Fluid motion and solute distribution around sinking aggregates II: Implications for remote detection by colonizing zooplankters

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INTRODUCTION

Large, fast-sinking marine snow aggregates form by coagulation and other processes in the water column and are believed to be the main vehicles for vertical material transport in the ocean (Fowler & Knauer 1986). However, aggregates are also degraded in the water column. Bacteria hydrolyse (Smith et al. 1992, Grossart & Simon 1998) and remineralise (Ploug et al. 1999) aggregated material, and fish and zooplankters feed on aggregates (Lampitt et al. 1993, Larson & Shanks 1996). Thus, turnover rates of aggregated material may be high, and a substantial fraction of aggregated material may be remineralized before the aggregate leaves the upper mixed layer (Kiørboe 2000).

Degradation processes continue below the mixed layer, and lead to a decline in particle flux with depth (Martin et al. 1987, Pace et al. 1987). Banse (1990) proposed that zooplankton plays a pivotal role in the degradation of aggregates.

Grazing on large aggregates can occur in 2 ways, viz., fish and large zooplankters engulf intact snow aggregates (Larson & Shanks 1996, Dilling et al. 1998), or smaller zooplankters colonize aggregates and feed on their constituents (Alldredge 1976, Shanks & Walters 1997). Recent observations collected by divers and submersibles demonstrate that aggregates may house a very abundant fauna of colonizing zooplankters, both within the euphotic zone, and at depth, and that these contribute significantly to the degradation of aggregates (reviewed by Kiørboe 2000). This fauna is dominated by small, mm-sized crustaceans, mainly harpacticoid copepods and poecilostomatoid copepods of...
the genus *Oncaea* (Fig. 1). Although occurring in the water column, some members of these genera appear to be adapted to a life on solid surfaces (see Paffen-höfer 1993) such as that offered by aggregates. The quantity of colonizing zooplankters observed on aggregates suggest that the zooplankters will have to search substantial volumes of water in order to account quantitatively for the observed association and, therefore, that copepods need to be able to remotely detect aggregates (Kiørboe 2000).

Remote detection may be by chemical or hydro-mechanical cues, and a sinking aggregate leaking hydrolysation products potentially provides both. A sinking aggregate generates velocity gradients in the ambient water that may be detected by copepods (Fields & Yen 1997, Kiørboe et al. 1999). Bacteria solubilize aggregated particulate material at a higher rate than at which they assimilate the solutes (Smith et al. 1992, Vetter et al. 1998), and solutes therefore leak out of an aggregate and paint a chemical trail in its wake. Copepods are known to respond behaviorally to organic solutes, particularly amino acids (Poulet & Ouellet 1982, Gill & Poulet 1988).

The purpose of this study is to examine whether hydromechanical and/or chemical signals generated by a sinking aggregate may allow remote detection by copepods in a way that can account quantitatively for the observed abundance of colonizing zooplankters on aggregates. We use a model derived from first principles to describe the fluid motion and solute distribution around a sinking sphere and combine model results with what is known about amino acid leakage rates from aggregates and the sensitivity of copepods to amino acids and hydromechanical signals.

### THE FLOW AND CONCENTRATION FIELDS

To characterize the hydrodynamic and chemical signals generated by a sinking aggregate we modeled the aggregate as a rigid sphere sinking at constant speed. The flow is described by solving the Navier-Stokes’ equations numerically, and the solute distribution is described by subsequently solving the governing diffusion-advection equation. A full description of the model and of the numerical approach is given in Kiørboe et al. (2001, in this issue). The symbols used here are defined in Table 1.

### Flow field

Following the convention of Karp-Boss et al. (1996) we take the radius of the sphere (a) as the characteristic length, and the sinking velocity of the sphere (U) as the characteristic velocity. Normalization of the Navier-Stokes’ equations show that the flow field is then solely a function of the Reynolds number

\[
Re = \frac{aU}{v}
\]

where v is the kinematic viscosity. We have made computations at Re up to 20, which cover the range of Re characteristic of marine snow (Alldredge & Gotschalk 1988). We computed streamlines and velocity distributions around the falling sphere (Kiørboe et al. 2001). Of particular relevance for hydrodynamic detection of the sinking aggregate is the spatial distribution of the

### Table 1. Definition of symbols and standard parameter values used in computations

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
<th>Unit</th>
<th>Default value</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>Aggregate radius</td>
<td>cm</td>
<td>–</td>
</tr>
<tr>
<td>A</td>
<td>Cross-sectional area of plume with concentrations exceeding C*</td>
<td>cm^2</td>
<td>–</td>
</tr>
<tr>
<td>C</td>
<td>Solute concentration</td>
<td>M</td>
<td>–</td>
</tr>
<tr>
<td>C*</td>
<td>Threshold concentration for behavioral response</td>
<td>M</td>
<td>4 × 10^{-8}</td>
</tr>
<tr>
<td>D</td>
<td>Diffusion coefficient</td>
<td>cm^2 s^{-1}</td>
<td>10^{-5}</td>
</tr>
<tr>
<td>Q</td>
<td>Leakage rate of solutes from aggregate</td>
<td>mol s^{-1}</td>
<td>Eq. (8)</td>
</tr>
<tr>
<td>N_a</td>
<td>Normalized abundance of zooplankters on an aggregate</td>
<td>ml aggregate^{-1}</td>
<td>–</td>
</tr>
<tr>
<td>r</td>
<td>Copepod equivalent radius</td>
<td>cm</td>
<td>0.05</td>
</tr>
<tr>
<td>S</td>
<td>Signal strength</td>
<td>cm s^{-1}</td>
<td>–</td>
</tr>
<tr>
<td>S*</td>
<td>Threshold signal strength</td>
<td>cm s^{-1}</td>
<td>4 × 10^{-3}</td>
</tr>
<tr>
<td>U</td>
<td>Aggregate sinking velocity</td>
<td>cm s^{-1}</td>
<td>Eq. (7)</td>
</tr>
<tr>
<td>v</td>
<td>Copepod swimming velocity</td>
<td>cm s^{-1}</td>
<td>0.1</td>
</tr>
<tr>
<td>β</td>
<td>Encounter volume rate</td>
<td>cm^3 s^{-1}</td>
<td>Eq. (6)</td>
</tr>
<tr>
<td>Δ</td>
<td>Maximum deformation rate</td>
<td>s^{-1}</td>
<td>–</td>
</tr>
<tr>
<td>ν</td>
<td>Kinematic viscosity</td>
<td>cm^2 s^{-1}</td>
<td>10^{-2}</td>
</tr>
</tbody>
</table>
maximum deformation rate (= maximum component of the stress tensor; see Kiørboe et al. 1999), which was computed numerically. The spatial distribution of deformation rate for $Re < 1$ (Stokes’ flow), 1, and 10 are shown in Fig. 2. With increasing $Re$, the deformation field becomes compressed upstream of the sphere, elongated downstream, but retains approximately the same width.

Concentration field

Given the flow field, the concentration field is determined by the Peclet number ($Pe$)

$$Pe = \frac{Ua}{D}$$

where $D$ is the diffusion coefficient, and by the leakage rate of solutes from the aggregate ($Q$, mol s$^{-1}$). The diffusion-advection equation was solved with the outer boundary condition, that far from the sphere the solute concentration approaches the ambient concentration $C_\infty$:

$$C_{|r=\infty} = C_\infty$$

(3)

We examined 2 alternative inner boundary conditions by assuming either that the solute concentration is constant over the surface of the sphere (Dirichlet boundary condition):

$$C_{|r=a} = c_0$$

(4)

or that the flux is constant over the surface of the sphere (Neumann condition):

$$\frac{\partial C}{\partial r}_{r=a} = c_0$$

(5)

Observations suggest that neither of the 2 alternative boundary conditions accurately apply to marine snow aggregates, but that these describe extremes (Kiørboe et al. 2001).

All computations were made with dimensionless variables and were converted to absolute numbers by inserting absolute (case-specific) values of $U$, $a$, and $Q$ and in all cases assuming $v = 10^{-2}$ cm$^2$ s$^{-1}$ and $D = 10^{-5}$ cm$^2$ s$^{-1}$ (characteristic of seawater and amino acids at 20°C).

Examples of computed concentration fields at $Pe = 0$ (pure diffusion), and $Pe = 100$, and 10000 (all at $Re = Pe/1000$) are shown in Fig. 3. With increasing $Pe$, the plume becomes increasingly elongated. By $Pe = 100$ the plume is very long and slender, and concentrations both upstream and to the sides of the sphere drop almost immediately to ambient concentrations.
Before examining potential detection and encounter mechanisms, we first summarize observations of colonization rates with which model predictions can be compared. Kiørboe (2000) compiled the available information on the abundance of zooplankters sitting on aggregates (no. aggregate–1) in surface waters and found that normalized abundances (\(N_A, \text{ml aggregate}^{-1}\)), which were obtained by dividing absolute abundances with ambient animal concentrations (no. \(\text{ml}^{-1}\)), scaled approximately with aggregate radius (\(a\)) square—

\[N_A (\text{ml aggregate}^{-1}) = 2240 a^{2.27}\]

Most colonizing crustaceans appear to reside on individual aggregates only for short periods of time before they swim away to find a new aggregate (Alldredge 1972, Shanks & Walters 1997). Shanks & Walters (1997) estimated an average residence time of 3 min. By dividing normalized abundance by average residence time, an average ‘encounter volume rate’ (\(b, \text{ml s}^{-1}\)) can be estimated:

\[b (\text{ml s}^{-1}) = N_A/180 s = 12.5 a (\text{cm})^{2.27}\] (6)

Thus, the encounter rate of a 0.1 cm radius aggregate is about (0.067 ml s\(^{-1}\)) = 0.25 l h\(^{-1}\) and of a 1 cm aggregate (12.5 ml s\(^{-1}\)) = 45 l h\(^{-1}\) (Table 2). Depending on the encounter mechanism, the encounter volume rate can be interpreted as the volume of water that a sinking aggregate scavenges for zooplankters per unit time, or as the volume of water that an individual zooplankter searches for aggregates per unit time (akin to clearance rate). The rate at which an individual zooplankter encounters aggregates is given by \(\beta C_A\), and the colonization rate of an aggregate (no. zooplankters arriving per unit time and aggregate) as \(\beta C_Z\), where \(C_A\) and \(C_Z\) are the concentrations of aggregates and zooplankters.

### ZOOPLANKTON COLONIZATION RATES

Knowing the flow and concentration fields around a sinking aggregate, we can now examine different detection and encounter mechanisms and compare their predictions with Eq. (6) (Table 2). The simplest mechanisms of encounters between sinking marine snow and colonizing zooplankters is either that the sinking aggregate ‘scavenges’ colonizers as it sinks and that the colonizers can be considered inert (behaviorless) particles, or that a cruising colonizer simply bumps into the aggregate. For the latter mechanism, the encounter volume rate is simply \(\pi a^2v\), where \(v\) is the cruising velocity of the colonizer. For a typical \(v\) of a 1 mm sized copepod of 1 mm s\(^{-1}\), this yields estimates that are 1 to 2 orders of magnitude less those estimated by Eq. (6) (Table 2).

To estimate encounter volume rates by scavenging we need to know aggregate sinking velocities (\(U\)). All dredge & Gotschalk (1988, 1989) made \textit{in situ} observations of sinking velocities of aggregates of various sizes, and provided the following empirical relationships between sinking velocity and equivalent radius:

\[U (\text{cm s}^{-1}) = 0.13a (\text{cm})^{0.26}\] (Alldredge & Gotschalk 1988) (7a)

or

\[U (\text{cm s}^{-1}) = 0.2a (\text{cm})\] (Alldredge & Gotschalk 1989) (7b)

These relations differ mainly in their scaling, but provide sinking rate estimates of similar magnitudes for aggregates in the size range typical for (most) marine snow, 0.1 to 1 cm radius. Other relations mainly yielding lower sinking velocities have been reported in the literature. For example, direct \textit{in situ} estimates provided by Asper (1987) and Pilskaln et al. (1998) for

### ENCOUNTERS WITHOUT REMOTE DETECTION

<table>
<thead>
<tr>
<th>Mechanism</th>
<th>Equation</th>
<th>Estimated encounter volume rate (ml s(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Observed encounter rate</td>
<td>(6)</td>
<td></td>
</tr>
<tr>
<td>No perception, cruising copepod</td>
<td>(\pi a^2v)</td>
<td>6.7 \times 10^{-2} 12.5</td>
</tr>
<tr>
<td>No perception, scavenging aggregate</td>
<td>(1.5\pi r^2U)</td>
<td>3.1 \times 10^{-3} 0.31</td>
</tr>
<tr>
<td>Hydromechanical perception, stationary copepod</td>
<td>(\pi R^2U)</td>
<td>8.4 \times 10^{-4} 1.5 \times 10^{-3}</td>
</tr>
<tr>
<td>Chemical perception, cruising copepod</td>
<td>(\pi a^2v)</td>
<td>9.3 \times 10^{-3} 0.72</td>
</tr>
<tr>
<td></td>
<td>(\pi R^2U)</td>
<td>2.6 \times 10^{-3} 1.1</td>
</tr>
<tr>
<td></td>
<td>(\pi a^2v)</td>
<td>4 \times 10^{2} 12.7</td>
</tr>
<tr>
<td></td>
<td>(\pi R^2U)</td>
<td>7.4 \times 10^{-2} 14.3</td>
</tr>
</tbody>
</table>

\(a = 0.1 \text{ cm} \)                  \(a = 1.0 \text{ cm} \)

\textsuperscript{a}Computed for a Neuman boundary condition; a Dirichlet boundary condition yields practically identical results
1 to 5 mm diameter aggregates are 3 to 50 times lower than those predicted by Eq. (7). We shall use the above relations and, by sensitivity analysis, demonstrate that the below conclusions are conservative if sinking velocities are in fact lower.

A first, upper, estimate of the encounter volume rate due to scavenging is \( \pi a^2 U \), which combined with Eq. (7a) yields approximately the correct scaling but again gives estimates that are 1 to 2 orders of magnitude less those predicted by Eq. (6). These are upper estimates, however, because hydrodynamic effects have been ignored. A sinking aggregate pushes the ambient water around itself, and taking this effect into account, significantly changes the scaling and reduces the estimated encounter volume rate, particularly for the larger particles. The encounter volume rate in this situation is the volume flow of water passing the aggregate within a distance \( r \), where \( r \) is the radius of the colonizer. If we assume Stokes’ flow around the sinking aggregate (i.e., low \( Re \)), an analytical solution exists, and the encounter volume rate becomes \( \sim 1.5 \pi r^2 U \) (Spielman 1977, Kiørboe & Titelman 1998).

For a 1 mm sized colonizer, this yields estimates that are 2 to 4 orders of magnitude less those predicted by Eq. (6) (Table 2). At higher \( Re \), numerically estimated rates are slightly higher than those estimated for Stokes’ flow, e.g., a factor of 4 at \( Re = 20 \), a 0.1 cm copepod and a 1 cm (radius) aggregate, but still 3 orders of magnitude less that predicted by Eq. (6). Lower aggregate sinking velocities than those assumed (Eq. 7) yield lower encounter volume rates.

The above considerations thus suggest that remote detection is required to account for the observed abundances of colonizing zooplankters on marine snow aggregates.

HYDROMECHANICAL PERCEPTION

Copepods are equipped with mechanosensory setae extending in particular from the antennae. Neurophysiological studies have revealed that bending of these setae may elicit a neurophysiological response, that the setae are velocity rather than displacement sensors, and that a threshold velocity difference (signal strength) between the copepod and the ambient fluid of about 20 \( \mu \text{m s}^{-1} \) is required to elicit neurophysiological response (Yen et al. 1992). Experiments have demonstrated that signal strength several orders of magnitude higher is required to elicit escape responses in copepods (Kiørboe et al. 1999). Attack responses appear to be elicited at signal strengths closer to the neurophysiological limit, e.g., 40 \( \mu \text{m s}^{-1} \) in Oithona similis, a copepod with particularly long setae (Svensen & Kiørboe 2000), and about an order of magnitude higher in the predatory copepod Centropages typicus (Titelman 2001). A copepod will be entrained in the fluid flow generated by a moving particle if this is much larger than the copepod. Velocity gradients in this flow will cause a velocity difference between the copepod and the ambient fluid that, in turn, will cause extending setae to bend. Specifically, the copepod responds to the fluid deformation rate (\( \Delta \)), and the signal strength is \( S = \Delta r \), where \( r \) is the equivalent radius of the copepod (Kiørboe & Visser 1999, Kiørboe et al. 1999).

Consider now a mm-sized copepod. If we conservatively assume a threshold signal strength for a behavioral response of \( S^* = 40 \mu \text{m s}^{-1} \), then the threshold deformation rate required for perception is \( S^*/r = 40/500 \text{ s}^{-1} = 0.08 \text{ s}^{-1} \). Fig. 4A shows the 0.08 \( \text{s}^{-1} \) isoline for an aggregate at \( Re = 5 \), corresponding to an aggregate of radius 0.5 cm with a settling velocity of 0.1 cm \( \text{s}^{-1} \) (Eqs. 7a,b). Assume now that the colonizing copepod utilizes an ambush strategy, i.e., it sits motionless in the water column waiting for fluxing particles to pass by within a distance at which they can be perceived. The horizontal extension of the deformation field then becomes what matters, and the encounter volume rate is given by \( \pi R^2 U \), where \( R \) is the perception distance or half the horizontal extension of thresh-

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**Fig. 4.** (A) Deformation field for a of 0.5 cm radius sphere sinking at 0.1 cm s\(^{-1}\) (\( Re = 5 \)). The isolines are 0.002, 0.008, 0.02, and 0.08 s\(^{-1}\) and depict the distance at which a 1 mm copepod can perceive a sinking aggregate, provided it has a threshold sensitivity, \( S^* \), of 1, 4, 10, or 40 \( \mu \text{m s}^{-1} \), respectively. (B) The detection distances required to account for encounter volume rates computed from observed abundances of zooplankters on aggregates (solid lines), as well as those computed assuming a threshold sensitivity of 40 \( \mu \text{m s}^{-1} \) for a 1 mm copepod (dotted lines), both as a function of aggregate size. Computations have been made assuming 2 different size-sinking velocity relations.
old deformation isoline. (For simplicity and to be conservative, in computing the encounter volume rate we ignore that the fluid motion generated by the sinking aggregate will move the colonizer slightly.) In our example, this corresponds to a detection distance (measured from the center of the particle) of 0.7 cm, i.e., only slightly more than the particle’s radius. The detection distance required to account for the encounter volume rate predicted by Eq. (6) is 2.9 cm, or 4 times higher. A detection distance of this order would require a threshold deformation rate of 0.002 s\(^{-1}\), or a threshold signal strength of 1 \(\mu\)m s\(^{-1}\), i.e., more than an order of magnitude less than the neurophysiological limit. A similar analysis for a range of particle sizes (Fig. 4B, Table 2) generalizes this conclusion. Decreasing the encounter volume rate estimated from Eq. (6) by an order of magnitude relaxes the sensitivity requirement to 25 \(\mu\)m s\(^{-1}\), i.e., close to the neurophysiological limit. Decreasing the particle sinking velocity by a factor of 10 decreases the required threshold sensitivity to 10\(^{-2}\) \(\mu\)m s\(^{-1}\). Even a copepod with a cruising strategy of 10 decreases the required threshold signal strength of 1 \(\mu\)m s\(^{-1}\), i.e., more than an order of magnitude less than the neurophysiological limit. A similar analysis for a range of particle sizes (Fig. 4B, Table 2) generalizes this conclusion. Decreasing the encounter volume rate estimated from Eq. (6) by an order of magnitude relaxes the sensitivity requirement to 25 \(\mu\)m s\(^{-1}\), i.e., close to the neurophysiological limit. Decreasing the particle sinking velocity by a factor of 10 decreases the required threshold sensitivity to 10\(^{-2}\) \(\mu\)m s\(^{-1}\). Even a copepod with a cruising strategy would not benefit much from the deformation rate generated by the sinking particle, because the deformation rate attenuates rapidly in all directions to values unlikely to elicit behavioral responses (Fig. 2).

Copepods that colonize aggregates, mainly harpacticoid copepods and Pocillostomatoid copepods of the genus *Oncaea*, are typically characterized by very short antennae (Fig. 1). This morphology appears inconsistent with a high sensitivity to hydrodynamic signals. Our considerations thus suggest that copepods that colonize marine snow aggregates are unlikely to utilize hydrodynamic cues to detect these.

**CHEMICAL PERCEPTION**

The long slender plume of solutes left behind the leaking and sinking aggregate (Fig. 3) may be detected by a horizontally cruising copepod. Such a detection mechanism has in fact been demonstrated for the planktonic shrimp *Acetes sibogae* (Hamner & Hamner 1977). Shrimps encountering the scent trail from a sinking food particle follow the chemical track and capture the particle. A similar mechanism has been described for male copepods *Calanus marshallae* searching for females (Tsuda & Miller 1998). The slowly sinking female releases pheromones, thus leaving a plume of male-attractant similar to that shown in Fig. 3. On encountering the pheromone plume, the male swims down the chemical trail to find the female. These situations resemble the one we have in mind here.

Aggregates do not release pheromones, but leak DOM. In order to examine the potential for chemical perception we need first to consider likely components of DOM that copepods may detect, at what rate these components leak out of an aggregate, and at what threshold concentration a copepod may detect and respond to the solute.

**Sensitivity to chemical stimuli**

Several studies have demonstrated that crustaceans, including copepods, are very sensitive and respond behaviorally to dissolved amino acids, with response thresholds to specific amino acids being often much less than typical background concentrations of total amino acids. Bulk background concentrations of dissolved amino acids are on order \(<10^{-7}\) to \(10^{-9}\) M in surface waters, less at depth (Mopper & Lindroth 1982, Carlucci et al. 1984, Poulet et al. 1991). Behavioral responses in crabs have been recorded at amino acid concentrations as low as \(10^{-12}\) to \(10^{-20}\) M (Fuzessery & Childress 1975, Pearson & Olla 1977, Thomson & Ache 1980). In copepods, neurophysiological responses in *Undinula* sp. have been observed at concentrations of the amino acid serine down to \(10^{-9}\) M (Yen et al. 1998). Amino acids at concentrations of \(10^{-5}\) M may induce swarming behavior in *Eurytemora herdmani* (Poulet & Ouellet 1982), and Gill & Poulet (1988) reported that enhanced beat frequency of the first maxilla—a more direct index of feeding behavior—occurred at concentrations of \(10^{-6}\) M, but not at \(10^{-8}\) M, in *Temora longicornis*. There are likely to be major species-specific differences in sensitivity, and motor responses in the more sensitive copepods may therefore be expected at amino acid concentrations down to \(10^{-7}\) to \(10^{-9}\) M or less.

**Leakage of amino acids from aggregates**

Marine snow aggregates do leak amino acids (Smith et al. 1992, Grossart & Simon 1998, Vetter et al. 1998). Bacteria attached to marine snow aggregates hydrolyse particulate substances to solute monomers faster than these are taken up by the bacteria (Smith et al. 1992) and solutes hence leak out of the aggregate. Based on Smith et al. (1992), an estimate of the leakage of free amino acid-N as a fraction of aggregate N is about 0.2 d\(^{-1}\). Alldredge (1998) gives the size-dependent nitrogen content of marine snow aggregates as PON (mol N) = 6.3 \(\times\) \(10^{-7}\) \(r^{1.5}\) (cm). Because 1 mol N \(\equiv\) 0.75 mol amino acid, the size-dependent amino acid leakage (Q) rate becomes:

\[
Q (\text{mol amino acid s}^{-1}) = 10^{-12} r^{1.5} (\text{cm})
\]

(8)

The actual exudation rates measured by Smith et al. (1992) in ‘larger than 2 mm diameter’ aggregates, 0.3 to 0.8 nmol aggregate\(^{-1}\) h\(^{-1}\), are somewhat higher than
that estimated by Eq. (8) for 2 mm diameter aggregates, 0.11 nmol aggregate$^{-1}$ h$^{-1}$. Grossart & Simon (1998) provided the only other measurements of amino acid leakage rates in sinking aggregates, and found that 0.27 cm (radius) aggregates leak between 0.45 and 2.25 nmol amino acid h$^{-1}$. This is consistent with that predicted by Eq. (8), 0.6 nmol h$^{-1}$. (The numbers given in Grossart & Simon were wrong, but have been corrected here; Grossart pers. comm.)

We may first examine whether the solute concentration produced by a leaking and sinking aggregate is sufficiently high to elicit motor responses. It can be shown from the diffusion-advection equation that the surface concentration of solutes is

$$C_{r,a} = \frac{Q}{4\pi Da Sh}$$  \hspace{1cm} (9)

where $Sh$ is the dimensionless Sherwood number and $C_{r,a}$ the surface concentration. Combining Eqs. (8) & (9) and the $Sh$ numbers computed by Kiørboe et al. (2001), yields surface concentration as a function of particle size (Fig. 5A; here employing a Dirichlet boundary condition). For aggregates with a radius less than ca 0.02 cm, surface concentration drops rapidly with decreasing particle size, suggesting that there is a minimum particle size for chemical detection. For bigger particles, however, the surface concentration is largely independent of particle size and close to $5 \cdot 10^{-7}$ M. Employing instead a Neumann boundary condition yields even higher surface concentrations in the downstream direction. This suggests that there is a potential for chemical detection.

**Encountering the plume**

The rate at which a cruising copepod encounters chemical plumes from aggregates depends on the vertical cross-sectional area of the plume with concentrations exceeding the threshold ($A$) multiplied by the horizontal swimming velocity of the copepod ($v$) and the concentration of aggregates. The encounter volume rate is $A \times v$. Assuming $v = 1$ mm s$^{-1}$ we have computed the area required to account for the encounter volume rates predicted by Eq. (6) and in Fig. 5B depicted the threshold concentration that yields that area. Almost independent of particle size, the threshold concentration required is on order $4 \cdot 10^{-8}$ M. Varying the default parameter values of swimming velocity, leakage rate, and encounter volume rate over an order of magnitude (i.e., from $\times 1/3$ to $\times 3$ the default values) yield required threshold concentrations of the same order, with required threshold concentration being most sensitive to variation in leakage rate (Table 3). Decreasing the aggregate sinking velocity by a factor of 10 increases the required threshold concentration by 50%. The independence of the estimated threshold of particle size implies that the scaling is correct.

The length of the plume holding concentrations exceeding the threshold increases with aggregate size (Fig. 5C) and reaches about 1 m in length (~100 particle radii) for a 1 cm radius aggregate.
Table 3. Effect of varying parameter values from $\times 1/3$ to $\times 3$ of their default values (horizontal swimming velocity 0.1 cm s$^{-1}$, leakage rate: Eq. 8; encounter volume rate: Eq. 6) on the estimated threshold concentration ($C^*$) of amino acids required to elicit a behavioral response to a 0.5 cm radius sinking aggregate. The effect is expressed as the estimated range in $C^*$ over the applied change in parameter value.

<table>
<thead>
<tr>
<th>Parameter changed</th>
<th>None</th>
<th>Swimming velocity</th>
<th>Leakage rate</th>
<th>Encounter volume rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Range in estimated $C^*$, M</td>
<td>$4 \times 10^{-8}$</td>
<td>$2.4 - 7 \times 10^{-8}$</td>
<td>$1.1 - 11.4 \times 10^{-8}$</td>
<td>$2.4 - 7 \times 10^{-8}$</td>
</tr>
</tbody>
</table>

DISCUSSION AND CONCLUSIONS

All the encounter mechanisms examined here, except for scavenging, yield scalings of encounter volume rates with aggregate size that are consistent with the observed scaling of the abundances of colonizers on aggregates. The encounter volume rates inferred from observed abundances suggest that remote detection is required for colonizers to encounter aggregates. This suggestion hinges, however, on the estimated short average residence of zooplankters on the aggregates. Longer residence times would relax the requirements for remote detection. The 3 min residence time reported by Shanks & Walters (1997) is based on laboratory experiments, but is consistent with observations of Alldredge (1972), who noted that *Oncaea mediterranea* feeding on aggregates were highly active but spent only little of their time (5%) actually on the aggregates. One may ask why a small zooplankter that has just succeeded to locate a rich feeding environment in the form of an aggregate would bother to leave this so soon again, just to spend time finding a new one? This would at first appear inconsistent with predictions from optimal foraging theory, unless different aggregates provide very different feeding environments, and the animals would need to test many aggregates until eventually finding a nutritive one. However, in any particular environment, aggregates are probably more similar than they are different, because one source of aggregates would typically dominate (e.g., diatom aggregates during diatom blooms; marine snow formed from abandoned larvacean houses when larvaceans are dominant) (Alldredge & Gotshall 1990).

While aggregates provide luxurious feeding microhabitats, they may also constitute a risky environment, because fish, euphausids, and other macrophageous predators may feed on aggregates and their inhabitants. If the predation risk is higher for a zooplankter sitting on an aggregate than for one swimming in the ambient water, then the zooplankter should seek refuge away from the aggregate as soon as it has filled its gut. Filling the gut may in fact only take a few minutes when food is plentiful. For this strategy to be viable, however, the chance of finding a new aggregate when ready to fill the gut again should be high. To be more exact, the encounter frequency with aggregates should be higher than or of the same order as gut turnover time. In surface waters, abundances of >3 mm aggregates are typically 1 to 10 l$^{-1}$ (Alldredge & Silver 1988), and encounter frequencies estimated as encounter volume rate (Eq. 6) multiplied by aggregate concentration would therefore be >0.6 to 6 h$^{-1}$ (to be conservative). This is in fact very similar to gut turnover times for small copepods, e.g., 3 h$^{-1}$ in *Acartia tonsa* at 14°C (Kiørboe & Tiselius 1987). Thus, the short residence time may make sense. Since the optimal residence time is a trade-off between differential mortality rates, gut filling and turnover times, and aggregate encounter frequency, residence times are likely to vary between environments. At depth, there are typically fewer aggregates, yielding lower encounter frequencies. This may be (partly?) compensated by the lower temperatures at depth, which would increase gut turnover times ($Q_{10} = 3$, Kiørboe et al. 1982, Dam & Peterson 1988). Thus, the optimal residence time may not change much with depth.

If we accept that remote detection is required, we shall finally consider the nature of the cue that allows colonizers to locate sinking aggregates at distance. Our considerations suggest that hydromechanical signals are too weak to provide remote cues to colonizers. This leaves us with chemical cues. As evident from Fig. 3, the volume of water with elevated solute concentrations—or the plume transsectional area—decreases as a function of the sinking velocity of the aggregate. For the 3 Pe shown in Fig. 3 (0, 100, 10 000), for example, the relative transsectional areas of the 0.005 (× $Q/Da$) isoline (dark red area) are 1, 0.02, and 0.005, i.e., sinking washes the chemical signal away! On the other hand, sinking is a prerequisite for remote chemical detection, because the time scales associated with setting up the plumes are radically different when solute transport is solely by diffusion ($Pe = 0$) or facilitated by advection ($Pe > 0$). The diffusion time scale is $-L^2/D$, where $L$ is the diffusion distance. Thus, the time required for the chemical signal to reach 1 radius outside the particle by diffusion is $-10^5$ s for a 1 cm aggregate, and it increases with the square of the distance, e.g., to $-2.5 \times 10^8$ s = 4 wk = generation time...
for a copepod at 5 particle radii. The time scale for the chemical signal to be set up by a sinking aggregate depends mainly on the ratio of plume length to sinking velocity. Thus, the times required to establish the gradient out to 1 and 5 radii downstream in a sinking aggregate are about 5 and 25 s (cf. Eq. 7). Thus, chemical detection of large aggregates requires that these sink!

The estimated amino acid threshold concentrations required to account for the estimated encounter volume rates ($10^{-7}$ to $10^{-8}$ M) are not inconsistent with neurophysiological observations in copepods and other crustaceans (see also Yen et al. 1998). Some crustaceans have demonstrated neurophysiological and motor responses to concentrations of specific amino acids that are many orders of magnitude less than the threshold concentration estimated here. Also, precise tracking of trails of pheromones (Yen et al. 1998) and amino acids (authors’ unpubl. obs.) have been demonstrated in copepods, and the general mechanism suggested here, that cruising copepods encounter a plume of elevated solute concentration left by a sinking aggregate, has actually been documented to work for copepods seeking mates (Tsuda & Miller 1998). Tsuda & Miller (1998) found that males could detect sinking females at a distance of 50 cm or more downstream, equivalent to ca 130 bodylengths. This is consistent with the plume lengths suggested by the present analysis. However, 2 problems need consideration, namely the resistance (or possible lack thereof) of such long plumes to ambient turbulence, and the apparently low signal-to-noise ratio of the chemical signal.

The time scale to set up the chemical gradient in the 1 m long plume of a 1 cm aggregate is about 10 min. One could imagine that strong turbulence would lead to the rapid dissipation of such long plumes. Direct observations in the ocean of plumes left by small sinking particles leaking fluorescent dye suggest that plumes may in fact be that long (or longer) and persistent during calm conditions, in the thermoline, and at depth (Woods 1968, 1971). Moderate shear may cause the plume to meander, and moderate turbulence would cause the plume to become filamentous. Because of the coherent nature of turbulence, this would not necessarily reduce plume concentrations much, but rather cause the chemical cue to be more variable in both time and space than Fig. 3 indicates. Crabs may still be able to locate prey from the chemical signals carried by a meandering and filamentous plume (Zimmer-Faust et al. 1995). The observations of colonizer abundances on aggregates used to generate Eq. (6) were collected under calm enough conditions to allow diving. It is conceivable, however, that strong turbulence in the surface layer would dissipate the plume and prevent remote chemical detection. If zooplankters do use chemical cues to localize sinking aggregates, one would therefore expect lower colonization rates in the upper ocean during turbulent conditions.

The threshold amino acid concentrations estimated here are of the same order as background concentrations of bulk amino acids, at least in the upper ocean of rich environments. High background concentrations may mask the chemical signals from a sinking aggregate. Bulk amino acid concentrations may, however, not be the only parameter of interest, but the quality of the chemical signal may be relevant as well (Ache 1988). That is, the composition or the occurrence of specific amino acids may allow discrimination from background, even though bulk amino acid concentrations are not very different. This is likely the way the extremely low response threshold concentrations of specific (combinations of) amino acids—orders of magnitude less than bulk background concentrations—that have been recorded in several crustaceans should be understood. Crabs and lobsters, for example, appear to be particularly sensitive to taurine, an amino acid characteristic of animal tissue (response threshold <10$^{-20}$ M, Thomson & Ache 1980). The amino acid signature of an aggregate is likely to be different from the background signature. Important components of marine snow aggregates are phytoplankton cells, fecal pellets and detritus. While the amino acid composition of copepod fecal pellet largely reflects the composition of the food, both may be radically different from the background (Poulet et al. 1986).

This study has demonstrated that chemosensory localization of sinking aggregates by colonizing copepods is, in principle, possible. However, observations are required to test the hypothesis. The biology of the kinds of copepods, such as Oncaea spp., that colonize snow aggregates is very poorly known. The Oncaea genus is widespread and abundant and sometimes even numerically dominates the mesozooplankton, particularly at depth and below the euphotic zone (e.g., Pancucci-Papadopoulou et al. 1992, Böttger-Schnack 1997). They may play a major role in the degradation of marine snow and, hence, in regulating vertical material fluxes in the ocean. Yet, biological and behavioral observations are very few (Paffenhold 1993, Go et al. 1998, Metz 1998). This study suggests that observations of sensitivities to hydromechanical and chemical cues, gut filling and gut throughput times, and swimming behavior are required to further explore the association between these copepods and marine snow aggregates and their significance in regulating vertical material transport in the ocean.

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