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Potential application and beneficial effects of a marine microalgal biomass produced in a high-rate algal pond (HRAP) in diets of European sea bass, *Dicentrarchus labrax*

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Abstract

Microalgae have been used as live food in aquatic species. In recent years, the interest in microalgae has considerably increased, thanks to the evolution of production techniques that have identified them as an ecologically attractive aquafeed ingredient. The present study provides the first data about the effects of dietary inclusion of a microalgae consortium grown in a high-rate algal pond system on zootechnical performance, morphometric indices, and dietary nutrient digestibility as well as morphology and functionality of the digestive system of European sea bass, *Dicentrarchus labrax*. A dietary treatment including a commercial mono-cultured microalgae (*Nannochloropsis* sp.) biomass was used for comparison. Six hundred and thirty-six European sea bass juveniles (18 ± 0.28 g) were randomly allotted into 12 experimental groups and fed 4 different diets for 10 weeks: a control diet based on fish meal, fish oil, and plant protein sources; a diet including 10% of *Nannochloropsis* spp. biomass (100 g/kg diet); and two diets including two levels (10% and 20%) of the microalgal consortium (100 and 200 g/kg diet). Even at the highest dietary inclusion level, the microalgal consortium (200 g/kg diet) did not affect feed palatability and fish growth performance. A significant decrease in the apparent digestibility of dry matter, protein, and energy was observed in diets including 10 and 20% of the microalgal consortium, but all fish exhibited a well-preserved intestinal histomorphology. Moreover, dietary inclusion with the microalgal consortium significantly increased the enzymatic activity of maltase, sucrase-isomaltase, and β-glutamil transpeptidase in the distal intestine of the treated European sea bass. Algal consortium grown using fish farm effluents represents an attempt to enhance the utilization of natural biomasses in aquafeeds when used at 10% as substitute of vegetable ingredients in diet for European sea bass.

Keywords Microalgae consortium · *Nannochloropsis* spp. · *Oocystis* sp. · Gut physiology · Diet digestibility · Water treatment · European sea bass

Introduction

Aquaculture plays a key role in supporting human nutrition (Olsen 2011), and the increased availability of raw materials for feed formulation is required to support its rapid and continuous growth. For this reason, research has been focused, for a long time, on finding alternative ingredients to the traditional ones used by the feed industry to reduce pressure on natural
resources while addressing the growing market demand for aquaculture products. Management of sustainable feeding practices for aquatic organisms involves, both from a technical and an economic point of view, identifying alternative resources that consider the nutritional profile and the effects on animal welfare (resistance to stress and disease), while preserving the nutritional quality of the seafood product. It also implies the management of livestock activities waste into the environment (FAO 2018).

The use of microalgae as a potential ingredient of aquafeeds could represent an ecologically attractive alternative not only to traditional ingredients of marine and plant origin (Becker 2007) but also to innovative ingredients such as insects, seaweeds, and yeasts. In addition to their basic nutritional value (Spolaore et al. 2006), the inclusion of microalgae in aquafeeds is becoming popular as feed supplements in the aquaculture sector (Chu 2012; Priyadarshani and Rath 2012), thanks to the functional properties of their pigments and bioactive compounds. In addition, they may have a high content of proteins (30–70%), lipid (10–20%), and essential fatty acids (Becker 2007; Nasir et al. 2015; Shah et al. 2018; Cardinaletti et al. 2018). The main microalgal cultured genera are *Chlorella*, *Nannochloropsis*, *Scenedesmus*, *Arthrospira*, *Tisochrysis*, and *Tetraselmis* (Sirakov et al. 2015; Bleakley and Hayes 2017), thanks to their nutritional and health properties and consolidated cultivation technology. Recently, Castro et al. (2020) have proved that the inclusion up to 15% of *Nannochloropsis* sp. in diets for European sea bass has decreasing effects on the liver and intestinal antioxidant activity, while Abdelghany et al. (2020) have demonstrated that dietary *N. oculata* significantly improves growth parameters and resistance to pathogens such as *Aeromonas veronii* in Nile tilapia (*O. niloticus*).

However, the use of microalgae as an ingredient of aquafeeds may imply a few drawbacks such their high costs. In the recent decades, many biotechnological processes have emerged that are based on microalgal cultures and significant both from the environmental and industrial point of view. Such processes include biogas enrichment and purification; wastewater treatment (Quijano et al. 2017); CO2, NOx, and SOx removal from flue gas (Yen et al. 2015); and recovery of added-value products such as pigments, nutraceuticals, fertilizers, and biofuels (Bahr et al. 2013). The traditional processes for wastewater treatment are very expensive due to the chemical additives required during each phase. However, the cost could be minimized by using the microalgae biomass obtained by this technology as a feed for aquaculture. Some researchers have thus studied the potential value of multiple applications of microalgae to contribute to a circular economy approach (Valente et al. 2019) through their use in wastewater treatment (Velichkova et al. 2014; Nasir et al. 2015) and the sustainable production of biofuels (Rawat et al. 2011; Oliveira et al. 2020).

Numerous studies have been conducted on the characteristics of the microalgae obtained by a phycoremediation process (Yaakob et al. 2014; Nasir et al. 2015; Badre et al. 2019; Apandi et al. 2019; Michelon et al. 2021). Phycoremediation is a biotechnological process to remove contaminants from wastewater and is considered a simpler method than the conventional one (Raskin et al. 1997; Atiku et al. 2016). Microalgae have already been used to remove inorganic molecules and improve water quality (Ruiz-Martinez et al. 2012). Moreover, wastewater from the fish farm and the fresh market has also been used as a medium for a non-axenic microalgae culture (Apandi et al. 2019; Andreotti et al. 2017; Michelon et al. 2014).

In general, wastewater contains a high level of nutrients (nitrogen, phosphorus, and carbon) and organic matter, which act as elements to support microalgae biomass (Riaño et al. 2016); nitrogen availability has been shown to improve biomass production (Maizatul et al. 2017), thus modulating their nutritional value. Michels et al. (2014) used the wastewater obtained from a fish farm as a culture medium for the non-axenic production of *Tetraselmis suecica* biomass that in turn was used in juvenile shellfish culture resulting in increased productivity and constant quality in the hatchery phase. Several processes at pilot or industrial scale are actually based on non-axenic microalgae cultures from wastewater treatment, biogas purification/upgrading, or flue gas treatment. Moreover, recent studies have proved that microalgae biomass and composition, such as lipid composition, can be adjusted under physiological stress conditions, namely nitrogen depletion with increased salinity and/or increased salinity with temperature shock (Markou et al. 2016; Anitha et al. 2018). In non-axenic production processes, both microalgal and bacterial communities play key roles (Coronado-Apodaca et al. 2019), and the combination of different microalgal species can provide a balanced diet and improve animal growth and welfare (Spolaore et al. 2006; Cardinaletti et al. 2018). Cultivating microalgae in a high-rate algal pond (HRAP) system is a simple and economic way to produce valuable biomass to be included in aquafeeds. It provides fish farm wastewater treatment allowing the re-use of water for aquaculture while providing free nutrients for microalgae biomass production (Craggs et al. 2014). In addition, the technological treatment of microalgae biomass could also represent an important source of proteins, n-3 rich lipids, antioxidants, and natural bioproducts.

In this context, this study aimed to test the effects of dietary inclusion of a microalgae consortium grown in a HRAP system on zootechnical performance, morphometric indices, and dietary nutrient digestibility, as well as on the morphology and functionality of the digestive system of European sea bass. A dietary treatment including the commercial mono-cultured microalgae *Nannochloropsis* sp. was used for comparison,
based on its nutritional properties and, in particular, n-3 PUFA content.

**Materials and methods**

**Microalgae consortium production and characterization**

The trial was conducted at the Ifremer experimental station in Palavas les Flots, France. The microalgae consortium was cultivated in a conventional oval-shaped raceway HRAP. Water mixing in the HRAP (140 m² and 60 m³) was maintained at 0.2 m³/s using a vacuum airlift column developed and patented by COLDEP® (Barrut et al. 2012; Barrut et al. 2013). The column was connected to the HRAP and consisted of a central tube, the top of which was hermetically closed and connected to a vacuum pump. Water was raised to the top of the central tube with a vacuum and allowed to flow over the central tube so that it could be returned to the HRAP (Fig. 1). The raceway was initially filled with natural marine water filtered at 30 μm and supplied with an effluent profile corresponding to European sea bass (Dicentrarchus labrax) breeding tanks providing 80 g N/day and 30 g P/day for 75 days. The total biomass profile consisted of 2000 fish of 80±2.3 g (average body weight) fed with a fixed daily rate (1.2% of the biomass). The experimental natural consortium of marine microalgae was grown under a natural irradiance directed by the local weather at 43° 31′ 59.98″ N, 3° 55′ 59.99″ E in autumn 2017 in France on the western Mediterranean coast. CO₂ addition flow was adjusted by an automatic pH detection device which was adjusted to photosynthetic demand based on pH level monitoring (Galès et al. 2020).

Chlorophyll a concentrations were measured (Lorenzen 1967) twice a week during the exponential period of growth, corresponding to the sample collection period. The data showed an increasing concentration from 0.8 to 3.5 mg/L of chlorophyll a. Culture productivity calculated on the sampling period was 2.53 g/m²/day. Algal consortium biomass was weekly sampled until 5 kg of dried biomass was obtained. Natural algae concentration was pre-concentrated using COLDEP® column (around 10- to 20-fold depending on the culture stage) and centrifuged using a plate divider Alf Laval “Clara15” to obtain a paste featuring an approximate 7% dryness. Residual water was extracted by freeze-drying, and the final product was ground to obtain a meal mesh comparable to the industrial fish meal. The dried consortium biomass (5 kg) was defined in terms of nutrients before being used at graded levels in formulated feeds satisfying the European sea bass nutritional requirements (Peres and Oliva-Teles 1999a).

The species composition of the consortium was determined using 18S rRNA gene analysis. For each experimental run, 10mL samples were filtered through 0.2μm membranes (PALL ALL Supor® 200 PES), the membranes being stored at −20 °C for subsequent DNA extractions. The DNA was extracted using DNeasy PowerWater Kit (Qiagen) according to the manufacturer’s instructions. The V4 region of the 18S rRNA gene was amplified over 30 amplification cycles at an annealing temperature of 65°C, with forward and reverse primers (5′-CTTTCCCTAACCAGCTCTTCGGATCTGCCGTAATTCCAGCTCCAA-3′ and 5′-GGAGTTCACTTCCAGCTCCTTGGCCAATGCCTTGC-3′, respectively). The resulting products were purified and loaded onto an Illumina MiSeq cartridge for sequencing, paired with 300bp reads following the manufacturer’s instructions (v3 chemistry). Sequencing and library preparation steps were carried out at the Genotoul Lifescience Network Genome and Transcriptome Core Facility in Toulouse, France (get.genotoul.fr). A modified version of the standard operation procedure for MiSeq data (Kozich et al. 2013) in Mothur version 1.35.0 (Schloss et al. 2009) was used for alignment and taxonomic outline. Mothur was also used to identify representative sequences of operational taxonomic units (OTUs).

**Test ingredients and diets**

Four diets were formulated to be isoproteic (48.5±0.8) and isolipidic (18.3±0.5). As a control diet (C), a formulation was used that simulated a commercial diet containing fish meal and oil and vegetable-derived protein mix, including solvent-extracted soybean meal, pea protein concentrates, and wheat meal. The microalgae consortium was included to partially replace the vegetable-derived protein mix in diet...
MC10 (10% replacement) and diet MC20 (20% replacement) (Table 1).

A diet (N10) including 10% of commercial *Nannochloropsis* sp. dry biomass was used for comparison. *Nannochloropsis* sp. cells were cultivated in a photobioreactor using industrial chemical fertilizer. The spray-dried *Nannochloropsis* biomass was provided by an industrial algae farm (GREENSEA, Meze-Fr).

Dietary composition was supplemented with L-methionine so that the sulfur amino acid levels met the requirements of the European sea bass (Tulli et al. 2010). Yttrium oxide (20 mg/100 g diet) was added as an indigestible marker to assess nutrients and energy digestibility of the test diets. The diets were manufactured by INRA in Donzacq (F) as standardized 2 mm pellets. The ingredients and proximate composition of the experimental diets are shown in Table 1.

Experimental animals and feeding trial

Six hundred thirty-six European sea bass (*Dicentrarchus labrax*) juveniles were purchased from a commercial hatchery (Poissons du Soleil, Balaruc les-Bains, France).

After 3 weeks of acclimation to the experimental conditions, the fish (average initial body weight 18.0 ± 0.28 g) were randomly allotted among 12 cylindrical tanks, featuring a volume of 1 m³ each (53 fish per tank) and equipped with a collection tube for feces and uneaten pellets in a recirculation aquaculture system (RAS), thus ensuring optimal water conditions for European sea bass (Table 2), and were fed a commercial diet. At the beginning of the feeding trial, the fish were individually implanted with a microchip (Biomark Inc., ID, USA) under moderate anesthesia (90 ppm benzocaine) (Topic Popovic et al. 2012).

The fish were assigned to fish groups/tanks according to a completely random design with diets as the main factor and three replicates per treatment and hand-fed the experimental diets starting on 28th March 2018 over 75 days to apparent satiation in 3 daily meals from 8 am to 2 pm. The fish were group-weighted every 4 weeks and at the end of the feeding trial, under moderate anesthesia after 40 hours fasting. Relative feed intake (RFI= feed intake/[(Initial body weight +Final body weight) × 0.5 × days]), specific growth rate (SGR=100 × (ln Final body weight – ln Initial Body Weight)/days), feed conversion ratio (FCR= feed intake/weight gain), protein efficiency ratio (PER= weight gain/protein intake), and gross protein retention (GPR=100 × [(final body protein content-initial protein content)/protein intake]) were calculated.

At the end of the feeding trial, after 40 hours fasting, 3 fish per tank (9 fish per dietary treatment) were sacrificed with a lethal solution of benzocaine (200 ppm; Vignet et al. 2014). Individual weight and length and viscera, liver, and mesenteric fat weight were recorded. The intestinal tract was excised for histological and physiological evaluations: Fulton’s condition index (K= body weight/standard length³), viscerosomatic index (VSI=100 × viscera weight/body weight), phosphorus and energy content of the test diets

| Table 1 | Ingredients (g/kg) and proximate composition, phosphorus and energy content of the test diets |
|---------|------------------------------------------------------------------------------------------|
| CTRL | MC10 | MC20 | N10 |
| Fishmeal Chile prime | 25.25 | 25.25 | 25.25 | 25.25 |
| Vegetable mix§ | 37.87 | 35.05 | 34.03 | 36.02 |
| Wheat gluten meal | 4.04 | 5.05 | 5.05 | 0.00 |
| Wheat meal | 17.17 | 9.09 | 0.00 | 14.74 |
| Fish oil | 13.94 | 13.73 | 13.73 | 12.12 |
| Microalgal consortium | 0.00 | 10.10 | 20.20 | 0.00 |
| *Nannochloropsis* sp.# | 0.00 | 0.00 | 0.00 | 10.10 |
| Min. and Vit. supplement§ | 1.00 | 1.00 | 1.00 | 1.00 |
| Yttrium oxide | 0.02 | 0.02 | 0.02 | 0.02 |
| Binder | 0.20 | 0.20 | 0.20 | 0.20 |
| L-Methionine | 0.50 | 0.50 | 0.50 | 0.50 |
| Chemical composition | | | | |
| Dry matter (%) | 96.94 | 96.87 | 97.17 | 97.12 |
| Protein (% DM) | 49.17 | 49.26 | 47.61 | 48.24 |
| Lipids (% DM) | 18.14 | 17.84 | 18.18 | 19.12 |
| Ash (% DM) | 7.84 | 12.80 | 18.33 | 10.29 |
| Phosphorus (% DM) | 1.03 | 1.07 | 1.11 | 1.13 |
| Gross energy (MJ/kg) | 23.10 | 22.20 | 21.20 | 22.80 |

§Vegetable mix: including soy protein concentrate, pea protein concentrate, solvent extracted soybean meal in a 4:1:4 ratio

#*Nannochloropsis* sp. from GREENSEA, Meze-Fr

$Mineral supplement composition (% mix): CaHPO4*2H2O, 78.9; MgO, 2.725 g; KCl, 0.005; NaCl, 17.65; FeCO3, 0.335; ZnSO4*H2O, 0.197; MnSO4*H2O, 0.094; CuSO4*5H2O, 0.027; Na2SeO3, 0.067

Vitamin supplement composition (% mix): thiamine HCL Vit B1, 0.16; riboflavin, Vit B2, 0.39; pyridoxine HCL Vit B6, 0.21; cyanocobalamine B12, 0.21; niacin Vit PP, 2.12; calcium pantotenate, 0.63; folic acid, 0.10; biotin Vit H, 1.05; myoinositol, 3.15; stay C Roche, 4.51; tocopherol Vit E, 3.15; menadione Vit K3, 0.24; Vit A (2500 UI/kg diet) 0.026; Vit D3 (2400 UI/kg diet) 0.05; choline chloride, 83.99

| Table 2 | Average and range values of physico-chemical water parameters over 75 days |
|---------|--------------------------------------------------------------------------------|
| Parameter | Average | Min. | Max. |
| Salinity (g/L) | 35.3 | 27.8 | 39.1 |
| Temperature (°C) | 22.8 | 19.7 | 23.2 |
| pH | 7.1 | 6.5 | 7.6 |
| Dissolved oxygen (mg/L) | 7.6 | 5.6 | 6.7 |
| N-NH3 (ppm) | 0.3 | 0.1 | 0.8 |
| N-NO2 (ppm) | 0.0 | 0.0 | 0.0 |
| N-NO3 (ppm) | 1.4 | 0.4 | 3.3 |
| P-PO4 (ppm) | 0.1 | 0.0 | 0.2 |
weight), hepatosomatic index (HSI= 100 × liver weight/body weight), mesenteric fat index (MFI= 100 × mesenteric fat/ body weight), and carcass yield = 100 × carcass weight/ body weight were calculated.

**Diet digestibility evaluation**

To evaluate the in vivo nutrient digestibility of the test diets, fish feces were daily collected from each tank during the last 3 weeks of the feeding trial and preserved at −20°C until used. Fecal biomass was centrifuged (10 min at 3000× g at 4°C), freeze-dried, and stored (− 20 °C) until analyzed. Feed and feces were analyzed for dry matter (AOAC 934.01), protein (AOAC 2001.11), lipids (AOAC 2003.05), and energy (ISO 9831-1998) (IKA – C7000) content. Yttrium concentration in feed and feces was determined by inductively coupled plasma mass spectrometry (ICP-MS) according to Carignan et al. (2001). Apparent digestibility coefficients (ADCs) of dry matter, protein, lipid, and energy of the diets were calculated according to the following formula:

\[
ADC = 1 - \left[ \frac{F}{D} \times \frac{Di}{Fi} \right]
\]

where D = % of the nutrient or kJ/g of the energy in the diet; F = % of the nutrient or kJ/g of the energy in the feces; Di = % Y in the diet; and Fi = % Y in the feces (Cho et al. 1982).

**Gut histology**

Two fish from each tank were used for histologic analyses. Fish gut was dissected, and proximal intestine samples were collected from below the pyloric caeca (0.5cm fragment). Samples were fixed in 4% neutral-buffered formaldehyde and embedded in paraffin. Cross sections of each sample were cut (3 μm thick) in a semi-automated rotary microtome (Leica RM 2245). Slides were then dewaxed and stained with specific Alcian Blue/PAS (pH=2.5). Micrographs of each section were measured in two sections of each sample by using imaging software Olympus cellSens Dimension Desktop: cross-sectional area; muscularis externa thickness (inner circular and outer longitudinal muscle layers); fold length and width and goblet cell presence, as previously described (Batista et al. 2020a, b) (Fig. 2). Briefly, the muscularis externa was measured in eight points of each cross section, and the mean value was considered; the eight highest folds in each section were selected to measure their length and width. Goblet cells (mucus-producing cells) were counted in the eight selected folds (blue and magenta cells), and the average number of goblet cells per fold was determined.

**The activity of the brush border membrane (BBM) enzymes**

One fish per tank was used to obtain the digestive tract that was divided into pyloric caeca (PC), proximal intestine (P, the section from below the PC to the increase in diameter indicating the start of the distal intestine), and distal intestine (D, the terminal part of the intestine with a larger diameter, reaching the anus). When necessary, the remaining feed residues were gently squeezed out. Tissue samples were lightly blotted with absorbent paper, put in individual plastic tubes, and stored at −20°C until the analysis of the BBM enzyme activity was performed. The extraction of the BBM enzymes and the analysis of maltase, sucrase-isomaltase (SI), γ-glutamyl transpeptidase (γ-GT), and alkaline phosphatase (ALP) were carried out as reported by Messina et al. (2019). One unit (U) of enzyme activity corresponded to the amount of enzyme that transforms or hydrolyses 1 μmol of substrate/mL/min. The specific enzymatic activity was calculated as U = μmol/min/ mg of supernatant protein for maltase and sucrase-isomaltase and mU for ALP and γ-GT.

The amount of total protein in the supernatant was determined according to Bradford et al. (1976) by using Bradford reagent (Sigma-Aldrich, Milan, Italy) and bovine serum albumin (Sigma-Aldrich, Milan, Italy) as a standard.
Statistical analysis

Data are expressed as average ± standard deviation. Zootechnical and digestibility data were analyzed by one-way ANOVA to test statistical significance within the main factor. BBM enzyme activity data were analyzed by a two-way ANOVA test, considering the dietary treatment and the intestinal section as main factors. If appropriate, Duncan’s post hoc test was applied at a significant level of 95%. IBM-SPSS statistical package (release 17.0) was used to carry out data analysis.

Results

Marine consortium characterization

The consortium biomolecular characterization identified 34 assignments with 6 dominant algal species (Table 3). The main species of the natural consortium were as follows: *Oocystis* sp., *Chlorella stigmatophora, Tetraselmis* sp. Depending on the open pond culture cycle and the season, *Isochrysis* sp. and *Phaeodactilum tricornutum* were observed in the minority.

The chemical composition of the dried consortium biomass is shown in Table 4. The consortium was characterized by 2.8% nitrogen and 3.2% total lipid. Oleic (16.8% FAMEs) and linolenic (12.4% FAMEs) acids were the main fatty acids. The free amino acid fraction was dominated by proline, alanine, arginine, and glutamate (42.0, 21.3, 16.0, 16.5 nmol/mg, respectively). Natrium, iron, and boron were the most abundant elements in the mineral fraction.

Fish growth performance

During the experimental period, the fish easily accepted the experimental diets, and mortality was negligible. Growth performance, RFI, FCR, and PER of the European sea bass juveniles fed with the experimental diets over 75 days are shown in Table 5. The fish fed with diet MC10 exhibited a significantly higher final body weight as compared to those fed with the control diet (64.9 vs 61.0 g; P<0.05), while the SGR value in MC10, though being the highest, did not reach statistical significance (P = 0.066). The fish fed with diet MC20 exhibited the highest relative feed intake (18.4 g/kg ABW/day), a significantly different value from the RFI of the C group that showed the lowest one (16.7 g/kg ABW/day) (P<0.05). Feeding diet MC20 also resulted in a significant increase in FCR values as compared to the other dietary treatments (1.25 vs 1.15 respectively, P = 0.0052). On the contrary, PER was significantly lowered by the microalgae inclusion in the MC20 group compared to diet C and Nannochloropsis-including diet (1.73 vs 1.83, P<0.05). GPR was not affected by the experimental diets.

The biometric morphometric index values calculated at the end of the trial on European sea bass did not reveal any significant effects of the experimental diets (Table 6).

Diet digestibility

The dry matter, protein, lipid, and energy ADCs of the experimental diets are shown in Table 7. Diet N, including 10% of *Nannochloropsis* sp., resulted in ADC values similar to the ones observed in the reference C diet (74.2, 92.2, 85.4, and 85.8 %, respectively for dry matter, protein, lipid, and energy).

Table 3 Taxonomic composition of the marine consortium

| Taxonomic composition of the marine consortium | Quantity (%) | Properties | Reference |
|-----------------------------------------------|--------------|------------|-----------|
| Algae                                         |              |            |           |
| *Oocystis* sp.                                | 80.32        | High EPA   | Anthony and Stuart 2015 |
| *Tetraselmis* sp.                             | 6.06         | High EPA and ARA | Vizcaíno et al., 2016 |
| *Chlorella stigmatophora*                     | 2.06         | High EPA   | Anthony and Stuart 2015 |
| *Clamydomonas* sp.                            | 1.22         | Mineral (boric acid and calcium) | Kliphuis et al. 2012 |
| *Nannochloropsis gaditana*                    | 0.06         | 15% EPA    | Anthony and Stuart 2015 |
| *Nannochloris* sp.                            | 0.06         | 35% EPA    | Anthony and Stuart 2015 |
| Others                                       |              |            |           |
| *Rotifera* (ciliated)                         | 5.84         | Monounsaturated fatty acid | Awaiss et al. 1992 |
| *Lacrmaridiaceae* (ciliated)                  | 2.09         | x          |           |
| *Cinerochilidae* (Phylasterides)              | 2.07         | x          |           |
| *Chytridiomycotina* (Chytridiomycota)         | 0.18         | x          |           |
| *Strombidiidae* (ciliated)                    | 0.02         | x          |           |
| *Isochrysis*                                  | Traces       | x          |           |
| *Phaeodactylum tricornutum*                   | Traces       | x          |           |
On the contrary, increasing the dietary inclusion of the marine consortium resulted in a significant decrease in dry matter, protein, lipid, and energy ADCs as observed in MC10 (65.9, 90.0, 82.8, 80.1 % respectively) and MC20 (57.7, 85.2, 84.9, 75.2 % respectively) diets (P<0.05).

**Intestine morphology**

The fish fed with the experimental diets did not exhibit major alterations in intestinal morphology, as shown in Table 8. Most of the traits considered did not vary significantly among the dietary treatments (P>0.05), and the intestine from all sampled fish showed a well-preserved morphology. A significant reduction in the total number of acid goblet cells per fold was registered in the fish fed with the highest consortium dietary inclusion level as compared to those fed with the control diet (36.77 vs 64.29 n. GC/fold respectively for MC20 and C, P<0.05) (Fig. 3).

**The activity of the intestinal brush border membrane enzymes**

The activity of maltase, SI, γGT, and ALP varied depending on the intestinal tracts (Fig. 4). The PC was the major site of activity for all enzymes studied. In this tract, the diet containing 10% of *Nannochloropsis* sp. resulted in a significant decrease of the maltase activity as compared to the control diet (3.83 vs 6.41 U, P<0.05).

In the distal portion of the intestine, the activity of SI and maltase showed a similar pattern and their activity resulted significantly enhanced in the fish fed with the consortium-including diets as compared to diet C (2.05 vs 0.96 U and 3.63 vs 1.76, respectively, P<0.05).

The dietary treatment considerably affected the activity of γ-GT in the PC. The highest value was observed in the fish fed with the MC10 diet (P=0.001). The activity of ALP was not affected by the dietary treatments (P > 0.05).

The effect of the two main factors (diet and intestinal tract) was tested on the brush border membrane enzyme activity of the European sea bass fed with the experimental diets. The two-way ANOVA results are summarized in Table 9.
significant interaction between the main factors was revealed for maltase and γ-GT.

### Discussion

The interest in the use of microalgae dry biomass in aquafeeds is recent; several studies already tested the effects of their dietary inclusion in the in vivo trials on different fish species (Hussein et al. 2013; Eryalçın and Yıldız 2015; Haase et al. 2016; Kissinger et al. 2016; Vizcaíno et al. 2016a; Sarker et al. 2020).

Different microalgae species such as *Gracilaria gracilis*, *Nannochloropsis oceanica*, *Tisochrysis lutea*, and *Tetraselmis suecica* have been used as partial replacement of fish meal in diets for European sea bass with no adverse effects on zootechnical performance and intestinal physiology (Cardinaletti et al. 2018; Messina et al. 2019; Valente et al. 2019; Batista et al. 2020a). *Isochrysis* sp. has also been proposed as a source of n-3 PUFA in partial substitution of fish meal in diets for European sea bass without any effects on feed intake and zootechnical performance (Tibaldi et al. 2015). Similar results in the same fish species have been obtained by Haas et al. (2016) when part of fish oil was substituted by *Pavlova viridis* and *Nannochloropsis* sp. Moreover, treatment with microalgae in this trial did not affect the histological aspect of the liver and intestine. Studies carried out on rainbow trout have demonstrated the effectiveness of microalgae when included at low levels in the diets. Sarker et al. (2020) demonstrated that *Isochrysis* sp. and *Schizochytrium* sp. are possible candidates for DHA supplementation in rainbow trout diet formulations, while Sheikhzadeh et al. (2012) showed that dietary *Haematococcus pluvialis* might enhance the antioxidant system when added at 0.3 %.

### Table 5  Zootechnical performance of European sea bass fed the experimental diets over 75 days

| Biometric index | CTRL | MC10 | MC20 | N10 | P value |
|-----------------|------|------|------|-----|---------|
| Mortality (%)   | 1    | 0    | 1    | 0   |         |
| Initial body weight (g) | 18.00 ± 1.10 | 18.50 ± 0.90 | 18.50 ± 1.20 | 18.40 ± 1.00 | 0.088 |
| Final body weight (g) | 61.0 ± 11.00b | 64.9 ± 13.40a | 63.4 ± 12.80ab | 63.70 ± 12.70ab | 0.024 |
| RFI (g/kg ABW/day)¹ | 16.70 ± 0.60b | 17.4 ± 0.10ab | 18.4 ± 0.30a | 17.30 ± 0.30ab | 0.018 |
| SGR (%)² | 1.65 ± 0.02 | 1.70 ± 0.01 | 1.66 ± 0.03 | 1.68 ± 0.02 | 0.066 |
| FCR³ | 1.15 ± 0.03b | 1.16 ± 0.01b | 1.25 ± 0.03a | 1.16 ± 0.03b | 0.005 |
| PER (%)⁴ | 1.82 ± 0.04a | 1.80 ± 0.02ab | 1.73 ± 0.04b | 1.84 ± 0.05a | 0.007 |
| GPR (%)⁵ | 32.09 ± 1.75 | 31.33 ± 0.87 | 30.05 ± 1.76 | 29.10 ± 1.28 | 0.075 |

Data are presented as mean ± standard deviation; values with different letters on the same row are significantly different (P < 0.05), n= 3

¹ RFI (Relative Feed Intake) = \[
\frac{\text{feed intake}}{\text{Initial Body Weight} + \text{Final Body Weight} \times 0.5 \times \text{days}}
\]

² SGR (Specific Growth Rate) = \[
\frac{100 \times \log \frac{\text{Final Body Weight}}{\text{Initial Body Weight}}}{\text{days}}
\]

³ FCR (Feed Conversion Ratio) = \[
\frac{\text{feed intake}}{\text{weight gain}}
\]

⁴ PER (Protein Efficiency Ratio) = \[
\frac{\text{weight gain}}{\text{protein intake}}
\]

⁵ GPR (Gross Protein Retention) = \[
\frac{100 \times \left(\text{Final Body protein content} - \text{Initial Body protein content}\right)}{\text{protein intake}}
\]

### Table 6  Biometric morphometric indices of European sea bass fed the experimental diets over 75 days

| Biometric index | CTRL | MC10 | MC20 | N10 | P value |
|-----------------|------|------|------|-----|---------|
| K¹ | 1.72 ± 0.11 | 1.73 ± 0.11 | 1.70 ± 0.10 | 1.73 ± 0.12 | 0.108 |
| HSI (%)² | 1.18 ± 0.31 | 1.03 ± 0.27 | 1.02 ± 0.28 | 1.27 ± 0.70 | 0.539 |
| VSI (%)² | 10.0 ± 1.20 | 9.54 ± 1.00 | 9.64 ± 1.90 | 11.0 ± 1.90 | 0.162 |
| MFI (%)² | 5.93 ± 1.12 | 5.18 ± 0.90 | 4.85 ± 1.13 | 5.77 ± 2.13 | 0.344 |
| Carcass yield (%)² | 82.89 ± 2.28 | 84.26 ± 1.90 | 83.75 ± 2.29 | 81.19 ± 3.76 | 0.101 |

Data are presented as mean ± SD; values with different letters in the same row are significantly different (P < 0.05) n=9

¹ K (Fulton’s Condition Index) = \[
\frac{\text{body weight}}{\text{std length}^3}
\]

² HSI, VSI, MFI, Carcass Yield = \[
100 \times \frac{\text{weight of liver viscera mesenteric fat carcass}}{\text{body weight}}
\]
However, contrary to the vast majority of studies with microbial biomass, the marine consortium tested in the present study consisted of a certain number of different organisms, namely *Oocystis* sp. (80%), *Tetraselmis* sp. (6%), *Chlorella stigmaphora*, *Chlamydomonas* sp., *Nannochloropsis gaditana*, *Nannochloris* sp., Rotifera, Laciromariidae (ciliated), Cinerochilidae (phylasterides), Chytridiomycotina (Chytridiomycota), and Strombidiidae (ciliated). The analysis method cannot precisely define the *Oocystis* species, but whereas *Oocystis* sp. is generally associated with a freshwater genus of green microalgae, oceanic strains can be found in marine or brackish water such as *Oocystis submarina* (Śliwińska-Wilczewska and Latala 2018) or *Oocystis borgei* which could inhibit harmful microalgae by expressing allopathic effects (Wang et al. 2020). *Oocystis heteromucosa* belongs to strains of marine algae consortium found in marine aquaculture pond wastewater with a high ammonia tolerance (Katayama et al. 2020). In a HRAP, algae predators such as rotifers can have a negative effect on the consortium growth to the point of cultural annihilation. That was not the case in our experiment: we hypothesize that continuous CO2 delivery linked to the photosynthetic demand maintains a pH value around 7, which could be uncomfortable for organism reproduction adapted to marine pH at 8.2. In addition, the culture did not collapse, probably because of the high culture volume and dynamic algae growth. Previous experiments showed that the algae cells’ reproduction rate has to be higher than the total grazers’ reproduction rate in order to keep a culture alive (Strom and Morello 1998). That was the case in the exponential culture growth phase, which is the sample period for biomass extraction. Algae predators also sequester compounds of some interest from algae ingestion and their tolerated presence in the culture contributes to final powder value.

The use of a non-axenic culture of a blend of *Tetraselmis suecica*, *Isochrysis galbana*, and *Dunaliella tertiolecta* has been evaluated in its remediation potential for the nutrient assimilation of fish farm wastewater (Andreotti et al. 2017). Dallaire et al. (2007) have previously described the effect on trout fry of the dietary inclusion of a freshwater photosynthetic microorganism consortium (mainly *Scenedesmus* sp. and *Chlamydomonas* sp.) derived from the sedimentation pond of a fish farm. The results showed that a maximum of 12.5% of consortium could be included in the feed formulation without affecting growth or whole-body fish composition. Anyway, to the best of our knowledge, there seem to be no other studies considering the dietary inclusion of a non-axenic multi-species marine consortium in fish feeds. For these reasons, the comparison of the present results with previous research studies is not straightforward and should be done with caution. In any case, the results of the present study are consistent with the ones of Dallaire et al. (2007) as the dietary inclusion of microalgae biomass generally improved fish performance and feed intake, although only the fish fed with the 10% microalgae consortium (MC10) reached a significantly

### Table 7  Nutrient and energy apparent digestibility coefficients (%) of the experimental diets

|                        | CTRL       | MC10       | MC20       | N10        | P value |
|------------------------|------------|------------|------------|------------|---------|
| Dry matter             | 76.7 ± 1.0a| 65.9 ± 2.0bc| 57.7 ± 6.2c| 74.2 ± 1.1ab| 0.000   |
| Protein                | 92.7 ± 0.3a| 90.0 ± 0.5b | 85.2 ± 1.2c| 92.2 ± 0.1a | 0.000   |
| Lipid                  | 86.8 ± 0.2a| 82.8 ± 0.8b | 84.9 ± 2.3ab| 85.4 ± 0.3ab| 0.023   |
| Energy                 | 87.4 ± 0.9a| 80.1 ± 1.3b | 75.2 ± 3.4c| 85.8 ± 0.8a | 0.000   |

Data are presented as mean ± SD; values with different letters in the same row are significantly different (P < 0.05), n=3

### Table 8  Intestinal morphology of European sea bass fed the experimental diets over 75 days

|                        | CTRL       | MC10       | MC20       | N10        | P value |
|------------------------|------------|------------|------------|------------|---------|
| Cross-sectional area (mm²) | 11.2 ± 2.97| 11 ± 4.41 | 9.5 ± 2.29 | 10.2 ± 1.62| 0.761   |
| Villus length (μm)      | 1414.7 ± 256.7| 1416.5 ± 368.5| 1327.6 ± 207.9| 1240.0 ± 126.6| 0.592   |
| Villus width (μm)       | 226.9 ± 34.76| 225.1 ± 32.41| 222.3 ± 29.50| 225.9 ± 14.85| 0.993   |
| Muscularis externa (μm) | 131.7 ± 20.45| 104.4 ± 21.69| 106.5 ± 30.88| 129.3 ± 17.68| 0.103   |
| Inner circular layer (μm) | 85.1 ± 13.23| 67.4 ± 14.23| 65.6 ± 18.58| 81.4 ± 10.69| 0.069   |
| Outer longitudinal layer (μm) | 46.6 ± 8.26| 37 ± 7.83 | 40.9 ± 13.23| 47.9 ± 9.40 | 0.226   |
| Goblet cells (no. GC/fold) | 88.8 ± 16.94| 67.2 ± 20.55| 56.7 ± 13.05| 82.2 ± 29.64| 0.061   |
| Acid GC (no. GC/fold)   | 64.3 ± 13.46a| 46.4 ± 8.14ab| 36.8 ± 6.26b| 54.7 ± 17.44ab| 0.006   |
| Neutral GC (no. GC/fold) | 24.6 ± 15.54| 20.8 ± 14.92| 19.9 ± 8.13 | 27.6 ± 15.21| 0.760   |

Data are presented as mean ± SD; values with different letters in the same row are significantly different (P < 0.05), n=6
higher final body weight than the control diet. MC20 diet had a growth performance that did not differ from other treatments but resulted in a significant increase of FCR. Similarly, the rainbow trout fed with diets including 9.5% of a *Nannochloropsis* and *Isochrysis* blend or including also *Schizochytrium* exhibited significantly poorer FCR, a result not unlike those obtained by Walker and Berlinski (2011) on cod or Cardinaletti et al. (2018) on European sea bass.

For the above-mentioned reason, in the present study, a comparison with a test diet including monospecific dried biomass of *Nannochloropsis* spp. has been considered in the experimental design. *Nannochloropsis* sp. is a unicellular microalgae with a polysaccharide cell wall (Hibberd 1981) and a promising ingredient in aquafeeds both as a successful fish oil substitute (Eryalçın and Yıldız 2015; Gbadamosi and Lupatsch 2018; Lozano-Muñoz et al. 2020) and the form of the defatted meal as an alternative to fish meal (Sørensen et al. 2017). Moreover, *N. oceanica* became better digested by European sea bass than other microalgae marine species (Batista et al. 2020b). In the present study, the replacement of terrestrial plant source by *Nannochloropsis* sp. dried biomass did not significantly affect diet palatability, fish growth performance, or biometric indices compared with C diet after a 75-day feeding period, confirming previous observation in the European sea bass fed with 8% of *N. oceanica* (Batista et al. 2020a). A recent study carried out by Valente et al. (2019) showed that the dietary inclusion up to 15% of defatted *Nannochloropsis* sp. biomass can replace fish meal in European sea bass diets without affecting fish growth performance and biometric indices. Moreover, Haas et al. (2016) showed that a 50% fish oil replacement by *Nannochloropsis* sp. biomass did not hamper the growth performance of juvenile European sea bass. Other studies considering different dietary inclusion of *N. oceanica* indicate that moderate inclusion levels (<15 g/kg diet) do not affect growth and feed performance in other fish species like spotted wolffish (*Anarhichas minorhas*) and Atlantic salmon (*Salmo salar*) (Sørensen et al. 2017; Knutsen et al. 2019). On the contrary, higher dietary inclusion levels of *Nannochloropsis* sp. biomass hampered growth and feed conversion in Nile

![Fig. 3](image)

Anterior intestine histological sections (Alcian Blue/PAS staining, pH = 2.5) of European sea bass at the end of 75-day feeding trial. Blue points represent the acid goblet cells. A, CTRL diet and B, MC20 diet.

![Fig. 4](image)

Specific activity of SI (sucrase-isomaltase), maltase, ALP (alkaline phosphatase), and γ-GT (gamma glutamyl transpeptidase) in PC (pyloric caeca), P (proximal intestine), and D (distal intestine) of European sea bass fed the experimental diets over 75 days. Data are presented as means ± SD (n=3). Different letters indicate significant differences among the treatment diets (lower case superscript *P* < 0.05, capital superscript *P* < 0.001).

Table 9 ANOVA of the main factors which affect the activity of the BBM enzymes

| Diet | Tract | Diet × Tract |
|------|-------|-------------|
| Maltase | * | ** | ** |
| Sucrase | * | ** | NS |
| ALP | * | ** | NS |
| γ-GT | ** | * | ** |

*P* < 0.05; **P** < 0.001
Electronic supplementary material

Figure S1. Histogram of wet samples from the different treatments. 

Figure S2. Enzyme activity of different treatments. 

Table S1. Nutrient content of the different diets. 

Table S2. Digestibility of the different treatments. 

Table S3. Food intake of the different treatments. 

Table S4. Weight gain of the different treatments. 

Table S5. Survival rate of the different treatments. 

Table S6. Liver weight of the different treatments. 

Table S7. Total protein of the different treatments. 

Table S8. Total lipid of the different treatments. 

Table S9. Total energy of the different treatments. 

Table S10. Total carbohydrate of the different treatments.

Table S11. Total starch of the different treatments.

Table S12. Total dietary fiber of the different treatments.

Table S13. Total ash of the different treatments.

Table S14. Total crude protein of the different treatments.

Table S15. Total crude lipid of the different treatments.

Table S16. Total crude energy of the different treatments.

Table S17. Total crude fiber of the different treatments.

Table S18. Total crude ash of the different treatments.

Table S19. Total dietary protein of the different treatments.

Table S20. Total dietary lipid of the different treatments.

Table S21. Total dietary energy of the different treatments.

Table S22. Total dietary fiber of the different treatments.

Table S23. Total dietary ash of the different treatments.

Table S24. Total dietary crude protein of the different treatments.

Table S25. Total dietary crude lipid of the different treatments.

Table S26. Total dietary crude energy of the different treatments.

Table S27. Total dietary crude fiber of the different treatments.

Table S28. Total dietary crude ash of the different treatments.

Table S29. Total dietary protein of the different treatments.

Table S30. Total dietary lipid of the different treatments.

Table S31. Total dietary energy of the different treatments.

Table S32. Total dietary fiber of the different treatments.

Table S33. Total dietary ash of the different treatments.

Table S34. Total dietary crude protein of the different treatments.

Table S35. Total dietary crude lipid of the different treatments.

Table S36. Total dietary crude energy of the different treatments.

Table S37. Total dietary crude fiber of the different treatments.

Table S38. Total dietary crude ash of the different treatments.

Table S39. Total dietary protein of the different treatments.

Table S40. Total dietary lipid of the different treatments.

Table S41. Total dietary energy of the different treatments.

Table S42. Total dietary fiber of the different treatments.

Table S43. Total dietary ash of the different treatments.

Table S44. Total dietary crude protein of the different treatments.

Table S45. Total dietary crude lipid of the different treatments.

Table S46. Total dietary crude energy of the different treatments.

Table S47. Total dietary crude fiber of the different treatments.

Table S48. Total dietary crude ash of the different treatments.

Table S49. Total dietary protein of the different treatments.

Table S50. Total dietary lipid of the different treatments.

Table S51. Total dietary energy of the different treatments.

Table S52. Total dietary fiber of the different treatments.

Table S53. Total dietary ash of the different treatments.

Table S54. Total dietary crude protein of the different treatments.

Table S55. Total dietary crude lipid of the different treatments.

Table S56. Total dietary crude energy of the different treatments.

Table S57. Total dietary crude fiber of the different treatments.

Table S58. Total dietary crude ash of the different treatments.

Table S59. Total dietary protein of the different treatments.

Table S60. Total dietary lipid of the different treatments.

Table S61. Total dietary energy of the different treatments.

Table S62. Total dietary fiber of the different treatments.

Table S63. Total dietary ash of the different treatments.

Table S64. Total dietary crude protein of the different treatments.

Table S65. Total dietary crude lipid of the different treatments.

Table S66. Total dietary crude energy of the different treatments.

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Table S75. Total dietary crude lipid of the different treatments.

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Table S87. Total dietary crude fiber of the different treatments.

Table S88. Total dietary crude ash of the different treatments.

Table S89. Total dietary protein of the different treatments.

Table S90. Total dietary lipid of the different treatments.

Table S91. Total dietary energy of the different treatments.

Table S92. Total dietary fiber of the different treatments.

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Table S94. Total dietary crude protein of the different treatments.

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Table S96. Total dietary crude energy of the different treatments.

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Table S99. Total dietary protein of the different treatments.

Table S100. Total dietary lipid of the different treatments.

Table S101. Total dietary energy of the different treatments.

Table S102. Total dietary fiber of the different treatments.

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Table S104. Total dietary crude protein of the different treatments.

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Table S131. Total dietary energy of the different treatments.

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Table S138. Total dietary crude ash of the different treatments.

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Table S141. Total dietary energy of the different treatments.

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Table S143. Total dietary ash of the different treatments.

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Table S146. Total dietary crude energy of the different treatments.

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Table S148. Total dietary crude ash of the different treatments.

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Table S152. Total dietary fiber of the different treatments.

Table S153. Total dietary ash of the different treatments.

Table S154. Total dietary crude protein of the different treatments.

Table S155. Total dietary crude lipid of the different treatments.

Table S156. Total dietary crude energy of the different treatments.

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Table S158. Total dietary crude ash of the different treatments.

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Table S160. Total dietary lipid of the different treatments.

Table S161. Total dietary energy of the different treatments.

Table S162. Total dietary fiber of the different treatments.

Table S163. Total dietary ash of the different treatments.

Table S164. Total dietary crude protein of the different treatments.

Table S165. Total dietary crude lipid of the different treatments.

Table S166. Total dietary crude energy of the different treatments.

Table S167. Total dietary crude fiber of the different treatments.

Table S168. Total dietary crude ash of the different treatments.
Conclusion

This is so far the first study aimed at evaluating the dietary utilization of multispecific marine microalgae consortium biomass originated from a HRAP for a commercially relevant species. The results support a possible substitution of up to 10% of terrestrial vegetable ingredients by the microalgae consortium dried biomass with a significant increase of European sea bass final body weight, though impairing nutrient and energy digestibility. *Nannochloropsis* sp. biomass also has the potential to partially substitute terrestrial plant ingredient up to 10% of the diet without affecting growth performance, dietary nutrient utilization, and gut enzymatic activities. Algal consortium and *Nannochloropsis* sp. biomass could undergo specific processing techniques before being included in fish feed formulation to improve nutrient bioavailability. To increase aquaculture sustainability, this study using fish farm effluents to produce a multispecific marine non-axenic valuable biomass represents the first attempt to enhance a circular use of natural biomasses aquafeeds. Such an approach still needs further efforts, and the safety issues connected with their utilization need specific evaluations.

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Availability of data and materials

All data generated or analyzed during this study are included in this published article.

Author contribution

CP and GD provided the microalgae consortium and controlled the sea bass rearing plant. MM, BO, LP, and GP performed the laboratory examinations. FT performed the analysis of the data and supervised the project. LMPV provided the funding. GP, MM, and FT wrote the original draft. All the authors reviewed and approved the final manuscript.

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Declarations

Ethics approval and consent to participate

The handling procedures and sampling methods involving fish were in accordance with the guidelines of the European Union Directive 2010/63/EU on the protection of animals used for scientific purposes (Ethical authorisation: APAFIS#12871-2018091215242876).

Consent for publication

Not applicable

Competing interests

The authors declare no competing interests.

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