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Dietary vitamin mix levels influence the ossification process in European sea bass (Dicentrarchus labrax) larvae


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Submitted 12 September 2007; accepted in final form 19 November 2007

Dietary vitamin mix levels influence the ossification process in European sea bass (Dicentrarchus labrax) larvae. Am J Physiol Regul Integr Comp Physiol 294: R520–R527, 2008. First published November 21, 2007; doi:10.1152/ajpregu.00659.2007.—The influence of dietary vitamins on growth, survival, and morphogenesis was evaluated until day 38 of posthatching life in European sea bass larvae (Dicentrarchus labrax). A standard vitamin mix (VM), at double the concentration of the U.S. National Research Council’s recommendations, was incorporated into larval feeds at 0.5%, 1.5%, 2.5%, 4.0%, and 8.0% to give treatments VM 0.5, VM 1.5, VM 2.5, VM 4.0, and VM 8.0, respectively. The group fed the VM 0.5 diet all died before day 30. At day 38, the larvae group fed VM 1.5 had 33% survival, while the other groups, with higher vitamin levels, showed at least 50% survival. The higher the percentage VM in the diet, the lower the percentage of column deformities. High dietary vitamin levels positively influenced the formation of mineralized bone in larvae: the higher the dietary vitamin level, the higher the ossification status. In the larvae group fed at the highest vitamin levels, we observed a temporal sequence of coordinated growth factor expression, in which the expression of bone morphometric protein (BMP-4) preceded the expression of IGF-1, which stimulated the maturation of osteoblasts (revealed by high osteocalcin expression levels). In groups fed lower proportions of vitamins, elevated proliferator peroxisome-activated receptors (PPAR-γ) expression coincided with low BMP-4 expression. Our results suggest that high levels of PPAR-γ transcripts in larvae-fed diets with a low VM content converted some osteoblasts into adipocytes during the first two weeks of life. This loss of osteoblasts is likely to have caused skeletal deformities.

MATERIALS AND METHODS

Animals and diets. Three-day old European sea bass (Dicentrarchus labrax) larvae were obtained from the Ecloserie Marine de Gravelines (Gravelines, France). The fish were acclimated and divided into fifteen 35-liter cylindrical fiberglass tanks (2,100 larvae/tank) at an initial density of 60 larvae/liter. Tanks were supplied with artificial light for 12 hours/day. The larvae were fed a competitive diet, and the diet was supplemented with VM at a basal level of 0.5% to achieve a final concentration of 0.5% VM. The VM was formulated to meet the dietary needs of the larvae as determined by the U.S. National Research Council (NRC) (17), and it was adjusted to meet the basic nutritional requirements of fish larvae in terms of protein, lipids, and carbohydrates. The VM was incorporated into experimental larvae feeds on the basis of experience, the most common level used being 8 times the requirement of juvenile fish (6, 7). This strategy allowed the basic nutritional requirements of fish larvae in terms of proteins, lipids, and carbohydrates to be established, avoiding any possible vitamin deficiency risks (3, 19, 28).

Over the last decade, many studies have reported the important influence that dietary vitamins have on the appearance of larval deformities. In particular, high dietary vitamin A was seen to negatively affect larval morphogenesis during the first weeks of life in Japanese flounder and European sea bass, through one of its active metabolites: retinoic acid (4, 23, 25), which induces skeletal malformations. According to Ville-neuve et al. (25, 26), this vitamin A effect on morphogenesis is mediated by retinoid nuclear receptors that disrupt the normal ontogenic expression pattern of genes involved in bone differentiation. Other dietary vitamins, such as vitamin D3, can also induce vertebral deformities and hypermelanosis in Japanese flounder (10).

These results clearly demonstrate the necessity to accurately define dietary vitamin needs during the larval period. The development of a compound diet, which can totally and efficiently replace live prey, now allows studies to be made on the optimization of vitamin supply in feeds for marine fish larvae. The aim of this study was therefore to test different levels of the same dietary VM during the development of European sea bass larvae, to observe the influence on their morphogenesis process. The first objective was the comparison of different morphological processes, to identify bone differentiation mechanisms that could be affected by dietary vitamins. The second objective was to determine the most appropriate level at which the standard VM should be incorporated into larvae feeds to induce good growth and survival but also harmonious morphological development.
with through-flowing seawater, which had been previously filtered through a sand filter, then passed successively through a tungsten heater and a degassing column packed with plastic rings. Throughout the experiment, salinity was 35%, and the oxygen level was maintained above 6 mg/l by setting the water replacement in the tank at up to 30% per hour (flow rate: 0.18 l/min). Photoperiod was 24-0 h ligh-dark cycle, and light intensity was 9 W/m² maximum at the water surface. All procedures concerning the animals and their handling were conducted in compliance with the Guide for the Care and Use of Laboratory Animals (16). The study was performed under the licence no. 29.021 by the French Department of Veterinary Services (Direction Départementale des Services Vétérinaires) to conduct experimental protocols and samplings on fish.

To provide a wider range of dose levels and have more flexibility in the formula, we doubled the concentration of the standard VM (17). Consequently, 1% concentrated-vitamin mix used in this experiment corresponds to 2% standard VM. Microdiets were prepared in our experimental unit as previously described (3), and the pellet size was 200–400 μm.

Five experimental groups (three replicates per group) of sea bass larvae were reared at 20°C and fed, from day 6 until day 38 posthatching, on microparticulate diets incorporating 0.5% (VM 0.5), 1.5% (VM 1.5), 2.5% (VM 2.5), 4% (VM 4.0), and 8% (VM 8.0) of the concentrated VM. The composition of the VM is described in Table 1. Diet formulas were very close to that of a patented diet (WO 00064273), which includes 8% of the standard VM, equivalent to 4% of our concentrated VM. Throughout the experimental period, to allow the larvae to find the diet microparticles easily in the water column, they were continuously fed in large excess (24 h per day) using automatic belt feeders. As it is not possible to quantify the amount of food ingested by fish larvae, food ingestion was monitored under a binocular microscope by observing the filling of the digestive tract with dietary microparticulates (1 h after feed distribution started).

Sampling. Thirty larvae were collected from each tank for weight measurement (formalin preserved) every week, and at the end of the experiment. Larvae were sampled from each tank at 27 and 38 days posthatching (n = 30–50 larvae, depending on wet body wt) and kept at −20°C for future enzyme assays. For each treatment, another 50 larvae were collected at days 9, 15, and 38, and total RNA was immediately extracted to measure the expression of some of the genes involved in larval development.

The incidence of skeletal malformations (head and vertebral column deformities) was determined by sampling n = 20 38-day-old larvae per tank. These larvae were stained with Alcian blue and Alizarin red S to color cartilage and bone areas, respectively. Survival was evaluated by counting the individuals in each tank at the end of the experiment.

Image analysis. Colored larvae were put on a glass plate containing glycerol and directly scanned using a desk scanner (Epson Perfection 4990 Photo). A 2,500-kb picture was then created.

Six readers made separate overall analyses of the deformities of the spinal column (cyphosis, lordosis, number of vertebrae) and the head (prognatism). The results were compiled and statistically analyzed as described below.

Individual size and the surfaces corresponding to cartilage and bone in whole larvae (day 38) were visualized and quantified using a computerized image analysis package (IMAQ Vision Builder, National Instruments, Austin, TX) after staining with Alcian blue and Alizarin red S. The larvae from the various groups (VM 1.5, 2.5, 4.0, 8.0) were treated simultaneously during the coloration to avoid any bias due to technical variability. The scripting feature of IMAQ Vision Builder was used to record a series of image-processing steps and their specific parameters, so that the computerized image analyses were also performed simultaneously for all samples (batch processing). The script used a list of image-processing commands encompassing the selection of pixel color range and quantification. Selecting ranges of pixel values in color images (threshold operations) allowed the pixels associated with red (bone) or blue (cartilage) colorations to be distinguished. The number of selected pixels was then quantified using a particle analyzes operation. Larval size was quantified by calculating the surface area (in pixels) covered by whole stained larvae.

Analytical methods. Total RNA was extracted using TRIzol and reverse-transcribed in duplicate (iScript cDNA Synthesis Kit; Bio-Rad Laboratories, Hercules, CA). These duplicates were then pooled. Quantitative PCR analyses for each gene were performed in triplicate in a total volume of 15μl containing 5 μl cDNA (dilution: 10⁻³), 0.5 μl primers (10 μmol/l), 7.5 μl iQ SYBR Green supermix 2X (Bio-Rad Laboratories), and 2 μl sterile water. For each target gene [bone

Table 1. Composition (in %) of the diets

<table>
<thead>
<tr>
<th>Ingredients*</th>
<th>VM 0.5</th>
<th>VM 1.5</th>
<th>VM 2.5</th>
<th>VM 4.0</th>
<th>VM 8.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Defatted fish meal†</td>
<td>52.0</td>
<td>52.0</td>
<td>52.0</td>
<td>52.0</td>
<td>52.0</td>
</tr>
<tr>
<td>Fish meal hydrolysate (CPSP 90)</td>
<td>14.0</td>
<td>14.0</td>
<td>14.0</td>
<td>14.0</td>
<td>14.0</td>
</tr>
<tr>
<td>Soy lecithin</td>
<td>7.0</td>
<td>7.0</td>
<td>7.0</td>
<td>7.0</td>
<td>7.0</td>
</tr>
<tr>
<td>Marine lecithin</td>
<td>14.0</td>
<td>14.0</td>
<td>14.0</td>
<td>14.0</td>
<td>14.0</td>
</tr>
<tr>
<td>Concentrated vitamin mix‡</td>
<td>0.5</td>
<td>1.5</td>
<td>2.5</td>
<td>4.0</td>
<td>8.0</td>
</tr>
<tr>
<td>Mineral mix§</td>
<td>4.0</td>
<td>4.0</td>
<td>4.0</td>
<td>4.0</td>
<td>4.0</td>
</tr>
<tr>
<td>Betaine</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
</tr>
<tr>
<td>Cellulose</td>
<td>7.5</td>
<td>6.5</td>
<td>5.5</td>
<td>4.0</td>
<td>4.0</td>
</tr>
</tbody>
</table>

Proximal composition

<table>
<thead>
<tr>
<th></th>
<th>VM 0.5</th>
<th>VM 1.5</th>
<th>VM 2.5</th>
<th>VM 4.0</th>
<th>VM 8.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dry matter</td>
<td>87.0</td>
<td>86.7</td>
<td>87.2</td>
<td>87.1</td>
<td>87.7</td>
</tr>
<tr>
<td>Proteins</td>
<td>62.1</td>
<td>64.0</td>
<td>64.7</td>
<td>64.8</td>
<td>65.8</td>
</tr>
<tr>
<td>Lipids</td>
<td>21.0</td>
<td>20.8</td>
<td>21.3</td>
<td>20.4</td>
<td>21.1</td>
</tr>
<tr>
<td>Neutral lipids</td>
<td>5.6</td>
<td>5.4</td>
<td>5.6</td>
<td>4.7</td>
<td>5.3</td>
</tr>
<tr>
<td>Phospholipids</td>
<td>14.1</td>
<td>14.0</td>
<td>14.3</td>
<td>14.2</td>
<td>14.4</td>
</tr>
</tbody>
</table>

*All dietary ingredients were obtained commercially. Fish meal hydrolysate CPSP 90:10% lipids; Soluble Fish Protein Concentrate (Sopropêche, Boulogne sur Mer, France); soy lecithin (Ets Louis François, St. Maur des Fossés, France); marine lecithin LC 60 (Phosphotech, St. Herblain, France). †Defatted in the laboratory using Norse LT 94 fish meal (La Lorientaise, Lorient, France). ‡Composition per kilogram of the vitamin mixture: choline chloride 60%, 335 g; vitamin A acetate, (500 000 UI/g) 1 g; vitamin E (500 UI/g) 20 g; vitamin D3 (500 000 UI/g) 0.96 g; vitamin B3 2 g; vitamin B5 4 g; vitamin B1 200 mg; vitamin B2 80%, 1 g; vitamin B6 600 mg; vitamin C 35%, 28.6 g; vitamin B9 80%, 250 mg; vitamin concentrate B12 (10 kg/g), 0.2 g; biotin, 1.5 g; vitamin K3 51%, 3.92 g; meso-inositol 60 g; cellulose, 542.4 g. §Composition per kilogram of the mineral mixture: 90 g KCl, 40 mg KIO3, 500 g CaHPO4 2H2O, 40 g NaCl, 3 g CuSO4 5H2O, 4 g ZnSO4, 7H2O, 20 mg CoSO4, 7H2O, 20 g FeSO4, 7H2O, 3 g MnSO4·H2O, 215 g CaCO3, 124 g MgSO4·7H2O, and 1 g NaF. VM, vitamin mix.
morphogenetic protein 4 (BMP-4), IGF-1, osteocalcin, retinoid X receptor (RXR\(\alpha\)), retinoic acid receptor alpha (RAR\(\alpha\)), and peroxisome proliferator-activated receptor gamma (PPAR\(\gamma\)), specific complementary primers designed from previously cloned sequences are listed in Table 2. The housekeeping gene Ef1 was chosen as a reference since it did not exhibit any significant variation in expression among the samples. Thermal cycling was initiated with incubation at 95°C for 3 min to activate iTaq DNA polymerase. After this initial step, 45 cycles of PCR were performed. Each PCR cycle consisted of heating for 30 s at 95°C for denaturation, then for 1 min at 60°C for annealing and extension. Cycle threshold (CT) values corresponded to the number of cycles at which the fluorescence emission monitored in real time exceeded the threshold limit. Melting curve analysis was performed to confirm production of a single product in these reactions, and the products were sequenced by MilleGen (Labe`ge, France). Standard curves were established for each gene by plotting CT values against the log\(_{10}\) of five different dilutions (in triplicate) of cDNA sample solutions. Real-time PCR efficiencies were determined for each gene from the slopes obtained with Bio-Rad software, applying the equation \(E = 10^{-\frac{1}{slopel}}\), where \(E\) is PCR efficiency. To determine the relative quantity of target gene-specific transcripts present in the different samples, expression ratios (\(R\)) were calculated according to the following formula:

\[ R = \frac{\langle E_{\text{target gene}} \rangle^{\text{ACT target gene}} - \text{mean sample} - \text{mean ref sample}}{\langle E_{\text{Ef1}} \rangle^{\text{ACT Ef1}} - \text{mean sample} - \text{mean ref sample}} \]

where Ef1 is the housekeeping gene, mean sample corresponds to triplicate average, and ref sample was chosen in the VM 4.0 group.

Statistics. Results are expressed as means \(\pm\) SD. Survival rates, malformation rates, and bone/cartilage ratios were normalized by arcsin square root transformation and were analyzed by one-way ANOVA with Statview followed by the Newman-Keuls test when significant differences were found at the \(P < 0.05\) level. Gene expression data, excluding the VM 0.5 group, were analyzed by two-way ANOVA followed by Newman-Keuls tests when significant differences were found at the \(P < 0.05\) level.

To analyze VM 0.5 results for specific genes, we performed one-way ANOVAs on day 9, day 15, and day 38.

RESULTS

Larval performance. All of the experimental feeds were efficiently ingested by the larvae. The larvae fed at the lowest vitamin level, VM 0.5 diet, had all died by day 30. At day 38, the groups fed VM 2.5, VM 4.0, and VM 8.0 diets all had survival rates higher than 50\% (Table 3) and significantly higher from that of the VM 1.5 group (33\%). In terms of weight, the VM 1.5 group also had the lowest performance (29.8 mg), while groups fed diets VM 4.0 and VM 8.0 showed 30\% higher weight (Table 3). The same trend was observed with larval length (Table 3), although the differences were less pronounced (only a 14\% difference in length between the VM 1.5 and VM 8.0 groups).

All of the experimental groups exhibited a statistically similar percentage of head deformities, mostly because of the very high variability (Fig. 1). The higher the dietary level of VM, the lower the percentage of column deformities (\(P < 0.0001\)), although no significant difference was observed between VM 4.0 and VM 8.0. The percentage of column deformities decreased by a factor of 3.5 between the groups fed VM 1.5 and 8.0 diets (\(P < 0.001\)).

Bone/cartilage ratio was analyzed to evaluate the ossification process in larvae (Fig. 2). The high dietary vitamin levels increased the formation of mineralized bone in larvae; the higher the dietary vitamin level, the higher the ratio (\(P < 0.001\)).

Gene expression. BMP-4 expression was effectively halved between day 9 and day 38 (\(P < 0.0001\)). This decrease was much more pronounced (2.9 times; \(P < 0.0001\)) in larvae fed the low-vitamin diets, i.e., lower than 4.0\% (Fig. 3). One-way ANOVAs of BMP-4 expression at days 9, 15, and 38 indicated that the larvae groups fed diets with low vitamin mix levels exhibited the lowest values (\(P = 0.017, P < 0.006, P = 0.011\) for days 9, 15, and 38, respectively). In contrast, there was at least a 10-fold increase in IGF-1 expression at day 38 compared with day 9 (\(P < 0.001\)). The only significant influence of dietary vitamin level was noted at day 38 in group VM 4.0, where IGF-1 showed a relative expression 1.8 times higher (Fig. 4, effect of diet \(P < 0.0001\) and diet \(\times\) age interaction \(P = 0.0001\)) than the three other groups. Similarly, an elevation of osteocalcin expression (by at least 20 times) occurred during the experiment (\(P < 0.001\)), although the greatest increase happened after day 15 (\(P < 0.001\)) (Fig. 5). The highest osteocalcin expression values were noted in larval groups fed the highest dietary vitamin levels (\(P = 0.0009\)), but this influence of diet was only shown at day 38 (diet \(\times\) age interaction \(P = 0.0003\)). One-way ANOVA analysis revealed that the osteocalcin expression in group VM 8.0 was 1.5, 1.8, and 2 times higher than in groups VM 4.0, 2.5, and 1.5, respectively (\(P = 0.0086\)). It should be added that, at day 38, bone/cartilage ratios were highly correlated with osteocalcin expression data (\(Y = 0.534X + 4.756; R^2 = 0.961; P = 0.019\)).

RXR\(\alpha\) expression levels increased with age (\(P = 0.0002\)) but, once again, more markedly after day 15 (\(P < 0.0005\)) (Fig. 6). The influence of diet was clearly shown at days 9 and 38 (\(P = 0.0002\)) when the highest RXR\(\alpha\) values were measured in larvae groups fed VM 4.0 and VM 8.0. One-way ANOVA analysis showed that the expression of RXR\(\alpha\) in both the VM 4.0 and 8.0 groups was over 2.5 and 5 times higher than the other groups, at days 9 and 38, respectively.

A sharp decrease in RAR\(\alpha\) expression occurred between day 9 and 38 (\(P < 0.0001\)) (Fig. 7), which was more marked in groups VM 1.5 and 2.5 (50\% decrease) than in groups VM 4.0 and 8.0 (30\% decrease), revealing an effect of dietary vitamin level on this parameter (\(P = 0.0001\)).

PPAR\(\gamma\) expression increased with age after day 9, doubling by the end of the experiment (\(P < 0.0001\)) (Fig. 8). The expression of PPAR\(\gamma\) was also positively influenced by the

Table 2. Oligonucleotide primers used in real-time PCR

<table>
<thead>
<tr>
<th>Genes</th>
<th>Accession Number</th>
<th>Forward and Reverse Primers</th>
</tr>
</thead>
<tbody>
<tr>
<td>BMP(\gamma)</td>
<td>AJ5673451</td>
<td>P: CGTCTCTCTCTCAGGCTGAA</td>
</tr>
<tr>
<td>IGF-1</td>
<td>AY800248</td>
<td>R: GGCTACACAGGACAGGCTCA</td>
</tr>
<tr>
<td>Osteocalcin</td>
<td>AY663813</td>
<td>R: RGGGCAGGAGCTGACAGCA</td>
</tr>
<tr>
<td>RXR(\alpha)</td>
<td>AJ567907</td>
<td>R: TGGCCCTAGGTCGTGTGGCT</td>
</tr>
<tr>
<td>RAR(\alpha)</td>
<td>AJ496189</td>
<td>F: CGTCTTGGTGAGCACCCTGCA</td>
</tr>
<tr>
<td>PPAR(\gamma)</td>
<td>AY590303</td>
<td>R: CAGCATACGAGCAGCGAGTGC</td>
</tr>
<tr>
<td>Ef1</td>
<td>AJ866727</td>
<td>R: CGTGAACGAGGAGCAAGCAC</td>
</tr>
</tbody>
</table>
of certain indicators of bone differentiation can be used to
parameters other than just weight or survival. The measurement
period and refine vitamin dosing by taking into account pa-
sanding of true dietary vitamin requirements during the larval
26). These results underline the need to improve our under-
clear evidence for the influence of dietary nutrients, and more
is not satisfactory. Recent studies on fish larvae have provided
marine fish hatcheries (15) reveals that this empirical approach
spinal disorders occur with a high frequency and variability in
(though an additional safety margin of extra vitamins is gen-
tional interrelationships, fairly wide ranges of requirement
values have been reported for the same species (17) because of
differences in experimental conditions. Moreover, these same
wide value ranges established for juveniles have also been
considered empirically valid for earlier developmental stages
(though an additional safety margin of extra vitamins is gen-
spinal disorders and vertebral and
organ systems occur with a high frequency and variability in
marine fish hatcheries (15) reveals that this empirical approach
not satisfactory. Recent studies on fish larvae have provided
clear evidence for the influence of dietary nutrients, and more
particularly vitamins, on their developmental processes (10,
26). These results underline the need to improve our under-
standing of true dietary vitamin requirements during the larval
period and refine vitamin dosing by taking into account pa-
rameters other than just weight or survival. The measurement
of certain indicators of bone differentiation can be used to
monitor for the appearance of deformities, so as to reduce these
as much as possible. Moreover, such studies should also use
feed ingredients that do not contain significant amounts of
vitamins. Larval feeds traditionally contain fish meal, which
mainly provides a source of protein but which also contains
significant and variable amounts of other nutrients, such as
fatty acids, minerals, and vitamins (particularly liposoluble
vitamins A, D, E, and K). To overcome possible vicariance
(underestimation of one vitamin requirement when others are
present in excess), we decided to produce a defatted fish meal
that improved our control over the liposoluble vitamin supply,
particularly that of vitamins A and D, which are known to
fluence bone differentiation. Essential fatty acids were sup-
plied using purified marine phospholipids, containing only
traces of vitamins A and E. The experimental diet formula used
in this work was very similar to that of a patented diet known
to induce good development in fish larvae (3), even though a
certain and variable percentage of deformities are still ob-
served, and this diet allowed us to examine the physiological
effects of variation in dietary vitamin levels.

Studies testing the effect of different levels of vitamin mix
on fish larvae are scarce. Escaffre et al. (6) showed that a diet
with 5% standard VM led to better growth performance than
one with 2% VM in common carp larvae. However, our present
study is the first to test such a large range of incorporation
levels of the standard VM into larval feeds. Past studies have
described the effects of vitamins A (4, 5, 6, 24, 25), C (7), or
D (10) on fish larval development. Among these, vitamin A has
been the most studied, not only for growth and survival but
also to identify the molecular processes leading to the appear-
ance of deformities. Consequently, comparisons could be made
between these studies and our own by examining this particular
vitamin. Villeneuve et al. (25) reported a negative effect of
high and low dietary vitamin A level on growth and survival in
European sea bass larvae that were fed diets incorporating
retinol acetate at 1,000 mg/kg diet or less than 50 mg/kg diet.

VM 1.5 and 2.5 diets \((P < 0.0001)\), with the maximum effect at
day 15 (diet \( \times \) age interaction \( P < 0.0001)\).

One-way ANOVA of PPAR\(\gamma\) expression at day 9 and at day
15 indicated that the larvae groups fed diets with low VM
levels exhibited values 1.9 and 2.9 times higher than those
found in groups fed the two highest dietary vitamin mix levels
\((P = 0.0002 \text{ and } P < 0.0001 \text{ for days 9 and 15, respectively})\).
At day 38 this effect, induced by low dietary vitamin mix, is
less marked.

DISCUSSION

In fish, the dietary vitamin requirement has classically been
determined in juvenile fish on the basis of their weight gain and
absence of deficiency signs. As vitamin requirements are
affected by various environmental, physiological, and nutri-
tional interrelationships, fairly wide ranges of requirement
values have been reported for the same species (17) because of
differences in experimental conditions. Moreover, these same
wide value ranges established for juveniles have also been
considered empirically valid for earlier developmental stages
(though an additional safety margin of extra vitamins is gen-
"
(500,000 USP U/g), respectively. The highest level tested by these authors would correspond to 10 times our highest level of vitamin A (diet VM 8.0), and their lowest levels would correspond to our diets VM 1.5 and VM 2.5. Villeneuve et al. (25) only found a slight negative effect on larval survival when feeding diets with an intermediate level of vitamin A: 250 mg retinol acetate/kg diet. Dedi et al. (4) observed negative effects on growth of flounder larvae fed live prey enriched with different vitamin A-palmitate levels, above 416 IU/g, which correspond approximately to the vitamin A level found in VM 4.0. Other authors (5, 18) also described adverse effects of high dietary vitamin A levels on growth. The upper range of dietary VM levels tested in our experiment was obviously too low, compared with the extremely high values reported in the literature, to evidence any adverse effect of hypervitaminosis on growth and survival. In our experiment, growth (length and weight) and survival were only negatively influenced by diets incorporating less than 2.5% VM, corresponding to a hypovitaminosis. Considered together, these data indicate that growth and survival parameters are not sensitive enough to reveal any effect of the dietary VM when this varied within a “normal range,” as it can do in commercial feeds.

The elevation of the vitamin A level in larvae feeds generally induced a higher percentage of larval deformities. Villeneuve et al. (25) found almost 20% head deformities with a 50 mg retinol acetate/kg diet. Dedi et al. (4) and Takeuchi et al. (24) found that high dietary vitamin A levels adversely affected the vertebral development of fish. In the present experiment, this adverse effect of high dietary vitamin levels on the column was not noted since the upper levels of vitamin tested in our experiment corresponded to medium/normal levels of vitamins in the studies reported by the literature. We found a high level of head deformities whatever the dietary vitamin level, which could suggest that vitamins do not influence head morphogenesis (although this is unlikely considering the data in the literature) or that there was an inadequate proportion of certain vitamins in the mixture. However, we simultaneously observed a lower occurrence of column deformities and a more intense ossification process in larvae fed the diets with the higher percentages of vitamin mix. This result constitutes an original finding and suggests that dietary VM level affects the differentiation of vertebral column bone cells at early stages of fish larval development. The formation of the vertebral column in
...include cartilage and bone tissue types, with three types of more than 500 genes either directly or indirectly (2).

Target genes and thereby suppress or enhance the transcription response elements of vitamin D or to retinoid acid localized on the signaling network (29). These heterodimers can bind to the then involved in nearly all processes associated with development by forming obligate heterodimers with RXR, which are on the vitamin D receptors (VDR) (12). VDR receptors function by controlling chondrocyte function. Vitamin C is a cofactor in connective tissue synthesis (15). The physiological role of structural components such as bone matrices, collagen, and immunological responses, including a role in the formation of many biological processes of cells involved in hormonal and developmental processes. For example, a possible effect of vitamin A on chondrocyte differentiation has already been observed in fish larvae. Suzuki et al. (22, 23) reported that Japanese flounder larvae exposed to retinoic acid at a concentration above 10−7 M show a depressed expression of sonic hedgehog (shh) and Hoxd-4 and reduce the scale of expansion of shh expression domains in the pharyngeal area. Pharyngeal cartilages that formed in these larvae were malformed. The period when shh expression domains expand corresponds to the stage of active proliferation of the cartilage precursor cells and their differentiation into chondrocytes (23). The malformations observed by Suzuki et al. (23) were attributed to a misregulation of the shh signaling pathway in the head area.

Osteoblasts are derived from multipotent mesenchymal stem cells that give rise to osteoblastic progenitor cells. These then undergo proliferation/amplification before final differentiation and expression of specific osteoblastic markers (11).

In the present experiment, we found a temporal sequence of coordinated growth factor expression. Indeed, the expression of BMP-4, a potent osteoblast differentiation factor, was more elevated during the earliest larval developmental stages than later on, and its expression was positively modulated by the level of dietary vitamins. This pattern of expression is in agreement with a primary effect of BMP-4 on the multipotent cells, and the high expression of BMP-4 can cause these cells to commit to an osteoblastic pathway. At a later developmental stage (from day 15 onward), we observed an elevation in the expression of IGF-1, which was more marked when the diets contained high levels of vitamins. Considering that IGF increases proliferation and plays a major role in stimulating mature osteoblast function (11), our result suggests that IGF promoted the late-stage differentiation of sea bass larvae osteoblasts in a dose-dependent manner, according to the level of dietary vitamins. Indeed, IGF expression preceded the expression of osteocalcin by several days. Osteocalcin, which is probably the most, if not the only osteoblast-specific gene (13),

teleost fish occurs by intramembranous ossification, in which bone is formed as a bone matrix and directly ossified (9). The cranial bone is formed by endochondral ossification, however, in which bone is first formed as cartilage and then ossified into mineralized bone during larval development (23). These physiological differences between dermal head bone cells and vertebral column bone cells may explain the fact that the development of bone in the head and the vertebral column is not affected in the same way by the dietary vitamin mix.

Among the vitamins contained in the vitamin mix, vitamins A, C, D, and K particularly influence the formation of bone (15). The biological activity of vitamin A is mainly mediated by its active metabolite retinoid acid, obtained by the dehydrogenation of vitamin A. Some important functions of vitamin A include regulation of cellular differentiation and proliferation and regeneration of rhodopsin that is necessary for vision. Vitamin A regulates skeletogenesis and cartilage development by controlling chondrocyte function. Vitamin C is a cofactor in many biological processes of cells involved in hormonal and immunological responses, including a role in the formation of structural components such as bone matrices, collagen, and connective tissue synthesis (15). The physiological role of vitamin D3 (1,25-dihydroxy vitamin D) in fish is not clearly defined although, as in mammals, it seems to be involved in intestinal calcium and phosphate absorption and also to have a direct effect on bone (15). Vitamin K is specially known for its effect on blood clotting, but it also has a role in stimulating bone formation and inhibiting resorption (15).

The biological effects of vitamin A metabolites are mainly exerted through the activation of two groups of nuclear receptors: RARs and RXRs, with different retinoid acid isomer specificity (20). Similarly, vitamin D3 signaling is dependent on the vitamin D receptors (VDR) (12). VDR receptors function by forming obligate heterodimers with RXR, which are then involved in nearly all processes associated with development, emphasizing the pivotal role of the retinoid pathway in the signaling network (29). These heterodimers can bind to the response elements of vitamin D or to retinoid acid localized on target genes and thereby suppress or enhance the transcription of more than 500 genes either directly or indirectly (2).

Skeleton cells are derived from different embryonic lineages and include cartilage and bone tissue types, with three types of cells: chondrocyte in the cartilage, osteoblasts or bone-forming cells, and osteoclasts or bone-resorbing cells in the bone (13). It is now well established that cell differentiation into chondrocytes or osteoblasts and subsequent cartilage and bone formation are processes governed by several growth factors and their intracellular signals (27), both of which can, in turn, be modulated by nutritional factors. For example, a possible effect of vitamin A on chondrocyte differentiation has already been observed in fish larvae. Suzuki et al. (22, 23) reported that Japanese flounder larvae exposed to retinoic acid at a concentration above 10−7 M show a depressed expression of sonic hedgehog (shh) and Hoxd-4 and reduce the scale of expansion of shh expression domains in the pharyngeal area. Pharyngeal cartilages that formed in these larvae were malformed. The period when shh expression domains expand corresponds to the stage of active proliferation of the cartilage precursor cells and their differentiation into chondrocytes (23). The malformations observed by Suzuki et al. (23) were attributed to a misregulation of the shh signaling pathway in the head area.

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was not regulated by the dietary vitamin level until day 15. Osteocalcin, moreover, displayed a constant low expression level until day 28 (result not shown). The positive impact of the high dietary vitamin levels on osteoblast differentiation is shown clearly at day 38 by both osteocalcin gene expression and red alizarin coloration of bones. It is interesting to note that the high correlation between osteocalcin expression and red alizarin coloration of mineralized bone tissue at day 38 demonstrates that this gene is a good indicator of bone differentiation and can be used to investigate ossification status in whole body larval homogenates.

We also found that larval groups fed a low vitamin level and exhibiting a low expression of the BMP-4 gene had a concomitantly elevated expression of PPAR-γ. This result is interesting since it has been shown that osteoblastic cells, when transfected with PPAR-γ, can be converted to express an adipocytic phenotype (1). The possibility that a differentiated osteoblastic cell subsequently redifferentiates as a distinct phenotype is supported by the existence of bipotent osteoblast-adipocyte cells from bone marrow in several species (11). It is also interesting to note that, in our experiment, the high expression levels of BMP-4 were associated with high expression levels of RARα and RXRα in larvae groups fed the diets with elevated vitamin content. This result reflects a higher retinoid acid content in these larvae, since the expression of retinoid receptors is known to be modulated by the level of retinoid acid (25) that was obviously higher in groups fed the diets with elevated vitamin levels. Considering that retinoid acid inhibits adipose conversion and acts with BMP to promote osteoblast differentiation, our findings suggest that larvae that were fed diets with low VM content and expressing high levels of PPAR-γ transcripts, have had part of their osteoblast potential converted into adipocytes during the first two weeks of life. This loss of osteoblasts is a likely cause of skeletal deformities.

In conclusion, this study has indicated that fish larvae require higher dietary VM levels than juvenile fish to achieve their developmental processes correctly. The National Research Council standard VM that incorporated into larval feeds at 8 times the content recommended for juveniles gave the best results in terms of growth, survival, and also morphogenesis. However, the fact that the percentage of head and column deformities remained significant demonstrates the need to further refine the proportions of certain vitamins (particularly those known to be involved in bone and collagen synthesis, i.e., Vitamins A, D, and C) in the standard vitamin mix. Our results showed a temporal sequence of coordinated growth factor expression, involving BMP-4 and IGF-1, controlling the differentiation of osteoblasts and also revealed how this sequence could be disrupted by differences in the dietary level of certain vitamins (that need to be precisely identified), leading to the appearance of deformities.

Perspectives and Significance

The normal programmed development of a multicellular organism is a synchronized series of events driven by genetic instructions. An organism however, has the ability to respond to environmental/nutritional situations by adaptations during the critical early period of its life, which will permanently affect its physiology and/or morphology in a positive or negative way. This plasticity is of particular importance in cultured marine fish larvae, especially concerning their bone metabolism and the high frequency of skeletal deformities. One of the major bottlenecks for understanding skeletal deformities in fish is the lack of effective methods to characterize bone anomalies induced by nutrition early in life. This study initiates the characterization of the basis of fish skeletal abnormalities and brings into view new elements that should be considered for fish larval feed composition.

ACKNOWLEDGMENTS

The authors are grateful to Viviane Verhac from DSM Nutritional Products-France, who kindly provided all of the vitamins used in this work. The authors also thank Hervé Le Delliou for the feeds analyses and Helen McCombie for correcting the English.

GRANTS

This work was, in part, supported by FINEFISH, a Collective Research Project of the sixth Framework Programme of the European Union (Contract 012451).

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