INTRODUCTION

Polycyclic aromatic hydrocarbons (PAHs) are toxic and often also carcinogenic organic contaminants that enter the marine environment by incomplete combustion of fuels, industrial pyrolysis, or spills of crude oils and refined products. The PAHs are readily accumulated in marine sediments due to high hydrophobicity and very low degradation rates. Microbial mineralization of PAHs is considered the most important process leading to PAH decontamination and permanent removal from the environment (Neff 2002). PAH-degrading bacteria are present in most PAH-polluted environments although in very different numbers.

Effects of the polychaetes Arenicola marina and Nereis diversicolor on microbial pyrene mineralization

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ABSTRACT: The effects of 2 polychaetes, Nereis diversicolor and Arenicola marina, on the microbial mineralization of the organic contaminant pyrene, a polycyclic aromatic hydrocarbon (PAH), were followed over 44 d. We also examined whether the effect of the polychaetes was caused by enhanced oxygen supply, altered pyrene bioavailability and/or a changed abundance or activity of pyrene-degrading bacteria. The presence of polychaetes enhanced microbial pyrene mineralization by 180 to 200 % compared with defaunated sediment. Collectively, the replicates of the different treatments showed that mineralization rates were positively correlated with the amount of oxidized sediment, which comprised mainly the 3 mm surface layer and zones around burrows (burrow sediment). The biogenic sediment structures had similar mineralization potential and abundance of pyrene-degrading bacteria as surface sediments. Pyrene mineralization potential in bulk (reduced and presumably anoxic) sediment was significantly lower than for surface and burrow sediments. However, when the bulk sediments were oxidized, mineralization rates increased rapidly. Collectively, these data indicate that oxygen availability controlled pyrene mineralization in these experiments. On the other hand, the presence of the polychaetes significantly reduced the bioavailability of pyrene to the microbial degraders. Pyrene bioavailability in burrow sediment was always lower than the bioavailability in both surface and bulk sediments. In addition, N. diversicolor and especially A. marina decreased the bioavailability of pyrene in surface sediments compared with that of surface sediments in the non-bioturbated control. In conclusion, these polychaetes enhanced microbial pyrene mineralization significantly and this enhancement seemed to be caused by the increased oxygen supply due to burrow construction and irrigation. In contrast, these worms decreased pyrene bioavailability and, hence, counteracted to some extent the stimulating effect of irrigation.

KEY WORDS: Oxygen · Bioavailability · PAH · Biogenic structures · Bioturbation · Mineralization potential · MPN
depending on factors such as pollution history (Johnsen & Karlson 2005), redox status and nutrient availability (Carmichael & Pfaender 1997, Joner et al. 2002).

Oxygen is a key variable for microbial PAH degradation and mineralization, determining both the degradation rate (McNally et al. 1999, Lei et al. 2005), pathway (Meckenstock et al. 2004) and the composition of bacteria involved in PAH degradation (Roslev et al. 1998, Eriksson et al. 2003). In general, aerobic PAH degradation is orders of magnitude higher than degradation under both nitrate- and sulphate-reducing conditions (Coates et al. 1997, Rockne & Strand 1998). PAH degradation may also be limited by the low bioavailability. The amount, quality and diagenetic status of the sediment organic matter is known to affect PAH dissolution rates (Huang & Weber 1997, Kukkonen et al. 2003) and, hence, the bioavailability of PAHs (Bosma et al. 1997, Johnsen et al. 2005).

Bioturbating infauna influence the availability of oxygen by their burrow construction and irrigation, which extends the sediment–water interface and oxidizes sediments at depth (e.g. Wenzhofer & Glud 2004, Timmermann et al. 1995), allowing aerobic processes to occur in otherwise reduced sediments (Nielsen et al. 2004). Irrigated burrows often show enhanced activity of biogeochemical processes (Kristensen et al. 1985, Mayer et al. 1995, Papaspyrou et al. 2006) and both bacteria and meiofauna are often more abundant in burrows and tubes compared with the surrounding sediment (Reise 1981, Papaspyrou et al. 2006). The significance of burrow structures as sites of enhanced capacity for microbial PAH mineralization have also been documented (Chung & King 1999, 2001, Granberg et al. 2005).

The stimulating effect of sediment-dwelling infauna on microbial processes and abundance is well recognised (Kristensen 2000). The few studies examining the effects of macrobenthos on total microbial PAH mineralization indicate that this stimulation also applies to PAH degradation (Bauer et al. 1988, McElroy et al. 1990, Christensen et al. 2002). However, decreased PAH degradation in the presence of polychaetes has also been observed due to transport of PAH-containing material from the surface into deeper anoxic zones (Kure & Forbes 1997, Banta & Andersen 2003).

The aim of the present study was to examine the effect of 2 common polychaetes, Arenicola marina and Nereis diversicolor, on the microbial mineralization of the tetracyclic model PAH pyrene in sediment, and furthermore, to determine whether an observed effect was caused by increased oxygen availability, changed microbial pyrene bioavailability, changed activity or abundance of pyrene degraders, or combinations of these factors.

**MATERIALS AND METHODS**

The experiment was conducted in 3 steps: (1) a pre-exposure phase where pyrene contaminated sediment was incubated for 25 d to allow growth of pyrene degraders, (2) an exposure phase and (3) an incubation phase. During the exposure phase, sediment was exposed to different treatments (see Table 1) consisting of sediment contaminated with either pyrene (+) or 14C-pyrene (++ combined with the addition of either Arenicola marina (Am) or Nereis diversicolor (Nd), or no fauna (C). During the exposure phase, 14CO2 production was measured in treatments containing 14C-pyrene. After the exposure phase, sediment samples were collected from surface, burrow and bulk sediment from each treatment and transferred to 14C-respirometric flasks. Sediment samples originating from non-radioactive pyrene treatments were incubated with freshly added 14C-pyrene to measure the microbial pyrene mineralization potential, whereas sediment samples originating from treatments with 14C-pyrene were incubated with an inoculum of known pyrene degraders to measure the bioavailability of pyrene in the different sediment compartments.

**Collection of sediment and polychaetes.** Sediment and polychaetes were collected at a pristine, near-coastal sandflat in Roskilde Fjord, Denmark (55° 41’ 51.9” N, 12° 5’ 47.4” E). The fjord is brackish and, at the time of sampling, salinity was measured as 13‰. The water depth at the sampling site was 0.3 m during sample collection, but varied between 0 and 1.5 m depending on wind speed and direction. Sediment was collected in March from the top 10 cm of the seafloor, sieved (1 mm) in situ to remove stones and macrofauna, and frozen for 1 wk to kill meiofauna and juvenile macrofauna. The sediment was characterised as fine to medium sand (dominated by 125 to 500 µm grain size; data not shown) with a mean porosity of 0.35 and an organic matter content of ~1.5%. Polychaetes were collected in April by digging and kept in aquaria with sediment and water from the sampling site until use.

**Table 1. Exposure phase treatments.** Microcosms contained sediment contaminated with either pyrene (+) or pyrene and 14C-pyrene (++). Total nominal pyrene concentration was 10 µg kg⁻¹ in all treatments. Each microcosm contained either Arenicola marina (Am), Nereis diversicolor (Nd) or no fauna (control sediment, C). Four replicate microcosms were used in each treatment (i.e. n = 4)

<table>
<thead>
<tr>
<th></th>
<th>Pyrene</th>
<th>Pyrene + 14C-pyrene</th>
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<tbody>
<tr>
<td>No fauna (control)</td>
<td>C+</td>
<td>C++</td>
</tr>
<tr>
<td>Arenicola marina</td>
<td>Am+</td>
<td>Am++</td>
</tr>
<tr>
<td>Nereis diversicolor</td>
<td>Nd+</td>
<td>Nd++</td>
</tr>
</tbody>
</table>
**Sediment contamination.** Contaminated sediment was prepared by mixing (24 h) wet sediment with pyrene dissolved in acetone to a final concentration of 20 mg kg⁻¹. The contaminated sediment was then distributed in boxes in an approximately 1 cm layer covered with water that was continuously aerated with air stones. The boxes were incubated for 25 d at 12°C.

After this pre-exposure phase, the pyrene contaminated sediment was further mixed (24 h) with clean sediment and either [¹⁴C-4,5,9,10]-pyrene (58.7 mCi mmol⁻¹, purity >98%, Sigma-Aldrich) or nonradioactive pyrene. The specific activity of the [¹⁴C]-pyrene labelled sediment was 5000 disintegrations per minute (dpm) g⁻¹ and the total nominal concentration of pyrene was 10 mg kg⁻¹ in all treatments.

**Experimental set-up.** The experiment was set up in polyacrylic core tubes. The dimensions of the core tubes used in treatments with *Nereis diversicolor* and controls (groups Nd and C) were 5 cm in diameter (Ω) and 30 cm high (I). These core tubes were filled with 1.3 kg of contaminated sediment after which 100 ml of filtered (0.2 μm) seawater, diluted with deionized water to a salinity of 15‰, was carefully added to each core. One *N. diversicolor* (0.47 ± 0.09 g wet weight [mean ± SD]) was added to each core in group Nd and Nd++, equivalent to a density of ~500 worms m⁻². Larger core tubes (Ω = 8.2 cm, I = 40 cm) were used for *Arenicola marina* (Am) due to greater space requirements needed for its feeding strategy and lower natural density. These core tubes were filled with 3.2 kg of contaminated sediment overlaid by 250 ml of filtered seawater. One *A. marina* (1.7 ± 0.5 g wet weight) was added to each core, equivalent to a density of ~190 worms m⁻². Worms that failed to burrow within 15 min were replaced. The cores were incubated for 44 d at 12°C in the dark to prevent photo-oxidation of pyrene and photoautotroph fixation of [¹⁴CO₂] (produced during microbial [¹⁴C]-pyrene mineralization).

To trap [¹⁴CO₂] cores with [¹⁴C]-pyrene contaminated sediment (i.e. groups C++, Am++, Nd++) were sealed with rubber stoppers, and out-flowing air was passed through 2 serially connected CO₂ traps consisting of gas-wash bottles each containing 10 ml of 0.5 M NaOH. Trap efficiency was >92% (tested with [¹⁴C]-NaHCO₃). Cores with non-radioactive pyrene were also sealed with stoppers, but the out-flowing air was not connected to CO₂ traps.

The radioactivity in CO₂ traps and overlying water was measured every 4th day when the CO₂ trapping tubes and overlying water was changed. Radioactivity in the CO₂ traps was quantified by liquid scintillation counting on a Wallac LKB 1219 counter using 4 ml of sample and 10 ml Packard Ultima Gold XR. Radioactive CO₂ in overlying water was measured before each water change by transferring 20 ml of overlying water to a closed vessel containing a CO₂ trap with 5 ml of 0.5 M NaOH. Water was acidified with 1 ml of 1 M HCl and after 4 h, 4 ml NaOH from each trap was transferred to 10 ml Packard Ultima Gold XR and the radioactivity from [¹⁴CO₂] was quantified.

**Collection of surface, bulk and burrow sediment.** After incubation, the cores were sectioned into different sediment pools containing surface sediment, anoxic bulk sediment, burrow sediment (only Am and Nd groups) and oxidized non-burrow sediment when observed. Surface sediment was collected by cutting off the top 3 mm of each sediment core. In all cores, this layer was completely oxidized. Burrow sediment was defined as the inner 2 mm of sediment lining the burrows and collected by carefully scraping the burrows with a spatula. When oxidized non-burrow sediment was observed during sectioning, that sediment was also collected. In the present study, oxidized sediment is defined by the light grey–brown colour (i.e. the presence of oxidized metals). Although oxygen is not necessarily present in oxidized sediments, oxidation of metal species is ultimately linked to molecular oxygen through a series of coupled oxidation–reduction reactions and we use oxidized sediment as an indicator of sediment that is continuously or periodically subjected to oxygen. Bulk sediment, which was almost black, was defined as reduced (and presumably anoxic) sediment.

**Microbial pyrene mineralization potential.** Pyrene mineralization potential was measured as the ability to mineralize freshly added [¹⁴C]-pyrene under oxic conditions in surface, bulk and burrow sediment samples collected from treatments without radioactive pyrene (C+, Nd+ and Am+). Sediment samples (3 to 5 g) were transferred to Erlenmeyer flasks coated with [¹⁴C]-pyrene (0.07 μg, 0.02 μCi) and 5 ml of seawater was added. The sediment slurries were incubated on an orbital shaker (60 revolutions [rev.] min⁻¹) in the dark for 99 d. The [¹⁴CO₂] was produced was collected in 5 ml of 0.5 M NaOH contained in 6 ml plastic vials suspended from the silicone stoppers and quantified by mixing 4 ml of the NaOH with 6 ml of Ultima Gold scintillation cocktail and counted. Trapping efficiency of the incubation system was >95% (tested with [¹⁴C]-NaHCO₃).

**Microbial pyrene bioavailability.** The bioavailability of [¹⁴C]-pyrene aged during the exposure phase (44 d) was measured in surface, burrow and bulk sediment samples from treatments C++, Nd++ and Am++ using the set-up described for mineralization potential with the following changes. Non-coated Erlenmeyer flasks with sample material were inoculated with 2 bacterial pyrene degraders (DSM 44346: Mycobacterium Frederiksborgense Fan9 and DSM 7251: Mycobacterium vanbaalenii PYR-1; 10⁶ and 10⁷ CFU [colony forming units] g⁻¹ sediment, respectively) so that
pyrene bioavailability, and neither pyrene degradation capacity nor oxygen, was the limiting factor for pyrene mineralization. The inoculum was grown in Luria-Bertani (LB) medium and washed twice in MgSO$_4$ (50 mM) before addition to the sediment. Bioavailability was measured both as the flux of pyrene from the sediment into the bacteria and as the amount of bioaccessible pyrene, i.e. the total amount of pyrene that becomes bioavailable over time (Reichenberg & Mayer 2006). The flux of bioavailable pyrene was estimated as the mineralization rate of aged 14C-pyrene immediately after the addition of the pyrene degraders, whereas the total amount of bioavailable pyrene (bioaccessibility) was determined as the total cumulative 14CO$_2$ production at the end of the incubation period.

**Microbial populations.** The most probable number (MPN) of aerobic bacteria growing on pyrene in surface and burrow sediments was determined using a microplate method based on the respiration indicator WST-1 (Johnsen et al. 2002) modified for marine conditions. In short, microbial cells were extracted from 5 g (wet weight) of sediment with 45 ml of pyrophosphate buffer (1.2 mM tetrasodium pyrophosphate, pH 7.0, salinity 15‰). The sediment–buffer suspension was shaken for 10 min, after which a 3-fold dilution series was made in phosphate minimal medium (Johnsen et al. 2002) adjusted to a salinity of 15‰. A dilution of 1:270 of the sediment was used as the lowest dilution, giving a detection limit of 150 pyrene degraders g$^{-1}$. The pyrene was dissolved in hexane, added to the wells and the hexane was allowed to evaporate leaving the microplate wells coated with crystalline pyrene. Six replicates (200 µl each) of sediment–buffer suspension dilution were transferred to the wells. Pyrene was added as the sole source of carbon and energy to the wells. Therefore, positive growth in the wells was attributed to bacteria-degrading pyrene. Negative control plates (with no carbon or energy source) were treated with the hexane solvent only. The plates were incubated for 4 wk in a fume hood at approximately 20°C. To test the wells for growth, the potential respiration of each well was assayed by adding a carbon mixture (glucose, succinate and pyruvate) and Cell Proliferation Reagent WST-1 (Roche Molecular Biochemicals). The respiratory reduction of WST-1 (absorbance at 450 nm with a reference wavelength at 630 nm) was measured at time zero and after 6 h incubation on an orbital shaker (300 rev. min$^{-1}$) at room temperature. Wells were considered positive (i.e. microbial growth on pyrene) when the absorbance change was higher than 0.020 compared with the initial absorbance. The absorbance change in negative control plates was always much less than 0.020.

MPN estimates were calculated using the DOS program ‘MPN Calculator’ version 2.70 (Klee 1993). Negative wells at low dilutions were considered false negatives when all wells were positive in one or more following dilutions. The total bacterial population that was culturable was determined as the number of heterotrophic colony forming units (CFU) by plating 10-fold serial dilutions on marine agar 2216 (Difco) followed by incubation at 22°C for 10 d.

**Statistical analysis.** Rates of microbial pyrene mineralization, measured as 14CO$_2$ production, were calculated for each replicate using the slope of regression on the linear part of cumulative CO$_2$ production versus time plots. Differences in calculated slope values were tested using ANOVA after assuring that the ANOVA assumptions (normal distribution of residuals and homogeneity of variances) were fulfilled. ANOVA main effects were further analysed using a post hoc Tukey’s honest significant difference (HSD) test. Total cumulative 14CO$_2$ production was defined as the last data point of the cumulative data sets. Differences in cumulative production were tested using ANOVA followed by the post hoc HSD test. The estimated MPNs of pyrene degraders in different sediment compartments were analysed using the non-parametric Kruskal-Wallis test, since the ANOVA assumptions were not fulfilled.

**RESULTS**

**Pyrene mineralization in microcosms**

Pyrene mineralization started at maximum rates in all microcosms (Fig. 1). Mineralization rates were nearly constant throughout the incubation period in the Nd++
treatment, whereas mineralization rates in C++ and especially in Am++ treatments decreased over time, resulting in a non-linear cumulative $^{14}$CO$_2$ production (Fig. 1). The maximum rate of pyrene degradation was estimated using linear regression performed on data from Days 2 to 23 (sampling every 4 to 5 d), whereas total pyrene degradation was estimated as the cumulative $^{14}$CO$_2$ production at the end of the experiment (44 d).

The presence of macrofauna significantly enhanced the maximum rate of $^{14}$CO$_2$ production compared with control sediment by approximately 200% (Table 2). No difference was detected between macrofauna. After 44 d, significant differences were detected in total cumulative $^{14}$CO$_2$ production (1-way ANOVA, $p = 0.007$). Total $^{14}$CO$_2$ production was higher in treatments with *Nereis diversicolor* compared with control sediment, whereas no significant difference was detected between sediment with *Arenicola marina* and control sediment or between the 2 polychaetes (Table 2).

At the end of the 44 d exposure phase, we determined the amount of oxidized sediment in all the replicates of the different treatments. The calculated pyrene mineralization rates in the different replicates are plotted against the amount of oxidized sediment (Fig. 2). Considering all the replicates of the different treatments collectively, Fig. 2 shows a strong link between the amount of oxidized sediment and the pyrene mineralization rate (positive correlation; $r = 0.93$, $p < 0.0001$) substantiating the strong dependence of pyrene mineralization on the availability of oxygen.

### Pyrene mineralization potential

In all oxidized sediment samples (i.e. burrow and surface sediments), production of $^{14}$CO$_2$ started at maximum rates immediately after the addition of $^{14}$C-pyrene (Fig. 3). In samples with bulk (previously anoxic) sediment, the mineralization rates increased during the first 6 to 10 d, indicating growth of microbial pyrene degraders.

<table>
<thead>
<tr>
<th>Exposure phase</th>
<th>Mineralization rate (% d$^{-1}$) (HSD test)</th>
<th>Total mineralization after 44 d (%) (HSD test)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>C++</td>
<td>0.0060 ± 0.0012 A</td>
<td>0.29 ± 0.05 A</td>
<td>0.003</td>
</tr>
<tr>
<td>Am++</td>
<td>0.0125 ± 0.0024 B</td>
<td>0.38 ± 0.05 B A</td>
<td>0.0007</td>
</tr>
<tr>
<td>Nd++</td>
<td>0.0109 ± 0.0011 B</td>
<td>0.48 ± 0.04 B A</td>
<td>0.0001</td>
</tr>
</tbody>
</table>

Table 2. Pyrene mineralization rates (±SD) and total cumulative pyrene mineralization in microcosms containing $^{14}$C-pyrene-contaminated sediment (+++) and either *Arenicola marina* (Am), *Nereis diversicolor* (Nd) or no fauna (C). ANOVA tests indicated significant differences in mineralization rates ($p = 0.003$) and total mineralization ($p = 0.007$) among treatments. Significant differences between treatments detected by the post hoc Tukey’s honest significant difference (HSD) test are indicated with capital letters.

Pyrene mineralization potential, measured as the initial rate of $^{14}$CO$_2$ production, was equally high in oxidized sediments, regardless of origin (burrow and surface sediments) whereas significantly lower mineralization potentials were observed in bulk sediments ($p < 0.0001$, Table 3). There was no apparent effect of fauna on degradation rates.

### Microbial populations

The number of heterotrophic colony forming units (CFU) ranged from $1.2 \times 10^6$ to $7.6 \times 10^7$ CFU g$^{-1}$ (Fig. 4) and no significant differences in CFU were detected between the different types of sediment ($p > 0.2$). Aerobic bacteria growing on pyrene were detected in all the oxidized sediment samples (Fig. 4). The estimated number ranged from 150 cells g$^{-1}$ detected in surface sediment to more than 20 000 cells g$^{-1}$ in burrow sediments of *Arenicola marina* and *Nereis diversicolor*; however, MPN estimates were highly variable within treatments. The average numbers of bacteria growing on pyrene in burrow sediments were 12 000 (range: 6700 to 20 000) cells g$^{-1}$ (*A. marina*) and 9670 (range: 3800 to 21 000) cells g$^{-1}$ (*N. diversicolor*). In surface sediment, the estimated number was 1300 (range: 150 to 2800) cells g$^{-1}$. These differences between burrow and surface sediments could not be confirmed to be statistically significant, however, according to a Kruskal-Wallis test ($p = 0.061$). The contribu-
tion of bacteria able to grow solely on pyrene to the total number of (colony forming) bacteria was significantly higher (Kruskal-Wallis, $p < 0.011$) in burrow sediment from *A. marina* ($0.51 \pm 0.24\%$) compared with surface sediment ($0.036 \pm 0.03\%$) and *N. diversicolor* burrow sediment ($0.068 \pm 0.04\%$). Potential pyrene mineralization rates measured in the different sediment samples were significantly correlated with the estimated number of bacteria growing on pyrene ($p = 0.026$, $r = 0.64$).

**Microbial pyrene bioavailability**

Pyrene bioavailability, determined as mineralization of $^{14}$C-pyrene, aged in the sediment for 44 d (Fig. 5), was significantly different among treatments (1-way ANOVA, $p < 0.0001$).

The initial mineralization rates (i.e. flux of bioavailable pyrene from sediment to bacteria) were higher in surface sediments compared with bulk and burrow sediments (Table 4). Furthermore, the pyrene flux in surface sediment was significantly inhibited by fauna, with the highest flux in surface sediments from control treatments, followed by surface sediments from *Nd++*

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Pyrene mineralization potential (% d$^{-1}$)</th>
<th>Group (HSD test)</th>
</tr>
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<tbody>
<tr>
<td>C+</td>
<td>Surface 14.47 ± 0.66</td>
<td>A</td>
</tr>
<tr>
<td></td>
<td>Bulk 1.48 ± 1.07</td>
<td>B</td>
</tr>
<tr>
<td>Am+</td>
<td>Surface 14.91 ± 1.29</td>
<td>A</td>
</tr>
<tr>
<td></td>
<td>Burrow 13.44 ± 0.43</td>
<td>A</td>
</tr>
<tr>
<td></td>
<td>Bulk 2.34 ± 0.96</td>
<td>B</td>
</tr>
<tr>
<td>Nd+</td>
<td>Surface 11.58 ± 2.52</td>
<td>A</td>
</tr>
<tr>
<td></td>
<td>Burrow 16.67 ± 0.48</td>
<td>A</td>
</tr>
<tr>
<td></td>
<td>Bulk 1.83 ± 0.71</td>
<td>B</td>
</tr>
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</table>

Table 3. Pyrene mineralization potential ($\pm$SD) measured as the ability to mineralize freshly added $^{14}$C-pyrene under oxic conditions in surface, burrow and bulk sediment. Sediment samples originate from exposure phase treatments with non-radioactive pyrene and with either *Arenicola marina*, *Nereis diversicolor* or no fauna. Prior to the incubation $^{14}$C-pyrene was added. An ANOVA test indicated that there were significant differences in mineralization potential among sediment types ($p < 0.0001$). Significant differences between sediment types detected by the post hoc Tukey’s HSD test are indicated with capital letters.

Fig. 4. Abundance of total culturable bacteria (as colony forming units, CFU) and most probable number (MPN) of bacteria growing on pyrene in surface sediment and sediment from burrows of *Arenicola marina* (Am) and *Nereis diversicolor* (Nd). Error bars represent SD
and Am++ treatments. There were no significant differences in mineralization rates between bulk sediments and burrow sediments from the 2 polychaetes (Table 4).

The total bioavailable amount (bioaccessability) of the aged 14C-pyrene, determined as total cumulative 14CO2 production, was also highest in surface sediments followed by sediments and burrow sediments (Fig 5). Pyrene bioaccessibility in surface sediment from treatment C++ was significantly higher than in surface sediment from treatment Am++, but not different from treatment Nd++ (Table 4). Contrary to the initial mineralization rates, bulk sediment showed a significantly higher total cumulative 14CO2 production compared with sediments from burrows made by Nereis diversicolor (Table 4). No significant differences in bioaccessibility were detected between bulk sediment and burrow sediment from the Am++ treatment or between burrow sediment from the Nd++ and Am++ treatments (Table 4).

**DISCUSSION**

The production of 14CO2, indicative of 14C-pyrene mineralization, was enhanced approximately 2-fold in microcosms inhabited by polychaetes compared with defaunated sediment (Table 2). Although only a few studies have examined the effects of infauna on microbial PAH degradation, most of these show that the presence of infauna such as Nereis virens, Arenicola marina and Capitella sp. stimulate the degradation of various PAHs, and enhancement factors of between 2 and 3 have been reported (Bauer et al. 1988, McElroy et al. 1990, Christensen et al. 2002). In contrast, Kure & Forbes (1997) observed decreased mineralization of fluoranthene in a sediment containing A. marina. In that case, however, the fluoranthene was added as a contaminated top layer that was rapidly buried into anoxic sediment due to the conveyor belt feeding behaviour of A. marina (Kure & Forbes 1997). Our results are consistent with other sediment studies where the pollutants were uniformly distributed throughout the sediment.

Collectively, the replicates of the different treatments showed that mineralization rates were positively correlated with the amount of oxidized sediment (surface plus burrow sediment), which clearly supports the hypothesis that pyrene mineralization in microcosms is controlled by the availability of oxygen (Fig. 2). Furthermore, the increase of oxidized sediments due to the formation and irrigation of burrows is the main effect of macrofauna leading to the stimulation of microbial PAH mineralization. Oxygen has been shown to be the controlling factor for microbial PAH degradation in aquatic sediments (Boyd et al. 2005) as well as in low permeability soils and soil aggregates (Nocentini & Pinelli 2001), and low oxygen availability, especially for the bulk sediment, was probably limiting for PAH mineralization in this study.

The fact that biogenic structures are important sites for microbial PAH mineralization is further confirmed by the high pyrene mineralization potential measured in sediment from both Arenicola marina and Nereis diversicolor burrows (Fig. 3, Table 3). Pyrene mineralization was significantly higher in burrows than in the ambient bulk sediment and equalled the high rates observed in surface sediments. Hence, pyrene mineralization in macrofauna burrows not only contribute to
significant differences between sediment types detected by the post hoc Tukey’s (p < 0.0001) and bioaccessible pyrene (p < 0.0001) among sediment types. ANOVA tests indicated significant differences in the flux of bioavailable pyrene, i.e. the total amount of bioavailable pyrene (see text). The measurements of pyrene mineralization potential in burrow and surface sediments (Table 3). Although anaerobic degradation of low molecular weight PAHs (≤3 fused benzene rings) has been observed under both nitrate- and sulphate-reducing conditions, anaerobic degradation rates are orders of magnitude lower than aerobic degradation (Coates et al. 1997, Rockne & Strand 1998). Furthermore, the few cases where anaerobic degradation of high molecular weight PAHs, such as pyrene, have been reported were from sediment with a long history of heavy PAH contamination (e.g. Rothermich et al. 2002, Bach et al. 2005), which is not the case for the pristine sediment we investigated.

The measurements of pyrene mineralization potential were performed under oxic conditions and the low initial rate of pyrene mineralization in the bulk sediment indicates that the bacteria from the reduced sediment were less adapted to oxic conditions when compared with the bacteria from oxidized sediments, and that the abundances of aerobic pyrene-degrading bacteria were low. However, mineralization rates increased when the reduced sediment was exposed to oxygen. After >1 wk, rates approached, but never equalled, that of oxidized sediments, presumably due to the establishment of a community of aerobic bacteria that could degrade pyrene. These results provide insights into the potential importance of bioturbation for pyrene degradation in bulk sediments. Temporary or short-term (<1 wk) oxygenation of the bulk sediment is not likely to enhance pyrene degradation significantly. For example, establishment of permanent or semi-permanent burrows in bulk sediments should enhance degradation rates, while sporadic oxygenation of bulk sediments, e.g. due to the passage of errant polychaetes, will not have much of an effect.

The pyrene mineralization rates in sediment with *Nereis diversicolor* and without worms were almost constant throughout the experimental period, whereas mineralization rate decreased over time in sediment inhabited by *Arenicola marina*. The explanation for the
decreasing pyrene mineralization rates is probably related to the decreased pyrene bioavailability in oxidized sediments, which was most pronounced in the presence of A. marina. Since PAH degradation in oxic environments is often limited by the amount of bioavailable substrate, the decreased pyrene bioavailability probably leads to the observed decreasing degradation rates.

The bioavailability of pyrene was determined as mineralization of 14C-pyrene after addition of a surplus of pyrene degrader cells (Semple et al. 2006) so that pyrene bioavailability was the limiting factor for mineralization. Two measures of bioavailability (Reichenberg & Mayer 2006) can be inferred from these mineralization curves (Fig. 5). The initial mineralization rates are a relative measure of the ability of the environment to supply the bacteria with pyrene, i.e. a measure of the flux of substrate to the bacteria. In contrast, the maximum amount of pyrene mineralized is a measure of the bioaccessible amounts, i.e. the amounts that may potentially be mineralized over time. Both estimates are relative measures as some of the degraded pyrene is incorporated into the bacterial biomass. Intriguingly, both aspects of microbial pyrene bioavailability were strongly affected negatively by the presence of macrofauna and furthermore the effect of fauna seemed to be species specific (Table 4). In all treatments the bioavailability in surface sediments was higher than in bulk sediment. Generally, surface layers have a higher porosity compared with subsurface sediment (Kure & Forbes 1997), leading to higher apparent diffusion coefficients, and hence, a potentially higher transfer of PAH to the bacteria (Liu et al. 2001). The presence of macrofauna did, however, decrease pyrene bioavailability in surface sediments compared with surface sediments of defaunated microcosms. Since both polychaetes defecate at the sediment surface, it is possible that gut passage of PAH-contaminated sediment may have reduced PAH bioavailability in the defecated sediment. Giessing & Mayer (2004) showed that gut fluid from marine deposit-feeding polychaetes catalyzed oxidative coupling of pyrene metabolites to organic matter and subsequently decreased the bioavailability. The authors suggested that this reaction, which leads to covalent binding of PAHs, represents a sink for PAHs. This process may explain the macroinfauna-induced reduction in PAH bioavailability in surface sediment observed in this study. Microbial pyrene bioavailability in sediment from both Nereis diversicolor and Arenicola marina burrows was lower than in surface sediments, and the amounts of bioaccessible pyrene in burrows were also lower than in the surrounding bulk sediment. Most polychaetes, including A. marina and N. diversicolor, are known to produce mucus to stabilise their burrows, which leads to an increased organic carbon content of burrow lining and walls compared with other sediment compartments. Organic matter is one of the key factors controlling microbial pyrene bioavailability in soil and sediments, and increases in organic content may decrease PAH bioavailability substantially (e.g. Bogan & Sullivan 2003). Whatever the mechanism, the polychaetes studied clearly decreased microbial PAH bioavailability in surface sediment and biogenic structures, which probably influenced the overall PAH degradation, especially in sediments with A. marina. Since the effects of macroinfauna on microbial PAH bioavailability have not previously been examined, it is not clear to what extent other polychaetes, or macroinfauna in general, affect microbial PAH bioavailability.

CONCLUSIONS

Nereis diversicolor and Arenicola marina stimulated microbial pyrene mineralization, but the effects of A. marina decreased over time. The mechanism responsible for this stimulation was most likely the increased oxygen availability caused by the construction and irrigation of macrofaunal burrows. Burrows and surface sediments were important sites for microbial pyrene mineralization, characterised by high mineralization potentials and high abundance of bacteria able to grow on pyrene, whereas the capacity for aerobic PAH mineralization in bulk sediment was low. The polychaetes, especially A. marina, decreased pyrene bioavailability to bacteria in surface and burrow sediments, and thereby reduced the otherwise stimulating effect of bioturbation on the mineralization of pyrene.

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