Does otolith structure reflect the nutritional condition of a fish larva? Comparison of otolith structure and biochemical index (RNA/DNA ratio) determined on cod larvae

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ABSTRACT: Cod larvae from laboratory rearing experiments aged from 1 to 12 d after hatching, both fed and deprived of food, were analysed. The number of increments on the otolith and the width of these increments were determined together with the RNA/DNA ratios on the same individual larva. Alizarin marking of the otoliths was performed to confirm the formation of daily increments. Cod larvae reared at 6°C formed the first ring right after hatching and deposited increments on a daily basis. A comparison of the measurements between the right and the left lapilli showed that these can differ, if the radius is taken. The lapillae core showed especially high individual variability, whereas the sum of the increments did not differ between both lapilli. Until Day 10 after hatching, while the larvae were still feeding on their yolk, the external food situation did not affect the increment width of the lapilli or the RNA/DNA ratios. In larvae older than 10 d the width of the daily increments was dependent on the nutritional situation and RNA/DNA ratios decreased in starving larvae in comparison to feeding larvae. RNA/DNA ratios and increment widths were correlated.

KEY WORDS: Nutritional condition · RNA/DNA ratio · Otolith microstructure · Recruitment · Cod larvae

INTRODUCTION

Fluctuations in the size of fish populations may occur as a consequence of changes in the annual influx of young or recruiting fish. Recruitment variations are compounded by the effects of human exploitation and often attributed to the effects of environmental variations on the survival of egg and larval stages. The success or failure of annual recruitment can have a number of different abiotic and biotic causes. High mortality rate during early stages is considered one of the major factors causing stock fluctuations. The lack of food or a mismatch in the distribution of larval fish and food organisms are principal causes of poor year class strength (Hjort 1914, Hunter 1976, Lasker 1978, Hewitt et al. 1985).

The relationship between RNA and DNA is an index of a cell's metabolic intensity and has been used to measure recent growth in fish (Buckley 1984, Buckley & Lough 1987, Bulow 1987, Hovenkamp 1990, Hovenkamp & Witte 1991) and has proved to be a useful indicator of nutritional condition, as shown in several larval fish studies (Buckley 1980, 1984, Martin et al. 1985, Fukuda et al. 1986, Buckley & Lough 1987, Clemmesen 1987, 1994, Raee et al. 1988, Robinson & Ware 1988).

Otolith structures reflecting daily patterns were described by Panella (1971, 1974) and experimentally proven by Struhsacker & Uchiyama (1976). Brothers et al. (1976) used otolith increment structure to determine the age of larvae and juvenile fish. The environment experienced by a larva will influence its otolith structure, as was shown in several laboratory experiments (Panella 1980, Neilson & Geen 1982, Radtke & Dean 1982, Berghahn & Karakiri 1990, Mugiya & Oka 1991).

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It is known that environmental factors such as constant darkness (Dale 1984) or too high temperatures (Mosegaard et al. 1988) can affect the otolith increment structure. Based on the assumption that somatic growth is reflected in the growth of the otolith, a larva that is growing well should deposit a wider daily increment than a starving larva with slower growth. Maillet & Checkley (1990) and Zhang & Runham (1992) determined changes in the otolith structure in starving laboratory-reared Atlantic menhaden and Oreochromis niloticus. Studies on Norwegian spring-spawning herring and North Sea autumn-spawning herring have shown the potential of using otolith microstructure in recruitment research (Fossum & Moksness 1993).

All condition indices determine the nutritional condition at the time of catch, reflecting the situation of the last hours, days or weeks before catching, depending on the methods used. It is sometimes difficult to assess if the larva's condition is improving or deteriorating. RNA/DNA ratios are commonly used to backcalculate and to forecast the growth and survival potential of larvae in order to help predict recruitment. The validity of the RNA/DNA ratio can be improved by incorporating otolith increment structure studies (daily increment studies) on some subsamples, in order to have the growth history of the larvae as well as the condition at catch.

The aim of this study was to combine otolith microstructure analysis and RNA/DNA ratio determination on the same individual larva to compare the effect of food deprivation on the width of the daily increment and the RNA/DNA ratio and evaluate the potential use of this combination for recruitment research.

MATERIALS AND METHODS

Cod larvae (Gadus morhua) were reared in the Havsfiskelaboratoriet Lysekil, Sweden, from 3 to 27 May 1994. Adult cod caught in November 1993 in the Bornholm Basin were transported to Lysekil, adapted to the higher salinity (30 ppm) and kept in 10,000 l tanks at 6 to 7°C as brood stock for the experiments. Cod spawned naturally. The fertilized eggs were transferred into 100 l tanks using a 300 μm plankton net. The eggs floated on the surface and were moved by a gentle water flow (0.5 l min⁻¹). Temperature in the incubation tank was 6 to 7°C. Larvae hatched after 13 d and were transferred to 100 l rearing tanks. Larvae were fed starting on Day 4 after hatching with the rotifer Brachionus plicatilis reared on the flagellate Isochrysis galbana. Temperature throughout the experiment varied from 6 to 7.5°C and salinity was between 33 and 34 ppm. Tanks were illuminated on a 16:8 h day/night cycle. Larvae were fed 3 times a day at 08:00, 12:00 and 16:00 h at a density of 0.65 Brachionus ml⁻¹. The daily food ration amounted to 2 Brachionus ml⁻¹. Due to unexpectedly high mortality rates the experiments had to be terminated after Day 12 after hatching.

For marking of the otoliths 100 larvae were transferred to a 5 l bucket containing Alizarin (50 mg l⁻¹; Tsukomoto 1988, Blom et al. 1994, Geffen 1995) on Day 4 after hatching and kept there for 16 h to let the Alizarin set a fluorescent mark on the increment formed that day. After that the larvae were carefully transferred to clean sea water. Samples of the larvae were taken daily starting on Day 4 at 11:00 h after the larvae had been fed at 08:00 h. Five larvae per sample were taken, transferred into Eppendorf vials and stored in liquid nitrogen until the end of the experiment. After that the samples were stored in a −70°C freezer and left there until otolith and RNA/DNA analysis was performed.

The analyses of the RNA and DNA content of the larva and the dissection of the otolith were performed simultaneously. Larvae were thawed, and the standard length was measured. Sagittae and lapilli were identified using a polarisation filter attached to a binocular. Since the lapilli in cod larvae are initially the larger otoliths and therefore easier to extract and read than the sagittae, they were dissected from the individuals. After age 25 d, as the rate of otolith growth of the sagitta increases, it then becomes the most accurate otolith to us for estimating age and growth rates (Bergstad 1984). The lapilli were located using a dissecting microscope with cross-polarized light and dissected from the larvae using fine insect-needles. Adhering tissue was carefully scraped away from the otolith, which was then rinsed with distilled water. After 5 to 10 min of drying at room temperature the lapilli were mounted on glass slides using fingernail polish. A circle was drawn on the slide around the otolith to facilitate future manipulation and analysis (Stevensen & Campana 1992). It was not necessary to polish the otoliths due to the young age of the sampled larvae.

The larva without the lapilli was put into an Eppendorf vial kept on ice. After the lapilli of 5 larvae had been fed at 08:00, 12:00 and 16:00 h at a density of 0.65 Brachionus ml⁻¹, the daily food ration amounted to 2 Brachionus ml⁻¹. Due to unexpectedly high mortality rates the experiments had to be terminated after Day 12 after hatching.

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cence microscope (Zeiss Axioplan) at 1000× magnification. Both lapilli were used for the analysis, and the radii, number of increments and width were measured 4 times in 4 different directions on the lapilli and the means were calculated.

RESULTS

To determine the accuracy of the age determination using the number of otolith increments, the results of the readings on the lapilli were compared with the known age of the laboratory-reared cod larvae (Fig. 1). Out of 150 analysed lapilli, 147 showed the deposition of a daily increment starting on the day of hatch. Only 2% of the analysed lapilli showed 1 ring less than expected. A 16 h treatment with Alizarin resulted in an orange increment deposited on the day of marking. It could later be identified under the fluorescence microscope. The number of increments formed after the Alizarin-marked increment was counted and compared with the known number of days passed since the marking. The rings after marking are deposited on a daily basis and give further confirmation of the formation of the first increment on the day of hatching (Fig. 2).

A comparison of left and right lapilli showed that the radius or diameter of the lapilli not only depends on the growth of the daily increments but is very much affected by the size of the lapilli's core. The size distribution of the lapilli cores between the left and the right side varied between 4.5 and 13 μm, but did not show a statistically different distribution between left and right side (t-test, p < 0.05). The comparison of the lapilli core of the left and the right lapilli measured on the same individual larva revealed that there were differences in the size of the core (Fig. 3). A similar picture appeared when the radius of the left and the right lapilli on the same larva was compared (Fig. 4). In comparison the sum of the increments deposited on the left and the right lapilli did not differ (Fig. 5). The variability between the size of the left and right lapilli was mainly caused by the difference in the size of the core.

To evaluate the effect of food availability or food withdrawal on the width of the daily otolith increments a comparison of the size of the increments of all analysed fed and starved larvae showing mean values and standard deviations is presented (Fig. 6). The variability in the size of the daily increments is high in both groups but a comparison of the mean values shows a trend of increasing increment width in fed larvae starting on Day 7 and decreasing width of the increments in the starved group.

From Day 4 to Day 7 the RNA/DNA ratio decreased from a value of 7 to below 2 (Fig. 7). This seems to be due to yolk absorption causing a decrease in condition. Between Day 7 and Day 12 the RNA/DNA ratio in the starved group decreased further, in contrast to that of the fed group, which slightly increased. The effect of food availability started to be visible on Day 11 (Fig. 7). For analysis of the relationship between mean RNA/
DNA ratios and mean relative lapilli growth the values for fed and starved larvae are given in Fig. 8. The relative lapillus growth was calculated by setting the width of increment 4 (Day 5) to a value of 1 and calculating the relative growth of the following increments in relation to that value. Every data point represents the mean of 4 or 5 larvae. (The dataset for this figure is

Fig. 3. *Gadus morhua*. Correlations between radii of otolith cores of left and right lapilli. Values are means calculated from 4 measurements in 4 different directions on the lapilli of 4 to 12 d old cod larvae. A linear regression model was fitted to the data.

Fig. 4. *Gadus morhua*. Correlations between radii of left and right lapilli. Values are means calculated from 4 measurements in 4 different directions on the lapilli of 4 to 12 d old cod larvae. A linear regression model was fitted to the data.

Fig. 5. *Gadus morhua*. Correlations between sums of increments of left and right lapilli. Values are means calculated from 4 measurements in 4 different directions on the lapilli of 4 to 12 d old cod larvae. A linear regression model was fitted to the data.

Fig. 6. *Gadus morhua*. Mean increment width of 59 fed cod larvae compared to 27 starved cod larvae. Error bars give the standard deviation.
Fig. 7 Gadus morhua. RNA/DNA ratios of cod larvae in relation to age and feeding situation. Starving larvae (n = 45) were deprived of food starting on Day 4 and Day 7 and compared to fed larvae (n = 86). Error bars give the standard deviation based on the Alizarin marking experiment and is reduced compared to the dataset shown in Fig. 7. The reason for taking the width of increment 4 is that the switch from yolk absorption to external feeding in cod larvae at the given temperature starts at that age (Lau-rence 1978, Radtke & Waiwood 1980, Solberg & Tilketh 1984, Fossum 1986). The growth observed in the following increments therefore should be reacting to external food supply. The relative growth of the lapilli of the starved larvae was reduced compared to the fed group. The RNA/DNA ratios of fed larvae doubled between Day 6 and Day 12, whereas the value in the starved group was reduced to half of the ratio of starved larvae at the beginning (Fig. 8). It can be seen that the feeding larvae have higher RNA/DNA ratios and a higher growth of the daily increments, showing that the RNA/DNA ratio and the increment growth are coupled.

DISCUSSION

The study showed that cod larvae form the first increment on the otolith on the day of hatching. Age determination based on increment numbers in comparison to known laboratory age as well as Alizarin marking results confirmed the deposition of daily increments. These results are in agreement with increment formation studies in cod larvae by Dale (1984) and Geffen (1995). Neilson & Geen (1982) showed differences in the size of the otoliths taken from the left or right labyrinth in salmonids (Oncorhynchus tscha-wytscga). Comparison of left and right lapilli in this study clearly showed that care has to be taken when measuring the total size (radius), since differences between left and right otoliths occurred. These differences did not result from the growth of the daily increments, but depended on the size of the otolith core at the time of hatch. Individual egg development as well as environmental factors might affect the size of the core. Influences on the formation of the otolith core should be further analysed.

The growth of the increments was affected by the availability of external food sources and differed between feeding and starving larvae starting on Day 7. The RNA/DNA ratio started to increase on Day 7 in the feeding group, becoming more pronounced on Day 11, whereas the RNA/DNA content in the starved group decreased and also reflected the situation found in the otolith microstructure. During the yolk sac phase no clear differences between feeding and starving larvae could be found, which is in agreement with results discussed in Clemmesen (1994). Results for RNA/DNA ratios shown here are in agreement with studies on cod larvae by Buckley (1979) and results for herring larvae by Clemmesen (1987, 1994).

Alizarin marking did not affect the RNA and DNA measurements, since the concentrations of nucleic
acids determined on marked and unmarked cod larvae did not differ. Unfortunately, sampling of the larvae could not be performed after Day 12. It is postulated that the trends shown in this study would have been much more significant if samples from older larvae had been available. Future studies should increase the sampling protocol up to an age of 21 d at least.

Biochemical indicators (RNA/DNA ratios) have been used to demonstrate starvation mortality in the field, and correlations between food availability and larval condition have been found (Setzler-Hamilton et al. 1987, Frank & McRuer 1989, Canino et al. 1991, Theilacker et al. 1996). A positive relationship of RNA/DNA ratio and prey abundance has been shown for striped bass larvae Morone saxatiis (Martin et al. 1985) and Atlantic cod Gadus morhua and haddock Melanogrammus aeglefinus larvae (Buckley & Lough 1987). In field studies of condition indices in wild juvenile cod, recent otolith growth was not highly correlated with fish growth, but it was significantly related to zooplankton biomass, therefore presumably responding to feeding condition (Suthers et al. 1992).

First observations showing the relationship between growth rates over the last 5 d (as determined from the width of otolith daily increments) and protein growth rates based on nucleic acid determinations are available for North Sea plaice (Hovemkamp 1990), but have not been performed on the same individual larva. To our knowledge this study is one of the first to determine otolith microstructure and RNA/DNA ratio analysis on the same larva. After yolk absorption the effect of an external food supply could be measured. The trend started to be visible on Day 7 and increased on Day 11 for RNA/DNA ratios as well as otolith microstructure analysis. High RNA/DNA ratios were coupled with greater increment width in fed larvae. Lower RNA/DNA ratios and smaller increment widths were found in starved larvae. Mosegaard et al. (1988) found uncoupling between somatic and otolith growth rates at hyperoptimal temperatures and therefore suggested that metabolic activity, not necessarily somatic growth rate, governs otolith growth rate. Somatic growth rate results mainly from the balance between protein synthesis and degradation, and hyperoptimal temperatures would accelerate both components, especially degradation, resulting in no somatic growth (Houlihan et al. 1988). However since somatic growth is comparable with components from metabolic rates within the range of optimal temperatures, the RNA/DNA ratio, an index of protein biosynthesis, will be a reflection of metabolic components at an appropriate temperature. Therefore it appears reasonable to use this ratio for examining the relationship between somatic and otolith growth rates, even if otolith growth is a function of metabolic rate.

By coupling RNA/DNA ratio determinations and otolith increment structure analysis on the same larva, it should be possible to determine whether the larva's condition is improving or deteriorating. The validity of these studies for the recruitment problem could be further improved. Whether the findings in this study, determined on laboratory-reared larvae, can also be found in field-caught larvae has to be further evaluated. Results by Cemmensen (1996) on field caught anchovy Engraulis anchoita larvae revealed that the sum of the last increments was correlated with the RNA/DNA ratio, meaning that it should be possible to determine the effect of lack of food in the wild by a combination of biochemical and otolith studies.

Acknowledgements. We thank the staff of the fishery department of the University of Marine Sciences in Kiel and the staff at the Institute of Marine Research in Lysekil for their support. Special thanks to Jana Pickova for rearing of the cod larvae and sharing her expertise. We thank Karin Burkert for helping in the laboratory. We are indebted to Prof. Dr. D. Schnack for his support. Partial funding was provided by the Institute in Kiel, the Deutsche Forschungsgemeinschaft (DFG, NE 99/23-1) and the European Union (EU-AIR 2 96 1226).

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