Environmental gradients and physical barriers drive the basin-wide spatial structuring of Mediterranean Sea and adjacent eastern Atlantic Ocean prokaryotic communities

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Abstract

The Mediterranean Sea is a miniature ocean divided by the Sicily Strait into two basins with a marked west to east trophic gradient and separated of the nearby eastern Atlantic Ocean by the Strait of Gibraltar. Here, we test the hypothesis that these physical and environmental barriers favor the development of specific prokaryotic assemblages, leading to changes in community structure both in the vertical and horizontal spatial scales. By analyzing taxonomic and phylogenetic diversity using amplicon sequence variants (ASVs) of the 16S rRNA gene, we show that there is indeed marked vertical segregation of prokaryotic groups, similar to that found in other areas of the ocean, but also a clear horizontal structuring among the two Mediterranean basins and the adjacent Atlantic waters. Prokaryotic diversity increased with depth and toward the Atlantic, whereas the easternmost stations displayed more phylogenetically diverse phylotypes, despite harboring globally less diverse communities. Basin-indicator taxa (ASVs) accounted for a large fraction of the community (between 50% and 80%) in each of the basins at the surface and bathypelagic layers, being associated with different environmental variables. The existence of biogeographic and environmental barriers in the Mediterranean Sea is likely related to the trophic gradient at the surface and the isolation of water bodies in depth due to the Gibraltar and Sicily straits. Our work highlights the importance of studying microbial regional biogeography and provides the basis for future studies on the impact of this regionalization in the function of Mediterranean Sea prokaryotic communities.

With a global abundance of ca. $1.2 \times 10^{29}$ cells (Whitman et al. 1998), marine prokaryotes (bacteria and archaea) drive most biogeochemical transformations in our planet (Falkowski et al. 2008). The advent of molecular methods, allowing to obtain large datasets at relatively low costs, has shown that marine microbes comprise a vast diversity and a myriad of metabolic capabilities. Marine prokaryotes are key mediators in a broad range of elemental cycles and thus, knowing their biogeography is critical to develop a predictive understanding of microbial processes in the ocean and how they will evolve in future global change scenarios. Despite the profusion of microbial biogeographic studies (Herlemann et al. 2011; Sunagawa et al. 2015; Milici et al. 2016) there are still ocean areas that are far from being well-known. For instance, studies looking at prokaryotic composition and its variability in the dark ocean are limited (but see Agogué et al. 2011; Salazar et al. 2015), even though dark ocean waters (> 200-m depth) host 70% of ocean’s prokaryotic cells (Arístegui et al. 2009).

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Microbial biogeographic patterns result from speciation, extinction, and dispersal processes that lead to a combination of cosmopolitan and endemic species in different regions (Hanson et al. 2012). In ocean ecosystems, dispersal limitation is often determined by the existence of physical barriers such as straits or islands, different water masses, currents, or fronts, which highly determine prokaryotic community structure (Baltar and Arístegui 2017; Morales et al. 2018). On the other hand, environmental factors such as temperature, inorganic and organic nutrients, or sunlight highly constrain prokaryotic community structure by imposing niche differentiation (Coleman and Chisolm 2010; Gilbert et al. 2011; Sunagawa et al. 2015).

The Mediterranean Sea, with a total area of $2.5 \times 10^6 \text{ km}^2$, is the largest semi-enclosed basin on Earth. It is frequently treated as a miniature ocean because it has its own overturning circulation, and relevant oceanographic processes, such as dense water formation, occur in this basin (Bethoux et al. 1999). Globally oligotrophic, it presents a west to east gradient of increasing oligotrophy (Krom et al. 1991), with growth and activity of microbial communities inhabiting the euphotic Mediterranean Sea being generally limited by phosphate availability (Thingstad et al. 1998; Pinhassi et al. 2006). Exclusive from the Mediterranean Sea are also the high salinity (38–39 relative to ca. 35 in the open ocean) and the relatively high temperatures in the meso- and bathypelagic layers (about 10°C warmer than deep waters of the global ocean) that appear to lead to higher remineralization rates compared to other deep ocean basins (Christensen et al. 1989; Santinelli et al. 2010). One of the main physical features of this basin is that it is connected to the adjacent Atlantic Ocean only by the shallow and narrow Strait of Gibraltar (300 m deep, 13 km wide). This physical barrier reduces water mixing and promotes an inflow of surface nutrient-poor waters from the Atlantic to the Mediterranean Sea that outweighs the outflow of nutrient-rich deep waters in the opposite direction (Sammartino et al. 2015). In addition, the strait of Sicily (500 m deep) divides the Mediterranean Sea into two basins, the Western and Eastern Mediterranean, physically isolating the deep water bodies from both basins. The existence of these barriers led us to hypothesize that prokaryotic community structure in the Mediterranean Sea must be impacted by a combination of these physical barriers (i.e., prokaryotic communities will be more disconnected in the meso- and bathypelagic basins than in the euphotic layers) together with other environmental parameters, namely the west-to-east and vertical gradients in substrate availability.

Most published studies looking at biogeographic patterns of prokaryotic communities using state-of-the-art high-throughput sequencing of the 16S rRNA gene in the Mediterranean Sea have focused in a few localized areas (reviewed in Pulido-Villena et al. 2012; Luna 2014): the coastal Tyrrenian (Thiele et al. 2017), the NW Mediterranean Sea (Laghdass et al. 2010; Crespo et al. 2013; Severin et al. 2016; Mestre et al. 2017), the Ligurian Sea (Celussi et al. 2018), the Adriatic (Korlević et al. 2015), and, to a lesser extent, the Eastern Mediterranean Sea (De Corte et al. 2009; Yokokawa et al. 2010; Techtmann et al. 2015). To date, the broad-scale patterns of prokaryotic diversity and community composition in this enclosed sea have only been studied once using automated ribosomal intergenic spacer analysis (ARISA) fingerprinting (Mapelli et al. 2013, which found a longitudinal and vertical structuring of prokaryotic communities, but lacked information on the taxonomic composition of these communities.

To explore the role of the geographic barriers and the oceanographic gradients in structuring prokaryotic communities in the Mediterranean Sea, we present a quasi-synoptic view of the basin-wide distributions of archaeal and bacterial communities using 16S rRNA gene illumina sequencing. We collected samples on a transect that covered the most representative regions and water masses of the Mediterranean Sea in the opposite direction to the surface Atlantic water flow, from east to west, and from surface to near-bottom waters, including the adjacent Eastern North Atlantic Ocean. Complementing these data with a large dataset of ancillary environmental parameters, we investigated the main physical and environmental factors driving prokaryotic diversity as well as the determinants of spatial differentiation. The main hypotheses addressed were: (1) the west–east gradient in oligotrophy is reflected by a west to east decreasing trend in prokaryotic diversity; and (2) the combination of physical barriers (i.e. the Gibraltar and Sicily straits) and the sharp changes in environmental parameters segregate prokaryotic communities by depth and basin.

**Methods**

Seawater sampling was carried out during the HOTMIX 2014 cruise on board the R/V Sarmiento de Gamboa, from 29 April to 28 May 2014. Seawater was collected at 29 stations along a Mediterranean Sea section from east to west, also extending to the adjacent subtropical northeast (NE) Atlantic Ocean reaching the Canary Islands (Fig. 1). Detailed information about the distributions of salinity, potential temperature and chlorophyll $a$ (Chl $a$) concentrations as well as the water masses intercepted during the cruise is provided in Catalá et al. (2018). Samples were collected using a rosette sampler holding 24 Niskin bottles (12 liters each), coupled to a Seabird SBE 9-11 plus conductivity-temperature-pressure probe, complemented with a SBE43 oxygen sensor and a SeaTech fluorometer. Up to 13 depths were sampled covering the entire water column, from 3 m down to 10 m above the seafloor.

Samples from the epipelagic layer (surface water; down to 200 m) were systematically collected at four depths: 3 m, the depth receiving 20% of the surface photosynthetically active radiation, the depth of the deep chlorophyll maximum (DCM), and between 10 and 45 m below the DCM. The depth of the DCM was determined after visual inspection of the
vertical profiles of Chl a fluorescence. Sampling depths in the meso- and bathypelagic layers were chosen according to the full-depth potential temperature, salinity, and dissolved oxygen profiles, to ensure that all the water masses of the Mediterranean Sea and their respective mixing zones were sampled (for more details, see Catalá et al. 2018). In this study, we discriminated mesopelagic from bathypelagic samples using the global criterion of 1000 m depth (sample from 200 to 1000 m were considered mesopelagic and samples deeper than 1000 m were considered as bathypelagic).

Environmental variables
Dissolved oxygen, inorganic nutrients (nitrogen, phosphorus, and silicate), Chl a, total and particulate organic carbon, particulate organic nitrogen, colored and fluorescent dissolved organic matter (DOM), transparent exopolymer particles, heterotrophic prokaryotic abundance (PA), and the percentage of high nucleic acid cells were determined in discrete samples using standard protocols as described elsewhere (Catalá et al. 2018; Martínez-Pérez et al., 2019; Ortega-Retuerta et al. 2019). For a brief description on the methodologies, see Supporting Information Methods.

Nucleic acids extraction
Samples for nucleic acids extraction (4–12 liters) were sequentially filtered through 47-mm 3-μm polycarbonate membrane filters (Merk Millipore, Isopore polycarbonate) and 0.2-μm pore size Sterivex units with a peristaltic pump. Total nucleic acids from the 0.2–3-μm size fraction were extracted using the PowerWater Sterivex™ DNA isolation Kit (MO BIO Laboratories) following the manufacturer instructions. DNA was quantified using a Qubit fluorometer assay (Life

Fig 1. (A) Study area and stations sampled during the HOTMIX2014 cruise, different symbols correspond to the different basins (dots: Atlantic; triangles: Western Med. basin; asterisks: Eastern Med. basin). (B) Vertical profile showing the Gibraltar and Sicily straits that act as physical barriers in the dark Mediterranean Sea. Dots represent the samples collected for prokaryotic diversity. The color contour plot shows the vertical distribution of phosphate concentration along the transect.
Technologies). The V4-V5 region of the 16S gene was amplified with the primers 51SF and 926R (Parada et al. 2016) and sequenced in an Illumina MiSeq platform using 2 × 250 bp paired-end approach at the RTLGenomics facility (https://rلالロンジモニゴス.com/). The collected samples for nucleic acids are shown in Fig. 1B.

Data analyses

Primers and spurious sequences were trimmed using cutadapt (Martin 2011). DADA2 v1.8 was used to differentiate exact sequence variants (Callahan et al. 2016). DADA2 resolves amplicon sequence variants (ASVs) by modeling the errors in Illumina-sequenced amplicon reads. Taxonomic assignment was performed using the function “assignTaxonomy” against SILVA v.132 through the RDP naive Bayesian classifier method described in (Wang et al. 2007). ASVs assigned to chloroplasts or eukaryotes were removed for subsequent analyses. All raw sequences used in this study are publicly available at the European Nucleotide Archive (PRJEB44474). The ASV table was randomly subsampled down to the minimum number of reads per sample using the “rarefy” function in the Vegan 2.5-5 package (Oksanen et al. 2015) for the nmds ordination, the diversity index estimates, and the indicator ASV analyses (see below).

Data treatment and statistical analyses were performed with the R (version 3.6.0) and RStudio software (version 1.2.1335). The phylogenetic diversity (PD) was calculated considering the evolutionary relationships among ASVs using the computed phylogeny for each sample, as the sum of the lengths of all the branches in the phylogeny. For this, we used the “DECHIPER” 2.14.0 and “phangorn” 2.5.5 packages (Schliep 2011; Wright 2016). Prokaryotic richness (the number of ASV per sample) and sample evenness (using the Pielou index: \( J = H/\ln(n_{ASV}) \), where \( H \) is the Shannon index and \( n_{ASV} \) is the richness in every sample), were calculated using the “Vegan” 2.5-5 package.

In order to assess to which extent environmental conditions and the physical barriers affect the assembly of the prokaryotic communities, we investigated if there were indicator ASVs from each different basin: North East Atlantic, Western Mediterranean, and Eastern Mediterranean Sea. Indicator taxa were identified using the IndVal index from the “labdsv” 2.0-1 package (Roberts 2019), which takes into account the fidelity and relative abundance of the taxa in the different basins. \( p \)-Values were adjusted using the false discovery rate approach (Benjamini and Hochberg 1995). Given the vertical stratification of the prokaryotic communities, we calculated the basin-indicator taxa separately for each of the four depth layers studied: surface, DCM, mesopelagic, and bathypelagic.

To test for significant differences in richness, PD, the ratio PD/richness, and evenness, we used one-way ANOVA and post hoc Tukey tests using depth layer and basin as factors. The same statistical tests were applied using the proportion of sequences of each ASV per layer and basin.

Diversity metrics were correlated against the main environmental variables using pairwise Pearson correlations on log10-transformed data, with posterior Bonferroni corrections. The figures were constructed using “ggplot2” 3.2.1 in R.

The association of the basin-indicator ASVs with biotic and environmental variables was explored using a sparse partial least squares (sPLS) regression analysis using the “mixOmics” package (Rohart et al. 2017) in R in “regression” mode. We included the following variables: temperature, salinity, apparent oxygen utilization, nitrate and phosphate concentration, fluorescent dissolved organic matter (FDOM) peaks C (humic-like), and T (protein-like), the spectral slope of chromophoric dissolved organic matter (CDOM) between 275 and 295 nm, total organic carbon (TOC), particulate organic carbon, prokaryotic cell abundance, % high nucleic acid prokaryotes, transparent exopolymer particles concentration, and photic layer integrated (upper 200 m) nitrate and phosphate concentration values. Leave-one-out cross-validation was performed to evaluate model performance (Lê Cao et al. 2008). sPLS allowed the simultaneous selection of meaningful variables in both the basin-indicator ASVs and the environmental dataset, and the resulting pair-wise associations were visualized using clustered image maps (González et al. 2012).

Results

The environmental and biotic parameters monitored along our transect exposed the existence of marked gradients in different regions and depths. While the overall sea surface temperature ranged from 16.3°C to 19.3°C and all stations were thermally stratified (Fig. S1A), Mediterranean deep waters were increasingly warmer toward the Eastern basin, up to 9°C warmer than the Atlantic ones (Fig. S1A). This marked spatial gradient was also observed for salinity, which increased from west to east at all depths of the water column (Fig. S1B), and for inorganic nutrients, where a clear west to east increase in oligotrophy (i.e., decrease in inorganic nutrient concentrations) was evidenced (Figs. 1, S1C). TOC decreased from surface to deep waters and, while TOC in deep waters decreased from west to east, TOC accumulation in the surface occurred in the Eastern basin (Martínez-Pérez et al. 2019; Fig. S1D). PA ranged from 0.21 to 10.2 × 10^5 cells mL^-1 and decreased with depth, from an average of 5.12 × 10^5 cells mL^-1 in epipelagic waters to an average of 0.42 × 10^5 cells mL^-1 in the bathypelagic. PA was lower in bathypelagic waters of the Atlantic and Eastern Mediterranean basins than in bathypelagic Western Mediterranean waters (Fig. S1E).

Besides the clear differences in environmental conditions among the three main basins (i.e., East and West Mediterranean and Atlantic Ocean; Fig. S1), there were sharp differences in some of the subbasins, particularly the Tyrrenian Sea, which had higher overall temperature, lower salinity, lower nutrients, and higher mesopelagic PA abundances than the westernmost stations in the Mediterranean Sea. Given that
Prokaryotic diversity was only sampled in two stations of this subbasin (Sta. #15 and #16), we did not formally compare the different subbasins of the Western Mediterranean.

Prokaryotic community structure and diversity
The taxonomic similarity between samples was assessed by placing them in a two-dimensional space based on nonmetric multidimensional scaling (Fig. 2). Prokaryotic communities were significantly structured by depth layer (PERMANOVA test, $R^2 = 0.43$, $p < 0.001$; Fig. 2; Table S1), and then clustered according to the different basins (PERMANOVA test, $R^2 = 0.07$, $p < 0.001$; Table S1). Differences between basins were also significant when treating the different water layers as separate datasets (PERMANOVA test, $p < 0.001$, $R^2 = 0.39$ for surface, $R^2 = 0.14$ for DCM, $R^2 = 0.30$ for mesopelagic, and $R^2 = 0.41$ for bathypelagic; Table S1).

Prokaryotic richness was on average 396 ASVs per sample, and increased with depth, showing significantly higher values in the meso- and bathypelagic layers than in the euphotic layers (surface and DCM, ANOVA + Tukey HSD post hoc pairwise test, $p < 0.001$; Fig. 3). On the horizontal scale, a significant west to east decrease in richness was also observed (Fig. 3; Table 1). These spatial patterns in richness were also followed by PD (Fig. 3), which ranged widely from 12.2 (Eastern Mediterranean, surface) to 102.2 (Western Mediterranean, mesopelagic; Table 1). In fact, richness and PD were positively correlated (Pearson correlation, $n = 186$, $R = 0.94$, $p < 0.001$). Patches of relatively high richness and PD were observed in NE Atlantic waters around 1300 m, in bottom Western Mediterranean waters next to the Strait of Gibraltar, and in the Ionian sea (Sta. 11 and 12) in the Eastern Mediterranean (Fig. S2). By contrast, PD normalized by richness (PD/rich, an indication of how phylogenetically diverse are the communities normalized by overall richness) was higher in the Eastern Mediterranean (Fig. 3), particularly at the easternmost stations of the Levantine basin (Fig. S2). Sample evenness showed no clear longitudinal or vertical trends, but displayed overall higher values in the mesopelagic (Fig. 3).

The different diversity indices correlated significantly with several environmental variables, particularly in the bathypelagic. In this layer, both richness and PD presented significant positive correlation coefficients with inorganic nutrients, salinity, and the humic-like FDOM peak C (Table S2). In contrast, the PD/richness ratio was significantly negatively correlated with PA and TOC. In the other depth layers, diversity indexes were less influenced by environmental variables, although we found that humic-like FDOM peak C was negatively correlated with the PD/richness ratio in the mesopelagic layer, and positively correlated with evenness at the DCM (Table S2).

Looking at the fraction of shared taxa throughout the water column (considering all ASVs in every basin), we found that overall, around half of the ASVs were exclusively detected in the dark ocean (meso- and bathypelagic layers; Fig. 4), and only 1% of the ASVs was shared among all layers. Microbial communities from the Atlantic displayed a large degree of isolation in the bathypelagic, with 45.2% of the basin ASVs

![Fig 2. Nonmetric multidimensional scaling (NMDS) ordination based on Bray Curtis similarities of the prokaryotic communities. Color codes separate depth layers. B, Bathypelagic; M, Mesopelagic; DCM, deep chlorophyll maxima; S, surface; symbols denote basins.](image-url)
exclusively found at this layer. This isolation was less evident for the mesopelagic, where unique ASVs represented 17% of the ASVs. In contrast, the proportion of ASVs exclusively found in meso- and bathypelagic layers were similar in the Mediterranean basins (~25% of the ASVs) (Fig. 4). Meso- and bathypelagic layers shared on average 14% of the ASVs, whereas the proportion of ASVs shared between the photic layers (surface and DCM) was on average 4%. In terms of sequences, these unique ASVs were quite rare, as they represented overall a very small proportion of the community (<7%) (Fig. S3). A clear exception was the Atlantic bathypelagic unique ASVs, which represented on average ~40% of bathypelagic communities. This high contribution of unique ASVs, however, was partly due to the deeper waters (>3000 m) sampled in this basin, where unique ASVs represented up to 80% (data not shown). Considering similar depths to the Mediterranean basins, the contribution of unique ASVs to the Atlantic bathypelagic communities was still significant, around 30%, but more similar to values found in the deeper bathypelagic waters of the Eastern Mediterranean (~20%, data not shown).

**Spatial changes in the prokaryotic community composition**

The bulk taxonomic composition of surface, mesopelagic, and bathypelagic communities was relatively similar throughout the
Table 1. Average (± standard deviation) richness (number of ASVs), phylogenetic diversity, phylogenetic diversity normalized by richness (PD/richness), and evenness (Pielou index) in the different basins and depth layers. *n* = number of samples. Letters denote significant differences (*p* < 0.05) between basins after one-way ANOVA and post hoc Tukey tests.

|                | Surface | DCM    | Mesopelagic | Bathypelagic | All layers |
|----------------|---------|--------|-------------|--------------|------------|
| **Richness**   |         |        |             |              |            |
| Atlantic       | 313±116a| 380±130a| 602±86a     | 650±188a     | 524±203a   |
| Western Med    | 246±74a | 312±124a| 489±186ab   | 480±146b     | 401±177b   |
| Eastern Med    | 243±59a | 280±65a | 405±192b    | 407±151b     | 345±156b   |
| Global         | 256±78  | 308±102| 467±189     | 483±181      |            |
| **Phylogenetic diversity** |         |        |             |              |            |
| Atlantic       | 31.5±13.3a| 36.4±10.2a| 55.9±5.7a  | 70.9±14.8a | 53.6±20.5a |
| Western Med    | 25.5±7.4a| 34.7±12.6a| 45.6±17.9a | 55.7±13.9b| 42.0±18.0b |
| Eastern Med    | 24.5±5.5a| 32.1±5.1a | 45.6±16.7a | 50.3±16.5b| 38.5±16.6b |
| **PD/richness**|         |        |             |              |            |
| Atlantic       | 0.105±0.013a| 0.112±0.012a| 0.094±0.009a | 0.118±0.012a | 0.103±0.014a |
| Western Med    | 0.101±0.013a| 0.098±0.009a| 0.094±0.006a | 0.118±0.015a | 0.107±0.019a |
| Eastern Med    | 0.102±0.012a| 0.118±0.016a| 0.109±0.019b | 0.127±0.018b | 0.114±0.015b |
| **Evenness**   |         |        |             |              |            |
| Atlantic       | 0.70±0.02a | 0.76±0.05a | 0.83±0.06ab | 0.81±0.04a | 0.78±0.06a |
| Western Med    | 0.79±0.04b | 0.82±0.02a | 0.82±0.02a  | 0.78±0.03b | 0.80±0.04ab |
| Eastern Med    | 0.80±0.03b | 0.78±0.06a | 0.85±0.02b  | 0.81±0.04a | 0.81±0.04b |
| **n**          |         |        |             |              |            |
| Atlantic       | 8       | 4      | 8           | 13          | 33         |
| Western Med    | 16      | 9      | 20          | 19          | 64         |
| Eastern Med    | 24      | 12     | 25          | 28          | 89         |
transect, except for surface Atlantic communities (Fig. 5). However, communities were remarkably heterogeneous among stations at the DCM layer. SAR11 clade bacteria and cyanobacteria dominated both the surface and the DCM communities (Fig. 5), although sharp transitions in the dominance of cyanobacterial genera were observed between the different basins. Whereas Synechococcus dominated surface cyanobacterial communities of both Mediterranean basins and the DCM of the Western Mediterranean, Prochlorococcus dominated both surface and DCM in the Atlantic and in the DCM of the Eastern basin (Figs. 5, S4). The mesopelagic and bathypelagic layers were dominated by archaeal ASVs, which comprised roughly half of the 16S sequences (52% and 44% of the communities for the meso- and bathypelagic, respectively; Fig. 5). From these, the majority of sequences corresponded to Thaumarchaeota. The most abundant bacterial groups in the bathypelagic were alpha- and gammaproteobacteria (representing ~10% each), followed by members of the phylum Chloroflexi and the SAR406 clade (Marinimicrobia), which represented around 8% of the reads. Specific physical features of the Mediterranean Sea appeared to have a notable impact on the prokaryotic community structure: in Sta. #13–14, located in the Sicily Strait, and #24, at the Eastern side of the Gibraltar Strait, a higher relative abundance of archaea, mostly Thaumarchaeota, could be detected in surface and DCM waters (Fig. 5), reflecting the upward displacement of waters due to the shallow straits.

ANOVA tests using depth layer and basin as factors confirmed the high spatial structuring of many of the taxonomical groups found (Table 2, Table S3). In fact, most of the groups showed significantly different distributions among depth layers (Table 2). For instance, Archaea were predominantly located in the mesopelagic layer, but some archaeal groups were more dominant in the bathypelagic, like the Euryarchaeota Marine Group III or the Woesearchaeota. In contrast, members of the Marine Group I of Thaumarchaeota, like Candidatus nitrosopumilus, were enriched at the DCM.

Some bacterial phyla were predominantly found in the euphotic layers (Table 2), like Actinobacteria and Planctomycetes (enriched in DCM waters), Bacteroidetes (both surface and DCM), and Verrucomicrobia, with the exception of the Artic97B-4 marine group. Alphaproteobacteria overall dominated the prokaryotic communities of the surface layer, but there were contrasting patterns within this class. The orders Rhodobacterales, Rickettsiales, and SAR11 bacteria showed a preference for the surface (Table 2), except the deep SAR11 clade that was predominantly found in the mesopelagic, whereas Caulobacterales, Rhodospirillales, and
Sphingomonadales were more frequently found in the bathypelagic. Within the Gammaproteobacteria, the orders Cellvibrionales, the KI89A clade, and the SAR86 were mostly found in surface waters, whereas Alteromonadales, E01-9C-26 marine group, Pseudomonadales, and Xanthomonadales increased their relative abundance in bathypelagic waters. Other phyla were predominantly observed in the deep-water layers, such as Marinimicrobia, Gemmatimonadetes, Lentisphaerae, Nitrospina (higher at the mesopelagic), and Nitrospira (higher at the bathypelagic).

Besides the clear vertical structuring, we also detected horizontal structuring (i.e., differences among basins) for some groups. For instance, SAR324 bacteria and Lentisphaerae displayed higher relative abundances in the Eastern North Atlantic (Tables 2, S3), whereas the relative abundance of Acidobacteria, Chloroflexi, and phylum SBR1093 and the group E01-9c-26 from Gammaproteobacteria increased significantly in the Eastern Mediterranean (Table 2; Fig. S5). Nitrospina showed comparatively higher abundances in the Atlantic and Western Mediterranean basins than in the Eastern Mediterranean basin, while Nitrospira displayed the opposite pattern (Table 2; Fig. S5). Overall, many of the groups that presented changes in abundance among basins were significantly enriched in the dark ocean (Table 2), although there were also some clear segregations in the surface, as for example Surface 2 group of the SAR11 clade, which was over-represented in the Eastern Mediterranean basin (Table 2; Fig. S6).

Biogeographical barriers in the Mediterranean define different prokaryotic communities in surface and bathypelagic waters

In order to further explore to which extent each basin contained specific prokaryotic taxa, we performed an indicator taxa (at the ASV level) analysis (see the Methods section) for each of the depth layers separately. The number of indicator ASVs and their contribution to the prokaryotic communities was notably higher in the surface and bathypelagic waters of the Eastern North Atlantic basin, decreasing sharply at the Mediterranean Sea waters interface. The transition between Mediterranean Sea basins was less abrupt, with Western ASVs gradually being replaced by Eastern Mediterranean ASVs (Fig. 6). Atlantic indicator taxa accounted for ~60% and up to 80% of the communities in surface and bathypelagic waters of the Eastern North Atlantic basin, respectively, decreasing sharply at the Mediterranean Sea waters interface. The transition between Mediterranean Sea basins was less abrupt, with Western ASVs gradually being replaced by Eastern Mediterranean ASVs (Fig. 6). Yet, the contribution of Western Mediterranean and Eastern Mediterranean indicator ASVs to surface and bathypelagic communities of the Western and Eastern basins, respectively, was still high, representing up to 50% of the communities. In contrast to surface and bathypelagic layers, there were few indicator ASVs in the
Table 2. Vertical and basin segregation of prokaryotic groups in the Mediterranean Sea. Relative abundance (in %) and results of ANOVA + post hoc Tukey tests looking for significant differences among depth layer or basins. Only those taxa with a different spatial structuring than the group they belong to are listed. Color code denotes the results of the post hoc Tukey tests (see legend below) with darker colors representing significantly higher relative abundances. Light blue shading indicates those groups that displayed abundances higher than 1%.

| Layer effect | Average % | ANOVA P | Basin effect | ANOVA P | ATLAS | Warm | Cold |
|--------------|-----------|---------|--------------|---------|-------|------|------|
| **Archaea**  |           |         |              |         |       |      |      |
| Euryarchaeota| 4.21      | <0.0001 | 0.016        |         |       |      |      |
| Thermoplasmata| 4.19     | <0.0001 | 0.015        |         |       |      |      |
| Marine_Group_II| 3.16    | <0.0001 | 0.002        |         |       |      |      |
| Marine_Group_III| 1.03   | <0.0001 | ns           |         |       |      |      |
| Thaumarchaeota| 26.45     | <0.0001 | *            |         |       |      |      |
| Marine_Benthic_Group_A| 0.47   | <0.0001 | *            |         |       |      |      |
| Marine_Group_I| 25.91    | <0.0001 | *            |         |       |      |      |
| Candidatus_Nitrosopelagicus| 2.86   | <0.0001 | *            |         |       |      |      |
| Candidatus_Nitrosopumilus| 0.76   | <0.0001 | *            |         |       |      |      |
| Woesearchaeota_(DHVEG-6)| 0.26  | <0.0001 | *            |         |       |      |      |
| **WSA-2**   | 0.01      | <0.0001 | *            |         |       |      |      |
| **Bacteria**|           |         |              |         |       |      |      |
| Acidobacteria| 1.10      | <0.0001 | 0.036        |         |       |      |      |
| Actinobacteria| 1.59     | <0.0001 | ns           |         |       |      |      |
| Bacteroidetes| 4.13      | <0.0001 | ns           |         |       |      |      |
| Flavobacteria| 3.45      | <0.0001 | ns           |         |       |      |      |
| Flavobacteriales| 3.45    | <0.0001 | ns           |         |       |      |      |
| Chloroflexi| 730.00     | 0.0001  |              |         |       |      |      |
| SAR202_clade| 3.49      | <0.0001 | 0.005        |         |       |      |      |
| Cyanobacteria| 9.68      | <0.0001 | 0.038        |         |       |      |      |
| Pochlorococcus| 5.64     | <0.0001 |              |         |       |      |      |
| Synechococcus| 4.00      | <0.0001 | 0.019        |         |       |      |      |
| Gemmatimonadetes| 0.39    | <0.0001 | ns           |         |       |      |      |
| Lentisphaeraceae| 0.07    | <0.0001 | 0.0000       |         |       |      |      |
| Marinimicrobia_(SAR406_clade)| 4.90   | <0.0001 | ns           |         |       |      |      |
| Nitrospinae| 1.27      | <0.0001 | *            |         |       |      |      |
| Nitrospina| 0.76      | <0.0001 |              |         |       |      |      |
| Nitrospira| 0.45      | <0.0001 |              |         |       |      |      |
| PAUC4f| 0.27      | <0.0001 | *            |         |       |      |      |
| Planctomycetes| 1.91      | <0.0001 | *            |         |       |      |      |
| Proteobacteria| 36.01     | <0.0001 | *            |         |       |      |      |
| AEGEAN-245| 0.07      | <0.0001 | *            |         |       |      |      |
| Alphaproteobacteria| 24.12     | <0.0001 | *            |         |       |      |      |
| Caulobacteriales| 0.06     | <0.0001 | 0.015        |         |       |      |      |
| Rhizobiales| 0.60      | <0.0001 | ns           |         |       |      |      |
| Rhodobacteriales| 1.90     | <0.0001 | ns           |         |       |      |      |
| Rhodospirillales| 1.66     | <0.0001 | 0.005        |         |       |      |      |
| Rickettsiales| 2.84      | <0.0001 | *            |         |       |      |      |
| SAR16_clade| 2.26      | <0.0001 | *            |         |       |      |      |
| SAR11_clade| 15.62     | <0.0001 | *            |         |       |      |      |
| Deep_1| 2.40      | <0.0001 | *            |         |       |      |      |
| Surface_1| 11.20     | <0.0001 | *            |         |       |      |      |
| Surface_2| 1.23      | <0.0001 | 0.049        |         |       |      |      |
| Surface_3| 0.46      | <0.0001 | ns           |         |       |      |      |
| Sphingomonadales| 0.43     | <0.0001 | ns           |         |       |      |      |
| Betaproteobacteria| 0.27     | 0.0044  |              |         |       |      |      |
| Deltaproteobacteria| 3.90     | <0.0001 | 0.071        |         |       |      |      |
| SAR324_clade(Marine_group_B)| 3.46   | <0.0001 | 0.003        |         |       |      |      |
| Gammaproteobacteria| 9.55     | 0.0054  |              | <0.0001 |       |      |      |
| Alteromonadales| 0.29      | <0.0001 | ns           |         |       |      |      |
| Cellibacteriales| 0.52     | <0.0001 | *            |         |       |      |      |
| E09-9C-26_marine_group| 2.35     | <0.0001 | 0.035        |         |       |      |      |
| K19A_clade| 0.31      | <0.0001 | ns           |         |       |      |      |
| Oceanospirillales| 3.40     | <0.0001 | *            |         |       |      |      |
| SAR66_clade| 2.34      | <0.0001 | 0.077        |         |       |      |      |
| Pseudomonadales| 0.93     | 0.017   | 0.015        |         |       |      |      |
| Salinisphaeraceae| 0.87     | <0.0001 | 0.021        |         |       |      |      |
| Thiorthiales| 0.33      | <0.0001 | ns           |         |       |      |      |
| Vibrionales| 0.01      | 0.0015  | 0.037        |         |       |      |      |
| Xanthomonadales| 0.29      | <0.0001 | ns           |         |       |      |      |
| SBR1093| 0.36      | <0.0001 |              |         |       |      |      |
| Spirichaeae| 0.03      | <0.0001 | ns           |         |       |      |      |
| Verrucomicrobiaceae| 1.32     | <0.0001 | ns           |         |       |      |      |
| Arctic97B-4_marine_group| 0.35     | <0.0001 | 0.040        |         |       |      |      |

A: S, DCM, M, B. B: S, DCM, M, B. C: S, DCM, M, B. D: S, DCM, M, B.
DCM and mesopelagic layers (Fig. 6) that generally accounted for < 10% of the communities. An exception was the mesopelagic Western Mediterranean indicators, which comprised around 25% of mesopelagic communities in this basin.

sPLS regression analysis unveiled strong associations between abundant (representing 1% of the sequences in at least one sample) surface and bathypelagic basin indicator ASVs and some environmental variables, particularly in the Western and Eastern Mediterranean (Figs. S7, S8). Surface Western Mediterranean indicators were associated with comparatively higher values of photic integrated inorganic nutrients, FDOM (Peaks T and C), the spectral slope of CDOM between 275 and 295 nm (i.e., lower molecular weight of CDOM compounds; Helms et al. 2008), dissolved oxygen, particulate carbon, and PA, and negatively associated to temperature and the proportion of labile to recalcitrant compounds (FDOM peak T to peak C ratio). Conversely, Eastern Mediterranean indicators displayed the opposite associations. Surface Atlantic indicator ASVs were associated with relatively warmer waters. These Atlantic indicator ASVs were mostly represented by Prochlorococcus, which decreased sharply in relative abundance in the Western basin (Figs. S7, S9). In contrast, Western indicators included Synechococcus, Flavobacteriales and SAR11 surface clade 1 ASVs. Eastern indicators comprised other Synechococcus and SAR11 surface clade 1 ASVs, as well as several SAR116 ASVs and members of the SAR86 clade (Figs. S7, S9).

Fig 6. Contribution of basin indicator ASVs to the prokaryotic communities in each of the depth layers. Basins are separated by a vertical dashed line. Left panel: Atlantic indicators. Mid-panel: Western Mediterranean indicators. Right panel: Eastern Mediterranean indicators. The n value within each plot represents the number of indicator ASVs per layer and basin.

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In the bathypelagic layer, the basin indicators were also associated with different environmental variables (Fig. S8). Bathypelagic Atlantic indicators were linked to cold temperatures, and increasing values of inorganic nutrient concentrations, humic-like FDOM (peak C), the spectral slope of CDOM between 275 and 295 nm, high apparent oxygen utilization, and the proportion of high nucleic acid bacteria, whereas Eastern indicators displayed the opposite trend (Fig. S8). Indicators from the Western basin showed in general weaker associations with environmental variables but were linked to overall comparatively higher PAs. Thaumarchaeota represented a large fraction of bathypelagic indicators in the three basins, but in the Western Mediterranean they accounted for most of the indicator sequences (Fig. S9).

Discussion

Our study represents the first quasi-synoptic overview of prokaryotic diversity and community composition along an east–west transect in the Mediterranean Sea. This miniature ocean is characterized by a marked horizontal gradient in surface nutrient availability, decreasing from west to east (Figs. 1B, S1C). Moreover, the Gibraltar and Sicily straits act as physical obstacles that reduce circulation and water mass mixing across basins in the bathypelagic. Taken together, these characteristics lead to a high spatial structuring of prokaryotic communities. This structuring is reflected by the differences in diversity, community structure, and composition in both the vertical and horizontal dimensions.

Prokaryotic communities are highly structured along the vertical axis

The marked vertical structuring observed agrees with previous findings, showing that the Marine Group I Thaumarchaeota are more prevalent in the mesopelagic layer (Santoro et al. 2019), and Marine Group III in the bathypelagic layer (Martin-Cuadrado et al. 2008; Li et al. 2015). Marine Group II archaea were significantly enriched in the mesopelagic in our study, although they have been reported to be most abundant in sunlit waters in other regions of the ocean, as the North Pacific (Massana et al. 1997; DeLong et al. 2006). Different environmental conditions in the Mediterranean could lead to the presence of different Marine Group II ecotypes (Pereira et al. 2019) with a preference for mesopelagic waters. Woesearchaeota are found in low abundance in a wide variety of environments (Liu et al. 2018) but in our study they were significantly enriched in the bathypelagic. The abundance of WSA2 Archaea, which have been reported to possess energy conservation mechanisms (Nobu et al. 2016), also increased in the bathypelagic. Besides the observed changes in the vertical distribution of prokaryotic groups, our broad-scale spatial resolution allowed us to explore how this vertical structuring changed from east to west and how it compares with the adjacent Atlantic waters, looking for the particularities of each basin.

Prokaryotic richness increased with depth (Fig. 3), in line with previous studies performed in the Atlantic Ocean (Agoué et al. 2011; Frank et al. 2016), the Mediterranean Sea (Pommier et al. 2010; Luna 2014; Techtmann et al. 2015; Severin et al. 2016; Mestre et al. 2017), or in global ocean waters (Sunagawa et al. 2015). This increase in richness was caused by a high number of ASVs exclusively found in deep layers (Fig. 4), and to an increased contribution of rare members in the deep ocean communities (Fig. S10). Conversely, lower diversity and few exclusive ASVs were found in the euphotic layer, which were dominated by several SAR11 and cyanobacterial ASVs and displayed comparatively a reduced rare biosphere in all three basins (Fig. S10). Rare prokaryotes have been shown to guarantee the metabolic potential of deep ocean microbial communities in the absence of fresh carbon inputs (Sebastián et al. 2018), which may explain why deep waters harbor a larger rare biosphere. The existence of a diverse microbial community ensures the potential to quickly exploit sudden increases in substrates associated to active transport by migrating animals (Steinberg et al. 2000; Calleja et al. 2018; Hernández-León et al. 2019), sinking particles arriving to deep waters (Smith et al. 2018), or deep-water formation events (Tambrurini et al. 2013; Luna et al. 2016; Severin et al. 2016).

Richness and PD decreased from the Atlantic to the Eastern Mediterranean regardless of the depth layer. However, while in the epipelagic and the mesopelagic, they did not correlate with any of the environmental variables tested (Table S2), in the bathypelagic they were significantly correlated with the concentration of organic carbon, inorganic nutrients, and humic-like FDOM peak C. This suggests that deep waters with increased resource availability (i.e., in the Atlantic) sustain more diverse communities, or alternatively that richer communities might transform the available organic carbon into less available humic by-products, in what is called the microbial carbon pump (Jiao et al. 2010). The reason why we did not observe any correlation between diversity and environmental variables in the mesopelagic may be the fact that this layer spans a transition from epipelagic waters (hence illuminated and influenced by primary production) to deep waters (thus dark, nutrient-enriched, cold, and high-pressure dominated), and therefore, likely comprises a wide variety of habitats. In fact, in the Mediterranean Sea, samples that are globally defined as mesopelagic (from 200 to 1000 m) comprise not only intermediate but also deep-water masses (Catalá et al. 2018). To explore the hypothesis that the mesopelagic layer comprised a wide variety of niches, we calculated the coefficient of variation of several environmental...
parameters along the four depth layers (surface, DCM, meso- and bathypelagic; Fig. S11) and observed that mesopelagic waters displayed the highest variability in the variables related to particle organic matter pools (POC, PON, and TEP), suggesting a broad variation range in particle abundance and quality. Despite the lack of correlation with environmental variables, the high richness observed in this layer agrees with previous findings (Sunagawa et al. 2015) and with the idea that the mesopelagic is a hotspot for microbial diversity and activity (Aristegui 2005; Calleja et al. 2018).

The most oligotrophic waters (i.e., the Eastern Mediterranean) displayed communities with higher PD/richness ratios (Fig. S2), which were also negatively correlated with the concentration of organic carbon in deep waters (Table S2). One could hypothesize that oligotrophic waters favor the coexistence of more phylogenetically diverse phylotypes despite harboring globally less diverse communities, implying less microdiversity. Although not many studies have addressed this issue, these findings contrast with previous results obtained in a mesocosm study showing that taxa tended to be more distantly related at higher ecosystem productivities (Horner-Devine and Bohannan 2006). Given that closely related taxa seem to be more functionally similar than distantly related taxa (Bryant et al. 2012), the relative diversification we observed in Eastern waters may be a consequence of competitive exclusion between closely related taxa (Cavender-Bares and Wilczek 2003) facing poor trophic conditions. Nonetheless, further studies looking at the genetic repertoire and functional redundancy of prokaryotic communities could help to gain a clearer insight into how microbial communities structure across distinct trophic gradients.

**Prokaryotic communities are horizontally structured across the Mediterranean Sea**

Besides the high structuring in the vertical dimension, our analyses revealed also a clear spatial structuring in the horizontal dimension, particularly in the surface and the bathypelagic layers (Table 2; Fig. 6).

In surface waters, Atlantic indicator ASVs were abruptly substituted by Mediterranean ASVs, and we could document a gradual exchange of ASVs across the W–E transect (Fig. 6). The sharp decrease in the contribution of Atlantic indicator taxa at the interface of Mediterranean waters was mostly driven by the disappearance of Prochlorococcus in surface waters (Fig. S9). Eastern indicators were associated with lower values of photic-zone integrated nutrients (Fig. S7), which agrees with the known west to east gradient of increasing oligotrophy, and the strong phosphorus limitation in Eastern Mediterranean waters (Thingstad et al. 2005). This gradient was also reflected in the distribution of SAR11 populations: even though SAR11 are the most abundant taxa everywhere in the epipelagic, there was a spatial structuring of different surface clades. For instance, the surface 1 clade of SAR11 was mostly present in the Western Mediterranean while the surface 2 clade was mostly seen in the Eastern Mediterranean (Fig. S6). Even within surface 1 clade, different ASVs appear as indicators of the different basins (Fig. S7). Indeed, previous studies have already pointed out the importance of P limitation as a selective force shaping the biogeography of distinct SAR11 clades (Coleman and Chisholm 2010; Salter et al. 2014). Cyanobacteria also presented a high degree of spatial structuring within the Mediterranean: while Synechococcus were mostly present in the surface of the Levantine basin (Eastern Mediterranean), Prochlorococcus appeared deeper, at around 100 m deep (Fig. S4). This vertical segregation in the Levantine basin is in line with previous studies (Techtmann et al. 2015), and agrees with the idea that Synechococcus grow better at low nutrients and high light levels (Agawin et al. 1998; Nunes et al. 2018), while Prochlorococcus dominates at lower light levels. Notably, Cyanobacteria were less abundant overall in the Western Mediterranean, coinciding with those stations with higher Chl a concentration (Martinez-Perez et al. 2017), which may indicate that these waters were dominated by eukaryotic phytoplankton (Aristegui, Gasol et al. unpubl.). However, these groups may display strong seasonality (Malmstrom et al. 2010), so the patterns observed here could be different in other times of the year (Mena et al. 2020).

Furthermore, these groups may display strong seasonality (Malmstrom et al. 2010), so the patterns observed here could be different in other times of the year (Mena et al. 2020). The practically absence of basin DCM indicators may be due to the fact that this structure is highly heterogeneous at a fine vertical spatial scale (Latasa et al. 2016; Haro-Moreno et al. 2018), as reflected by the variable taxonomic composition observed along the transect (Fig. 5). Nonetheless, despite the lack of horizontal structuring between the different basins, we found some groups to be preferentially enriched in this layer. One clear example is the OM1 clade of Actinobacteria (Table 2). This clade was first described in the Western Mediterranean (Candidatus Actinomarina; Ghai et al. 2013), but in our study it also accounted for a large share of microbial community composition in the Eastern basin near the strait of Sicily and in the Atlantic basin (Fig. S12).

Mesopelagic prokaryotic communities were also poorly structured throughout the basins, as indicator ASVs were few and always showed low relative abundances along the transect, except for a slight increase in the Western basin (Fig. 6). Besides the presence of different intermediate and deep-water masses in our mesopelagic samples, the ubiquitous distribution of mesopelagic ASVs may be due to the fact that the Mediterranean mesopelagic is connected through the Levantine Intermediate Water (LIW), a water mass formed in the Eastern basin that reaches 200–500 m depth (Malanotte-Rizzoli and Hecht 1988; Catala et al. 2018), and travels west, flowing cyclonically in the Western basin while being progressively mixed with surrounding waters until it exits the Mediterranean Sea through the strait of Gibraltar (Emelianov et al. 2006). Hence, the LIW connects all Mediterranean Sea mesopelagic waters, presumably also connecting their prokaryotic communities and thus erasing biogeographical barriers. The LIW also transports DOM that is progressively consumed by prokaryotes (Martinez-Perez
et al. 2017), thus Western Mediterranean mesopelagic waters hold more refractory low-molecular weight DOM that could select for specific ASVs degrading it. This may be a reason for the increase in the relative abundance of mesopelagic indicators observed in this basin (Fig. 6).

As opposed to DCM and mesopelagic communities, there was a marked horizontal structuring of benthopelagic communities, mostly between the Atlantic and the Mediterranean basins, as indicated by the high percentage of exclusive ASVs in the Atlantic (Fig. 4) and the sharp changes in the contribution of basin indicator ASVs when crossing the strait of Gibraltar (Fig. 6). This shows that the shallow Gibraltar and Sicily straits act as physical barriers that prevent the direct mixing and spreading of the deep-water bodies in the Mediterranean (Astraldi et al. 1999), creating unique environments that lead to the ecological selection of specialized taxa. In fact, closely related phylotypes appeared as indicators of the different basins (Fig. S8), suggesting a fine-tuning of metabolic capacities among closely related microbes facing the different environmental conditions of the isolated water masses within each basin, as has been proposed to explain prokaryote niche partitioning in other systems (Carlson et al. 2008; Hehemann et al. 2016). Temperature has been found to be one of the most determinant factors in the ecological selection of prokaryotes (Sunagawa et al. 2015; Milici et al. 2016), and benthopelagic Atlantic waters are characterized by temperatures ~9°C colder than benthopelagic Mediterranean waters. Thus, it is likely that the segregation of taxa between the Atlantic and the Mediterranean is due to the drastic change in temperature, as hinted by the sPLS analyses (Fig. S8). Between the Mediterranean basins, other environmental factors such as the amount of organic carbon that reaches deep-water layers or the quality of the DOM may play a role, as depicted by the association of Eastern basin benthopelagic indicators with comparably lower amount of TOC and higher molecular weight of its compounds (lower S275–295 CDOM spectral slope, Helms et al. 2008; Català et al. 2018) (Fig. S8).

Zooming in the meso- and bathypelagic layers, some prokaryotic groups were particularly enriched in the Atlantic Ocean (Table 2), such as Lentisphaerae, or the Arctic 97B-4 marine group of Verrucomicrobia. Lentisphaerae have been described to be exopolymer particle producers (Cho et al. 2004), and higher proportions of exopolymer particles have been reported in the deep Atlantic Ocean compared to the deep Mediterranean Sea (Ortega-Retuerta et al. 2019). SAR324 bacteria were also enriched in the Atlantic meso- and bathypelagic, but were abundant in the mesopelagic layer of the Eastern Mediterranean basin (Fig. S5). The later observation is in line with a previous study that proposed SAR324 as indicative prokaryotes of the LIW in this area (Techtmann et al. 2015). This group is characterized by high metabolic versatility, being able to fix carbon with the energy derived from sulfur oxidation but also being capable of hydrocarbon oxidation (Swan et al. 2011; Sheik et al. 2014). Remarkable trends were observed in the distribution of the nitrite oxidizers across the Mediterranean Sea, with Nitrospina showing higher relative abundances throughout the dark waters of the Atlantic and the mesopelagic of the Western Mediterranean, and Nitrospira displaying higher relative abundances in the dark waters of the Eastern Mediterranean basin (Fig. S5). Members of these groups play an important role in carbon fixation in the deep ocean (Pachiadaki et al. 2017), but their contrasting biogeography suggests they are governed by different selective forces (Nunoura et al. 2015; Hou et al. 2018).

Among the taxonomic groups that were relatively enriched in the dark waters of the Eastern Mediterranean Sea (Table 2; Fig. S5), we underline the Gammaproteobacteria group E01-9C-26, with no cultured representatives but potential versatile methylo trophs and dissimilatory sulfate reducers as hinted by single cell genomics (Landry et al. 2018) and the Chloroflexi-related SAR202 group, frequently observed in deep-water samples (Severin et al. 2016) and in the Eastern Mediterranean Sea (Mehrshad et al. 2018). SAR202 are potential sulfite oxidizers that may use recalcitrant carbon substrates (Landry et al. 2017, 2018; Liu et al. 2020). Given the oligotrophic character of Eastern Mediterranean waters, one could hypothesize that the “hostile” conditions in the carbon-limited bathypelagic would promote a higher diversification of prokaryotic metabolisms, something that will need to be examined in future studies.

Our study presents a quasi-synoptic view of prokaryotic diversity across the Mediterranean Sea and the nearby Atlantic Ocean. It clearly identifies a strong biogeographical spatial structuring, and a large differentiation not only between depth layers but also between basins. This structuring is stronger in the surface and in the bathypelagic layers, pointing to the existence of biogeographic barriers within these layers that are likely caused by the oligotrophic gradient in the surface and the isolation of the bathypelagic water bodies at the Gibraltar and Sicily straits. Our sampling was performed during mid spring (May 2014) when stratification starts to strengthen and the temperature and light increase culminates in the early spring phytoplankton blooms that vary strongly in timing and magnitude in different basins of the Mediterranean Sea (Lazzari et al. 2012). It is thus plausible that the basin and vertical structuring of prokaryotic communities is even stronger in late spring and summer, when oligotrophy becomes more intense and organic matter production feeding the deep-sea microbes is scarcer. Likewise, mixing and deep-water formation events in winter could lead to DOC export to the mesopelagic and sometimes down to the bathypelagic (see Santinelli et al. 2015 and references therein). This, together with seasonal changes in the intensity of the inflows of LIW through the Strait of Sicily and Atlantic water through the Strait of Gibraltar (Manzella and La Violette 1990), could also drive changes in the spatial and vertical structuring of prokaryotic communities. Hence, the biogeographic patterns evidenced here should serve as basis for further detailed
studies on prokaryotic community structure in the Mediterranean and other ocean basins, the environmental factors driving this structure, and how it varies over time, as well as on the impact of this regionalization in the function of prokaryotic communities.

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Conflict of Interest
None declared.