Ontogenetic vertical distribution patterns in small copepods: field observations and model predictions

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ABSTRACT: We investigated fine-scale (5m, 4h) species- and stage-specific (N1 to C6) distributions of common copepods at an anchor station in a Swedish fjord during two 24 h periods in October 1997. Generally, both calanoid and cyclopoid nauplii were found near the surface, while copepodids stayed deeper in the water column. No diurnal migration was observed. To analyze the observed distributions, we combined mechanistic models of predation risk from fish and copepods, formulations of temperature-dependent growth and a habitat optimization model, maximizing expected lifetime reproductive output. Motility pattern has implications for encounter rates with copepod predators, and therefore affected optimal vertical positioning in the model. By applying species- and stage-specific motility, and accounting for the ambient copepod predator field, we computed depth profiles of the mortality risk for the observed field situation. Predicted diel and ontogenetic vertical distribution patterns for various levels of fish concentrations were compared with observed distributions, and much of the patterns in the field were explained by the model. While the risk of fish predation governs the deeper habitat selection of the larger copepodids, the risk of copepod predation is probably more important for nauplii and small copepods. In addition, the vertically homogeneous growth profile and dense layers of copepod predators may wipe out potential benefits of diurnal migration.

KEY WORDS: Nauplii · Oithona spp. · Habitat optimization · Predation risk · Encounter · Behavior

INTRODUCTION

The pelagic realm provides few physical refuges from predators. However, vulnerable zooplankton may reduce predation risk from visual predators by resting at depths where ambient light levels limit search efficiency (Zaret & Suffern 1976, Aksnes & Giske 1993). If subject to rheotactic predators, such as larger copepodids, zooplankters may limit predation risk by adopting a ‘silent’ motility pattern that minimizes predator detection and encounter rates (cf. Tiselius et al. 1997). Alternatively, plankters may behave according to the concept that ‘an enemy’s enemy is a friend’, and reside near the surface when large invertebrate predators face high risk from fish, and descend at night when their predators come to the surface (Ohman et al. 1983). Migration patterns vary with size; large zooplankters are very visible to fish and may therefore leave the surface during the day to reduce predation risk (De Robertis et al. 2000, De Robertis 2002), while small zooplankton are less visible but more susceptible to predation from invertebrates (e.g. Landry & Fagerness 1988).

Theoretical studies suggest that younger stages maximize fitness by selecting habitats supporting high growth rates, even when risk is high, in order to minimize lifetime exposure to predators (Fiksen & Giske 1995). Later stages with higher reproductive value should instead pay more attention to survival (Clark 1994). In addition, zooplankton habitat selection is flexible, and may change with the abundance of dominant predator (Ohman 1990, Frost & Bollens 1992, Loose & Dawidowicz 1994). If growth is maximized near the surface, small or juvenile copepods may maximize their fitness by staying there for 2 reasons; viz. (1) the large and potentially dangerous zooplankton leave the surface waters during daytime, and (2) faster growth allows prey to grow out of the predation size...
window of many larger zooplankters more quickly (Fiksen & Giske 1995, Fiksen 1997).

There has been a considerable research effort describing and quantifying the behavior of zooplankton, and small-scale predator-prey interactions in the laboratory (e.g. Tiselius & Jonsson 1990, Paffenhofer et al. 1996, Titelman & Kiorboe 2003a). Many copepods are omnivorous and feed effectively on younger developmental stages (e.g. Landry & Fagerness 1988, Sell et al. 2001). Recently, theoretical and experimental studies have demonstrated how nauplii and copepodid prey may influence detectability and encounter rates with tactile predators through their motility patterns (e.g. Tiselius et al. 1997, Kiorboe et al. 1999, Caparroy et al. 2000, Titelman 2001). Prey size also has direct and indirect influences on predation by both copepods and fish. Size affects both the magnitude of the hydrodynamic signals generated by motility and signal perception (Kiorboe & Visser 1999, Kiorboe et al. 1999, Titelman & Kiorboe 2003b). In addition, escape velocity often increases with size (Mauchline 1998, Titelman & Kiorboe 2003b). Prey may also simply outgrow the predators’ prey-size spectrum. When compared to similarly sized copepodids, nauplii are generally more susceptible to copepod predation (e.g. Landry & Fagerness 1988). The susceptibility to visual predators typically increases with prey size (Brooks & Dodson 1965).

It is not yet known how well the small-scale mechanisms involved in motility, signal perception and generation, and predator-prey interactions that we observe in the laboratory scale up to field situations and the distribution of zooplankters. Unfortunately, of the many reports of copepod vertical distributions, only few consider naupliar stages, and even fewer separate nauplii into species or stages (e.g. Lagadeuc et al. 1997, Durbin et al. 2000, Incze et al. 2001). This contrasts with the fact that nauplii make up the numerical bulk of copepods. One may hypothesize that species- or stage-specific differences in behavior (e.g. Titelman & Kiorboe 2003a,b) are also reflected in field distributions. Here we attempt a step towards exploring this by examining the distribution of nauplii and small copepodids in a Swedish fjord.

The vertical distribution of all stages (N1 to C6) of common species was recorded during 2 d of intense sampling. We modeled growth and predation rates from the physical and biological environment observed in the fjord. We implemented the motility patterns of each stage and species into mechanistic models of both copepod and fish predator-prey interactions. Then, by use of a dynamic optimization model, we calculated the sequence of habitat selections that maximize reproductive value for all size categories, in increments of 1 h. Finally, we compared the model results with the field observations.

**MATERIALS AND METHODS**

**Sampling scheme.** We sampled at an anchor station (58° 16.00’ N, 11° 28.34’ E, depth ~60 m) in Gullmarsfjorden, western Sweden, during two 24 h periods between 26 and 29 September 1997. Sampling was conducted at 4 h intervals, starting at 09:30 h on 26 and 28 September and ending after the 09:30 h sampling the following day. Bottle (5 l Hydrobios PWS) samples for zooplankton were collected at 5 m depth intervals between the surface and 55 m, simultaneously with CTD (Neil Brown MK III) and relative fluorescence measurements (Sea Tech). Additional CTD casts were carried out at 2 h intervals. Surface light was measured nearby at Kristineberg Marine Research Station, Fiskebäckskil. Profiles of chlorophyll a (chl a) were obtained at the 13:30 and 01:30 samplings. Water samples for chl a were GF/F filtered, the filters extracted in alcohol and the extract measured on a fluorometer (Turner Designs AU-10).

**Zooplankton.** Zooplankton were collected from the bottle samples by screening the water through a 40 µm mesh, whereupon the plankton were preserved in 4 % formalin. We considered 12 taxa, of which all the calanoids and oithonids were identified to stage while Coryceaus anglicus, Onceaea borealis, and Microsetella norvegica were only separated into nauplii or copepodids. Nauplii were identified to species and stage where possible under an inverted microscope (Leica DMIL equipped with a relief segment), following Oberg (1906) and Ogilvie (1953) for the calanoids, Lovegrove (1956) for M. norvegica, Haq (1965), Oberg (1906), Gibbons & Ogilvie (1933) and Lovegrove (1956) for Oithona similis and Oithona nana, Gibson & Grice (1978) for C. anglicus and Malt (1982) for Onceaea borealis. Oceaea spp. and Centropages spp. nauplii were assumed to belong to the same species as the observed adults. Other copepod nauplii (normally less than 5 %) were counted but not identified. In each sample, all or ca. 200 nauplii per sample were identified. For samples containing more than 200 nauplii, the species and stage composition were corrected to the total count.

Initially we had difficulty separating N1 and N2 of Oithona nana and O. similis. Therefore, lengths from later identified samples were used to construct size cut-off points for the separation of O. similis and O. nana stages N1 and N2 for the early samples (10 samples). The cut-off points were verified by applying the constructed size limits to the already analyzed data, and then comparing the outcome of the procedure with the numbers resulting from real identifications (Table 1). The same procedure was used to separate uncertain cases, mostly stages N1 and N2, of Microcalanus pusillus and Paracalanus parvus (data not shown).
Table 1. Oithona nana and O. similis. Verification of the separation of stages N1 and N2. Cross-table comparison of numbers resulting from assigning nauplii to groups based on body length (x, µm) with numbers of actually observed identifications based on taxonomic characters (see ‘Materials and methods’). The percentage values (in parentheses) represent the fraction of the total observed nauplii of a specific species and stage (each of columns 3 to 6) being assigned to a length group (groups defined in column 1). Percentage values sum up to 100% in the vertical direction, i.e. all observed nauplii of a certain species and stage are assigned to 1 of the 4 possible size groups. For easy comparison the total sums of observed and assigned numbers are indicated in the 2 bottom rows. Gray shading indicates that the assigned group matches the observed group.

<table>
<thead>
<tr>
<th>Assigned group</th>
<th>Body length (µm)</th>
<th>Observed counts based on taxonomic characters</th>
<th>O. nana N1</th>
<th>O. nana N2</th>
<th>O. similis N1</th>
<th>O. similis N2</th>
</tr>
</thead>
<tbody>
<tr>
<td>O. nana N1</td>
<td>65.1–83.7</td>
<td>180 (88.7)</td>
<td>27 (4.1)</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td></td>
</tr>
<tr>
<td>O. nana N2</td>
<td>83.7–102.3</td>
<td>23 (11.3)</td>
<td>621 (95.1)</td>
<td>2 (1.7)</td>
<td>0 (0)</td>
<td></td>
</tr>
<tr>
<td>O. similis N1</td>
<td>102.3–120.9</td>
<td>0 (0)</td>
<td>5 (0.8)</td>
<td>94 (78.3)</td>
<td>20 (11.4)</td>
<td></td>
</tr>
<tr>
<td>O. similis N2</td>
<td>120.9–155.0</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>24 (20)</td>
<td>156 (88.6)</td>
<td></td>
</tr>
<tr>
<td>( \Sigma_{\text{observed}} )</td>
<td></td>
<td>203</td>
<td>653</td>
<td>120</td>
<td>176</td>
<td></td>
</tr>
<tr>
<td>( \Sigma_{\text{assigned}} )</td>
<td></td>
<td>207</td>
<td>646</td>
<td>119</td>
<td>180</td>
<td></td>
</tr>
</tbody>
</table>

Model. We formulated a habitat selection model based on sub-models of predation risk from visually foraging planktivores (i.e. juvenile sprat and herring that dominate in this area) and zooplankton predators (i.e. ambush copepods), and of temperature-dependent growth rate.

Growth and reproduction: Food, in terms of phytoplankton (chl a) and nauplii, was available in high concentrations (see Fig. 1). The mean chl a concentration of ~3 µg l⁻¹ at the peak corresponds to ~150 µg C l⁻¹. Most adult copepods have a maximum or near maximum ingestion at such concentrations, both when feeding on phytoplankton (e.g. Calanus pacificus, Frost 1972; Oithona nana, Lampitt & Gamble 1982; Acartia tonsa, Saiz & Kiorboe 1995) and ciliates (e.g. A. tonsa, Saiz & Kiorboe 1995). Younger developmental stages require less food (reviewed in Mauchline 1998). Given that carbon is also available from nauplii (see Fig. 2) and other microzooplankton (not quantified), as well as from the chl a, we assumed that food is plentiful. Therefore, growth (g) is controlled here by temperature \( T(z) \) at depth (z) only (cf. Huntley & Lopez 1992). Thus:

\[
g(z) = g_{15} \times Q_{10}^{\frac{T(z) - 15}{10}}
\]

where \( g_{15} \) is the maximum specific growth rate (g g⁻¹ s⁻¹) at 15°C, and \( Q_{10} \approx 3.0 \) for calanoids (Huntley & Lopez 1992). Each species was specified by its egg and adult weight and the weights of intermediate stages were computed assuming isochronal development (e.g. O. similis, Sabatini & Kiorboe 1994). Kiorboe & Sabatini (1995) compiled \( g_{15} \), \( Q_{10} \), and weights for the relevant species. The model currency is µg C, but for several processes the body length (L) is more relevant. We applied published species-specific length-weight relationships, separate for nauplii and copepodids, to switch between length and weight (regressions compiled by Mauchline 1998), and applied a conversion factor of 0.4 µg C µg⁻¹ dry weight (Parsons et al. 1984) when necessary. We assumed that adult copepods allocate all surplus energy to reproduction such that:

\[
b(w, z) = \frac{g(z) \times w_{\text{adult}}}{w_{\text{egg}}}
\]

where \( b \) is the number of eggs produced per unit time and \( w_{\text{adult}} \) and \( w_{\text{egg}} \) are female and egg weights, respectively.

Predation risk: Predation risk is a function of predator-prey encounter rates and post-encounter escape probabilities. Volume encounter rate (\( \beta_{\text{fp}} \), cm³ predator⁻¹ s⁻¹) with a predator, be it a fish (f) or a copepod (A), depends on the predator detection distance (\( R_{f} \), cm) and the relative velocity between predator and prey (\( \Delta v \), cm s⁻¹) such that:

\[
\beta_{\text{fp}} = \pi R_{f}^{2} \Delta v
\]

Ultimately, the reactive distance \( R \) is the key to understanding the susceptibility of zooplankton to both visual and tactile predators. In interactions with fish, predator velocity (\( u_{f} \)) exceeds that of the prey (\( v_{b} \)), i.e. \( u_{f} >> v_{b} \). Thus, \( \Delta v \) simplifies to \( u_{f} \) and:

\[
\beta_{f} = q_{f} R_{f} u_{f}
\]

where \( q_{f} \) is the fraction of the cross sectional area of the perceptive field that is efficiently scanned (~50%, Rosenthal & Hempel 1970). For small planktivorous fish, e.g. larvae, prey to predator size ratios (\( \lambda \)) typically range from 2 to 8% (Munk 1995), with smaller prey being ignored or undetected, and larger prey being too evasive to be captured. We let each individual prey be subject to a fish predator of length about 1/\( \lambda \) (= 20) times the prey’s own body length (L). This implies that predators that are larvae or small fishes were considered. The search velocity of the fish is 1 body length s⁻¹, or L/\( \lambda \) s⁻¹. Predator visual ability, prey size, prey pigmentation, and light all affect \( R_{f} \) (Aksnes & Giske 1993, Aksnes & Utne 1997). If \( R_{f} \) approximates 1 fish body length under light saturated conditions (or L/\( \lambda \)) and light limitation is modeled as suggested by Aksnes & Utne (1997), then Eq. (3) becomes:
where $K_c$ is the light level ($5 \mu\text{mol m}^{-2} \text{s}^{-1}$, Aksnes & Utne 1997). $R_b$ is half of its light satiated value (i.e. $L/2\lambda$) and $E_b$ is the ambient light. Our choice of $K_c$ is representative of fishes adapted to strong light (Aksnes & Utne 1997). $E_b$ depends on depth ($z$) and irradiance at the surface $S(h)$ at any given hour of day ($h$) such that:

$$E_b(z, h) = S(h) \sum_{0}^{z} e^{-k(z)}$$

where the local light attenuation coefficient is a function of chlorophyll $a$ concentration [$\Psi(z)$] (see Fig. 1E,F) (Riley 1956):

$$k(z) = 0.14 + 0.054\Psi(z)^3 + 0.0088\Psi(z)$$

The solar irradiance at the surface was modeled as a function of $h$ from the measured light level at midday (see Rosland & Giske 1994).

We assumed that all encounters with fish are lethal. In summary, the rate of clearance by fish increases exponentially with prey body size, decreases non-linearly with depth and varies over the diel light cycle. Because both the prey predator size-ratio and capture success are constant, the predation rate from fish $\mu_z(L, h, L, h)$ is a function of total fish concentration ($C_t$), prey length, depth and time of the day:

$$\mu_z(L, h, L, h) = \beta_z(z, L, h)C_t$$

We did not sample the fish community during the cruise, but evaluated the sensitivity of our predictions to different levels of $C_t$, assuming that $C_t$ is constant over depth and prey size.

As with fish, volume encounter with other copepods $\beta_3$ is given by Eq. (1). Nauplii and copepodids typically engage in 3 types of behavior; sinking, swimming, and jumping, here denoted by subscript s, w, and j respectively (e.g. Tiselius & Jonsson 1999, Titelman & Kiorboe 2003a). In addition, behavior is highly species- and stage-specific; also for nauplii (Titelman & Kiorboe 2003a). Because $R_\Lambda$ strongly depends on prey motility behavior, we calculated $R_\Lambda$, and $\Delta v$ using behavior-specific speeds for each of the various species and stages.

For each prey species and stage we thus obtained volume encounter kernels for each of these 3 behaviors; $\beta_3s$, $\beta_3w$, and $\beta_3j$ (Titelman & Kiorboe 2003b). Many copepods are capable of switching between an ambush and a feeding current or a cruising foraging strategy (e.g. Jonsson & Tiselius 1990, Tiselius & Jonsson 1990). We here assumed that all copepod predators are ambush foragers. This assumption generates conservative volume encounter rates because it minimizes $\Delta v$.

A copepod detects a (spherical) prey of radius $a$, or $0.5L_i$ sinking passively with a speed $v_{bs}$ at a distance $R_{bs}$:

$$R_{bs} = \frac{Kv_{bs}a}{u^*}$$

where $u^*$ is the critical velocity required for the predator to respond, and $K$ is a non-dimensional parameter of ~1 (Kiorboe & Visser 1999). The distance at which a self-propelled prey with velocity $v_p$ can be detected is better approximated as:

$$R_{Awj} = \left(\frac{3\pi v_{ps}a^2[1 + 3\cos(2\theta)]}{2u^*}\right)^{0.5}$$

where $\theta$ is the detection angle (Svensen & Kiorboe 2000, Visser 2001) and $v_p$ is jumping ($v_{ps}$) or swimming ($v_{sw}$) velocity. We always set $\theta = \pi/2$. Because Eq. (2) assumes that $R_\Lambda$ is measured from the center of the detection sphere, we corrected by adding half a predator width to the estimates. For swimming and jumping prey we assumed random arrival of encounters, $\Delta v = (v_{ps}^2 + v_{sw}^2)^{0.5}$, and for sinking prey we assumed differential settling, $\Delta v = |v_{ps} - v_{sw}|$, where $u_p$ is predator sinking speed. The total predator volume encounter rate is now computed as:

$$\beta_\Lambda = \beta_s\tau_s + \beta_w\tau_w + \beta_j\tau_j$$

where $\tau_s$, $\tau_w$, and $\tau_j$ are the time fractions spent sinking, jumping and swimming, respectively (Titelman & Kiorboe 2003b).

Once encountered, prey may escape. We modeled the prey escape probability $P_e$ as an empirical (sigmoid) function of prey ($L$) to predator ($L_\Lambda$) length ratio:

$$P_e(L_\Lambda, L) = \left[1 + \exp\left(-\frac{(L/L_\Lambda) - k_b}{b}\right)\right]^{-1}$$

where $k_b$ is the prey-predator size ratio where $P_e$ equals 0.5, and $b$ is a shape-parameter. The parameters were adopted from the size-dependent capture probabilities reported in Caparroy et al. (2000, their Fig. 4), yielding $k_b = 0.2$ and $b = 0.05$.

So far, the predation risk from copepodids depends on prey motility and prey and predator size. To evaluate the spatial profile of mortality risk at our field station, we first needed to summarize the size-structured concentrations of predators $C_\Lambda(z, L_\Lambda)$ at each depth. We averaged the observed copepodid concentrations in length intervals ($L_\Lambda$) of 20 $\mu$m at each sampling depth, and used linear interpolation to get a concentration for each meter in the water column. Then, the rate of deadly encounters with copepod predators, $\mu_\Lambda(z, L)$ (h$^{-1}$), for a prey of length $L$ at depth $z$ becomes a size-dependent function of volume encounter, predator concentration and escape:

$$\mu_\Lambda(z, L) = \sum_{L_\Lambda} \beta_\Lambda(L_\Lambda, L)C_\Lambda(L_\Lambda, z)[1 - P_e(L_\Lambda, L)]$$

Species-specific motility parameters and behavioral time budgets are generally only available for adult fe-
males (e.g. Tiselius & Jonsson 1990) and some stages of nauplii (e.g. Titelman & Kiørboe 2003a) (see Fig. 5). We therefore converted speeds to body lengths (bl) s⁻¹ and applied them to 3 stage groups for each species; N1–N2, N3–N6, and C1–C6. For nauplii, we approximated sinking speeds with the relationship \( v = 1.68L^{1.38} \) (Titelman & Kiørboe 2003a) and for copepods with \( v = 0.88L^{1.36} \) \((R^2 = 0.47, n = 6, \text{fitted from data in Tiselius & Jonsson 1990, Svensen & Kiørboe 2000)}\), where units of \( v \) and \( L \) are mm s⁻¹ and mm, respectively.

**Fitness and optimal depth selection:** We applied a dynamic programming model (Mangel & Clark 1988) to compute the optimal habitats for each stage over time. We assumed that animals are free to move 1 m up or down every hour, and select habitats such that the number of offspring produced over the lifetime is maximized. This implies that an organism in any state \( w \) maximizes its expected future reproductive value \( V(w, h, H) \) from time \( h \) to the time horizon \( H \) by picking the optimal habitat \( z^* \). Our general dynamic programming (sensu Mangel & Clark 1988) equation is:

\[
V(w, h, H) = \max[b(w, z) + \exp[-\mu_z(z, L_w, h)] - \\
\mu_A(z, L_w)] \times V[w(1 + g(z)), h + 1, H]
\]

The terms \( b(w, z) \), \( g(z) \), \( \mu_z(z, L_w, h) \) and \( \mu_A(z, L_w) \) are the contributions at time \( h \) to fitness from reproduction, growth and fish and copepod predation, respectively.

The model was run over at least 30 d in time intervals of 1 h, always using the environment recorded in the field program. ‘Terminal effects’ are behaviors that maximize the fitness criterion specified at \( H \) rather than the reproductive value (Mangel & Clark 1988). We set terminal fitness \( V(w, H, H) = w \), and always checked that \( H \) was sufficiently large to avoid any influence from this assumption.

**RESULTS**

**Field observations**

Environmental conditions

Hydrographical conditions and zooplankton distributions were largely invariant during the cruise (Fig. 1A–F). We therefore averaged the profiles across sampling times, i.e. both days, to get environmental forcing for the model, including the size-structured copepodid predator field (Figs. 1J & 2). We considered 2 temperature scenarios; viz. one obtained from averaging the first 10 profiles, i.e. until 17:30 h on Day 2, and one obtained from averaging the remaining 4 profiles, i.e. from 21:30 h Day 2 (Fig. 1A,B,G).

**Zooplankton distribution patterns**

The water column was strongly stratified in terms of zooplankton abundance, with maximum at the less saline surface, minimum at 25 to 30 m and increasing concentrations towards the bottom (Figs. 1C–F & 3). Concentrations reached ~220 and 70 ind. l⁻¹ for nauplii and copepodids respectively. The small cyclopoids *Oithona nana* and *Oithona similis* dominated numerically, *Oithona nana* being most abundant with concentrations up to 40 l⁻¹ for individual nauplius stages and 10 adult females l⁻¹ (Figs. 3 & 4). Most calanoid nauplii were located close to the surface, while *Microcalanus pusillus*, *Oncaea borealis* and *Microsetella norvegica* were found towards the bottom (Figs. 3 & 5). Nauplii of *Oithona similis* and *Pseudocalanus elongatus* had an intermediate distribution (Figs. 3 & 5). Copepodid distribution varied more. Generally, most calanoids concentrated around the deeper pycnocline, the oithonids higher in the water column, and *Microcalanus pusillus*, *Oncaea borealis*, and *Microsetella norvegica* in deeper waters (Figs. 3 & 5).

Two general features characterized the distribution patterns. First, most species exhibited ontogenetic habitat shifts, with nauplii near the surface and copepodids deeper in the water column (Fig. 5). The calanoid N1 were generally more widely distributed over depth than N2 to 6 (Fig. 5). Curiously, the later copepodid stages of *Oithona nana* were found in 2 distinct layers, one near the surface and one at depth, while most of the females were found in the surface layer (Fig. 4). Although *O. similis* concentrations peaked near the surface, the distribution was not as confined to this layer as that of *O. nana* (Figs. 3 & 5).

Secondly, we could not recognize any patterns of diel vertical migration (DVM) in any stage or species, even when pooling day and night samples separately (cf. *Oithona nana*, Fig. 4, others not shown). However, because of the high resolution of the data, small sampling volumes, and thus low specific counts, a thorough analysis was only possible for *O. nana* (Fig. 4). We applied the model to examine factors involved in governing the (lack of) temporal dynamics in the observed distributions, focusing mainly on the nauplii and on the dominating small *O. nana*.

**Model predictions**

We computed optimal ontogenetic distribution patterns for different predator situations; (1) copepods only, (2) fish only, and (3) a combination of fish and copepod predation (Fig. 5), at different levels of \( C_i \) (Fig. 6). When including both fish and copepod predation, the model generally predicts an ontogenetic migration reasonably
Fig. 1. Environmental forcing and model inputs. (A,B) Isolines of density (σt), and shadings of relative fluorescence, (C,D) total counts of nauplii and (E,F) copepodids. (A–F) Left hand panels represent Day 1, and right hand panels Day 2. (G–J) Vertical profiles of model inputs. (G) Temperature. The high T profile is the average from 09:30 h Day 1 to 17:30 h Day 2, and the low T the average of the remaining profiles. (H) Chlorophyll a, (I) light, (J) copepodid predator field, here depicted in groups based on lengths in 10 µm intervals.
similar to that observed (Fig. 5). Without risk of fish predation most species remain in the surface layer throughout the ontogeny (Fig. 5). However, some of the younger nauplii (e.g. Temora spp.) are at high risk from the small Oithona copepodids in the surface layer and may thus benefit from a deeper distribution, also in the absence of fish predators (Fig. 5).

The tendency of larger plankters to remain in the surface layer increases with decreasing fish abundance (Figs. 5 & 6). Similarly, small prey that are relatively more exposed to copepod predators, i.e. nauplii that jump and sink at high frequencies with a noisy motility pattern (e.g. Acartia spp.), require comparatively higher fish concentrations to leave the surface than do silent cruisers (e.g. late Centropages spp.) and cyclopid nauplii that only jump 1 to 2 times min^{-1} (Fig. 6). Hydrodynamic conspicuousness also increases with size (Eqs. 9 & 10), and therefore smaller nauplii may at times benefit from being deeper in the water column than larger nauplii (e.g. Temora spp., Fig. 6). The relative risk from fish and copepod predators thus depends on prey behavior (Fig. 6). Similarly sized nauplii may therefore benefit from different depth distributions, despite similar growth profiles (Figs. 5, 6 & 7).

The model best matches the field observations for those species where an adequate description of behavior is available for all size groups (e.g. Oithona, Acartia, Temora, Centropages). Concordant with the observations, the model predicted DVM to be an unprofitable strategy for all nauplii, as well as for oithonid copepodids under the prevailing environmental conditions, even at very high fish concentrations (Oithona nana, Fig. 4, others not shown).

**DISCUSSION**

**Model structure and process formulations**

Our model differs from other models of optimal distribution patterns in zooplankton by its high content of mechanistic process formulations of predation risk. While several papers have modeled the implications of both fish and invertebrate predation on optimal vertical distributions, few have coupled contemporary predator-prey theory with actual distributions of zooplankton in the field and derived optimal habitat selection over time. De Robertis (2002) considered mechanistic models of visual predation and modeled the timing of vertical migration by allowing euphausiids of different sizes to select depth habitats by minimizing the ratio of mortality and energy gain. Ohman et al. (1983) and Ohman (1990) used life-history theory, estimated mortality and growth rates, and observed diel distributions of predators to evaluate fitness of different migration strategies. Fiksen & Giske (1995) and Fiksen (1997) modeled non-visual predation risk from an empirical size-dependent function, and showed how the relative contribution to mortality from different predators could shape distribution patterns.

Factors such as physical mixing, buoyancy, or UV-radiation may also influence the vertical positioning of zooplankton. Our optimization model ignores these, as well as density-dependent factors, such as ‘safety in numbers’ or competition for food. Giske et al. (1997) modeled a theoretical ideal free distribution of copepods based on such density-dependent processes. They found that food limitation generates dispersion, and that otherwise predation risk and temperature govern distribution patterns and generate aggregation. In the Gullmarsfjord, food was available in high concentrations (Fig. 1H). Although we cannot rule out the role of aggregation as a predator avoidance strategy, we therefore think that the assumption of no density-dependent intra-species interactions is justified.

Implicit in our approach is the assumption of static predator distributions, i.e. we do not consider any response of predators to prey distributions. If predators relocate depending on prey distributions, and prey act as risk-sensitive foragers, the resulting equilibrium distributions of both predators and prey need to be modeled using game theory (e.g. Iwasa 1982). Similarly, we have ignored potential flexibility in motility patterns in response to altered food, predator, or turbulence regimes. Adjustment of motility pattern poten-
Fig. 3. Vertical profiles of nauplii (left hand panels) and copepodids (right hand panels) of various species. Concentrations are expressed relative to the depth-specific total count of nauplii and copepodids shown in the top panel. Species are indicated in panels and values are averages over all profiles. Error bars are SD.
sultantly alters the profitability of the habitats. However, in nauplii, differences in motility between species and stages have larger implications for food and predator encounter than do realistic changes in their respective motility (cf. Titelman & Kiørboe 2003a,b).

We applied a simple measure of fitness, namely maximization of expected future reproductive output (reproductive value). For *Oithona nana*, the computed reproductive values are high, from about 5 in N1 to >30 in adults (Fig. 7C,D). This implies that the population of *O. nana* was growing rapidly at the time. Alternatively, the model overestimates growth (reproduction) or underestimates mortality rates, or else there are other predators in the system that control population growth.

**Distribution patterns of nauplii and Oithona nana**

The vertical profile of risk from copepods and fish at midday for stages N1, N6, and C6 reveal that visual predators contribute little to the mortality risk for *Oithona nana* (Fig. 8A). Also, the model predicts that the small and silently moving *O. nana* nauplii are comparatively less susceptible to copepod predators than later copepodids (Fig. 8A) or nauplii of other species (Figs. 6 & 8). The minimum predation risk from copepods is at 30 m for the nauplii and at 5 m for C6 (Fig. 8C). This can be interpreted in terms of size-dependent predation risk. The largest copepods, which belong to a size category where risk from fish is significant, were found deeper in the water column than the small and intermediated sized copepods (Fig. 1J). The copepodid stages (~0.03 cm) of *O. nana* are small enough to be vulnerable to the larger copepods (Fig. 8A), and are hence predicted to stay near the surface (Fig. 8C,D). The high concentration of young copepodid stages of *O. nana* around 5 m (Figs. 3, 4 & 5) in turn generates a higher risk for small nauplii, explaining the predicted predation minimum at 30 m (Fig. 8A). The specific growth rate depth profile changes after the shift in temperature (Fig. 8B). When running the model with the initial temperature profile (i.e. high °T), the temperature peak at 5 m (Fig. 1G) makes this depth the most profitable one for all stages (Fig. 4). However, when forcing the model with the other temperature profile (i.e. low °T, Fig. 1G), the deep (30 m) alternative becomes the most profitable depth for nauplii, while C6 should remain in the surface habitat (Fig. 8D). Contrary to that of the nauplii, the growth of C6 does not decrease sufficiently to offset the benefit of reduced predation risk near the surface. This situation parallels other reports of optimal distributions shaped by size-dependent predation risk. For example, Ohman et al. (1983) found that *Pseudocalanus* sp. displayed reverse DVM when its invertebrate predators left the surface to avoid fish predation. Although the observed situation in Gullmarsfjord was apparently more static, it may still be interpreted in terms of size-dependent predation chains.

The bimodal distribution of the older copepod stages in *Oithona nana* is less intuitively understood (Fig. 4). The predicted optimal depth for all stages of *O. nana* is near the surface under the initial temperature regime (Figs. 4 & 5). The modeled rate of predation by other copepods peaks near 15 m (Fig. 8A), which is also the depth that *O. nana* avoids (Fig. 4). Given the initial temperature profile (Fig. 1G), the fitness of all stages peaks near the surface, but profiles are relatively flat.
with local minima around 20 m (Fig. 8C). After the disappearance of the sub-surface temperature peak, the optimal depth for stages <C2 is 30 m (Fig. 8D). The bimodal distribution pattern could therefore result from frequent switches in the hydrographic regime, and corresponding alterations in optimal habitats over short time spans. Alternatively, the bimodal pattern may result directly from predation. If *O. nana* was initially uniformly or randomly distributed above the thermocline, then predation from large copepods could potentially reduce the concentration of the copepodids substantially at intermediate (15 to 25 m) depths. A third possibility is that the feeding habits of *O. nana* may affect its fitness profile and optimal depth. Oithonids are true ambush feeders, feeding predominately on mobile prey and sinking fecal pellets (Sabatini & Kiørboe 1994, Svensen & Kiørboe 2000). A deeper distribution of *O. nana* C3 to C5, below the bulk of other zooplankton, may secure a steady supply of sinking fecal pellets. Similarly, the phytoplankton composition below the deep chlorophyll maximum might be the motivator of this distribution. Bimodal dis-

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Fig. 5. Stage-specific depth distribution for *Acartia clausii*, *Calanus finmarchicus*, *Centropages typicus*, *Microcalanus pusillus*, *Oithona nana*, *Paracalanus parvus*, *Temora longicornis*, *Pseudocalanus elongatus* and *O. similis*. The box plots were constructed from mean vertical percentage distributions obtained by combining all sampling times. The boxes represent the 25 and 75%, the line the median, the whiskers the 5 and 95 %. Overlaid lines are model simulations with different predator regimes: (○) copepods (ambush) only, (□) fish only, and (△) both copepods and fish. We applied velocities, time budgets and jump frequencies from the literature. For nauplii: *Oithona* spp (*Oithona* sp., Paffenhöfer 1993); *Pseudocalanus* (Landry & Fagernes 1988); *Paracalanus* (*P. aculeatus*, Paffen-höfer et al. 1996); all others (Titelman & Kiørboe 2003a). For copepods: *Oithona* spp. (*O. similis*, Svensen & Kiørboe 2000); *Temora* (Van Duren & Videler 1995); *Calanus* (Mauchline 1998); all others (Jonsson & Tiselius 1990, Tiselius & Jonsson 1990). No behavioral data was available for *Microcalanus*. We assumed $u^* = 0.004$ cm s$^{-1}$ (*O. similis*, Kiørboe & Visser 1999), $C_f = 0.001$ l$^{-1}$, and the initial temperature regime (high $T$) in all depicted simulations. Note that the bimodal distribution of *O. nana* is disguised in this graph (cf. Fig. 4).

Sizes for N1 and females (F) used in the model are indicated in the panels. N = nauplius, C = copepodid, M = male
Distributions of copepods in the same developmental stages (Durbin et al. 1995, Hays et al. 2001) may be explained by internal energy conditions, with fatty individuals in safe, deep habitats and less fatty individuals in risky, shallow habitats (Fiksen & Carlotti 1998). However, *O. nana* possesses no lipid sac and suffers only a minor predation risk from fish.

**Fig. 6.** (A) Critical fish concentration $C_{f}^{*}$ required for an animal with various motility patterns to descend from the surface habitat to below 20 m in the water column. Behaviors are representative of the 2 extremes of jump-sink motility in nauplii (i.e. *Oithona* spp. and late *Acartia tonsa*) and of a continuous cruiser (e.g. late *Temora longicornis*) (Titelman & Kiørboe 2003a). The motility parameters indicated in the panel are applied to an *O. similis* (Sabatini & Kiørboe 1994) in order to isolate the impact of motility. Simulations are assuming no ontogenetic changes in motility. All other model parameters are as in Figs. 3, 4 & 5. Small nauplii of the noisier types cannot descend because their optimal habitat is already deeper (see text). (B) Predation mortality (black line) from fish and copepods for an animal with various motility patterns at 5 m depth. Temperature and predation regime are as in Fig. 1. Symbols and explanations as in (A)

**Fig. 7.** *Oithona nana*, *Acartia clausii* and *Calanus finmarchicus*. Model predictions of predation rates from fish and copepods at midday. Behavior is as in Fig. 5, all other model parameters and assumptions as in Figs. 4, 5 & 6.
For nauplii and copepodids of small species such as *Oithona* spp., the predation risk from copepods dominates over that from fish (Figs. 7 & 8A), and depends strongly on motility (Fig. 6). Hydrodynamically 'noisy', frequently jumping nauplii (e.g. *Acartia* spp.) experience higher risk of encounter with copepod predators compared to cruising nauplii (e.g. *Centropages* spp.) or nauplii with a low frequency jump-sink type of motility (*Oithona* spp.), and therefore require comparatively higher concentrations of fish to descend to deeper waters (Fig. 6).

Changes in the physical environment may also alter distributions. Incze et al. (2001) pointed to behavioral differences between nauplii based on their vertical distributions in response to wind forcing. Some species aggregated at depth during periods of wind and closer to the surface upon cessation of wind, while other species resisted downward mixing and remained at the surface (Incze et al. 2001). Our field data reveal that nauplii of most species reside near the surface (Figs. 1, 3 to 5). Exceptions are *Oithona similis*, *Oncaea borealis*, and *Coryceaus anglicus* nauplii (Figs. 3 & 5), whose behavior generates very low volume encounter rates with copepod predators (Titelman & Kørboe 2003b). These nauplii remain virtually motionless most of the time and relocate occasionally by a brief jump, a few times per minute (Paffenhöfer 1993, Paffenhöfer et al. 1996). Such a motility pattern may allow a higher degree of vertical overlap between nauplii and larger copepodids. This accords with the dispersed distribution patterns of *Oithona similis* and *C. anglicus* nauplii, and the deeper distribution of *Oncaea borealis* and *Microsetella norvegica* nauplii (Figs. 3 & 5). In addition, copepodids of the latter 2 genera, both of which are egg-bearing, are commonly associated with feeding on marine snow (Green & Dagg 1997), and may therefore benefit from a deeper distribution.

**Lack of DVM**

As pointed out recently in 2 reviews of DVM, it is also possible that a low indication of DVM results from individuals moving asynchronously; feeding in food-rich habitats until their stomachs are filled and then...
returning to safer habitats (Hays 2003, Pearre 2003). However, as we had no way of following individuals, we could not evaluate this hypothesis.

The weak gradient in temperature and growth opportunities (Fig. 8B) partially explains the general lack of DVM that we observed (Oithona nana, Fig. 4). This means that the trade-off between growth and predation risk is weak or absent, such that an animal can move to a deeper, safer habitat without losing much in terms of growth. Another factor that limits the profitability of both normal and inverse DVM in situations such as the one presented here is the static distribution and relatively high concentrations of larger copepods at intermediate depths (Fig. 1J). Nauplii and small copepods, like O. nana, would have to pass through a danger zone to migrate between shallow and deeper habitats (Fig. 8A). Jellyfish, chaetognaths and other zooplankton predators, which were not sampled here, may similarly constitute layers of danger.

Migration requires increased cruising speeds or jump frequencies, which leads to much higher risk of being detected and encountered by copepods or other rheotactic predators (Kiørboe et al. 1999, Titelman 2001, Saito & Kiørboe 2001). Thus, moving through a zone with high predator densities may not be strategic — despite potential benefits on the other side. We therefore suggest that sometimes, and especially if you are small or hydrodynamically inconspicuous to begin with, it is better to remain calm and stay where you are.

**Summary and conclusions: influence of motility on distributions**

We have tried to bridge the traditional approaches of descriptive field work and theoretical modeling to examine the vertical distribution and habitat selection of copepod developmental stages. Naturally, distributions of animals result from trade-offs between eating and growing on the one hand, and avoiding being eaten on the other. Despite the limitations of both the field sampling and the model, our results suggest that the distribution of small copepods, especially the nauplii and small species such as Oithona spp., differs from that of the better studied large species and later life history stages. Nauplii and small oithonids are less sensitive to visual predators and more sensitive to invertebrate predators such as larger copepods. While the relative sensitivity to fish predators increases with size, the risk of getting killed by larger copepods is, in addition to size, very sensitive to motility (our Fig. 6B, Titelman & Kiørboe 2003b). One may therefore expect that small nauplii and copepodids adjust their motility and distribution in relation to non-visual predators. For the nauplii, which are generally most sensitive to copepod predation, one may further expect that the relatively silent motility types, which spend most of their time motionless (e.g. Oithona spp. and Oncaea spp.) or slowly swimming (e.g. N3 to N6 Temora spp., Centropages spp.) can handle more overlap with predators, than larger (late Calanus spp. nauplii) or faster moving hop-sink types (e.g. N1 to N6 Acartia spp.), which generate strong hydrodynamic signals. Albeit that factors such as food layers, density stratification and advection all interplay to determine the distribution of plankton, we suggest that individual behavior may be an important determinant of a zooplankter’s optimal depth location, mediated by its effects on the relative susceptibility to functionally different predators such as visually hunting fish or rheotactic copepods. (see also Fiksen et al. in press)

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