The Lipid-Lowering Effect of Dietary Taurine in Orange-Spotted Groupers (Epinephelus coioides) Involves Both Bile Acids and Lipid Metabolism

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An 8-week feeding trial was conducted to investigate how dietary taurine supplementation attenuates the lipid deposition induced by dietary high lipid in juvenile orange-spotted grouper (Epinephelus coioides). Three isonitrogenous (47% crude protein) semipurified diets were formulated to contain two levels of lipid and termed as 10% lipid diet, 15% lipid diet, and 15% lipid with 1% taurine (namely, diet 10L, diet 15L, and diet 15L + T, respectively). Groupers fed diet 15L + T showed higher weight gain and feed efficiency compared with diet 15L. Groupers fed diet 15L showed higher liver lipid contents, plasma total cholesterol (TC), and leptin contents vs. those fed diet 10L. Diet 15L + T decreased hepatosomatic index, liver lipid content, and plasma TC and adiponectin contents, and increased liver 3-hydroxy-3-methylglutaryl coenzyme A reductase content compared with diet 15L. Fish fed diet 15L had higher contents of chenodeoxycholic acid, deoxycholic acid, and lithocholic acid, and lower contents of glycodeoxycholic acid (GDCA), glycolithocholic acid, glycodeoxycholic acid, taurodeoxycholic acid (TDCA), and β-, γ-, and ω-muricholic acid (MCA) when compared with fish fed diet 10L. Diet 15L + T downregulated the contents of glycocholic acid, glycochenodeoxycholic acid, tauromuricholic acid, TDCA, ursodeoxycholic acid, GDCA, and β-MCA vs. diet 15L. Diet 15L upregulated expression of peroxisome proliferator-activated receptor α (pparaα) gene but downregulated expression of acyl-CoA carboxylase (acc), fatty acid synthase (fas), and glucose-6-phosphate dehydrogenase (g6pd) genes in comparison with diet 10L. The gene expression level of fas and g6pd was downregulated and the pparaα gene expression level was upregulated in fish fed diet 15L + T compared with those in fish fed diet 15L. Overall, this study indicated that dietary taurine supplementation can attenuate the liver lipid deposition of groupers caused by feeding 15% lipid through accelerating lipid absorption of taurine-conjugated bile acids and fatty acid β-oxidation and inhibiting lipogenesis.

Keywords: taurine, high lipid diet, bile acid profile, lipid metabolism, liver lipid deposition, Epinephelus coioides
INTRODUCTION

Taurine (2-aminoethanesulfonic acid), as a free amino acid, does not participate in protein synthesis but is known to play a wide range of key roles in animal physiology, such as immune regulation, osmoregulation, antioxidation, nervous system development and regeneration, and bile acid (BA) conjugation (Huxtable, 1992; Salze and Davis, 2015; Wu, 2020; Xu et al., 2020). BAs are regarded as biosurfactants, which emulsify lipid into micelles to facilitate its digestion and absorption by allowing for more cleavage sites for lipase (Suga et al., 2019; Romano et al., 2020). However, the emulsification of long-chain triglycerides (TGs) is achieved by conjugated BAs, which are better emulsifiers than BAs due to their hydrophilic and lipophilic properties (Gupta and Kim, 2003). It was reported that most primary BAs are linked with taurine or glycine to form conjugated BAs (Chiang and Ferrell, 2020), and taurocholic acid (TCA) or taurodeoxycholic acid (TDCA) formed by taurine conjugating with BAs is predominant in BAs for most teleost fish except common carp (Cyprinus carpio) (Kim et al., 2007, 2015). Previous studies demonstrated that dietary taurine addition increased taurine-conjugated BAs contents in rockfish (Sebastes schlegeli) and tiger puffer (Takifugu rubripes) (Kim et al., 2015; Xu et al., 2020). Thus, taurine may regulate lipid metabolism through accelerating BAs to form taurine-conjugated BAs.

It is clear that dietary taurine supplementation can reduce peripheral cholesterol and visceral fat contents of diabetic rats (Tsuboyama-Kasaoka et al., 2006) and obese humans (Brons et al., 2020). Thus, taurine may regulate lipid metabolism through accelerating BAs to form taurine-conjugated BAs and even fatty liver syndrome caused by the widespread use of high-lipid feed in intensive aquaculture for protein-sparing and system development and regeneration, and bile acid (BA) conjugation (Huxtable, 1992; Salze and Davis, 2015; Wu, 2020; Xu et al., 2020). BAs are regarded as biosurfactants, which emulsify lipid into micelles to facilitate its digestion and absorption by allowing for more cleavage sites for lipase (Suga et al., 2019; Romano et al., 2020). However, the emulsification of long-chain triglycerides (TGs) is achieved by conjugated BAs, which are better emulsifiers than BAs due to their hydrophilic and lipophilic properties (Gupta and Kim, 2003). It was reported that most primary BAs are linked with taurine or glycine to form conjugated BAs (Chiang and Ferrell, 2020), and taurocholic acid (TCA) or taurodeoxycholic acid (TDCA) formed by taurine conjugating with BAs is predominant in BAs for most teleost fish except common carp (Cyprinus carpio) (Kim et al., 2007, 2015). Previous studies demonstrated that dietary taurine addition increased taurine-conjugated BAs contents in rockfish (Sebastes schlegeli) and tiger puffer (Takifugu rubripes) (Kim et al., 2015; Xu et al., 2020). Thus, taurine may regulate lipid metabolism through accelerating BAs to form taurine-conjugated BAs.

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of 270 juvenile groupers (36.30 ± 0.09 g/fish) were randomly allocated into nine fiberglass tanks of recirculating system, and three replicates (30 fish per replicate) were assigned to each treatment. Groupers were hand-fed to apparent satiation two times daily at 8:30 and 18:00 over 8 weeks. Feces and excess feed were removed by siphoning 30 min after each meal. Excess feed was collected, dried at 65°C, and weighed for use in the calculation of FI. During the experimental period, the water temperature was 28.0 ± 0.5°C, dissolved oxygen level was 6.1 ± 0.2 mg/l, and ammonia nitrogen level was 0.21 ± 0.03 mg/l.

At the end of the feeding trial, groupers in each tank were batch-weighed and recorded on a wet weight basis after fasting for 24 h to determine the weight gain (WG) and feed efficiency (FE). Blood samples were taken from the caudal vasculature of 10 fish per tank by a sterile 2 ml heparinized syringe and centrifuged at 1,027 g for 10 min at 4°C to obtain plasma. The plasma samples in each tank were pooled and stored at −80°C for biochemical indices. After drawing blood, the bile of gallbladders was then collected and pooled by tank, and frozen immediately in liquid N2 for BA profiles analysis. Livers were collected from the same fish and weighed to calculate hepatosomatic index (HSI), pooled by tank and frozen immediately in liquid N2, and stored at −80°C for the analysis of lipid and enzyme contents, and the expression of genes related to lipid metabolism.

Bile Acid Profile Analysis
An aliquot of 500 µl precooled methanol and 20 µl internal standard solution (Sigma-Aldrich, Mosby, MO, United States) were added into the mixed solution of 10 µl aliquot of BAs sample and 90 µl precooled water and then mixed homogeneously via the vortex. After centrifugation (14,000 g, 4°C, 15 min), the supernatant was collected and concentrated by vacuum drying. The concentrates were re-dissolved in 50% methanol-water solution, and resultant supernatants were collected after centrifugation (14,000 g, 4°C, 15 min) for subsequent analysis of BAs.

Waters ACQUITY UPLC I-Class (Waters Corporation, Milford, Massachusetts, United States) and 5500 QTRAP mass spectrometer (SCIEX Corporation, Foster City, California, United States) with an electrospray ion source (ESI) source were used in the BA profile analysis. Briefly, the supernatants were separated using an ACQUITY UPLC BEH C18 (1.7 µm, 2.1 mm × 100 mm column), and the samples were placed in an 8°C automatic sampler with column temperature of 45°C, and the flow rate was 300 µl/min and the injection volume was 2 µl. The 0.1% formic acid-water solution and methanol were used as mobile phases A and B, respectively. The correlation gradient was programed as follows: 0–6 min from 60% B to 65% B, 6–13 min from 65% B to 80% B, 13–13.5 min from 80% B to 90% B, and 13.5–15 min, B phase maintained at 90%. The mass spectrometer was operated in the ESI negative mode under the following conditions: source temperature: 550°C; ion source gas1 (Gas1): 55; ion source gas2 (Gas2): 55; curtain gas (CUR): 40; Ion Sapary Voltage Floating (ISVF): −4,500 V. The ion pair was detected by the multiple reaction monitoring mode. The chromatographic peak area and retention time were extracted using multiquant 3.02 (SCIEX Corporation, Foster City, California, United States). To determine the stability of data, a QC sample was run and analyzed every ten samples. A standard substance of BAs was used to correct retention time and identify metabolites. The stability of the system and reliability of data was proved by extracted-ion chromatogram spectra of the standard mixture and relative SD distribution of quality control (QC) samples (Supplementary Figures 1, 2). The contents of BAs were determined by a standard curve (Supplementary Figure 3).

Principal component analysis (PCA) and orthogonal partial least squares discriminant analysis (OPLS-DA) were performed after all BAs were scaled with unit variance (UV). PCA was performed to examine the intrinsic variation in the data set. OPLS-DA was applied to maximize the separation between samples. The confidence level for membership probability was considered to be 95% (observations at < 5% were considered outliers). The farther the distance between components, the greater the difference. The potential overfitting of the OPLS-DA model was checked by performing 200 permutation tests and visualization using a validation plot (Supplementary Figure 4). The biplots were used to present the correlation between BA molecule species and different dietary treatments. A threshold of corrected p-value < 0.05 and variable important in projection value > 1 were used to determine the differential BAs.

Proximate Composition Determination
The moisture, ash, crude protein, and lipid contents of diets were determined according to the method of the Association of Official Analytical Chemists (AOAC, 2006). The moisture content was determined by drying the samples to a constant weight at 105°C. The crude protein contents (N × 6.25) were assayed by the Kjeldahl method using Kjeltec™ 8400 Auto Sample Systems (Foss Tecator AB, Hoganas, Sweden). Crude lipid was measured via the Soxtec extraction method using a solvent extraction system (Soxtec Avanti 2050 Auto System, Foss Tecator AB, Hoganas, Sweden), and the ash content was determined after incineration in a muffle furnace at 550°C for 8 h. The liver lipid was extracted by chloroform-methanol (2:1, vol/vol) solution and determined gravimetrically after drying a 5 ml aliquot under nitrogen (Folch et al., 1957).

Plasma and Liver Biochemical Indices Analysis
High-density lipoprotein cholesterol (HDL-C), low-density lipoprotein cholesterol (LDL-C), TG, and TC of plasma were measured using commercial assay kits. Leptin (LEP) and adiponectin (ADPN) contents in plasma and 3-hydroxy-3-methylglutaryl coenzyme A reductase (HMGR) content in the liver were determined by using ELISA kits according to the manufacturer’s instructions. All the commercial kits were purchased from Jiancheng Bioengineering Institute (Nanjing, China).

Real-Time Quantitative PCR Analysis of Lipid Metabolism Genes in the Liver
Total RNA was extracted using Trizol reagent (Invitrogen, Carlsbad, United States) with quantity and quality assessed...
via NanoDrop ND-2000 spectrophotometer (NanoDrop Technologies, Waltham, United States) and 1.2% denaturing agarose gel electrophoresis. RNA samples with 260/280 absorbance ratio of 1.86–2.00 were treated with 30 µl recombinant DNase I (RNase-free) (Takara, Tokyo, Japan), after which reverse transcription was conducted immediately on 1,000 ng RNA sample to synthesize cDNA using the PrimeScript™ RT Reagent Kit (Takara, Tokyo, Japan) following the manufacturer's instructions. The cDNAs were stored at −80°C until quantitative real-time (qRT)-PCR analysis.

Real-time quantitative-PCR was performed to determine relative mRNA levels of four genes related to lipid metabolism, namely, fatty acid synthase (fas), acyl-CoA carboxylase (acc), glucose-6-phosphate dehydrogenase (g6pd), and peroxisome proliferator-activated receptor α (ppara) with primers designed by Primer Premier 5.0 (Supplementary Table 1). β-actin was used as a house-keeping gene after confirming its stability across the experimental treatments according to the previous study (Pfaffl et al., 2004). The amplification efficiency of primers was approximately 100% according to the specific genes standard curves generated from serial dilutions (Hanaki et al., 2014). The qRT-PCR was performed using a Light Cycler™96 (Roche, Rotkreuz, Switzerland) with a reaction system containing 10 µl of TB Green premix Ex Taq™ (Tli RNaseH Plus) (2×), 2 µl cDNA template, 0.8 µl forward and reverse primers (10 µM), and 6.0 µl diethyl pyrocarbonate-treated water. The program parameters were set as follows: 30 s at 95°C, 40 cycles at 95°C for 10 s, and 60°C for 30 s. The data were optimized using the comparative Ct (2−ΔΔCt) value method as described previously (Livak and Schmittgen, 2001).

Calculations and Statistical Analysis
Data transformation was performed the prior statistical analysis. Student’s t-test was applied for comparison of diet 10L vs. diet 15L and diet 15L vs. diet 15L + T. PCA and OPLS-DA were used to analyze the data and find the changes of metabolites using SIMCA-P + 14.1 software package. The plot of correlation analysis was drawn by EXCEL 2019. All the results are presented as means ± SEM (n = 3).

RESULTS

Growth Performance, Feed Utilization, Lipid Content, and Biometric Indices
As shown in Figure 1, FI was not affected by dietary treatments. There was no difference in WG, FE, and HSI between groupers fed diet 10L and diet 15L. Groupers fed diet 15L + T had significantly higher WG and FE, and lower HSI and liver lipid content when compared with those fed diet 15L. The liver lipid content was significantly higher in fish fed diet 15L than in those fed diet 10L.

Bile Acid Profile
Figure 2A showed that the TCA, TDCA, andtaurochenodeoxycholic acid (TCDCA) were the predominant BAs among groupers fed diet 10L, 15L, and 15L + T. The PCA score plot (Figure 2B) showed that components of BA molecule species in the bile were clustered as three groups and separated from each other with no overlap. The OPLS-DA score plots and biplots showed that the models of comparison of diet 10L vs. diet 15L and diet 15L vs. diet 15L + T were well-established (Figures 3A,D). Biplot showed the best contributor to different treatments (Figures 3B,E). All differential BAs in diet 10L vs. diet 15L and diet 15L vs. diet 15L + T are presented in Figure 3C. The contents of chenodeoxycholic acid (CDCA), deoxycholic acid (DCA), and lithocholic acid (LCA) significantly increased with dietary lipid increasing from 10 to 15%, and the glycglycodeoxycholic acid (GDCA), glycolithocholic acid (GLCA), glycoursodeoxycholic acid (GUDCA), TDCA, and β-, γ-, and ω-muricholic acid (MCA) contents were totally opposite. The diet taurine downregulated the contents of all differential BAs, namely, GDCA, glycocholic acid (GCA), glycochenodeoxycholic acid (GCDC), taurohydrodeoxycholic acid (THDCA), tauromuricholic acid (TMCA), taouroursodeoxycholic acid (TUDCA), Ursodeoxycholic acid (TUDCA), and β-MCA in comparison with taurine free diet at 15% lipid (Figure 3F). The GDCA and β-MCA contents were downregulated by increasing dietary lipid and taurine levels.

Plasma and Hepatic Biochemical Components
The plasma contents of TG, HDL-C, and LDL-C were not affected by dietary treatments (Figure 4). However, the plasma TC content was increased when the dietary lipid level was increased from 10 to 15%, and the value was decreased when dietary taurine was increased from 0 to 1% in the case of dietary 15% lipid level. About 1% dietary taurine significantly decreased plasma ADPN content and increased liver HMGR content at 15% lipid diet. There was no difference in plasma ADPN and liver HMGR contents between diet 10L and diet 15L. The plasma LEP content was increased by increasing dietary lipid from 10 to 15%, but not affected by dietary taurine levels.

Expression of Genes Related to Liver Lipid Metabolism
Groupers fed diet 15L showed a significantly lower expression level of acc when compared to those fed diet 10L (Figure 5). The expression levels of fas and g6pd genes were lower in groupers fed diet 15L than in those fed diet 10L, and higher than in those fed diet 15L + T. However, the ppara gene showed the opposite expression change. The dietary taurine addition in 15% lipid feed did not affect acc gene expression.

DISCUSSION
Many studies have shown that dietary taurine has a positive effect on the growth performance of fish (de Moura et al., 2018; Liu et al., 2018; Candebat et al., 2020). In this study, compared to groupers fed taurine free diet (15L), fish fed diet containing 1% taurine (diet 15L + T) showed a significantly higher growth rate, as observed in juvenile Cobia (Rachycentron canadum), European
Results in yellowtail (Seriola quinqueradiata) and red sea bream (Pagrus major) showed that dietary taurine addition improved the FI (Matsunari et al., 2005; Matsunari et al., 2008). Although FI was not affected by dietary taurine levels at 15% lipid in this study, the trend of FE in response to dietary taurine addition was similar to WG. This finding indicated that taurine promotes the growth rate of grouper via improving feed utilization. In addition, the growth rate and feed utilization were not affected by dietary lipid levels in the absence of taurine. These results disagreed with those that occurred in meager (Argyrosomus regius) and silver barb (Puntius goniotous), i.e., high lipid diet led to a reduced growth rate (Chatzifotis et al., 2010; Nayak et al., 2018). The contradictory result may be due to different fish species and rearing environments, etc.

Previous studies showed that liver lipid content in Nile tilapia (Oreochromis niloticus) increased with increasing dietary lipid levels (He et al., 2015; Zhou et al., 2019), which was consistent with our current study results. We observed no variations in HSI and plasma TG, HDL-C, and LDL-C contents between groupers fed diet 10L and diet 15L. Plasma components could reflect the absorption, transportation, utilization, and deposition of lipids (Xu et al., 2016). Thus, maybe the grouper has a relatively high tolerance to dietary high lipid under the conditions of this experiment, which supported our result of growth performance. However, decreased HSI and liver lipid content were observed in 15% lipid feed with 1% taurine addition in comparison with 15% lipid feed without taurine addition. This finding supported previous observations on turbot (Scophthalmus maximus), Nile tilapia, and yellowfin seabream (Acanthopagrus latus) (Yun et al., 2012; Michelato et al., 2018; Dehghani et al., 2020), indicating a lipid-lowering effect of taurine in grouper. Furthermore, we also determined plasma hormones contents to find clues to the change of liver lipid content in the presence of taurine in high lipid feeds. Both ADPN and LEP are hormones secreted...
by adipose tissue and released into the bloodstream. LEP can reduce lipid accumulation by directly inhibiting lipid synthesis in adipocytes (de Gortari et al., 2020). ADPN can enhance the decomposition and energy consumption of fatty acids, and reduce the TG contents in the muscle and liver of large yellow croaker (Larimichthys crocea) (Ji et al., 2020). However, it was also reported that ADPN acts as the feedback signaling regulator of energy expenditure and metabolism (O’Brien et al., 2018). In the current study, groupers fed diet 15L + T showed lower plasma ADPN content than those fed diet 15L, indicating a decrease in liver lipid deposition in groupers. Meanwhile, consistent with our result, the increased plasma LEP content of mice was found to be accompanied by HSI, as a result of the high-lipid FI (Zhao et al., 2020). Dietary taurine reduced the secretion of LEP in rats and plasma LEP content in M. albus (Kim et al., 2012; Li, 2016), which was not observed in this study. However, the associations of dietary taurine with hormone secret and transport were not well-clarified so far, which needs further study.

Cholesterol is an important component of blood lipids. In this study, plasma TC content was upregulated by a 15% lipid diet without taurine addition but downregulated by a 15% lipid diet with taurine addition, as evidenced with increased liver lipid content in this study and previous studies in tiger puffer and European sea bass (Xu et al., 2020; Martins et al., 2021). Thus, taurine could reduce the risk of atherosclerosis by removing cholesterol (Gupta and Kim, 2003). We also observed an upregulated of liver HMGR content of groupers fed a 15% lipid diet with taurine in comparison with a 15% lipid diet without taurine. It was demonstrated that HMGR is the key rate-limiting enzyme that catalyzes the synthesis of a precursor of cholesterol in the regulation of cholesterol synthesis (Mahboobi et al., 2018). Our results may indicate the promotion of cholesterol biosynthesis by taurine via HMGR (Xu et al., 2020). The liver lipid content was found to have a positive association with the plasma content of TC, HMGR, LEP, and ADPN (Figure 6), indicating lipid accumulation in the liver was the result of the combined action of enzymes, hormones, and plasma lipids (Zhao et al., 2020). Furthermore, cholesterol is an integral component of cell membranes and acts as precursors of some hormones and BAs (Abramo, 1989; Suga et al., 2019), which combines with taurine to form taurine-conjugated BAs, the major BAs in most teleost fish (Kim et al., 2015). Thus, The BA profile of groupers was determined in this study.

Bile acids (BAs), amphiphilic (hydrophilic and lipophilic) molecules, are steroids biosynthesized from cholesterol in the liver and stored in the gallbladder, and released into the intestinal lumen after food ingestion (Nguyen et al., 2018; Suga et al., 2019). BAs can effectively emulsify lipids to form small chylous particles, which increase the contact surface between lipid molecules and lipase to accelerate the hydrolysis of lipid molecules in vivo (Ding et al., 2020). The BAs synthesis and its composition vary among animal species (Chiang and Ferrell, 2020). For example, 25 and 75% of primary BAs were conjugated with taurine and glycine in humans, respectively; while 95 and 5% of primary BAs are taurine-conjugated and glycine-conjugated in mice, respectively (Hofmann et al., 2010). However, it was previously believed that almost all BAs secreted by finfish were taurine-conjugated BAs (Kim et al., 2007; Kim et al., 2015). Here, a total of 29 BAs molecules in the bile of the grouper were detected and identified. The majority of them were taurine-conjugated BAs, but a small part of BAs are glycine-conjugated which was consistent with

![FIGURE 2](image-url)
FIGURE 3 | Orthogonal partial least squares discriminant analysis (OPLS-DA) score plots (A,D), biplot (B,E), and significantly different bile acids (C,F) of juvenile grouper fed the experimental diets. Values are means ± SEM (n = 3). Student’s t-test was applied for comparison of diet 10L vs. diet 15L and diet 15L vs. diet 15L + T. *, **, and *** represent significant differences with $P < 0.05$, $P < 0.01$, and $P < 0.001$. 
previous studies in other fish species (Garcia-Organista et al., 2019; Xu et al., 2020). It was reported that taurine-conjugated BAs have a high ability to remove cholesterol by enhancing serum HDL and suppressing LDL and very low-density lipoprotein concentration (Gupta et al., 2009). In this study, the three taurine-conjugated BAs such as TCA, TCDCA, and TDCA were the main components of BAs, and the contents of TDCA, GDCA, GLCA, and GUDCA were downregulated by the dietary lipid increasing from 10 to 15%, indicating an inhibiting of cholesterol metabolism, which agreed with plasma TC content discussed before. Interestingly, we also observed downregulation of GDCA, GCA, and GCDCA contents by dietary taurine addition at 15% lipid, which may indicate the higher competition of taurine combining with BAs than glycine in grouper and mice (Hofmann et al., 2010). In a study with Swiss female mice, CA promotes cholesterol absorption, but CDCA does not (Reynier et al., 1981). In our current study, CA content was not affected by dietary lipid level, but CDCA content was promoted by high lipid feed. Similar results were also observed in a recent study (Lu et al., 2018). The findings indicated that cholesterol absorption was limited. Moreover, CA and CDCA are efficacious endogenous ligands, whereas UDCA, MCA, and TMCA are antagonists of farnesoid X receptor (FXR) (Sayin et al., 2013; Su et al., 2019), which could inhibit TG synthesis by inducing small heterodimer partner in mice and grass carp (Ctenopharyngodon idella) (Li et al., 2013; Tian et al., 2021). In this study, β-MCA, γ- and ω-MCA contents were lower in groupers fed diet 15L than that in those fed diet 10L. Moreover, UDCA, β-MCA, and TMCA contents continued to decline when taurine content was increased from 0 to 1% in a 15% lipid diet. This means the activation of FXR
via reducing their antagonists could be triggered by taurine too. Notably, compared to taurine free diet at a 15% lipid level, taurine supplementation reduced the contents of THDCA and TUDCA. A previous study in humans demonstrated that the ratio of taurine to glycine-conjugated BAs is about 3:1 and this could be reduced by low taurine supply (Haslewood, 1978). Our results suggested that taurine addition at a 15% lipid diet could promote the taurine-conjugated BAs forming and play more important roles in lipid-lowering effects. Study results showed that most BAs through re-absorption in the intestinal lumen was routed to the liver where they were recycled to the gallbladder (Dawson, 2018). However, the re-absorption of BAs in the intestinal lumen was inhibited by dietary taurine in rats (Chen et al., 2003; Lam et al., 2006; Nishimura et al., 2009). This finding further supported the reduction of BAs caused by dietary taurine addition in our current study. Thus, taking liver lipid content and BAs profile into consideration (Figure 6), we speculate that taurine-conjugated BAs have a stronger ability to emulsify and absorb lipids than glycine-conjugated and other BAs in the fish species, which might be one of the reasons...
for the growth improvement. Interestingly, DCA and LCA, are hydrophobic cholic acids that have strong toxicity to colonic cells, they induce cell death involving either death receptor or mitochondrial (Rodrigues et al., 2004; Nguyen et al., 2018). The DCA and LCA contents were higher in groupers fed diet 15L than that in those fed diet 10L in this study and were found a positive correlation with liver lipid content (Figure 6), indicating the negative effect of high dietary lipid intake on the lipid metabolism of fish.

The mRNA expression levels of several genes related to both lipid metabolism and lipid accumulation to elucidate the molecular mechanisms underlying the taurine-mediated lipid-lowering effect on groupers. Acyl-CoA carboxylase is a cytosolic enzyme-producing malonyl-CoA, the first step in the biosynthesis of long-chain fatty acids (Qian et al., 2015). G6PD is a key enzyme related to the production of NADPH, and essential for de novo fatty acid biosynthesis catalyzed by FAS (Chen et al., 2013; Zheng et al., 2013). PPARα can modulate the expression of genes encoding several mitochondrial fatty acid-catabolizing enzymes in addition to mediating inducible mitochondrial and peroxisomal fatty acid β-oxidation (Aoyama et al., 1998). In this study, diet 15L upregulated the expression of the ppara gene but downregulated the expression of acc, fas, and 6pgd genes compared with diet 10L. The findings were consistent with previous results in blunt snout bream (Megalobrama amblycephala), hybrid grouper (Epinephelus lanceolatus♀ × Epinephelus fuscoguttatus♂), and large yellow croaker (Yan et al., 2015; Cao et al., 2019; Tan et al., 2019). Moreover, the expression levels of fas and 6pgd genes were decreased, and the expression level of the ppara gene was promoted with the dietary taurine increase. This indicated that liver lipid deposition reduction caused by dietary taurine addition may be the result of lipogenesis inhibition and fatty acid β-oxidation acceleration.

CONCLUSION

Under the present experimental conditions, the growth performance of the grouper was not affected by dietary lipid level increased from 10 to 15%, but significantly increased by dietary taurine addition at 15% lipid. High lipid diet increased the liver lipid content and decreased the contents of glycine-conjugated BAs and TDCA. Dietary taurine supplementation in a 15% lipid diet could decrease the lipid deposition in the liver of groupers through forming more taurine-conjugated BAs and triggering the activation of FXR, and inhibiting the expression of genes involved in lipid anabolism and accelerating fatty acid β-oxidation (Figure 6). Taurine-conjugated BAs have a higher ability to accelerate lipids emulsification and absorption than glycine-conjugated and other BAs in the fish species.

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 Animal Care and Use Committee of Fujian Agriculture and Forestry University.

AUTHOR CONTRIBUTIONS

XW: conceptualization, data curation, formal analysis, investigation, methodology, project administration, validation, visualization, and writing-review. FB: data curation, investigation, methodology, project administration, validation, visualization, and writing-review. JY: conceptualization, formal analysis, funding acquisition, resources, supervision, and writing-review. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

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