INTRODUCTION

Biotic mineralization of dissolved organic material (DOM) in aquatic ecosystems is carried out by a diverse community of heterotrophic microbes, potentially consisting of representatives from over 50 divisions (or phyla) of bacteria and Archaea (Rappé & Giovannoni 2003). In aerobic water columns of many aquatic ecosystems, the heterotrophic assemblage is dominated by only a few bacterial groups from 2 of these divisions (Giovannoni & Rappé 2000, Kirchman 2002); Archaea appear to be abundant in the deep ocean, but not in surface waters (Karner et al. 2001).

The abundant heterotrophic bacterial groups in surface waters include 3 subdivisions of Proteobacteria (alpha-, beta-, and gamma-proteobacteria) and a subgroup of the Bacteroidetes, often referred to as Cytophaga-Flavobacterium or Cytophaga-like bacteria. Actinobacteria can be another abundant group in freshwaters (Glöckner et al. 2000), although these waters are often dominated by beta-proteobacteria (Glöckner et al. 1999). Alpha-proteobacteria, including the SAR11 clade (Morris et al. 2002), are abundant in the oceans (Glöckner et al. 1999) whereas Cytophaga-like bacteria can be abundant in both freshwater and marine systems (Kirchman 2002).
The abundances of these bacterial groups are probably affected by several factors, including the composition and concentration of DOM. These other DOM parameters often control community-level rates of bacterial growth and biomass production, and there is evidence of DOM affecting the structure of natural bacterial communities (Rossello-Mora et al. 1999, van Hannen et al. 1999, Eilers et al. 2000a, Covert & Moran 2001, Fandino et al. 2001, Muylaert et al. 2002). That DOM affects the abundance of individual ‘species’ or ribotypes (e.g. van Hannen et al. 1999, Fandino et al. 2001, Muylaert et al. 2002) would not be surprising, because bacteria in pure cultures are well known to differ in their capacity to use various DOM components (Martinez et al. 1996). What is less clear is the impact of DOM on the abundance of major phylogenetic groups such as the subdivisions of Proteobacteria and Cytophaga-like bacteria. Variation in DOM may affect the abundance of these phylogenetic groups if there are systematic differences in how these groups use or respond to DOM or if these groups are consistently dominated by a few species that differ in DOM utilization Cottrell & Kirchman (2000) found differences in DOM use among subdivisions of Proteobacteria and Cytophaga-like bacteria, supporting the common generalization that Cytophaga-like bacteria are adept at using biopolymers and high molecular weight (HMW) DOM (Kirchman 2002). Davey et al. (2001) noted that high Cytophaga-like bacterial abundance and proteolytic activity co-occurred in surface waters of the northeast Atlantic Ocean. However, it is still unclear whether generalizations about DOM use at this phylogenetic level will prove useful, because each of these groups contains several subgroups and ribotypes (Giovannoni & Rappé 2000) that probably vary greatly over time and space.

An important step in DOM mineralization which community structure may affect is the hydrolysis of HMW DOM by cell-associated extracellular enzymes (ectoenzymes). Although not necessarily a rate-limiting step in DOM degradation (Arnosti 2003), it is often the first step because the molecular size of labile DOM is frequently large enough (Amon & Benner 1994, Amon & Benner 1996) to require extracellular hydrolysis before transport and ultimate mineralization. Previous work with pure cultures demonstrated that ectoenzyme activity varies greatly among bacterial species, and that no single species expresses all ectoenzymes with high activity (Martinez et al. 1996), and so variation in bacterial community may be linked with variation in enzymatic activity. Little is known about the ectoenzyme activity of uncultured bacterial groups, except for the possible special role of Cytophaga-like bacteria in HMW DOM hydrolysis (see above). It is unclear if all bacterial communities are equally capable of expressing all enzymes necessary for hydrolyzing the range of HMW DOM components found in natural aquatic ecosystems, or if a new community must develop before new DOM components are hydrolyzed and ultimately mineralized.

Lakes and rivers may be ideal systems to explore relationships between community structure and DOM properties, since the amount and composition of DOM can vary greatly in these aquatic systems (Cotner et al. 2000, Mulholland et al. 2001, Cole et al. 2002). In the Hudson River, for example, concentrations of dissolved organic carbon (DOC) vary spatially by nearly 2-fold, and several direct and indirect indices suggest even greater variation in DOM composition (Findlay et al. 1998). This variation is caused by input of DOM from both plankton and terrestrial sources. Plankton-derived DOM is enriched in protein and labile polysaccharides, whereas terrestrial DOM contains humic material and structural polysaccharides, such as cellulose and lignin, which are relatively resistant to mineralization by microbial processes (Benner 2002, 2003). DOM in rivers, which is often dominated by terrestrial organic material, appears to be relatively less labile than DOM in marine systems, which is mostly from planktonic sources. In long-term bioassays, ca. 12% of DOC from rivers is consumed on average, only half as much as the average fraction (26%) consumed in bioassays of marine DOC (del Giorgio & Davis 2003).

We examined relationships among ectoenzyme activity, bacterial growth, and bacterial community structure in the Hudson River and its tributaries. Our main approach was to mix bacterial communities and water from various locations within the Hudson River system and then to monitor bacterial biomass production, ectoenzyme activity, and community structure. The data from these experiments help us explore bidirectional controls: the influence of DOM on bacterial composition and activity, and the role of bacterial community structure in controlling DOM degradation. We found that both ectoenzymatic activity and bacterial community structure changed in response to DOM composition and concentrations and that the activities of several ectoenzymes were associated with changes in microbial community structure.

MATERIALS AND METHODS

The water and microbial assemblages used in these experiments were from locations spanning the tidal Hudson River, which extends for 240 km from north of Albany (Peebles Island State Park, above the dam, but referred to here as ‘Albany’) to New York City (Dykeman Street RM10 in Manhattan) (Fig. 1). Water was taken from these locations and transported to the laboratory, where processing began within 4 h. In addition to the
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Dissolved organic carbon in ambient waters and from the experimental flasks was measured with a Shimadzu 5050 TOC Analyzer. Fluorescence characteristics of ambient waters were measured with a Perkin Elmer spectrofluorometer. Concentrations of dissolved inorganic nitrogen (DIN) and soluble reactive phosphorus (SRP) in ambient waters were measured by standard automated wet chemistry.

**Bacterial growth experiments.** Two types of growth experiments were conducted. One set used water and microbial assemblages (the inoculum) from 2 endmembers of the Hudson River, Albany and New York City (‘switch experiments’). In brief, 900 ml of 0.22 µm-filtered water (mixed esters of cellulose, Millipore GS) from each site was inoculated with 100 ml of an unfiltered inoculum derived from either the same location or the opposite endmember. Each treatment had 3 replicates. The experiments were then subsampled daily for 3H-leucine incorporation (see below) to track growth, and the experiment was terminated when rates were relatively constant.

Another type of bacterial growth experiment (‘tributary’ experiment) consisted of adding one inoculum to water from 3 tributaries of the Hudson River and from a 4th location, the upper Hudson River near Waterford, above the tidal portion. The inoculum was from Norrie Point (Fig. 1), which is near the longitudinal mid-point of the main stem of the tidal Hudson. The 3 tributaries were Mohawk River, Esopus Creek and Wallkill River. As with the switch experiment, an unfiltered inoculum (100 ml) was added to 0.22 µm-filtered water (900 ml). Each treatment had 3 replicates, which were subsampled over time as described above.

Bacterial growth was assayed at 5 time points (0, 24, 48, 54 and, 72 h) as the rate of incorporation of 3H-leucine (Smith & Azam 1992, Kirchman 2001). At the end of the experiment, ectoenzyme activity, bacterial community structure (see below), and the rate of incorporation of 3H-thymidine into DNA was measured using the procedure routinely used for field samples in the Hudson River (Findlay et al. 1991).

**Ectoenzyme activity.** The activity of various ectoenzymes was measured both in the experiments and at the locations in the Hudson system where we collected water and microbes for the experiments. The procedure followed that already described elsewhere (Findlay et al. 1998). In brief, ectoenzyme activity was assayed by observing the release of fluorescent methylumbelliferone (MUF) from MUF-linked analogues. The following substrates were used for the indicated enzyme (‘substrate for enzyme’): 4-MUF-acetate for esterase; 4-MUF-phosphate for phosphatase; L-leucine 7-amido-4-methyl-coumarin for aminopeptidase; 4-MUF-alpha-glucoside for alpha-glucosidase; 4-MUF-beta-glucoside for beta-glucosidase; 4-MUF-beta-xyloside for xylase; 4-MUF-beta-N-acetylglucosaminide for beta-N-acetylglucosaminidase; 4-MUF-4-guanidinobenzoate for trypsin. Enzyme activity was measured at a single concentration (400 µM, except for MUF-acetate which was added at 100 µM). These concentrations resulted in maximal enzyme rates or were at the solubility limit of the substrate (Findlay et al. 1998). Fluorescence was monitored over 1 to 4 h in a 96-well microtiter plate spectrofluorometer (Perkin-Elmer LS50B) and the rate was calculated from the linear part of the fluorescence versus time relationship. Fluorescence quenching was corrected by addition of internal standards (addition of known amounts of MUF to actual samples) and fluorescent units were converted to molar rates with a standard curve.

**Fluorescence in situ hybridization (FISH).** Bacterial community structure was examined by FISH with oligonucleotide probes (Amann et al. 1995). Water for this analysis was preserved in 2% formaldehyde and...
frozen (–20°C) until filtration through 0.2 µm polycarbonate filters was possible. The number of cells detectable by FISH after this preservation appears to be similar to that in samples preserved in paraformaldehyde without freezing (Cottrell & Kirchman unpubl. data).

The relative abundance of major phylogenetic groups was determined using CY3-labeled (MWG Biotech) Probe Eub338 for bacteria (Amann et al. 1990), Alf968 for alpha-proteobacteria (Glöckner et al. 1999), Bet42a for beta-proteobacteria (Manz et al. 1992), Gam42a for gamma-proteobacteria, CF319a for the Cytophaga-like bacterial group (Manz et al. 1996) and a negative control probe for non-specific binding (Karner & Fuhrman 1997). Samples were probed for 18 to 20 h at 42°C. The hybridization solution contained 0.9 M NaCl, 20 mM Tris-HCl (pH 7.4), 0.01 % sodium dodecyl sulfate, and the appropriate concentration of formamide (Zarda et al. 1997, Eilers et al. 2000b). After hybridization, the sample was transferred to a wash solution containing 20 mM Tris-HCl (pH 7.4), 5 mM EDTA, 0.01 % sodium dodecyl sulfate, and a concentration of NaCl appropriate for the probe (Zarda et al. 1997, Eilers et al. 2000b). After staining with DAPI, the sample was mounted with Vectashield and Citifluor (1:4) and examined with a semi-automated image analysis system coupled to an Olympus epifluorescence microscope (Cottrell & Kirchman 2003).

**Statistical analysis.** The data were analyzed by standard ANOVA and regression techniques. The FISH data, which were expressed as a percentage of total prokaryotes (DAPI-positive cells), were arcsin transformed before analysis. A principal components analysis (PCA) was used to simplify the 8 enzyme activities or the abundance of 4 bacterial groups into 2 new linear variables that encompassed most of the variability in the original data. The PCA uses the correlation among variables rather than absolute rates, so all variables have equal weight in the analysis.

**RESULTS**

**Ambient DOM and bacterial communities**

DOM characteristics and the bacterial communities differed greatly among the sampling sites examined by this study. There was a 2-fold difference in bulk DOC concentration among the tributaries and a ca. 25 % difference between New York City (NYC) and Albany, which served as sites for the switch experiments (Fig. 1). These concentration differences parallel previously observed gradients in DOC in the Hudson River (Findlay et al. 1996). DOC composition, as revealed by in vivo fluorescence (McKnight et al. 2001), also varied among sites; the lowest ratios of fluorescence to DOC were in the Wallkill (fluorescence = 1.82 ± 0.03 [SD] and DOC = 6.2 mg C l⁻¹) and the highest in NYC water (fluorescence =2.0 ± 0.07 and DOC = 3 mg C l⁻¹), implying less of a contribution of humic material in the latter. Inorganic nutrient concentrations also varied spatially, with much higher DIN and SRP concentrations in NYC (1.03 ± 0.2 mg N l⁻¹; 0.13 ± 0.05 mg P l⁻¹) than near Albany (0.31 ± 0.17 mg N l⁻¹; 0.008 ± 0.0003 mg P l⁻¹). Tributaries were more similar in DIN (0.4 to 0.6 mg N l⁻¹) but had larger differences in SRP (0.02 ± 0.01 mg P l⁻¹ for Mohawk; 0.04 ± 0.04 mg P l⁻¹ for Waterford; 0.006 ± 0.006 mg P l⁻¹ in Esopus; 0.062 ± 0.003 mg P l⁻¹ for Wallkill).

The relative amount of labile DOM differed among some of the locations we examined. Labile DOM was estimated from the decrease in DOC concentrations in experiments where we added a bacterial inoculum to water from the various locations (see below). Absolute loss of DOC (change in mgC l⁻¹) over the experimental period did not differ among water sources, but the percent loss was significantly higher (F<sub>5,40</sub> = 4.58; p = 0.002) for the Albany and NYC waters than for 3 (Waterford, Mohawk and Wallkill) of the 4 tributaries we examined. The relative amount of labile DOM from Albany and NYC was similar; 25 to 27 % of the initial DOC was depleted over the 3 to 4 d duration of the experiments compared with 9.6 to 14.2 % for the 3 tributaries. The labile DOM fraction at Albany and NYC was not significantly different from that of the 4th tributary, Esopus Creek (18.4 %).

Bacterial community structure, ectoenzyme activity and biomass production (leucine incorporation) were examined in unmanipulated bulk water at 3 locations in the main stem of the Hudson River and in 3 tributaries in late summer or fall of 1999, 2000, and 2001. Nearly all variables differed significantly, due to both location and time (p < 0.05; ANOVA). Ectoenzyme activity varied from 2-fold (N-acetylglucosaminidase) to nearly 6-fold (aminopeptidase) among the Hudson River locations and tributaries (Table 1). Concomitant with this variation, we observed differences in community structure as revealed by FISH.

The fraction of cells detectable by FISH (Eub338-positive cells) varied nearly 2-fold, due to location with the Albany and Waterford sites having the lowest (40 ± 12 %) and highest (76 ± 7 %) percentage, respectively. Of the 4 bacterial groups we examined, the most abundant group was either 1 of 2 proteobacterial subdivisions (alpha- or beta-proteobacteria) or the Cytophaga-like bacteria; the least abundant group was always the gamma-proteobacteria (Fig. 2), which usually accounted for <10 % of total abundance. Alpha-proteobacteria were quite abundant (45 % of total) and beta-proteobacteria relatively rare (ca. 5 %) in the Esopus tributary, which is surprising
since beta-proteobacteria often dominate freshwater systems (Glöckner et al. 1999). At the other extreme, beta-proteobacteria dominated the Wallkill tributary (on average 35% of total abundance), whereas alpha-proteobacteria were only ca. 10% of total prokaryotic abundance. In the entire data set, there was no significant correlation between alpha- and beta-proteobacterial abundances (see below).

### Upstream-downstream switch experiments

We hypothesized that some of the differences in ambient ectoenzyme activity and community structure observed in the Hudson River system were due to differences in DOM composition. To explore this hypothesis, we conducted ‘switch’ experiments in which a bacterial inoculum from one site was added to

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Waterford</th>
<th>Mohawk</th>
<th>Albany</th>
<th>Esopus</th>
<th>Wallkill</th>
<th>NYC</th>
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<tr>
<td>Alpha-glucosidase</td>
<td>3.66 ± 1.34</td>
<td>6.35 ± 1.84</td>
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<td>N-acetylglucosaminidase</td>
<td>12.0 ± 3.1</td>
<td>24.5 ± 7.9</td>
<td>19.2 ± 5.0</td>
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<td>Xylosidase</td>
<td>5.83 ± 1.61</td>
<td>6.99 ± 1.82</td>
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<td>10.22 ± 0.56</td>
<td>3.7 ± 1.65</td>
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<td>Phosphatase</td>
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<td>0.170 ± 0.005</td>
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<td>Aminopeptidase</td>
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<td>3.49 ± 0.30</td>
<td>1.09 ± 0.30</td>
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</table>

Table 1. Summary of in situ enzyme activities in the Hudson River system. Means (±SE) of rates measured in triplicates on 3 (New York City [NYC] and Albany) or 2 dates (all other sites). The 6 locations are arranged from the northernmost station on the left. Units for alpha-glucosidase, beta-glucosidase, N-acetylglucosaminidase and xylosidase are pmol l⁻¹ h⁻¹, whereas the units for the remaining enzymes are nmol l⁻¹ h⁻¹.

Fig. 2. Box and whisker plots of bacterial community structure (fluorescence in situ hybridization; FISH) at the locations used in the experiments (see Fig. 1). The data for Albany and New York are averaged over 3 yr, whereas the data presented for the other sites are from a single date (October 2000) with the statistics reflecting the within-sample variation of the FISH method. Wat = Waterford; Mo = Mohawk; Al = Albany; Es = Esopus; Wk = Wallkill; NYC = New York City
microbe-free water of another site. The 2 sites were at Albany (high allochthonous DOM) and NYC (lower terrestrial DOM), and 3 experiments were conducted in 1999, 2000, and 2001. These 2 sites significantly differed (p < 0.05; ANOVA) in ambient activity of most ectoenzymes (Table 1) and in the 2 most abundant bacterial groups, the alpha- and beta-proteobacteria (Fig. 2). Bacterial abundance at these 2 sites did not differ substantially, and any difference in abundance in the inoculum was small compared to growth during the experiments.

As with the in situ data, nearly all variables differed significantly due to the inoculum, source of water, and/or date of the experiment (Table 2). Both source of the water and of the inoculum affected ectoenzyme activity, with the exception of N-acetylglucosaminidase, which was not significantly affected by either the inoculum or the water. In addition, xylosidase activity was not affected by the water source, whereas 3 enzymes (alpha-glucosidase, esterase and trypsin) were not affected by source of the inoculum. Some of this enzyme activity may have been in the dissolved phase, but any difference in the initial dissolved enzyme activity would not affect our results because bacteria grew and enzyme activity increased several fold during the experiments. In short, 5 enzymes were affected by the water source, whereas 4 were affected by the inoculum source in the switch experiments (Table 2).

Principal component analysis (PCA) indicates that water source is the predominant factor driving variability in enzyme activity (Fig. 3A). There was a significant difference in PC1 scores among water sources (1-way ANOVA, p = 0.038) but no significant effect of inoculum (Fig. 3A). Enzyme activity in NYC water (regardless of inoculum) differed significantly from the Albany water source, with both NYC water treatments scoring lower on PC1 than the Albany water. There was no significant difference among water sources for scores on PC2, and the inoculum source had little effect on PCA scores; the reciprocal inocula overlapped on both axes. The PCA analyses suggest that the water was more important than the inoculum in shaping overall ectoenzyme activity.

Bacterial community structure, as revealed by FISH, also changed in the switch experiments. Table 3 summarizes the ANOVA analysis for the upstream-downstream switch experiment conducted during August/November of 1999, 2000 and 2001. The probability of a Type II error is given (p-values). The 3 independent variables include source of the water and inoculum, and date of the experiment. Eub338 is the general bacterial probe.
marizes the changes in all 3 experiments. The relative number of bacteria detectable by FISH increased in all 3 experiments; the number of Eub338-positive cells often increased by as much as 50% (Table 3) and approached 80%, regardless of the water and inoculum. Neither water nor the inoculum had a significant impact, however, on the relative number of Eub338-positive cells, although date of the experiment did (Table 2).

As with enzyme activity, source of water appeared to have a greater impact than the inoculum on bacterial community structure, although the influences varied among the 4 bacterial groups we examined. The water source had a significant effect on the alpha-proteobacterial abundance, but the inoculum source and experiment date did not (Table 2). Beta-proteobacteria were affected significantly by all 3 variables, whereas only the water source had a significant impact on gamma-proteobacterial abundance. In contrast, the Cytophaga-like bacteria were affected significantly by both water source and experiment date, but not by the inoculum. In short, the water source had a significant effect on all 4 bacterial groups, whereas the inoculum source had a statistically significant effect on only beta-proteobacterial abundance.

The PCA analysis suggested that sources of both the water and inoculum had roles in shaping the bacterial community growing up in these experiments. Scores on PC2 differed significantly (p < 0.001) among source waters (Fig. 3B), but the inoculum effect was not significant. There was a significant interaction term in the 2-way ANOVA (p = 0.001), with the Albany inoculum in NYC water significantly separated from the 2-way ANOVA (p = 0.056), whereas the other enzymes did not differ significantly (p > 0.05). There was no significant difference in the final abundance of the 4 bacterial groups when either Albany or NYC bacteria were added to NYC water (data not shown). Similarly, in the experiments with Albany water, ultimate activities for phosphatase and aminopeptidase differed significantly for the assemblages grown from the 2 inocula (NYC and Albany) differed significantly in beta-glucosidase, xylosidase, and peptidase activities and in leucine incorporation at the end of the experiment (Fig. 4; p < 0.05; ANOVA). The activity of N-acetylglucosaminidase was marginally significant in leucine incorporation at the end of the experiments (p = 0.056), whereas the other enzymes did not differ significantly (p > 0.05). There was no significant difference in the final abundance of the 4 bacterial groups whether ectoenzyme activity and bacterial community structure developing in incubations with bacteria from 2 different endmembers added to the same water converged to the same levels of activity and community structure. In the NYC water experiments, the bacterial assemblages resulting from the 2 inocula (NYC and Albany) differed significantly in beta-glucosidase, xylosidase, and peptidase activities and in leucine incorporation at the end of the experiment (Fig. 4; p < 0.05; ANOVA). The activity of N-acetylglucosaminidase was marginally significant (p = 0.056), whereas the other enzymes did not differ significantly (p > 0.05). There was no significant difference in the final abundance of the 4 bacterial groups when either Albany or NYC bacteria were added to NYC water (data not shown). Similarly, in the experiments with Albany water, ultimate activities for phosphatase and aminopeptidase differed significantly for the assemblages grown from the 2 inocula (p < 0.05; ANOVA); activity of the other enzymes was the same at the end of the experiments in which either Albany or NYC bacteria were added to Albany water (data not shown). Both leucine and thymidine incorporation rates also differed significantly for the 2 inocula.

Table 3. Changes in community structure as revealed by fluorescence in situ hybridization (FISH) in bacteria (‘inoculum’) added to filter-sterilized water (‘water’) from the indicated locations. The initial relative abundance in each bacterial group observed in the inoculum was subtracted from the final relative abundance (fraction of total direct counts). A positive number indicates that the group increased in abundance relative to the inoculum; a negative number indicates that it decreased. Eub338 is the general bacterial probe. Alpha, beta, and gamma are subdivisions of the proteobacteria. Cytophaga refers to a complex group of microbes identified by the CF319a probe often referred to as the Cytophaga-Flavobacter or Cytophaga-like group. AL: Albany; NYC: New York City. SE in parentheses

<table>
<thead>
<tr>
<th>Date (mm/yy)</th>
<th>Water</th>
<th>Inoculum</th>
<th>Change in community composition (% of total abundance)</th>
<th>Eub338</th>
<th>Alpha</th>
<th>Beta</th>
<th>Gamma</th>
<th>Cytophaga</th>
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</table>
cantly with inocula for the Albany water experiments. Unlike the NYC water experiments, the abundance of both alpha- and beta-proteobacteria remained different in the Albany water experiments with both inocula, even after 3 to 4 d of incubation (data not shown).

Enzyme activity in incubations with the Albany inoculum was generally higher than for the NYC inoculum incubations, even for Albany bacteria added to NYC water (Fig. 4). Likewise, leucine incorporation rates were 5-fold higher in 2 of 3 experiments for Albany bacteria added to NYC water (Fig. 4); thymidine incorporation rates were 2-fold higher for Albany bacteria than for NYC bacteria added to Albany water in 2 of 3 experiments (data not shown). The higher level of activity for the Albany bacteria reflects the initial difference between the 2 sites; rates of most of the enzymes and leucine incorporation were higher in the ambient water at Albany than at NYC (Table 1). In short, some differences in enzyme activity, biomass production, and community structure remained even after several days of incubation of different inocula added to the same water.

Tributary experiments

To explore further the role of water source (and thus DOM characteristics) in shaping bacterial activity and community structure, we added a single inoculum from the Hudson River main stem at Norrie Point to water from 3 tributaries and 1 location in the upper Hudson above the tidal influence (Waterford). Of the 8 ectoenzymes, 3 varied significantly due to the water source: beta-glucosidase, esterase, and trypsin (p < 0.001; ANOVA). Although the impact on individual ectoenzymes was relatively minor, the overall effect was large, according to the PCA analyses (Fig. 5A) with significantly different scores on PC1 (ANOVA, p = 0.02). The first PC score for the Waterford source water (upper Hudson, above tidal portion) was significantly different from that of the 3 other locations, and the score for the Mohawk source water was significantly different from the Wallkill (Fig. 5A). None of the sources had significantly different scores on the second principal component.

The water source also had a large overall impact on bacterial community structure, as revealed by FISH,
although the absolute change in most of the bacterial groups was small. The beta-proteobacteria were the most abundant group in these experiments (Fig. 6), but water source did not have a significant effect on their abundance ($p > 0.05$; ANOVA). Alpha-proteobacteria were the least abundant group (<5% of total abundance) and were also not affected by the water source. The water source had a marginally significant effect on gamma-proteobacterial abundance ($p = 0.055$; ANOVA), and a post-hoc test revealed that abundance of this group in the Esopus and Waterford waters differed significantly ($p = 0.017$; Tamhane’s test). Cytophaga-like bacteria, which, along with gamma-proteobacteria, were second in abundance to beta-proteobacteria (Fig. 6), were significantly affected by the water source ($p < 0.001$; ANOVA). The post-hoc test indicated that final Cytophaga-like bacterial abundance in these experiments differed significantly in water from the Esopus and Mohawk; abundance of this group was higher and lower, respectively, than the abundances in water from the other tributaries ($p < 0.001$).

While the effect of water source on the abundance of individual bacterial groups was rather minimal in this tributary experiment, the overall effect was large, as revealed by significant differences in PC2 scores ($p = 0.005$). The Mohawk and Waterford sources differed significantly in PC2 scores from the Wallkill and Esopus sources (Fig. 5B), although none of the water sources had significantly different PC1 scores. The PCA analyses indicated that the community growing up in the Esopus water was similar to that in the Wallkill water, and both differed greatly from the Waterford and Mohawk waters (Fig. 5B).

**Ectoenzyme activity and bacterial community structure**

We used 2 statistical approaches to explore the relationships between ectoenzyme activity and bacterial community structure. These analyses were applied to
the entire data set (ambient and experiments) in order to increase our sample size and to help ensure the generality of our results. Not unexpectedly, the activities of several ectoenzymes were positively correlated with each other (Table 4). A notable exception is trypsin activity, which was not significantly correlated with any of the other ectoenzymes. It is surprising how few of the ectoenzymes correlated significantly with the 2 measures of biomass production (leucine and thymidine incorporation).

Only a few of the bacterial groups we examined covaried significantly with each other (Table 4). Two of the proteobacterial subdivisions, alpha- and gamma-proteobacteria, correlated significantly (r = 0.477; p < 0.001), whereas the beta-proteobacteria strongly covaried with Cytophaga-like bacteria (r = 0.775; p < 0.001). There was no significant correlation between alpha- and beta-proteobacterial abundances (r = −0.032; p > 0.05).

Several of the ectoenzymes and the 2 measures of bacterial production covaried significantly with the abundance of 1 or more of the 4 bacterial groups examined here (Table 4). The abundance of alpha-proteobacteria was positively correlated with 5 of the 8 ectoenzymes; alpha-glucosidase, N-acetylglycosaminidase, and esterase correlations all exceeded 0.5 (Table 4). Of the 2 ectoenzymes that Cytophaga-like bacteria were significantly correlated with (Table 4), the strongest correlation was with N-acetylglycosaminidase activity (r = 0.503; p < 0.0001). Gamma-proteobacterial abundance was significantly correlated with only esterase activity (r = 0.399; p < 0.001).

Leucine incorporation was positively correlated with the abundance of alpha- and gamma-proteobacteria, but thymidine incorporation was only negatively correlated with beta-proteobacteria.

Perhaps the most interesting correlation is that between beta-proteobacteria and phosphatase activity. This bacterial group was significantly correlated with only 1 ectoenzyme (phosphatase), but the correlation coefficient was the highest of any of the pairwise comparisons between the bacterial groups and ectoenzyme activity (r = 0.718). The correlation appears to be determined by 2 groups of samples, one with low phosphatase activity and relative betaproteobacterial abundance, and another with high values (Fig. 7). With only 2 exceptions, the samples with high phosphatase activity and high relative beta-proteobacterial abundance were from the switch experiment with Albany water, to which Albany or NYC bacteria were added.

To further explore the relationship between community structure and ectoenzyme activity, we performed a multi-variate linear regression using the various ectoenzymes as the dependent variable (one at a time) and the 4 bacterial groups as the independent variables (all together). Nearly all of the regression models explained a significant amount of variation, although only a few explained more than 40% for a particular ectoenzyme activity (Table 5). As revealed by the correlation analysis, beta-proteobacterial abundance was the sole bacterial group accounting for (in a positive sense) a significant amount of the variation in phosphatase activity; alpha- and gamma-proteobacteria

<table>
<thead>
<tr>
<th>α-glcase</th>
<th>β-glcase</th>
<th>NAGase</th>
<th>Xylosidase</th>
<th>P-ase</th>
<th>Pep-ase</th>
<th>Est-ase</th>
<th>Trypsin</th>
<th>Leu</th>
<th>TdR</th>
<th>Eub338</th>
<th>Alpha</th>
<th>Beta</th>
<th>Gamma</th>
<th>CF-like</th>
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</thead>
<tbody>
<tr>
<td>α-glcase</td>
<td>1</td>
<td>0.813*</td>
<td>0.523*</td>
<td>0.678*</td>
<td>−0.255</td>
<td>0.596*</td>
<td>0.792*</td>
<td>−0.137</td>
<td>0.553*</td>
<td>−0.326</td>
<td>0.054</td>
<td>0.532*</td>
<td>−0.291</td>
<td>0.323</td>
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<td>0.790*</td>
<td>−0.314</td>
<td>0.448*</td>
<td>0.653*</td>
<td>−0.267</td>
<td>0.419*</td>
<td>−0.313</td>
<td>0.210</td>
<td>0.479*</td>
<td>−0.213</td>
<td>0.362</td>
<td>0.062</td>
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<tr>
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<td>Est-ase</td>
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<td>0.560*</td>
<td>−0.376</td>
<td>−0.089</td>
<td>0.335*</td>
<td>−0.289</td>
<td>0.399*</td>
<td>−0.102</td>
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<td>0.531*</td>
<td>0.138</td>
<td>0.471*</td>
<td>0.004</td>
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<td>TdR</td>
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<td>−0.193</td>
<td>−0.424*</td>
<td>−0.037</td>
<td>−0.356</td>
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<td>Eub338</td>
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<td>0.455*</td>
<td>0.374</td>
<td>0.362</td>
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<tr>
<td>Alpha</td>
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<td>−0.032</td>
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<td>−0.005</td>
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<tr>
<td>Beta</td>
<td>1</td>
<td>−0.097</td>
<td>0.775*</td>
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Table 4. Correlation coefficients for the entire data set. Missing values were eliminated list-wise, resulting in N = 57. *p = 0.05/15 = 0.003 (or better), using the Bonferroni adjustment. Correlation coefficients > 0.5 in bold. Abbreviations: α-glcase = alpha-glucosidase; β-glcase = beta-glucosidase; NAGase = N-acetylglycosaminidase; P-ase = phosphatase; Pep-ase = aminopeptidase; Est-ase = esterase; Eub338 = fraction of all cells hybridized by the general bacterial probe; Alpha, Beta, and Gamma = fraction of cells in these 3 proteobacterial subdivisions. CF-like refers to the Cytophaga-Flavobacterium cluster. Fluorescence in situ hybridization (FISH) data were arcsine transformed before analysis.
had weak, negative relationships with phosphatase activity. The regression model explained over 60% ($r^2 > 0.6$) of the variation in phosphatase activity (Table 5).

The abundance of *Cytophaga*-like bacteria, however, was the dominant variable in the regression models for 3 other ectoenzymes: beta-glucosidase, N-acetylglucosaminidase, and xylosidase (Table 5). In particular, for N-acetylglucosaminidase, the standardized coefficient for *Cytophaga*-like bacterial abundance (0.878) was over twice that of the next most-important bacterial group (0.383). The regression model with the 4 bacterial groups explained 61% of the variation in N-acetylglucosaminidase activity (Table 5). The alpha-proteobacterial abundance was second in importance for these ectoenzymes, but it was the dominant variable in explaining the variation in alpha-glucosidase activity. The abundance of beta-proteobacteria had a negative effect on all ectoenzymes, except phosphatase activity, according to this analysis (Table 5). This regression analysis implies an important role for *Cytophaga*-like bacteria in determining the activity of several ectoenzymes.

Community structure did not explain any significant variation in the activity of the 2 proteases, aminopeptidase and trypsin, the only 2 enzymes for which community structure appears to have no role (Table 5).

**DISCUSSION**

Concentrations and composition of DOM, bacterial community structure and, undoubtedly, other factors interact in complicated ways to determine ectoenzymatic activity, which drives DOM hydrolysis and eventually mineralization of DOM by heterotrophic bacteria in aquatic ecosystems. Our approach for exploring these issues was to examine ectoenzyme activity and community structure in experiments with water and inocula from locations in the Hudson that differ in DOM concentrations and composition. Although the water sources differed in other respects, DOM was likely the main factor affecting bacteria in these experiments. Inorganic nutrients were probably not important factors, because concentrations were sufficiently high to prevent N or P limitation and because bacterial growth is primarily limited by the supply of organic carbon in the Hudson River (Roland & Cole 1999). The impact of particulate organic carbon and detritus was

Table 5. Linear regression analysis of the contribution of the major bacterial groups (alpha-, beta, and gamma-proteobacteria and *Cytophaga*-like bacteria [CF]) to the variation in bacterial production (thymidine [Tdr] and leucine [Leu] incorporation) and enzyme activity. The entire data set was used in the data analysis. 'NS' indicates the coefficient is not statistically significant ($p > 0.05$). Standardized coefficients measure the contribution by the bacterial group to explaining variation in the dependent variables. Parameters higher than 0.60 are in **bold**. NS: not significant

<table>
<thead>
<tr>
<th>Dependent variables</th>
<th>Alpha</th>
<th>Standardized coefficients</th>
<th>Adjusted</th>
<th>$r^2$</th>
<th>$p$</th>
</tr>
</thead>
<tbody>
<tr>
<td>TdR incorporation</td>
<td>NS</td>
<td>-0.388</td>
<td>NS</td>
<td>0.165</td>
<td>$9.2 \times 10^{-3}$</td>
</tr>
<tr>
<td>Leu incorporation</td>
<td>0.382</td>
<td>0.436 0.281 NS</td>
<td>NS</td>
<td>0.353</td>
<td>$5.8 \times 10^{-4}$</td>
</tr>
<tr>
<td>Alpha-glucosidase</td>
<td>0.436</td>
<td>-0.449 NS</td>
<td>NS NS</td>
<td>0.317</td>
<td>$2.6 \times 10^{-3}$</td>
</tr>
<tr>
<td>Beta-glucosidase</td>
<td>0.307</td>
<td>-0.561 0.262 0.536</td>
<td>0.878</td>
<td>0.606</td>
<td>$5.3 \times 10^{-12}$</td>
</tr>
<tr>
<td>N-acetylglucosaminidase</td>
<td>0.383</td>
<td>-0.450 0.227 0.878</td>
<td>NS</td>
<td>0.624</td>
<td>$1.3 \times 10^{-12}$</td>
</tr>
<tr>
<td>Phosphatase</td>
<td>-0.183</td>
<td><strong>0.872</strong> -0.183 NS</td>
<td>NS</td>
<td>0.447</td>
<td>$7.3 \times 10^{-4}$</td>
</tr>
<tr>
<td>Xylosidase</td>
<td>0.312</td>
<td>-0.535 0.271 <strong>0.768</strong></td>
<td>NS NS</td>
<td>0.332</td>
<td>$1.4 \times 10^{-5}$</td>
</tr>
<tr>
<td>Esterase</td>
<td>0.415</td>
<td>-0.387 NS</td>
<td>NS</td>
<td>0.080</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>Aminopeptidase</td>
<td>NS</td>
<td>-0.402 NS</td>
<td>NS NS</td>
<td>0.043</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>Trypsin</td>
<td>NS</td>
<td>NS NS NS</td>
<td>NS</td>
<td>&gt;0.05</td>
<td>&gt;0.05</td>
</tr>
</tbody>
</table>
not examined here, although this form of carbon could impact the structure of in situ bacterial communities (DeLong et al. 1993) and of the inocula for our experiments.

Our experimental design (2 to 3 d incubations in bottles with diluted microbial assemblages) has limitations, such as possibly favoring the growth of microbes quite different from the initial community. Both community structure and enzymatic activity changed from initial values, even in incubations with the water and inoculum from the same source (the ‘controls’). These changes complicate attempts to understand the impact of terrestrial DOM on community structure and enzyme activity, but the comparison among DOM and inoculum sources still allowed examination of the relative strength of these factors in affecting community structure and activity. Even if some of the changes in controls were due to bottle effects, and even though it is difficult to relate effects observed in bottle incubations to in situ processes, the data can still be used to explore interactions between DOM and bacterial community structure, and to examine basic questions about the controls on activity and community composition in the Hudson.

Our experiments illustrate the impact of DOM on bacterial community structure, and suggest that community structure (at the phylogenetic level we assayed) did not constrain the activity of some of the ectoenzymes examined here. Activity of some ectoenzymes of communities derived from disparate inocula converged to similar rates, even when grown in the same water from either NYC or Albany. Thus, the water and, presumably, DOM generally overrode differences in initial community structure in determining final enzyme activities. The PCA analysis indicated that the water, i.e. DOM, affected community structure more strongly than it affected ectoenzyme activity, since there were significant differences in both principal components for the FISH data but only in one component for the ectoenzyme data in the tributary experiment. These data suggest that the activity of some ectoenzymes and, thus, the hydrolysis of some DOM components may not depend strongly on bacterial community structure.

However, other analyses point to a dependence of some ectoenzymes on bacterial community structure. The rates of some ectoenzymes did not converge when NYC and Albany inocula were added to either NYC or Albany water. The rates of some ectoenzymes (and of leucine and thymidine incorporation) were higher in incubations with Albany bacteria than with NYC bacteria, even in NYC water. This difference parallels the initial difference in activity of the in situ bacterial communities; like in the experiments, metabolic activity of Albany bacteria was generally higher than that of NYC bacteria in situ. Since the 3 to 4 d incubations should have been long enough for any phenotypic response by the bacterial community, the persistent difference in activity in our incubations suggest that bacterial communities are not equally capable of expressing all ectoenzymes necessary for hydrolyzing DOM. Changes in community structure may sometimes be required before new DOM is hydrolyzed, and perhaps ultimately mineralized.

Other data further illustrate possible impacts of community structure on ectoenzyme activity. We found several significant correlations between enzyme activity and the abundances of the 4 bacterial groups we examined, with the strongest being between beta-proteobacterial abundance and phosphatase activity. If beta-proteobacteria do in fact have more active phosphatases, as suggested by the correlation analysis, it may explain their high abundance in freshwater systems (Glöckner et al. 1999), which are often limited by phosphorus (Wetzel 2001). Also, the multi-variate regression models using bacterial community structure explained a significant amount of variation in the activity of all ectoenzymes, except the 2 proteases. Although it is difficult to extrapolate from these statistical analyses to causal relationships, the data imply that the major bacterial groups we examined differ in ectoenzyme activity. Other evidence also points to differences in DOM uptake among the bacterial groups we examined, that is, the alpha-, beta-, and gamma-proteobacteria and the Cytophaga-Flavobacterium cluster. These 4 groups appear to differ in their response to changing DOM supply and composition in field studies and experiments (Rossello-Mora et al. 1999, Eilers et al. 2000a, 2001). Also, uptake of several DOM components differs among these 4 bacterial groups (Cottrell & Kirchman 2000, Malmstrom et al. 2004). These differences may explain why alpha- and beta-proteobacteria and Cytophaga-like bacteria responded differently in mesocosms that varied in primary production (Horner-Devine et al. 2003).

It is surprising that the relationships between DOM and the major bacterial groups appear to differ and, more specifically, that the relationship between abundance and ectoenzyme activity varied among the bacterial groups we examined. Each of these groups is potentially diverse, consisting of many ‘species’ or ribotypes (Giovannoni & Rappé 2000). One may expect that these bacterial groups would be sufficiently large and diverse so that all ectoenzymes are potentially expressed by a single group. Consequently, enzyme activity should not differ between broad bacterial groups, even if individual ribotypes do (Martinez et al. 1996). It is possible, however, that each broad group may actually be represented by a small number of dominant ribotypes in any given environmental
sample at a particular time. Differences in ectoenzymatic activity at the ribotype level would become apparent even when measured at the division and sub-division level if the bacterial groups in our samples were not very diverse. In support of this hypothesis, preliminary data from denaturing gradient gel electrophoresis (DGGE) analyses (Muyzer & Smalla 1998) indicate that the bacterial communities had on the order of 10 dominant ribotypes in our experiments. If divided equally among the 4 bacterial groups examined here, the DGGE and FISH data suggest that each group is represented by only ca. 2 to 3 dominant ribotypes, although several other less abundant ribotypes (<10%) may also be present.

Regardless of exactly how bacterial community structure is linked to DOM use, the existence of any link suggests that community structure could have an impact on DOM mineralization. The nature and extent of this impact are likely to vary with the factors that control community structure, and the effect may be substantial when heterotrophic bacteria are limited by factors other than DOM. These other factors include inorganic nutrients such as phosphate and iron (Pakulski et al. 1996, Cotner et al. 1997), grazing (Šimek et al. 1997, Suzuki 1999, Riemann et al. 2000, Pernthaler et al. 2001, Beardsley et al. 2003, Cottrell & Kirchman 2004), viral lysis (Fuhrman 2000), temperature (Pomeroy & Wiebe 2001), and other abiotic factors (del Giorgio & Bouvier 2002). Previous studies have also shown how factors other than DOM can affect community structure (Gasol et al. 2002, Šimek et al. 2003). If a bacterial group specializing in using some DOM components is limited by one of these other factors, then degradation of those components may also be affected indirectly.

Even the composition of a DOM-controlled bacterial community could have short-term (days) impacts on DOM degradation when DOM concentrations or composition change enough to induce changes in bacterial metabolism. Our results suggest that not all bacterial communities are equally capable of maximal ecto-enzyme activity and perhaps of effective DOM mineralization when the DOM pool changes. Even after several days, the activity of some ectoenzymes was not as high for the NYC inoculum, for example, as observed for the Albany inoculum in the same water. For those ectoenzymes, a change in the bacterial community appeared necessary before hydrolysis of those targeted DOM components could be maximal. These results suggest that community structure could control degradation of some DOM components.

Such control by bacterial community structure is one level at which DOM mineralization is regulated, with the other extreme being the kinetic response of already synthesized enzymes (Sinsabaugh & Findlay 2003). Rates of constitutive enzymes can respond quite rapidly, on the order of minutes to hours, whereas days may be required for a rare ribotype to increase sufficiently in abundance before appreciably affecting DOM mineralization at the community level (Findlay 2003). In between these 2 extremes is induction and synthesis of new enzymes, which occur within hours. All of these levels undoubtedly contribute to regulating DOM mineralization, and the contributions probably vary with the seasons and habitats. How DOM mineralization is regulated will set the time scale for DOM mineralization and biomass production, which in turn will affect storage of C in DOC and transport of DOM from more productive to less productive regions, such as from rivers and estuaries to the oceans.

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LITERATURE CITED

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