- 2 <u>Dear Dr. Kirsten Kuesel,</u>
- 3 <u>Co-Editor of Biogeosciences</u>

We would like to thank you and the reviewers for the positive and constructive reviews that have
improved the manuscript significantly. We have accepted all comments and addressed them in
detail below (our response is in bold).

7

8 <u>Main comments:</u>

9 1) The archaeal sequences belonging to uncultured lineages (MCG, MBG-B, MBG-D) represent

10 less than 5% of the sequences. It is impossible in this case to infer on their potential metabolisms

11 based solely on the geochemical characteristics of the environment in which they were detected. I

12 would suggest deleting all paragraphs pertaining to these groups in the discussion and discuss the

13 dominant archaeal groups which are also known described cultured organisms.

14 It is correct and the sequences of archaeal classes that were below 5% of the community 15 (MCG, MBG-B) were not discussed, as it impossible to infer their potential metabolisms 16 based only on the geochemical characteristics of the environment in which they were detected. 17 However MBG-D (as important group in methane cycle mechanism) was represented by

- 18 about 5% of the sequences in the deep sediment and thus discussed in the manuscript.
- 19

20 2) Part 4.2 is long and descriptive, whereas part 4.3 contains very interesting information. Please

21 delete part 4.2 and integrate the most relevant data from this part in part 4.3.

## 22 Part 4.2 was deleted as suggested and the relevant information was integrated to part 4.3.

23 3) Most of the discussion in part 4.3 needs to be checked for English and the authors need to

24 clearly write what data type they are discussing. Here are some examples:

# The revised version was edited as requested and additional round of language editing wasdone.

L. 543 states 'mcrA functional gene of methanogens/anaerobic methanotrophs shows', which
should be written 'xxx analysis of mcrA functional genes ... shows.'

## 29 The sentence was changed according to the advice.

- L. 546-548 'is likely because that the mcrA observed ...', what are the authors referring to here?
  qPCR data or mcrA gene sequences?
- 32 The sentence was clarified in the text.
- L. 547 'methanogens which maybe cause the reverse methanogenesis', methanogens possibly
  perform reverse methanogenesis but they don't cause it. Etc ...
- 35 This sentence was corrected in the text.

- 36
- 37 Specific comments
- 38 L. 82 : Please change 'microbial pathways' to 'microbial communities'
- 39 The change was made.
- 40 L. 187: 'Chimeras?'

41 Using the position window in the denoising pipeline of MOTHUR helps us to look at the same

42 region in the 16S rRNA gene in our sequenced dataset. In this way we can better compare the

43 data set between other samples of the same position window. After using the position window

- 44 we also use chimera check and discard all the chimeras. This was explained better in the text.
- L. 266-271: This paragraph has information that is repeated after in more detail, please delete. Same
  comment for paragraph L. 280-286.
- 47 The paragraph was deleted as suggested.
- 48 L. 285-286: Recent data suggests that the MCG and MBG-B are separate phyla and are not
- 49 considered to belong to the Thaumarchaeota anymore. Hence some of the further discussions need50 to be amended.
- 51 The MCG and MBG-B were almost not mentioned in the revised manuscript, however the 52 recent change in phylogenetics of MCG and MBG-B was added to the revised version.
- L. 304: MBG-D and DHVEG-1 belong to the same cluster, they can all be labelled MBG-D.
- 54 This was corrected.
- L. 384-385: please explain this sentence, what was analysed exactly?
- 56 The sentence was clarified.
- 57 L. 403: 'microbial populations', 'methane cycling'

#### 58 This paragraph was deleted.

- 59 L. 515-517: 'which creates ... in deep sediments', please rewrite this sentence
- 60 The sentence was rewritten and clarified.
- 61 L. 526: there is a verb missing in this sentence
- 62 The sentence was fixed.
- 63 L. 533: 'can occur by one microorganism?' Please rephrase.
- 64 The sentence was rephrased.
- L. 534-536: 16S rRNA gene data is not enough to infer on metabolism. Please delete this sentence.

66	The possible direct	mechanism that	could be per	formed by uni	ique methanotrophy	v like MBG-
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- 67 D (Thermoplasmata class) or MCG/MBG-B, which some of our sequences were similar was
- 68 suggested.
- 69

Revised version to Biogeosciences

## 71 Methane related changes in prokaryotes along geochemical profiles in sediments

72 of Lake Kinneret (Israel)

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- 89

90 Keywords: anaerobic methane oxidation, iron reduction, archaeal classification, bacterial91 classification, Lake Kinneret.

92

Abstract. Microbial methane oxidation is the primary control on the emission of the greenhouse gas methane into the atmosphere. In terrestrial environments, aerobic methanotrophic bacteria are largely responsible for this process. In marine sediments a coupling of anaerobic oxidation of methane (AOM) with sulfate reduction, often carried out by a consortium of anaerobic methanotrophic archaea (ANME) and sulfate reducing bacteria, consumes almost all methane produced within marine sediments. Motivated by recent evidence for AOM with iron(III) in Lake

Kinneret sediments, the goal of the present study was to link the geochemical gradients in the lake 99 porewater to the microbial community structure. Screening of archaeal 16S rRNA gene sequences 100 revealed a shift from hydrogenotrophic to acetoclastic methanogens with depth. The observed 101 changes in microbial community structure suggest possible direct and indirect mechanisms for the 102 AOM coupled to iron reduction in deep sediments. Members of the Burkholderiales and 103 Nitrospirales orders increased with depth, suggesting their involvement in iron reduction together 104 with Geobacter genus and "reverse methanogenesis". An indirect mechanism through sulfate and 105 ANMEs seems less probable to explain the data due to the absence of ANME sequences. This is 106 despite the abundant sequences related to sulfate reducing bacteria (Deltaproteobacteria) together 107 with the occurrence of dsrAB in the deep sediment that could indicate the production of sulfate 108 (disproportionation) from  $S^0$  for sulfate-driven AOM. The presence of the functional gene *pmoA* in 109 the deep anoxic sediment together with sequences related to Methylococcales suggests the existence 110 of a second unexpected indirect pathway - aerobic methane oxidation pathway in an anaerobic 111 112 environment.

113

#### 115 **1. Introduction**

116 Chemical profiles in the porewater of aquatic sediments reflect the sequence of microbially 117 mediated redox reactions that are driven by the availability of both, electron donors and of suitable 118 electron acceptors. The latter are depleted in the order of decreasing chemical potential, beginning 119 with oxygen and proceeding through nitrate, manganese and iron oxides, and then sulfate. Below 120 the main zone of sulfate reduction, the fermentation of organic carbon leads to the formation of 121 methane (CH<sub>4</sub>) by the process of methanogenesis (Froelich et al., 1979).

The produced methane is isotopically depleted in  ${}^{13}C$ , with values of ~-50 to -110‰ (Schoell, 122 1988), and the residual dissolved inorganic carbon (DIC) pool is enriched by an isotopic 123 fractionation factor (ɛ) of 50 to 70‰ (e.g., (Borowski et al., 2000; Whiticar, 1999)). When the 124 produced methane diffuses into a zone with a suitable electron acceptor, it can be consumed by 125 microbial oxidation (methanotrophy), the main process by which the important greenhouse methane 126 is prevented from escaping into the atmosphere. While in the terrestrial environment, aerobic 127 methanotrophy is the dominant process (Chistoserdova et al., 2005), in anaerobic marine sediments 128 archaea are found to consume the majority of upward diffusing methane coupled to sulfate 129 reduction (Knittel and Boetius, 2009; Thauer, 2010; Valentine, 2002). 130

131 Evidence from lipids and from fluorescence in situ hybridization (FISH) showed that a consortium of archaea and sulfate reducing bacteria are involved in this anaerobic methane 132 133 oxidation (AOM) (Boetius et al., 2000; Hinrichs et al., 1999; Orphan et al., 2001). To date, three groups of anaerobic methanotrophic archaea (ANME), named ANME-1, ANME-2, and ANME-3, 134 135 are known to perform sulfate driven AOM (Niemann et al., 2006; Orphan et al., 2002). However Recently Milucka et al. (2012) demonstrated AOM mediated solely by archaea, where the archaea 136 was shown to oxidize the methane and reduce the sulfate to elemental sulfur. Disproportionating 137 bacteria, also involved in this mechanism, oxidize and reduce this elemental sulfur to sulfate and 138 sulfide, respectively. The carbon isotopic fractionation factor (ɛ) for this methanotrophic process 139 was shown to be in the range of 4-30% (Kinnaman et al., 2007; Whiticar, 1999). Enrichment 140 cultures of ANME from different environments showed a carbon isotopic fractionation of 12-39‰ 141 (Holler et al., 2009). 142

Other electron acceptors were recently shown to drive AOM. Nitrite driven AOM by oxygenic bacteria was observed in two different freshwater ecosystems in Netherlands (Ettwig et al., 2009; Raghoebarsing et al., 2006) and also in peatlands (Zhu et al., 2012). Beal et al. (Beal et al., 2009) showed the potential of manganese and iron-driven AOM in marine sediments, and Egger et al., (2014) showed it in brackish costal sediments. In our recent study (Sivan et al., 2011), we provided *in situ* geochemical evidence for AOM coupled to microbial iron reduction below the main

methanogenesis zone in Lake Kinneret (LK) sediments, where dissolved sulfate and nitrate are 149 absent. However, the mechanism that is responsible for this process was not investigated. The goal 150 of the present study was is to explore the possible microbial pathways communities that may be 151 involved in the iron driven AOM in LK sediments. This is accomplished by combining chemical 152 153 and isotope analyses of porewater samples along a depth profile from LK sediments with molecular biological techniques. By using specific functional genes related to dissimilatory sulfate reductase 154 (dsr) for sulfate reducers (Klein et al., 2001), methyl-coenzyme M reductase (mcr) for methanogens 155 and anaerobic methanotrophs (Hallam et al., 2003) and particulate methane monooxygenase 156 (pmmo) for aerobic methanotrophs (McDonald et al., 2008) we pinpoint the most likely candidate 157 for this relatively unknown process. 158

#### 159 **1.1. Study site**

Located in northern Israel, Lake Kinneret (LK, Fig.1) is a warm monomictic subtropical lake. 160 161 Typical concentrations of major electron acceptors in the water column during the mixed period are 35–50 µM nitrate and 600 µM sulfate (Adler et al., 2011; Nishri et al., 2000; Serruya et al., 1974). 162 In the spring, the newly formed epilimnion is characterized by increasing temperatures and 163 enhanced phytoplankton development, while in the hypolimnion heterotrophic microorganisms 164 gradually deplete oxygen and then nitrate. Organic matter degradation by bacterial iron and 165 manganese reduction takes place below the thermocline in the summer, and in the end of the 166 stratification period, sulfate reduction starts in the bottom water. In the upper part of the sediment, 167 sulfate reduction is the dominant microbial process year round, and below depth of 5 cm this 168 process is mostly replaced by methanogenesis (Adler et al., 2011; Eckert and Conrad, 2007). Total 169 iron content (Fe<sub>(tot)</sub>) in the sediment increases with depth till 18 cm and then remains uniform 170 around 550 µmol/g Dry Weight (Eckert, 2000). Mn(II) concentration is about 13 µmol/g Dry 171 Weight in the sediment (Serruya, 1971). The total carbon (C<sub>tot</sub>) in the solid phase shows a decrease 172 from 14 µmol/g Dry Weight in the top part of the sediment to 8 µmol/g Dry Weight in the deep part 173 174 of the sediment (~30 cm) (Eckert, 2000).

175

#### 176 2. Material and Methods

#### 177 **2.1.** Sampling

Several sediment cores were collected from the center of the lake (Station A, Fig. 1) at a water depth of ~42 m (maximum lake water column depth) at different times using Perspex tubes, measuring 55 cm long by 5 cm in diameter, with a gravity corer. The cores were stored in the dark at 4°C until they were sliced (on the same day or the day after). Core sampling for the microbial 182 community study took place in December 2009. Over a dozen of porewater chemical and isotope 183 profiles were conducted seasonally (every 3-4 months) from 2007 to 2013. The slight seasonal 184 changes allowed using typical geochemical profiles (Adler et al., 2011) in order to sample for the 185 microbiology communities in the different electron acceptors zones and to correlate between the 186 microbial sampling and the geochemical profiles.

#### 187 **2.2.** Geochemical analyses

Cores were divided into 2-cm slices from top to bottom under a constant flow of N<sub>2</sub> using a slicing device. About 1.5 ml of each sediment slice was transferred into N<sub>2</sub>-flushed crimp bottles containing 5 ml of 1.5 N NaOH for the headspace measurements of CH<sub>4</sub> and  $\delta^{13}C_{CH4}$  (after Sivan et al., 2011). CH<sub>4</sub> concentrations were measured on a SHIMADZU gas chromatograph (GC 8IF) equipped with a FID detector at a precision of 2 µmol·L<sup>-1</sup>.

193 Porewater was extracted immediately from each slice by centrifugation at 27,000 g at 4°C in a  $N_2$  atmosphere, and the supernatant was filtered through 0.45-µm filters. A 1-ml subsample was 194 fixed with Ferrozine and analyzed for dissolved Fe(II) (Stookey, 1970). Four millimeters of 195 subsample were poured into an acidified vial (1 ml of 0.5N nitric acid) to measure dissolved Mn(II). 196 197 The sample was analyzed via an Inductively Coupled Plasma Mass Spectrometer (ICP-MS, Elan DRC II, Perkin Elmer) at a precision of  $\pm 10\%$ . For sulfide profiles, 1 ml of subsample was added to 198 199 zinc acetate and hydrogen sulfide concentrations were determined by the methylene blue method with MDL of the method is 1 µM (Cline, 1969). For sulfate measurements, 5 ml of the subsamples 200 were analyzed with a Dionex DX500 high-pressure liquid chromatograph with a precision of  $\pm 3\%$ . 201 Iron isotope analysis was done by acidifying the subsamples with 10% HCl for one week (to 202 dissolve any precipitated iron) and then purifying them by anion exchange chromatography (Borrok 203 et al., 2007). δ<sup>56</sup>Fe was measured on a Neptune multi-collector ICP-MS in high resolution mode 204 according to standard methods and standardized against isotopic reference material (IRMM-014) 205 with a precision of  $\pm 0.1\%$  (John and Adkins, 2010). Total lipids were extracted using the Bligh-206 (Bligh Dyer, 1959) with 207 Dyer procedure and solvent mixture of 2:1:0.8 (methanol:dichloromethane:buffer).  $\delta^{13}$ C of the total lipid extraction (TLE) was measured on an 208 Elemental Analyzer Isotopic Ratio Mass Spectrometer (EA-IRMS) with a precision of 0.1%. 209

210 **2.3**.

#### DNA extraction and quantitative PCR (qPCR) amplification from sediment samples

Part of the sediment slices were kept frozen at -20°C for the microbial work. Samples from three different depths (0–3 cm representing the sulfate reduction zone, 6–9 cm representing the methanogenesis zone, and 29–32 cm representing the deep AOM zone) were defrosted. Those depth zones were chosen based on the geochemical profiles sampled at different times, showing quasi

steady state and stabilization of the sediments (Adler et al., 2011; Sivan et al., 2011). Therefore, 215 microbial community sampling of the three depth zones at a single time point should represent the 216 different microbial habitats as they pertain to the respective geochemical zones. Total genomic 217 DNA was extracted from the sediment samples using the MoBio Power Soil DNA isolation kit 218 219 (MoBio Laboratories, Solana Beach, CA). Genomic DNA was eluted using 60 µl of elution buffer and stored at -20°C. Concentrations of DNA were determined via UV-Vis spectrophotometry (ND-220 1000 NanoDrop Technologies, Wilmington, DE) (sulfate reduction zone 22ng/µl, methanogenesis 221 zone 35.8ng/ $\mu$ l and AOM zone 14ng/ $\mu$ l). 222

Quantification of functional genes were performed using the mcrA primers (Luton et al., 2002), 223 F 'GGTGGTGTMGGATTCACACARTAYGCWACAG' R 224 and 'TTCATTGCRTAGTTWGGRTAGTT', dsrA primers, DSR1F 'ACSCACTGGAAGCACG' 225 (Wagner et al., 1998) and RH3-dsr-R 'GGTGGAGCCGTGCATGTT' (Ben-Dov et al., 2007) and 226 pmoA primers, a189F 'GGNGACTGGGACTTCTGG' and mb661R 227 'CCGGMGCAACGTCYTTACC' (Yan et al., 2006) with a ABI prism 7000 Sequence Detection 228 System (Applied Biosystems). The qPCR reaction consisted of 10 µL of Absolute Blue SYBR 229 230 Green ROX, 150 nM each of forward and reverse primers and 5 µL of each DNA template. Thermal cycling conditions were as follows: 15 minutes at 95°C for enzyme activation, followed by 231 40 rounds of 15 seconds at 95°C for denaturation and 1 minute at 60°C for annealing/extension. To 232 verify that the used primer pair produced only a single specific product, a dissociation protocol was 233 234 added after thermo cycling, to determine dissociation of the PCR products from 60°C to 95°C. Standards for the calibration curves for quantification were made using pGEM-T Easy plasmid 235 236 cloned with 1.9 kb dsrAB and amplicons of the mcrA and pmoA encoding for functional genes at a known concentration with six serial dilution points (in steps of tenfold). All runs included a no-237 template control. Plasmid standards and environmental samples were simultaneously assayed in 238 triplicates. The ABI prism 7000 Sequence Detection System and SDS Software were used for data 239 analysis. qC values were exported into a Microsoft Excel Work sheet for further statistical analysis. 240

#### 241 **2.4.** Sequence analysis

454 tag-encoded FLX amplicon pyrosequencing was performed by the Research and Testing
Laboratory (Lubbock, Texas, USA) as previously described (Dowd et al., 2008). The bacterial and
archaeal 16S rRNA gene primers that were used are 28F 'GAGTTTGATCNTGGCTCAG' and
519R 'GTNTTACNGCGGCKGCTG' and Arch349F 'GYGCASCAGKCGMGAAW' and Arch806R
'GGACTACVSGGGTATCTAAT' respectively.

Data analysis was made using two different methodologies (MOTHUR and SILVA ngs). The initial 247 trimming of the sequences was made by MOTHUR v1.33 (Schloss et al., 2009) and generated 248 around 300 bp length of sequences. The trimmed sequences were taken to further analysis by 249 MOTHUR (Schloss et al., 2011) using SILVA.nr\_v119 database. MOTHUR 454 pipeline filtration 250 251 and denoising remove from the analysis sequences that were <150 bp, when they contained homopolymers longer than 8 bp, ambiguous bases, more than one mismatch to barcode sequences 252 or more than two mismatches to the forward primer sequence. We further removed sequences that 253 did not align in the same nucleotide position on the reference database. 254

255 Most of the bacterial sequences were in the same region, however the archaeal sequences were spared spaced between two different regions. We used one position window that was dominant in 256 the top sample and implied it on the other archaeal analysis in order to better compare the dataset 257 between other samples of the same position window. Identical sequences were grouped and then 258 were aligned against SILVA.nr\_v119. Chimeras were removed using MOTHUR CHIMERA 259 UCHIME (Edgar et al., 2011). The lengths of the remaining sequences were around 200 bp. A 260 further screening step (pre-cluster) was applied to reduce sequencing noise by clustering reads 261 differing by only one base every 100 bases (Huse et al., 2010). In addition, the rest of the sequences 262 263 were classified in order to remove eukaryote, mitochondria and chloroplast classified sequences. The remaining sequences were used to generate a distance matrix and clustering into operational 264 265 taxonomic units (OTUs) defined at 97% cutoff using the average neighbor algorithm. The OTUs were classified using SILVA.nr v119 database with a confidence threshold of 80%. Alpha diversity 266 267 was calculated by MOTHUR using the remaining sequences at 97% similarity. The beta diversity was calculated using comparable data, the number of sequences per sample was made equal through 268 subsampling (Table 1 and 2). 269

The second program which we used for analysis was SILVA ngs pipeline (Quast et al., 2013). 270 The trimmed sequences from mother were Aligner against the SILVA SSU rRNA seed. Sequences 271 shorter than 50 aligned nucleotides and with more than 2% of ambiguities or 2% of homopolymers, 272 respectively, were removed. The sequences that were not aligned as being putative 273 contaminations/artifacts were removed. Then sequences were clustered to OTUs with 98% 274 similarity and classified by local nucleotide BLAST search against SILVA database v119. To filter 275 out low identity and artificial BLAST hits, hits for which the function '(%sequence identity 276 277 +%alignment coverage)/2' did not exceed the value of 93% were discarded. Sequences with weak low score were classified as 'No Relatives' and will be disregarded from now on. Standard 278 279 deviation between the percentages of the two classification showed that at the phylum level there 280 are small differences between the two pipelines (supplementary Table 1). In the order classification, the standard deviations increased but the sequences still showed close similarity. Estimates of phylotype richness, diversity coverage and similarity were calculated according to the abundancebased coverage estimate (ACE), Chao's estimator (Chao, 1984; Chao and Ma, 1993), the Shannon diversity index, Good's coverage (Good, 1953) were calculated in MOTHUR (Table 1 and 2). The distribution and abundance matrix of the OTUs was normalized to the sample with the smallest number of reads by randomly resampling of the MOTHUR data set for statistical comparisons. Raw sequencing data was deposited in the MG-RAST (metagenomics.anl.gov) archive.

#### 288 2.5. Microbial community structure related to the environment conditions

To estimate community similarity among samples, we used PC-ORD 6 software. Subsample 289 OTU data from MOTHUR was used to create a distance matrix based on the Sørensen (Brav-290 Curtis) dissimilarities of the OTU composition of the samples. The data was normalized to 291 292 percentages before the analysis. Community relationships were visualized using principal coordinate analysis (PCoA) based on this distance matrix. Different environmental variables were 293 added as well (CH<sub>4</sub>,  $\delta^{13}C_{CH4}$ , Fe(II),  $\delta^{56}$ Fe(II), Mn(II), H<sub>2</sub>S, SO<sub>4</sub><sup>2-</sup>, dissolved organic carbon (DOC), 294  $\delta^{13}C_{TLE}$ , NH<sub>4</sub><sup>+</sup> and PO<sub>4</sub><sup>3-</sup>) in order to identify potential explanatory variables. The environmental 295 vector were applied and projected to the PCoA ordination. In addition, Venn diagrams for graphical 296 descriptions of unshared and shared OTUs between the three samples were constructed using 297 298 MOTHUR.

299

#### 300 **3. Results**

#### 301 **3.1.** Porewater profiles

This study focused on microbial community shifts along the porewater profiles of electron 302 acceptor gradients related to the methane cycle. To characterize shifts in main electron acceptors 303 with depth, Oover- a dozen of porewater profiles were performed seasonally from station A (Fig. 1) 304 to characterize shifts in main electron acceptors with depth. The shown profiles (Fig. 2) are 305 representatives of the slight seasonal trends. Sulfate and sulfide profiles show depletion from ~100 306 µM at the top part of the sediment, to below detection limits within the upper 15 cm (Fig. 2A). This 307 typical concave curvature profile of sulfate profile in the porewater indicates intensive sulfate 308 309 reduction in the upper few centimeters of the sediment throughout the year. A typical LK methane concentration profile (Fig. 2B) shows increase from 250 µM at the water-sediment interface to a 310 maximum of about 1.25 mM in the depth range of 7 to 15 cm, and then a decrease below 15 cm 311 depth. The profile of  $\delta^{13}C_{CH4}$  (Fig. 2B) shows a decrease from -60‰ at a depth of 1 cm to about 312

-65% at a depth of 7 cm and then an increase in the deeper sediments to a maximum value of -53.5% at a depth of 25 cm. Also, the profile of  $\delta 13CTLE$  (Fig. 2C) shows a decrease in this deepest part of the sediment.

Manganese and iron oxides are the most probable electron accepters acceptors in the deep part 316 317 of the sediment where methane is decreased. Dissolved Mn(II) concentration (Fig. 2D) increased from 5  $\mu$ M at the top of the sediment to a plateau of about 23  $\mu$ M from depths of 23 to 36 cm. The 318 dissolved Fe(II) concentration profile also showed an increase with depth (Fig. 2D), however 319 however infollowing a different pattern. In the upper 15 cm, dissolved Fe(II) concentrations were 320 below the detection limit, and gradually increased below 15 cm to about 90 µM at a depth of 36 cm. 321 The  $\delta^{56}$ Fe profile (Fig. 2C) showed a decrease with depth from 0.5% in the upper part to -1.7% to 322 -2.3% in the deep part of the sediment. 323

### 324 **3.2.** Sediment microbial communities

To study the sediment depth-dependent microbial community shifts, 16S rRNA gene sequences 325 326 of the prokaryotic community from three different depth zones (0-3, 6-9 and 29-32 cm, see arrows)in Fig. 2a) were extracted and analyzed. As mentioned above, these three zones represent the main 327 sulfate reduction zone, the methanogenesis zone, and the deep iron driven AOM zone, respectively. 328 The diverse bacterial and archaeal communities at these depth zones varied in composition and 329 richness throughout the sediment. A total of 13227 and 4881 bacterial and archaeal sequences 330 representing 3852 and 705 OTUs (cut-off value of 98%), respectively, were identified by SILVA 331 database project. The normalized Shannon and Chao index were used as proxies for diversity and 332 richness, respectively (Table 1 and 2). The upper layer showed lower diversity (Shannon index, 333 3.91 and 1.8 for bacteria and archaea) and richness (Chao index, 925.7 and 83.4 for bacteria and 334 archaea) than the other depths. Moreover, the bacterial coverage of the upper layer was the highest 335 (91%), while that of the deeper layers was about 80%. The archaeal coverage was almost the same 336 (~88%) in all the sampled layers. Most of our sequences were affiliated with uncultured 337 microorganisms mainly from various sediment environments. 338

Although the bacterial OTUs were distributed over 43 phyla, we represent here only the phyla that were over 1% sequences of the 454 library. The most abundant phyla were: *Proteobacteria*, *Chloroflexi*, *Nitrospirae*, *Bacteroidetes*, *Firmicutes* and *Chlorobi* observed in at least one of the libraries (Fig. 3). Dominant phyla had different trends; *Chloroflexi* and *Nitrospirae* showed increase with depth while *Proteobacteria* showed decrease with depth. *Firmicutes* were highly abundant only in the top layer. *Bacteroidetes* and *Chlorobi* were more abundant in the middle layer of the sediment.

The communities of microorganisms in the top layer (0-3cm) of the sediment had less sequence 346 overlap with those of the deeper layers (70 and 33 shared bacterial sequences with the middle and 347 bottom layers respectively) and of the deeper layers (Supplementary Fig 1and 2). The most 348 dominant (~48%) phylum in this bacterial community was Proteobacteria; 349 class 350 Gammaproteobacteria (91%) and the genera Acinetobacter (28%) and Pseudomonas (14%) (Fig. 3). The second most abundant (38%) phylum was Firmicutes; class Clostridia (21%) and Bacilli 351 (17%), family *Clostridiaceae* (15%) and genus *Bacillus* (14%). Further sequences were related to 352 other phyla and order (Fig 3 and supplementary Table 1). 353

In the archaeal community the sequences number were much lower, however the coverage 354 calculation shows about 90% coverage. The dominant phylum in the upper layer was 355 Euryarchaeota (98%), and the rest of the 2% was Thaumarchaeota. Methanomicrobia (92%) was 356 the dominant class of Euryarchaeota divided between genera of Methanoregula (58%), 357 Methanosaeta (22%) and Methanolinea (11%). The other abundant classes of Euryarchaeota were 358 Halobacteria (2%) and Thermoplasmata (3%). Thaumarchaeota was divided between two classes: 359 Marine Benthic Group B (MBG-B 0.7%) and Miscellaneous Crenarchaeotic Group (MCG 1%) 360 (Fig. 4 and supplementary table 1). 361

The middle layer (6-9cm) was the richest and most diverse in its bacterial and archaeal 362 populations comparing to the other layers, but with the lowest community coverage (bacterial 77% 363 and archaeal 86%). The dominant phyla in this depth were divided to Proteobacteria (40%), 364 Chloroflexi (14%), Bacteroidetes (7%), Chlorobi (5%) and Nitrospirae (4%). The dominant classes 365 in Proteobacteria were *Deltaproteobacteria* (68%), *Betaproteobacteria* (14%)366 and Gammaproteobacteria (14%). Sva0485 (8%), Syntrophobacterales (7%), Desulfarculales (3%) and 367 Desulfuromonadales (2%) were the dominant affiliated order in Deltaproteobacteria. 368 369 Methylococcaceae family of Gammaproteobacteria had 1% affiliated sequences. The dominant family in Chloroflexi was Anaerolineaceae (6%). Ignavibacteriales (4%) order was the dominant in 370 Chlorobi. Nitrospirae increased from the upper layer and was mainly represented by the 371 Nitrospiraceae (4%) family. Further sequences were related to other phyla and order (Fig. 3, 372 supplementary Table 1). 373

The archaeal community was the richest and most diverse in the middle zone. The dominant 374 phyla were Euryarchaeota (96%) while Thaumarchaeota has the rest of the 4%. Euryarchaeota was 375 divided between 3 dominant classes: Methanomicrobia (78%), Thermoplasmata (11%) and 376 Halobacteria (6%). Methanomicrobia was divided between Methanoregula (19%), 377 Methanosaeta(50%) and Methanolinea (10%) genus's. Thermoplasmata was divided between 378 Marine Benthic Group D-and DHVEG-1 (5%) and Terrestrial Miscellaneous Gp(TMEG) (3%). 379 Halobacteria dominant family was Deep Sea Hydrothermal Vent Gp 6(DHVEG-6) (6%). 380

*Thaumarchaeota* was divided mainly between two classes: Marine Benthic Group B (MBGB 1.5%)
 and Miscellaneous Crenarchaeotic Group (MCG 1.5%) (Fig. 4 and supplementary table 1).

The bottom layer (29-32 cm) bacterial community dominant phyla included: Proteobacteria 383 (39%), Chloroflexi (19%) and Nitrospirae (8%). Proteobacteria were divided to three main classes: 384 385 Deltaproteobacteria (20%), Gammaproteobacteria (10%) and Betaproteobacteria (6%). Sva0485 (11%) and Syntrophobacterales (4%) were the dominant affiliated order in Deltaproteobacteria. 386 The dominant family in Chloroflexi was Anaerolineaceae (7%) and Ignavibacteriales (4%) order is 387 the dominant *Chlorobi*. *Nitrospirae* increased from the upper layer and was mainly represented by 388 Nitrospiraceae (8%) family. Further sequences were related to other phyla and order (Fig. 3, 389 supplementary Table 1) 390

At this depth, the dominant archaeal phyla were Euryarchaeota (98%) while Thaumarchaeota 391 has the rest of the 2%. Euryarchaeota was divided between 3 dominant classes: Methanomicrobia 392 (86%), Thermoplasmata (6%) and Halobacteria (5%). Methanomicrobia was divided between 393 394 Methanoregula (16%), Methanosaeta (58%) and Methanolinea (11%) genus's. Thermoplasmata was divided between Marine Benthic Group D (MBG-D)and DHVEG-1 (5%) and Terrestrial 395 Miscellaneous Gp(TMEG) (3%) family. Halobacteria dominant family was Deep Sea 396 Hydrothermal Vent Gp 6(DHVEG-6) (1%). Thaumarchaeota was divided mainly between two 397 classes: Marine Benthic Group B (MBG-B 1%) and Miscellaneous Crenarchaeotic Group (MCG 398 0.6%) (Fig. 4 and supplementary table 1). The detailed description of the major communities in the 399 sediment is described at the discussion and the appendix. No ANME sequences were detected even 400 though specific primers (ANME2C-AR468F, ANME3-1249, ANME1-395F, ANME1-1417, 401 ANME3-140F, ANME3-1249, ANME2a-426 and ANME2a-1242R, see Miyashita et al., 2009) 402 were used. 403

404

#### 405 **3.3.** Depth comparison of microbial communities

An ordination plot from the 454 pyrosequencing by MOTHUR subsample OTUs dataset was 406 derived from Principal coordinate's analysis (PCoA) of the bacterial and archaeal. The PCoA 407 display the similarities and differences between the bacterial and archaeal communities varied with 408 depth (Fig. 5a and 5b respectively). The PCoA of bacteria and archaea show very similar separation 409 between the communities of different layers related to the different environment conditions of each 410 layer. The two deeper layers are on the same distant on the X axis showing relatively more 411 similarity than the upper layer. The vectors of the sulfate and sulfide are correlative with 412 communities of the top layer, while methane,  $\delta^{13}C_{TLE}$  and  $\delta^{56}Fe(II)$  were more associated to the 413 communities of the middle layer. The communities in the bottom layer were more correlated to 414

Fe(II), Mn(II) and  $NH_4^+$ . The Venn diagram shows also that more OTUs from the deep layers were shared than between the upper layer and the deeper layers (Supplementary Fig 1 and 2).

#### 417 **3.4.** Functional gene profile

In order to better understand the abundance of microbial functionality in the different depth we used qPCR for different functional genes. The functional genes *mcrA* and *dsrA* had very similar pattern - low concentrations in the upper layer  $(9*10^5\pm6.4*10^4 \text{ and } 1*10^6\pm1.7*10^5 \text{ respectively})$  and highest concentrations  $(6.9*10^6\pm6.7*10^5 \text{ and } 6.9*10^6\pm9.9*10^4)$  in the middle layer. *pmoA* gene showed also the same pattern, however with lower concentration  $(2.3*10^5\pm9.7*10^3 \text{ to} 1.6*10^6\pm1.7*10^5)$  (Fig. 6).

424

## 425 **4. Discussion**

In this study of LK sediments we investigated changes in the microbial diversity associated 426 with porewater geochemistry and the transition of the dominant electron acceptors with depth (0-427 40 cm). The geochemical porewater profiles in LK (Fig. 2) suggest that the sediment can be broadly 428 divided into three different regions of microbial processes: the upper 5 cm which is dominated by 429 sulfate reduction, the methanogenesis zone between 5 to 17 cm, and the deep sediments, dominated 430 by anaerobic oxidation of methane coupled to iron reduction. This division of the sediment and the 431 deep iron-driven AOM processes were confirmed by *in situ* profiles of methane,  $\delta^{13}C_{CH4}$ , sulfate, 432 and ferrous iron and subsequent use of a numerical mass conservation model (based on the 433 geochemical profiles of DIC and  $\delta^{13}C_{DIC}$  (Adler et al., 2011) and a set of geochemical incubation 434 experiments conducted in our previous work (Sivan et al., 2011). 435

The mechanisms that enable the process of AOM via iron reduction can be characterized by the 436 existence of specific prokaryotic populations at each depth and their resemblance to similar 437 environments with a distinct characterization. There are only few studies that analyzed microbial 438 communities relative to geochemical zones in fresh water sediments (Deutzmann and Schink, 2011; 439 Koizumi et al., 2004; Ye et al., 2009). Therefore, this study can deepen the information on 440 microbial community shifts under different electron acceptors conditions, especially those related to 441 the methane cycle. It is also the first study in LK that divides the sediment to different layers and 442 attribute significantly to the diversity of the microbial ecology in the different zones. 443

444 Pipeline of SILVA ngs was used for the alpha-diversity to get more classified sequences and 445 better coverage of the classification and population. However, for the beta-diversity the data from 446 the MOTHUR pipeline applied to get more statistical results without the bias of different sample size. The changes between the pipeline results for the alpha diversity were not large in the phyla
level (standard deviation of ~0.5%) but increased with higher taxonomy levels.

#### 449 **4.1. Sediment geochemical conditions**

In the upper layer of sediment (0-3 cm), sulfate concentrations were the highest and decreased 450 with depth as particulate organic matter from the water column accumulates, making organic carbon 451 degradation available (Hadas and Pinkas, 1992). Adler et al. (2011) calculated that the bulk of 452 bacterial sulfate reduction occurs in the top 1 cm of sediment, a finding that was supported by the 453 microbial work of Hadas and Pinkas, (1992). Using sulfide and particulate organic carbon 454 measurements, Eckert and Conrad. (2007) showed that sulfate reduction accounts for most of the 455 mineralization of organic matter in the upper 15 cm of sediment. Methane concentrations in the 456 upper layer were relatively low and probably reached from upward diffusion as methanogens are 457 outcompeted by sulfate reducers there (Lovley and Klug, 1983). The depletion in methane 458 concentrations and the decline  $\delta^{13}C_{TLE}$  values and the heavier isotopic values of  $\delta^{13}C_{CH4}$  in the 459 upper layer (Fig 2) compared to the methanogenic zone may be explained by AOM via sulfate 460 461 reduction, although no ANME sequences were found using specific primers and Fluorescent in situ hybridization methods or -by our phylogenetic alignment of 16S rRNA gene at libraries at any depth 462 in the sediment. 463

464 Below the sulfate reduction zone, in the middle layer of sediment (6-9 cm), methane reached its maximum concentrations (Fig. 2B). The low values of  $\delta^{13}C_{CH4}$  in the methanogenesis zone are 465 typical due to the large carbon isotope fractionation that occurs during methane production, thereby 466 leaving the produced methane isotopically light and the DIC isotopically heavy. Below the 467 maximum methane production zone (around 20 cm), methane concentrations began to decrease 468 while  $\delta^{13}C_{CH4}$  values increased (Fig. 2B), an outcome that may be due to the AOM process, which 469 leaves residual methane isotopically heavier. Depleted  $\delta^{13}C_{TLE}$  (Fig. 2C) in the deep part of the 470 sediment also supports the presence of AOM with the production of light biomass from methane 471 oxidation. In addition, although all the suitable electron acceptors at this depth were depleted, both 472 Fe(II) and Mn(II) increased (Fig. 2D). Also,  $\delta^{56}$ Fe values (Fig. 2C) of dissolved iron in the deep 473 sediment were isotopically negative, similar to sediments with active dissimilatory bacterial iron 474 reduction (Severmann et al., 2006) and are an indication of active iron reduction in the deep 475 sediments and not just diffusion. The increase in Fe(II) concentrations below a depth of 15 cm was 476 probably supported by the absence of sulfide. The potential of highly reactive Fe(III) oxides to drive 477 AOM in LK was also supported by our set of mesocosm incubation studies (Sivan et al., 2011). 478 Since manganese oxide concentrations were very low (~0.04%) throughout the LK sediment 479 column (Serruya et al., 1974), iron oxide seemed to play a bigger role in this AOM process. 480

#### 481 **4.2.** Microbial population that could be involved in methane cycle

482 *4.2.1. Deltaproteobacteria* 

Deltaproteobacteria were the most abundant in the middle and deep sediment samples. 483 Many members of the families of Syntrophobacterales are sulfate reducers or as fermentative 484 485 organism, depending on the environmental conditions (McInerney et al., 2007). Syntrophaceae, the dominant family of Syntrophobacterales, are acetate degrading sulfate reducing bacterium (Jiang et 486 al., 2009). Desulfarculales are strictly anaerobic mesophilic sulfate-reducing bacterium with the 487 capability to oxidize acetate and fatty acids (Sun et al., 2010). Desulfobacteralesare is sulfate 488 reducer, capable of oxidizing hydrogen in soils and sediments (Burow et al., 2014). 489 Desulfuromonadales was shown as S<sup>0</sup> respiring (Pjevac et al., 2014) and may also reduce Fe(III) 490 and Mn(IV) in marine surface sediments (Lovley, 2006). About 1% of the sequences in the middle 491 and bottom layers or the cores were affiliated to the Geobacter genus members which are involved 492 493 in iron reduction. In the deep layer the most abundant class was Sva0485. Sequences of the SVa485 order were similar to those found in different aquatic environments, but the metabolic functions of 494 members of this order are not clear. Pelobacter carbinolious (Lovley et al., 1995), a member of 495 SVa485 is capable of Fe(III) and sulfur reduction. Schwarz et al. (2007a) showed that in the upper 496 part of LK sediment Deltaproteobacteria was one of the dominant classes, and that most of them 497 were affiliated with acetate oxidizing sulfate reducing bacteria, that outcompete the acetoclastic 498 methanogens. Even though Deltaproteobacteria are best known for their sulfate reduction 499 500 metabolism, they can shift their metabolism in response to depleted sulfate concentrations (Plugge et al., 2011). 501

#### 4.2.2. Betaproteobacteria

503 *Betaproteobacteria* percentages increased slightly with depth. These percentages are comprised of 504 chemoheterotrophs and chemoautotrophs that derive nutrients from decomposition of organic 505 material. *Burkholderiales* was the dominant order in *Betaproteobacteria* and increased with depth 506 (1.5% to 4%). (Schwarz et al., 2007b) showed that *Burkholderiales* is active in the upper sediment 507 of LK. Members of this group are able to carry out anaerobic oxidation of acetate with (per)chlorate 508 as electron acceptor (Yoshida et al., 2005) or oxidize hydrogen in deep subserface(Orcutt et al., 509 2011).

#### 510 *4.2.3. Gammaproteobacteria*

502

511 Our *Gammaproteobacteria* affiliated sequences have different phylogenetic classification 512 distribution in each layer. In the middle and bottom layers members of the *Methylococcales* order 513 were observed (~1%). *Methylococcales* are aerobic methanotrophs which some members were 514 present within the sediments and overlying water column from dysoxic, methane-rich vent and seep 515 systems (Tavormina et al., 2008). The most abundant order in the deep layer was HOC36 (5%)

- 516 which is an uncultured *Gammaproteobacteria*. however when compared to NCBI data base it was
- 517 found to be closely related to uncultured LK clones (99%) and to cultured *Methylocaldum* sp. (94%)
- 518 (Bodrossy et al., 1997) which is a Thermophilic methanotroph isolated from landfill cover soil.

519 4.2.4. *Nitrospirae* 

520 Members of the Nitrospirae phylum, increased with depth. Sequences from our samples were classified to Nitrospirales order in Nitrospirea and were most abundant in the bottom layer of the 521 522 sediment (7%). Part of our sequences were similar to those found in aprevious study (Schwarz et al., 2007b). . Using SIP-RNA, (Schwarz et al., 2007b) showed t not only that Nitrospirae were 523 524 present, but also that they were functionally bioactive. Although Nitrospirae is a known nitrate 525 oxidizer (Ehrich et al., 1995), the conditions of this environment suggest that it utilized another 526 metabolic pathway. Nitrospirae also include the iron reducing candidates such as Magnetobacterium bavaricum (Spring et al., 1993) and sulfur reducers (Sonne-Hansen and Ahring, 527 528 1999). Part of our sequences were aligned to uncultured clone (98%) from freshwater sediment which enhanced degradation of phenanthrene and pyrene by amorphous ferric hydroxide (Yan et 529 530 <del>al., 2012).</del>

531

#### 4.2.5. Methanomicrobia

The majority of our sequences were classified within the Euryarchaeota, in the 532 Methanomicrobia class. Our 16S rRNA sequences were divided between the Methanomicrobiales 533 and Methanosarcinales order. Methanosaeta genus of Methanosarcinales increases with depth. 534 Methanosaeta are acetoclastic methanogens which only able to grow on acetate even at low 535 concentrations (Jetten et al., 1990). Methanosaeta was also shown in previous studies of LK 536 (Schwarz et al., 2007a, 2007b) and in other meso to eutrophic freshwater lakes (Glissman et al., 537 538 2004; Koizumi et al., 2003). Yamada et al. (2014) showed that Methanosaetacan also have the 539 ability to reduce ferrihydrite with H<sub>2</sub> as the electron source. The Methanosaeta concilii was the most similar cultured acetoclastic methanogen (96%) to our sequences which was observed also in 540 541 the previous study. The Methanolinea genus of Methanomicrobiales a hydrogenotrophic methanogen was observed in constant percentages for all depths. Members of the Methanoregula 542 543 genus, a genus that is also hydrogenotrophic methanogen, decreased with depth in the core. Methanoregula, Methanosaeta and Methanolinea represent 60% of sequences of methanogens 544 545 retrieved from freshwater lakes (Borrel et al., 2011).

546 *4.2.6. Thermoplasmata* 

547 The dominant family in *Thermoplasmata* is The Marine Benthic Group D and DHVEG-1
548 (MBG-D). Members of the MBG-D have been shown to exist in a variety of freshwater and marine
549 environments (Beal et al., 2009; Borrel et al., 2012), and it is the most widely encountered,
550 uncultured lineage in freshwater lake sediments. Even though their metabolism is unknown,

hypotheses about their functionalities are based on the environments in which they were found. 551 Methanogenesis was suggested, as they were found in deep lake sediments with high methane 552 concentrations (Borrel et al., 2012), and they were also hypothesized to be involved in AOM, as 553 they were found in AOM zones (Schubert et al., 2011) and in marine seep sediment (Beal et al., 554 555 2009). However, in other environments, in which methane concentrations were low, the utilization of waste products, intermediates, or dead cells by MBG-D was also suggested (Smith et al., 1975). 556 557 Recently, a single cell genomics study showed that members of MBG-D were capable of exogenous protein degradation in cold anoxic environments (Lloyd et al., 2013). 558

#### 559 *4.2.7. Thaumarchaeota*

560 Thaumarchaeota are mostly known as archaeal ammonia oxidizers but in the sediments they may 561 contribute significantly to the reservoir of nitrogen oxides in ocean waters and thus to productivity, including the assimilation of carbon(Pester et al., 2011). The enzyme ammonia monooxygenase, 562 563 which belongs to the enzyme family of copper containing membrane bound monooxgenases that possess wide substrate ranges. Closely related enzymes, which can often perform the same function 564 but with different substrates, include ammonia oxidizing bacteria with ammonia (in 565 Gammaproteobacteria) and methane as a substitute (Lontoh et al., 2000). However, a 566 microorganism's metabolic energy is mainly defined by its downstream enzyme machinery 567 (Tavormina et al., 2011). In our sediment Thaumarchaeota observed in low percentages with 2 568 main classes: Marine Benthic Group B and Miscellaneous Crenarchaeotic Group. 569

#### 570

#### 4.2.8. Miscellaneous Crenarchaeotic Group

571 The Miscellaneous Crenarchaeotic Group (MCG) is a cosmopolitan class assigned to the 572 *Thaumarchaeota* phylum whose members are found mostly in anoxic habitats and have the 573 capability to take up organic carbon (Biddle et al., 2006). Because of their huge environmental 574 range and their complex phylogeny, the MCG collectively possess great metabolic diversity(Jiang 575 et al., 2008). Current evidence for members of the MCG lineage suggests that they may obtain 576 energy from the anaerobic oxidation of methane, but they do so via a dissimilatory methane-577 oxidizing process and they do not assimilate its carbon (Biddle et al., 2006)

#### 578

#### 4.2.9. Marine Benthic Group B

579 Members of the MBGB were detected in a growing number of benthic marine environments, 580 including Atlantic deep sea sediments, marine mud volcanoes, marine carbonate crusts from the 581 Black Sea, organic poor deep subsurface sediments of the central oceanic basins, and organic rich 582 methane or methane hydrate containing sediments near continental margins(Sørensen and Teske, 583 2006). Recent results from Ocean Drilling Program showed that no known methantroph was 584 detected however methane oxidation appeared to be mediated by Marine Benthic Group B and the 585 Miscellaneous Crenarchaeotal Group(Biddle et al., 2006).

#### 4.2 Concept for methane cycle mechanisms

A few possible mechanisms for the novel process of iron driven AOM in the deep part of the 587 sediment can be suggested based on the geochemical profiles and the microbial communities. A 588 possible direct process could be through new, currently unknown bacteria/archaea that reduce iron 589 590 and utilize methane. Possible iIndirect processes could include: 1) Reduction of Fe(III) oxides by oxidation of sulfide (in pyrite or FeS minerals) to sulfur intermediates, and then disproportionation 591 to sulfide and sulfate (such as in (Holmkvist et al., 2011)), and/or sulfate driven AOM but without 592 ANME. 2) Reduction of Fe(III) by utilizing H<sub>2</sub> (Lovley, 1991), could consume most of produced 593  $H_2$  in the deep sediment. Thus which createsing a low concentration of  $H_2$  and which drives 594 methanogenesis backwards ("reverse methanogenesis") (Hallam et al., 2004; Hoehler et al., 1994) 595 by the methanogens archaea in the deep sediment. 3) An oxygenic methane oxidation pathway in an 596 anaerobic environment as described by Ettwig et al. (2010), when methane is oxidized by oxygen 597 that is released from iron oxides. 598

Changes in tThe microbial population shown in the present study changes can be used may to 599 600 support the deep iron driven AOM mechanism (presented by the sample from 29-32cm). The bacterial diversity in this depth was lower than in the methanogenesis zone (6-9 cm) but higher than 601 in the upper layer (0-3 cm) of the sediment. The changes in bacterial diversity with depth could be 602 603 related to the availability of different electron acceptors and of organic matter (Nam et al., 2008). However, there is an overlap between bacterial and archaeal sequences in the middle and the bottom 604 605 layers (PCoA and Venn diagram), indicating that the environmental conditions are affecting the processes. Alpha diversity indexes (Table 1and 2) were compared to previous studies conducted 606 607 only in the upper layer of the sediment of LK using different methods (Schwarz et al., 2007a; Wobus et al., 2003), and showed a similarity in the upper parts of the sediment. However, the 608 deeper layers in this study showed higher diversity and richness than the previous studies (Nusslein 609 et al., 2001; Schwarz et al., 2007a, 2007b) and was close to other studies using 454 sequencing 610 (Hollister et al., 2010; Röske et al., 2012). 611

The direct mechanism of iron reduction coupled with methane oxidation could be performed by 612 a single microorganismean occur by one microorganism, as methanogens have the ability to reduce 613 iron (Van Bodegom et al., 2004; Bond et al., 2002) however not with methane as the electron donor. 614 According to our results, this kind of This direct mechanism could be performed by unique 615 methanotrophy could be fromlike MBG-D (Thermoplasmata class) or- by MCG/MBG-B 616 (Thaumarchaeota phylumwhichthat MCG may represent as new archaeal phylum (Lloyd et al., 617 2013) or as a sister lineage with Thaumarchaeota (Meng et al., 2014). However MBG-B is still 618 classified as Thaumarchaeota (Marlow et al., 2014)), to which some of our sequences were similar 619

to. The representation of sequences similar to About 5% of the archaeal sequences in the middle and 620 deep layers were affiliated to MBG-D, were close to 5% of the archaeal community in the middle 621 and deep layers, however MCG and MBG-B showed much lower representation and will not be thus 622 are not discussed. the Members of the MBG-D have been shown to exist in a variety of freshwater 623 and marine environments (Beal et al., 2009; Borrel et al., 2012), and it is the most widely 624 encountered, uncultured lineage in freshwater lake sediments. Even though their metabolism is 625 unknown, hypotheses about their functionalities are based on the environments in which they were 626 found. Methanogenesis was suggested, as they were found in deep lake sediments with high 627 methane concentrations (Borrel et al., 2012), and they were also hypothesized to be involved in 628 629 AOM, as they were found in AOM zones (Schubert et al., 2011) and in marine seep sediment (Beal et al., 2009). However, inIn other environments, in which methane concentrations were low, the 630 utilization of waste products, intermediates, or dead cells by MBG-D was also suggested (Smith et 631 al., 1975). Recently, a single cell genomics study showed that members of MBG-D were capable of 632 633 exogenous protein degradation in cold anoxic environments (Lloyd et al., 2013). Although their 634 metabolisms are unknown, these groups were found in methane rich environments.

Another option is a consortium of-<u>iron reducing microbes microbes including those belonging</u>
 *Geobacter* genus, which are known for iron reduction, or *Nitrospirales*, known to reduce iron,
 together with methanotrophs or with methanogens capable of "reverse methanogenesis".

The well-known iron reducers are Geobacter genus, however only about 1% of the sequences 638 639 in the middle and bottom layers or the cores were affiliated to the Geobacter genus members. In the deep layer the most abundant class was Sva0485. Sequences of the SVa485 order were similar to 640 641 those found in different aquatic environments, but the metabolic functions of members of this order 642 are not clear. Pelobacter carbinolious (Lovley et al., 1995), a member of SVa485 is capable of Fe(III) and sulfur reduction. Desulfuromonadales, which was found in our sequences in the deep 643 parts of the sediment, was shown as  $S^0$  respiring (Pjevac et al., 2014) and may also reduce Fe(III) 644 and Mn(IV) in marine surface sediments (Lovley, 2006). Schwarz et al. (2007a) showed that in the 645 upper part of LK sediment Deltaproteobacteria was one of the dominant classes, and that most of 646 them were affiliated with acetate oxidizing sulfate reducing bacteria, that outcompete the 647 acetoclastic methanogens. Even though Deltaproteobacteria are best known for their sulfate 648 reduction metabolism, they can shift their metabolism in response to depleted sulfate concentrations 649 (Plugge et al., 2011). 650 In addition, our The results suggest that Desulfuromonadales of Deltaproteobacteria class 651 could be also involved in indirect mechanisms of disproportionation of sulfur together with other 652

653 Deltaproteobacteria sulfate reducer. The dsrAB functional gene shows that sulfate reducers are

654 present at the same level of the core where mcrA functional gene (Fig. 6) of methanogens or

655 methanotrophs are found in the deep part of the sediment. Additionally, the presence of the 656 *Deltaproteobacteria* at the deepest sediment depth and the observed accumulation of acetate with 657 depth (data not shown) could indicate their exploitation of a different metabolic path in the deep

658 <u>sediment than in the upper part of the sediment.</u>

659 AnoOther options for functioning iron reducers areinclude members of the Nitrospirae phylum which that increased with depth. Indeed, Sequences from our samples were classified to 660 Nitrospirales order in Nitrospirea and were most abundant in the bottom layer of the sediment 661 (7%). Part of our sequences were similar to those found in a previous study (Schwarz et al., 2007b). 662 Using SIP-RNA, (Schwarz et al., (2007c2007b) showed not only that Nitrospirae were present, but 663 also that they were functionally bioactive. Although Nitrospirae is a known nitrate oxidizer (Ehrich 664 et al., 1995), the conditions of this environment suggest that it utilized another metabolic pathway. 665 Nitrospirae also include the iron reducing candidates such as Magnetobacterium bavaricum (Spring 666 et al., 1993) and sulfur reducers (Sonne-Hansen and Ahring, 1999). Part of our sequences were 667 aligned to uncultured clone (98%) from freshwater sediment which enhanced degradation of 668 phenanthrene and pyrene by amorphous ferric hydroxide (Yan et al., 2012). 669

The "reverse methanogenesis" pathway could be carried out by the dominant deep layer 670 methanogens Methanosaeta, Methanosaeta are acetoclastic methanogens, which are only able to 671 grow only on acetate, even at low concentrations (Jetten et al., 1990). Methanosaeta was also 672 shown asto be the dominant active methanogens in the upper layer in previous studies of LK 673 (Schwarz et al., 2007a, 2007b) and in other meso to eutrophic freshwater lakes (Glissman et al., 674 2004; Koizumi et al., 2003). Yamada et al. (2014) showed that Methanosaetacan also have the 675 ability to reduce ferrihydrite with H<sub>2</sub> as the electron source. The Methanosaeta concilii was the 676 most similar cultured acetoclastic methanogen (96%) to our sequences which was observed also in 677 678 the previous study.which have been shown in a previous study as the dominant active methanogens in the upper layer (Schwarz et al., 2007b).\_ Methanosaeta could perform a different metabolic 679 680 process or may have been inhibited. The other methanogens that could perform the "revers methanogenesis" are from the Methanomicrobiales order. The Methanolinea genus of 681 Methanomicrobiales a hydrogenotrophic methanogen was observed in constant percentages for all 682 depths. Members of the Methanoregula genus, a genus that is also hydrogenotrophic methanogen, 683 684 decreased with depth in the core. qPCR analysis of mcrA functional gene of methanogens/anaerobic methanotroph shows that in the middle layer methanogens were more abundant than in the upper 685 and bottom layers (Fig 6). ANME were not found in the sediment using specific primers and 686 687 Fluorescent in situ hybridization methods. An additional reason for not finding any ANME sequences in our samples is likely because 454 sequences related to methanogens were found in 688

high percentages and they are probably that dominant in the <u>qPCR analysis of mcrA gene</u> observed
 in the deep layer belongs to the methanogens which maybe cause the reverse methanogenesis.

The results suggest that Desulfuromonadales of Deltaproteobacteria class could be involved in 691 indirect mechanisms of disproportionation of sulfur together with other Deltaproteobacteria sulfate 692 693 reducer. The dsrAB functional gene shows that sulfate reducers are present at the same level of the core where mcrA functional gene (Fig. 6) of methanogens or methanotrophs are found in the deep 694 part of the sediment. Additionally, the presence of the Deltaproteobacteria at the deepest sediment 695 depth and the observed accumulation of acetate with depth (data not shown) could indicate their 696 exploitation of a different metabolic path in the deep sediment than in the upper part of the 697 sediment. 698

699 The third additional indirect mechanism of anaerobic methane oxidation via an oxygenic pathway was shown clearly by the *pmoA* functional gene. The *pmoA* concentration in the deep part 700 701 of the sediment was higher than in the upper part, indicating an oxygenic pathway (Fig. 6). However, pMMO is a homologue enzyme of ammonia monooxygenase and might be sequenced 702 together with pMMO (Tavormina et al., 2011). The source of pMMO could be Methylococcales 703  $(\sim 1\%)$ , of *Gammaproteobacteria*). which were observed in the middle and bottom sediment layers. 704 Methylococcales are aerobic methanotrophs, which some members were present within the 705 sediments and overlying water column from dysoxic, methane-rich vent and seep systems 706 (Tavormina et al., 2008). The most abundant order in the deep layer was HOC36 (5%), which is an 707 uncultured Gammaproteobacteria. hHowever, when compared to NCBI data base, it was found to 708 be closely related to uncultured LK clones (99%) and to cultured Methylocaldum sp. (94%) 709 (Bodrossy et al., 1997), which is a Thermophilic methanotroph isolated from landfill cover soil. 710 711 (Gammaproteobacteria), which is an aerobic methanotroph found in the deep part of our samples. 712 However, bias of *pmoA* with ammonia monooxygenase could be cased caused by *Thaumarchaeota*, Betaproteobacteria or Nitrospirae, which were found in all depth. 713

714 Thaumarchaeota comprise not only all known archaeal ammonia oxidizers, but also several clusters of environmental sequences representing microorganisms with unknown energy 715 716 metabolisms (Pester et al., 2011). Members of *Thaumarchaeota* phylum could have monooxgenases 717 like enzymes that are able to capture methane due to the enzymes' phylogenetic proximities to 718 methane monooxgenases, but that activity requires the necessary downstream metabolic pathway. If they could have that ability, they would have been good candidates for group of methanotrophic 719 720 archaea. This indirect pathway can be similar to that found in the NC10 phylum (Zhu et al., 2012), 721 which produces oxygen via the reduction of nitrite and the oxidation of methane, but with iron oxides. Also many bacterial ammonia oxidizer sequences were found in our environment. 722 723 Ammonium profiles show increase with depth mainly due to decomposition of the organic matter,

and theoretically the large amount of ammonium could be oxidized by ferric iron minerals and produce nitrite (Clement et al., 2005). Maybe ammonia monooxygenase can function for ammonium up take for iron reduction and for methane up take for oxidation, but that is only a speculation. However, no nitrate or nitrite was detected in the deep sediment, moreover no NC10 phylum (Ettwig et al., 2010) was observed in our sequences.

To summarize, this study attempted to correlate between the performed geochemical and 729 microbial profiles in lake sediments. The geochemical data suggest three main depth related zones 730 of electron acceptor activities in the sediment: sulfate reduction, methanogenesis and a novel, deep 731 732 iron-driven AOM. The prokaryotic analysis provided clues regarding the microorganisms that may be involved in this novel process and the metabolic paths that occur throughout the microbial 733 assemblage. For AOM via iron reduction to occur, a number of potential pathways and their 734 combinations have been suggested. Orders that become enriched (Sva0485, Methanosarcinales and 735 736 *Nitrospirales*) with depth can be assumed to participate in the AOM process either directly or indirectly. A possible direct process could be through new, currently unknown bacteria/archaea that 737 reduce iron and utilize methane, which may be carried out by a MBG-D MCG-as a methanotroph in 738 a consortium with an iron reducer like Geobacter. Possible indirect processes could be Fe(III) 739 reduction by sulfide, oxidation of the sulfide to elemental sulfur and other sulfur intermediates and 740 then disproportionation to sulfide and sulfate and sulfate driven AOM. However, this process is less 741 likely because ANMEs were not found. It could also be via reduction of Fe(III) by utilizing H<sub>2</sub>, 742 creating a low concentration of H<sub>2</sub> and driveingdriving "reverse methanogenesis". Fe(III) reduction 743 processes could be carried out by Nitrospirae and/or Deltaproteobacteria. Members of both groups 744 can reduce iron while in a consortium with methanogenic *Methanosarcinales*/ *Methanomicrobiales*, 745 or can reduce the iron with sulfur minerals, creating sulfate (see above) that Deltaproteobacteria 746 747 can utilize while in a consortium with a MBG-D MCG as a methanotrophic archaea (Schubert et al., 2011). An oxidation of methane coupled to iron reduction pathway as described by Ettwig et al., 748 749 (2010) could occur by Thaumarchaeota or Nitrospirae with a monooxygenase enzyme that can utilize methane while using iron oxides to generate the oxygen needed to oxidize the methane. 750 751 Further research involving even larger sample of the microbial community and characterization of more diverse functional genes will provide better indication for the composition of the microbial 752 753 communities at the different depths. Also, microbial and geochemical experiments have the potential to give more clues on the mechanism of this novel iron driven AOM. 754

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## 1008 Tables

**Table 1**. Bacterial sequences used for classification in SILVA ngs and for subsample in MOTHUR for thealpha diversity.

Bacteria	# seq SILVA ngs	OTUs SILVA ngs	# seq MOTHUR	<b>OTUs MOTUR</b>	coverage	invsimpson	chao	ace	shannon
0-3 cm	3631	599	2588	388	0.92	10/10.8/11.7	805/925.7/1092.8	1246.5/1384.0/1546.7	3.83/3.91/3.98
6-9 cm	4641	1735	3365	1337	0.77	286.6/315.9/351.8	2437.9/2673.0/2958.4	3876.8/4135.2/4420.0	6.48/6.53/6.57
29-32 cn	5038	1516	3615	1214	0.81	127.8/142.4/160.6	2104.4/2318.7/2583.1	3047.7/3259.5/3495.4	6.13/6.19/6.24

#### 1012 **Table 2**. Archaeal sequences used for classification in SILVA ngs and for subsample in MOTHUR for the

alpha diversity.

1014

Archaea	# seq SILVA ngs	OTUs SILVA ngs	# seq MOTHUR	OTUs MOTHUR	coverage	invsimpson	chao	ace	shannon
0-3 cm	288	48	268	32	0.90	2.5/2.9/3.5	53.9/83.4/164.5	232.8/344.6/520.4	1.6/1.8/2.0
6-9 cm	1382	184	1408	200	0.86	8.4/10.2/13.2	39.0/131.4/220.4	144.7/195.2/277.4	2.87/3.05/3.22
29-32 cm	3110	383	2829	245	0.88	7.9/9.2/10.9	73.5/104.8/181.7	133.2/190.9/289.6	2.58/2.74/2.91

#### 1015 **Figure Captions:**

Figure 1: Schematic location of Lake Kinneret. Numbers correspond to altitude measured in meters
from the level of Lake Kinneret. Cores were taken from station A (taken from Hambright et al.,
2004).

Figure 2: Geochemical porewater profiles in LK sediment. A. profile of  $SO_4^{-2}$  (black circles) and 1019 sulfide (gray diamonds) in the porewater **B**. Headspace measurements of methane (gray circles) and 1020  $\delta^{13}C_{CH4}$  (black squares) in the sediments C.  $\delta^{13}C$  of total lipids extraction (TLE) (black diamonds) 1021 from the sediment, and  $\delta^{56}$ Fe (gray circles) of the dissolved iron in the porewater. **D**. Electron 1022 acceptor profiles of dissolved Fe(II) (gray triangles) and Mn (II) (black squares) in the porewater,. 1023 Black arrows indicate the sampled sections for 16S rRNA gene analysis. The presented methane 1024 profile was taken two weeks prior to the sampling for the microbial communities on December 1025 2009. Sampling for Fe(II), Mn(II) and  $SO_4^{-2}$  profiles was carried out between 2007 and 2011 and 1026 for sulfide from 2007 to 2013. The  $\delta^{13}C_{CH4}$  and  $\delta^{56}$ Fe profiles were performed four months prior to 1027 the microbial sampling on August 2009. 1028

**Figure 3**: Classification of bacterial sequences using SILVA ngs pipeline. Phyla and classes distribution of sequences of the 454 sequencing at the different depths above 1% in the communities.

**Figure 4**: Classification of archaeal sequences using SILVA ngs pipeline. Phyla and orders distributions of sequences of the 454 sequencing at the different depths above 1% in the communities.

Figure 5: Principal coordinate analysis (PCoA) of the A. bacterial and B. archaeal community
matrix (based on operational taxonomic units) and vector fitting of the environmental geochemical
conditions in the different depth. The bacterial community from the upper layer is labeled with red
triangle, the middle layer is labeled with green triangle and bottom layer is labeled with blue
triangle.

Figure 6: profiles of functional genes from the 3 different depths. Green bar represent *pmoA*functional gene of aerobic methanotroph. Red bar represent *dsrAB* functional gene of sulfate
reducers and blue bar represent *mcrA* functional gene from methanogens and anaerobic
methanotrophs.

1044



Figure 1.



















Figure 6

## 1076 Supplementary





Figure 2

**Supplementary Fig. 2.** Venn diagram showing bacterial sequences overlap between the different depths. The green circle represents the top sample (0-3 cm), the blue circle represents the middle sample (6-9 cm) and the red circle represents the bottom sample (29-32 cm).

1090 Supplementary Table 1. The classification percentage and number of sequences of archaea A.

1091 phylum, B. class and C. order by SILVA ngs.

Archaea	0-3 cm		6-9	cm	29-32 cm		
Phylum	# seq	%	# seq	%	# seq	%	
Euryarchaeota	283	98.3	1430	96.4	3048	98.0	
Thaumarchaeota	5	1.7	53	3.6	62	2.0	

1052
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Archaea	0-3	0-3 cm		6-9 cm		29-32 cm	
Class	#seq	%	# seq	%	# seq	%	
AK59					1	0.0	
AK8			1	0.1	4	0.1	
Group C3			2	0.1	2	0.1	
Halobacteria	7	2.4	96	6.5	165	5.3	
Marine Benthic Group B	2	0.7	24	1.6	30	1.0	
Marine Group I					2	0.1	
Methanobacteria	1	0.3					
Methanomicrobia	266	92.4	1164	78.5	2679	86.1	
Miscellaneous Crenarchaeotic Group	3	1.0	21	1.4	19	0.6	
pSL12			2	0.1	3	0.1	
South African Gold Mine Gp 1(SAGMCG-1)			3	0.2	1	0.0	
Thermoplasmata	9	3.1	170	11.5	204	6.6	

1093

Archaea	0-3	0-3 cm 6-9		cm	29-3	2 cm
Order	# seq	%	# seq	%	# seq	%
20a-9			5	0.3	3	0.1
ANME-1					1	0.0
Cenarchaeales					2	0.1
Halobacteriales	7	2.4	96	6.5	165	5.3
Methanobacteriales	1	0.3				
Methanomicrobiales	202	70.1	422	28.5	878	28.2
Methanosarcinales	64	22.2	742	50.0	1800	57.9
Thermoplasmatales	8	2.8	165	11.1	201	6.5
WCHA1-57	1	0.3				
Uncultured	5	1.7	53	3.6	60	1.9

1094

**Supplementary Table 2.** The classification percentage and number of sequences of bacterial A.

1096 phylum, B. class and C. order by SILVA ngs.

Bacteria	0-3	cm	6-9 cm		29-32 cm	
Phylum	# seq	%	# seq	%	# seq	%
Acidobacteria	33	0.9	112	2.4	123	2.4
Actinobacteria	56	1.5	61	1.3	90	1.8
Armatimonadetes	21	0.6	12	0.3	14	0.3
Bacteroidetes	80	2.2	349	7.5	126	2.5
BD1-5	2	0.1	2	0.0	2	0.0
Caldiserica					2	0.0
Candidate division BRC1	2	0.1	14	0.3	8	0.2
Candidate division JS1	1	0.0	2	0.0		
Candidate division KB1					3	0.1
Candidate division OD1			1	0.0	18	0.4
Candidate division OP11	2	0.1	17	0.4	1	0.0
Candidate division OP3	1	0.0	108	2.3	84	1.7
Candidate division OP8	9	0.2	81	1.7	106	2.1
Candidate division SR1			4	0.1		
Candidate division TM7	6	0.2	5	0.1		
Candidate division WS3	15	0.4	70	1.5	103	2.0
Chlorobi	38	1.0	235	5.1	206	4.1
Chloroflexi	65	1.8	672	14.5	969	19.2
CKC4	1	0.0				
Cyanobacteria	18	0.5	69	1.5	21	0.4
Deferribacteres	1	0.0	122	2.6	126	2.5
Deinococcus-Thermus	1	0.0	8	0.2	9	0.2
Elusimicrobia			7	0.2	13	0.3
Fibrobacteres			20	0.4	9	0.2
Firmicutes	1376	37.9	140	3.0	152	3.0
Gemmatimonadetes	2	0.1	15	0.3	15	0.3
Hyd24-12	4	0.1	8	0.2		
GOUTA4					4	0.1
JL-ETNP-Z39			1	0.0	3	0.1
Lentisphaerae	12	0.3	38	0.8	10	0.2
Nitrospirae	31	0.9	195	4.2	409	8.1
NPL-UPA2	1	0.0	33	0.7	5	0.1
OC31			1	0.0		
Planctomycetes	22	0.6	140	3.0	172	3.4
Proteobacteria	1762	48.5	1837	39.6	1954	38.8
SHA-109	6	0.2	6	0.1	1	0.0
Spirochaetae	1	0.0	102	2.2	134	2.7
Synergistetes					1	0.0
TA06	5	0.1	66	1.4	102	2.0
Tenericutes	13	0.4	1	0.0	1	0.0
Thermotogae			8	0.2	1	0.0
TM6	19	0.5	27	0.6	21	0.4
Verrucomicrobia	3	0.1	39	0.8	20	0.4
WCHB1-60	22	0.6	13	0.3		

Bacteria	0-3 cm		6-9 cm		29-32 cm	
CLASS	# seq	%	# seq	%	#seq	%
Acidimicrobiia	5	0.1	4	0.1	2	0.0
Acidobacteria	26	0.7	80	1.7	83	1.6
Actinobacteria	/	0.2	5	0.1	5	0.1
	23	0.0	20	6.4	375	7.4
Ardenticatenia	51	0.5	1	0.4	5/5	7.4
ARKDMS-49			2	0.0		
ARKICE-90			11	0.2		
Bacilli	611	16.8	23	0.5	49	1.0
Bacteroidia			5	0.1	1	0.0
BD2-2	2	0.1	16	0.3	8	0.2
Betaproteobacteria	65	1.8	268	5.8	344	6.8
BS5	3	0.1	7	0.2		
BSV13			9	0.2	1	0.0
	1	0.0	8	0.2	3	0.1
Caldisericia	6	0.2	42	0.0	<u> </u>	0.0
Chloroplast	3	0.2	42	0.9	10	0.4
Clostridia	761	21.0	116	2.5	99	2.0
Coriobacterija	3	0.1	4	0.1	7	0.1
Cvanobacteria	13	0.4	22	0.5	7	0.1
Cytophagia	1	0.0	4	0.1	8	0.2
Deferribacteres	1	0.0	122	2.6	126	2.5
Dehalococcoidia	2	0.1	276	5.9	453	9.0
Deinococci	1	0.0	8	0.2	9	0.2
Deltaproteobacteria	57	1.6	1257	27.1	1022	20.3
DEV055	6	0.2	9	0.2		
Elusimicrobia			7	0.2	13	0.3
Epsilonproteobacteria			12	0.3	1	0.0
Erysipelotrichia			ļ		3	0.1
FFCH16263	1	0.0			2	0.0
Fibrobacteria			20	0.4	9	0.2
Flavobacteriia	9	0.2	34	0.7	3	0.1
Gammaproteobacteria	1614	44.5	257	5.5	536	10.6
Gemmatimonadetes	2	0.1	15	0.3	15	0.3
Holophagae	5	0.1	24	0.5	29	0.6
	32	0.9	193	4.2	188	3.7
	0	0.2	12	0.3	6	0.1
Ktedonobacteria	9	0.2	5	0.1	4	0.1
LD1-PB3		0.0	1	0.0	1	0.0
Lentisphaeria	1	0.0	3	0.0		0.0
MB-A2-108	6	0.2	4	0.1	14	0.3
MBMPE71	-		1	0.0	1	0.0
Melainabacteria	2	0.1	1	0.0	4	0.1
ML635J-21			4	0.1	6	0.1
Mollicutes	13	0.4	1	0.0	1	0.0
MSB-5B2					4	0.1
MSBL3			6	0.1		
Negativicutes	4	0.1	1	0.0	1	0.0
Nitrospira	31	0.9	195	4.2	409	8.1
OM190	1	0.0	4	0.1	2	0.0
Oligosphaeria	-	0.4	7	0.2	5	0.1
	3	0.1	25	0.5	11	0.2
OPB41 Opituteo	20	0.6	21	0.5	35	0.7
	2	0.1	12	0.3	1	0.1
Phycisphaerae	11	0.3	101	22	130	26
Pla3 lineage		0.0	10	0.2	7	0.1
Pla4 lineage			9	0.2	. 8	0.2
Planctomycetacia	10	0.3	15	0.3	19	0.4
Proteobacteria Incertae Sedis			1	0.0	1	0.0
S085					1	0.0
SB-1			7	0.2	10	0.2
SB-5	19	0.5	33	0.7	12	0.2
S-BQ2-57 soil group					1	0.0
SHA-26			7	0.2	14	0.3
Spartobacteria			1	0.0	1	0.0
Sphingobacteriia	30	0.8	95	2.0	30	0.6
Spirochaetes	1	0.0	102	2.2	134	2.7
Subgroup 22	2	0.1	8	0.2	11	0.2
	2	0.1	Л	0.1		0.0
Thermoleophilia	3 14	0.1	4 22	0.1	25	0.5
Thermotogae	14	0.4	2 	0.0	20	0.5
TK10			0	0.2	2	0.0
vadinHA17	18	0.5	145	31	51	1.0
vadinHA49	10	0.0	1-5	0.1	5	0.1
VC2.1 Bac22			1	0.0		0.1
Verrucomicrobiae			1	0.0		
WCHB1-41			1	0.0	2	0.0
WCHB1-32	1	0.0			1	0.0
uncultured	138	3.8	538	11.6	585	11.6

Bacteria	0-3 cm		6-9 cm		29-32 cm	
Order	# seq	%	# seq	%	# seq	%
10bav-F6					2	0.0
43F-1404R			2	0.0	3	0.1
Acidimicrobiales	5	0.1	4	0.1	2	0.0
Alteromonadales			9	0.2	14	0.3
Amsterdam-1B-07			1	0.0	4	0.1
Anaerolineales	31	0.9	296	6.4	375	7.4
B1-7BS			1	0.0		
B276-D12			1	0.0		
Bacillales	602	16.6	23	0.5	47	0.9
Bacteroidales			5	0.1	1	0.0
BD2-11 terrestrial group			2	0.0	5	0.1
Bdellovibrionales			34	0.7	13	0.3
BP-U1C-1g10			1	0.0		
Burkholderiales	56	1.5	86	1.9	208	4.1
C86					2	0.0
Caldilineales	1	0.0	8	0.2	3	0.1
Caldisericales					2	0.0
Campylobacterales			12	0.3	1	0.0
Caulobacterales	12	0.3	1	0.0	1	0.0
CCM11a			9	0.2	6	0.1
Chlorobiales	6	0.2	42	0.9	18	0.4
Chromatiales	3	0.1	11	0.2	21	0.4
Chthoniobacterales			1	0.0	1	0.0
Clostridiales	760	20.9	107	2.3	99	2.0
Coriobacteriales	3	0.1	4	0.1	7	0.1
Corynebacteriales	1	0.0	1	0.0	3	0.1
Cytophagales	1	0.0	3	0.1		
D8A-2	1	0.0				
Deferribacterales	1	0.0	122	2.6	126	2.5
Dehalococcoidales			6	0.1	2	0.0
Deltaproteobacteria Incertae Sedis	2	0.1	55	1.2	2	0.0
Desulfarculales	12	0.3	132	2.8	62	1.2
Desulfobacterales	13	0.4	118	2.5	88	1.7
Desulfuromonadales			80	1.7	41	0.8
Erysipelotrichales					3	0.1
Fibrobacterales			18	0.4	6	0.1
Flavobacteriales	9	0.2	34	0.7	3	0.1
Frankiales	1	0.0	2	0.0	3	0.1
FS117-23B-02			2	0.0	4	0.1
FW22	1	0.0	2	0.0	2	0.0
Gaiellales	11	0.3	19	0.4	20	0.4
Gammaproteobacteria Incertae Sedis			4	0.1	4	0.1
Gastranaerophilales			1	0.0	4	0.1
Gemmatimonadales	2	0.1	13	0.3	10	0.2
GIF3			2	0.0	7	0.1
GIF9			117	2.5	251	5.0

GR-WP33-30	1	0.0	14	0.3	34	0.7
Haloplasmatales	13	0.4				
HOC36	2	0.1	72	1.6	267	5.3
Holophagae Incertae Sedis			5	0.1	12	0.2
Holophagales			2	0.0	2	0.0
Hydrogenophilales			48	1.0	44	0.9
Ignavibacteriales	32	0.9	193	4.2	188	3.7
KCLunmb-38-53	1	0.0	9	0.2		
KD3-62	1	0.0	8	0.2	9	0.2
KI89A clade			4	0.1	5	0.1
Lactobacillales	2	0.1			2	0.0
Legionellales	5	0.1	17	0.4	15	0.3
Lineage IIb					4	0.1
Lineage IIc			5	0.1	4	0.1
Lineage IV			1	0.0	1	0.0
MD2894-B20			1	0.0	3	0.1
Methylococcales	9	0.2	70	1.5	69	1.4
Methylophilales			27	0.6	10	0.2
Micrococcales	1	0.0	2	0.0	1	0.0
mle1-8	6	0.2	5	0.1	8	0.2
MSB-3A7 sediment group			3	0.1	3	0.1
MSBL8			1	0.0		
MSBL5	1	0.0	139	3.0	169	3.4
MSBL9	1	0.0	22	0.5	59	1.2
MVP-21	1	0.0				
MVP-88					1	0.0
Myxococcales	1	0.0	82	1.8	45	0.9
NB1-n			1	0.0	1	0.0
Nitrosomonadales	2	0.1	36	0.8	22	0.4
Nitrospirales	31	0.9	195	4.2	409	8.1
ODP1230B30.09					1	0.0
Oceanospirillales			1	0.0		
Oligosphaerales			7	0.2	5	0.1
Opitutales			12	0.3	7	0.1
Order II (Bacteroidetes)			1	0.0	5	0.1
Order Incertae Sedis			2	0.0		
PBS-18			3	0.1	1	0.0
PeM15	1	0.0				
Phycisphaerales	2	0.1	10	0.2	10	0.2
Pla1 lineage			18	0.4	17	0.3
Planctomycetales	10	0.3	15	0.3	19	0.4
Propionibacteriales	3	0.1				
possible order 07			2	0.0	3	0.1
Pseudomonadales	1523	41.9	1	0.0		
Rhizobiales	6	0.2	16	0.3	45	0.9
Rhodobacterales	1	0.0	1	0.0		
Rhodocyclales	4	0.1	43	0.9	29	0.6
Rhodospirillales			3	0.1	1	0.0
Rickettsiales	3	0.1	4	0.1		

S15A-MN16	1	0.0	18	0.4	14	0.3
S-70			3	0.1		
SAR324 clade(Marine group B)			2	0.0		
SC-I-84	2	0.1	27	0.6	31	0.6
Sh765B-AG-111					2	0.0
Sh765B-TzT-29			1	0.0	3	0.1
SHA-43			2	0.0	6	0.1
Selenomonadales	4	0.1	1	0.0	1	0.0
Solirubrobacterales	3	0.1	4	0.1	5	0.1
Sphingobacteriales	30	0.8	95	2.0	30	0.6
Sphingomonadales	1	0.0			3	0.1
Spirochaetales	1	0.0	98	2.1	133	2.6
Streptomycetales					1	0.0
Subgroup 17	1	0.0	3	0.1		
Subgroup 18	3	0.1	27	0.6	21	0.4
Subgroup 23	2	0.1	6	0.1	7	0.1
Subgroup 4	2	0.1				
Subgroup 6	20	0.6	31	0.7	25	0.5
Subgroup 7	2	0.1	3	0.1		
Subgroup 13			5	0.1	5	0.1
Subgroup 19			1	0.0	13	0.3
Subgroup 2			1	0.0	6	0.1
Subgroup 21			9	0.2		
Subgroup 25			1	0.0	6	0.1
Subgroup 3			1	0.0	1	0.0
Subgroup 11					1	0.0
Subgroup 9					1	0.0
SubsectionII (Cyanobacteria)					1	0.0
SubsectionI (Cyanobacteria)	6	0.2	8	0.2	3	0.1
SubsectionIV (Cyanobacteria)	7	0.2	14	0.3	3	0.1
Sva0485	20	0.6	390	8.4	546	10.8
Synergistales					1	0.0
Syntrophobacterales	8	0.2	347	7.5	183	3.6
SZB30	4	0.1	9	0.2	7	0.1
Thermoanaerobacterales			9	0.2		
Thermotogales			8	0.2	1	0.0
Thiotrichales	3	0.1	5	0.1	3	0.1
TPD-58	1	0.0	6	0.1	8	0.2
TRA3-20	1	0.0				
vadinBA26			8	0.2	16	0.3
Verrucomicrobiales			1	0.0		
Vampirovibrionales	2	0.1				
VAN12	7	0.2				
Victivallales	1	0.0	3	0.1		
WD2101 soil group			1	0.0		
Xanthomonadales	65	1.8	50	1.1	129	2.6
uncultured	237	6.5	951	20.5	816	16.2

#### 1100 Appendix

1101 The major bacterial populations in the sediment:

1102 Proteobacteria

*Proteobacteria* was the most dominant bacterial phylum in all the depths. Their percentage was slowly decreases with depth however the composition within its classes were changing with depth. *Proteobacteria* is also among the most abundant phyla in LK (Schwarz et al., 2007a) and in other freshwater sediments (Tamaki et al., 2005; Wobus et al., 2003). *Proteobacteria* is a very diverse phylum, divided in to number of classes which most of them have been observed in LK sediments.

1108 Betaproteobacteria

Betaproteobacteria percentages were slowly increasing with depth. They are comprised of 1109 1110 chemoheterotrophs and chemoautotrophs which derive nutrients from decomposition of organic material. Burkholderiales was the dominant order in Betaproteobacteria and increased with depth 1111 1112 (1.5% to 4%). Schwarz et al. (2007a) showed that Burkholderiales is active in the upper sediment of 1113 LK. Members of this group are able of anaerobic oxidation of acetate with (per)chlorate as electron acceptor (Yoshida et al., 2005) or oxidize hydrogen in deep subserface(Orcutt et al., 2011). 1114 Hydrogenophilales and Nitrosomonadales orders were observed in lower abundant but in the 1115 middle and deep layer of the sediment. Hydrogenophilales can utilize hydrogen (Ontiveros-1116 Valencia et al., 2013) and *Hydrogenophilaceae* family are obligatory chemolithoautotrophic, 1117 aerobic or facultative anaerobic sulfur oxidizers and reduction of nitrate (Kelly and Wood, 2000) 1118 which are often found also in wastewater treatment systems(Luo et al., 2011). Nitrosomonadales 1119 are freshwater bacteria which presumed to have a role in ammonia oxidation and/or denitrification 1120 processes (Edlund et al., 2008). McBeth et al. (2013) also showed that Nitrosomonadales are 1121 commonly found as freshwater Fe oxidizers. It should be noted that the upper sediment may 1122 1123 contain denitrifying bacteria that arrived from the water column where nitrification conditions exists throughout the year. 1124

#### 1125 Gammaproteobacteria

Our sequences which were affiliated to the *Gammaproteobacteria* have different phylogenetic
classification distribution in each layer. The upper layer was dominated by *Pseudomonadales* (42%)
which were divided to 2 main genera: *Acinetobacter* (28%) and *Pseudomonas* (14%). *Acinetobacter*was shown to have the ability to degrade organic matter. Our sequences which similar to *Acinetobacter genus were closest to Acinetobacter lwoffii* (99%) found in sewage and growndwater
(Nazina et al., 2000). our sequences classified as *Pseudomonas were* similar to *Pseudomonas*

1132 *stutzeri* (99%) which has the ability to oxidize a large variety of organic compounds, and some

members also involved in nitrification and denitrification, degradation of aromatic compounds, and 1133 nitrogen fixation (Chen et al., 2011). In the middle and bottom layers *Methylococcales* order was 1134 observed (~1%). *Methylococcales* are aerobic methanotrophs which some members were present 1135 within the sediments and overlying water column from dysoxic, methane-rich vent and seep 1136 1137 systems (Tavormina et al., 2008). The most abundant order in the deep layer was HOC36 (5%) which is an uncultured Gammaproteobacteria. however when compared to NCBI data base it was 1138 closly related to uncultured LK clones (99%) and to cultured *Methylocaldum* sp. (94%) (Bodrossy 1139 et al., 1997) which was a Thermophilic methanotroph isolated from landfill cover soil. 1140

#### 1141 *Deltaproteobacteria*

1142 *Deltaproteobacteria* were the most abundant in the middle and deep sediment samples. In the

1143 middle layer the abundant classes were *Syntrophobacterales* (8%), *Desulfarculales* (3%),

1144 *Desulfobacterales* (2%) and *Desulfuromonadales* (2%). Many members of the families of

1145 *Syntrophobacterales* are sulfate reducers or as fermentative organism, depending on the

1146 environmental conditions (McInerney et al., 2007). Syntrophaceae the dominant family of

1147 *Syntrophobacterales* are acetate degrading sulfate reducing bacterium (Jiang et al., 2009).

1148 *Desulfarculales* are strictly anaerobic are mesophilic sulfate-reducing bacterium with the capability

1149 to oxidize acetate and fatty acids (Sun et al., 2010). *Desulfobacteralesare* sulfate reducers capable

1150 to oxidize hydrogen in soils and sediments (Burow et al., 2014). Desulfuromonadales was shown as

1151  $S^{\theta}$  respiring (Pjevac et al., 2014) and may also reduce Fe(III) and Mn(IV) in marine surface

sediments (Lovley, 2006). About 1% of the sequences in the middle and bottom layers were

1153 affiliated to Geobacter genus which involve in iron reduction. In the deep layer the most abundant

1154 class was Sva0485. Sequences of the SVa485 order were similar to those found in different aquatic

- 1155 environments, but the metabolic functions of members of this order are not clear. *Pelobacter*
- 1156 *carbinolious* (Lovley et al., 1995), a member of SVa485 capable of Fe(III) and sulfur reduction.

1157 Schwarz et al. (2007a) showed in the upper part of LK sediment that *Deltaproteobacteria* was one

1158 of the dominant classes, most of which were affiliated with acetate oxidizing sulfate reducing

1159 bacteria, which outcompete the acetoclastic methanogens. Even though *Deltaproteobacteria* are

1160 best known for their sulfate reduction metabolism, they can shift their metabolism in response to

1161 depleted sulfate concentrations (Plugge et al., 2011).

#### 1162 *Chloroflexi*

1163 *Chloroflexi* was found to be one of the most abundant bacterial phyla in LK and represented diverse

1164 functional groups that increased with depth. The abundant classes that increase with depth were

1165 Anaerolineae and Dehalococcoidia. Although they are found in a variety of anaerobic

- 1166 environments, only a few representatives have been cultivated so far (Yamada and Sekiguchi,
- 1167 2009). The metabolisms of the cultured representatives showed that they utilized organic matter, but
- 1168 no specific electron acceptor has as yet been reported (Yamada et al., 2006). *Chloroflexi* was one of
- 1169 the most dominant phyla in contaminated soil environment which had a lot of polycyclic aromatic
- 1170 hydrocarbons (Winderl et al., 2008). Members of *Chloroflexi* were found throughout the entire core,
- 1171 indicating that Anaerolineae representatives could be the main aromatic organic matter
- 1172 decomposers in LK. *Dehalococcoidia* are widely distributed in the marine subsurface, yet metabolic
- 1173 properties of the many uncultivated lineages are completely unknown. *Dehalococcoidia* could be
- 1174 capable of oxidizing various fatty acids and/or structurally related substrates (Wasmund et al.,
- 1175 2013) or reductive dechlorination of chlorinated hydrocarbon(Maymo et al., 1995).

#### 1176 *Nitrospirae*

Nitrospirae phylum, were increased with depth. Sequences from our samples were classified to 1177 Nitrospirales order in Nitrospirea which were most abundant in the bottom layer of the sediment 1178 (7%). Part of our sequences were similar to previous study (Schwarz et al., 2007b). (Schwarz et al., 1179 2007b) showed by SIP-RNA not only that Nitrospirae was present, but also that it was functionally 1180 bioactive. Although Nitrospirae is a known nitrate oxidizer (Ehrich et al., 1995), the conditions of 1181 1182 this environment suggest that it utilized another metabolic pathway. Nitrospirae also include the 1183 iron reducing candidates Magnetobacterium bavaricum (Spring et al., 1993) and sulfur reducers (Sonne Hansen and Ahring, 1999). Part of our sequences were aligned to uncultured clone (98%) 1184 1185 from freshwater sediment which enhanced degradation of phenanthrene and pyrene by amorphous ferric hydroxide (Yan et al., 2012). 1186

#### 1187 *Chlorobi*

The Chlorobi, green sulfur bacteria, are a group of anoxygenic photosynthetic bacteria observed 1188 frequently in anoxic microbial mats and meromictic lakes. Chlorobi in found in deeper layers of the 1189 sediment since they support higher H<sub>2</sub>S concentrations and require less light intensity. They 1190 photosynthesize using H<sub>2</sub>S as electron donor, although they can also grow photoheterotrophically, 1191 1192 and some of them form symbiotic consortia with heterotrophic partners that are widely distributed in chemoclines of meromictic lakes throughout the world (Briée et al., 2007). Chlorobi metabolic 1193 1194 features include fixing CO<sub>2</sub>, fixing N<sub>2</sub> and oxidize sulfide or other reduced sulfur compounds (Liu et al., 2012). Most our sequences were classified to Ignavibacteria order which capable of 1195 1196 organoheterotrophy under both oxic and anoxic conditions.

1197 *Firmicutes* 

*Firmicutes* are mostly abundant in the upper part of the sediment. Our sequences were classified to 1198 2 main orders: Bacillales and Clostridia. Bacillales have been observed in diverse environments 1199 and with diverse metabolic functions. Our sequences were closely related to a few species: Bacillus 1200 thioparans (99%) which grew chemolithoautotrophically by oxidation of thiosulfate to sulfate and 1201 1202 was found in deep sea sediment (Pérez-Ibarra et al., 2007). Bacillus subterraneus (99%) utilize amorphous iron(III), Mn(II)(IV), nitrate, nitrite and fumarate as electron acceptors (Kanso et al., 1203 2002). *Clostridia* are anaerobic bacteria which play an important role in the biogeochemical cycling 1204 of carbon, sulfur and iron. Clostridia are key microbial drivers in iron reduction in estuarine 1205 sediments and acid mine drainage environments(Shah et al., 2014). Part of our sequences were 1206 closely related to *Clostridium tunisiense* (96%) which uses elemental sulfur but not sulfate, 1207 1208 thiosulfate, nor sulfite as terminal electron acceptors(Thabet et al., 2004).

1209 Bacteroidetes

1210 Sequences of *Bacteroidetes* were mostly abundant in the middle layer of the LK sediment core.

1211 Bacteroidetes are known as hydrolytic fermentative bacteria, they are major utilizers of high-

1212 molecular-mass dissolved organic matter in marine ecosystems (Cottrell and Kirchman, 2000).

1213 Therefore, fresh organic matter (high in high molecular mass dissolved organic matter) that

descends from the water column could be utilized by the *Bacteroidetes* present in the upper part ofthe sediment.

1216 The major archaeal populations in the sediment:

1217 The distribution of the population was described above. However the environmental conditions and 1218 metabolic functionality of our sequences were not shown. The uncultured clones <u>similersimilar</u> to 1219 our sequences might help indicate on microbial process of the dominant sequences in our samples.

1220 *Euryarchaeota* 

1221 *Methanomicrobia* 

1222 The majority of our sequences were classified within *Euryarchaeota*, in *Methanomicrobia* class.

1223 Our 16S rRNA sequences were divided between *Methanomicrobiales and Methanosarcinales* 

1224 order. Methanosaeta genus of Methanosarcinales increases with depth. Methanosaeta are

acetoclastic methanogens which only able to grow on acetate even at low concentrations(Jetten et

al., 1990). *Methanosaeta* was also shown in previous studies of LK (Schwarz et al., 2007a, 2007b)

and in other meso to eutrophic freshwater lakes (Glissman et al., 2004; Koizumi et al., 2003).

1228 Yamada et al. (2014) showed that *Methanosaetacan* also have the ability to reduce ferrihydrite with

1229 H<sub>2</sub> as the electron source. The closest relative cultured acetoclastic methanogen to our sequences

1230 was *Methanosaeta concilii* (96%) similar results to the previous studysstudies. *Methanolinea* genus

1231 of *Methanomicrobiales* which is hydrogenotrophic methanogen was observed in constant

1232 percentages for all depths. *Methanoregula* genus which is also hydrogenotrophic methanogen is

decreasing with depth. *Methanoregula*, *Methanosaeta and Methanolinea* represent 60% of

sequences of methanogens retrieved from freshwater lakes (Borrel et al., 2011).

#### 1235 Thermoplasmata

The dominant family in Thermoplasmata is The Marine Benthic Group D and DHVEG-1 (MBG-1236 1237 D). Members of the MBG-D have been shown to exist in a variety of freshwater and marine environments (Beal et al., 2009; Borrel et al., 2012), and it is the most widely encountered, 1238 uncultured lineage in freshwater lake sediments. Even though their metabolism is unknown, 1239 hypotheses about their functionalities are based on the environments in which they were found. 1240 Methanogenesis was suggested, as they were found in deep lake sediments with high methane 1241 concentrations (Borrel et al., 2012), and they were also hypothesized to be involved in AOM, as 1242 they were found in AOM zones (Schubert et al., 2011) and in marine seep sediment (Beal et al., 1243 2009). However, in other environments, in which methane concentrations were low, the utilization 1244 of waste products, intermediates, or dead cells by MBG-D was also suggested (Smith et al., 1975). 1245 1246 Recently, a single cell genomics study showed that members of MBG-D were capable of exogenous protein degradation in cold anoxic environments (Lloyd et al., 2013). 1247

In addition Terrestrial Miscellaneous Gp(TMEG) family was observed only in the middle and bottom layers of the sediment. This lineage was includes clones from the terrestrial subsurface and from soils, marine sediments and freshwater lakes (Teske and Sørensen, 2008). Sequences closely related to this group were found in sediment influenced by sulfer-rich, hypoxic groundwater and aquatic sediment samples.

#### 1253 *Thaumarchaeota*

1254 *Thaumarchaeota* are mostly known as archaeal ammonia oxidizers but in the sediments they may

1255 contribute significantly to the reservoir of nitrogen oxides in ocean waters and thus to productivity,

1256 including the assimilation of carbon(Pester et al., 2011). The enzyme ammonia monooxygenase,

1257 which belongs to the enzyme family of copper containing membrane bound monooxgenases that

- 1258 possess wide substrate ranges. Closely related enzymes, which can often perform the same function
- 1259 but with different substrates, include ammonia oxidizing bacteria with ammonia (in
- 1260 *Gammaproteobacteria*) and methane as a substitute (Lontoh et al., 2000). However, a
- 1261 microorganism's metabolic energy is mainly defined by its downstream enzyme machinery
- 1262 (Tavormina et al., 2011). In our sediment *Thaumarchaeota* observed in low percentages with 2
- 1263 main classes: Marine Benthic Group B and Miscellaneous Crenarchaeotic Group.

- Miscellaneous Crenarchaeotic Group
- 1265 The Miscellaneous Crenarchaeotic Group (MCG) is a cosmopolitan class assigned to the
- 1266 *Thaumarchaeota* phylum whose members are found mostly in anoxic habitats and have the
- 1267 capability to take up organic carbon (Biddle et al., 2006). Because of their huge environmental
- 1268 range and their complex phylogeny, the MCG collectively possess great metabolic diversity(Jiang
- 1269 et al., 2008). Current evidence for members of the MCG lineage suggests that they may obtain
- 1270 energy from the anaerobic oxidation of methane, but they do so via a dissimilatory methane-
- 1271 oxidizing process and they do not assimilate its carbon (Biddle et al., 2006)
- 1272 Marine Benthic Group B
- 1273 Members of the MBGB were detected in a growing number of benthic marine environments,
- 1274 including Atlantic deep-sea sediments, marine mud volcanoes, marine carbonate crusts from the
- 1275 Black Sea, organic-poor deep subsurface sediments of the central oceanic basins, and organic-rich-
- 1276 methane or methane hydrate containing sediments near continental margins(Sørensen and Teske,
- 1277 2006). Recent results from Ocean Drilling Program showed that no known methantroph was-
- 1278 detected however methane oxidation appeared to be mediated by Marine Benthic Group B and the-
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