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# Accepted Manuscript

Possible effect of hala extract (*Pandanus tectorius*) on immune status, anti-tumour and resistance to *Yersinia ruckeri* infection in rainbow trout (*Oncorhynchus mykiss*)

Elham Awad, Dawn Austin, Alastair Lyndon, Amani Awaad



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1 **Possible effect of hala extract (*Pandanus tectorius*) on immune status, anti-**  
2 **tumour and resistance to *Yersinia ruckeri* infection in rainbow trout**  
3 **(*Oncorhynchus mykiss*)**

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7 Elham Awad<sup>a,b\*</sup>, Dawn Austin<sup>a</sup>, Alastair Lyndon<sup>a</sup>, Amani Awaad<sup>c</sup>.

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10 <sup>a,b</sup> Institute of Life and Earth Sciences, Heriot-Watt University, Edinburgh, UK

11 <sup>b</sup> Department of Hydrobiology, National Research Center, Cairo, Egypt

12 <sup>c</sup> Pharmacognosy Department, College of Pharmacy, Salman Bin Abdulaziz University, Al-  
13 Kharj, Saudi Arabia

14  
15  
16 **\*Corresponding author:** E. Awad, Department of Hydrobiology, National Research Centre, Cairo,  
17 Egypt.

18 E-mail address: elhamsawad@yahoo.com

19 Tel.: +20-1003060046

20 Fax: +20-233370931

21

**Abstract:**

The possible effect of dietary administration of hala extract (*Pandanus tectorius*) on rainbow trout (*Oncorhynchus mykiss*) immune status as well as its effect as an anti-tumour agent was studied. Fish were divided into 4 groups before feeding with commercial diet (0%, control; 0.5%, 1% and 2% of hala extract) for 2 weeks. The effect of diet on the humoral immune parameters, ie total protein, myeloperoxidase content, antiproteases, lysozyme and bactericidal activities were studied. Also, the effect of the diets on the expression of some immune-related genes in rainbow trout head-kidney (*TNF*, *LYZ2*, *IL-8* and *CD-4*) as well as tumour suppressor gene (*WT-1a*) was investigated. At the end of the feeding trial fish groups were challenged with *Yersinia ruckeri*. The results demonstrated enhancement in all the immune parameters in fish fed hala extract diets compared to control fish especially with the highest dose (2%) which recorded the highest significant increase ( $p < 0.05$ ) in some parameters (total protein, myeloperoxidase content, antiproteases, and bactericidal activities) compared to the control. The results obtained from challenge with *Y. ruckeri* revealed reduction in the mortalities in fish groups fed with 1% and 2% doses of hala extract. Feeding with hala extract provoked upregulation in all immune-related genes. Again, the highest dose of hala extract showed a significant upregulation in *WT1a* expression ( $p < 0.05$ ). The current study suggest that the hala extract, especially the highest dose, could be considered a good food additive to improve the immune status, resist tumour formation and to resist or control infectious diseases of rainbow trout.

42

43

**Key words:**

Rainbow trout, *Pandanus tectorius* , immune response, *Yersinia ruckeri*, anti-tumour.

## 46 1. Introduction:

47 Aquaculture is considered to provide a valuable source of essential protein required for  
48 human health. However, the intensive and extensive aquaculture industry is subject to disease  
49 outbreaks [1]. However, controlling fish diseases by antibiotics and chemotherapeutics has  
50 caused the development of drug resistant pathogens in addition to accumulation of residues in  
51 environment and fish tissue and subsequently in humans [2]. On the other hand, medicinal  
52 plants provide a promising, alternative method for resisting and/or controlling fish diseases  
53 [3]. The world tends to use medicinal plants for treatment not only because they are cost  
54 effective, biodegradable, and safe but also for long lasting effects than the synthetic drugs  
55 provide and which have faster recovery rates [4]. It is worth mentioning that many medicinal  
56 plants have antioxidant properties which delay or prevent oxidative damage, therefore  
57 playing a vital role in disease prevention. [5].

58 Previous studies showed a marked enhancement of the fish immune system after  
59 administration of different parts of medicinal plants (roots, leaves, seeds, and flowers).  
60 Moreover, various levels of immune response depended upon plant concentrations, time and  
61 method of administration [3]. For example, dietary supplement with three doses of fenugreek  
62 seeds improved the immune status of gilthead seabream (*Sparus aurata* L.) especially with  
63 the highest dose (10%) [6]. Moreover, common carp (*Cyprinus carpio*) showed enhancement  
64 in immune parameters after administration of a diet supplemented with a 2% dose of *Achillea*  
65 *wilhelmsii* leaf extract [7]. Interestingly, catfish (*Clarias gariepinus*) injected with 50 mg/kg  
66 of leek leaf extract showed an increase in humoral immune response one month post-  
67 injection [8].

68

69 Hala tree (*Pandanus tectorius*) belongs to the family Pandanaceae that comprises around 600  
70 members. [9]. This tree was initially cultured in Asia and extends to tropical northern  
71 Australia and Pacific islands of Oceania [10]. It contains triterpenoids and flavonoids [11],  
72 thus it was successfully used in folk medicine in many countries. In Kiribati, the leaves are  
73 used as therapy for cold, influenza, asthma, hepatitis, boils and cancer, while the roots are  
74 used to treat haemorrhoids. Fruits, flowers and aerial roots are used to treat digestive and  
75 respiratory disorders in Hawaii. While, In Palau, roots and leaves are used to alleviate  
76 stomach cramps and vomiting, respectively [10].

77 Previous studies recorded antioxidant, antibacterial, anticoagulant, anti-inflammatory,  
78 hepatoprotective, antidiarrheal, anticonvulsant, diuretic and anti-cancer activities for leaf  
79 extract [10, 12]. However, there has been no investigation carried out on the effect of hala  
80 leaf extract on the immune system of fish. Thus the current study was carried out to  
81 investigate the possible effects of dietary supplement of hala leaf extract on rainbow trout  
82 immune response either in serum or in cell by examine the expression of some immune-  
83 related genes (*TNF*, *LYZ2*, *IL-8* and *CD-4*) in head kidney as well as study its effect as anti-  
84 tumour agent by examination of the expression of *WT-1a* gene (tumour suppressor gene).

85

## 86 2. Materials and methods:

### 87 2.1. Preparation of plant extract and diets:

88 Hala (*Pandanus tectorius*) leaves were collected from a local market in Saudi Arabia. About  
89 one kilogram of dried powder of hala was extracted using 95% alcohol (Merck, Germany) by  
90 percolation till exhaustion (4 X 4 l) and filtered off by filter paper. The combined filtrates of  
91 the plant were evaporated under reduced pressure and low temperature using rotator

92 evaporator. The obtained residue (250 g) was used in preparing the diets. Four different  
93 concentrations were prepared; commercial diet non-supplemented (0%, control), commercial  
94 diet supplemented with 0.5 g (0.5%), 1 g (1 %) and 2 g (2%)/ 100 g of hala extract.

95

## 96 2.2. Fish, experimental design and sampling:

97 Rainbow trout (*Oncorhynchus mykiss*) of average weight  $18 \pm 1$  g were obtained from a  
98 commercial fish farm in Scotland, and acclimatized in aerated free flowing freshwater ( $14 \pm$   
99  $2^{\circ}\text{C}$ ). During acclimatization, fish were fed three times daily with a commercial diet  
100 (Biomar). Fish were distributed randomly into 4 groups each with 30 fish (10 per replicate)  
101 and fed for 14 days with 0.5 g (0.5%), 1g (1%) and 2 g (2%)/100 g of *hala* extract. Controls  
102 were fed with commercial diet only to examine the possible mode of action and effect on  
103 immune status. Blood was collected from fish anaesthetised using 3- amino benzoic acid  
104 ethyl ester; Sigma-Aldrich, Basingstoke, U.K.) by syringe before transfer to Vacuettes  
105 without heparin (Greiner, Stonehouse, U.K.) and left to clot for 2 h at  $4^{\circ}\text{C}$ , prior to  
106 centrifugation (1600 g, 25 min,  $4^{\circ}\text{C}$ ), and stored at  $-20^{\circ}\text{C}$  until use. The fish were then  
107 sacrificed using an overdose of the above anaesthetic.

## 108 2.3. Humoral immune parameters:

### 109 2.3.1. Lysozyme activity:

110 Serum lysozyme activity was measured according to [13]. Briefly, 60  $\mu\text{L}$  of serum was  
111 added to 2 mL of a suspension of *Micrococcus lysodeikticus* ( $0.2 \text{ mg ml}^{-1}$  in a 0.05 M sodium  
112 phosphate buffer (pH 6.2) and absorbance was measured at 530 nm after 0.5 and 4.5 min on a  
113 spectrophotometer. A unit of lysozyme activity was defined as the sample amount causing a  
114 decrease in absorbance of  $0.001 \text{ min}^{-1}$ .

## 115 2.3.2. Total protein content

116 Total protein was measured by Bradford assay using bovine serum albumin (BSA) as the  
117 standard. Briefly, 2 mg ml<sup>-1</sup> solution of BSA was prepared and serial dilutions made with  
118 phosphate buffer saline (PBS). Around 20 µl of each dilution was added to 1 ml of Bradford  
119 reagent (Sigma-Aldrich) before incubated at room temperature for 15 min. The standard  
120 curve was prepared by measuring the absorbance of each sample at 595 nm verses the sample  
121 concentration. Serum samples were diluted (1: 100) in PBS before 20 µl of each serum  
122 dilution was added to 1 ml of Bradford reagent. After incubation for 15 min, the absorbance  
123 of the unknown samples was taken and plotted onto the standard curve to obtain the total  
124 protein content for each sample [14].

## 125 2.3.3 Antiproteases activity

126 The serum anti-trypsin activity was measured according to Lange, Guðmundsdottir [15].  
127 Thus, 20 µl of standard trypsin solution (Sigma-Aldrich, 5 mg ml<sup>-1</sup>) was incubated with 20 µl  
128 of serum for 10 min at 22°C. Subsequently, 200 µl of 0.1 M PBS (PH 7.2) and 250 µl of 2%  
129 azocasein solution (Sigma-Aldrich, 20 mg ml PBS<sup>-1</sup>) were added. The mixture was incubated  
130 for 1 h at 22°C before stopping with the addition of 500 µl of 10 % (v/v) trichloro acetic acid  
131 (TCA). Then, the mixture was incubated for 30 min at 2°C before centrifuging at 6000 x g for  
132 5 min. About 100 µl of the supernatant was transferred to a 96 microwell flat bottom plate  
133 containing 100 µl of 1 N NaOH well<sup>-1</sup>. The absorbance was read in the ELISA reader at 410  
134 nm. Positive control (100%) was prepared by replacing the serum with buffer. For a negative  
135 control, buffer replaced both serum and trypsin. The percentage inhibition of trypsin activity  
136 was calculated by comparing with a positive control sample.

137



#### 138 2.3.4. Myeloperoxidase content:

139 The myeloperoxidase content of serum was measured according to [16]. Briefly, 50 µl serum  
140 was diluted with 135 µl of Ca<sup>+2</sup> and Mg<sup>+2</sup> free HBSS (Sigma-Aldrich) in flat-bottomed 96-  
141 well plates. Then, 50 µl of 20 mM 3,3',5,5'-tetramethylbenzidine hydrochloride (TMB,  
142 Sigma-Aldrich) and 5 mM H<sub>2</sub>O<sub>2</sub> (Sigma-Aldrich) were added (both substrates of peroxidase).  
143 The reaction was stopped by adding 50 µl of 4 M sulphuric acid (H<sub>2</sub>SO<sub>4</sub>) after 2 min. The  
144 absorbance was read at 450 nm by ELISA reader. Blank sample was without serum.

#### 145 2.3.5. Bacterial culture and bactericidal activity:

146 *Yersinia ruckeri* was identified and obtained from Heriot Watt University before inoculation  
147 into TSA media (Oxoid) for 48 h at 25 °C. The culture was centrifuged for 10 min at 3000 ×g  
148 at 4 °C and the pellet resuspended in 0.9% of saline. The bacterial suspensions were counted  
149 using a hemocytometer slide at a magnification of 400× on a light microscope.

150 Serum bactericidal activity was done according to Kajita, Sakai [17] using *Yersinia ruckeri*.  
151 Briefly, 100 µl of serum was mixed with 100 µl of bacterial suspension, before incubation for  
152 1 h at 25 °C. A blank control was also prepared by replacing serum with sterile PBS. The  
153 mixture was then diluted with 0.05 M sodium phosphate buffer, PBS (pH 6.2) at a ratio of  
154 1:10. Around 50 µl of mixture was plated onto the nutrient agar plates and incubated for 48 h  
155 at 25 °C before the number of colonies was counted.

#### 156 2.4. Challenge with *Yersinia ruckeri*:

157 All challenge experiments were done under a UK Home Office Project License (held by A.R.  
158 Lyndon) and a UK Home Office Personal License (held by D. A. Austin) and approved by  
159 the Heriot-Watt University Ethics Committee.

160 *Yersinia ruckeri* was grown in nutrient broth (Oxoid) for 24 h at 25°C. The culture was  
 161 centrifuged at 3000 x g for 10 min at 4°C, before the supernatants were discarded, and the  
 162 pellets resuspended in 0.9% (w/v) saline. The challenge test was carried out on fish by  
 163 intraperitoneal injection with 0.1 ml volumes containing 10<sup>3</sup> cells/fish, as preliminary  
 164 experiments had determined this to be the LD80 for the challenge strain of *Y. ruckeri*.  
 165 Mortalities were recorded for up to 10 days [18].

#### 166 2.5. Gene expression by real time PCR:

167 Total RNA was extracted from the head kidney and liver using RNA extraction kit (Applied  
 168 biosystem, UK). The quantity of RNA was measure using nanodrop. Around 1 ug of RNA was used  
 169 in Reverse transcription by using cDNA kit from Fermentas (York, UK).

170 The expression of *TNF*, *LYZ2*, *IL-8*, *CD-4*,  $\beta$ -*actin*, and *WT-1a* genes (Table 1) in head kidney were  
 171 analysed by real-time PCR machine (ABI PRISM 7500 instrument, Applied Biosystems) using SYBR  
 172 Green PCR Core Reagents (Applied Biosystems). Mixture (comprised of 10 ml of SYBR Green  
 173 supermix, 5 ml of primers (0.6 mM each) and 5 ml of cDNA template) were incubated for 2 min at 95  
 174 °C, followed by 40 cycles of 15 s at 95 °C, 1 min at 60 °C, and finally 15 s at 95 °C, 1 min at 60 °C  
 175 and 15 s at 95 °C. Gene expression was corrected by the reference gene,  $\beta$ -actin in each sample. The  
 176 primers used are shown in Table 1. Gene expression of the samples compared to the controls was  
 177 calculated according to the following equation:

$$178 \quad (E_{\text{target}})^{\Delta C_{T \text{ target (control - sample)}}}$$

179 Ratio =  $\frac{\quad}{(E_{\text{EF1}\alpha})^{\Delta C_{T \text{ target (control - sample)}}}}$

181

#### 182 2.6. Statistical analysis

183 Data were expressed as fold increase (mean  $\pm$  standard error, SE), obtained by dividing each sample  
184 by the mean control value. Values higher than 1 express an increase while values lower than 1 express  
185 a decrease in the indicated gene. Data were analyzed by one-way analysis of variance (ANOVA).  
186 When differences were found among treatments, Tukey's test was used to compare means by  
187 Minitab statistical software (Minitab, Coventry, UK). Differences were considered significant at P  
188  $<0.05$ .

### 189 3. Results:

#### 190 3.1. Humoral immune parameters:

191 In Lysozyme activity and Myeloperoxidase content (Fig. 1 & 2), the highest significant values ( $p <$   
192  $0.05$ ) were reported in fish groups fed with 0.5% and 2% of hala extract with respect to the values  
193 found in the control group (0%). Although all doses of hala extract showed highly serum total protein  
194 compared to the control (0%) none of them showed significant differences compared to the values  
195 found in control group (Fig.3). The antiprotease activity (Fig.4) was increased in specimens fed with  
196 0.5% or 2% doses of hala extract (compared to control), although only in the group fed 2% dose the  
197 increases were statistically significant, compared to the values found in control ( $p < 0.05$ ). Regarding  
198 bactericidal activity (Fig. 5), all groups fed with hala extract showed high activity compared to the  
199 control group (especially with 2% dose) but without significant differences.

#### 200 3.2. Challenge with *Yersinia ruckeri*

201 Fish groups fed with diets supplemented of hala extract for 2 weeks resulted in reduction in  
202 mortalities after challenge with *Y. Ruckeri* (Fig.6). The resistance to *Y. ruckeri* infection was increased  
203 in fish groups fed for 2 weeks with 2% and 1% doses of hala extract where the survival percent was  
204 26.67% and 21.43 %, respectively, compared to control group (33.3 %). However, the difference was  
205 not significant.

#### 206 3.3. Gene expression:

207 The results revealed an increase in the expression of immune related genes (*TNF*, *LYZ2*, *IL-8*  
208 and *CD-4*) in the head kidney of rainbow trout after administration of diets enriched with  
209 0.5% and 1% of hala extract for 2 weeks compared to the expression recorded in the control  
210 group (Fig. 7). Interestingly, the lowest dose (0.5%) showed the highest expression compared  
211 to other groups and control group while only being statistically significant ( $p < 0.05$ ) for *TNF*  
212 and *IL-8* compared to the control.

213 Moreover, the results showed significant up- regulation in *WT-1a* gene of head kidney ( $p <$   
214  $0.05$ ), in fish fed with 1% and 2% doses of hala extract compared to control. The highest  
215 expression was recorded in the dose of 2%.

216 Discussion:

217 Lysozyme is an important non-specific immune parameter which plays a vital role in fish defence  
218 mechanisms against diseases. It is responsible for opsonin, and thus activates the complement system  
219 and phagocytes [19]. Our results revealed a significant enhancement in serum lysozyme activity of  
220 the fish group fed 0.5% and 2% of hala extract, respectively, for 2 weeks compared to the control  
221 group (0%). This could be attributed to a dose dependent effect of hala extract on rainbow trout.  
222 Similar observation was reported in lysozyme activity of rainbow trout fed for 2 weeks with 0.5% and  
223 1% of tetra (*Cotinus coggyria*) [20], and 1% & 2% of black cumin oil (*Nigella sativa*) [21], . Also, the  
224 highest activity depended mainly on the dose administration.

225 Myeloperoxidase is an important enzyme expressed mainly in neutrophils, which have the ability to  
226 produce hypochlorous acid from one of the oxidative radicals ( $H_2O_2$ ) [22]. This process has a great  
227 benefit to kill invading microorganisms [23]. The results showed an increase in myeloperoxidase  
228 content in all treatment groups with hala extract, especially in those fed with 0.5 % and 2% where a  
229 significant value ( $p < 0.5$ ) was recorded compared to control group. Similar to our study,  
230 myeloperoxidase was increased significantly in rainbow trout fed for two weeks with 1% and 2% of  
231 lupin (*Lupinus perennis*), mango (*Mangifera indica*), and nettle (*Urtica dioica*) [3]. Also, the highest

232 myeloperoxidase content and lysozyme activity values were recorded in rainbow trout fed 0.1% and  
233 0.5% of caper leaf extract for 4 weeks [19]. It is worth mentioning that time and dosage are two  
234 important factors which control the efficiency of plant immunostimulants. For example,  
235 Christyapita, Divyagnaneswari [24] noticed a significant increase in common tilapia fed with diets  
236 containing different concentrations of false daisy leaf for 1 week, while feeding for 2 or 3 weeks  
237 didn't showed any significant increase.

238

239 Serum protein is an important parameter in humeral immune system in fish. Its composition plays a  
240 vital role in keeping fish healthy. Moreover, the most important role played by acute phase proteins is  
241 in limiting the spread of infectious agents through repairing tissue damage and killing micro-  
242 organisms [25, 26]. Present results also demonstrated enhancement in total protein value in all groups  
243 that received different concentration of hala extract groups as compared to the control (especially with  
244 the dose of 2%). This is in agreement with Dügenci, Arda [27] who reported increases in serum  
245 protein levels in rainbow trout fed with 0.1% and 1% of ginger (*Zingiber officinale*), nettle (*Urtica*  
246 *dioica*) and mistletoe (*Viscum album*). Several studies reported an increase of serum protein levels in  
247 fish species after using dietary supplement with plants as immunostimulants [6, 7, 28, 29]. Moreover,  
248 they suggested that elevation in fish total protein were probably a result of enhancement of the non-  
249 specific immune response.

250 Antiproteases or protease inhibitors are active molecules in the non-specific immune system that  
251 inhibit the action of proteases either by binding to their active sites or by 'trapping' the protease to  
252 prevent protein hydrolysis [30] and thus limiting the growth of invading bacteria in fish [31]. In the  
253 present study, the highest dose of hala extract recorded the highest significant enhancement in serum  
254 antiproteases compared to the control. In agreement with our result several studies recorded  
255 enhancement in fish species after administration of diets supplemented with medicinal plants [6, 32,  
256 33]. It is worth emphasizing that the increase in fish immune response depend mainly on dose and  
257 time of administration. Moreover, the response also depends on fish species. For example, 0.5% dose

258 of garlic reported the highest antiprotease activity in rainbow trout after feeding for 2 weeks [33].  
259 Although, the highest antiprotease in Asian seabass fed for 2 weeks was reported at 1.5% dose of  
260 garlic [34].

261 Various humoral molecules involved in non-specific immune response have a power to protect the  
262 fish from invading microbes [35]. Serum bactericidal activity is a lysin mechanism known for the  
263 killing and clearing of pathogenic organisms in fish [31]. In our study, *Y. ruckeri* was used as a model  
264 to examine the activity of hala extract to kill the bacterial infection. The strength of immune  
265 molecules in fish serum to kill *Y. ruckeri* can be detected by the lowest number of bacterial colonies  
266 grown on media. Fish groups fed with hala extract revealed higher bactericidal activity compared to  
267 control, especially in the group fed with the highest dose (2%). Similarly, using the highest dose of  
268 black cumin seed (3%) as food supplement in rainbow trout diet caused higher bactericidal activity against  
269 *Aeromonas hydrophila*, [21]. Also, rohu (*Labeo rohita*) recorded an enhancement in serum  
270 bactericidal activity against *A. hydrophila* after feeding for 2 weeks with doses of prickly chaff-flower  
271 seed and the activity was elevated with higher concentration of seeds [35].

272 Challenge with target pathogen is one of the most valuable tests to evaluate the efficiency of  
273 immunostimulant to resist microbes. The resistance level of fish can be recognized from the survival  
274 percent after a bacterial infection [36]. The results revealed that dietary supplement with hala extract  
275 relatively increased the resistance of rainbow trout against *Y. ruckeri*, where the highest doses (2%  
276 and 1% respectively) recorded the highest resistance. In agreement with our study using stinging  
277 nettle (*Urtica dioica*) as food supplement of rainbow trout diet increased the resistance to *Y. ruckeri*,  
278 especially with the highest dose [37]. However, some immunostimulant can enhance the resistance of  
279 fish against some bacteria but failed to resist the other. For example an improvement in survival  
280 percent of rainbow trout fed with probiotic for 2 weeks was recorded following challenge with  
281 *Aeromonas salmonicida*, *Vibrio ordalii*, and *Y. ruckeri*, but not so with *V. anguillarum* [38].  
282 Previous studies showed reduction in mortality against bacterial infections in fish after using plant  
283 immunostimulant [3, 39, 40].

284

285 *TNF- $\alpha$* , is one of pro-inflammatory cytokines, that mainly produced by activated  
286 monocytes/macrophages and regulate the expression of many cytokines [41]. Many studies revealed  
287 that using plant immunostimulants in fish can induce pro-inflammatory responses. For example, using  
288 0.1% of caper as supplement in rainbow trout diet caused up-regulation in the expression of *TNF- $\alpha$*  of  
289 head kidney [42]. Also, an up-regulation in *TNF- $\alpha$*  expression was observed in common carp treated  
290 with *Rehmannia glutinosa* in spleen, head kidney and gut [43]. Moreover, common carp fed with  
291 different concentrations of guava leaf powder showed an increase in *TNF- $\alpha$*  expression in the head-  
292 kidney, hepatopancreas, and intestine [44]. Similarly our study demonstrated up-regulation of *TNF- $\alpha$*   
293 expression of head kidney in fish groups fed with hala extract compared to control. Increasing in the  
294 *TNF- $\alpha$*  levels could be attributed to the activity of the compounds in hala extract like flavonoids and  
295 antioxidant [12].

296 *IL-8* is another pro-inflammatory cytokine, that produced in response to many stimulation factors like  
297 cytokines, LPS and viruses [45]. It plays an important role in attract T-lymphocytes and neutrophils to  
298 sites of inflammation [46]. Similar to *TNF- $\alpha$*  expression, result demonstrated an increase in *IL-8* in  
299 rainbow trout head kidney in groups received 0.5% and 1 % doses of hala extract. Although, only 0.5%  
300 dose recorded significant difference with the control. In agreement with this study, *IL-8* expression  
301 increased in rainbow trout head kidney after fed diets supplemented with 1% and 2% of stinging  
302 nettle [47] and 0.1% of caper [42]. Interestingly, *IL-8* expression in rainbow trout spleen was  
303 unaffected by the green tea supplementation, while in head kidney the expression showed significant  
304 increase especially in fish fed with 500 mg kg<sup>-1</sup> of green tea [48]. Similarly, gilthead seabream fed  
305 dietary supplement with 10% fenugreek showed enhancement in *IL-8* expression of head kidney after  
306 4 weeks [6].

307 *CD4* is an important co-receptor has been reported in teleosts, expressed on T- helper cell,  
308 monocytes and macrophages [49]. *CD4* is binding to major histocompatibility complex (MHC) class I  
309 molecules on the antigen-presenting cells, stabilizing the interaction between the T cell receptor

310 complex (TCR) and the MHC [50]. The results demonstrated an increase in *CD4* expression in head  
311 kidney in group fed with hala extract compared to control group. Similar observations have been in  
312 other fish species after administration immunostimulant. For example; Atlantic salmon (*Salmo salar*)  
313 injected with lipopolysaccharide (LPS) and  $\beta$ -glucan as immunostimulant, showed an enhancement in  
314 the expression of *CD4* in head kidney [51]. Also, an increase in *CD4* expression of posterior intestine  
315 of European sea bass (*Dicentrarchus labrax*) has been reported after feeding diets contain low level of  
316 synbiotic additive (mannan oligosaccharides) [52]. The slightly higher expression values of *CD4* of  
317 fish group fed with hala extract may indicate an early adaptive immune response.

318 Lysozyme gene is a bactericidal enzyme that mainly present in lymphoid tissue like head kidney and  
319 thymus as well as serum, mucus, gills [53]. There is two types of lysozyme (types I and II), have been  
320 identified in the kidney of rainbow trout. Particular, lysozyme type II (*LYZ2*) showed a potential  
321 antibacterial activity against four gram-negative bacteria. Such finding supports the role which  
322 lysozyme plays in non-specific immune defence in fish [54]. Our study reported the highest level of  
323 *LYZ2* expression in head kidney of fish group fed with the lowest dose of hala extract (0.5%), this is  
324 agreement with the result obtained from analysis serum lysozyme level for the same group. Similar  
325 observation have been reported in common carp fed for 8 weeks with diet supplements with date palm  
326 fruit (200 ml kg<sup>-1</sup>) where showed a remarkable increase in the expression level of lysozyme gene in  
327 head kidney (*LYZ2*) and serum lysozyme compared to control [55].

328 The kidney is plays important functions in fish, not only in immune system for production the  
329 leucocytes but also as osmoregulatory function [56]. Thus any malfunction in this organ in really  
330 preferable. Wilms' tumour suppressor *WT1* is a tumor suppressor gene, any mutation can led to  
331 Wilms' tumour, a pediatric kidney cancer [57]. *WT1* is a modulatory gene involved in cell growth and  
332 development of the urogenital system development (kidney and gonad) [58]. *WT1* have been  
333 identified in fish species [59-61]. The suppression of *WT1* led to appear the edema in zebrafish which  
334 suggest that it is involved in pronephros development [62]. *WT1a* and *WT1b* are two types have been  
335 reported from *WT1* in zebrafish. Inactivation of *wt1a* leads to the absence of glomeruli while targeting  
336 of *wt1b* resulted in the formation of renal cysts [60]. The result showed an increase in *WT1a*



337 expression in head kidney of fish group treatment with hala extract, especially in the group fed with  
338 2% which recorded the highest value. This could be contributed to the activity of ROS in plant extract  
339 compounds to activate the tumour suppressor gene and suggest the role of this plant to work as anti-  
340 tumour agent in kidney.

341 In conclusion, our results demonstrated that dietary supplement with hala extract to rainbow trout for  
342 two weeks stimulates the non-specific immune response and increase its resistance toward *Y. ruckeri*  
343 infection. The current results suggest that the hala extract, specially the highest dosage (2%) could be  
344 considered as a good fish food supplements to enhance the immune system and resist and/or control  
345 pathogenic bacterial, in addition to its potential effect to resist the tumour formulation. Future  
346 investigations could focus on the effects of long feeding time on the immune status and general health  
347 status of fish.

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Table 1. Primers used for real-time PCR.

<b>Gene name</b>	<b>Primer sequences (5' - 3')</b>	<b>Products size</b>	<b>GenBank number</b>
Cluster of differentiation (CD4)	GCCCTGCAGAGGACAAATCT TACAAAGGCCACTGGAGCTG	171	NM_001124539.1
Lysozyme II (LYZ2)	TCCAGATCAACAGCCGCTAC GATTCCGTTCCGGTCCAACA	149	NM_001124716.1
Interleukin 8 receptor (IL-8)	CGGTGCCGTCATATTCCTGT GGGTCAGGGACTGTTGACTG	110	NM_001124279.1
Beta-actin ( $\beta$ -actin)	ATGGGCCAGAAAGACAGCTACGTG CTTCTCCATGTCGTCACAGTTGGT	186	AJ438158.1
Tumor necrosis factor TNF	CAAGAGTTTGAACCTCATTAG GCTGCTGCCGCACATAAAG	130	NM_001124374
Wilms' tumor suppressor 1a (WT-1a)	ATGTTTCAGCAACGCACCCTA GAACTGGGAGGAGTGGTGTG	129	NM_001124294.1

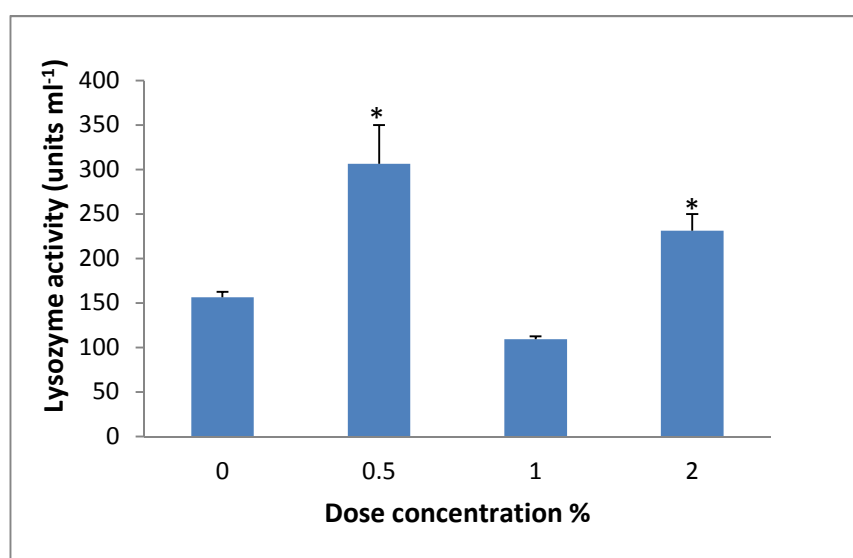


Figure 1. Serum lysozyme activity of rainbow trout fed for 2 weeks with dietary hala extract supplemented doses: 0% (control), 0.5%, 1% and 2%. Data are presented as mean  $\pm$  S.E. Asterisk represents significant difference from control  $p \leq 0.05$ . Bars =mean  $\pm$ S.E.

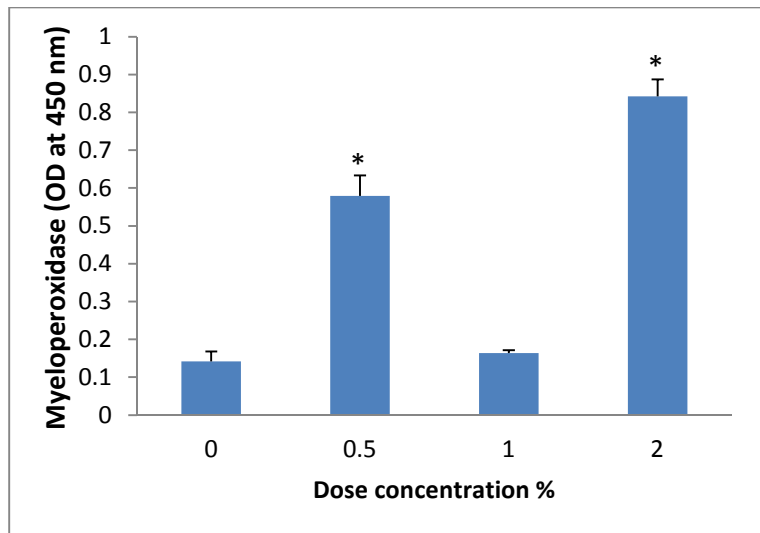


Figure 2. Serum Myeloperoxidase content of rainbow trout fed for 2 weeks with dietary hala extract supplemented doses: 0% (control), 0.5%, 1% and 2%. Data are presented as mean  $\pm$  S.E. Asterisk represents significant difference from control  $p \leq 0.05$ . Bars = mean  $\pm$  S.E.



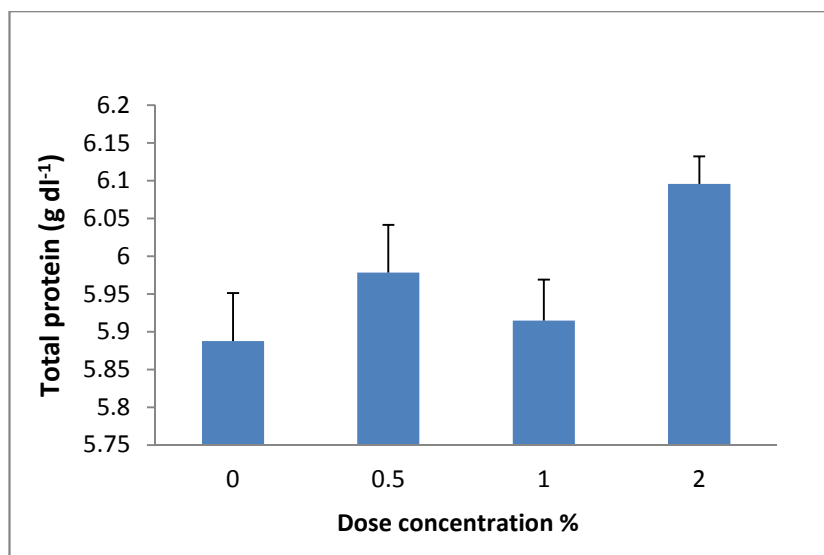


Figure 3. Serum total protein of rainbow trout fed for 2 weeks with dietary hala extract supplemented doses: 0% (control), 0.5%, 1% and 2%. Data are presented as mean  $\pm$  S.E. Asterisk represents significant difference from control  $p \leq 0.05$ . Bars =mean  $\pm$ S.E.

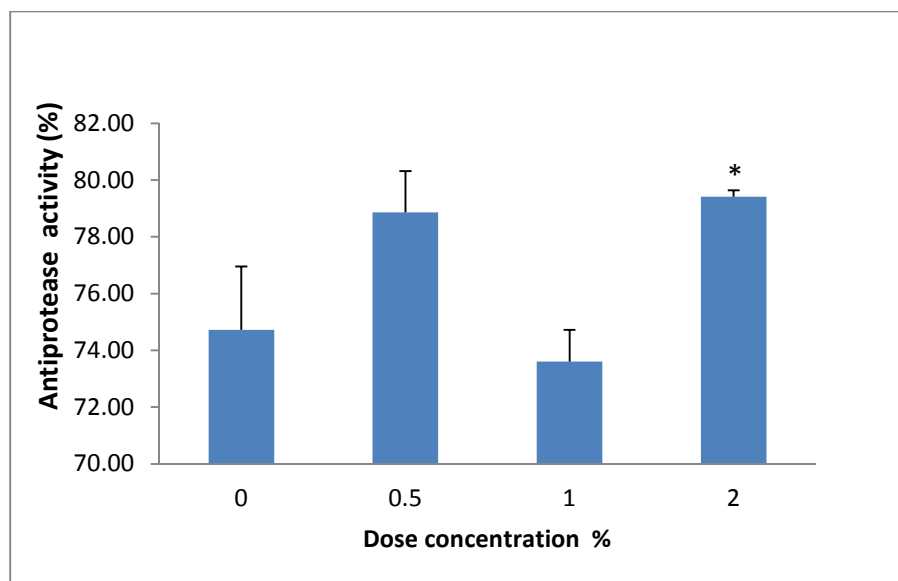


Figure 4. Serum Antiprotease of rainbow trout fed for 2 weeks with dietary hala extract supplemented doses: 0% (control), 0.5%, 1% and 2%. Data are presented as mean  $\pm$  S.E. Asterisk represents significant difference from control  $p \leq 0.05$ . Bars =mean  $\pm$ S.E.

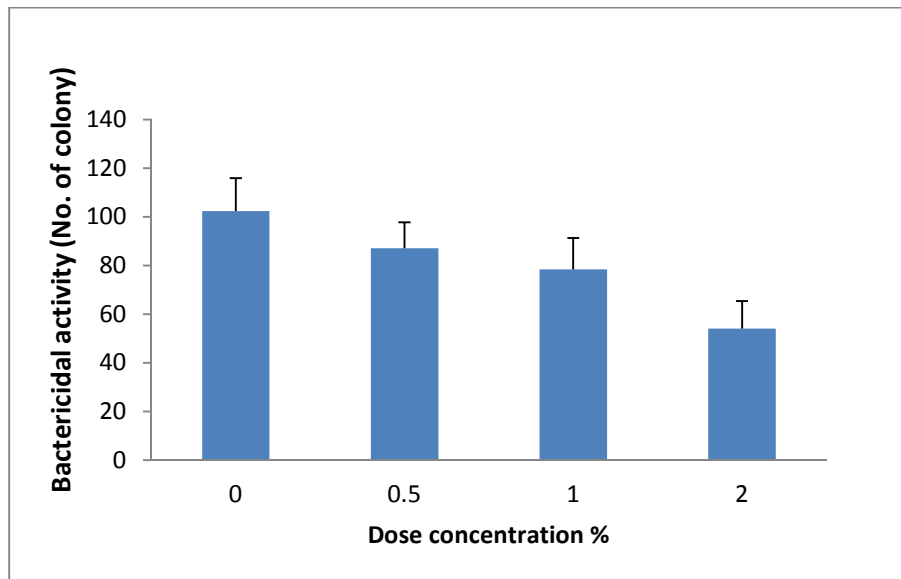


Figure 5. Serum bactericidal activity of rainbow trout fed for 2 weeks with dietary hala extract supplemented doses: 0% (control), 0.5%, 1% and 2%. Data are presented as mean  $\pm$  S.E. Asterisk represents significant difference from control  $p \leq 0.05$ . Bars =mean  $\pm$ S.E.

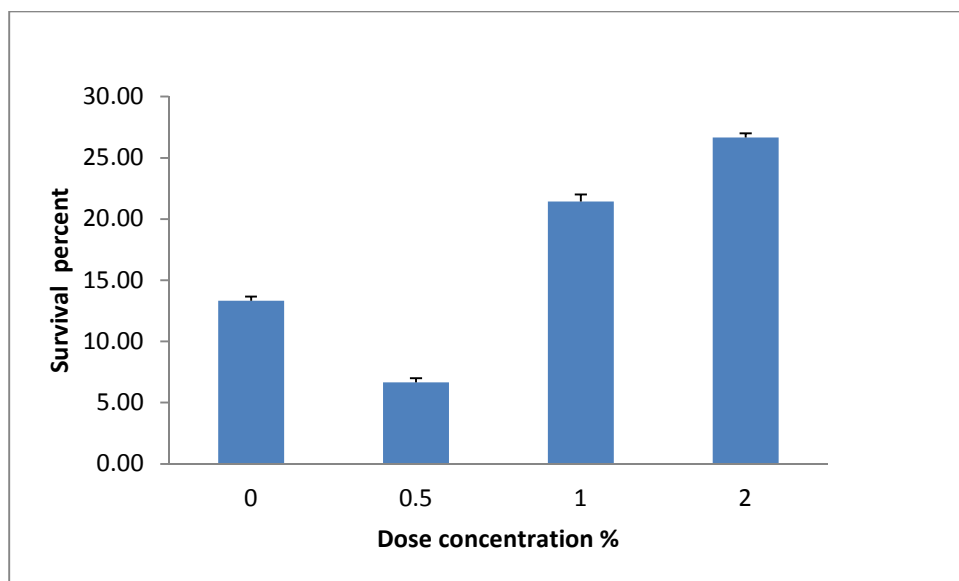


Figure 6. Survival of rainbow trout fed for 2 weeks with dietary hala extract supplemented doses: 0% (control), 0.5%, 1% and 2% followed by challenge with *Yersinia ruckeri*. Data are presented as mean  $\pm$  S.E. Asterisk represents significant difference from control  $p \leq 0.05$ . Bars =mean  $\pm$ S.E.

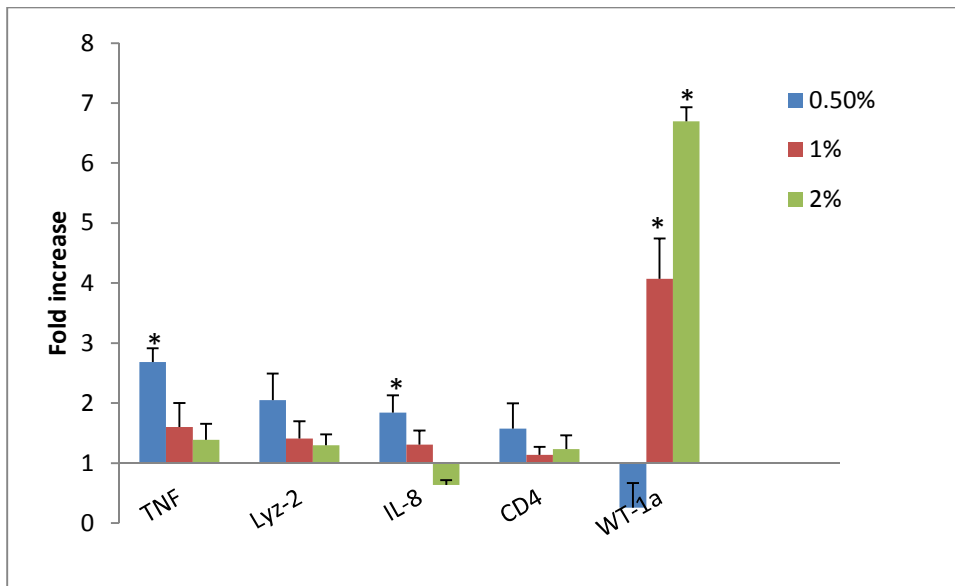


Figure 7. Expression of immune related genes (TNF, LYZ2, IL-8 and CD-4) and tumour suppressor genes (WT-1a ) in the head kidney of rainbow trout fed dietary hala extract supplemented doses of 0.5%, 1% and 2% for 2 weeks. Data are expressed as fold increase (mean  $\pm$  standard error, SE), obtained by dividing each sample value by the mean control value at the same sampling time. Values higher than 1 express an increase while values lower than 1 express a decrease in the indicated gene. Asterisks denote significant differences between control and treatment groups ( $P < 0.05$ ).

- Effects of hala extract (*Pandanus tectorius*) on the immune status of rainbow trout
- Hala extract provoked significant up-regulation in most of immune-related.
- Hala extract can use to resist tumour formation.
- Feeding with hala extract increased resistance toward *Yersinia ruckeri* infection

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