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Published in:
Environmental Microbiology

Document Version:
Peer reviewed version

Queen's University Belfast - Research Portal:
Link to publication record in Queen's University Belfast Research Portal

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Microbial community of the deep-sea brine Lake Kryos seawater-brine interface is active below the chaotropicity limit of life as revealed by recovery of mRNA.

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Running title: Prokaryotes within the interface of deep-sea Lake Kryos
Key words: Messinian evaporites, Deep-sea anoxic brine lakes, Halophiles, Limits of life, Chaotropicity, Habitability, Mars
Summary

Within the complex of deep, hypersaline anoxic lakes (DHALs) of the Mediterranean Ridge we identified a new, unexplored DHAL and named it "Lake Kryos" after a nearby depression. This lake is filled with MgCl$_2$-rich, athalassohaline brine (salinity >470 practical salinity units), presumably formed by the dissolution of Messinian bischofite. Compared to the DHAL Discovery, it contains elevated concentrations of kosmotropic sodium and sulfate ions, which are capable of reducing the net chaotropicity of MgCl$_2$-rich solutions. The brine of Lake Kryos may therefore be biologically permissive at MgCl$_2$ concentrations previously considered incompatible with life. We characterized the microbiology of the seawater-Kryos brine interface and managed to recover mRNA from the 2.27-3.03 M MgCl$_2$ layer (equivalent to 0.747-0.631 water-activity) thereby expanding the established chaotropicity window-for-life. The primary bacterial taxa present there were KB1 candidate division and DHAL-specific group of organisms, distantly related to Desulfohalobium. Two euryarchaeal candidate divisions MSBL1 and HC1, detected in minority in the overlaying layers, accounted for more than 85% of the rRNA-containing archaeal clones analyzed in 2.27-3.03 M MgCl$_2$ layer. These findings shed light on the plausibility of life in highly chaotropic environments, geochemical windows for microbial extremophiles, and have implications for habitability elsewhere in the Solar System.
Introduction

In the eastern part of Mediterranean seafloor, an accretionary complex, named the Mediterranean Ridge, is formed by subduction of the African plate under the Eurasian and Anatolian plates. During the Messinian salinity crisis (late Miocene epoch, 5.33 - 5.96 million years ago) the repeated desiccations and re-fillings of the Mediterranean Sea resulted in the formation of enormous deposits of layered evaporites, that attain the thickness of up to 3.5 km in some places of eastern Mediterranean (Cita, 2006). In contrast to other tectonically active ridges, the deformational activity of the Mediterranean Ridge accompanied with presence of huge subsurface salt deposits appears to control the creation of peculiar submarine hydrological formations within confined depressions.

The peculiar hydrology and chemistry of such lakes, which are named deep-sea hypersaline anoxic lakes (DHALs), discourages mixing of their brines with the overlying seawater (Raup, 1970). Seven such lakes, L’Atalante, Bannock, Discovery, Medee, Thetis, Tyro and Urania have been discovered and studied in the deep eastern Mediterranean over the last 20 years (De Lange and Ten Haven, 1983; MEDRIF Consortium, 1995; Wallmann et al., 1997; Chamot-Rooke et al., 2005; La Cono et al., 2011; Yakimov et al., 2013). The surfaces of these brine lakes lie between 3.0 and 3.5 km below sea level and the salinity of their brines ranges from five to 13 times higher than that of seawater. Although these DHALs lie geographically close to each other (Fig. 1a), their hydrochemical diversity suggests that the processes leading to their formation were qualitatively different. As is generally accepted, during the desiccation/re-flooding cycles the salt deposition implied the simultaneous existence of early- and late-stage primary brines and evaporites. Seawater can be evaporated 10-fold without salt precipitation, resulting in formation of brine with salinity $\leq$ 330 PSU. This brine is named as “thalassohaline early-stage primary brine” (ESPB) and has proportions of all major ions characteristic to that of seawater. When the evaporation of seawater continues, salinity increases and the salts begin to precipitate changing the proportion of dissolved ions, thus forming the “athalassohaline late-stage primary brine” (LSPB). The insoluble calcium minerals precipitated first,
followed by precipitation of halite (NaCl), kieserite (MgSO₄·KCl·3H₂O), carnallite (KMgCl₃·H₂O), kainite (MgSO₄·KCl·3H₂O) and ending with formation of bischoffite (MgCl₂·6H₂O), which is the most soluble of all marine evaporite salts (Wallmann et al., 1997; Cita, 2006). Due to favorable climatic and geological conditions, both brines and solid stratified evaporite suites were stored in the subsurface for millions of years until tectonic activities would squeeze them on the seabed. For some Mediterranean DHALs, their idiosyncratic geomorphology implies the formation mechanisms other than simple outcropping of the Messinian evaporites followed by accumulation of high-density brines in the nearby depression. As has been proposed elsewhere, the evaporite dissolution could occur in sub-bottom deposits without direct exposure on the seafloor (Camerlenghi, 1990; Camerlenghi and McCoy, 1990; Cita, 1991, 2006). Tectonic activity in the Mediterranean Ridge leads to tensional stress and formation of seabed fractures and through these seawater can penetrate into deeper sediment layers ultimately reaching the subsurface layer of the Messinian evaporites. Osmotic pressure encourages movement of seawater towards the solid evaporites, dissolving the most soluble salts and increasing the volume of internal brine lenses. Notably, this movement of seawater is almost unidirectional, because the argillaceous Plio-Quaternary superficial sediments overlying the Messinian evaporites are “salt-rejecting”, effectively behaving as a semipermeable membrane (Cita, 1991; 2006). Such continuing enrichment by evaporite dissolution leads to interstitial hydrologic formations, which in turn causes the overlying sediments to collapse and form a brine lake with characteristic confined, negative topography enriched by simple or complex morphologies ranging from sub-circular to elliptical, arc- or U-shaped basins, frequently including mounds and small, deeper depressions (Camerlenghi and McCoy, 1990).

Among all Mediterranean DHALs explored so far, only the Discovery Lake is filled with near-saturated MgCl₂-brine (5.05 M), suggesting that it derived via dissolution of bischoffite, which is located within the uppermost layer of the evaporitic suite as explained above. Hence, the Discovery Lake is one of the saltiest athalassohaline water bodies on Earth (Table 1 and 2; Wallmann et al., 1997; Cita, 2006).
Due to the exceptionally high concentration of the divalent salt MgCl₂, this lake is approaching an anhydrous condition and is, simultaneously, an exceptionally chaotropic system with the lowest water activity (A_w) value registered for any hydrological formation on our planet (Marion et al., 2003; Hallsworth et al., 2007). In our previous study we demonstrated that exceptional chaotropicity of MgCl₂, rather than water activity reduction, is the window-of-life-determining parameter (Hallsworth et al., 2007). We suggested that in the absence of compensating (e.g. kosmotropic) ions, such as sodium and sulfates, the upper concentration of MgCl₂ permissible for life, is about 2.3 M. This finding is consistent with the apparent MgCl₂ limit for microbial activity in the Dead Sea (Oren, 1999; 2010). As observed by Harrison et al. (2013), there have been relatively few studies on the way in which multiple stress parameters interact to determine the habitability of specific environments. A number of studies have, however, explored the way in which factors such as water activity, chaotropicity, nutrient availability and temperature can interact to determine biological permissivity of high-solute environments (Daffonchio et al., 2006; Williams and Hallsworth, 2009; Chin et al., 2010; Cray et al., 2013a; 2013b; Lievens et al., 2014).

Here, we present the results of the first oceanographic, geochemical and microbiological explorations of Lake Kryos, a second Mediterranean athalassohaline DHAL filled with nearly saturated MgCl₂-brine. Aside from slightly elevated concentrations of Na⁺ and SO₄²⁻, the hydrochemistry of this novel lake was found to share commonalities with that of the Lake Discovery (Table 1). As revealed by a comprehensive analysis of the vertical distribution of major prokaryotic groups along the seawater-brine interface, the Kryos prokaryotic community forms sharply stratified and dense ecosystem, operating at the very edge of Earth’s biosphere. In order to decipher the stratification of principal metabolic pathways within this environment and, considering that DNA may be effectively conserved under highly chaotropic conditions, comparative analysis of recovered rRNA and mRNA transcripts were performed for three layers of the interface.
Results and discussion

Geomorphological and geochemical characterization of Lake Kryos.

During the cruise MIDDLE08 on RV Urania in September 2008, while on transit from the Anoxic Lakes Region West of Crete to the Lake Medee, we surveyed by 3.5 Chirp kHz swath-bathymetry profiling (SBP) confined depressions deeper than or similar to known seawater-brine lake interfaces (Fig. 1a). The candidate targets were localized by the morpho-bathymetric analysis of MEDIMAP data with resolution of 500 m (Loubrieu et al., 2008). Given that the strong density contrast at the seawater-brine lake interface would have produced a straight line on acoustic swath bathymetrical profiling (SBP) data, we expected to be able to identify some yet unexplored DHALs. Approximately 20 nautical miles from the Urania Lake, we moved over a narrow North-South, elongated fracture (22°01'E 35°02'N – 22°02'E 34°53'N) and a sharp crisp line, hinting at the existence of a brine lake, was detected with maximum depth of about 3500 m. This was confirmed by direct conductivity-temperature-dissolved oxygen (CTD) profiling, brine sampling and bottom coring. Using the SBP data of the RV Urania DEEPRESSURE cruise in 2013 and correcting the depths in the brines with the pressure data of the CTD casts, we obtained a map of the Kryos Lake with 20 to 25 m resolution. The Lake Kryos (named after the neighboring oxic depression) has the seawater-brine interface at 3387 dB (3337 m) and fills a steep, narrow basin approximately 18 km long and 1.7 km wide, oriented N-S and bending N-N-E at its northern tip with two arms oriented E-N-E (Fig. 1b). The bottom of the basin is 300-400 m below the depth of the surrounding region and has a well defined, continuous and very steep slope to the west, while in the opposite direction the seabed rises more moderately. The southern part of Kryos basin is characterized by several N-S oriented mounds and depressions,
presumably indicating the existence of isolated brine pools. Similar small pools may be detected at the northernmost part of the lake. Lake Kryos, including these small polar pools, has an area of about 100 km$^2$ and a volume of about 10 km$^3$. The central area of the lake is slightly deeper than 3500 m below sea level, implying that the depth of brine within the lake is approximately 160-170 m. The temperature measured at the seawater : brine interface was 13.98°C and slightly increased to 14.66°C within the brine, close to the seabed.

Chemical characterization of the Kryos brine revealed its extremely high salinity (471 g [kg H$_2$O]$^{-1}$) mainly due to extreme, close to saturation, concentration of Mg$^{2+}$ (4.38 M) and Cl$^-$ (9.04 M). As shown in the Table 1, the Kryos hydrochemistry is quiet similar to that of the Discovery brine with the exception of elevated concentrations of Na$^+$ and SO$_4^{2-}$, which are present in the former. The Kryos basin is filled with almost 10 km$^3$ of MgCl$_2$-rich brine which compares to Lake Discovery volume of nearly 0.2 km$^3$ (Wallmann et al., 1997, 2002); the DHAL Kryos is thus the largest deep-sea athalassohaline formation on Earth. Moreover, Lake Discovery, the CaCl$_2$-saturated Don Juan Pond and Lake Kryos together form a triad of the saltiest aquatic systems on our planet (Table 2). Previously made equilibrium calculations with the PHRQPITZ model (Wallmann et al., 1997) have indicated that a LSPB similar in composition to those of the MgCl$_2$-rich athalassohaline brines may be produced when seawater is evaporated to the point of bischoffite precipitation, i.e. until only 5 g of initial 1000 g of H$_2$O remained in solution. A similar composition, termed as a secondary brine (SB), may also be formed when seawater is equilibrated with solid bischoffite and kainite (K$_4$Mg$_4$Cl$_4$(SO$_4$)$_4$11H$_2$O) (Table 2). Therefore, the major ion composition of both brine lakes is consistent with either a primary (evaporated seawater) or secondary origin (dissolution of the most soluble marine evaporite salts). As it generally accepted, concentrations of lithium could be used to differentiate between the primary and secondary brines, because this cation is conserved during seawater evaporation path and does not co-precipitate with evaporites in the presence of high Mg$^{2+}$ concentrations (Carpenter, 1978; McCaffrey et al., 1987; De Lange et al., 1990; Wallmann et al., 1997, 1997)
By comparison with the LSPB values, lithium concentration in the *Discovery* and *Kryos* brines was 20-25 times less, indicating that both lakes have evidenced an extreme evaporation of the eastern Mediterranean, which is likely to have taken place during the late Messinian. As was proposed for Lake *Discovery*, the upmost layer of evaporite suite, represented by precipitated and lithium depleted bischoffite, was subsequently re-dissolved and has migrated to form a deep-sea brine pool. As it was shown by analysis of $^3$He concentrations, before it entered the *Discovery* basin, the re-dissolved MgCl$_2$-saturated brine was initially stored for unknown period of time inside the sediments as interstitial brine pool (Wallmann *et al.*, 1997, 2002). We hypothesized that this scenario of the origin is equally applicable to the *Kryos* Lake.

The *Kryos* and *Discovery* Lakes are the most chaotropic large-scale aquatic systems on Earth

Earlier we have measured the water activities ($A_w$) in various MgCl$_2$–dominated solutions and evidenced that this salt is one of the most powerful $A_w$-reducing agents known (Hallsworth *et al.*, 2007); see also Winston and Bates (1960). Due to its high solubility and divalency, MgCl$_2$ is able to depress the water-activity values much below the limit observed for cell division or metabolic activity (Fig. 2a; Pitt, 1975; Marion *et al.*, 2003; Grant, 2004; Williams and Hallsworth, 2009). The water activity of saturated MgCl$_2$ is 0.340 in the range 10 - 15°C (Fig. 2a; Winston and Bates, 1960); we empirically determined the value for the *Discovery* brine; which was 0.382 $A_w$ at 14.5°C (Hallsworth *et al.*, 2007). However, the established water-activity limit for (active) life (0.605$^2$; Pitt and Christian, 1968) is equivalent to 3.7 M MgCl$_2$ (Hallsworth *et al.*, 2003a, 2007). The brine of *Kryos* contains a considerably higher concentration of MgCl$_2$ (4.38 M), which corresponds to 0.399 $A_w$. Another

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1. Whereas there have been a number of unsubstantiated claims of germination and growth of *Streptomyces* and *Micromonospora* strains at 0.500 $A_w$ from one research group (Doroshenko *et al.*, 2005; 2006; Zvyagintsev *et al.*, 2007; 2009; 2012; Kurapova *et al.*, 2012), the limit for such Actinobacteria has recently been determined empirically at 0.890 $A_w$ with a theoretical lower limit which was derived by construction of isopleths growth profiles of approximately 0.870 $A_w$ (Stevenson and Hallsworth, 2014).
harmful feature of MgCl$_2$-rich solutions, incompatible with existence of actively metabolizing organisms, is their exceptional chaotropicity (Hallsworth et al., 2007; Cray et al., 2013a), and it is this property, rather than any other activity of the solute, which can limit the microbial biosphere in high-MgCl$_2$ (and presumably other highly chaotropic) environments (Hallsworth et al., 2007). Supporting this, our previous study on microbial communities of the Discovery Lake and recovery of unstable biomarkers, such as messenger RNA, suggested that in almost pure solutions of MgCl$_2$ representing the Discovery brine, the active life, as we currently know it, is not likely at MgCl$_2$ concentrations of > 2.3 M (Hallsworth et al., 2007), which corresponds to < 0.790 A$_w$ (Fig. 2a). We are aware that the equation between these specific chaotropicity and water-activity values might be true only for the Discovery brine, almost depleted by sodium and sulfates. However, various sources of evidence suggest that this limit can also be expected for other habitats because chaotropes can to some extent be compensated by kosmotropes (Oren, 1983; Hallsworth et al., 2003b, 2007; Williams and Hallsworth, 2009; Bhaganna et al., 2010; Bell et al., 2013), so the presence of other anions, like sodium and sulfate, can reduce the net chaotropicity of a hypersaline environment and widen the chaotropicity windows of life. Compared with the Lake Discovery, the Lake Kryos brine is slightly impoverished with MgCl$_2$ and simultaneously enriched with kosmotropic sodium and sulfate ions (Table 1), thus representing a unique opportunity to explore and test this assumption.

As shown in Figure 3, there is a sharp, ~2.5 m halocline at the seawater : brine lake interface characterized by a steep Mg$^{2+}$ gradient ranging in concentration from 55 mM at its upper layer to 4.38 M in proximity to brine. Using our previous approach for A$_w$ measurements of MgCl$_2$ solutions applied to Discovery samples (Hallsworth et al., 2003b; 2007), we measured the A$_w$ and chaotropicity levels of the Kryos interface (Fig. 2). The current window for chaotropicity equivalent (A$_w$ = 0.790), established for the interface of Discovery, and the current window for xerophilic cellular life (A$_w$ = 0.605) embrace only upper two-thirds of the Kryos interface. An A$_w$ value of 0.399 was determined for the Kryos brine itself and it is far below a minimal level of water activity, essential
for cellular function. The Kryos brine is thus an exceptionally chaotropic and low-water-activity environment, possibly the most large-scale, MgCl$_2$-saturated, aquatic system on Earth.

One year after the discovery of Lake Kryos, we begun exploring the extent to which cellular systems have been able to adapt to the Kryos interface conditions. Following this aim, the Kryos interface was sampled and fractionated using our previously established methodology to sample the DHAL interfaces (Daffonchio et al., 2006; Borin et al., 2009; Hallsworth et al., 2007; Yakimov et al., 2007a, 2007b, 2013; La Cono et al. 2011). Immediately after the rosette recovery, initial measurements of salinities of the bottommost content of Niskin bottles were performed. Obtained values were plotted over the reconstructed salinity profile (Fig. S1). We were aware that accurate capturing from elevated depths of in situ patterns of extremely unstable mRNA is potentially biased by changes in environmental conditions during the sample recovery (Feike et al., 2012). To diminish this concern, all interface layers carrying similar biases, were sampled during the same cast and were processed in parallel. Due to the favorable weather conditions, little or no mixing had occurred during the sampling. Five Niskin bottles, exhibiting equivalent salinities at their bottoms, were used for further biological analyses (Fig. S1). Their contents were carefully fractionated anaerobically by slowly recovering 0.5-litre, 1-litre or 2-litre fractions from bottom tap. The subsamples collected from the bottommost part of these Niskin bottles (range of MgCl$_2$ 2.27 - 3.03 M) were pooled and hereafter termed as the AWW layer. As shown in Fig. 2a, the calculated A$_w$ values for this layer (from 0.747 to 0.631) extend beyond both the established chaotropicity limit fro life$^3$ and close to the established water-activity limit for cell division of prokaryotes (for references, see Grant, 2004; Stevenson et al., 2014). As anticipated, the presence of kosmotropic substances in the Kryos brine, such as sulfates, has a mitigating effect on the chaotropicity of MgCl$_2$ (Fig. 2b). For example, at ~0.760 A$_w$ agar-gel point temperatures for both the Kryos brine and synthetic Kryos brine were ~6°C

$^3$Based on the chaotropicity - A$_w$ equivalence for the closest comparator brine, that of the Discovery lake (Hallsworth et al., 2007).
higher than that of a MgCl$_2$ solution (Fig. 2b). This temperature difference is equivalent to a kosmotropicity of 25 kJ kg$^{-1}$ (Hallsworth et al., 2013a; Cray et al., 2013a); the kosmotropic activity that is exerted by 2.3 M NaCl (Hallsworth et al., 2007). The overlaying layer, hereafter termed CHW, had the range of MgCl$_2$ concentrations 1.30 – 2.27 M, corresponding to the established chaotropicity boundary (CHW) (Fig. S1), so the Kryos brine potentially represents habitable high-MgCl$_2$ environment thus far identified; equivalent to a chaotropicity of between 143 and 296 kJ kg$^{-1}$ (Fig. S2). The upper interface (UIF) layer with salinity of 50-140 PSU was additionally analyzed to affirm the occurrence of stratified and metabolically active microbial populations thriving in deeper AWW and CHW compartments.

**Characterization of dissolved organic matter in the Kryos brine**

After dissolved organic matter (DOM) isolation by means of solid phase extraction, the desalted eluate was analyzed with ultrahigh resolution mass spectrometry (ion cyclotron resonance Fourier transform mass spectrometry, ICR-FT/MS) enabling a direct depiction of the DOM compositional space with a few thousands of assigned elemental formulas of this complex organic mixture. The mass spectra show a near Gaussian signal distribution typical of natural organic matter, with recognizable main mass spacings of methylene ($\Delta m = 14.056$ amu), double bond equivalents ($\Delta m = 2.0157$ amu) and a splitting of $\Delta m = 0.0024$ amu, denoting closely spaced CHO and CHOS compounds (Schmitt-Kopplin et al., 2010a) indicative of a highly processed organic matter with appreciable extent of sulfurization at a relatively small overall molecular weight (<500 amu, Fig. 4a). Conversion of the signals into elemental compositions revealed a high abundance of sulfur compounds (700 CHOS and 250 CHNOS were assigned molecular formulas) reflecting the remarkable diverse sulfur chemistry in these particular extreme sulfide rich conditions.
environments (Fig. 4b,c). Neither organochlorines nor organomagnesium compounds were indicated by these datasets. The ratios of CHOS/CHO and CHNOS/CHNO molecular compositions in DOM were different, reflecting divergent mechanisms of sulfurization resulting in CHOS and CHNOS molecules. In contrast, purely abiotic reactivity of reactive sulfur species of presumably mineral origin with CHO and CHNO compounds led to comparable ratios (Schmitt-Kopplin et al., 2010b). Hence, a biotic origin of the sulfur compounds observed in the Kryos brine seems highly likely. The van Krevelen diagrams show compounds with rather pronounced aliphaticity (elevated H/C ratio) and especially remarkable extent of oxygenation (O/C ratio > 0.6), extending even further than the previously described carboxyl-rich alicyclic materials (CRAM) (Fig. 4d). Further research is needed to elucidate the structural diversity of these compounds resembling the condensed alicyclic structures of biogenic origin such as sterols and hopanoids, which offer nominal unsaturation without overly abundance of sp² carbon (Hertkorn et al., 2006).

Prokaryotic abundance and community composition of the Kryos interface using CARD-FISH

At the depth of ~3338 m, where the UIF Kryos sample was taken, total prokaryote number (DAPI-stained cells) increased six-fold compared to DAPI values for overlaying deep-sea water (Table 3). The number of cells in the CHW interface layer increased to $5.57 \pm 0.45 \times 10^5$ cell ml⁻¹ and then gradually decreased to $2.47 \pm 0.11 \times 10^5$ cell ml⁻¹ in the AWW layer, which was below the CHW. CARD-FISH indicated that while almost all DAPI-stained cells from the overlaying seawater contained 16S rRNA (89%), the numbers of CARD-positive microorganisms in the UIF interface layer dropped almost to a half of those visualized by DAPI (53 %). The gradual increase of CARD-positive fraction from 68 to 81% of all DAPI-stained cells was observed in deeper layers (Table 3). This phenomenon of increase in cell density likely reflects trapping and effective conservation under highly chaotropic conditions of stable biological macromolecules, such as DNA and rRNA, albeit in an inactivated form.
(Duda et al., 2004; Hallsworth et al., 2007; Cray et al., 2013a). Nevertheless, the existence of as-yet-
undiscovered life forms, that have evolved greater chaotropicity and water activity tolerances than
presently known, cannot be ruled out (Hallsworth et al., 2007).

As revealed by taxon-specific CARD-FISH analysis (Table 3 and Table S1), bacteria dominated
all studied layers of the Kryos interface. Previously we have shown that bacterial community
thriving in the low interface of Lake Discovery was characterized by overwhelming dominance of
members of KB1 candidate division and organisms, distantly related to Desulfohalobium (Hallsworth
et al., 2007). Application of KB1-specific FISH-probes (Yakimov et al., 2013) revealed that, being
absent in UIF community these extremely halophilic prokaryotes are gradually dominating the CHW
and AWW populations. Distribution of DHAL-specific deltaproteobacteria was found be more
homogeneous in both saltiest layers. Thaumarchaeota and Euryarchaeota exhibited opposing
distribution patterns in relation to depth. The absolute dominance of Euryarchaeota in hypersaline
and anoxic habitats is a characteristic feature for all currently studied DHALs interfaces (Daffonchio
et al., 2006; Borin et al., 2009; Hallsworth et al., 2007; Yakimov et al., 2007a, 2007b, 2013; La Cono et
al., 2011). Noteworthy, below the established $A_w$-limit of life (0.605) the CARD-FISH analysis with the
universal archaeal probe ARCH915 detected more than 40000 ribosome-containing cells ml$^{-1}$,
whereas none of them were visualized there by more specific EURY806 probe (Table 3). This
observation can be explained by the presence in the AWW samples of organisms, such as the
members of MSBL1 candidate division, whose 16S rRNA sequences are out of the EURY806 probe
specificity range.

Stratification of the Kryos interface prokaryotic 16S rRNAs across the chaotropicity limit of life

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To survey the distribution of ribosome-containing Bacteria and Archaea, total RNA (50.42, 35.37 and 32.18 ng µl⁻¹) was respectively extracted from UIF, CHW and AWW samples. Total cDNA was further obtained by reverse transcription with hexa-random primers, PCR amplified with 16S rDNA-specific primers (Table S2), cloned and a total of 464 and 386 archaeal and bacterial inserts were partially sequenced. Phylogenetic analysis of the resulting reads revealed a pronounced stratification of prokaryotes thriving in the extremely chaotropic and salty compartments of the Kryos interface just above and below the established chaotropicity boundary of life (Fig. 5a). Remarkably, presence of layer-specific groups of 16S rRNA sequences in all three samples indicated that accurate fractionation of the Kryos interface was successful and reciprocal mixing had not occurred during recovery and subsequent processing of gradient samples. As for the overlaying deep seawater, members of Marine Group I Thaumarchaeota dominate the UIF archaeal community. In concordance with CARD-FISH analysis, ribosomal RNA-containing dormant thaumarchaeal cells have been settled in deeper interface compartment CHW from above layers, thus resulting in a significantly distorted indication of diversity of autochthonous metabolically active archaeal population. Two groups of halophilic Euryarchaeota, the MSBL1 and HC1 candidate divisions, detected in minority in the CHW layer, became dominant euryarchaeal groups in AWW layer, accounting for more than 85% of all archaeal clones (Fig. 5b). 16S rRNA sequences of extremely halophilic haloarchaea and methylotrophic methanogens, also being presented in CHW by singletons, were remainders of the AWW archaeal community. Remarkably, together with clones retrieved from the Lake Discovery, the MSBL1- and Halobacteriales-related Kryos clones formed separate clusters, which constitute evidence for the existence of MgCl₂-adapted species or genera within these candidate divisions (Fig. 6).

The empirically determined water-activity value for the 3.03 M MgCl₂ Kryos layer is 0.631, which is considerably close to the established limit for growth of halophilic prokaryotes (i.e.
Recent studies, however, have demonstrated cell division of more than 10 halophilic prokaryotes, including members of the Halobacteriales in the range 0.717 to 0.6011 A_w (JE Hallsworth et al., unpublished data) and the findings of the current study are consistent with their water-activity minima. These include empirically determined A_w values of 0.693 for Halococcus salifodinae, 0.687 for Halobacterium noricense, and 0.681 for Natrinema pallidum as well as values derived by extrapolation of 0.680 for Halorhodospira halochloris, 0.675-0.670 for halophilic bacteria belonging to the Salinibacter assemblage from crystallizer pond CR-30 (Braç del Port, Alicante), 0.668 for Haloanaerobium lacusrosei, 0.660 for Actinopolyspora halophila, 0.658 for Halobacterium strain 004.1, 0.647 for Halorhabdus utahensis, 0.623 for Halorhodospira halophila, 0.615 for Halobacterium strain GN-5, and 0.611 for Halobacterium strain GN-2.

Phylogenetic composition of the bacterial fraction recovered from all three analyzed layers is shown in Figure 5a and in Supplementary Material (Figure S3 and S4). Compared with the UIF and CHW 16S rRNA libraries, much lower diversity of bacterial phylotypes was recovered from the AWW layer of the Kryos interface. This included members of KB1 candidate division (54% of all clones sequenced) and yet unknown hyperhalophilic groups of the class Deltaproteobacteria (rest of the AWW clones) (Fig. 5b). Whereas coherent KB1-related organisms thrived also in the upper CHW layer, two phylogenetic clusters of Deltaproteobacteria, probably representing different extremely halophilic genera, were detected exclusively in the ultimate layer of the water-activity window for life (AWW) (Fig. 7). The less chaotropic and less salty CHW layer of the Kryos interface was inhabited by completely distinct population of Deltaproteobacteria, consisting of sulfate reducing bacteria (SRB) distantly related to the genera Desulfotignum and halophilic Desulfosalsimonas (Fig. 7). Noteworthy, all bacterial AWW phylogenetic groups have close relatives recovered from sediments of the Mediterranean solar salterns and surficial hypersaline lakes Aran-Bidgol (Iran) and Tebenquiche (Chile) (Demergasso
et al., 2008; Baati et al., 2010; Makhdoumi-Kakhki et al., 2012), thus considerably reducing the sampling efforts needed for their eventual culturing and the study of their physiology and metabolism.

As we have shown previously (Hallsworth et al., 2007), bacterioplankton from overlaying compartments once entered by sedimentation in the sterile Discovery brine, is accumulating there at such highly conserved state that DNA from these organisms can be amplified. Consequently, DNA-based methodologies seem inaccurate approaches to study the “signatures of active life” under highly chaotropic conditions. Indeed, phylogenetic analysis of total DNA sampled at the depth of 3370 m revealed the occurrence in the Kryos brine of 16S rDNA signatures belonging to both Bacteria and Archaea dominating the deep-sea seawater and surficial layers of the interface, but missing in AWW layer (Figures 5a and S3-S5). Namely, almost 30% and 15% of all bacterial and archaeal clones recovered from the Kryos brine were respectively attributed to the Epsilonbacteria and Marine Group I Thaumarchaeota, the groups of prokaryotic organisms which dominated the UIF and CHW layers but were undetectable in the AWW layer. Similar distribution patterns, i.e. lack in AWW but occurrence in the Kryos brine, were observed for the members of Bacteroidetes, Gammaproteobacteria, Planctomycetes and archaeal candidate division SA1. None of brine-specific archaeal 16S rRNA sequences different from that of UIF, CHW and AWW libraries was detected in the Kryos brine, suggesting that all prokaryotic diversity detected in the Kryos brine derived from the overlaying deep seawater column and the interface. UniFrac PCA analysis affirmed that the microbial community of the Kryos interface exhibited notable stratification, mediated by a succession of different groups of organisms. Whereas being marginally different from the intermediate CHW layer ($P = 0.039$), the AWW bacterial population resulted statistically different from the less salty UIF sample ($P = 0.001$) (Fig. S6a). Consistently with the statement that the Kryos brine acts as a trap for descending allochthonous bacterioplankton, no statistical significant distance was found between BB (DNA-
based survey) and AWW layers; and only small difference was observed between BB and CHW layers ($P = 0.033$). The archaeal community behaved in similar manner, although the detected stratification was found to be less pronounced due to aforementioned influence of Marine Group I Thaumarchaeota. Both statistically allied AWW and BB layers resulted only slightly different form the UIF sample (corresponding $P$ values of 0.12 and 0.18) and no statistical significant distances between the AWW, BB and CHW layers were detected (Fig. S6b).

Evidence that metabolic activity occurs in the AWW layer; i.e. below the established chaotropicity window for life

As mentioned above, the majority of the AWW archaeal community comprised of the MSBL1 and HC1 candidate divisions. Previously we speculated, that on basis of phylogenetic relatedness to methanogens and the lack of other groups that might be responsible of the detected methane production in some of Mediterranean DHALs, the MSBL1 members might be involved in methanogenesis at high salinity (van der Wielen et al., 2005; Daffonchio et al., 2006; Borin et al., 2009; Yakimov et al., 2013). Due to the fact that genetic determinants for methanogenesis in MSBL1 organisms remain unknown, we cannot examine their metabolic activities. Nevertheless, the phylogenetic lineage related to the genus Methanohalophilus was detected in AWW interface layer as considerable fraction of clones (5%), thus making feasible the assessment of their methanogenic activity via the recovery of alpha subunit of methylcoenzyme M reductase ($mcrA$) gene transcript. Unlike the $mcrA$ diversity of the Discovery interface, where only Methanohalophilus-related sequences were detected (Hallsworth et al., 2007), the Kryos interface possessed two distinct phylogenetic clusters of this gene (Fig. 5a). The CHW-specific $mcrA$ group was found be distantly related to methylcoenzyme M reductase of Methanomassiliicoccus
luminyensis. This methylotrophic methanogenic Thermoplasmata archaeon carries a reduced methanogenesis pathway, restricted by reduction in the presence of H$_2$ of methanol and other methylated compounds to methane (Dridi et al., 2012; Grolas et al., 2012; Borrel et al., 2013).

Although this type of metabolism was never sought in the DHAL ecosystems, the eventual occurrence of this obligate H$_2$-dependent methylotrophic type of methanogenesis should be taken into account in future studies and cultivation attempts. Coherently with the *Discovery* mcrA gene expression survey, the diversity of the AWW mcrA transcripts was extremely low and all sequences were found be almost identical to that retrieved from the deepest, populated layer of the *Discovery* interface (2.23 M of MgCl$_2$) (Hallsworth et al., 2007). This observation confirmed the assumption that the AWW layer of the *Kryos* interface is inhabited by a distinct archaeal population, which is able to thrive at high concentrations of Mg$^{2+}$.

The bacterial community of the AWW layer characterized by an extremely low diversity, with only two major taxa of hyperhalophilic organisms present. Similarly to MSBL1, lack of genomic information of the KB1 candidate division precludes any of metabolic gene expression surveys. However, the metabolic activity of sulfur reducing deltaproteobacteria in the AWW layer of the *Kryos* interface was indicated by the recovery and analysis of *dsrAB* gene transcripts. It is important to note that the AWW layer contained both the highest H$_2$S concentration and the number of SRBs-related sequences in all three layers analyzed, pointing out to an important ecological role of their members in the sulfur cycle of the *Kryos* ecosystem. This statement is also coherent with the analysis of DOM in the sterile *Kryos* brine, where the biotic sulfur compounds, obviously originated from the overlaying interface, were observed. The existence of hitherto unknown hyperhalophilic groups within SRBs was subsequently corroborated by the phylogenetic attribution of *dsrAB* gene transcripts recovered from the AWW layer (Fig. 7). Remarkably, the recovery and further analysis of *dsrAB* gene transcripts revealed the presence in the AWW layer of the sequences distantly related to that of *Desulfotignum balticum* and
halophilic *Desulfosalsimonas propionicica*. This observation let us to an assumption, that once immersed in the AWW layer, these organisms can likely withstand the high concentrations of Mg$^{2+}$ and remain, albeit briefly, metabolically active.

Concluding remarks

The results obtained in this study portray a very stratified indigenous prokaryotic community thriving at the edge of life in the MgCl$_2$-rich DHAL *Kryos* interface under highly chaotropic conditions. The 25-cm thick *Kryos* interface layer AWW was sampled in range from 2.27 to 3.03 M of MgCl$_2$, which corresponds to salinity of 245 – 330 PSU and water activity values from 0.747 to 0.631. Despite lying beyond the established chaotropicity and prokaryotic life boundaries, the AWW layer seems inhabited by a highly specific community of prokaryotes far different from the thriving above communities. The majority of our AWW archaeal clones (85%) were affiliated to the candidate divisions MSBL1 and HC1 that branched deeply within the Euryarchaeota. These divisions were proposed recently to comprise the majority of the archaeal clones retrieved from the deep-sea Mediterranean Sea Brine Lakes (MSBL) and surficial salt-saturated anoxic lakes (van der Wielen et al., 2005; Jiang et al., 2007). The divisions are equivalent in genetic depth and breadth to Halobacterales and likely represent new orders of yet-to-be-cultivated taxa (van der Wielen et al., 2005). The haloarchaeal *Kryos* AWW clones together with the clones retrieved from the Lake *Discovery* formed a distinct, deeply branched cluster within Halobacterales, thus eventually inferring the existence of new, MgCl$_2$-adapted species or genera. Similarly to archaeal community, the AWW bacterial phylotypes belonged to hitherto uncultured hyperhalophilic organisms, present exclusively in the DHALs and in highly reduced sediments of some surficial hypersaline lakes.

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habitats. Noteworthy, the *Kryos* microbial community, thriving below the established chaotropicity boundary for life, previously established for the Lake *Discovery*, was found be very similar, at least at the level of 16S rRNA phylogeny, to that of the other most hypersaline anoxic environments sampled worldwide. It is plausible that obligate anaerobic hyperhalophiles, adapted to thrive in salt-saturated habitats under low-A$_w$ conditions, possess hitherto uncharacterized mechanisms to resist chaotropicity and to be metabolically active in such harsh athallasohaline habitats (e.g. exceptional levels of cellular desiccation, or unusual and/or highly kosmotropic compatible solutes; Potts, 1994; Cray *et al.*, 2013a; Wyatt *et al.*, 2014a; 2014b). The relevance of our findings encourages digging into the genetic and metabolic diversity of these MgCl$_2$-adapted hyperhalophiles. We are, therefore, conducting additional culturing and metagenomic approaches to obtain a better understanding of the functioning of the *Kryos*-interface ecosystem.

Thus, at present, we must conclude that the question of the window of tolerance of life (i.e. cellular division and/or metabolic activity) for chaotropic activity remains yet open. Compared to Lake *Discovery*, the *Kryos* lake contains slightly elevated concentrations of the kosmotropic ions Na$^+$ and SO$_4^{2-}$. These ions, via their compensating effect against extreme chaotropicity of MgCl$_2$ solutions, are likely to enable cellular activities at MgCl$_2$ concentrations which hitherto considered incompatible with life (Hallsworth *et al.*, 2007). In concordance with our previous statement, we may conclude that life in environments with extremely high concentrations of MgCl$_2$ is unlikely. Nevertheless, the simultaneous presence of kosmotropic ions in Mg-rich environments decreases their chaotropicity and thus, turns them inhabitable for diverse hyperhalophilic microbes. This assumption also has implications for hypersaline Mg-rich milieu, which are known to be located in extraterrestrial environments. However, chaotropic substances such as MgCl$_2$ can be beneficial at low temperatures (those below 10°C, and most especially sub-zero temperatures) by enhancing the flexibility of macromolecule systems, which permits cellular function and thereby reduces the
temperature minimum for cell division of psychrotolerant or psychrophilic microbes (Chin et al., 2010). Ironically, therefore, the possibility remains that high concentrations of MgCl₂ (or other chaotropic salts) on moons or other planetary bodies, which are colder than Earth may potentially increase habitability of aqueous milieu.

Experimental procedures

Oceanographic and geophysical characterization of Kryos basin

The morphobathymetric analysis of the Mediterranean Sea at 500 m resolution (Loubrieueu et al., 2008) was used to locate confined depressions deeper or equal than known DHALs interfaces depths (on average 3200-3300m). The target areas were therefore investigated with the RV Urania hull mounted 16 transducer Benthos 3.5 KHz Chirp SBP looking at any perfectly straight line of reflection, produced by the sharp salinity : density contrast at the seawater : brine interface. Multibeam swath bathymetry was obtained by the R/V Urania Kongsberg-Simrad EM-302 and processed with Neptune, CARIS and GMT packages (Wessel et al., 2013).

Sampling of halocline and brine in the Kryos Lake

Sampling of the Kryos Lake was conducted from the RV Urania at location (22°01'E 35°02'N – 22°02'E 34°53'N) during two oceanographic cruises in September-October 2008 and September 2009 (Fig.1b). Samples were collected using 12-litre Niskin bottles housed on a rosette (General Oceanics, Inc., Miami, FL, USA) equipped with SBE-911plus conductivity-temperature-depth (CTD) sensors (Sea-Bird Electronics, Inc., Bellevue, WA, USA). Determination of oxygen concentration at chosen depths was carried out using the Winkler method (Carpenter, 1965) with an automatic endpoint detection
burette 716 DNS Titrino (Metrohm AG, Herisau, Switzerland). Samples for determining major ion concentrations were collected in 1000 ml dark polyethylene (DPE) vials and stored at room temperature. Alternatively, 110 ml of the samples were diluted with double volume of 0.1 M of HNO$_3$ and stored in 500 ml DPE vials under room temperature prior the chemical analyses. Samples for determining nutrient concentrations were collected in 20 ml DPE vials, quickly frozen in liquid nitrogen and then stored at -20°C. Nutrient concentrations were determined within a few weeks of the end of each cruise using SEAL QuAAtro Microflow Analyzer (SEAL Analytical, ltd, Hampshir, UK).

All running standards were prepared with Low Nutrient Seawater and calibrated against Ocean Scientific Standards (OSIL, Hampshir, UK). Sample analyses were performed at least twice using the same set of equipment. The interface was captured and fractionated as described elsewhere (Daffonchio _et al._, 2006; Hallsworth _et al._, 2007; Yakimov _et al._, 2013). Briefly, 12-L Niskin bottles housed on a rosette with a CTD sensors were closed when a large increase in conductivity, indicating that the interface had been entered, was observed. This was confirmed on-board by measuring the refractive index of the top and bottom of the brine in the Niskin bottles using a hand refractometer (Atago, Tokyo, Japan). Fractions (about 0.5 - 2 l) of the captured interface were sub-sampled and preserved in sealed bottles. Redox potentials (Eh) of subsamples were measured immediately according to the procedure described by Pearson and Stanley (1979). The samples possessing the equal values of salinities were pooled for further treatments as reported below. Among all fractionated samples, the interface layers UIF, CHW and AWW, with salinities of 50-140 PSU, 140-245 PSU and 245-330 PSU, respectively, were subjected to comprehensive analysis of autochthonous microbial life. Moreover, 5 l of the _Kryos_ brine was sampled for comparative purposes from the depth 3370 m.

_Geochemical analyses_
Dissolved cations, anions and organic acids were quantified from diluted samples using standard ion chromatographic techniques, as described below. Conductivity measurements of the samples, determined by a Conductivity meter HI 9818 (Hanna Instruments, Italy), were used to program the dilution. Na⁺, K⁺, Mg²⁺ and Ca²⁺ concentrations were measured using ion exchange chromatography, with a 761 Compact IC ion chromatography system (Metrohm AG, Switzerland) fitted with Metrosep C4 column used without chemical suppression (direct conductivity and reverse polarity modality). Components were separated using a phosphoric acid (5mM) gradient, with a flow of 1 ml min⁻¹.

Volatile fatty acids and sulfate concentrations were measured by ion exchange chromatography using an ICS-2000 ion chromatography system (Dionex®, UK) fitted with two AS15-HC 4 mm columns in series, and a Dionex® Anion Self-Regenerating Suppressor (ASRS-ULTRA II 4-mm) unit in combination with a Dionex®DS6 heated conductivity cell. Components were separated using a potassium hydroxide gradient program as follows: 6.0 mmol KOH (38 min isocratic), 16.0 mmol KOH min⁻¹ to 70 mmol (17 min isocratic). For chloride (Cl⁻) concentrations, column was exchanged with a Ionpac AS9-SC. Chloride were separated using a sodium carbonate (Na₂CO₃) at 2 mmol and sodium bicarbonate (NaHCO₃) at 0.75 mmol with a flow of 1.0 ml min⁻¹.

**Extraction of dissolved organic matter**

Untreated brine (200 ml) was filtered through pre-combusted Whatman GF/F glass fiber filters. The pH was adjusted to 2.0 by using high purity grade formic acid (98 %). Solid-phase extraction (SPE) was followed using Agilent Bond Elut PPL SPE cartridges filled with highly functionalized styrene-divinylbenzene (SDVB) polymer that has been modified with a proprietary non-polar surface. The SPE cartridge was activated using methanol (Sigma-Aldrich Chromasolv LC-MS grade methanol), washed with acidified (pH 2.0) high purity water (Sigma-Aldrich Chromasolv LC-MS grade water). Then, the acidified sample was gravity-fed through the SPE cartridge. The cartridge was washed again with acidified pure water to replace the last remaining inorganic ions from the...
SPE cartridge. After washing, the cartridge was dried under high purity grade nitrogen gas and eluted with methanol.

**Ultrahigh resolution mass spectrometry**

Ultrahigh-resolution mass spectra were acquired on a Bruker (Bremen, Germany) APEX 12 Qe Fourier transform ion cyclotron resonance mass spectrometer equipped with a 12 T superconducting magnet and a APOLLO II electrospray source. The SPE methanol eluate was diluted 1:20 into methanol and introduced into the micro electrospray source at a flow rate of 120 mL/h with a nebulizer gas pressure of 20 psi (138 kPa) and a drying gas pressure of 15 psi (103 kPa) at 250°C through an Agilent sprayer. Spectra were externally calibrated on clusters of arginine (5 mg l⁻¹ in methanol) and systematically internally calibrated with appropriate reference mass list reaching accuracy values lower than 100 ppb in routine day-to-day measurements. Data acquisition was performed using DataAnalysis associated software (Bruker Daltonics, version 4.0). The possible elemental formulas were calculated from the exported masses list for each peak in batch mode by a software tool written in-house (Netcalc). The generated formulas were validated by setting sensible chemical constraints (N rule, double bond equivalent non-negative integers, O/C ratio ≤1, H/C ratio ≤2+2/n (where n is the number of carbon). Van Krevelen diagrams (H/C vs O/C) and (H/C vs m/z) diagrams were used to visualize these datasets.

**Quantitation of water activity and chaotropic activity**

Besides the natural Kryos brine, an artificial analogue brine was also used for water activity and chaotropicity determination. This synthetic, analogue 'Lake Kryos' brine was made up by dissolving following salts: MgCl₂ (4.1841 M), MgSO₄ (0.2183 M), Na₂SO₄ (62 mM) K₂SO₄ (42.2 mM), CaCl₂ (1 mM), (NH₄)₂SO₄ (0.4 mM) which was stored for one week at 14.3°C prior to water-activity determinations.
Water activities were determined over a range of concentrations at 14.3°C using a Novasina IC II water activity machine fitted with an alcohol-resistant humidity sensor and eVALC alcohol filter (Novasina, Pfäffikon, Switzerland), as described previously (Hallsworth and Nomura, 1999). This brine was super-saturated as a fine, powdery precipitate could be seen by eye. For quantification of chaotropic activity, agar gel-points were determined over a range of salt or brine concentrations (see Hallsworth et al., 2003a; 2007) using a Cecil E2501 spectrophotometer fitted with a thermoelectrically controlled heating block (Milton Technical Centre, Cambridge, England) as described previously (Cray et al., 2013a).

CARD-FISH analysis

CARD-FISH samples (50 ml) were collected from overalying seawater, the interface layers UIF, CHW, AWW and the Kryos brine. Samples were fixed with 2% formaldehyde (v/v, final concentration) at room temperature for 1 hour and stored at -20°C in the dark until laboratory analysis. Subsamples (from 1 to 10 ml, according to cell concentrations) were filtered through polycarbonate membranes (Ø25 mm, 0.22 µm pore size, NTG). Cells were permeabilized with lysozyme (10 mg ml$^{-1}$, 1 h) and achronopeptidase (5 mg ml$^{-1}$, 30 min) at 37°C. Intracellular peroxidase was inhibited by treatment with HCl (0.01 mmol l$^{-1}$) at room temperature for 20 min. We used the following horseradish peroxidase labeled probes: EUB338 I-III, ARCH915, CREN537, EURY806, KB1, and Delta-DHAL. Detailed information about the probes shown in Table S1. The nonspecific probe NON338 did not detect any cells. The filters sections were counter-stained with DAPI (2 mg ml$^{-1}$) in a 4:1 ratio of Citifluor (Citifluor Ltd, Leicester, UK) and Vectashield (Linaris GmbH, Wertheim- Bettingen, Germany). At least 200 DAPI cells, in a minimum of 10 fields, were counted in the AXIOPLAN 2 Imaging microscope (Zeiss). Negative control counts were performed with HRP-Non338 probe, always amounting to < 1% of DAPIstained cells.
**Nucleic acid purification and cDNA synthesis**

For DNA/RNA extraction, 2-5 l of the fractionated interface and brine samples were filtered through sterile Sterivex capsules (0.2μm pore size, Millipore) using a peristaltic pump. After filtration, filters were treated with 400μl of TE buffer (pH 8.0) containing lysozyme (5 mg ml⁻¹), vortexed for 5 sec and incubated 10 min at room temperature. 1600μl of lysis buffer QRL1 (containing β-mercaptoethanol) were added and filters were than stored at -20°C until processing. Total DNA and RNA were extracted using Qiagen RNA/DNA Mini Kit (Qiagen, Milan, Italy). The extraction was carried out according to the manufacturer’s instructions. DNA and RNA samples were examined by agarose gel electrophoresis and concentrations were determined using the NanoDrop ND-1000 Spectrophotometer (Wilmington, DE, USA). RNA-containing extracts were purified from DNA by Turbo DNA-free kit (Ambion, Austin, TX, USA). Each RNA sample was immediately converted into cDNA with SuperScript II Reverse Transcriptase (Invitrogen, Carlsbad, CA, USA) and hexa-random primers according to the manufacturer instructions.

**PCR-amplification, gene cloning and sequencing**

Bacterial 16S rRNA, archaeal 16S rRNA and key genes involved in sulphur respiration (dsrAB) and methanogenesis (mcrA), were amplified by PCR using primers listed in the Table S1. All reactions were carried out in a MasterCycler 5331 Gradient PCR (Eppendorf, Hamburg, Germany). The conditions for PCR and cloning were performed as described elsewhere (La Cono et al., 2011, Yakimov et al., 2013). Positive clones from each library were randomly selected by PCR amplification. The PCR products were further purified and sequenced at Macrogen (Amsterdam, Netherlands).

**Phylogenetic trees**
Pintail software (Ashelford et al., 2005) was used to check sequences for possible chimeric origin. 16S rRNA gene amplified sequences and close relatives identified with BLAST (Altschul et al., 1997) were aligned using the SILVA alignment tool (Pruesse et al., 2007) and manually checked with ARB (Ludwig et al., 2004). MEGA 5 (Tamura et al., 2007) was used to align functional genes nucleotides sequences. After alignment, the neighbor-joining algorithm of ARB and MEGA 5 program packages were used to generate the phylogenetic trees based on distance analysis for 16S rRNA and functional genes, respectively. The robustness of inferred topologies was tested by bootstrap re-sampling using the same distance model (1,000 replicates). Significant difference of the microbial assemblages derived from different samples depths was detected via the $P$-test significance and principal coordinates analysis (PCA) using UniFrac program (http://bmf2.colorado.edu/fastunifrac (Hamandy et al., 2009; Lozupone et al., 2007) for comparison of the microbial communities using phylogenetic information.

### Nucleotide sequence accession numbers

The nucleotide sequences produced in the present study have been deposited in the DDBJ/EMBL/GenBank databases under accession numbers: KJ922395 to KJ922487 for the bacterial and archaeal 16S rRNA gene sequences, KJ922632 to KJ922638 for the archaeal $mcrA$ gene sequence, KJ922623 to KJ922631 for the bacterial $dsrA$ gene sequences.

### Acknowledgements

This work was performed with the financial support of CNR in frames of the EU FP7 Projects MAMBA (KBBE-2009-2B-226977) and MicroB3 (OCEAN 2011-2-287589). JAC was supported by funding received from the Department of Agriculture and Rural Development (Northern Ireland). We thank the Master, crew of RV Urania and all participants to the cruises for their valuable professionalism and
support during the cruises. A. Modica, M. Catalfamo are indebted for geochemical analyses. The
discoveries were possible only because of good topographic data made available by other
Institutions, and we thank Dr. Benoit Loubrieu of IFREMER for providing the 500m-resolution DTM
data.

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16S rRNA sequence records currently held in public repositories is estimated to contain


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Borin, S., Brusetti, L., Mapelli, F., D'Auria, G., Brusa, T., Marzorati, M., et al. (2009) Sulfur cycling and
methanogenesis primarily drive microbial colonization of the highly sulfidic Urania deep


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*Nature* **305**: 797-798.


Table 1. Chemical composition of the three saltiest DHALs on Earth.

<table>
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<th>DISCOVERY*</th>
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<th>L’ATALANTE</th>
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<td>6.8</td>
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These data were partially taken from Wallmann *et al.* 1997, 2002.
Table 2. Chemical compositions of the most anhydrous \((A_w < 0.700)\) athalassohaline lakes on Earth and primary (LSPB) and secondary (SB) brines. All concentrations are in mM \((\text{kg H}_2\text{O})^{-1}\) unless otherwise stated.

<table>
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<tr>
<th>Parameters</th>
<th>LSPB*</th>
<th>SB*</th>
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<th>DON JUAN PONDd</th>
<th>DISCOVERY</th>
<th>KRYOS</th>
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<td>459</td>
<td>5830</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>SO(_4)</td>
<td>173</td>
<td>122</td>
<td>6</td>
<td>&lt;1</td>
<td>110</td>
<td>320</td>
</tr>
<tr>
<td>Cl</td>
<td>10100</td>
<td>10926</td>
<td>6824</td>
<td>12192</td>
<td>10150</td>
<td>9043</td>
</tr>
<tr>
<td><strong>Parameters</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>pH</td>
<td>~5.6(^a)</td>
<td>~5.6(^a)</td>
<td>7.7</td>
<td>~5.4(^a)</td>
<td>~4.5(^a)</td>
<td>~5.4(^a)</td>
</tr>
<tr>
<td>Water activity, (A_w)</td>
<td><strong>0.420</strong></td>
<td><strong>0.380</strong></td>
<td>0.690</td>
<td><strong>0.411</strong></td>
<td><strong>0.382</strong></td>
<td><strong>0.399</strong></td>
</tr>
<tr>
<td>Salinity, g kg(^{-1})</td>
<td>513</td>
<td>515</td>
<td>359</td>
<td>670</td>
<td>510</td>
<td>471</td>
</tr>
<tr>
<td>Density, kg L(^{-1})</td>
<td>1.33</td>
<td>1.33</td>
<td>1.22</td>
<td>1.39</td>
<td>1.33</td>
<td>1.32</td>
</tr>
</tbody>
</table>

\(^a\) As it described elsewhere (Wallmann et al., 1997, 2002), the late stage primary brine (LSPB) was produced by evaporation of seawater and precipitation of anhydrite \((\text{CaSO}_4)\), halite \((\text{NaCl})\), kieserite \((\text{MgSO}_4 \cdot \text{H}_2\text{O})\) and carnallite \((\text{K}_2\text{MgCl}_3 \cdot 6\text{H}_2\text{O})\). The evaporation was performed at atmospheric pressure and 30°C and continued until only 5g of the initial 1 kg H\(_2\)O remained in solution.

\(^b\) Secondary brine (SB) produced by equilibrating of calcite-saturated seawater with the evaporite minerals bischofite \((\text{MgCl}_2 \cdot 6\text{H}_2\text{O})\), kainite \((\text{K}_2\text{Mg(SO}_4)\text{Cl} \cdot 3\text{H}_2\text{O})\), halite, and gypsum \((\text{CaSO}_4 \cdot 2\text{H}_2\text{O})\) at 14°C and 1 atm (Wallmann et al., 2002).

\(^c\) Composition of the Dead Sea (Israel) and the Don Juan Pond (Antarctica) were taken from Marion et al. 2003. Water activity values below the window of cellular life \((A_w < 0.605)\) are highlighted in bold.
Table 3. Abundance of general and specific phylogenetic groups within *Bacteria* and *Archaea* in the Kryos interface layers and overlaying seawater.

The total cell numbers are given as $10^5$ cells ml$^{-1}$ unless otherwise stated. Cells were collected from the indicated layers and hybridized with the specific CARD-FISH probes (Yakimov et al., 2013).

<table>
<thead>
<tr>
<th>Interface layer, (Mg$^2+$, M / salinity, PSU)</th>
<th>DAPI</th>
<th>EUB338 I-III</th>
<th>KB1</th>
<th>Delta-DHAL</th>
<th>ARCH915</th>
<th>CREN537</th>
<th>EURY806</th>
</tr>
</thead>
<tbody>
<tr>
<td>SW*</td>
<td>0.16±0.02</td>
<td>0.12±0.02</td>
<td>0</td>
<td>0</td>
<td>0.02±0.008</td>
<td>0.02±0.004</td>
<td>0</td>
</tr>
<tr>
<td>UIF (0.16/52)</td>
<td>0.93±0.10</td>
<td>0.32±0.08</td>
<td>0</td>
<td>0</td>
<td>0.17±0.05</td>
<td>0.16±0.02</td>
<td>0.02±0.01</td>
</tr>
<tr>
<td>CHW (1.55/195)</td>
<td>5.57±0.45</td>
<td>2.97±0.56</td>
<td>0.17±0.03</td>
<td>0.14±0.04</td>
<td>0.81±0.77</td>
<td>0.05±0.01</td>
<td>0.22±0.01</td>
</tr>
<tr>
<td>AWW1 (3.03/327)</td>
<td>4.60±0.43</td>
<td>2.45±0.21</td>
<td>0.69±0.12</td>
<td>0.14±0.03</td>
<td>0.43±0.07</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>AWW2 (3.41/370)</td>
<td>2.47±0.11</td>
<td>1.53±0.07</td>
<td>1.22±0.11</td>
<td>0.13±0.01</td>
<td>0.47±0.05</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

*See Figure 2 for exact positioning of sampling points within three layers of the Kryos interface.

These data correspond to the seawater column sampled from the depth of 2850 m twenty nautical miles NE from the Lake Kryos during the same cruise as it described elsewhere (La Cono et al. 2011).
Figure 1. Location of currently known DHALs in the Eastern Mediterranean Sea (a) and the detailed swath bathymetry at the Kryos Lake area (b). The map to the left was constructed using the Ocean Data View software (Schlitzer et al., 2010). On the right, the shape of the anoxic lake and small polar satellite pools are colored in blue starting from the seawater : brine lake interface (3337 m depth). The sampling sites are highlighted by asterisks.

Figure 2. Physicochemical activities of MgCl$_2$ solutions, the Lake Kryos brine and a synthetic Kryos-brine analogue: (a) water-activity reduction over a range of MgCl$_2$ concentrations at 14.3°C (for comparative purposes all values are expressed according to their MgCl$_2$ content) and the upper dotted line indicates the lower boundary of the previously established chaotropicity limit of life (CHW, equivalent to 2.3 M MgCl$_2$; Hallsworth et al., 2007) and the lower dotted line denotes the established water-activity limit for xerophilic fungi (AWW; Pitt and Christian, 1968); and (b) agar gel-point depression (a measure of chaotropic activity; Cray et al., 2013a). The solid lines with arrows indicate the water-activity values corresponding to the AWW layer (MgCl$_2$ 2.27 - 3.03 M).

Figure 3. Depth profiles of geochemical markers through the Lake Kryos and the established boundary for chaotropicity and xerophilic cellular life occurred in the Kryos gradient. As far as all conventional on-line CTD sensors were not functional in MgCl$_2$-rich ambience, chemical analysis of fractionated interface samples were performed in the in-land laboratory. The brine was collected at the depths of 3340 and 3370 m bsl. The layers of interface collected for the molecular
analyses are highlighted in green, blue and red. Following the $A_w$ calculations (Fig. 2a), the boundary for chaotropicity ($A_w 0.790$) and xerophilic cellular life ($A_w 0.605$) occurred in the $Kryos$ gradient are shown. Abbreviations used: AWW, the interface layer corresponding to lower boundary of estimated xerophilic cellular life; BB, body brine; CHW, the interface layer corresponding to lower boundary of chaotropicity life; UIF, upper interface. Data points are mean ± standard error (n=3).

Figure 4. Ultrahigh resolution mass spectrometry of the $Kryos$ brine DOM showing hundreds of low molecular weight organic compounds (a) with $m/z <$500 amu. The van Krevelen diagrams (b, c) illustrate the high proportion of largely saturated structures and the remarkable extent of oxygenation of the CHNO and poly-sulfur compounds (d). The blue line refers to any fully saturated open chain aliphatic (poly)carboxylic acid, (comparable to polymaleic acid or polyacrylic acid as model structures) and the red line to the compositional range of CRAM molecules as described in Hertkorn et al. (2006).

Figure 5. Overview on prokaryotic diversity, stratification (a) and relative abundance (b) of phylogenetic groups recovered from the different compartments of Lake $Kryos$.

(a) Stratification and relative abundance of each phylogenetic group found in different layers of the Lake $Kryos$ is shown as number of cloned and analysed sequences related to the indicated group. The clones recovered from the $Kryos$ brine, the upper interface (UIF), the layer of chaotropicity (CHW) and the water activity (AWW) windows are shown in black, green, blue and red, respectively. Scale bar corresponds to 10% estimated difference in nucleotide sequence positions.

(b) Extent of recovery of 16S crDNA AWW clone sequences in overlaying layers UIF and CHW. Scale white bar corresponds to 20% of all cloned sequences analyzed separately in UIF, CHW and AWW clone libraries. Exact percentages of clones corresponding to each indicated phylogenetic group are given for clarity.
Abbreviations of candidate division used: BRC1, Bacterial Rice Cluster; DP, Deltaproteobacteria; HA, haloarchaea; HC1, Halophilic Cluster 1; KB1, Kebrit Deep Bacteria 1; MH, Methanohalophilus; MSBLx, Mediterranean Sea Brine Lakes; OM27, Ocean Margins 27; SA1, Shaban Deep Archaea 1; SARx, Sargasso Sea Clusters.

Figure 6. Phylogenetic analyses of clone sequences of Archaea and mcrA gene transcripts recovered from the AWW interface layer.

The 16S rRNA phylogenetic analysis indicates the relationship between AWW archaeal clone sequences and related sequences recovered from the CHW interface layer, the Kryos brine and other DHALs and surficial hypersaline lakes. The analysis of sequences derived from mRNA coding for methyl co-M reductase (mcrA) indicates that methanogens similar to both the lake Discovery organisms and Methanohalophilus halophilus are active in the AWW layer. The white and solid cycles at the nodes indicate the percentages of recovery in 1,000 bootstrap resamplings of < 75% and ≥ 75%, respectively. Only relevant bootstrap values of ≥70% are shown. Scale bar corresponds to 5% estimated difference in nucleotide sequence positions. Trees were respectively rooted with Desulfotignum balticum 16S rRNA (AF233370) and Methanobrevibacter arboriphilus DSM 1125 mcrA (AF414035) gene sequences.

Figure 7. Phylogenetic analysis of bacterial clone sequences and dsrAB gene transcripts recovered from the AWW interface layer.

The phylogenetic analysis indicates the relationship between AWW bacterial clone sequences and related sequences recovered from the CHW interface layer, the Kryos brine and other DHALs and surficial hypersaline lakes. It also demonstrates that a taxonomic (16S rRNA) and a functional (dsrAB) marker give largely congruent phylogenies and the main taxa identified were
*Desulfobacteracaea* and *Desulfohalobiaceae*. The white and solid cycles at the nodes indicate the percentages of recovery in 1,000 bootstrap resamplings of < 75% and ≥75%, respectively. Only relevant bootstrap values of ≥70% are shown. Scale bar corresponds to 5% estimated difference in nucleotide sequence positions. Trees were respectively rooted with *Halorhabdus tiamatea* 16S rRNA (NR_113213) and *Thermodesulforhabdus norvegica* dsrAB (AF334597) gene sequences.