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Conversion of uric acid into ammonium in oil-degrading marine microbial communities: a possible role of halomonads

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Abstract

Uric acid is a promising hydrophobic nitrogen source for biostimulation of microbial activities in oil-impacted marine environments. This study investigated metabolic processes and microbial community changes in a series of microcosms using sediment from the Mediterranean and the Red Sea amended with ammonium and uric acid. Respiration, emulsification, ammonium and protein concentration measurements suggested a rapid production of ammonium from uric acid accompanied by the development of microbial communities containing hydrocarbonoclastic bacteria after 3 weeks of incubation. About 80% of uric acid was converted to ammonium within the first few days of the experiment. Microbial population dynamics were investigated by Ribosomal Intergenic Spacer Analysis and Illumina sequencing as well as by culture-based techniques. Resulting data indicated that strains related to *Halomonas* spp. converted uric acid into ammonium, which stimulated growth of microbial consortia dominated by *Alcanivorax* spp. and *Pseudomonas* spp. Several strains of *Halomonas* spp. were isolated on uric acid as the sole carbon source showed location specificity. These results point towards a possible role of halomonads in the conversion of uric acid to ammonium utilized by hydrocarbonoclastic bacteria.

Introduction

Microbial hydrocarbon degradation in the sea occurs on a global scale in a large variety of latitudinal zones and at all ocean depths. More than five decades of research have shown that marine oil degrading microbes are highly specialised with regard to their substrate spectrum and climate zone (Head et al., 2006; Yakimov et al., 2007). Cold-adapted microbes such as *Oleispira* spp. prevail in cold polar zones and in the deep sea whereas oil degrading consortia in temperate zones are dominated by *Alcanivorax* species (Kube et al., 2013) in the aftermath of major oil spills. Many members of *Oceanospirillales* and *Alteromonadales* (e.g. *Thalassolituus* spp. and *Marinobacter* spp.) become predominant in the early stages of oil-degrading microbial consortia due to their highly adapted metabolism (Kasai et al., 2002; Yakimov et al., 2004). Late stages of marine hydrocarbonoclastic consortia on the other hand are characterized by bacteria such as *Cycloclasticus* spp. specialized in aromatic and polyaromatic hydrocarbons that are degraded in a slower process (Head et al., 2006).

To perform hydrocarbon degradation, microbes depend upon nitrogen and, to a lesser extent, phosphorus sources (Atlas and Bartha, 1972). Both elements are scarce in seawater in most parts of the oceans and over most of the warmer months due to competition for nutrients by phototrophs that use sunlight and CO₂ as energy and carbon sources. In fact, dissolved organic nitrogen, ammonium and nitrate exist in mere trace amounts after phytoplankton spring blooms from March to June and are recycled in winter months when low temperatures prohibit fast growth of heterotrophic microbes (Garcia et al., 2010). The artificial addition of nitrogen and phosphorus sources into an oil impacted environment on the other hand leads to a quick growth of hydrocarbonoclastic microbes and substantial microbial hydrocarbon degradation. An example of the successful implementation of this approach was the M/V Exxon Valdez mitigation trial (Lindstrom et al., 1991). Studies of marine oil-degrading microbial communities growing in dispersed oil did not show a significant increase of oil degrading microbes beyond 10⁵ cells/mL and no significant biogenic hydrocarbon degradation, either (Hazen et al., 2010).

A variety of different nitrogen sources can be used to support growth of hydrocarbonoclastic bacteria (HCB). Oil spill remediation trials commonly use nitrogen in forms of ammonium, nitrate, urea, uric acid and amino acids (Fava et al., 2004). Each nitrogen source has its advantages and disadvantages (Nikolopoulou and Kalogerakis, 2008): ammonium and nitrate are water-soluble and universally accepted by microbes, but they are toxic at high concentrations as well as prone to leaching and losses by denitrification or precipitation in seawater (Wrenn et al., 2001). Urea is water-soluble and widely accepted by microbes but easily leached. Uric acid is less widely accepted by microbes. Its oleophilic nature allows it to attach to oil droplets (Koren et al., 2003) but it is largely insoluble in water. Lecithin is a

complex nitrogen and phosphorus source that has recently been used in aromatic hydrocarbon degradation experiments (Fava, et al., 2004). While it is readily used and can be supplied in large quantities, the mechanism of lecithin metabolism in PAH-degrading microbial consortia remains unknown. Furthermore, large amounts of lecithin must be applied due to the high C:N ratio of the lecithin molecule. Certain industrial processes can overcome the disadvantages of some nitrogen sources e.g., sulphur coating of urea (Lee et al., 1993) or using vegetable oil coated ammonium/nitrate in slow release fertilizers (Gertler et al., 2009). This reduces the cost efficiency of such nitrogen sources in bioremediation trials, though.

Uric acid could be used without prior treatment, but its synthesis is relatively expensive. However, uric acid is the main nitrogen excretion product of reptiles, birds and certain mammals. It is therefore readily available in poultry litter, in which it makes up to 55-63% of total nitrogen and to 3% (w/w) of the total weight (Nahm, 2003). Uric acid enters marine ecosystems in substantial amounts via faeces of seabirds. A recent study detected measurable effects of uric acid fertilisation in the vicinity of Great Cormorant (*Phalacrocorax carbo sinensis*) colonies in the Baltic Sea (Gagnon et al., 2013). More than 50% of the nitrogen taken up by algae at the observed sites originated from faecal uric acid deposition. While it is obvious that valuable nitrogen sources such as uric acid are quickly taken up by marine food webs, the mechanisms and microbial uric acid utilization and the microorganisms involved in this process remain largely unknown (Capone et al., 2009)

Trials in sediments and soil revealed that microbes related to *Acinetobacter baumannii* can use uric acid as a nitrogen source during hydrocarbon degradation (Koren et al., 2003). When uric acid was applied to marine sediments in form of guano, microbes related to the genera *Alcanivorax*, *Halomonas* and *Alteromonas* could be enriched and isolated (Knezevich et al., 2006). Uric acid is an intermediate of the purine metabolism and is oxidised by uricase enzymes. Members of the *Bacillus*, *Pseudomonas*, *Alcaligenes* and *Microbacterium* genera have been identified as being able to use uric acid as carbon and nitrogen source (Koren et al., 2003; Vogels & van der Drift, 1976). It is therefore highly probable that marine oil degradation with uric acid as main nitrogen source requires a consortium of currently unknown uric acid-degrading microbes and highly adapted HCBs. This knowledge gap is very surprising as uric acid-based bioremediation approaches have been investigated in trials at different scales (Koren et al., 2003; Knezevich et al., 2006; Nikopoulou, 2008, 2013).

This study aimed at the identification of marine microbial consortia members involved in the metabolism of uric acid in marine oil-degrading and uric acid utilizing consortia.

Materials and Methods

Study sites

The sampling sites were located along the Northern and Southern shores of the Mediterranean Sea and the Red Sea. In particular: (1) the site of El Max (31°9'31.20"N, 29°50'28.20"E) located west of the city of Alexandria, Egypt, the El Max seashore is the most contaminated in the Alexandria area exceeding the legal limits for heavy metals, PAHs and other crude oil-derived pollutants; (2) the Bizerte lagoon (37°16'7.72"N, 9°53'19.61" E) in the North of Tunisia, is polluted by petroleum components in the area adjacent to STIR refinery, the Tunisian Company for Refining Industries; (3) the Jordanian coast in the Gulf of Aqaba (30°22'42''N, 25°24'57''E) in the north-west end of the Red Sea. The Gulf of Aqaba is the northernmost tropical marine ecosystem but hosts an oil terminal that yearly moves up to 30 million tons and is subjected to accidental oil spills at the oil terminal as well as spills (with high sulphur concentrations) during loading and bunkering operations at the industrial jetty; (4) the harbour of Ancona, Italy, (43°37'N; 13°30'15"E), which is a major ferry terminal and industrial port on the Adriatic Sea and heavily contaminated with PAHs and heavy metals.

Microcosm design and set-up

Two different sizes of microcosms were set up for this trial (Table 1). For the analysis of microbial population dynamics, respiration and other parameters, a series of 16 microcosm flasks (two parallels each for a uric acid and an ammonium treatment for sediments from the Ancona, Aqaba, Bizerte and El Max sites, respectively) was set up in 500 mL Erlenmeyer flasks (Microcosms 1-8). Seventy-five grams of sand, 5 mL of crude oil and 150 mL of the ONR7 medium mentioned above were used. Overall nutrient concentrations and C:N:P ratios were identical to the ones in the large microcosms mentioned above. All microcosms were incubated at 20°C for 28 days in a shaking incubator at 120 rpm. The small microcosms were sampled on days 0, 2, 4, 6, 8, 10, 12, 15, 18, 21, 24 and 28. On these days, 1 mL of sediment was taken for DNA extraction, 1 mL of supernatant was taken for photometric measurements and 6 ml were taken for emulsification measurements. Fifty millilitres of each microcosm were taken on days 0, 4, 8, 12, 15, 18, 21, 24 and 28 to measure respiration activity.

The sample volume taken for each sampling day was replaced by an equivalent volume of modified ONR7a medium. For ammonium treatments, the equivalent amount of ammonium chloride and disodium hydrogen phosphate was added to compensate for loss of water-soluble nitrogen sources during sampling. As uric acid in the respective uric acid treatments is

insoluble in water, sample volumes were replaced with ONR7a containing disodium hydrogen phosphate but no ammonium chloride.

Four control treatments with a total of eight control flasks were set up. A negative control contained only sterile sand and ONR7a. Two further controls contained sand, ONR 7a, crude oil and either uric acid or ammonium chloride solution but no sediment sample. Finally, one control contained oil, sterile sand, ONR7a medium and a sediment sample from Ancona harbour, but no additional nitrogen source or phosphorus source was provided (Microcosms 9 – 12).

To generate high DNA yields required for sequencing analysis, a series of up-scaled microcosms was set up (Microcosms 13-20). 1 L Erlenmeyer flasks were filled with 150 g of sand (Sigma-Aldrich, St. Louis, USA), sterilized and spiked with 10 mL of sterile filtered Arabian light crude oil. One gram of sediment from each sampling site was mixed into the oil-spiked sand as the inoculum. For the Aqaba site, 2 g of sediment were used, as this sediment was very coarse. Three hundred millilitres of modified ONR7a medium (Dyksterhouse et al., 1995) (omitting ammonium chloride and disodium hydrogen phosphate) were added. We added 5 ml of Arabian light crude oil, which based upon average literature values for density and molecular weight equals about 300 mM of C (Wang et al., 2003), 5 mM of NH_4Cl and 0.5 mM of Na_2HPO_4 resulting in a molar N:P ratio of approximately 10:1. For each uric acid treatment microcosm (UA), 0.21 g (1.25 mmol = 5 mmol N) of uric acid was provided as nitrogen source while the ammonium treatment microcosms (NP) were each supplied with 2.5 mL of a 2M ammonium chloride solution (5 mmol; pH 7.8). Both treatments also contained 2.5 ml of a 0.2M disodium hydrogen phosphate solution (0.5 mmol; pH 7.8). Excess amounts of crude oil were added to compensate for the 35% carbon losses due to evaporation of volatile hydrocarbons over the course of the experiment (Wang et al., 2003). Including losses due to evaporation the C/N/P ratio was approximately 400:10:1. One hundred and four millilitres of mesocosm water were replaced by an equal volume of modified ONR 7 containing uric acid or ammonium as described above on days 0, 4, 8, 12, 15 and 18. The microcosms were destructively sampled on day 21.

Emulsification measurements

Six millilitres of sample were pipetted into screw cap glass centrifuge tubes (15x150 mm). Six millilitres of *n*-hexane (Sigma-Aldrich, St. Louis, USA) were added and each tube was vortex-shaken at maximum rate for 60 s. The resulting emulsion was left to settle for 24 h at room temperature and the height of the interface which settled between hexane and the aqueous phase was measured with a calliper.

Respiration measurements

Oxygen consumption was measured using a Micro-Oxymax Respirometer (Columbus Instruments, Columbus, OH, USA) to serve as a proxy for total aerobic metabolic activity. Fifty millilitre aliquots were transferred to 100 mL screw top glass bottles (VWR, Lutterworth, UK) and connected to the respirometer. Automatic volume determination was performed for each glass bottle prior to respiration measurement. Measurements were performed every 2 h for a total of 72 h. For each measurement, sensors and bottles were automatically purged with sterile air to avoid inhibitory effects of oxygen depletion.

Photometric determination of ammonium, nitrate and protein concentration

Due to the presence of emulsified oil in most microcosms, all samples underwent a pre-treatment before photometric measurement. Briefly, each 1 mL sample was sonicated using a Soniprep 150 sonicator (MSE, London, UK) for 20 s in an ice bath. Subsequently, samples were centrifuged at 16,000g in an Eppendorf centrifuge to separate biomass incorporating oil at the bottom of the centrifuge tube from free oil at the surface of the aqueous phase. Sub samples were taken from the centre of the aqueous phase, diluted and applied in photometric analysis. A nitroprusside-based assay was used (Mulvaney et al., 1996) for the determination of ammonium concentrations and a vanadium chloride based method (Miranda et al., 2001) was used to measure nitrate concentration. Protein concentrations were determined as a proxy for microbial growth using the BioRad Protein assay (BioRad, Carlsbad, USA) according to the manufacturer's guidelines.

Isolation, 16S rRNA gene sequencing and phylogenetic analysis of uric acid-utilising microorganisms

Samples of one hundred microliters were taken from microcosms 13 to 20 on days 6 and 12 of the experiment. A serial dilution was performed using sterile modified ONR7a medium that contained neither nitrogen nor phosphorus source. 100 µL of each dilution were plated on modified ONR7a agar plates containing 0.3% (w/v) uric acid and 0.45 mM disodium hydrogen phosphate. Due to its low solubility in water, uric acid forms a precipitate, which makes the agar plates appear milky. The plates were incubated at 20°C for up to 5 d. Isolates with capabilities to degrade uric acid were identified by clear “halos” around the individual

colonies. Eight colonies of each variant were re-plated twice on modified ONR7a agar plates to obtain pure isolates. Each isolate was tested for the ability to use uric acid as the sole source of carbon and nitrogen in 15 mL polypropylene tubes containing 5 ml ONR7 liquid medium supplemented with uric acid and disodium hydrogen phosphate, as described above. Tubes were incubated at 20°C for 5 to 7 d. Colony PCR of each isolate was performed using primers F530 and R1492 to amplify the 16S rRNA gene. Each 20 µL PCR reaction contained 2 µL of PCR Buffer B (Roboklon, Berlin, Germany), 2.5 mM MgCl₂, 200 µM of each dNTP (Promega, Madison, WI), 10 nM each of primer F530 (5'- GTG CCA GCM GCC GCG G-3') and primer 1492r (5'- GGT TAC CTT GTT ACG ACT T-3'), 0.5 U of Taq DNA Polymerase (Roboklon, Berlin, Germany) and 4 µL of a PCR-enhancing mixture (3 M betaine and 1% Tween 20). PCR products were visualised on 1% agarose gels. Sequencing was performed at Macrogen (Amsterdam, Netherlands) with primers F530 and R1492. The processing and assembly of DNA sequences as well as testing for chimeric sequences was conducted using BioEdit as described before (Ashelford et al., 2005; Gertler et al., 2012). Sequence alignment was performed against a database of validly published reference DNA sequences of typed strains using ClustalW (Larkin et al., 2007). The sequences were clustered by maximum parsimony according to the recommendation of the MODELTEST software (Posada & Crandall, 1998) and phylogenetic tree was constructed using MEGA 4.0 as previously described (Gertler et al., 2010, 2012). DNA sequences of *Halomonas* isolates were submitted to the European Nucleotide Archive (ENA) under the accession numbers HG803097 to HG803140.

Nucleic acid extraction

As both Illumina sequencing and RISA performed in this study require different qualities and quantities of DNA, nucleic acids were isolated using two approaches. For Illumina sequencing (Microcosms 13-20), a modified CTAB-phenol-chloroform extraction was performed. Briefly, for the initial cell lysis, 20 g of sediment, 1 g of 2-3 mm glass beads (5 to 6 glass beads) and 10 mg of lysozyme were placed in a 50 mL centrifuge tube. Fifteen millilitres of extraction buffer containing 10% (w/v) sucrose, 20% (v/v) modified CTAB solution (5% CTAB; 0.35 M NaCl, 120 mM Na₂HPO₄ pH 8.0; to improve solubilisation of CTAB, this solution is a 1:1 mixture of 10% CTAB/0.7 M NaCl and 240 mM Na₂HPO₄ solutions), 100 mM NaEDTA pH 8.0 and 100 mM Tris/HCl pH 8.0 were added to the sample, and all tubes were incubated on a shaker at 220 rpm and 30°C for at least 30 min. Twenty milligrams of proteinase K were added to each tube, and all samples were incubated on a shaker at 220 rpm and 50° C for at least 30 min. Two millilitres of a 20% SDS (w/v) solution were added and all samples were incubated on a shaker at 100 rpm and 60° C for 20 min. Finally, 0.5 g of activated charcoal powder was added to the mixtures, and the samples were

incubated on a shaker at 100 rpm and 60° C for 30 min. Resulting mixtures were centrifuged at 17,000g for 10 min.

Resulting supernatants were transferred to another 50 mL centrifuge tube. The pellets were extracted again by adding 5 mL of extraction buffer to the pellet, mixing for 5 min and centrifugation as described above. The resulting supernatant was combined with the supernatant from the previous step. To remove organic contaminants, the combined supernatants were treated with 0.5 g of activated charcoal, which was subsequently removed by centrifugation at 20,000g for 10 min. Supernatants were transferred to new 50 mL centrifuge tubes and 0.5 M NaEDTA solution was added to the supernatants for receive a final EDTA concentration of 2.5 mM. Deproteination and DNA extraction were performed by adding one supernatant volume of phenol/chloroform/isoamylalcohol (PCI) solution (25:24:1 (v/v); Sigma-Aldrich, St. Louis, USA). The mixtures of PCI and supernatant were inverted manually for 2 min and subsequently centrifuged at 3,700g for 10 min. The aqueous phase was transferred to a new centrifuge tube and extracted again with chloroform/isoamylalcohol (24:1 (v/v); Sigma-Aldrich, St. Louis, USA). The resulting supernatants were precipitated by adding 0.8 volumes of isopropanol and incubating at 4°C overnight. Precipitates were collected by centrifugation at 18,000 rcf for 20 min at 4° C. Pellets were washed once with 70% ethanol and centrifuged at 18,000 rcf for 10 min. The supernatants were discarded, and remaining traces of ethanol were evaporated by air-drying. The pellets were dissolved in 100 µL of sterile deionised water. Nucleic acid extraction for RISA (Microcosms 1-12) was performed with a similar but less time-consuming and less waste-producing protocol due to the large amount of samples. Briefly, a phenol-chloroform extraction (Anderson & MacKay, 1983) was used with an additional bead-beating step using one sterile 2 mm glass bead and bead beating at a setting of 4.0 for 20 s in a FP120 Cell Disruptor (Qbiogene, Carlsbad, USA) prior to the addition of SDS buffer.

PCR, Ribosomal Intergenic Spacer Analysis

PCR amplification of the intergenic spacers was performed using the primer set ITSF/ITSReub (Cardinale et al., 2004). Each 20 µL reaction contained 2.5 mM MgCl₂, 200 µM of each dNTP (Promega, Madison, WI), 10 nM each of primer ITSF (5'- TCG TAA CAA GGT AGC CGT A-3') and ITSReub (5'- GCC AAG GCA TCC ACC-3') (Cardinale et al., 2004), 2 µL of PCR Buffer B (Roboklon, Berlin, Germany), 0.5 U of Taq DNA Polymerase (Roboklon, Berlin, Germany) and 4 µL of a PCR-enhancing mixture (3 M betaine and 1% Tween 20). One nanogram of template DNA was used in each reaction. PCR was performed as described before (Cardinale et al., 2004) using a DNA Engine Tetrad 2 Thermal Cycler

(BioRad, Hercules, USA). Gel electrophoresis was conducted on an Ingeny PhorU gel electrophoresis apparatus using 1x TAE 6% acrylamide gels with 1x TAE buffer at 110 V for 16 h. The gels were stained with 40 mL 1x SYBR Gold (Life Technologies, Carlsbad, USA) staining solution and visualised on a ChemiDoc XRS gel documentation system (BioRad, Hercules, USA).

RISA fingerprinting analysis, multivariate statistical analysis

Quantity One software (BioRad, Hercules, USA) was used for band identification and band matching, with a total of 112 band classes assigned. Following square root transformation, principal coordinate analysis was conducted with PAST using the Bray-Curtis dissimilarity (Hammer et al., 2001). Analysis of similarities (ANOSIM) was performed using PRIMER 6 (Clarke and Gorley, 2006). One-way ANOSIMs of the whole data set were conducted for the parameters “Location (Inoculum)”, “Nitrogen source” and “Sampling time”. To investigate effects of different nitrogen sources on individual subsets of the data for each location, a two-way nested ANOSIM was conducted.

DNA sequencing, assembly, gene prediction and annotation

Sequencing was performed by pair-end sequencing (Illumina Hiseq 2000) at BGI – Beijing Genomics Insititute (Hong Kong, People’s Republic of China). A total of 20,000,000 sequences with a mean read length of 170 nts were obtained per sample. For gene prediction, the software MetaGeneMark (version 2.10, default parameters; Noguchi et al., 2006) was used to predict the open reading frames (ORFs) based on the assembly results. The predicted amino acid sequences were then aligned using the KEGG database (Kanehisa et al., 1997, 2004, 2006) through BLAST (version 2.2.23), and the relevant information was extracted and summarised with self-developed scripts. For Illumina Hiseq 2000 sequencing and data processing, the DNA samples were sequenced following standard pipelines of the Illumina platform. Data filtration was conducted with the following in-house scripts (1) Removal of reads with 3 N bases and removal of reads contaminated by adapters (15 bases overlapped by reads and adapter), (2) removal of reads with 20 bp low quality (20) bases and (3) removal of duplication contamination. The read removal process is simultaneously conducted for the read1 and read2 operation. The resulting data set can be used for subsequent analysis of quality data (Clean Data) (Li et al., 2010). SOAPdenovo (Version 1.0, <http://soap.genomics.org.cn/soapdenovo.html>) was used for the assembly of Illumina Hiseq 2000 sequences and to assemble filtered data. Assembly results with the best N₅₀ contig length were optimized by in-house scripts (Li et al., 2008).

Rarefaction curves of the observed species were established for each sample to analyse the species sampling coverage. Known bacteria, fungi and archaea sequences were extracted

from the Nucleotide database by an in-house script. Filtered reads were mapped to these sequences by SOAPaligner (version 2.21) (Li et al., 2009). Mapped reads were classified in different taxonomic levels (including domain, phylum, class, order, family, genus, and species) and corresponding abundance was also calculated by in-house scripts. All rarefaction curves (data not shown) indicated closeness to saturation in each of the samples, which suggests that biases during the comparative analysis within the metagenomes herein reported were not introduced due to differences in microbial coverage.

Accession numbers

The projects have been registered as umbrella BioProjects at NCBI with the IDs PRJNA222667 [for MGS-AQ(UA)], PRJNA222664 [for MGS-ANC(UA)], PRJNA222665 [for MGS-BIZ(AMM)], PRJNA222666 [for MGS-EIMAX(UA)], PRJNA222663 [for MGS-ANC(AMM)]. These Whole Genome Shotgun projects have been deposited at DDBJ/EMBL/GenBank under the accession numbers AZIG000000000 [for MGS-AQ(UA)], AZIH000000000 [for MGS-ANC(UA)], AZII000000000 [for MGS-BIZ(AMM)], AZIJ000000000 [for MGS-EIMAX(UA)] and AZIK000000000 [for MGS-ANC(AMM)]. The versions described in this paper are versions AZIG0100000 [for MGS-AQ(UA)], AZIH0100000 [for MGS-ANC(UA)], AZII0100000 [for MGS-BIZ(AMM)], AZIJ0100000 [for MGS-EIMAX(UA)] and AZIK0100000 [for MGS-ANC(AMM)]. Abbreviations used for database deposition as follows: MGS, Metagenome Sequences; AQ(UA), Aqaba uric acid (microcosm 16); ANC(UA), Ancona uric acid (microcosm 14); EIMAX(UA), El Max uric acid (microcosm 18); ANC(AMM), Ancona ammonium (microcosm 13); BIZ(AMM), Bizerte ammonium (microcosm 19).

Results

Microbial metabolic activity in microcosms

All microcosms showed a strong increase in respiration rate and emulsification immediately after the start of the experiment corresponding to high aerobic microbial activity (Fig.1). However, total rates of respiration for all uric acid treatments were 8-fold higher at its peak compared to the initial values. These values were higher than the peak values for the ammonium treatments, which, in turn, were 5-fold higher than the initial values. The peak values for respiration were reached on day 12 in the uric acid-containing microcosms (2, 4, 6, 8) and between days 8 and 15 in the ammonium-containing microcosms (1, 3, 5, 7). Following their respective peaks, respiration dropped to base levels after day 18 in uric acid

treatments but remained elevated in the Ammonium treatments due to a continuous influx of ammonium during sampling procedures. The analysis of total respiration showed that uric acid treatments had, on average, a 1.5-fold higher total respiration (620 mg/L O₂) than ammonium treatments (376 mg/L O₂).

The emulsification assay served as a proxy for the activity of hydrocarbon degrading microbes. Emulsification occurred in both treatments after day 4 of the experiment and peaked at day 10 in both cases. Following the maximum, emulsification dropped to 50% of the peak value but remained constant until the end of the experiment. The error bars for all curves are relatively high, which can be explained by temporal shifts in metabolic activity in individual microcosms by ± 2 days.

The photometric quantitation of ammonium and protein concentrations shows an anticyclic trend in both treatments. Ammonium becomes readily available at high concentrations in both the uric acid and ammonium treatments. The ammonium concentration increased immediately after the start of the experiment (with ammonium values not detectable in uric acid treatments at day 0 of the trial) and peaked at day 4. Following the peak, ammonium concentrations rapidly decreased from day 6 to 18 in both microcosms and remained low in uric acid treatments in contrast to NP microcosm which were resupplied with ammonium during sampling procedures. Small residual amounts of ammonium could be detected in both treatments at the end of the experiment.

Conversely, protein concentrations, which served as a proxy of total biomass, increased in both treatments from day 6 of the experiment onward. However, the concentrations dropped after day 18 in all uric acid treatments but increased constantly in Ammonium treatments.

Dynamic changes in microbial community composition

Microbial community changes and composition were analysed using Ribosomal Intergenic Spacer Analysis (RISA) and Pair-end sequencing. Figure 2 shows the results of the Principal Coordinate Analysis (PCoA). The individual fingerprints used for this analysis are presented in the Supplementary Figure 1. The PCoA plot shows no distinctively separated clusters, indicating that the microbial communities in the microcosms are essentially similar over the course of the experiment. This similarity can be explained by the choice of samples, which come from sites with a legacy of hydrocarbon pollution as well as from the same body of water and climate zone. The data points for both the NP and UA treatments overlap for all sampling sites, indicating that the treatments ultimately produce identical microbial communities. This result is supported by the ANOSIM results presented in Supplementary Table 1a.

Statistically significant similarities could be detected when analysing the factor “nitrogen source”. High similarities (R value -0.156) were detected when using two-way ANOSIM to investigate the similarities between the UA and NP treatments for samples at each location. Data points from individual locations, on the other hand, show subtle differences separating the Ancona and Aqaba samples, although there is some degree of congruence for the Northern African samples; however, the ANOSIM analysis did not indicate significant similarities between the microbial communities from individual locations (Supplementary Table 1b). Most obvious is the separation of the early sampling points (days 0 – 10) in the upper right section of Figure 2 from the bulk of the late sampling points (days 12 – 28). This trend coincides with the peaks of both emulsification (days 10 – 12 in both treatments) and protein production (days 12 – 20), which is strongly supported by the ANOSIM results presented in Supplementary Table 1c. Significant similarities could be detected in the RISA profiles for samples taken between days 4 and 8, as well as for samples taken between days 12 to 28.

Metagenome analysis of marine oil-degrading consortia: bacterial diversity and composition

DNA isolated from each microbial community sampled from microcosms 13-20 at the day 21 was sequenced using Illumina HiSeq 2000. The data and annotation features resulting from the assemblies of each sample are shown in Supplementary Table 2. Three samples from uric acid enrichments and two samples from ammonium enrichments were chosen to ensure that all sampling sites were covered and to allow a direct comparison of at least one site. The samples for sequencing were selected based upon the results of RISA and whether the ANOSIM analysis showed high similarities between the ammonium and uric acid treatments for a particular site, omitting redundant samples. However, to confirm these similarities, both treatments of the Ancona microcosms were analysed. The results of this analysis, presented in Figure 3, show compositional similarities between the uric acid and ammonium treatments of the Ancona sample when the 16S rRNA partial gene sequences obtained in the non-assembled Illumina reads were surveyed. For this purpose, we used only sequences with a length of >100 nucleotides. Similarities could be observed between the uric acid treatments of the Ancona and El Max site (microcosms 14, 18), as well as between the ammonium treatments of the Ancona and Bizerte sites (microcosms 13, 19). The Aqaba uric acid treatment (microcosm 16) showed relatively little similarity to any other treatments and was characterised by a high percentage of *Pseudomonadaceae*. Table 2 indicates that *Pseudomonas* spp. were the most abundant microbes in every treatment. They were more abundant in the uric acid treatments, and their percentage increased along a North-South gradient for these treatments. Specialised hydrocarbon-degrading microbes belonging to the genera *Alcanivorax*, *Cycloclasticus*, *Oleiphilus*, *Oleispira* and *Thalassolituus* were detected in all treatments. Most prominent among these microbes were the *Alcanivorax* spp., which were more abundant after NP (from

4.99 to 16.51% total reads) rather than uric acid treatments (from 0.44 to 3.94% total reads). The analysis of the metagenomic data showed that most *Alcanivorax*-like DNA sequences belonged to *Alcanivorax borkumensis*.

In parallel to this result, the ammonium treatments contained a higher percentage (~9.0-fold increment) of *Alphaproteobacteria*, consisting mostly of *Roseovarius* and *Sulfitobacter* spp. Conversely, uric acid treatments were more likely to contain *Marinobacter* spp. (up to ~2.0-fold) and *Halomonas* spp. (up to ~3.8-fold). Direct comparison of both Ancona treatments showed a 1.5-fold higher percentage of *Halomonas* spp. and a 2-fold higher percentage of *Marinobacter* spp. but an 8-fold lower abundance of *Alphaproteobacteria* in the uric acid treatments. Furthermore, the percentage of *Psychrobacter* spp. in the Ancona uric acid treatment was more than 30-fold elevated in comparison to the corresponding ammonium treatment. The percentage of *Pseudomonas* spp. and *Alcanivorax* spp., however, remained similarly high in both ammonium and uric acid treatments.

Genomic signatures associated with uric acid metabolism.

In the metagenomes we identified 176,914 potential protein-coding genes (cut-off of ≥ 20 amino acid-long sequences), with El Max containing the highest number (61,277). Functional assignment of the predicted genes was made on the basis of BLASTP analysis against a reference dataset for Clusters of Orthologous Groups (COGs) and Kyoto Encyclopedia of Genes and Genomes (KEGG) assignments. Circa 63.1-67.9% could be assigned to COG, and 54.0-63.7% could be assigned to KEGG pathways (Supplementary Table 2). On average, 6.6 to 12.8 genes belonged to COG and from 5.6 to 11.2 genes belonged to KEGG, depending on the sample.

All annotated data sets were examined for gene homologs involved in uric acid utilisation pathways. In both eukaryotes and prokaryotes, two urate hydroxylases (EC 1.7.3.3 and EC 1.14.13.113) have previously been shown to convert uric acid into 5-hydroxyisourate, which is ultimately broken down into urea and finally into ammonium. However, no genes with similarity to genes coding for either of these two enzymes could be detected in any of the datasets. In contrast, genes coding for enzymes that catalyse downstream reactions of uric acid utilisation, including allantoinase, allantoicase, allantoate amidohydrolase, ureidoglycolate hydrolase, 5-hydroxyisourate hydrolase, and 2-oxo-4-hydroxy-4-carboxy-5-ureidoimidazoline (OHCU) decarboxylases, could be found (Supplementary Table 3a - e). The percentages of genes involved in uric acid metabolism were up to 3-fold higher in UA enrichments; they were particularly enriched (~0.11% protein-coding genes) in Aqaba uric

acid treatments while being particularly depleted in the Bizerte ammonium treatments (0.02% total protein-coding genes). However, we note that the NP-treated microcosm from Ancona port also contained a significant amount of those genes (0.06% total protein-coding genes). Binning analysis by BLASTP search further suggests that at least 15 (for Aqaba – uric acid), 11 (for Ancona - ammonium), 14 (for Ancona – uric acid), 2 (for Bizerte - ammonium) and 15 (for El Max- uric acid) distinct microorganisms may be potentially involved in the metabolism of uric acid and/or its degradation products. A direct comparison between the Ancona ammonium and uric acid treatments revealed that only 4 out of 25 of such microorganisms were shared, suggesting that most of the uric acid-utilization microbes are distinct to those using ammonium. The possibility that the microbes in the Ancona - ammonium enrichment possess genes encoding pathways for uric acid utilization could not be ruled out; however, the data presented in this study suggests that in the presence of uric acid such proteins do not support bacterial growth and thus under these conditions other microbes are enriched. Supplementary Table 3a - e show genes for uric acid catabolism with similarity to the genes found in members of *Aeromonadaceae*, *Halomonadaceae*, *Pseudomonaceae* and *Rhodobactereaceae* within the three metagenomes from the uric acid enrichments, whereas the metagenomes from the ammonium enrichments predominantly contained uric acid catabolism genes from *Alteromonadaceae*, *Oceanospirillaceae* and *Rhodobacteraceae*. To further investigate which of these potential degrading organisms may play a role in uric acid metabolism, a cultivation-approach was used.

Isolation of uric acid-degrading bacteria

Uric acid-degrading bacteria were picked based upon the formation of a halo around the colony and purified to a single colony by triple repetitive plating on the same type of agar. As mentioned above, all isolates were successfully grown in ONR7a liquid media containing uric acid as sole source of nitrogen and carbon.

The 16S rRNA gene sequencing results (Supplementary Table 4) revealed that 44 of the 47 isolates belonged to the genus *Halomonas*. Two further isolates from the Bizerte enrichment belonged to the genus *Marinobacter*, and one isolate from the El Max enrichment was identified as a member of the genus *Pelomonas*. Phylogenetic analysis (Figure 4) showed that isolates were closely related to *Halomonas alkaliphila*, *H. cupida*, *H. salfondinae*, *H. titanicae*, *H. venusta* and *H. ventosae*. With a few exceptions, the majority of the isolates showed location specificity. Most of the isolates from Ancona were closely related to *H. alkaliphila*; however, the isolates from Aqaba were predominately related to *H. venusta* and *H. hydrothermalis*. The majority of isolates from the Bizerte enrichments clustered with *H. salfondinae* and *H. pacifica*. Finally, all but two isolates from the El Max/Alexandria enrichments were distantly related to *H. cupida* and *H. shangliensis*.

The analysis of the metagenomic data mentioned above showed that genes with a high similarity to genetic material from *Halomonas* spp. were present in all five metagenomes, albeit in low percentages, ranging from 1.4% to 6.1% as revealed by abundance of SSU rRNA signatures. The percentage of *Halomonas*-like sequences was elevated in the uric acid enrichments.

Discussion:

Comparison of uric acid and ammonium as nitrogen source in sediment microcosms

Ammonium concentrations were measured as a proxy for nitrogen uptake in ammonium treatments and uric acid utilization in uric acid treatments, as it was the sole nitrogen source in the ammonium treatments but also an intermediate of uric acid conversion in the uric acid treatments. The ammonium treatments did show a relatively low ammonium concentration at the beginning of the experiment with a subsequent increase. This result indicates a general problem with ammonium and phosphate additions to seawater. As seawater is a saline solution, the addition of further ions may lead to precipitation. The addition of phosphate to artificial seawater leads to precipitation of calcium or magnesium ammonium phosphate or phosphates of trace elements that could affect microbial growth (Atlas, 1975). In this study, phosphate was added to artificial seawater that contained Fe and other trace elements that precipitate with both ammonium and phosphate (e.g. as a ferrous ammonium phosphate). The secondary increase of ammonium concentrations could be explained by consumption of both iron and phosphate for microbial growth and a subsequent release of ammonium into the medium.

The ammonium concentrations in the uric acid treatments increased rapidly to approximately 270 mg /L, which corresponds to 80% of the total uric acid nitrogen introduced into the microcosms. As uric acid was the only nitrogen-containing substance in these microcosms, it stands to reason that it is quickly and almost entirely converted into ammonium. Photometric measurements of nitrate showed no significant increase in its concentrations (data not shown) in either treatment throughout the course of the experiment; it can therefore be assumed that uric acid was entirely transformed into ammonium.

Several recent studies have focused on the use of uric acid as a stand-alone nitrogen source or as a bioremediation strategy in combination with other substances. Koren and co-authors (2003) described the use of pure uric acid in hydrocarbon-polluted sediments and succeeded in isolating a strain of *Acinetobacter baumannii* that directly used uric acid for hydrocarbon degradation. In a further study, Knezevich et al. (2006) described the application of guano as a complex source of uric acid and a soluble nitrogen source for bioremediation using both *A. borkumensis* and mixed oil-degrading microbial communities in an open system. While the uric acid-fed communities showed rapid growth, with cell numbers of up to 10^8 cells/mL and degradation of 70% of total oil hydrocarbons, the cell numbers in the NPK-fertilised experiment remained two orders of magnitude lower and oil degradation was negligible. The data presented in our study is in concordance with a number of previous publications (Knezevich et al., 2006; Nikolopoulou and Kalogerakis, 2008; Nikolopoulou et al., 2013) which report enhanced microbial growth similar to what could be observed for the uric acid

treatments. Microbial communities grew more rapidly and developed cell numbers three orders of magnitude higher than comparable treatments with soluble nitrogen sources (Nikolopolou et al., 2013). Furthermore, aliphatic hydrocarbon degradation and PAH degradation were both significantly accelerated in presence of uric acid (Nikolopoulou and Kalogerakis, 2008; Nikolopoulou et al., 2013). Comparison of growth rates furthermore indicated that addition of molasses and rhamnolipids may speed up the microbial growth in the experiment, but is not stringently required for oil-degrading microbes to reach comparable cell numbers (Nikopoulou and Kalogerakis, 2008).

Microbial population dynamics of microbial communities utilising uric acid

Two recent studies have investigated the use of fertilisers containing uric acid by oil-degrading microbial communities (Knezevich et al., 2006; Nikopoulou et al., 2013). The findings from both studies indicate the presence of *Alcanivorax borkumensis* in enrichments or microcosms supplied with uric acid. In the study of Knezevich et al. (2006), *Alcanivorax* spp., *Alteromonas* and *Halomonas* spp. were enriched and isolated. The results of our study also show abundance of *Alcanivorax* spp. in all samples. However, all uric acid enrichments showed a significantly lower abundance of these microbes in uric acid enrichments compared to ammonium enrichments. Despite the extensive research into the *Alcanivorax* genus, only one strain able to use uric acid as nitrogen source has been described to date (Knezevich et al., 2006). The data from our study suggests that the degradation of uric acid could be performed by other members of the microbial consortium present in uric acid-amended hydrocarbon spiked enrichments. Uric acid enrichment also showed a high conversion rate into ammonium and elevated respiration rates in contrast to ammonium enrichment from days 0 to 9. However, the use of uric acid from days 4 to 6 did not lead to an increase in protein or biomass production. This result could be due to the extremely low CN ratio of uric acid, which limits the total amount of biomass microbes can produce when grown on uric acid alone. Figures 1 a and 1 c show that both emulsification and protein production begin to increase from day 6 onward. At this time, large quantities of uric acid have already been turned over into ammonium.

Therefore, microbial hydrocarbon degradation using uric acid as a nitrogen source could be described as a two-stage process involving the primary conversion of uric acid into ammonium and a secondary stage in which marine hydrocarbonoclastic microbes use ammonium as the primary nitrogen source.

This two-stage process can also be observed in Figure 2, as most DNA fingerprints converged in a large cluster towards the end of the experiment. Similar trends have been observed in previous studies with similar experimental conditions (Gertler et al., 2009a,b; 2012).

Microbial climax communities with high similarity formed within 14 days of the experiment, yet a subtle location specificity of the communities can be observed. While these differences can be observed within the DNA fingerprints from Aqaba, Ancona, Bizerte, and El Max (Figure S1), they are not statistically significant, as shown by multivariate analysis. The results of ANOSIM analysis indicated that the microbial communities showed significant similarities at two stages during the experiment, within days 4 to 8 and from day 12 to the end of the experiment, supporting the hypothesis of a two-stage process. The formation of a highly similar climax community within 14 to 21 days in different locations has been observed in a previous study involving the Irish Sea, the North Sea and the Mediterranean (Gertler et al., 2012). While obligate marine oil-degrading bacteria, most notably *Alcanivorax* species, form the backbone and most active hydrocarbon degraders of these consortia (Gertler et al., 2009a,b), there is a certain degree of variability in the composition of other consortium members. In contrast to the intense clustering, the data points from days 0 to 6 of the experiment are located outside of this cluster, indicating a community change. The ANOSIM results (Supplementary Table 1c) indicate that within this period a highly similar community that differs from the oil-degrading climax community established after day 12 formed in all microcosms.

Metagenome sequencing was conducted for five samples from Aqaba, Ancona, Bizerte and El Max (Microcosms 13, 14, 16, 18, 19), which were selected after the RISA fingerprinting. One sample from each Aqaba, Bizerte and El Max was chosen to cover location-specific characteristics of the sediment, while two samples from Ancona (one for the uric acid and one for the ammonium treatment) were selected to directly compare the effects of both treatments in one location. The sample material was based upon single biological replicate to preserve maximum coverage and sequencing depth as well as for other technical reasons. This procedure however does not take in account possible culture variability. A solution to this problem could be pooling of replicate samples, which reduce this bias, it would also reduce the overall coverage. Multiplexing of individual samples in a single sequencing run on the other hand bears the risk of adapter misidentification while true replication of samples (e.g. for a time series of microcosm samples) is still financially prohibitive. To ensure the validity of metagenome data in this study, rarefaction curves were established for each of the metagenome data sets and that were close to saturation and exhibited low standard deviation. Metagenome sequencing confirmed the composition of the microbial consortia to be dominated by *Pseudomonas* spp. and to some extent by *Alcanivorax* spp. in all cases. *Pseudomonas* spp. were the most abundant species in all consortia, while the Ancona - uric acid, Ancona - ammonium and Bizerte - ammonium samples contained 4 - 16% of *Alcanivorax* spp. -related 16S rRNA reads in the whole SSU rRNA dataset. The samples from El Max and Aqaba showed an overall similar species composition to the other samples,

but a significantly higher percentage of *Pseudomonas* spp.. This data is only apparently in contrast to previous studies conducted with similar microcosms that showed a significantly higher percentage of *Alcanivorax* spp. (Gertler et al., 2010a, 2012) considering that the current study used the sediment rather than seawater as the inoculum. Furthermore, the genomes of *Pseudomonas* spp. are known to possess six or more rRNA operons, in contrast to the three rRNA operons present in the genome of *Alcanivorax borkumensis*. This difference in the number of operons could result in an overestimation of the numbers of pseudomonads from 16S rRNA-related reads affiliated with *Pseudomonas* spp. in comparison to *Alcanivorax* spp. Most notably, the composition of uric acid and ammonium-based enrichments from Ancona proved to be similar, as both communities were enriched from identical inocula and effectively supplied with ammonium, in the latter case directly added to the medium while in the former case deriving from uric acid conversion. In addition to the clustering of uric acid and ammonium enrichment fingerprints in Figure 2, these findings indicated that given identical experimental parameters, uric acid supplementation could effectively produce identical oil-degrading consortia due to the quick conversion of uric acid into ammonium. This suggestion is statistically supported by nested two-way ANOSIM analysis of the RISA location data sets.

Uric acid conversion and uric acid-degrading isolates

The analysis of the five metagenomes annotated within this study remained inconclusive regarding uric acid metabolism. Neither of the genomes showed any genes coding for uricase enzymes. However, urate oxidase activity has also been detected in other classes of enzymes, e.g., a laccase from a *Lysobacter* species (Tamaki et al., 2010). It is therefore possible that the actual genes responsible for urate oxidation in the analysed samples could not be detected because they were annotated differently. Genes coding for ureidoglycolate hydrolase (EC 5.3.19), 5-hydroxisoureate hydrolase (EC 3.5.2.17), allantoinase (EC 3.5.3.4.), allantoinase (EC 3.5.2.5) and OHCU decarboxylase (EC 4.1.1.-), on the other hand, were identified within the genomes. These genes are involved in uric acid metabolism and appeared three times more frequently in the metagenomes of the uric acid-based enrichments in relation to the total number of sequences. Furthermore, uric acid catabolism genes in ammonium-based enrichments were mostly similar to the genes of *Oceanospirillaceae* and *Rhodobacteriaceae*, whereas the uric acid catabolism genes in the uric acid-based enrichments were predominantly similar to genes found in members of *Pseudomonadaceae* and *Halomonadaceae*. The analysis of 16S rRNA data in the metagenomes showed comparable percentages of *Halomonas* spp. and *Pseudomonas* spp. in the uric acid treatments compared to ammonium treatments (Figure 3, Table S3). However, the direct comparison of the ammonium and uric acid treatments for the Ancona site revealed higher percentages of *Halomonas* spp. in the presence of uric acid and comparable percentages of *Pseudomonas* spp. in both treatments. The evidence from the metagenome analysis, therefore, indicates that the *Halomonas* spp. rather than the *Pseudomonas* spp. are the main utilisers of uric acid in this experiment.

In addition to the metagenome analysis, major degraders of uric acid were identified by cultivation and isolation at various points during the experiment. Ninety-three per cent of the isolates belonged to the genus *Halomonas*, a ubiquitous, oligotrophic, highly salt-tolerant and metabolically versatile group of microbes (Llamas et al., 2006; Mata et al., 2002). *Halomonas* strains capable of biopolymer and bioemulsifier production have been previously cultured (Gutierrez et al., 2013). Members of this genus with hydrocarbon-degradation capabilities have also been isolated (Mnif et al., 2009) and detected in oil-degrading consortia using culture-independent techniques (Kleinstaub et al., 2006; Gutierrez et al., 2011). In contrast, the strains grown in this study were isolated in absence of hydrocarbons with uric acid as sole source of carbon, nitrogen and energy. While the results of this approach are subject to the limitations of our cultivation techniques, the presence of similar *Halomonas* species is confirmed by their detection within the metagenomic data of all five samples mentioned

above. Sequences of *Halomonas* spp. made up between 1.4 and 6.1% of the total amount of rRNA gene sequences.

The *Halomonas* isolates displayed a certain location specificity, with most isolates from each sampling site clustering together independently of experiment time and nitrogen source. This clustering could be because the genus *Halomonas* is understudied, and relatively few genomes and DNA sequences of few isolates are present in the databases. Another possibility is that the metabolic versatility of this genus (Mata et al., 2002) may have lead to the selection of specific *Halomonas* strains with uricase activity adapted to the unique abiotic parameters at the specific sampling site, suggesting a biogeographic distribution of this genus.

Prospects for uric acid application in bioremediation techniques

This study investigates the population dynamics of oil-spiked marine sediment microcosms supplied with ammonium or uric acid in equal amounts and C:N:P ratios using identical crude oil batches. These procedures enable the direct comparison of microbial communities utilising either nitrogen source. The use of uric acid as a nitrogen source has been extensively discussed, experimented on and even patented, yet the mechanism of uric acid utilisation in an actual microbial community remains unknown. In our study, marine microbial communities dominated by *Pseudomonas* spp. and *Alcanivorax* spp., the most prominent, ubiquitous and competitive obligate hydrocarbon marine microbes, have been found. It was shown that uric acid could be quickly transformed into ammonium by members of the *Halomonas* genus. This observation was made for sediments originating from four independent sites located around the Mediterranean (Ancona, Bizerte, El Max) and the Red Sea (Aqaba), which implies that a similar pattern likely occurs on a more global scale.

As each uric acid molecule contains four nitrogen atoms, such a nitrogen source can be applied in lower molarities compared to ammonium. While problems with leaching can be successfully overcome by paraffin coating (urea) or the use of slow-release fertilisers (ammonium, nitrate) (Gertler et al., 2009), these treatments require industrial processes that greatly reduce the cost-efficiency of bioremediation. Uric acid, on the other hand, can be found in concentrations of 3-6% (w/w) in poultry faeces, making up 60% of the total nitrogen. The global production of up to 50 billion chickens produces 50 million tons of faecal matter every year with an estimated content of 1.5 to 3 million tons of uric acid (Nahm, 2003). Poultry litter is currently used as an agricultural fertiliser after sterilisation through heat treatment and drying and are well established on the fertiliser market. This study has shown that uric acid can be quickly transformed into ammonium, which in turn is utilised by highly efficient oil-degrading microbes in marine sediments. This process was observed in sediment collected from various locations in different water bodies without the addition of allochthonous

microbial communities, suggesting that cost-effective uric acid-based biostimulation techniques can be successfully achieved in a wide range of marine sediments.

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Figures

Figure 1a – d: Observations of protein and ammonium concentrations, and microbial respiration in microcosms. **Figure 1a** and **c** show data collected from uric acid enrichments (Microcosms 1, 3, 5, 7), and **1b** and **d** from ammonium enrichments (Microcosms 2, 4, 6, 8). Filled circles represent respiration rates; open triangles indicate emulsification rates. Filled squares indicate total protein concentrations, whereas open diamonds show ammonium concentrations. All data points represent the mean of triplicates from each sampling point. Bars represent standard error.

Figure 2: Principal coordinate Analysis of RISA fingerprints. Light colours indicate ammonium enrichments; dark colours indicate uric acid enrichments. Blue circles represent enrichments from Ancona, Italy (Microcosms 1, 2); red squares show samples from Aqaba, Jordan (Microcosms 3, 4). Grey triangles show samples from El Max, Alexandria, Egypt (Microcosms 5, 6), whereas green diamonds represent enrichments from Bizerte Lagoon, Tunisia (Microcosms 7, 8). Numbers correspond to the sampling day of each data point.

Figure 3: Relative abundances of microbial families within annotated metagenomes based on the 16S small subunit rRNA data taken on day 21 from the following samples: Ancona Uric Acid (Microcosm 14), Ancona Ammonium (Microcosm 13), Bizerte Ammonium (Microcosm 19), Aqaba Uric Acid (Microcosm 16) and El Max Uric Acid (Microcosm 18). Only lineages with abundance of reads >1 % (SSU rRNA tags) are shown. Members of families belonging to Actinobacteria and Firmicutes were combined.

Figure 4: Rooted phylogenetic tree of 16S rRNA gene sequences from *Halomonas* spp. isolates grown on ONR7 agar containing uric acid. The prefix codes B (diamonds), X (triangles) Q (squares) and A (circles) represent the experimental sites Bizerte, El Max, Aqaba and Ancona, respectively. The prefix codes U and N represent the experimental treatment: 'UA' or 'NP,' respectively. The colour coding of symbols corresponds to the colours used in Figure 2. Sequences were clustered by maximum parsimony according to the recommendation of the MODELTEST software (Posada & Crandall, 1998). The 16S rRNA gene sequences of the most closely related sequences from type strains of *Halomonas* species were acquired from the NCBI nucleotide database. The evolutionary history was inferred using the Maximum Parsimony method (Eck & Dayhoff, 1966). The bootstrap consensus tree inferred from 500 replicates is taken to represent the evolutionary history of the taxa analysed (Felsenstein, 1985). Branches corresponding to partitions reproduced in less than 50% bootstrap replicates are collapsed. The MP tree was obtained using the Close-Neighbor-Interchange algorithm (Nei & Kumar, 2000) with search level 3 (Felsenstein, 1985; Nei & Kumar, 2000) in which the initial trees were obtained with the random addition of sequences (10 replicates). The tree is drawn to scale, with branch lengths calculated using the average pathway method (Nei & Kumar, 2000). Scale bar reflects the total nucleotide mismatch numbers in the whole sequence. All positions containing gaps and missing data were eliminated from the dataset (Complete Deletion option). There were a total of 952 positions in the final dataset, out of which 97 were parsimony informative. Phylogenetic analyses were conducted in MEGA4 (Tamura et al., 2007).

Table 1: Composition of microcosms established in this study.

Table 2: Percentages of microbial genera within metagenomes of samples from day 21 from the following microcosms: Ancona Uric Acid, Ancona Ammonium, Bizerte Ammonium, Aqaba Uric Acid and El Max Uric Acid. The data are based on the frequency of the appearance of reads of SSU rRNA. Genera were combined into families, classes or phyla where necessary.

Supporting Information

Supplementary Figure 1: RISA fingerprinting profiles used for the multivariate analysis presented in Figure 2 of the main text. Full DNA profiles of two out of three microcosm replicates were used to avoid mismatching in the band matching required for statistical analysis. All RISA fingerprints were run using the O'gene Ruler Plus (Thermo Scientific, Lutterworth, UK) as indicated by the letter M under the marker lanes. Gel images show DNA fragments of 300 to 5000 bp in length. Numbers under individual lanes represent sampling days. Fingerprints from microcosm replicates are displayed as blocks of profiles with increasing sampling time (0 to 28 days).

Supplementary Table 1a: ANOSIM results of bacterial community RISA fingerprinting profiles on the basis of Bray-Curtis distance measure, showing the univariate partition of factors "Location", "Nitrogen source" and "Sampling day" and the multivariate partition of factors "Location" within the factor "Nitrogen source". The R-value was calculated based on the Null hypothesis of no similarity between samples. The values highlighted in bold are statistically significant ($P < 0.05$).

Supplementary Table 1b: R-values derived from ANOSIM pairwise comparisons of factor "(sampling) Location" values using Bray-Curtis values. The R-value was calculated based on the Null hypothesis of no similarity between samples. The values highlighted in bold are statistically significant ($P < 0.05$).

Supplementary Table 1c: R-values derived from ANOSIM pairwise comparisons of factor "Sampling day" values using Bray-Curtis values. The R-value was calculated based on the Null hypothesis of no similarity between samples. The values highlighted in bold are statistically significant ($P < 0.05$).

Supplementary Table 2a: Data statistics for samples as obtained by Illumina sequencing

Supplementary Table 2b: Data statistics for the best assembly results

Supplementary Table 2c: Gene prediction results

Supplementary Table 3a: Annotation of the metagenomic data from the Aqaba UA sample from day 21, displayed in Figure 3 of the main text. The subset consists of 30 genes involved in uric acid metabolism from a total of 26,866 ORFs identified in the metagenome. Gene sequences with similarity to *Halomonas* spp. are printed in bold.

Supplementary Table 3b: Annotation of the metagenomic data from the Ancona UA sample from day 21, displayed in Figure 3 of the main text. The subset consists of 15 genes involved in uric acid metabolism from a total of 27,893 ORFs identified in the metagenome. Gene sequences with similarity to *Halomonas* spp. are printed in bold.

Supplementary Table 3c: Annotation of the metagenomic data from the Ancona NP sample from day 21, displayed in Figure 3 of the main text. The subset consists of 18 genes involved in uric acid metabolism from a total of 32,180 ORFs identified in the metagenome. Gene sequences with similarity to *Halomonas* spp. are printed in bold.

Supplementary Table 3d: Annotation of the metagenomic data from the Bizerte NP sample from day 21, displayed in Figure 3 of the main text. The subset consists of 6 genes involved in uric acid metabolism from a total of 28,698 ORFs identified in the metagenome. Gene sequences with similarity to *Halomonas* spp. are printed in bold.

Supplementary Table 3e: Annotation of the metagenomic data from the El Max UA sample from day 21, displayed in Figure 3 of the main text. The subset consists of 29 genes involved in uric acid metabolism from a total of 61,277 ORFs identified in the metagenome. Gene sequences with similarity to *Halomonas* spp. are printed in bold.

Supplementary Table 4: Closest relatives of 16S rRNA gene sequences of uric acid-degrading isolates. The first letter of the isolate code designates the sediment sampling location (A – Ancona; Q – Aqaba; X – El Max; B – Bizerte), and the second letter indicates the nitrogen source (N – Ammonium; U- Uric Acid). The first number in the isolate code represents the sampling date, and the last number indicates the individual isolate.

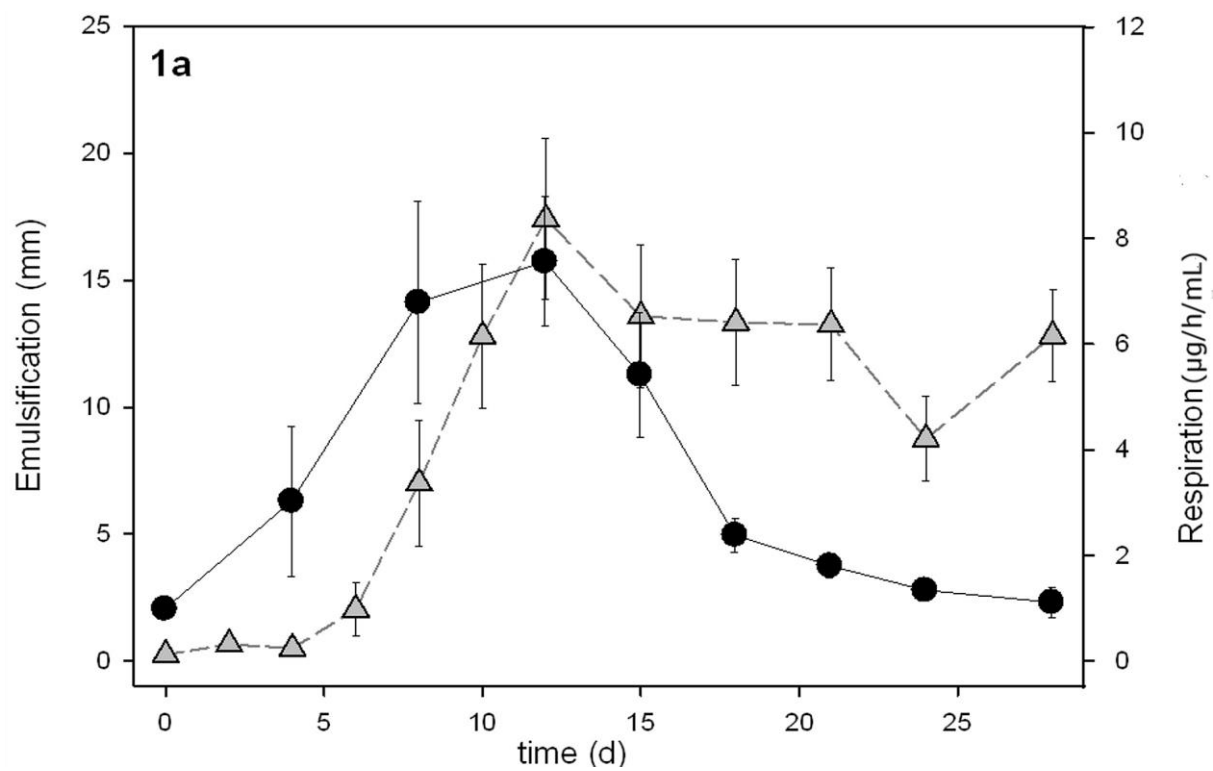


Figure 1a.

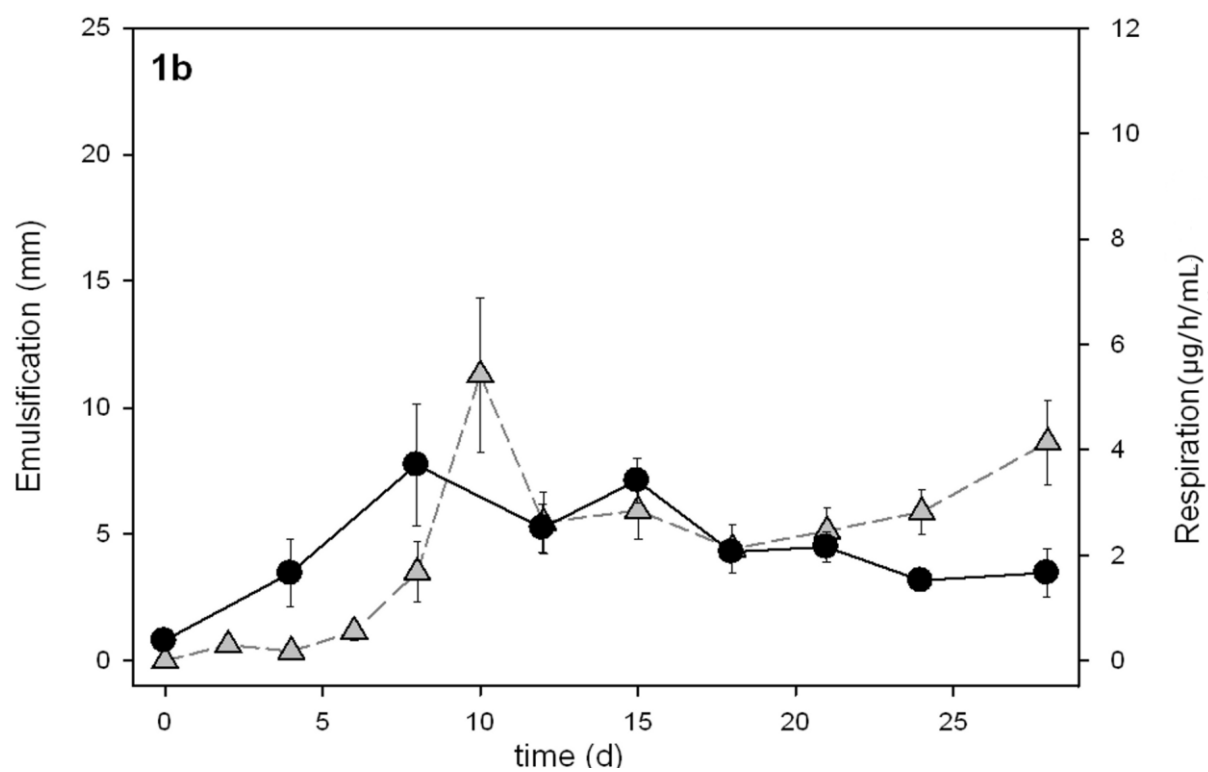


Figure 1b.

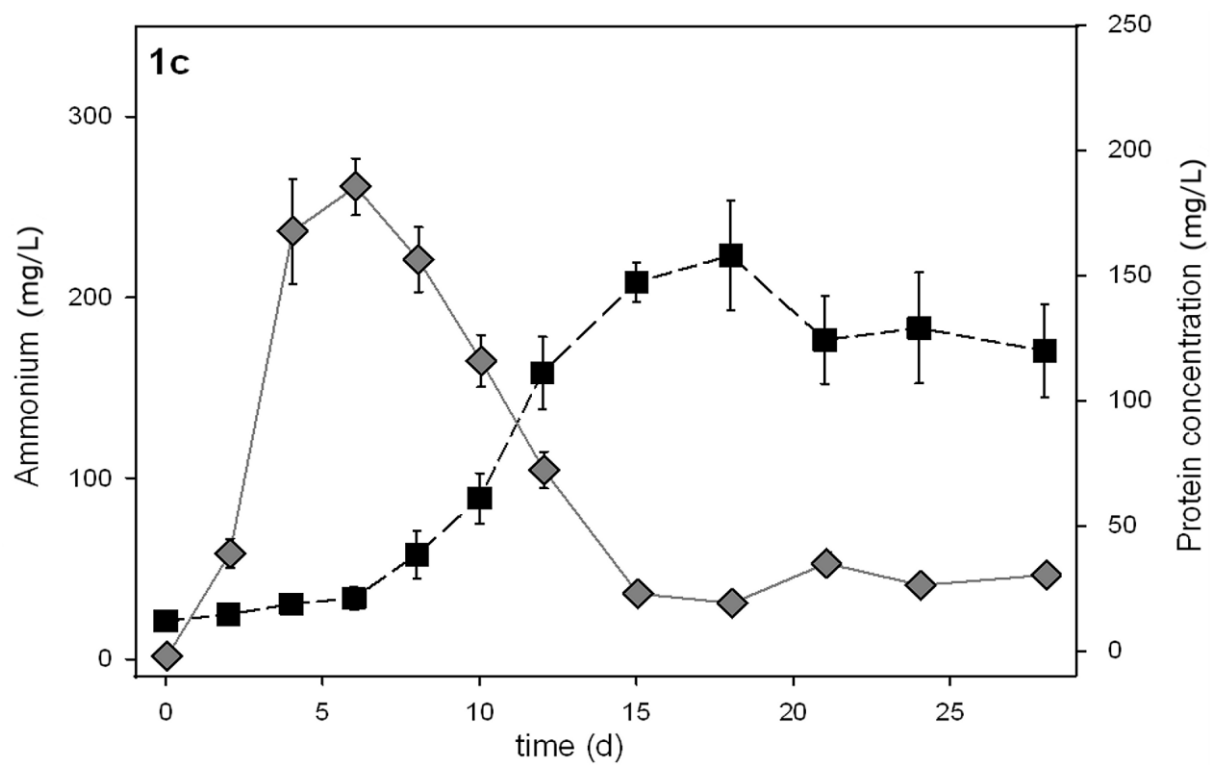


Figure 1c.

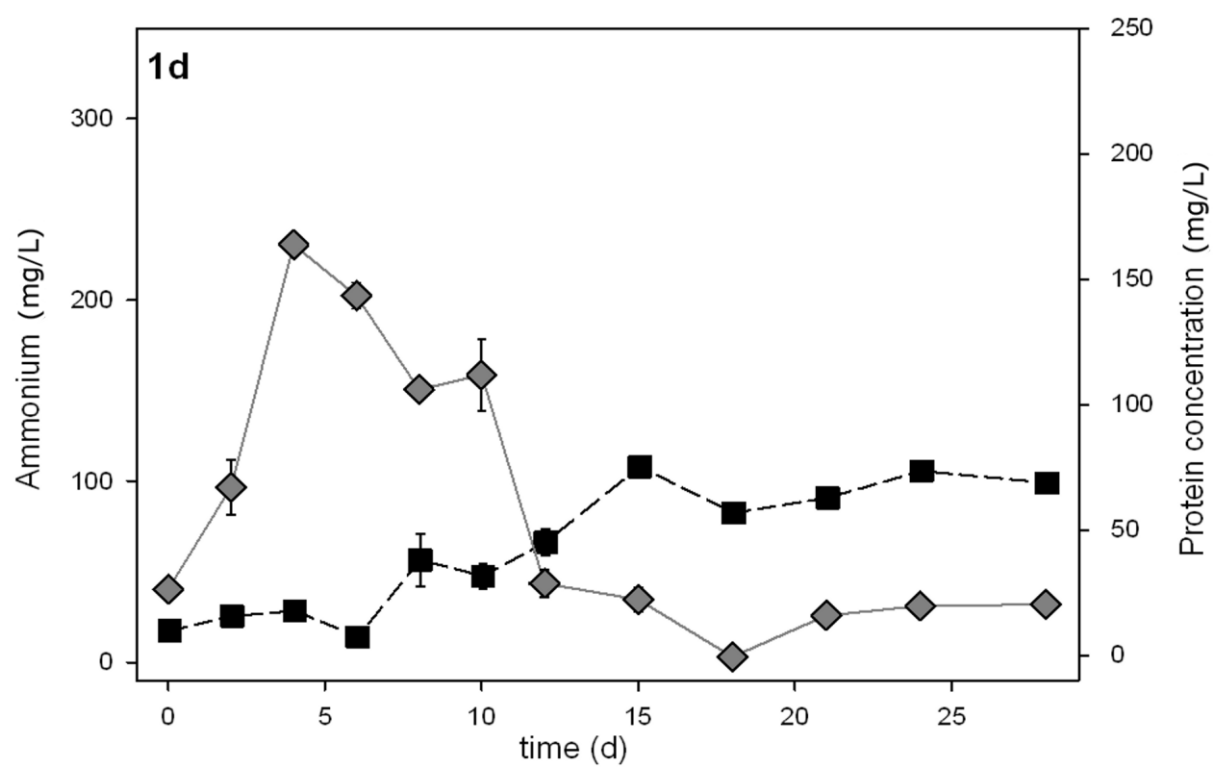


Figure 1d.

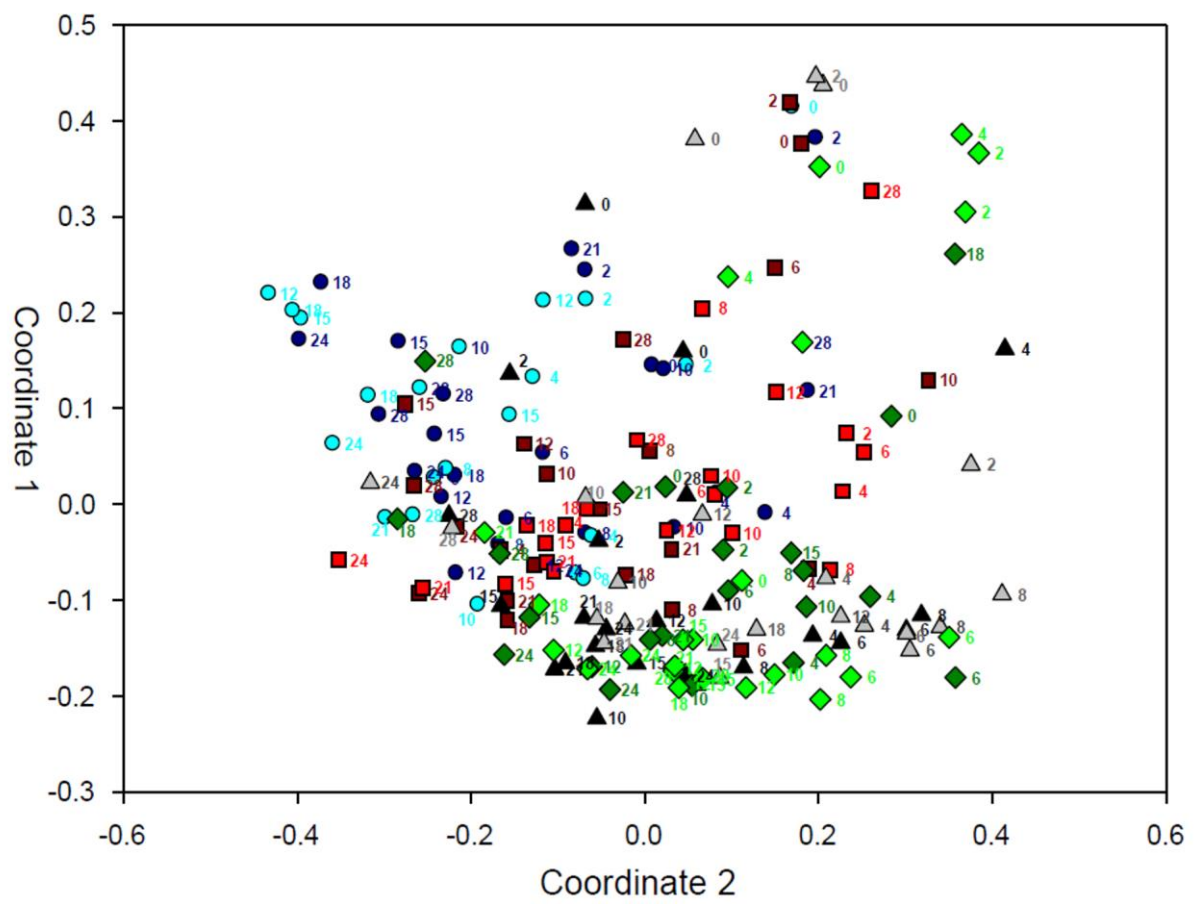


Figure 2.

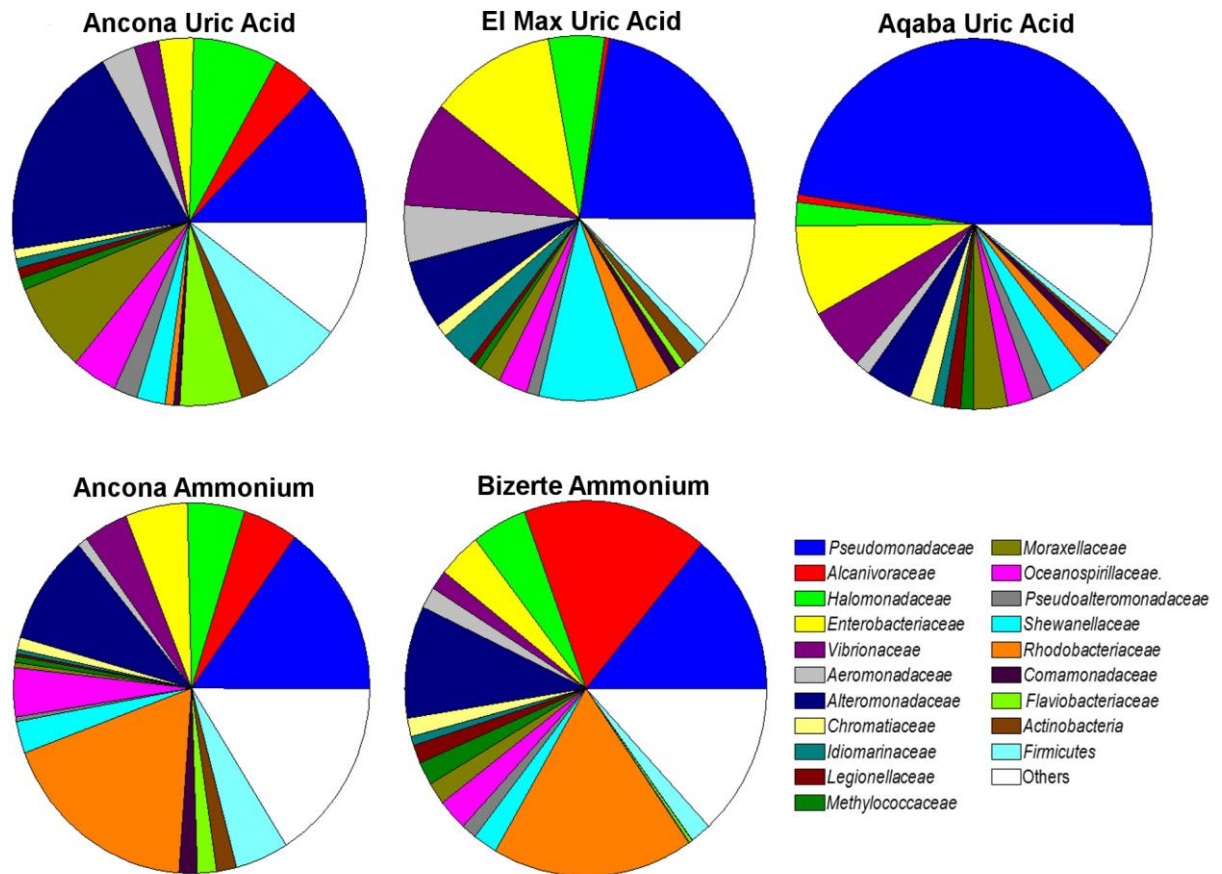


Figure 3.

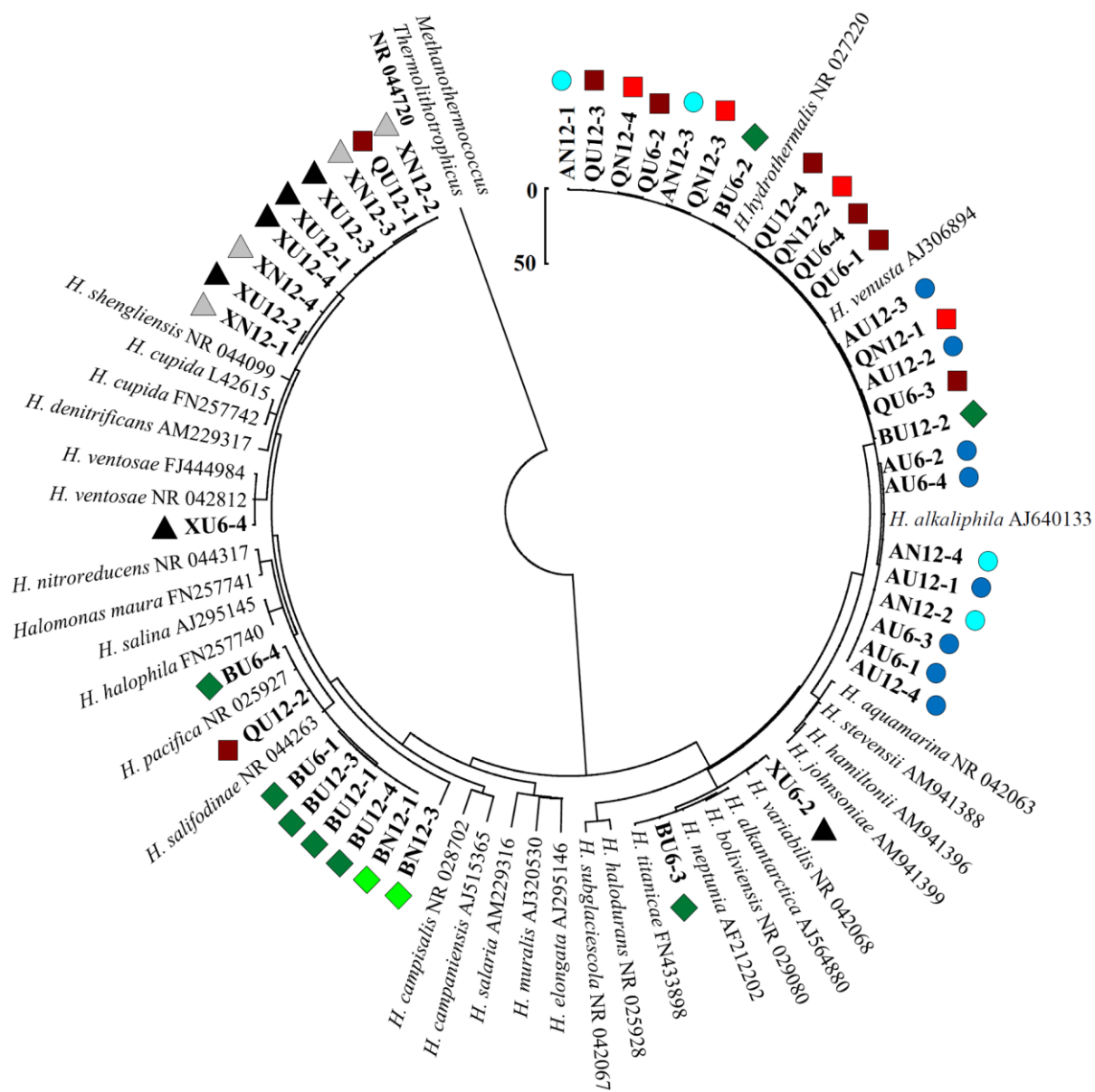


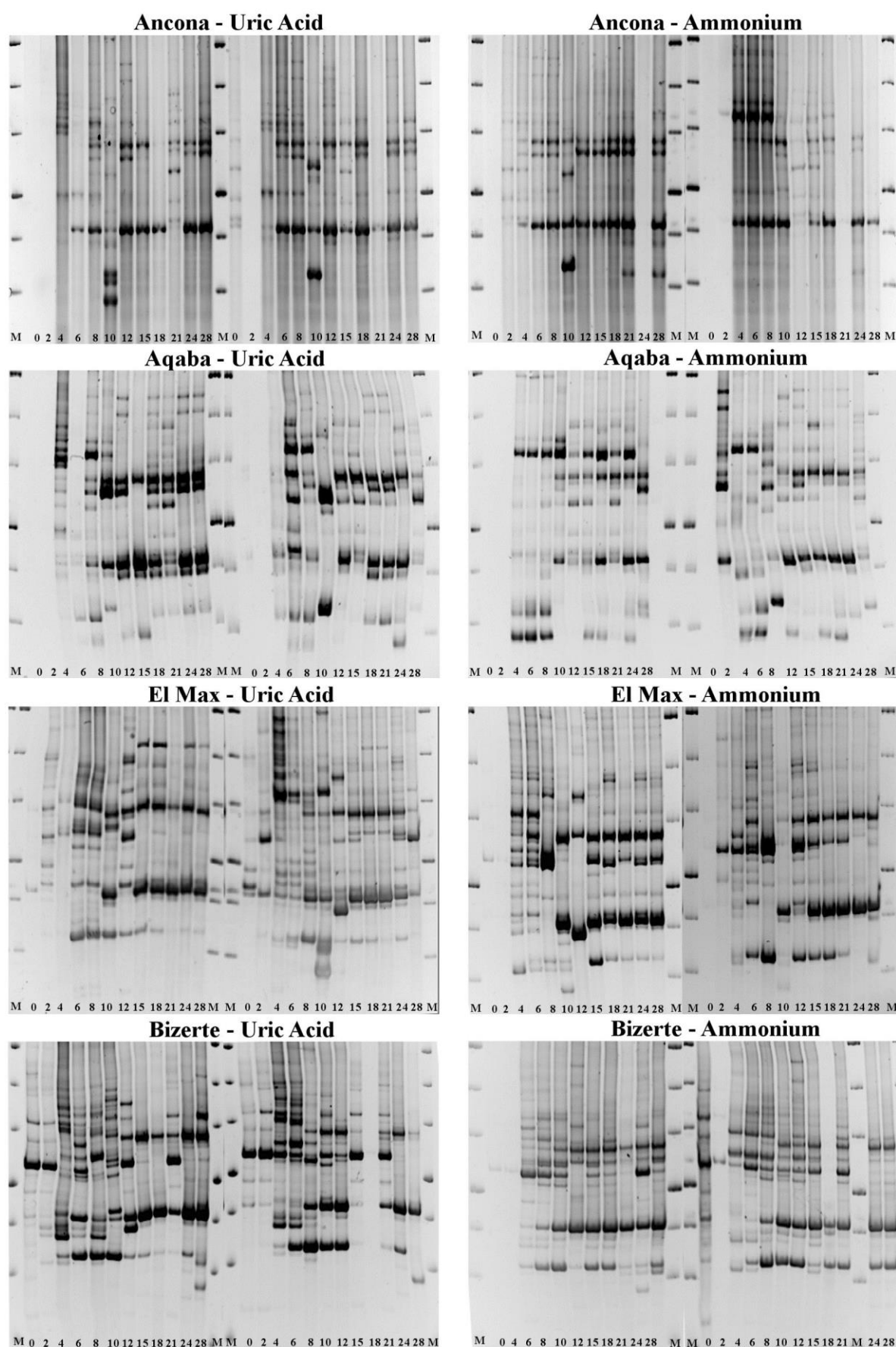
Figure 4.

Table 1.

Microcosm nr.	Microcosm name	Flask size (ml)	Inoculum (location)	Sterile sand (g)	ONR7 medium (ml)	Uric Acid (mmol)	Crude oil (ml)	NH ₄ Cl (mmol)	Na ₂ HPO ₄ (mmol)
1	Ancona - ammonium	500	Ancona	75	150	0	5	5	0.5
		500	Ancona	75	150	0	5	5	0.5
2	Ancona - uric acid	500	Ancona	75	150	1.25	5	0	0.5
		500	Ancona	75	150	1.25	5	0	0.5
3	Aqaba - ammonium	500	Aqaba	75	150	0	5	5	0.5
		500	Aqaba	75	150	0	5	5	0.5
4	Aqaba - uric acid	500	Aqaba	75	150	1.25	5	0	0.5
		500	Aqaba	75	150	1.25	5	0	0.5
5	El Max - ammonium	500	El Max	75	150	0	5	5	0.5
		500	El Max	75	150	0	5	5	0.5
6	El Max - uric acid	500	El Max	75	150	1.25	5	0	0.5
		500	El Max	75	150	1.25	5	0	0.5
7	Bizerta - ammonium	500	Bizerta	75	150	0	5	5	0.5
		500	Bizerta	75	150	0	5	5	0.5
8	Bizerta - uric acid	500	Bizerta	75	150	1.25	5	0	0.5
		500	Bizerta	75	150	1.25	5	0	0.5
9	Control - sterile	500	none	75	150	0	0	0	0
		500	none	75	150	0	0	0	0
10	Control - ammonium	500	none	75	150	0	5	5	0.5
		500	none	75	150	0	5	5	0.5
11	Control - uric acid	500	none	75	150	1.25	5	0	0.5
		500	none	75	150	1.25	5	0	0.5
12	Control - non sterile	500	Ancona	75	150	0	5	0	0
		500	Ancona	75	150	0	5	0	0
13	Ancona - ammonium	1000	Ancona	150	300	2.5	10	10	1
14	Ancona - uric acid	1000	Ancona	150	300	2.5	10	10	1
15	Aqaba - ammonium	1000	Aqaba	150	300	2.5	10	10	1
16	Aqaba - uric acid	1000	Aqaba	150	300	2.5	10	10	1
17	El Max - ammonium	1000	El Max	150	300	2.5	10	10	1
18	El Max - uric acid	1000	El Max	150	300	2.5	10	10	1
19	Bizerta - ammonium	1000	Bizerta	150	300	2.5	10	10	1
20	Bizerta - uric acid	1000	Bizerta	150	300	2.5	10	10	1

Table 2.

		Bizerte NP	Ancona NP	Ancona UA	El Max UA	Aqaba UA
Alphaproteobacteria	<i>Jannaschia</i> spp.	0,41	0,49	0,01	0,08	0,05
	<i>Loktanella</i> spp.	0,71	0,84	0,04	0,13	0,10
	<i>Oceanicola</i> spp.	0,48	0,65	0,03	0,13	0,05
	<i>Paracoccus</i> spp.	0,80	0,62	0,01	0,28	0,12
	<i>Rhodobacter</i> spp.	0,80	0,67	0,01	0,21	0,10
	<i>Roseobacter</i> spp.	0,20	0,16	0,01	0,03	0,02
	<i>Roseovarius</i> spp.	1,44	1,39	0,08	0,28	0,13
	<i>Ruegeria</i> spp.	1,10	1,08	0,05	0,16	0,11
	<i>Sulfitobacter</i> spp.	2,01	1,45	0,08	0,19	0,28
	<i>Thalassospira</i> spp.	0,00	0,51	0,00	0,06	0,02
Betaproteobacteria	<i>Acidovorax</i> spp.	0,00	0,21	0,09	0,11	0,44
	<i>Alcaligenes</i> spp.	0,01	0,00	0,00	0,05	0,14
	<i>Polaromonas</i> spp.	0,00	0,08	0,04	0,05	0,12
Gammaproteobacteria	<i>Aeromonas</i> spp.	0,96	0,54	1,52	2,77	1,09
	<i>Alcanivorax</i> spp.	16,35	4,94	3,94	0,35	0,64
	<i>Acinetobacter</i> spp.	0,66	0,05	0,93	1,13	1,31
	<i>Azotobacter</i> spp.	0,34	0,50	0,45	0,48	0,95
	<i>Citrobacter</i> spp.	0,20	0,29	0,11	0,76	0,54
	<i>Chromohalobacter</i> spp.	0,75	0,48	0,59	0,13	0,12
	<i>Colwellia</i> spp.	0,19	0,23	0,11	0,44	0,28
	<i>Cronobacter</i> spp.	0,14	0,00	0,07	0,70	0,48
	<i>Cycloclasticus</i> spp.	0,52	0,01	0,04	0,01	0,00
	<i>Enterobacter</i> spp.	0,31	0,42	0,17	0,70	0,48
	<i>Gallibacterium</i> spp.	0,06	0,07	0,06	0,09	0,22
	<i>Haemophilus</i> spp.	0,10	0,11	0,08	0,16	0,25
	<i>Hahella</i> spp.	0,19	0,12	0,23	0,16	0,44
	<i>Halomonas</i> spp.	3,64	4,18	6,05	4,18	1,40
	<i>Idiomarina</i> spp.	0,68	0,26	0,76	2,67	0,97
	<i>Legionella</i> spp.	1,59	0,23	0,91	0,63	1,52
	<i>Marichromatium</i> spp.	0,18	0,15	0,14	0,16	0,31
	<i>Marinobacter</i> spp.	6,40	6,32	16,76	3,14	1,97
	<i>Marinobacterium</i> spp.	0,78	1,48	1,09	1,00	1,09
	<i>Marinomonas</i> spp.	0,48	0,94	0,47	0,96	0,60
	<i>Methylophaga</i> spp.	0,19	0,19	0,22	0,14	0,30
	<i>Microbulbifer</i> spp.	0,69	0,82	0,60	1,10	0,52
	<i>Moraxella</i> spp.	0,42	0,17	0,06	0,23	0,37
	<i>Oceanimonas</i> spp.	0,44	0,16	0,43	1,15	0,10
	<i>Oceanospirillum</i> spp.	0,12	0,29	0,16	0,23	0,18
	<i>Oleiphilus</i> spp.	0,05	0,03	0,15	0,04	0,08
	<i>Oleispira</i> spp.	0,05	0,14	0,13	0,11	0,05
	<i>Pantoea</i> spp.	0,30	0,52	0,36	1,33	0,63
	<i>Photobacterium</i> spp.	0,41	0,86	0,52	2,15	0,83
	<i>Pseudoalteromonas</i> spp.	1,27	0,37	1,92	0,85	1,63
	<i>Pseudomonas</i> spp.	12,93	14,08	12,07	21,14	45,57
	<i>Psychromonas</i> spp.	0,21	0,22	0,26	1,12	0,34
	<i>Psychrobacter</i> spp.	0,64	0,21	6,81	0,51	1,26
	<i>Rheinheimeria</i> spp.	0,42	0,13	0,10	0,27	0,45
	<i>Salmonella</i> spp.	0,44	0,38	0,19	1,68	1,14
	<i>Shewanella</i> spp.	2,19	2,78	2,59	9,12	3,38
	<i>Thalassolituus</i> spp.	0,24	0,80	0,04	0,04	0,02
	<i>Thalassomonas</i> spp.	0,24	0,19	0,10	0,40	0,29
	<i>Thiomicrospira</i> spp.	0,26	0,40	0,63	0,66	0,40
	<i>Thiorhodococcus</i> spp.	0,17	0,13	0,11	0,11	0,23
	<i>Vibrio</i> spp.	0,96	2,38	1,14	5,08	3,85
	<i>Xenorhabdus</i> spp.	0,18	0,60	0,32	0,33	0,49
	<i>Methylococcaceae</i>	2,00	0,54	1,05	0,63	1,16
	<i>Xanthomonadaceae</i>	2,43	0,20	0,15	1,63	0,25
Deltaproteobacteria	<i>Deltaproteobacteria</i>	0,33	0,68	0,29	0,43	0,84
Flavobacteria	<i>Chryseobacterium</i> spp.	0,00	0,53	0,01	0,03	0,00
	<i>Flavobacterium</i> spp.	0,00	0,37	0,79	0,10	0,04
Actinobacteria	<i>Actinobacteria</i>	0,05	1,81	2,56	1,51	0,34
Firmicutes	<i>Firmicutes</i>	1,86	4,86	7,42	1,04	0,89
Spirochaeta	<i>Leptospira</i> spp.	0,00	0,03	0,03	0,06	0,17
	Others	28,02	35,68	23,87	24,43	17,90



Suppl. Fig 1.

Suppl. Tab. 1a.

<i>One way ANOSIM</i>	<i>R</i>	<i>Significance %</i>
Location	0,168	0,1
Nitrogen source	0,012	2,3
Sampling day	0,141	0,1
<i>Nested two way ANOSIM</i>	<i>R</i>	<i>Significance %</i>
Location within N source	-0,156	85,7

Suppl. Tab. 1b.

	<i>Ancona</i>	<i>Aqaba</i>	<i>Bizerte</i>	<i>El Max</i>
<i>Ancona</i>		0,164	0,25	0,254
<i>Aqaba</i>	0,164		0,14	0,132
<i>Bizerte</i>	0,25	0,14		0,076
<i>El Max</i>	0,254	0,132	0,076	

Suppl. Tab. 1c.

	<i>0</i>	<i>2</i>	<i>4</i>	<i>6</i>	<i>8</i>	<i>10</i>	<i>12</i>	<i>15</i>	<i>18</i>	<i>21</i>	<i>24</i>	<i>28</i>
<i>0</i>		-0,02	0,198	0,262	0,28	0,302	0,28	0,313	0,292	0,27	0,356	0,217
<i>2</i>	-0,02		0,11	0,162	0,186	0,23	0,181	0,227	0,197	0,168	0,275	0,121
<i>4</i>	0,198	0,11		-0,022	0,045	0,214	0,158	0,271	0,255	0,227	0,33	0,166
<i>6</i>	0,262	0,162	-0,022		-0,044	0,092	0,101	0,211	0,209	0,168	0,308	0,183
<i>8</i>	0,28	0,186	0,045	-0,044		0,056	0,081	0,191	0,186	0,149	0,268	0,181
<i>10</i>	0,302	0,23	0,214	0,092	0,056		0,068	0,08	0,089	0,061	0,188	0,113
<i>12</i>	0,28	0,181	0,158	0,101	0,081	0,068		0,006	-0,001	-0,006	0,049	0,009
<i>15</i>	0,313	0,227	0,271	0,211	0,191	0,08	0,006		-0,047	-0,032	-0,003	-0,002
<i>18</i>	0,292	0,197	0,255	0,209	0,186	0,089	-0,001	-0,047		-0,039	-0,062	-0,04
<i>21</i>	0,27	0,168	0,227	0,168	0,149	0,061	-0,006	-0,032	-0,039		-0,037	-0,008
<i>24</i>	0,356	0,275	0,33	0,308	0,268	0,188	0,049	-0,003	-0,062	-0,037		0,003
<i>28</i>	0,217	0,121	0,166	0,183	0,181	0,113	0,009	-0,002	-0,04	-0,008	0,003	

Suppl. Tab. 2a.

Sample name	Insert size (bp)	Sequence type (bp)	Raw reads (Mbp)	High quality reads (Mbp)	Clean data rate (%)
MGS-AQ (UA)	350	Index 91 PE	1.163	1.100	94,6
MGS-EIMAX (UA)	350	Index 91 PE	1.149	1.100	95,73
MGS-ANC (UA)	350	Index 91 PE	1.189	1.100	92,51
MGS-ANC (AMM)	350	Index 91 PE	1.160	1.100	94,83
MGS-BIZ (AMM)	350	Index 91 PE	1.266	1.100	86,88

Suppl. Tab. 2b.

Sample name	Contig num	Contig length (bp)	N50 (bp)	N90 (bp)	Max (bp)	Min (bp)	Map to own contigs (%)
MGS-AQ (UA)	8.371	21.504.196	12.101	753	273.210	500	83,38
MGS-ANC (AMM)	12.469	24.752.834	3.617	697	358.781	500	71,11
MGS-ANC (UA)	12.691	19.364.101	2.076	622	62.304	500	58,77
MGS-BIZ (AMM)	12.814	20.253.283	2.117	667	73.362	500	71,81
MGS-EIMAX (UA)	23.220	45.489.734	3.627	698	431.244	500	59,16

Suppl. Tab. 2c.

Sample name	ORFs	Avg. length (bp)	Assigned to COG	Assigned to KEGG	Nr of to COG	Nr of KEGG
MGS-AQ (UA)	26.866	712,98	18.242	17.111	2.719	2.780
MGS-ANC (AMM)	32.180	690,62	21.101	19.029	2.855	2.856
MGS-ANC (UA)	27.893	619,19	18.012	15.065	2.719	2.596
MGS-BIZ (AMM)	28.698	619,65	18.261	15.796	2.433	2.790
MGS-EIMAX (UA)	61.277	666,25	39.915	36.320	3.115	3.231

Suppl. Tab. 3a.

Gene Name	Definition	Ec number	ID Best Hit	Organism	Comments
gene_ULIXES1_12295	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	WP_009825456.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES1_19221	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	WP_004579155.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae or Alteromonadaceae - some similarity to Halomonas
gene_ULIXES1_21068	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	WP_016855753.1	Gammaproteobacteria	Likely similar to Halomonadaceae (Halomonas genus)
gene_ULIXES1_21700	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	NP_746404.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae (Pseudomonas genus)
gene_ULIXES1_2453	5-Hydroxyisourate hydrolase	3.5.2.17	WP_005850231.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES1_2499	5-Hydroxyisourate hydrolase	3.5.2.17	WP_009828007.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES1_12911	5-Hydroxyisourate hydrolase	3.5.2.17	WP_022962137.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae or Alteromonadaceae
gene_ULIXES1_14824	5-Hydroxyisourate hydrolase	3.5.2.17	WP_003298230.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae (Pseudomonas genus)
gene_ULIXES1_21697	5-Hydroxyisourate hydrolase	3.5.2.17	NP_746401.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae (Pseudomonas genus)
gene_ULIXES1_23813	5-Hydroxyisourate hydrolase	3.5.2.17	YP_004715528.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae (Pseudomonas genus)
gene_ULIXES1_7891	5-Hydroxyisourate hydrolase	3.5.2.17	YP_001173592.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae (Pseudomonas genus)
gene_ULIXES1_13633	Allantoicase (EC 3.5.3.4)	3.5.3.4	WP_003298235.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae (Pseudomonas genus)
gene_ULIXES1_19220	Allantoicase (EC 3.5.3.4)	3.5.3.4	WP_004579156.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae or Alteromonadaceae - some similarity to Halomonas
gene_ULIXES1_4670	Allantoate amidohydrolase	3.5.1.6	WP_009825998.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES1_14474	Allantoate amidohydrolase	3.5.1.6	WP_019966333.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae or Rhizobiaceae
gene_ULIXES1_21968	Allantoate amidohydrolase	3.5.1.6	WP_019436796.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae (Pseudomonas genus)
gene_ULIXES1_12294	Allantoin catabolism protein	3.5.3.-	WP_005850227.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES1_21067	Allantoin catabolism protein	3.5.3.-	YP_006557201.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae or Alteromonadaceae - some similarity to Halomonas
gene_ULIXES1_25528	Allantoin catabolism protein	3.5.3.-	NP_745667.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae or Alteromonadaceae - some similarity to Halomonas
gene_ULIXES1_2451	Urate catabolism protein/Allantoinase		WP_005850229.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES1_2498	Urate catabolism protein/Allantoinase		WP_008886954.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES1_14709	Urate catabolism protein/Allantoinase		WP_005861136.1	Alphaproteobacteria	Ambiguous
gene_ULIXES1_14823	Urate catabolism protein/Allantoinase		WP_017246039.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae (Pseudomonas genus)
gene_ULIXES1_19219	Urate catabolism protein/Allantoinase		WP_004579157.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae or Alteromonadaceae - some similarity to Halomonas
gene_ULIXES1_21066	Urate catabolism protein/Allantoinase		WP_022963453.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae or Alteromonadaceae - some similarity to Halomonas
gene_ULIXES1_21698	Urate catabolism protein/Allantoinase		WP_021783015.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae (Pseudomonas genus)
gene_ULIXES1_24466	Urate catabolism protein/Allantoinase		WP_017246039.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae (Pseudomonas genus)
gene_ULIXES1_12912	OHCU decarboxylase/allantoate amidohydrolase		WP_022962138.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae or Alteromonadaceae
gene_ULIXES1_13631	OHCU decarboxylase/allantoate amidohydrolase		YP_001173574.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae (Pseudomonas genus)
gene_ULIXES1_21699	OHCU decarboxylase/allantoate amidohydrolase		WP_021783014.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae (Pseudomonas genus)

Suppl. Tab. 3b.

Gene Name	Definition	Ec number	ID Best Hit	Organism	Comments
gene_ULIXES3_9556	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	YP_006557202.1	Gammaproteobacteria	Likely similar to Halomonadaceae (Halomonas)
gene_ULIXES3_18871	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	YP_005093662.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae or Alteromonadaceae
gene_ULIXES3_19460	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	YP_002799286.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae
gene_ULIXES3_26746	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	WP_008329785.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES3_5999	5-Hydroxyisourate hydrolase	3.5.2.17	WP_008295599.1	Gammaproteobacteria	Ambiguous
gene_ULIXES3_17905	5-Hydroxyisourate hydrolase	3.5.2.17	YP_005093665.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae or Alteromonadaceae
gene_ULIXES3_27361	5-Hydroxyisourate hydrolase	3.5.2.17	WP_009828007.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES3_19459	Allantoicase (EC 3.5.3.4)	3.5.3.4	WP_020682643.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae or Alteromonadaceae
gene_ULIXES3_13829	Allantoate amidohydrolase	3.5.1.6	WP_010138457.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES3_10994	Urate catabolism protein/Allantoinase		WP_022989072.1	Gammaproteobacteria	Likely similar to Alteromonadales or Halomonadaceae (Halomonas)
gene_ULIXES3_17906	Urate catabolism protein/Allantoinase		WP_019936009.1	Gammaproteobacteria	Likely similar to Aeromonadaceae
gene_ULIXES3_19238	Urate catabolism protein/Allantoinase		WP_007152518.1	Gammaproteobacteria	Likely similar to Alteromonadaceae
gene_ULIXES3_19461	Urate catabolism protein/Allantoinase		YP_572141.1	Gammaproteobacteria	Likely similar to Halomonadaceae (Halomonas)
gene_ULIXES3_27360	Urate catabolism protein/Allantoinase		WP_009572478.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES3_26108	OHCU decarboxylase/allantoate amidohydrolase		YP_006571999.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae

Suppl. Tab. 3c.

Gene Name	Definition	Ec number	ID Best Hit	Organism	Comments
gene_ULIXES-2_7300	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	WP_008329785.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES-2_10738	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	WP_017641447.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae or Alteromonadaceae
gene_ULIXES-2_27693	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	WP_009572476.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES-2_30028	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	YP_007683906.1	Gammaproteobacteria	Likely similar to Oceanospirillales (Thalassolituus)
gene_ULIXES-2_7302	5-Hydroxyisourate hydrolase	3.5.2.17	WP_009828007.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES-2_13190	5-Hydroxyisourate hydrolase	3.5.2.17	WP_008295599.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae or Alteromonadaceae
gene_ULIXES-2_30029	5-Hydroxyisourate hydrolase	3.5.2.17	YP_007683907.1	Gammaproteobacteria	Likely similar to Oceanospirillales (Thalassolituus)
gene_ULIXES-2_31605	5-Hydroxyisourate hydrolase	3.5.2.17	WP_009572479.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES-2_10737	Allantoicase (EC 3.5.3.4)	3.5.3.4	WP_020682643.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae or Alteromonadaceae
gene_ULIXES-2_30031	Allantoicase (EC 3.5.3.4)	3.5.3.4	YP_007683909.1	Gammaproteobacteria	Likely similar to Oceanospirillales (Thalassolituus)
gene_ULIXES-2_27694	Allantoin catabolism protein	3.5.3.-	WP_009572477.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES-2_7301	Allantoinase	3.5.2.5	WP_008207416.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES-2_27695	Allantoinase/bifunctional allantoinase/OHCU decarboxylase		WP_009572478.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES-2_10739	Urate catabolism protein/Allantoinase	3.5.2.5	YP_572141.1	Gammaproteobacteria	Likely similar to Halomonadaceae
gene_ULIXES-2_10740	Urate catabolism protein/Allantoinase	3.5.2.5	WP_017891710.1	Proteobacteria	Ambiguous
gene_ULIXES-2_23917	Urate catabolism protein/Allantoinase	3.5.2.5	WP_017891710.1	Gammaproteobacteria	Ambiguous
gene_ULIXES-2_30025	Urate catabolism protein/Allantoinase		YP_007683903.1	Gammaproteobacteria	Likely similar to Oceanospirillales (Thalassolituus)
gene_ULIXES-2_13189	OHCU decarboxylase/allantoate amidohydrolase		WP_020560622.1	Gammaproteobacteria	Ambiguous
gene_ULIXES-2_30030	OHCU decarboxylase/allantoate amidohydrolase		YP_007683908.1	Gammaproteobacteria	Likely similar to Oceanospirillales (Thalassolituus)

Suppl. Tab. 3d.

Gene Name	Definition	Ec number	ID Best Hit	Organism	Comments
gene_ULIXES4_4504	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	YP_007683906.1	Gammaproteobacteria	Likely similar to Oceanospirillales (Thalassolitus genus)
gene_ULIXES4_4506	5-Hydroxyisourate hydrolase	3.5.2.17	YP_007683907.1	Gammaproteobacteria	Likely similar to Oceanospirillales (Thalassolitus genus)
gene_ULIXES4_10647	5-Hydroxyisourate hydrolase	3.5.2.17	WP_008225166.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES4_4508	Allantoicase (EC 3.5.3.4)	3.5.3.4	YP_007683909.1	Gammaproteobacteria	Likely similar to Oceanospirillales (Thalassolitus genus)
gene_ULIXES4_4501	Urate catabolism protein/Allantoinase		YP_007683903.1	Gammaproteobacteria	Likely similar to Oceanospirillales (Thalassolitus genus)
gene_ULIXES4_4507	OHCu decarboxylase/allantoate amidohydrolase		YP_007683908.1	Gammaproteobacteria	Likely similar to Oceanospirillales (Thalassolitus genus)

Suppl. Tab. 3e.

Gene Name	Definition	Ec number	ID Best Hit	Organism	Comments
gene_ULIXES5_16814	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	WP_007482915.1	Gammaproteobacteria	Likely similar to Pseudomonadales (Acinetobacter genus)
gene_ULIXES5_22395	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	NP_746404.1	Gammaproteobacteria	Likely similar to Pseudomonadales (Pseudomonas genus)
gene_ULIXES5_29775	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	YP_001534537.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES5_30682	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	WP_016855753.1	Gammaproteobacteria	Likely similar to Halomonadaceae (Halomonas)
gene_ULIXES5_47410	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	YP_005093662.1	Gammaproteobacteria	Likely similar to Aeromonadaceae - some similarity to Halomonas
gene_ULIXES5_47641	Ureidoglycolate hydrolase (EC:3.5.3.19)	3.5.3.19	WP_003299751.1	Gammaproteobacteria	Likely similar to Pseudomonadales (Pseudomonas genus)
gene_ULIXES5_17480	5-Hydroxyisourate hydrolase	3.5.2.17	WP_007481680.1	Gammaproteobacteria	Likely similar to Pseudomonadales (Acinetobacter genus)
gene_ULIXES5_29777	5-Hydroxyisourate hydrolase	3.5.2.17	WP_007803838.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES5_29869	5-Hydroxyisourate hydrolase	3.5.2.17	YP_004715528.1	Gammaproteobacteria	Likely similar to Pseudomonadales (Pseudomonas genus) - some similarity to Halomonas
gene_ULIXES5_41564	5-Hydroxyisourate hydrolase	3.5.2.17	YP_004715528.1	Gammaproteobacteria	Likely similar to Pseudomonadales (Pseudomonas genus) - some similarity to Halomonas
gene_ULIXES5_45441	5-Hydroxyisourate hydrolase	3.5.2.17	WP_008566757.1	Gammaproteobacteria	Likely similar to Pseudomonadales (Pseudomonas genus)
gene_ULIXES5_47413	5-Hydroxyisourate hydrolase	3.5.2.17	YP_005093665.1	Gammaproteobacteria	Likely similar to Aeromonadaceae - some similarity to Halomonas
gene_ULIXES5_49758	5-Hydroxyisourate hydrolase	3.5.2.17	WP_004880295.1	Gammaproteobacteria	Likely similar to Pseudomonadales (Acinetobacter genus)
gene_ULIXES5_56324	5-Hydroxyisourate hydrolase	3.5.2.17	YP_005091502.1	Gammaproteobacteria	Likely similar to Aeromonadaceae - some similarity to Halomonas
gene_ULIXES5_16815	Allantoicase (EC 3.5.3.4)	3.5.3.4	WP_007482917.1	Gammaproteobacteria	Likely similar to Pseudomonadales (Acinetobacter genus)
gene_ULIXES5_28263	Allantoin catabolism protein	3.5.3.-	YP_005090755.1	Gammaproteobacteria	Likely similar to Aeromonadaceae - some similarity to Halomonas
gene_ULIXES5_29772	Allantoin catabolism protein	3.5.3.-	WP_009816826.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES5_30681	Allantoin catabolism protein	3.5.3.-	YP_006557201.1	Gammaproteobacteria	Likely similar to Pseudomonadaceae or Alteromonadaceae - some similarity to Halomonas
gene_ULIXES5_29776	Allantoinase/bifunctional allantoicase/OHCu decarboxylase		YP_001534538.1	Alphaproteobacteria	Likely similar to Rhodobacteraceae
gene_ULIXES5_16808	Urate catabolism protein/Allantoinase		WP_004880270.1	Gammaproteobacteria	Likely similar to Pseudomonadales (Acinetobacter genus)
gene_ULIXES5_30680	Urate catabolism protein/Allantoinase	3.5.2.17	WP_019019214.1	Gammaproteobacteria	Likely similar to Halomonadaceae (Halomonas)
gene_ULIXES5_45440	Urate catabolism protein/Allantoinase		WP_017246039.1	Gammaproteobacteria	Likely similar to Pseudomonadales (Pseudomonas genus)
gene_ULIXES5_45741	Urate catabolism protein/Allantoinase		WP_004579157.1	Proteobacteria	Ambiguous
gene_ULIXES5_47412	Urate catabolism protein/Allantoinase		YP_005093664.1	Gammaproteobacteria	Likely similar to Aeromonadaceae
gene_ULIXES5_48158	Urate catabolism protein/Allantoinase	3.5.2.17	WP_004579157.1	Gammaproteobacteria	Likely similar to Alteromonadaceae or Halomonadaceae (Halomonas)
gene_ULIXES5_22394	OHCu decarboxylase/allantoate amidohydrolase		WP_021783014.1	Gammaproteobacteria	Likely similar to Pseudomonadales (Pseudomonas genus)
gene_ULIXES5_47411	OHCu decarboxylase/allantoate amidohydrolase		YP_005093663.1	Gammaproteobacteria	Likely similar to Aeromonadaceae
gene_ULIXES5_49757	OHCu decarboxylase/allantoate amidohydrolase		WP_007482940.1	Gammaproteobacteria	Likely similar to Pseudomonadales (Acinetobacter genus)
gene_ULIXES5_53891	OHCu decarboxylase/allantoate amidohydrolase		WP_017246038.1	Gammaproteobacteria	Likely similar to Pseudomonadales (Pseudomonas genus)

Suppl. Tab. 4.

Sequence	Closest relative	Accession nr.	Phylogeny	Seq. identity (%)
AN12-1	<i>Halomonas venusta</i>	NR_042069	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	96%
AN12-2	<i>Halomonas alkaliphila</i>	NR_042256	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
AN12-3	<i>Halomonas venusta</i>	AY669165	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
AN12-4	<i>Halomonas alkaliphila</i>	NR_042256	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	100%
AU6-1	<i>Halomonas venusta</i>	JF710982	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
AU6-2	<i>Halomonas alkaliphila</i>	NR_042256	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	100%
AU6-3	<i>Halomonas venusta</i>	JF710982	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
AU6-4	<i>Halomonas alkaliphila</i>	JQ680444	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
AU12-1	<i>Halomonas alkaliphila</i>	NR_042256	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
AU12-2	<i>Halomonas venusta</i>	AY669165	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
AU12-3	<i>Halomonas venusta</i>	AY669165	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
AU12-4	<i>Halomonas alkaliphila</i>	NR_042256	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	100%
BN12-1	<i>Halomonas salifodinae</i>	NR_044263	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
BN12-2	<i>Marinobacter adhaerens</i>	NR_074765	Bacteria; Proteobacteria; Gammaproteobacteria; Alteromonadales; Alteromonadaceae	87%
BN12-3	<i>Halomonas denitrificans</i>	EU541350	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	98%
BN12-4	<i>Marinobacter lipolyticus</i>	NR_025671	Bacteria; Proteobacteria; Gammaproteobacteria; Alteromonadales; Alteromonadaceae	99%
BU6-1	<i>Halomonas salifodinae</i>	NR_044263	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
BU6-2	<i>Halomonas venusta</i>	NR_042069	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
BU6-3	<i>Halomonas neptunia</i>	KC534295	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
BU6-4	<i>Halomonas salifodinae</i>	NR_044263	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	98%
BU12-1	<i>Halomonas salifodinae</i>	NR_044263	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
BU12-2	<i>Halomonas venusta</i>	JF710984	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
BU12-3	<i>Halomonas salifodinae</i>	NR_044263	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
BU12-4	<i>Halomonas salifodinae</i>	NR_044263	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
QN12-1	<i>Halomonas venusta</i>	AY669165	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
QN12-2	<i>Halomonas venusta</i>	AY669165	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
QN12-3	<i>Halomonas venusta</i>	NR_042069	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
QN12-4	<i>Halomonas venusta</i>	AY669165	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
QU6-1	<i>Halomonas venusta</i>	AY669165	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
QU6-2	<i>Halomonas venusta</i>	AY669165	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
QU6-3	<i>Halomonas venusta</i>	AY669165	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
QU6-4	<i>Halomonas venusta</i>	AY669165	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
QU12-1	<i>Halomonas nitroreducens</i>	NR_044317	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	98%
QU12-2	<i>Halomonas salifodinae</i>	JN645859	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
QU12-3	<i>Halomonas venusta</i>	AY669165	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
QU12-4	<i>Halomonas venusta</i>	AY669165	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
XN12-1	<i>Halomonas nitroreducens</i>	NR_044317	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	98%
XN12-2	<i>Halomonas ventosae</i>	FJ444984	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	98%
XN12-3	<i>Halomonas ventosae</i>	FJ444984	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	98%
XN12-4	<i>Halomonas ventosae</i>	FJ444984	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	98%
XU6-1	<i>Pelomonas aquatica</i>	NR_042614	Bacteria; Proteobacteria; Betaproteobacteria; Burkholderiales; Comamonadaceae	99%
XU6-2	<i>Halomonas neptunia</i>	KC354707	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
XU6-4	<i>Halomonas ventosae</i>	NR_042812	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	99%
XU12-1	<i>Halomonas ventosae</i>	FJ444984	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	98%
XU12-2	<i>Halomonas denitrificans</i>	EU541350	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	97%
XU12-3	<i>Halomonas ventosae</i>	FJ444984	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	97%
XU12-4	<i>Halomonas ventosae</i>	FJ444984	Bacteria; Proteobacteria; Gammaproteobacteria; Oceanospirillales; Halomonadaceae	98%