

Assessment of a high protein distillers dried grain (HP-DDG) augmented with phytase in diets for European sea bass, *Dicentrarchus labrax* fingerlings on growth performance, haematological status, immune response and related gut and liver histology

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25 **Abstract**

26 Inclusion of high levels of plant protein ingredients in fish feeds induces the presence of
27 undesirable compounds such as **Anti-Nutritional Factors**, including non-starch polysaccharides and
28 phytates. **The present study evaluated the effect of partial replacement of dietary soybean meal by a high
29 protein distillers dried grains (HP-DDG) a co-product of corn based ethanol production.** We evaluated
30 HP-DDG in experimental diets with a supplemented commercial phytase on growth performance,
31 physiological parameters and histological changes of the intestine and liver of European sea bass,
32 *Dicentrarchus labrax* fingerlings. **The experiment was conducted for six weeks. A total of 240 *D. labrax*
33 fingerlings was randomly divided to four experimental treatments (each in triplicates groups) and fed to
34 apparent the satiation six days a week for a six weeks' period. Four dietary treatments: containing 0, 30,
35 40 and 50% HP-DDG supplemented with enzyme phytase 0.5g kg⁻¹ diet respectively, were tested.** The
36 results showed that growth performance and feed utilization efficiency of sea bass was significantly
37 higher (P≤0.05) with increasing levels of HP-DDG-and phytase supplementation. Superior phosphorous
38 utilization was also observed with respect to whole body retention for each incremental level of HP-DDG
39 inclusion. Hematology and serum biochemistry (hemoglobin (Hb), red blood cells (RBCs), white blood
40 cells (WBCs) and **humoral immune** parameters including total protein, globulin, cholesterol, **lysozyme**
41 activity and total antioxidant capacity (TAC) were improved (P≤0.05) in fish fed diet with increased
42 levels of HP-DDG and supplemented phytase. The findings suggest that the use of HP-DDG
43 supplemented with phytase in sea bass diets enhanced growth, physiological and immunological
44 responses, and evidenced a cost benefit advantage for European sea bass production compared to the use
45 of a diet without either **HP-DDG** or phytase incorporation.

46
47 **Keywords:** **HP-DDG, phytase, growth, P utilization, liver & intestine histology, hematology**

48 **Introduction**

49 Sea bass, *Dicentrarchus labrax* is the most economically relevant marine fish species produced in Egypt,
50 representing 2% of the total marine production in the country (GAFRD, 2018). Unlike, European seabass
51 production, where fries are obtained from large scale commercial hatcheries, Egyptian production still rely on wild
52 collected fries. Thus, intensive aquaculture is of great importance, as their production represents about 60% of
53 animal protein used for human consumption, but as rapidly expanding sector, it places enormous pressure on the
54 aquaculture industry to find sustainable and cost-effective ingredients in fish diets (Tacon & Metian, 2015; Goda
55 et al., 2020) Plant ingredients are now mainly used to reduce costs and improve sustainability of ingredients used
56 in feeds for high valuable carnivorous species like sea bass and sea bream (*Sparus aurata*). However, plant
57 ingredients have some restrictions, including competitive demand with human consumption and international
58 market availability with escalating prices and questionable sustainability with respect to environmental
59 stewardship (Matos et al., 2017). Thus, not surprisingly, the manufacturing process wastes have gained
60 considerable interest as direct protein sources in fish feeds. High Protein Distillers Dried Grains (HP-
61 DDG) is a co-product from the fermentation of milled corn and distillation of bioethanol in the increasing
62 bio-refinery markets. HP-DDG is relatively low in lysine but nonetheless contains high crude protein
63 content (43 to 49%). This makes HP-DDG a good candidate for aquafeeds when formulated with
64 complementary protein concentrates. However, Anti-Nutritional Factors (ANFs) as NSP's (Non-Starch
65 Polysaccharides) in HP-DDG may diminish or even inhibit the digestibility of organic matter, energy
66 and protein due to their higher viscosity in the intestinal tract. The addition of exogenous enzymes could
67 mitigate the negative impacts of ANFs and represent an innovative strategy to improve nutrient
68 availability of plant-based diets (Castillo & Gatlin, 2015; Dalsgaard et al., 2016; Hassaan et al 2020).
69 Previous research succeeded to prove the usefulness of using phytase to enhance phosphorus availability
70 (Kumar et al., 2012). The use of exogenous enzymes is a common procedure to improve ANFs effectively
71 and enhance nutrient digestibility in swine and poultry (Bedford & Cowieson, 2012), and its use has also
72 been reported to increase the bioavailability of amino acids and nutrient utilization in aquaculture diets
73 (Castillo & Gatlin, 2015). The anti-nutritive effects of phytate are highly influential on dietary amino

74 acid and energy digestibility, raising the value of phytase to the end user beyond being just a contributor
75 to phosphorus (and calcium) nutrition. The information on HP-DDG efficiency in combination with
76 exogenous enzymes is still scarce since many potential enzymes are available including those based on
77 SSF (solid state fermentation products) as mentioned by Bowyer et al. (2020). An identical HP-DDG
78 product was significantly improved in diets for sea bass as reported recently by Goda et al. (2020)
79 augmented with a commercial protease. This success required validation of phytase under similar
80 conditions for sea bass. Therefore, the present study aimed to evaluate the effects of including a
81 commercial phytase supplemented HP-DDG as a relatively new ingredient source on the growth
82 performance, feed utilization efficiency, humoral immune parameters, liver and intestinal morphology
83 in European sea bass, *Dicentrarchus labrax* and to assess its economic relevance.

84 **Materials and methods**

85 **Feed preparation**

86 Four isonitrogenous and isocaloric experimental diets were formulated (Table 1). The control diet
87 (C₀%) had no high protein distillers dried grains (HP-DDG). Tested diets were formulated to contain 30%,
88 40%, and 50% of HP-DDG added at the expense of the soybean content. All the diets were equally
89 supplemented supplemented with 0.5 g/kg of Phytase (Aextra PHY[®]). The patented phytase, isolated from
90 *Buttiauxella*, Danisco Animal Nutrition, Dupont Industrial Biosciences, Marlborough, UK. HP-DDG
91 and phytase in the present study are considered as a single ingredient complex. HP-DDG is one of the
92 most competitive sources of protein due to its lower moisture content, higher levels of lysine (3%) and
93 longer shelf life. HP-DDG (a co-product of bioethanol distillation from corn fermentation with high
94 levels of residual gluten protein after yeast separation and recovery) was supplied from POET Nutrition
95 Dakota Gold; POET Nutrition, LLC 4506 N. Lewis Ave, Sioux Falls, SD 57104 (USA).

96 The chemical composition of HP-DDG used in the feed formulation was crude protein 47%, crude fat 4%,
97 crude fiber 4%, Ash 7% and moisture 7%.

98 **Feeding protocol**

99 During the 42 -day experimental period, all fish were fed with their respective diets at 5% of body
100 weight d^{-1} for 6 days week^{-1} . Every 14 days, fish were weighed and the daily ration was adjusted
101 accordingly. The daily ration was divided into three equal amounts and offered three times a day (09:00,
102 12:00 and 15:00 h). Experimental diets were individually prepared by mixing the dry ingredients with
103 200 ml of water per kg diet. Two grams of commercial phytase enzyme contain the enzymatic activity
104 of 2000,000 Units was dissolved into the 200 mL water at 37 °C (Yoo et al., 2005). Commercial phytase
105 enzyme product (Aextra® PHY) was purchased from Gloray Vet COMPANY, USA.

106 The solution was incubated for 24 hours at room temperature according to the method of von Danwitz
107 et al. (2016) prior to its addition to the experimental diets. The mixture was blended, turned into a paste
108 and pelleted by passing the blended mixture through a laboratory pellet machine with a 1mm diameter
109 matrix. The resulting wet pellets were dried at room temperature for two days and then stored in plastic
110 bags and kept refrigerated (-2°C) until use.

111 **Fish and experimental facilities**

112 European sea bass, *D.labrax* fingerlings with an average initial body weight of $7.5 \pm 0.5 \text{ g fish}^{-1}$
113 were obtained from a commercial fish farm “El-Shref farm”, Wady Marriott, Alexandria” Egypt and
114 acclimated to the experimental conditions for 15 days. During this period, fish were fed a standard commercial
115 diet (Biomar) (45% protein). Then, fish were randomly distributed into twelve glass aquaria measuring
116 ($70 \times 40 \times 30 \text{ cm}$ each) representing four treatments (each in triplicate) at a stocking density of 20 fish
117 per aquaria.

118 On a daily basis, 50% of the water volume of each tank was exchanged to maintain adequate water
119 quality. Environmental parameters throughout the experiment were; salinity (37 ppt), temperature ($18 \pm 1^{\circ}\text{C}$),
120 and pH (7.0 ± 0.50) under a photoperiod regime of 12:12 hr (light: dark). The experimental protocols were all
121 approved by the local Institutional Animal Care Committee (IACC) meeting ethical standards and
122 legislation and statutes for animal studies.

123 **Growth Indices**

124 The mean final body weight (FBW) in experimental treatment was determined by dividing the
125 total fish weight in each aquarium by the number of fish. Weight gain (WG), specific growth rate (SGR),
126 feed conversion ratio (FCR), protein efficiency ratio (PER), protein productive value (PPV), fat retention
127 (FR), energy retention (ER), Phosphorus retention, economical conversion rate (ECR) and survival (%)
128 were calculated using the following equations:

129 $WG = \text{final body weight (g)} - \text{initial body weight (g)}$.

130 $SGR = 100 \times [(\ln \text{ final body weight (g)} - \ln \text{ initial body weight (g)}) / \text{duration of feeding (day)}]$

131 $FCR = \text{feed intake (g)} / \text{weight gain (g)}$.

132 $PER = \text{weight gain (g)} / \text{protein intake (g)}$.

133 $PPV = (\text{protein gain (g)} / \text{protein intake (g)}) \times 100$.

134 $FR = (\text{fat gain (g)} / \text{fat intake (g)}) \times 100$.

135 $ER = (\text{energy gain (kJ)} / \text{energy intake (kJ)}) \times 100$.

136 $ECR = \text{cost of diet (\$ kg}^{-1}\text{)} \times \text{Feed Conversion Ratio (FCR)}$

137 $\text{Survival (\%)} = 100 \times (\text{initial number of the fish} / \text{final number of fish})$.

138 Phosphorous Retention (PR %) was calculated according to Morales et al. (2018):

139 $PR = 100 \times (BW_{\text{final}} \times P_{\text{final}} - BW_{\text{initial}} \times P_{\text{initial}}) / (\text{Feed intake} \times P\%_{\text{diet}})$

140 Where: $P\%_{\text{diet}}$ is the content of phosphorous in the diet; P_{initial} and P_{final} represent the initial and final
141 concentration of phosphorous in fish.

142 **Diet and feed analysis**

143 Diets were analysed as described by Davies et al., (2019) in accordance with standard proximate
144 composition AOAC (2000) methods for crude protein $N \times 6.25$, crude lipid, ash, crude fibre and
145 moisture/Dry Matter DM.

146 **Phosphorous analysis**

147 250 mg of sample were accurately weighed into glass tubes. After that, 5 mL of the digestion solution (a

148 mixture of nitric and perchloric acid at the ratio of 2:1, 3:1, or 4:1 v/v) were added. The tubes were then
149 heated at 200°C until the solution became translucent and a brownish smoke stopped being released,
150 which indicated the complete digestion of the organic matter. Digested samples were quantitatively
151 transferred to 50 mL volumetric flasks using ash-free quantitative filter paper (Whatman No. 41,
152 Whatman International Ltd, Springfield, Kent, England, UK). The volume of the solutions was made up
153 to 50 mL using deionized water. Aliquots of the solutions were transferred to polyethylene flasks and
154 kept cool (4°C).

155 Total phosphorous (P) in feeds and fish was determined according to the following principle.
156 When ammonium molybdate solution is added to a solution of phosphate containing concentrated H₂SO₄
157 it produces a yellow crystalline precipitation of ammonium phospho-molybdate. Phospho-molybdate
158 reacts with amino-naphthol-sulphonic acid and produces a molybdenum complex which forms an intense
159 blue- coloured solution. A standard curve was produced from KH₂PO₄ solution and the absorbance
160 intensity of the colour of the reaction mixture is measured by colorimeter at 570nm. The colour generated
161 from suitably diluted extracts from individual samples of digested fish diets and fish was measured
162 against the standard curve according to the method of Palma et al., (2015)

163 **Blood Samples and Haematological Analysis**

164 Blood samples were collected at the end of the experiment. From each of the dietary treatments,
165 five fish were used for hematological indices analysis and five for plasma content analysis. The fish were
166 anesthetized with Tricaine Methanesulfonate (MS-222) and the blood samples were taken by puncturing
167 the caudal vessels. Blood samples were collected into two tubes, one containing heparin as anticoagulant
168 agent for haematological assessment and the other was anticoagulant free for biochemical estimation.
169 The haematological parameters are expressed in international units (SI). The total red and white blood
170 cell counts (RBC; 10⁶ mm⁻³ and WBC; 10³ mm⁻³, respectively) were obtained by using a standard
171 Neubauer-hemocytometer chamber using Shaw's solution as diluting fluid (Stoskopf, 1993). Hemoglobin

172 (Hb; g dL⁻¹) was determined colorimetrically using commercial kits (Diamond, Egypt) according to the
173 cyan- methemoglobin procedure (Drabkin, 1945). Hematocrit (Hct) were determined by using
174 microhematocrit-heparinized capillary tubes and a microhematocrite centrifuge (10000 g for 5 min)
175 Levels of serum aspartate aminotransferase (AST), alanine aminotransferase (ALT) and *alkaline*
176 *phosphatase* (ALP) were estimated according to the method described by Reitman & Frankel (1957).

177 **Biochemical and immune parameters**

178 The total protein (g dL⁻¹) was determined in plasma samples of fish from the different
179 experimental groups by the Biuret method according to Doumas et al. (1981). Albumin (g dL⁻¹) was
180 determined by the bromocresol green method (Reinhold, 1953) and globulin (g dL⁻¹) was calculated as
181 the difference between total protein and albumin, and cholesterol was measured by a commercial kit
182 (Pasteur, Lab, France, Egypt) (Yousefi et al. 2011). Triglycerides were determined according to MGowan
183 et al., (1983). Lysozyme activity (U mg⁻¹ protein) in serum was determined according to the method of
184 Ellis (1990) based on the lysis of the lysozyme sensitive gram-positive bacterium *Micrococcus*
185 *lysodeikticus* (Sigma, St. Louis, MO). Lysozyme acts upon susceptible bacteria by combining with and
186 breaking down a mucopolysaccharide. This mucopolysaccharide, which is the lysozyme substrate, has
187 been shown to be situated in the bacterial cell wall (Salton, 1952) and can be characterised chemically.

188 Total antioxidant capacity (TAC) level was estimated spectrophotometrically at 552 nm
189 following the method with Tween 80 oxidation (Galaktionova et al. 1998). Briefly, 0.2 ml of tissue
190 homogenate was added to 2 ml of 1% Tween 80. Instead of the sample, the blank assay included 0.2 ml
191 of distilled water. The mixture was incubated for 48 hours at 37 °C. After cooling, 1 ml of 40 % TCA
192 was added. The mixture was centrifuged at 3,000 g for 10 min. After centrifugation, 2 ml of supernatant
193 and 2 ml of 0.25% TBA reagent were mixed in. The mixture was heated in a boiling water bath at 100
194 °C for 15 minutes. The TAC level was expressed in (%).

195 **Histological examination**

196 Four fish from each replicate of *D. labrax* were randomly collected and dissected for tissue removal.
197 The distal section of the intestine were removed, thoroughly washed with a physiological saline (0.9%
198 Nacl) solution and fixed in Bouin's fluid. The material was dehydrated, cleared and finally embedded in
199 paraffin wax. Serial sections were cut to the thickness of 5-6µm. The sections were stained with
200 haematoxylin counterstained with eosin and mounted in DPX (Yano, 1988). The sections were examined
201 with an Olympus light microscope and photographed with a digital camera as required. The histological
202 examination was carried out according to Culling (1983).

203 **Economic evaluation**

204 The economic evaluation was calculated according to the following equations (Salama et al., 2010).
205 Feed cost per kg fresh fish (LE) = Cost / kg diet (LE) * consumed feed to produce 1 kg fish .
206 Relative feed cost/kg fresh fish = Values of feed cost/kg fresh fish / the minimum value of the same
207 parameter
208 Feed cost / 1 kg gain (LE) = Feed intake per kg gain (FCR) * cost/kg diet (LE).
209 ECR = cost of diet (\$ kg⁻¹) x Feed conversion ratio (FCR)

210 **Statistical analysis**

211 One-way ANOVA and Duncan's multiple rank test (Duncan, 1955) were used to test were
212 calculated to test effects with a probability of $P \leq 0.05$ that were considered significant. The data from
213 the experiments were statistically analyzed using GLM (general linear model) procedure according to
214 Statistical Analysis System (SAS, Institute 2003, SAS User's Guide: Statistics. SA Institute, Cary, NC.
215). However, data is presented untransformed to facilitate comparisons.

216 **Results**

217 **Growth performance and feed utilization**

218 Growth parameters are presented in (Table 2). At the end of the experiment, fish fed diets with either 40%
219 HP-DDG or 50 % HP-DDG supplemented with phytase, grew significantly more ($P < 0.05$) with improved growth
220 parameters (higher final body weight, weight gain, specific growth rate and better feed conversion ratio) than those

221 fed either the control or 30% HP-DDG supplemented with phytase diets. As for the phosphorus retention, fish fed
222 50% HP-DDG supplemented with phytase recorded the superior values ($P \leq 0.05$), while the control group
223 recorded the lowest value (Table 3). Significantly higher whole body P level was noticed in sea bass with
224 the incremental inclusion of HP-DDG with phytase and this reflected dietary P levels. The best relative
225 feed cost/kg per fresh fish was recorded for fish fed diet containing 50% HP-DDG supplemented with
226 phytase, in opposition to fish fed the control diet (Table 8).

227 **Blood parameters**

228 Fish fed 50% HP-DDG recorded significantly lower values ($P \leq 0.05$) of ALT, AST, and ALP, while fish
229 fed the control diet recorded significantly higher values ($P \leq 0.05$) in all blood parameters (Table 3). Moreover,
230 fish fed 50% HP-DDG recorded significantly higher ($P \leq 0.05$) triglyceride values than fish fed the
231 remaining dietary treatments. Fish fed either 30% or 50% HP-DDG supplemented with phytase recorded
232 the highest ($P \leq 0.05$) values of RBCs, WBCs and PCV, while fish fed the control diet recorded the lowest
233 value ($P \leq 0.05$) of WBCs, RBCs, Hb and PCV (Table 4). In addition, compared to the control diet, fish
234 fed diets 50% HP-DDG supplemented with phytase recorded significantly higher values ($P \leq 0.05$) of
235 total protein, albumin and globulin (Table 5). In terms of immune parameters, it was observed that fish
236 fed diets with increasing inclusion rates of HP-DDG with supplemented phytase, evidenced a significant
237 ($P \leq 0.05$) and correspondent increase of cholesterol, lysozyme and TAC levels compared to fish fed the
238 control diet (Table 6).

239 **Histological studies**

240 The histological changes of intestinal and hepatic tissue were assessed by light microscopy.
241 Observations revealed that intestine and liver of fish fed the control diet showed normal, intact intestinal
242 wall, intestinal villi as well as goblet cells and distribution (D_{0%}, Figure 1a). Conversely, fish fed 30%
243 HP-DDG supplemented with phytase showed moderate improvement in length and width of intestinal
244 villi as well as goblet cells (D_{30%}, Figure 1b); fish fed 40 and 50% HP-DDG showed an improvement

245 in length and width of intestinal villi as well as an increase in the number of goblet cells (D40%, Figure
246 1c and D50%, Figure 1d). The histopathological analysis also revealed a significant increase in villus
247 length, villus width and area of absorption in fish fed diets containing HP-DDG with phytase (Figure 2).
248 Sea bass fed the control diet showed normal organization of the hepatic cell and blood capillaries (Figure
249 3a). Hepatopancreas of European sea bass, *D. labrax*, fingerlings fed (30% HP-DDG with phytase)
250 showed activation of melano-macrophage centers and normal pancreatic acini and normal hepatocytes
251 (Figure 3b), in addition with increasing dietary HP-DDG supplemented with phytase up to 50% necrosis
252 in pancreatic tissue (blue arrows) with activation of melano-macrophage centers (black arrows) (Figure
253 3c & d), was also observed.

254 Discussion

255 The addition of HP-DDG with a supplementation of phytase resulted in an improved growth and
256 feed efficiency sea bass under experimental conditions. The current results are in accordance with
257 previous studies on different species including *L. rohita* (Bano and Afzal, 2017), hybrid grouper
258 (Anthonius et al., 2018), and Nile tilapia (Abo Norag et al., 2018). Furthermore, Goda et al. (2019)
259 reported that fed HP-DDG enriched with phytase resulted in higher growth rate of *D. labrax*. Ranjan, et
260 al. (2017) found that 0.01 % phytase supplementation in basal diet significantly improved ($P \leq 0.05$) the
261 weight gain, SGR and FCR of *L. rohita*. However, the current results are inconsistent with the findings
262 of Hu et al. (2016) and Yigit et al. (2018) who reported no differences ($P \geq 0.05$) in WG, FCR and gut
263 health respectively, Nile tilapia and rainbow trout fed diets supplemented with dietary microbial phytase.
264 The discrepancies between these studies may be due to several factors such as the dosage and phytate
265 sources, types of feed ingredients, fish species and the pH of the stomach (Dersjant-Li et al., 2015; Yigit
266 et al., 2018). It should be noted that in the current investigation our source of phytase was a novel
267 product, Axtra[®] PHY that offers unprecedented phytate degradation and phosphorus digestibility when
268 compared with *E. coli* phytases that are more commonly employed in the industry. It has exceptionally

269 rapid activity in the stomach. The activity of Aextra® PHY, a *Buttiauxella* phytase, at pH 4.0 is almost
270 double that at pH 5.5, the level at which all commercial phytases have their activity standardized, and
271 much higher than other phytases. Aextra® PHY also improves sodium - and therefore also protein, glucose
272 and nutrient - absorption from the gut, with positive effects on growth performance (Danisco Technical
273 Report).

274 The present results could be attributed to several factors i) the inclusion of phytase eliminates the
275 negative impact of phytate to reduce the availability of minerals particularly calcium, magnesium, iron,
276 and zinc (Shah et al., 2016) and negatively affect the absorption of lipids and proteins (Jacob 2015) thus,
277 the addition of the phytase in the diets enhances mineral availability and utilization of dietary energy and
278 amino acids (Bowyer et al., 2020; Sharawy et al., 2020); ii) inclusion of phytase modulates the gut
279 microbiota by hydrolysis of the phytate and may thus positively influence the intestinal health
280 (Rachmawati et al., 2017); iii) the exogenous phytase improves nutrient digestibility and consequent
281 availability by destruction of insoluble cell wall complexes and subsequent release of low-molecular-
282 weight carbohydrates as sources of available energy for growth (Jacob, 2015); v) Phytase is also capable
283 of converting the inactive form trypsinogen into the active form trypsin which degrades protein and
284 oligopeptides into amino acids that consequently improves overall protein utilization (Hagbayan and
285 Mehdi 2015).

286 The superior values of hematology and immune parameters were observed in fish fed the diet
287 containing 50% HP-DDG, which is in agreement with the results obtained by Peatman and Beck (2016)
288 who found that channel catfish fed with phytase supplemented diets significantly elevate RBC's, WBCs,
289 PCV% and Hb levels. Also, the most elevated ALT, AST, ALP activities decrease with increasing HP-
290 DDG dietary inclusion. The present results are also consistent with Shelby et al. (2007) who found that
291 Nile tilapia fed DDGS diets showed enhanced immune system, liver function and disease resistance.
292 Furthermore, Ghaly et al. (2017) revealed that for broiler chickens fed dietary DDGS supplemented with

293 different levels of Avizyme enzymes, there was a significant ($P \leq 0.05$) decrease in the values of ALT and
294 AST activity in blood of birds. In the present study, enhancement of the immune response of sea bass
295 fed diets contain HP-DDG supplemented with phytase could be due to i) the presence of significant
296 amounts of biologically active compounds (mannans, β -glucans and nucleotides) derived from yeast,
297 which comprises about 10 percent of total DDGS mass (Shurson, 2018; Kim et al., 2008).

298 In terms of Phosphorus (P) retention, the present study showed that increased phosphorus retention
299 was detected in sea bass fed diet levels up to 50% HP-DDG with phytase compared to the control group
300 (Table 7). This is in agreement with the findings of von Danwitz et al., (2016) who found the lowest
301 phosphorus retention in fish fed a diet without phytase supplementation (46.5%) and increased by
302 addition of 1000 FTU or 2000 FTU phytase to 52.2% and 67.2%. Furthermore, Totok Yudhiyanto et al.
303 (2017) found that phosphorus retention increases with increasing phytase supplementation when added to
304 the diet of Asian Seabass, *Lates calcarifer*.

305 Light microscopy revealed a normal and healthy morphology of the intestinal tract of European sea
306 bass fed HP-DDG enriched with phytase. The fish intestine enterocytes displayed healthy brush borders
307 and no signs of damage. The present results are consistent with Adeoyea et al., 2016 who found that
308 inclusion of exogenous enzymes in tilapia diets improved ($P \leq 0.05$) the intestinal morphology, goblet
309 cells abundance and microvilli diameter surface than those fed the control diet. These results are also in
310 line with those of Adeoyea et al. (2016). In terms of liver histology, sea bass fed diets supplemented with
311 phytase showed a marked decrease in the hepatic and pancreatic lesions (Figure 3 a, b and c). The present
312 findings are consistent with the studies of Abo Norag et al., 2018, who recorded a noticeable decrease in
313 the hepatic and pancreatic lesions of fish fed diets supplemented with phytase in experiments with tilapia.

314 The current results show that the best relative feed cost per kg fish gain was observed in fish fed
315 50% HP-DDG supplemented with Phytase (Table 8). These are consistent with Nehad et al., (2019) who
316 found that the feed cost to produce one kg fish gain was reduced by inclusion of phytase in the diet by

317 41.35%. In addition, Khan et al., (2006) stated that enzyme supplementation is a more realistic and cost-
318 effective strategy to achieve maximum profitability.

319 In conclusion, using phytase in aquafeeds improves growth performance and minimizes the
320 environmental impact of fish production when incorporating higher levels of vegetable protein sources
321 in the diet for E. sea bass. The effective use of a phytase supplement in feed formulations can have a
322 significant positive impact in terms of improving performance, reducing costs and boosting production
323 efficiencies. In this example, our work demonstrates the potential of HP-DDG as a novel and sustainable
324 ingredient with unique characteristics and versatility in compound feed formulations in a marine species
325 such as sea bass. The combination of HP-DDG with phytase is synergistic and provides added value to
326 the plant ingredient. More research is warranted to explore the optimum rate of enzyme feed application
327 with considerations to thermal stability in extruded diets, post pelleting spraying technology and the
328 different temperature and feeding strategies of various fish species. It will be imperative to examine the
329 range of options best suited to the different production stages where diet formulations can be adjusted to
330 incorporate novel feed ingredients like HP-DDGS especially in grower diets for larger fish approaching
331 harvest weights where economic considerations prevail.

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Table (1): The composition (g/kg) and chemical analysis (% on dry matter basis) of the experimental diets (HP-DDG + phytase).

Ingredient	Control	30%+ Phytase	40%+ Phytase	50%+ Phytase
Fish meal 68 %	300	300	300	300
Soy bean meal 47%	375	262.5	225	187.5
Corn gluten 60%	90	89.5	89.5	89.5
Rice bran 12%	65	50	50	50
Wheat middlings 13%	69.5	83.8	84.8	85.8
HP-DDG ⁺	0	112.5	150	187.5
Soy bean oil	41	41	40	40
Fish oil	48	49	49	48
Dicalcium phosphate	8	8	8	8
Vitamin/Mineral Premix	2	2	2	2
Vitamin C	0.2	0.2	0.2	0.2
Antytocsec	1	1	1	1
Phytase*	0.5	0.5	0.5	0.5
Total	1000	1000	1000	1000
Dry matter (DM)	93.75	93.77	93.8	93.6
Crude protein (CP)	44.88	44.4	44.97	45.2
Ether extract (EE)	13.1	15.2	14.6	15.0
Nitrogen free extract	25.7	25.2	26.5	27.6
Crude fiber (CF)	4.32	4.3	3.4	3.1
Ash	12	9.8	10	11
Gross energy (GE: MJ/Kg DM)	20.93	21.56	21.52	21.87

Gross energy (GE) = (CP×5.6) + (EE ×9.44) + (NFE× 4.1) Kcal/ 100g (NRC, 1993)

*Axta[®] PHY (Danisco Animal Nutrition, Dupont Industrial Biosciences, Marlborough, UK + POET Nutrition Dakota Gold; POET Nutrition, LLC 4506 N. Lewis Ave, Sioux Falls, SD (USA)

Table (2): Growth performance and feed efficiency of European sea bass, *Dicentrarchus labrax* fed various levels of high protein distillers dried grains (HP-DDG) supplemented with phytase

Diets	IBW (g fish⁻¹)	FBW (g fish⁻¹)	WG (g fish⁻¹)	FCR (Feed: gain)	Feed intake (g fish⁻¹)	SGR (%/d)
Control	7.50±0.04	15.51±0.21 ^b	8.1±0.12 ^c	1.85±0.05 ^a	15.05±0.77 ^a	1.11±0.05 ^b
Diet (30%) + Phytase	7.50±0.06	17.60±0.21 ^b	9.80±0.15 ^b	1.40±0.05 ^b	13.95±0.40 ^b	1.37±0.01 ^b
Diet (40%) + Phytase	7.60±0.01	17.80±0.15 ^{ab}	9.95±0.15 ^{ab}	1.43±0.03 ^b	13.25±0.17 ^b	1.45±0.02 ^{ab}
Diet (50%) + Phytase	7.57±0.03	18.92±0.21 ^a	10.73±0.20 ^a	1.20±0.03 ^b	13.10±0.06 ^b	1.58±0.02 ^a

Values are mean ±SD of triplicate analyses. Means in the same column bearing different superscript differ significantly ($P \leq 0.05$). HP-DDG, high protein distillers dried grains; IW, initial weight; FBW, Final body weight; BWG, body weight gain ; FCR, feed conversion ratio and SGR specific growth rate.

Table (3) Effect of various dietary levels of high protein distillers dried grains (HP-DDG) supplemented with phytase on triglyceride levels, and AST, ALT and ALP activity of European sea bass, *Dicentrarchus labrax* fingerlings.

Diets	Triglyceride (mg dL ⁻¹)	AST (IU/L)	ALT (IU/L)	ALP (IU/L)
Control	95.88±1.22 ^c	39.77±0.36 ^a	36.87±0.60 ^a	18.48±0.05 ^{ab}
Diet (30%) + Phytase	140.35±2.20 ^b	36.64±1.50 ^{ab}	28.89±1.62 ^b	18.37±0.01 ^{ab}
Diet (40%) + phytase	171.70±2.23 ^b	31.49±1.26 ^b	23.73±0.40 ^b	18.39±0.12 ^{ab}
Diet (50%) + phytase	188.38±3.10 ^a	29.33±1.91 ^c	21.96±1.41 ^c	17.78±0.45 ^b

Values are mean ± SD of triplicate analyses. Means in the same column bearing different superscript differ significantly ($P \leq 0.05$). ALP: Alkaline phosphatase; ALT: Alanine aminotransferase; AST: Aspartate transaminase.

Table (4) Effect of various dietary levels of high protein distillers dried grains (HP-DDG) supplemented with phytase on hematological parameters of European sea bass, *Dicentrarchus labrax* fingerlings.

Diets	WBCs (10^3mm^{-3})	RBCs (10^6mm^{-3})	Hb (g dL^{-1})	PCV (%)
Control	21.030±0.140 ^d	1.360±0.020 ^d	7.675±0.055 ^d	18.590±0.070 ^c
Diet (30%) + phytase	22.700±0.520 ^c	1.500±0.030 ^c	8.015±0.105 ^b	19.905±0.125 ^{b,c}
Diet (40%) + phytase	25.225±0.055 ^{ab}	1.815±0.035 ^{ab}	8.800±0.340 ^{ab}	21.215±1.055 ^{ab}
Diet (50%) + phytase	26.400±0.040 ^a	1.915±0.025 ^a	9.275±0.015 ^a	22.400±0.010 ^a

Values are mean ±SD of triplicate analyses. Means in the same column bearing different superscript differ significantly ($P \leq 0.05$). HP-DDG, high protein distillers dried grains; WBCs, white blood cells; RBCs, red blood cells; Hb, Hemoglobin; PCV%, Packed cell volume.

Table (5) Effect of various dietary levels of high protein distillers dried grains (HP-DDG) supplemented with phytase on biochemical parameters of European sea bass, *Dicentrarchus labrax* fingerlings.

Diets	Total protein (g dL⁻¹)	Albumin (g dL⁻¹)	Globulin (g dL⁻¹)
Control	3.360±0.020 ^d	2.160±0.010 ^b	1.200±0.030 ^d
Diet (30%) + phytase	3.405±0.015 ^{cd}	2.125±0.005 ^b	1.280±0.020 ^d
Diet (40%) + phytase	3.535±0.055 ^b	1.895±0.025 ^c	1.640±0.030 ^b
Diet (50%) + phytase	3.785±0.055 ^a	2.285±0.045 ^a	1.940±0.060 ^a

Values are mean ±SD of triplicate analyses. Means in the same column bearing different superscript differ significantly ($P \leq 0.05$). HP-DDG, high protein distillers dried grains.

Table (6) Effect of various dietary levels of high protein distillers dried grains (HP-DDG) supplanted with phytase on immune parameters of European sea bass, *Dicentrarchus labrax* fingerlings.

Diets	Cholesterol (mgdI)	Lysozyme (U mg⁻¹ protein)	TAC (%)
Control	140.115±2.575 ^d	1.785±0.085 ^d	7.1195±0.035 ^c
Diet (30%) + phytase	159.885±3.655 ^c	2.315±0.045 ^c	8.100±0.130 ^b
Diet (40%) + phytase	175.230±2.020 ^b	2.810±0.060 ^b	9.555±0.185 ^a
Diet (50%) + phytase	191.695±2.555 ^a	3.190±0.070 ^a	9.855±0.265 ^a

Values are mean ±SD of triplicate analyses. Means in the same column bearing different superscript differ significantly ($P \leq 0.05$). HP-DDG, high protein distillers dried grains.

Table (7) Effect of various dietary levels of high protein distillers dried grains (HP-DDG) supplemented with phytase on dietary phosphorus utilization of European sea bass, *Dicentrarchus labrax* fingerlings (P values on a DM basis)

Diets	P in Diet (g /Kg)	P (Initial fish) (g/Kg)	P (final fish) (g/Kg)	P (feed intake) (g/Kg)	P (fish gain) (g/kg)	P retention (%)*
Control	6.42±50 ^c	3.78±0.11	15.31±1.31 ^c	46.24±1.13 ^c	11.55±1.27 ^c	24.98±0.44 ^c
Diet (30%) + Phytase	7.38±12 ^{bc}	3.78±0.11	22.79±0.02 ^b	55.49±0.60 ^b	19.00±0.57 ^b	34.24±0.71 ^b
Diet (40%) + phytase	9.94±60 ^b	3.80±0.11	26.21±0.04 ^{ab}	53.85±0.56 ^b	22.41±0.56 ^b	41.61±1.24 ^b
Diet (50%) + phytase	11.25±25 ^a	3.78±0.11	35.85±0.04 ^a	62.95±0.34 ^a	32.06±0.31 ^a	50.93±1.12 ^a

Values are mean ±SD of triplicate analyses. Means in the same column bearing different superscript differ significantly ($P \leq 0.05$). HP-DDG, high protein distillers dried grains.

*Phosphorus retention (%) = $100 \times (BW_{\text{final}} \times P_{\text{final}} - BW_{\text{initial}} \times P_{\text{initial}}) / (\text{Feed intake} \times P\%_{\text{diet}})$. Where: $P\%_{\text{diet}}$ is the content of phosphorous in the diet; P_{initial} and P_{final} represent the initial and final concentration of phosphorous in fish.

Table (8): Cost of feed required for producing one Kg gain when seabass were fed various levels of high protein distillers dried grains (HP-DDG) supplemented with phytase on European sea bass (*Dicentrarchus labrax*).

Diets	Feed cost per Kg (\$^a)	FCR (Feed : gain)	ECR (\$^a)	Cost / Kg fresh fish (\$^a)	Relative feed cost/Kg
Control	0.86	1.58	1.36	0.66	1.31
Diet (30%) + phytase	0.93	1.40	1.33	0.62	119
Diet (40%) + phytase	0.95	1.43	1.32	0.62	119
Diet (50%) + phytase	1.10	1.20	1.32	0.59	114

^a1\$ = 16.15 L.E. (Egyptian pound).

FCR: feed conversion ratio; ECR: economic conversion rate. (0, 30, 40 and 50%) levels of HP-DDG.

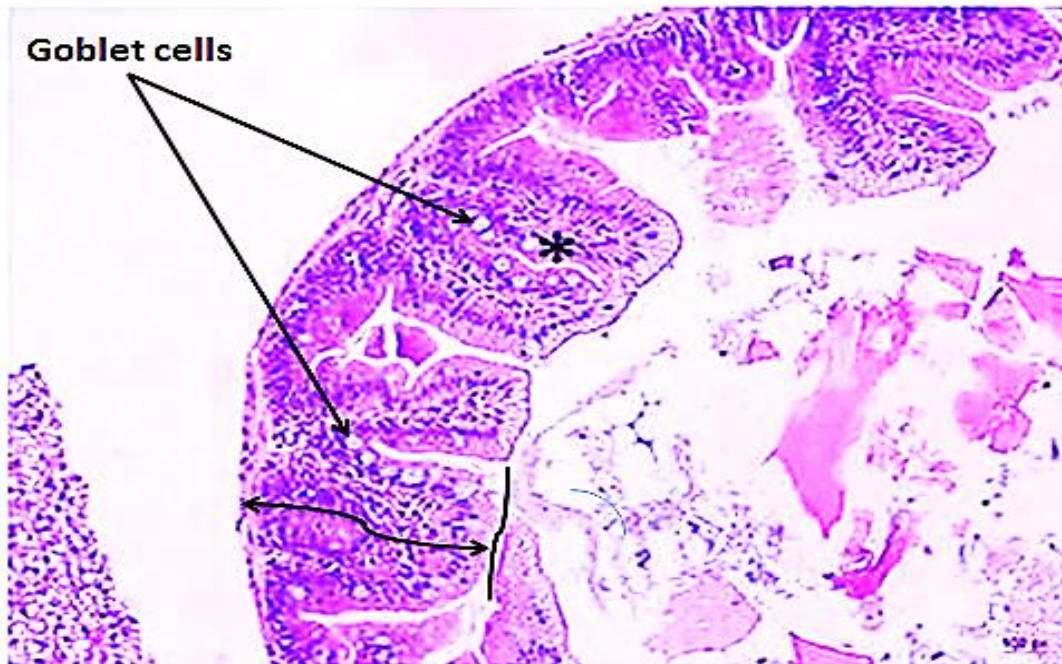


Figure (1a) Transverse section of Intestine of European sea bass, *D.labrax*, fingerlings fed the control diet (feed on basal diet + Phytase) for six weeks showing normal, intact intestinal wall intestinal villi as well as goblet cells.

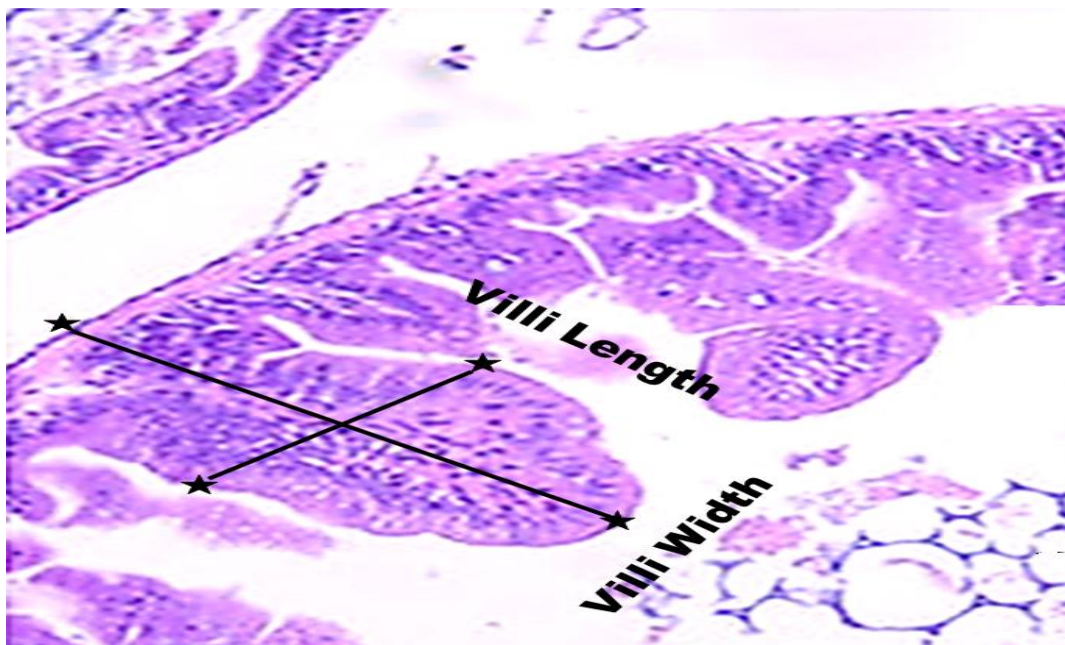


Figure (1b) TS of Intestine of European sea bass, *D.labrax*, fingerlings fed the (30% HP-DDG + phytase) for six weeks showing moderate improvement in length and width of intestinal (villi) as well as goblet cell density.

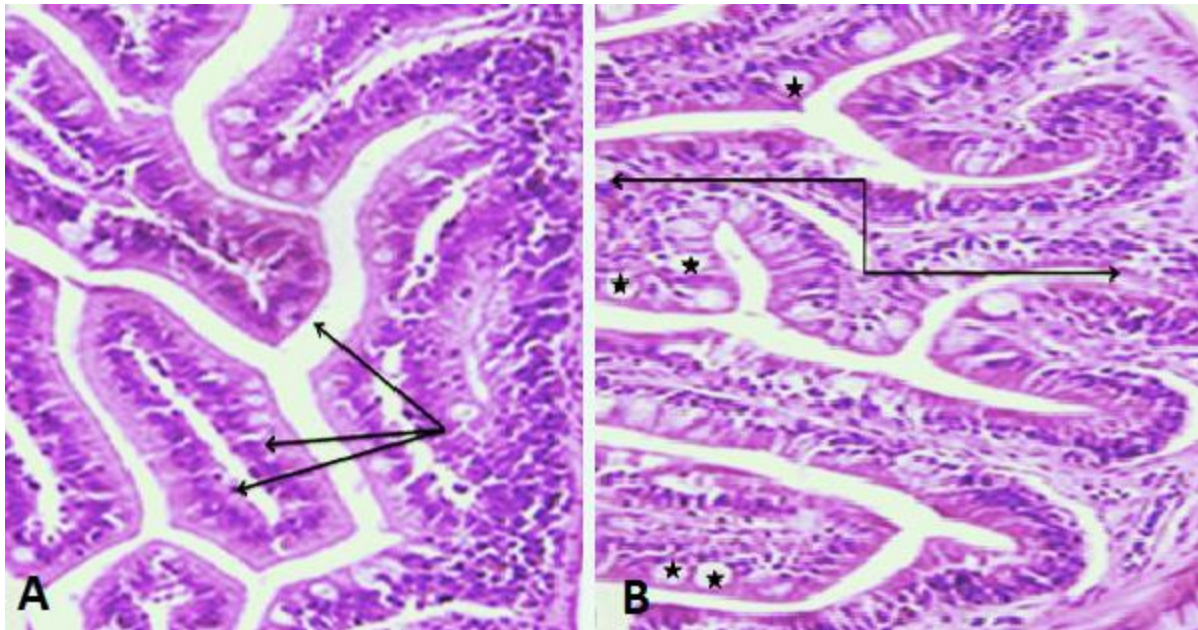


Figure (1c) TS of Intestine of European sea bass, *D.labrax*, fingerlings fed (40% HP-DDG + phytase) for six weeks showing highly increased length and width of the intestinal villi (arrows) and increase number of goblet cells. Stars show intracellular vacuolization indicative of lipid accumulation

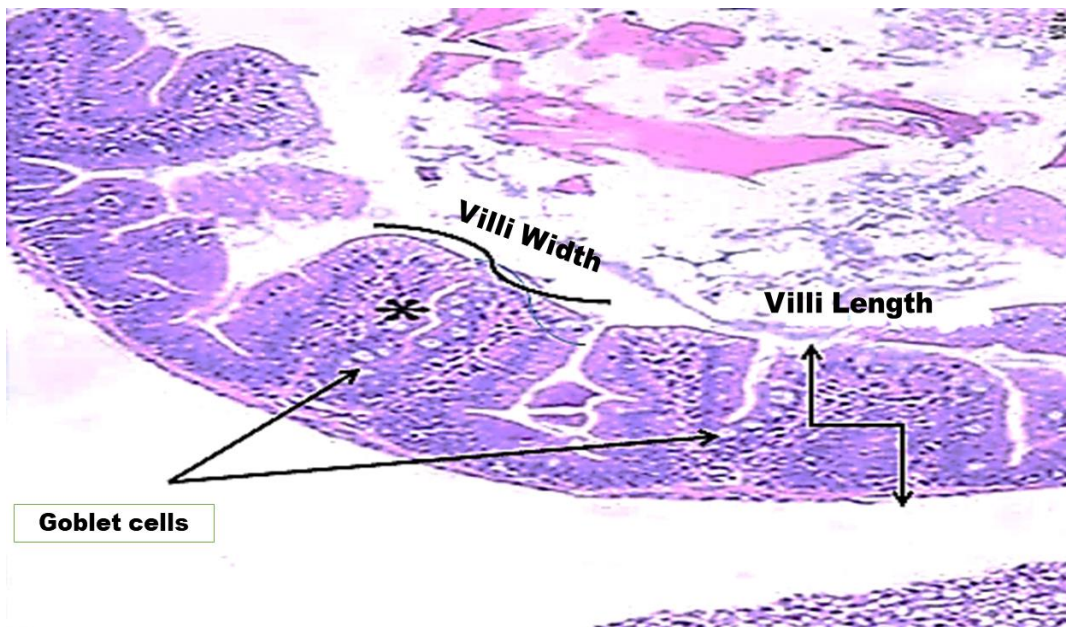


Figure (1d) TS Intestine of European sea bass, *D.labrax*, fingerlings fed (50% HP-DDG + phytase) for six weeks showing improvement in length and width of intestinal villus as well as increase number of goblet cells.

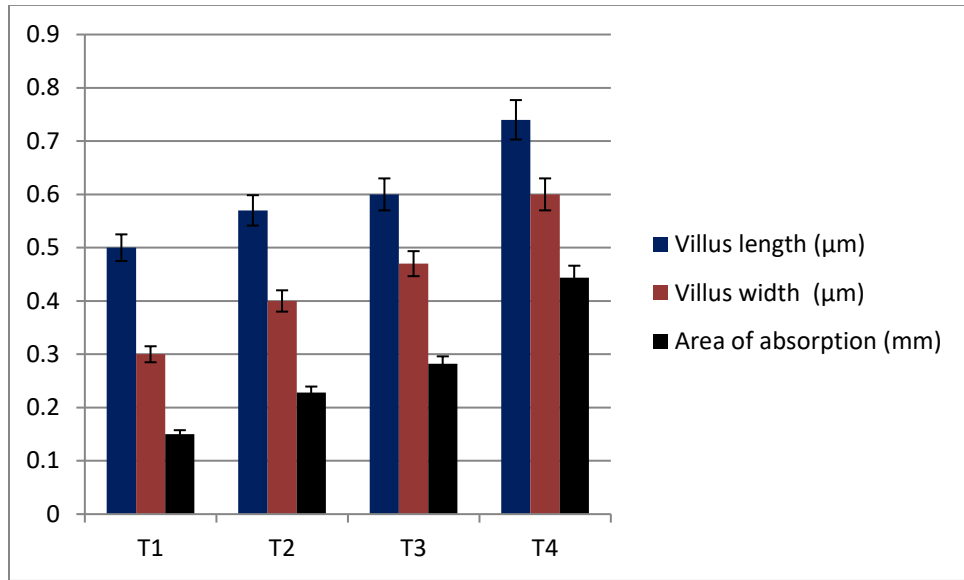


Figure (2) : Villi length and width and area of absorption in the (proximal? distal?) intestine of European sea bass fingerlings fed control (C), 30% HP-DDG (30), 40% HP-DDG (40), 50% HP-DDG (50) at the end of the 6-weeks feeding period.

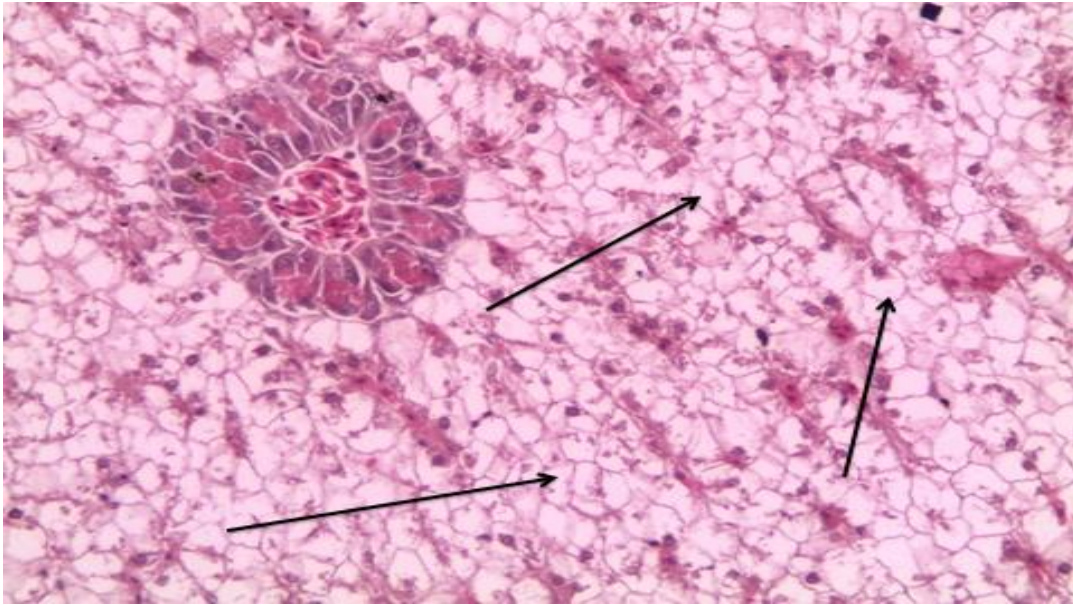


Figure (3a) Hepatopancreas of European sea bass, *D. labrax*, fingerlings (fed on the basal diet with phytase) for six weeks showing normal organization of the hepatic cell and blood capillaries (Arrows). (H&EX 400).

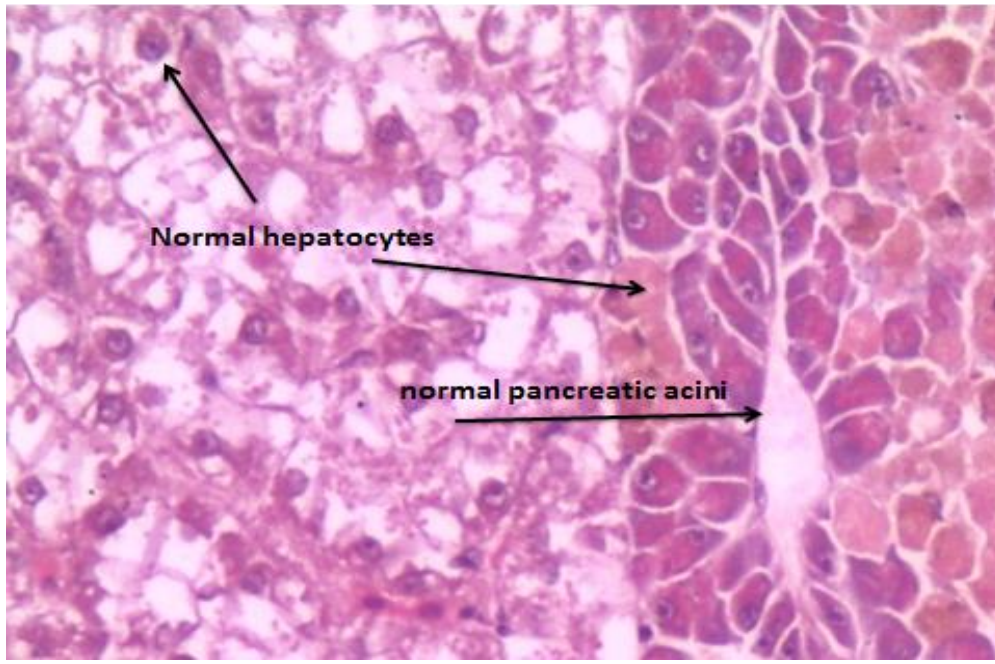


Figure (3b) Hepatopancreas of European sea bass, *D. labrax*, fingerlings fed (fed on 30% HP-DDG + Phytase) for six weeks showing activation of melano-macrophage centers and normal pancreatic acini and normal hepatocytes. (H&E X 400).

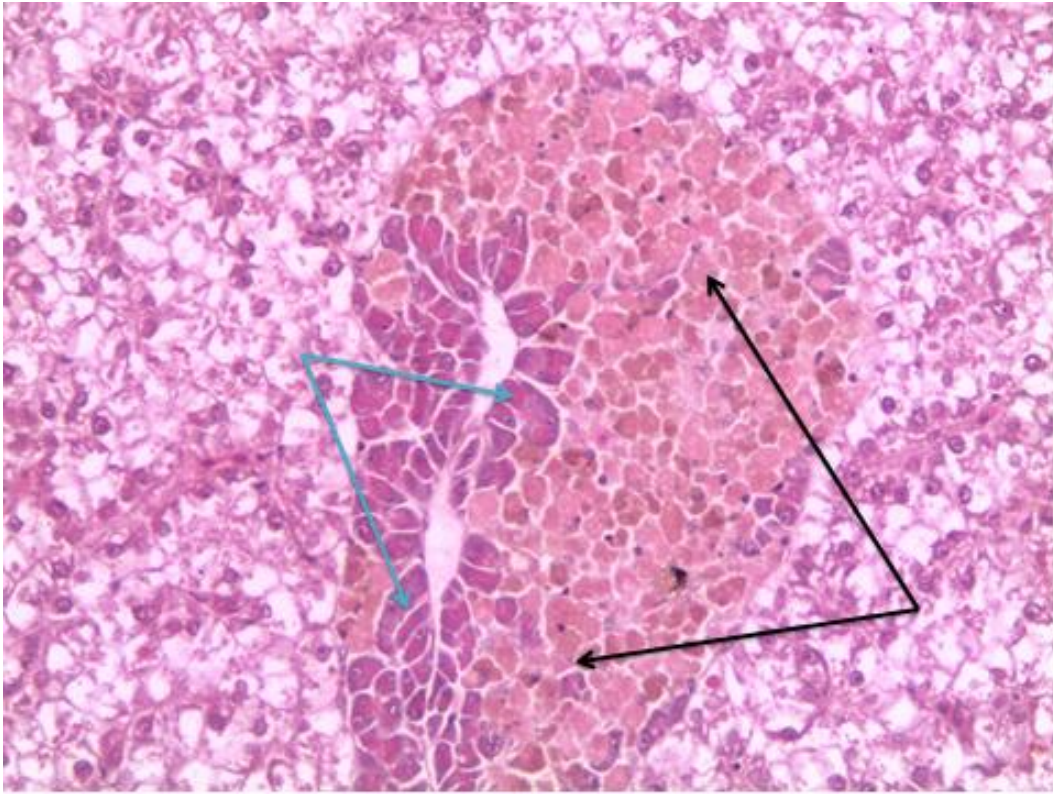


Figure (3c) Hepatopancreas of European sea bass, *D.labrax*, fingerlings (fed on 40% HP-DDG + Phytase) showing necrosis in pancreatic tissue (blue arrows) with activation of melanomacrophage centers (black arrows) (H&E X 20).

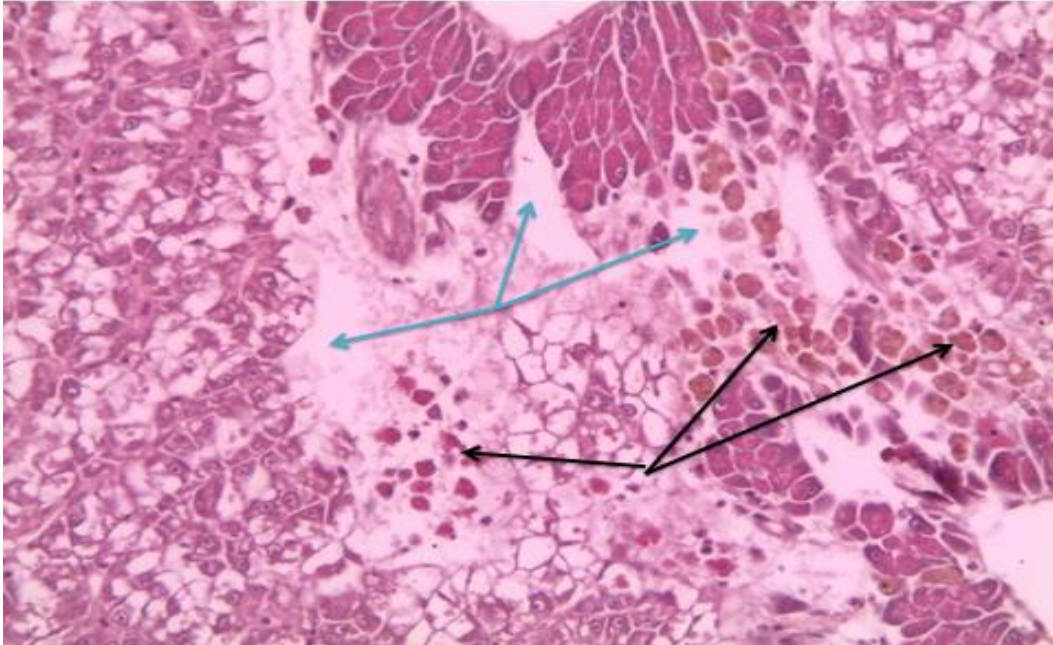


Figure (3d) Hepatopancreas of European sea bass, *D.labrax*, fingerlings (fed on 50% HP-DDG + Phytase) showing wide area of necrosis in hepatopancreas (blue arrows) with activation of melanomacrophage centers (black arrows) (H&E X 20).