Decomposition and release of elements from zooplankton debris

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ABSTRACT: In experiments examining the retention of trace elements in decomposing zooplankton debris, the rates of C degradation and metal release from radiolabeled copepod fecal pellets and carcasses were measured for up to 1 mo using radiotracers. Fecal pellets incubated at 18 and 2 °C retained 35 to 40% and 80% of their 14C, respectively, after 30 d. Carcasses retained only 13 to 18% and 28 to 38% of their 14C after 18 d at 18 and 2 °C, respectively. Leaching of 14C from fecal pellets and carcasses as DO14C accounted for about half of the 14C loss, even in the absence of microbial activity. Proportionately more of the 14C from carcasses was microbiologically oxidized to 14CO2 than 14C from fecal pellets. Release of the particle-reactive transuranic element 241Am from fecal pellets was unaffected by microbial activity while release of 75Se and 65Zn from both fecal pellets and carcasses increased with microbial activity and closely followed 14C loss. Release rates of all elements decreased exponentially over time, with the most pronounced decreases occurring within the first 6 d. Retention half-times (t1/2's) of 241Am were ≥ 48 d in fecal pellets. By contrast, the t1/2's for 65Zn ranged from 2.1 to 13.5 d in fecal pellets and both 65Zn and 75Se had t1/2's of only about 1 d in carcasses. The results help explain oceanographic observations that C, Se and Zn are recycled in surface waters while scavenged elements like Am are enriched in fecal pellets and have short residence times in surface waters.

INTRODUCTION

Numerous studies have shown that large biogenic particles are important vectors for the vertical transport of organic compounds, radionuclides and metals from the surface ocean to the deep ocean (McCave 1975, Knauer et al. 1979, Wakeham et al. 1984, Fowler & Knauer 1986, Fisher et al. 1988). These particles include phytoplankton aggregates, zooplankton fecal pellets, eggs, molts, animal carcasses, larvacean houses, and marine snow. Among these particles, fecal pellets in particular have received considerable scrutiny due to their ubiquity, abundance and rapid sinking rates. The geochemical cycling of many elements is strongly influenced by sinking particles during their descent in the water column, either through oxidation or dissolution of the particles or by scavenging of particle-reactive elements from ambient water.

The flux of particles measured by sediment traps indicates that particles are rapidly regenerated beneath the euphotic zone (Knauer et al. 1979, Hargrave 1985, Martin et al. 1987). Although information on the degradation of these organic particles is essential to understanding the geochemical cycling of nutrients and metals in the ocean, the fate of settling biogenic particles is not well understood and explanations of processes responsible for degradation are controversial (Cho & Azam 1988, Karl et al. 1988, Jumars et al. 1989, Allredge et al. 1990, Banse 1990, Lampitt et al. 1990). Several processes are involved in the degradation of particles. One of these, leaching (i.e. loss of soluble material) of organic compounds and elements from biogenic particles (Gardner et al. 1983, Collier & Edmond 1984, Knauer et al. 1984), is potentially very important but remains poorly understood.

While the release of some elements from biogenic particles may be influenced by degradation processes, no attempt has been made to combine the elemental release rates with particle degradation rates, with the exception of phytoplankton (Lee & Fisher 1992, Fisher &
pH of the water was 8.0 after the isotope additions. The initial concentration gradients. Periodically, the radio-
samples were shaken gently daily to minimize poten-
tial concentrations (37 kBq 14C, 37 kBq 75Se, 37 kBq 65Zn, or 14C. During the incubation,
the dark for up to 1 mo in GF/C-filtered seawater, or
fed 14C-labeled food. The fecal pellets and carcasses were incubated in
given initial cell density of 5.0 × 10^4 ml^-1. One flask of
and Cu, Zn and EDTA. Two of the flasks were inocu-
lized with T. pseudonana, and a third with I. galbana to
ment 8.4). The isotope additions resulted in total metal
additions of 290 ng Am (1.20 nM), 29 ng Se (0.38 nM)
and 4.0 ng Zn (64 pM).

The cultures were incubated on a 14 h light:10 h
dark cycle at 18 ± 1 °C for 4 d. Periodically, the quanti-
ty of each isotope associated with cells was determined by
filtering a 10 ml aliquot from each culture through a 0.2 µm Nuclepore filter, following estab-
lished procedures (Fisher et al. 1983a), and a 1 ml aliquot was taken for cell counting by hemacytometer.
The cells underwent 5 to 6 divisions, yielding final cell
densities of 1.7 to 2.0 × 10^5 cells ml^-1. After the 4 d
incubation period, each culture was resuspended via
Nuclepore filtration (Thalassiosira pseudonana) or cen-
trifugation (Isochrysis galbana) into 3 different feeding
vessels containing 2 l of 0.2 µm-filtered seawater to
give cell densities of 1 × 10^5 ml^-1. This corresponds to
6.1 × 10^8 µm^-3 l^-1 or 0.8 mg C l^-1 of T. pseudonana, and
5.8 × 10^9 µm^-3 or 1.1 mg C l^-1 of I. galbana.

About 200 copepods (Acartia tonsa) were transferred to
each feeding vessel containing either Thalassiosira
pseudonana labeled with 14C, T. pseudonana with
241Am, 75Se, 65Zn, or Isochrysis galbana with 14C.
Copepods were separated by a 210 µm Nitex mesh
above the bottom of the vessel so that any fecal pellets
produced were separated from the copepods above.
The fecal pellets produced in the first 2 h of feeding on
radiolabeled food were discarded to eliminate fecal
pellets composed of unlabeled food. During the feed-
ing, fecal pellets were collected from the feeding con-
tainer within 2 h of production and rinsed several times
with 0.2 µm filtered seawater. One hundred fecal pel-
lets were transferred via wide-bore pipet into 50 ml
snap-top plastic vials. After allowing the copepods to
empty undigested radioactive food by feeding on un-
labeled food (same species and cell density) for 12 h,
c copepods fed radiolabeled food for 5 d were killed by
drying in air on a Nitex mesh for about 5 min. Twenty
dead copepods from each food source were transferred to
each of 3 other vials. The vials, containing fecal
pellets or carcasses, were measured for initial total
radioactivity (see below) and were then filled with
GF/C filtered seawater. Similarly, radiolabeled fecal
pellets and carcasses were collected by feeding the
copepod Temora longicornis with 241Am, 75Se, 65Zn
labeled T pseudonana cells. T longicornis were not
fed 14C-labeled food.
The fecal pellets and carcasses were incubated in
the dark for up to 1 mo in GF/C-filtered seawater, or
GF/C-filtered seawater poisoned with 0.9 mM HgCl2,
at either 18 ± 1 °C or 2 ± 1 °C. During the incubation,
samples were shaken gently daily to minimize potential
concentration gradients. Periodically, the radio-

MATERIALS AND METHODS

The calanoid copepods Acartia tonsa and Temora
longicornis were collected from coastal waters off Long
Island, New York, USA, using a 63 µm mesh. Adults
were separated and maintained in the laboratory in
35 % glass-fiber filtered (GF/C) seawater at 18 °C and fed the diatom Thalassiosira pseudonana (clone 3H) and
the haptophyte Isochrysis galbana (clone iso) prior to experiments.

Radiolabeled copepod fecal pellets and carcasses were collected after feeding on radiolabeled phyto-
plankton which had been prepared as follows. The 2
algal species were maintained in unialgal, clonal culture in sterile-filtered (0.2 µm Nuclepore filter), 35 % surface seawater (collected 8 km off Southampton, New York)
enriched with f/2 nutrients (Guillard & Ryther 1962). Ex-
perimental inocula, taken from each algal stock culture in late log phase, were resuspended off of 1 µm Nucle-
 pore filters (Thalassiosira pseudonana) and centrifuged out of their media (Isochrysis galbana) and added to
three 11 Erlenmeyer flasks. Each flask contained 400 ml
sterile-filtered seawater, enriched with f/2 nutrients minus Cu, Zn and EDTA. Two of the flasks were inocu-
lized with T. pseudonana, and a third with I. galbana to
give an initial cell density of 5.0 × 10^4 ml^-1. One flask of
T. pseudonana received 37 kBq 241Am (t1/2 = 433 yr, from Amersham; 37 kBq = 1 µCi) in 251 µl of 3N HNO3,
37 kBq 75Se (t1/2 = 120 d, from Amersham) in 39 µl of
0.5N HCl, and 148 kBq 65Zn (t1/2 = 244 d, from Amers-
ham) in 127 µl of 0.1N HCl. Since the isotopes were in
dilute acids (Ultrex), immediately prior to addition of the
isotopes, the flask received appropriate amounts (µl quantities) of dilute Suprapur NaOH to ensure that the
pH of the water was 8.0 after the isotope additions. The
remaining flasks of T. pseudonana and I. galbana each
received 370 kBq 14C (as NaH14CO3 in distilled water,
ph 8.4). The isotope additions resulted in total metal
additions of 290 ng Am (1.20 nM), 29 ng Se (0.38 nM)
and 4.0 ng Zn (64 pM).

Wente 1993). Such information is essential for quantifying and modeling the impact of sinking biogenic parti-
cles on elemental transport and cycling in the ocean.

To address this deficiency, we have conducted labora-
tory radiotracer experiments to provide a qualitative
assessment of the degradation and elemental release rates from zooplankton carcasses and fecal
pellets. We have assessed the fate of C in this debris
and also compared 3 elements – Zn, an essential tran-
sition metal, Se, an essential metalloid, and Am, a non-
essential actinide. All of these elements are concen-
trated out of seawater by phytoplankton and can be transferred to herbivorous zooplankton (Fisher et al.
1991a, Reinfielder & Fisher 1991). Their retention in
zooplankton debris was compared with that of C. Further, the influence of poisons on elemental release
rates was determined to distinguish between biologi-
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ally mediated and physical/chemical processes.
activity of gamma-emitting radioisotopes ($^{241}$Am, $^{75}$Se, $^{65}$Zn) retained by the particles was measured by counting each vial (typically a 5 min count was made) after the upper 45 ml of seawater was gently removed by pipetting. Appropriate corrections were made for the radioactivity associated with the 5 ml of water remaining in the vial. Immediately after the radioactivity was determined, a fresh batch of 45 ml GF/C-filtered seawater for each temperature (pre-equilibrated at the incubation temperatures and containing the appropriate amount of HgCl$_2$, if necessary) was added to each vial and the vials were returned to their incubators.

$^{14}$C activity was partitioned into 3 fractions as in Lee & Fisher (1992): particulate (>0.2 μm), dissolved organic $^{14}$C (<0.2 μm), and remineralized $^{14}$C ($^{14}$CO$_2$). Briefly, at each sample time, 1 vial from each treatment was removed, gently shaken, and filtered through a 0.2 μm Nuclepore filter. The particles retained on the filter were transferred to a scintillation vial containing 10 ml of Aquasol scintillation cocktail. $^{14}$C activity associated with the filtered particles was considered particulate. To separate DO$^{14}$C from $^{14}$CO$_2$, the filtrate was transferred into a 125 ml flask and bubbled vigorously with N$_2$ gas for 15 min after acidification with concentrated HCl (pH <2). Liberated $^{14}$CO$_2$ was collected in series in 2 scintillation vials each containing 10 ml of CO$_2$ absorber (Carbamate 1). $^{14}$C activity in the second vial was not significantly different from background, indicating that almost all of the liberated $^{14}$CO$_2$ was trapped in the first vial. The radioactivity remaining in the solution after acidification and purging of $^{14}$CO$_2$ was assumed to be activity associated with DO$^{14}$C (<0.2 μm). The DO$^{14}$C was measured in 10 ml of filtrate transferred into a scintillation vial containing 10 ml of Aquasol.

HgCl$_2$ was used to suppress microbial activity in these experiments. However, some poisons can accelerate elemental release from diatom cells (Lee & Fisher 1992), possibly as a result of displacement or complexation of metals bound to debris. To further examine this question, the effects of 2 poisons, Na$_2$O$_4$ and HgCl$_2$, on the release of $^{241}$Am, $^{75}$Se and $^{65}$Zn by glass beads and egg albumin particles were tested; glass beads served as representative of end member mineral surfaces, while chicken egg albumin particles served as a model for protein (Fisher & Teyssie 1986). A boiled egg white was dried for 48 h at 65 °C, ground with a mortar and pestle, and sieved to produce particles of about 5 to 10 μm diameter. Acid-washed glass beads (5 to 10 μm diameter) and egg albumin particles were labeled with $^{241}$Am, $^{75}$Se and $^{65}$Zn by exposing particles to radioisotopes for 2 d; the suspended particle load was 300 mg l$^{-1}$ for both particle types. The experimental vessels, water, and radioisotope handling and concentrations were identical to those used for labeling the diatoms. The radiolabeled particles were resuspended into 1.0 μm filtered seawater to give the following treatments: sterilized (by autoclaving) seawater incubated at 18 and at 2 °C; HgCl$_2$ (0.9 mM) treated seawater at 18 and at 2 °C; Na$_2$O$_4$ (154 mM) treated seawater at 18 °C; and unsterilized seawater without poisons (microbial treatment) at 18 °C. Retention of radioactivity by the particles was monitored for 4 h for the glass beads and for 10 d for the albumin particles.

The radioactivity of $^{241}$Am, $^{75}$Se and $^{65}$Zn was determined in a Pharmacia-Wallac LKB CompuGamma with a well-type NaI(Tl) detector and spectra software to correct for spillover of each isotope into another's counting 'window.' The gamma emissions of $^{241}$Am were determined at 60 keV, $^{75}$Se at 264 keV and $^{65}$Zn at 1115 keV. A calibration of the detector was made for its energy and counting efficiency with appropriate standards for each of the geometries used. The beta emissions of $^{14}$C-containing samples were measured using an external standards ratio to correct for quenching. Counting times for all samples were 5 to 10 min and propagated counting errors were generally <5 %.

### RESULTS

Partitioning of $^{14}$C activity from decomposing copepod carcasses and fecal pellets over the 30 d incubation period is shown in Figs. 1 & 2. Retention of $^{14}$C, $^{241}$Am, $^{75}$Se and $^{65}$Zn by zooplankton debris over the incubation period is shown in Figs. 3 to 5.

Retention of $^{14}$C, $^{241}$Am, $^{75}$Se and $^{65}$Zn by the unpoisoned samples generally decreased according to the equation:

$$y = 100(t + 1)^{-b}$$  \[(1)\]

where $y$ = percent element remaining in the particles; $t$ = time (d); and $b$ = release rate coefficient. Values for $b$, specific for each element and treatment, were estimated from a linear regression analysis of log-transformed data (log percent element retained by particles vs log time); all regressions were significant ($p < 0.001$). Consequently, the instantaneous release rate of each element ($R$) decreased exponentially over time:

$$\frac{dy}{dt} (\% d^{-1}) = -100bt(t + 1)^{(b - 1)}$$  \[(2)\]

where negative values represent decrease in percent element retained by particles. Values and 95 % confidence intervals for $b$, $R^2$ values for the regression lines, and retention half-times ($t_{1/2}$'s) of each element estimated from Eq. 1 are given in Table 1. Table 2 shows rate constants ($R$) estimated from Eq. 2 for release of $^{14}$C, $^{241}$Am, $^{75}$Se and $^{65}$Zn from copepod fecal pellets and carcasses.
Fig. 1. *Acartia tonsa* fed *Isochrysis galbana* and *Thalassiosira pseudonana*. Partitioning of total $^{14}$C activity among (■) particulate, (□) $^{14}$CO$_2$ and (△) DOC$^{14}$C fractions during 18 d incubations of copepod carcasses at (a) 18 °C, (b) 2 °C, and (c) 18 °C with HgCl$_2$.

Fig. 2. *Acartia tonsa*. Partitioning of total $^{14}$C activity among (■) particulate, (□) $^{14}$CO$_2$ and (△) DOC$^{14}$C fractions during 30 d incubations of *T. pseudonana*-fed copepod fecal pellets at (a) 18 °C, (b) 2 °C, and (c) 18 °C with HgCl$_2$, and (d) *Isochrysis galbana*-fed copepod fecal pellets at 18 °C.
Results show that degradation of copepod fecal pellets and carcasses was faster at 18 than at 2 °C and that the release of fecal pellet $^{14}$C was slower than that of carcasses (Fig. 3, Table 1). Altering the diet of the copepods (Thalassiosira pseudonana vs Isochrysis galbana) had a negligible effect on the degradation rates of fecal pellets and carcasses. After 30 d of incubation with microorganisms, about 59 to 65 % of the total fecal pellet $^{14}$C was lost (as $^{14}$CO$_2$ and $^{14}$CO$_2$) at 18 °C (20 % at 2 °C); about 82 to 87 % of total copepod carcass $^{14}$C was lost after 18 d at 18 °C (62 to 72 % at 2 °C) (Figs. 1 & 2). The greater microbial remineralization to $^{14}$CO$_2$ at 18 °C largely accounted for the higher degradation rates at 18 °C. No attempts were made, however, to enumerate bacteria in these samples.

The greatest changes in the $^{14}$C release rates ($R$), both over time and among treatments, occurred during the first 6 d of incubation. The instantaneous $^{14}$C loss rate for Days 1 to 4 for carcasses and fecal pellets at 18 °C decreased from 22.1 to 4.5 % d$^{-1}$ (14.3 to 4.0 % d$^{-1}$ at 2 °C) and from 10.4 to 3.3 % d$^{-1}$ (3.6 to 1.3 % d$^{-1}$ at 2 °C), respectively (Table 2). $^{14}$C loss rates for Days 6 to 30 decreased from ca 2 to ca 0.2 % d$^{-1}$, regardless of incubation temperature and particle type.

Microbial activity was effectively suppressed by 0.9 mM HgCl$_2$, as indicated by the lack of released $^{14}$CO$_2$ (Figs. 1c & 2c). Even without microbial activity, however, the carcasses and fecal pellets incubated with 0.9 mM HgCl$_2$ released up to half the $^{14}$C lost by particles incubated with microorganisms. This leach-
Table 1. Calculated retention half-times ($t_{1/2}$), release rate coefficients ($b$), 95% confidence limits of $b$, and $r^2$ values of regression lines [$y = 100(t + 1)^b$] describing elemental release from particles. CC: copepod carcass, FP: fecal pellet; A: Acartia tonsa; T: Temora longicornis; Ig: fed with Isochrysis galbana; Tp: fed with Thalassiosira pseudonana.

<table>
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<th>Isotope</th>
<th>Particle type</th>
<th>°C</th>
<th>$t_{1/2}$ (d)</th>
<th>$b$</th>
<th>95% confidence limits of $b$</th>
<th>$r^2$</th>
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<td>$^{14}$C</td>
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<td>18</td>
<td>1.6</td>
<td>0.733</td>
<td>0.700, 0.767</td>
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Table 2. Elemental release rates (% d$^{-1}$) estimated from the equation: $dy/dt (R) = -100b(t + 1)^b$ for periods up to 30 d. CC: copepod carcass; FP: fecal pellet; A: Acartia tonsa; T: Temora longicornis; Ig: fed with Isochrysis galbana; Tp: fed with Thalassiosira pseudonana.

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<th>3</th>
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ing of 14C from Hg-treated fecal pellets as DO14C was gradual and proceeded at a rate of 1.25 % d−1 over the 30 d incubation period. In contrast, in Hg-treated samples about 35 % of total carcass 14C leached out within the first 24 h, after which there was little change. In unpoisoned samples, most of the DO14C released by leaching from carcasses was remineralized to 14CO2 (Fig. 1), whereas only a small fraction of the released DO14C from fecal pellets was remineralized to 14CO2 (Fig. 2b). Thus, by 30 d, the fraction of unutilized DO14C increased to about 40 % in the unpoisoned fecal pellet samples.

The C degradation data of Acartia tonsa fecal pellets and carcasses were combined with metal and metalloid data by plotting C vs trace elements retained by particles for each sample throughout the incubation periods (Fig. 6). Least squares fits of the data for metals and C were calculated from unpoisoned samples and are also presented in Fig. 6. A linear model was applied to the 65Zn and 75Se data and for all other data a power function was used (Fig. 6). These lines provide comparisons of metal retention normalized to C in a way that is independent of temperature or microbial activity. Generally, 241Am was retained for longer periods than C by fecal pellets, whereas 65Zn was lost at approximately the same rates as C, and 75Se was lost more rapidly than C (Fig. 6).

Neither temperature nor HgCl2 treatment had a large effect on the 241Am release rate. Because 241Am showed no assimilation in the copepods and 65Se was almost completely assimilated by the copepods, the radioactivity of 241Am in the carcasses and of 75Se in the fecal pellets was very low, precluding long-term monitoring.

The release rates of both 75Se and 65Zn from copepod carcasses were generally faster at 18 than at 2 °C (Fig. 4). There was no significant difference between release rate coefficients (b) for 75Se and 65Zn from carcasses of Acartia tonsa and Temora longicornis. Release of 75Se from A. tonsa carcasses was initially faster than the release of C but eventually slowed to less than that of C (Fig. 6). Release of 65Zn from the same carcasses was slightly faster than 14C release, with a slope of 1.2 and a y intercept of −17 %, indicating that the 65Zn in the carcasses retained 65Zn less effectively than did fecal pellets, where the 65Zn release rate was slower than the 14C loss rate.

The changes in 241Am, 75Se and 65Zn release rates from both types of debris were similar to those changes seen with 14C in that the most pronounced changes occurred within the first 6 d of incubation, with no appreciable differences in rates thereafter (Table 2). Instantaneous loss rates from Days 1 to 4 decreased from 7.9 to 2.1 % d−1 for 241Am in fecal pellets, from 26.3 to 3.3 % d−1 for 75Se in carcasses, from 25.5 to 1.4 % d−1 for 65Zn in carcasses and from 20.1 to 3.4 % d−1 in fecal pellets (Table 2). The calculated t1/2's ranged from 48 to 215 d for 241Am in fecal pellets (although differences in retention curves for 241Am were relatively small and inconsistent in response to temperature; Fig. 5). These t1/2's were 1 to 2 orders of magnitude greater than for 75Se and 65Zn, which ranged from 0.7 to 1.6 d for 75Se in carcasses, and from 2.1 to 4.2 d for 65Zn in fecal pellets at 18 °C (3.3 to 13.5 d at 2 °C) and 0.9 to 1.6 d for 65Zn in carcasses (Table 1).

The retention curves of 241Am, 75Se and 65Zn by glass beads and egg albumin particles are shown in Fig. 7. About 20 to 30 % more 75Se and 65Zn were lost from egg albumin particles treated with HgCl2 than with any other treatment, including NaN3. The release rates of 241Am from both particle types and of 65Zn from glass beads showed no appreciable differences among all treatments. Desorption of 75Se from glass beads was not determined since essentially no 75Se sorbed to the glass beads during the 2 d exposure period.
DISCUSSION

Carbon release

The rate of loss of $^{14}$C from unpoisoned zooplankton debris was both time and temperature dependent, decreasing over time according to a power function. This inverse relationship of loss rate of organic C and incubation time is consistent with previous findings for phytoplankton detritus (Otsuki & Hanya 1972, Cole et al. 1984, Biddanda 1988, Lee & Fisher 1992) and fecal pellets (Pomeroy et al. 1984, González & Biddanda 1990).

Rapid leaching of organic $^{14}$C was largely responsible for the initial higher $^{14}$C loss rate observed in this study. It can be inferred that microbial activity enhanced this rate, especially at 18 °C, since greater quantities of particulate $^{14}$C were lost from unpoisoned particles than from poisoned samples during the first few days of incubation. Other studies have also shown that microbial biomass and activity on decomposing biogenic debris are greatest at the beginning of particle decomposition and decline over time (Andrews et al. 1984, Pomeroy et al. 1984, Peduzzi & Herndl 1986, González & Biddanda 1990). These results indicate that a C loss rate calculated from 2 end point determinations, assuming linear decay rate, is likely to underestimate the rate for initial decomposition periods and to overestimate it for later periods of decomposition. Similarly, C flux rates assessed with unpoisoned sediment traps, assuming linear decay rate, may be overestimates for long deployments and underestimates for short deployments.

The $^{14}$C loss rates from fecal pellets measured here were comparable to those (6 % d$^{-1}$ at 18 °C, 1.1 % d$^{-1}$ at 5 °C) reported for fecal pellets collected from natural assemblages (Roy & Poulet 1990), but higher than those (1 to 1.8 % d$^{-1}$) reported for fecal pellets of the copepod *Calanus pacificus* (Jacobson & Azam 1984). The decay rates in this study also are comparable to the 6 % d$^{-1}$ measured *in situ* for bulk material (mostly herbivore fecal pellets) collected in a sediment trap in the upper 1000 m (Lorenzen et al. 1983), and the 0.1 to 1 % d$^{-1}$ estimated for a variety of biogenic particles in unpoisoned sediment traps deployed at depths >2700 m (Gardner et al. 1983). The rates of C loss observed here suggest that degradation of particles sinking in the water column will decrease substantially from surface to deep waters, partially in response to temperature differences and partially to the increasingly refractory nature of the remaining particulate matter after initial decomposition. Flux measurements determined with sediment traps indicate that particles are, in fact, rapidly regenerated in the upper 1000 m and that C flux decreases exponentially with depth (Knauer et al. 1979, Hargrave 1985, Martin et al. 1987).

Other studies have noted the significance of protozoans and metazoans in mediating the decomposition of zooplankton fecal pellets (Jacobson & Azam 1984, Pomeroy et al. 1984, González & Biddanda 1990, Lampitt et al. 1990). Lampitt et al. (1990), for example, demonstrated that copepods can enhance the degradation rates of fecal pellets by breaking up their own pellets, and Jacobson & Azam (1984) reported that microzooplankton doubled the rate of decomposition of *Calanus pacificus* fecal pellets held at 18 °C. Therefore, the estimates of C loss reported here should be considered conservative, since the effects of animal activities were not determined in our experiments. Moreover, the leaching rates in the experiments reported here may be less than those from sinking particles because boundary layer effects on diffusion would be greater in the laboratory, although the model of Jumars et al. (1989) suggests that these differences would be relatively small.
The greater C degradation of both types of particles at 18 than at 2 °C may be attributable to greater microbial activity at 18 °C, since the initial leaching of soluble organic C is reported to be relatively independent of temperature (Cole et al. 1984). Similarly, other studies focusing on the degradation of zooplankton (Turner 1979, Roy & Poulet 1990) and phytoplankton debris (Iturriaga 1979, Cole et al. 1984, Lee & Fisher 1992) reported higher particle degradation rates with increasing temperature. The much greater 14C loss from copepod carcasses than fecal pellets (at both temperatures) was due to greater leaching of DO14C from carcasses. This may reflect the fact that fecal pellets are composed of unassimilated, presumably refractory phytoplankton fragments, the labile fraction of phytoplankton having been absorbed by animals. Thus, in seawater those zooplankton carcasses not eaten should rapidly decompose. Harding (1973) reported that carcasses of Calanus finmarchicus at 22 °C were almost completely disintegrated within 3 d, a rate 4 times as rapid as at 4 °C.

Cho & Azam (1988) and Karl et al. (1988) concluded from studies in the Pacific that large sinking particles are poor substrates for bacterial growth; rather, particles appear to degrade indirectly by free-living bacteria utilizing DOC released from larger particles. Cho & Azam suggested that bacteria attached to particles could solubilize particles via enzymatic hydrolysis. The greater 14C loss rate at 18 than at 2 °C could be the result of enhanced leaching of DO14C caused by microbiologically mediated solubilization and subsequent utilization by free-living bacteria. Nevertheless, it can be concluded that leaching, even in the absence of microbial activity, could account for a significant amount of the DOC released from biogenic particles.

Lee & Fisher (1992) reported that phytoplankton cells treated with either HgCl2 or Formalin lost 26 to 29 % of their total cellular C as DOC within 24 h. The observed DO14C loss patterns in the poisoned samples could have resulted from differences in leaching mechanisms. Leaching from phytoplankton and zooplankton carcasses was presumably due to cellular 14C loss resulting from damages to cell membranes and/or cell lysis. Leaching of DO14C from fecal pellets was probably due to the gradual diffusion of solubilized organic material entrapped in fecal pellet matrices.

Jumars et al. (1989) concluded from model calculations that most of the organic solute in fecal pellets would diffuse out of the pellets within minutes of ingestion. This was observed with fecal pellets produced by the pontellid copepod Anomalocera patersoni (Fisher et al. 1991a), but not with the fecal pellets in this study. The most likely explanation for the lack of rapid release of DOC from the fecal pellets is that the chitinous peritrophic membrane encapsulating the pellets was relatively impermeable and hindered the diffusion of solute from the particles. We did not examine the degradation of the peritrophic membranes, however, several studies have shown that they degrade over periods of several hours to several weeks, depending on temperature (Small & Fowler 1973, Honjo & Roman 1977, Turner 1979, Bathelet & Schelske 1983). It is also possible that diffusion of solute from the fecal pellets occurred before collection or during rinsing, in which case the observed leaching represents solubilization of the remaining material after unassimilated digestive solute was lost. In any case, leaching was a major process responsible for C loss from the fecal pellets.

There is also an indication that DOC released from carcasses and fecal pellets differ in quality. Most of the DO14C leached from carcasses at both temperatures was rapidly remineralized to 14CO2, and only 10 % of the total 14C released from carcasses persisted as unutilized DO14C throughout the incubation. In contrast, only a small fraction of the released DO14C from fecal pellets at both temperatures was converted to 14CO2, and unutilized DO14C steadily increased to about 40 % of the total fecal pellet 14C by Day 30. Thus, up to 40 % of the material excreted by these copepods and ultimately released as DOC from fecal pellets appears to be refractory and a poor food source for microorganisms. This may explain why sinking particles collected from deep waters are sparsely colonized by microorganisms (Ducklow et al. 1985, Wiebe & Pomeroy 1972).

**Metal/metalloid release**

This is the first study to describe the kinetics of metal/metalloid release from zooplankton debris in the context of C release. The results are relevant to understanding processes that affect the remineralization of decomposing planktonic debris during its descent in the water column.

It appears from the data that microbial activity had a negligible effect on the release of 241Am from fecal pellets, since neither temperature nor HgCl2 appreciably affected release rates. Conversely, the release rate of 65Zn from fecal pellets was faster at 18 than at 2 °C; HgCl2 treatment had negligible effects on 65Zn retention. The y-intercepts of the lines relating retention of 241Am and 65Zn with C in decomposing fecal pellets were 53 and 20 %, respectively (Fig. 6), indicating greater particulate retention of these metals than of C.

Differential retention times of these elements were probably due to differences in the compounds to which
the metals were bound in the fecal pellets and different binding strengths of the elements to individual compounds. Reinfelder & Fisher (1991) found that 93% of Am, 52 to 74% of Zn, and 10% of Se in *Thalassiosira pseudonana* were bound to cell surfaces. They also showed that copepods assimilate the cytoplasmic fraction of diatoms, including those elements associated with this fraction. The more refractory material in cells, including cell walls, are not assimilated (Reinfelder & Fisher 1991) and are, in fact, found as major components of zooplankton fecal pellets (Silver & Bruland 1981, Gowing & Silver 1983). Thus, the $^{241}$Am and $^{65}$Zn in the fecal pellets were presumably bound to these refractory materials. $^{241}$Am was retained more effectively in the fecal pellets, being largely unaffected by leaching or microbial action, whereas $^{65}$Zn retention was clearly influenced by these processes.

The initial rapid loss of $^{241}$Am and $^{65}$Zn from fecal pellets was primarily attributable to chemical desorption, since about 15% of the $^{241}$Am and 20% of the $^{65}$Zn were lost during the initial period when essentially no $^{14}$C was lost from the fecal pellets (Fig. 6). Thus, $^{241}$Am bound to cell fragments in the fecal pellets rapidly desorbed to equilibrate with the $^{241}$Am-free ambient seawater, after which any $^{241}$Am release was probably due to both desorption and fecal pellet fragmentation leading to decreased particulate surface area. Some of the initial loss of $^{65}$Zn may also have resulted from isotope exchange of radioactive $^{65}$Zn on the particles with stable Zn in the seawater (there is no stable Am available for isotope exchange). After the initial $^{65}$Zn loss, any subsequent loss probably was due to leaching of dissolved organic compounds to which the $^{65}$Zn was bound and to microbial decomposition of cellular C.

$^{65}$Zn was retained more effectively by copepod fecal pellets than by carcasses, as would be expected given the rapid loss of DO$^{14}$C from carcasses and the faster degradation of organic $^{14}$C from the carcasses than from the fecal pellets. Analysis of euphausiid fecal pellets together with the animals which produced them and the food that they ate has shown that the fecal pellets were more enriched than the food or the zooplanktons in a variety of transition (Fowler 1977) and transuranic (Higgo et al. 1977, Fowler et al. 1983) metals. Greatest differences occurred for those metals (e.g. Am, Pu, Ce, Pb, Eu) which are least assimilated by the zooplankton, and which are most enriched in the fecal pellets (Fowler 1977).

The effects of incubation temperature on $^{75}$Se and $^{65}$Zn retention in copepod carcasses suggest that microbial activity had a significant effect on their release into the ambient seawater. However, $^{75}$Se was effectively retained in carcasses when treated with HgCl$_2$, while $^{65}$Zn was lost most rapidly from HgCl$_2$-treated carcasses. The experiment assessing the influence of poisons on elemental release from different particle surfaces showed that elements bound to mineral (SiO$_2$) surfaces were not displaced but $^{75}$Se and $^{65}$Zn bound to protein particles were displaced by HgCl$_2$ but not NaN$_3$. The displacement by Hg was probably due to the greater affinity of Hg than of Se or Zn for S ligands in the protein (Sillén & Martell 1964). The negligible difference observed in $^{65}$Zn release rates between samples treated with and without HgCl$_2$ may have been due to 2 opposing processes acting simultaneously. Suppression of microbial activity by HgCl$_2$, which would depress the release rate of $^{65}$Zn, could have been offset by displacement of the particulate $^{65}$Zn by Hg. $^{75}$Se was not displaced from carcasses by Hg$^{2+}$, suggesting that the $^{75}$Se in these particles was covalently bound in protein or selenoamino acids deriving from ingested phytoplankton (Wrench 1978, Fisher & Reinfelder 1991). The negligible association of $^{75}$Se with the glass beads probably reflects the fact that Se can behave analogously to S and binds to proteins and amino acids.

Loss of $^{75}$Se was faster than that of $^{14}$C from carcasses in which > 60% of the initial $^{14}$C was still particulate, but slower than $^{14}$C after > 40% of the initial $^{14}$C was lost. Similarly, release of $^{65}$Zn from carcasses was slightly faster than that of $^{14}$C. Since 35% of total carcass $^{14}$C leached out within 24 h and $^{75}$Se and $^{65}$Zn closely followed $^{14}$C, it appears that these radioisotopes were associated with soluble organic C. Therefore, leaching of soluble organic material together with microbial activity appear to be the mechanisms most responsible for release of these elements and organic C from decomposing biogenic particles. Other studies reported rapid release of Cd, Se and Zn from phytoplankton cells (Fisher & Wente 1993, Lee & Fisher 1992) and P, Cd, Ni and Mn from net plankton assemblages (mainly diatoms and copepods) (Collier & Edmond 1984).

The release rates of $^{241}$Am, $^{75}$Se and $^{65}$Zn from zooplankton debris decreased exponentially over time. Comparisons with other studies are difficult because few data exist on metal release from degrading zooplankton debris, and these are generally expressed as linear rates. Combining instantaneous release rates for $^{241}$Am, $^{75}$Se and $^{65}$Zn after 1 wk of < 1.4 % d$^{-1}$ (Table 2) with sinking rates of zooplankton fecal pellets of ≥ 50 m d$^{-1}$ (Komar et al. 1981) yields estimations that these elements still retained by particles which sank to depths of 300 m below the euphotic zone are likely to be released very slowly and continue to sink to deep waters. The release rates of these elements, very different in the early stages of loss, eventually converge to similar values.
The $t_{1/2}$'s of $^{241}$Am in the copepod fecal pellets were somewhat greater than values for this radioisotope in euphausiid fecal pellets (41 and 51 d at 5 and 13 °C, respectively) produced on a diet of radiolabeled Thalassiosira pseudonana cells (Fisher et al. 1983b). The $t_{1/2}$'s for $^{241}$Am were 1 to 2 orders of magnitude greater than the $t_{1/2}$'s of $^{65}$Zn and $^{75}$Se. The $t_{1/2}$'s of the elements in the debris in this study (labeled via ingestion) are $t_{1/2}$'s for debris labeled by scavenging. Thus, $t_{1/2}$'s for scavenged $^{241}$Am (3.5 to 32 d) and $^{65}$Zn (0.8 to 4 d) in various types of zooplankton fecal pellets and marine snow (at 15 °C) (Fisher et al. 1991b) were shorter ($^{241}$Am) than or comparable to ($^{65}$Zn) values observed here. The $t_{1/2}$'s for $^{75}$Se (4.5 d at 18 °C and 6 d at 4 °C) and $^{65}$Zn (7.5 d at 18 °C and 13 d at 4 °C) in decomposing diatom cells (Lee & Fisher 1992) were generally greater than values given here.

Marine phytoplankton can concentrate many metals very appreciably out of seawater, with concentration factors exceeding 10$^{5}$ for some metals (IAEA 1985). Zooplankton feeding on this material could influence the cycling of many elements in the ocean (Fisher et al. 1991a, Reinfelder & Fisher 1991). These studies demonstrated that Se and Zn are efficiently assimilated in zooplankton and should be recycled in surface waters while particle-reactive, unassimilated metals such as Am are largely packaged into fecal pellets. Given the elemental release rates and $t_{1/2}$'s reported here, Se and Zn associated with copepod carcasses and fecal pellets would be expected to be rapidly released and recycled in surface waters, consistent with oceanographic observations (Whitfield & Turner 1987). Particle-reactive metals such as Am, which show minimal association with the organic cycle and negligible assimilation in herbivores, are effectively retained in sinking fecal pellets during their descent in the water column, resulting in shorter oceanic residence times (Cherry et al. 1978) and rapid transport to deep water. Scavenging on sinking debris (Fowler et al. 1983, Fisher et al. 1991b) could result in further removal of these metals from the water column.

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