A high frequency time series at Weathership M, Norwegian Sea, during the 1997 spring bloom: the reproductive biology of *Calanus finmarchicus*

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ABSTRACT: The reproductive biology of *Calanus finmarchicus* was investigated at a permanent station in the Norwegian Sea (Weathership Stn M, 66°N, 2°E) during a time series between March and June 1997. The temporal development of female abundance, egg production rate and gonad development stage in relation to the phytoplankton production cycle is described. Abundance of females, copepodite stage 5 and males as well as female gonad morphology were examined from WP2 net samples taken daily from the upper 100 m. Daily egg production rate and number of spawning females were determined from 50 individual females placed in multiwells or beakers. Once a week a multinet haul from 1000–500–250–100–50–0 m was performed to study the depth distribution of females and gonad development stages. Results show that the reproductive period of *C. finmarchicus* in the Atlantic region of the Norwegian Sea can be subdivided in 3 periods in relation to phytoplankton development:

1. During the prebloom over a period of 49 days mean egg production rate was 8 eggs female⁻¹ d⁻¹ and an average of 47% of the females were mature.
2. Coincident with the bloom in mid May the egg production rate increased up to a maximum of 44 eggs female⁻¹ d⁻¹ while more than 80% of the females were mature.
3. After the bloom at the beginning of June, egg production decreased, and mature females were only rarely found. Feeding experiments indicate that food quantity limited egg production prior to the bloom, while presumably food quality was not sufficient during postbloom. However, due to high female abundance the total population egg production prior to the bloom was the same as during the bloom. This implies that the reproduction of *C. finmarchicus* in the Norwegian Sea is to some extent decoupled from the phytoplankton bloom.

KEY WORDS: *Calanus finmarchicus* Norwegian Sea Egg production Gonad development Phytoplankton

INTRODUCTION

*Calanus finmarchicus* is one of the key species in northern pelagic ecosystems. Detailed knowledge of its life cycle is essential for understanding the population dynamics and quantifying its role in the Norwegian Sea, one of the main centres of distribution (Skjoldal & Rey 1989). Gonad maturation and egg production rate are key parameters for studying reproductive strategy, which plays an important role in adaptation to highly variable environments. Also, the use of egg production rates of herbivorous copepods as a measure of community secondary production has been discussed (McLaren & Leonard 1995, Poulet et al. 1995). Despite its significance there is still a lack of knowledge of the environmental factors controlling reproduction. Several time series focusing on reproductive activity and environmental conditions have been carried out since the initial work of Marshall & Orr (1953) in the Tromsø Sound, Norway, but samples were taken in weekly or biweekly intervals. This sampling frequency does not permit the assessment of short-term changes in reproduction as reproductive activity responds within 2 to 3 days to changes in ambient conditions.
conditions such as temperature or food supply (Hirche et al. 1997). Moreover, previous time series were conducted in coastal areas (Sea of Clyde, UK, Marshall & Orr 1955; lower estuary of the Gulf of St. Lawrence, Canada, Plourde & Runge 1993) or in North Norwegian fjords (Tromsø Sound, Marshall & Orr 1953; Balsfjord, Tande 1982; Malangenfjord, Diel & Tande 1992). In such locations, reproduction and development of a new generation of *C. finmarchicus* is closely related to the phytoplankton bloom (Diel & Tande 1992, Runge & Plourde 1996). In the absence of phytoplankton *C. finmarchicus* is also capable of maintaining egg production based on microzooplankton feeding as shown by Ohman & Runge (1994) in the Gulf of St. Lawrence. Information from the open ocean is scarce and available only from sporadic measurements (e.g. Greenland Sea, Hirche 1990; Fram Strait, Smith 1990; Norwegian Sea, Hirche et al. 1997) or derived from the occurrence of nauplii and eggs (Ostvedt 1955, Skjoldal et al. 1987). To fill this gap, we present here a study of the reproduction of *C. finmarchicus* performed in the Norwegian Sea including field observations and laboratory experiments. The goals of this study were (1) to investigate gonad development and to determine the timing of spawning and the magnitude of egg production at the open sea Weathership Stn Mike, (2) to relate reproduction to phytoplankton development, and (3) to compare the reproductive cycle of *C. finmarchicus* in the open ocean with that in coastal areas and fjords.

**MATERIAL AND METHODS**

**Sampling.** The time series was conducted from 22 March to 9 June 1997, at the Weathership Stn Mike (Stn M) in the Norwegian Sea (66°N, 2°E). Due to drifting, samples were taken in an area of about 20 nautical miles in diameter (Fig. 1), mainly between 2000 and 2500 m bottom depth. During the whole investigation period of 80 d no samples were taken on 17 days due to bad weather conditions or due to monthly port calls.

Two vertical hauls with a WP2 net (mesh size 50 μm) from the upper 100 m were taken daily at about 10:00 h. One haul was fixed in 4% formalin buffered with hexamethylenetetramin for determination of abundance and gonad development stage. From these samples copepodite stage V (CV), males and females were sorted. From the second haul, live females were sorted immediately for egg production experiments. Once a week a vertical multinet (Hydrobios, Kiel; mesh size 150 μm) haul in depth intervals from 1000–500 m, 500–250 m, 250–100 m, 100–50 m and 50–0 m was conducted to study the depth distribution of females.
and gonad development stages (GS). CTD profiles were performed daily from 200 m, and weekly from 1000 m, to the surface. Chlorophyll a concentration (chl a) was measured daily from rosette samples taken at standard depth intervals from 100 m to the surface (for details see Irigoien et al. 1998).

Classification of gonad maturity. Gonad maturity was determined from preserved females from both WP2 net and multinet hauls. From the multinet hauls up to 50 females from each sample were stained with borax carmine, dehydrated and stored in glycerine. GS was classified according to the classification schema of Niehoff & Hirche (1996). Young oocytes stain red, whereas oocytes undergoing final maturation stain pink. This method was slightly modified for the females from WP2 net hauls. Due to excellent fixation, their gonad development stage was determined without previous staining using the natural colour of the oocytes as a criterion. Young oocytes appeared opaque, mature oocytes brown. GS1, 2 and 3 describe immature females of increasing maturity; GS4 characterises mature females ready to spawn. Senescent females were characterised by abnormal oocytes and gaps in the gonads (Niehoff & Hirche 1996) which have ceased reproductive activity.

Egg production. To determine the egg production rate 50 single females were placed at in situ temperature either in multiwell trays (10 ml volume) or in Plexiglas cylinders with net false bottoms suspended in beakers containing 200 ml seawater to separate females and eggs. In the multiwells, eggs were counted and removed at 4 to 12 h intervals. Females were removed from the beakers after 24 h incubation, and the eggs were counted.

Frequently, both multiwells and beakers were used for egg production experiments to test the influence of the experimental set-up on the egg production rate. On the other days only multiwells were used. Runge & Plourde (1994, 1996) suggested that cannibalism occurs in beakers due to wave movement keeping the eggs in suspension. During our study, only 6 out of 48 parallel experiments showed significant differences between egg production rates (eggs female⁻¹ d⁻¹) in multiwells and beakers (ANOVA: 0.05 > p > 0.005). In these cases, only data from the multiwells were used since we supposed them to be more precise due to small control intervals. Otherwise, we pooled the data from the multiwells and the beakers.

Feeding experiments. Feeding and starvation experiments were conducted on 6 occasions to examine the influence of food on the reproduction of *Calanus finmarchicus*. In each experiment 50 single females were placed in Plexiglas cylinders suspended in beakers for 5 d at in situ temperature. One half of the females was kept in filtered seawater (0.45 μm GFC filters), and the other half was fed with the diatom *Thalassiosira weiss-flogii* at concentrations >30 μg chl a l⁻¹. Water was changed daily and egg production checked. At the end of the experiments females were preserved for gonad analysis.

RESULTS

Hydrography

Stn M is situated on the slope of the Norwegian Shelf in the North Atlantic Current. The hydrography is only briefly described here. Details are presented in Irigoien et al. (1998). In the study area, the surface water is of Atlantic origin (Hopkins 1991). According to temperature and salinity measurements, the Atlantic layer mostly extended to 300–400 m depth and did not reduce to less than 150 m depth during the investigation period. Intermediate Water of Atlantic and Polar origin (temperature < 3°C; salinity < 35 psu) underlay the Atlantic Water at depths between 150 and 700 m. The deep water in this region is formed by Arctic Water (review by Hopkins 1991). According to the hydrography, samples in the upper 100 m (WP2 net, Rosette, upper multinet samples) were taken in Atlantic Water whereas multinet samples below 250 m were taken in Intermediate Water, and below 500 m in Deep Water.

At the beginning of this study, the upper 200 m was well mixed, with temperatures between 5 and 6°C. The warming of the surface water to close to 8°C at the beginning of May led to stratification at about 30 to 50 m depth and consequently a phytoplankton bloom developed.

Population development and female maturation

Considering only CV, males and females, the *Calanus finmarchicus* population in the upper 100 m was mainly dominated by females during spring until the end of May (Fig. 2A). During this period the abundance of females varied from 72 to 29,069 females m⁻², with a median of 2513 females m⁻² (Fig. 2B). The mean female size ranged from 2.4 to 2.7 mm prosome length and remained constant during the study. The proportion of CV varied between 20 and 40% in March and April. By the end of May, the proportion of CV had increased up to 95%, which was due to both an increase in the abundance of CV and the low abundance of females, with a median of 567 females m⁻² (min. 15 and max. 989 females m⁻²). The relative abundance of males reached its maximum in April with over 25% and decreased to 2–8% in May and June. Conse-
Weekly vertical sampling showed that females were concentrated in the upper 50 m from the end of March to the beginning of May, except in mid April, when the majority (40%) was caught between 250 and 100 m (Fig. 3). On May 17, females were found in equal proportion in the 50 to 0 and 100 to 50 m depth intervals. At the end of the sampling period (May 25 and 31, June 7), most of the females were found between 30 and 100 m. In depths below 250 m (Fig. 3) females were absent or very rare. The temporal development of the gonads determined from the multinet samples corresponds to the results from daily WP2 net sampling (compare Figs. 2C & 3). However, there were no significant differences in GS of females from different depth intervals: the proportions of the different development stages (GS 1 to 4, senescent) were similar from 250 m to the surface. Below 250 m, often too few females were found to determine the proportions of GS. However, sometimes the GS were evenly distributed through the whole water column, and sometimes there was a tendency for proportionately more immature stages in deeper waters.

Reproduction

According to Irigoien et al. (1998), the investigation period can be divided into 3 periods in relation to phytoplankton development (Fig. 4): (1) During the prebloom period from March until mid May, the chlorophyll a values (integrated over the upper 100 m of the water column). (C) Gonad development (proportion of total population in GS [gonad stage] 1 to 4 and senescent [S]) values varied between 16 and 64 mg m⁻², with an average of 26 mg m⁻². (2) During the phytoplankton bloom (May 10 to 29), the average chlorophyll a concentration was 87 mg m⁻², with a maximum 133 mg m⁻². (3) During the postbloom (May 30 to June 9), the chlorophyll a concentration had decreased to 53 mg m⁻².

The reproductive parameters, egg production rate, clutch size, proportion of spawning females and proportion of mature females (%) GS4 were clearly related to phytoplankton development (Fig. 4). From the end of March until the onset of the phytoplankton bloom, the average egg production rate over 49 d was 8 eggs female⁻¹ d⁻¹ (SD 4 eggs female⁻¹ d⁻¹). Mean clutch size was 19 eggs (SD 7 eggs clutch⁻¹). Clutches <10 eggs were relatively frequent (15%) (Fig. 5). Between 15 and 57% of the females spawned per day (mean 38%), and an average of 47% of mature females (GS4) was found in the WP2 net samples (Fig. 4). With the onset of the spring bloom in mid May, the daily egg production rate increased up to the maximum of 44 eggs female⁻¹ d⁻¹ measured on May 25. During the whole phytoplankton bloom period (19 d), the mean egg production was
24.5 eggs female\(^{-1}\) d\(^{-1}\) (SD 11 eggs female\(^{-1}\) d\(^{-1}\)). Clutch size shifted to higher values; clutches <10 eggs were only rarely found (Fig 5). Hence, the mean clutch size increased to 41 eggs (SD 11 eggs clutch\(^{-1}\)), with a maximum of 56 eggs (Fig. 4). The average proportion of spawning females reached 56%, varying from 83% in the middle of the bloom to 30% at the end. An average of 83% females in GS4 was found, with a maximum of 100% on May 18. With the end of the bloom at the end of May, the reproductive activity of *Calanus finmarchicus* decreased: egg production rate decreased to an average of 3 eggs female\(^{-1}\) d\(^{-1}\) (SD 2 eggs female\(^{-1}\) d\(^{-1}\)) and mean clutch size to 21 eggs clutch\(^{-1}\) (SD 11 eggs clutch\(^{-1}\)) in the postbloom period. Relatively many clutches of 20 to 30 eggs were found. However, the frequency of clutches <10 eggs was low (Fig. 5). Only 12.5% of the females spawned on average in the first week of June. Correspondingly, just 11% of the samples were mature females.

Statistical analysis (ANOVA) confirmed that egg production rate and clutch size were significantly higher during the phytoplankton bloom \((p < 0.0001)\). Significant linear relations were found between the reproduction parameters and the chl a concentration but variance was always very high \((n of each analysis = 40; egg production vs chl a: p < 0.0001, R^2 = 0.422; clutch size vs chl a: p < 0.0001, R^2 = 0.513; % spawning females vs chl a: p = 0.0041, R^2 = 0.202; % GS4 vs chl a: p < 0.0001, R^2 = 0.354)\). This was also the case when chlorophyll values from the previous days were used to...
Fig. 4. *Calanus finmarchicus*. Reproduction at Stn M in relation to phytoplankton development. (A) Integrated total chl a standing stock for the upper 100 m of the water column. (B) *In situ* egg production rate. Points represent mean of 50 replicates. (C) Clutch size. Points represent means of 10 to 56 replicates. (D) Proportion of females spawning within 24 h after capture. (E) Mature females (% GS4) estimated from preserved samples.

Consider the response time of reproduction to changes in food availability. Longer time lags were not expected according to experiments by Hirche et al. (1997). When the analysis was performed with data split up according to prebloom, bloom and postbloom periods, no significant correlation was found between egg production and chlorophyll a concentration in any of the 3 periods. The proportion of spawning females \((n = 17, p = 0.034, R^2 = 0.292)\) and the proportion of GS4 \((n = 17, p = 0.002, R^2 = 0.480)\) were significantly related to chl a concentration only during the prebloom while clutch size versus chl a was significant only during the bloom \((n = 14, p = 0.027, R^2 = 0.345)\).

**Feeding experiments**

As typical examples of the prebloom, bloom and postbloom situations, 3 out of 6 experiments are presented here in detail, to demonstrate the influence of feeding and starvation on reproduction in relation to *in situ* feeding conditions. The results of the other experiments (23 April, 4 May and 23 May) show the same response of egg production to experimental conditions.

During the prebloom period, on April 28 the *in situ* egg production rate was 12 eggs female\(^{-1}\) d\(^{-1}\) (SD 13 eggs female\(^{-1}\) d\(^{-1}\)), about half of the females were mature (GS4; Fig. 6A). After 5 d of feeding, egg production rate increased to 48 eggs female\(^{-1}\) d\(^{-1}\) (SD 30 eggs female\(^{-1}\) d\(^{-1}\)). During this period, the percentage of mature females increased. Non-fed females stopped egg production within 2 d and less than 10% remained mature.
In females sampled during the bloom on May 17 (in situ egg production rate 35 eggs female\(^{-1}\) d\(^{-1}\), SD 27 eggs female\(^{-1}\) d\(^{-1}\); GS 4 100%), the egg production rate increased to 50 eggs female\(^{-1}\) d\(^{-1}\) (SD 29 eggs female\(^{-1}\) d\(^{-1}\)) after 5 d of feeding; 75% of the females were found to be in GS4 after 5 d (Fig. 6B). Under starvation conditions, the egg production decreased dramatically within 2 d to 15 eggs female\(^{-1}\) d\(^{-1}\) (SD 16 eggs female\(^{-1}\) d\(^{-1}\)) and finally ceased on Day 5. Correspondingly, just 12% of the females were in GS4, 35% were senescent, and the rest were immature.

Females collected during the postbloom period on June 2 were mainly in GS3, only 35% were in GS4. The in situ egg production rate was 3 eggs female\(^{-1}\) d\(^{-1}\) (SD 7 eggs female\(^{-1}\) d\(^{-1}\)). After 5 d of feeding the egg production rate increased to 45 eggs female\(^{-1}\) d\(^{-1}\) (SD 28 eggs female\(^{-1}\) d\(^{-1}\)), and 95% of the females were in GS4 (Fig. 6C). In starved females the egg production rate remained as low as the in situ rate. The proportion of mature females had decreased and more senescent females and females in GS2 were found.

**DISCUSSION**

**Reproductive cycle of *Calanus finmarchicus***

The reproductive cycle of *Calanus finmarchicus* in the Atlantic Water of the Norwegian Sea followed 3 periods of phytoplankton development: (1) the pre-bloom with relatively low egg production rates, (2) the bloom with high egg production rates, and (3) the post-bloom with egg production close to zero. This agrees with the prevailing knowledge of the close relationship between the reproduction of *C. finmarchicus* and the phytoplankton concentration (e.g. Marshall & Orr...
short since the phytoplankton bloom occurred shortly

1996). St. Lawrence (Ohman & Runge 1994, Runge & Plourde 1994, Runge & Hirche et al.: mean ca 2.8, range 1, 5-10 (1 wk) at 8°C with surplus food measured in laboratory experiments by Hirche et al. (1997). Since these 2 studies were conducted with larger females (Plourde & Runge: see Table 1, Hirche et al.: mean ca 2.8, range 2.7-3.0 mm [unpubl. data]), the higher maximum rates could be due to female size.

The decrease of egg production and clutch size during postbloom at Stn M matches earlier observations in the Malangenfjord (Diel & Tande 1992). In contrast, significant postbloom reproduction, based mostly on microzooplankton, has been described for the Gulf of St. Lawrence (Ohman & Runge 1994, Runge & Plourde 1996).

Considerable differences exist in the temporal development of the reproductive activity of Calanus finmarchicus between the Norwegian Sea, North Norwegian fjords and the lower estuary of the Gulf of St. Lawrence (Table 1) which are apparently due to the local development of the phytoplankton. In these areas different hydrographical and latitudinal factors control phytoplankton development. In the Norwegian Sea the phytoplankton bloom does not develop before mid May (Halldal 1953, Dale 1995, Irigoien et al. 1998), when warming of the surface water leads to stratification. In the North Norwegian fjords, the spring bloom is driven by the light regime and starts earlier, usually in mid April (Eilertsen & Taasen 1984). In contrast, in the Gulf of St. Lawrence the phytoplankton bloom starts in mid June and persists usually until September (Plourde & Runge 1993, and references therein). Consequently, the spawning activity of C. finmarchicus starts in different months and has a variable duration.

Prior to the spring bloom, mature gonads and spawning activity have been observed occasionally by several authors (Sømme 1934, Marshall & Orr 1955, Davis 1976, Tande & Hopkins 1981). At Stn M, however, more than 50% of female Calanus finmarchicus were mature and maintained low egg production over a period of more than 9 wk. In contrast, in the Gulf of St. Lawrence, Plourde & Runge (1993) found only a few mature females and no egg production for 6 wk during the prebloom, whereas the period of low egg production (<10 eggs female\(^{-1}\) d\(^{-1}\)) lasted for 2 wk only. In the North Norwegian fjords, the prebloom period was short since the phytoplankton bloom occurred shortly after the females had developed from CV (Tande 1982, Diel & Tande 1992). During the prebloom period at Stn M, egg production was strongly food limited, as shown during the feeding and starvation experiments (Fig. 6). Irigoien et al. (1998) suggested that internal lipid reserves were also used for reproduction as ingestion of phytoplankton was not sufficient to cover the costs of egg production. However, the experiments indicate that the food was essential for gonad maturation and egg production of C. finmarchicus and that no eggs are produced from internal reserves alone. This is in accordance with earlier experiments (e.g. Hirche et al. 1997).

Suboptimal production was observed at Stn M during the bloom as well. The maximum egg production rate in this period differed only slightly from that in the Malangenfjord, although temperature at Stn M was about 4°C higher (Table 1). Since the females had the same size at both locations (Table 1), we expected higher rates at the warmer Stn M. The maximum rate reported by Plourde & Runge (1993, see Table 1) is in the range of the egg production rate of 73 egg female\(^{-1}\) d\(^{-1}\) at 8°C with surplus food measured in laboratory experiments by Hirche et al. (1997). Since these 2 studies were conducted with larger females (Plourde & Runge: see Table 1, Hirche et al.: mean ca 2.8, range 2.7-3.0 mm [unpubl. data]), the higher maximum rates could be due to female size.

The decrease of egg production and clutch size during postbloom at Stn M matches earlier observations in the Malangenfjord (Diel & Tande 1992). In contrast, significant postbloom reproduction, based mostly on microzooplankton, has been described for the Gulf of St. Lawrence (Ohman & Runge 1994, Runge & Plourde 1996).
According to our hydrographic data, the results described here represent the reproductive cycle of *Calanus finmarchicus* in the Atlantic Water masses of the North Atlantic Current (NAC), as all of the measurements in the upper 100 m were carried out in Atlantic Waters (Irigoien et al. 1998). Stn M is situated in the central part of a large eddy, where the renewing of the water masses takes place very slowly (Halldal 1953). There was no indication of tongues of coastal water advecting phytoplankton and zooplankton populations from the Norwegian shelf, which was observed during earlier time series studies (Dale 1995). Due to its close relationship with food availability, the reproductive biology of *C. finmarchicus* should strongly react to regional variability in phytoplankton development. In the Norwegian Sea both longitudinal and latitudinal gradients were noted. Phytoplankton development starts in March along the Norwegian coast, where the salinity range maintains stratification throughout the year (Rey 1981). In contrast, in the open ocean bloom development depends on heating of the surface layers and takes place much later. Accordingly, Niehoff & Hirche (1997) registered maximum reproductive activity of *C. finmarchicus* 2 mo earlier than at Stn M. Within the NAC the bloom occurred in the southern part at about 62° N 2 or 3 wk earlier than in the northern part at about 69° N (Braarud et al. 1958). This delay towards the north should also affect the reproductive cycle.

**Food supply and egg production**

The importance of food supply for the reproductive cycle of *Calanus finmarchicus* at Stn M is also supported by the significant relation of ingestion rate and egg production (Irigoien et al. 1998). However, high variance of the correlation between chl a concentration and egg production rate suggests that the chl a concentration is not an ideal parameter for predicting egg production. The significant correlation is presumably due to the large differences between prebloom and bloom values. When data were split up into prebloom, bloom and postbloom periods, no significant correlations were found between the egg production rate and chl a. Plourde & Runge (1993) found no significant correlation either; once the phytoplankton bloom had started in the Gulf of the St. Lawrence. However, at Stn M other reproductive parameters were significantly affected by food concentration in different periods: during prebloom, the proportion of both mature (GS4) and spawning females was correlated with chl a, whereas during the bloom clutch size was related to food concentration. This leads to the hypothesis that the increase of the mean egg production of a population due to increasing food supply occurs stepwise due to (1) an increase of mature females at low food concentration and (2) an increase of clutch size when most females are mature. If this holds true, it might help to explain why chl a is such a poor indicator for egg production rate. Other possible explanations include the patchy distribution of chlorophyll in the water column and the fact that it does not cover heterotrophic microorganisms which can provide an important food resource for reproduction (Ohman & Runge 1994). During our study in the Norwegian Sea the micrzooplankton abundance was very low and contributed only a few percent to the food of *C. finmarchicus* (Irigoien et al. 1998).

Apart from food concentration, food quality is important for reproductive success of *Calanus finmarchicus* (e.g. Marshall & Orr 1955, Runge & Plourde 1996, Ban et al. 1997). Factors such as particle size and species composition influence feeding (for review see Harris 1996). In addition, algal species might have different nutritional values. Thus Runge & Plourde (1996) suggested that dinoflagellates might be more nutritious than diatoms. At Stn M, the cell size and composition of the phytoplankton, and thus the food quality, changed considerably. *C. finmarchicus* feeds on particles > 5 μm (Frost 1972). The contribution of this size fraction to the total phytoplankton stock was below 20% during the prebloom, increased up to 60% during the bloom, and decreased again after the bloom to below 20% (Irigoien et al. 1998). Egg production rate was significantly correlated with the concentration of this size fraction (ANOVA, p = 0.0001), with considerably lower variance (R² = 0.823) compared to the correlation between egg production rate and total phytoplankton stock (R² = 0.577). The prebloom community was dominated by green algae and prymnesiophytes, the bloom by diatoms, and the postbloom by cryptophytes and again prymnesiophytes (Irigoien et al. 1998). Meyer-Harms et al. (1999) observed an evident preference for dinoflagellates during the prebloom, and for diatoms (mainly *Rhizosolenia* spp.) during the bloom. Statistical analysis shows significant correlations between egg production and concentrations of dinoflagellates and diatoms (Table 2). However, dinoflagellates were found only at low concentrations (2 to 3% of the chl a) at Stn M, while diatoms contributed up to 33% during the bloom and constituted an important food component of *C. finmarchicus* (Meyer-Harms et al. 1999). A reduction of egg production rate by diatoms, as discussed recently (Ianora et al. 1995, Ban et al. 1997) could not be proven in this study.

In the postbloom period, low egg production rates were observed, although the phytoplankton concentration in the water column was relatively high, and the ingestion of phytoplankton carbon exceeded the
Table 2. Statistical analysis (linear regression, ANOVA) of the correlation between the egg production rate of *Calanus finmarchicus* and the concentration of different algal species (data of pigment concentration from Irigoien et al. 1998) during a time series at Stn M (Norwegian Sea)

<table>
<thead>
<tr>
<th>Algal group</th>
<th>p</th>
<th>R²</th>
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<tbody>
<tr>
<td>Dinoflagellates</td>
<td>0.003</td>
<td>0.601</td>
</tr>
<tr>
<td>Diatoms</td>
<td>0.042</td>
<td>0.352</td>
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<tr>
<td>Chlorophyceae</td>
<td>0.051</td>
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<tr>
<td>Prymnesiophytes</td>
<td>0.938</td>
<td>0.0062</td>
</tr>
</tbody>
</table>

Costs for egg production (Irigoien et al. 1998). Food quality presumably limited egg production during the postbloom period when prymnesiophytes, mainly *Phaeocystis pouchetii*, dominated the phytoplankton stock (Meyer-Harms et al. 1999) since females fed in the laboratory during postbloom matured and commenced spawning. This indicates that no internal regulation reduced egg production after the bloom as suggested by Diel & Tande (1992). Recent field studies (Plourde & Runge 1993, this study) and laboratory observations (Hirche 1990, Niehoff unpubl, data) have shown that the duration of the reproductive period of *Calanus finmarchicus* is extremely variable and depends on food conditions rather than on genetic determination.

**Reproductive strategy of *Calanus finmarchicus***

The capability of *Calanus finmarchicus* to mature and to maintain egg production at low food concentrations enhances the adaptation to a highly variable environment in the following ways:

1. The final maturation processes with respect to metabolic pathways, including the processes of vitellogenesis 1 and 2 and maturation division (Niehoff & Hirche 1996), are established and enable the females to rapidly respond to either sudden local food maxima or to developing phytoplankton blooms. Thus, in feeding experiments during prebloom, egg production increased after 3 d. The high frequency sampling during our study shows that *Calanus finmarchicus* reacted immediately to the spring bloom. Runge & Plourde (1996) described a time lag of 8 d between the onset of the phytoplankton bloom and the increase of egg production from zero to maximum values, and suggested that this time was needed for oocyte maturation. In the Gulf of St. Lawrence, the majority of these females was immature during the prebloom period (Plourde & Runge 1993), indicating that the food supply was not sufficient for maturation and thus causing a longer lag phase compared to Stn M.

2. Egg production during the prebloom extends the reproductive period of *Calanus finmarchicus* considerably. Thus, at Stn M it was more than twice as long as in North Norwegian fjords (Tande 1982, Diel & Tande 1992). It is also important to point out that due to the high abundance of females during the prebloom period, the egg production of the population was high in spite of low individual egg production rates (Fig. 7). During the prebloom period, *C. finmarchicus* females produced on average 34 187 eggs m⁻² d⁻¹, which is as high as during the bloom (34 855 eggs m⁻² d⁻¹). Assuming an egg carbon content of 0.23 pg (Ohman & Runge 1994), the secondary production of the females is 7.86 mg carbon m⁻² d⁻¹. Since the prebloom was more than twice as long as the bloom period, more eggs were actually produced when phytoplankton was rare than during the period of high reproductive activity during the bloom.

The decoupling of the reproduction of *Calanus finmarchicus* from the phytoplankton bloom may enhance the chance for reproductive success of the population. The relevance of early production for the population has to be revealed by data on abundance of nauplii and young copepodes giving information on hatching, recruitment success and mortality. However, eggs and nauplii of *C. finmarchicus* are known to be an important food resource for fish larvae (e.g. Runge 1988, Lough & Mountain 1996). Hence, the high secondary production by *C. finmarchicus* could provide food for the larvae of the herring (Hardy 1924, cited in Raymont 1983) migrating through this area (Misund et al. 1997) months before the phytoplankton bloom starts.

![Fig. 7 *Calanus finmarchicus*. Egg production rate of the population during the time series between 22 March and 9 June 1997](image-url)
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