Feeding and growth by ephyrae of scyphomedusae  
Chrysaora quinquecirrha

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ABSTRACT: Clearance rates of Chrysaora quinquecirrha ephyrae were quantified in the laboratory using monocultures of different plankton organisms as prey, as well as mixed natural zooplankton assemblages. Further, growth of ephyrae was measured at different densities of rotifers or ctenophores as prey. The ciliate Strobilidium sp. (strain CRE) and the rotifer Brachionus plicatilis were cleared in monocultures at medium rates: 4 ml h⁻¹ ind⁻¹ and 1 ml h⁻¹ ind⁻¹ respectively. Clearance of copepod nauplii was low (0.5 ml h⁻¹ ind⁻¹) and the dinoflagellates Gymnodinium sanguineum were not eaten by the ephyrae. In a mixed zooplankton assemblage, rotifers and copepod nauplii were cleared at rates similar to those obtained in monocultures, and tintinnids also were eaten. In contrast, larvae of the ctenophore Mnemiopsis leidyi were cleared at a much higher rate (30 ml h⁻¹ ind⁻¹) than microzooplankton. When the rotifer B. plicatilis were offered in excess to ephyrae, a maximum specific growth rate of 0.3 d⁻¹ was measured. Growth rate more than doubled (0.7 d⁻¹) when ctenophore larvae were offered. The growth rate obtained on ctenophore larvae is the highest rate reported so far for any scyphomedusan species. We suggest that M. leidyi may be of critical importance for the high growth of C. quinquecirrha in natural populations during springtime in Chesapeake Bay, USA.

KEY WORDS: Scyphomedusae, Chrysaora quinquecirrha, Ctenophore, Predation, Growth

INTRODUCTION

Scyphomedusae are large conspicuous predators of zooplankton in many coastal areas. The sea nettle Chrysaora quinquecirrha is found from the southern coast of New England (USA) to the tropics along the east coast of North America (Mayer 1910). In the mesohaline region of Chesapeake Bay (USA), C. quinquecirrha becomes extremely abundant in late spring and summer (Purcell 1992, Purcell et al. 1994a, b) and negatively affects aquatic recreational activities there. Observed changes in the plankton community (Feigenbaum & Kelly 1984), a model of Chesapeake Bay food webs (Baird & Ulanowicz 1989), and recent trophic studies (Purcell 1992, Purcell et al. 1994a, b) all suggest that C. quinquecirrha medusae are important in the plankton dynamics of the bay.

Most previous studies of this species have been done on the benthic polyp stage (e.g. Cargo & Schultz 1966, 1967, Cones 1969, Loeb 1972, Loeb & Blanquet 1973, Cargo & Rabenold 1980), or on the abundance and feeding ecology of the medusa stage (Clifford & Cargo 1978, Feigenbaum & Kelly 1984, Cargo & King 1990, Purcell 1992). The adult medusa of Chrysaora quinquecirrha is known to feed on a wide variety of zooplankton, with copepods, ichthyoplankton and ctenophores being key components of the diet (Purcell 1992, Purcell et al. 1994a, b).

Ephyrae of Chrysaora quinquecirrha are budded from the polyps in spring when water temperatures rise above 17°C. There is little information on feeding of the ephyrae of any jellyfish species. Purcell (1992) summarizes data from Haven & Morales-Almo (1973) showing that protozoa and rotifers were 63 and 23% of the prey caught by ephyrae of C. quinquecirrha in natural water samples.
Similarly, very few measurements exist on growth rates of ephyrae of jellyfish species. Olesen et al. (1994) measured that ephyrae of Aurelia aurita attained a maximum growth rate of 0.3 d⁻¹, when feeding on the rotifer Brachionus plicatilis in the laboratory. This value was close to the maximum values reported for field populations of A. aurita (Hernroth & Groendahl 1983, Van der Veer & Oorthuysen 1985), and 3 times higher than the maximum growth rate of a food limited natural population in the shallow fjord Kerting Nor, Danmark (Olesen et al. 1994). In contrast, when Chrysaora quinquecirrha ephyrae were fed Artemia salina larvae, polychaetous annelids, or strained ctenophores in the laboratory, growth rates were low compared to apparent rates in situ (Cargo & Schultz 1966, 1967).

This paper reports on the feeding rates of Chrysaora quinquecirrha ephyrae on a variety of zooplankton organisms, offered in monocultures and in mixed zooplankton assemblages. Growth rates of ephyrae feeding on rotifers and on ctenophores are compared, and the relative importance of the different zooplankton organisms as food sources for the ephyrae of C. quinquecirrha are discussed.

**MATERIALS AND METHODS**

**Production of ephyrae.** Ephyrae of Chrysaora quinquecirrha for experimental purposes were raised in the laboratory. Oyster shells bearing sessile stages (cysts and polyps) of C. quinquecirrha were collected from Broad Creek, a tributary of Chesapeake Bay, in March 1993 and kept in estuary water (12% S) at 14°C. Water was changed and the polyps were fed twice a week with newly hatched Artemia salina nauplii. Excystment and strobilation were initiated by raising the temperature to 21°C. The ephyrae used in experiments were from 3 to 10 d old and 1 to 4 mm in diameter.

**Clearance rates of microplankton in monoculture.** Five different prey types were used to estimate clearance rates of Chrysaora quinquecirrha: a dinoflagellate, Gymnodinium sanguineum (50 μm), a ciliate, Strobilidium sp. (35 to 40 μm), and a rotifer, Brachionus plicatilis (200 μm), which were cultured in the laboratory, and copepod nauplii (150 to 300 μm) and ctenophores Mnemiopsis leidyi (1 to 2 mm larvae), which were collected in a zooplankton net from the Choptank River tributary of Chesapeake Bay. The dinoflagellate G. sanguineum, a common species in Chesapeake Bay (Bockstahler & Coats 1993), was cultured in the medium t/2 minus silica (Guillard 1975) made with estuarine water (12 to 15%) at 20°C on a 14:10 h light:dark cycle. The ciliate Strobilidium sp. (strain CRE) was isolated from Choptank River, and grown under the same temperature and light conditions in estuarine water amended with 0.1 ml l⁻¹ of t/2 iron-EDTA trace metal solution with algae (Isochrysis galbana and Heterocapsa triquetra) added as prey.

From 2 to 15 ephyrae were incubated for 8 to 24 h in 0.5 or 1.0 l polycarbonate bottles with filtered (20 μm) estuary water (12% S). Prey was added from a stock solution. The prey density of this stock solution was determined after settling of subsamples as described by Hasle (1978). An inverted microscope was used for counting. Ten or 12 bottles were used in each experiment, and the decrease in prey organisms as a function of time were followed by processing 2 to 6 bottles at appropriate intervals. When copepod nauplii and rotifers were used as prey, water was filtered through a 20 μm plankton gauze. The retained prey organisms were preserved in 5% Bouins solution, and later counted by using a dissecting microscope. When ctenophores were used as prey, they were counted live at the end of the incubation. When ciliates or dinoflagellates were used as prey, we stopped the incubation by adding 30 ml of 100% Bouins solution directly into the experimental bottle, after which subsamples (10 ml for dinoflagellates and 25 ml for ciliates) were taken with a Hensen stempel pipette. These subsamples were then settled and counted as described above.

The incubation periods were 8 to 24 h. At the end of each experiment, ephyrae were placed with the sub-umbrellar side down, and the interradial diameter was measured. Further, a subsample of 30 ephyrae from each experiment was examined in order to measure the average number of prey present in the gut of each individual by the end of the incubation. From 2 to 4 bottles in each experiment served as controls (without ephyrae). The experimental conditions are summarized in Table 1.

**Clearance (C)** was calculated from the equation:

\[ C = \frac{V}{n} \ln \left( \frac{C_0}{C_1} \right) \]  

where \( V \) is volume of the experimental bottle (ml); \( n \) is number of ephyrae; \( t \) = incubation time (h); \( C_0 \) and \( C_1 \) are number of prey organisms at times 0 and \( t \), respectively.

All bottles were incubated in dim light and a test experiment revealed that there was no aggregation of prey items: when bottles were rotated on a plankton wheel (1 rpm) this did not affect the outcome of the experiments.

**Clearance on mixed zooplankton.** The clearance by Chrysaora quinquecirrha ephyrae of natural microzooplankton in mixed assemblage was also estimated. Microzooplankton was collected from the Choptank River in August 1993. Twelve liters of estuary water was back-filtered (20 μm) to concentrate plankton organisms in order to obtain an adequate number at
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Table 1. Chrysaora quinquecirrha. Experimental conditions during clearance measurements using monoculture microplankton organisms. Salinity was 12%.

<table>
<thead>
<tr>
<th>Prey</th>
<th>Temp (°C)</th>
<th>Container volume (ml)</th>
<th>Incubation period (h)</th>
<th>Number of ephyrae</th>
<th>Ephyrae diameter (mm ± SD)</th>
<th>Prey density (ind. ml⁻¹)</th>
<th>Prey size (mm ± SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gymnodinium sanguineum</td>
<td>24</td>
<td>500</td>
<td>8</td>
<td>15</td>
<td>1.92 ± 0.38</td>
<td>40</td>
<td>0.05</td>
</tr>
<tr>
<td>Strobilidium sp. (strain CRE)</td>
<td>23</td>
<td>500</td>
<td>8</td>
<td>15</td>
<td>2.05 ± 0.42</td>
<td>8.8</td>
<td>0.035</td>
</tr>
<tr>
<td>Brachionus plicatilis</td>
<td>23</td>
<td>500</td>
<td>8</td>
<td>15</td>
<td>1.77 ± 0.45</td>
<td>2</td>
<td>0.2</td>
</tr>
<tr>
<td>Copepod nauplii</td>
<td>24</td>
<td>500</td>
<td>23</td>
<td>10</td>
<td>1.74 ± 0.54</td>
<td>0.2</td>
<td>0.15–0.3</td>
</tr>
<tr>
<td>Ctenophore (1)</td>
<td>23</td>
<td>500</td>
<td>12</td>
<td>2</td>
<td>3.27 ± 0.51</td>
<td>0.04</td>
<td>1.5 ± 0.4</td>
</tr>
<tr>
<td>Ctenophore (2)</td>
<td>23</td>
<td>1000</td>
<td>24</td>
<td>2</td>
<td>3.60 ± 0.57</td>
<td>0.02</td>
<td>1.6 ± 0.4</td>
</tr>
<tr>
<td>Ctenophore (3)</td>
<td>23</td>
<td>1000</td>
<td>24</td>
<td>2</td>
<td>2.80 ± 0.44</td>
<td>0.01</td>
<td>1.6 ± 0.4</td>
</tr>
</tbody>
</table>

the start of the experiment. The concentrate was added to 6 l of unfiltered water from Choptank river (12.5% S). The 4 dominant types of microzooplankton were identified and counted: aloricate ciliates, tintinnids, copepod nauplii and rotifers.

Fourteen incubation polycarbonate bottles (0.5 l) were filled with the microplankton sample, and kept in dim light at 24°C. Thirty ml of 100% Bouins was then added to 4 of the 14 bottles, and two 20 ml subsamples were taken from each bottle. These subsamples were then settled and counted as above to determine initial densities. Of the remaining 10 bottles, 20 ephyrae were added to each of 6 bottles (experimental bottles), and the last 4 bottles served as controls without ephyrae (control bottles). After 12 h, 2 of the experimental bottles were stopped by adding 30 ml of 100% Bouins and 10 ephyrae from each were examined using a dissecting microscope in order to count the average number of prey present in the gut of each individual. After 24 h, the remaining 4 experimental bottles and the 4 control bottles were stopped by adding 30 ml 100% Bouins. Two 20 ml subsamples were finally taken from each of the experimental and control bottles in order to count the final number of prey organisms. Before Bouins was added to 2 experimental bottles, 20 ephyrae from each were isolated and the umbrella diameter was measured as previously described. The experiment was initiated within 2 h of when the water sample was collected. Clearance was calculated according to Eq. (1), where \( C_0 \) = total number of prey at \( t = 0 \) and \( C_t = C_0 - (\text{total number of prey in control bottles at } t = 24) - \text{total number of prey in experimental bottles at } t = 24 \).

**Gut contents of ephyrae sampled in Chesapeake Bay.** Thirty ephyrae were collected in tributaries of Chesapeake Bay for analyses of their gut contents. Four stations (the Tred Avon River, Broad Creek, and 2 stations in the Choptank River) were sampled every second week from 21 May to 25 August. Ephyrae were collected in surface tows of a 0.75 m diameter plankton net (333 μm mesh size) with a 0.5 l cod end. The live samples were resuspended in a 20 l bucket with estuarine water, and the ephyrae removed individually. The ephyrae were carefully rinsed in 20 μm filtered water (12.5% S), and then preserved in 5% Bouins for later analysis using a dissecting microscope.

**Growth rates of ephyrae.** Growth of ephyrae was measured in 3 separate experiments using 3 l plastic buckets. In Expt 1, 5 buckets with different densities of rotifers (94 to 1500 ind. l⁻¹) and 20 μm filtered estuary water were used. In Expt 2, ctenophore larvae Mnemiopsis leidyi were added to each of 6 buckets at densities of 0.3 to 23 ind. l⁻¹, and in Expt 3, both rotifers and ctenophores were added at densities of 94 to 1500 rotifers l⁻¹ and 3 to 50 ctenophores l⁻¹.

In all 3 experiments, the incubation period was 4 d. The ephyrae were transferred daily to new water with new prey organisms at the same initial density. After each 24 h period, the water in each bucket was filtered through a 20 μm plankton gauze and the retained prey organisms were preserved in 5% Bouins solution to be counted later by using a dissecting microscope.

A control for Expt 3 was conducted in order to test if rotifers were eaten by the ctenophores. Three buckets with rotifers Brachionus plicatilis (750 ind. l⁻¹) were incubated for 24 h together with ctenophore larvae (10 ind. l⁻¹). The numbers of rotifers were unchanged by the end of the incubation period.

Clearance (C) was calculated according to Eq. (1), and the instantaneous specific growth rate (μ), ingestion (I), production (P) and net growth efficiency (NGE) calculated according to the following equations:

\[
\mu = \ln(W_t/W_0) / t^{1} \quad (2)
\]

\[
I = (C_0 - C_t)/n \quad (3)
\]

\[
P = B_0 - B_t \quad (4)
\]

\[
NGE = P/A = P/I.A_{\text{eff}} \quad (5)
\]

where \( W_0 \) and \( W_t \) are mean dry weight (mg) of ephyrae at time 0 and time t respectively; \( t \) is incubation time (h); \( C_0 \) and \( C_t \) are numbers of prey organisms at time 0 and time t; \( n \) is number of ephyrae; \( B_0 \) and \( B_t \) are the mean carbon content of medusae at time 0 and time t.
respectively; \( A \) is assimilation; and \( A_{ass} \) is the assimilation efficiency. The biometric conversion factors and the assimilation efficiency used in the calculations are given in Table 2.

### Biometric conversion factors for ctenophores.

In order to calculate net growth efficiency (NGE), the relationship between ctenophore length and dry weight was established; 40 ctenophore larvae were allowed to empty their guts, and the length of each individual (1 to 13 mm) was then measured with an optical micrometer on a dissecting microscope. The ctenophores were then dried to constant weight in pre-weighed aluminium pans at 60°C and finally weighed to the nearest 1 μg on an electronic balance. In order to determine the carbon content as a percentage of dry weight, 6 groups of 5 ctenophores (2 to 12 mm) with empty guts were carefully rinsed in filtered water (10% S), placed on pre-ashed, pre-weighed GF/C filters, dried to constant weight at 60°C and weighed to the nearest 1 μg. The carbon content was measured using an elemental C/N analyzer. These values for *Mnemiopsis leidyi* and conversions for *Chrysaora quinquecirrha* are given in Table 2.

### Digestion rates.

The digestion times for *Chrysaora quinquecirrha* ephyrae feeding on rotifers *Brachionus plicatilis* and on ctenophore larvae *Mnemiopsis leidyi* were measured. Ephyrae were maintained in 500 ml glass bowls for 12 h at ambient water temperature (23 to 24°C) in 20 μm filtered estuary water (12% S) with Artemia salina nauplii as prey, so that their guts would empty of rotifer or ctenophore prey. *B. plicatilis* or *M. leidyi* were then added to the glass bowls with ephyrae. After a 15 min feeding period, each ephyra was carefully examined for prey. If prey were present, the ephyra was isolated in a small beaker with 20 μm filtered estuary water (12% S) at 23°C, and prey in the gut examined at 15 min intervals until prey could not be identified. The digestion times were averaged to give a digestion time of *C. quinquecirrha* ephyrae feeding on *B. plicatilis* and on *M. leidyi* (accuracy = ±30 min).

## RESULTS

### Clearance rates of microplankton in monoculture

The dinoflagellate *Gymnodinium sanquineum* was not eaten by ephyrae of *Chrysaora quinquecirrha* (Fig. 1A). The numbers of prey left in the experimental bottles (with ephyrae) compared with the control bottles (without ephyrae) were not significantly different at the end of incubation (\( p > 0.05 \), t-test). Moreover, no prey were found in the guts of ephyrae.

In contrast, both ciliates *Strobilidium* sp. and rotifers *Brachionus plicatilis* were cleared by *Chrysaora quinquecirrha* ephyrae (Fig. 1B, C) (\( p < 0.001 \)). Clearance on *Strobilidium* sp. and *B. plicatilis* was estimated to be 4 and 1 ml h\(^{-1}\) ind.\(^{-1}\), respectively. The feeding rates on rotifers were calculated by 2 different methods: (i) according to Eq. (1), and (ii) from the average gut contents of ephyrae by the end of incubation (10 ± 5 rotifers ephyrae\(^{-1}\) d\(^{-1}\)) and the digestion time of the prey (5.2 h, Table 3). By using method (i), it was found that each ephyra had removed an average of 15 rotifers during 8 h incubation. This value is close to the value of 11 obtained when using method (ii). Gut contents were not examined with *Strobilidium sp.* as prey.

Clearance rates of copepod nauplii by *Chrysaora quinquecirrha* ephyrae were low (0.55 ml h\(^{-1}\) ind.\(^{-1}\)) (Fig. 1D). However, after 24 h of incubation, the experimental bottles contained significantly fewer copepod nauplii than the control bottles without ephyrae (\( p < 0.005 \)). Based on the differences between control and experimental bottles, the feeding rate was 2.3 nauplii ingested ephyrae\(^{-1}\) d\(^{-1}\) calculated from the average gut content of 0.3 ± 0.5 nauplii ephyrae\(^{-1}\).

Ephyrae of *Chrysaora quinquecirrha* also fed on larvae of the ctenophore *Mnemiopsis leidyi* (Fig. 1E). The numbers of prey remaining in experimental and

### Table 2. Biometric conversions. C: carbon (mg); DW: dry weight (mg); Diam: live diameter (mm); L: live length (mm)

<table>
<thead>
<tr>
<th>Ctenophore Species</th>
<th>DW = 0.0021 Diam(^{0.94}), ( r^2 = 0.94 ) (Purcell 1992)</th>
<th>Assimilation efficiency = 90% (size range 14–100 mm diam)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Mnemiopsis leidyi</em></td>
<td>( C = 0.000215 \text{Diam}^{0.93}, r^2 = 0.92 ) (Purcell 1992)</td>
<td>(This study)</td>
</tr>
<tr>
<td><em>Brachionus plicatilis</em></td>
<td>( C/\text{DW} = 0.088 \pm 0.007, n = 30 ) (size range 2–12 mm)</td>
<td>(This study)</td>
</tr>
</tbody>
</table>

### Table 3. *Chrysaora quinquecirrha*. Digestion time (h) for ephyrae fed on ctenophore *Mnemiopsis leidyi* and rotifer *Brachionus plicatilis*. n = number of prey followed

<table>
<thead>
<tr>
<th>Prey</th>
<th>Size of ephyrae (mm)</th>
<th>Size of prey (mm)</th>
<th>Digestion time</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>B. plicatilis</em></td>
<td>3.1 ± 1.2</td>
<td>0.2</td>
<td>5.28 ± 2.48</td>
<td>21</td>
</tr>
<tr>
<td><em>M. leidyi</em></td>
<td>2.8 ± 0.7</td>
<td>1.5 ± 0.3</td>
<td>2.22 ± 0.72</td>
<td>10</td>
</tr>
</tbody>
</table>
control bottles were significantly different in all 3 experiments (p < 0.001). The ephyrae could ingest ctenophores one-half their own size (Table 1). If the ctenophore was too large to enter the gut, it was digested in the oral lips or even outside the ephyrae. Feeding by the ephyrae was saturated when the initial prey concentration was 0.04 and 0.02 ind. ml⁻¹, but no saturation was observed at 0.01 ind. ml⁻¹. At that prey density clearance was 30 ml h⁻¹ ephyra⁻¹ (Fig. 1E).

**Clearance on mixed zooplankton**

The number of prey in natural zooplankton assemblages decreased in all incubation bottles over 24 h, including the controls without ephyrae (Table 4). However, the decrease in prey organisms was more pronounced in the bottles with ephyrae than in control bottles without ephyrae, except for ciliates. The difference between the number of prey in grazed bottles and the number in control bottles at t = 24 h was interpreted as removal due to predation by ephyrae. Significant grazing by ephyrae was observed on rotifers *Synchaeta* sp. (p < 0.01, t-test), copepod nauplii, and loricate ciliates (tintinnids) (p < 0.02), but grazing was not detected on aloricate ciliates. Clearance was estimated to be 1.3 ml h⁻¹ ind⁻¹ on rotifers and 0.3 ml h⁻¹ ind⁻¹ on nauplii and tintinnids. Rotifers were the predominant prey type in gut contents of ephyrae at 12 h, but all 4 prey types were found in the guts (Table 4).
Table 4. *Chrysaora quinquecirrha*. Removal of prey from natural zooplankton assemblages by ephyrae. Eight control bottles were incubated (4 were stopped at t = 0 and 4 at t = 24). Six experimental bottles were incubated and stopped at t = 24. Prey densities and total number of prey are means ± SD. Differences in average prey densities between control and grazed treatments at the end of each incubation are presented. Negative differences indicate removal of prey by ephyrae; difference significant at p < 0.01 (*), at p < 0.02 (**)

<table>
<thead>
<tr>
<th>Prey type</th>
<th>No. of prey ephyrae (^{-1})</th>
<th>SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aloricate ciliates</td>
<td>1.43</td>
<td>±0.17</td>
</tr>
<tr>
<td>Rotifers</td>
<td>0.77</td>
<td>±0.77</td>
</tr>
<tr>
<td>Copepods</td>
<td>0.70</td>
<td>±0.26</td>
</tr>
<tr>
<td>Tintinnids</td>
<td>0.23</td>
<td>±0.50</td>
</tr>
<tr>
<td>Copepod nauplii</td>
<td>0.17</td>
<td>±0.45</td>
</tr>
<tr>
<td>Fish larvae</td>
<td>0.07</td>
<td>±0.36</td>
</tr>
<tr>
<td>Polychaete larvae</td>
<td>0.03</td>
<td>±0.18</td>
</tr>
<tr>
<td>Bivalve veligers</td>
<td>0.03</td>
<td>±0.18</td>
</tr>
</tbody>
</table>

Gut contents of ephyrae sampled in Chesapeake Bay

Rotifers, aloricate ciliates, tintinnids and copepod nauplii were found in gut contents of ephyrae collected in Chesapeake Bay (Table 5). Those prey, together with adult copepods, were the dominant prey items (96%) found in ephyrae 4 mm in diameter. The high number of ciliates was due to the presence of 40 ciliates in 1 ephyra. No attempt was made to quantify phytoplankton, but the numbers were low.

Growth experiment

Ephyrae of *Chrysaora quinquecirrha* showed increasing growth rate as the number of rotifers *Bra- chionus plicatilis* available increased (Fig. 2A). A maximal growth rate (\(\mu_{\text{max}}\)) of 0.32 d\(^{-1}\) was obtained when about 375 rotifers \(1^{-1}\) were added each day, and the growth rate did not increase further at higher prey levels. At the maximal growth rate, the mean diameter of ephyrae increased from 2.2 mm to 3.6 mm during the 4 d incubation, corresponding to an increase in mean dry weight from 0.02 to 0.08 mg (Table 6).

When ctenophore larvae were used as prey, the growth rate also increased with increased prey level, and the \(\mu_{\text{max}}\) obtained was considerably higher. At 23 ctenophore \(1^{-1}\), a \(\mu_{\text{max}}\) of 0.66 d\(^{-1}\) was measured, and the mean diameter of ephyrae increased from 3.3 to 8.8 mm, corresponding to an increase from 0.06 to 0.85 mg dry weight (Table 6). When both rotifers and ctenophores were added, the growth pattern was similar to the experiment where ctenophores were added alone, and the maximal growth rate obtained was only slightly higher (0.7 d\(^{-1}\)) (Fig. 2A).

Ingestion \((I)\) of rotifers increased linearly with prey concentration (Fig. 2B), regardless of the presence of ctenophores in the incubation bottles. We did not calculate a relationship between ingestion of ctenophores and prey density because all ctenophores in most growth experiments were consumed.

Clearance on rotifers was constant with increasing prey level averaging about 22 ml h\(^{-1}\) ind\(^{-1}\), both when rotifers were added alone and when rotifers were added with ctenophore larvae (Fig. 2C) (slope of line, \(b = 0, p = 0.56\)). When ctenophore larvae were added at low densities (with or without rotifers), all larvae were eaten by the end of 24 h incubation (Table 6). Therefore clearance could only be calculated at the highest prey concentration. Clearance rates on ctenophore larvae decreased from 55 ml h\(^{-1}\) ind\(^{-1}\) at 3 prey \(1^{-1}\) to 20 ml h\(^{-1}\) ind\(^{-1}\) at 50 prey \(1^{-1}\) (Fig. 2C).

For ephyrae showing positive growth, the net growth efficiency (NGE) varied between 2 and 25% with the highest NGE at medium prey levels (Table 6). On average, NGE was lower with ctenophores added alone (4.9 ± 1.8%) than in the experiment with both ctenophores and rotifers (10.0 ± 3.8%), or in the experiment with only rotifers offered as prey (13.6 ± 8.4%).
DISCUSSION

Clearance rates of microplankton in monoculture

Adult medusae of Chrysaora quinquecirrha, which occur up to 18 cm in diameter in Chesapeake Bay, feed extensively on both copepods (Purcell 1992) and ctenophores (Miller 1974, Feigenbaum et al. 1982, Feigenbaum & Kelly 1984), but it has been suggested that clearance rates might be higher on ctenophores than on copepods (Larson 1986). In this study, when small ephyrae (3 mm diameter) of C. quinquecirrha fed on larvae of the ctenophore Mnemiopsis leidyi (1.5 mm length), the clearance was much higher than on any other prey offered (Fig. 1). The relatively large size of ctenophore larvae might increase the predator-prey encounter rate and hence increase clearance on ctenophore larvae compared with the other prey types. Further, in contrast to copepod nauplii and the ciliate Strobilidium sp. (Buskey et al. 1993), ctenophore larvae may lack an escape response. The results suggest that ctenophore larvae when abundant are a very important food source for ephyrae of C. quinquecirrha.

Clearance on copepod nauplii (0.55 ml h⁻¹ ind⁻¹) was 7% of clearance on ctenophore larvae (30 ml h⁻¹ ind⁻¹). Copepod nauplii have vigorous escape responses (Buskey et al. 1993), which may explain in part the low clearance. Density of nauplii was 200 ind. l⁻¹. Although this is a high value, such numbers are occasionally found in nature (Roman et al. 1993). The low clearance obtained was not due to saturation, as an average of only 0.3 nauplii was found in the gut contents of the ephyrae at the end of the experiment.

The dinoflagellate Gymnodinium sanguineum was not cleared by Chrysaora quinquecirrha (Fig. 1A). This is in accordance with the generally accepted role of jellyfish as carnivores, and it is also in agreement with the results of Haven & Morales-Almo (1973), who found very few phytoplankters in gut contents of ephyrae fed on concentrated water samples from Sara Creek, Chesapeake Bay, USA.

In contrast, the ciliate Strobilidium sp. was cleared by the ephyrae, but the clearance rates were low (4 ml h⁻¹ ind⁻¹; Fig. 1B) compared to the clearance on ctenophores. The density of Strobilidium sp. used in this study (8.8 ind. ml⁻¹) was similar to the average in situ density of ciliates (11 ind. ml⁻¹) reported by Dolan & Gallegos (1991) in the Rhode River tributary of Chesapeake Bay.

The rotifer Brachionus plicatilis was also cleared by ephyrae (1.7 mm diameter), but clearance rates were even lower (1 ml h⁻¹ ind⁻¹; Fig. 1C). However, we might have underestimated clearance on rotifers as the density of B. plicatilis in our experiment (2 ind. ml⁻¹) was...
Table 6. Chrysaora quinquecirrha. Production (P), ingestion (I) and net growth efficiency (NGE) of ephyrae feeding on: rotifer Brachionus plicatilis (Expt 1); ctenophore Mnemiopsis leidyi (Expt 2); or B. plicatilis + M. leidyi (Expt 3). Incubation period = 96 h.

<table>
<thead>
<tr>
<th>Expt no.</th>
<th>CO Rotifer/cten (mm)</th>
<th>C24 Rotifer/cten (mm)</th>
<th>D0 (mm)</th>
<th>D50 (mm)</th>
<th>P (µg C ind.⁻¹)</th>
<th>I (µg C ind.⁻¹)</th>
<th>Total I</th>
<th>NGE (%)</th>
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<tr>
<td>1</td>
<td>4500/0</td>
<td>-</td>
<td>2.9 ± 0.8</td>
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<tr>
<td></td>
<td>2250/0</td>
<td>-</td>
<td>2.1 ± 0.6</td>
<td>3.5 ± 0.9</td>
<td>9.7</td>
<td>107.9</td>
<td>117.7</td>
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<tr>
<td></td>
<td>1125/0</td>
<td>-</td>
<td>2.2 ± 0.7</td>
<td>3.6 ± 1.1</td>
<td>11.7</td>
<td>85.6</td>
<td>117.7</td>
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<tr>
<td></td>
<td>563/0</td>
<td>-</td>
<td>2.5 ± 0.6</td>
<td>3.8 ± 0.9</td>
<td>7.3</td>
<td>7.3</td>
<td>14.0</td>
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<tr>
<td></td>
<td>281/0</td>
<td>-</td>
<td>2.6 ± 0.7</td>
<td>3.2 ± 0.6</td>
<td>2.8</td>
<td>14.0</td>
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<td>3.3 ± 0.6</td>
<td>8.0 ± 1.3</td>
<td>83.1</td>
<td>-1986.1</td>
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<td>54.8</td>
<td>-903.0</td>
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<td></td>
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<td>544 ± 435 / 27.3 ± 22.5</td>
<td>2.2 ± 0.5</td>
<td>6.1 ± 2.0</td>
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<td>237.4/894.5</td>
<td>1131.8</td>
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<td>366 ± 152 / 70.15</td>
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<td>165 ± 99 / 77.50</td>
<td>7.9 ± 0.9</td>
<td>6.1 ± 1.8</td>
<td>36.2</td>
<td>57.6/138.5</td>
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<td>58 ± 33 / 70.15</td>
<td>3.0 ± 0.6</td>
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<td>31 ± 9 / 70.15</td>
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<td>4.1 ± 1.9</td>
<td>9.1</td>
<td>13/ 65.6</td>
<td>80.6</td>
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</table>

high compared with the average natural density of rotifers in Rhode River tributary (0.7 ind. ml⁻¹). For example, clearance by Aurelia aurita ephyrae decreased approximately 30% when density of B. plicatilis increased from 0.7 to 1.5 ind. ml⁻¹ (Olesen et al. 1994). Mean clearance on rotifers in the growth experiment was 22 ml h⁻¹ ind⁻¹ at 1.5 ind. ml⁻¹ (Fig. 2C), but the ephyrae used were slightly bigger (2.9 to 4.6 mm diameter). We believe that the true clearance rates for a 2 mm ephyrae of B. plicatilis is in the range 1 to 22 ml h⁻¹ ind⁻¹. In natural waters it is generally Synchaeta spp, which are dominant rather than Brachionus, and Synchaeta spp, might be captured at different rates.

However, the results suggest that both ciliates and rotifers may also play an important role as food sources for ephyrae of Chrysaora quinquecirrha.

It is well established that increased container volume tends to increase clearance (reviewed by Olesen 1995), and some of the differences between clearance rates obtained in monoculture experiments and clearance rates measured in growth experiments can also be explained from differences in container volume. In a 0.5 l container Brachionus plicatilis was cleared at 1 ml h⁻¹ ind⁻¹ by 1 to 7 mm diameter ephyrae (Fig. 1C), whereas the rate was 22 ml h⁻¹ ind⁻¹ in 3 l containers by 2.9 to 4.6 mm diameter ephyrae (Fig. 2C, Table 6). Likewise, Mnemiopsis leidyi larvae was cleared at 30 ml h⁻¹ ind⁻¹ by 2.8 mm diameter ephyrae in 1 l containers (Fig. 1E), whereas the rate was 60 ml h⁻¹ ind⁻¹ in 3 l containers by 3.6 to 7.1 mm diameter ephyrae.

**Clearance on mixed zooplankton**

The clearance of rotifers Synchaeta sp. by ephyrae of Chrysaora quinquecirrha in mixed zooplankton culture was approximately the same (1.3 ml h⁻¹ ind⁻¹) (Table 4) as clearance on monoculture of another rotifer Brachionus plicatilis (1.0 ml h⁻¹ ind⁻¹) (Fig. 1C). The decreasing number of rotifers in control bottles without ephyrae might be due to predation by the adult copepods, which were also present in the mixed zooplankton, as it is well established that copepods feed on rotifers (reviewed in Stoecker & Capuzzo 1990, Gifford 1991).

The clearance of copepod nauplii by ephyrae in mixed zooplankton (0.3 ml h⁻¹ ind⁻¹) (Table 4) also was similar to the rate obtained in monocultures (0.5 ml h⁻¹ ind⁻¹) (Fig. 1D). The decreased number of nauplii in control bottles without ephyrae were not statistically significant (p > 0.05, t-test), but some consumption by adult copepods may have occurred.

Acanthate ciliates were also eaten by ephyrae in mixed zooplankton as they were identified in the gut contents of ephyrae at the end of incubation. However,
even though ciliates were eaten by ephyrae, the numbers were higher in bottles with ephyrae than in those without ephyrae by the end of incubation (Table 4). Both copepods (Stoecker & Egloff 1987, Stoecker & Capuzzo 1990, Gifford 1991) and rotifers (Pourriot 1977, Stoecker & Capuzzo 1990, Gifford 1991) are known to feed on aloricate ciliates and it is possible that total predation pressure on ciliates were lower in the bottles with ephyrae compared with the control bottles as the density of rotifers and copepods was reduced due to predation by the ephyrae. Moreover, food conditions for the ciliates may have been improved in the experimental bottles as predation by ephyrae on rotifers and copepods may have led to reduced grazing on phytoplankton and bacteria.

Tintinnids were also eaten by ephyrae of *Chrysaora quinquecirrha* although clearance was low (0.3 ml h⁻¹ ind⁻¹). The low clearance on tintinnids (Table 4), which are loricate ciliates, and the relatively high clearance on the aloricate ciliate Strobilidium sp. (Fig. 1D) are in agreement with the findings of Stoecker et al. (1987) for ephyrae of *Aurelia aurita*. Whereas microzooplankton may be important as a food source for ephyrae, *C. quinquecirrha* clearance rates obtained in this study were in general low, and it seems unlikely that ephyrae has any significant impact on microzooplankton populations in *situ*.

**Gut contents of ephyrae sampled in Chesapeake Bay**

The gut contents given in Table 5 should be considered as minimum estimates, as some material might have been lost during net collection of the ephyrae. However, the relatively high numbers of ciliates and rotifers found in the guts are in agreement with the results of Haven & Morales-Almo (1973). They found that protozoa were the most frequently found components in the gut of ephyrae collected from May to June in Sarah Creek, Chesapeake Bay. Further, when ephyrae were fed in the laboratory on concentrated water samples, it was demonstrated that rotifers were ingested with higher efficiency than for any other microplankton component. Thus, the gut content analysis present in this study and others support the idea of ciliates and rotifers as significant food sources for ephyrae of *Chrysaora quinquecirrha*.

**Growth of ephyrae**

Several species of scyphomedusae and hydromedusae are known to eat other species of gelatinous competitors and intraguild predation may confer several advantages (reviewed by Purcell 1981). In this study, we found that growth rates of *Chrysaora quinquecirrha* ephyrae in the laboratory were greatest with Mnemiopsis leidyi larvae as prey, and it is likely that *M. leidyi* is critically important for the growth of *C. quinquecirrha* also *in situ*. Following the onset of a ctenophore bloom, the mean diameter of *C. quinquecirrha* in the Tred Avon River increased dramatically (Fig. 3), and the same maximal growth rate (0.6 d⁻¹) was found *in situ* as in the laboratory experiments with ctenophore larvae as prey. This growth rate is, to our knowledge, the highest ever reported for any scyphomedusan species. Earlier laboratory growth studies on ephyrae showed lower growth rates than rates observed *in situ*. Ephyrae fed newly hatched *Artemia salina* larvae and other small organisms lived less than 7 d, and small medusae (8.8 mm) grew at only 0.11 d⁻¹ when fed polychaetous annelids (Cargo & Schultz 1966). Ephyrae fed strained ctenophores lived for 49 d, but the growth rate was only 0.28 d⁻¹ compared to a rate of 0.39 d⁻¹ in St. John Creek, Chesapeake Bay, USA (Cargo & Schultz 1967). Larson (1986) obtained a biomass doubling in 6 d for one *C. quinquecirrha* medusa (65 mm diameter) (fed pieces of *Cassiopea frondosa* medusae), corresponding to an instantaneous specific growth rate of 0.1 d⁻¹. In the present study, ephyrae (3.3 mm diameter) increased their biomass (dry weight) by a factor of 13 in only 4 d corresponding to a growth rate of 0.6 d⁻¹, when live ctenophore larvae were offered in excess. In contrast, when ephyrae (2.9 mm) were fed the rotifer *Brachionus plicatilis*, the maximal growth rate was only 0.32 d⁻¹ (Fig. 2A). That is similar to the value of 0.22 d⁻¹ obtained by Olesen et al. (1994) for *Aurelia aurita* ephyrae (4 mm diameter) with *B. plicatilis* as prey. In *situ* populations of scyphomedusae usually show growth rates in the range of 0.1 to 0.4 d⁻¹ (*C. quinquecirrha*: Cargo & Schultz 1967, *A. aurita*: Moller 1980, Hernroth & Groendahl 1983, Van der Veer & Oorthuysen 1985, *Cyanea* sp.: Brewer 1989).
The digestion time for Mnemiopsis leidyi larvae was less than half the digestion time for Brachionus plicatilis (Table 6), and this may contribute to the relative high growth rate of ephyrae feeding on M. leidyi. However, it is surprising that NGE for ephyrae feeding on ctenophores only averaged 5%. Larson (1986) found an NGE of approximately 11% for a 65 mm medusae of Chrysaora quinquecirrha feeding on another medusa, Cassiopea frondosa (assuming an assimilation efficiency of 90%; Purcell 1983). However, we believe that NGE in both studies may have been underestimated due to superfluous feeding. Reeve et al. (1989) found that the ctenophore M. mccradyi continued to feed at high food concentrations, displacing partially digested material. Therefore, NGE calculated as production/ingestion decreased at high food concentration due to reduced assimilation efficiency. In the present study NGE was calculated from ingestion by assuming a constant assimilation efficiency of 90% regardless of food concentration and thus, the low NGE at high food concentration may be due to an overestimation of the assimilation efficiency. Ctenophores had diameters nearly as large as the ephyrae, and could not be completely ingested upon capture. Therefore, part of digestion occurred before complete ingestion, and some ctenophore tissue could have been lost.

Clearance by Chrysaora quinquecirrha ephyrae appeared constant with increasing concentrations of rotifers in the present study, and as consequence a positive and linear relationship between food concentration and ingestion was observed (Fig. 2B, C). There was no sign of saturation of ephyrae even at the highest prey concentration (clearance remained at the same level). A similar linear relationship between food concentration and feeding rate for C. quinquecirrha medusae has previously been reported in field studies with copepods and ichthyoplankton as prey (Purcell 1992, 1994b). These results demonstrate a high predation potential of C. quinquecirrha, even at high zooplankton densities. Linear relationships between prey density and feeding rate are generally observed for gelatinous predators, for example the hydromedusan Nemopsis bachei (Purcell & Nemazie 1992), the scyphomedusan Aurelia aurita (Olesen et al. 1994) and the ctenophore Mnemiopsis mccradyi (Reeve et al. 1989). Feeding by ephyrae of C. quinquecirrha was saturated at 20 ctenophore larvae 1⁻¹ but not at 10 ind. 1⁻¹ (Figs. 1E & 2C). However, maximum abundance of M. leidyi larvae in natural populations is usually below 1 ind. 1⁻¹ (Olney & Houde 1993). Therefore, saturation of ephyrae by M. leidyi larvae in situ is unlikely.

Larvae of the ctenophore Mnemiopsis leidyi may be very important prey for ephyrae of Chrysaora quinquecirrha, as clearance on this species was several times higher than on any other zooplankton organisms tested in the laboratory. Larvae of M. leidyi were easily caught by newly budded ephyrae, even if the larvae were about the same size as the ephyrae, and the digestion time of ctenophore larvae was less than half the digestion time of rotifers, on which ephyrae also feed. Moreover, the maximum growth rate for ephyrae of C. quinquecirrha feeding on ctenophores was the highest rate reported for any scyphomedusae, and exceeded the rate obtained when rotifers were offered as prey by more than a factor of 2. Clearance rates of adult medusae feeding on ctenophores >2 cm long are also high (Miller 1974, Purcell & Cowan 1995) and Purcell (1992) calculates that ctenophores could contribute more nitrogen to the diet than crustacean zooplankton. Predation by C. quinquecirrha medusae can reduce ctenophore population to zero in tributaries of Chesapeake Bay (Purcell et al. 1991). Therefore, we suggest that the high growth rates of C. quinquecirrha in natural populations may be possible due to the abundance of M. leidyi in Chesapeake Bay.

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