Aquaculture contributes to global food security but sustainable aquaculture development in terms of biodiversity impacts requires the establishment of viable solutions in replacement of the fisheries-based components in aquafeeds. In the current work, pit-tagged Atlantic salmon individuals were grown, from fresh water (18 g body weight) to salt water in tanks (up to 800 g body weight), on diets with low fish meal (FM) (10%) and 1–1.25% total n-3 LC-PUFA levels balanced across the experimental diets. Dietary n-3 LC-PUFAs were supplemented by (1) fish oil (FO), (2) *Schizochytrium limacinum* biomass (ScB), or (3) a mix of the two (FO/ScB). Further, the fish from all treatments were mixed and redistributed in sea cages reared to slaughter (ca. 3 kg body weight) on either FO or ScB. As FO was rich in both EPA and DHA and ScB was rich in DHA and nearly devoid of EPA, the experimental diets differed significantly in DHA/EPA ratio (0.8 vs. 10.5 in average for FO and ScB, respectively). ScB treatment fish grew to significantly higher body weight in the end of the experiment (2.8 kg vs. 3.3 kg, for FO and ScB, respectively) but similar FCR, survival rate, and biometric indexes compared to the FO groups. ScB fish contained higher levels of EPA + DHA in the filet but lower in the liver, and better filet pigmentation already from the tank phase of the experiment as determined chemically, by salmonfan and a trained sensory panel, and lower prevalence of melanin spots at slaughter. The trained sensory panel found no differences in flavor or odor in the filets from the different dietary groups; however, filets in the FO group were perceived as softer and juicier compared to ScB. The pre-diets up to 800 g body weight had minor effects on fish performance. Global transcriptomics in liver and intestinal tissues revealed significant dietary effects on the expression of immune modulating, as well as ion, lipid, protein, and xenobiotic metabolism genes.

**Keywords:** DHA, Atlantic salmon, n-3 LC-PUFA, filet quality, *Schizochytrium limacinum*
INTRODUCTION

Novel sustainable sources of high-quality proteins and long chain n-3 polyunsaturated fatty acid (n-3 LC-PUFA) rich oils are needed, as feed grade fisheries resources exploited for fish oil (FO) and FM production reach sustainability limits (Pauly et al., 2002; Hilborn et al., 2003; Pike and Barlow, 2003; Heal and Schlenker, 2009). In salmon feeds, FM is largely substituted by different plant protein sources (Ytrestøy et al., 2015; Aas et al., 2019) whereas fewer alternatives are commercially relevant when it comes to dietary n-3 LC-PUFA sources. Genetically modified plants can be used for the production DHA and EPA rich oil, such as in the recently tested omega-3 canola oil successfully replacing FO in juvenile salmon diets (Ruyter et al., 1994b; Bowles et al., 1999), for both biofuel and other food and feed applications including fish feeds (Barclay, 1994a; Barclay and Zeller, 1996; Barclay et al., 2005). S. limacinum biomass contains high levels of fat (55–75% in dry matter) and docosahexaenoic acid (DHA) (up to 49% of total lipids) (Nakahara et al., 1996; Rent et al., 2010), and is reported as a good DHA-rich lipid source for farmed seabream Sparus aurata (Ganuza et al., 2008) and Atlantic salmon, even at 100% replacement of the supplemental dietary FO (Carter et al., 2003; Miller et al., 2007; Kousoulaki et al., 2015).

Based on records of fish growth, feed efficiency, intestine histology, survival and liver cell transcriptomics, S. limacinum biomass appears to attribute benefits on fish filet quality and n-3 LC-PUFA retention efficiency without indications of negative physiological impacts for the fish in short-term studies (Kousoulaki et al., 2015, 2016).

In the current study we present results of a lifelong study on Atlantic salmon fed practical diets with low levels of FM and either FO or whole S. limacinum biomass as supplementary dietary n-3 LC-PUFA source, at 1–1.25% total dietary EPA + DHA and similar dietary n-3/n-6 ratio and total saturated fatty acids (SFAs) levels among the experimental diets. Production performance, tissue composition, filet quality, intestinal histology, and liver and intestine microarray results are discussed.

MATERIALS AND METHODS

Formulation and Chemical Analyses in Experimental Feeds

Three commercially relevant Atlantic salmon diets were formulated (Table 1), based on the trends on average raw material levels and nutrient values used in salmon feeds by the three major salmon feed producers in Norway in 2012 and 2013 (Ytrestøy et al., 2015). The experimental diets were formulated to contain 1–1.25% EPA + DHA (in the diet) and were further balanced for crude protein, crude lipid, digestible energy, total SFAs, and n-3/n-6 fatty acid ratio, using different oil blends and plant protein mixes. FO were the control diets containing low FM level (FM in diet FO: 10%), and FO as supplementary n-3 LC-PUFA source. FO/ScB, used only in the tank phase trials, also contained 10% FM and a mix of FO and S. limacinum biomass, the latter providing 50% of total EPA + DHA in the diets, whereas ScB diets contained only S. limacinum biomass as supplementary n-3 LC-PUFA source, besides the respective dietary contribution of FM. The S. limacinum biomass used was whole and spray dried (Alltech Inc., United States). The chemical composition of the biomass is as described in Kousoulaki et al. (2016). The experimental diets were produced at the Feed Technology Center of Nofima in Bergen, Norway, in the same production series, using a Wenger TX-52 co-rotating twin-screw extruder with 150 kg h⁻¹ capacity. The settings of the extruder were “normal,” i.e., the production can be up scaled to a feed factory [extruder settings considered: screw configuration (D), die opening (4.5 mm), knife speed (1494–1981 rpm), Specific Mechanical Energy (7.7–9.5 kW), feed rate (150 kg/h), and amount of steam (0 kg/h) and water (0.15–0.41 kg/min) added to the process]. The levels of S. limacinum biomass used in the experimental diets FO/ScB and ScB were within the range that allows the production of good hot extruded pellets under similar conditions (Samuelsen et al., 2018) with high nutrient digestibility (Kousoulaki et al., 2015, 2016).

Feeds were analyzed for proximate composition according to standard methods: Crude protein: (Kjeldahl method N × 6.25; ISO 5983, 1997), moisture (ISO 6496, 1999), ash (ISO 5984, 2002), and lipid (Bligh and Dyer, 1959). Preparation of fatty acid methyl esters (FAMES) for the determination of fatty acid profile in raw materials and feeds was realized according to the AOCS Official Method Ce 1b-89 using a trace GC gas chromatograph (Thermo Fisher Scientific) with flame ionization detector (GC–FID), equipped with a 60 m × 0.25 mm BPX-70 cyan propyl column with 0.25 μm film thickness (SGE, Ringwood, VIC, Australia). Helium 4.6 was used as a mobile phase under the pressure of 2.60 bar. The injector temperature was 250°C and the detector temperature was 260°C. The oven was programmed as follows: 60°C for 4 min, 30°C min⁻¹ to 164°C, and then 1.0°C min⁻¹ to 213°C, and 100°C min⁻¹ to 250°C where the temperature was held for 10 min. The sample solution (3.0 μl) was injected split-less and the split was opened after 2 min. The FAMES were identified by comparing the elution pattern and relative retention time with the reference FAME mixture (GLC-793, Nu-Chek Prep Inc., Elysian, MN, United States). Chromatographic peak areas were corrected by empirical response factors calculated from the areas of the GLC-793 mixture. FA composition was calculated by using 23:0 FAME as internal standard and reported on a sample basis as g/100 g FAMES. Dietary gross energy was determined in a Parr adiabatic bomb calorimeter. All aforementioned analyses were performed at Nofima laboratories in duplicate. If differences between parallels exceeded standardized values, new duplicate analyses were carried out. Yttrium in feeds and feces was analyzed at an external laboratory (Eurofins, Molde, Norway).

Fish Feeding Trials

The trial period commenced in tanks in February 2015, and lasted ca. 11 months, i.e., until December 2015. During the tank phase, individually pit-tagged fish were fed one of three experimental
**TABLE 1 |** Experimental diets’ formulations (raw material inclusion levels in %) and chemical composition.

| Growing phase | Fresh water phase (parr) | Saltwater tank phase 1 (smolt) | Saltwater tank phase 2 (post-smolt) | Sea cage phase (slaughter) |
|---------------|--------------------------|-------------------------------|-------------------------------------|---------------------------|
| Diet name     | FO1 FO/ScB1 ScB1 | FO2 FO/ScB2 ScB2 | FO3 FO/ScB3 ScB3 | FO4 ScB4 |
| DHA/EPA ratio | 0.72 0.94 7.20 | 1.12 3.56 10.50 | 0.71 2.00 12.33 | 0.69 14.73 |
| DHA + EPA (%) | 0.95 0.99 0.87 | 0.77 0.87 1.00 | 0.97 0.98 1.01 | 1.25 1.25 |
| Fish meal Super Prime | 9.00 9.00 9.00 | 10.00 10.00 10.00 | 10.00 10.00 10.00 | 10.00 10.00 |
| Soluble fish protein concentrate | 8.45 8.45 8.45 | 3.50 3.50 3.50 | 2.50 2.50 2.50 | 2.50 2.50 |
| Krill hydrolysate | | | | |
| Cod fish powder | 2.25 2.25 2.25 | | | |
| Schizochytrium limacinum | 2.62 5.24 | 2.75 5.50 | 2.75 5.50 | 6.25 |
| Fish oil | 3.83 1.92 | 3.70 1.85 | 3.70 1.85 | 4.30 |
| Camelina oil | 6.34 6.78 7.28 | 2.50 2.50 2.50 | | |
| Rapeseed oil | 4.29 6.19 7.05 | 4.78 5.88 6.89 | 7.65 8.79 9.87 | 14.34 17.50 |
| Linseed oil | 3.56 3.75 3.98 | 6.80 7.05 7.29 | 9.25 10.60 | |
| Palm oil | 1.88 0.94 | 2.77 1.90 1.04 | 2.77 1.82 0.94 | 4.55 1.00 |
| Soy protein concentrate | 23.5 23.5 23.5 | 26.50 26.50 26.50 | 23.50 23.50 23.50 | 25.00 25.00 |
| Wheat/wheat gluten | 21.25 20.32 19.21 | 17.00 17.00 17.00 | 21.00 21.06 21.11 | 5.15 5.45 |
| Horse beans | 11.00 10.33 9.80 | 16.39 15.07 13.79 | 12.78 11.38 9.99 | 16.56 13.35 |
| Vitamin mix | 3.04 3.04 3.04 | 3.00 3.00 3.00 | 3.00 3.00 3.00 | 3.00 3.00 |
| Phosphate | 2.00 2.00 2.00 | 2.70 2.70 2.70 | 2.70 2.70 2.70 | 2.00 2.00 |
| Choline chloride + soy lecithin | 1.00 1.00 1.00 | 1.00 1.00 1.00 | 1.00 1.00 1.00 | 1.00 1.00 |
| Lys | 0.80 0.80 0.80 | 0.75 0.75 0.75 | 0.75 0.75 0.75 | 0.50 0.50 |
| Mineral mix | 0.61 0.61 0.61 | 0.64 0.64 0.64 | 0.64 0.64 0.64 | 0.64 0.64 |
| Cholesterol | 0.40 0.50 0.50 | 0.50 0.50 0.50 | 0.50 0.50 0.50 | 0.40 0.40 |
| Ethanol | 0.23 0.23 0.23 | 0.25 0.25 0.25 | 0.25 0.25 0.25 | 0.25 0.25 |
| Carop. Pnp (10%) | 0.05 0.05 0.05 | 0.05 0.05 0.05 | 0.05 0.05 0.05 | 0.05 0.05 |
| Yttrium oxide | 0.05 0.05 0.05 | 0.01 0.01 0.01 | 0.01 0.01 0.01 | 0.01 0.01 |
| Inositol | 0.03 0.03 0.03 | | | |

**Table 1 continued...**
diets (FO, FO/ScB, or ScB) in triplicate tanks and were weighed and samples were taken for tissue analyses from 10 fish per tank four times: at start (ca. 18 g fish mean body weigh BW), soon after smoltification (at ca. 60 g BW), at ca. 200 g BW and at ca. 800 g BW. At the end of the tank phase of the trial all fish were mixed and transferred to the cage facilities of Marine Harvest at Averøy, Norway, where they were distributed by equal representation of fish from each tank into each of six trial cages and reared on either FO or ScB in triplicate, for over one more year (Supplementary Figure S1). This was realized in order to detect early life and dilution effects of variable dietary levels of DHA, EPA, and feeding salmon relatively high levels microalgae biomass as a replacement for FO. In January 2017, the trial was ended, and fish had cage mean BW of ca. 2800–3300 g. During samplings, fish that would go further in the trial were sedated using low doses of tricaine methane-sulfonate (MS-222 at 0.01–0.02 g/l) prior to further handling (weighing and pit-tag reading). Sample fish were euthanized with an overdose of the same anesthetic (MS-222 0.05–0.08 g/l) before tissue sampling.

**Tank Phase (Fresh and Saltwater)**

Each one of the experimental feeds was given to triplicate populations of Atlantic salmon parr kept in tanks at the land-based facilities of Nofima at Sunndalsøra, Norway. The start BW of the experimental fish was 18 g and the final BW was ca. 800 g. During the freshwater phase the fish were kept in 150-L circular fiberglass tanks, with 2.3 kg initial and 4.8 kg final biomass per tank (15.3 and 32.0 kg/m$^3$ density, respectively). During the saltwater phase the fish were kept in 2.3 m$^3$ fiberglass tanks with 60 cm water depth, at 4.8 kg initial biomass and 56.2 kg per tank final biomass (3.1 and 40.7 kg/m$^3$ density, respectively). Feeding was continuous using automatic feeders. The daily feed amounts given were calculated to provide 120% of the fish’s satiation levels. Uneaten feed was collected and weighed daily for the estimation of total daily feed intake in each tank only during the salt water stage of the tank trial period, i.e., after smoltification from ca. 60 to 800 g body weight, but not during the fresh water phase as feed pellets were too small to be collected in a representative way. The mean water temperature during the trial was ca. 9°C during fresh water and saltwater phase 1, and 11.5°C during tank saltwater phase 2. In the saltwater phases, sea water with a salinity ranging between 32 and 33 ppt was used. The water system was flow-through using UV treated and filtrated fresh or sea water from 40 m depth. At the end of the tank trial phase, fish from each tank were pit-tag scanned, weighed, measured for biometric indexes, sampled for tissues, and stripped for feces, the latter were separated from urine and collected in a pre-weighed box per tank and frozen immediately at −20°C until further analyses. At the end of the tank trial phases, whole fish were used for lipid and fatty acid profile analysis and fish filets were used for sensory, lipid, and fatty acid profile analyses.

**Sea Cage Phase**

Atlantic salmon from the end of the tank trial [section “Tank phase (fresh and saltwater)"], at ca. 800 g average body weight, were transferred from the in-door tanks in Nofima Sunndalsøra, Norway, to six seawater net-cages at Marine Harvest’s facilities at Averøy, Norway. The fish were transported by a truck with six separate tanks containing 81 fish each dedicated for each one of the six-destination net-pens. Fish from all three pre-treatments were equally distributed to each one of the six net-cages, 27 fish from each pre-treatment (FO, FO/ScB, and ScB) in each net-cage, approximately 71 kg initial biomass per cage. Diets FO and ScB were randomly assigned to three cage each and fed until slaughter resulting in six pre-end-feed dietary treatments in total (FO:FO, FO/ScB:FO, and ScB:FO in the FO fed cages and FO:ScB, FO/ScB:ScB, and ScB:ScB in the ScB cages). The net-cages (125 m$^3$) were equipped with automatic feeders (Betten, Vågland, Norway), feeding three times × 30 min each day, and equipment for collection of feed waste. The fish were raised to a weight of ca. 3 kg, before final harvest and sampling. Sea transfer took place in December 2015, when water temperature was 7.4°C, but declining. Minimum temperatures were recorded during the winter and were 3.8°C, and the maximum water temperature recorded was 17.9°C during summer 2016.

Lice removal was required seven times during the months from February to October 2016 and was realized by hand. Slice was also given during 1 week in September 2016. Cleaner fish Labrus wrasse (Labrus bergylta) were introduced in the cages in September 2016, at ratios up to 24% of the salmon individual.
numbers. Nets were changed six times during the cage trial. In October 2016, sexually mature fish and obvious “losers” were removed, as such fish tend to attract even more lice, and their predominance was similar in the two treatments. At the end of the trial all fish were pit-tag scanned, weighed, and measured for standard length. Fish were sampled for analyses in three batches, where each batch consisted of three randomly selected fish per pre-treatment group (nine fish in total per sea cage). Batches 1 and 2: Fish were anesthetized to death before sampling of (i) fish for whole body analyses and (ii) liver, skeletal muscle (the cutlet between the posterior end of the dorsal fin and the intestine, termed Norwegian Quality Cut, NQC), and intestine tissue, respectively. Batch 3: The fish were cut by the gills after anesthetization, bled for 10 min in iced seawater before filets were sampled for filet technical quality, color scoring, and analyses by a trained sensory panel. All sampled fish were visually scored for intestinal fat and liver color. Feces were collected from the rest of the fish in each net pen. Whole filets (Batch 3) were stored on ice for 1 week before they were subjected to quality analyses of raw filets and cutting into 2.5 cm × 2.5 cm pieces, vacuum packed, and frozen before sensory assessment of raw and cooked samples.

**Fish Filet and Whole-Body Composition and Filet Technical Quality**

Fish filet and whole-body nutritional quality were evaluated in terms of fatty acid, technical, and sensory quality. Melanin discoloration (presence of melanin spots) and gaping were evaluated on whole filet sides, while mechanical properties (firmness) and filet color were analyzed in duplicate on the dorsal (anterior) and NQC filets part (posterior). Astaxanthin and idoxanthin levels were analyzed in the NQC cutlet. NQC cutlets from nine fish per tank and three fish per dietary treatment per cage at slaughter were pooled, homogenized, and analyzed for fatty acid composition. Lipid extraction was realized according to Bligh and Dyer (1959) and the fatty acid composition according to AOCS Ce 1b-89 as described in section “Formulation and Chemical Analyses in Experimental Feeds.” Filet color was evaluated visually using DSM SalmonFan™ (score 21–34). Instrumental analyses of fish filet firmness were performed perpendicular to the muscle fibers using a Texture analyzer, TA-XT2 (Stable Micro System Ltd., Surrey, United Kingdom) equipped with a flat-ended cylindrical probe (12.5 mm diameter, type P/0.5) at a 1 mm/s traveling speed and a 30 kg load cell. Firmness predicted using this method correlates well with sensory assessment of firmness of raw and smoked salmon filets (Mørkore and Einen, 2003). Melanin discoloration was evaluated visually as the number of dark, hyperpigmented myotomes (Mørkore et al., 2015).

Homogenized filets from individual fish were thawed, and carotenoids were extracted using a 1:1:3 mixture of distilled water, methanol (containing 500 mg l⁻¹ BHT), and chloroform according to Bjerkeng et al. (1997). The solvent was removed from an aliquot under reduced pressure and re-dissolved in mobile phase (acetone/n-hexane/methanol 20:80:0.1), filtered through a 0.45-µm filter (Minisart SRP 15, Göttingen, Germany), and analyzed isocratically by HPLC. A Spherisorb S5-CN nitrile column (PhaseSep, Queensferry, Clywd, United Kingdom; length 250 mm; internal diameter 4.6 mm; particle size 5 mm), using 20% acetone in n-hexane as the mobile phase was used to determine astaxanthin and 3′,4′-cis and 3′,4′-trans glycolic isomers of idoxanthin in muscle samples. The retention times (R_f) of astaxanthin and the 3′,4′-cis and 3′,4′-trans glycolic isomers of idoxanthin on this system were about 4, 6.5, and 8.5 min, respectively. The Shimadzu LC-10AS Liquid Chromatograph was connected to a Shimadzu SPD-M6A Photodiode array UV–VIS detector and the detection wavelength was set to 470 nm. Standards of known concentration were prepared from crystalline all-E-astaxanthin (Hoffmann-La Roche Ltd., Basel, Switzerland), and the concentration of the standard solution was measured spectrophotometrically (UV-260, Shimadzu, Japan) using molar absorptivity E₁%₁ cm = 2100 at absorbance maximum (λ_max = 470 nm) in n-hexane containing 4.5% chloroform. The percentages of the different isomers were calculated from chromatogram areas and corrected for differences in extinction coefficients (E₁%₁ cm). Astaxanthin concentrations were calculated using E₁%₁ cm-values of 2100 for all-E-astaxanthin (Britton, 1995), and 1350 and 1750 for 13Z- and 9Z-astaxanthin, respectively.

**Sensory Analysis**

A trained panel of 10 assessors (10 women, aged 37–64 years) at Nofima (Ås, Norway), performed a sensory descriptive analysis according to “Generic Descriptive Analysis” as described by Lawless and Heymann (2010) and (ISO 13229, 2016). Sensory analysis – Methodology – General guidance for establishing a sensory profile (2016). The assessors had been tested, selected, and trained according to ISO standards (ISO 8586, 2012), and the sensory laboratory follows the practice of ISO standards (ISO 8589, 2007). Raw salmon was evaluated for odor (sour, marine, cucumber, algae, feed, fish, odor) and color [yellowish color (hue), intensity, whiteness]. Cooked salmon was evaluated for flavor (sour, bitter, marine, cucumber, feed, fish, rancid) and texture (hardness, juiciness, and cohesiveness). Salmon filets were thawed overnight at +0–1°C before sample preparation the following day. The muscle was baked in a combi-oven (Electrolux Air-o-steam, Model AOS061EANQ; +75°C and 50% steam/50% heat) until a core temperature of +58°C was reached. Samples were then cooled immediately to +0–1°C. Both raw and heated samples, coded at a three-digit number, were served at the same time in white plastic container with a lid at +18°C. All attributes were evaluated on unstructured line scales with labeled endpoints ranging from “no intensity” (1) to “high intensity” (9). Each assessor evaluated the samples at individual speed on a computer system for direct recording of data (EyeQuestion, Software Logic8 BV, Utrecht, Netherlands). The coded samples were evaluated in duplicate per dietary per-treatment and served in a balanced block experimental design over 2 days and 10 sessions. Tap water and unsalted crackers were available for palate cleansing.

**Skin Coloration**

Non-invasive skin color analyses were performed at the end of the cage trial, using a Minolta Chroma Meter (CR-400 Minolta,
Konica Minolta Sensing, Inc., Japan). The tristimulus CIE $L^*a^*b^*$ 1976 color space was used, where the $L^*$ variable represents lightness ($L^* = 0$ for black, $L^* = 100$ for white) and the $a^*$ and $b^*$ indicate color directions: $+a^*$ is the red direction, $-a^*$ is the green direction, $+b^*$ is the yellow direction, and $-b^*$ is the blue direction. Analyses were performed anterior and posterior to the dorsal fin, below and above the lateral line.

**Microarray Analyses**

Sample tissue (mid intestine and liver) RNA was extracted using PureLink RNA Mini kits according to the manufacturer’s protocol (Invitrogen). Concentration of total RNA (NanoDrop 1000 Spectrometer, Thermo Scientific, Waltham, MA, United States) and RNA integrity was measured (Agilent 2100 Bioanalyzer with RNA Nano kits, Agilent Technologies, Santa Clara, CA, United States). Samples with RNA integrity number (RIN) > 8 were accepted for analysis. Multiple gene expression profiling was performed with the Nofima’s Atlantic salmon 15k SIQ6 array produced by Agilent Technologies, Santa Clara, CA, United States. One-color hybridizations were performed. Analyses included 12 replicates from each study group, totally 48 microarrays were used [two tissues $\times$ 2 dietary groups (FO and ScB from start to slaughter) $\times$ 12 fish]. RNA amplification, labeling, and fragmentation were performed using the One-Color Low Input Quick Amp Labeling kit and Gene Expression Hybridization kit following the manufacturer’s instructions (Agilent Technologies, Santa Clara, CA, United States). The input of total RNA in each reaction was 100 ng. The samples were labeled with Cy3. Overnight hybridization (17 h, 65°C, and rotation speed of 10 rpm) was executed in an oven (Agilent Technologies, Santa Clara, CA, United States). The slides were washed with Gene Expression Wash Buffers 1 and 2 and scanned with Agilent scanner. Data were processed with Nofima’s pipeline STARS (Krasnov et al., 2011) as described in Pellizzari et al. (2013). Global normalization was performed by equalizing the mean intensities of all microarrays. Next, the individual values for each feature were divided to the mean value of all samples thus producing expression ratios (ER). The log2-ER were calculated and normalized with the locally weighted non-linear regression (Lowess). Differentially expressed genes (DEG) were selected by the following criteria: ER > 1.5-fold and $p < 0.05$ ($t$-test). Results were submitted to NCBI GEO (GSE140558).

**Histological Observations in the Intestine**

The mid and hind intestine from 12 fish from each one of two dietary groups (ScB:ScB and FO:FO) in the present trial was evaluated histologically. Samples of mid and hind intestine were fixed in 4% buffered formalin and stored at 4°C until processing. Paraplast-embedded samples were cut with a Leitz 1208 microtome (Ernst Leitz Wetzlar GmbH) (5 µm) and stained with standard hematoxylin–eosin (Merck KGaA). Stained slides were examined using a standard light microscope (Nikon Optiphot, Nikon). Images were captured by a Micropublisher 3.3 RTV camera and QCapture 2.9.13 software (QImaging). A total of 48 samples (from 24 fish) were processed blind, 12 samples per end diet. Further on, observations were realized in coded samples in order to characterize systematic differences.

**Calculations and Statistics**

Apparent digestibility coefficient (ADC) of dietary nutrients in the test diets was calculated from the following formula: $ADC = (d - d_{wt}) / d_{wt}$, where $d$ is the feces, $Y$ yttrium content, and $N$ nutrient content. Feed conversion ratio (FCR) is feed consumed/biomass increase; protein efficiency ratio (PER) is the fish weight gain/protein consumption; thermal growth coefficient (TGC) is (w1/3 − w1/3) $\times$ 1000/$\Sigma$(t $\times$ feeding days), where $\Sigma$(t $\times$ feeding days) is the sum of water temperatures (°C) for every feeding day in the experiment (Cho, 1992); condition factor (CF) is bw (g) $\times$ fish fork length-3 (cm) × 1000; dress out percentage (D%) is gutted fish weight/bw × 100; hepatosomatic index (HSI) is liver weight/bw × 100. Data were tested for normality using a Kolomogorov–Smirnov test and homogeneity of variance using Levene’s test. Biological and analytical data were subjected to analysis of variance (ANOVA) tests (univariate and repeated measures GLM) using IBM SPSS statistics 24. When differences among groups were identified, pairwise comparisons among means were made using Duncan’s post hoc test. Significant treatment effects were considered at a significance level of $P < 0.05$ and tendencies at $P < 0.1$. For sensory performance, ANOVA using a two-way mixed model with the assessor and interaction effects considered random and samples as a fixed effect was performed on the descriptive sensory data to identify the sensory attributes that discriminated between samples. Least significance differences were calculated by Tukey’s test ($P < 0.05$). A principal component analysis (PCA) analysis on the average of the sensory descriptive data was performed with mean centered data and no standardization. The statistical software used for the sensory analysis was EyeOpenR (Logic8 BV). For the multivariate data analysis, Unscrambler X Version 10.4.1 was used for the PCA.

**RESULTS**

**Experimental Diets**

The experimental diet’s fatty acid and proximate composition is presented in Table 1. The diets per trial segment were similar in all respects where intended and as expected FO diets had the lowest, FO:ScB intermediate, and ScB diet the highest DHA/EPA ratio.

**Fish Performance and Biometrics**

**Fish Group Performance**

During the tank salt water phase and using mean values of measurements for all fish in each tank as unit, fish growth rate was high and similar for all treatments (TGC of 3–3.8), and FCR was low (0.67–0.79 on feed dry matter basis) with no significant difference between the three dietary treatments (Table 2). Likewise, CF, D%, and HSI showed no significant differences between the dietary treatments during the tank trial phase. In the cage trial phase, body growth was significantly higher for the ScB compared to FO treatment. FCR was elevated compared to the tank phase and was similar in all treatments during the cage trial phase, ranging from 1.27–1.43 for the FO
and 1.32–1.41 for the ScB treatments, respectively. At 2.9–3.2 kg mean fish body weight, at the end of the cage trial, all ADC values measured were reduced in both end treatments, compared to earlier growth stages in this trial (in particular for lipids, then total energy and less for protein) or previously reported studies (Kousoulaki et al., 2015, 2016). In the smolt phase ADC of protein tended to be higher in the ScB compared to the FO diet whereas there was a gradual decrease in lipid ADC with increasing inclusion level of *S. limacium* biomass in the diets. No dietary effects were noticed in terms of ADC of energy in the smolt phase. At slaughter, protein ADC was similar for the FO and ScB treatments and the results on ADC of lipids were inversed compared to the smolt phase and previous studies (Kousoulaki et al., 2015, 2016), with the ScB rich diet having significantly higher lipid ADC compared to FO. ADC of energy also tended to be higher for the ScB feed.

The final mean body weights of the fish in the FO end treatment varied compared to that of the ScB (mean fish BW in FO cage 1: 2609 g, FO cage 2: 2691 g and FO cage 3: 3154 g; ScB cage 1: 3187 g, ScB cage 2: 3355 g, and ScB cage 3: 3353 g) thus most of the observed effects, though large, were not statistically significant when analyzed with one-way ANOVA and cage as unit.

### Individual Fish Performance

The observed effects were more pronounced when using the data from individually pit tagged fish analyzed for pre-diet and end diet treatment effects (Table 3). Repeated measured ANOVA was conducted to evaluate the effect of time and (pre)-feeding on fish body weight and TGC in the tank phase, and the effect of time, pre-feeding, and end-feeding on fish body weight and CF during the cage phase of the trial, using individual fish measurements and the results are summarized in Table 4. Equality of error variances was tested with Levene’s test and was found not significant in all cases (P > 0.05). The results of the ANOVA indicated a significant effect of time (P < 0.001) and tank (P < 0.001) on both BW and TGC and also of (pre)feed on TGC during the tank phase. Pairwise comparisons with Duncan’s *post hoc* test show that BW increased with time whereas TGC was lowest close to smoltification, highest just after, and decreased before transfer to the sea. TGC was significantly higher in ScB compared to the FO/ScB groups whereas in FO was not significantly different compared to the two other treatments during the tank trial phase. In the sea cage phase of the trial, ANOVA showed significant effect of cage (P < 0.001) and an indication of effect of pre-feeding (P = 0.054) and end-feeding (P = 0.062) on fish BW. The groups fed the FO diet in the cages had at slaughter significantly lower body weight compared to the ScB groups (P < 0.001), however, within these groups the fish that were fed previously with FO were the largest, followed by FO/ScB and ScB, but the effect of pre-feeding was not statistically significant (P > 0.1). Body weight was lowest in two of three FO cages compared to the 3rd one and all three ScB cages (P < 0.001). Fish CF decreased significantly from the end of the tank phase to slaughter (P = 0.000) and was affected significantly by pre-diet (P = 0.000) but not the cage or end diet (P > 0.1). Pairwise comparisons showed that CF in fish previously fed the FO feed was higher compared to those fed FO/ScB (P = 0.096) or ScB (P = 0.069), and the two latter were not different from each other (P > 0.1).

### ADC of Dietary Fatty Acids

In terms of ADC of individual fatty acids (Table 4), the dietary treatments differed significantly nearly in all both phases (smolt and slaughter). In general, at smolt phase the ADC of nearly all analyzed fatty acids was highest in the FO, intermediate in the FO/ScB, and lowest in the ScB treatments. At slaughter the FO dietary group had higher ADC of SFA and lower ADC of MUFA, n-3 and n-6 LC-PUFA compared to ScB.

### NQC Filet Lipid Level, Fatty Acid Composition, and Organ Scores

The NQC filet lipids generally mirrored the dietary composition and were also influenced by the apparent digestibility of the different dietary fatty acids (Table 5). Thus, the FO (end feed) fish had higher levels of total SFA and specifically 14:0, 16:0, and 18:0, total MUFA and specifically 16:1, 18:1, and 22:1, as well as 22:5 n-3 and 20:5n-3 (EPA) in their filet and lower levels of total n-6 PUFA fatty acids and specifically 18:2 n-6 and 20:2 n-6, total n-3 PUFA fatty acids, and specifically 18:3 n-3 (ALA, a-linolenic acid) and 22:6 n-3 (DHA), and total EPA + DHA compared to the ScB treatment. The respective levels in fish previously fed a mix of FO/ScB or FO and then ScB and *vice versa*, were in between. Liver score regarding coloration was significantly affected by the end diet. The used method for liver color scoring is developed to grade livers in terms of coloration most commonly related to lipid levels providing numbers corresponding to the different gradients of liver “paleness” where more “healthy” looking livers have a more dark red hue whereas fatty livers have a more yellowish – orange hue. Livers in ScB fish at slaughter had a dark green tone not occurring in FO fish. The authors have no previous experience regarding this observation. No differences were found in the intestinal fat score of the sample fish at the end of the sea cage trial.

### Liver Lipid Level and Fatty Acid Profile

Total liver lipid levels were higher in the ScB treatment at the end of the sea cage phase (Table 6). Nevertheless, within the ScB fish, one group was distinct, namely that previously fed the FO diet, having higher levels of liver lipids, the lowest levels of SFA, sum n-3 and EPA + DHA, and the highest levels of MUFA and N-6 compared to all other pre-treatments, both from the FO and ScB end-feed groups (not in all cases significantly compared to the two remaining pre-treatment groups within ScB). Pre-diet alone had no significant effect on the total lipids or liver fatty acid profile. End diet, as in the case of the filets, affected significantly nearly all analyzed liver fatty acids. Livers had higher relative levels of total SFA and N-3 PUFA and lower levels of MUFA and N-6 PUFA compared to the filet. Again, as in the filet, FO fish had significantly higher levels of SFA and N-6 PUFA in the liver compared to the ScB end diet treatment. However, unlike in the filet the levels of MUFA, N-3 PUFA, and EPA + DHA...
### TABLE 2 | Growth, feeding performance, and biometrics of all Atlantic salmon individuals participating each one of the successive trials, fed diets with different levels of fish oil (FO), and *Schizochytrium limacinum* biomass (ScB).

| Diet name | Fresh water phase | Salt water tank smolt phase | Salt water tank post-smolt phase | Sea water cage phase |
|-----------|------------------|-----------------------------|---------------------------------|---------------------|
|           | FO1 | FO/ScB1 | ScB1 | FO2 | FO/ScB2 | ScB2 | FO3 | FO/ScB3 | ScB3 | FO4 | ScB4 |
| Start fish number/tank | 127 | 124 | 126 | 110 | 110 | 110 | 127 | 124 | 126 | 110 | 110 | 110 |
| Mortality (%) | 0.00 | 0.00 | 0.00 | 0.00 | ns | 0.30 | 0.10 | ns | 0.30 | 0.10 | ns | 0.30 | 0.10 |
| Initial weight (g) | 18.3 | 18.2 | 18.1 | 64.7 | 65.5 | 63.4 | 0.48 | ns | 0.50 | 0.50 | 0.30 | ns | 11.8 |
| Final weight (g) | 64.4 | 65.3 | 63.1 | 270 | 277 | 273 | 2.59 | ns | 889 | 11.28 | ns | 2818 |
| Total feed intake (g DM/fish) | 142 | 139 | 138 | 1.46 | ns | 489 | 446 | 475 | 10.08 | ns | 2784 | 196.0 |
| Feed intake (%bw/day) | 1.52 | 1.43 | 1.45 | 0.02 | ns | 1.15 | 1.04 | 1.08 | 0.02 | ns | 1.51 | 1.42 |
| FCR (DM) | 0.69 | 0.66 | 0.66 | 0.01 | ns | 0.82 | 0.79 | 0.77 | 0.15 | ns | 1.51 | 1.42 |
| TGC (*1000) | 0.90 | 0.92 | 0.89 | 3.78 | 3.84 | 3.86 | 0.02 | ns | 3.07 | 2.92 | 3.16 | 0.16 |
| PER | 3.08 | 3.16 | 3.15 | 0.03 | ns | 2.50 | 2.60 | 2.67 | 0.03 | ns | 0.11 | 1.37 |
| CF | 1.11 | 1.11 | 1.09 | 0.01 | ns | 1.37 | 1.36 | 1.40 | 0.01 | ns | 1.51 | 1.24 |
| D% | 88.5 | 89.0 | 88.8 | 86.7 | 86.1 | 86.2 | 0.11 | ns | 87.6 | 87.8 | 88.3 | 0.21 |
| HSI | 0.85 | 0.84 | 0.82 | 1.29 | 1.24 | 1.33 | 0.03 | ns | 1.32 | 1.18 | 1.28 | 0.03 |
| ADC\(_{protein}\) (%) | 88.3\(^A\) | 88.7\(^{aB}\) | 89.5\(^B\) | 0.21 | 0.055 | 87.3 | 87.2 | 0.30 | ns |
| ADC\(_{protein}\) (%) | 94.6\(^c\) | 92.6\(^p\) | 90.8\(^a\) | 0.56 | 0.000 | 84.5\(^p\) | 87.1\(^d\) | 0.67 | 0.019 |
| ADC\(_{energy}\) (%) | 77.2 | 77.3 | 78.3 | 0.31 | ns | 75.7\(^a\) | 78.2\(^{B}\) | 0.71 | 0.055 |

Values are mean (n = 3 tanks or cages per treatment) and standard error of the mean (SEM). At the end of the fresh water and salt water smolt phase, a number of fish were sacrificed each time for the performance of biological studies and chemical analyses and the rest continued further in the next phase of the trial. *Values of the same line and phase, not sharing common small superscript letter are significantly different (P ≤ 0.05) or not sharing common capital superscript letter tend to be different (P ≤ 0.1) as determined by one-way ANOVA followed by Duncan's post hoc test; ns, non-significant.
TABLE 3 | Individually pit-tagged fish body weight (BW), TGC, and condition factor (CF) evolution from trial start to slaughter.

| Pre-diet | FO   | FO/ScB | ScB   | FO   | FO/ScB | ScB   | SEM | Time | Time * Tank | Time * Cage | Time * pre-feed | Time * end-feed |
|----------|------|--------|-------|------|--------|-------|-----|------|------------|------------|----------------|----------------|
| End diet | FO   | FO/ScB | ScB   | FO   | FO/ScB | ScB   |     |      |            |            |                |                |
| BW1      | 18.3 | 18.1   | 18.3  | 18.8 | 18.0   | 17.7  | 0.12| 0.000|             |            |                |                |
| BW2      | 66.6 | 66.1   | 63.2  | 66.4 | 67.0   | 64.7  | 0.49|      |             |            |                |                |
| BW3      | 277  | 281    | 274   | 277  | 283    | 277   | 1.36|      |             |            |                |                |
| BW4      | 871  | 852    | 881   | 889  | 832    | 892   | 6.78|      |             |            |                |                |
| BW5      | 2889 | 2940   | 2609  | 3309 | 3291   | 3278  | 75.1|      |             |            |                |                |
| TGC1     | 0.92 | 0.94   | 0.89  | 0.91 | 0.95   | 0.93  | 0.01|      |             |            |                |                |
| TGC2     | 3.85 | 3.87   | 3.88  | 3.81 | 3.86   | 3.86  | 0.01|      |             |            |                |                |
| TGC3     | 3.02 | 2.92   | 3.07  | 2.98 | 2.83   | 3.09  | 0.03|      |             |            |                |                |
| CF4      | 1.49 | 1.40   | 1.41  | 1.47 | 1.39   | 1.41  | 0.01|      |             |            |                |                |
| CF5      | 1.22 | 1.21   | 1.16  | 1.23 | 1.22   | 1.23  | 0.01|      |             |            |                |                |

Values are means (n = 3 tanks/cages per feeding treatment) and standard error of the mean (SEM). Wilks’ Lambda multivariate test significance (P) values are following repeated measures ANOVA for BW (BW1–4) and TGC (TGC1–3) during the tank phase (T: n = 9) and BW (BW4–5), TGC (TGC3–4), and CF (CF3–4) from the end of the tank phase to slaughter in the cages (C: n = 6); Pairwise comparisons by Duncan’s post hoc test (P < 0.05). *Coded values in the same cell not sharing common superscript letter are significantly different as determined by repeated measures ANOVA and separated by pairwise comparisons by Duncan’s post hoc test (P < 0.05); ns, non-significant.
TABLE 4 | Dietary fatty acid apparent digestibility coefficient (ADC) (%) in Atlantic salmon fed diets with fish oil (FO) or ScB as supplementary marine omega 3 fatty acid sources at slaughter (fish mean body weight 2.8–3.2 kg).

|                | Smolt phase (tank trial) | Slaughter phase (cage trial) |
|----------------|--------------------------|-----------------------------|
|                | FO | FO/ScB | ScB | SEM | One-way ANOVA (P) | FO | ScB | SEM | One-way ANOVA (P) |
| 14:0           | 97.5<sup>c</sup> | 88.2<sup>d</sup> | 69.4<sup>a</sup> | 4.12 | 0.000 | 84.5 | 58.3 | 5.87 | 0.000 |
| 16:0           | 96.1<sup>c</sup> | 86.1<sup>b</sup> | 76.0<sup>a</sup> | 2.90 | 0.000 | 79.2 | 59.7 | 4.40 | 0.000 |
| 18:0           | 96.7<sup>c</sup> | 95.1<sup>tu</sup> | 92.8<sup>b</sup> | 0.58 | 0.000 | 83.7 | 78.8 | 1.15 | 0.004 |
| 20:0           | 97.3<sup>b</sup> | 94.4<sup>a</sup> | 94.7<sup>a</sup> | 0.51 | 0.005 | 84.5 | 77.2 | 1.70 | 0.003 |
| 22:0           | 94.6<sup>c</sup> | 92.6<sup>b</sup> | 90.8<sup>b</sup> | 0.56 | 0.000 | 84.5 | 78.7 | 1.52 | 0.009 |
| 16:1 n-7       | 99.0<sup>c</sup> | 98.8<sup>b</sup> | 96.9<sup>a</sup> | 0.33 | 0.000 | 88.7 | 94.3 | 1.42 | 0.023 |
| 18:1 (n-9) + (n-7) + (n-5) | 99.0 | 98.8 | 99.1 | 0.13 | ns | 84.3 | 92.5 | 1.90 | 0.002 |
| 20:1 (n-9) + (n-7) | 98.6<sup>c</sup> | 97.8<sup>b</sup> | 97.9<sup>a</sup> | 0.11 | 0.000 | 81.0 | 89.3 | 1.90 | 0.002 |
| 22:1 (n-11) + (n-9) + (n-7) | 98.2<sup>c</sup> | 97.6<sup>b</sup> | 96.9<sup>a</sup> | 0.33 | 0.000 | 75.0 | 82.8 | 2.01 | 0.023 |
| 24:1 n-9       | 94.6 | 96.3 | 93.9 | 0.58 | ns | 53.5 | 74.3 | 4.74 | 0.000 |
| 18:2 n-6       | 98.4 | 98.4 | 98.3 | 0.05 | ns | 89.9 | 95.4 | 1.25 | 0.001 |
| 20:2 n-6       | 98.2<sup>c</sup> | 97.5<sup>b</sup> | 95.4<sup>a</sup> | 0.42 | 0.000 | 84.5 | 84.5 | 0.64 | ns |
| 18:3 n-3       | 99.3<sup>c</sup> | 99.3<sup>b</sup> | 99.3<sup>a</sup> | 0.03 | 0.009 | 92.2 | 96.7 | 1.05 | 0.001 |
| 18:4 n-3       | 99.4 | 99.0 | 96.9 | 0.99 | ns | 94.8 | 100.0 | 1.16 | 0.000 |
| 20:5 n-3 (EPA) | 99.1<sup>c</sup> | 98.3<sup>ab</sup> | 96.9<sup>a</sup> | 0.39 | 0.045 | 92.7 | 87.1 | 1.25 | 0.000 |
| 22:6 n-3 (DHA) | 98.0<sup>c</sup> | 97.7<sup>ab</sup> | 97.5<sup>a</sup> | 0.08 | 0.014 | 80.5 | 62.7 | 3.99 | 0.000 |
| Sum SFA        | 96.5 | 87.2<sup>d</sup> | 77.5<sup>a</sup> | 2.71 | 0.000 | 84.9 | 94.0 | 2.44 | 0.039 |
| Sum MUFA       | 98.9<sup>c</sup> | 98.7<sup>ab</sup> | 98.5<sup>a</sup> | 0.05 | 0.001 | 84.1 | 92.3 | 1.89 | 0.002 |
| Sum PUFA (n-6) | 98.4<sup>b</sup> | 98.3<sup>ab</sup> | 98.1<sup>a</sup> | 0.07 | 0.093 | 89.8 | 95.3 | 1.25 | 0.001 |
| Sum PUFA (n-3) | 99.3<sup>c</sup> | 99.9<sup>b</sup> | 98.8<sup>a</sup> | 0.07 | 0.000 | 91.7 | 95.8 | 0.95 | 0.002 |
| Total PUFA     | 98.9<sup>c</sup> | 98.7<sup>ab</sup> | 98.5<sup>a</sup> | 0.06 | 0.004 | 91.0 | 96.6 | 1.06 | 0.001 |
| Total fatty acids total | 98.4<sup>c</sup> | 96.4<sup>b</sup> | 94.1<sup>a</sup> | 0.63 | 0.000 | 86.2 | 88.5 | 0.60 | 0.024 |

Values are means (n = 3) and standard error of the mean (SEM). Values in the same row with different superscript lowercase letter are significantly different (<0.05), whereas values in the same row with different superscript uppercase letter tend to be different (<0.1).

were significantly lower in the liver of the ScB compared to the FO end diet fish.

Whole Body Lipid Level and Fatty Acid Profile

Whole body lipids and fatty acid composition did not differ among the two dietary treatments during the sea cage period, and all effects of pre-dietary treatments which were developed during the tank-phase of the experiment were not present at the end of the cage trial phase (Table 7).

Filet Pigmentation, Color, and Technical Quality

The fish in the ScB treatment had in the end of the cage trial significantly higher levels of astaxanthin but not idoxanthin compared to FO (Table 8). Pre-feeding had no effect on the final pigmentation results. ScB had also higher visual color scores in the filets compared to the FO groups. The filet color evaluated by SalmoFan in the three filet segments from head to tail were P1 = 27.85 ± 0.46, P2 = 27.00 ± 0.48, and P3 = 28.63 ± 0.55 for FO and P1 = 28.56 ± 0.11, P2 = 27.30 ± 0.45, and P3 = 29.07 ± 0.45 for ScB. Paired sample t-test showed significantly higher filet color both in the anterior part of the filet and the posterior for the fish fed the ScB compared to the FO supplemented feeds (P = 0.001; n = 3) in the end of the cage trial, but the differences were nevertheless small. Colorimetric analyses of the skin showed more greenish color (lower a<sup>+</sup> values) of the ScB group and a tendency to more bluish color (lower b<sup>+</sup> values) color compared with FO group. Moreover, pre-dietary treatment affected skin darkness. Filets of salmon fed ScB before harvesting had significantly less segments (myomera) with melanin discoloration compared to FO, but no effect was observed on filet gaping or firmness. However, filet firmness was affected by pre-diet as well as interaction of pre- and end diet.

Sensory Analysis

Parr-Post Smolt – 0.8 kg Mean Body Weight Atlantic Salmon

The results from the sensory evaluation of A. salmon post smolt filets of 0.8 kg approximate body weight at the end of tank phase and beginning of cage trial phase showed that there were some significant differences in coloration (hue) between the different feeding treatments. Fish that were fed diets supplemented with ScB had more red filets (P < 0.05) stronger coloration (hue) compared to FO group. Moreover, pre-dietary treatment affected skin darkness. Filets of salmon fed ScB before harvesting had significantly less segments (myomera) with melanin discoloration compared to FO, but no effect was observed on filet gaping or firmness. However, filet firmness was affected by pre-diet as well as interaction of pre- and end diet.

Slaughter – 4.15 kg Mean Body Weight Atlantic Salmon

Similar dietary effects as early in the trial were also observed at final sampling where the ScB feeding regime gave filets with
TABLE 5 | Fatty acid composition in filets of A. salmon fed life-long diets supplemented with FO, ScB, or a mix of the two until 0.8 kg body weight and either FO or ScB to slaughter.

| Lipid %          | FO     | FO/ScB | ScB  | SEM | P value* |
|------------------|--------|--------|------|-----|----------|
| C 14:0           | 11.23  | 10.52  | 11.53| 0.26| ns       |
| C 16:0           | 13.23  | 14.40  | 13.30| 0.20| 0.000    |
| C 18:0           | 3.04   | 3.37   | 3.16 | 0.06| 0.017    |
| C 22:0           | 0.05   | 0.05   | 0.05 | 0.05| ns       |
| SUM SFA          | 17.88  | 19.01  | 17.34| 0.26| 0.001    |
| C 16:1 n-7       | 1.93   | 1.43   | 1.00 | 0.14| 0.000    |
| C 18:1 n-9       | 33.64  | 33.70  | 34.04| 0.09| ns       |
| C 20:1 n-11      | 0.96   | 0.80   | 0.70 | 0.04| 0.002    |
| C 20:1 n-9       | 2.48   | 2.49   | 2.20 | 0.05| 0.012    |
| C 22:1 n-7       | 1.20   | 1.17   | 1.10 | 0.03| ns       |
| SUM MUFA         | 40.22  | 39.60  | 39.04| 0.18| 0.002    |
| C 18:2 n-6       | 14.35  | 14.73  | 15.39| 0.16| 0.003    |
| C 18:3 n-6       | 0.14   | 0.13   | 0.13 | 0.01| ns       |
| C 20:2 n-6       | 1.28   | 1.42   | 1.46 | 0.03| 0.036    |
| C 20:3 n-6       | 0.48   | 0.49   | 0.38 | 0.02| 0.003    |
| C 20:4 n-6       | 0.21   | 0.18   | 0.12 | 0.01| 0.000    |
| SUM N-6 PUFA     | 16.45  | 16.93  | 17.47| 0.15| 0.000    |
| C 18:3 n-3       | 13.86  | 13.57  | 15.30| 0.28| 0.001    |
| C 20:3 n-3       | 1.30   | 1.41   | 1.53 | 0.04| 0.020    |
| C 20:5 n-3       | 2.77   | 1.09   | 1.09 | 0.24| 0.000    |
| C 22:5 n-3       | 1.01   | 0.46   | 0.46 | 0.08| 0.000    |
| C 22:6 n-3       | 3.38   | 5.46   | 5.49 | 0.31| 0.000    |
| SUM N-3 PUFA     | 22.32  | 22.11  | 23.87| 0.29| 0.002    |
| EPA + DHA        | 6.15   | 6.49   | 6.58 | 0.09| ns       |
| Liver score      | 2.76   | 3.00   | 2.92 | 2.08| 0.008    |
| Intestinal-fat score | 2.58  | 2.83   | 2.75 | 2.50| 0.081    |

One-way ANOVA was performed to compare data at the end of the tank phase using mean values from sample fish per tank as units (n = 3). Univariate GLM ANOVA with diet and pre-diet as fixed factors was used to compare data at the end of the cage trial using mean values from analyses performed in fish of same pre-diet history per cage as units (n = 3). Values are % in identified fatty acids and standard error of the mean (SEM). *Values with different superscript letter, separately for cage trial start and end, are significantly different separated by Duncan’s post hoc test (P < 0.05); ns, non-significant.
TABLE 6 | Fatty acid composition in liver of A. salmon fed life-long diets supplemented with FO, ScB, or a mix of the two until 0.8 kg body weight and either FO or ScB to slaughter.

| Pre-diet: end diet | Cage trial end (pre-diet: end diet) | ANOVA-P value* |
|--------------------|------------------------------------|-----------------|
| FO:FO              | FO/ScB:FO                          | FO/ScB:ScB      | ScB:ScB | SEM |
| Foch %             | 4.30**                            | 3.89a           | 4.43a   | 6.98b  | 4.82a  | 4.94a  | 0.312 |
| 14:0               | 0.87                              | 0.68            | 0.80    | 0.47   | 0.36   | 0.33   | 0.070 |
| 16:0               | 13.24**                           | 14.05b          | 12.99b  | 9.50a  | 11.82ab | 11.41ab | 0.450 |
| 18:0               | 4.65**                            | 4.61b           | 4.33b   | 2.13a  | 3.27ab  | 3.31ab  | 0.262 |
| 22:0               | 0.09                              | 0.40            | 0.20    | 0.16   | 0.21   | 0.12   | 0.048 |
| SUM SFA            | 18.85**                           | 19.74d          | 18.31cd | 12.25a | 15.66bc | 15.17ab | 0.708 |
| 16:1 n-7           | 0.79p                             | 0.72bc          | 0.81b   | 0.50a  | 0.44a   | 0.49a   | 0.040 |
| 16:1 n-5           | 0.46                              | 0.47            | 0.25    | 0.30   | 0.41   | 0.36   | 0.076 |
| 18:1 n-11          | 0.19bc                            | 0.23c           | 0.20bc  | 0.07a  | 0.13ab  | 0.13ab  | 0.016 |
| 18:1 n-9           | 22.31ab                           | 20.96a          | 23.20ab | 30.63c | 25.69ab | 28.51bc | 0.932 |
| 18:1 n-7           | 1.80                              | 1.69            | 1.79    | 1.91   | 1.77   | 1.82   | 0.028 |
| 20:1 n-9           | 1.56                              | 1.30            | 1.50    | 2.22   | 1.85   | 2.03   | 0.113 |
| 22:1 n-7           | 1.90                              | 1.98            | 1.79    | 1.56   | 1.83   | 1.86   | 0.055 |
| 22:1 n-11          | 0.17                              | 0.00            | 0.13    | 0.11   | 0.09   | 0.06   | 0.021 |
| SUM MUFA           | 29.18ab                           | 27.34a          | 29.68ab | 37.30c | 32.21abc| 33.26bc | 0.982 |
| 18:2 n-6           | 9.77a                             | 9.91a           | 10.52ab | 13.18a | 11.62bc | 11.42ab | 0.310 |
| 18:3 n-6           | 0.07                              | 0.10            | 0.08    | 0.08   | 0.03   | 0.07   | 0.014 |
| 20:2 n-6           | 1.26                              | 1.17            | 1.30    | 1.52   | 1.47   | 1.49   | 0.053 |
| 20:3 n-6           | 0.88ab                            | 0.84cd          | 0.86d   | 0.49a  | 0.63ab  | 0.68bc  | 0.099 |
| 20:4 n-6           | 1.47b                             | 1.49p           | 1.43p   | 0.60a  | 0.72ab  | 0.75a   | 0.097 |
| SUM N-6 PUFA       | 13.44a                            | 13.51a          | 14.19b  | 15.87b | 14.47a  | 14.40a  | 0.240 |
| 18:3 n-3           | 8.01a                             | 8.04a           | 8.46a   | 11.56a | 10.13b  | 10.22b  | 0.352 |
| 20:4 n-3           | 0.16                              | 0.28            | 0.27    | 0.18   | 0.17   | 0.16   | 0.024 |
| 20:3 n-3           | 1.07                              | 1.05            | 1.13    | 1.83   | 1.60   | 1.01   | 0.113 |
| 20:5 n-3           | 7.95c                             | 8.10f           | 7.59f   | 1.26a  | 2.58b   | 2.65p   | 0.713 |
| 22:5 n-3           | 2.24p                             | 2.31b           | 2.17b   | 0.44a  | 0.62a   | 0.57a   | 0.208 |
| 22:6 n-3           | 11.93ab                           | 12.40ab         | 11.60a  | 11.38a | 15.88c  | 15.13bc | 0.555 |
| SUM N-3 PUFA       | 31.36b                            | 32.19b          | 31.21b  | 26.66a | 31.00b  | 29.74ab | 0.578 |
| EPA + DHA          | 19.88b                            | 20.50b          | 19.19b  | 12.64a | 18.47b  | 17.78b  | 0.762 |

Univariate GLM ANOVA with diet and pre-diet as fixed factors was used to compare data at the end of the cage trial using mean values from analyses performed in fish of same pre-diet history per cage as units (n = 3). Values are % in identified fatty acids and standard error of the mean (SEM). *Values with different superscript letters are significantly different separated by Duncan’s post hoc test (P < 0.05); ns, non-significant.

redder coloration compared to FO. Moreover, FO gave samples which were softer and juicier compared to ScB (Figure 2).

There were no significant effects identified on the sensory quality of the fillets from pre-feeding within fish fed the FO and ScB during the cage phase.

Effect of Pre-feeding on Cage Level – 4.15 kg Mean Body Weight Atlantic Salmon

Principal component analysis was used to identify similarities and differences between the sensory attributes and the 18 different treatments (six cages × three treatments per cage) (Figure 3). The first and second principal component (PCs) explained 42 and 17% of the observed variation, respectively. The biplot (scores and loadings) showed what we have seen from other results, namely that pre-feeding has had minor effects on the end sensory quality of the fillets. Pre-diet samples are spread all over in the figure, whereas samples from ScB end diet (marked as bold) are positioned above or just under the middle line, correlating on PC2 with color intensity, hue, and hardness, and samples from FO end diet were positioned from the middle to the lower side of PC2 correlating with juiciness and whiteness.

Microarray Analyses of Liver and Mid Intestine

Microarray (transcriptome) analysis was realized in liver and mid intestine tissues from fish in the FO:FO and ScB:ScB dietary treatments. Only a few genes were differentially expressed in the liver of the two treatments, but some of those are worth mentioning. For instance, significant upregulation was found of the hsp (heat shock protein), gpx (glutathione peroxidase), and gadd45 (growth arrest and DNA-damage-inducible protein) genes in the ScB compared to the FO group. More immune system-related genes were downregulated in the liver and mid intestine in the ScB compared to the FO groups (Table 9). There were just a few differences in gene
| Treatment          | Cage trial start | ANOVA-P value | Cage trial end (pre-diet: end diet) | ANOVA-P value |
|--------------------|------------------|---------------|------------------------------------|---------------|
| Fat (%)            | FO               | FO/ScB        | ScB                                | SEM           |
| 14:0               | 1.71            | 1.27          | 0.88                               | 0.12          |
| 16:0               | 13.63           | 14.77         | 14.62                              | 0.21          |
| 18:0               | 3.52            | 3.34          | 3.22                               | 0.06          |
| 20:0               | 0.21            | 0.21          | 0.28                               | 0.02          |
| 22:0               | 0.14            | 0.14          | 0.25                               | 0.03          |
| Sum SFA            | 19.22           | 19.72         | 18.25                              | 0.25          |
| 16:1 n-7           | 1.60            | 1.12          | 0.67                               | 0.13          |
| 18:1 n-9 + n-7 + n-5 | 35.50       | 35.94         | 35.48                              | 0.11          |
| 20:1 n-9 + n-7     | 2.84            | 2.74          | 2.48                               | 0.06          |
| 22:1 n-11 + n-9 + n-7 | 1.46        | 1.23          | 0.85                               | 0.09          |
| 18:2 n-6           | 14.57           | 14.84         | 15.56                              | 0.17          |
| 18:3 n-6           | 0.21            | 0.21          | 0.11                               | 0.02          |
| 18:4 n-6           | 1.31            | 1.44          | 1.66                               | 0.06          |
| 20:3 n-6           | 0.46            | 0.46          | 0.39                               | 0.02          |
| 20:4 n-6           | 0.21            | 0.21          | 0.11                               | 0.02          |
| 22:4 n-6           | 0.04            | 0.00          | 0.00                               | 0.01          |
| Sum MUFA           | 41.61           | 41.24         | 39.69                              | 0.30          |
| 18:2 n-6           | 14.57           | 14.84         | 15.56                              | 0.17          |
| 18:3 n-6           | 0.21            | 0.21          | 0.11                               | 0.02          |
| 18:4 n-6           | 1.31            | 1.44          | 1.66                               | 0.06          |
| 20:3 n-6           | 0.46            | 0.46          | 0.39                               | 0.02          |
| 20:4 n-6           | 0.21            | 0.21          | 0.11                               | 0.02          |
| 22:4 n-6           | 0.04            | 0.00          | 0.00                               | 0.01          |
| Sum N-6 PUFA       | 16.81           | 17.16         | 17.65                              | 0.17          |
| 18:3 n-3           | 13.38           | 13.05         | 15.07                              | 0.33          |
| 18:4 n-3           | 0.85            | 0.81          | 0.53                               | 0.05          |
| 20:3 n-3           | 1.24            | 1.27          | 1.70                               | 0.08          |
| 20:4 n-3           | 1.10            | 1.09          | 1.02                               | 0.03          |
| 20:5 n-3 (EPA)     | 1.74            | 1.20          | 0.64                               | 0.16          |
| 21:5 n-3           | 0.07            | 0.00          | 0.00                               | 0.02          |
| 22:5 n-3           | 0.85            | 0.53          | 0.25                               | 0.09          |
| 22:6 n-3 (DHAA)    | 2.95            | 3.83          | 5.02                               | 0.30          |
| Sum N-3 PUFA       | 22.14           | 21.77         | 24.23                              | 0.40          |
| Sum total-PUFA FAs | 39.16           | 39.03         | 42.06                              | 0.52          |

One-way ANOVA was performed to compare data at the end of the tank phase using mean values from sample fish per tank as units (n = 3). Univariate GLM ANOVA with diet and pre-diet as fixed factors was used to compare data at the end of the cage trial using mean values from analyses performed in fish of same pre-diet history per cage as units (n = 3). Values are % in identified fatty acids and standard error of the mean (SEM). *Values with different superscript letter, separately for Cage trial start and end, are significantly different separated by Duncan’s post hoc test (P < 0.05); ns, non-significant.
expression between the two dietary groups in mid intestine (Table 10). It is worth mentioning that in the mid intestine of fish from the ScB group there was upregulation of several genes involved in xenobiotic metabolism, whereas there was also seen upregulation of a zinc transporter. Downregulation of some few more genes related to inflammatory response in the intestine, such as chemokine/cytokines, was also observed in the ScB group.

Histological Evaluation of Intestinal Tissues

Hind Intestine

No diet-related differences were found in the histology of hind intestine between fish from FO:FO and ScB:ScB groups at the end of the trial. However, the hind guts of all 24 fish studied were judged to be abnormal (Figure 4) compared to normal hind gut histology in A. salmon (e.g., as in Knudsen et al., 2007). The intestinal folds were hyperplastic, abnormally tall, and large with extensively developed branches, which locally coalesced into networks. Side-branches were observed also in the simple folds, which should normally have none. The enterocytes lacked the vacuolization which is normal for mucosa in this region. In the center of the folds, the normally delicate strands of connective tissue were replaced by thick layers of connective tissue, and there was a prominent infiltration of inflammatory cells in connective tissue, both within and below the intestinal folds. These deviations were consistent throughout the material, although some variations were present, in particular in the degree of elongation and branching of folds as well as in cell infiltration.

Mid Intestine

The mid intestine was found normal in all 24 fish observed (Figure 5). Mucosal folds were of regular heights, and the enterocytes appeared normal and well developed. Some variation was observed, e.g., in the number of goblet cells, but all were considered to be within normal range. In particular, there was no sign of any difference in tissue morphology between the two dietary groups.

DISCUSSION

The main findings of this study are related to the: (a) significantly positive effect on fish growth and ADC by the ScB diet during the sea phase of the trial compared to the FO treatment, (b) differential dietary physiological effects on body lipid fatty acid composition, liver composition, and microarray and intestine microarray, and (c) positive ScB effect on Atlantic salmon filet coloration.

Atlantic Salmon Performance Replacing Dietary Fish Oil by Schizochytrium limacinum Biomass

The experimental design included replacement of supplemental dietary FO with S. limacinum biomass to achieve equal levels of total EPA + DHA in the diet. Due to the high DHA and very low level of EPA in S. limacinum biomass, the experimental diets had very different DHA/EPA ratio (0.69 in FO and 14.73 in the ScB diet – cage phase). Further formulation adjustments included balancing: (a) total dietary SFA levels, n-3/n-6 ratio, and digestible energy, by adjustments in the inclusion level of different plant oils, and (b) macronutrients by modifications in the inclusion levels of dietary binders. We chose a total dietary level of EPA + DHA as low as 1.25% in the diet during the sea cage phase, which was recently shown to be too low for farming Atlantic salmon robust enough to deal with very stressful conditions (Bou et al., 2017a). The experimental groups revealed growth performance differences only during the last phase of the study, namely during the sea cage phase, when ScB fish reached significantly higher final body weight compared to the FO fish. DHA, which ScB diet was richer of, compared to FO, is reported to be more essential than EPA in some fish species, in particular to be more essential than EPA in some fish species, in particular at high growth larval stages (Watanabe et al., 1989; Takeuchi et al., 1990; Copeman et al., 2002). Moreover, improved growth and non-specific immunity immune-related gene expression and disease resistance following natural infestation of parasites was reported in juvenile large yellow croaker (Larmichthys crocea) fed diets with moderately higher DHA/EPA levels (Zuo et al., 2012). In our study the best performing diet (ScB) had much higher DHA/EPA ratio and we did not see any significant differences in total mortalities or fish robustness (histology and microarray data). Improved growth was observed also in farmed Tilapia
fed diets where FO was replaced by *S. limacinum* biomass, however, unlike in our present study, dietary EPA + DHA levels were not balanced and increased with increasing levels of *S. limacinum* supplementation (Sarker et al., 2016). Other than salmon lice infestation and treatment-related mortalities, none of the experimental populations suffered from any other disease during the experimental period, and fish survival was identical in both FO and ScB groups. Thus, improved growth in the ScB treatment appears to be related to metabolic and behavioral effects leading to higher feed intake rates. In ScB diets we used whole cell *S. limacinum* biomass, which in contrast with extracted oils and plant concentrates, is expected to contain a great number of functional components. Although diets were balanced to cover the known fish’s basic needs, additional contributions of nucleotides, minerals, phospholipids, vitamins, and cell wall components from *S. limacinum* biomass may have contributed to improved intestine function during the most stressful growing period for the fish, which included several handling incidences for the removal of lice. Handling of fish may result in periods of low dissolved oxygen (DO) levels which may reduce the intestinal barrier function of Atlantic salmon, as for instance measured by reduced electrical resistance and increased paracellular permeability in the distal region of the fish’s intestine in the study by Sund et al. (2010). The hypothesis of improved intestine function in the ScB groups is also supported by the observed increased ADC of dietary nutrients in this treatment, as also found in mammals fed other unicellular organism based raw materials such as yeast culture in the diet for pigs (Shen et al., 2009). Such unicellular biomass components are commonly used as dietary supplements for farmed animals at rather low inclusion levels (pro- and prebiotics) and expected to improve intestine health (Anderson et al., 1999) both in terms of morphology (van der Peet-Schwering et al., 2007) and microflora composition (van Heugten et al., 2003). Accordingly, Lyons et al. (2016) found body growth improvement and greater level of microbial diversity in the microbial communities of rainbow trout (*Oncorhynchus mykiss*) fed low levels of *S. limacinum* in the diet compared to a control diet without, with significantly elevated bacterial taxa belonging to the genera *Streptococcus*, *Leuconostoc*, *Lactobacillus*, *Lactococcus*, and *Weissella*. This is also in accordance with our previous study where we saw growth improvement in Atlantic salmon post smolt by lower than in the present study dietary supplementation of *S. limacinum* biomass combined with yeast extract (1% in the diet) (Kousoulaki et al., 2015). However, little experience and evidences exist on intestine microbiome effects of higher life-long dietary inclusion levels of *S. limacinum* biomass, comparable to those used in this study. High inclusion levels of *S. limacinum* biomass combined with yeast extract induced a strong promotion of goblet cell, iNOS, and F-actin production in the intestinal epithelium of Atlantic salmon, compared to the FO control diet, suggesting immune system reaction without...
any histological evidence of enteritis or any other negative physiological effects (Kousoulaki et al., 2015).

Assimilation of Fatty Acids in Whole Fish and Filet

Atlantic salmon filet lipid fatty acid composition mostly reflected the composition of the experimental diets. Nevertheless, though the diets were balanced for total SFA and EPA + DHA, at slaughter FO fish had higher levels of SFA and lower levels EPA + DHA compared to the ScB fish. As recently identified, _S. limacinum_ biomass contains high levels tripalmitin (16:0/16:0/16:0) with high melting point (Bogevik et al., 2018) and low digestibility, explaining the differences in the whole-body SFA levels and retention efficiency of the dietary SFA. Early in the trial both dietary DHA and EPA were slightly more digestible in the FO diet, and at slaughter ADC of EPA but not that of DHA had higher ADC compared to that of the ScB diet. In terms of total retention efficiency FO-diet fed fish had much lower retention efficiency of EPA and higher DHA compared to ScB-fed fish. The FO diet had a DHA/EPA ratio of 0.69 against 14.73 in the ScB diet which would stimulate elongation and desaturation of EPA to DHA to ensure adequate amounts of DHA in the membranes of vital tissues of the fish (Glencross et al., 2014; Kuah et al., 2015).

During all the trial, ScB fish had higher levels of EPA + DHA in their filets compared to FO fish. This may be explained by breakdown of the more available EPA in the FO diet and preservation of the more plentiful DHA in the ScB fish (Tocher, 2003). This effect may have been more evident during the stressful period in sea cages where EPA with anti-inflammatory action is consumed whereas DHA with more structural function in cell membranes is retained, even more in ScB fish which exhibited higher ADC of DHA, and higher body growth compared to FO fish. Xu et al. (2014) also reported in the Japanese sea bass (_Lateolabrax japonicus_) increased tissue levels of n-3 LC-PUFA and increased gene expression of Δ6 fatty acyl desaturases (FADS2) when fish were fed diets rich in palmitic acid in the form of tripalmitin, compared to diets rich in either n-3 LC-PUFA, stearic, oleic, or ALA. It is generally common to see inhibition of ALA conversion to EPA and reduced retention of EPA in fish whole body by increasing dietary DHA levels (e.g., Bou et al., 2017b). Some retro-conversion of DHA to EPA may also occur in Atlantic salmon, which is nevertheless found to be rather limited, though increasing with essential fatty acid deficiency (Ruyter and Thomassen, 1999). Our data may thus indicate that the chosen dietary levels of EPA + DHA in this study were low. There are also indications that retroconversion of supplemented DHA to EPA occurs in humans, rats, and cell cultures but as reviewed by Metherel and Bazinet (2019) it is more likely that the reason
behind this response is a feedback inhibition of EPA elongation to DPA n-3 (e.g., Metherel et al., 2017).

**Dietary *Schizochytrium limacinum* Biomass Effects on Atlantic Salmon Filet Color**

Dietary *S. limacinum* biomass provided during the whole study a dose-related positive effect on Atlantic salmon filet pigmentation. Those effects were demonstrated in terms of increased redness, reduced paleness, and reduced melanin spots in the filets of the fish at slaughter. Intestinal uptake facilitation and antioxidant effect provided by the whole unicellular organism’s components, such as vitamins, organic Se, and DHA, may have contributed to these effects (Kanazawa, 1991). Bjerkeng et al. (1999) found a small positive effect on Atlantic salmon filet pigmentation (up to 14% increased deposition efficiency of astaxanthin) by dietary supplementation of *α*-tocopheryl acetate (vitamin E acetate). Likewise, in a study with turbot larvae, normal pigmentation was positively correlated with DHA/EPA ratio in live feed lipids and the authors concluded that DHA requirement for this purpose could not be replaced by EPA (Reitan et al., 1994). Assimilation of astaxanthin in salmon is rather poor increasing the need for dietary supplementation, which in turn increases significantly the formulation costs (Torrissen, 1995). Improved deposition of the supplemented astaxanthin by use of *S. limacinum* biomass in the diet, in combination with the observed positive effect on growth, may render the use of this natural alternative source of marine omega 3 fatty acids economically viable. When it comes to melanin discoloration, Mørkøre et al. (2015) reported that every fifth commercially produced salmon has black spots which occur early in the seawater phase increasing in prevalence and size toward slaughter. In the aforementioned report, melanin spots were associated with among other chronic inflammation and increased levels of minerals and microorganisms, which may have been alleviated by the use of high levels of *S. limacinum* biomass acting as prebiotic in the diet of the ScB fish groups (Komiyama et al., 2011).

**Liver and Mid Intestine Transcriptome**

Transcriptome analyses are a useful tool to widely identify potential health effects. The observed upregulation of the hsp, gpx, and gadd45 genes in the liver of ScB compared to the FO fish, in combination with the high incidence of pale livers, higher total lipid level, increased relative level of 18:1 n-9, and low EPA (EPA is known to have anti-inflammatory effects) in the liver of the ScB group may be an indication of mild stress in the liver of ScB fish. Even though we cannot from this study conclude with certainty, potential mild stress in the liver may have partly been due to specific lack of EPA in the ScB treatment. The *Schizochytrium* biomass rich diet was characterized by high DHA/EPA ratio. When the present trial was initiated it was considered that 1–1.3% EPA + DHA was adequate to cover salmon’s requirement (Ruyter et al., 2000). Nevertheless, a later study by Bou et al. (2017a) shows that 1% EPA of DHA in the diet long-term results in increased mortality under challenging conditions, as well as...
TABLE 9 | Microarray analysis results showing genes with expression change in the liver of salmon fed ScB compared to the FO diet (n = 12).

| Cell folding, protein modification                                      | Log-ER |
|------------------------------------------------------------------------|--------|
| Heat shock cognate 70 kDa protein                                      | 0.78   |
| Heat shock cognate 71 kDa protein                                      | 0.77   |
| Heat shock cognate 70 kDa protein                                      | 0.75   |
| Heat shock cognate 70 kDa protein                                      | 0.62   |
| **Cell redox**                                                         |        |
| Glutathione peroxidase                                                 | 0.82   |
| **Cell stress**                                                        |        |
| Growth arrest and DNA-damage-inducible protein GADD45 gamma            | 0.81   |
| **Metabolism retinoid**                                                |        |
| Retinol-binding protein 2                                              | 1.54   |
| **Metabolism xenobiotic**                                              |        |
| UDP-glucuronosyltransferase 2A2 precursor                              | 1.03   |
| **Immune**                                                            |        |
| Complement C1q tumor necrosis factor-related protein                   | −0.67  |
| **Immune adhesion**                                                   |        |
| Vascular cell adhesion protein 1                                       | −0.62  |
| **Immune antigen presentation**                                       |        |
| HLA class II histocompatibility antigen, DP alpha chain precursor      | −0.65  |
| HLA class II histocompatibility antigen, DP alpha chain precursor      | −0.69  |
| L77086_1 MHC class II alpha                                            | −0.72  |
| MHC class I antigen                                                    | −0.85  |
| BOLA class I histocompatibility antigen, alpha chain BL3-7 precursor  | −0.89  |
| **Immune chemokine**                                                  |        |
| C–C motif chemokine 19-2                                               | −0.72  |
| **Immune complement**                                                 |        |
| Complement factor H1 protein                                           | −0.62  |
| C1Q- and TNF-related protein                                           | −0.64  |
| **Immune IFN-virus response**                                         |        |
| Proteasome subunit beta type-8 precursor                               | −0.82  |
| **Immune Ig**                                                         |        |
| Ig kappa chain V–IV region J1 precursor                               | −1.17  |
| **Immune lectin**                                                     |        |
| Toll-like leucine-rich repeat [Salmo salar] Toll-like receptor 5       | 0.88   |
| Lectin receptor 5                                                      | −0.67  |
| **Immune TNF**                                                        |        |
| Metalloproteinase STEAP4                                               | 0.67   |
| TNF receptor member 11B                                                | −0.63  |
| C1Q- and TNF-related protein                                           | −0.64  |

Data are log-ER (relative gene expression between the dietary groups). Significant up- and down-regulated genes in the ScB compared to the FO groups (log-ER > 0.6; P < 0.05) are shown with red and green color shades, respectively.

TABLE 10 | Microarray analysis results showing genes with expression change in the mid intestine of salmon fed ScB compared to the FO diet (n = 12).

| Metabolism ion                                                                 | Log-ER |
|--------------------------------------------------------------------------------|--------|
| Zinc transporter ZIP4                                                          | 0.84   |
| **Metabolism lipid**                                                           |        |
| Perilipin                                                                      | −0.71  |
| Endothelial lipase precursor                                                   | −1.72  |
| **Metabolism protease**                                                        |        |
| Trypsin-2 precursor                                                            | 1.02   |
| Trypsin-1 precursor                                                            | 0.62   |
| **Metabolism xenobiotic**                                                      |        |
| Cytochrome P450 monooxygenase CYP2K1v2                                         | 1.02   |
| Sulfotransferase 6B1                                                           | 0.86   |
| Cytochrome P450                                                                 | 0.65   |
| **Immune Antigen presentation**                                               |        |
| MHC class I alpha 2; PCR product                                              | −0.68  |
| **Immune chemokine**                                                           |        |
| Chemokine CCL–C11b                                                             | 0.71   |
| **Immune effector**                                                            |        |
| Myeloperoxidase precursor                                                     | −0.93  |
| **Immune eicosanoid**                                                          |        |
| Leukothriene b4 12-hydroxydehydrogenase/15-ketoreductase                      | −0.62  |
| **Immune IFN-virus response**                                                 |        |
| Suppressor of cytokine signaling 1                                             | −0.62  |
| Patched domain containing 3                                                    | −0.63  |
| **Immune Ig**                                                                  |        |
| Ig kappa chain V region K29-213                                               | −0.63  |
| **Immune regulator**                                                           |        |
| Suppressor of cytokine signaling 1                                             | −0.62  |

Data are log-ER (relative gene expression between the dietary groups). Significant up- and down-regulated genes in the ScB compared to the FO groups (log-ER > 0.6; P < 0.05) are shown with red and green color shades, respectively.

have shown that feeding zebrafish microalgae rich diets can lead to gene expression changes in the liver. Microalgae supplements containing a species rich in arachidonic acid (ARA) lead to upregulation of immune system-related genes and antioxidant defense (Nayak et al., 2018) in contrast to our study where we saw mostly downregulation. EPA and DHA are known to have anti-inflammatory effects, whereas ARA is known to be pro-inflammatory, which can be part of the explanation for the differential responses. The fact that only a few differences were identified in gene expression between the two dietary groups in mid intestine agrees in general with the fact that mid intestine in both groups was found to be normal through histological evaluation.

Regarding the observed upregulation of a zinc transporter gene in the fish from the ScB group, recent unpublished results (FHF project no. 901265) also show that increased levels of EPA and DHA in the feed lead to increased levels of Zn in the intestinal tissue which indicates better Zn uptake. It appears thus that in our study higher level of DHA from the Schizochytrium biomass in the diet may have caused upregulation of the ZIP4 Zn transporter. More animal model studies as well as studies in human cell cultures has shown an increased incidence of hypervacuolization in the intestine and fatty liver which indicates that the requirement for these fatty acids is higher than what it was assumed to be earlier. The experimental fish were fed diets with ca. 1% EPA + DHA early on in the fresh water, and 1.3% EPA + DHA in the sea water phase. Higher growth in the ScB compared to the FO groups can also be the reason why signs of potential EPA deficiency first appeared in the ScB groups. The authors have no adequate explanation for the differential expression of other immune system-related genes which were also downregulated in the liver and mid intestine in the ScB compared to the FO group. More studies
The observed downregulation of other genes related to inflammatory response in the intestine, such as chemokine/cytokines may also indicate potential anti-inflammatory effects in the mid-intestine of the ScB compared to the FO dietary groups. Mammalian studies refer to observations of anti-inflammatory effects of beta-glucans in the intestine (Raa, 2015) which are compounds present in the cell walls of unicellular organisms such as yeast and microalgae.

**Intestinal Histology**

No diet-related differences in intestinal histology were observed between the ScB and FO dietary groups, indicating no specific adverse effect on intestinal development in relation to long-term feeding of ScB compared to FO. Mid-intestinal mucosa displayed no signs of abnormal vacuolization, as was previously observed in A. salmon fed diets which were low in EPA + DHA (Bou et al., 2017a). In distal intestine, the observations painted a more complicated picture, although it is important to note that there were no observations indicating differences between the dietary groups. The fact remains that all samples displayed a deviant morphology, with hyperplastic intestinal folds and non-vacuolized mucosa. Despite these deviations, the tissues were seemingly viable and functional, as judged by fish performance. This unknown condition had no apparent likeness to the intestinal inflammation associated with soybean meal-induced enteritis (Knudsen et al., 2007). Although having some common features, the hyperplastic growth of the mucosal folds in the present study contrasts the markedly reduced development of these structures in the soybean meal-associated condition. The cause of the deviant state of the distal intestinal mucosa in the present study remains unknown.

**CONCLUSION**

*Schizochytrium limacinum* biomass appears to be a good alternative source of marine omega 3 fatty acids for the fish in our study, in particular during stressful conditions, resulting in improved growth, indications of anti-inflammatory effects in the intestine based on gene expression data, and improved filet pigmentation tested both chemically and by sensory panel. However, it is important to cover the fish's requirement for total LCn-3HUFA, and in the present study there were indications that the dietary levels of EPA + DHA used (ca. 1.3%) were inadequate for both groups, but in particular for the faster growing *S. limacinum* dietary group showing increased liver fat content. It appears that *S. limacinum* biomass can be used in the feed for salmon throughout the whole production cycle from parr to slaughter without any negative effects on growth or product quality if the dietary requirement for EPA is covered. Lastly, the life-long use of *S. limacinum* biomass affects texture but not the taste or smell of the salmon filet.

**DATA AVAILABILITY STATEMENT**

The microarray data are available at GEO accession GSE140558.
ETHICS STATEMENT

The four feeding experiments followed the Norwegian animal welfare act guidelines, in accordance with the Animal Welfare Act of 20 December 1974, amended 19 June 2009. The facility at Sunndalsora, Nofima (division 60) was granted permission by the Norwegian Food Safety Authority (FOTS ID 8060) to run the experiments. The decision was made on the basis of Regulations 18 June 2015 on the use of animals in experiments.

AUTHOR CONTRIBUTIONS

KK, MT, KA, CM, SJ, and RB: planning and preparing experiment. KK, GB, MT, YT, CM, and RB: participated in carrying out experiment. KK, GB, MT, KA, BG, YT, and CM: analysis. KK, MT, KA, BG, YT, CM, SJ, and RB: writing and editing.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2020.00057/full#supplementary-material

FIGURE S1 | Atlantic salmon were kept in tanks and fed in triplicate groups from parr through smoltification and further on to body weight of ca. 800 g three diets, containing fish oil (FO), S. limacium biomass (ScB) or a mix of the two as EPA + DHA source. Fish were then mixed and moved to sea cages where there were fed in two triplicate groups either FO or ScB to slaughter.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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