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In vitro effects of Italian Lavandula multifida L. leaf extracts on gilthead seabream (Sparus aurata) leucocytes and SAF-1 cells

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12	In vitro effects of Italian Lavandula multifida L. leaf extracts on gilthead seabream
13	(Sparus aurata) leucocytes and SAF-1 cells
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### 31 Abstract

Lavandulla multifida is very appreciated by pharmaceutical and cosmetic industries. In 32 Italy is only found in Calabria and Sicily and, at present, urge its valorization due to its 33 high extinction and genetic erosion risks. Possible applications of L. multifida extracts 34 on fish were assayed by using gilthead seabream (Sparus aurata) as a marine fish 35 model, due to its importance in fish aquaculture. The in vitro effects of both aqueous 36 and ethanolic leaf extracts obtained from two Italian populations of L. multifida on head 37 kidney leucocyte activities (viability, phagocytosis, respiratory burst and peroxidase 38 content) where compared with those originated for similar extracts obtained from a 39 widely-distributed species of L. angustifolia. Furthermore, the possible cytotoxic effects 40 of all extracts on SAF-1 cells and bactericidal effects for three pathogenic bacteria for 41 fish (Vibrio harveyi, Vibrio anguillarum, Aeromonas salmonicida) were also evaluated. 42

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45 Keywords: Lavandula multifida L.; Leaf extracts; Innate immune response; Sparus
46 aurata; SAF-1 cell line.

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### 49 **1. Introduction**

Lavandula (lavender, belongs to the family Labiatae, Lamiaceae) is a genus of 39 50 species, which has been largely used (either dried or as an essential oil) in folk and 51 52 traditional medicine (Cavanagh and Wilkinson, 2002; Denner, 2009). Lavender has been used for centuries. Several studies revealed that Lavandula species are rich in a wide 53 range of secondary metabolites such as phenolic compounds (Areias et al, 2000; Upson 54 et al, 2000), monoterpenes, diterpenes (Politi et al, 2002), triterpenes (Topcu et al, 55 2001), sesquiterpenes (Ulubelen et al, 1988), and coumarins (Shimizu et al, 1990) 56 which are supposed to be responsible for their antimicrobial, antifungal and antioxidant 57 58 properties. Because of the high content of these secondary metabolites, Lamiaceae family accounts for beneficial and therapeutical properties which allow a wide spectrum 59 of applications in food industry, cosmetic, perfumery and pharmaceutical preparations 60 61 with high industrial and commercial value (Wilkinson et al, 2003).

Among Lavandula species, Lavandula multifida is a short-living plant 30 to 100 cm 62 high, with a diploid genetic pool, equipped with triangular pinnatisect leaves, and blue 63 or white purple flowers which give off a strong smell. It spontaneously grows along the 64 Mediterranean coast of Egypt, Tunisia, Marocco, Algeria, Portugal and Spain, while in 65 66 Italy it has been found only in Calabria and in Sicily (Galesi et al, 2005). The great interest for this plant is due to its essential oils and metabolites (whose antimicrobial, 67 antifungal and antioxidant properties are well-documented) and to the wide possibilities 68 of application in pharmaceutical and cosmetic industries (Ramchoun et al, 2009; Sosa et 69 70 al, 2005; Zuzarte et al, 2012). Unfortunaltely, the human impact on its natural habitat caused the reduction and fragmentation of the populations of this plant in Southern Italy, 71 72 and therefore, due to the rarity and the threats to this species, it was included in the "Regional Red Lists of Italian Plants" under the status IUCN of "critically endangered" 73

in Calabria region and "endangered" in Sicily (Conti et al, 1997). For this important
reason, the valorization of *L. multifida* is strictly necessary due to its high extinction and
genetic erosion risks.

77 Currently, there is a growing interest in screening medicinal plants extracts for their bactericidal, fungicidal and even immunostimulant properties, in order to exploit new 78 biocompounds of natural origin which could be employed in the prevention and/or 79 80 control of fish diseases in aquaculture (Reverter et al, 2014). The importance of using immunostimulants in aquaculture is known since many years (Sakai, 1999) and, at 81 present, it is a topic of great interest in research. In the intensive aquaculture system, 82 83 application of antibiotics and chemotherapeutics as prophylactic measures has been widely criticized for their negative impacts on the environment and also on fish, such as 84 immunodepression or residue accumulation in tissues, and besides this leads to the 85 development of drug resistant pathogens (Rijkers et al, 1980; Harikrishnan et al, 2009; 86 FAO, 2003; Smith et al, 1994). The use of plant extracts as immunostimulants has 87 recently received increasing attention not only because they combine lower costs with 88 the low impact on the environment, but also because they may have additional 89 physiological effects since they contain many nutrients, micronutrients as well as other 90 immunostimulant substances (Cuesta et al, 2005). Taking in considerations the needed 91 92 of looking for new applications of L. multifida extracts, and taking into account the importance of gilthead seabream (Sparus aurata) in Mediterranean and marine fish 93 aquaculture, we developed the present study. The in vitro effects of leaf extracts 94 obtained from two Italian populations of L. multifida on head kidney leucocytes 95 activities (viability, phagocytosis, respiratory burst and peroxidase activity) were tested, 96 97 making a comparison with similar extracts obtained from the widely-distributed species L. angustifolia. Furthermore, the cytotoxic activity on SAF-1 cells (a cell line obtained 98

99 from gilthead seabream) and the bactericidal activity of the extracts on Vibrio harveyi,

100 *Vibrio anguillarum* and *Aeromonas* salmonicida were also checked.

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### 103 **2. Materials and Methods**

104 *2.1. Plant extracts* 

Leaves from Sicilian *L. multifida* L. (collected from Brucoli, Siracusa, indicated in the figures as *L. multifida* S), from Calabrian *L. multifida* L. (collected from Capo dell'Armi, indicated in the figures as *L. multifida* C), and from *L. angustifolia* Miller were collected and air-dried at 50 °C until the weight was stable.

For preparation of aqueous extracts, air-dry leaves were macerated and shaked with boiling water for 4 h at 25°C, then the mixture was filtered twice using nylon net filter with a 100 $\mu$ m pore size, and evaporated in a rotary evaporator until dryness. Prior to be used in the assays, extracts were filtered using sterile filters of 0.22  $\mu$ m of diameter. For preparation of ethanol extracts, air-dry leaves were macerated and shaked with pure ethanol (1:40, 48 h, 25 °C). The resulting mixture was then filtered-twice as described above, and concentrated by vaporizing using a rotary evaporator.

116 *2.2. Animals* 

117 Thirty specimens (40.51  $\pm$  1.47 gr weight) of the seawater teleost gilthead seabream (*S. aurata* L.), obtained from a local farm (Murcia Spain), were kept in re-circulating 119 seawater aquaria (250 L) in the Marine Fish Facility at the University of Murcia. The 120 water temperature was maintained at 20  $\pm$  2 °C with a flow rate of 900 L h<sup>-1</sup> and 28‰ 121 salinity. The photoperiod was 12 h light: 12 h dark. Fish were allowed to acclimatise for 122 15 days before the start of the trial, where they were fed with a commercial pellet diet 123 (Skretting, Spain) at a rate of 2% body weight day<sup>-1</sup>. The fish were killed after starving for 24 h by using an overdose of MS-222 (Sandoz, 100 mg ml<sup>-1</sup> water). All
experimental protocols were approved by the Ethical Committee of the University of
Murcia (Permit Number A13150104).

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### 128 2.3. Head-kidney leucocyte isolation and incubation with extracts

Before the dissection of the head-kidney (HK), the specimens were bled. Blood was 129 collected from the caudal vein and afterwards fish were dissected to obtain HK 130 131 fragments. For isolation of leucocytes to carry out the assays, HK fragments were transferred to 8 ml of sRPMI [RPMI-1640 culture medium (Gibco) supplemented with 132 0.35% sodium chloride (to adjust the medium's osmolarity to gilthead seabream plasma 133 osmolarity of 353.33 mOs), 3% fetal calf serum (FCS, Gibco), 100 i.u. ml<sup>-1</sup> penicillin 134 (Flow) and 100 mg ml<sup>-1</sup> streptomycin (Flow)] (Esteban et al., 1998). Cell suspensions 135 136 were obtained by forcing fragments of the organ through a nylon mesh (mesh size 100 μm), washed twice (400 g, 10 min), counted (Z2 Coulter Particle Counter) and adjusted 137 to  $2 \times 10^7$  cells ml<sup>-1</sup> in sRPMI. Cell viability was higher than 98%, as determined by the 138 139 trypan blue exclusion test.

To study the possible effects of water and ethanolic extracts on leucocyte activities, 140 aliquots of 50 µl of the obtained leucocytes suspension containing  $2 \times 10^7$  cells ml<sup>-1</sup> were 141 142 dispensed into flat-bottomed 96-well microtitre plates (Nunc). Then aliquots of 50 µl well<sup>-1</sup> of water or ethanolic extracts ranging from 10, 100, 500,to 1000  $\mu$ g ml<sup>-1</sup> prepared 143 in sRPMI were added. The extract aliquot was replaced by sRPMI for control samples 144 145 for assays with water extracts, or by 0,1% dymethil sulfoxide (DMSO, Sigma) in RPMI in case of control samples for assays with ethanolic extracts. Cells were incubated in 146 presence of the extracts for 24 h at 20 °C in an incubator with 5% CO<sub>2</sub> and 85% 147

humidity. After incubation, leucocyte viability and phagocitic, respiratory burst andperoxidase activities were determined as described below.

150 *2.4. Leucocyte viability* 

Aliquots of 100 µl of leucocytes previously incubated for 24h without (control) or with 151 the plant extracts were placed in 5 ml glass tubes (Falcon, Becton–Dickinson) and 40 µl 152 of propidium iodide (PI) (400µg ml<sup>-1</sup>, Sigma) were added to each sample. The tubes 153 were gently mixed before analysis in a FACScan (Becton–Dickinson, Madrid, Spain) 154 155 flow cytometer with an argon-ion laser adjusted to 488 nm. Analyses were performed on 5000 cells, which were acquired at a rate of 300 cells s<sup>-1</sup>. Data were collected in the 156 form of two-parameter side scatter (granularity, SSC) and forward scatter (size, FSC), 157 and green fluorescence (FL1) and red fluorescence (FL2) dot plots or histograms were 158 made on a computerised system. Dead cells were estimated as the percentage of cells 159 160 with propidium iodide (red-PI fluorescent cells).

### 161 *2.5. Phagocitic activity*

162 The phagocytic activity of gilthead seabream HK leucocytes was studied by flow 163 cytometry according to Esteban et al. (1998). Heat killed (30 min, 60 °C) lyophilised Saccharomyces cerevisiae, strain S288C, were washed twice, counted and adjusted to 164 108 yeast cells/ml in sRPMI-1640. To label yeast cells with fluorescein isothiocyanate 165 (FITC, Sigma) yeast cells were incubated with 5  $\mu$ g ml<sup>-1</sup> FITC at 22 °C with constant 166 stirring (40 cycles min<sup>-1</sup>) and in darkness for 15 min (Rodriguez et al, 2003). After 167 labeling, free FITC was removed by washing twice in phosphate buffer saline (PBS) 168 and the yeast cells were resuspended in sRPMI-1640. FITC-labeled yeast cells were 169 acquired for flow cytometric study. The staining uniformity was examined and then the 170 171 yeast cell suspensions were aliquoted and stored at 4 °C.

Samples of 125 µl of medium containing FITC-labeled-yeast cells and 100 µl of HK 172 leucocytes in sRPMI were mixed, centrifuged (400 g, 5 min, 25°C), resuspended and 173 174 incubated at 25 °C for 60 min in dark conditions. At the end of the incubation time, the samples were placed on ice to stop phagocytosis and 400 µl of cold PBS was added to 175 176 each sample. The fluorescence of the extracellular yeast cells (i.e. free yeast cells and yest cells adhered to phagocytes but not ingested) was quenched by adding 50 µl of cold 177 Trypan Blue (0.4% in PBS) per sample. Standard samples of FITC-labelled S. 178 179 cerevisiae yeast cells or HK leucocytes were included in each phagocytosis assay. Immediately, the samples were mixed gently, acquired, and analysed in a FACScan. 180 Data were collected in the form of two-parameter side scatter (SSC) and forward scatter 181 (FSC), and green fluorescence (FL1) dot plots or histograms were made on a 182 computerized system. Fluorescence histograms represented relative fluorescence on a 183 184 logarithmic scale. Flow cytometer was set to analyze the phagocytic cells gated from all the leucocytes because of their higher SSC and FSC values. Phagocytic ability was 185 186 defined as the percentage of cells with one or more ingested yeast cells (green-FITC 187 fluorescent cells) within the phagocytic cell population. The relative number of ingested yeast cells per cell (phagocytic capacity) was assessed in arbitrary units from the mean 188 fluorescence intensity of the phagocytic cells. 189

## 190 2.6. Respiratory burst activity

191 The respiratory burst activity of seabream HK leucocytes was studied by a 192 chemiluminescence method (Bayne and Levy, 1991). Samples of 100  $\mu$ l of a PMA/ 193 luminol solution [1 ng ml<sup>-1</sup> phorbol myristate acetate (PMA, Sigma) and 10<sup>-4</sup>M luminol 194 (Sigma) in HBSS with Ca<sup>2+</sup> and Mg<sup>2+</sup>] were added to the HK leucocytes (previously 195 incubated as described above). The plate was shaken and immediately read in a 196 chemiluminometer (BMG, Fluoro Star Galaxy). Measurements were performed in 30

197 cycles of 2 minutes each. The kinetics of the reactions were analyzed and the maximum
198 slope of each curve calculated. Control samples containing leucocytes that had not been
199 incubated with the extracts were also analyzed.

200 2.7. Peroxidase content

201 The total peroxidase content of HK leucocytes was measured according to Quade and 202 Roth (1997). To do this, 5 µl of HK leucocytes (previously incubated as described above) were incubated for 10 min with 0.02% cetyltrimethylammonium bromide 203 204 (CTAB, Sigma) at 60 rpm. Afterwards, 100 µl of 10mM 3,3',5,5'-tetramethylbenzidine hydrochloride (TMB, Sigma) and 5mM H<sub>2</sub>O<sub>2</sub> (both substrates prepared daily) were 205 206 added and after 2 min, 50 µl of 2M sulfuric acid was also added to stop the reaction. 207 The absorbance of the samples was measured at 450 nm in a microplate reader (BMG Fluostar Omega, USA). Control samples containing leucocytes that had not been 208 209 incubated with extracts were also analyzed.

210 2.8. SAF-1 cell culture

The established cell line SAF-1 (ECACC n 00122301) was seeded in 25 cm<sup>2</sup> plastic 211 212 tissue culture flasks (Nunc, Germany) cultured in L-15 Leibowitz medium (Life Technologies, UK), supplemented with 10% fetal bovine serum (FBS, Life 213 Technologies), 2mM L-glutamine (Life Technologies), 100 i.u. ml<sup>-1</sup> penicillin (Life 214 Technologies) and 100 µg ml<sup>-1</sup> streptomycin (Life Technologies). Cells were grown at 215 25 °C and with 85% humidity. Exponentially growing cells were detached from culture 216 flasks by brief exposure to 0.25% of trypsin in PBS, pH 7.2-7.4, according to the 217 218 standard trypsinization methods. The detached cells were collected by centrifugation (1000 x g, 5 min, 25 °C) and the cell viability was determined by the trypan blue 219 220 exclusion test.

221 2.9. Cytotoxicity assay on SAF-1 cell line

Cytotoxicity assay was performed in quadruplicates. When SAF-1 cell lines were 222 approximately 80% confluent, they were detached from flasks culture with trypsin (as 223 described before), and aliquots of 100 µL containing 50000 cells well<sup>-1</sup> were dispensed 224 in 96-well tissue culture plates and incubated (24 h, 25 °C). This cell concentration was 225 previously determined in order to obtain satisfactory absorbance values in the cytotoxic 226 assav and avoided cell overgrowth. After that, the culture medium was replaced by 100 227  $\mu$ L well<sup>-1</sup> of the extracts to be tested at the appropriate dilution. Tested concentrations of 228 water and ethanol extracts ranged from 1 to 1000  $\mu$ g mL<sup>-1</sup> (1, 10, 100, 1000). Cells were 229 then incubated for 24 h. Control samples received the same volume of culture medium 230 (for water extracts) or of DMSO 0,1% (for ethanolic extracts). Cells were incubated for 231 232 24h at 25°C and then their viability determined using the MTT assay.

The MTT assay is based on the reduction of the yellow soluble tetrazolium salt (3-(4,5-233 234 dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) (MTT, Sigma) into a blue, insoluble formazan product by the mitochondrial succinate dehydrogenase (Berridge 235 236 and Tan, 1993; Denizot and Lang, 1986). After incubation with the leaf-extracts, SAF-1 cells with phosphate buffer saline solution (PBS) and 200 µLwell<sup>-1</sup> of MTT (1 mg mL<sup>-</sup> 237 <sup>1</sup>) were added. After 4 h of incubation, the cells were washed and the formazan crystals 238 solubilized with 100µL well<sup>-1</sup> of DMSO. Plates were shacked for 5 minutes in dark 239 240 conditions and the absorbance at 570 nm and 690 nm determined in a microplate reader.

241 *2.10. Bacteria* 

Three pathogenic bacteria for fish (*V. harveyi*, *V. anguillarum* and *A. salmonicida*) and *Escherichia coli*, as control, were used in the bactericidal assays. All bacterial strains were grown from 1 mL of stock culture that had been previously frozen at -80°C. *V. harveyi*, *V. anguillarum* and *A. salmonicida* were cultured for 48 h at 25 °C in Triptic Soy Agar (TSA, Difco Laboratories), and then inoculated in Triptic Soy Broth (TSB,

Difco Laboratories), both supplemented with NaCl to a final concentration of 1% (w/v). 247 Bacteria in TSB medium were then cultured at the same temperature, with continuous 248 shaking (100 rpm) during 24 h. E. coli was cultured in Luria Bertani Agar (LB Agar, 249 Difco) for 48 h at 37 °C and then inoculated in Luria Bertani Broth (LB Broth, Difco). 250 E. coli bacteria in LB broth medium were then cultured for 24 h at 37 °C with 251 continuous shaking (100 rpm). Exponentially growing bacteria were resuspended in 252 sterile Hank's balanced salt solution (HBSS) and adjusted to  $1 \times 10^{8}$  colony forming 253 units (cfu)  $ml^{-1}$ . 254

255 2.11. Bactericidal assay

Bactericidal activity was determined following the method of Stevens et al (1991) with 256 some modifications. Samples of 20 µl of water or ethanolic leaf-extracts previously 257 adjusted to 20, 100, 200, 500, 1000 µg ml<sup>-1</sup> were added in quadruplicate wells of a U-258 259 shaped 96-well plate (Nunc). Hank's balanced solution was added to some wells instead 260 of the extracts and served as positive control. Aliquots of 20 µl of the bacteria 261 previously cultured were added and the plates were incubated for 2.5 h at 25°C (in case 262 of V. harveyi, V. anguillarum and A. salmonicida), or at 37°C (in case of E. coli). After that, 25 µl of MTT (1mg ml-1) were added to each well and the plates were newly 263 incubated for 2 h (at the appropriate temperature taken into account the assayed 264 265 bacteria) to allow the formation of formazan. Plates were then centrifuged (2000 x g, 10 min), being the precipitates dissolved in 200 µl of DMSO and transferred to a flat-266 bottom 96 well-plate. The absorbance of the dissolved formazan was measured at 560 267 268 nm. Bactericidal activity was expressed as percentage of no viable bacteria, calculated as the difference between absorbance of bacteria surviving compared to the absorbance 269 270 of bacteria from positive controls (100%).

271 2.12. Statistical analysis

Results showed are representative of at least three independent experiments and are expressed as Mean  $\pm$  Standard Error. All assays related to leucocyte activities were performed in duplicate and results were expressed as Mean  $\pm$  Standard Error for each group (three fish per group). Data were analyzed by one-way analysis of variance (ANOVA), and Tukey post-hoc test was performed in order to make a multiple comparison between experimental groups. Differences were considered statistically significant when P<0.05.

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280 3. Results
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282 *3.1. Effects of leaves extracts on gilthead seabream head kidney leucocyte* 

Head kidney leucocyte viability was tested after 24 h of incubation with water or ethanolic extracts of the three tested plants in the range of concentrations from 10 to 1000  $\mu$ g ml<sup>-1</sup> (10, 50, 100, 250, 500, 750, 1000  $\mu$ g ml<sup>-1</sup>). Results demonstrated that any significant effect was observed on HK leucocyte viability after incubation with the cited extracts, respect to the values recorded for control HK leucocytes (Fig. 1).

Regarding phagocytosis, incubation of HK leucocytes with water or ethanolic extracts 288 of L. multifida (from both Calabrian and Sicilian populations) significantly enhanced 289 290 their phagocitic ability when using the range of extract concentrations from 10 to 500  $\mu$ g ml<sup>-1</sup> and 10 to 100  $\mu$ g ml<sup>-1</sup>, respectively (Fig. 2). However, after incubation of HK 291 292 leucocytes with L. angustifolia water extracts significant enhancements of the 293 phagocitic ability were only recorded when using highest concentrations (from 100 to 1000 µg ml<sup>-1</sup>) (Fig. 2a). On the other hand, incubation of leucocytes with ethanolic 294 295 extracts from L. angustifolia did not affect their phagocytic ability.

On the contrary, significant decreases were observed in the phagocitic ability of HK leucocytes after being incubated with 500 and 1000  $\mu$ g ml<sup>-1</sup> of ethanolic extracts from all plant tested, except for 500  $\mu$ g ml<sup>-1</sup> ethanolic extract of *L. multifida* S which increased their phagocytic ability (Fig. 2b).

A similar trend was observed in the phagocytic capacity because HK leucocytes incubated with 10  $\mu$ g ml<sup>-1</sup> and 10 to 100  $\mu$ g ml<sup>-1</sup> water and ethanolic extracts from *L. multifida* S, respectively, showed an increased phagocytic capacity (Fig. 3). Instead, water and ethanolic extracts from *L. angustifolia* did not affect significantly the phagocitic capacity of gilthead seabream HK leucocytes, respect to control samples.

Regarding respiratory burst, incubation of HK leucocytes with water extracts from all 305 306 populations of Lavandula tested increased this activity at all concentrations tested in a significant manner, in comparison with the control group, except for L. multifida S 307 water extract which decreased significantly this activity when using at 1000  $\mu$ g ml<sup>-1</sup> of 308 concentration (Fig. 4). Also, ethanolic extracts from both populations of L. multifida 309 used at 10 and 100 µg ml<sup>-1</sup> increased the HK leucocyte respiratory burst activity, while 310 the ethanolic extracts from all plants tested at 1000  $\mu$ g ml<sup>-1</sup> significantly decreased the 311 respiratory burst activity of HK leucocytes compared to the control samples (Fig. 4). 312 The observed enhancements of the respiratory burst activity of leucocytes was always 313 314 higher after incubation with water extracts than with ethanolic extracts.

No significant variations in the peroxidase content of HK leucocytes were observed after being incubated neither with aqueous nor with ethanolic extracts from the three plants tested in the present work, respect to the values recorded for control samples (Fig. 5).

319

## 320 *3.2. Effects of leaves extracts on SAF-1 cell line*

The effects of water and ethanolic extracts of *L. multifida* on cytotoxicity of SAF-1 cells were also evaluated. Results from the cytotoxicity test showed that water extracts from all plants tested did not alter significantly the cell viability, respect to values for control samples (Fig. 6). Interestingly, incubation of SAF-1 cells with *L. multifida* S water extract increased the cell viability in a significant manner at the higher concentration tested (1000 µg ml<sup>-1</sup>) (Fig. 6a).

On the contrary, incubation of SAF-1 cells with ethanolic extracts significantly affect their viability, indeed showing high levels of cytotoxicity at the higher concentration tested (1000  $\mu$ g ml<sup>-1</sup>) (Fig. 6b). Particularly, ethanolic extracts from the two populations of *L. multifida* had a significantly higher level of cytotoxicity on SAF-1 cells if compared to the cytotoxicity provoked on these cells after being incubated with *L. angustifolia* ethanolic extracts (Fig. 6).

333

## 334 *3.3. Bactericidal activity of leaves extracts*

335 In this study L. multifida plant extracts were also tested (in comparison with the extracts 336 from the commercial species L. angustifolia) for their bactericidal activity against some fish pathogens relevant in fish aquaculture: V. harveyi, V. anguillarum and A. 337 salmonicida, being E. coli used as control bacteria. Results from bactericidal assays 338 revealed that any of the assayed plant extracts neither water nor ethanolic affect bacteria 339 viability in a significant manner (Figs. 7-10). Only a slight difference in bacteria 340 viability (80% cell viability) was observed when ethanolic extracts from L. multifida C 341 342 were used to incubate V. harveyi bacteria cells (Fig. 7).

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344

### 346 **4. Discussion**

347 The massive use of antimicrobials for disease control of farmed animals is considered, at present, a real hazard. Indeed food-producing animals are one of several potential 348 sources of antibiotic-resistant bacteria which may spread from animals to man via the 349 food chain (Harikrishnan et al, 2011). Furthermore, pathogens may also transfer their 350 antibiotic-resistance genes into human pathogenic bacteria thus posing a threat to 351 352 human health and considered as one of the main environmental problems (Alderman and Hastings, 1998; Cabello, 2006; Abutbul et al, 2004; Smith et al, 1994; MacMillan, 353 2001). 354

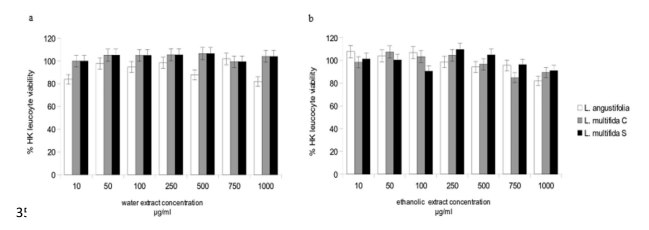


Fig. 1. Percentage of viability of S. aurata HK leucocytes after 24 h of incubation with water (a)
or ethanolic extracts (b) obtained from leaves of Calabrian and Sicilian Lavandula multifida. L.
angustifolia was considered as control plant. Results of cytotoxicity are given as the percentage of
viability, related to the control of untreated cells (the mean optical density of untreated cells was
set to 100% viability). Results showed are representative of at least three independent
experiments and are expressed as Mean ± Standard Error.

Regarding farmed fish, while some immunostimulants cannot be used due to their high cost and/or limited effectiveness (Ringo et al, 2012), treatment with medicinal plants is gaining attention due to the fact that they may be a potentially beneficial alternative to antibiotics. Firstly, these plants mitigate many of the side effects associated with the use of chemicals and antibiotics (Madhuri et al, 2012). Furthermore, treatment of bacterial diseases with different herbs has been safely used in organic agriculture, veterinary and human medicine (Madhuri et al, 2012; Bairwa et al, 2012).

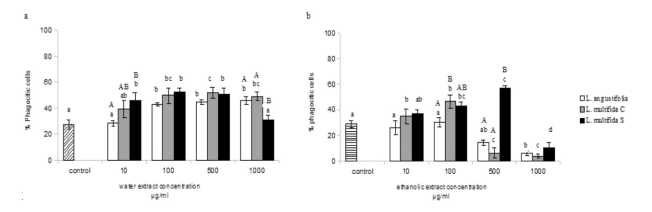
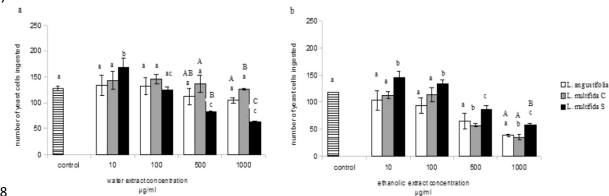
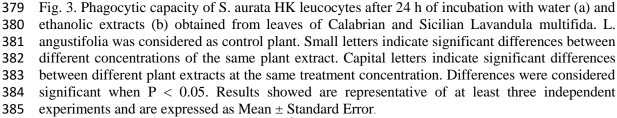


Fig. 2. Percentage of phagocytic cells of S. aurata HK leucocytes after 24 h of incubation with water (a) and ethanolic extracts (b) obtained from leaves of Calabrian and Sicilian Lavandula multifida. L. angustifolia was considered as control plant. Small letters indicate significant differences between different concentrations of the same plant extract. Capital letters indicate significant differences between different plant extracts at the same treatment concentration. Differences were considered significant when P < 0.05. Results showed are representative of at least three independent experiments and are expressed as Mean  $\pm$  Standard Error.



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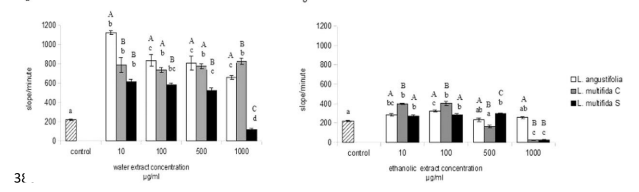


Fig. 4. Respiratory burst activity (expressed as slope/minute) of S. aurata HK leucocytes after
24 h of incubation with water (a) and ethanolic extracts (b) obtained from leaves of Calabrian
and Sicilian Lavandula multifida. L. angustifolia was considered as control plant. Small letters

indicate significant differences between different concentrations of the same plant extract. Capital letters indicate significant differences between different plant extracts at the same treatment concentration. Differences were considered significant when P < 0.05. Results showed are representative of at least three independent experiments and are expressed as Mean  $\pm$  Standard Error

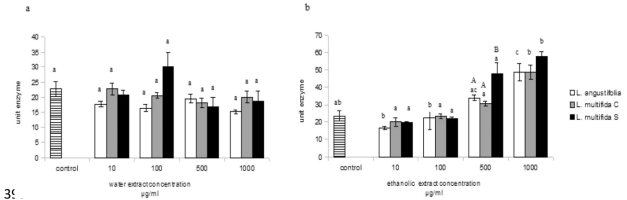


Fig. 5. Peroxidase activity (expressed as unit enzyme) of S. aurata HK leucocytes after 24 h of incubation with water (a) and ethanolic extracts (b) obtained from leaves of Calabrian and Sicilian Lavandula multifida. L. angustifolia was considered as control plant. Small letters indicate significant differences between different concentrations of the same plant extract. Capital letters indicate significant differences between different plant extracts at the same treatment concentration. Differences were considered significant when P < 0.05.

402 Results showed are representative of at least three independent experiments and are expressed
 403 as Mean ± Standard Error.

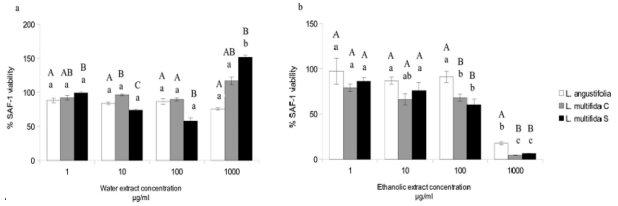
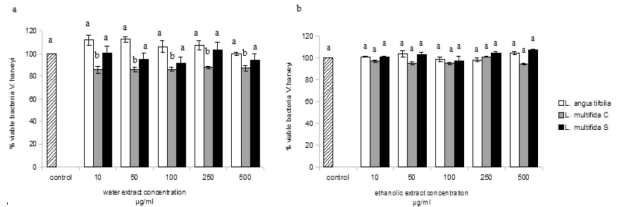
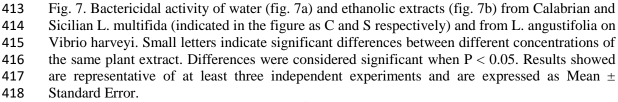


Fig. 6. Viability (expressed as percentage) of SAF-1 cell line after 24 h of incubation with water (a) and ethanolic extracts (b) obtained from leaves of Calabrian and Sicilian Lavandula multifida. L. angustifolia was considered as control plant. Small letters indicate significant differences between different concentrations of the same plant extract. Capital letters indicate significant differences between different plant extracts at the same treatment concentration. Differences were considered significant when P < 0.05. Results showed are representative of at least three independent experiments and are expressed as Mean  $\pm$  Standard Error.





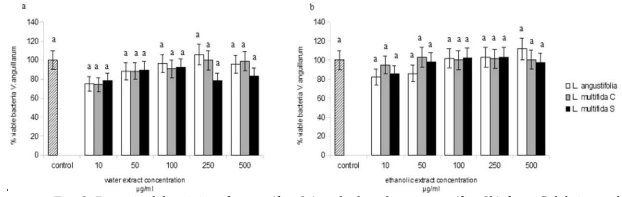
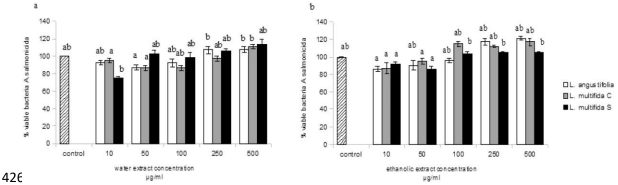
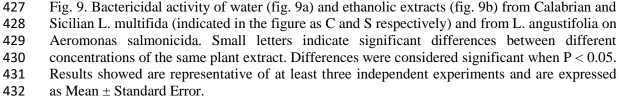
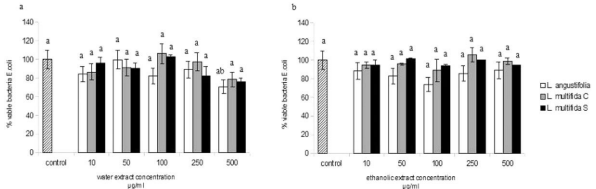


Fig. 8. Bactericidal activity of water (fig. 8a) and ethanolic extracts (fig. 8b) from Calabrian and Sicilian L. multifida (indicated in the figure as C and S respectively) and from L. angustifolia on Vibrio anguillarum. Small letters indicate significant differences between different concentrations of the same plant extract. Differences were considered significant when P < 0.05. Results showed are representative of at least three independent experiments and are expressed as Mean  $\pm$  Standard Error.









434 Fig. 10. Bactericidal activity of water (fig. 10a) and ethanolic extracts (fig. 10b) from Calabrian435and Sicilian L. multifida (indicated in the figure as C and S respectively) and from L.436angustifolia on Escherichia coli. Small letters indicate significant differences between different437concentrations of the same plant extract. Differences were considered significant when P < 0.05.</td>438Results showed are representative of at least three independent experiments and are expressed439as Mean ± Standard Error.

Lately, the use of new and non-purified immunostimulants has received many 441 442 attentions, and an essential prerequisite is that any new potential immunostimulant should always be respectful with the environment (Sakai et al, 1999). Currently 443 increased consumer demand of quality in farmed fish and shellfish, news efforts in order 444 445 to avoid the possible pollutants, use of antibiotics and/or carcinogens during the production process urged. In this scenario, plants or their byproducts are preferred since 446 they content numerous compounds like phenols, polyphenols, alkaloids, quinones, 447 terpenoids, lectines and polypeptides many of which have been shown to be very 448 effective alternatives to antibiotics, chemicals, vaccines, and other synthetic compounds 449 450 (Harikrishnan et al, 2011). Since herbs have little side effects and are easily degradable, various studies have investigated the effect of plant products on innate and adaptive 451 452 immune response and to prevent and control fish and shellfish diseases (reviewed by 453 Valladão et al, 2015). Fruits, leaves, seeds, flowers, roots and barks extracts from many plants have been reported to have immunomodulator porperties (NagaPreethi and 454 Rajeshwari, 2014). Concomitantly, natural plant products promote different activities in 455 456 animals like growth, appetite stimulation, immunostimulation, aphrodisiac, antistress and antimicrobial properties due to the already mentioned numerous active principles 457

458 present in them (Reverter et al, 2014). The necessity of looking for new natural 459 substances with potential application in fish aquaculture due to their immunostimulant 460 properties or biocides activities against potential pathogenic microbes move us to carry 461 out the present work.

Among plants belonging to Lavandula genus, L. multifida represents a remarkable species because of its wide utilization in traditional medicine since ancient times. Most of the studies performed to date focused on the antibacterial, antifungal and antioxidant properties of its essential oils, as well as on their biochemical constituents.

It is well known that biological and therapeutic properties of medicinal plants are 466 closely related to their chemical compounds; due to the differences in solubility in water 467 or in alcohol, each class of these chemical compounds has a preferred effective 468 469 extraction method which facilitate getting out the chemicals out of the plant and into the 470 herbal remedy that is being prepared, and this is the reason why water and ethanolic 471 extracts are obtained in order to analyze the different biological and therapeutic 472 properties (Hashemi et al, 2008). Many plant species are known to synthesize many 473 bioactive secondary metabolites like terpenoids, alkaloids, glycoside, phenolic compounds, polysaccharides, flavonols and tannins which possess various biological 474 properties. Furthermore, it is well-known that phytomedicines are inexpensive, eco-475 476 friendly and easy to be prepared (Hashemi et al, 2008).

Although actually there are no available data concerning the effects of lavender extracts on fish immunity and their possibility of use as immunostimulant in aquaculture, some scientific evidences exist about the effect of extracts from other plants belonging to Lamiaceae family on innate immune response. Extracts from *Thymus guyonii*, *Salvia verbenaca* and *Stachys circinata* (Lamiaceae family) have been tested for their immunostimulant potential on the phagocitic activity in adult male mice (*Mus* 

*musculus*) through the carbon clearance rate test, resulting in an appreciable increasing 483 in phagocitic index in animals when administered plant extracts by intraperitoneal 484 injection (Nassar et al, 2015). Also, leaf extracts from Ocimum sanctum (Lamiaceae 485 486 family) have been tested for their effect on the innate and specific immune responses and disease resistance against Aeromonas hydrophila in Oreochromis mossambicus, 487 resulting in an important immunostimulatory effect when leaf extracts were 488 administered orally and intraperitoneally (Logambal et al, 2000). Similarly, dietary 489 490 administration of peppermint (Menta piperita, Lamiaceae family) promoted growth performance and increased the main hematological and immune humoral parameters in 491 492 fry Caspian white fish (Rutilus frisii kutum) (Adel et al, 2015). Furthermore Zataria multiflora (Lamiaceae) exerted immunostimulatory effects on some immunological 493 factors such as antibody titers, total white blood cells and serum bactericidal activity 494 495 when its essential oils were dietary administered in Cyprinus carpio (Soltani et el, 496 2010). Present results corroborate, firstly that both aqueous and ethanolic extracts of L. 497 multifida have any negative impact on HK leucocyte viability at the concentrations and 498 incubation times tested in the present study. Furthermore, and more interesting, our results demonstrate that L. multifida L. extracts increased HK leucocytes phagocytosis 499 500 and respiratory burst activities, thus supporting the possibility of use of L. multifida L. 501 extracts as immunostimulant in the production of this important farmed marine fish 502 Also, respiratory burst activity showed a significant increase when HK species. leucocytes were treated with water and ethanolic extracts from all populations of 503 504 Lavandula, although a negative effect was observed when using ethanolic extracts at the maximum concentration tested (1000 mg mL<sup>-1</sup>), being this similarly observed for the 505 506 phagocytic ability. Present results show that L. multifida L. extracts positively affect phagocytosis and respiratory burst activity of gilthead seabream HK leucocytes, 507

without any significant effect neither on HK leucocytes viability nor on HK leucocytes
peroxidase, thus supporting the possibility of use of L. multifida L. extracts, in a welldefined range of concentrations, as immunostimulant in the production of this

511 important farmed marine fish species. Immunostimulants are capable of promoting a 512 greater and more effective sustained immune response to those infectious agents 513 (viruses, bacteria, fungi, and parasites), producing subclinical disease without risks of 514 toxicity, carcinogenicity or tissue residues (Muthusamy et al, 2013). Future in vivo 515 studies could demonstrate if similar immunostimulant effects are induced after dietary 516 administration of such extracts to gilthead seabream specimens.

Fish cell lines are increasingly important research tools as an alternative to the 517 experimental animals. The long-term SAF-1 cell line was established in 1996 and it is a 518 fibroblast-like culture derived from gilthead seabream (Béjar et al., 1997). Since then, 519 520 the SAF-1 cell line has proved useful in many applications and varied studies related to virology (Pérez-Prieto et al., 1999; Tafalla et al., 2004; Alonso et al., 2005; Bandín et 521 522 al., 2006; García Rosado et al., 2008; Cano et al., 2016), bacteriology (Acosta et al., 2009; El Aamri et al., 2012), immunology (Ray et al., 2002; Pelegrín et al., 2004), and 523 more recently in nanoparticles interiorization (Trapani et al., 2015), metal toxicity 524 (Morcillo et al., 2016) or freshness and freeze-thawing of seabream fillets (Diop et al., 525 526 2016) studies. Curiously, to the best of our knowledge, only one previous paper focused on the immune modulatory effects of a plant extract (Aloe arborescens) on SAF-1 cells 527 (Picchietti et al., 2013). In the study, SAF-1 cells were treated with Aloe extract at 528 different concentrations (1.2-4.8 mg ml<sup>-1</sup>) and times (24-72 h). The Aloe extract 529 (1.2 mg ml<sup>-1</sup>) induced a synergic effect when cells were also stimulated with LPS- or 530 531 poly I:C which was detected by studying some immune-related gene expression (Picchietti et al., 2013). 532

Present results from the cytotoxic assay developed on SAF-1 cell line after incubation 533 with L. multifida extracts demonstrate no significant effects on cell viability in a limited 534 range of concentrations tested. However, the maximum doses used in our experiments 535 (1000 mg mL<sup>-1</sup>) affected their viability in a significant manner, causing an increase or a 536 decrease in this parameter when SAF-1 cells were incubated with water or ethanolic 537 extracts, respectively. These data agreed with results obtained from the determination of 538 phagocytosis and respiratory burst activities: indeed ethanolic extracts from L. multifida 539 540 C and S provoked a significant decrease in the number of yeast cells ingested after 24 h of incubation at 1000 mg mL<sup>-1</sup> of concentration, furthermore also the percentage of 541 phagocytic cells showed a significant decrease when HK leukocytes were incubated 542

with ethanolic extracts from all plants tested during 24 h at 1000 mg mL<sup>-1</sup> of concentration, and a similar trend was observed also for the respiratory burst activity. Although the main components of the leaves extract have not been studied yet, the analysis of the chemical composition is one of our future next step. These could allow to know if differences in the biochemical components between water and ethanolic extracts exist, which may explain the different properties of the extracts.

Finally, present results demonstrated that L. multifida L. extracts had no bactericidal 549 activity against the fish pathogens tested. There are no many available data about the 550 551 antibacterial activity of L. multifida L. extracts until now, and the majority of antibacterial studies have been performed on its essential oils. As reported by Khadir 552 and colleagues (2016), L. multifida essential oils were tested against methicillin-553 554 resistant Staphylococcus aureus (MRSA) using disc diffusion method, revealing a good anti-MRSA activity, whereas the ethanolic extract was less active, thus suggesting that 555 556 different biological effects between leaf extracts and essential oils could be due to differences in their chemical composition. In fact the chemical composition of L. 557

multifida essential oils studied by GC and GC-MS (Khadir et al, 2016) revealed that 558 carvacrol is the main component (from 27.5% to 57%), being carvacrol a monoterpenic 559 phenol responsible for many biological activities such as antimicrobial, antitumor, 560 antimutagenic, antigenotoxic, analgesic, antiinflammatory, antiparasitic, antiplatelet 561 (Baser, 2008). Instead, in a recent work ethanolic leaf extract of L. multifida L. from 562 Southern Italy was screened by UV spectroscopic analysis (Fazio et al, 2014), revealing 563 the presence of vitexin and flavones glucoside derivates of hypolaetin, scutellarein, 564 565 luteolin, isoscutellarein, apigenin and chrysoeriol, which were reported to possess mainly antiinflammatory, anticancer and antioxidant properties (Alcaraz and Hoult, 566 1985; Lin et al, 2008; Saeed et al, 2015; Choi et al, 2005). On the other hand, opposite 567 results were obtained with ethanolic extracts from Lavandula officinalis, Melissa 568 officinalis, Ocimum basilicum, Origanum vulgare, Rosmarinus officinalis and Salvia 569 570 officinalis, all belonging to Lamiaceae family, which were tested for antimicrobial 571 activity, exhibiting a broad spectrum of inhibitory effects on some fish pathogens like L. 572 anguillarum, Y. ruckeri, P. damselae, L. garvieae (Bulfon et al, 2014).

To conclude, L. multifida L. water and ethanolic leaf extracts up to 100 mg mL<sup>-1</sup> of 573 concentration increased innate immune activities of S. aurata HK leucocytes, more 574 concretely, their phagocytic and respiratory burst activities, but did not exert any 575 576 bactericidal activity on the bacterial strains tested in the present study, and did not affect significantly SAF-1 cells viability up to 100 mg mL<sup>-1</sup> of concentration. Although the 577 extracts did not exert any bactericidal activity directly on the assayed bacteria strains, 578 579 present results suggest the possibility of use such extracts in in vivo studies in order to corroborate if could be possible using those extracts in aquaculture in order to achieve 580 581 protection against pathogenic infections through enhancement of the innate immunity of

582 fish.

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