

ORIGINAL ARTICLE

A new class of marine Euryarchaeota group II from the mediterranean deep chlorophyll maximum

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We have analyzed metagenomic fosmid clones from the deep chlorophyll maximum (DCM), which, by genomic parameters, correspond to the 16S ribosomal RNA (rRNA)-defined marine Euryarchaeota group IIB (MGIIIB). The fosmid collections associated with this group add up to 4 Mb and correspond to at least two species within this group. From the proposed essential genes contained in the collections, we infer that large sections of the conserved regions of the genomes of these microbes have been recovered. The genomes indicate a photoheterotrophic lifestyle, similar to that of the available genome of MGIIA (assembled from an estuarine metagenome in Puget Sound, Washington Pacific coast), with a proton-pumping rhodopsin of the same kind. Several genomic features support an aerobic metabolism with diversified substrate degradation capabilities that include xenobiotics and agar. On the other hand, these MGIIIB representatives are non-motile and possess similar genome size to the MGIIA-assembled genome, but with a lower GC content. The large phylogenomic gap with other known archaea indicates that this is a new class of marine Euryarchaeota for which we suggest the name *Thalassoarchaea*. The analysis of recruitment from available metagenomes indicates that the representatives of group IIB described here are largely found at the DCM (ca. 50 m deep), in which they are abundant (up to 0.5% of the reads), and at the surface mostly during the winter mixing, which explains formerly described 16S rRNA distribution patterns. Their uneven representation in environmental samples that are close in space and time might indicate sporadic blooms.

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Introduction

Marine Euryarchaeota group II (MGII) are widely distributed in the global ocean (Massana *et al.*, 2000) and are the dominant archaeal community within the euphotic zone of temperate waters (DeLong, 1992; Massana *et al.*, 1997, 2000; Karner *et al.*, 2001; Herndl *et al.*, 2005; DeLong *et al.*, 2006; Belmar *et al.*, 2011). They have been classified into two major groups, MGIIA and MGIIIB, by their 16S ribosomal RNA (rRNA) (Massana *et al.*, 2000; Martín-Cuadrado *et al.*, 2008; Galand *et al.*, 2009; Belmar *et al.*, 2011). Two other groups, MGIIIC and D, related to hydrothermal and deep samples have also been proposed (Belmar *et al.*, 2011). In addition, several subclusters have

been identified in each of these general groups (Massana *et al.*, 2000; Galand *et al.*, 2010). For example, Euryarchaeota MGIIA is separated in subclusters K, L and M, and MGIIIB is separated into subclusters O and N. Subcluster O comprises the WHARM subcluster (formerly phylotype II-CC) (Massana *et al.*, 2000), widely distributed in surface waters (Galand *et al.*, 2010; Hugoni *et al.*, 2013). MGII representatives have a cosmopolitan distribution and are relatively abundant during summer months in temperate and tropical waters (Pernthaler *et al.*, 2002; Herfort *et al.*, 2007). In Mediterranean surface waters, there are reports of different seasonal prevalence, with MGIIIB predominating in winter and MGIIA in summer (Galand *et al.*, 2010; Hugoni *et al.*, 2013). There is also evidence that indicates MGIIA is found more often at surface samples (Frigaard *et al.*, 2006), while MGIIIB is more frequently found in deeper waters (Massana *et al.*, 2000; Galand *et al.*, 2010; Hugoni *et al.*, 2013).

To date, there are no cultivated representatives of MGII and little is known about their physiology and ecological role in the oceans. For years, the only information available came from a handful of

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metagenomic clones containing phylogenetic markers (16S and 23S rDNA) (Beja *et al.*, 2000; Moreira *et al.*, 2004; Frigaard *et al.*, 2006; Martin-Cuadrado *et al.*, 2008). In 2012, by using the *de novo* assembly from a metagenome from superficial estuarine waters, a composite genome sequence grouping 4–6 strains of Group II archaea (MG2-GG3) was published (Iverson *et al.*, 2012). Recently, two single amplified genomes (SAGs) classified as MGII were released to the Genbank database (SCGC-AAA-288-C18, isolated from the North-Pacific Subtropical Gyre at 700 m deep, and SCGC-AB-629-J06 from Lake Washington, USA). However, both SAGs lack the ribosomal operon and their similarity to the MG2-GG3-assembled genome is very low (the euryarchaeal affiliation could be ascertained only for SCGC-AAA-288-C18). MG2-GG3, similar to many other marine prokaryotes, seems to be a photoheterotroph, possessing a rhodopsin to collect light energy but with the hallmarks of a heterotrophic lifestyle. The existence of many large peptidases, similar to the ubiquitous thermoacidophilic archaeon *Aciduliprofundum boonei*, a known protein degrader from deep-sea hydrothermal vents (Reysenbach *et al.*, 2006), suggested that proteins may be important substrates for this microbe. Moreover, the presence of a complete fatty acid degradation pathway, together with proteins with a variety of adhesion domains and a type II/IV secretion systems to transport such proteins to the cell surface, led the authors to suggest a particle-associated lifestyle. The analysis of the MGII transcripts from coastal surface samples in northern California (Ottesen *et al.*, 2013) also support the ability to metabolize large and complex polymers.

In most of the global ocean when the photic zone becomes thermally stratified (in summer, at temperate latitudes and permanently at tropical ones), phytoplankton biomass concentrates at the deep chlorophyll maximum (DCM) (Estrada *et al.*, 1993; Huisman *et al.*, 2006). The DCM is a layer, usually between 50 and 150 m deep, which is free from the strong hydrodynamism and damaging UV light of the surface and, on the other hand, is connected to the nutrient-rich deep waters. Thus, the DCM has a good compromise of available light and inorganic nutrients availability for planktonic photosynthetic microbes. There is some evidence in the literature, indicating that MGII could be relatively abundant at DCM depths. The maximum abundance of MGII in the Hawaii Ocean Time-Series permanent DCM by 16S rRNA gene amplification from a metagenomic fosmid library (DeLong *et al.*, 2006) was found at 130 m. Ghai *et al.* (2010) found rRNA fragments of MGII in proportions close to 4% in the Mediterranean DCM using raw metagenomic reads. In addition, in a metagenomic fosmid library from the same sample, up to 22 fosmids (from a total of 197 larger than 10 kb) were classified as belonging to MGII.

Sequencing large metagenomic clones provides a powerful strategy for obtaining valuable information about the structure and ecology of uncultured

microorganisms (Martin-Cuadrado *et al.*, 2008; Ghai *et al.*, 2013; Deschamps *et al.*, 2014). From a metagenomic library constructed from the biomass recovered at the DCM in the Mediterranean, we sequenced about 7000 fosmid clones (Ghai *et al.*, 2010, 2013; Mizuno *et al.*, 2013) and we have analyzed the MGII genome fragments present in these fosmids. MGII contigs assembled from different DCM metagenomes obtained from the same location at different times were also included in this study. Here we present the first genomic data about a new class of abundant low GC MGII, which has been named *Thalassoarchaea*, a distant relative to the class represented by the MG2-GG3 assembly (Iverson *et al.*, 2012).

Materials and methods

Sampling and sequencing

A metagenomic fosmid library (~13 000 clones) was constructed with biomass recovered from the Mediterranean DCM (50 m, 38°4'6.64"N 0°13'55.18"W) on 15 October 2007. A metagenome (MedDCM-OCT2007) directly sequenced by 454 from the same filter and the sequencing results of nearly 1000 fosmids have been described previously in Ghai *et al.* (2010). The sequencing of additional metagenomic fosmids (~6000) substantially extended these data sets and have been partially published in Ghai *et al.* (2013) and Mizuno *et al.* (2013). Three more samples were collected at the same location and at the chlorophyll maximum depth (as determined by a Seabird SBE 19, Sea-Bird Electronics, Bellevue, WA, USA, multiprobe profiler including fluorometers) in different years: June 2009 (65 m), July 2012 (75 m) and September 2013 (55 m). On each of these dates, sea water was collected at DCM depth and sequentially filtered on board using a positive pressure system. Nylon filters of 20 µm were used as prefilters, followed by 5 µm polycarbonate and finally 0.22 µm Sterivex filters (Durapore; Millipore, Billerica, MA, USA). Filters retaining the 20–5 µm (large fraction (LF), enriched in particle attached bacteria) and the 5–0.2 µm (small fraction, enriched in free-living planktonic cells) were conserved in lysis buffer (40 mM EDTA, 50 mM Tris/HCl and 0.75 M sucrose) at –20 °C until DNA extraction. Filters were thawed on ice and then treated with 1 mg ml⁻¹ lysozyme and 0.2 mg ml⁻¹ proteinase K (final concentrations). Nucleic acids were extracted with phenol–chloroform–isoamyl alcohol and chloroform–isoamyl alcohol, and then concentrated using a microconcentrator (Centricon 100; Amicon, Millipore). DNA integrity was checked by agarose gel electrophoresis and quantified with Quant-iT PicoGreen dsDNA Assay Kit (Invitrogen, Carlsbad, CA, USA).

Sequencing was done using Illumina HiSeq 2000 with 100 bp paired ends (Macrogen, Seoul, South Korea). For the 2009 sample, a total of 6.8 Gb of

sequence data was produced for the large-size fraction (metagenome MedDCM-JUN2009-LF). A total of 13.6 Gb was obtained for the free-living fraction of the 2012 sample (MedDCM-JUL2012). In addition to the 2012 sample, a mate-paired library of 3 kb insert size was constructed from DNA previously amplified using the Illustra GenomiPhi V2 DNA amplification kit (GE Healthcare, Piscataway, NJ, USA) (50 ng to a final amount of 5.2 μ g). As a result, an extra 6.3 Gb of sequence was obtained (metagenome MedDCM-JUL2012-3 kb). For the sample of September 2013, the sequencing of both size-fraction filters, 20–5 and 5–0.22 μ m, generated 7.7 and 10.5 Gb, respectively (metagenomes MedDCM-SEP2013-LF and MedDCM-SEP2013, respectively).

Assembly and annotation

A total of 146 genomic fragments (>5 kb) with a GC content of 34–40% could be classified as Thalamoarchaea by annotation and tetranucleotide frequencies. Their origin and other characteristics are summarized in Supplementary Table 1.

The assembly of the DCM metagenomic fosmid from October 2007 were previously described in (Ghai *et al.*, 2010, 2013; Mizuno *et al.*, 2013). Briefly, 7152 fosmids were sequenced using 454-pyrosequencing and Illumina in two different rounds, each divided in 12/24 different batches containing 96 to ~250 fosmids each. Using the contigs (>5 kb) assembled from this sample, a second assembly was performed using Geneious v.6.1.7 (<http://www.geneious.com>) with strict parameters (overlapping of >100 bp at 99% identity with no gaps). A total of 69 DNA fragments classified as Thalamoarchaea were obtained from this sample.

Sequences from metagenomes MedDCM-JUN2009-LF, MedDCM-SEP2013-LF and MedDCM-SEP2013 were quality trimmed and assembled independently using meta-iterative de Bruijn *de novo* assembler (Peng *et al.*, 2012). A combined assembly of the MedDCM-JUL2012 metagenome plus the sequences from the 3-kb library was also performed using the iterative de Bruijn *de novo* assembler (Peng *et al.*, 2012). A total of 68 DNA fragments were classified as Thalamoarchaea from MedDCM-JUL2012 and 9 more from the metagenome MedDCM-SEP2013. From the LF metagenomes (MedDCM-SEP2009-LF and MedDCM-SEP2013-LF), no Euryarchaeota contigs >5 kb were obtained, probably due to the low coverage of this taxa present and the high percentage of eukaryotic reads obtained (based on the 18S to 16S rRNAs ratios found, only ~1% of the sequences belonged to prokaryotes).

Gene predictions of the assembled fosmids were done using Prodigal (Hyatt *et al.*, 2010). Ribosomal genes were identified using *ssu-align* (Nawrocki 2009) and *meta_rna* (Huang *et al.*, 2009), and transfer RNAs were predicted using tRNAscan-SE (Lowe and Eddy 1997). Functional annotation was

performed by comparison of predicted protein sequences against the NCBI NR database, Pfam (Bateman *et al.*, 2004), COGs (Tatusov *et al.*, 2001) and TIGRfams. Local BLAST searches against local databases were performed whenever necessary. The identification of MGII contigs was based on the condition that >50% of the open reading frames (ORFs) contained in the DNA fragment had their best hit to Euryarchaeota, and for this manual examination of each contig was performed. The contigs were named according to the time of sampling, that is, MedDCM-OCT2007-CX, MedDCM-JUL2012-CX and MedDCM-SEP2013-CX, where X is the number of the contig.

Oligonucleotide frequency analysis

The tetranucleotide and pentanucleotide frequencies of the contigs were computed using the *wordfreq* program from the EMBOSS package (Rice *et al.*, 2000). Principal components analysis was performed using the *FactoMineR* package in 'R' (Lê *et al.*, 2008). The genomes of *A. boonei* T469, MG2-GG3, *Nitrosopumilus maritimus* and the single-cell genome of SCGC-AAA288-C18 were included in the analysis as references of marine archaea. The frequencies of randomly generated 30 kb fragments of these reference genomes were also included in the analysis. From the initial 154 genomic fragments, a total of 146 (69 from the fosmid collection MedDCM-OCT2007, 68 from MedDCM-JUL2012 and 9 from MedDCM-SEP2013) were clustered together and are described here.

Phylogenetic analysis

Archaeal 16S rRNA and 23S rRNA gene sequences were detected in the genomic fragments and aligned using MUSCLE (Edgar 2004) with their closest relatives as identified by BLAST (Altschul *et al.*, 1990), those from available Euryarchaeota genomes and selected genome fragments present in Genbank and those from the Global Ocean Time Series (GOS) data set (available from <http://camera.calit2.net/>). Assembled site-specific GOS scaffolds were screened for the presence of ribosomal genes using a stringent cutoff of 98% identity in at least 97% of their length, ensuring that they belonged to the same lineage as the Mediterranean thalamoarchaeal 16S and 23S rRNA sequences assembled from the fosmids and metagenomes. Using the same stringent criteria, the entire Ribosomal Database Project (Cole *et al.*, 2009) was also searched to identify related sequences. 16S rRNA sequences were screened and trimmed using *ssu-align* (Nawrocki, 2009). Phylogenetic reconstructions were conducted by maximum-likelihood and neighbor-joining methods using MEGA5 (v.5.2, GTR 1 CAT model and a gamma approximation with 1000 bootstraps) (www.mega-software.net) (Hall, 2013), first using those sequences longer than 700 bp as reference and later

including smaller metagenomic reads. In both cases, tree topologies were consistent.

For the rhodopsin tree and the geranylgeranyl-glyceryl phosphate synthase proteins, sequences were selected based on existing literature, PFAM domain searches and BLAST searches against NCBI-NR and the GOS data set metagenomic reads. Sequences were aligned using MUSCLE (Edgar, 2004) and a maximum-likelihood tree was constructed using MEGA5 (v.5.2, GTR 1 CAT model and a gamma approximation with 1000 bootstraps) (Hall, 2013).

Fluorescence in situ hybridization

For fluorescence *in situ* hybridization (FISH) detection of the Thalamoarchaea cells, a DCM water sample was recovered in June 2009 and fixed with a 1% paraformaldehyde solution. Subsamples of 15 ml were filtered at low pressure (100 mbar) through 0.2 μ m pore size filters (Nucleopore, Whatman, Piscataway, NJ, USA) and kept at -20°C . A new probe was used, specifically targeting low GC Thalamoarchaea (ThaMar 5'-GTAGTGAACTATGG ATCATTA-3'). The general euryarchaeal probe EURY806 (5'-CACAGCGTTTACACCTAG-3') was used as a control. Oligonucleotide probes for FISH analysis were purchased from Isogen (Utretch, The Netherlands) and 5' monolabeled with the indocarbocyanine dye, Cy3. The same filters were cut in quarters for hybridization with the different probes. FISH analysis was carried out by the standard protocol (Glockner *et al.*, 1999) using $5\text{ ng}\mu\text{l}^{-1}$ of probe. The hybridization solution was prepared as described in DeLong (1992) and DeLong *et al.* (1999), with the following variations: 50% (vol/vol) formamide, 10% (wt/vol) dextran sulfate in $5 \times \text{SET}$ ($1 \times \text{SET}$ is 150 mM NaCl, 1 mM Na_2EDTA , 20 mM Tris-HCl, pH 7.8). Hybridization mixtures were incubated overnight in a hybridization oven at 65°C . The filters were washed for 2 h at 45°C in a solution containing $0.2 \times \text{SET}$ and 50% (vol/vol) formamide. After incubation in a pre-warmed washing solution, filter sections were dipped in 80% ethanol, dried on Whatman 3M paper (Whatman Ltd, Jenbach, Austria) and placed on a glass slide. Subsequently, they were mounted with a drop of DAPI (4',6-diaminodino-2-phenylindole) at a final concentration of $1\ \mu\text{g}\text{ml}^{-1}$.

Metagenomic recruitment

Recruitment plots of the virtual genome reconstructed with the thalamoarchaeal contigs, the MG2-GG3 and SAG SCGC-AAA-288-C18 were performed using BLASTN (a hit was considered only when the alignment was at least 50 nucleotides long). For estimating the abundance of these MGII cells in the metagenomes studied, only the best hits with an identity over 95% in at least 50 bp were considered. To compare the results among the different data sets, the number of reads was

normalized considering the size of the contig and of the collection (number of reads per kilobase per gigabase of the collection). The meso and bathypelagic metagenome collections screened were: Marmara Sea-1000 m (Quaiser *et al.*, 2011), Mapan-Vavilov Deep in the Mediterranean Sea-4900 m (Smedile *et al.*, 2012), North Pacific subtropical gyre-ALOHA station-4000 m (Konstantinidis *et al.*, 2009), Guaymas Basin hydrothermal vent in the Gulf of California (Lesniewski *et al.*, 2012) and a metagenome obtained at 800 m in the South Atlantic Gyre (Swan *et al.*, 2014).

Genome and proteome comparisons

For the genomic comparison showed in Figure 4, the thalamoarchaeal contigs were aligned with the genomes of MG2-GG3 and *A. boonei* T469 by TBLASTX, and only those alignments longer than 200 amino-acids were shown. Contigs with no similarities are not shown. A total of 4070 ORFs were detected within the 146 marine thalamoarchaeal contigs identified. Due to the occurrence of several overlaps among the DNA fragments, some genes were represented more than once. The proteins were clustered using CD-HIT at 50% similarity and 50% of coverage, resulting in a non-redundant proteome of 2435 ORFs. This set of proteins was compared separately to the proteins of the MG2-GG3 (1698), *A. boonei* T469 (1544) and SCGC-AAA288-C18 (398) genomes using a reciprocal best blast hit analysis to identify putative homologs. The same set was also compared with the proteins detected in the bathypelagic fosmids from the KM3 (9946 proteins), AD1000 (4215) and SAT1000 (1301) collections (Deschamps *et al.*, 2014). Previously, all the sequences used in the comparison were annotated using the same methodology as the thalamoarchaeal contigs to avoid methodological discrepancies.

Accession numbers

The assembled contigs and the metagenomic data have been submitted to NCBI SRA and are accessible under the BioProject identifier PRJNA257723 (SRA run identifiers: MedDCM-SEP2013-LF SRR1539385; MedDCM-SEP2013 SRR1539645; MedDCM-JUL2012-3kb SRR1539382; MedDCM-JUL2012 SRR1539383; MedDCM-JUN2009-LF SRR1539203).

Results and discussion

Retrieval of rRNA-containing fosmids belonging to MGII
Metagenomic fosmid clones provide discrete natural contigs that can be efficiently assembled to obtain genomic fragments from all members in the community, even from those that are less prevalent and hence less accessible by direct sequencing and assembly (Ghai *et al.*, 2013). During a systematic search of a Mediterranean DCM metagenomic

fosmid library (Ghai *et al.*, 2010) for rRNA genes, we identified four fosmids (6.2 to 36 to kb in size) containing rRNA sequences classified as MGII, two of them containing 16S rRNA and the other two 23S rRNA genes (Supplementary Figure 1). Interestingly, they had an atypical low GC content (34–40%) compared with the 52% of the assembled MG2-GG3 genome (Iverson *et al.*, 2012) or the MGII fosmids previously described (DeLong *et al.*, 1999; Moreira *et al.*, 2004; Frigaard *et al.*, 2006; Martin-Cuadrado *et al.*, 2008; Rich *et al.*, 2008; Ghai *et al.*, 2010; Deschamps *et al.*, 2014). MGII have the two rRNA genes separated in the genome (Moreira *et al.*, 2004; Martin-Cuadrado *et al.*, 2008), a feature observed also in the Thermoplasmatales and in *A. boonei* (Ruepp *et al.*, 2000; Moreira *et al.*, 2004). Hence, it was not surprising that the 16S and 23S genes were not contained in one single contig. The 16S rRNA genes detected in the DCM fosmids belonged to group IIB, hitherto known only by 16S rRNA sequences. The two sequences were only 94% similar (well below the species threshold) but both clustered in the large subcluster O of MGIIB (Figure 1). One of them was affiliated with the surface WHARN subcluster (Massana *et al.*, 2000). Fortunately, one large ribosomal protein cluster (spectinomycin or *spc* operon) is found in the proximity of the 16S rRNA gene (Moreira *et al.*, 2004; Martin-Cuadrado *et al.*, 2008), providing a very robust phylogenetic anchor. We generated maximum-likelihood trees for the alignments of the 16S rRNA genes (Figure 1) and also for a concatenated alignment of the 22 ribosomal proteins of the *spc* operon (Supplementary Figure 2). All analyses produced consistent results and unambiguously placed the sequences in a group that corresponds to the 16S rRNA group IIB (MGIIB). It is important to emphasize that the only genome available for MGII (Iverson *et al.*, 2012) belongs to the 16S rRNA clade MGIIA. The two fosmids containing 23S rRNA genes were also clearly euryarchaeal. Although the number of 23S rRNA genes available for MGII is much smaller, a maximum-likelihood tree also clustered them separately from the 23S rRNA of the MG2-GG3 (Supplementary Figure 3), and belonging to a separate euryarchaeal clade. By GC content and other genomic parametric approaches, the 23S rRNA containing fosmids belonged to the same group as the 16S rRNA containing ones (see below). Confirming these results, the phylogenetic relationship of the typically archaeal geranylgeranylglycerol phosphate synthase (GGGP) genes identified in fosmids from the same group, clustered together with *A. boonei* T469 and the Thermoplasmatales single cell genome SCGC AB-539-N05, isolated from marine sediments and clearly separated from the cluster of MG2-GG3 (Supplementary Figure 4). The affiliation of this gene also suggests that MGIIB may share the tetraether membrane lipids characteristic of the Thermoplasmatales (DeLong, 2006).

We examined whether similar sequences had been assembled before from other metagenomic data sets by searching the 16S rRNA and 23S rRNA genes in the entire collection of assembled scaffolds from the GOS data set (Venter *et al.*, 2004; Rusch *et al.*, 2007). Five GOS scaffolds were retrieved with a 16S rRNA 100% identical over 97% of the gene to either of our MGIIB 16S rRNA genes (species threshold level) (Supplementary Figure 1), and an additional five at 95% identity at 95% coverage. Therefore, it seems that the microbes represented by the MGIIB fosmids described here are geographically widespread and not an artifact of assembly. The GOS scaffolds were short and contained mostly only the rRNA sequence. However, when they also presented a few more genes they were syntenic to our fosmids. It is interesting to note that most of them were from temperate or tropical regions. When a similar search was performed using the MGIIA 16S rRNA found in the assembled MG2-GG3 (Iverson *et al.*, 2012), no 16S rRNA sequence was found at >97% identity, suggesting less prevalence of this clade in open ocean or coastal metagenomes. In an effort to visualize the abundance of these MGIIB cells at the DCM we carried out FISH with a lineage-specific probe designed with the sequence of the 16S rRNA present in our fosmids. The plankton sample used came from the DCM of June 2009 (see Methods). A specific probe was designed to detect MGIIB and we could see a significant number of DAPI stained cells that hybridized with the probe (1.8%, $n = 10$). Cells were very small, less than a 1 μm (Figure 2). Considering that we are providing an image for these microbes and on the grounds of the phylogenetic distance we would like to propose the taxonomic rank of class for the members of this group that overlaps with the 16S rRNA defined group IIB. We suggest the name *Candidatus* Thalassoarchaea marina and *Candidatus* Thalassoarchaea mediterranei, from the greek name for the sea ‘thalassa’, for the microbes represented by the 16S rRNA sequences described here and stained by the probe (Figure 1).

Genomic reconstruction of MGIIB representatives

Reconstructing genomes from metagenomes is very challenging. First of all, there is the problem of the concurrent diversity. There are always several clones coexisting at any given time that although similar are by no means identical (Rodriguez-Valera *et al.*, 2009; Gonzaga *et al.*, 2012; Lopez-Perez *et al.*, 2013; Kashtan *et al.*, 2014). Secondly, some regions of these genomes are hypervariable, i.e. they vary sharply from one clone to another and are very poorly covered even in high coverage metagenomes. All these caveats prevent the reconstruction of complete and closed genomes from environments in which a significant diversity of microbes is present. Using metagenomic fosmids help in the process by providing long natural contigs that can be compared and joined together by genomic binning (Ghai *et al.*, 2013).

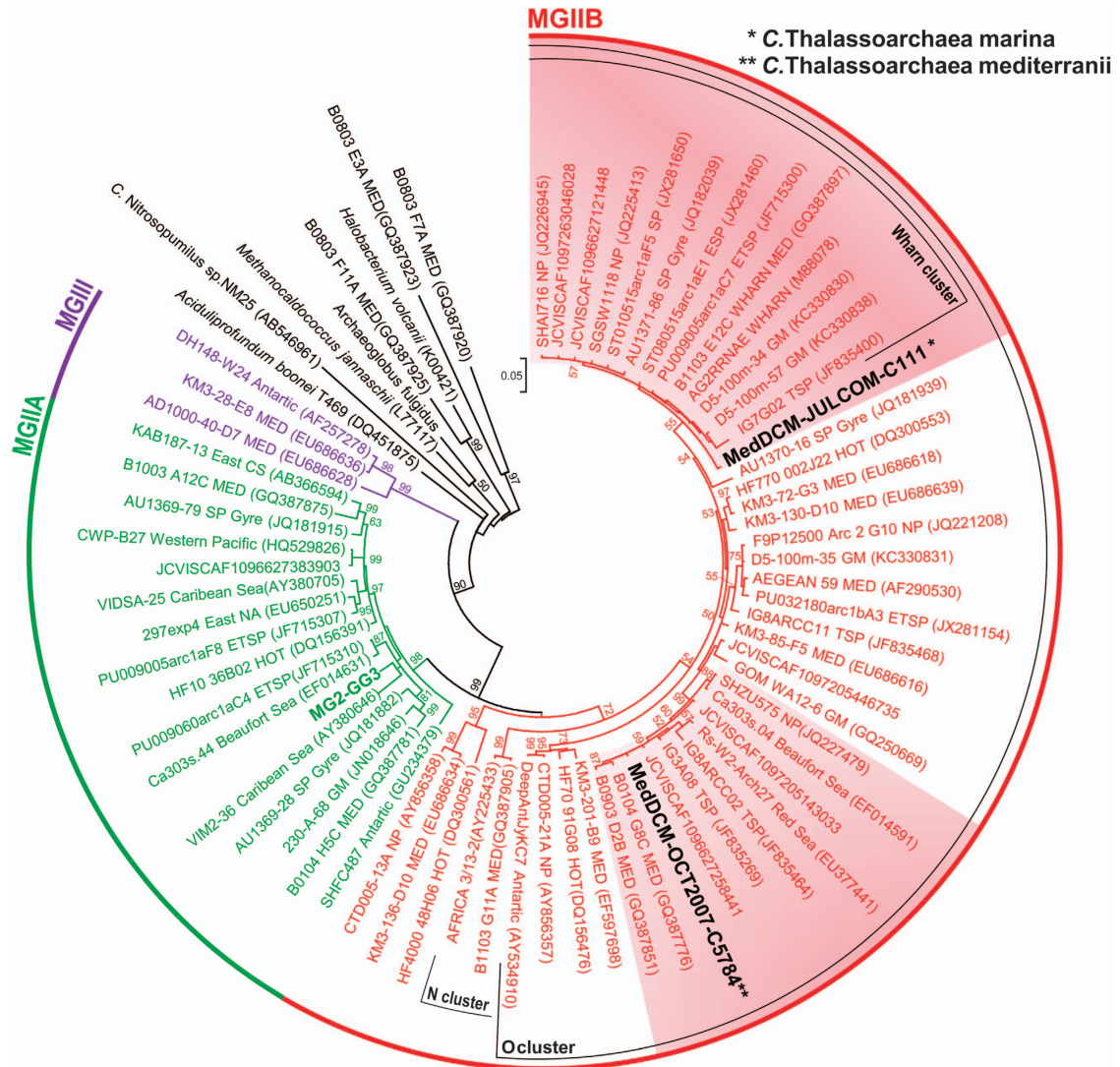


Figure 1 16S rRNA phylogeny. Maximum-likelihood 16S rRNA gene tree (739 unambiguously aligned nucleotides) showing the relationship of the Thalassoarchaea (bold and black) with other MGII (MG2-GG3-assembled genome in bold and green). The clusters where thalassoarchaeal sequences appear are red shadowed. Numbers at nodes indicate bootstrap support (shown as percentages and only those >50%) by ML in MEGA 5.10. Scale bar represents the estimated number of substitutions per site. The different clusters are named following Galand *et al.*, 2010. Sampling locations: MED, Mediterranean Sea; HOT, Hawaii Ocean Time-Series, North Pacific Gyre (ALOHA station); SP, South Pacific; ETSP, Eastern Tropical South Pacific; ESP, Eastern South Pacific; WP, western Pacific; SA, South Atlantic; NA, North Atlantic; GM, Gulf of Mexico; NP, North Pacific; CS, China Sea; NAEC, North American East Coast; GI, Galapagos Islands; CS, Caribbean Sea; TSP, Tropical South Pacific. The GOS data set identifiers are shown next to each GOS scaffold.

We identified a total of 146 fosmids (total sequence ~4 Mb) that were affiliated to MGIIB by binning the fosmids containing similar tetranucleotide frequencies to the MGIIB 16S and 23S rRNA containing fosmids described above (Figure 3) (the main features of the contigs can be seen in Supplementary Table 2). They represent the first extensive collection of genomic fragments from this elusive marine euryarchaeal group. All the fosmids were manually curated and we checked that most hits of the ORFs belonged to Euryarchaeota. The comparison with the assembled genome MG2-GG3 (Iverson *et al.*, 2012) or to the cultivated euryarchaeon *A. boonei* T469 indicated only a distant relationship to either, and

very little synteny (Figure 4). The clustered fosmids had lower and homogeneous GC content (range 34.1–40.4%, average 37.03%), much lower than the one described for the MG2-GG3-assembled genome (52%) (Iverson *et al.*, 2012) and closer to *A. boonei* T469 (34%). The MGII fosmids described up to now in the literature have also higher GC content (DeLong 1992; DeLong *et al.*, 1999; Moreira *et al.*, 2004; Frigaard *et al.*, 2006; Martin-Cuadrado *et al.*, 2008; Rich *et al.*, 2008; Ghai *et al.*, 2010). The low similarity to other available genomes concurs with the rRNA and the ribosomal protein concatenate phylogenetic trees (see above) supporting that these microbes belong to a new class.

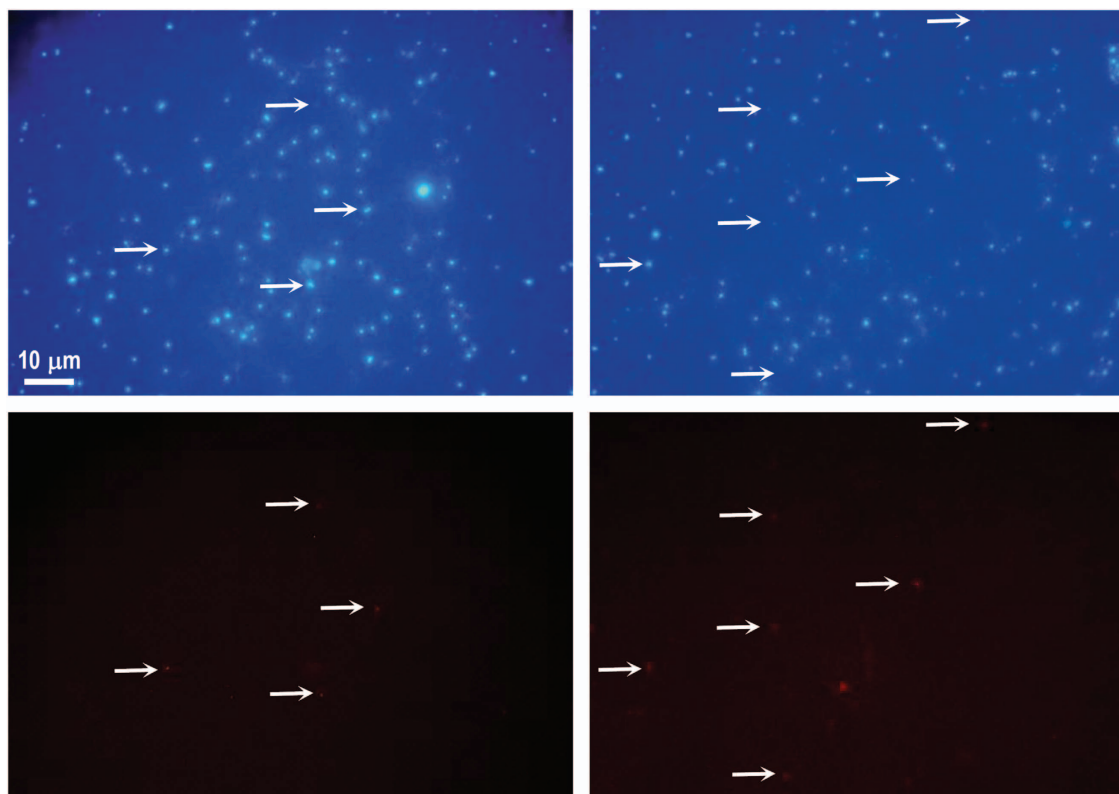


Figure 2 Microscopy of MGII cells. DCM seawater sample from Mediterranean Sea hybridized with fluorescein-labeled 16S rRNA group II archaeal probes (left panel, ‘ThaMar’ probe and right panel, ‘Eury 806’ probe). Images of the same field were captured using the fluorescein filter set (lower panels) and the DAPI filter set (upper panels). Scale bars, 10 μm .

The identity of the two thalassoarchaeal 16S rRNAs (94%) indicates that more than one species were present simultaneously in the fosmid collection. In addition, we found several contigs from overlapping genome regions that were largely syntenic but which average nucleotide identity (ANI) ranged from 98% to 80% (Figure 4), i.e. consistent with more than one species being present. To establish how many we searched among the DNA fragments for different versions of 36 housekeeping genes (Supplementary Table 3). The results indicate that although there are clearly more than one abundant species, their number is likely not very high, probably two, and diverging less than 20% ANI, i.e. within a single genus. The use of other sequence parameters such as codon usage, pentanucleotide frequencies, %GC or the coverage in the metagenomes did not permit to separate them.

We analyzed the contigs for the presence of 35 (Raes *et al.*, 2007) or 100 (Albertsen *et al.*, 2013) previously defined orthologous markers to estimate the completeness of the assembled genomes. Using these criteria, the thalassoarchaeal genomic fragments retrieved represent between 78 and 100% of a complete genome. By the same extrapolation, the estimated genome length of an individual genome of this group would be $\sim 2\text{Mb}$, very similar to the estimated size of the MG2-GG3 genome (Iverson *et al.*, 2012). A total of 4074 ORFs could be identified in the thalassoarchaeal fosmids. Most hits (61%) against the

nr database (Genbank) were to Euryarchaeota, 16% to Bacteria and 2% to eukaryotes (the remaining 21% were unclassified). The corresponding proteins were clustered using CD-HIT at 50% similarity and coverage resulting in a smaller data set of 2435 non-redundant proteins. This set was compared to the 1698 proteins of the MG2-GG3 genome (using a reciprocal best blast hit analysis) and 1427 proteins were found to share more than 50% similarity. However, the average similarity was low (65%). Only 639 homologs were found within the 1544 proteins of *A. boonei* T469 (average similarity 60%). The SAG SCGC-AAA-288-C18 from 700 m deep in the central North Pacific also had a low similarity (average 70%). The highest similarities were found with the proteins of the large set of MGII fosmids retrieved from bathypelagic Mediterranean samples (Ionian Sea 3000 m, 324 fosmids; and Adriatic Sea 1000 m, 139 fosmids) (Deschamps *et al.*, 2014). Of the non-redundant thalassoarchaeal proteins 1599 had a hit with Adriatic (38% of their total) and 1850 with the Ionian fosmids. However, synteny was seldom conserved and average similarity was of 74 and 73% respectively, suggesting that the deep MGII belong to very different taxa.

Inferred metabolic and ecological features

We found genes coding for enzymes for glycolysis, the tricarboxylic acid cycle and oxidative

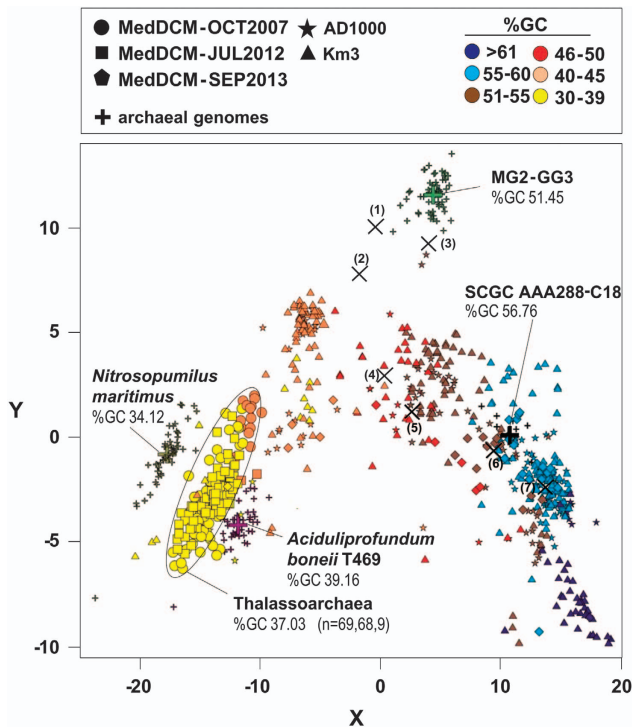


Figure 3 Comparison of MGII fosmid and contigs with known archaeal genomes. Principal component analysis of tetranucleotide frequencies of the low-GC MGII MedDCM-OCT2007 fosmid and the assembled low-GC MGII contigs from the MedDCM-JUL2012 and MedDCM-SEP2013 data sets. Reference genomes are shown as crosses (*N. maritimus*, *A. boonei* T469, MG2-GG3 and the SAG SCGC_AAA288-C18). Smaller crosses around the larger ones represent the tetranucleotide frequencies of 35 kb fragments from the same genome. Colors correspond to the %GC accordingly with the legend. The contigs classified as Thalassoarchaea are marked by a circle and the number of contigs from each of the data sets (MedDCM-OCT2007, MedDCM-JUL2012 and MedDCM-SEP2013) is indicated. The length of the assembled DNA fragments and %GC average are also shown. Numbers in brackets indicate MGII fosmid containing 16S rRNA sequences previously described: (1) *HF10-29C11*, %GC 45.56; (2) *HF10-3D09*, %GC 46.84; (3) *HF70-59C08*, %GC 51.36; (4) *HF70-19B12*, %GC 48.75; (5) *EF100-57A08*, %GC 50.66; (6) *DeepAnt-15E7*, %GC 56.03; (7) *HF4000-APKG2H5*, %GC 56.08).

phosphorylation. However, due perhaps to the incomplete nature of these genomes, not all genes could be found, e.g. Thalassoarchaea cells appear to contain many of the enzymes of the Embden–Meyerhof–Parnas (EMP) pathway for the metabolism of hexose sugars, with the exception of the first (glucokinase) and the last (pyruvate kinase). For the first, several carbohydrate kinases of unknown specificity found could serve as alternatives. For the second, a pyruvate phosphate dikinase was found that could operate in the catabolic direction, as have been described in some archaea (Tjaden *et al.*, 2006). Similar proteins were also found among the fosmids from the deep Mediterranean libraries (Deschamps *et al.*, 2014) but they were absent in the MG2-GG3 or any other MGII genomic fragment. Therefore, further investigations must be performed in this direction to clarify if the EMP functions

in the gluconeogenic direction rather than the glycolytic pathway, as has been proposed for other Archaea (Hutchins *et al.*, 2001; Hallam *et al.*, 2006). Along these lines, typical gluconeogenesis enzymes such as pyruvate carboxylase (subunits A and B) and a gene coding for a phosphoenolpyruvate carboxykinase, both typical gluconeogenic enzymes, were found. As in other Euryarchaeota (Makarova *et al.*, 1999; Makarova and Koonin 2003; Hallam *et al.*, 2006) glucose 1-dehydrogenase, gluconolactonase and 2-keto-3-deoxy gluconate aldolase homologues are absent, suggesting that the Entner–Duodoroff hexose catabolic pathway is not present. In addition, we identified a complete non-oxidative pentose phosphate pathway, but the irreversible oxidative branch was missing. The reactions of the oxidative branch are important for generating NADPH, which is a source of reducing energy required by many enzymes in central biosynthetic pathways. However, we found genes for enzymes such as a 2,5-dihydroxygluconate reductase, or a malate dehydrogenase that could act as alternatives for reducing NADP⁺ to NADPH.

One of the metabolic features that we can infer is that these representatives of group IIB are facultative photoheterotrophs as seems to be the case of MG2-GG3. Three different rhodopsin genes were found among our contigs, sharing a similarity between 76 and 92% (Figure 5). Rhodopsin genes are widespread in the open oceans (Beja *et al.*, 2000) and are used as back up for heterotrophic energy generation using sunlight. These rhodopsin genes were neither adjacent to the 16S rRNA nor to the archaeal geranylgeranyl glyceryl phosphate synthase genes (Figures 4 and 5) as found previously for other MGII fosmids (Frigaard *et al.*, 2006), including those assembled previously from the Mediterranean DCM (Ghai *et al.*, 2010). Two different genomic contexts were found for the thalassoarchaeal rhodopsins, likely corresponding to different species (Figure 5). Several GOS scaffolds were found to be syntenic to these rhodopsin containing fosmids, but none larger than 5 kb. The phylogenetic tree (Supplementary Figure 5) shows the rhodopsin of the thalassoarchaeal representatives to be closely related to the one found in MG2-GG3. Both are in a cluster (clade B) separate from other rhodopsins from bacterial or eukaryotic origin (clade A). We would like to suggest the term thalassorhodopsin to designate this group. Their key residues (listed herein with EBAC31A08 numbering), indicate some important differences with other related rhodopsins (Supplementary Figure 6). Residue 105, involved in spectral tuning, was a methionine, characteristic of green absorbing rhodopsins from shallow depths (Fuhrman *et al.*, 2008) of the Bacterioidetes proteorhodopsin isoform, such as in the marine flavobacterium *Dokdonia donghaensis* MED134 (Gomez-Consarnau *et al.*, 2007; Riedel *et al.*, 2010). It is known that different amino acid substitutions in residues aspartic-97 (D) and glutamic-108 (E), which function as Schiff base proton acceptor and donor in

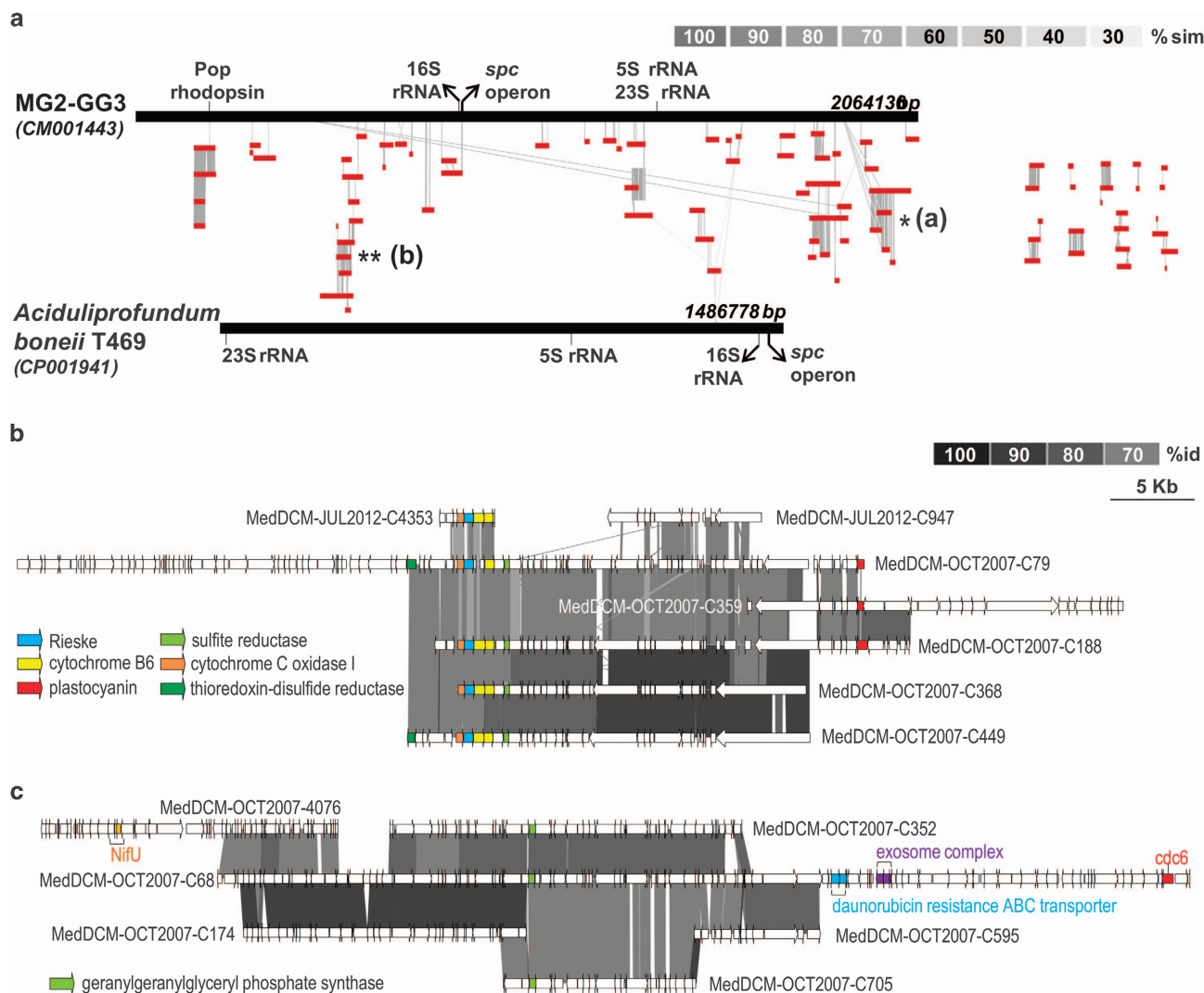


Figure 4 Genomic comparisons. **(a)** Linear representation of the synteny of thalassoarchaeal contigs with MG2-GG3-assembled genome and *A. boonei* T469. **(b and c)** Comparative genomic organization of thalassoarchaeal contigs. Conserved genomic regions between contigs are indicated by gray shaded areas, gray intensity being a function of sequence similarity by BLASTN. Specific ORFs mentioned in the text are highlighted by a graphic code.

many proteorhodopsins, provide different capacities to pump outward different ions, varying from H⁺ (Gushchin *et al.*, 2013), Na⁺ (Inoue *et al.*, 2013) and even lithium (Inoue *et al.*, 2013). These residues are partially conserved in the thalassorhodopsins and instead of the E, the basic aminoacid lysine (K) is found, a very unusual residue in proton pumping rhodopsins found previously only in *Exiguobacterium sibiricum*, a permafrost soil Gram-positive (Balashov *et al.*, 2013; Gushchin *et al.*, 2013). This suggests that thalassorhodopsins are proton pumps and add even more diversity to these ubiquitous photoreceptors.

In one of the contigs containing the rhodopsin gene we found a gene for a plastocyanin (69% similar to its homolog in MG2-GG3). A manual inspection of the predicted proteins showed their best similarities to be with halocyanins (Supplementary Figure 10). These are archaeal blue

copper redox proteins (type I) found in halophilic archaea such as *Natronomonas pharaonis* (Mattar *et al.*, 1994) that serve as mobile electron carrier at a peripheral membrane protein. The presence of the typical copper binding domain in our thalassoarchaeal subunits was confirmed. Close to the halocyanin, we found in two contigs a gene cluster coding for a Rieske/cytochrome b (Rieske/cytb) complex (Figure 4) also with their best similarities with haloarchaeal subunits. This complex was found in four more contigs (Figure 2), suggesting that this is widespread electron transport chain among these archaea. The Rieske/cytb complexes are energy-converting enzymes that operate between the initial electron donors and the terminal electron acceptors of respiratory chains. In halophiles, two distinct clusters of Rieske/cytb encoding genes have been described, (i) the Rieske/cytb complex cluster along with the NAR-encoding genes implied in the

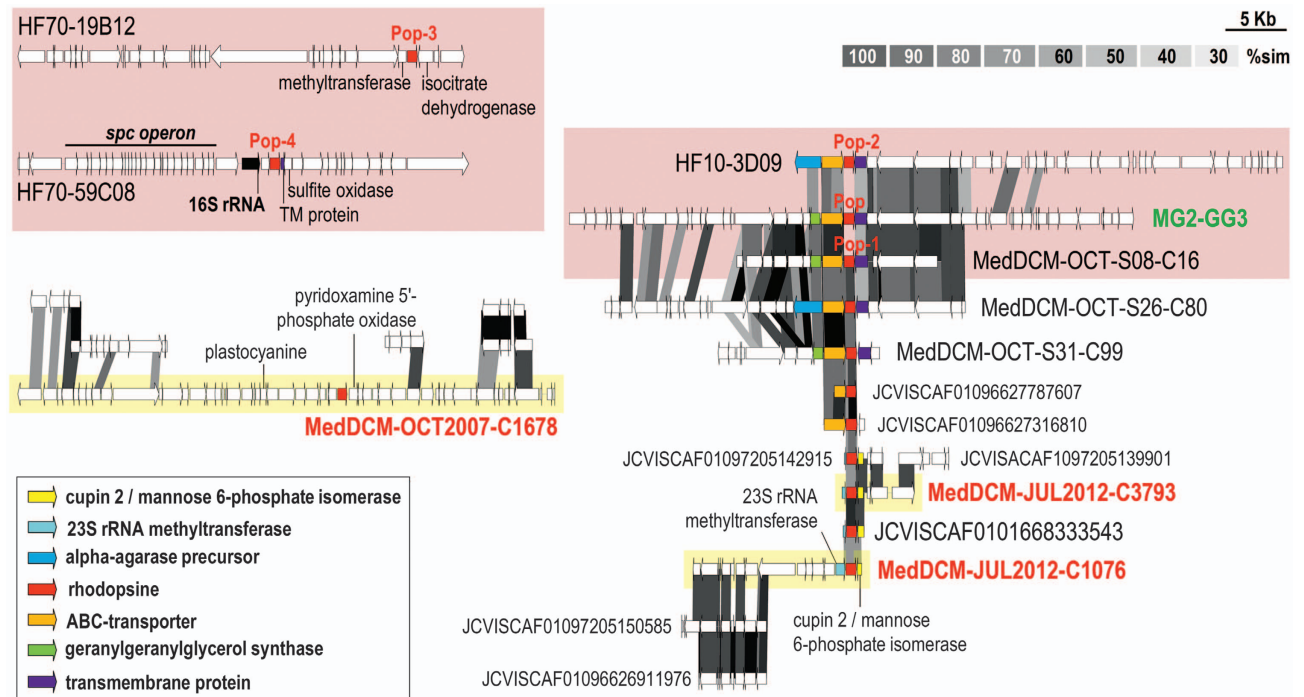


Figure 5 Comparison of rhodopsin genes. Comparative genomic organization of thallassoarchaeal contigs containing rhodopsin genes (yellow-shadowed rectangles) in context with other genomic fragments containing the MGII Pop, Pop-1, Pop-2, Pop-3 and Pop-4 rhodopsins (pink-shadowed rectangles) (Iverson *et al.*, 2012). Scaffolds from the GOS collections similar to the thallassoarchaeal sequences where also included in the comparison.

denitrifying chain (Martinez-Espinosa *et al.*, 2007) and (ii) the Rieske/cytb complex cluster with a gene coding for an halocyanin followed by a cytochrome b (Nitschke *et al.*, 2010). The implication of this second type of gene cluster in aerobic respiration seems the most likely function, where the halocyanin will work as an electron acceptor of the Rieske/cytb complex and the donor to a cytochrome oxidase (Baymann *et al.*, 2012). The MGIIB Rieske/cytb complexes found have the canonical order as in the haloarchaeal ones, but the halocyanin is found in a separate gene cluster nearby (Figure 4). Two extra genes not present in haloarchaea were conserved among the thallassoarchaeal contigs, a cysteine rich protein and a gene for a NADPH sulfite reductase. Other difference is that the haloarchaeal Rieske subunits are predicted to be of high potential, while these thallassoarchaeal homologs have the same residues than the actinobacterial medium potential subunits (Lebrun *et al.*, 2006; Baymann *et al.*, 2012). Previously, it has been suggested that Haloarchaea acquired their Rieske/cytb complexes by different lateral gene transfer events from the Actinobacteria or Thermus/Deinococcus group (Baymann *et al.*, 2012). The similarity in the thallassoarchaeal subunits to residues of the Actinobacterial ones, together with the delocalization of halocyanin also supports this hypothesis. We can speculate that the presence of the Rieske/cytb complex in MGIIB might optimize the energy yield of respiratory metabolism due to the efficient proton

pumping capacity of the complex. This could reflect the necessity to answer to an extra demand of energy in conditions of fast growth, typical of bloomers or *r*-strategists (see below).

The assembly mechanism of many archaeal surface structures is related to the one of bacterial type IV pili (Lassak *et al.*, 2012). The genome of MG2-GG3 contains these genes together with a complete cluster of flagellum biosynthesis (Iverson *et al.*, 2012). In our case, neither genes related with type IV secretion systems nor with flagellum assembly were identified. Therefore, thallassoarchaeal cells are likely to be non motile.

Five percent (137 genes) of the total number of ORFs identified in the contigs were transporters. Unfortunately, 37% of them could not be assigned a substrate. We found four contigs containing a complete set of genes for the uptake of phosphonate, indicating the capacity to utilize this refractory form of phosphorus abundant in marine habitats (Martinez *et al.*, 2010). Additionally, two clusters of genes were found coding for high affinity inorganic phosphate ABC transporters. Transporters for branched chain amino acids, peptides and oligopeptides were 10.2% of the total. Interestingly, 26.3% of the transporters were related with drug/multidrug transport systems, e.g. a daunorubicin resistance ABC transporter (Figure 4). These are ATP-dependent transport proteins which efflux a variety of compounds as a defense against toxic compounds or just transporting endogenous metabolites out of cells. Also a bleomycin

hydrolase was found. This abundance of resistance mechanisms may indicate a defensive lifestyle, typical of organisms exposed to high natural toxin concentrations, such as blooms of cyanobacteria producers of toxins.

Although no sugar transporters could be identified, a putative alpha-agarase like protein was detected (34% similarity to the experimentally tested agarase of *Thalassomonas agarivorans*) (Hatada *et al.*, 2006) (Figure 5). Other agarases were also detected near the 16S rRNA gene in other MGII fosmids already published (Figure 5). Breaking into oligosaccharides agarose or agaropectin present in the cell wall of many algae, the presence of this enzyme would make accessible an abundant food source in the ocean.

Recently, the metatranscriptomes of a serial samples from a coastal upwelling in the North California coast has been published (Ottesen *et al.*, 2013). About 5% of the transcripts of each collection were classified as MGII archaea, which synchronized their transcription with *Pelagibacter* and SAR86 marine groups. We confirmed the presence of Thalamsoarchaea in this metatranscriptomic collection finding a similar number of transcripts than for *P. ubique* HTCC7211 and SAR86 (Supplementary Figure 7). In order to find out which genes were metabolically active at the sampling time, we calculated the reads per kilobase per gigabase of the collection values for the thalamsoarchaeal ORFs (cutoff 95% identity) for each of the metatranscriptomes. Only 316 ORFs (<8% of the total) had transcripts, indicative of a low coverage. In addition

to the expected hits to ribosomal proteins, elongation factors and translation initiation factors (that is, SUI1), which were abundant across all the data sets, we also found a putative collagen-like cell surface protein in all the collections. It has been described that in blooming bacterioplankton, genes predicted to increase cell surface adhesiveness are highly transcribed, having an important role in microbial aggregation (Rinta-Kanto *et al.*, 2012). Along these lines, we found a putative thalamsoarchaeal flotillin being highly expressed (61% similar to its homolog in MG2-GG3). Flotillins are proteins that form membrane microdomains implicated in signal transduction, vesicle trafficking and cytoskeleton rearrangements (Bach and Bramkamp, 2013). They trigger changes in cell surface properties in response to environmental signals inducing the formation of polysaccharides (D'Argenio and Miller, 2004). Therefore, the high frequency of this flotillin-like protein suggests that aggregation was happening at the time of sampling. Other highly expressed transcripts were rhodopsins, cold shock proteins, amino acids ABC transporters and the previously mentioned bleomycin hydrolase.

Presence of thalamsoarchaeal reads in metagenomic collections

In the MedDCM-OCT2007 454-metagenome (Ghai *et al.*, 2010), a total of 4% of the 16S rRNA sequences were attributed to Euryarchaeota. However, when recruitment assays were performed using

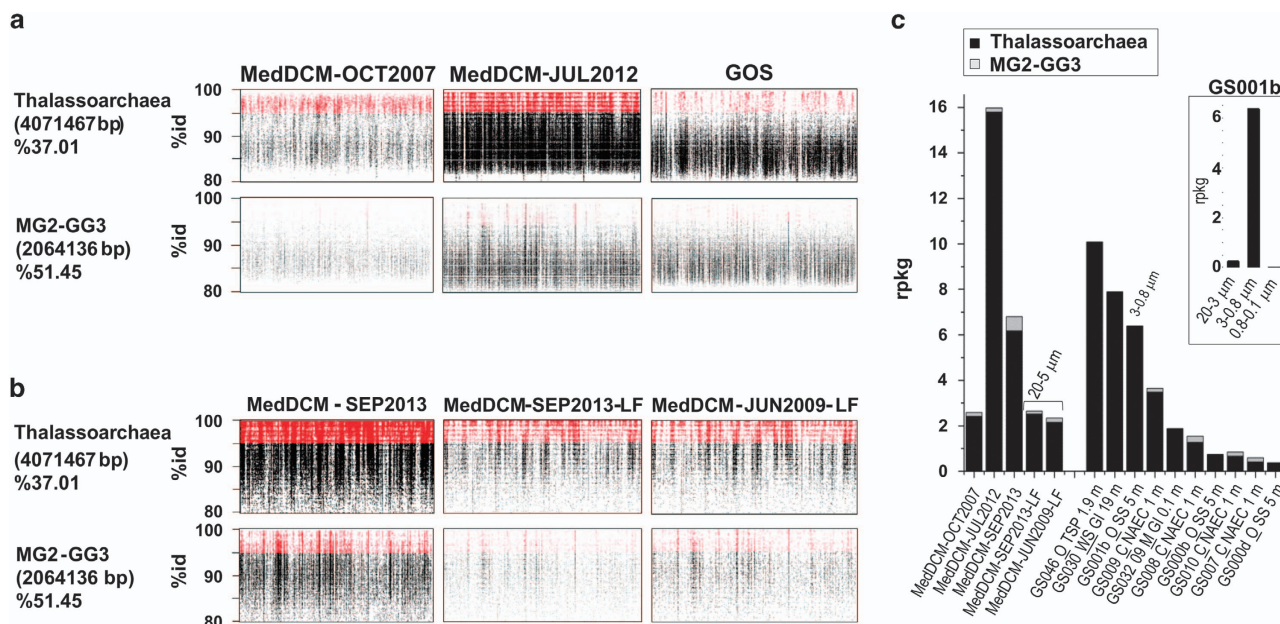


Figure 6 Metagenome recruitments. (a) Metagenomic reads recruited (BLASTN) by the thalamsoarchaeal contigs and MG2-GG3 genome in metagenomes from the Mediterranean deep chlorophyll maximum MedDCM-OCT2007, MedDCM-JUL2012 and from the GOS. (b) Metagenomic reads recruited (BLASTN) by the thalamsoarchaeal contigs and MG2-GG3 genome in metagenomes from the Mediterranean DCM recovered from different pore-size filters, small fraction (5–0.22 μm) MedDCM-SEP2013 and LF (20–5 μm) MedDCM-JUN2009-LF and MedDCM-SEP2013-LF. (c) Recruitment comparison of the Mediterranean data sets and GOS collections. Metagenomes obtained with large-fraction filter size are indicated.

a virtual genome reconstructed with the thalassoarchaeal contigs, only 0.5% of the reads in MedDCM-OCT2007 and MedDCM-JUL2012 metagenomes were recruited over 95% of identity (species cutoff) (Figure 6). However, the MG2-GG3 genome recruits one order of magnitude less (0.019% and 0.002%, respectively) and no hits were detected for SAG SCGC-AAA-288-C18. We examined the geographic distribution of the thalassoarchaeal genomes, analyzing their presence in the GOS database (Venter *et al.*, 2004; Rusch *et al.*, 2007) that contains mainly surface samples. In general, the collections with more hits came from the tropics (for example, tropical Pacific, Galapagos Island and Sargasso Sea), with high temperature (22.9–26.9 °C range in the metadata provided). However, they were also detected in GS009, Block Island, and in GS008, Newport Harbor, where temperatures decrease to 11–9.4 °C (Figure 6). They were very abundant at GS046, 300 miles from Polynesia, but not in other collections recovered in consecutive days at similar depths in the same transect (every 100/200 miles). In summary, the abundance of this group appears to be very uneven including in warm waters. One possible explanation is the heterogeneity of surface waters, which are submitted to a strong hydrodynamism. Microbial communities coming from deeper waters may upwell eventually. A different but non-exclusive explanation is that the thalassoarchaeal cells produce localized blooms. In this sense, one of the collections where they were better represented was a warm seep from the Galapagos Island (GS030; 19 m, 26.9 °C). Such a high temperature could promote the growth of bloomers.

Analyzing the few available metagenomic depth profile collections (subtropical gyres of North Atlantic (Bermuda Atlantic Time Series) and north Pacific (Hawaii Ocean Time-Series, Station ALOHA); (Coleman and Chisholm 2010; DeLong *et al.*, 2006)), we found evidence of the preference of thalassoarchaeal cells for the photic zone (above 100 m) (Supplementary Figure 8). However, their abundance was one or two orders of magnitude lower than in the Mediterranean DCM (Supplementary Figure 8). Thalassoarchaeal sequences were also scarce along the colder waters of the coast of the northeast subarctic Pacific NESAP collections (Allers *et al.*, 2013) and also in the metagenomes from the eastern tropical south Pacific (Stewart *et al.*, 2012), where reads matching thalassoarchaeal sequences peaked in abundance in the small-size fraction (1.6–0.22 µm) upper water column samples (70 m). These authors reported that 50% of the sequences matching Euryarchaeota peaked in abundance at this depth on prefilters (3–1.6 µm). Of these sequences, those matching MGII constituted 40% of the total, indicating that still an important fraction of MGII remains in the smaller fraction.

A high prevalence of group IIA in particle-rich coastal waters has been described (Galand *et al.*, 2010). In this respect, the presence of genes for the degradation of proteins and lipids suggested that the MG2-GG3-assembled genome may have a particle-

attached lifestyle (Iverson *et al.*, 2012), although no direct evidence was provided. To clarify this issue, three more metagenomes were sequenced from Mediterranean DCM samples recovered in 2009 and 2013 (same location the DCM-2007 and 2012) but from the large-size filter. MedDCM-JUN2009-LF and MedDCM-SEP2013-LF were obtained from biomass recovered from the 20 to 5 µm fraction. MedDCM-SEP2013 was obtained from the smaller 5- to 0.22-µm fraction. When the corresponding recruitments were done using the thalassoarchaeal genomic fragments, homologous sequences were preferentially found (three to eight times more) in the small-fraction data sets than in the large ones (Figure 6), suggesting that these microbes are free living. The MG2-GG3-assembled genome was not significantly present in either fraction. However, for the sample from the Sargasso Sea GS001, most of the homologous sequences to Thalassoarchaea were found in the 3- to 0.8-µm filter (GS001b) instead of the smaller fraction (inset in Figure 6). Recently, some data sets have been published that come from river estuaries or plumes (similar to the environment from which MG2-GG3 metagenome was obtained) (Smith *et al.*, 2013; Satinsky *et al.*, 2014). These metagenomes have large numbers of reads homologous to both MG2-GG3 and Thalassoarchaea. In the Columbia River plume (very close to the site of origin of MG2-GG3), this assembled genome was found almost equally in the small (0.8–0.1 µm) than in the larger fraction (3–0.8 µm) (Supplementary Figure 9), although Thalassoarchaea recruited scarcely. In the metagenomes of the Amazon plume, both microbes were found in one LF collection (156–2 µm) (Supplementary Figure 8). Interestingly, Thalassoarchaea did not recruit from any of the other collections from the same study, including the smaller size fraction retrieved at the same location. In this case, dramatic differences were found even in collections taken only 1 h apart, that is, ACM25 and ACM1. One plausible explanation is the blooming of both Euryarchaea in the sample.

Several meso and bathypelagic metagenome collections were also screened with negative results (see Materials and Methods). They were also absent in marine collections from polar regions (Alonso-Saez *et al.*, 2012) and from the Baltic collections (Larsson *et al.*, 2014). The Red Sea DCM metagenome (365 Mb) (Thompson *et al.*, 2013) was also analyzed, and although the sample was described as recovered at the DCM (50 m), no significant recruitment was found.

Conclusions

Owing to cultivation bias, Archaea were considered extremophilic or anaerobic microbes for many years. Still, the number of isolates of non-extremophilic archaea is very small. The reasons for this reluctance to grow in pure culture are still obscure but, in any case, our knowledge remains restricted to culture-

independent approaches such as metagenomic assemblies. In fact, the number of marine planktonic microbes brought into culture is small for any group and the difficulties of cultivating members of the Pelagibacterales illustrate this point. Here, by sequencing metagenomic fosmids and assembling high-coverage metagenomes, we are providing information about a group of Euryarchaeota hitherto known only from 16S rRNA sequences. Using specific FISH probes, we could visualize the cells and assess their abundance at *ca.* 2% of the DAPI-stained cells. The genomic information indicates that (at least some representatives of this group) are planktonic photoheterotrophic microbes similar to other known components of the open ocean planktonic microbiota such as the Pelagibacterales or the Actinomarinales. They are all non-motile small cells, with streamlined genomes, and with diverse heterotrophic capabilities, as should be expected from the diverse composition of dissolved organic matter in the open ocean. However, Thalassoarchaea appear very unevenly in metagenomic data sets. This is true also of the available genome of MGIIA, MG2-GG3, which seems to be more abundant in estuarine systems or river plumes. Thalassoarchaea are present in all the available Mediterranean DCM samples and other temperate and tropical seas. They seem to be also detectable at the DCMs at Bermuda Atlantic Time Series (Sargasso) and Hawaii Ocean Time-Series (Central Pacific), and therefore the DCM seems to be its main habitat. In addition, they appear at the surface during sporadic blooms. For example, MGII B 16S rRNA representatives have been found before at the surface during winter mixing, and many of them cluster together with Thalassoarchaea. Their preference for the DCM can explain this apparently paradoxical finding. They could bloom after hydrodynamic mixing with deeper nutrient-rich waters as happens in temperate latitudes during winter, or more sporadic phenomena due to currents, river plume fertilization and so on. They might use light as an additional energy source and start growing as soon as nutrient concentrations increase after water column mixing.

This bloomer strategy is more often found in particle-associated microbes. However, the data presented seem inconclusive regarding the known representatives of MGIIA or B. Although MG2-GG3 genome indicated a particle-attached lifestyle, it recruits in different size fractions depending on the sample and mostly appears in the small filter sizes, like the Thalassoarchaea. This might be an artifact of the filtration method (larger size filters collect smaller microbes as they become clogged) but it is also possible that MGII representatives are present in both habitats. These hypotheses remain open to be tested in the future with metagenomes obtained with different filter sizes or alternative methodologies. However, it is clear that Thalassoarchaea are an abundant component of microbial plankton in the Mediterranean DCM and also in other oceans.

Further work of genome reconstruction coupled to single-cell genomics or development of cultivation methods to obtain laboratory cultures will allow a better understanding of this widespread and important marine microbial group.

Conflict of Interest

The authors declare no conflict of interest.

Acknowledgements

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