INTRODUCTION

Pelagic copepods pass through 6 naupliar and 5 copepodite stages prior to reaching the adult stage. Nauplii generally have an oval-shaped body, and they use the second antennae and mandibles for feeding and swimming (Björnberg 1986, Paffenhöfer & Lewis 1989, Mauchline 1998). Copepodites resemble adults with complete sets of mouth parts, but exhibit progressive development of the appendages and segmentation with increase of body size. Juvenile copepods often exceed adults in terms of numbers and biomass, and play an important role in the food web both as grazers and as prey. This role changes during development, and the size and type of prey captured as well as vulnerability to predation may also change (e.g. Buskey 1994, Swalding & Marcus 1994). Our understanding of the ecology of juvenile copepods is still limited compared to what we know of adults.

Knowledge of the swimming behaviour of copepods is essential to explain prey selectivity and susceptibility to predation as well as copepod migration and distribution patterns (e.g. Jonsson & Tiselius 1990, Osgood & Frost 1994, Kiørboe et al. 1996, Tiselius et al. 1997, Titelman & Fiksen 2004). Much attention has recently been paid to the behaviour of nauplii and their importance as prey organisms (e.g. Titelman 2001, Titelman & Kiørboe 2003a,b, Jiang & Paffenhöfer 2004). In contrast, development of behaviour throughout the copepodite stages (CI copepodite to adults) has been less well examined, probably because they visually resemble adults, varying only in size (Allan et al.

ABSTRACT: Swimming behaviour of copepodite stage CI to adults of the planktonic copepod Acartia clausi were observed at different food levels (0, 60, 500 and 2000 cells ml⁻¹) of the diatom Thalasiosira weissflogii. Concurrent measures of clearance in bottle incubations were performed for CI and adult females. All individuals exhibited typical jump–sink behaviour. At low food concentrations early copepodites filtered 36 % of the time, and later stages 18 %, yet the increased suspension-feeding effort did not enhance their clearance rate. Older stages and adults did not show feeding-bouts at low food concentrations, which is interpreted as a switch to the ambush-feeding mode, whereby the copepods are assumed to be searching for potentially abundant larger and motile prey, for instance, ciliates. When offered no food, frequent and long jumps were observed in all stages except for CI. Search volumes during ambush feeding were estimated, and the slow sinking of early stages resulted in a small search volume as compared to that in later stages. Early copepodite stages depended to a higher degree on suspension feeding, regardless of food concentration, due to their undeveloped food searching ability. High food availability is therefore crucial for the growth and survival of early copepodite stages of A. clausi. The heavier body of later stages enhanced foraging efficiency both during suspension feeding (by gravity tethering) and in ambush feeding (by faster sinking).

KEY WORDS: Swimming behaviour · Suspension feeding · Ambush feeding · Switching behaviour · Lower feeding threshold · Life history · Calanoid

Ontogenetic change of foraging behaviour during copepodite development of Acartia clausi

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Previous observations have shown that the swimming pattern changes little throughout copepodite development (e.g. Van Duren & Videler 1995, Paffenholzer et al. 1996) or that, in some species, early copepodite stages have different swimming behaviours (Paffenholzer & Lewis 1989, Paffenholzer et al. 1996).

The swimming behaviour of copepods is constrained by the physical properties of water. Strickler (1982) found that copepods during suspension feeding are ‘tethered’ by gravity, allowing them to create a feeding current. In small zooplankton (<0.6 mm), however, hydrodynamic theory predicts that drag replaces gravity as a tether force (Emlet & Strathmann 1985). Tiselius & Jonsson (1990) suggested that the energetic cost to remain suspended should be small for copepods <0.6 mm and that different sizes would affect efficiency in the ambush-feeding mode, since it involves sinking during prey search (e.g. Jonsson & Tiselius 1990, Kiørboe et al. 1996, Svensen & Kiørboe 2000). The size at which marked changes occur in these models are within the range of copepodes of most coastal calanoid copepods. The models also predict an ontogenetic modification of the optimal foraging strategy, but this prediction has not been investigated. Feeding mode, starvation tolerance and susceptibility to predation are all key factors in the foraging strategy.

In this study we investigated the ontogenetic change of foraging behaviour in copepoides and adults of Acartia clausi. This species and its congeners are ubiquitous in coastal temperate water, and the adult female is known to show a switching behaviour between suspension- and ambush-feeding modes depending on food availability (Kiørboe et al. 1996, Gismervik & Andersen 1997). Our study consists of behavioural observations and feeding experiments and demonstrates that the relative time allocation to each feeding mode varies during copepodite development. Possible causes for and implications of ontogenetic shifts in foraging strategies are discussed.

**MATERIALS AND METHODS**

**Collection and maintenance.** Experiments were conducted from July to October in 2003 and 2004. Acartia clausi were collected from the Gullmar fjord close to the Kristineberg Marine Research Station, on the morning of the day of the experiments using a plankton net (90 or 200 µm mesh). In the laboratory, copepods were sorted and placed in filtered seawater for 1 to 2 h and then acclimated to the experimental conditions for at least 1 h before the filming or the feeding experiments. All observations and experiments were conducted in a dark room at 18°C.

**Observation of behaviour.** The swimming behaviour was investigated by filming individual copepods in small Plexiglas aquaria (50 × 50 × 200 mm high). Four different treatments were used: filtered seawater and suspensions of 60, 500 and 2000 cells ml⁻¹ of the diatom Thalassiosira weissflogii (12.6 µm equivalent spherical diameter, ESD). The T. weissflogii strain was provided by the Marine Algal Culture Centre at Göteborg University (GUMACC) and grown at 18°C in batch cultures using B1 medium + vitamins under a 12 h light: 12 h dark illumination regime provided by cool white fluorescent lamps. T. weissflogii was selected since it is captured with 100% efficiency by all copepodite stages of Acartia clausi (Nival & Nival 1976, Dexter 1986). Only 1 copepod in each aquaria was observed, and each individual received a fresh suspension of algae. Filming was performed with a CCD-camera (OSCAR-458) fitted with a macro-lens (Vivitar 100 mm). The camera was connected via a time–date generator (Panasonic, WJ-810) to a VCR (Panasonic, NV-FS200) and a monitor. All observations were made in a dark room, and illumination consisted of an array of 880 nm infrared (IR) diodes. After observation, copepods were fixed with formaldehyde for subsequent determination of stage and body (prosome) length with a stereomicroscope and micrometer. The filming was conducted between 12:00 and 16:00 h to avoid effects of endogenous rhythms.

Swimming behaviour was recorded from the video tapes by frame-by-frame analysis with a transparency sheet mounted on the monitor. Behaviours were divided into sequences of: (1) sinking (no appendage movements), (2) feeding-bouts (slow swimming with movements of feeding appendages) and (3) jumps (swift displacements lasting 0.04 to 0.12 s). Two types of jumps were distinguished: long jumps (>0.5 body length) and feeding-bout-associated short jumps (<0.5 body length). The short jumps occurred at the end of a feeding-bout and resulted in the reorientation of the body from a vertical to a horizontal position (Saiz & Alcaraz 1992, Tiselius 1992, Saiz 1994, Tiselius et al. 1997). Frequencies of jumps and feeding-bouts were estimated for the sequences analysed. Distance and velocities of sinking and jumping events were calculated by tracking successive copepod positions (the tip of the head; 0.02 s resolution). Finally, time budgets were calculated for the 3 behaviours of sinking, swimming and jumping. Only sequences in which the copepod trajectory could be determined with good focus were used for the analysis. Periods when the copepods were close to the surface, bottom, or walls of the aquarium were excluded from the analysis. For each copepodite stage and each food treatment, 3 to 7 individuals were observed for at least 30 s ind⁻¹, resulting in 118 copepods analysed for 4945 s in total.
Feeding experiments. Feeding experiments were carried out in order to compare the time budgets to actual food intake. Clearance was estimated by incubating copepodite CI and adult females in glass bottles with a suspension of algae. A total of 10 adult females or 40 CI copepodites were placed into each of 3 to 4 screw-cap glass bottles (400 ml for adult females, 100 ml for CI) containing a suspension of Thalassiosira weissflogii at 60, 500, or 2000 cells ml⁻¹. Three bottles without copepods served as controls. The bottles were incubated standing in a dark room for 4 h (12:00 to 16:00 h) to make the conditions similar to the filming conditions. The number of cells was determined with an Elzone 5380 electronic particle counter with threshold settings to avoid electronic noise (active window: 9.948 to 18.47 µm ESD, peak at 12.58). Microscopic inspection of cell cultures, as well as samples drawn from the end of experiments, showed no particles other than algal cells. The clearance and ingestion rates were calculated using the equations of Frost (1972). Since T. weissflogii is retained with 100% efficiency, clearance equals the volume of water filtered by Acartia clausi.

Estimation of search volume in ambush feeding. When Acartia spp. sink, they display the ambush-feeding mode, i.e. searching for larger and/or moving prey (e.g. ciliates, Jonsson & Tiselius 1990). Copepods detect prey by means of mechanoreceptors on the first antennae (Paffenhöfer & Stearns 1988, Jonsson & Tiselius 1990, Kiørboe et al. 1996). The potential searched volume (F, ml ind.⁻¹ h⁻¹) during ambush feeding can be calculated by the model of Jonsson & Tiselius (1990): F = A × U, where A is the cross-sectional area (cm²) of the capture volume in which the copepod detects and attacks the prey and U is the copepod sinking velocity (cm h⁻¹). In adult female A. tonsa the parameter A is mainly dependent on prey size and includes various detection abilities of the copepod and various escape capabilities of the prey (Jonsson & Tiselius 1990). We defined A as a rectangle with length equals twice the length of the first antenna and width equals the distal, longest, forward-projected seta on the first antenna (A = 2 × antenna length × maximal setal length). The points where A. tonsa have been found to successfully capture prey are concentrated anterior to the first antennae (Jonsson & Tiselius 1990), which is why we chose only the forward-projecting setae. This estimation is conservative because it excludes the area sensed by remote detection (Jonsson & Tiselius 1990, Broglio et al. 2001). The lengths of the first antenna, the distal seta and the prosome of 5 to 7 individuals at each copepodite stage were measured under a stereomicroscope. Sinking velocity was determined from the video recordings.

Statistical analysis. Two-way factorial ANOVAs followed by Tukey-Kramer post hoc comparisons were used for testing the effect of copepodite stage and food condition (food concentration and presence/absence of food) on the behavioural components. Mean distance or duration time displayed by single individuals in a given treatment were used as the dependent variable. Proportions of time spent feeding at various food concentrations were arcsine transformed (Sokal & Rohlf 1995) prior to analyses. Analysis of covariance (ANCOVA) was used to compare mean jumping distance for each of the food conditions (food concentration or presence/absence of food), using prosome length as the covariate. Effects of food treatments on sinking velocity were also tested with ANCOVA, using log₁₀-transformed prosome lengths as the covariate. All statistical analyses were performed using the SPSS (v.10) package.

RESULTS

Behavioural observations

All copepodite stages of Acartia clausi exhibited a jump–sink motility pattern, spending most of their time (60 to 97%) sinking, but with varying degrees of feeding activity in response to food conditions (Table 1). The suspension-feeding behaviour appeared to be the same throughout development: the thoracic legs extended in association with the movement of mouth parts as observed for adult females by Rosenberg (1980). The time spent suspension feeding is therefore equivalent to the time spent swimming. We observed that the feeding-bouts in young copepodites (CI and CII) often occurred directly after a jump, whereas in older stages they usually occurred following passive sinking events.

Filtered seawater rarely induced feeding-bouts in late stages (0 to 1%; CIV to adults), whereas early stages (CI to CIII) exhibited clear feeding-bouts, 6 to 16% of their time (Table 1). When offered food, all copepodite stages showed feeding-bouts, but the time allocated to feeding varied significantly among stages and food concentrations (Table 2). Feeding time was significantly longer in 500 and 500 cells ml⁻¹, and CI copepodites spent significantly more time feeding than adult females (Table 2) at 60 cells ml⁻¹. CV copepodites and adults tended to reduce feeding activity in 60 cells ml⁻¹.

Feeding-bout frequencies were significantly affected by food concentration, but not by stage (Table 2). The frequency was highest in 500 cells ml⁻¹ and lowest in 2000 cells ml⁻¹, and there was a tendency of lower frequencies at 60 cells ml⁻¹ compared to 500 cells ml⁻¹ for CV copepodites and adults (Fig. 1a). Mean duration of feeding-bouts was significantly longer at lower food concentrations.
concentrations due to longer feeding-bouts in younger stages at 60 and 500 cells ml\(^{-1}\), even though the interaction between stages and food concentration was not significant (Fig. 1b, Table 2). Feeding-bout duration declined with body weight when all data from 60 and 500 cells ml\(^{-1}\) were pooled, but this relationship was not found in 2000 cells ml\(^{-1}\) (Fig. 2).

There were no significant differences in the frequency of long jumps between stages (Table 1). Long jumps served 2 different purposes depending on the stage. In late stages (CIV to adults) the jumps helped the individual to maintain its vertical position by countering the larger sinking distance. In early stages (CI to CIII), which did not sink rapidly, jumps were horizontally oriented and moved the copepod forward. The jumping distance was significantly related to body (prosome) length of copepods (Fig. 3). The distance was significantly longer for copepods in filtered seawater compared to those in algal suspension, while food concentration had no significant effect on the jumping distance (Table 3). The jumping distances corresponded to 1.2–1.4 body lengths in filtered seawater and to 0.7–1.2 in food suspensions, excluding CI and adult males (Fig. 4a). The jumping distance of CI copepodes was significantly shorter (0.6 to 0.7 body lengths) and that of adult males significantly longer (1.3 to 1.9 body lengths) than those of other stages, regardless of the presence/absence of food (Fig. 4a; Tukey-Kramer test, \(p < 0.05\)).

The jump frequency was significantly affected by food treatments, but not by stage (Fig. 5a, Table 2). Although adult males showed frequent jumps in the 60 cells ml\(^{-1}\) treatment, they generally jumped less than other stages, including females (Fig. 5a; Tukey-Kramer test, \(p < 0.05\)). Except for adult males, the jump frequencies were consistently higher in filtered seawater than in the treatments of 60 and 500 cells ml\(^{-1}\) (Tukey-Kramer test, \(p < 0.05\)). Short jumps were limited to copepodite stages older than CIV, and were significantly more common in adult females (Fig. 5b; Tukey-Kramer test, \(p < 0.05\)).

**Table 1.** *Acartia clausi*. Time budgets of the motility pattern of copepodes and adults in filtered seawater and suspension of the diatom *Thalassiosira weissflogii* at 60, 500 and 2000 cells ml\(^{-1}\). \(n\): number of individuals observed.

<table>
<thead>
<tr>
<th>Food treatment</th>
<th>Stage or sex of <em>A. clausi</em></th>
<th>Mean prosome length (µm)</th>
<th>(n)</th>
<th>Mean percentage of each behavioural event (±SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Male</td>
<td>782</td>
<td>6</td>
<td>97.1 (±0.6) 2.7 (±0.6) 0.1 (±0.3)</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>826</td>
<td>5</td>
<td>95.9 (±1.1) 4.1 (±1.1) 0.0</td>
</tr>
<tr>
<td></td>
<td>CV</td>
<td>697</td>
<td>5</td>
<td>96.5 (±0.9) 3.5 (±0.9) 0.0</td>
</tr>
<tr>
<td></td>
<td>CIV</td>
<td>577</td>
<td>4</td>
<td>96.0 (±1.4) 3.2 (±1.1) 0.8 (±0.6)</td>
</tr>
<tr>
<td></td>
<td>CIII</td>
<td>477</td>
<td>5</td>
<td>89.2 (±1.1) 3.4 (±0.9) 7.4 (±1.2)</td>
</tr>
<tr>
<td></td>
<td>CII</td>
<td>423</td>
<td>4</td>
<td>90.8 (±4.9) 2.8 (±0.8) 6.4 (±4.1)</td>
</tr>
<tr>
<td></td>
<td>CI</td>
<td>310</td>
<td>4</td>
<td>80.7 (±11.3) 3.3 (±2.0) 16.0 (±11.1)</td>
</tr>
<tr>
<td>60 cells <em>T. weissflogii</em> ml(^{-1})</td>
<td>Male</td>
<td>776</td>
<td>4</td>
<td>78.5 (±8.2) 3.7 (±0.9) 17.8 (±9.0)</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>788</td>
<td>5</td>
<td>78.6 (±12.2) 3.4 (±0.8) 18.0 (±12.5)</td>
</tr>
<tr>
<td></td>
<td>CV</td>
<td>744</td>
<td>4</td>
<td>79.5 (±7.1) 2.9 (±0.6) 17.6 (±6.6)</td>
</tr>
<tr>
<td></td>
<td>CIV</td>
<td>646</td>
<td>5</td>
<td>69.2 (±8.1) 2.8 (±0.5) 28.0 (±8.2)</td>
</tr>
<tr>
<td></td>
<td>CIII</td>
<td>526</td>
<td>4</td>
<td>74.7 (±8.4) 2.9 (±0.4) 22.5 (±8.4)</td>
</tr>
<tr>
<td></td>
<td>CII</td>
<td>423</td>
<td>3</td>
<td>74.2 (±6.8) 2.4 (±1.5) 23.4 (±7.1)</td>
</tr>
<tr>
<td></td>
<td>CI</td>
<td>359</td>
<td>4</td>
<td>61.8 (±11.3) 2.5 (±0.8) 35.7 (±10.6)</td>
</tr>
<tr>
<td>500 cells <em>T. weissflogii</em> ml(^{-1})</td>
<td>Male</td>
<td>846</td>
<td>3</td>
<td>65.4 (±5.0) 2.6 (±0.6) 32.0 (±4.5)</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>851</td>
<td>5</td>
<td>74.0 (±5.8) 4.6 (±1.7) 21.4 (±6.5)</td>
</tr>
<tr>
<td></td>
<td>CV</td>
<td>790</td>
<td>5</td>
<td>69.7 (±3.9) 2.7 (±1.0) 27.6 (±3.7)</td>
</tr>
<tr>
<td></td>
<td>CIV</td>
<td>673</td>
<td>4</td>
<td>69.3 (±11.6) 2.6 (±0.4) 28.1 (±11.5)</td>
</tr>
<tr>
<td></td>
<td>CIII</td>
<td>574</td>
<td>5</td>
<td>75.4 (±5.4) 2.8 (±0.6) 21.8 (±5.5)</td>
</tr>
<tr>
<td></td>
<td>CII</td>
<td>496</td>
<td>3</td>
<td>69.5 (±9.6) 2.3 (±0.7) 28.2 (±9.3)</td>
</tr>
<tr>
<td></td>
<td>CI</td>
<td>374</td>
<td>5</td>
<td>60.1 (±8.6) 2.4 (±0.8) 37.5 (±9.1)</td>
</tr>
<tr>
<td>2000 cells <em>T. weissflogii</em> ml(^{-1})</td>
<td>Male</td>
<td>776</td>
<td>4</td>
<td>87.4 (±7.0) 2.4 (±0.9) 10.3 (±6.4)</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>782</td>
<td>5</td>
<td>89.3 (±4.1) 3.8 (±1.1) 6.9 (±3.9)</td>
</tr>
<tr>
<td></td>
<td>CV</td>
<td>744</td>
<td>4</td>
<td>84.7 (±2.7) 3.9 (±1.6) 11.4 (±3.6)</td>
</tr>
<tr>
<td></td>
<td>CIV</td>
<td>610</td>
<td>5</td>
<td>87.5 (±4.6) 3.7 (±0.7) 8.7 (±4.7)</td>
</tr>
<tr>
<td></td>
<td>CIII</td>
<td>528</td>
<td>5</td>
<td>85.5 (±2.7) 2.5 (±0.7) 12.0 (±2.5)</td>
</tr>
<tr>
<td></td>
<td>CII</td>
<td>427</td>
<td>3</td>
<td>82.5 (±5.6) 2.6 (±0.5) 14.9 (±6.0)</td>
</tr>
<tr>
<td></td>
<td>CI</td>
<td>359</td>
<td>3</td>
<td>86.3 (±8.9) 2.4 (±1.1) 11.3 (±8.1)</td>
</tr>
</tbody>
</table>
Sinking velocity increased with body length (Fig. 6), but no effect of food was evident (Table 3). Relative sinking distances (normalised to prosome lengths), however, varied significantly between food treatments and between stages (Table 2). Adult males sank significantly further than other stages (Fig. 4b; Tukey-Kramer test, p < 0.05), and filtered seawater induced the furthest sinking.

**Feeding experiments**

Clearance was highest in 500 cells ml⁻¹ and decreased both in 60 and 2000 cells ml⁻¹ (Table 4). Weight-specific ingestion approached 100% d⁻¹ in 500 to 2000 cells ml⁻¹ for both stages (70 to 120% d⁻¹). In both stages, food-gathering efficiencies varied according to food concentration. Although CI copepodes spent a longer time feeding than adult females at all food concentrations, the food-gathering efficiencies of CI copepodes were at the same or below the level of adult females (Table 4).

**Estimation of search volume in ambush feeding**

The lengths of antennae and distal setae were both linearly related to body length, and slightly shorter in males (Fig. 7). Multiplying the cross-sectional area estimated from Fig. 7 with sinking velocities (Fig. 6) for

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**Table 2. *Acartia clausi*. Results of 2-way ANOVAs for effects of copepodite stage (Stage), food concentration (Diatom), and food treatment (Food: no food and food) on various behavioural parameters. *Significant effects, p < 0.05**

<table>
<thead>
<tr>
<th>Variable</th>
<th>Effect</th>
<th>df</th>
<th>F</th>
<th>p</th>
<th>Post hoc test (Tukey-Kramer test, p &lt; 0.05)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proportion spent in feeding-bouts*</td>
<td>Stage 6</td>
<td>2.97</td>
<td>0.0126*</td>
<td>0.0126*</td>
<td>CI &gt; adult female</td>
</tr>
<tr>
<td></td>
<td>Diatom 2</td>
<td>41.09</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
<td>60, 500 cells ml⁻¹ &gt; 2000 cells ml⁻¹</td>
</tr>
<tr>
<td></td>
<td>Stage × Diatom</td>
<td>1.10</td>
<td>0.38</td>
<td>0.38</td>
<td></td>
</tr>
<tr>
<td>Frequency of feeding-bouts</td>
<td>Stage 6</td>
<td>0.33</td>
<td>0.92</td>
<td>0.92</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Diatom 2</td>
<td>26.34</td>
<td>&lt;0.00001*</td>
<td>&lt;0.0001*</td>
<td>500 cells ml⁻¹ &gt; 60 cells ml⁻¹ &gt; 2000 cells ml⁻¹</td>
</tr>
<tr>
<td></td>
<td>Stage × Diatom</td>
<td>0.73</td>
<td>0.72</td>
<td>0.72</td>
<td></td>
</tr>
<tr>
<td>Duration of feeding-bouts</td>
<td>Stage 6</td>
<td>8.38</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
<td>CI &gt; CIII–CV, adult female and male, CI &gt; adult female</td>
</tr>
<tr>
<td></td>
<td>Diatom 2</td>
<td>9.39</td>
<td>0.003*</td>
<td>0.003*</td>
<td>60, 500 cells ml⁻¹ &gt; 2000 cells ml⁻¹</td>
</tr>
<tr>
<td></td>
<td>Stage × Diatom</td>
<td>0.08</td>
<td>0.08</td>
<td>0.08</td>
<td></td>
</tr>
<tr>
<td>Frequency of long jump</td>
<td>Stageb 5</td>
<td>0.34</td>
<td>0.89</td>
<td>0.89</td>
<td>Filtered seawater &gt; 60, 500 cells ml⁻¹</td>
</tr>
<tr>
<td></td>
<td>Food 3</td>
<td>5.74</td>
<td>0.0013*</td>
<td>0.0013*</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Stage × Food</td>
<td>0.79</td>
<td>0.68</td>
<td>0.68</td>
<td></td>
</tr>
<tr>
<td>Relative sinking distance</td>
<td>Stage 6</td>
<td>3.12</td>
<td>0.002*</td>
<td>0.002*</td>
<td>Male &gt; all other stages</td>
</tr>
<tr>
<td></td>
<td>Food 3</td>
<td>22.6</td>
<td>&lt;0.0001*</td>
<td>&lt;0.0001*</td>
<td>Filtered seawater &gt; 2000 cells ml⁻¹ &gt; 60, 500 cells ml⁻¹</td>
</tr>
<tr>
<td></td>
<td>Stage × Food</td>
<td>1.40</td>
<td>0.15</td>
<td>0.15</td>
<td></td>
</tr>
</tbody>
</table>

*Values were arcsine-transformed  
*Adult males were excluded from analysis (see 'Results')
each stage gives the potential search volume in ambush-feeding mode for the different stages (Fig. 8, Line a). The model shows that the potential search volume increases linearly with body weight, using the length–weight regression of Durbin & Durbin (1978). The weight-specific search volume, however, ranges from 1.35 to 2.05, with the highest values for adult females (Fig. 8, Line b). The weight-specific search volume in CI copepodites was <70% of that in adult females.

![Graph showing relationships between prosome length and feeding-bout duration](image1)

**Fig. 2. Acartia clausi.** Relationships between prosome length (PL) and feeding-bout duration (FBD) in 2000 cells ml−1 or 60 and 500 cells ml−1 (pooled) of *Thalassiosira weissflogii*. The regression for the 60 and 500 cells ml−1 is FBD = 0.403 − 0.00026 PL (n = 58, R² = 0.42, p < 0.0001).

![Graph showing relationships between prosome length and jumping distance](image2)

**Fig. 3. Acartia clausi.** Relationships between prosome length (PL) and jumping distance (JD) of copepodites and adults in filtered seawater (FSW) and various concentrations of *Thalassiosira weissflogii*. Vertical bars are ±SD. The regression line for FSW is JD = −501.0 + 2.25 PL (n = 7, R² = 0.83, p = 0.0046). The regression line for the pooled algal suspensions is JD = −308.8 + 1.44 PL (n = 21, R² = 0.86, p < 0.001) and includes observations for food concentrations of 60, 500 and 2000 cells ml−1. No statistical difference in jumping distance was found among food concentrations (see Table 3).

<table>
<thead>
<tr>
<th>Source</th>
<th>SS</th>
<th>df</th>
<th>F</th>
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</thead>
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<tr>
<td><strong>Effects of presence/absence of diatom on jumping distance</strong></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Presence/absence of diatom</td>
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<td>1</td>
<td>16.59</td>
<td>0.0004*</td>
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<td>1</td>
<td>132.64</td>
<td>&lt;0.0001*</td>
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<td>Error</td>
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<td>25</td>
<td></td>
<td></td>
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<tr>
<td><strong>Effects of diatom concentration on jumping distance</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diatom concentrations</td>
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<td>2</td>
<td>0.51</td>
<td>0.683</td>
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<tr>
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<td>1</td>
<td>138.11</td>
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<tr>
<td>Error</td>
<td>193050.0</td>
<td>17</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Effects of food treatments on sinking rate</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Food treatments (FSW, 60, 500, 2000 cells ml−1)</td>
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<td>3</td>
<td>1.931</td>
<td>0.129</td>
</tr>
<tr>
<td>Prosome length</td>
<td>3.352</td>
<td>1</td>
<td>337.58</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>Error</td>
<td>1.172</td>
<td>113</td>
<td></td>
<td></td>
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</table>

Table 3. *Acartia clausi*. Results of ANCOVA test for effects of food conditions on behavioural parameter of copepodite stages. Prosome length used as a covariate. Interaction terms have been removed from the analysis, since no significant differences in slopes among food conditions were found. *Significant effects, p < 0.05

* Sinking velocity and prosome length were log₁₀-transformed.
DISCUSSION

All copepodite stages of *Acartia clausi* exhibited a jump–sink behaviour, which agrees with previous reports for females of *Acartia* spp. including *A. clausi* (Gauld 1966, Jonsson & Tiselius 1990, Tiselius & Jonsson 1990, Saiz & Alcaraz 1992, Tiselius 1992). The jump–sink behaviour has also been reported in nauplii of *A. tonsa* (Buskey 1994, Titelman & Kiørboe 2003a), and, although the jump frequency is higher in nauplii (*A. tonsa*; 88 to 183 min⁻¹ in filtered seawater) than that in copepodites (30 to 45 min⁻¹ in filtered seawater, Fig. 5a), it appears to be a consistent feature throughout the development of *Acartia* spp. When offered a suspension of diatoms, all copepodite stages reduced jumping distance and generally also jump frequency (Figs. 3 & 5). A decreased jumping distance in food was also reported in adult female *A. clausi* (Piontkovskii & Petipa 1976); this probably serves as an adaptation to remain in food patches and also to lower predation risk. In contrast, longer and more frequent jumps in filtered seawater is a behaviour adaptive to searching for food patches in the field.

The time allocated to suspension feeding varied depending on the food concentration and stage (Table 1). Time spent feeding was highest in 500 cells ml⁻¹ and lowest in 2000 cells ml⁻¹, regardless of copepodite stage, and the feeding activity reflected the clearance rates estimated in feeding experiments (Table 4), as also reported for adult female *Acartia tonsa* (Kiørboe et al. 1996). At low food concentrations, however, clear ontogenetic differences were observed. CI to CIV copepodites engaged longer in suspension feeding than latter stages (CV to adults). Clearance at 60 cells ml⁻¹ was lower both in CI copepodites and adult females (Table 4), indicating the existence of a lower feeding threshold throughout copepodite development. A feeding threshold has been found in adult females of *Acartia* spp. (Reeve & Walter 1977, Kiørboe et al. 1985, Paffenhofer & Stearns 1988, Wlodarczyk et
al. 1992), as well as in other calanoid copepods (Corner et al. 1972, Frost 1975, Frost et al. 1983, Price & Paffenhofer 1986, Paffenhöfer 1988). In *A. tonsa*, the lower feeding rate — below the threshold — is the result of a decrease in mouthpart movements (time spent feeding) and a lower fraction of encountered cells being captured (Paffenhöfer & Stearns 1988). Paffenhöfer & Stearns' suggestion agrees with the results for adult females in the present study (Table 4). CI copepodites, in contrast, did not decrease their time spent feeding, and the energy conserving hypothesis might not be applicable to early copepodite stages of *A. clausi*.

Feeding-bouts were slightly longer for CI copepodites at 60 and 500 cells ml\(^{-1}\), than at 2000 cells ml\(^{-1}\) (Fig. 1b). Small copepodites may need to make longer feeding-bouts to collect sufficient diatoms for a single ingestion event if food concentration is low. Adult females ingested the same number of cells at 500 and 2000 cells ml\(^{-1}\) and were able to compensate for the lower food concentration by increasing activity. Specific ingestion of CI copepodites at 500 cells ml\(^{-1}\) was only 60% of that in 2000 cells ml\(^{-1}\) (Table 4), suggesting that CI copepodites had an inefficient suspension-feeding ability. Overall, feeding-bout duration at 60 and 500 cells ml\(^{-1}\) was negatively correlated with body size (Fig. 2), and the efficiency of suspension feeding seems to improve during growth.

Efficient suspension feeding requires a tethering force to anchor the animal when it feeds. The shift from drag to gravity as the tethering force during growth allows a stronger current in heavier animals (Emlet & Strathmann 1985, Tiselius & Jonsson 1990). Frequent short jumps associated with feeding-bouts were observed only in later copepodite stages, CIV to adults (Fig. 5b), and they reoriented themselves with respect to the gravitational force to maximise the efficiency of their suspension-feeding current. CI swam more horizontally, probably using drag rather than gravity as a tether force, and were less efficient at anchoring themselves to support a strong feeding current.

The short feeding-bouts in early copepodites (CI to CIII) at 60 cells ml\(^{-1}\) probably indicate that they are just sampling the water for food (Tiselius 1992), and these bouts are also performed in filtered seawater (Table 1, Fig. 1). Late copepodite stages in contrast showed no feeding-bouts in the filtered seawater and may instead search for motile prey in the ambush-feeding mode (Jonsson & Tiselius 1990, Tiselius & Jonsson 1990, Kiørboe et al. 1996). The presence of the sample feed-

![Fig. 6. *Acartia clausi*. Relationships between prosome length (PL) and sinking velocity (SV) in filtered seawater and various concentrations of *Thalassiosira weissflogii*. Note that x- and y- axes are logarithmic. All data from the 4 food treatments (60, 500 and 2000 cells ml\(^{-1}\), and filtered seawater) were pooled for the regression. The regression is log\(_{10}\) transformed SV = \(-4.325 + 1.41 \times \log_{10} \text{PL}\) (n = 118, \(R^2 = 0.746, p < 0.0001\))

Table 4. *Acartia clausi*. Clearance, ingestion rates and specific ingestion of adult female and CI copepodites, determined by feeding experiments. Carbon contents of copepods were assumed to be 43% of body dry weight. *Thalassiosira weissflogii* was 79 pg C cell\(^{-1}\) (estimated from Strathmann 1967). Specific ingestion was calculated by multiplying the 4 h feeding experiment by 6 (= 24 h). Proportion spent feeding is based on data from behavioural observations (see ‘Results’ and Table 1). Food-gathering efficiency calculated from specific ingestion divided by fraction of time spent for feeding.
Foraging behaviour of juvenile copepods in early copepodite stages is probably due to their low efficiency of ambush feeding. Weight-specific search volume in CI copepodites was only about 70% of that in adult females (Fig. 8), and the smaller individuals apparently depended more on suspension feeding regardless of food availability.

Predatory feeding on ciliates or copepod nauplii has been shown in nauplii and early copepodite stages of _Acartia_ spp. (Landry 1978, Stoecker & Egloff 1987), but Landry (1978) reported that early copepodite stages (CI and CII) of _A. clausi_ did not prey on conspecific nauplii when alternate food (phytoplankton) was offered. In contrast, later stages switched to predatory feeding under the same conditions (Landry 1978). Constant feeding bouts for sampling in early copepodite stages _Acartia clausi_ indicate a costly foraging strategy, which may not be sustainable unless food conditions are favourable, e.g. in a phytoplankton patch. Young copepodites of _Acartia_ spp. tend to occur in the upper part of the water column (Landry 1978, Ueda 1987), and _A. clausi_ CI copepodites have been observed to maintain a close association with the chlorophyll maximum throughout the diel cycle (Harris 1988, Titelman & Fiksen 2004). In the chlorophyll maximum, early copepodite stages can suspension feed continuously and sustain their relatively high metabolic requirements.

Our results confirm that foraging behaviour in _Acartia clausi_ is different in early and late copepodite stages. Based on a food encounter model, Titelman & Kierboe (2003a) concluded that nauplii of _A. tonsa_ rely on motile food. Predation on motile prey is well established in adult female _Acartia_ spp. (e.g. Stoecker & Egloff 1987, Wiadnyana & Rassoulzadegan 1989), as is their switching behaviour, using either suspension or ambush feeding according to food availability (Jonsson & Tiselius 1990, Kierboe et al. 1996, Gismervik & Andersen 1997). Judging from the jump frequency and sinking velocity, ambush feeding can be divided into 2 phases. Frequent jumps of nauplii serve to avoid local food depletion (Titelman & Kierboe 2003a), and they search the water column by means of jumping. Copepodite stages, in contrast, scan the water column while...
they are sinking and jump much less frequently (Jons-
son & Tiselius 1990). This indicates that the ambush-
feeding efficiency of Acartia spp. is greatly improved
through the transformation from nauplius to cope-
podite. This efficiency relies on 2 factors, both of which
change during ontogeny: sinking velocity, which in-
creases due to increased body weight, and jumping
capability, using the first antenna and urosome, which
compensates for the larger sinking distance. Since CI
copepodites jump less and sink slowly, these small ani-
mals largely have to depend on suspension feeding.

Based on the balance between gravitational and
drag force on zooplankton in the water column, Emlet
& Strathmann (1985) predicted that behaviour and
the energetic cost in swimming of zooplankton should
change at a body length of about 0.6 mm. Tiselius &
Jonsson (1990) found a strong shift in the efficiency of
suspension feeding in the same size range. Our obser-
vations agree with these predictions, since many be-
avoural parameters such as feeding-bouts and jump
frequency notably change between CIII and CIV cope-
podites, corresponding to about 0.6 mm (in prosome
length). Therefore, we suggest that increased body
weight will enhance the foraging efficiency during
both suspension (by tethering force) and ambush feed-
ing (by sinking) and that body weight is an important
factor for optimal feeding in Acartia clausi.

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LITERATURE CITED

Allan JD, Richman R, Heinle DR, Huff R (1977) Grazing in
juvenile stages of some estuarine calanoid copepods. Mar
Biol 43:317–331

Björnberg TKS (1986) Aspects of the appendages in develop-
ment. Sylogeus 38:51–66

Broglio E, Johansson M, Jonsson PR (2001) Trophic interaction
between copepods and ciliates: effects of prey swimming

Buskey EJ (1994) Factors affecting feeding selectivity of visual
predators on the copepod Acartia tonsa: locomotion, visibil-

Corner EDS, Head RN, Kilvington CC (1972) On the nutrition
and metabolism of zooplankton. VIII. The grazing of Bid-
dulphia cells by Calanus helgolandicus. J Mar Biol Assoc
UK 52:847–861

Dexter BL (1986) Developmental grazing capabilities of
Pseudocalanus sp. and Acartia clausi (Cl to adult): a com-
parative study of feeding. Sylogeus 58:547–551

Durbin EG, Durbin AG (1978) Length and weight relations-
ships of Acartia clausi in Narragansett Bay, Rhode Island.
Limnol Oceanogr 23:958–969

Emlet RB, Strathmann RR (1985) Gravity, drag and feeding
currents of small zooplankton. Science 228:1016–1017

Frost BW (1972) Effects of size and concentration of food par-
ticles on the feeding behavior of the marine planktonic copepod Calanus pacificus. Limnol Oceanogr 17:805–815


Frost BW, Landry MR, Hassett RP (1983) Feeding behavior of
large calanoid copepods Neocalanus cristatus and N. plumchrus from the subarctic Pacific Ocean. Deep-Sea Res 30:1–13

Gauld DT (1966) The swimming and feeding of planktonic copepods. In: Barns H (ed) Contemporary studies in ma-


Harris RP (1988) Interactions between diel vertical migratory

Jiang H, Paffenholzer GA (2004) Relation of behavior of cope-
pod juveniles to potential predation by omnivorous cope-
pods: an empirical-modeling study. Mar Ecol Prog Ser 278:
225–239

Jonsson PR, Tiselius P (1990) Feeding behaviour, prey detec-
tion and capture efficiency of the copepod Acartia tonsa
feeding on planktonic ciliates. Mar Ecol Prog Ser 60:35–44

Kierboe T, Møhlenberg F, Hamburger K (1985) Bioenergetics
of the planktonic copepod Acartia tonsa: relation between feeding, egg production and respiration, and composition
of specific dynamic action. Mar Ecol Prog Ser 26:85–97

Kierboe T, Saiz E, Viitasalo M (1996) Prey switching behav-
ior in the planktonic copepod Acartia tonsa. Mar Ecol
Prog Ser 143:65–75

Landry MR (1978) Population dynamics and production of a
planktonic marine copepod, Acartia clausi, in a small tem-

Mar Biol 33:1–710

Nival P, Nival S (1976) Particle retention efficiencies of an
herbivorous copepod, Acartia clausi (adult and copepodite

Osgood KE, Frost BW (1994) Ontogenetic diel behaviors of the

Paffenholzer GA (1988) Feeding rates and behavior of zoo-

Paffenholzer GA, Lewis KD (1989) Feeding behavior of nauplii
of the genus Eucalanus (Copepoda, Calanoida). Mar Ecol
Prog Ser 57:129–136

Paffenholzer GA, Stearns DE (1988) Why is Acartia tonsa (Copepoda: Calanoida) restricted to nearshore environ-

Motion behavior of nauplii and early copepodite stages of

Price JH, Paffenholzer GA (1985) Quantitative description of
the behaviour of copepod Acartia clausi during feeding on

Reeve MR, Walter MA (1977) Observations on the existence of
lower threshold and upper critical food concentrations
29:211–221


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