is more efficient to inactivate fungi, Gram-negative and 313 Gram-positive bacteria than its structural congeners: cinnamaldehyde > cinnamic acid > 314 cinnamyl alcohol > cinnamyl acetate (Chang et al, 2001; Wang et al., 2005), and as such its 315 contribution to the antimicrobial effect of cinnamon EO is large. Of the compounds found in 316 significant amounts in oregano and thyme EOs, thymol and carvacrol induce the strongest 317 antimicrobial effect as compared to (p-cymene, γ -terpinene etc.) (Bagamboula et al., 2004; Burt 318 et al., 2005; Sokovic et al., 2006). As they are also the compounds with the highest relative 319 abundance in these EOs, the contribution of thymol and carvacrol towards the antimicrobial 320 effect of oregano and thyme EOs is large. Nonetheless, there are some indications that synergy 321 among EO components could occur (Lambert et al., 2002; Periago et al., 2004; Burt et al., 322 2005), and as such the antimicrobial efficiency of an EO cannot be solely attributed to one or 323 a few of its major compounds without explicit evidence.

324 Considerable research is published on the use of EOs on meat and fish products in order to
325 extend the microbial shelf life. Chicken breast fillet has been treated with *Oreganum* EOs,
326 mostly *Origanum vulgare* (Chouliara et al., 2007; Khanjari et al., 2013; Fernandez-Pan et al.,

327 2014 ;Radha Krishnan et al., 2014), Thymus vulgaris EO (Giatrakou et al., 2010; Thannissery 328 & Smith, 2014a) and *Cinnamomum cassia* (Radha Krishnan et al., 2014). Lean pork meat has 329 been treated with thymol and *Thymus vulgaris* EO (Carramiñana et al., 2008; Tao et al., 2014) 330 and pork back-fat sausages with thymol (Mastromatteo et al., 2011). The published information 331 concerning preservation of salmon (Salmo salar) with EOs is limited. However, some research 332 been published on the preservation of the closely related (both belong to the Salmonidae 333 family) rainbow trout (Onchorynchus mykiss). Rainbow trout fillet has been treated with 334 Origanum vulgare EO (Mexis et al., 2009) and Cinnamomum zeylanicum EO (Andevari & 335 Rezaei, 2011). Shrimps (Palaemon serratus) have been treated with thymol (Mastromatteo et 336 al., 2010), and precooked peeled shrimps (Penaeus spp.) with Thymus saturoïdes EO and (E)-337 cinnamaldehyde (Ouattara et al., 2001). In most of the aforementioned studies, the potential of 338 these EOs to slow the growth of some of the analyzed groups of spoilage microorganisms for 339 a certain period of storage time has been observed, given a sufficient dose of EO. The collective 340 goal of these antimicrobial studies is to gain understanding concerning the dose-response of 341 the EO treatment on the spoilage microorganisms on these foodstuffs. Ultimately the actual EO 342 dose is the pick-up and herein lies the current problem. For virtually all the aforementioned 343 studies, it is unknown how much of the EO actually remained on the food matrix after 344 treatment, which can consist of EO being i) massaged in the food matrix, ii) added to the food 345 matrix, iii) added to the minced food matrix, iv) pipetted on the food matrix, v) the food matrix 346 can be immersed in EO emulsion etc. The results in the current study could be compared with other studies by the pick-up values. In the current study this was done by multiplying the 347 348 concentration of EO in the marinade with the pick-up values (Table 3). The EO pick-up is a 349 rough estimation because i) not all (EO) components of the marinade are expected to be 350 transferred to the same extent to the food matrix, ii) variance in the pick-up due to transfer of 351 some solid matter from the tissue to the EO + marinade emulsion during the marinating process,

352 iii) variance in the pick-up due to transfer of water from the tissue to the marinade emulsion 353 because of the high salt content in the marinade emulsion (osmotic effects). These issues were 354 reflected in the relatively high standard deviation in pick-up values for each food matrix. A 355 more accurate method would consist of actually determining the quantity of the adsorbed EO components, through e.g. GC-MS analysis. In order to gain understanding concerning the use 356 357 of EOs on foods in order to extend the shelf life it is of paramount importance that a method to 358 measure the pick-up is developed and adopted by researchers, because at the moment very little 359 quantitative conclusions can be drawn from the ample collection of generated antimicrobial 360 data.

361 When EOs are applied in food formulations, the sensorial impact of these EOs is a limitation 362 towards the quantity of EO that can be applied. In this study, baking improved the perception 363 of the odor coming from the baked meat and fish, probably in part due to volatilization of EO 364 compounds during the baking process as well as the mix of the EO odor with generated odorous 365 compounds from the baked matrices. The results suggest that the antimicrobial treatment with 366 1% EO + marinade could be increased to 3% EO + marinade without compromising the odor 367 of the food matrices. An increase to 5% EO + marinade seems to result in less well perceived 368 odors on baked salmon, as does the use of oregano on baked pork LTL. In this study, only the 369 odor after 1 day of storage was assessed, mainly to detect possible detrimental influences on 370 the fish/meat as this is critical information for valorization of this EO application. As such, the 371 possible beneficial influence of the EOs on the sensorial quality of the meat/fish during storage 372 was not assessed explicitly, only indirectly through microbial enumerations. With an estimated 373 sensorial acceptable concentration in the range between 3 and 5 % EO + marinade immersion 374 treatments, an acceptable pick-up concentration between 0.09 and 0.15 w/w % on pork LTL 375 and between 0.05 and 0.09 w/w % on salmon can be expected. The acceptable EO 376 concentrations are quite diverse when comparing studies. When applying Origanum vulgare

377 EO on meat, the added concentrations that resulted in acceptable odor and taste were in the 378 range of 0.1 to 1 (w/w or v/w) % (Sánchez-Escalante et al., 2003; Skandamis & Nychas, 2001; 379 Chouliara et al., 2007; Govaris et al., 2010; Karabagias et al., 2011; Petrou et al., 2012), while 380 unacceptable added concentrations were in the range 0.2 to 1% (Chouliara et al., 2007; 381 Ntzimani et al., 2010; Karabagias et al., 2011). When applied on fish, acceptable concentrations 382 were in the range 0.1 to 0.4 % (Giatrakou et al., 2008; Mexis et al., 2009; Frangos et al., 2010), 383 while 0.4 % was considered unacceptable on rainbow trout fillet (Frangos et al., 2010). Use of 384 Thymus vulgaris EO on meat was acceptable concerning odor and taste in the range of 0.2 to 385 0.6 % (Solomakos et al., 2008; Giatrakou et al., 2010) but unacceptable at 0.9 % on minced 386 beef (Solomakos et al., 2008). For fish, acceptability was in the range 0.1 to 0.4 % (Kostaki et 387 al., 2009; Kykkidou et al., 2009; Abdollahzadeh et al., 2014) but unacceptable at 0.8 % on 388 minced silver carp (Abdollahzadeh et al., 2014). Cinnamon EO as an antimicrobial on meat 389 and fish has been studied much less than oregano or thyme EO. Treatment of sheep patties by 390 immersion in 0.25 % Cinnamomum cassia (Luo et al., 2007) and chicken breast fillet by 391 immersion in 1 % Cinnamomum cassia (Radha Krishnan et al., 2014) were found to be 392 acceptable concerning odor and taste. The observed substantial range of acceptable EO 393 concentrations is explained by the actual concentration of EO that remains on/in the meat/fish 394 tissue after treatment, the variation in compatibility between a certain EO and a certain 395 meat/fish product, and the inherent subjectivity that arises when applying small, moderately 396 trained sensory panels (sensory acceptability is a. o. function of age, gender and cultural 397 background) (Samant et al., 2015). Acceptability of EO treated meat/fish does not imply that 398 the EO does not influence the taste and odor. In the current study, the presence of 0.030 ± 0.002 399 % EO on pork LTL and 0.018 ± 0.002 % EO on salmon (both due to a 2 min dipping treatment 400 in 1% EO + marinade) resulted in observable but acceptable odors after 24 h storage (and 401 cooking). Treatment through addition of 0.1% Origanum vulgare to swordfish fillet, 0.2% to

402 rainbow trout fillet, submerging of chicken breast fillet in 1% Origanum vulgare, and addition 403 of 0.2 % Thymus vulgaris to chicken kebab and sea bass fillet, all resulted in an acceptable but 404 very noticeable taste and odor (Giatrakou et al., 2008, 2009; Frangos et al., 2010; Kostaki et 405 al., 2009; Khanjari et al., 2013). The use of an active compound instead of the EO (e.g. 406 cinnamaldehyde instead of cinnamon EO) would reduce the total amount of added compounds 407 that have sensorial impact on the foodstuff. Although this would not rule out the sensorial 408 limitations, it could potentially improve the usability of these antimicrobials and is worth 409 investigating.

410 **5. Conclusion**

411 Marinade (10% NaCl, 2% lactic acid, pH 4.5) in itself did not inhibit microbial growth on the 412 food matrices. Cinnamon, oregano and thyme EOs, applied at low concentrations, show 413 potential to slow the growth (extend the microbial shelf life) of some spoilage microorganisms 414 on meat/fish products when applied in a marinade. Of particular interest is cinnamon EO, which 415 is especially efficient for inhibition of fungal growth on meat and fish. Combinations of EOs 416 or specific compounds could be a strategy to increase the antimicrobial spectrum. Comparison 417 of research on the effects of EOs on the shelf life of foodstuffs is hampered by the lack of the 418 use of a method that determines the pick-up (or otherwise stated the active dose). As long as 419 such a method is not adopted, quantitative understanding of these antimicrobial treatments 420 remains limited to the applied experimental setup. Besides the antimicrobial effects, the results 421 in this and other studies also show that the sensorial properties of the meat/fish are inevitably 422 affected (positively, neutrally or negatively) when the necessary EO concentrations to extend 423 the microbial shelf life are applied. This implies that the sensorial effect that results from 424 combining a certain EO with a certain meat/fish product is virtually always a significant factor 425 and not all combinations will be acceptable in commercial use.

426 Acknowledgments

The research leading to these results has been facilitated by the Flemish Agency for Innovation by Science and Technology (IWT) under grant agreement IWT TETRA nr. 130214. The authors want to thank Joël Hogie and Yannick Verheust for supplying the needed equipment and assistance, and the thesis students Jens Beernaert, Stefanie Carpentier and Lisa Vandenberghe for their work.

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646Table 1. Composition of the essential oils Cinnamomum zeylanicum, Origanum

compactum, Thymus zygis ct. thymol (expressed as % of the ion signal area) (n=3)

retentio	n time (min) compound	Cinnamomum zeylanicum	Origanum compactum	<i>Thymus zygis</i> ct. Thymol
7.13	α-thujene	0.18	0.48	0.60
7.41	α-pinene	0.98	0.43	0.81
8.06	camphene	0.45	0.06	0.65
9.40	β-pinene	0.28	0.06	0.15
10.19	β-myrcene		0.91	0.97
10.83	α -phellandrene	0.99	0.17	0.13
11.49	α-terpinene	0.94	1.70	1.12
11.94	p-cymene	2.43	8.53	20.61
12.14	sylvestrene/limonene		0.38	0.53
12.15	β-phellandrene	3.94		
12.26	eucalyptol	0.30	0.05	0.32
13.86	γ-terpinene	0.12	13.44	5.59
15.54	terpinolene	0.12	0.08	0.27
16.26	linalool	1.60	1.05	3.30
18.62	L-camphor			0.35
19.88	borneol		0.13	1.40
19.90	hydrocinnamic aldehyde	0.46		
20.58	terpinen-4-ol	0.66	0.43	0.76
21.39	α-terpineol	0.65	0.19	0.18
23.04	(Z)-cinnamaldehyde	0.64		
23.71	hydrocinnamyl alcohol	0.22		
24.42	thymyl methyl ether		0.12	0.62
25.94	(E)-cinnamaldehyde	66.28		
27.28	thymol		21.41	55.91
27.76	carvacrol	0.12	47.80	2.90
30.51	eugenol	2.25		
31.33	α-copaene	0.38		
33.50	β-caryopyllene	2.40	1.56	1.13
34.95	cinnamyl acetate	10.54		
35.19	α-caryophyllene	1.97	0.07	
41.35	caryophylllene-oxide	0.56		
	Not identified	0.61	0.95	1.74

650Table 2. Necessary ratio of Tween 80:EO to emulsify 10 w/w % of the studied essential

EO	NaCl (m%)	lactic acid buffer (w/w %)	sunflower oil (w/w %)	Tween 80:EO	Particle size (µm)	Span
cinnamon	0	0	0	1:100	0.40 ^A	2.75
	0	0	0	1:10	0.26	2.62
	0	2	0	1:10	ND^{B}	
	10	0	0	2:10	ND	
	10 ^C	2	0	2:10	0.23	3.22
	10	2	0	12:10	0.19	3.29
	10	2	5	1:10	0.52	2.16
oregano	0	0	0	1:10	0.24	6.14*
	0	2	0	7:10	ND	
	10	0	0	12:10	ND	
	10	2	0	12:10	0.20	2.61
	10	2	0-15	>7:10	ND	
thyme	0	0	0	1:10	0.41	13.0*
-	0	2	0	7:10	ND	
	10	0	0	10:10	ND	
	10	2	0	12:10	0.20	2.53
	10	2	5	7:10	0.21	123.0*

oils in the presence of NaCl, lactic acid buffer, and sunflower oil (n=2)

 A Sauter mean diameter (d₃₂), ^B ND: not determined, ^C lines in boldface denote the EO +

marinade emulsions used in the sensorial and antimicrobial experiments, * a bimodal particle

654 size distribution was observed.

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	pick-up (marinade)	estimated pick- up (EO)	fat	protein	water
food matrix	g/100 g	g/100 g	g/100 g	g/100 g	g/100 g
chicken skin	9.0±1.1 ^B	0.090±0.011	44.9	9.6	42.9
chicken filet	4.9±0.5	0.049 ± 0.005	1.3	22.8	74
pork back fat	4.2 ± 0.4	0.042 ± 0.004	53.3	10.6	34
pork LTL	3.0±0.2	0.030 ± 0.002	1.9	20.5	76
salmon	1.8 ± 0.2	0.018 ± 0.002	16.5	18.4	63
scampi	0.9 ± 0.4	0.009 ± 0.004	0.1	17.5	79

Table 3. Pick-up of EO + marinade on the studied food matrices^A (n=20)

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^Afat, protein and water content were acquired from the food producer and www.internubel.be and for chicken skin from (Bonifer et al., 1996; Badr, 2005), ^B standard error of mean

Table 4. Microbial counts (log CFU/g) of selected microbial parameters during storage

of treated chicken breast filet and chicken skin (n=3)

		chicken	breast filet	chicken skin		
storage time (o	storage time (days)			1	6	
total coliforms	blank	2.1±0.3	2.7±0.6	3.8±0.5	5.2±0.4	
	marinade	1.9 ± 0.1	2.3 ± 0.8	3.6 ± 0.2	4.9 ± 0.1	
	1% cinnamon + marinade		2.9±1.5	3.5±0.1	4.1 ± 0.5^{A}	
	1% oregano + marinade	1.9 ± 0.1	3.4±0.4	4.6±0.7	4.6±0.7	
	1% thyme + marinade	1.9 ± 0.0	2.8 ± 1.3	3.8 ± 0.4	$4.4{\pm}1.1$	
E. coli	blank	<2	<2	3.2±0.2	2.8 ± 0.5	
	marinade	<2	<2	3.3±0.1	3.0±0.4	
	1% cinnamon + marinade	<2	<2	3.1±0.2	2.6 ± 0.4	
	1% oregano + marinade	<2	<2	$2.9{\pm}0.6$	2.7 ± 0.3	
	1% thyme + marinade	<2	<2	3.2±0.2	2.7 ± 0.4	
Y&M	blank	2.2 ± 0.3	3.7±0.4	2.8 ± 0.4	4.1±0.2	
	marinade	1.9 ± 0.1	3.5±0.3	2.7 ± 0.1	4.1 ± 0.2	
	1% cinnamon + marinade	2.2 ± 0.1	$2.7{\pm}0.8^{\mathrm{A},\mathrm{B}}$	2.7 ± 0.3	$3.4{\pm}0.4^{\text{A},\text{B}}$	
	1% oregano + marinade	1.9 ± 0.1	3.1±0.1	3.0±0.3	4.0 ± 0.1	
	1% thyme + marinade	2.0 ± 0.2	3.1±0.2	3.2±0.4	4.4±0.3	
LAB	blank	1.8 ± 0.6	3.6±0.2	3.5 ± 0.4	5.6±0.3	
	marinade	1.6 ± 0.2	3.1±0.4	3.5 ± 0.2	5.3±0.5	
	1% cinnamon + marinade	1.7 ± 0.9	$2.6\pm0.2^{A,B}$	3.9±0.3	4.9 ± 0.3^{A}	
	1% oregano + marinade	1.4 ± 0.6	3.0±1.1	3.4±0.3	5.0 ± 0.6	
· · · · · · · · · · · · · · · · · · ·	1% thyme + marinade	2.0±0.7	3.0±1.0	4.3±0.7	5.3±0.5	

 A significant reduction (p < 0.05) compared to the untreated (blank) sample, ^B significant

reduction (p < 0.05) compared to the marinated (without EO) samples.

Table 5. Microbial counts (log CFU/g) of selected microbial parameters during storage of treated pork LTL and pork back-fat (n=3)

	pork LTL						pork b	ack-fat	
storage time (days)		1	6	10	16	1	6	10	16
Total	blank	<2	<2	<2	<2	3.8±0.5	5.9±0.4	5.7±1.1	5.9±0.9
	marinade	<2	<2	<2	<2	2.4 ± 0.2^{A}	5.3±0.7	5.1±1.3	5.3 ± 1.0
	1% cinnamon + marinade	<2	<2	<2	<2	2.1 ± 0.2^{A}	$3.1 \pm 1.0^{A,B}$	$4.0{\pm}1.8$	3.3 ± 2.3^{A}
	1% oregano + marinade	<2	<2	<2	<2	2.3 ± 0.6^{A}	3.5 ± 1.5^{A}	5.5 ± 1.3	5.5 ± 0.7
	1% thyme + marinade	<2	<2	<2	<2	2.3 ± 0.6^{A}	3.2 ± 2.1^{A}	3.8 ± 1.9	4.6 ± 2.3
E. coli	all treatments	<2	<2	<2	<2	<2	<2	<2	<2
Y&M	blank	2.3 ± 0.6	4.7±0.5†	6.6 ± 0.4	6.2 ± 0.5	4.1±0.2	6.3±0.1†	6.3±0.4	7.0 ± 0.4
	marinade	2.1±0.2	4.9±0.2†	6.5 ± 0.6	7.3±0.6	4.2 ± 0.2	6.4±0.1†	6.9 ± 0.1	7.3±0.3
	1% cinnamon + marinade	2.0 ± 0.0	$2.3 \pm 0.5^{A,B}$	$3.5 \pm 0.6^{A,B}$	4.9±1.1 ^{A,B} †	$2.4 \pm 0.6^{A,B}$	$2.7 \pm 0.9^{A,B}$	$3.0{\pm}1.0^{A,B}$	3.4±2.1 ^{A,B} †
	1% oregano + marinade	2.2 ± 0.2	$3.8{\pm}0.6^{A,B}$	5.7±0.1 ^{A,B} †		$3.2{\pm}1.1$	5.3±1.7†	6.1 ± 0.6	6.2 ± 1.1
	1% thyme + marinade	2.0 ± 0.0	$3.8{\pm}0.8^{A,B}$	5.6±0.4 ^{A,B} †	6.7 ± 0.5	3.3±1.1	5.0±2.0†	6.0 ± 0.7^{B}	6.9 ± 0.5
LAB	blank	1.8 ± 0.6	5.1 ± 0.5	6.8±0.3	7.2 ± 0.5	2.8 ± 0.4	5.1±0.2	5.7 ± 0.6	5.8 ± 0.9
	marinade	1.2 ± 0.2	4.6 ± 0.4	6.5 ± 0.7	7.0 ± 0.6	2.6 ± 0.1	5.3±0.8	5.7 ± 0.5	5.9 ± 0.6
	1% cinnamon + marinade	1.3±0.3	4.7 ± 0.4	6.1±0.3	7.2±0.7†	2.1 ± 0.4^{A}	$4.0{\pm}1.3$	5.6 ± 0.7	6.4±0.3
	1% oregano + marinade	1.4 ± 0.3	4.6±0.3	5.9 ± 0.5^{A}	6.5 ± 0.4	2.0 ± 0.3^{A}	4.5 ± 0.9	5.9 ± 0.5	5.8 ± 1.5
	1% thyme + marinade	$1.0{\pm}0.1$	4.5 ± 0.6	6.1±0.7	7.1±0.6	2.1 ± 0.5^{A}	5.1 ± 0.5	4.9 ± 1.4	5.4±1.5

^A significant reduction (p < 0.05) compared to the untreated (blank) sample, ^B significant reduction (p < 0.05) compared to the marinated (without

EO) samples, † the end of shelf life is reached due to the value of this microbial parameter.

Table 6. Microbial counts (log CFU/g) of selected microbial parameters during storage

			salmon			scampi	
storag	ge time (days)	1	3	6	1	3	6
Y& M	blank	3.1±0.3	3.7±0.1	4.7±0.2	<2	<2	<2
	marinade	3.2±0.2	3.8±0.2	5.0±0.1†	<2	<2	<2
	1% cinnamon marinade	+ $2.1\pm0.2^{A}_{,B}$	3.4±0.6	$2.9{\pm}0.7^{A,B}$	<2	<2	<2
	1% oregano marinade	+ 2.9±0.2	3.8±0.4	4.8±0.2†	<2	<2	<2
	1% thyme marinade	+ 3.0±0.3	3.8±0.2	4.7±0.1	<2	<2	2.2±0.4
LAB	blank	<1	3.1±0.2	2.9±0.3	1.1±0.1	1.8±0.1	2.6±0.5
	marinade	<1	2.9±0.1	3.3±0.1	1.7±0.6	$2.2{\pm}1.0$	2.6±0.8
	1% cinnamon marinade	+ <1	2.4±0.2	3.1±0.1	1.2±0.2	1.4±0.4	1.8±0.7
	1% oregano marinade	+ <1	2.9±0.4	3.1±0.3	1.0±0.0	1.3±0.5	1.2±0.2 ^{A,B}
	1% thyme marinade	+ <1	3.1±0.1	3.0±0.1	1.3±0.2	1.1±0.2 ^A	1.0±0.1 ^{A,B}
TAP	blank	5.5±0.3	7.3±0.4†	9.3±0.4	5.1±0.2	5.7±0.4	8.0±2.3†
	marinade	5.3±0.3	6.7±0.3†	8.7 ± 0.7	4.7±0.5	5.5±0.1	6.5 ± 0.4
	1% cinnamon marinade	+ $4.5\pm0.2^{A}_{,B}$	6.2±0.3 ^A	8.9±0.6†	4.2±0.2 A	4.8±0.4 ^A	6.0±0.3
	1% oregano marinade	+ 5.1±0.3	6.5±0.1 ^A	9.2±0.5†	4.2±0.3	$3.7 \pm 0.7^{A,B}$	7.4±2.1†
	1% thyme marinade	+ 5.1±0.2	7.0±0.2†	9.5±0.0	4.0±0.5 A	$3.9{\pm}0.5^{A,B}$	$5.6\pm0.3^{A,B}$

of treated salmon and scampi (n=3)

^A significant reduction (p < 0.05) compared to the untreated (blank) sample, ^B significant reduction (p < 0.05) compared to the marinated (without EO) samples, † the end of shelf life is reached due to the value of this microbial parameter.

Table 7. Results of triangle tests for detecting a difference between raw and fried pork

raw pork LTL	correct	α-risk ^A
sunflower oil 1% VS cinnamon 1%	10/10	<0.1%
sunflower oil 1% VS oregano 1%	9/10	<0.1%
sunflower oil 1% VS thyme 1%	10/10	<0.1%
cinnamon 1% VS cinnamon 3%	3/10	>20%
oregano 1% VS oregano 3%	4/10	>20%
thyme 1% VS thyme 3%	6/10	8%
cinnamon 1% VS cinnamon 5%	7/10	2%
oregano 1% VS oregano 5%	6/10	8%
thyme 1% VS thyme 5%	6/10	8%
sunflower oil 1% VS EO 1%	29/30	<0.1%
EO 1% VS EO 3%	13/30	17%
EO 1% VS EO 5%	19/30	<0.1%
raw salmon	correct	α-risk
sunflower oil 1% VS cinnamon 1%	8/10	0.3%
sunflower oil 1% VS oregano 1%	10/10	<0.1%
sunflower oil 1% VS thyme 1%	7/10	2%
cinnamon 1% VS cinnamon 3%	5/10	>20%
oregano 1% VS oregano 3%	3/10	>20%
thyme 1% VS thyme 3%	5/10	>20%
cinnamon 1% VS cinnamon 5%	7/10	2%
oregano 1% VS oregano 5%	5/9	>20%
thyme 1% VS thyme 5%	3/10	>20%
sunflower oil 1% VS EO 1%	25/30	<0.1%
EO 1% VS EO 3%	13/30	17%
EO 1% VS EO 5%	15/29	3%
fried pork LTL	correct	a-risk
sunflower oil 1% VS cinnamon 1%	7/8	0.3%
sunflower oil 1% VS oregano 1%	6/8	2%
sunflower oil 1% VS thyme 1%	7/8	0.3%
cinnamon 1% VS cinnamon 5%	4/8	>20%
oregano 1% VS oregano 5%	6/8	2%
thyme 1% VS thyme 5%	5/8	9%
sunflower oil 1% VS EO 1%	20/24	<0.1%
EO 1% VS EO 5%	15/24	0.3%
fried salmon	correct	a-risk
sunflower oil 1% VS cinnamon 1%	7/8	0.3%
sunflower oil 1% VS oregano 1%	5/8	9%
sunflower oil 1% VS thyme 1%	4/8	>20%
cinnamon 1% VS cinnamon 5%	5/8	9%
oregano 1% VS oregano 5%	6/8	2%
thyme 1% VS thyme 5%	4/8	>20%
sunflower oil 1% VS EO 1%	16/24	<0.1%
EO 1% VS EO 5% Aprobability of false positive result	15/24	0.3%

LTL and salmon treated with sunflower oil/EO+marinade

EO 1% VS EO 5%15/240.3%Aprobability of false positive result, determined via the binomial distribution

	-	pork LTL hedonic value		llmon nic value
raw	number of tests	mean	number of tests	mean
sunflower oil1%+marinade	45	6.6±2.2	30	6.6±2.7
cinnamon 1%+marinade	45	$5.0\pm 2.3^{\text{A}}$	45	5.4 ± 2.4 ^A
cinnamon 3%+marinade	22	4.6 ± 2.0^{A}	14	5.2 ± 2.6^{A}
cinnamon 5%+marinade	25	4.4 ± 2.6^{A}	15	2.5 ± 2.1^{A}
oregano 1%+marinade	44	5.1±2.2 ^A	43	$5.4 \pm 2.5^{\text{A}}$
oregano 3%+marinade	25	3.7 ± 2.4^{A}	16	6.0 ± 2.5
oregano 5%+marinade	25	3.3 ± 2.9^{A}	14	2.6 ± 2.4^{A}
thyme 1%+marinade	45	4.3 ± 2.5^{A}	43	$5.1 \pm 2.6^{\text{A}}$
thyme 3%+marinade	22	$4.0{\pm}2.0^{A}$	16	4.8±3.3 ^A
thyme 5%+marinade	20	3.8 ± 2.1^{A}	13	3.6 ± 2.4^{A}
fried	number of tests	mean	number of tests	mean
sunflower oil1%+marinade	36	6.5±2.3	34	6.5 ± 2.7
cinnamon 1%+marinade	24	5.5 ± 2.0	20	6.0 ± 2.5
cinnamon 5%+marinade	12	5.7±2.4	11	$4.8{\pm}2.7$ ^A
oregano 1%+marinade	24	$4.5 \pm 2.2^{\text{A}}$	23	5.7±2.6
oregano 5%+marinade	12	$4.7 \pm 3.0^{\text{A}}$	12	$4.7 \pm 3.0^{\text{A}}$
thyme 1%+marinade	24	5.4 ± 2.6	24	5.9 ± 2.8
thyme 5%+marinade	12	5.4±2.7	12	$4.5 \pm 2.4^{\text{A}}$

Table 8. Summary of hedonic values for each treatment and food matrix

Asignificant difference (p < 0.05) from the hedonic value of sunflower oil 1% + marinade