Sinking rates of organic particles

Abstract—The flux of organic particles collected by sediment traps decreases in a non-linear fashion with depth. This implies a loss of organic carbon with time as the particles sink. Samples of such particles held in the laboratory or left in situ do decay, showing a significant loss of carbon.

Particles were collected in the upper 1,000 m of the water column with sediment traps south of the Hawaiian Islands. Aliquots from the traps were placed in combusted glass bottles and returned to the depths from which they were collected. After 5 days the bottles were recovered and the carbon content determined. On the average, about 30% of the particulate carbon was lost during the experiment.

With this knowledge of the decay rate, one can estimate the expected change in "apparent carbon flux" with time. This in combination with the measured vertical flux can be used to calculate the depth interval required for this change to occur. Average sinking rates of 92 m d⁻¹ were thus estimated.

The flux of organic matter, measured as organic carbon, shows a nonlinear decrease with increasing depth (Suess 1980). The processes involved in this apparent loss of organic carbon have not been documented, but we will assume that the loss is related to decomposition rather than to solubilization or fragmentation because a summary of the published data (Menzel 1974) shows no marked increase in dissolved or particulate organic carbon with depth. Most likely this loss of carbon is the result of biological activity, either microbial or possibly via reingestion and repackaging by midwater animals.

We here report the results of an exper-
iment conducted in the oligotrophic waters south of Oahu in which we investigated the sinking rates of organic particles (RV Thomas G. Thompson cruise TT-157).

Floating sediment trap arrays were deployed in two parts consisting of VLS Sampson braided rope (9.5 mm) and stainless steel wire (3.9 mm) (Fig. 1). The traps on the braided rope portion were PVC cylinders (15.2-cm diam × 45.7 cm high: Lorenzen et al. 1981) placed at 250, 400, 600, and 900 m. The other traps, half as large, were placed on the wire at 20-m intervals starting at 20 m and going to a depth of 200 m. These traps were closed by a messenger at a predetermined time before the arrays were recovered. Previous comparisons showed that both traps captured and retained material with equal efficiency (Welschmeyer 1982). The arrays were deployed twice: the first time for 4 days and the second time for 5 days. After the first deployment, each trap was sampled for particulate organic carbon, POC, and plant pigments. Samples were filtered either on combusted silver filters (0.45-μm pore size) for POC determination by elemental analysis or Whatman GF/F glass-fiber filters for pigment analysis (Lorenzen 1966). For the decay experiments, aliquots from the larger traps were placed in combusted glass bottles (1 liter) which were attached to the array when it was redeployed so that these samples were returned to the depths from which they were captured. After the second recovery, POC was measured in each bottle.

Primary production was measured in situ by the 14C technique (Strickland and Parsons 1972). Exposure was for 24 h, sunrise to sunrise. Plant pigments were measured repeatedly in the water of the euphotic zone by a fluorometric technique; samples for those measurements were collected with 2.5-liter Niskin bottles.

Flux rates were calculated from the sediment trap samples for both POC (Fig. 2) and plant pigments (Fig. 3). Flux rates were maximum in the upper 250 m of the
water column. Below the euphotic zone, at the 250-m level, flux rates of both POC and pigments are markedly reduced; actual values are 45% (POC), 10% (pheopigments), and 0.1% (chlorophyll) of those at the maximum nearer the surface.

The change in POC $[C_p]/[C_o]$ over the course of the in situ "rotting" experiment was about 30% (Table 1).

The loss of POC during the in situ rotting experiments, the change measured in the glass bottles suspended on the second deployment, shows that some portion of the organic matter sinking out of the surface layers is labile. The pigment-laden portion of this material appears to be fecal pellets from herbivores, since the ratio of pheopigments to chlorophyll was 9.2 which is characteristic of fecal pellets of herbivores feeding on algae (Lorenzen unpubl.). On the other hand, the carbon : pheopigment ratio is entirely too high for herbivore fecal pellets (>400), suggesting that a relatively minor portion of the

[Table 1. Carbon flux, change in carbon concentration during the rotting experiments, $[C_p]/[C_o]$, apparent carbon flux, depth interval necessary to account for change in carbon flux, and calculated sinking rates derived from rotting experiments.]

| Z (m) | $C_{max}$ | $[C_p]/[C_o]$ | Apparent C Flux (mg m$^{-2}$ d$^{-1}$) | $\Delta Z$ (m d$^{-1}$) |
|-------|-----------|---------------|----------------------------------;|-----------------|
| 250   | 76.2      | 0.73          | 55.9                            | 393             | 78   |
| 400   | 63.0      | 0.66          | 41.6                            | 656             | 131  |
| 900   | 51.6      | 0.71          | 36.6                            | 335             | 67   |
| $\bar{Z}$ | 0.70     | -             | -                               | -               | 92   |

Below 100 m, pheopigment flux decreases markedly. Past experience in this laboratory shows that reingestion of pheopigments (in fecal pellets) by crustaceans brings about alterations in the pigments so that they are lost to our analytical technique. Coprophagy is probably of major significance in the upper layers of the water column. On the other hand, as pointed out below, coprophagy may only be a minor process in the deeper portions of the water column.

Chlorophyll flux is substantially lower than pheopigment flux below 50 m. At 100 and 400 m the daily flux of chlorophyll represents only 0.8 and 0.08% of the integrated standing crop of chlorophyll above these depths, suggesting that sinking of intact algae out of the euphotic

[Table 2. Tabulation of carbon flux, apparent carbon utilization (ACU), and percentage change in carbon per meter with depth.]

<table>
<thead>
<tr>
<th>Z (m)</th>
<th>$C_{max}$</th>
<th>ACU (mg m$^{-2}$ d$^{-1}$)</th>
<th>$\Delta$C% m$^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>250</td>
<td>76.2</td>
<td>0.088</td>
<td>0.11</td>
</tr>
<tr>
<td>400</td>
<td>63.0</td>
<td>0.051</td>
<td>0.081</td>
</tr>
<tr>
<td>600</td>
<td>52.8</td>
<td>0.004</td>
<td>0.0075</td>
</tr>
<tr>
<td>900</td>
<td>51.6</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
zone is of minor importance at this location.

Knowledge of the vertical distribution of organic carbon flux does not yield the information needed to estimate a sinking rate, but it does permit estimation of carbon utilization in the water column. In this case, the vertical distribution of organic carbon flux between 250 and 900 m (Fig. 2) is described by the statistically significant relationship \((n = 4, r^2 = 0.82):\)

\[
Z = 6,276 - 1,400 \times \ln(C_{\text{flux}}) \tag{1}
\]

(where \(C_{\text{flux}}\) is carbon flux in mg C·m\(^{-2}\)·d\(^{-1}\), and \(Z\) is depth in meters).

The apparent carbon utilization over this depth range, \(\text{ACU}\), calculated by

\[
\text{ACU} = \frac{\text{flux}_1 - \text{flux}_2}{Z_2 - Z_1} \tag{2}
\]

is tabulated for the sediment traps at 250 m and below (Table 2). The ACU decreases with depth both in absolute terms and as a percentage of the flux. In the first interval it is 0.088 and in the last interval 0.004 mg C·m\(^{-3}\)·d\(^{-1}\). This is also reflected in the percentage loss. For the same intervals the values are 0.59 and 0.26 ΔAC%·m\(^{-1}\), perhaps indicative of the material becoming more refractory deeper in the water column.

Deriving an estimate of the sinking rates of these particles is not straightforward. The results from the rotting experiments (Table 1) show that the collected material is indeed labile. The actual kinetics of the reactions involved are unknown, but a first-order chemical reaction may be assumed. If we use the measured change to reduce the flux at the level from which the material was collected and calculate the "apparent carbon flux" 5 days later, we can use Eq. 1 to calculate the depth at which this flux would be found. For example, in the case of the 400-m sample: 63.0(\(C_{\text{flux}}\)) × 0.66(\(C_a/\ C_0\)) = 41.6 (apparent carbon flux). The depth where a carbon flux of 41.6 mg·m\(^{-2}\)·d\(^{-1}\) would be found is 1,056 m. The interval 1,056 to 400 m, or 656 m covered in 5 days, would suggest an average sinking rate of 131 m·d\(^{-1}\). The calculations yield sinking rates of 78, 131, and 67 m·d\(^{-1}\) for material collected at 250, 400, and 900 m. The average sinking rate is calculated as 92 m·d\(^{-1}\).

The fate of these organic particles as they sink is unclear and the mechanisms of their destruction may vary with depth. We suggest that in and just below the euphotic zone the major loss results from reingestion, i.e. coprophagy. At greater depths this seems less important: between 400 and 900 m, carbon flux decreases 18%, while pheopigment flux is unchanged. This suggests that coprophagy is relatively insignificant at these depths.

Carl J. Lorenzen
Nicholas A. Welschmeyer
Andrea E. Copping
Maria Vernet

School of Oceanography WB-10
University of Washington
Seattle 98195

References


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2 Present address: The Biological Laboratories, Harvard University, Cambridge, Massachusetts 02138.