Temperature Affects Musculoskeletal Development and Muscle Lipid Metabolism of Gilthead Sea Bream (Sparus aurata)

Sara Balbuena-Pecino, Natàlia Riera-Heredia, Emilio J. Vélez, Joaquim Gutiérrez, Isabel Navarro, Miquel Riera-Codina and Encarnación Capilla*

Departament de Biologia Cel·lular, Fisiologia i Immunologia, Facultat de Biologia, Universitat de Barcelona, Barcelona, Spain

World population is expected to increase to approximately 9 thousand million people by 2050 with a consequent food security decline. Besides, climate change is a major challenge that humanity is facing, with a predicted rise in mean sea surface temperature of more than 2°C during this century. This study aims to determine whether a rearing temperature of 19, 24, or 28°C may influence musculoskeletal development and muscle lipid metabolism in gilthead sea bream juveniles. The expression of growth hormone (GH)/insulin-like growth factors (IGFs) system-, osteogenic-, myogenic-, and lipid metabolism-related genes in bone and/or white muscle of treated fish, and the in vitro viability, mineralization, and osteogenic genes expression in primary cultured cells derived from bone of the same fish were analyzed. The highest temperature significantly down-regulated igf-1, igf-2, the receptor igf-1ra, and the binding proteins igfbp-4 and igfbp-5b in bone, and in muscle, igf-1 and igf-1ra, suggesting impaired musculoskeletal development. Concerning myogenic factors expression, contrary responses were observed, since the increase to 24°C significantly down-regulated myod1 and mrf4, while at 28°C myod2 and myogenin were significantly up-regulated. Moreover, in the muscle tissue, the expression of the fatty acid transporters cd36 and fabp11, and the lipases lipa and lpl-lk resulted significantly increased at elevated temperatures, whereas β-oxidation markers cpt1a and cpt1b were significantly reduced. Regarding the primary cultured bone-derived cells, a significant up-regulation of the extracellular matrix proteins on, op, and ocn expression was found with increased temperatures, together with a gradual decrease in mineralization along with fish rearing temperature. Overall, these results suggest that increasing water temperature in this species appears to induce unfavorable growth and development of bone and muscle, through modulating the expression of different members of the GH/IGFs axis, myogenic and osteogenic genes, while accelerating the utilization of lipids as an energy source, although less efficiently than at optimal temperatures.

Keywords: thermal exposure, bone cells, white muscle, mineralization, GH/IGFs system, lipid catabolism
INTRODUCTION

Nowadays, society is facing one of the greatest world challenges: how to feed 9 thousand million people by 2050 in the context of global change and economic and financial uncertainty (1, 2). In this situation, aquaculture has a relevant role, satisfying the growing need of fish, and gilthead sea bream (Sparus aurata, L.) has become over the last 30 years one of the most important fish species farmed in the Mediterranean area (3). Besides food insecurity, climate change is also a major global challenge that concerns humanity. Warming of the climate system is unequivocal and particularly relevant for this study, the mean sea surface temperature may increase more than 2°C by the end of this century (4). Increased water temperature is known to directly influence several biochemical and physiological processes in ectothermic fish (5), including growth or metabolic rate (6).

Temperature can influence the growth hormone (GH)/insulin-like growth factors (IGFs) system, the main endocrine axis controlling growth in vertebrates. Previous studies have reported a link between environmental temperature and plasma levels of IGF-1 and GH, independently of the nutritional status (7, 8). The GH/IGFs system includes peptides (IGF-1 and IGF-2), IGF-1 receptors (IGF-IRα and IGF-IRβ), as well as is composed of six IGF binding proteins (IGFBPs) that can exert different effects on IGFs function depending on the cellular context (9, 10). Interestingly, IGFBP-2 in teleosts is the main circulatory binding protein and shows a physiological regulation similar to the most relevant one in mammals, IGFBP-3 (11). Furthermore, elevated temperatures, along with other risk factors such as mineral and vitamin deficiencies, light regimes or fast growth, have been linked in fish with increased occurrence of skeletal anomalies (12). In fact, fast muscle growth can exert high mechanical pressure on the developing bone, hence, synchronicity between bone and muscle growth is required for proper musculoskeletal development [reviewed by Ytteborg et al. (13)]. The higher prevalence of vertebral deformities as a result of increased temperature during the early stages of development has been documented in gilthead sea bream (14), as well as in other species including S. senegalensis (15), Salmo salar (16), or Dicentrarchus labrax (17). In the case of Sparids, the presence of abnormalities is more evident in larvae reared below 15°C and above 22°C (18), and recent studies have demonstrated in this species that thermal imprinting during embryogenesis causes long-term effects on bone physiology (19, 20). In this sense, the increase in temperature can be recognized as one important problem for aquaculture and animal welfare in a global climate change context.

Cellular and molecular mechanisms for musculoskeletal development in teleost fish have been demonstrated to be similar than in mammals. Osteoblasts, as well as myocytes, arise from mesenchymal stem cells (MSCs), precursor cells that are also able to differentiate into other cell types like chondroblasts or adipocytes after the coordinated induction of key transcription factors expression. Recently, morphological and molecular characterization of a bone-derived cell culture of gilthead sea bream has been reported (20, 21), and the ability of those MSCs to differentiate into other cell types such as adipocyte-like cells has been demonstrated (22). However, these multipotent cells have not been deeply characterized at a structural/functional level. Concerning the main regulators of bone development at a transcriptional level, Runt-related transcription factor 2 (Runx2) is the one required for commitment toward the osteogenic lineage. Afterwards, osteoblasts express molecules of the extracellular matrix (ECM), which include structural fibers as collagen or fibronectin but also non-collagenous proteins that regulate mineralization of the ECM such as osteonectin (ON), osteopontin (OP), and osteocalcin (OCN) (20). In the case of muscle, abundant studies using a satellite cell model system have properly characterized the process of myogenesis in gilthead sea bream (23). The coordinated expression of myogenic regulatory factors (MRFs) is also required for myogenesis to properly occur. Among these transcription factors, Myf5 and MyoD are involved in myocytes activation and proliferation, whereas Myogenin and MRF4 act later allowing myotube formation and maturation (24).

In addition to growth, increased water temperature is also known to directly affect energy demand in ectotherms, and consequently to exert an impact in lipid metabolism and the use of fat depots (25). Lipids are an important energy source for fish skeletal muscle. Lipases such as lipoprotein lipase (LPL) and lipase A (LIPA) can provide fatty acids from triglycerides circulating in the form of chylomicrons and very low-density lipoproteins. Then, fatty acid transporters such as CD36 and FATP1, which are nutritionally and hormonally regulated in fish muscle (26, 27), facilitate the entry of these fatty acids into the cell. Endogenous stored triglycerides, when necessary, can also be hydrolyzed by other lipases as the hormone-sensitive lipase (HSL). Then, non-esterified fatty acids undergo β-oxidation in the mitochondrial matrix (28).

In this framework, the aim of the present study was to evaluate the effects of three increasing rearing temperatures (19, 24, and 28°C) in gilthead sea bream juveniles through an in vivo/in vitro approach. First, the in vivo expression of GH/IGFs axis-, osteogenic-, myogenic- and lipid metabolism-related genes in bone and/or white muscle was determined, and then, the in vitro development and expression of osteogenic genes in primary cultured bone-derived cells. All this performed to extend the knowledge of the possible impacts of global climate change on musculoskeletal growth and the physiology in this important aquaculture marine species.

MATERIALS AND METHODS

Animals and Experimental Trial

Gilthead sea bream juveniles (50 g body weight), were obtained from Piscimar fish farm (Andromeda Group, Burriana, Spain) and maintained at the animal facilities of the Faculty of Biology at the University of Barcelona (Spain). After 2 weeks acclimation period, fish were randomly distributed into three 200 L glass tanks (11 fish per tank and condition) under a 12 h light/12 h dark photoperiod, at room temperature (19 ± 1°C). The experiment was performed in January. Fish were daily fed ad libitum twice with a commercial diet (Skretting, Burgos, Spain). At the beginning of the trial, all three tanks started at the same temperature of 19°C and then, two of them went from 19 to 24°C.
or 28°C, with a rate of Δ1°C each day following the protocol of Hevrøy et al. (6) with a 250 W thermostatic heater (EHEIM, Deizisau, Germany). Once the water temperature required was achieved, fish were held for 3 more days and sampled on the day fourth. A schema of the experimental trial is shown in Figure 1. The temperature of the tanks was registered with a precision thermometer (Sera®) three times a day to ensure the corresponding temperature was maintained. Before sampling, fish were fasted for 24 h and then were anesthetized (MS-222 150 mg/L) and subsequently sacrificed by a blow to the head. Samples of white muscle and vertebral bone were collected and immediately frozen in liquid nitrogen and stored at −80°C until further analyses, and small fragments of bone were also used to perform the primary cultures just after sampling as explained in section Primary Culture of Bone-Derived Cells.

All animal handling procedures were approved by the Ethics and Animal Care Committee of the University of Barcelona, in accordance with the guidelines of the European Union Council (86/609/EU), and the Spanish and Catalan Government assigned principles and legislation (permit numbers DAAM 6759 and 9336 for the in vitro and in vivo experiments, respectively).

Primary Culture of Bone-Derived Cells
Bone fragments of 10 fish per temperature condition were used per culture, each one considered an independent replicate, following the protocol of Capilla et al. (21). Briefly, the vertebrae were removed, cleaned of all adherent tissues and washed twice in phosphate buffered saline with 1% antibiotic/antimycotic solution (A/A) prior to manually obtain with a scalpel small (<1 mm) fragments. After that, two digestions of 30 and 90 min, respectively, were done with 0.125% Type II collagenase in Hank’s balanced salt solution at 18°C with gentle agitation. Next, the fragments were washed twice with Dulbecco’s Modified Eagle Medium (DMEM) supplemented with 1% A/A solution and finally cultured in complete growth medium composed of DMEM supplemented with 10% fetal bovine serum and 1% A/A solution. Cells were seeded into 6 or 12-well plates and incubated at 23°C and 2.5% CO₂. Medium was changed every 2 days. As indicated in the following sections, the fragments were removed from the plates at different days after seeding in order to perform the corresponding assays with the cells attached. To investigate whether temperature could affect the phenotype and differentiation of bone-derived cells in culture, pictures at days 8 and 15 of development were taken with a Canon EOS 1000D digital camera. All cell-culture reagents were purchased from Sigma–Aldrich (Tres Cantos, Spain) and all plastic items were obtained from Nunc (Labclinics, Barcelona, Spain).

Viability Assay
The methylthiazolyldiphenyl-tetrazolium bromide (MTT) assay was used to evaluate cell viability as previously done in gilt head sea bream bone cells (21) and muscle cells (29). This method was selected since comparative studies of different viability assays regularly used revealed that it is the most sensitive one in terms of detecting cytotoxicity (30). Briefly, on days 8 and 15 cell samples of two duplicate wells of the 12 well-plates were incubated for 3 h in DMEM with a final concentration of 5 mg/mL of MTT (Sigma–Aldrich). Then, cells were washed with phosphate buffered saline and resuspended in 150 µL of dimethyl sulfoxide (DMSO) per well. The viability values were obtained from the absorbance measured at 570 nm in duplicate 96-wells, with correction at 650 nm, using a microplate reader (Tecan Infinite 200). Data from day 15 cells are presented as fold change relative to each corresponding day 8 of culture (n = 10).

Mineralization Assay
Culture differentiation was evaluated according to mineralization of the ECM. The deposition of minerals was analyzed in day 20 cultured cells by alizarin red S (ARS) staining, following the protocol of Capilla et al. (21). Cells were fixed for 15 min with 10% formalin and stained with 2% ARS (pH 4.1–4.3) during 20 min. After washing excessive dye with water, quantification of the staining was done by means of acid extraction of the ARS stain with 10% acetic acid. The monolayer was then scrapped and transferred to a 1.5 ml tube. After vortex, the slurry was overlaid with mineral oil (Sigma–Aldrich), heated to 85°C for 10 min, cooled on ice and centrifuged at 16,000 g for 15 min. At this point, 10% ammonium hydroxide was added to the supernatant to neutralize the acid and finally, aliquots of the different samples were read at 405 nm in duplicate 96-wells, using a microplate reader (Tecan Infinite 200). Data are presented as optical density arbitrary units (n = 10).

Gene Expression Analyses
RNA Extraction and cDNA Synthesis
Total RNA was extracted from ~100 mg of vertebral bone and white muscle tissues, or from cell samples of two duplicate wells of the 6 well-plates at day 15 with 1 mL of TRI Reagent Solution (Applied Biosystems, Alcobendas, Spain) following the manufacturer’s instructions. Total concentration and purity were determined using a NanoDrop 2000 (Thermo Scientific,
Alcobendas, Spain), and integrity of the different samples was confirmed in a 1% agarose gel (w/v) stained with SYBR-Safe DNA Gel Stain (Life Technologies, Alcobendas, Spain). Next, 1,000 ng of total RNA were treated with DNase I (Life Technologies) to remove all genomic DNA, and reverse transcribed with the Transcriptor First Strand cDNA Synthesis Kit (Roche, Sant Cugat del Valles, Spain). The cDNA obtained was stored at −20°C for real-time quantitative PCR analyses (qPCR).

Real-Time Quantitative PCR

The mRNA transcript levels of the target genes plus three reference genes were examined in a CFX384TM real-time system (Bio-Rad, El Prat de Llobregat, Spain). All the analyses were performed in triplicate wells using 384-well plates with 2.5 µL of iTaq Universal SYBR Green Supermix (Bio-Rad), 250 nM final concentration of forward and reverse primers (Table 1) and 1 µL of diluted cDNA for each sample, in a final volume of 5 µL. As described before (31, 32), reactions were performed with an initial activation step of 3 min at 95°C, 40 cycles of 10 s at 95°C and 30 s at 55–68°C (primer-dependent, see Table 1) followed by an amplicon dissociation analysis from 55 to 95°C at 0.5°C increase each 30 s. Before the analyses, a dilution curve with a pool of samples was run to confirm primer efficiency, specificity of the reaction, absence of primer-dimers, and to determine the appropriate cDNA dilution for each assay. Negative controls [no template control (NTC), no reverse transcriptase control (RT C) and MilliQ water (PCR)] were included and run in duplicate. The expression level of each target gene analyzed was calculated using the Pfaffl method (33), relative to the geometric mean of the two most stable reference genes determined for each tissue by the geNorm algorithm, both implemented in the Bio-Rad CFX manager 3.1. software.

Statistical Analysis

Data were analyzed using IBM SPSS Statistics v. 22 (IBM, Armonk, USA) and are presented as Mean ± SEM. Data normality and homoscedasticity were tested among temperature groups using Shapiro–Wilks and Levene’s tests, respectively. Statistical significance was assessed by one-way analysis of variance (one-way ANOVA) followed by Tukey’s post-hoc test. When homoscedasticity was not observed Dunnett T3 test was applied. Statistical differences were considered significant for all analysis when p < 0.05.

RESULTS

HSPs and Proliferation Marker Genes Expression in Bone and White Muscle Tissues

In bone, heat shock proteins, hsp30 and hsp90b mRNA levels were similar among the three groups (Figure 2A), while the temperature of 24°C caused a significant increase in the gene expression of the proliferating cell nuclear antigen (pcna) respect to the other groups (Figure 2B).

In white muscle, hsp30 remained also unaltered, but hsp90b gene expression was highest at 28°C (Figure 2C). Concerning pcna, a significant up-regulation at 24°C compared with the 19°C condition as in bone was observed (Figure 2D).

GH-IGFs Axis-, Osteogenic-, and Myogenic-Related Genes Expression in Bone and White Muscle Tissues

In bone, the mRNA levels of igf-1, igf-2, igfbp-4, igfbp-5b, and igf-1ra were significantly down-regulated at 28°C compared with the 19°C reared fish and, in most cases, compared to the 24°C condition as well (Figures 3A,B). Contrarily, igfbp-1a, gh-1, gh-2, and igf-1b did not show differences among groups, although the former presented a tendency to gradually increase along with temperature. Concerning the osteogenic genes analyzed, none of them showed significant differences in response to temperature treatment under the experimental in vivo conditions tested (Figure 3C).

In white muscle, igf-1 mRNA levels were lower at 28°C, although differences were only significant compared to 24°C-exposed fish (Figure 3D). Moreover, igfbp-1a showed the same pattern as observed in bone tissue, significantly increasing its expression along with temperature. igf-2 and igfbp-4 did not revealed differences among groups, and igfbp-5b was not detectable in this tissue. Furthermore, the levels of expression of gh and igf-1 receptors were unaltered for gh-1 compared with the group of 19°C but were significantly lower for gh-2 and igf-1b at 24°C and for igf-1ra at 24 and 28°C (Figure 3E). With regards to the myogenic-related genes, the expression of myod1 and mrf4 was significantly lower in the 24°C-held fish compared to the other two groups, while myod2 and myogenin mRNA levels were significantly higher in the fish kept at 28°C, and neither muscle growth inhibitor, mstn1 nor mstn2, were affected by the rearing temperature (Figure 3F).

In addition, the expression of igfbp-2b was analyzed in both tissues, although none of them showed detectable levels.

Lipid Metabolism-Related Genes Expression in White Muscle Tissue

The fatty acid transporter cd36 was significantly up-regulated with the temperature rise, showing the fish at 28°C the highest mRNA levels. The expression of fatp1 remained unaltered, but fabp11 was significantly enhanced at 24 and 28°C compared with the 19°C group (Figure 4A). Concerning lipases, although hsl mRNA levels were not different among groups, lipa and lpl-ik transcript levels were significantly increased in 28°C, or 24 and 28°C-exposed fish, respectively, compared with the 19°C group. In parallel to this, significant up-regulation of the lipase maturation factor (lmtf1, an essential gene for the folding and assembly of LPL) was detected in the 28°C-reared fish (Figure 4B). Regarding β-oxidation markers, the gene expression of mitochondrial carnitine palmitoyltransferases (cpt1a and cpt1b) was significantly lower in juveniles maintained at 24 and 28°C when compared with fish reared at 19°C, while hydroxyacyl-CoA dehydrogenase hadh expression did not show significant differences among groups. Contrarily, the mitochondrial uncoupling protein ucp2 showed significantly higher mRNA levels with increased temperature (Figure 4C).
| Gene     | Primer sequences (5′-3′)                      | Tm, °C | Accession number |
|----------|-----------------------------------------------|--------|------------------|
| rps18    | F: GGGTGGTGGCGAGCGTTAC                       | 60     | AM490061.1       |
|          | R: CTCTGCGCTGGAGGAACCA                      |        |                  |
| ef1a     | F: CTCAACGGCTAGGTCATCAT                     | 60     | AF184170         |
|          | R: GCACAGGGAGAAGCGGAAGAGGGA                  |        |                  |
| rp27a    | F: AAAGGGAACACACATCTGCGCCCCAC               | 68     | AY188520         |
|          | R: GCTTGCCTTGGGCCAGAAACTTGTAG               |        |                  |
| hsp30    | F: GTGTGACGGGAAAAAGAGA                     | 60     | GU60312          |
|          | R: CTGAGGAGGAGGTGCTGTTC                     |        |                  |
| hsp90b   | F: TTAACGAGGAAGAGAQTGTG                    | 56     | DO012949         |
|          | R: GTGCCACACACACAGCATGAA                    |        |                  |
| pcna     | F: TGTTTGGCGACGCTGTGGT                     | 58     | NM_131404.2      |
|          | R: TGGCTAAGTTTGTCTGCG                      |        |                  |
| igf-1    | F: ACAGAATGAGGGACGGAGCAATGGAC               | 60     | EF688016         |
|          | R: TGGCCACTTGAGTGACCCTGCTGTG                |        |                  |
| igf-2    | F: TGGGATCTAAGGAGGAGGTGTTGT                | 60     | AY996778         |
|          | R: CTGAGGAGGAGGCGCGAC                      |        |                  |
| igfbp-1a | F: AGTGCGAGTCTCCTGQAT                     | 60     | KM522771         |
|          | R: TCTCCTTTAAGGCGACTCGGC                   |        |                  |
| igfbp-2a | F: CGGCCTCTCTGQACATCGC                     | 60     | AF377998         |
|          | R: GTCCCTGCGACCTCATTGT                    |        |                  |
| igfbp-4  | F: TCGCAAAACACAGAAAGCAGAA                   | 68     | F5T95C02JM29K    |
|          | R: GGAGTATGGGAGTTGTGAAAGA                   |        |                  |
| igfbp-5a | F: TTTGTTCTCCTGQGTGC                      | 60     | AM963285         |
|          | R: TCAAGATGCGGCTCCAG                      |        |                  |
| ghr-1    | F: ACCTGTCAGCCACACATGA                    | 60     | AF438176         |
|          | R: TCGTGCAGACCTGGGCTGTAG                   |        |                  |
| ghr-2    | F: GAGTAGAACCGGCCGTQACAG                   | 60     | AY573601         |
|          | R: GCCTGQGTATCTGTATTCTGAT                   |        |                  |
| igf-1ra  | F: AGCATCAAACACAGACCTGQG                   | 55     | KT156846         |
|          | R: CTCTGCTGAGAAGAAAGC                      |        |                  |
| igf-1rb  | F: GCTAATGCGAATGGTGTGQG                     | 55     | KT156847         |
|          | R: CGCTCTTATTGCTGCTGTAG                    |        |                  |
| runx2    | F: ACCGTCCTACCTGAGCTC                     | 60     | JX2352063        |
|          | R: AGAGAAGACCTGGCAATCGTC                   |        |                  |
| fb1a     | F: CGTGAAACTCAAGAATCGGTGAG                 | 60     | FG262933         |
|          | R: CGCATTTGTGACCTGCGCTTGG                 |        |                  |
| bmp2     | F: GAGAGAAGCGCGGCTGGTTAAGACQAAT             | 65     | AY500244         |
|          | R: GGCTCCTGCGCTGCAAACATATT                |        |                  |
| bmp4     | F: CAAGCCATTTCAGAGACT                   | 60     | FJ436409         |
|          | R: CCCTCCGACTACCTTCT                     |        |                  |
| mgp      | F: TGTTGAATTTATGTAGGTTGGCTTGCGATCTCC      | 68     | AY065652         |
|          | R: CGGCGGATAGTGAGAAATGGTTAGA               |        |                  |
| on       | F: AGAAGAGGGCTCAGCGAGAAGC                  | 68     | AY239014         |
|          | R: GTGCTGCGATGCGCGGATCTCTCA               |        |                  |
| op       | F: AAAACCCAGGGATAACTCTCAAGAGAAGCCA         | 68     | AY651247         |
|          | R: AGAACCGTGCCAAAGAGCAGAAGCAGAA           |        |                  |
| ocn      | F: TCCTGAGTGAGAACAGAGA                    | 56     | AF048703         |
|          | R: CGCTCCTGCTGAGAGGCGGTGAG               |        |                  |
| pax7     | F: ATGAAACCTGCTGGAAGCAGAGA                | 64     | JN034418         |
|          | R: AGCGCTGCGACACTTCTGAG                    |        |                  |

(Continued)
TABLE 1 | Continued

| Gene   | Primer sequences (5′-3′)                            | Tm, °C | Accession number |
|--------|-----------------------------------------------------|--------|------------------|
| myf5   | F: CTACGAAGACGAGTGGAAGACT                         | 64     | JN034420         |
|        | R: TGTTCTATCGCCACAAAGTCT                           |        |                  |
| myod1  | F: TTGGAGAAGCGGGGAGAGGACC                         | 60     | AF478568.1       |
|        | R: CTTCTCGGGGTGATAGGACC                           |        |                  |
| myod2  | F: CACTACGGGGGAGCTCGACC                           | 60     | AF478569         |
|        | R: CGTGTGCTCTGGACTCT                              |        |                  |
| myogenin | F: CAGAGGTGGCCAGAAGGTGAAGGAG                         | 68     | EF462191         |
|        | R: CAGGTGGCTGGCCAGAAGGTGAAGGACC                     |        |                  |
| mrf4   | F: CATCCACAGCTTTAAAGGCA                            | 60     | JN034421         |
|        | R: GAAGACGACCGAGAGATCT                              |        |                  |
| mstn1  | F: GTAGAGCTGCTGGAGAGACG                            | 60     | AF258448.1       |
|        | R: CGTAGAAGCTGCGAGCTAC                              |        |                  |
| mstn2  | F: ACCTGGTGAAACAAAGGAACAC                          | 60     | AY046314         |
|        | R: TGCCGGTAAAGACACATG                              |        |                  |
| cd36   | F: GCTGGTGGCTAGCTCCCA                               | 60     |                  |
|        | R: TTCCCGGGCTGCTGATTCC                              |        |                  |
| fatp1  | F: CAACAGAGGTGGAGAGGACAT                           | 60     |                  |
|        | R: GGGAAGTACAGGCAGAAACAC                          |        |                  |
| fabp11 | F: CATTGGAGAGACCGACCAGCT                          | 60     |                  |
|        | R: ACTTGAAGTGTGGCTACAT                             |        |                  |
| hsl    | F: GCTTTGCGCTTAGTTACACCTATTC                       | 60     | EU254478         |
|        | R: GATTGAGCGACCTCTGGAGATGT                        |        |                  |
| lipa   | F: TACCTACGGGAGACTCTCAAAGGAAC                      | 60     | JQ308831         |
|        | R: GTGGAAGAAGCTGATGACTGATC                         |        |                  |
| lpl-ik | F: CAGAGATGGAGCCTGACTC                            | 60     | JQ390609         |
|        | R: TGTGTCACAGCAGAGGAAAGATG                        |        |                  |
| lmf1   | F: CGGCTGGACTGCTGTACAT                             | 60     | JK975718         |
|        | R: CTGACACGCTGCTGTAGCTGAGGAGGAGG                  |        |                  |
| cpt1a  | F: GTGCGCTGCTCAGCAGGACC                          | 60     | JQ308822         |
|        | R: TGATGCTTAGCTGCTGGCTGAGGAGG                     |        |                  |
| cpt1b  | F: CCACGAGCGAGCTACAGCAG                           | 60     | DQ868621         |
|        | R: CACCACGCGCCACACATATTAG                         |        |                  |
| hadh   | F: GAACTCGCAGAAGGCCAAGCAGAAGG                      | 60     | JQ308829         |
|        | R: CTAAGAGGGGCTGACAAGTGACAT                       |        |                  |
| ucP2   | F: CGGCGCGCTCCTCAATGG                            | 60     | JQ599959         |
|        | R: AACCAAGGTGGCTCCCTTTGCTCAT                      |        |                  |

F, forward; R, reverse. rps18, ribosomal protein s18; elf1a, elongation factor 1 alpha; rps27a, ribosomal protein 27a; hsp30, heat shock protein 30; hsp90b, heat shock protein 90b; pcna, proliferating cell nuclear antigen; igf-1, insulin-like growth factor 1; igf-2, insulin-like growth factor 2; igf1bp-1a, insulin-like growth factor binding protein 1a; igf1bp-2b, insulin-like growth factor binding protein 2b; igf2bp-4, insulin-like growth factor binding protein 4; igf2bp-5b, insulin-like growth factor binding protein 5b; ghr-1, growth hormone receptor 1; ghr-2, growth hormone receptor 2; igf-1ra, insulin-like growth factor 1 receptor a; igf-1rb, insulin-like growth factor 1 receptor b; runx2, runt-related transcription factor 2; fib1a, fibronectin subunit 1a; bmp2, bone morphogenetic protein 2; bmp4, bone morphogenetic protein 4; mmp, matrix metalloproteinase; mmp1, matrix metalloproteinase 1; mmp3, matrix metalloproteinase 3; mmp4, matrix metalloproteinase 4; mmp9, matrix metalloproteinase 9; matrix gla protein; op, osteopontin; oss, osteocalcin; px, paired box 7; myf5, myogenic factor 5; myod1, myogenic differentiation 1; myod2, myogenic differentiation 2; myogenin, myogenin; mrf4, myogenic regulatory factor 4; mstn1, myostatin 1; mstn2, myostatin 2; cd36, cluster of differentiation 36; fatp1, fatty acid transport protein 1; fabp11, fatty acid binding protein 11; hsl, hormone sensitive lipase; lipase; lipase a; lipase b; lipoprotein lipase-1; lmf1, lipase maturation factor 1; cpt1a, carnitine palmitoyltransferase 1a; cpt1b, carnitine palmitoyltransferase 1b; hadh, hydroxacyl-CoA dehydrogenase; ucP2, uncoupling protein 2.

Characterization of the Bone-Derived Cells Culture

The effects of temperature during the differentiation of gilthead sea bream bone-derived cells are presented in Figure 5A. Morphologically, in all three groups, the cells showed at day 8 a spindle-like phenotype that changed to a polygonal one at day 15 as the cells differentiated spontaneously into osteoblasts. Moreover, deposits of minerals started to accumulate in the ECM, although mineral nodules were apparently in greater proportion in cells derived from the 19°C-reared fish compared to the other two groups. Besides, as shown in Figure 5B, significantly lower viability values were found in cells coming from fish reared at 24 and 28°C, compared with those at 19°C; and the same was observed concerning the deposition...
of minerals in the ECM (Figure 5C), which agreed with the visual observation.

**HSPs and Proliferation Marker Genes Expression in Bone-Derived Cells**

Exposure of gilthead sea bream juveniles to 24°C of temperature significantly increased the expression of hsp30 and hsp90b transcript levels in bone-derived cultured cells compared to those coming from the other fish (Figure 6A). In addition, a significant down-regulation of the proliferation marker pcna gene expression was detected in the 28°C-cells compared with those obtained from 24°C-held fish (Figure 6B).

**GH-IGFs Axis- and Osteogenic-Related Genes Expression in Bone-Derived Cells**

The analysis of GH-IGFs system-related genes expression in cultured bone-derived cells revealed no differences among groups (Figures 7A,B). Regarding the expression of osteogenic genes, significant differences were neither observed for most of them but significant up-regulation of on, op, and ocn gene expression was found in cells coming from gilthead sea bream maintained at elevated temperatures, compared with those cells from 19°C-reared fish (Figure 7C). To corroborate the determination of the bone-derived cultured cells toward the osteogenic lineage, the gene expression of pax7, one of the main transcription factors shaping the fate of MSCs into the muscular lineage, was analyzed, resulting undetectable.

**DISCUSSION**

As existing literature reports, an increment of water temperature has been proved to be a valid approach to evaluate impact of global climate change on physiological responses in various fish species (6, 15, 25, 34, 35). Besides, we have recently described in gilthead sea bream temperature-dependent differential expression of genes involved in osteogenesis, indicating a
modulation of bone formation caused by this abiotic factor (20). In the current study, the aim was to characterize the effects of increased temperature in gilthead sea bream juveniles’ musculoskeletal growth, muscle lipid metabolism and, in the in vitro development of primary cultured bone-derived cells to test the hypothesis that global climate change modulates the expression of key genes locally in bone and muscle, which might increase the occurrence of skeletal anomalies in this species.

Effects of Temperature on Cell Culture Development

The bone-derived cultured cells from gilthead sea bream vertebrae gave an initial homogenous population of cells. At the first stages, cells were mostly triangular as previously reported for this species. At day 15, cell differentiation became more evident, with the change into a polygonal shape, characteristic of the osteoblast phenotype. These changes followed the profile also reported for gilthead sea bream in vitro model when similar temperature changes were applied directly into the cells (20). Previous studies in larvae of sole and grass carp (Ctenopharyngodon idella) also described an increased expression of hsp90 in response to a temperature rise (39, 40), as a protective mechanism against thermal stress.

Although the gene expression of hsp90b was increased in vivo only in the muscle tissue under the highest temperature tested, the in vitro experiment showed direct evidence of water temperature up-regulating hsp30 and hsp90b; thus, supporting the stressful condition induced to the animals. A similar response was reported for both genes in the same gilthead sea bream in vitro model when similar temperature changes were applied directly into the cells (20). Previous studies in larvae of sole and grass carp (Ctenopharyngodon idella) also described an increased expression of hsp90 in response to a temperature rise (39, 40), as a protective mechanism against thermal stress. In fact, an increase in hsp30 mRNA levels was observed only after 3 h of an in vivo temperature increase in rainbow trout (Oncorhynchus mykiss) (41). Similar activations of chaperones involved in protein folding have been reported in muscle tissue of gilthead sea bream (42) and rainbow trout (43) facing another stressful situation such as fasting.

The proliferation marker pcna showed the same pattern of expression in vivo in bone and muscle than in vitro in bone-cultured cells, with increased mRNA levels in 24°C-reared fish respect to the other groups. Thus, despite the cells showed reduced viability with the increase in temperature, this up-regulation of pcna could be considered a compensatory response, attempting these cells to recover from their initial heat stress-related situation. Accordingly, that could be considered a compensatory growth mechanism in vivo. Thus, it appears that 24°C would be the limiting temperature to properly
FIGURE 4 | In vivo effects of temperature over the expression of lipid metabolism-related genes in white muscle. Relative mRNA expression normalized to rps18 and rpl27a of (A) fatty acid transporters cd36, fatp1, fabp11, (B) lipases hsl, lipa and lpl-k, lmf1, and (C) β-oxidation markers cpt1a, cpt1b, hadh, and ucp2. Data are shown as Mean ± SEM (n = 8). Significant differences among fish held at different temperatures were determined by one-way ANOVA and are indicated by different letters (p < 0.05). N.S., non-significant.

Effects of Temperature on GH/IGFs Axis-Related Genes Expression in vivo and in vitro

The expression of igf-1 was significantly decreased in both bone and muscle tissues of 28°C-held fish when compared with fish reared at 24 or 19°C, while its expression was not detectable in cultured osteoblasts. IGF-1 plays an important role inducing not only muscle differentiation and hypertrophy, but also bone matrix production (16, 45). The down-regulating effect of high temperature on igf-1 observed in gilthead sea bream has been also reported in muscle of different fish species (i.e., Atlantic salmon and southern flounder), as well as, in IGF-1 plasma levels (6, 46, 47), suggesting restricted growth. Notwithstanding, similar studies in rainbow trout showed contrarily, an increase in plasma GH and IGF-1 levels with high temperature (7, 8), overall suggesting that the response to temperature increase of the major growth factors in fish could be species-specific. Regarding igf-2, changes were not observed among groups in muscle and cultured bone cells, although the same response as igf-1 was observed in the bone in vivo, indicating that this tissue appears to be more sensitive to changes in temperature. This data is in agreement with that observed previously in unresponsive rainbow trout muscle, both at mRNA and plasma levels (8); however, in Atlantic salmon, Hevrøy et al. (6) found that igf-2 mRNA levels were significantly down-regulated in muscle and liver after 45 days of exposition to warm temperature but not after only 15 days. Thus, it cannot be excluded that a prolonged trial time could have affected the expression of this gene in the present study as well.

Concerning the GH and IGF-1 receptors, juveniles held at 24 and 28°C presented in muscle significantly lower levels of expression of igf-1ra than at 19°C; those reared at 24°C also had decreased igf-1rb and ghr-2, and in bone those at 28°C also showed reduced igf-1ra mRNA levels. Wargelius et al. (16) revealed that an increase in the gene expression of igf-1ra relates with an increase in bone density in Atlantic salmon. Thus, the decrease in igf-1ra expression observed in gilthead sea bream could lead to reduced mineralization in the long-term caused by the high rearing temperature, which would be in agreement with that observed by Ytteborg et al. (48) in the former species. Therefore, in this context, the results of the present study suggest that the GH/IGFs axis is influenced in gilthead sea bream by elevated temperature to locally decrease the expression of ghrs, igfs, and igf-1rs in bone and muscle in order to delay musculoskeletal growth. Interestingly, although differences exist between the in vivo and in vitro data, which could be due to modulation of the gene expression by systemic factors in the whole animal, the results obtained in the bone-derived cultured cells reflect this impaired situation as well.

With regards to the IGFBPs of major local action in the musculoskeletal tissues, the same increasing pattern in expression with temperature was observed concerning igfbp-1a in white
FIGURE 5 | (A) Representative images of cells derived from vertebrae bone of gilthead sea bream reared at different temperatures, at (a,c,e) day 8 and (b,d,f) day 15 of culture development. Magnification, 10x. Arrowheads indicate the presence of mineral nodules. (B) Quantification of viability in gilthead sea bream cultured bone-derived cells using an MTT assay presented as fold change of day 15 relative to day 8 of culture. (C) Quantification of mineralization in gilthead sea bream cultured bone-derived cells at day 20 determined by ARS staining. Data are shown as Mean ± SEM (n = 10). Different letters among temperature groups indicate significant differences, calculated by one-way ANOVA (p < 0.05).

FIGURE 6 | In vitro effects of fish rearing temperature over the gene expression of heat shock proteins and a proliferation marker in bone-derived cells at day 15 of culture development. Relative mRNA expression normalized to ef1a and rps18 of (A) hsp30 and hsp90 and (B) pcna. Data are shown as Mean ± SEM (n = 6–7). Significant differences among temperature groups gene were determined by one-way ANOVA and are indicated by different letters (p < 0.05).

muscle and bone, as previously found in Atlantic salmon muscle (6). Previous studies in zebrafish (Danio rerio) revealed that elevated expression of igfbp-1a limits cellular actions of IGF-1, being an important growth and developmental inhibitor (49). Moreover, this binding protein has been associated with stressful or negative conditions, since a strong relation with elevated serum cortisol levels has been reported (50, 51). In this context, the gradual increase of igfbp-1a along with temperature observed in this study, suggests impaired growth conditions in agreement with the reduced expression of igfs observed at
28°C. Furthermore, *igfbp-4* and *igfbp-5b* mRNA levels were also significantly decreased in bone tissue of 28°C-held fish, although remained unaltered in muscle and bone cells. In previous studies, these binding proteins have been reported as positive regulators of IGF-1 actions, with *igfbp-5b* promoting bone differentiation (43); and its mRNA levels being highly correlated with *igf-1* and *igf-2* in muscle (10, 32, 52, 53). Therefore, altogether, the down-regulation of *igfbp-4* and *igfbp-5b* in bone tissue and the increase of *igfbp-1a* in muscle of gilthead sea bream maintained at 28°C, indicates that such high temperature is an unfavorable condition, leading to reduced musculoskeletal growth and differentiation in this species.

**Effects of Temperature on Osteogenic- and Myogenic-Related Genes Expression in vivo and in vitro**

In the present study, although none of the osteogenic-related genes analyzed showed differences in vivo, increasing water temperature was sufficient to induce an up-regulating response on some genes involved in the mineralization of the ECM (*on*, *op*, and *ocn*) in the cultured bone-derived cells, suggesting that the thermal history can influence the developmental plasticity of the osteogenic process in vitro. Interestingly, the ECM glycoprotein ON has been reported as a heat shock protein having chaperone-like properties to prevent collagen denaturation (54, 55). Therefore, as it was observed in rainbow trout by Currie et al. (41) and in a previous study by our group using the same cellular system (20), the increase in *on* expression caused by changes in temperature could potentially represent an initial response of bone cells to stressful conditions. According to this, the increase in *on* mRNA levels may also suggest a negative effect for ECM production and mineralization, which agrees with the reduced number of deposited minerals in the ECM along with temperature in the bone-cultured cells of the present study. Moreover, the elevated expression of *op*, a well-known inhibitor of matrix mineral deposition (56) supports this improper mineralization of the bone when fish are maintained at high temperatures. However, the ability of OP to regulate this process depends on its state of phosphorylation (57), thus, further analyses should be done to confirm this hypothesis.

Concerning in vivo studies, after a long-term high-temperature treatment in Atlantic salmon, Ytteborg et al. (38) reported that the mRNA levels of *runx2* (the key transcription factor of osteogenesis), decreased when fish reached 15 g but not at 2 g of body weight; whereas other non-collagenic ECM molecules such as *ocn, on,* or *colla1* were down-regulated already at the 2 g stage. These authors proposed that these results might suggest a defect in the late maturation of osteoblasts, which agrees with the lower mineral density and shorter length-height proportion observed in the vertebrae of these animals (38). It also agrees with the subsequent significantly increased incidence of malformations found in the fish reared at high temperature at body weights of 15 and 60 g. Moreover, the same authors observed, in an in vitro study with precursor muscle cells differentiated into osteoblasts and cultured at an elevated temperature, a reduced expression of *ocn* and *colla1* (48). In gilthead sea bream, expression of osteogenic genes was modified by temperature in both embryo and larval stages, but in the juveniles, differences were only observed after producing a temperature challenge (20). In the same study, lower transcript levels of most of the osteogenic genes analyzed in cultured osteoblasts in response to a long-term treatment of increased temperature were reported. Overall, these data suggest that at a transcriptional level, the deleterious effects of temperature on bone development could depend on the time of exposure. Therefore, it cannot be discarded that a prolonged temperature treatment could have also affected the expression of osteogenic genes in gilthead sea bream in vivo, pointing out that osteoblast differentiation and bone ECM mineralization could be impaired.

The coordinated expression of MRFs to properly control muscle development can be modulated by temperature in teleost fish, thus affecting muscle growth (42, 58). In the present study, *myod1* transcript levels were reduced in the fish maintained at 24°C, while *myogenin* and *mrf4* expression was highest in the 28°C-reared fish, suggesting potentially slackened cell proliferation but enhanced myocyte differentiation with the increase in water temperature, overall uncoupling the myogenic process. In this framework, with high rearing temperatures, gilthead sea bream musculoskeletal growth would not be under harmonic conditions, which could be leading in the long-term to increased prevalence of bone deformities.
Effects of Temperature on Muscle Lipid Metabolism-Related Genes Expression in vivo

The increase of temperature, up-regulated in the present work the muscle expression of the fatty acid transporter and binding protein cd36 and fabp11, respectively, suggesting elevated fatty acid uptake and intracellular transport, upon high-temperature conditions. These data are in accordance with a recent study by Zoladz et al. (59), which reported enhanced protein expression of CD36 in rat skeletal muscle under hyperthermia. Nevertheless, the function of FABP11, which is probably an isof orm restricted to fishes, is not completely known (60). With regards to the endothelial enzymes with a triglycerides lipase activity, gilthead sea bream juveniles held at 28°C presented significantly higher mRNA levels of lipa and lpl-ik, an exclusive fish lineage isoform of LPL (59). Regulation of the latter enzyme is far from being established yet, although it is known that in skeletal muscle of gilthead sea bream, changes in LPL-like are correlated with LMF1 (61), an endoplasmic reticulum membrane protein involved in the post-translational folding and assembly of LPL, among other proteins (62). This agrees with the results of this study, where lmf1 mRNA levels were also increased as temperature was raised. Contrarily, the increase of temperature did not induce significant changes in hsl gene expression among groups, suggesting that fatty acids are mostly being uptaken by the muscle from circulating triglycerides, or non-esterified fatty acids provided by adipose tissue. Recent studies in Atlantic salmon and catfish (Pelteobagrus vachellii) demonstrated a reduction in hepatic triglycerides and relative viscera weight during exposure to elevated temperatures (6, 63), which illustrated that lipid metabolism may have increased, accelerating utilization of lipids as an energy source in peripheral tissues. In accordance with this, in rainbow trout exposed to high temperatures, endogenous lipid stores remained the most important energy source contributing up to 55% of total demand (64).

Concerning β-oxidation, an association between water temperature and fatty acid catabolism has been shown, but with inconsistent results in the literature. A recent study in Atlantic salmon showed an increase of β-oxidation in white muscle with increased temperature (25), while changes were not observed by Hevroy et al. (47), and another study in rainbow trout reported increased capacities for oxidizing lipids at cold temperatures (65). In any case, it has been generally accepted that liver and red muscle, but not white muscle, are the most important tissues involved in fatty acid catabolism in fish; thus, an increase in water temperature has been shown to induce increased β-oxidation primarily in those tissues (66, 67). Other study in salmon found that elevation of water temperature was responsible for reduced β-oxidation in liver (47). These last data would be in agreement with the present study, where cptla and cpt1b were down-regulated at 24 and 28°C compared with the low-temperature group, considering the flux of β-oxidation is primarily determined by CPT1, which allows long chain fatty acids to enter into the mitochondria (25).

Notwithstanding, temperature can also affect mitochondrial uncoupling. The increase of temperature from 19 to 28°C led in gilthead sea bream muscle to the up-regulation of the uncoupling protein ucpc2, indicating a higher proton leak, which is in agreement with previous studies in skeletal muscle in mammals (68, 69). UCPS are known to be activated not only by free fatty acids, but also by reactive oxygen species (70, 71). According to this, ucpc2 may play a role as a mechanism for attenuating the possible increase in reactive oxygen species associated with elevated temperature; therefore, overall indicating a less efficient use of fatty acids to obtain energy in this species, upon these environmental conditions.

CONCLUSIONS

To sum up, the present work reports that an increase in water rearing temperature from 19 to 24 and specially 28°C causes in gilthead sea bream juveniles unfavorable growth conditions for the musculoskeletal system due to reduced gene expression of members of the GH/IGFs system and specific MRFs. In white muscle as well, as energetic demand is increased along with temperature, the uptake of fatty acids is enhanced, although apparently, their use as an energy source is less efficient. In addition, the high temperatures applied in vitro, induced changes in vitro in the expression of several key osteogenic genes, suggesting reduced osteoblasts development and matrix production, consistent with the decrease observed in the deposition of minerals. Overall, the present study provides new insights into the possible impact of global climate change in this important Mediterranean species, demonstrating that temperature is a key environmental factor whose increase can lead to unbalanced muscle and bone growth, which should be considered to take preventive measures to reduce production losses and guarantee the sustainability and success of aquaculture.

AUTHOR CONTRIBUTIONS

MR-C and EC conceived and designed the experiments. SB-P, NR-H, and EV performed the experiments. JG, IN, MR-C, and EC contributed reagents and analysis tools. SB-P, NR-H, EV, JG, IN, MR-C, and EC drafted and critically reviewed the paper.

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

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