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Long lifetimes of β-glucosidase, leucine aminopeptidase, and phosphatase in Arctic seawater

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Abstract

The active lifetime of extracellular enzymes is a critical determinant of the effectiveness of enzyme production as a means for heterotrophic marine microbes to obtain organic substrates. Here, we report lifetimes of three classes of extracellular enzyme in Arctic seawater. We also investigated the relative importance of photochemical processes and particle-associated processes in inactivating extracellular enzymes. Enzyme inactivation in filtered seawater was slow, with apparent half-lives of enzyme activities on the order of hundreds of hours. The presence of particles (including cells) did not significantly change inactivation rates, suggesting that the long half-lives observed in filtered seawater were realistic for enzymes in unfiltered seawater. Phosphatase and leucine aminopeptidase were susceptible to photoinactivation, but only under high intensity UV-B and UV-C illumination; there was no evidence for increased inactivation rates under natural illumination at our study site in Ny Ålesund, Svalbard. Comparison of inactivation rates of commercially-obtained enzymes from non-marine sources with the extracellular enzymes naturally present in Arctic seawater suggests that the natural enzymes contain structural features that confer longer lifetimes, consistent with observations reported by others from a range of field sites that cell-free enzymes can contribute a substantial fraction of total hydrolytic activity in the water column.

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1. Introduction

Heterotrophic microbes rely on extracellular enzymes to perform the initial step of organic matter remineralization by transforming high molecular weight (HMW) organic matter into molecules small enough to cross the cell membrane (Benz and Bauer, 1988). The dynamics of extracellular enzymes in solution therefore affect the nature and patterns of organic matter remineralization by aquatic microbial communities. Extracellular enzymes may be anchored to the bacterial outer membrane; they may also be released from the cell into solution, depending on microbial growth phase and nutrient state (Christ, 1991, and references therein). Enzymes may also be released into solution as a result of viral lysis, or of zooplankton grazing (Bochdansky et al., 1995; Karner et al., 1994). Extracellular enzymes released into solution can provide sufficient substrate for cellular growth in pure culture (Vetter and Deming, 1999), and can at times constitute a considerable fraction of total enzyme activity in seawater (Baltar et al., 2010; Karner and Rassoulzadegan, 1995; Keith and Arnosti, 2001; Obayashi and Suzuki, 2008a,b). The active lifetime, as well as the physical location of extracellular enzymes thus influence the conditions under which microbial communities may “profitably” remineralize high molecular weight organic matter (Allison, 2005; Vetter et al., 1998). Enzymes with longer lifetimes would be more valuable to the organisms that produce them (Allison, 2005; Vetter et al., 1998), and might catalyze degradation of organic matter farther from the organism that produced them. Despite the importance of enzyme activities to heterotrophic microorganisms and to the marine carbon cycle as a whole, relatively little is known about the active lifetime of extracellular enzymes in seawater. Most studies of extracellular enzyme lifetimes have been conducted on purified enzymes from marine organisms in sterile buffer solution, or on ensembles of enzymes naturally present in fresh water. For example, a cold-active extracellular protease isolated from an Arctic marine bacterium had a half-life of approximately 12 h in sterile buffer solution, and retained activity for 45 h when stabilized with extracellular polymeric substances (Huston et al., 2004). Commercially-available phosphatase and glucosidase had half-lives of 4 to 8 h in humic-rich lake water (Scully et al., 2003). The principal decay pathway in that case was an acid-catalyzed reaction with iron, a pathway that is not likely relevant at the higher pH of seawater. Interactions between ultraviolet (UV) radiation and humic substances also may have complex effects on enzyme inactivation in lake water (Boavida and Wetzel, 1998; Tietjen and Wetzel, 2003). Humic substances and UV radiation each initially decreased the activity of extracellular enzymes, but enzymes complexed with humic substances were apparently protected from
inactivation via other pathways, and some humic-bound enzymes could apparently be re-activated by exposure to UV.

The rates and pathways of enzyme inactivation in seawater can be expected to differ from humic-rich and acidic freshwater environments described above. The goals of this study were to investigate decay rates of cell-free extracellular enzymes naturally present in Arctic seawater, and to assess the importance of photochemical processes on those decay rates. Extracellular enzymes from marine microbes might be structurally adapted for maximal lifetime in seawater (Vetter et al., 1998), so the use of naturally present extracellular enzymes may yield more realistic lifetimes than an investigation using commercially available enzymes, from non-marine sources, added to seawater. Assays of the lifetime of enzymes found naturally in seawater also avoid potential artifacts stemming from the fact that families of enzymes carrying out the same function can differ greatly in structure (Domäń-Pytko and Bardowski, 2004).

In order to prevent production of new enzymes by seawater microbes, which would mask inactivation of the enzymes present at the time of sampling, we removed microbes by filtration (0.2 μm) and measured the decay rates of enzymes naturally present in the filtrate. Incubations were kept short (<36 h) to minimize the importance of enzyme production by cells that may have passed through the filter. The role of visible and ultraviolet light in inactivating seawater enzymes was assessed by exposing natural enzymes to the full spectrum of natural sunlight, only the visible portion of the solar spectrum, and to UV light from an artificial source. An experiment comparing inactivation rates of commercially-obtained phosphatase in filtered and unfiltered seawater was conducted to investigate the extent to which inactivation rates measured in filtered water can be compared to rates in unfiltered seawater. These experiments provide an assessment of the lifetime of several extracellular enzymes in Arctic seawater, as well as the extent to which sunlight and particles act to influence those lifetimes. These results will help to define the contexts in which cell-free extracellular enzymes represent a viable strategy for resource acquisition by heterotrophic microbes in the cold waters.

2. Methods

2.1. Sample collection

Surface water samples from Kongsfjord, Svalbard (78.93°N, 11.95°E), were collected by bucket from the end of a pier in Ny Ålesund between August 16 and 22, 2007, immediately prior to performing the experiments. Water temperature was between 4 °C and 7 °C at the time of sampling. Samples were processed in an environmental chamber at 7 °C. In experiments using filtered seawater, seawater was filtered through 0.22 μm pore-size Durapore filters under a light vacuum (~50 mm Hg).

2.2. Measurement of enzyme activities

Enzyme activities were measured using the fluorogenic substrate proxies 4-methylumbelliferyl-phosphate (MUF-PO₄, to measure phosphatase activity), 4-methylumbelliferyl-α-D-glucopyranoside and 4-methylumbelliferyl-β-D-glucopyranoside (MUF-α-gluc and MUF-β-gluc, to measure α-glucosidase and β-glucosidase activities respectively), and L-leucine-7-amido-4-methylcoumarin hydrochloride (leu-MCA, to measure leucine aminopeptidase activity) following King (1986) and Somville and Billen (1983). All substrates were obtained from Sigma-Aldrich. Fluorescence was measured using a Turner Biosystems TBS-380 fluorometer, with excitation/emission channels set to the UV setting (365 nm excitation, 440–470 nm emission). In all experiments, the procedure to measure enzyme activities was as follows: 30 μL fluorogenic substrate stock solution was added to a disposable methacrylate cuvette followed by 2 mL sample water for a final concentration of 300 μM L⁻¹ (previously determined to be saturating concentrations for MUF-PO₄, MUF-β-glu, and leu-AP) and mixed with a micropipetor. Fluorescence was measured immediately afterwards, and one to three more times over the course of 30 min. Enzyme activity in each replicate was taken as the slope of the regression line of fluorescence versus incubation time. The number of replicates for each experiment is given in each figure legend. Abiotic hydrolysis of the fluorogenic substrate was assessed using autoclaved seawater; this value was subtracted from the measured enzymatic activities and was in all cases <20% of the total hydrolysis rate. The fluorimeter was calibrated using standard solutions of 4-methylumbelliferyl-β-D-glucosidase and 4-methylcoumarin in filtered seawater. Detector stability was monitored using a sealed standard (Turner Biosystems). Statistical analysis was carried out using the R statistical platform. No α-glucosidase activity was detected in the saturation curve measurements, so α-glucosidase activity was not assayed in any further experiments.

To assess the contribution of cell-free leucine aminopeptidase activity to total activity, leucine aminopeptidase activity was measured at 7 °C in 0.2 μm-filtered water and in unfiltered water.

2.3. Inactivation of natural enzymes under natural light

To assess the inactivation rates of the three classes of natural extracellular enzymes under sunlight, UV-free sunlight, and in the dark, nine sterilized polyethylene bags were filled with 250 mL 0.2 μm-filtered seawater and placed in low-walled tubs. Three bags were left uncovered in these tubs (full-spectrum treatment), three bags were covered in Lexan (visible light only treatment) and three were covered in aluminum foil (dark treatment). Polyethylene absorbs light weakly from 240–280 nm, and is transparent above 280 nm. Lexan is opaque below 400 nm and fully transparent above 440 nm. The absorbance spectra of the specific materials used here were confirmed by spectrophotometry (data not shown). Temperature of the tubs was stabilized at 4–7 °C by filling the tubs with 4 °C water, constantly supplied from a cooler. The bags were periodically subsampled through a sampling port by withdrawing ~10 mL from each bag into a sterile syringe. The subsamples were transferred to a glass scintillation vial.

Phosphatase, β-glucosidase, and leucine aminopeptidase activities were measured as described previously, after warming the samples to 20 °C by immersing the scintillation vials in a 20 °C bath for eight minutes. The purpose of this brief warming was to increase enzyme activity by a constant factor at each timepoint, in order to more precisely measure changes in enzyme activity as a function of time. As in other studies, the assumption is that Q₁₀ values (the dependence of enzyme activity on temperature) are constant, so this warming should not change the rate constant for enzyme inactivation that had occurred during incubation at 7 °C. Because hydrolysis rates measured at 20 °C do not reflect in situ rates, relative fluorescence units are reported rather than moles of monomer freed. Our focus is on determining the activity (under constant conditions) of the functional pool of enzymes at each time point, rather than on determining the in situ hydrolysis rates. Calibration curves indicated a conversion factor of 287 PM MUF- (fluorescence unit)⁻¹, and 610 PM MCA- (fluorescence unit)⁻¹.

The weather conditions were cloudy during the experiment. Average global radiation in Ny Ålesund was 65 W m⁻², of which 3.8 W m⁻² was UV. Meteorological data was obtained from the Ny Ålesund observatory maintained by the Alfred Wegener Institute. The relative contributions of UV-A and UV-B were not reported, but UV-B typically accounts for 1–2% of the total UV radiative power at Ny Ålesund (Hanelt et al., 2001); UV-B power was therefore 0.04–0.08 W m⁻².

2.4. Inactivation rates of commercial enzymes in artificial seawater

Commercially-obtained phosphatase (Sigma P4252, from Escherichia coli), β-glucosidase (Sigma 49290, from almonds) and leucine
aminopeptidase (Sigma L5006, from porcine kidney) were added to three replicate beakers of 100 mL autoclaved, artificial seawater (Sigma sea salts 59883, 35.0 g L$^{-1}$). Sigma-Aldrich confirmed that no secondary enzymes (e.g. proteases) were added to the purified enzymes as part of the production process (pers. comm.). Final enzyme concentrations were 3 ng phosphatase mL$^{-1}$, 2 μg β-glucosidase mL$^{-1}$, and 12 μg leucine aminopeptidase mL$^{-1}$. Initial enzyme activities were 28.9 nmol L$^{-1}$ min$^{-1}$ (phosphatase), 866 nmol L$^{-1}$ min$^{-1}$ (β-glucosidase) and 7.12 μmol L$^{-1}$ min$^{-1}$ (leucine aminopeptidase). The experiment was carried out in an environmental room at 7 °C, and enzyme activity was measured periodically as described above, but at 7 °C. Because the activity of added commercial enzymes was much greater than that of cell-free enzymes naturally present in filtered seawater, activity was sufficiently high to be measured with precision at 7 °C.

2.5. Role of particles in inactivating commercial phosphatase

To assess the importance of the presence of particles for the lifetime of dissolved extracellular enzymes, inactivation of commercially-obtained phosphatase (see previous section) was assessed in both unfiltered and filtered seawater collected from Kongsfjord as described above. Three sterile 50-mL centrifuge tubes were each filled with 50 mL unfiltered seawater, and three others were filled with 50 mL filtered seawater. Phosphatase was added to each sample to a concentration of 12.2 μg protein L$^{-1}$. This quantity of phosphatase yielded an activity about 20 times greater than in situ phosphatase activity in unfiltered seawater, thus minimizing the relative contribution of any phosphatase enzymes produced by microbes in the unfiltered seawater, which could mask phosphatase inactivation. Centrifuge tubes with added phosphatase were incubated in the dark in an environmental room at 7 °C, and enzyme activity was measured periodically (as described above) at 7 °C.

2.6. Inactivation of natural enzymes in the presence of artificial UV illumination

Filtered seawater was used to assess the susceptibility of natural dissolved enzymes to photochemical degradation. A 100 mL volume of filtered seawater was dispensed into each of six sterile Pyrex beakers, with UV-transparent polyethylene stretched across the beaker mouth (Pyrex itself is transparent in the visible but largely opaque in UV). Three beakers were wrapped completely in aluminum foil. All six beakers were placed in an environmental room at 7 °C directly under UV- and visible-emitting fluorescent lamps (UV: Q-Panel UVA-340, Cleveland, OH, USA; visible: Osmar L65/25S, Munich, Germany). Visible photon flux was 18.6 μmol photon m$^{-2}$ s$^{-1}$ (approximately 4.0 W m$^{-2}$), UV-A power was 4.6 W m$^{-2}$, and UV-B power was 0.4 W m$^{-2}$.

Approximately 15 mL water was removed via sterile syringe from each beaker at each timepoint and placed in a clean 20 mL glass scintillation vial. Phosphatase, β-glucosidase, and leucine aminopeptidase activities were measured as described above, at 20 °C.

3. Results

3.1. Inactivation of natural enzymes under natural light

Natural cell-free enzymes, incubated in filtered seawater at 7 °C (and assayed at 20 °C) lost very little activity over the course of 36 h (Fig. 1). None of the treatments of phosphatase or leucine aminopeptidase showed a significant relationship between activity and time, whereas all of the treatments of β-glucosidase did (Table 1). Calculated half-lives of β-glucosidase centered on a range of 100–200 h. No effect of light treatment was clearly distinguishable, indicating that natural sunlight had at most a modest effect on inactivation rates.

3.2. Inactivation rates of commercial enzymes in artificial seawater

All three commercially-obtained enzymes lost substantial activity over the course of 12–48 h when incubated at 7 °C in the dark in artificial seawater (Fig. 2). Phosphatase decay fit an exponential decay model well ($r^{2}=0.82$), with a half-life of 6.8 h. The decay of leucine aminopeptidase and β-glucosidase did not correspond to first-order kinetics, with an initial rapid rate of inactivation and a later, slower phase, so half-lives were not calculated for those enzymes.

3.3. Inactivation rates of commercial phosphatase in filtered and unfiltered seawater

The activity of commercial phosphatase incubated in both filtered and unfiltered seawater at 7 °C in the dark decreased substantially during a 5-h incubation (Fig. 3). Unlike the incubation in artificial seawater, phosphatase inactivation appeared linear with time in both filtered and unfiltered seawater. Decay rates, calculated from a linear least-squares fit, were 12.0 nmol L$^{-1}$ h$^{-1}$ ($r^{2}=0.98$) in unfiltered water, two times faster than the rate of 6.0 nmol L$^{-1}$ h$^{-2}$ ($r^{2}=0.85$) in 0.2 μm-filtered water, although the difference in slopes was not significant (one-way analysis of covariance, p = 0.42).

3.4. Inactivation rates of natural enzymes under artificial UV light

Activities of natural phosphatases and leucine aminopeptidases decreased when exposed to artificial UV illumination (incubated at 7 °C, assayed at 20 °C; Fig. 4). Natural β-glucosidases activities, however, did not change systematically. In order to isolate the effect of light from other causes of enzyme inactivation, results are reported as the ratio of activity in the light-exposed samples to activity in the dark-exposed samples, $\text{time}_{\text{r}}=\text{UV}_{\text{r}}/\text{dark}_{\text{r}}$. The clear effect of artificial illumination on phosphatases and aminopeptidases contrasts with the weak or non-existent effect of natural radiation (compare Fig. 4 to Fig. 1 a,b,c). Initial enzyme activities in the dark samples were 49.8 ± 0.5 fl unit h$^{-1}$ (phosphatase), 3.63 fl unit h$^{-1}$ (β-glucosidase), and 31.5 fl unit h$^{-1}$ (leucine aminopeptidase). Initial activities in the light-exposed samples were all within the range of error of the dark values.

3.5. Contribution of cell-free enzymes to total enzyme activities

The only naturally present enzyme assayed in both filtered and unfiltered seawater at 7 °C was leucine aminopeptidase. Activity in unfiltered water was 563 nmol L$^{-1}$ h$^{-1}$, with 10 ± 2.0% of that activity measured in filtered water.

4. Discussion

4.1. Inactivation of natural and commercial enzymes in the dark

Over 36-h dark incubations, all three commercially-obtained enzymes decayed much faster than their natural counterparts (Figs. 1–3). Natural enzymes – which are in fact enzyme classes, and may be represented by tens or more distinct enzymes (Arrieta et al., 1998) – exhibited no detectable loss of activity over this time period, with only β-glucosidase unambiguously exhibiting any decay (28% at the final timepoint, Fig. 1). The greater stability (longer half-lives) of the natural compared to the commercially-obtained enzymes may be due to intrinsic structural stability of the naturally-occurring enzymes. Enzymes intended to function in the extracellular environment, a dilute aqueous medium compared to the crowded interior of a cell, likely have structural features that stabilize them relative to their intracellular counterparts.

Extracellular enzymes may also be protected against inactivation by association with DOM (Keil and Kirchman, 1994) and/or...
extracellular polymeric substance (EPS; Huston et al., 2004). Such associations with DOM or EPS could potentially also stabilize the commercially-available phosphatase against inactivation, although the relatively rapid inactivation of these enzymes in filtered and in unfiltered seawater (44% loss of activity in unfiltered water and 21% loss of activity in filtered water over a 5-time course; Fig. 3) suggests that this mechanism is either not as effective for the commercial as for the natural enzymes (decay undetectable over 36 h; Fig. 2), or that the protective effect of DOM/EPS was overwhelmed by the addition of the extra phosphatase enzymes to the seawater solution. Since the stoichiometry and mechanism of this protective effect are both unknown, however, the degree of saturation cannot be assessed. A definitive test of stabilization mechanisms (and measurement of the nature and extent of protection afforded by specific fractions of DOM and EPS) awaits considerable improvements in capabilities of determining the solution structure of enzymes, of DOM, and of EPS.

4.2. Pathways of enzyme decay: light and particles

Photochemical reactions have been implicated in the inactivation of extracellular enzymes (Espeland and Wetzel, 2001) and degradation of bulk DOM (Moran and Zepp, 1997) and DON (Vähätalo and Zepp, 2005). The increased inactivation rates of natural phosphatase and leucine aminopeptidase in the presence of bright, artificial UV demonstrate that phosphatase and leucine aminopeptidase were in principle susceptible to photochemical inactivation (Fig. 4). However, neither full spectrum natural sunlight nor the visible fraction of sunlight elicited a decrease in natural enzyme activity over 36 h (Fig. 1). Intensities of visible and UV-A light from the artificial light source were comparable to the outdoor incubation. The major difference between the natural and artificial light spectra was at the shorter UV wavelengths: UV-B (280–215 nm) was roughly 5–10 times more intense from artificial lamps than outdoors. Fluorescent lamps also typically emit some UV-C (<290 nm) radiation, whereas essentially no radiation shorter than 300 nm reaches the Earth’s surface, even under an ozone-depleted atmosphere (Caldwell and Flint, 1994). The rapid inactivation of leucine aminopeptidase and phosphatase under artificial illumination was therefore likely due to shorter wavelengths than those that are relevant to natural systems. Photochemical reactions therefore do not appear to be a relevant pathway for the decay of these broad enzyme classes in Arctic seawater on the timescale of days.

In addition to intrinsic (structural) factors, interaction with DOM and EPS, and light intensity, interactions with particles—including live cells—may affect the inactivation rates of dissolved enzymes. This interaction can be envisioned in several forms: sorption of enzymes to cell or particle surfaces may extend the active lifetime of the enzymes (Ziervogel et al., 2007). Dissolved enzymes may also be the target of

<p>| Table 1 | Results of model fits for natural enzymes incubated under natural light. Slope is linear least-squares fit. n.s. indicates Bonferroni-adjusted $p &gt; 0.05$. |</p>
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<tr>
<th>Phosphatase</th>
<th>Leucine aminopeptidase</th>
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<td>Dark</td>
<td>Vis</td>
<td>Full spectrum</td>
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<td>Slope, fl unit hr$^{-1}$</td>
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<td>Half-life, hr (95% conf limits)</td>
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Inactivation of commercially-obtained phosphatase, leucine aminopeptidase and \(\beta\)-glucosidase in artificial seawater, incubated at 7 °C in the dark and measured at 7 °C. Activity is expressed relative to initial activity for each enzyme. Error bars represent standard deviation of 3 replicates.

Fig. 3. Inactivation of commercially-obtained phosphatase incubated in filtered and unfiltered seawater at 7 °C in the dark and measured at 7 °C. The line is a linear least-squares regression of the data. Error bars represent standard deviation of 3 replicates.

4.3. Implications of multi-day half-lives of extracellular enzymes in cold seawater

In cold water, cell-free extracellular enzymes are stable for days or weeks, implying that enzymes may contribute to substrate availability at considerable distances (potentially meters to kilometers; see supplemental information) from the producing cell. The cost–benefit analysis of enzyme production under such circumstances might need to include the possibility that cell-free enzymes may benefit a community of organisms working in concert to hydrolyze substrates. The observation that enzyme production may be linked to quorum-sensing mechanisms (Chernin et al., 1998; Givskov et al., 1997; Gram et al., 2002; Withers et al., 2001) supports such a perspective. In any case, the extent to which enzyme production would be ‘profitable’ for a microbial community would depend upon the return of substrate to the cells while they are still in proximity to the enzymes (Vetter et al., 1998).

Field evidence clearly shows that cell-free enzymes can contribute substantially to total hydrolytic activity. A range of 5% to nearly 100% has been reported for the contribution of cell-free extracellular enzymes to total enzyme activity in diverse aquatic environments, including estuarine, coastal, and deep-sea sites (Baltar et al., 2010; Karner and Rassoulzadegan, 1995; Keith and Arnosti, 2001; Obayashi and Suzuki, 2008a,b). The 10% contribution of cell-free leucine aminopeptidase to total leucine aminopeptidase activity is on the low end of that range. To the best of our knowledge, these are the first data on cell-free extracellular enzyme activity from the Arctic. A recent report, however, found that the contribution of cell-free enzymes was particularly large in the deep sea, another permanently cold environment (Baltar et al., 2010). This latitudinal transect of the Atlantic showed cell-free enzymes consistently contributing >60% of total activity in samples deeper than 100 m, with contributions often close to 100%. Such a high fraction is something of a puzzle, since model results suggest that cell-free enzymes would only be ‘profitable’ in the context of aggregates or other particles (Allison, 2005; Vetter et al., 1998). Baltar et al. (2010) suggested that disruption of colloidal material during sampling might have been responsible for the high fraction of cell-free activity. The results presented here suggest a second possibility: extracellular enzymes diffusing or advecting from aggregates (Ziervogel and Arnosti, 2008) would be stable over long timescales, and might make a substantial contribution to total enzyme activity against a low background of cell-associated enzyme activity in the deep ocean.

Stability of enzymes – both cell-free enzymes and those bound to particles – is an important parameter for the microbial loop, as well as for efforts to determine factors constraining the turnover rates of organic carbon in marine systems (Boudreau et al., 2008). Further investigation is required to understand more precisely how the specifics of enzyme structure, biological activity, and enzyme physical state (e.g. sorbed enzymes versus dissolved, DOM interactions, nature of sorbing surface) influence enzyme stability. More precise modeling efforts require accurate measurements of enzyme inactivation rates.
The data presented here indicate that commercially-obtained enzymes may be a poor model for marine enzymes in such studies. The use of natural marine enzymes found in situ as in this study also has some significant complications: enzyme activities cannot easily be manipulated, and activity may be close to the detection limit. Further progress may be achieved by using enzymes released by pure cultures of organisms isolated from the relevant environment (e.g. Boavida and Wetzel, 1998; Huston et al., 2004). A thorough understanding of the dynamics of extracellular enzymes after they are produced by marine microbes will allow a more predictive understanding of the marine microbial loop.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at doi:10.1016/j.marchem.2010.10.006.

References


