

ORIGINAL ARTICLE

Global diversity and biogeography of deep-sea pelagic prokaryotes

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The deep-sea is the largest biome of the biosphere, and contains more than half of the whole ocean's microbes. Uncovering their general patterns of diversity and community structure at a global scale remains a great challenge, as only fragmentary information of deep-sea microbial diversity exists based on regional-scale studies. Here we report the first globally comprehensive survey of the prokaryotic communities inhabiting the bathypelagic ocean using high-throughput sequencing of the 16S rRNA gene. This work identifies the dominant prokaryotes in the pelagic deep ocean and reveals that 50% of the operational taxonomic units (OTUs) belong to previously unknown prokaryotic taxa, most of which are rare and appear in just a few samples. We show that whereas the local richness of communities is comparable to that observed in previous regional studies, the global pool of prokaryotic taxa detected is modest (~3600 OTUs), as a high proportion of OTUs are shared among samples. The water masses appear to act as clear drivers of the geographical distribution of both particle-attached and free-living prokaryotes. In addition, we show that the deep-oceanic basins in which the bathypelagic realm is divided contain different particle-attached (but not free-living) microbial communities. The combination of the aging of the water masses and a lack of complete dispersal are identified as the main drivers for this biogeographical pattern. All together, we identify the potential of the deep ocean as a reservoir of still unknown biological diversity with a higher degree of spatial complexity than hitherto considered.

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Introduction

The pelagic dark ocean (the water column >200 m deep) contains 70% of the ocean's microbial cells and 60% of its heterotrophic activity, with a pivotal regulatory role in planetary biogeochemical cycles (Aristegui *et al.*, 2009). Yet, current knowledge of the pelagic microbial community structure of the dark ocean, the largest biome in the biosphere, is based on a pool of samples collected at specific locations (DeLong *et al.*, 2006; Martín-Cuadrado *et al.*, 2007; Brown *et al.*, 2009; Galand *et al.*, 2010; Agogué *et al.*, 2011; Eloë *et al.*, 2011; Quaiser *et al.*, 2011; Smedile *et al.*, 2012; Wang *et al.*, 2013; Wilkins *et al.*, 2013; Ganesh *et al.*, 2014) (Supplementary Figure S1) and

thus are dwarf in comparison with the analyses of upper ocean microbial communities, which have indeed been assessed at global scales (Rusch *et al.*, 2007; Yooseph *et al.*, 2007; Zinger *et al.*, 2011; Sunagawa *et al.*, 2015). Whereas the deep ocean is often considered to be a rather uniform environment, the connectivity of pelagic microbial communities may be reduced by the limited mixing between water masses (Agogué *et al.*, 2011; Hamdan *et al.*, 2013) or modulated by advection (Wilkins *et al.*, 2013) imposing limitations on the dispersion of marine microbes in this low-turbulence environment. In addition, the spatial structure of the bathypelagic ocean, organized in partially isolated basins created by the emergence of submarine mountains, has not been tested as a potential factor affecting the biogeography of pelagic microbial communities, as happens for specialized deep-sea fauna (Moalic *et al.*, 2012) and bacteria inhabiting deep-sea surface sediments (Schauer *et al.*, 2010), either by imposing limits to deep-ocean connectivity or by delineating different environments that select for distinct

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microbial communities. Therefore, the deep pelagic ocean may present a mosaic of biogeographical domains with distinct microbial assemblages, a hypothesis not yet fully tested.

We created a global collection of samples retrieved during the Malaspina 2010 circumnavigation expedition (cf. Irigoien *et al.*, 2014) and we have used high-throughput sequencing of the 16S rRNA genes jointly with automated ribosomal intergenic spacer analysis (ARISA) profiles and metagenomic data of the prokaryotes present in bathypelagic waters of the main world's oceans to describe their diversity, community structure and biogeographical distribution and identify the cosmopolitan and/or abundant prokaryotes in the dark ocean at a global scale. Moreover, we aimed to test whether deep-sea pelagic prokaryotic communities are uniform or present biogeographical patterns delineated by water mass and/or deep-oceanic basins.

Material and methods

A total of 60 water samples were taken during the Malaspina 2010 expedition (<http://scientific.expedicionmalaspina.es/>) corresponding to 30 different sampling stations globally distributed across the world's oceans (Figure 1). We focused on the samples at the depth of 4000 m, although a few samples were taken at lower depths, all well within the bathypelagic realm.

Sample collection and processing

Two different size fractions were analyzed in each station representing the free-living (0.2–0.8 μm) and

particle-attached (0.8–20 μm) prokaryotic communities (Crump *et al.*, 1999; Ghiglione *et al.*, 2009; Allen *et al.*, 2012). For each sample 120 l of sea water were sequentially filtered through a 200 and a 20 μm mesh to remove large plankton. Further filtering was done by pumping water serially through 142-mm polycarbonate membrane filters of 0.8 μm (Merk Millipore, Darmstadt, Germany, Isopore polycarbonate) and 0.2 μm (Merck Millipore, Express Plus) pore size with a peristaltic pump (Masterflex, EW-77410-10). The filters were then flash-frozen in liquid N_2 and stored at -80°C until DNA extraction. The time span from bottle closing to filter freezing was ~ 4 h and except for the time needed to empty the rosette bottles, the water was kept at 4°C . DNA extractions were performed using the standard phenol–chloroform protocol (see Supplementary Information), and prokaryotic diversity was assessed by amplicon sequencing of the V4 region of the 16S rRNA gene with the Illumina MiSeq platform (iTags) using paired-end reads (2×250 bp) and primers F515/R806 (details in Supplementary Information) targeting both Archaea and Bacteria (Caporaso *et al.*, 2011). Sequence data processing included the paired-end reads assembly, end-trimming, sequence quality control and chimera checking process integrated in the Joint Genome Intitute pipeline. Operational taxonomic units (OTUs) were obtained by clustering the processed data at a 97% identity and the taxonomic annotation of consensus sequences was performed using the SILVA v111 database (details in Supplementary Information). ARISA and metagenomic data analyses were also applied to the same samples as an independent validation of the iTag approach (see Supplementary Information).

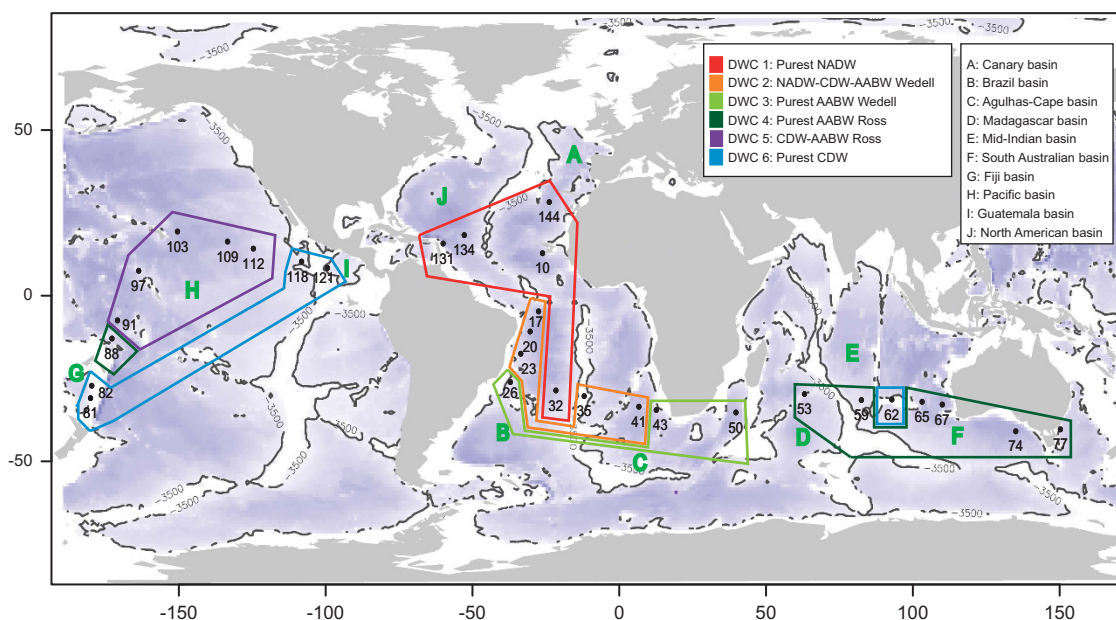


Figure 1 World map showing the location of the Malaspina sampling stations in the present study. The deep-water cluster derived from dominant water masses found at each station are color-coded, and the deep-oceanic basins defined according to bathymetry below 3500 m depth (see Materials and methods for details) are indicated with letters.

Statistical data analyses

Statistical analyses (see details below) included richness estimation and rarefaction curves. The analysis of differences in community composition among samples and their relation to potential drivers assessed by means of a combination of multivariate exploratory techniques based on Bray–Curtis similarities (non-metric multidimensional scaling) and hypothesis testing methods (permutational multivariate analysis of variance). The novelty of the obtained 16S rRNA gene sequences was checked against the SILVA, NCBI and RDP public databases using BLAST. All the sequences used in this study are publicly available at the NCBI Sequence Read Archive (<http://www.ncbi.nlm.nih.gov/Traces/sra>) under accession ID SRP031469. All statistical analyses and data treatment were conducted with the R Statistical Software (R Core Team, 2014) using version 3.0.1 and the following packages: *BiodiversityR*, *ecodist*, *gdistance*, *marelac* and *vegan*. The iTags were used as the primary data set for the whole study. ARISA and metagenomic data were used in specific analyses to compare with the iTag-derived data.

Novelty of the deep-ocean 16S rRNA gene sequences

To evaluate the novelty of the obtained 16S rRNA gene sequences, the 3507 representative OTU sequences were compared with RDP (Cole *et al.*, 2014), SILVA v111 (Pruesse *et al.*, 2007) and NCBI RefSeq (Pruitt *et al.*, 2012) public databases. The nucleotide subsets of the databases were downloaded (January 2013) and served as reference for a BLAST (Altschul *et al.*, 1990) comparison. A cutoff *e*-value of 1E-05 was used, a maximum of five target sequences were allowed for each query sequence and only the matches with coverage >90% were considered. When more than one match existed with an acceptable coverage and *e*-value, the one with the highest identity was chosen and identity values to the closest match for each sequence were collected.

Beta-diversity patterns of prokaryotic community composition

To infer the variation of the prokaryotic assemblages in space and along environmental gradients (that is, beta-diversity), the Bray–Curtis dissimilarity index was used on community composition. The OTU-abundance table obtained from the sequence clustering was sampled down to the lowest sampling effort (10 617 reads per sample), and dissimilarities between all pairs of samples were calculated using Bray–Curtis dissimilarity coefficient to obtain a beta-diversity matrix. The resulting dissimilarity matrix was used to perform a non-metric multidimensional scaling (Minchin, 1987) analysis using random starts. Permutational multivariate analysis of variance using 1000 permutations was used to test for significant differences and to partition the

beta-diversity matrix variance between groups of samples (Anderson, 2001; Anderson and Walsh, 2013).

The differential contribution ($D_{i,b}$) of a specific basin to the total abundance of a specific OTU was computed for the 30 most abundant OTUs. This was calculated for each OTU and each basin following the formula:

$$D_{i,b}(\%) = (X_{i,b} - N_b) \times 100$$

where $X_{i,b}$ is the contribution of the basin b to the total abundance of OTU i (that is, the number of reads of OTU i in the samples belonging to the basin b divided by the total number of reads of OTU i) and N_b is proportion of samples in the data set belonging to basin b . Thus, an OTU with a percentage of reads coming from a specific basin higher/lower than would be expected under an even distribution across samples would have a positive/negative $D_{i,b}$ value.

Processes shaping prokaryotic biogeography

To infer the relative importance of the processes shaping the biogeographical patterns, the relative contribution of environmental drivers and geographical distance to the beta-diversity of deep-ocean prokaryotic communities was assessed by means of a combined statistical strategy applied separately to both size fractions. First, the best subset of environmental drivers was selected using the BIOENV approach (Clarke and Ainsworth, 1993). Second, permutation-based multiple regression on matrices was used to partition the variance of the Bray–Curtis dissimilarity into (i) pure environmental variation, (ii) pure geographical variation, (iii) spatially structured environmental variation and (iv) and the unexplained variation (see details in Supplementary Information).

In addition, the scale of geographical variation was studied by means of Mantel correlograms (Oden and Sokal, 1986), which assesses the spatial correlation of multivariate data by computing a Mantel statistic (r) between the Bray–Curtis dissimilarity matrix and a matrix where pairs of sites belonging to the same geographic distance class receive value 0 and the other pairs, value 1. The process is repeated for each distance class and each r -value can be tested for significance by permutation. Distance classes of 1500 km were used. Mantel correlograms were run for each size-fraction separately.

Dominant phylum level analysis

To study the composition of prokaryotic communities at a broad taxonomic level a phylum-abundance table was derived from the OTU-abundance table by adding up all the OTUs belonging to the same phyla based on their SILVA taxonomy affiliation. For comparison with similar studies the phylum Proteobacteria was divided into its Classes. OTUs that could not be assigned to any

phyla were included into an extra category (named as others). Only the phyla represented by >0.5% of the reads in the whole data set were considered. Differences in abundance for every Phylum between Oceans (categorized as North Atlantic, South Atlantic, Indian, South Pacific and North Pacific), deep-oceanic basins and 'deep-water clusters'—see Supplementary Information—were statistically tested using multivariate analysis of variance. *P*-values were then Bonferroni-corrected for the effect of multiple comparisons.

Results and discussion

We examined pelagic prokaryotic diversity from two size fractions considered free-living and particle-attached (see Materials and methods), in 30 deep-ocean sites distributed in the North Atlantic (4 sites), South Atlantic (8), Indian (6) and South (5) and North Pacific Oceans (6), and an additional set of samples that was taken from the Southern Ocean in waters close to Australia (Figure 1). We targeted 4000 meters as the water depth of study taken as representative of the bathypelagic ocean, yet some samples were taken from shallower depths (always >2000 m). The sites were assigned to each of six different deep-water clusters according to their water mass composition (Supplementary Figure S2 and Supplementary Table S1), as well as to 'deep-oceanic basins' based on the global ocean's bathymetry (Figure 1; details in Supplementary Information). Using Illumina sequencing of the V4 region of the 16S rRNA gene, we obtained a final data set of 1 789 427 sequences (iTags) that could be constrained into a total of 3695 OTUs, which represents a minimum estimate of the richness of prokaryote taxonomic units in the deep-ocean (Supplementary Table S2). The iTag data was compared with ARISA-derived and metagenomic data revealing a good consistency between techniques (see Supplementary Information and Supplementary Figures S3 and S4). Hereafter, the downstream analyses were performed using iTags as the default data set.

Richness of bathypelagic prokaryotic communities

Two kinds of rarefaction curves were computed to check whether prokaryotic richness was close or far from saturation, both at the local (individual site/sample) and at the global (all samples) scale. For each sample, a rarefaction curve (or individual sample-based rarefaction curves) was drawn by sequentially computing the number of OTUs for an increasing number of reads. In addition, a sample-based rarefaction curve was drawn by randomly accumulating an increasing number of samples for the whole data set. Rarefaction curves for individual samples showed that prokaryotic diversity at the OTU level (97% identity cutoff) was far from saturation locally with the sequencing effort used

(Figure 2a). In contrast, when considering the global set of samples, the sample-based rarefaction curves reached a considerably flat plateau at

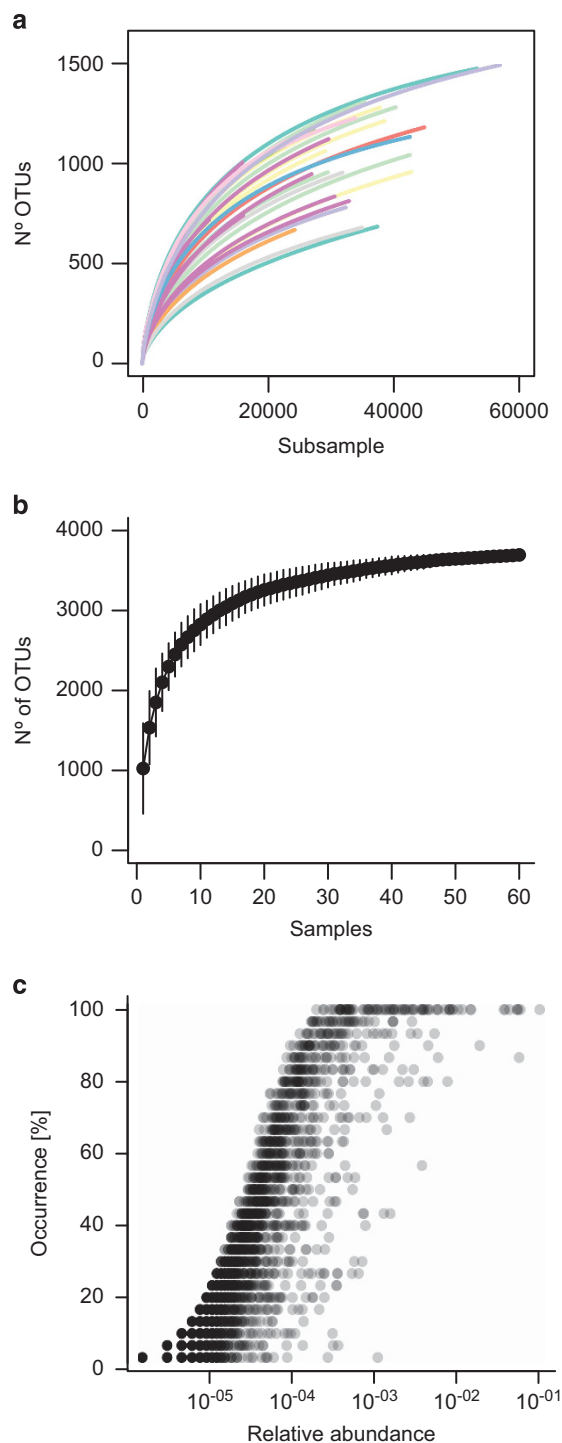


Figure 2 Rarefaction curves. (a) within samples, individual-based and (b) sample-based. Global data set relative abundance vs occurrence (that is, the percentage of samples in which an OTU occurs) for all the OTUs (c). The sample-based rarefaction curve has been calculated for the entire data set. The deep-oceanic basins to which each station belongs are indicated with different colors in (a). (legend in Figure 5). No significant differences were detected for richness/diversity (neither OTU number, Chao1 nor Shannon index) between basins.

~3500 OTUs (Figure 2b). The number of OTUs increased rapidly with the addition of the first 10 samples but once ~20 samples were considered, the addition of extra samples resulted in a small additional discovery of new OTUs. In fact, on average 42.0% of the OTUs present in one sample were shared with a second one taken at random from our data set (min = 15.7%, max = 76.2%), being these shared OTUs the ones with higher abundances (Figure 2c). This indicates that the global deep ocean contains a relatively modest number of prokaryotic phylotypes, likely in the order of a few thousands. The total number of OTUs identified here, 3695, represents, however, a minimum estimate, as strict data-cleaning criteria have been used in the data processing (specially, the removal of possible chimeric sequences and singletons) and as additional OTUs are likely to be present in areas not sampled in this study, such as the Arctic or Antarctica.

Each deep-sea prokaryotic community sampled here can be thus considered to be composed of (i) a set of dominant species shared with the rest of the stations in varying proportions, which we estimate at ~42% of the OTUs identified and (ii) a set of low abundant and relatively sample-specific (that is, highly unshared) set of taxa comprising a 'rare biosphere' (Sogin *et al.*, 2006; Pedrós-Alió, 2012) of the global pool of deep-sea prokaryotes.

To prevent artifacts during diversity/richness estimations owing to uneven sampling efforts among samples, the data set was randomly sampled down to the lowest sequencing effort (10 617 reads per sample). This resulted in 637 020 reads corresponding to a total of 3543 OTUs. The number of OTUs in each community ranged from 248 to 896 (mean = 659.1, s.d. = 146.0), comparable to the mean local richness reported before in the Atlantic Ocean (mean = 835, s.d. = 421; Agogué *et al.*, 2011). Slightly higher values (mean = 1037.3, s.d. = 173.6) had previously been reported in bathypelagic samples from polar and mid-latitudes (see samples below 1000 m from Supplementary Table 1 in Ghiglione *et al.*, 2012). Thus, the global assessment of bathypelagic prokaryotic communities that we report delivers local richness values comparable to those observed in previous regional-scale bathypelagic surveys. However, our estimate of the total prokaryotic richness in the bathypelagic ocean estimated at ~3600 OTUs is consistent with previous estimates that found a total richness of 10 846 OTUs with half of them corresponding to singletons (Zinger *et al.*, 2011). This represents a small fraction (~3% and 5.5%) of the total oceanic plankton bacterial richness found by recent surveys with comparable methodologies: a previous study combining 509 benthic and pelagic marine samples ranging from 0 to 5400 m depth found a total richness of ~120 000 OTUs (Zinger *et al.*, 2011), whereas a total richness of ~65 500 OTUs was detected in a different study using data from 277 epipelagic samples (243 of

which were also included in the previous one) from the Arctic, Atlantic, Pacific and Southern Oceans (Sul *et al.*, 2013). This would suggest that only a small fraction of all oceanic microbes are found in the deep ocean.

Novelty of bathypelagic prokaryotic lineages

We assessed the degree of novelty of bathypelagic prokaryotic diversity by comparing the detected 16S rRNA gene sequences to those present in public databases. OTU representative sequences were compared with the RDP, SILVA and NCBI databases using 95% and 97/99% identity values as proxies for genus and 'species' level, respectively. The three databases provided comparable identity distributions with two clear peaks, the first one at ~95% identity and a second peak near the 100% identity (Supplementary Figure S5). Interestingly, around one-third of the OTUs had identity values lower than 95%, half of the OTUs lower than 97% and 2/3 of the OTUs had values lower than 99%. Although the 97% identity is widely used in microbial ecology studies as a broad proxy for 'species' cutoff (Stackebrandt and Goebel, 1994; Cohan, 2002; Hagström *et al.*, 2002), it is well known that this value may integrate different species and overlook putative ecotypes within species with different ecological roles (Fox *et al.*, 1992; Acinas *et al.*, 2004; Stackebrandt, 2006). Therefore, it is safe to assume that we detected at least between 1687 (at 97%) and 2385 (at 99%) putative new prokaryotic OTUs as well as 986 OTUs belonging to putative new genera (at 95%) not present in the standard prokaryotic rRNA gene databases (corresponding to 45.7%, 64.6% and a 26.7% of the total OTUs, respectively). However, these novel lineages represented a minor fraction of the reads (4.5%, 9.1% and 2.2%, respectively) and thus they are likely members of the bathypelagic 'rare biosphere'. This pattern had already been observed in a single sample from a hydrothermal vent (Sogin *et al.*, 2006) where novel sequences belonged to very low abundant OTUs. This result suggests that the prokaryotic assemblages in the bathypelagic ocean are composed of a combination of a set of relatively abundant and widely distributed species, already detected in previous environmental surveys, and a set of rare species with limited distributions where most of the genetic novelty accumulates.

Dominant prokaryotes in the bathypelagic ocean

We aimed at identifying the most abundant prokaryotes present in the deep ocean at a global scale. The relative abundance (that is, proportion of reads) of every phylum (except proteobacteria, which were divided into classes) was highly similar among samples (Supplementary Figure S6): Gammaproteobacteria, Alphaproteobacteria, Actinobacteria, Thaumarchaeota and Deltaproteobacteria

dominated in all the stations in terms of relative abundance. Gammaproteobacteria was the most abundant group in all the sites, consistent with the previously described increase of their contribution to the total number of bacteria with depth (López-García *et al.*, 2001). Here, the proportion of members of Archaea ranged from 2.2% to 16.3% of the total reads in both fractions combined. This estimation is considerably lower than previous studies in which Archaea had been reported to be between 20 and 30% of the total of bacterioplankton in the deep ocean (DeLong, 1992,2003; Massana *et al.*, 1997) or even higher (39%) (Karner *et al.*, 2001). Yet, Archaea in the free-living samples reached up to 25–30% of the total in specific locations (stations 10, 81, 112, 118 and 121 located in the North Atlantic and North Pacific). Our findings are not attributable to PCR biases or primer mismatches as relative abundances of both Euryarchaeota and Thaumarchaeota obtained from the metagenomic data set were similar (Supplementary Figure S4). In all samples, Thaumarchaeota dominated over Euryarchaeota

(1.7–14% vs 0.29–3.7%) as described before for bathypelagic waters (Herndl *et al.*, 2005; Teira *et al.*, 2006). Only the Actinobacteria phylum had differences in relative abundance between oceans and between deep-oceanic basins (as tested by analysis of variance with bonferroni correction: $F=5.8$, $P=0.001$, $P_{\text{corrected}}=0.016$ and $F=4.9$, $P=0.003$, $P_{\text{corrected}}=0.045$ for Ocean and Basins, respectively) being more abundant in the North and South Pacific and in particular the Pacific and Guatemala Basins.

Despite the invariant composition of prokaryotic communities at a the phylum level, the distribution of the 30 most abundant OTUs (Figure 3, Supplementary Table S3) included only a few cosmopolitan organisms that were relatively evenly distributed along the whole data set: for example, the first, second and fifth most abundant OTUs, representatives of the *Alteromonas* genus, the Marine Group I Thaumarchaeota (MGI) and the SAR324 clade, respectively. This is consistent with current knowledge on the ecology of these three

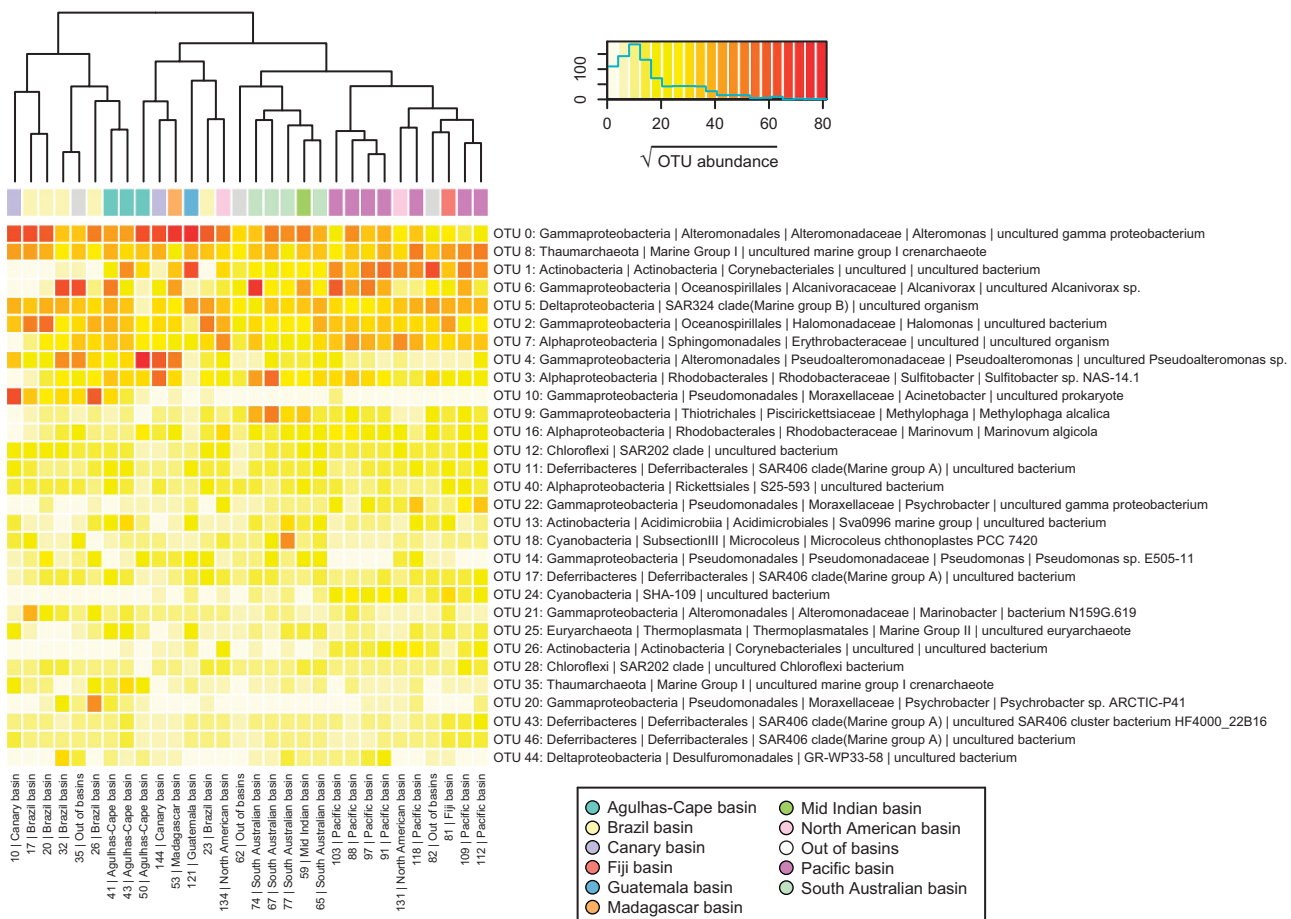


Figure 3 Heatmap representing the square root of abundances (number of reads) of the 30 most abundant OTUs (rows) along the 30 stations (columns). Subsampled abundances to the minimum sequencing depth (10 617 reads per sample) have been used for comparison and data from the two size fractions within a station was summed after subsampling. The deep-oceanic basins to which each station belongs are indicated at the top (see color legend). Taxonomical annotation for each OTU is based on the SILVA taxonomic assignment of each OTU representative sequence. OTUs are ordered top to bottom based on their global abundance in the whole data set.

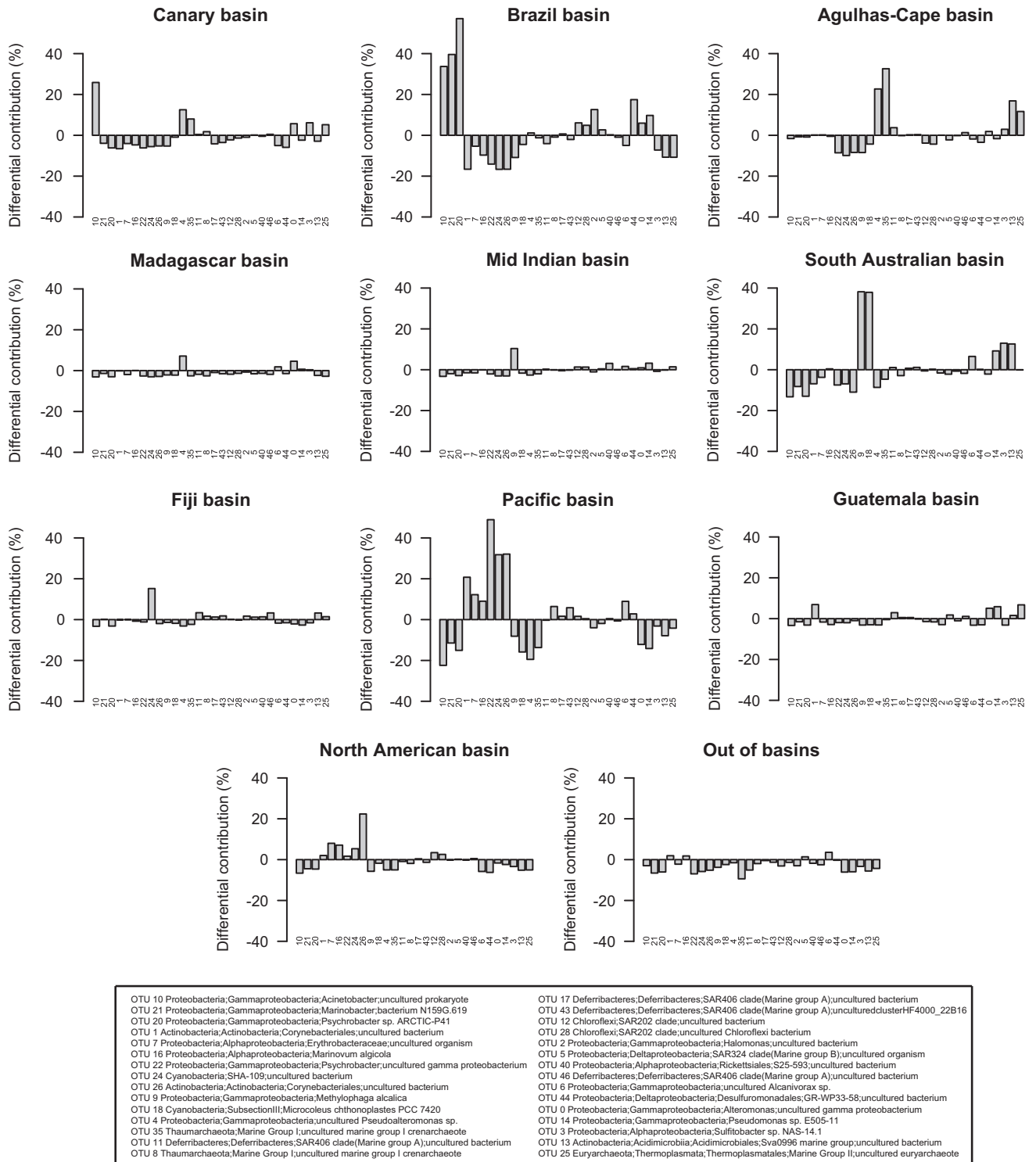


Figure 4 Differential contribution ($D_{i,b}$; in %) of each basin to the total abundance of each of the 30 most abundant OTUs (see Supplementary Information for calculation details). Numbers below each bar represent each OTU, whose taxonomical affiliation is described in the legend, based on SILVA taxonomy. OTUs are the same as in Figure 3 but ordered using a clustering based on $D_{i,b}$ values (details not shown) for a clearer visualization.

groups: the existence of a deep *Alteromonas macleodii* ecotype (identical at 16S rRNA gene sequence level to our most abundant OTU) with specific adaptations to deep-ocean conditions is well known (López-López *et al.*, 2005; Ivars-Martinez *et al.*, 2008) and

the MGI archaeal group, jointly with Marine Group II Euryarchaeota, are the most abundant Archaea in the ocean (Massana *et al.*, 2000). The SAR324 Deltaproteobacteria clade has also been described as a typical deep-sea group (López-García *et al.*, 2001; Agogué

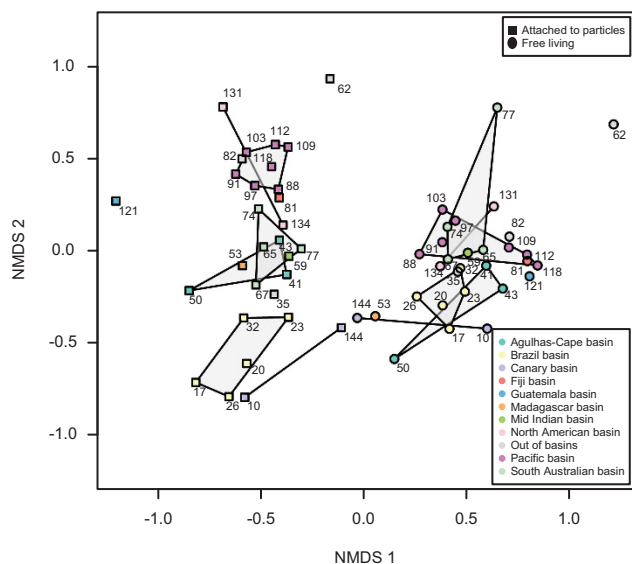


Figure 5 Non-metric multidimensional scaling (NMDS) analysis of beta-diversity (Bray–Curtis distances) for the 60 samples in the data set based on iTags. Size-fraction is coded with point style (squares, attached and circles, free-living) and deep-oceanic basins following color codes (see legends). Numbers close to each sample represent the station number (see Figure 1).

et al., 2011). Most of the rest of the dominating OTUs exhibited uneven abundances throughout the world's deep oceans, with most of them restricted to a specific geographical region: for example, *Alcanivorax* sp. and an uncultured Actinobacteria representatives were nearly absent from the Atlantic, whereas a *Pseudoalteromonas* sp. representative was nearly absent from the Pacific but abundant in the rest of the sites. As a result of the heterogeneity in the distribution of the most abundant organisms, the samples tended to cluster with other geographically close samples (Figure 3).

Differential OTU distribution through deep-oceanic basins

In addition, we calculated the differential contribution (in %) of a specific basin to the abundance of a specific OTU for the 30 most abundant OTUs (Figure 4). Within these, only a few did not have a clear differential contribution associated to a specific basin and thus, and as mentioned before (see Figure 3), these OTUs were equally distributed among basins (that is, were cosmopolitan): for example, members of the SAR406 clade and a representative of MGI and SAR324 clade. For the rest of the 30 most abundant OTUs, some were consistently overrepresented or underrepresented in each deep-ocean basin indicating that at least a fraction of the community exhibited an uneven distribution across basins. Representatives of the *Acinetobacter* and *Pseudoalteromonas* genus and MGI Thaumarchaeota were overrepresented in the Canary basin and underrepresented in the Pacific

basin. Despite their proximity, the Brazil basin was characterized by a different combination of over-represented OTUs: the same representative of the *Acinetobacter* genus was overrepresented in this basin but in combination with two OTUs assigned to the *Marinobacter* and *Psychrobacter* genus. A different OTU also assigned to the *Psychrobacter* genus was underrepresented in this basin but was especially abundant in the Pacific basin, together with two OTUs assigned to the Corynebacteriales order. The South Australian basin was characterized by the overrepresentation of two OTUs assigned to the *Methylophaga* and *Microcoleus* genus while the Agulhas-Cape basin had above-average contributions of the two OTUs assigned to *Pseudoalteromonas* genus and MGI that were abundant in the Canary basin but without the co-presence of *Acinetobacter*. Although the differential contribution was computed correcting for the different number of samples in each basin (details in the Supplementary Information), the deviation from an even distribution was higher for the basins with a higher number of samples (Brazil, South Australian and Pacific basin). Future studies with a higher spatial detail and sampling size within each basin would allow to define these basins in terms of prokaryotic community composition and to describe, if they exist, indicator OTUs or clades for the distinct basins.

Beta-diversity patterns of bathypelagic prokaryotic communities

Non-metric multidimensional scaling was applied in order to represent the Bray–Curtis dissimilarities (that is, beta-diversity patterns) of the 60 samples (Figure 5) based on the relative abundance of all the OTUs. The samples belonging to different size fractions were clearly separated along the first axis. Detailed analysis of these differences is in the process (Salazar *et al.* in prep). Particle-attached samples within a deep-oceanic basin tended to have similar community composition, and thus clustered together in the non-metric multidimensional scaling. The seven samples located in the Pacific basin formed a tight cluster together with stations 81 (Fiji basin) and 88 (located also at the Pacific Ocean but at 2150 m depth and thus out of the basins defined below 3500 m). Samples belonging to the Brazil basin also clustered together and close to the samples from the Canary basin, both in the Atlantic Ocean. In contrast, the samples from stations 131 and 134 located also in the Atlantic Ocean and belonging to the same deep-water cluster but in a different basin, the North American basin, were more similar to the Pacific group. A third group of samples was composed by the stations situated in the Indian Ocean, in the South Australia basin, Madagascar basin and Mid Indian basin. This geographical ordering of the samples was not as evident for the free-living group of samples (Figure 5 and Figure 1 for sample location).

A deep-ocean study has recently emphasized the role of distinct deep-sea water masses as potential bio-oceanographical islands for prokaryotic communities (Agogué *et al.*, 2011). In addition, physical transport processes, such as advection, have been proved to act as ecological drivers of marine bacterial communities (Wilkins *et al.*, 2013) but the effect of the deep ocean's floor morphology over the composition of microbial communities was only explored in few locations such as the Walvis Ridge or the Challenger Deep (Schauer *et al.*, 2010; Nunoura *et al.*, 2015). Here we test, at a global scale, the possibility that submarine mountains that divide the deep ocean into deep-oceanic basins may act as 'ecological barriers' for prokaryotic communities: either by (i) imposing a reduced dispersion between basins or (ii) by compartmentalizing the bathypelagic ocean into contrasting environments that exert an ecological selection on the prokaryotes that inhabit it. In addition, we tested the relative role of water masses and deep-oceanic basins in shaping the biogeography of bathypelagic prokaryotes by examining the significance, tested using permutational multivariate analysis of variance, of the sample's clustering in accordance to the size-fraction, the deep-water cluster and deep-oceanic basins they belonged to (Supplementary Table S4). There were significant differences in the three factors (all $P < 0.0001$) and they together explained almost 70% of the variance (size fraction, $R^2 = 0.31$; deep-water cluster, $R^2 = 0.18$; deep-oceanic basins, $R^2 = 0.18$). None of the interactions between factors was significant. As most of the variance was explained by the size fraction, we split the analyses by size fraction to further test the deep-water cluster and basin as explanatory variables. Both factors were significant for the particle-attached fraction, and differences in oceanic basin origin explained 35% of the variance, even when taking the effect of deep-water clusters into account. For the free-living fraction there were no significant differences between oceanic basins once the deep-water clusters were considered. The date of sampling was included in the analyses to take into account seasonal differences as a possible confounding factor, as it has been shown that dark-ocean prokaryotic communities can be as dynamic as those of the surface ocean (Winter *et al.*, 2009) where the seasonal patterns are extremely relevant (Brown *et al.*, 2005; Fuhrman *et al.*, 2006; Gilbert *et al.*, 2009, 2012; Giovannoni, 2012). In all cases the date of sampling appeared to be a significant factor but its inclusion in the analyses did not modify the variance explained by the other factors (Supplementary Table S5). As the particulate matter in which particle-attached communities develop may ultimately come from the surface ocean through sinking, the Longhurst provinces were also tested as a potential factor structuring the beta-diversity of free-living and particle-attached communities. In none of the two cases the grouping of the stations in the corresponding Longhurst

provinces was significant (details not shown). Thus, in summary, particle-attached prokaryotic communities exhibited a significant basin-specific composition, whereas this basin specificity was not observed for free-living prokaryotes. Consequently, different processes need to be structuring the particle-attached and free-living beta-diversity and thus generating differential biogeographical patterns.

Ecological processes shaping the biogeography of deep-ocean prokaryotic communities

The bathypelagic free-living and particle-attached prokaryotic communities clustered according to the water masses while only particle-attached communities exhibited a significant basin specificity. However, the biogeographical patterns observed can arise as the result of different ecological processes well established within a theoretical framework (Hanson *et al.*, 2012): (a) the existence of environmental differences between basins or water masses that exert a differential selection of prokaryotes, that is, 'environmental selection' or (b) a reduced dispersal of microbes between basins or water masses, that is, 'historical effects'. We estimated the relative contribution of both processes by relating community composition to a set of environmental variables, and to the geographical distance between sampling locations, using multiple regression on matrices (see Supplementary Information for details and a further explanation of the theoretical framework). The multiple regression on matrices analysis explained a total of 23.1% (particle-attached fraction) and 10.7% (free-living fraction) of the total variance in the Bray–Curtis dissimilarity matrix. For the free-living fraction, only the effect of the environmental variables on community composition was statistically significant ($P = 0.005$; explaining 6.4% of the variance) and the effect of the geographical distance was not ($P > 0.1$), indicating that dispersal limitation and historical processes are not relevant in shaping the biogeography of free-living prokaryotes. However, dispersal limitation seemed to have a minor, yet significant, role for the particle-attached communities, as the pure effect of the geographical distance between locations explained a small but significant fraction of the variance ($P < 0.005$; explaining a 5% of the variance). In addition, most of the variance was explained by the effect of the environmental variables ($P < 0.005$), that is, the pure and the spatially structure environmental variation. These accounted for a 6.3% and 11.8% of the variance, respectively. Thus, both environmental selection and historical effects appear to shape the biogeography of particle-attached communities, although spurious distance effects may arise as a result of unmeasured environmental variables (Hanson *et al.*, 2012).

The best subset of environmental drivers for the free-living fraction was temperature and depth of the sampling location (Table 1), pointing to a pure

environmental selection process exerted by these two drivers for these communities. Temperature has also been found to be the main environmental driver for upper ocean microbial communities explaining the spatial variation within the epipelagic ocean (Sunagawa *et al.*, 2015). In contrast, the best subsets of environmental drivers for the particle-attached fraction were the apparent oxygen utilization (AOU) and the prokaryotic biomass duplication time, although most of the effect was solely due to the AOU (Table 1). The AOU (that is, the difference between the saturation and measured dissolved oxygen) indicates the modification of oxygen concentration through the mixing of water masses and various biogeochemical processes and correlates with the aging of a water mass (Jenkins, 1982). In addition, in this case the AOU is clearly reflecting the deep-water clusters each sample belongs to (Supplementary Figure S2b). Thus, the fact that the AOU of the samples where particle-attached communities are found is the best explanatory variable suggest that water mass mixing and aging have an important role in the assembly of particle-attached bathypelagic communities.

Finally, the scale of geographical variation for the two size fractions was studied using Mantel correlograms. We tested how far in space the samples maintain a significant autocorrelation in community composition. For particle-attached communities, there was a significant spatial autocorrelation, which expanded until 7500 km (Figure 6a). These relatively short distances, considering the global scale of the data set (that is, the ship covered ~45 700 km), are consistent with the basin-specificity of particle-attached prokaryotic community composition described above (Figure 5). In fact, the mean and maximal distance between all the samples belonging to the same deep-oceanic basin is 4950 and 9800 km, respectively. These distances are also coherent with the only study with a similar approach, which explored the effect of the Walvis Ridge on the

bacterial communities in the deep-sea sediments at the Guinea, Angola and Cape basins, reporting an effect of the geographical distance on community composition detectable at distances >3000 km (Schauer *et al.*, 2010). Although the effect of the geographical distance on community composition was not significant for the free-living communities once the effect of the environmental drivers is considered, there was a significant autocorrelation when tested at short distances, which expanded until 4500 km (Figure 6b). This significant autocorrelation found at short distances for the free-living communities does not correspond to the basin organization of the deep ocean, as tested before (Supplementary Tables S4 and S5), and may be due to the effect of potentially relevant environmental variables structuring the free-living bathypelagic communities at shorter scales, which were not measured in this study.

Thus, in summary, it seems that although both, the free-living and the particle-attached prokaryotic communities, exhibited autocorrelation at short distances and differ between water masses, they appeared to be structured by contrasting processes and drivers. The free-living prokaryotic communities appears to respond to an environmental selection process exerted by temperature and depth variations, although a high proportion of the variance remains unexplained (89.3%), as in similar studies (Hanson *et al.*, 2012). In contrast, the particle-attached communities appear to respond to a more complex set of processes where the ageing and global circulation of the water masses and some degree of dispersal limitation create basin-specific communities not evident for the free-living fraction. This could be an indication that at least a fraction of the deep-oceanic particles where the prokaryotes are associated, instead of coming from the surface ocean through sinking, correspond to presumably buoyant or slow-sinking particles that are produced autochthonously at depth, as it has recently been hypothesized (Herndl and Reinthaler, 2013).

Table 1 Environmental drivers of free-living and particle-attached communities

Environmental variable	Particle-attached		Free-living	
	Mantel <i>r</i>	P-value	Mantel <i>r</i>	P-value
Depth	0.145	0.11	0.261	0.06
Salinity	0.332	<0.01*	0.060	0.24
Potential temperature	0.156	0.01*	0.124	0.07
Apparent oxygen utilization	0.426	<0.01*	0.077	0.17
Prokaryotic heterotrophic activity	-0.156	0.93	-0.170	0.94
Prokaryote abundance	-0.156	0.90	-0.145	0.86
Percentage of HNA-content prokaryotes	-0.100	0.78	-0.077	0.67
Prokaryote biomass duplication time	0.209	0.01*	0.100	0.15
Prokaryote abundance (0–200 m)	-0.145	0.95	-0.059	0.67
Best BIOENV model	0.427	<0.01*	0.280	0.02*
	(AOU+DT)		(depth+potential temperature)	

Abbreviation: HNA, high nucleic acid. Mantel correlation (using Pearson correlation) between the Bray–Curtis dissimilarity and the Euclidean distance for particle-attached and free-living prokaryotes of the environmental variables used in the BIOENV approach (see Supplementary Information). Single variables and the best BIOENV model for each fraction are tested. Significant *P*-values (<0.05) are indicated with an asterisk.

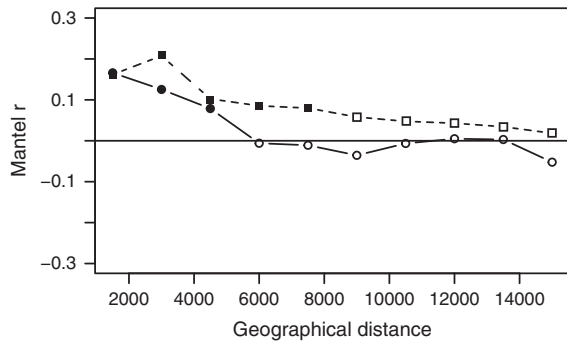


Figure 6 Mantel correlogram for particle-attached (squares) and free-living (circles) prokaryotic communities testing the autocorrelation on community composition by performing sequential Mantel tests between the Bray–Curtis dissimilarities and the grouping of samples using geographical distance classes set at 1500 m. Filled points represent significant correlations after Bonferroni correction. Mantel correlograms were run up to a maximal distance of 15 000 km.

This hypothesis would explain why particle-attached prokaryotes reflect the deep-water mass circulation and why a signal of reduced dispersal between basins is found only for particle-attached, and not for free-living communities.

Conflict of Interest

The authors declare no conflict of interest.

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