Effect of Activated Carbon Amendment on Bacterial Community Structure and Functions in a PAH Impacted Urban Soil

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Supporting Information

ABSTRACT: We collected urban soil samples impacted by polycyclic aromatic hydrocarbons (PAHs) from a sorbent-based remediation field trial to address concerns about unwanted side-effects of 2% powdered (PAC) or granular (GAC) activated carbon amendment on soil microbiology and pollutant biodegradation. After three years, total microbial cell counts and respiration rates were highest in the GAC amended soil. The predominant bacterial community structure derived from denaturing gradient gel electrophoresis (DGGE) shifted more strongly with time than in response to AC amendment. DGGE band sequencing revealed the presence of taxa with closest affiliations either to known PAH degraders, e.g. Rhodococcus jostii RHA-1, or taxa known to harbor PAH degraders, e.g. Rhodococcus erythropolis, in all soils. Quantification by real-time polymerase chain reaction yielded similar dioxygenases gene copy numbers in unamended, PAC-, or GAC-amended soil. PAH availability assessments in batch tests showed the greatest difference of 75% with and without biocide addition for unamended soil, while the lowest PAH availability overall was measured in PAC-amended, live soil. We conclude that AC had no detrimental effects on soil microbiology, AC-amended soils retained the potential to biodegrade PAHs, but the removal of available pollutants by biodegradation was most notable in unamended soil.

INTRODUCTION

Activated carbon (AC) addition is currently being investigated as in situ technology to remediate polluted sediments and soil. A critical consideration for the application in aerobic soil is the long-term impact of AC on intrinsic microbial community structure and functioning, including the potential to biodegrade pollutants. Adding AC to soil will reduce the soil porewater concentration of hydrophobic organic compounds (HOCs) which will be adsorbed and strongly bound by the AC (Figure 1, process i). Pollutants bound in the micropores of AC become less accessible for biouptake by soil-dwelling organisms and plants in comparison with weakly bound or dissolved pollutants. AC amendment may thereby reduce the transfer of HOCs from the soil matrix into the terrestrial food-chain and also reduce phytotoxicity (Figure 1, process iii). Addition of AC to soil will also reduce HOCs leaching (Figure 1, process iv) and volatilization risks (Figure 1, process ii). Strong binding of HOCs by AC will, however, reduce the availability of HOCs to microorganisms with the ability to metabolize these compounds. A potential consequence could be the disappearance of HOC degraders from the predominant soil microbial community and/or the down-regulation of HOCs metabolism in favor of other carbon substrates. In soils with abundant HOCs degrading microorganisms, this may impair the pollution attenuation via biodegradation (Figure 1, process v).

There are reports of differing effects of strong sorbent amendments on pollutant biodegradation processes. On the one hand, addition of AC to contaminated soils was shown to reduce spiked 14C phenanthrene metabolism to 14CO2 in laboratory batch experiments, and Karapanagioti et al. Receiving...
showed that slow sorption kinetics limited spiked phenanthrene biodegradation in sediment slurries containing coal particles. On the other hand, Vasilyeva et al.7 suggested that AC helped overcome toxicity of polychlorinated biphenyls to microbes, and Payne et al.11 also reported that addition of AC had a slight stimulatory effect on PCB dechlorination in sediment. Bushnaf et al.12 reported that reduced availability of monoaromatic hydrocarbons in biochar amended soil led to greater biodegradation of linear, branched and cyclic alkanes, and the total petroleum hydrocarbon vapor degradation was comparable in sandy soil with and without 2% biochar. In a direct comparison of AC amendment with biostimulation and bioaugmentation, Hale et al.13 found that 2% AC addition was more effective than biostimulation or bioaugmentation in further reducing the availability of an already strongly sequestered polycyclic aromatic hydrocarbon (PAH) pollution in River Tyne sediment.

The aim of this paper was to study effects of AC amendment on the predominant bacterial community structure, and its functioning with emphasis on the biodegradation of PAHs. To our knowledge, this is the first-ever investigation of the long-term effects of AC amendment on soil microbial communities under realistic field conditions. We report changes in the predominant bacterial community structure over a three year period following amendment of a PAH-polluted urban soil with granular or powdered activated carbon (GAC and PAC). For the samples collected in year 3, we compared the bacterial communities and their functioning in greater detail using molecular microbiological and chemical methods. We assess the combined effect of sorption and biodegradation on PAH availability in soil with and without GAC or PAC to better understand impacts on environmental risks at contaminated sites.

**Soil Characterization and Sampling.** Details of the field trials of AC-based soil remediation are reported by Hale et al.8 The soil used in the lysimeter experiment was excavated at a building site in Drammen, Norway, intermixed using an excavator bucket with 2% wet soil weight of PAC (SilCarbon TH90, average 20 μm grain size, with 80% < 45 μm) or GAC (SilCarbon 0.3–0.8, 300–800 μm grain size), and embedded in outdoor lysimeters with 25 m² surface area and between 2.5 and 3 m depth. The soil initially had a total organic carbon content of 2.50 ± 0.04% dry weight, and amended soils contained the intended AC dose, although with considerable variability, 2.0 ± 1.0% for GAC or 2.4 ± 1.9% for PAC.8 Two series of soil samples for microbial analysis were taken immediately after the AC amendment of the soil, after 6 months, and after 3 years, and frozen at −20 °C with and without the addition of absolute ethanol 1:1 (v/v). For the year 3 samples, additional soil samples were stored in the cold room at 4 °C without ethanol addition for the batch experiments.

**Total Microbial Cell Number.** Ten μL of the year 3 sample stored in ethanol was added to 990 μL of filter-sterilized phosphate buffer saline (PBS, Oxoid) and the cells were stained by adding 50 μL of SYBR Gold nucleic acid stain in 100× concentrate in dimethyl sulfoxide (DMSO) (Invitrogen Ltd., Paisley, UK), wrapped in foil and incubated at room temperature for 30 min, after which they were filtered using a sterile Millipore filter holder and a 0.2-mm-pore-size black polycarbonate filter (diameter 25 mm; Millipore). The filters were transferred to glass microscope slides containing a drop of Citifluor (Citifluor Ltd., Canterbury, United Kingdom) antifadent to help adhesion to the slide. A further drop of Citifluor was placed on top of the filter, and a coverslip was carefully placed on top of the preparation. Total bacteria were determined by direct count under an oil-immersion objective.
(100× magnification) using an Olympus BX40 Epifluorescence microscope; 20 randomly chosen fields of view were counted using a dilution that yielded between 30 and 300 fluorescent cells having a clear outline and finite cell shape. The error between samples was estimated using the standard deviation of the mean from three replicates measured per treatment. Cell number per g of soil was determined by considering the numbers and the areas of fields of view on the filter membrane, the area of the membrane and the original sample dilution factor.14

**Soil Respiration.** Soil respiration was measured for the year 3 samples by monitoring CO₂ concentrations in 50-mL crimp-top vials containing 15 g wet weight of either unamended, PAC-amended, or GAC-amended soil. For each soil type duplicate batches were monitored for 4 days at room temperature (20 °C). GC-MS analysis of CO₂ was performed on a Fisons 8060 GC linked to a Fisons MD800 MS with a HP- PLOT-Q capillary column.

**Bacterial Community Analysis.** A fingerprinting method, denaturing gradient gel electrophoresis (DGGE), was carried out to determine similarities and differences between the predominant bacterial communities in the experimental treatments, and to identify selected members of these communities. Total bacterial DNA was extracted from 0.5-g (wet weight) aliquots of the stored (without ethanol) soil samples, taken at times 0 and 6 months and 3 years after the start of the lysimeter experiments. The DNA extraction was carried out using Fast DNA Soil Kit (Qbiogene) with a cation step, modified from the method of Griffiths et al.15 added prior to the nucleic acid extraction in order to prevent the coextraction of compounds such as humic acids and clay minerals, which are known to inhibit PCR amplification. Briefly, the pretreatment consisted in the extraction of nucleic acids from the soil matrix by using hexadecyltrimethyammonium bromide (CTAB, Sigma-Aldrich) extraction buffer and phenol/ chloroform/isoamyl alcohol (25:24:1, Sigma-Aldrich), followed by phenol removal using chloroform/isoamyl alcohol (24:1, Sigma-Aldrich). Primers 2 and 3, targeting the bacteria, were used to PCR amplify the V3 region of bacterial 16S rRNA gene fragments, as previously described by Muyzer et al.,16 and the PCR products were analyzed by DGGE as described previously.16 The DGGE images were normalized and interpreted using the image analysis software BioNumerics (Applied Maths NV, St. Martens-Latem, Belgium). Primer 6 (Primer-E Ltd., Plymouth, UK) was used to perform cluster analysis using the Pearson product-moment correlation coefficient and Analysis of Similarity (ANOSIM).

**Band Sequencing.** Dominant DGGE bands were excised from the gel, PCR-amplified using primers 2/3, purified and sequenced with primer 2 or 3 (3.2 pmol/μL DNA) using the ABI prism Big Dye Terminator Cycle Sequencing Ready reaction Kit and an ABI Prism 377 DNA sequencer (Applied Biosystems, USA) as previously described.15 Sequences were compared against the Ribosomal Database Project (RDP10) and GenBank databases using the BLAST algorithm to determine the closest matching sequence identity.

**Quantitative Real-Time PCR.** The abundances of specific ring-hydroxylating dioxygenase α-subunit (RHDα) genes were measured in each sample by real-time PCR using the following primer sets: (i) P6B (forward, 5′-TGGCGAATCCTGTGCAGGGAC-3′; reverse, 5′-CGTCCAGRCAACCGGADAYC -3′), targeting a pdoA2/phdA clade that includes mostly Mycobacterium species and have *M. vanbaelenii* as a reference strain; (ii) P4 (5′- CCGGAGACCTTCTTGACCG-3′; 5′-GACGCGGGT-3′) targeting a *ebd*A1/ *ebd*A1/ *akk*A1b clade that only includes *Rhodococcus* species and have *R. jostii* RHA1 as a reference strain; and (iii) P7B (5′- CACBGTGACGTAY-CACG -3′; 5′- CATGTTGCCCTAGTAGAAC -3′) targeting a *ipbA1/bphA1* clade that includes mostly *Rhodococcus* and some *Pseudomonas* species and, in particular, includes several *R. erythropolis* strains. Real-time PCR experiments were conducted on a BioRad CFX96 (Hercules, CA) with a C1000 thermal cycler iCycler and software version 1.6 (BioRad CFX Manager). Ten μL of reaction mixture contained 3 μL of template DNA (or filtered sterile molecular biology water, Sigma-Aldrich, as negative control), and 15 pmol of each primer, with the SsoFast EvaGreen Supermix for the CFX96 (Biorad Laboratories Ltd.). The following temperature profile was used in the amplifications: step one heated to 98 °C (2 min), followed by 40 cycles of 2 s denaturation at 98 °C, 5 s at the primer specific annealing temperature (58 °C for P6B, 60 °C for P4, and 59 °C for P7B). At the end of the real-time PCR, a melting curve was performed as a final step that consisted of the measurement of the SYBR Green signal intensities during a 0.2 °C temperature increment every 10 s from 65 to 95 °C. The corresponding plot of change in fluorescence versus temperature shows a single peak for every amplicon at its specific melting temperature. For every set of primers, calibration curves were obtained from standard DNA prepared from plasmids containing the cloned target sequences. The plasmid DNA concentration was quantified using Nanodrop 1000 spectrophotometer (Thermo Scientific). The copy number of standard plasmids was calculated according to plasmid size, insert lengths, and assuming a molecular mass of 660 Da per base pair. Stock solutions of standard DNA were prepared at a concentration of 10⁶ copies of plasmid μL⁻¹. For the calibration curves, DNA standards ranging from 10⁰ to 10⁷ target gene copies μL⁻¹ were prepared by diluting the stock solutions. All the standard curves were linear (R² > 0.99) over the 9 orders of magnitude (10⁰ to 10⁷ μL⁻¹) of gene copy number. Every sample was run in triplicate and each experiment was repeated at least twice. In every run, the accuracy in the detection of the target gene in the soil samples was confirmed by comparing the melt curves and the agarose analysis of the qPCR products to the ones obtained with DNA extracted from model organisms (positive control) in order to avoid false detection in the environmental samples.

**PAH Availability Assessed by Uptake by Polyethylene Samplers.** A series of batch experiments was set up to study the uptake of available PAHs in passive samplers.13 Passive samplers are able to passively accumulate HOCs from contaminated water until partitioning equilibrium is established. Pollutants taken up by passive samplers are potentially also available for uptake by critical receptors such as plants and earthworms (Figure 1, process iii). To study the effect of sorption alone (Figure 1, process i) compared to the combined effect of sorption and biodegradation (Figure 1, processes i and v) on pollutant availability, batches were set up with and without sodium azide addition which inhibits PAH biodegradation.17,18 The microcosms comprised the following: 5 g (wet weight) of unamended soil, PAC-amended, or GAC-amended soil and 40 mL of water. The water content of the samples was between 16 and 21% wet weight. For each soil type triplicate batches were set up with and without 1 g/L sodium azide (Sigma Aldrich). 0.15 ± 0.01 g clean
polyethylene (PE) passive sampler was added to each batch. Blank controls were run in parallel and consisted of triplicate batches containing PE samplers and water with and without sodium azide. Batches were plugged with cotton balls and mixed at 100 rpm on an orbital shaker (IKA labortechnik, Germany) in a secondary dark containment consisting of a cardboard box with some holes punched into the lid for aeration. PE samplers were removed after 3 weeks and extracted twice for 24 h in 10 mL of 50:50 v:v hexane/acetone. A surrogate standard was added to the first extraction to monitor and correct for recoveries, which were 78 ± 11% for d_{10} phenanthrene, 83 ± 6% for d_{10} pyrene, 92 ± 4% for d_{12} benzoanthracene, and 94 ± 8% for d_{12} benzpyr.

Sample Cleanup and PAH Analysis. Sample cleanup was performed with a silica gel column topped by sodium sulfate before GC-MS analysis with an Agilent 6850 Gas Chromatograph (DB-XLB column length 30 m, i.d. 25 mm, and 1 μm film thickness) coupled to an Agilent 5973 mass spectrometer as described by Hale et al. 8

RESULTS AND DISCUSSION

Total Microbial Cell Count and Soil Respiration. A statistically significant difference was observed for the total cell counts 3 years after the amendment, with GAC amended soil having the highest cell count of 1.8·10^{10} cells per gram of dry soil weight, 2.5 times higher than the unamended soil, which had the lowest cell count (Figure 2a). These values are in the higher range for most typical soil types (e.g., 19, 20). The soil respiration rate was also a statistically significant factor of 2.6 and 2.7 higher in the GAC-amended soil compared to the unamended and PAC-amended soils (Figure 2b). Soil respiration resulted in a linear increase in the CO₂ batch headspace concentration over the entire 4-day monitoring period for all soils (R² values >0.98), and soil respiration could be quantified as 0.19 ± 0.02, 0.18 ± 0.01, and 0.49 ± 0.03 μg CO₂ per g of dry soil per hour for unamended, PAC-, and GAC-amended soil, respectively. These observations coincide with field observations of better plant growth in GAC amended soil, see Jakob et al. 21 The soil respiration rates were similar to the values found in another study on a hydrocarbon contaminated site. 22 These results show that AC amendment overall was not detrimental to aerobic microbial activity despite previously reported enhanced DOC binding in AC amended soils. 8

Changes in the Predominant Bacterial Community Structure. The predominant soil bacterial community structure, as assessed by DGGE and cluster analysis of amended and unamended soil after 0, 6, and 36 months (Figure 3), showed that all the samples from each particular time period clustered closely with other samples from the same time period, regardless of treatment. This indicates that the predominant bacterial community structure shifted more strongly with time than in response to PAC or GAC amendment (ANOSIM test for differences between time groups, Global R = 0.852, p < 0.01; for differences between treatments, Global R = −0.317, p = 0.97). Overall, similarity between predominant bacterial communities, determined by the Pearson product-moment correlation coefficient, was greater than 65% (Figure 3b). Duplicate DNA samples from the same time point and treatment showed higher similarity (>77% overall and average similarity coefficient >87%, Figure 3b), indicating good method reproducibility. The most abundant microbial species are important drivers of carbon and nutrient cycling in soil. The fairly high similarities of predominant bacterial community structures in AC amended and unamended soil observed in this study alleviates concerns about strong side-effects of AC amendment. The findings agree with field community surveys of macroinvertebrates in sediments where AC amendment generally had only weak effects on community structure and diversity, 23, 24 although clear effects on species abundance were observed in one field study. 24

Band Sequencing Results. Identification of the nearest matching neighbor to sequenced DGGE bands was carried out using the Basic Local Alignment Search Tool (BLAST) and classification was confirmed using the Classifier tool in the Ribosomal Database Project (RDP 10). Twenty-three of the twenty-six most predominant taxa were sequenced and classified into three main phyla, Bacteroidetes, Actinobacteria, and Proteobacteria (identified by different symbols, Figure 3 and Table S1 in Supporting Information) that harbor common environmental bacteria known to play an important role in the decomposition of organic materials. 25–28 The three most abundant and prevalent taxa (bands N6, N10, and N17 respectively) belonged to the order Micrococcinae within the Actinobacteria phylum, whose nearest neighbors were identified as an Curtobacterium, Arthrobacter, and Microbacterium species although with varied similarities (89–99%; Table S1 in Supporting Information). These bacteria are common soil inhabitants. The nearest-matching neighbor of several predom-
inantly abundant (intense bands) taxa are of interest because of the reported widespread ability to degrade PAHs in members of these taxa (e.g., 29): band N7 was 96% identical to a *Mycobacterium* sp. and 92% similar to the versatile PAH-degrading *M. vanbaalenii* PYR-1; band N8 was 95% similar to the versatile PAH-degrading *Rhodococcus josttii* strain RHA1; band N16 was 95% identical to an uncultured *Sphingomonas* sp.; and band N23 was 99% similar to a strain of the species *Rhodococcus erythropolis*, members of which are known to degrade biphenyls and isopropylbenzene (e.g., refs 33, 34). These groups are often present in all soils (e.g., 35), but appear to dominate more frequently in contaminated soils (e.g., 36, 37), although to our knowledge it is unusual to obtain such a predominance of sequences matching mycolic acid-containing actinomycetes (e.g., rhodococci and mycobacteria).

These bands were present at all times in bacterial communities from unamended, PAC-, and GAC-amended soil, but their relative intensity varied over time (see Table 1). The relative intensity of bands can provide a rough proxy of the proportional abundance of those taxa in the community. The relative intensities of these bands by year 3 were on average stronger than for the beginning of the study, but comparable between treatments, which would indicate a similar relative abundance of these putative PAH-degrading taxa in the soils. The *R. josttii* RHA-1-like sequence (band N8) showed a particularly marked increase in intensity between the start and end of the study, where it increased from about 3% to nearly 10% of the predominant bacterial community as expressed by relative intensity in AC amended and unamended soil.
Real-Time PCR Results. Real-time PCR is a valuable tool for the quantification of the functional PAH ring-hydroxylating dioxygenases.38,39 Dioxygenase systems add both atoms of molecular oxygen to the aromatic ring as a first step in the aerobic degradation of lower molecular weight PAHs, and dioxygenase gene quantification is therefore of interest in the assessment of PAH bioremediation potential.40,41 The RHD₆₄ gene copy numbers reported in Table 2 indicate that those represented by the reference Rhodococcus RHA1 species (from here-on called etbARHA1-RHD₆₄) showed the highest concentrations (∼10⁷–10⁹ g⁻¹), while those represented by a Rhodococcus erythropolis reference species (from here-on called ipdAReryth-RHD₆₄) showed the lowest concentrations (∼10⁵–10⁶ g⁻¹) for every treatment. Those represented by the reference Mycobacterium vanbaalenii PYR-1 species (from here-on called pdoAR_PYR1-RHD₆₄) had intermediate concentrations (∼10⁶–10⁷ g⁻¹). Those putative aromatic ring-degrading bacteria (i.e., RHD₆₄ containing bacteria) analyzed accounted for between 0.003 and 1% of the total bacterial population if each bacterial cell is assumed to contain one gene copy. These values are similar to the absolute and relative abundance values reported by Cebron et al.39 who used quantitative PCR to target general dioxygenase gene populations in contaminated soils, but higher than those in studies that only targeted specific dioxygenase gene populations.42–44 The concentrations of each targeted gene fell within an order of magnitude between unamended soil, soil amended with PAC, and soil amended with GAC. Indeed, the predominant etbARHA1-RHD₆₄ gene population increased over the time of the trial, and to a greater extent in AC-amended compared to unamended lysimeters. This confirms that bacteria with the ability to degrade PAHs remain in PAC- and GAC-amended soil at levels comparable to the populations in unamended soil. The real-time PCR method measures gene numbers, or the potential to degrade PAHs, not gene expression, which may be down-regulated by some pollutant degraders.45 In other words, while the results indicate that the ability to synthesize dioxygenases was present in all bacterial communities isolated from the soil samples, regardless of AC amendment, this does not necessarily imply active synthesis of the enzymes. It is interesting to note that there was a significant correlation between targeted RHD₆₄ gene abundances and the concentration of those taxa found by DGGE analysis (relative intensity × total cell count) that had close affiliations to the reference species of those RHD₆₄ genes (Pearson’s correlation, r² = 0.83, P < 0.01, n = 9). However, despite the close affiliation of these 16S rRNA gene fragments with the reference species for each targeted clade of RHD₆₄, the similarities were often lower (e.g., 92–96%) than those cut-offs usually used for species level (97%) and it cannot be confidently concluded that these taxa would necessarily contain RHD₆₄ genes even if they were deemed to be similar taxa.

Assuming a carbon content of 310 fg per cell44 and one etbARHA1-RHD₆₄ gene copy per cell, the biomass carbon corresponding to the gene copy numbers in Table 2 would be between 40 and 70 μg per g of dry soil, which is greater than the solid-phase PAH concentration and much greater than the freely dissolved PAH concentration of below 1 ng per g of soil.8 The comparison suggests that microorganisms with dioxygenase genes utilized other main carbon substrates and could therefore grow and persist long-term in the soil with and without AC amendment. The findings alleviate concerns that enhanced PAH binding by AC may cause a significant decrease in the abundance of putative PAH degraders in AC amended soils.

### Table 1. Summary of the Relative Intensities of Selected DGGE Excised Bands for Every Duplicate DNA Extraction and Their Average for Different Treatments and Time Point

| soil | time | sample | N7 DGGE Mycobacterium-like sequence | N8 DGGE Rhodococcus RHA1-like sequence | N23 DGGE Rhodococcus erythropolis-like sequence |
|------|------|--------|-----------------------------------|---------------------------------------|---------------------------------------------|
|      |      |        | relative band intensity % | relative band intensity % | relative band intensity % |
|      |      |        | average | average | average |
| unamended | 0 | DNA1 | 1.87 | 1.52 ± 0.50 | 2.93 | 2.72 ± 0.30 | 1.59 | 1.43 ± 0.22 |
|         |    | DNA2 | 1.16 |               | 2.51 |               | 1.28 |               |
|         | 6 months | DNA1 | 4.50 | 4.46 ± 0.05 | 3.26 | 3.30 ± 0.06 | 1.84 | 1.87 ± 0.04 |
|         |         | DNA2 | 4.42 |               | 3.34 |               | 1.90 |               |
|         | 3 years | DNA1 | 3.00 | 2.96 ± 0.06 | 8.36 | 8.92 ± 0.80 | 2.21 | 2.35 ± 0.20 |
|         |         | DNA2 | 2.92 |               | 9.48 |               | 2.50 |               |
| PAC    | 0     | DNA1 | 3.48 | 3.15 ± 0.46 | 4.65 | 4.26 ± 0.55 | 2.22 | 2.16 ± 0.08 |
|         |    | DNA2 | 2.83 |               | 3.87 |               | 2.11 |               |
|         | 6 months | DNA1 | 3.42 | 3.04 ± 0.54 | 3.34 | 3.33 ± 0.004 | 1.94 | 2.00 ± 0.08 |
|         |         | DNA2 | 2.65 |               | 3.33 |               | 2.06 |               |
|         | 3 years | DNA1 | 3.25 | 3.49 ± 0.34 | 9.37 | 9.26 ± 0.16 | 3.32 | 3.03 ± 0.40 |
|         |         | DNA2 | 3.73 |               | 9.14 |               | 2.74 |               |
| GAC    | 0     | DNA1 | 1.96 | 1.84 ± 0.17 | 3.53 | 3.19 ± 0.50 | 1.66 | 1.57 ± 0.12 |
|         |    | DNA2 | 1.71 |               | 2.85 |               | 1.48 |               |
|         | 6 months | DNA1 | 4.11 | 2.64 ± 2.08 | 3.33 | 2.71 ± 0.89 | 2.11 | 1.82 ± 0.40 |
|         |         | DNA2 | 1.17 |               | 2.08 |               | 1.54 |               |
|         | 3 years | DNA1 | 3.73 | 3.34 ± 0.55 | 8.55 | 7.90 ± 0.92 | 2.77 | 2.67 ± 0.15 |
|         |         | DNA2 | 2.96 |               | 7.25 |               | 2.57 |               |

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### Table 2. Dioxynogenase Gene Copy Number per gram of Wet Soil for Every Duplicate DNA Extraction Sample, for Unamended and AC-Amended Soils at Different Time Points

| soil        | targeted genes | Mycobacterium vanbaalenii | Rhodococcus jostii RHA1 | Rhodococcus erythropolis |
|-------------|----------------|---------------------------|-------------------------|-------------------------|
|             | gene copy n/g wet soil | gene copy n/g wet soil | gene copy n/g wet soil | gene copy n/g wet soil  |
|             | average         | average                   | average                 | average                |
| Unamended   | DNA1            | 1.18 ± 0.13 × 10⁷         | 3.76 ± 0.35 × 10⁷       | 1.95 ± 0.70 × 10⁶      |
|             | DNA2            | 1.60 ± 0.17 × 10⁷         | 3.69 ± 0.68 × 10⁷       | 2.39 ± 0.48 × 10⁶      |
| 6 mths      | DNA1            | 3.11 ± 0.42 × 10⁶         | 8.45 ± 0.04 × 10⁷       | 2.85 ± 0.33 × 10⁶      |
|             | DNA2            | 3.11 ± 0.15 × 10⁶         | 7.12 ± 0.74 × 10⁷       | 2.77 ± 0.55 × 10⁶      |
| 3 yrs       | DNA1            | 3.12 ± 0.44 × 10⁶         | 1.20 ± 0.09 × 10⁷       | 2.69 ± 0.78 × 10⁶      |
|             | DNA2            | 3.45 ± 0.21 × 10⁶         | 1.17 ± 0.10 × 10⁷       | 0.71 ± 0.06 × 10⁶      |
| PAC         | DNA1            | 1.98 ± 0.31 × 10⁷         | 4.43 ± 0.54 × 10⁷       | 6.43 ± 0.40 × 10⁶      |
|             | DNA2            | 1.69 ± 0.04 × 10⁷         | 4.38 ± 0.36 × 10⁷       | 2.62 ± 0.52 × 10⁷      |
| 6 mths      | DNA1            | 3.60 ± 0.20 × 10⁶         | 7.75 ± 0.99 × 10⁷       | 6.34 ± 0.80 × 10⁶      |
|             | DNA2            | 3.51 ± 0.70 × 10⁷         | 7.80 ± 0.77 × 10⁷       | 6.79 ± 0.61 × 10⁶      |
| 3 yrs       | DNA1            | 4.47 ± 0.90 × 10⁶         | 1.79 ± 0.17 × 10⁷       | 7.24 ± 0.42 × 10⁷      |
|             | DNA2            | 5.22 ± 0.26 × 10⁶         | 1.71 ± 0.04 × 10⁸       | 1.55 ± 0.08 × 10⁸      |
| GAC         | DNA1            | 9.48 ± 0.40 × 10⁶         | 3.37 ± 0.12 × 10⁷       | 1.07 ± 0.03 × 10⁸      |
|             | DNA2            | 1.13 ± 0.90 × 10⁷         | 3.57 ± 0.76 × 10⁷       | 1.57 ± 0.25 × 10⁸      |
| 6 mths      | DNA1            | 2.56 ± 0.81 × 10⁷         | 8.43 ± 0.13 × 10⁷       | 1.37 ± 0.30 × 10⁸      |
|             | DNA2            | 2.45 ± 0.38 × 10⁷         | 7.49 ± 0.43 × 10⁷       | 1.78 ± 0.22 × 10⁸      |
| 3 yrs       | DNA1            | 5.29 ± 0.90 × 10⁷         | 1.72 ± 0.23 × 10⁴       | 2.50 ± 0.16 × 10⁸      |
|             | DNA2            | 6.12 ± 0.63 × 10⁷         | 1.82 ± 0.13 × 10⁸       | 2.37 ± 0.22 × 10⁸      |

*Each quantification was performed in triplicate real-time PCR runs. phdA: iron—sulfur protein large subunit; pdoA2: putative PAH ring-hydroxylating dioxynogenase large subunit; etbA1/ebdA1: ethylbenzene dioxygenase alpha subunit; akbA1b: alkylbenzene dioxygenase; ipbA1: isopropylbenzene-2,3-dioxygenase iron-sulphur protein subunit; bphA1: large subunit of biphenyl dioxygenase.*
pollution in soil,\(^4\) as one PE sampler had much higher uptake of free aqueous PAH concentrations. To account for the pollution attenuation which may result in down-regulation of PAH metabolism. However, unamended soil, alleviating concerns that AC amendment was the lowest for PAC-amended soil without sodium azide, \((\text{Table S2 in Supporting Information})\). Overall, PAH uptake was more readily biodegradable,\(^4\) was on average 99% lower in the unamended soil \((t\text{ test, one-tailed, } p < 0.01\) for PAC). The e-\(\text{field test, one-tailed, } p < 0.01\) for GAC and \(p < 0.05\) for PAC). The effect of biodegradation is illustrated by statistically significant 75% and 46% reductions of the PAH uptake by PE passive samplers for unamended and GAC-amended soil, respectively, when comparing soil slurries with and without sodium azide \((t\text{ test, one-tailed, } p < 0.01\) for unamended and \(p < 0.05\) for GAC). The PAH compound pattern was also indicative of biodegradation; the uptake of the smaller two- and three-ring PAH compounds, which tend to be more readily biodegradable,\(^3\) was on average 99% lower in the unamended soil without, as compared to with, sodium azide \((\text{Table S2 in Supporting Information})\). Overall, PAH uptake was the lowest for PAC-amended soil without sodium azide, and our observations suggest active PAH biodegradation despite low PAH availability in both AC amended and unamended soil, alleviating concerns that AC amendment may result in down-regulation of PAH metabolism. However, the benefit of AC amendment was less clearly pronounced without biocide addition. For soils slurries without sodium azide, only the 44% reduction for PAC amended soil compared to the unamended soil was statistically significant \((t\text{ test, one-tailed, } p < 0.05)\). Therefore, for a realistic assessment of the long-term benefits of AC amendment under field conditions, laboratory pilot-trials should be conducted with both sterile and live soils to account for the pollution attenuation which may occur in control soil due to biodegradation.

To put these results into context, we compare in Figure 5 the free aqueous PAH concentrations \(C_w\) estimated from the PAH concentrations in polyethylene, \(C_{pe}\), using PE-water partitioning coefficients \(K_{pe}\) and the relationship \(C_w = C_{pe}/K_{pe}\),\(^3\) with free aqueous PAH concentrations reported by Hale et al.\(^8\) which were derived from passive polyoxymethylene (POM) samplers embedded field lysimeters. In-situ and ex-situ assessments are not exactly comparable because of differences in temperature and mixing regimes, which are more optimal under laboratory conditions, and therefore tend to facilitate pollutant mass transfer and biodegradation.\(^4\) Nevertheless, the combined data seem to show a continuing trend of decreasing free aqueous PAH concentrations in both AC amended and unamended soil over the 3-year remediation trial period, eventually falling to very low levels below EU guidance values for drinking water set out in EU directive 200/60/EC in all systems. This overall trend is the combined effect of all attenuation processes which comprise PAH sorption, biodegradation, chemical reactions, leaching, and volatilization.\(^5\) The difference between the accelerated pollution attenuation in the AC-amended systems and the natural pollution attenuation become smaller with time, and may eventually converge, when natural attenuation has depleted the readily available PAH pool to the same extend as AC amendment. According to our results, the biodegradation of available PAHs is a significantly contributing factor to this reduction in free aqueous PAH concentrations, even though total PAH concentration of 23 ± 15 μg per g of untreated dry soil\(^3\) were not measurably changed during the field experiment.\(^8\) AC amendment benefits, such as reduced PAH uptake by earthworms and plants\(^3\) and reduced PAH leaching,\(^3\) and AC amendment costs, need to be assessed in comparison with the working of intrinsic attenuation mechanisms, which may also reduce risks in the long term, in particular in soils with a high abundance of pollutant degrading microorganisms.

**ASSOCIATED CONTENT**

\(\text{Supporting Information}\)

Compilation of closest relatives and their similarities with the sequence of dominant bands excised from the DGGE gels and the PAH compound specific uptake by PE in soil slurries without compared to with sodium azide addition. This information is available free of charge via the Internet at http://pubs.acs.org/.
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