The inclusion of a medicinal plant leaf extract (MPLE) from sage (Salvia officinalis) and lemon verbena (Lippia citriodora), rich in verbascoside and triterpenic compounds like ursolic acid, was evaluated in gilthead seabream (Sparus aurata) fed a low fishmeal-based diet (48% crude protein, 17% crude fat, 21.7 MJ kg⁻¹, 7% fishmeal, 15% fish oil) for 92 days. In particular, the study focused on the effect of these phytogenic compounds on the gut condition by analyzing the transcriptomic profiling (microarray analysis) and histological structure of the intestinal mucosa, as well as the histochemical properties of mucins stored in goblet cells. A total number of 506 differentially expressed genes (285 up- and 221 down-regulated) were found when comparing the transcriptomic profiling of the intestine from fish fed the control and MPLE diets. The gut transcriptome revealed an expression profile that favored biological mechanisms associated to the 1) immune system, particularly involving T cell activation and differentiation, 2) gut integrity (i.e., adherens and tight junctions) and cellular proliferation, and 3) cellular proteolytic pathways. The histological analysis showed that the MPLE dietary supplementation promoted an increase in the number of intestinal goblet cells and modified the composition of mucins’ glycoproteins stored in goblet cells, with an increase in the staining intensity of neutral mucins, as well as in mucins rich in carboxylated and weakly sulfated glycoconjugates, particularly those rich in sialic acid residues. The integration of transcriptomic and histological results showed that the evaluated MPLE from sage and lemon verbena is responsible for the maintenance of intestinal health, supporting gut
INTRODUCTION

Aquaculture will supply the majority of aquatic dietary protein by 2050 (1), playing a relevant role in food security and supply, and poverty alleviation (2). The sustained growth of aquaculture is highly dependent on the intensification of production (3), sustainable feed formulations (4) and generating farming conditions supporting fish health and welfare (5). Among the former concepts, disease is considered a main persistent threat to intensive fish farming, which represents an estimated US$6 billion loss per annum at a global scale (5). Under this scenario, aquaculture depends on the use of antibiotics to fight against infectious diseases that threatens production (6), with emerging infectious diseases forecast to increase with warmer temperatures (7). However, their use tend to result in the emergence of antimicrobial resistant bacteria, which may not represent a direct threat in terms of aquatic food consumption, but they could directly impact production itself by lowering drug efficacy, decreasing the animal’s immune system and selecting more virulent strains of pathogens (7). Considering the above-mentioned reasons, along with the increasing public awareness regarding food safety issues and the environmental impact linked to antibiotics’ use and animal welfare (7, 8), the development of functional feeds focused on promoting and modulating the host’s immune response has been encouraged during the last decade (9–12).

Functional feeds are recognized for promoting the growth, welfare and health of farmed animals coupled with an improvement and/or modulation of their immune system, as well as inducing physiological benefits beyond traditional feeding practices (13). In this sense, by preventive health management through the diet, fish can divert more energy to somatic growth and reduce biological energy reserves needed to fight disease or stress resistance (14). Furthermore, they can be used in addition to chemotherapeutic agents and vaccines (15). Among the long list of feed additives used in animal production (9, 16), phytopharmaceuticals derived from herbs, spices, medicinal or aromatic plants are residue-free, unlike synthetic antibiotics, and are safe ingredients for sustainable feeds (11, 17, 18). Although the mode of action of most phytopharmaceutical feed additives has not yet been fully elucidated, they are well-known for their antimicrobial, immunomodulatory, antioxidative, and growth-promoting effects in livestock (17) and aquatic animals (9, 18).

In a recent study from our research group, we showed that a phytopharmaceutical feed additive from sage (Salvia officinalis, Lamiaceae) and lemon verbena (Lippia citriodora, Verbenaceae) is an effective additive for aquafeeds since its inclusion at 0.1% in diets with low fishmeal (FM) content not only improved some key performance indicators (i.e., growth and feed efficiency performances), but also promoted fish systemic immunity. In particular, an ex vivo study with splenocytes from fish fed this phytopharmaceutical exposed to lipopolysaccharide (LPS) showed an up-regulation of genes involved in non-specific immune response, as well as pro- and anti-inflammatory cytokines, surface T-cell marker cd4, and antioxidative stress enzymes (14). However, the effects of this phytopharmaceutical additive still needs to be explored in terms of local immune response, especially at the intestinal level, since optimal health and functionality of the intestinal mucosa is essential for sustainable animal production (19, 20). This is of special relevance under the current scenario in which aquafeeds are formulated with low levels of fishmeal (FM) (21), since several studies have indicated that low FM diets compromised systemic immunity (22–25), as well as gut immune response (26, 27). In this context, intestinal immunity is a key factor in maintaining the general health of aquatic animals (20). The intestinal mucosa is a complex organ composed of the digestive epithelium with its specific structure, the gut-associated lymphoid tissue (GALT), and the mucus overlying the epithelium with its commensal microbiota (19). Furthermore, the intestinal epithelium acts as a selectively permeable barrier for dietary nutrients, electrolytes and water, while maintaining an effective defense against pathogens and tolerance towards dietary antigens. Thus, the GALT is reputed for mediating mucosal innate and adaptive immune responses, as well as being a key element for proper distinction between pathogens and commensal microbiota inhabiting the intestine (28–30).

Considering the importance of interaction between the diet and the gut, in the current study we evaluated the transcriptomic profile and histochemical properties of mucins stored in goblet cells of the intestine in juvenile gilthead seabream (Sparus aurata) fed a functional diet containing a medicinal plant leaf extract (MPLE) from sage and lemon verbena. This species is recognized as the most important Mediterranean aquaculture fish species in terms of volume and economic value (31). For this purpose, we focused on the modulation of the intestinal mucosal functionality and health by the above-mentioned phytopharmaceuticals when included in a diet with low fishmeal levels.

MATERIAL AND METHODS

Diets, Fish and Rearing Conditions

Two isoproteic (48% crude protein), isolipidic (17% crude fat) and isoenergetic (21.7 MJ kg⁻¹) experimental diets were formulated with a low FM content (7% FM) as described in Salomón et al. (14). Diets, named as control and MPLE, only
differed in their content of the feed additive evaluated, the MPLEx obtained from sage and verbena leaves, which was included in the MPLEx diet at 0.1% (Table 1). The MPLEx was obtained by NATACT Biotech SL using water/ethanol extraction (plant leaf extract ratio 5:1) and characterized as described in Arthur et al. (32) and Wójciak-Kosior et al. (33). The tested extract (proximate composition: 73% carbohydrates, 2% crude lipids, <1% crude proteins, 5% salts and 4% water) contained 10%, ursoic acid, 3% other triterpenic compounds, 2% verbascoside and <1% polyphenols. Thus, the content in plant-derived bioactive compounds in the MPLEx diet was 100 ppm ursoic acid, 30 ppm other triterpenic compounds, 60 ppm verbascoside and <10 ppm polyphenols. This phytogenic was incorporated in the mixture prior to extrusion. In brief, all powder ingredients were mixed in a double-paddle mixer (model RM90L, Mainca, Spain) and ground (below 250 μm) in a micro pulverizer hammer mill (model SH1, Hosokawa-Alpine, Germany). Diets were manufactured with a twin-screw extruder (model BC45, Clextral, France) with a screw diameter of 55.5 mm. Extruded pellets were dried in a vibrating fluid bed dryer (model DR100, TGC Extrusion, France). Oils were added by vacuum coating (model PG-10VCLAB, Dinnissen, The Netherlands). Immediately after coating, diets were packed in sealed plastic buckets and shipped by road to the facilities at IRTA Sant Carles de la Ràpita, Spain. Both extruded diets (pellet size: 2 mm) used in this trial were manufactured by SPAROS Ltda. (Portugal) and kept at 4°C until their administration. Proximate composition of the extract and experimental diets have been described in Salomón et al. (14).

A total of 300 gilthead seabream (body weight, BW = 5.0 ± 0.2 g; mean ± standard deviation) were obtained from a commercial fish farm, Piscicultura Marina Mediterránea SL (Andromeda Group, Burriana, Spain) and transported to the experimental facilities of IRTA in Sant Carles de la Ràpita (Tarragona, Spain). Then, fish were acclimatized for three weeks in 450 L tanks connected to a water recirculation system (IRTAmar™) at an initial density of 2 kg m⁻³. Acclimation was conducted in the same experimental tanks (450 L) where the nutritional experiment was carried out. Just before the start of the trial, all animals were gently anesthetized (50 mg l⁻¹ MS-222, Sigma-Aldrich, Madrid, Spain), individually measured in BW (26.0 ± 0.2 g) and distributed homogeneously among the eight experimental tanks (n = 35 fish per tank; 4 replicate tanks per experimental diet). During the trial that lasted 92 days, fish were fed at the daily rate of 3.0% of the stocked biomass, which approached apparent satiation as described in Salomón et al. (14). Feed ration was regularly adjusted by means of intermediate samplings along the trial. At the end of the trial, all fish were anesthetized as previously indicated and measured for individual BW. Fish performance in terms of survival (Scontrol diet = 98.0 ± 1.0%; S MPLEx diet = 99.0 ± 0.8%), growth (BWcontrol diet = 173.8 ± 8.2 g; BW MPLEx diet = 189.6 ± 5.0g) and feed conversion ratio (FCRcontrol diet = 1.25 ± 0.04; FCMPLEx diet = 1.10 ± 0.04) was already published in Salomón et al. (14). Twelve fish from each experimental diet (total= 3 fish per tank replicate) were randomly selected and sacrificed with an overdose of anesthetic (300 mg l⁻¹ MS-222) for gut transcriptomic and histological analyses. Sacrificed animals were eviscerated and mid-anterior intestine samples (1-1.5 cm length per specimen) were put in RNAlater™ (Invitrogen, Thermo Fisher Scientific, Lithuania), incubated overnight at 4°C, and stored at -80°C until RNA extraction. This region of the intestine (mid-anterior section) was chosen due to its specialized immunological functionality when compared with other intestinal sections (13). In addition, a similar piece of tissue was also dissected and fixed in 10% v/v neutral formaldehyde (pH: 7.2 ± 0.01) buffered with sodium phosphate (0.1M) for histological and histochemical purposes.

Water temperature (25.1 ± 1.5°C, range: 22-27°C), oxygen (6.1 ± 0.2 mg l⁻¹) (OXI330, Crison Instruments), and pH (7.5 ± 0.01) (pH meter 507, Crison Instruments, Barcelona, Spain) were daily controlled. Salinity (35%) (MASTER-20 T; ATAGO Co. Ltd), ammonia (0.13 ± 0.1 mg NH₄⁺ l⁻¹) and nitrite (0.18 ± 0.1 mg NO⁻ 1⁻ l⁻¹) levels (HACH DR9000 Colorimeter, Hach®, Spain) were weekly monitored. The trial was run under natural photoperiod according to the season of the year (August to November; 40°37’41” N).

**Transcriptional Analysis**

**RNA Isolation and Quality Control**

Total RNA was extracted from mid-anterior intestine of individual fish (n = 12 fish per dietary treatment) using the RNAsafe® Mini Kit (QiaGen, Germany). Total RNA from each individual sample was eluted in a final volume of 35 μl; nuclease-free water and treated with DNase using the DNA-free™ DNA Removal Kit according to manufacturer’s instructions.

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**Table 1** | List of ingredients and proximal composition of experimental diets; control and a basal diet supplemented with a medicinal plant leaf extract (MPLEx) obtained from sage (*Salvia officinalis*) and lemon verbena (*Lippia citriodora*).

| Ingredients, % | Control diet | MPLEx diet |
|----------------|--------------|------------|
| Fishmeal LT70 | 7.0          | 7.0        |
| Soy protein concentrate | 21.0 | 21.0 |
| Pea protein concentrate | 12.0 | 12.0 |
| Wheat gluten | 12.0         | 12.0       |
| Corn gluten | 12.0         | 12.0       |
| Soybean meal 48 | 5.0 | 5.0 |
| Wheat meal | 10.4         | 10.4       |
| Fish oil (Savinor) | 15.0 | 15.0 |
| Vitamin and mineral Premix PV01 | 1.0 | 1.0 |
| Soy lecithin – Powder | 1.0 | 1.0 |
| Binder (guar gum) | 1.0 | 1.0 |
| MCP | 2.0 | 2.0 |
| L-Lysine | 0.3 | 0.3 |
| L-Tryptophan | 0.1 | 0.1 |
| DL-Methionine | 0.2 | 0.2 |
| MPLEx | – | 0.1 |

Proximate composition

| Crude protein, % | 48.37 | 48.37 |
| Crude fat, % | 17.19 | 17.21 |
| Fiber, % | 1.52 | 1.52 |
| Ash, % | 5.88 | 5.88 |
| Gross Energy, MJ/kg | 21.62 | 21.62 |

MPLEx, medicinal plant leaf extract obtained from sage (*Salvia officinalis*) and lemon verbena (*Lippia citriodora*). Proximal compositions of the diets were according to Salomón et al. (14), following the AOAC guidelines.
NanoDrop ND-2000 dye incorporation and cRNA yield were checked with the samples were purified using an Agilent 2100 Bioanalyzer (Agilent Technologies, Spain). Pooled RNA analyzed in this study were selected by the criteria of a RIN value > 8.5 (Supplementary Figure 1). This methodological approach of pooling RNA from four different specimens in each replicate (n = 3 microarray replicates) allowed authors evaluating population’s variability (n = 4 replicate tanks; 1 fish per tank in each pooled RNA; N = 12 animals); however, the information regarding individual variability was lost with this choice.

Microarray Hybridization and Analysis
Transcriptional analysis was carried out using the Aquagenomics Sparus aurata oligonucleotide microarray v2.0 (4 x 44 K) (SAQ) platform. The detailed information about the platform and the transcriptionic raw data for all samples included in this current analysis is available through the public repository Gene Expression Omnibus (GEO) at the US National Centre for Biotechnology Information (NCBI) (accession number GPL13442 and GSE166558, respectively).

Transcriptomic analysis from both experimental groups were conducted as described by Reyes-López et al. (35). Briefly, 200 ng of total RNA from each sample pool was reverse transcribed along with Agilent One-Color RNA spike-in kit (Agilent Technologies, USA). Then, total RNA was used as template for Cyanine-3 (Cy3) labelled cRNA synthesis and amplification with the Quick Amp Labelling kit (Agilent Technologies). cRNA samples were purified using the RNeasy micro kit (Qiagen). Dye incorporation and cRNA yield were checked with the NanoDrop ND-2000® spectrophotometer. Then, 1.5 mg of Cy3-labeled cRNA with specific activity > 6.0 pmol Cy3/μg cRNA were fragmented at 60°C for 30 min, and then the samples were mixed with hybridization buffer and hybridized to the array (ID 025603, Agilent Technologies) at 65°C for 17 h using the Gene expression hybridization kit (Agilent Technologies). The microarray washes were conducted as recommended by the manufacturer using Gene expression wash buffers (Agilent Technologies) and stabilization and drying solution (Agilent Technologies). Microarrays slides were scanned with an Agilent Technologies Scanner (model G2505B); spot intensities and other quality control features were extracted with Agilent’s Feature Extraction software version 10.4.0.0 (Agilent Technologies). Quality reports were checked for each array. Although validation by mean of qPCR is required when there is a high risk of obtaining paralog genes and unspecific hybridization; in our study, we used an oligonucleotide-based microarray, which has probes with a reduced number of bases and high affinity, which avoids the necessity of conducting the validation of gene expression results by qPCR validation.

Transcripteractome
The Search Tool for the Retrieval of Interacting Genes (STRING) public repository version 11.0 (https://string-db.org) was used in order to obtain the gut transcripteractome for those differentially expressed genes (DEGs) from fish fed the MPLE diet in comparison to the control group (P- value < 0.05) (36). This functional network analysis has gained increasing attention because of the association between different genes sorted in different clusters that may have complementary functions into a biological context of response (35). A protein-protein interaction (PPI) network of DEGs was conducted with a high-confidence interaction score (0.9) using Homo sapiens as model organism. Furthermore, to confirm matches of the genes acronyms tag between both H. sapiens and gilthead seabream species Genecards (37) and Uniprot (38) databases were used. In order to confirm match of gene acronyms between both H. sapiens and gilthead seabream species, human orthology identification based on gene/protein name was accessed through the Genecards (www.genecards.org) (37) and Uniprot (www.uniprot.org) (UniProt, 2019) databases. Additionally, protein-protein BLAST (BLASTp) were analyzed (E-value < 10^-7; query cover > 95%). Gene ontology (GO) enrichment analysis for the DEGs were also performed by STRING (P < 0.05).

Histological and Histochemical Analyses
Samples (n = 12 fish per experimental diet) were embedded in paraffin and sagittally sectioned (5-6 μm). A total of 576 sections (2 per each sample x 24 samples x 12 techniques) were used for histological and histochemical purposes. Two sections per each sample were stained with hematoxylin–VOF for descriptive purposes; the rest were used for evaluating the histochemical properties of epithelial and mucous cells. In brief, Schiff, Periodic Acid Schiff (PAS), diastase-PAS and Alician Blue (AB) pH 2.5, 1 and 0.5 (carboxylated and sulphated glycoconjugates/glycoproteins) techniques were used for studying carbohydrate distribution. Furthermore, several horseradish peroxidase (HRP) conjugated lectins (Sigma-Aldrich, Spain) were used for proper characterization of different glucidic residues bound to the glycoconjugates; in particular, Canavalia ensiformis/ConA (Mannose and/or Glucose), Ulex europeus/UEA-I (L-Fucose), Triticum vulgaris/WGA (N-acetyl-D-glucosamine and/or N-acetyleneuraminic acid, NeuNac/sialic acid/NANA), Glycine max/SBA (α-N-acetyl-D-galactosamine) and Sambucus nigra/SNA (NeuNac/sialic acid/NANA). Lectin concentrations ranged between 15 μg ml−1 to 30 μg ml−1. Regarding negative controls, omission of the respective lectin, substitution of lectin-HPR conjugates by TBS and treatments with different enzymes were performed according to Sarasquete et al. (39). The peroxidase activity was visualized with 3,3-diaminobenzidine tetra hydrochloride/DAB and hydrogen peroxide (0.05%). All the techniques were performed according to Pearse (40) and following proper standardized techniques and protocols (41). All reagents were purchased from Sigma Chemical Co. St Louis, MO, USA.

Histological images were taken with a Leitz Wetzlar microscope with a built-in SPOT Insight Color camera (Ernst Leitz Wetzlar GmbH, Germany). Results were manually
registered using a semi-quantitative assessment scoring based on color intensity scores (0, negative; 1, weak; 2, moderate; 3, intense; 4, very intense) from four independent observers, comparing the sections of the control with the experimental diet. The mucous cell count was determined in four different sites of each histological section, and the number of cells expressed per length unit of the basal lamina of the mucosal epithelium (1 mm) according to Yamamoto et al. (42).

Ethics Statement
All animal experimental procedures were complied with the Guiding Principles for Biomedical Research Involving Animals (EU2010/63), the guidelines of the Spanish laws (law 32/2007 and RD 1201/2015), and authorized by the Ethical Committee of the Institute for Research and Technology in Food and Agriculture (IRTA, Spain) for the use of laboratory animals.

Statistics
Extracted raw data from microarrays were imported and analyzed with Genespring version 14.5 GX software (Agilent Technologies). The 75% percentile normalization was used to standardize arrays for comparisons and data were filtered by expression. An unpaired t-test was conducted without correction to identify those DEGs between fish fed control and MPLE diets. The Principal Component Analysis (PCA) was carried out using GeneSpring software, four eigenvectors were calculated to describe the aggregation of the MPLE and control groups in a 3D plot. Venn diagram and the hierarchical heatmap were all obtained also with Genespring (version 14.5 GX software, Agilent). Changes in the number of mucous cells between experimental diets were analyzed by means of an unpaired t-test assuming data homoscedasticity (Barlett’s test). All the analysis were performed using GraphPad PRISM 7.00. The level of significance was set at $P < 0.05$ for all statistical tests.

RESULTS

Organization of the Intestine and Mucins’ Histochemistry Produced by Goblet Cells
In both experimental groups, the intestinal mucosa was lined by a simple columnar epithelium with basal nuclei, basophilic cytoplasm and prominent brush border with scattered goblet cells. The organization of the lamina propria-submucosa and muscular layers was normal. No signals of histological alterations associated to inflammatory processes in the intestine were observed in fish fed the MPLE diet compared to the control group (Figures 1A, B). Fish fed the MPLE diet showed a higher density of goblet cells along the intestinal epithelium compared to fish fed the control diet (Figure 2).

Regarding the histochemical properties of mucins in goblet cells from the anterior intestine, results showed a variable richness of neutral glycoproteins (PAS and diastase-PAS positive) (Table 2). In addition, mucins from goblet cells showed a mixture of carboxylated (AB pH = 2.5) and sulphated acidic groups (weak and strongly ionized; AB pH = 1.0 and pH 0.5, respectively) (Table 2 and Figure 1C).

Furthermore, a specific affinity for WGA, SNA and SBA lectins was detected in the mucinous content of goblet cells (Table 2 and Figures 3A–L). Moreover, no variations were detected in the distribution of mucosal cell glycoconjugates between the upper and the bottom areas of the intestinal folds. When comparing both dietary groups, the dietary administration of the MPLE modified the composition of glycoproteins of mucins produced by goblet cells, with an increase in the staining intensity of neutral mucins, as well as in mucins rich in carboxylated and weakly sulphated glycoconjugates (Table 2 and Figures 1C, D). In addition, an increase in affinity for WGA and SBA lectins and a decrease in the affinity for the SNA lectin was found in the mucinous content of goblet cells, whereas no changes were detected regarding ConA and UEA-I lectins (Table 2 and Figures 3A–L).

Microarrays and Gut Transcriptome
A total number of 506 DEGs were found when comparing the transcriptomic profiling of the intestine from gilthead seabream fed the control and MPLE diets ($P < 0.05$; Figure 4A and Supplementary Table 1). Common segregation among the pool samples within the same dietary treatment was observed in the hierarchical clustering for the gut transcriptomic response based in similitude patterns of the DEGs response ($P < 0.05$; Figure 4B). The observed segregation among dietary treatments is supported by the PCA analysis for the analyzed samples (Figure 4C); in particular, four eigenvectors were calculated and three principal components were plotted, explaining the 83.2% of the total variability [component 1 (X-axis): 45.5%; component 2 (y-axis): 24.4%; component 3 (z-axis): 13.3%]. The detailed analysis of gene absolute fold-change (AFC) revealed that genes were mostly up-regulated in fish fed the MPLE diet (56.3% of DEGs), while its modulation was moderate in terms of AFC intensity (Figure 4). In particular, 285 of the above-mentioned DEGs were up-regulated with 219 of them within the 1.0 ≤ AFC ≤ 1.5 interval, 58 DEGs were grouped within the 1.5 ≤ AFC ≤ 2.0, and the last 9 DEGs up-regulated were grouped 2.0 ≤ AFC ≤ 3.0. In contrast, 221 genes were significantly down-regulated and grouped in the range -1.0 ≤ AFC ≤ -3.0. ($P < 0.05$). In particular, 164 DEGs were mainly concentrated in the -1.0 ≤ AFC ≤ -1.5 range. Among them, 42 DEGs were grouped within the -1.5 ≤ AFC ≤ -2.0 interval, 10 DEGs were felt within the -2.0 ≤ AFC ≤ -3.0 category, while only 5 were found to have an AFC higher than -3 (Figure 4).

From the whole set of DEGs, a functional network analysis was performed. The transcriptome showed 244 genes with 552 interactions (edges). The remaining 264 DEGs were classified as unknown genes; thus, they were excluded from the analysis. According to GO results and their respective annotation hierarchy, three main representative groups of genes were identified in the transcriptome among the totality of biological processes obtained from the enrichment analysis (Supplementary Table 2): (1) immune system processes, (2) cellular development and organization, and (3) cellular catabolism (Supplementary Tables 3–5). Table 3 summarizes the most relevant DEGs in fish fed the MPLE diet in relation to the above-mentioned biological processes.
Regarding the dietary regulation of biological processes related to gut immunity, 18 genes were up-regulated and 14 genes down-regulated in fish fed the MPLE diet. In particular, several relevant GOs related to immunity were obtained such as “T cell activation” (GO:0042110; 7 up-regulated genes; 7 down-regulated genes), “T cell differentiation” (GO:0030217; 6 up-regulated genes; 5 down-regulated genes), “T cell lineage commitment” (GO:0002360; 3 up-regulated genes; 2 down-regulated genes), “leukocyte differentiation” (GO:0023610; 8 up-regulated genes; 7 down-regulated genes), “lymphocyte activation” (GO:0046649; 7 up-regulated genes; 6 down-regulated genes), “lymphocyte differentiation” (GO:0030098; 6 up-regulated genes; 5 down-regulated genes), “CD4+ or CD8+ T cell lineage commitment” (GO:0043369; 2 up-regulated genes; 2 down-regulated genes), “T cell selection” (GO:0045058; 3 up-regulated genes; 2 down-regulated genes) and “intracellular receptor signaling pathway” (GO:0030522; 7 up-regulated genes; 2 down-regulated genes) (Figure 5 and Supplementary Table 3).

Furthermore, the MPLE diet promoted the regulation of biological processes associated with cellular development and organization with 19 up-regulated and 14 down-regulated genes. Among them, we found the terms “regulation of adherens junction organization” (GO:1903391; 3 up-regulated genes; 3 down-regulated genes), “regulation of anatomical structure morphogenesis” (GO:0022603; 14 up-regulated genes; 12 down-regulated genes), “negative regulation of cell size” (GO:0045792; 2 up-regulated genes; 1 down-regulated gene), “establishment of endothelial barrier” (GO:0061028; 3 up-regulated genes; 1 down-regulated gene), “regulation of cell junction assembly” (GO:1901888; 4 up-regulated genes; 2 down-regulated genes), “regulation of cell-substrate adhesion” (GO:0010810; 4 up-regulated genes; 5 down-regulated genes), “negative regulation of cell-substrate adhesion” (GO:0010812; 3 up-regulated genes; 2 down-regulated genes), “regulation of focal adhesion
genes; 3 down-regulated genes) and “ubiquitin-dependent protein catabolic process” (GO:0006511; 14 up-regulated genes; 3 down-regulated genes) were obtained (Figure 7 and Supplementary Table 5).

DISCUSSION

Considering the close relationship between diet and gut condition and the consequences on the organism and overall health, evaluating the interactions between dietary ingredients and the intestine is of special relevance due to the wide array of functions that have been associated to the gastrointestinal tract (34, 43). This is of special relevance when evaluating functional feed additives that are supposed to promote health and nutrition in farmed animals (9, 44). In this context, the present study aimed to evaluate the effects of a MPLE from sage and lemon verbena on the transcriptomic profiling, histological organization and lectin histochemistry of the intestine. Extracts from these medicinal plants are reputed for their beneficial pharmacological activities, including antiseptic, antioxidant, and anti-inflammatory properties among others (45–47). Moreover, in gilthead seabream we demonstrated that in addition to act as growth promoters, they modulated systemic immunity when splenocytes were incubated with a bacterial-type pathogen-associated molecular pattern (PAMP) like LPS (14).

MPLE From Sage and Lemon Verbena
Exert Controlled Immune and Pro-Inflammatory Responses in GALT

Fish possess innate and adaptive immune defense systems. The innate parameters are at the forefront of immune defense and are a crucial factor in disease resistance. The adaptive response of fish is essential for long-lasting immunity and is considered as a key factor in improving prophylactic strategies, and the use of functional feeds with immunomodulatory properties (20, 48). T cells are one of the main players of the adaptive immune response, being the intestine of teleosts an important site of T cells production among mucosal tissues (49, 50). In particular, T cells represent the major leucocyte population within the ileal gut (51) with relevant cytotoxic activity (52), as well as playing an important role in foreign antigen recognition and gut homeostasis (53, 54). Under the current experimental conditions, MPLE from sage and lemon verbena promoted T cell activation, differentiation and selection in gilthead seabream gut. Thus, the MPLE-supplemented diet induced up-regulation of the mammalian target of rapamycin (mtor) gene in the gut of gilthead seabream. The mTOR signaling pathway has been reported to promote the production of anti-inflammatory cytokines by myeloid immune cells with an important ability to limit the pro-inflammatory mediators (55) thus, playing a crucial role in intestine inflammation, epithelial morphogenesis (56, 57), as well as being an important central regulator of immune responses (58, 59). The increase in mtor expression may be associated to the down-regulation of nlr3, which is a negative regulator of the PI3K/AKT/mTOR pathway (60), assembly” (GO:0051893; 3 up-regulated genes; 2 down-regulated genes), “negative regulation of adherens junction organization” (GO:1903392; 1 up-regulated genes; 1 down-regulated genes), and “cell aging” (GO:0007569; 3 up-regulated genes; 1 down-regulated gene) (Figure 6 and Supplementary Table 4).

The tested functional feed additive resulted in the positive regulation of biological processes related to cellular proteolytic processes, showing 35 up-regulated and 11 down-regulated genes. In particular, “cellular macromolecule catabolic process” (GO:0044265; 23 up-regulated genes; 6 down-regulated genes), “proteolysis” (GO:0006508; 27 up-regulated genes; 8 down-regulated genes), “protein catabolic process” (GO:0030163; 20 up-regulated genes; 3 down-regulated genes), “proteolysis involved in cellular protein catabolic process” (GO:0051603; 17 up-regulated genes; 3 down-regulated genes) and “ubiquitin-dependent protein catabolic process” (GO:0006511; 14 up-regulated genes; 3 down-regulated genes) were obtained (Figure 7 and Supplementary Table 5).

**TABLE 2 |** Histochemical characteristics and lectin affinity of mucins produced by goblet cells from anterior intestine of gilthead seabream (Sparus aurata) fed a control or a medicinal plant leaf extract (MPLE)-supplemented diet. The MPLE included in the basal diet was obtained from sage (Salvia officinalis) and lemon verbena (Lippia citriodora). The asterisk indicates statistically significant differences among dietary treatments (t-test; P < 0.05).

| General histochemistry | Control diet | MPLE diet |
|------------------------|--------------|-----------|
| Neutral glycoproteins  | 1 - 2        | 2 - 3     |
| Carboxylated glycoproteins | 1 - 3     | 3        |
| Weakly ionised sulphated glycoconjugates | 2 - 3 | 3 |
| Strongly ionised sulphated glycoconjugates | 2 - 3 | 2 - 3 |
| Lectin histochemistry   |              |           |
| ConA (Man/Glu)          | 0            | 0         |
| WGA ([βGlcNAc]>>NeuNAc/sialic acids/NANA) | 2 - 3 | 3 |
| SNA (NeuAcα2; sialic acids/NANA) | 1 - 3 | 0 |
| SBA (α/β GalNAc)        | 0 - 3        | 1 - 3     |
| UEA-I (Fuc)             | 0            | 0         |

Semi-quantitative assessment scoring based on color intensity scores: 0, negative (non-detected); 1, weak; 2, moderate; 3, intense; 4, very intense.
thus suggesting a balanced inflammatory dietary-induced response in the gut of fish fed the MPLE diet.

The NF-κB pathway is involved in the transcriptional regulation of several cytokines, chemokines, transcription factors, antimicrobial peptides, and interferon-stimulated genes, playing a critical role in regulating the survival, activation and differentiation of innate and adaptive immune cells (61); thus, playing a key role in regulating gut inflammation (62). In this way, several genes associated with the NF-κB pathway (cyld, clec4e, nlrc3, chi3l1, bcl2) were differentially regulated by the dietary supplementation of MPLE. Although scarce information is available for teleosts, CYLD identified in rainbow trout (Oncorhynchus mykiss) would have a similar function as in mammals (63). In our study, cyld was up-regulated, which may indicate that in fish fed the MPLE diet this gene acted as an inhibitor of the NF-κB signaling pathway, thus acting as a negative regulator of gut inflammation (63, 64). Accordingly, the up-regulation of ubiquitin specific peptidase 4 (usp4) and down-regulation of Ubiquitin specific peptidase 7 (usp7) may also be associated to the inhibition of the activation of NF-κB and mitogen-activated protein kinases stimulated by innate immune receptor (65–67). Additionally, chi3l1 was also up-regulated in fish fed the MPLE diet, promoting cell survival and proliferation, but not acting as a first inflammation-responsive factor. Thus, the differential regulation of the above-mentioned genes provide evidence that MPLE might induced inflammatory mechanisms related to the NF-κB signaling pathway (68, 69), which involved the production of pro-inflammatory cytokines (70). In this sense, histological data allow discarding the hypothesis that the administration of the MPLE caused an exacerbated gut inflammation, which would have been potentially detrimental for the health and condition of the intestinal mucosa of the gilthead seabream.

The Caspase-8 is a crucial executor for apoptotic initiation in fish (71). In our study, casp8 was significantly up-regulated in the intestine of fish fed the MPLE-supplemented diet. Several studies have reported the advantageous outcomes of having the casp8 expressed in a controlled way, demonstrating the essential role of this gene in maintaining the gut barrier in response to mucosal pathogens by permitting inflammatory shedding and preventing necroptosis of infected cells (72, 73). In addition, stat3 (Signal transducer and activator of transcription 3) was down-regulated, which suggests a healthy intestinal mucosa as this gene is positively regulated by pro-inflammatory cytokines under mucosal damage (74). Furthermore, our enrichment analysis showed regulation of biological processes associated with cellular proteolysis, in particular several genes coding for de- and ubiquitination (cyld, uba1, uba2a, uba2g1, uba2d2, usp4, usp7) and proteasome (cops4, psmd2, psmd1, psmd5, sqstm1) that may be indirectly involved in NF-κB pathway. In addition, degradation of a protein via the ubiquitin-proteasome pathway and subsequent proteolysis are important mechanisms in the
regulation of cell cycle, cell growth, tissue regeneration, signal transduction, and gene transcription (75).

A successful immune response involves the tight control of a wide repertoire of processes including activators and regulators at cellular and molecular level. In this context, another up-regulated gene that deserves attention is the Chitinase 3 like 1 (chi3l1). This gene is reputed for regulating the AKT1 signaling pathway (76), which controls innate immune cell development and function (77). Additionally, we found down-regulation of the NOD-like family CARD domain containing 3 (nlrc3), which plays an important role in modulating T cell responsiveness and inhibition of the pro-inflammatory mechanism (78). Besides, galectin 1 (lgals1) was also down-regulated in the intestine of fish fed the MPL diet. LGALS1 is considered a master regulator of homeostatic signals to shut off T-cell effector functions (79), acting as an immunoregulator and in turn modulating the...
pro-inflammatory cytokines secretion (80). At receptor level, the C-type lectin domain family 4 member e (clec4e, also known as mincle) was down-regulated in the intestine of fish from the MPLE group. Mincle is a transmembrane germline-encoded pattern recognition receptors (PRRs) with a pivotal role in the activation of the immune response (81, 82). Thus, when considering the overall expression patterns of DEGs, the dietary-induced activation of the immune function by the tested MPLE seems to be a well-controlled process, according to the observed balance between the up- and down-regulated DEGs (18 and 14 genes, respectively) and also the functional role of the DEGs related to T cells. As mentioned above, these results are in agreement with those recently published by Salomo et al. (14) where the same feed additive showed a tightly controlled systemic immune response in an ex vivo assay using splenocytes stimulated by LPS.

It is also of interest to illustrate that fish fed the supplemented-MPLE diet resulted in an increase of goblet cell population in the intestinal epithelium. These secretory cells are responsible for producing a thick layer of mucus that forms the front line of innate host defense, among other functions (83). The increase in intestinal goblet cell number in fish fed MPLE-supplemented diet would benefit fish by providing an effective immune barrier against potentially pathogenic gut bacteria. In addition to changes in their number, goblet cells also changed their histochemical composition. Particularly, we observed an increment in staining intensity of neutral and acid carboxylated and sulphated glycoproteins, results that are of special relevance since O-glycosylation pattern of mucin glycoproteins and their polypeptide backbone structure in combination with the depth and viscosity of the mucus layer, are important factors that regulate gut health and condition (84). In this sense, the increase in acid carboxylated (AB pH = 2.5) and sulphated (weak and strongly ionized; AB pH = 1.0 and pH 0.5, respectively) mucin glycoproteins produced by goblet cells from fish fed the MPLE-supplemented diet would be associated to an increment in the viscous properties of the mucus (85, 86). Our results are in agreement with those reported in poultry where the

### TABLE 3 | List of the most relevant DEGs related to three main representative biological processes identified by the transcripteractome (1, immune system processes; 2, cellular development and organization; 3, cellular catabolism) in fish fed the MPLE diet.

| Gene description                                      | Gene acronym | Biological processes | AFC | P-value |
|-------------------------------------------------------|--------------|----------------------|-----|--------|
| CYLD lysine 63 deubiquitinase                         | cyld         | 1                    | 2,374 | 0.027  |
| Actinin Alpha 4                                        | actn4        | 2                    | 1,631 | 0.022  |
| Proteasome 26S subunit, non-ATPase 1                  | psmd1        | 1, 3                 | 1,582 | 0.015  |
| Ubiquitin like modifier activating enzyme 1           | uba1         | 1, 3                 | 1,570 | 0.020  |
| Chitinase 3 Like 1                                     | chi3l1       | 1                    | 1,549 | 0.046  |
| Proteasome 26S subunit, non-ATPase 5                  | psmd5        | 1, 3                 | 1,537 | 0.041  |
| Erythrocyte membrane protein band 4.1                 | epb41        | 2                    | 1,476 | 0.032  |
| Caspase 8                                              | casp8        | 1                    | 1,472 | 0.004  |
| Sequestosome 1                                         | sqstm1       | 1, 3                 | 1,413 | 0.031  |
| Proteasome subunit beta 5                             | psmb5        | 1, 3                 | 1,408 | 0.020  |
| Zinc and ring finger 3                                 | znf3         | 3                    | 1,336 | 0.038  |
| RAP1B, member of RAS oncogene family                  | rap1b        | 2, 3                 | 1,321 | 0.004  |
| LIM domain binding 1                                   | ldb1         | 2, 3                 | 1,311 | 0.019  |
| COP9 signalosome subunit 4                            | cops4        | 1, 3                 | 1,255 | 0.001  |
| Ubiquitin specific peptidase 4                        | usp4         | 1, 3                 | 1,231 | 0.037  |
| Ras homolog family member A                            | rhoa         | 2                    | 1,222 | 0.047  |
| Fibroblast growth factor 18                           | fgf18        | 3                    | 1,206 | 0.041  |
| Mechanistic target of rapamycin kinase                | mtor         | 1, 3                 | 1,176 | 0.021  |
| Proteasome subunit beta 2                             | psmb2        | 1, 3                 | 1,167 | 0.023  |
| Phosphoglycerase kinase 1                             | pgk1         | 3                    | 1,164 | 0.046  |
| Ubiquitin conjugating enzyme E2 A                       | ube2a        | 1, 3                 | 1,102 | 0.039  |
| Ubiquitin conjugating enzyme E2 G1                     | ube2g1       | 1, 3                 | -1,100 | 0.001  |
| C-Type lectin domain family 4 member e                 | clec4e       | 1                    | -1,154 | 0.013  |
| NLR family card domain containing 3                    | nlr3         | 1                    | -1,315 | 0.020  |
| Signal transducer and activator of transcription 3    | stat3        | 1, 3                 | -1,367 | 0.0005  |
| BCL2, apoptosis regulator                              | bcl2         | 1                    | -1,378 | 0.038  |
| Ubiquitin specific peptidase 7                        | ups7         | 1, 3                 | -1,385 | 0.044  |
| Galectin 1                                             | lgals1       | 1, 3                 | -1,452 | 0.020  |
| Ubiquitin conjugating enzyme E2 D2                     | ube2d2       | 1, 3                 | -1,457 | 0.021  |
administration of phytogenics from sage promoted the secretion of neutral and acidic mucins in gut as well as mucus layer thickness in the ileum (87). Regarding lectin histochemistry, mucins from fish fed the MPLE-supplemented diet showed an increase in affinity for the WGA and SBA lectins, while for SNA lectin affinity was diminished. These results indicated that the tested feed additive changed the glycosylation patterns of intestinal mucins, being of special relevance the increase in WGA lectin affinity, as they indicate an increase in the presence of sialic acid residues. Sialic acids are involved in recognition processes, and they participate in defense mechanisms against microorganisms (88). These results from lectin histochemistry reinforce the hypothesis that the MPLE-supplementation varied the composition of mucus produced by goblet cells; thus, modulating an effective immune barrier against potentially pathogenic gut bacteria.
The homeostatic balance between epithelial cell proliferation and apoptosis is essential for the maintenance of the epithelial function, including regulation of epithelial permeability, the inflammatory response and the absorption of nutrients (89). An imbalance in the intestinal barrier structure can trigger an uncontrollable immune reaction in the intestinal microenvironment, increasing the translocation of bacterial antigens and stimulating inflammation in the intestine (90). Thus, the maintenance of the integrity of the gut barrier is essential to counteract such imbalances and guarantee fish growth, health and welfare (91). Dietary MPLE from sage and lemon verbena modulated the expression of genes involved in biological processes related to cellular development and organization. In particular, Ras homolog family member A (rhoa) was

**FIGURE 6** Cellular development and organization related Protein-Protein Interactions Network (PPI) network of the differentially expressed genes (DEGs) in the mid-anterior intestine of juvenile gilthead seabream (Sparus aurata) fed the mixture of MPLE obtained from sage (S. officinalis) and lemon verbena (L. citriodora) (see also Supplementary Table 4). Nodes colors indicate the biological processes for each DEG represented. Gene Ontology (GO) definitions, count of DEGs within each biological processes and respective false discovery rate are described in the graphical figure legend. ▲ nodes represent up-regulated genes and ▼ nodes represent down-regulated genes. Graphic keys and network stats are indicated in the graphical figure legend. Network Stats: number of nodes: 243; number of edges: 550; average node degree: 4.53; avg. local clustering coefficient: 0.416; expected number of edges: 423; PPI enrichment p-value: 1.98e-09. The MPLE included in the basal diet was obtained from sage (Salvia officinalis) and lemon verbena (Lippia citriodora).
RhoA belongs the small GTPase protein family, thus having a functional role in regulating many cellular events, including cell migration, organization of the cytoskeleton, cell cycle progression and cell adhesion (92). In this sense, rhoa expression has been associated to the assembly, regulation and maintenance of adherens (AJs) and tight (TJs) junctions (93, 94). Adherens junctions are cadherin/catenin-containing adhesive structures located below the tight junctions on the bilateral membrane (95).

In particular, the Erythrocyte membrane protein band 4.1 (epb41) was another up-regulated gene in the intestine of fish fed the MPLE-diet. EPB41 regulates cell proliferation and adhesion (96), as well as the integrity of the AJ (95) and plays an important role in the organization and function of the TJs and AJs by establishing a link between the TJ and actin in the cytoskeleton (97). Furthermore, other DEGs related to the formation and stabilization of AJs and TJs (f11r, actn4, rap1b) were also up-regulated in the gut of fish fed the MPLE-diet.
The F11 receptor (F11R) also known as the Junctional adhesion molecule (JAM)-A is a cell-cell adhesion molecule of the immunoglobulin superfamily, which is expressed by a variety of tissues, regulating diverse processes such as epithelial and endothelial barrier formation, among others (98). Our results are in agreement with those found in gilthead seabream fed a phytogenic feed additive derived from olive oil (89). Alpha-actinin-4 (ACTN4) is a member of the superfamily of actin-binding, which is localized at AJs and more specifically as a component of the zonula occludens of the TJ (99) and/or belt desmosomes in the zonula adherens (100) in connection with α-catenin, regulating the actin cytoskeleton and increasing cellular motility (101, 102). Another gene that was up-regulated by the dietary administration of MPL from sage and lemon verbena was the rap1b. This gene is a member of RAS oncogene family being necessary for normal human endothelial cell function (103), as well as having a crucial role for T cell homeostasis in the intestine (104). Regarding the gut integrity barrier, rap1b controls epithelial permeability probably by regulation of PI3K/Akt and correct nectin localization (94).

The homoeostasis of the constantly renewing intestinal epithelium relies on an integrated control of proliferation, differentiation and apoptosis, as well as on the functional architecture of the epithelial cells. In this sense, different DEGs indicated that the MPL-supplemented diet regulated mostly processes involved cell proliferation (fgf18, pgk1, ldb1, znrf3, mtor, lgal1, stat3) and de- and ubiquitination (cyld, uba1, ube2a, ubc2g1, ube2d2, usp4, usp7) and proteasome (cops4, psmb2, psmd1, psmb5, psmd5, sqstm1) pathways. The Fibroblast growth factor 18 (FGF18) is a pleiotropic growth factor which stimulates the proliferation of mesenchymal and epithelial cells and tissues (105); thus, its up-regulation in the tissue of gilthead seabream fed the MPL-supplemented diet suggests that the tested phytogenic feed additive might promote cell proliferation. Similar conclusions may be drawn when considering the up-regulation ldb1 (LIM domain binding 1) and znrf3 (Zinc and ring finger 3), since these genes regulate cell proliferation in intestinal crypts by means of the regulation of the Wnt/β-catenin signaling (106, 107). In addition, the up-regulation of mtor (mammalian Target of Rapamycin) and pgk1 (Phosphoglycerate kinase 1) indicate the regulation of intestinal cell proliferation by means of controlling cell metabolism (108, 109). These results are supported by the down-regulation of lgal1 (galectin 1), since this gene product may act as a negative growth factor that regulates cell proliferation (110). Differential expression of genes involved in de- and ubiquitination and proteasome pathways supports the above-mentioned idea that the tested MPL-supplemented diet promotes and regulates cell cycle associated to intestinal cell proliferation (75) as well in cell junctions (111). All these results together suggested that the MPL from sage and lemon verbena might promote a positive effect on intestinal cell proliferation, a process that is essential for the maintenance of the integrity and health of the gastrointestinal tract (112). In line with this idea, no signs of tissue damage were observed at histological level either pathways involved in cell apoptosis were found in these animals.

**CONCLUSIONS**

The present study shows that phytogenics obtained from sage and lemon verbena included at 0.1% in diets with a low FM content (7%), promoted transcriptional innate and adaptive immune responses in gut, especially through the modulation of those processes involved in T cell activation, differentiation and selection. Furthermore, the evaluated feed additive increased the number of intestinal goblet cells and modified the glycosylation properties of lectins from mucins. These changes resulted in a moderate increase in sialic acid residues, which also supports the idea that phytogenics from sage and common verbena might enhance gut immunity. Overall, this study shows that the evaluated phytogenic can be used as a safe feed additive for gilthead seabream, since its immunomodulatory properties were observed without compromising gut homeostasis and integrity of the intestinal epithelium.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article [Supplementary Material](#).

**ETHICS STATEMENT**

The animal study was reviewed and approved by Guiding Principles for Biomedical Research Involving Animals (EU2010/63), the guidelines of the Spanish laws (law 32/2007 and RD 1201/2015), and authorized by the Ethical Committee of the Institute for Research and Technology in Food and Agriculture (IRTA, Spain).

**AUTHOR CONTRIBUTIONS**

Conceptualization, EG. Methodology, EV-V, FER-L, CS, JO-D, RS. Formal analysis, RS, FER-L, JF, EV-V. Resources, EG, CS. Writing original draft, RS. Writing review and editing, EV-V, RS, FER-L, EG, JF, LT. Visualization, RS, FER-L, EV-V, JO-D, EG. Supervision, EG, EV-V. Project administration, EG. Funding acquisition, EG. All authors contributed to the article and approved the submitted version.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2021.670279/full#supplementary-material

**REFERENCES**

1. Stentiford GD, Bateman IJ, Hinchliffe SJ, Bass D, Hartnell R, Santos EM, et al. Sustainable Aquaculture Through the One Health Lens. *Nat Food* (2020) 1:468–74. doi: 10.1038/s43016-020-0127-5

2. Béné C, Arthur R, Norbury H, Allison EH, Beveridge M, Bush S, et al. Contribution of Fisheries and Aquaculture to Food Security and Poverty Reduction: Assessing the Current Evidence. *World Dev.* (2016) 79:177–96. doi: 10.1016/j.worlddev.2015.11.007

3. Little DC, Young JA, Zhang W, Al Mamun A, Murray FJ. Strategies for Fish Diseases. *Rev Aquac* (2020) 12:943. doi: 10.1111/raq.12492

4. Ghamkhar R, Hicks A. Comparative Environmental Impact Assessment of Oil Free Diets. *Ann Anim Sci* (2018) 18:338–54. doi: 10.1016/j.annals.2017.12.033

5. Gisbert E, Fournier V, Solovyev M, Skalli A, Andree KB. Diets Containing Antibiotic Use in Aquaculture, Policies and Strategies for the development and improvement of fish larvae production in Ibero-America” (117RT0521) funded by the Ibero-American Program of Science and Technology for Development (CYTED, Spain).
66. Collieran RS, Collins PE, O’Carroll C, Ahmed A, Mao X, McManus B, et al. Deubiquitination of NF-xb by Ubiquitin-Specific Protease-7 Promotes Transcription. Proc Natl Acad Sci USA (2013) 110:618–23. doi: 10.1073/ pnas.1204461110
67. Li Z, Hao Q, Luo J, USP4 Inhibits p53 and NF-xb Through Deubiquitinating and Stabilizing HDAC2. Oncogene (2016) 35:2902–12. doi: 10.1038/ onc.2015.349
68. Regula KM, Enk S, Kirshenbaum LA. IKK Beta is Required for Bel-2-mediated NF-Kappa B Activation in Ventricular Myocytes. J Biol Chem (2002) 277:38676–82. doi: 10.1074/jbc.M206175200
69. Chang MX, Xiong F, Wu XM, Hu YY. The Expanding and Function of NLRC3 or NLRC3-like in Telost Fish: Recent Advances and Novel Insights. Dev Comp Immunol (2021) 114:103859. doi: 10.1016/j.devz.2020.103859
70. El Aidy S, Derrien M, Aardema R, Hooiveld G, Richards SE, Dane A, et al. Transient Inflammatory-Like State and Microbial Dysbiosis are Pivotal in Establishment of Mucosal Homeostasis During Colonisation of Germ-Free Mice. Benef Microbes (2015) 6:57–77. doi: 10.3920/BM2013.0018
71. dos Santos N, Vale A, Reis M, Silva M. Fish and Apoptosis: Molecules and Pathways. Curr Pharm Design (2008) 14:14–69. doi: 10.2174/13816120878378743
72. Günther C, Buchen B, He GW, Hornel M, Torow N, Neumann H, et al. Caspase-8 Controls the Gut Response to Microbial Challenges by Trif-α-Dependent and Independent Pathways. BMJ J Gut (2015) 64:601–10. doi: 10.1136/gutjnl-2014-307226
73. Sun S, Ge X, Zhu J, Zhang W, Zhang Q. Molecular Cloning, Immunohistochemical Localization, Characterization and Expression Analysis of Caspase-8 From the Blunt Snout Bream (Megalobrama amblepuscula) Exposed to Ammonia. Fish Shellfish Immunol (2015) 47:645–54. doi: 10.1016/j.fsi.2015.10.016
74. Neufert C, Pickert G, Zheng Y, Wittkopf N, Warntjen M, Nikolaev A, et al. Motif in Chitinase 3-Like 1 (CHI3L1/YKL-40) Specifically Activates Akt Signaling Pathway in Colonic Epithelial Cells. J Cell Sci (2013) 126:3739–48. doi: 10.1242/jcs.111335
75. Glickman MH, Chechanover A. The Ubiquitin-Proteasome Proteolytic Pathway: Destruction for the Sake of Construction. Physiol Rev (2002) 82:373–428. doi: 10.1152/physrev.00027.2001
76. Chen CC, Liado V, Eirich K, Tran HT, Mizoguchi E. Protein 4.1R Links E-Cadherin/Catenin and Modulates Adherens Junction Integrity. J Proteome Res (2006) 5:2123–8. doi: 10.1021/pr060216f
77. Yang S, Guo X, Debnath G, Mohandas N, An X. Protein 4.1R Links E-Cadherin/β-Catenin Complex to the Cytoskeleton Through Its Direct Interaction With β-Catenin and Modulates Adherens Junction Integrity. Biochim Biophys Acta (BBA) - Biomembr (2009) 1798:1458–65. doi: 10.1016/ j.bbamem.2009.03.022
78. Paria A, Deepika A, Sreedharan K, Makesh M, Chaudhari A, Purushothaman P. Dual Functions of Rap1 are Crucial for T-cell Homeostasis and Prevention of Spontaneous Colitis. In: M Grosell, AP Farrell, CJ Brauner, editors. Fish Physiology, London, UK: Academic Press (2011). p. 111–34. doi: 10.1016/S1546-5098(10)30003-7
79. Marjomar JR, Lessey EC, Burridge K. Regulation of RhoA Activity by Adhesion Molecules and Mechanotransduction. Curr Mol Med (2014) 14:199–208. doi: 10.2174/156652414666612018104341
80. Devash M, Nier M, Matter K, Balda MS. Rho Signaling and Tight Junction Functions. Physiol (2010) 25:16–26. doi: 10.1152/physiol00034.2009
81. Citala-Madrid AF, Garcia-Ponce A, Vargas-Robles H, Betanzos A, Schnoor M. Small GTPases of the Ras Superfamily Regulate Intestinal Epithelial Homeostasis and Barrier Function Via Common and Unique Mechanisms. Tissue Barriers (2013) 1:5x26938. doi: 10.4161/tiss.26938
82. Yang S, Guo X, Mohandas N, An X. Protein 4.1R Links E-Cadherin/β-Catenin Complex to the Cytoskeleton Through Its Direct Interaction With β-Catenin and Modulates Adherens Junction Integrity. Biochim Biophys Acta (BBA) - Biomembr (2009) 1798:1458–65. doi: 10.1016/j.bbamem.2009.03.022
83. Sun S, Ge X, Zhu J, Zhang W, Zhang Q. Molecular Cloning, Immunohistochemical Localization, Characterization and Expression Analysis of Caspase-8 From the Blunt Snout Bream (Megalobrama amblepuscula) Exposed to Ammonia. Fish Shellfish Immunol (2015) 47:645–54. doi: 10.1016/j.fsi.2015.10.016
84. Huang JY, Lee SM, Mazmanian SK. The Human Commensal Bacteroides Oligonucleotide Translocator (Bot) Regulates the Intestinal Microbiota. J Cell Biol (2018) 217:97529–54. doi: 10.1083/jcb.201804031
85. Chen VC, Li X, Perreault H, Nagy J, Li X, Huan SC, Hartenstein JS, Benz EJ. Characterization of the Interaction Between Protein 4.1R and ZO-2: A Possible Link Between the Tight Junction and the Actin Cytoskeleton. J Cell Biol (2000) 275:30573–85. doi: 10.1083/jcb.2004578200
86. Ebnet K. Junctional Adhesion Molecules (Jams): Cell Adhesion Receptors With Pleiotropic Functions in Cell Physiology and Development. J Cell Sci (2008) 121:101. doi: 10.1242/jcs.209361
87. Chen VC, Li X, Perreault H, Nagy J. Interaction Molecules and Pathways. Curr Pharm Design (2006) 12. doi: 10.1016/j.cps.2018.03.022
88. Devash M, Nier M, Matter K, Balda MS. Rho Signaling and Tight Junction Functions. Physiol (2010) 25:16–26. doi: 10.1152/physiol00034.2009
89. Citala-Madrid AF, Garcia-Ponce A, Vargas-Robles H, Betanzos A, Schnoor M. Small GTPases of the Ras Superfamily Regulate Intestinal Epithelial Homeostasis and Barrier Function Via Common and Unique Mechanisms. Tissue Barriers (2013) 1:5x26938. doi: 10.4161/tiss.26938
90. Paria A, Deepika A, Sreedharan K, Makesh M, Chaudhari A, Purushothaman P. Dual Functions of Rap1 are Crucial for T-cell Homeostasis and Prevention of Spontaneous Colitis. In: M Grosell, AP Farrell, CJ Brauner, editors. Fish Physiology, London, UK: Academic Press (2011). p. 111–34. doi: 10.1016/S1546-5098(10)30003-7
91. Cain K, Swan C. Barrier Function and Immunity. In: M Grosell, AP Farrell, CJ Brauner, editors. Fish Physiology, London, UK: Academic Press (2011). p. 111–34. doi: 10.1016/S1546-5098(10)30003-7
106. Dey-Guha I, Mukhopadhyay M, Phillips M, Westphal H. Role of Ldb1 in Adult Intestinal Homeostasis. *Int J Biol Sci* (2009) 5:686–94. doi: 10.7150/ijbs.5.686

107. Spit M, Koo BK, Maurice MM. Tales From the Crypt: Intestinal Niche Signals in Tissue Renewal, Plasticity and Cancer. *Open Biol* (2018) 8:180120. doi: 10.1098/rsob.180120

108. Nie H, Ju H, Fan J, Shi X, Cheng X, Cang X, et al. O-GlcNAcylation of PGK1 Coordinates Glycolysis and TCA Cycle to Promote Tumor Growth. *Nat Commun* (2020) 11:36. doi: 10.1038/s41467-019-13601-8

109. Fritsch SD, Weichhart T. Metabolic and Immunologic Control of Intestinal Cell Function by Mtor. *Int Immunol* (2020) 32:455–65. doi: 10.1093/intimm/dxaa015

110. Sundblad V, Quintar AA, Morosi LG, Niveloni SI, Cabanne A, Smecul E, et al. Galectins in Intestinal Inflammation: Galectin-1 Expression Delineates Response to Treatment in Celiac Disease Patients. *Front Immunol* (2018) 9:379. doi: 10.3389/fimmu.2018.00379

111. Cai J, Culley MK, Zhao Y, Zhao J. The Role of Ubiquitination and Deubiquitination in the Regulation of Cell Junctions. *Protein Cell* (2018) 9:754–69. doi: 10.1007/s13238-017-0486-3

112. Wong WM, Wright NA. Cell Proliferation in Gastrointestinal Mucosa. *J Clin Pathol* (1999) 52:321–33. doi: 10.1136/jcp.52.5.321

**Conflict of Interest:** JQ and JP are current NATAC BIOTECH S.L. employers. FER-L was Senior research associate by Consorcio Tecnológico de Sanidad Acuícola, Ictio Biotechnologies S.A.

The remaining 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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