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RNA:DNA ratios as indicators of nutritional condition in the copepod *Calanus finmarchicus*

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**ABSTRACT:** As part of an investigation of ribonucleic acid (RNA) content as an index of growth and nutritional condition of zooplankton in the field, we describe here a method for measuring RNA and DNA in individuals, Stage N5 through adult, of the copepod *Calanus finmarchicus*. We used the technique to compare total RNA and DNA content and RNA:DNA ratios of *C. finmarchicus* copepodite stages cultured at different food densities. Copepods reared at growth-limiting phytoplankton concentrations (25 µg carbon l⁻¹) were smaller, had lower RNA:DNA ratios, and contained less total RNA and DNA than did copepods reared in excess food (500 µg C l⁻¹). This was true for each stage from C1 to C5. Stages C5 and C4 *C. finmarchicus* collected from Georges Bank and the Gulf of Maine were compared to those from the laboratory experiment. While RNA-DNA ratios of C5 and C4 individuals collected in May and June 1994 were intermediate between the 2 lab treatments, Stage C5 *C. finmarchicus* collected in November 1993 had the lowest RNA-DNA ratios of all copepods sampled. This seasonality of RNA:DNA ratios is most likely related to food availability and changing metabolic activity. Our data show that RNA is a useful index of physiological condition for *C. finmarchicus*.

**KEY WORDS:** Copepods · *Calanus finmarchicus* · RNA:DNA ratios · Nutrition · Growth

**INTRODUCTION**

Measurement of zooplankton growth rates and assessment of their nutritional state are important in studies of secondary production. Zooplankton growth in the field is commonly estimated by incubating animals in ambient seawater and determining the increase of carbon or dry weight during the incubation period (Kimmerer & MacKinnon 1987). Other methods have used molting rates (Heinle 1966, Miller et al. 1984, Runge et al. 1985) or egg production (Checkley 1980, Durbin et al. 1983, Runge 1985) as indices of growth rate. Biochemical growth-rate indicators, such as RNA concentration or the RNA:DNA ratio, are routinely used for estimating growth rates and nutritional condition of larval fish in the field (Buckley 1984, Clemmesen 1988); however, only a few studies have applied these techniques to zooplankton. These studies have produced mixed results regarding the use of RNA as a growth-rate index.

The RNA content of several species of copepods and euphausiids reflects the seasonal pattern in food supply and reproductive activity (Bamstedt 1983, Nakata et al. 1994). A number of researchers have reported a correlation of RNA concentration with growth rate in copepods and other zooplankton. Some have attempted to formulate a general relationship which applies to most invertebrate zooplankton (Sutcliffe 1965, 1969, Bamstedt & Skjoldal 1976, 1980, Skjoldal & Bamstedt 1976), but others have found that the relationship is only valid under certain temperature and food conditions (Ota & Landry 1984), or during certain stages of the life history (Dagg & Littlepage 1972).

Because RNA is an essential component of protein synthesis, its concentration in tissue often reflects the rate of protein synthesis. The RNA:DNA ratio provides an index of protein synthetic capacity per cell since the amount of DNA per cell is assumed not to vary with

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condition or with growth rate (Bulow 1987). The RNA content of tissue is related to growth rate, food density, and temperature and may also be affected by gamete production and developmental stage.

This study is the beginning of a project examining the relationship of RNA content, temperature, and growth rate for *Calanus finmarchicus*, a dominant copepod of the Gulf of Maine and Georges Bank region of the Northwest Atlantic. The first step in this project was to evaluate how food availability affects nucleic acid levels in this species. We examined the variation in total RNA, total DNA, and the RNA:DNA ratio during copepodite stages C1 through C5 in *C. finmarchicus* reared at 2 different food concentrations. In order to measure RNA and DNA of very small individual copepods, we adapted the fluorometric assay of Bentle et al. (1981) to a 96-well microplate format. The resulting technique, the microplate fluorescent assay (MFA), is similar to that of Nacci et al. (1994), who first used a microplate-based RNA-DNA assay with marine animals. Our MFA is capable of measuring nucleic acids in individual copepodite stages and late naupliar stages of *C. finmarchicus*. Copepod samples from the laboratory experiment were used to intercalibrate the MFA with another fluorometric RNA-DNA assay, the flow-injection analysis (FIA) technique of Caldarone & Buckley (1991). Copepod samples collected in the field in 1993 and 1994 were analyzed by the FIA method and their mean RNA:DNA ratios were converted in order to be qualitatively compared with the laboratory experiment.

**METHODS**

**Measuring RNA and DNA: the microplate fluorescent assay.** Nucleic acids from laboratory-grown *Calanus finmarchicus* were measured by using a modification of the Bentle et al. (1981) DNA and RNA assay. This is an ethidium bromide fluorometric technique in which RNA, then DNA, are sequentially degraded with nuclease (RNase and DNase). The concentration of each nucleic acid is determined from the difference in fluorescence measurements taken before and after each enzyme treatment. We found that for *C. finmarchicus*, the DNase step could be eliminated, nearly doubling the number of samples which could be analyzed in 1 d. Adapting the technique to 96-well microplates enabled us to measure the small quantities of DNA and RNA found in individual copepodites and late naupliar stages of *C. finmarchicus* (detection limit 15 ng DNA or RNA per well). The modified Bentle et al. (1981) assay, when combined with the sarcosyl extraction technique described by Caldarone & Buckley (1991), allows 1 person to process up to 140 samples per 8-hour day.

**Working reagents:** Tris buffer: 10 mM Tris, 1 mM Na₂EDTA, pH 7.5. Extraction buffer: 1% N-lauroylsarcosine (sarcosyl) in Tris buffer (Sigma, catalog no. L-5125). Standard buffer: 0.2% sarcosyl in Tris buffer. Ethidium bromide (EB) solution: 2.0 µg ml⁻¹ in Tris buffer (Gene-Mate™ catalog no. 5515-10). Ribonuclease A (RNase): 50 Kunitz Units ml⁻¹ in Tris buffer (Sigma catalog no. R-6513).

**Standards and controls:** Standard DNA ( calf-thymus, Hoefer Scientific TKO-102) and RNA (eukaryotic 18S and 28S ribosomal RNA, Sigma R-0889) stocks were diluted with Tris buffer to approximately 40 µg ml⁻¹. Exact concentrations of the stock solutions were determined spectrophotometrically, and they were aliquotted and stored at -80°C. Working DNA and RNA preparations were diluted in Standard buffer. Working DNA standards ranged from 0.200 µg ml⁻¹ to 2.00 µg ml⁻¹. Standard RNA concentrations ranged from 0.200 µg ml⁻¹ to 8.00 µg ml⁻¹. A *C. finmarchicus* control homogenate was prepared by grinding 0.69 g (wet weight) of adult copepods in a glass homogenizer with 1.00 ml of distilled, deionized water. The homogenate was stored in 5 µl aliquots at -80°C. Each day, an aliquot of the control homogenate was extracted and analyzed along with the copepod samples.

**Protocol:** The copepod samples were extracted by a modification of the sarcosyl extraction technique described by Caldarone & Buckley (1991). The samples were homogenized by vigorous shaking with an appropriate volume of 1% sarcosyl Extraction buffer. The volume of Extraction buffer varied with stage according to Table 1. Samples were shaken for 1 h at room temperature on a vortex mixer equipped with a multiple-vial head. They were then diluted 1:4 with Tris buffer (volumes given in Table 1) to reduce the sarcosyl concentration to 0.2%. The samples were shaken again for 5 min, then centrifuged (12000 × g) for 10 min to sediment any insoluble copepod remains. The control homogenate was extracted in 200 µl of Extraction buffer and was treated as previously described for copepod samples.

Duplicate 75 µl aliquots of controls, unknowns, and working standards were transferred to 96-well microtiter plates (Falcon Pro-bind). Controls were included on each plate so that we could monitor variation between plates. Seventy-five microliters of ethidium bromide (EB) solution was added to each well. The plates were shaken gently at room temperature for 15 min then scanned on a BioTek FL-500 fluorescence plate reader with 530 nm (excitation) and 590 nm (emission) filters. We set the plate reader options so that the probe would stop on each well for 0.1 s and take 10 replicate measurements, presenting the average of the 10 as the fluorescence measurement for the well.
Table 1. Extraction and dilution protocol for the microplate fluorescent assay (MFA). Individual copepods of a given stage are extracted in the specified volume of Extraction buffer, then diluted 1:4 with Tris buffer to yield the final volume given below.

<table>
<thead>
<tr>
<th>Stage</th>
<th>Extraction buffer (µl)</th>
<th>Tris buffer (µl)</th>
<th>Final volume (µl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1</td>
<td>75</td>
<td>300</td>
<td>375</td>
</tr>
<tr>
<td>C2</td>
<td>100</td>
<td>400</td>
<td>500</td>
</tr>
<tr>
<td>C3</td>
<td>150</td>
<td>600</td>
<td>750</td>
</tr>
<tr>
<td>C4</td>
<td>200</td>
<td>800</td>
<td>1000</td>
</tr>
<tr>
<td>C5</td>
<td>300</td>
<td>1200</td>
<td>1500</td>
</tr>
<tr>
<td>C6</td>
<td>300</td>
<td>1200</td>
<td>1500</td>
</tr>
</tbody>
</table>

Following the first scan, 7.5 µl of RNase solution was added to each well, including all standards. The plate was shaken again at room temperature for 20 min and scanned a second time. RNA concentrations were calculated from the difference in fluorescence between the first and second scans and DNA concentrations were calculated from the fluorescence of the second scan (Fig. 1). Standard curves were plotted by using the average fluorescence of the duplicate wells for each concentration. DNA standard values were 0.0150, 0.0450, 0.0750, 0.105, and 0.150 µg well⁻¹; RNA standard values were 0.0150, 0.0750, 0.150, 0.300, 0.450, and 0.600 µg well⁻¹. Unknown values were converted from µg well⁻¹ to µg ind⁻¹ by the calculation

\[
\frac{\mu g \text{ nucleic acid}}{\text{copepod}} = \frac{\mu g \text{ nucleic acid}}{\text{well}} \times \frac{\text{final volume in µl}}{75.0 \mu l}
\]

where the final volume for each stage is that given in Table 1.

Laboratory experiment. *Culturing Calanus finmarchicus*: Copepods were reared in the laboratory from eggs produced by adult female *C. finmarchicus* collected in the Gulf of Maine. Zooplankton were collected near Portsmouth, New Hampshire, USA, in March 1994. Several hundred adult females were isolated and were placed in 20 l polycarbonate buckets at a density of 10 ind⁻¹. They were fed a mixture of cultured phytoplankton (*Prorocentrum micans*, *Gymnodinium sanguineum*, *Heterocapsa triquetra*, and *Tetraselmis* sp.). To start the experimental cultures, 180 females were placed in a 200 µm mesh inside a 20 l polycarbonate bucket containing a total of 20 l of filtered seawater. The tanks were maintained at 12 ± 1°C on a 12 h light:12 h dark light cycle and were gently aerated. The density of the copepod cultures was initially 200 nauplii l⁻¹, and the cultures were diluted as the animals developed to a final density of less than 10 ind. l⁻¹ at stage C5.

Starting at the third naupliar stage, the copepods were fed cultured phytoplankton at excess or growth-limiting cell densities. Tanks in the high-food treatment were fed the equivalent of 500 µg C l⁻¹; tanks in the low-food treatment were fed 25 µg C l⁻¹. Naupliar stages were fed a mixture of equal parts *Gymnodinium sanguineum*, *Heterocapsa triquetra*, and *Tetraselmis* sp. Copepodite stages were fed a mixture of 2 parts *G. sanguineum* and 1 part *H. triquetra*. *Tetraselmis* sp. was eliminated from the diet since the copepods preferred the larger dinoflagellates. Phytoplankton were grown in the same temperature and light conditions as the copepods. Phytoplankton densities in the tanks were adjusted once daily, about 2 h before dark. In the high food treatment, food densities remained at 50 to 75% of the nominal density after 24 h; in the low food treatment, all food was grazed within 24 h.

**Sampling**: Cultures of *Calanus finmarchicus* were sampled throughout the life cycle, but only copepodite stages and adults will be discussed here. Although the sex ratio of the adults was approximately 1 male to 3 females, only females were sampled for RNA and DNA analyses. Males were not sampled because we felt that they were more valuable if kept alive for breeding. The copepod cultures were sampled at intervals of 2 or more days, depending on their development rate; we attempted to sample at least twice during each stage. The high-food tanks were sampled at 2 or 3 d intervals throughout the experiment. Because
the food-limited copepods developed more slowly with time, the sampling interval of the low-food tanks was increased to 3, 4, 5, then finally to 6 d. The cultures were sampled at the same time each day, about 3 h after the lights came on. At each sampling, a volume estimated to contain at least 25 ind. was removed from each culture. The animals were anesthetized with MS-222 (tricaine methane sulfonate; 0.576 g l\(^{-1}\)) and kept in a petri dish on ice during all subsequent sampling steps.

Twenty-five individual copepods were chosen at random. For each copepod sampled, developmental stage was noted and a video image was recorded for later length measurement. Copepods were placed individually in 1.5 ml microcentrifuge tubes, and stored at \(-80^\circ\text{C}\). Prosome lengths for each copepod were determined from the video images by using NIH Image 1.47 image-analysis software. About one-half of the samples were analyzed by the MFA technique described in this paper, the other half were analyzed by a modification of the flow-injection analysis (FIA) technique described by Caldarone & Buckley (1991). The mean RNA:DNA ratios from each stage within each treatment, determined from data collected by both assays, were used to intercalibrate the 2 techniques so that laboratory data generated by the MFA technique could be compared to field data generated by the FIA method.

To check for effects of ingested food on RNA:DNA ratios, we placed 30 well-fed Stage C3 copepodites into filtered seawater for 48 h without food. Thirty animals were kept in excess food. Ten individuals from each treatment were sampled at 0, 24, and 48 h. After 24 h, there was no food visible in the guts of the starved animals. RNA:DNA ratios of the starved group averaged 2.444 (SD 0.108) compared to 2.889 \(\pm\) 0.161 for the controls. Although a 12 h incubation might have been more appropriate, it is clear that the decline over 24 h was too small for ingested food to account for the large treatment differences observed in the experiment.

**Statistics:** All statistical comparisons were done by SAS v 6.0 run on an IBM-compatible personal computer. Since sample sizes varied, analysis of variance was done by using the SAS procedure PROC GLM.

**Sampling *Calanus finmarchicus* in the field.** In order to compare the lab cultures with wild specimens, we collected *C. finmarchicus* in the field on 2 occasions at different phases of the life cycle. Stage C5 *Calanus* were collected in the Gulf of Maine (Wilkinson Basin and Jordan Basin) in November 1993, and Stages C4 and C5 individuals were collected in late May and early June 1994 in the Gulf of Maine (Georges Basin) and Georges Bank. Zooplankton were collected by vertical tows with a 1 m\(^2\) ring net (153 \(\mu\)m or 333 \(\mu\)m mesh size). Depths of tows varied with water depth and the location of *C. finmarchicus* within the water column. Once aboard ship, the zooplankton were anesthetized with MS-222 and sorted. C5 and C4 *C. finmarchicus* were placed individually in 1.7 ml cryovials and were stored over liquid nitrogen. Upon arrival at the lab, they were stored at \(-80^\circ\text{C}\). RNA and DNA in the field-collected copepod samples were measured by the FIA technique of Caldarone & Buckley (1991). This method was found to be unsuitable for nauplii and early copepodite stages with low total RNA content and low RNA:DNA ratios, but worked well for the C4 and C5 animals in this study. Length measurements were not taken for the field-collected copepods.

**RESULTS**

**Analytical results**

Linearity of standards was excellent. \(R^2\) values typically exceeded 0.9950 and often exceeded 0.9990 (Fig. 2). Well-to-well reproducibility was good, coefficients of variation for 6 replicate wells of 3 different samples averaged 3.15 % for DNA and 4.22 % for RNA. Plate-to-plate and day-to-day variability was reasonably low, coefficients of variation of the control homogenate were 6.50 % for DNA and 6.45 % for RNA over 5 mo (49 plates). Known amounts of DNA and RNA (spikes) were added to some of the samples and controls. Eight DNA spikes of 0.500 \(\mu\)g ml\(^{-1}\) averaged 97.0 % recovery (range 85 to 109 %) and 12 RNA spikes of 0.200, 0.500, or 1.00 \(\mu\)g ml\(^{-1}\) averaged 92 % (range 76 to 118 %). Endogenous fluorescence of copepod extracts was measured by replacing the ethidium bro-

![Image](https://via.placeholder.com/150)

**Fig. 2.** Standard curves generated by the FIA for (A) RNA and (B) DNA.
mide solution with Tris buffer and scanning the plate at the usual excitation and emission wavelengths. No endogenous fluorescence was detected in *Calanus finmarchicus* at any stage of development. Residual fluorescence, the fluorescence left over when samples were treated with both RNase and DNase, averaged 8% of the total original fluorescence. As a compromise between accuracy and efficiency, we chose to eliminate the DNase step. Because DNase requires a longer incubation at elevated temperatures, eliminating this step of the procedure nearly doubled the number of samples that could be analyzed in 1 d. It should be recognized that the small residual fluorescence is included in DNA concentrations calculated in this manner.

**Effect of food level on development rate, body size, and nucleic acids**

The well-fed copepods in the laboratory experiment matured and produced viable eggs, but the experiment was terminated before adults had appeared in the low-food treatment. Data were grouped by stage regardless of when they were collected, so that each 'stage' class was made up of individuals collected on more than 1 d. Data are presented for stages C1 through C5 in the low-food treatment, and for C1 through adult females in the high-food treatment. The food-limited copepods developed more slowly than the well-fed copepods, taking about 47 d for 50% of the animals to reach stage C5, compared to 23 d for the well-fed copepods.

Animals in the low-food treatment were smaller at every stage studied, as is shown by their lengths and total DNA contents. The mean prosome length of copepods in the low food treatment increased from 0.65 mm at Stage C1 to 1.75 mm at Stage C5, while the range for the high-food treatment over the same stages was from 0.90 mm to 2.14 mm (Fig. 3A). Total DNA was also different in the 2 treatments, increasing from 0.25 to 0.69 µg ind.\(^{-1}\) in the low food treatment and from 0.30 to 1.02 µg ind.\(^{-1}\) in the high food treatment (Fig. 3B).

Food limitation clearly affected the RNA content and RNA:DNA ratios of copepods in this experiment. Total RNA ranged from 0.11 µg ind.\(^{-1}\) for C1 to 1.05 µg for C5 in the low-food treatment, and from 0.32 to 3.19 µg in the high-food treatment (Fig. 3C). The difference in total RNA content cannot be accounted for solely by differences in body size, since the mean RNA:DNA ratios for each stage also differed among the treatments (Fig. 3D). RNA:DNA ratios increased from 0.46 to 1.48 in the low food treatment and from 1.05 to 3.37 in the high food treatment.

The effects of food treatments, developmental stage, and replicate tanks were compared for each of the 4 dependent variables length, total DNA, total RNA, and RNA:DNA by using 2-way ANOVA (Food × Stage with a third independent variable, Tank, nested within Food). Outcomes of the 4 analyses are presented in Table 2. The effects of food and stage were significant (p < 0.05) for each of the dependent variables. Significant differences between replicate tanks were found only for the dependent variable length. A significant
interaction of tank and stage for all 4 independent variables indicated that within each food-stage combination the replicate tanks sometimes differed even though there was not always an overall tank effect. The interaction of food and stage was significant for total DNA, total RNA, and RNA:DNA. Since for these variables there was a significant food-stage interaction, but no overall effect of replicate tanks, we examined the effect of food at each stage by using a series of t-tests; data from the 2 tanks at each food level were pooled. A significant effect of food level was found at each stage for each of the variables length, total DNA, total RNA, and RNA:DNA (p < 0.01).

RNA:DNA ratios of field-collected copepods

In order to compare field data generated by the FIA technique with laboratory data generated by the MFA, RNA:DNA ratios from the field samples analyzed by the FIA method were converted to predicted values for the microplate fluorometric assay. To calibrate the 2 methods, linear regression was performed on the mean values for each stage within each treatment (Table 3).

Table 3. Comparison of RNA:DNA ratios from the flow-injection analysis (FIA) method of Caldarone & Buckley (1991) with ratios from the MFA. Mean RNA:DNA ratios were used for each developmental stage in each food treatment because different sets of samples were analyzed by the 2 assays. Values in parentheses show sample sizes for each stage-food combination. RNA:DNA<sub>FIA</sub> = 1.333 RNA:DNA<sub>MFA</sub> - 0.0733; r² = 0.981

<table>
<thead>
<tr>
<th>Stage</th>
<th>RNA:DNA&lt;sub&gt;FIA&lt;/sub&gt;</th>
<th>RNA:DNA&lt;sub&gt;MFA&lt;/sub&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low food</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C1</td>
<td>0.252 (14)</td>
<td>0.459 (42)</td>
</tr>
<tr>
<td>C2</td>
<td>0.324 (40)</td>
<td>0.478 (89)</td>
</tr>
<tr>
<td>C3</td>
<td>0.537 (46)</td>
<td>0.783 (90)</td>
</tr>
<tr>
<td>C4</td>
<td>0.928 (27)</td>
<td>1.097 (64)</td>
</tr>
<tr>
<td>C5</td>
<td>1.304 (7)</td>
<td>1.479 (25)</td>
</tr>
<tr>
<td>High food</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C1</td>
<td>1.217 (5)</td>
<td>1.054 (2)</td>
</tr>
<tr>
<td>C2</td>
<td>1.388 (27)</td>
<td>1.723 (11)</td>
</tr>
<tr>
<td>C3</td>
<td>1.856 (22)</td>
<td>2.543 (34)</td>
</tr>
<tr>
<td>C4</td>
<td>2.347 (32)</td>
<td>3.005 (59)</td>
</tr>
<tr>
<td>C5</td>
<td>2.417 (26)</td>
<td>3.367 (83)</td>
</tr>
<tr>
<td>C6</td>
<td>4.026 (12)</td>
<td>5.33 (12)</td>
</tr>
</tbody>
</table>
Calanus finmarchicus in this experiment showed a marked difference between the mean total DNA content of copepods in the 2 treatments (Fig. 3B). The well-fed animals had, depending on stage, from 15% to 50% more DNA per individual than copepods in the food-limited treatment. In vertebrates, total DNA content is considered to represent the number of cells in the organism, regardless of growth rate or condition (Bulow 1987). That is, all cells should contain roughly the same amount of DNA, so variations in DNA content should reflect changes in the number of cells. If the amount of DNA per cell is in fact constant, then the differences in total DNA content observed between the food treatments in this experiment suggest that cell number is higher at each stage for well-fed compared to food-limited C. finmarchicus. There is some evidence that in Calanus and related copepods the total number of nuclei is the same among individuals at any given point in development (McLaren & Marcogliese 1983). If cell number in C. finmarchicus is determinate, then the differences in total DNA observed between the food treatments would indicate differences in the amount of DNA per cell rather than the total number of cells per animal, possibly reflecting different rates of cell division and cell growth in the 2 treatments. More study of the relationship among body size, cell number, and DNA content is needed in order to reconcile the difference in DNA content of copepods in the 2 treatments.

Although the concentration of DNA itself can vary with age and condition, DNA content appears to be an acceptable way of normalizing RNA content in copepodite stages of this species. Since the 2 treatments in this experiment exhibited dramatic differences in RNA:DNA ratios over the range of stages studied (Fig. 3D), we conclude that the RNA:DNA ratio is an acceptable indicator of nutritional condition for C. finmarchicus. For Stages C1 to C4, there was little or no overlap between RNA:DNA frequency distributions for the 2 treatments (data not shown), indicating that it may be possible to infer the nutritional condition of an individual animal given its RNA:DNA ratio and stage.

Comparing data from the laboratory with data from the Gulf of Maine and Georges Bank (Fig. 4) demonstrates that RNA:DNA ratios are useful in monitoring the physiological condition of zooplankton in the field. Based on the time of year and the location of the animals deep in the water column, we expected that the C5 Calanus finmarchicus collected in the Gulf of Maine (Wilkinson Basin and Jordan Basin) in November 1993 would be in a resting stage. Live animals from these samples were brought to the laboratory and did
not feed for more than a week despite the presence of excess phytoplankton in the cultures, confirming that these populations had low levels of physiological activity. RNA:DNA ratios of the C5 C. finmarchicus collected in the field were extremely low, even lower than those of the same stage in the food-limited laboratory cultures. Low RNA:DNA ratios are consistent with our expectation that these animals were in diapause. Stage C5 C. finmarchicus collected from the Gulf of Maine (Georges Basin) and from Georges Bank in May and June 1994 were expected to be active, again based on the time of year and also because they were located near the surface. RNA:DNA ratios of these animals were intermediate between the 2 laboratory food levels, consistent with food densities somewhere between the 2 extremes used in the experiment. C4 individuals from Georges Bank, like C5s from the same area, were also intermediate between the lab high- and low-food treatments. Although the laboratory and field populations could only be compared qualitatively since they were processed by different analytical techniques, the RNA:DNA ratios of C. finmarchicus copepodites in the Georges Bank-Gulf of Maine system follow a seasonal pattern that may be related to food availability or the reduced metabolic activity of diapause, or both.

The variation in RNA:DNA ratios observed in this study reflects the radical changes in growth and metabolic activity over the annual cycle and demonstrates that RNA:DNA ratios can be used as an indicator of nutritional status, and diapause, for Calanus finmarchicus and related species. Our future work will focus on quantifying the relationship between RNA, temperature, and growth rates and on monitoring the seasonal pattern of RNA and growth in greater detail.

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