ABSTRACT: Sediments sampled next to the Cíes Islands, a natural reserve of the National Park of Atlantic Islands in Ría de Vigo (Spain), 3 mo after the first tide of fuel from the ‘Prestige’ tanker arrived, were chemically and microbiologically characterized. The chemical analysis of the sediments, using oil fingerprinting techniques, showed the occurrence of fresh ‘Prestige’ fuel oil with a background of older hydrocarbon contamination. The cultured bacterial community of the contaminated sediments harbored a high population of total heterotrophs and alkane degraders and a small proportion of aromatic-degrading bacteria. Based on partial 16S ribosomal RNA gene sequence data, 37 different bacterial strains, isolated in diluted marine agar, were detected. Most of them were classified as members of the groups Gammaproteobacteria (59%) and Alphaproteobacteria (21%), although members of the Bacteroidetes (10%) and of Firmicutes (10%) were also found. Some of the different identified bacteria have previously been described as fuel oil–degrading species such as Alcanivorax, Shewanella, Vibrio, Pseudoalteromonas and Marinomonas. Seven independent isolates were able to grow with hexadecane. However, these 7 strains are grouped under the same phylotype, based on the 16S rRNA gene sequence, and were closely related to Alcanivorax borkumensis (Gammaproteobacteria). No aromatic-degrading activities were detected among the culturable bacteria. The presence of fresh fuel without a detected level of biodegradation suggests that Alcanivorax was an early colonizer after the ‘Prestige’ oil spill. To our knowledge, the isolation of Alcanivorax from environmental samples without the aid of an enrichment procedure has not been previously reported. Our results suggest the ability of Alcanivorax to compete and coexist with other heterotrophic marine bacteria in an oil polluted marine environment rich in the nutrients N and P.

KEY WORDS: ‘Prestige’ oil spill · Alcanivorax · Pollution · Hydrocarbon contamination
reserves, such as the National Park of the Atlantic Islands, have been affected since then.

The ecological importance of the islands prompted the present work, which was carried out in order to obtain information about the natural microbiota present in surface sediments and to study their potential for natural attenuation processes on heavy fuel oil biodegradation after the 'Prestige' oil spill. Chemical and microbiological analyses provide valuable data for assessing the intrinsic bioremediation capabilities of the sediments.

Bioremediation, defined as the act of adding nutrients or bacteria to contaminated environments to cause an acceleration of the natural biodegradation processes, is recognized as an appropriate oil spill response tool (Swannell et al. 1996). Adding nitrogen (N) and phosphorous (P) to the environment (biostimulation) has been a common bioremediation strategy to favor the growth of degrading bacteria. The use of allochthonous petroleum-degrading bacteria (bioaugmentation) obtained from enriched laboratory cultures has not yet been shown to be more effective than the stimulation of the autochthonous species of degraders (Swannell et al. 1996, Diez et al. 2005, Vinas et al. 2005a). Recently, an enhancement of biodegradation, especially of heavier alkanes and, even more so, of alkylated polycyclic aromatic hydrocarbons (PAHs), has been described in the treatment of the 'Prestige' oil using an oleophilic fertilizer in vitro and in field conditions (Jiménez et al. 2006, 2007).

Before the application of in situ bioremediation, it is advisable to search for indigenous hydrocarbon-degrading microorganisms. There is limited information about the vast majority of oil-degrading marine bacteria, which remain uncultured, and even less is known about microbial species present in marine sediments contaminated with heavy fuels.

To our knowledge, only the experiments conducted after the ‘Nakhodka’ oil spill retrieved information about microbial populations adapted to heavy fuel oil. In this case, the development of microbial communities in polluted samples from natural beaches and water was studied, and the microorganisms later subjected to enrichment in batch culture. Under these conditions, adapted microorganisms were more easily grown in isolation media (Kasai et al. 2002). Different molecular approaches, mainly involving 16S rRNA analysis such as PCR/DGGE (Kasai et al. 2001), clone libraries and specific oligonucleotide probes (Maruyama et al. 2003), were used in order to follow the changes in the degrading community. All these studies reached similar conclusions, which could be summarized as follows: (1) the total population of bacteria was almost stable and the accident had no long-term effects on its composition; (2) the degraders of oil components showed a maximum population level immediately after the oil spill; (3) the oil-degrading community was predominantly composed of bacteria that closely resembled the aliphatic hydrocarbon decomposer Alcanivorax borkumensis, followed by the aromatic-degrading bacteria Cycloclasticus pugetii and Sphingomonas.

Several genera of marine bacteria capable of degrading petroleum hydrocarbons have recently been described such as Alcanivorax, Oleiphilus, Oleispira, Cycloclasticus and Marinobacter. These genera and a few others that are obligate consumers of oil hydrocarbons have been named obligate hydrocarbonoclastic bacteria (OHCB), which bloom after a pollution event. However, the types of hydrocarbon-degrading bacteria that may bloom depend on the latitude/temperature, salinity, redox conditions, and other physicochemical factors (Harayama et al. 2004, Yakimov et al. 2007). Indeed, it has recently been shown that biological factors, such as bioturbation, may also favor the development of oil-degrading communities in polluted environments (Cuny et al. 2007).

Alcanivorax borkumensis was isolated for the first time in the North Sea using n-tetradeacne as the sole carbon and energy source (Abraham et al. 1998, Yakimov et al. 1998). In a relatively short period of time, the genus Alcanivorax has been isolated or detected by molecular techniques from samples taken from different places such as the Atlantic Ocean, the Mediterranean Sea, the North Sea, the Sea of Japan, the South China Sea and the Antarctic (Golyshin et al. 2003).

It is an aerobic organism that has been defined as a ‘professional hydrocarbonoclastic bacteria’, since it is an obligate consumer of alkanes and branched alkanes unable to grow on sugars or amino acids (Harayama et al. 2004, Head et al. 2006, Yakimov et al. 2007). Alcanivorax usually exists in low numbers in unpolluted waters and rapidly increases along coastlines and in oil-polluted waters (Harayama et al. 1999, Kasai et al. 2001, Syutsubo et al. 2001). Its capacity to grow on branched alkanes seems to be the reason why this genus becomes the major component of the bacterial population in oil-contaminated seawater in the presence of an adequate supply of N and P (Hara et al. 2003).

Like most hydrocarbon-degrading species, Alcanivorax spp. have been isolated using media enriched in fuel or any oil fraction (Liu & Shao 2005), or in a very oligotrophic medium (Fernandez-Martinez et al. 2003), in order to avoid competition with heterotrophic marine bacteria, which usually outcompete hydrocarbon-degrading species. Alcanivorax has also been isolated from heavy oil-polluted samples from the ‘Nakhodka’ oil spill in Japan. Again, Alcanivorax could only be isolated from oil-polluted sand, gravel and seawater after growing the polluted samples in nutrient-enriched cul-
tures (Kasai et al. 2001, 2002, Roling et al. 2002). These experiments were performed in vitro or under artificial enrichment conditions, so their results cannot be directly extrapolated to the real environment. However, to answer the question of which microorganisms have particular significance for the removal of hydrocarbons from the environment, it could be useful to obtain evidence from culture-based studies that define the catabolic capabilities of candidate organisms and indicate the qualitative and quantitative importance of particular organisms in situ (Head et al. 2006).

In the present work, we report on a chemical and culture-based microbiological study of marine sediments affected by the ‘Prestige’ heavy fuel oil spill, 3 mo after the accident. Chemical fingerprinting was used to detect polluted sediments, verifying that no cross contamination from different sources other than the ‘Prestige’ fuel occurred. The most recently polluted sediments were used directly for a microbiological examination using non-selective media for the isolation of degrading bacteria. As far as we know, no previous study has tried to isolate degrading strains directly from heavy oil–polluted environmental samples without any prior enrichment procedure.

MATERIALS AND METHODS

Sampling. Surface sandy-gravel sediments next to Cíes Islands (42° 13.50' N, 08° 53.57' W, National Park of Atlantic Islands, Vigo, Spain) (Station 1; Fig. 1) were collected by CIS (Centro de Investigaciones Submari-

gas, Vigo) SCUBA divers in the subtidal zone (4 m water depth), 3 mo after the first ‘Prestige’ black tide reached the islands. Samples were collected from the upper 10 cm of sediment by means of sterilized flasks.

Once collected, samples were stored at 4°C and sent to the laboratory, where those for chemical analyses were frozen (–20°C). Microbiological analysis of Station 1 was conducted immediately after sampling. Other samples for chemical analysis were collected at the same time from around the islands at the stations shown in Fig. 1, using a box-core dredge (10 × 16 cm) and recovering the surface layer (1 to 2 cm, approximately 300 g).

Chemical analysis. The sediment samples (about 5 g) were spiked with the surrogates (anthracene-\textsubscript{10}, pyrene-\textsubscript{10} and benzo[a]pyrene-\textsubscript{12}) and extracted with a pressurized solvent extraction apparatus (Applied Separations) with hexane-acetone (1:1) at 110°C for 10 min in 3 cycles. The recovered extracts were treated overnight with recently activated copper for elemental sulfur removal, then carefully evaporated to near dryness, and dissolved with 0.5 ml of n-hexane for further fractionation into aliphatic and aromatic hydrocarbons.

Fractionation was performed by column chromatography with 1 g anhydrous sodium sulfate (top), 6 g neutral alumina (middle) (activated at 350°C, 5% water deactivated) and 6 g silica gel (bottom) (activated at 120°C and 5% water deactivated). Two fractions were collected: (1) aliphatic hydrocarbons, eluted with 20 ml of n-hexane and (2) polycyclic aromatic hydrocarbons with 40 ml of n-hexane:dichloromethane (80:20). The
collected extracts were concentrated and analyzed by gas chromatography coupled to mass spectrometry (GC-MS) using a Thermo-Electron Corporation system in the electron impact (EI) mode at 70 ev. The gas chromatograph was equipped with a split/splitless injector (splitless time: 0.80 min, split flow 50 ml min⁻¹) and a HP-5 MS capillary column of 30 m × 0.25 mm i.d. × 0.25 mm film (J&W Scientific). The initial column temperature was held for 1 min at 70°C, then programmed to increase at a rate of 15°C min⁻¹ to 150°C and then at 6°C min⁻¹ to a final temperature of 320°C, which was held for 10 min. The carrier gas was helium at a constant flow of 1.2 ml min⁻¹. The injector temperature was 310°C, and the transfer line and ion source were held at 250 and 200°C, respectively. Data were acquired in the full scan mode from 50 to 490 amu (2.4 scans s⁻¹) with 5 min of solvent delay and processed by the X-calibur software.

The peak areas of the target analytes were measured in the reconstructed ion chromatograms at m/z 85 for aliphatics, m/z 217 and 218 for steranes, m/z 191 for hopanes and the molecular ion for the aromatic hydrocarbons. Recoveries ranged from 70 to 110%, except for naphthalene, where it was 40 to 60% due to its higher volatility. The relative standard deviation (RSD) of the molecular biomarker indices was <5%.

**Bacterial isolation.** Results from chemical analysis showed Station 1 as the most polluted. Sediments from this place were used to isolate bacteria. It seems that the sea currents or new input of pollutants, which still occurred at the time of sampling, made this part of the estuary hold a greater amount of pollution compared with neighboring areas.

Ten-fold dilutions of the sediments from Station 1 were spread directly onto agar plates containing 1/5 strength of marine broth (Cultimed) and 1.5% (w/v) agar. Colony-forming units (CFUs) were determined after incubation at 20°C for 7 d, in order to assess the most abundant culturable bacteria. For morphological and biochemical characterization, pure cultures were obtained from the most abundant culturable bacteria by streaking them on marine agar and storing them in marine broth with 15% glycerol at −80°C for subsequent analysis.

**Enumeration of heterotrophic and hydrocarbon-degrading microbial populations.** Bacterial counts from the surface sediments were performed using the miniaturized most probable number (MPN) method performed in 96-well microtiter plates with 8 replicate wells per dilution (Wrenn & Venosa 1996). Total heterotrophs were counted in tryptone soy broth (TSB), aliphatic degraders were counted in a mineral medium called BMTM (Hareland et al. 1975), containing 1 g n-hexadecane l⁻¹, and aromatic hydrocarbon degraders were counted in BMTM containing a mixture of phenanthrene (0.5 g l⁻¹), fluorene, anthracene and dibenzothiophene (each at a final concentration of 0.05 g l⁻¹). All the media were supplemented with 3% NaCl. Hydrocarbon stocks were previously filtered through Teflon filters (Fluoropore 0.22 µm PTFE, Millipore).

BMTM mineral medium was composed of 18.6 mM KH₂PO₄·3H₂O; 7.2 mM NaH₂PO₄·H₂O; 37 mM NH₄Cl; 0.8 mM MgSO₄·7H₂O; 43 µM FeSO₄·7H₂O; 17.8 µM MnSO₄·H₂O; 10.5 µM ZnSO₄·7H₂O; 4.2 µM CoCl₂·6H₂O and 0.5 mM nitrilotriacetic acid disodium salt.

**Isolation and identification of hydrocarbon-degrading bacteria.** All isolated bacteria were screened for hydrocarbon degradation capabilities by growth in different selective media. Two media were used: one for the detection of alkane degradation and another to detect aromatic degradation activity. Both were tested on each culture using solid medium (Petri plates) and liquid medium (96-well microtitre sterile plates). Solid medium was used for the detection of alkane degraders on parafilm-sealed Petri plates containing mineral medium agar (BMTM 3% NaCl solidified with purified agar) with sterile cotton soaked in hexadecane on their lids as the sole source of C and energy as previously described. For the isolation of aromatic degraders, naphthalene pebbles, instead of cotton, were used. Screening was carried out in duplicate for each. A negative control (bacteria growing only on purified agar) and a positive control (in marine broth) were also included in the analysis.

The second test used to detect hydrocarbon-degrading strains was carried out in liquid mineral medium (BMTM + 3% NaCl) containing either n-hexadecane or a PAH mixture at the same concentrations described above for MPN enumeration. The strains were cultured overnight at room temperature in TSB. Cells were harvested by centrifugation (4000×g for 15 min), washed twice, and finally suspended in mineral medium (BMTM + 3% NaCl) to reach an optical density (OD) of around 0.5, as determined using a multi-scan spectrophotometer (Labsystems) at 620 nm. A total of 20 µl of suspended cells was used for the inoculation of 2 wells per plate. Plates with BMTM without hydrocarbons were inoculated with each strain and were used as negative controls.

**DNA extraction and PCR amplification of the 16S rRNA gene.** Colonies were picked up from the pure cultures, suspended in 100 µl of sterile milliQ water (Sigma-Aldrich Co.), boiled for 10 min, and centrifuged for 5 min at 12,000 × g. Then, 2 µl of supernatant was used as template DNA for PCR analysis. The complete 16S rRNA gene was amplified using the primers F27 and R1492 as previously described (Edwards et al. 1989, Lane 1991). All PCR amplifications were performed with a Perkin-Elmer GeneAmp 2700 Thermoc-
cycler (Applied Biosystems). The PCR reaction (25 µl) included 10 mM Tris, 50 mM KCl (pH 8.3), 2.5 mM MgCl₂, 400 µM of each deoxynucleotide, 1.25 U of Taq DNA polymerase (Amplitaq; PE Applied Biosystems), 0.4 µM of each primer and 100 ng of template DNA. The reaction mixtures were subjected to a hot start (5 min at 95°C) and after that to the following thermal cycling parameters: (1) 5 min at 95°C; (2) 40 cycles, with 1 cycle consisting of 30 s at 96°C for denaturation, 30 s at 55°C for annealing and 1.5 min at 72°C for elongation; and (3) a final extension step of 10 min at 72°C. PCR products were visualized in a 1% (w/v) agarose gel in 1× TAE buffer stained with ethidium bromide (0.6 mg ml⁻¹).

Sequencing and phylogenetic analysis. In order to eliminate the excess of primers and dNTPs for sequencing reactions, the PCR products were digested at 37°C for 1 h using shrimp alkaline phosphatase (SAP) (1 U µl⁻¹) and Exonuclease I (Exol) (10 U µl⁻¹) (U.S.B. Corporation). The enzymes were afterwards inactivated by heating the samples at 80°C for 15 min.

Sequencing was accomplished using the ABI Prism Big Dye Terminator cycle-sequencing reaction kit (Version 3.1) and an ABI Prism 3700 automated sequencer (PE Applied Biosystems) following the manufacturer’s instructions. 16S rRNA genes were fully sequenced in both directions using a set of 6 universal primers F27, R1492, F341, R907 (Edwards et al. 1989, Lane 1991), PSL and PSR (Campbell et al. 1995). Sequences were inspected, corrected and assembled into a single consensus sequence for each phylotype. After that, the sequences were examined with the BLAST search alignment tool comparison software (BLASTN) (Altschul et al. 1990) to detect the closest bacterial group to each strain from the GenBank database.

Sequences from all phylotypes were aligned with reference sequences obtained from GenBank using ClustalX (Thompson et al. 1997). The alignment obtained was transferred to MacClade (Maddison & Maddison 2003) and was modified using the software GBlocks (Castresana 2000) to eliminate poorly informative regions of the DNA alignment, which are convenient to delete prior to phylogenetic analysis.

The alignment obtained was finally edited using the MacClade program and directly transferred to Version 4.0b10 of PAUP* software (Swofford 2000). ModelTest software Version 3.6 (Posada & Crandall 1998) was run as a guide to determine the best-fit maximum-likelihood (ML) model for the edited alignment. We incorporated the best-fit model of nucleotide evolution, calculated by ModelTest, into the software PAUP* and PHYML (Guidon & Gascuel 2003), which uses a single, fast and accurate algorithm to estimate large phylogenies by ML. Finally, the trees created by PHYML and PAUP were edited using the software TreeViewX (Page 1996).

**Nucleotide sequence accession numbers.** The 37 bacterial 16S rDNA sequences reported in this paper are available under GenBank Accession Numbers EU195920 to EU195956.

**RESULTS**

**Chemical analysis**

The GC profiles of the aliphatic fraction of the subtidal sediment of the Cíes Islands (Station 1; Fig. 2) exhibited clear evidence of a fresh petrogenic contamination, based on occurrence of the homologous series of C₁₇ to C₄₀ n-alkanes overlying an unresolved complex mixture of hydrocarbons. In contrast, the profiles corresponding to the other samples (e.g. Station 3; Fig. 2) exhibited the general features of rather pristine coastal environments, with the predominance of terrestrial biogenic (higher plant) C₂₅ to C₃₃ n-alkanes with odd-even carbon number predominance and an almost absent unresolved complex mixture of hydrocarbons (Tolosa et al. 1996b). Station 5 is an exception, located inside the Vigo estuary, possibly influenced by local hydrocarbon inputs.

Confirmation of the presence of ‘Prestige’ oil in the Cíes Islands samples was obtained by a detailed study of the fossil biomarkers, namely steranes and triterpanes, currently used for oil spill fingerprinting (Daling et al. 2002). The diagnostic molecular parameters (Fig. 3, Table 1) indicate a clear correspondence with...
those of the fuel oil for Station 1, whereas Station 5 (inside the estuary) exhibits a pattern clearly different from the others.

Concurrently, the aromatic fraction of the Cíes sediments was dominated by the series of petrogenic alkyl phenanthrenes and dibenzothiophenes (Table 2), whereas the other stations exhibited distributions consistent with mixed petrogenic–pyrolytic sources, common in coastal sediments of urban/industrial areas and attributed to chronic runoff inputs. These are characterized by similar proportions of 2 to 4 aromatic ring alkylated components and the series of parent 4- to 6-ring PAHs, encompassing the benzof[b]+[k]fluoranthenes and benzo[a]+[b]pyrenes (Tolosa et al. 1996a).

The PAH concentrations reported in Table 2 (with the exception of Station 5) indicated moderate pollution in the subtidal samples from the Cíes Islands and low levels in other areas (77 to 213 mg kg⁻¹ dry wt), on the order of those found on the continental shelf, inside the Vigo estuary (1858 mg kg⁻¹ dry wt). The ratios of C₂ and C₃ dibenzothiophenes and phenanthrene/anthracenes (D₂/P₂ and D₃/P₃), proposed for differentiating sources of spilled oils in sediments (Douglas et al. 1996), also support the presence of ‘Prestige’ oil in the Cíes sediment samples (Fig. 3).

Enumeration of heterotrophic, alkane-degrading and polyaromatic-degrading microbial populations

At Station 1, the MPN results (Fig. 4) suggest a bacterial community composed of a moderately high MPN of heterotrophs (10⁵ MPN g⁻¹ sediment) of which almost 70% (10⁴ to 10⁵ MPN g⁻¹) belonged to the alkane-degrading population. The aromatic-degrading bacteria were also present, although in a much lower proportion (around 10³ MPN g⁻¹).

Culturable bacteria isolation

Ten-fold dilutions of the sediments were spread onto marine agar 1/5. More than 60 different bacteria were randomly isolated from the most diluted plates in order to include the most frequent culturable bacteria present in the polluted sediments. Only 51 isolated bacterial strains were able to maintain their viability throughout the whole experiment.

Bacterial identification and phylogenetic analysis

A total of 51 16S rRNA gene sequences were first compared with each other to detect identical sequences. Only 10 different bacterial populations were identified, each one belonging to a different taxon, according to the literature (Table 3).

### Table 1. Diagnostic indices used as source and weathering indicators for the ‘Prestige’ oil samples in sediments

<table>
<thead>
<tr>
<th>Index</th>
<th>Definition</th>
<th>Structures</th>
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<tbody>
<tr>
<td>27Ts</td>
<td>100 × Ts/(Ts+Tm)</td>
<td>Ts: 18α(H)-22, 29, 30-trisnorhopane Tm: 17α(H)-22, 29, 30-trisnorhopane</td>
</tr>
<tr>
<td>29qβ</td>
<td>100 × 29qβ/(29qβ+30qβ)</td>
<td>29qβ: 17α(H), 21β(H)-30-norhopane × 30qβ: 17α(H), 21β(H)-hopane</td>
</tr>
<tr>
<td>27dia</td>
<td>100 × 27d(R+S)/(27d(R+S)+27ββ(R+S))</td>
<td>27d: 13β(H), 17α(H)-diacholestane (20S and 20R) 27ββ: 14β(H), 17β(H)-cholestane (20R and 20S)</td>
</tr>
<tr>
<td>29ααS</td>
<td>100 × 29ααS/(29ααS+29ααR)</td>
<td>29αα: 24-ethyl-14α(H), 17α(H)-cholestane (20S and 20R)</td>
</tr>
<tr>
<td>29ββ(R+S)</td>
<td>100 × 29ββ(R+S)/(29ββ(R+S)+29αα(R+S))</td>
<td>29ββ: 24-ethyl-14β(H), 17β(H)-cholestane (20R and 20S) 27ββ: 14β(H), 17β(H)-cholestane (20R and 20S)</td>
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<tr>
<td>27ββ</td>
<td>100 × [27ββ(R+S)]/[27ββ(R+S)+28ββ(R+S)+29ββ(R+S)]</td>
<td>27ββ: 14β(H), 17β(H)-cholestane (20R and 20S) 28ββ: 24-methyl-14β(H), 17β(H)-cholestane (20R and 20S)</td>
</tr>
<tr>
<td>29ββ</td>
<td>100 × 29ββ(R+S)/(29ββ(R+S)+28ββ(R+S)+29ββ(R+S))</td>
<td>29ββ: 24-ethyl-14β(H), 17β(H)-cholestane (20R and 20S)</td>
</tr>
<tr>
<td>D2/P2</td>
<td>100 × D2/(D2+P2)</td>
<td>Dimethylbenzothiophenes (D2) and phenanthenes (P2)</td>
</tr>
<tr>
<td>D3/P3</td>
<td>100 × D3/(D3+P3)</td>
<td>Trimethylbenzothiophenes (D3) and phenanthenes (P3)</td>
</tr>
</tbody>
</table>
sequences by using Clustal X. Of these, 37 were different (Table 3) and were used to construct a whole alignment with sequences of known type species. After the first modifications in MacClade (cutting off left and right ends of the alignment), the number of characters in the alignment (approximately 1300) was reduced to only 737 after using GBlocks to make the alignment more suitable for phylogenetic analysis. Type strains from genera belonging to the phyla **Proteobacteria** and **Firmicutes** and members of the **Bacteroidetes** were used. The total alignment of 87 sequences (37 phylotypes plus type strain sequences) was used to estimate the taxonomic position of phylotypes by maximum likelihood, following the Tamura-Nei (TrN) model recommended by ModelTest. Phylogenetic analysis showed that most of our bacteria belonged to the **Gammaproteobacteria** (29 out of 37), followed by the **Alphaproteobacteria** (7 out of 37), **Bacteroidetes** (4 out of 37) and **Firmicutes** (4 out of 37). The highest richness was found in the **Gammaproteobacteria** group with strains of the genera *Marinomonas*, *Shewanella*, *Vibrio*, *Psychrobacter*, *Alcanivorax* and *Pseudomonas*. Among the **Alphaproteobacteria**, strains similar to the genus *Roseobacter* and others of difficult adscription were detected. Four strains were similar to members of the **Bacteroidetes** group, e.g. the genera *Brumimicrobium*, *Dokdonia*, *Winogradskyella* and *Cytophaga*, whilst the rest showed high similarity to **Firmicutes**, e.g. *Bacillus* and *Staphylococcus*.

**Table 2. Average polycyclic aromatic hydrocarbon (PAH) concentration (in mg kg dry wt⁻¹) of sediments from the Cíes Islands and surrounding areas (stations as in Fig. 1).** Naph: naphtalene; Phen: phenanthrene; A: anthracene; Fl: fluoranthene; Py: pyrene; D: dibenzoanthiophene; BaA: benzo[a]anthracene; C: chrysene; B(b+k)Fl: benzo[b+k]fluoranthene; BePy: benzo[e]pyrene; BaPy: benzo[a]pyrene; Per: perylene; DBA: dibenz[a]anthracene; BPer: benzo[ghi]perylen; IPy: indeno[1,2,3-cd]pyrene; N1 to N3, P1 to P3, D1 to D3 and C1 to C3: the methyl to trimethyl derivatives

<table>
<thead>
<tr>
<th>PAH</th>
<th>Stn 1</th>
<th>Stn 2</th>
<th>Stn 3</th>
<th>Stn 4</th>
<th>Stn 5</th>
<th>Stn 6</th>
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<tbody>
<tr>
<td>Naph</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>–</td>
<td>20</td>
<td>2</td>
</tr>
<tr>
<td>Naph 1</td>
<td>2</td>
<td>1</td>
<td>10</td>
<td>2</td>
<td>83</td>
<td>6</td>
</tr>
<tr>
<td>Naph 2</td>
<td>5</td>
<td>3</td>
<td>24</td>
<td>5</td>
<td>125</td>
<td>17</td>
</tr>
<tr>
<td>Naph 3</td>
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<td>25</td>
<td>5</td>
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<td>2</td>
<td>46</td>
<td>3</td>
</tr>
<tr>
<td>A</td>
<td>2</td>
<td>–</td>
<td>1</td>
<td>2</td>
<td>11</td>
<td>–</td>
</tr>
<tr>
<td>Phen 1</td>
<td>237</td>
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<td>10</td>
<td>3</td>
<td>152</td>
<td>14</td>
</tr>
<tr>
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<td>1</td>
<td>7</td>
<td>15</td>
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**DISCUSSION**

The oil-polluted sediment selected for isolation of bacteria in the current study presented a high degree of fresh hydrocarbon pollution characterized by fingerprint indexes of unweathered fuel from the ‘Prestige’ spill and a high preservation of the n-alkane series (Fig. 2), which may still have supported the bloom of obligate hydrocarbonoclastic bacteria at the time of sampling.
Petroleum pollution may stimulate the growth of hydrocarbon-degrading microorganisms that usually exist in low abundance in sediments, shifting the community structure of the affected sites. Previous studies have reported dominance of Gammaproteobacteria in bacterial communities from oil-affected marine habitats. However, the samples in these experiments had been recently affected (Roling et al. 2002), or had been retrieved from places where natural attenuation proceeded slowly, such as the Sea of Japan (Kasai et al. 2001).

However, communities from samples enriched with nutrients, or from places where high nutrient levels (0.8 mg l⁻¹ of total nitrogen) naturally occur (Macnaughton et al. 1999), presented Gammaproteobacteria-dominated communities, which were rapidly substituted by Alphaproteobacteria due to higher rates of biodegradation. Disappearance of Gammaproteobacteria was linked to a loss of Alcanivorax dominance in the community, and this change in the community was faster when amended with nutrients (Roling et al. 2002, 2004). In most cases in which Alcanivorax spp.
have been found to be dominant in oil-impacted environments, samples were analyzed within days of the oil pollution or bioremediation event. In samples retrieved >8 wk after the start of the experiments (Kasai et al. 2001, Maruyama et al. 2003), no such dominance was detected. In these cases, it is possible that the Alcanivorax spp. bloom had already occurred (Macnaughton et al. 1999), or that the nutrient levels and alkane percentage of the spilled oil, critical for the growth of this genus, were too low (‘Nakhodka’, Sea of Japan) (Kasai et al. 2001, 2002). All this is consistent with the observation that, after an initial rapid increase in population size, Alcanivorax spp. decline to much lower numbers within weeks (Syutsubo et al. 2001, Roling et al. 2002, 2004).

Polluted samples considered in the present study were retrieved 3 mo after the last fuel input from the ‘Prestige’. Since low amounts of alkanes were present in the ‘Prestige’ oil and high biodegradation rates were expected in the ría (Medina-Bellver et al. 2005), such a length of time would have already produced the Alcanivorax bloom long before sampling. Indeed, degradation of the fuel present at the rest of the stations was quite advanced, compared to that at Station 1. Sea currents probably transported buried oil from other places to Station 1, supplying its sediments with unweathered oil (Fig. 3) rich in alkanes (Fig. 2), which enabled the growth and, thus, the isolation of Alcanivorax at that station.

Most of the isolated bacteria found in Cíes sediments were identified as members of the Gammaproteobacteria (59%) (Table 3). Members of Alteromonas, Shewanella, Pseudoalteromonas, Vibrio, Marinomonas, Psychrobacter and Alcanivorax were identified. Although only the 7 strains (Table 3) from the phylotype close to Alcanivorax borkumensis (Fig. 5) were able to degrade alkanes, hydrocarbon-degrading strains have been previously described in all the isolated genera. For example, phenanthrene- and chrysene-degrading activities have been observed among members of the genera Vibrio, Pseudoalteromonas and Marinomonas (Melcher et al. 2002). Shewanella spp. able to grow on crude oil, tetradecane and naphthalene (Gentile et al. 2003) have also been detected.

Concerning the Alphaproteobacteria group, among the 4 isolated strains, P73 corresponded to Roseobacter sp. (100%) and Strain P128 belonged to the genus Agrobacterium (99%) (Table 3). Some species of these 2 genera have been described as PAH degraders (Widdel & Rabus 2001). Finally, 4 non-hydrocarbon-degrading strains, belonging to the Bacteroidetes and Firmicutes groups, were isolated. Only some species of the genus Staphylococcus have been previously described as naphthalene degraders (Zhuang et al. 2003).

The PAHs present in the ‘Prestige’ fuel were mainly of high molecular weight (3 or more aromatic rings), leading to a low PAH bioavailability. The PAH concentrations measured at the time of sampling were very low (around 2.5 µg g⁻¹ for Station 1; Table1). The small population (10³ MPN g⁻¹) of PAH-degrading bacteria detected could be explained by this low concentration and bioavailability. The fact that no aromatic-degrading activity was detected among the isolated bacteria did not necessarily mean that they do not play a role in the biodegradation processes. Microbial communities are mainly naturally selected by their metabolic cooperation. Although several members of the microbial community could be secondary hydrocarbon degraders, their success could be related to co-metabolism, supply of growth factors for solubilization of substrates and/or elimination of toxic metabolites (Kanaly et al. 2002). In a previous study on a hydrocarbon-degrading
co-culture, the importance of a non-degrading strain that eliminates an intermediate metabolite produced by the hydrocarbon-degrading strain was demonstrated (Casellas et al. 1998). Obviously, taking into account that the specific richness of the present study has been detected on diluted marine agar-culturable bacteria, other non-culturable species could fulfill critical roles in the global microbial community, as recently described in a hydrocarbon-degrading consortium (Vinás et al. 2005b). All but one of the isolated genera exhibited an absence of hydrocarbon-degrading capabilities. The low amount of degrading bacteria isolated was expected from the beginning, since no selective medium was used to isolate degrading bacteria.

The Russian tanker ‘Nakhodka’ released heavy oil, with a composition very similar to that of the ‘Prestige’ spill, which covered >500 km of the Japanese coastline. The levels of N and P (0.1 and 0.01 mg l⁻¹, respectively) in the Sea of Japan are relatively small, which may have hindered Alcanivorax from dominating the microbial community. When N and P were added in adequate quantities to cultures of seawater with crude oil as the only source of C, Alcanivorax became dominant and the rate of biodegradation was strongly promoted (Kasai et al. 2001, 2002, Roling et al. 2002). However, this effect could only be shown in batch cultures and was not observed in situ, due to low levels of naturally occurring nutrients.

Since the oil from the ‘Nakhodka’ had a similar composition to that from the ‘Prestige’ (heavy fuel) and Alcanivorax phyotypes were isolated in high proportion without nutrient amendments in the Vigo estuary, it can be hypothesized that the environmental conditions along the Atlantic coast could be more suitable to biodegradation than those in the Sea of Japan (total N = 0.1 mg l⁻¹). In fact, ongoing natural bioremediation has already been observed to occur along Galician coasts (Medina-Bellver et al. 2005).

In the Ría de Vigo, in addition to the boundary conditions that usually determine the patterns of circulation in estuarine systems, an additional factor must be taken into account. The coastal upwelling, as a consequence of the wind regime over the adjacent shelf, induces the inflow of subsurface oceanic eastern North Atlantic Central Water (ENAW) into the estuaries of Galicia. This inflow has a major influence on their hydrography. During the upwelling and as a consequence of the circulation pattern, part of the biomass that is produced inside the estuary is transported offshore by the outgoing surface current. Part of this exported organic matter is remineralized either in the water column or on the bottom of the continental shelf, and therefore the incoming bottom current supplies the Galician estuaries not only with new nutrients but also with remineralized nutrients through a feedback mechanism (Alvarez-Salgado et al. 1993). This fertilizing process causes the Vigo estuary to be a highly productive ecosystem that processes a considerable amount of dissolved N and P (mean values around 0.6 and 0.06 mg l⁻¹, respectively, reaching even higher concentrations at certain times of the year; Nigeria et al. 1997), which is 6 times higher than the total amount of N and P in the Sea of Japan. Furthermore, the sediments from Station 1 where sandy gravel with a low organic matter content and thus well oxygenated (Vilas et al. 2005), enhancing the degradation capacity of the aerobic Alcanivorax (Yakimov et al. 1998).

Both the high proportion of the alkane-degrading population (=70% of the total MPN bacteria) and the presence of Alcanivorax as the only n-alkane degrader could reflect an initial shift in the microbial community towards a new assemblage more adapted to hydrocarbon contamination, as previously observed in microcosm experiments (Roling et al. 2002). Taking into account that 14% of the isolates were classified as Alcanivorax borkumensis, despite the fact that this is a ‘professional hydrocarbonoclastic bacteria’, it could be concluded that a high proportion of this genus should be present in the sediments sampled, being able to compete and grow on the marine agar we used for isolation. The same method for isolation was used to count growing cells of Alcanivorax in oil-contaminated seawater supplemented with N and P fertilizers (Kasai et al. 2002), and good correlation between colonies and direct cell counts by fluorescent in situ hybridization (FISH) (Syutsubo et al. 2001) was obtained. Different genotypes of Alcanivorax might be adapted to environments with different concentrations of nutrients (Head et al. 2006), suggesting that the Alcanivorax strain isolated in this study could be more adapted to competence under high nutrient levels. Although further studies are needed to confirm that hypothesis, such an ability would be a good feature for bioremediation purposes.

Our results are in agreement with the enrichment cultures made after the ‘Nakhodka’ oil spill in the Sea of Japan (Kasai et al. 2002, Maruyama et al. 2003), but, in our case, no enrichment was necessary. The presence of fresh fuel (preservation of the n-alkane series; Figs. 2 & 3) and the high nutrient availability normally present in the Ría de Vigo support the growth of Alcanivorax, allowing its presence in quantities sufficient to be isolated without enrichment, even when competing with other heterotrophic bacteria.

Many previous studies have stated that Alcanivorax could play a critical role in the natural cleaning of oil-polluted marine systems, since the natural attenuation processes were accelerated when this species began to become dominant (Kasai et al. 2002). Unfor-
fortunately, the present study did not yield data showing whether any acceleration is actually happening. However, the presence of *Alcanivorax* could be used as a bioindicator in correlation with high rates of biodegradation. Indeed, the capacity of indigenous bacteria from the shore samples of the Galician coastline for biodegradation of the ‘Prestige’ oil has also been recently confirmed by Medina-Bellver et al. (2005).

The isolation of *Alcanivorax* strains directly from polluted sediment containing hydrocarbons is the first ecological evidence of the high natural abundance of this group of bacteria. Since the addition of fertilizers to the seawater would have no effect on the concentration of N and P in this area, the use of bioremediation, understood as the artificial addition of nutrients, is not advisable. Hydrocarbon-degrading populations seem to grow quite well by themselves under the environmental conditions of the Ría de Vigo. However, there may be exposed areas where the conditions are not as favorable and where bioremediation amendments could be effective.

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**LITERATURE CITED**

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