Modification of methane oxidation pathways
during long-term incubations of methanic lake sediments

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Abstract

Anaerobic oxidation of methane (AOM) is one of the major processes limiting the release of the greenhouse gas methane from natural environments. In Lake Kinneret sediments, iron-coupled AOM (Fe-AOM) was suggested to play a substantial role (10-15% relative to methanogenesis) in the methanic zone (>20 cm sediment depth), based on geochemical profiles and experiments on fresh sediments. Apparently, the oxidation of methane is mediated by a combination of mcr gene bearing archaea and aerobic bacterial methanotrophs. Here we aimed to investigate the survival of this complex microbial interplay under controlled conditions. We followed the AOM process during long-term (~18 months) anaerobic slurry experiments of these methanic sediments with two stages of incubations and additions of 13C-labeled methane, multiple electron acceptors and inhibitors. After these incubation stages carbon isotope measurements in the dissolved inorganic pool still showed considerable AOM (3-8% relative to methanogenesis). Specific lipid carbon isotope measurements and metagenomic analyses indicate that after the prolonged incubation aerobic methanotrophic bacteria were no longer involved in the oxidation process, whereas mcr gene bearing archaea were most likely responsible for oxidizing the methane. Humic substances and iron oxides are likely electron acceptors to support this oxidation, whereas sulfate, manganese, nitrate, and nitrite did not support the AOM in these methanic sediments. Our results suggest in the natural lake sediments methanotrophic bacteria are responsible for part of the methane oxidation by the reduction of combined micro levels of oxygen and iron oxides in a cryptic cycle, while the rest of the methane is converted by reverse methanogenesis. After long-term incubation, the latter prevails without bacterial methanotropic activity and with a different iron reduction pathway.

Keywords
Anaerobic oxidation of methane (AOM), redox, lake, sediments, dissolved inorganic carbon, stable isotope, electron acceptor

1. Introduction

Methane (CH$_4$) is an effective greenhouse gas (Wuebbles and Hayhoe, 2002) with anthropogenic and natural origins. Natural methane contributes about 50% of the global methane emissions to the atmosphere (Saunois et al., 2020). Aerobic as well as anaerobic oxidation of methane (AOM) control the release of this greenhouse gas to the atmosphere from its natural sources (Conrad, 2009; Reeburgh, 2007; Knittel and Boetius, 2009). While sulfate-dependent AOM, which is catalyzed by ANaerobic MEthanotrophs ANMEs 1-3, is widespread in marine environments (Hoehler et al., 1994; Boetius et al., 2000; Orphan et al., 2001; Treude et al., 2005, 2014), methane oxidation could theoretically be coupled to other electron acceptors. AOM coupled to reduction of iron and manganese oxides has been experimentally confirmed in several instances (Beal et al., 2009; Egger et al., 2015; Sivan et al., 2011; Sivan et al., 2014; Segarra et al., 2013; Bar-or et al., 2017; Aromokeye et al., 2020; Su et al., 2020). Humic substances, which shuttle electrons in anaerobic environments, may act as the terminal electron acceptors for AOM by ANME-2 (Scheller et al., 2016; Valenzuela et al., 2017; 2019; Bai et al., 2019).

There is also evidence that humic substances and synthetic analogs can stimulate metal-coupled AOM (Bond and Lovley, 2002; He et al., 2019; Valenzuela et al., 2019). Humic substances are considered complex organic compounds rich with redox functional moieties, such as quinones (Scott et al., 1998; Newman and Kolter, 2000; Ratasuk and Nanny, 2007), which provide these substances high redox capabilities (Valenzuela and Cervantes, 2021). The commercial quinone 9,10-anthraquinone-2,6-disulfonate (AQDS) can be used as a terminal electron acceptor (Scheller et al., 2016; Valenzuela et al., 2017; Bai et al., 2019; Zhang et al., 2019; Fan et al., 2020) or an electron shuttle (Lovley et al., 1996; Newman and Kolter, 2000) for AOM. Nitrate dependent AOM have been demonstrated in a consortium of archaea and denitrifying bacteria (Raghoebarsing et al., 2006) and an enrichment culture of ANME-2d (Haroon et al., 2013; Arshad et al., 2015), whereas nitrite fuels AOM by Methylomirabilis (NC-10, Ettwig et al., 2010). ANME-2d and Methylomirabilis can couple AOM to selenite reduction (Luo et al., 2018). It has also been shown that Methylococcales, which usually require oxygen, may use methane to support denitrification activity under hypoxia (Kits et al., 2015), and may couple methane oxidation and iron reduction (Zheng et al., 2020).

In Lake Kinneret sediments, in situ pore water profiles (Sivan et al., 2011), diagenetic modeling (Adler et al., 2011) and incubation experiments with freshly collected sediment slurries (Bar-Or et al., 2017) suggested that iron reduction coupled to AOM (Fe-AOM) removes 10-15% of the produced
methane in the deep methanic zone (>20 cm below water-sediment interface). Analysis of the microbial community structure revealed that both methanogenic archaea and methanotrophic bacteria are potentially involved in the methane oxidation (Bar-Or et al., 2015). Analyses of 16S rRNA amplicons and metagenomics suggested that archaea capable for reverse methanogenesis (probably Methanothrix or ANME-1) and the bacterial type I aerobic methanotrophs, Methylococcales, and methylothrophs, Methyloptenera, play a role in methane cycling (Bar-Or et al., 2017; Elul et al., 2021). The metagenomics analysis together with the isotope enrichment of carbon in bacterial fatty acids following anoxic incubations of the fresh sediment slurries with 13C-labelled methane (Bar-Or et al., 2017), provided evidence for the involvement of Methylococcales in methane oxidation.

This activity of aerobic methanotrophs has been observed in several anoxic lakes’ hypolimnions and sediments (Beck et al., 2013; Oswald et al., 2016; Martinez-Cruz et al., 2017; Cabrol et al., 2020), and has been speculated by potential presence of micro levels of oxygen in the deep hypolimnion or sediments, even several meters below the oxycline. Methane oxidation by pure cultures of several different aerobic methanotrophs under hypoxia was attributed to an ability to survive by switching to iron reduction (Zheng et al., 2020) or by self-generation of oxygen by methanobactins (Dershwitz et al., 2021). The latter study also showed the ability of Methylocystis sp. Strain SB2, a specific alphaproteobacterial methanotroph, to reduce iron by methane in these unique conditions.

Here, we explored the role of methanotrophic activity in natural methanic lake sediments, its survival outside of the natural conditions during long-term anaerobic incubations, and whether there is a shift in the potential electron acceptors. To answer these questions, we diluted fresh methanic sediments from Lake Kinneret with porewater from the same depth twice and amended the sediment with 13C-labeled methane to follow its oxidation to dissolved inorganic carbon (DIC). These incubations were then also amended with several types of potential electron acceptors and different inhibitors. The results of these experiments were compared to batch and semi-bioreactor experiments that were set up with freshly collected sediments to follow the changes in methane oxidation pathways along the incubation period. We also calculated methane oxidation and production rates of representative pre-incubated long-term slurry experiments. Alongside the 13C-labeled DIC measurements, we investigated the structure of the microbial population using metagenomics and lipid biomarkers to identify the potential microbial players and their dynamics over various incubation periods.

2. Methods

2.1 Study site

Lake Kinneret is a warm monomictic freshwater lake, located in the North of Israel. Its maximum depth is ~42 m and the average depth is 24 m. The lake is thermally stratified from March until December, with the hypolimnion turning anoxic starting from April. The sediment is composed
mostly of carbonates (40-50%) and clays (20%; Hadas and Pinkas, 1995). The total iron content in the top 40 cm of the sediments is ~3 wt % (Serruya, 1971; Eckert, 2000; Bar-Or et al., 2017). The composition of the sediment at the deep methanic depth used in this study (~20 cm sediment depth) was similar with 50% carbonates, 30% clay and 7% iron (Table S1). The dissolved organic carbon (DOC) concentration in the porewater increases with depth, ranging from ~6 mg C L\(^{-1}\) at the sediment-water interface to 17 mg C L\(^{-1}\) at 25 cm depth (Adler et al., 2011).

### 2.2 Experimental set-up

This study compares three incubation strategies with Lake Kinneret sediments amended with \(^{13}\)C-labeled methane, different potential electron acceptors for AOM (NO\(_2\), NO\(_3\), metal oxides and humic substances) and inhibitors for sulfur cycling and methanogens’ activity (details below) (Fig. 1):

1) Two stage slurry incubations with 1:1 sediment - pore water ratio for three months, followed by a 1:3 ratio and the addition of different manipulations for up to 18 months.

2) Semi-continuous bioreactor experiments with freshly collected methanic sediments and porewater with 1:4 ratio, where porewater was exchanged regularly.

3) Our previous results gained from batch incubation experiments with freshly collected methanic sediments and porewater with 1:3 ratio and several manipulations (Bar-Or et al., 2017; Elul et al., 2021).

#### 2.2.1 Two stage incubations

The sediments for the slurries were collected between 2017 and 2019 from the central lake (Station A) and pooled from the methanic zone (25 - 40 cm). The sediment was diluted under continuous flushing of N\(_2\) gas with porewater extracted by centrifugation from the same zone to create a 1:1 sediment - pore water ratio slurry (Fig. 1) in 250 ml glass bottles with a headspace of 70-90 ml. The slurries were flushed with N\(_2\) (99.999 %, MAXIMA, Israel) for 30 minutes, after which methane gas was injected to reach 20 % of each bottle headspace, 10 % of the injected methane was \(^{13}\)C-labeled methane (99 %, Sigma-Aldrich) using a gas-tight syringe. After three months of pre-incubation, when \(^{13}\)C-labeled DIC was observed (Fig S1), subsamples (18 g each) were transferred with a syringe under continuous flushing of N\(_2\) gas into 60 ml glass bottles and diluted with fresh anaerobic porewater to achieve a 1:3 sediment - pore water ratio (Fig. 1), which leaves 24 ml of headspace in each experiment bottle. All pre-incubated experiment bottles were crimp-sealed, flushed with N\(_2\) gas for 5 minutes, shaken vigorously and flushed again (3 times).

To verify the role of different potential electron acceptor/s and inhibitors we conducted ten experiments as outlined in Table S2. The possible influence of sulfate reduction and sulfur disproportionation on AOM was investigated by adding Na-molybdate (Lovley and Klug, 1983), to an
already running experiment in case of an active cryptic sulfur cycle, even with the absence of detectable sulfate (Holmkvist et al., 2011). Other inhibitors added were 2-bromoethanesulfonate (BES, Nollet et al., 1997) and acetylene (Oremland and Capone, 1988) (Table S2). BES is a specific inhibitor for methanogens and ANME’s mcrA genes, and acetylene is a non-specific inhibitor for methanogens (among others, as discussed later). BES was added at the beginning of the experiment, while acetylene gas was injected during the experiment to two bottles at different timepoints. Electron acceptors were added either as powder (hematite, magnetite, clay, MnO$_2$, humic substances) or in dissolved form (KNO$_3$ and NaNO$_2$). AQDS and phenazine-1-carboxylate (PCA) were dissolved in double distilled water (DDW) and then added. Amorphous iron (Fe(OH)$_3$) was prepared in the lab, by dissolving FeCl$_3$ in DDW, which was then titrated with NaOH 1.5 N, until the solution reached pH 7. The Fe(OH)$_3$ was added to the bottles by injection. The final concentration of each addition is described in table S2. The $^{13}$C-labeled methane was injected into all experiment bottles using a gas-tight syringe from a stock bottle filled with $^{13}$C-labeled methane gas (which was replaced with saturated NaCl solution). Electron acceptors and $^{13}$C-labeled methane were added to the “killed” control bottles after they were autoclaved twice and cooled. The variations in the $\delta^{13}$C$_{DIC}$ values between the experiments are the result of different amounts of $^{13}$C-labeled methane injected at the start of each experiment. 2 ml of porewater were sampled anaerobically for $\delta^{13}$C$_{DIC}$ (duplicates were taken from each experimental bottle) and dissolved Fe(II) concentrations during each sampling point from all experimental bottles. Methane was measured from the headspace (duplicates from each experimental bottle) and the porewater concentrations were calculated using the volume of the bottles and the slurries. All live treatments were set up in duplicates or triplicates, except for the black coffee treatment, which only had one replicate as an attempt to check a close analog for humic substances. In 4 experiments only one “killed” control bottle was set up because these controls had been showing repetitive results (no activity) for numerous previous experiments. For the humic substrate experiment we received natural humic substance extracted from a lake by a colleague in the University of Alaska, Fairbanks. One experiment was set up without any additional electron acceptor in order to assess the rate of methanogenesis in the pre-incubated slurries.

2.2.2 Semi-bioreactor experiment

Two semi-bioreactors (Fig. 1) were set up with fresh sediments from the methanic zone (25 - 40 cm) of Lake Kinneret central station (Station A) immediately after their collection. Both reactors were filled headspace-free with a slurry of a 1:4 sediment - pore water ratio. One of the bioreactors was amended with 10 mM hematite. To dissolve $^{13}$C-labeled methane in the porewater, 15 ml of headspace was produced with only methane gas for 24 hours. The reactors were shaken repeatedly during those hours. After 24 hours, the gas was replaced with anoxic pore-water, so that there was no headspace at all. The oxidation-reduction potential was monitored continuously by a redox electrode (Metrohm, Herisau, Switzerland) throughout the incubation period to verify anoxic conditions and to
know the redox state of the slurry in the reactor. The bioreactors were subsampled weekly to bi-
weekly, and the sample volume (5-10 ml) was replaced immediately by preconditioned anoxic
flushed with N₂ gas for 15 minutes before the exchange) porewater from the methanic zone. Samples
were analyzed for dissolved Fe(II), CH₄ and δ¹³CₐDIC as outlined below. Additional subsamples for
metagenome analysis and lipid analysis were taken at the beginning of the experiment and on day
151, and day 382 respectively. The purpose of the semi-bioreactors was to set up an experiment that
can monitor the redox state regularly, to have a closer to natural conditions, and to have another
indication for the processes involving methane in freshly collected sediments.

2.3 Porewater analyses

About 0.3 ml of filtered (0.22 um) pore-water was injected to 12 ml glass vial with He atmosphere
and 10 µl of H₃PO₄ 85% to acidify all the DIC species to CO₂ (g). The autosampler takes a gas sample
from the vials and measures the δ¹³CₐDIC of the sample on the GasBench interface of a DeltaV
Advantage Thermo Scientific isotope-ratio mass-spectrometer (IRMS) at a precision of ±0.1 ‰. It
should be noted that to measure δ¹³C₃H₄ the gas sample must be combusted before this procedure,
which means that the δ¹³C measured is of the DIC only. Results are reported versus the Vienna Pee
Dee Belemnite (VPDB) standard. Dissolved Fe(II) concentrations were measured using the ferrozine method (Stookey, 1970) by a spectrophotometer at 562 nm wavelength with a detection limit of 1 µmol L⁻¹. A gas sample was taken from the experiment bottle’s headspace by a gas-tight syringe and was analyzed for methane and ethylene concentrations by a focus gas chromatograph (GC) equipped with a flame ionization detector (FID) with a detection limit of 50 µmol L⁻¹. Methanogenesis rate was derived from temporal changes in methane concentration in a representative pre-incubated slurry experiment (Fig. S2). The amount of methane oxidized was calculated by a simple mass balance calculation according to Eq. 1 and 2:

\[
x \times F^{13}CH_4 + (1 - x) \times FDI^{13}C_i = FDI^{13}C_f
\]

\[
[CH_4]_{ox} = x \times [DIC]_f
\]

Where x is the mixing fraction of two sources which compose the final DIC; the initial DIC pool and the oxidized \(^1\)C-CH\(_4\). The letter x denotes the fraction of oxidized \(^1\)C-CH\(_4\), while 1-x denotes the fraction of the initial DIC pool out of the final DIC pool. \(F^{13}CH_4\) is the fraction of \(^1\)C out of the total CH\(_4\) at t0, \(FDI^{13}C_i\) is the fraction of \(^1\)C out of the total DIC at t0, and \(FDI^{13}C_f\) is the fraction of \(^1\)C out of the total DIC at t-final. \([CH_4]_{ox}\) is the amount (concentration in pore water) of the methane oxidized throughout the full incubation period, and \([DIC]_f\) is the DIC concentration at t-final. We assumed that the isotopic composition of the labeled CH\(_4\) did not change significantly throughout the incubation period.

2.4 Lipid analyses

A sub-set of samples was investigated for the assimilation of \(^1\)C-labeled methane into polar lipid-derived fatty acids (PLFAs) and intact ether lipid-derived hydrocarbons. A total lipid extract (TLE) was obtained using a modified Bligh and Dyer protocol (Sturt et al., 2004). PLFAs in the TLE were converted to fatty acid methyl esters (FAMEs) using saponification with KOH/MeOH and derivatization with BF\(_3\)/MeOH (Elvert et al., 2003). Intact archaeal ether lipids in the TLE were separated from the apolar archaeal lipid compounds using preparative liquid chromatography (Meador et al., 2014) followed by ether cleavage with BBr\(_3\) in dichloromethane forming hydrocarbons (Lin et al., 2010). Both FAMEs and ether-cleaved hydrocarbons were analyzed by GC-mass spectrometry (GC-MS; Thermo Finnigan Trace GC coupled to a Trace MS) for identification and GC-IRMS (Thermo Scientific Trace GC coupled via a GC Isolink interface to a Delta V Plus) for the determination of \(\delta^{13}C\) values using the column and temperature program settings described by Aepfler et al. (2019). \(\delta^{13}C\) values are reported with an analytical precision better than 1 ‰ as determined by long-term measurements of an n-alkane standard with known isotopic composition of each compound. Reported fatty acid isotope data are corrected for the introduction of the methyl group during
derivatization by mass balance calculation similar to eq. 1 using the measured $\delta^{13}$C value of each FAME and the known isotopic composition of methanol as input parameters.

2.5 Metagenome analysis

Total genomic DNA was extracted from the semi-bioreactor experiment (duplicates a and b), pre-incubation 1:1 experiments ($^{13}$CH$_4$-only control, $^{13}$CH$_4$ + hematite) and their respective initial slurries (t0), using the DNeasy PowerLyzer PowerSoil Kit (QIAGEN). Genomic DNA was eluted using 50 µl of elution buffer and stored at −20°C. Metagenomics libraries were prepared at the sequencing core facility at the University of Illinois at Chicago using Nextera XT DNA library preparation kit (Illumina, USA). 19-40 million 2 × 150 bp paired-end reads per library were sequenced using Illumina NextSeq500. For each library, taxonomic diversity was determined by mapping the reads to Silva V138.1 database of the small subunit rRNA sequences using phyloFlash (Glöckner et al., 2017; Gruber-Vodicka et al., 2019). Metagenomes were co-assembled from concatenated reads of all metagenomic libraries with Spades V3.12 (Bankevich et al., 2012; Nurk et al., 2013), following decontamination, quality filtering (QV= 10) and adapter-trimming with the BBduk tool from the BBMap suite (Bushnell B, http://sourceforge.net/projects/bbmap/). Downstream analyses, including read coverage estimates, automatic binning with maxbin (Wu et al., 2014) and metabat2 (Kang et al., 2019) bin refining with DAS tool (Sieber et al., 2018), were performed within the SqueezeMeta framework (Tamames and Puente-Sánchez, 2019).

3. Results

In this study we followed the progress of the methane oxidation process in long-term incubations from Lake Kinneret methanic sediments. This is by quantifying the modifications between experiments conducted on fresh sediments from the methanic zone (batch slurries presented by Bar-Or et al. (2017) and Elul et al. (2021) and semi-bioreactor slurries) and pre-incubated long-term batch slurry experiments.

3.1 Geochemical trends

In the pre-incubated long-term experiments, similarly to the fresh incubations, there was a conversion of $^{13}$C-methane to $^{13}$C-DIC in all the natural non-killed slurries, indicating significant AOM (Figs. 2-3). The $\delta^{13}$C$_{\text{DIC}}$ values in the natural slurries (so called methane-only control) reached hundreds of permilles, even with the low abundance of microbial populations in these sediments (Elul et al., 2021).

The geochemical experiments tested the potential of several electron acceptors to perform and stimulate this considerable AOM process. It should be noted that the actual involvement of sulfur cycling can be quantified directly by inhibiting this cycle, while the rest can be tested for their
potential involvement by their addition to the slurries. First, metal oxides were added. The addition of
hematite as an electron acceptor did not change the δ¹³C DIC increase with time (the slope) compared to
the methane-only controls (Fig. 2). This is in contrast with the freshly collected sediment experiments,
where this addition stimulated the conversion of ^¹³C-methane to ^¹³C-DIC and thus the AOM (Fig. 2).
Magnetite amendments resulted in less increase in δ¹³C DIC values as compared to the methane-only
controls (to 290% and ~360%, respectively, Fig. 3A), and amorphous iron amendments showed even
lower values (Fig. 3A). The addition of nontronite (iron bearing clay) did not cause any increase in the
δ¹³C DIC values, however, it did result in an increase in the dissolved Fe(II) concentrations compared to
the natural control (Fig. 3F, Fig. S3). The δ¹³C DIC values of the bottles with the addition of MnO₂ also
did not show any indication for AOM after 200 days, whereas the δ¹³C DIC values of the methane-only
controls reached over 500‰ (Fig. 3B).

The actual involvement of sulfate was quantified directly by the addition of Na-molybdate, an
inhibitor of sulfate reduction and sulfur disproportionation, to the methane-only controls and to
slurries amended with magnetite (Fig. 3A). This addition did not change the slope of the δ¹³C DIC
increase with time, clearly indicating no AOM inhibition and no role for sulfate in the AOM process.
Nitrate was added in two different concentrations (0.2 and 1 mM Fig. 3C) to the long-term slurries
amended with hematite, as these concentrations were shown previously to promote AOM in other
settings (Ettwig et al., 2010). Hematite addition alone increased the δ¹³C DIC values by circa 200‰
during 306 days of the experiment. The δ¹³C DIC in the bottles with the addition of 1 mM of nitrate,
with and without hematite, decreased on the other hand from 43‰ at the beginning of the experiment
to 35‰ after 306 days. The δ¹³C DIC in the bottles with the addition of 0.2 mM nitrate and hematite
increased only slightly in the end. We also observed no increase in δ¹³C DIC during the first 222 days
following the addition of 0.5 mM of nitrate (Fig. 3D), while δ¹³C DIC increased by 19‰ afterward until
the incubation was terminated. Following the addition of 0.1 mM nitrite, δ¹³C DIC increased only after
130 days and reached 158‰ at day 493. In the methane-only controls, δ¹³C DIC values reached the
highest values (330‰).

We also amended long-term pre-incubated slurries with potential organic electron acceptors. No
¹³C DIC enrichment was observed with the addition of AQDS (an analog for humic substrate) to slurries
with and without hematite (Fig. 3E). Similar trends were observed in δ¹³C DIC following the addition of
PCA, an analog for methanophenazines that are found in some archaeal membranes and shuttle
electrons (Wang and Newman, 2008) (Fig. 3F). We further tested the effect of naturally occurring
humic substances, using those isolated from a different natural lake. The results show that in the
beginning the δ¹³C DIC values did not change (Fig. 3F), while a steep increase in their Fe(II)
concentrations was observed (Fig. S3). However, after 20 days, the δ¹³C DIC values of these slurries
started to increase dramatically with a steep slope, indicating high AOM activity (Fig. 3F). We also
tested the addition of black coffee, as another example of a complex natural organic substance. In this incubation, again, the $\delta^{13}C_{\text{DIC}}$ values decreased during the first 20 days, but then increased very steeply (from 102‰ to 596‰). In those additions there is in general a mirrored trend of the dissolved Fe(II) concentrations to that of $\delta^{13}C_{\text{DIC}}$ with a steep increase, from 65 to 170 µM, during the first 20 days and then a decrease (from 170 µM to 133 µM, Fig. S3).

Geochemical analysis of $\delta^{13}C_{\text{DIC}}$ was performed also on two experiments that tested the effect of inhibitors on methane metabolism. In one experiment, BES, a specific inhibitor for methanogens and ANME’s mcrA genes, was added, and in another experiment, acetylene, a non-specific inhibitor for methanogens, was added. Both cases showed a complete inhibition of labeled $^{13}$C-DIC production following the addition, similarly to the killed control (Fig. 4). Acetylene can also inhibit nitrogen cycling in some cases, however ethylene is produced then (Oremland and Capone, 1988). In our case no ethylene was detected, supporting the inhibition only of methanogens’ activity.
Figure 3: The potential of different electron acceptors for AOM in Lake Kinneret sediments. In these pre-incubated long-term slurry experiments, the following treatments have been applied: (A) with and without the addition of magnetite and amorphous iron (Fe(OH)$_3$). Dashed line represents addition of $^{13}$C-labeled CH$_4$. Back arrow represents addition of sodium molybdate as an inhibitor for sulfate reduction. (B) with and without the addition of MnO$_2$. (C) with the addition of hematite and two different concentrations of nitrate. (D) with the addition of hematite and two different concentrations of nitrite. (E) with the addition of AQDS. (F) with clay, natural humic acid, black coffee and PCA. Green arrow represents the time clay was added to the relevant bottles, the dashed line represents the time the headspace of the bottles was flushed again with N$_2$, and the black arrow represents the second injection of 1 mL of $^{13}$C-labeled methane. $^{13}$C-labeled methane was added to all the bottles (specific details on each experiment can be found in Table S2). Each data point is the average of duplicate samples that were taken from each bottle; the error bars are smaller than the symbol.
3.2 Metagenomic and lipid analyses

The metagenomic analysis points to the potential involvement of several archaea and bacteria in the AOM observed in the pre-incubated slurries. Bona fide ANME (ANME-1), as well as various methanogens and high abundance of Bathyarchaeia were present in all the samples (Table S3). Known sulfate reducing bacteria, including Desulfobacterota, Desulfuromonadota and Thermodesulfovibrio, but not seep sulfate reducing bacteria, were found, and some in large read abundances (Table S3). Only very few metagenomic reads mapped to Methyloirabilaceae (NC-10) (<1%) and no reads mapped to Methanoperedens. The number of metagenomic reads mapped to functional genes narH and narG, which encode subunits of the respiratory nitrate reductase in Methanoperedens decreased with time in the pre-incubated sediments (Table S4). Very few reads mapped to the nirS gene, which encodes the nitrite reductase, and its coverage did not increase over time (Table S4).

The δ13C values of the archaeol-derived isoprenoid phytane showed 13C-enrichment (between 15-27‰ enrichment), and no 13C-enrichment in the killed control, indicative of methane assimilation by archaea. This signal was also found for acyclic biphytane but less pronounced (between 5-10‰ enrichment) (Table 1).
Table 1: δ¹³C values (in ‰) of fatty acids and isoprenoid hydrocarbons from different 1:1 incubations and experiments compared to values obtained from the original sediment in the methanic zone.

| Description                                      | Temperature (℃) | Sampling (days) | C₁₆:₁ω₉/₈/₇ | C₁₆:₁ω₅   | Phytane | Biphytane |
|--------------------------------------------------|-----------------|----------------|--------------|------------|---------|-----------|
| Incubated bottle +¹³CH₄+hematite                 | 20              | 411            | -40          | -43        | -17     | -23       |
| Incubated bottle +¹³CH₄                          | 20              | 1227           | -36          | -41        | -5      | -38       |
| Incubated bottle +¹³CH₄ + Typical fresh sediment| 20              | 470            | 610          | 1600       | -14     | -28       |
| Bioreactors+¹³CH₄+hematite                       | 16              | 382            | n.d.         | n.d.       | n.d.    | n.d.      |
| Original sediment (28-30 cm)                     | 14              |                | -44          | -50.7      | -32     | -33       |

4. Discussion

Our many porewater profiles of Lake Kinneret indicate that microbial sulfate reduction dominates the anoxic hypolimnion and the surface sediments, while methanogenesis is confined to the sediments below the sulfate boundary (Adler et al., 2011; Sivan et al., 2011; Bar-Or et al., 2015; Elul et al., 2021). Our previous work on fresh sediments from the lake also provided evidence for Fe-AOM in the methanic zone based mainly on geochemical and microbiological profiles and models (Adler et al., 2011; Sivan et al., 2011; Bar-Or et al., 2015). It was combined also with measurements of stable carbon isotopes in specific lipids and microbial metagenomic analyses during ¹³C-labeled methane batch incubations on fresh sediments from the methanic zone (Bar-Or et al., 2017; Elul et al., 2021; Fig. 2). These showed the unexpected significant abundance of known aerobic bacterial methanotrophs together with anaerobic microorganisms (as methanogens and iron reducers).

The first noticeable observation from the current pre-incubated long-term slurries data is that the δ¹³CₐDIC values of the natural amendments (only with the addition ¹³C-labeled methane) increased dramatically. This indicates a clear AOM signal, even after the long-term incubations and the low abundance of the microbial populations. Below, we characterize this AOM process.

4.1 Potential electron acceptors for AOM in the long-term pre-incubated experiments

The pre-incubated long-term incubations data show a sharp increase in the δ¹³CₐDIC values of both natural and hematite amendments. However, as opposed to the freshly collected sediment experiments, there was no difference between the addition of hematite as the electron acceptor and the natural (methane-only) amendment. This means that hematite does not have a potential to stimulate the AOM activity or that there is enough natural Fe(III) in the sediments to sustain the maximum potential of Fe-AOM.
Following this observation, we quantified the effect of other metal oxides, such as magnetite, amorphous iron, and manganese oxide (Fig. 3A and B), which are present in Lake Kinneret sediments (Bar-or et al., 2017), on AOM in the long-term incubation slurries from the methanic zone. The results show that the addition of any of these iron oxides showed less increase in the δ¹³C_DIC values compared to the methane only controls (Figs. 2 and 3). This indicates not only that their addition did not stimulate AOM it might even inhibit it. The less increase in the δ¹³C_DIC values with their addition could result from their direct inhibition of the AOM process or by their reduction by organic compounds other than methane (organoclastic iron reduction), which added isotopically light carbon from the organics and not heavy carbon from the ¹³C-labeled methane (masking the natural AOM signal shown in the natural control). We further tested whether ferric iron from clays, which can act as terminal electron acceptors (Kostka et al., 2002; Liu et al., 2011; Liu et al., 2012), could support AOM. However, the addition of the clay minerals appears again to encourage only organoclastic iron reduction (Fig. 3F, Fig. S3). Like iron oxides, manganese oxide, did not support AOM and likely encouraged organoclastic manganese reduction. Given that manganese oxides are found in very low abundance in Lake Kinneret sediments (0.1 %, Table S1), their potential role in metal-AOM is likely low anyway.

Sulfate concentrations in the methanic Lake Kinneret sediments are low (< 5 µM, Bar-Or et al., 2015; Elul et al., 2021). Sulfide concentrations are accordingly minor (<0.3 µM, Sivan et al., 2011). However, since pyrite and FeS precipitate in the top sediments, cryptic cycling via pyrite or FeS may replenish sulfate accessible for AOM (Bottrell et al., 2000). The role of sulfate as an electron acceptor was tested directly by the addition of Na-molybdate as an inhibitor for sulfate reduction. This addition to the long-term pre-incubated slurries amended with and without magnetite (Fig. 3A) did not change the δ¹³C_DIC values, and they increased similar to the natural (methane only) control, as occurred also in the fresh batch sediment slurries (Bar-Or et al., 2017). This indicate clearly that sulfate is not involved in the AOM process in Lake Kinneret methanic sediments, as the inhibition of the sulfur cycling did not inhibit the AOM. This is despite the presence of potential sulfate-reducing bacteria as indicated by their relatively high abundance (Table S3) in the sediments.

The concentrations of both, nitrate and nitrite, are also below detection in the porewater of Lake Kinneret sediments (Nüsslein et al., 2001), but their occurrence as an intermediate product through ammonium oxidation coupled to iron reduction (Li et al., 2015; Shuai and Jaffé, 2019) cannot be excluded. Therefore, the potential role of nitrate and nitrite as electron acceptors in the pre-incubated slurries was quantified. The results indicate that rather than stimulating AOM, the addition of nitrate (Fig. 3C) delayed AOM and promoted organoclastic denitrification. Similarly, even low nitrite concentrations appeared to inhibit AOM, potentially facilitating denitrification (Fig. 3D).
Humic substances were also investigated as potential electron acceptors for the AOM process. They could also promote AOM by continuously shuttling electrons to metal oxides (Valenzuela et al., 2019). Humic substances were not measured specifically in Lake Kinneret sediments, but DOC concentrations in the pore-water at the methanic depth were high (~1.5 mM, Adler et al., 2011), suggesting their possible role in the AOM process. The addition of the synthetic humic analogs AQDS did not cause any enrichment in $^{13}$C of the DIC. This could be due to their high electron shuttling ability and encouraging organoclastic oxidation that adds light carbon isotope (as opposed to the labeled $^{13}$C-methane) and lowers the $\delta^{13}$C$_{DIC}$ values without AOM at all, or by masking its signal (Fig. 3E). Similar trends were observed in $\delta^{13}$C$_{DOC}$ following the addition of PCA, a synthetic analog for methanophenazines (Fig. 3F). Yet, the addition of natural humic acids or black coffee exhibited different behavior. At first, the natural humic substances promoted organoclastic iron reduction, probably by shutting electrons from organic compounds other than methane to natural iron oxides in the sediments (Figs. 3F, S3). Then, perhaps when the availability of the iron oxides or the organic matter decreased, humic substances were used as terminal electron acceptors for AOM, as was suggested by Valenzuela et al. (2017). In that study, AOM was coupled to the reduction of humic substances in the presence of inorganic electron acceptors simultaneously with methanogenesis.

Overall, our experiments with different electron acceptors indicate clearly that sulfate is not involved in the AOM process in Lake Kinneret methanic sediments, and that nitrate, nitrite and Mn-oxides are less likely. The potential electron acceptors are natural humic substrates with or without iron minerals that are abundant in the sediment and preferably react with methane rather than with other organics.

### 4.2 Main microbial players in the long-term pre-incubated experiments

As mentioned above, the pre-incubated long-term incubations data show a sharp increase in the $\delta^{13}$C$_{DOC}$ values of natural amendments. However, the addition of BES, a specific inhibitor for methanogens and ANME’s $mcrA$ genes, stopped immediately the AOM, similarly to the killed bottles, and to the fresh sediment experiments (Bar-Or et al., 2017), indicating methane oxidation by methanogens or ANMEs in all stages of incubations (Fig. 3). In addition, the complete inhibition of labeled DIC production following the addition of acetylene (Fig. 4) suggests the involvement of methane metabolizing microorganisms, also evidenced by the enrichment in $\delta^{13}$C values of phytane and biphytane (Table 1). Such a signal is generally indicative of active archaea, i.e. methanogens or ANMEs in this case, which assimilate $^{13}$C-carbon from an unknown intermediate or existing DIC.

The essential role of methanogens or ANMEs in the AOM in all stages of incubations suggest that this process is performed by reverse methanogenesis. Indeed, in metagenome-assembled genomes (MAGs) of ANME-1 and Methanothrix, all the seven genes ($mer$, $mtr$, $mer$, $mtt$, $mch$, $fir$, $fmd$) needed for the reverse methanogenesis (Meyerdierks et al., 2010; Wang et al., 2014; Wegener et al., 2021) were found. It should be noted that ANME-1 was found in very low abundance (< 1.5%) and other
ANMEs were not found at all. In addition, while the abundant Bathyarchaeia in all incubation stages might be involved in methane metabolism (Evens et al., 2015), the mcrA genes were not found in their Lake Kinneret MAGs, thus their role in AOM is questionable.

On the other hand, both the metagenomic and lipid isotopic analysis suggest that the role of aerobic type I methanotrophs (of the class gammaproteobacteria) in methane turnover in the long-term incubations is negligible (Table S3). This contrasts with the natural sediments and fresh incubations that show their presence in the sediments and their important role in oxidizing the methane.

4.3 Methane oxidation pathway in the long-term incubations

Our results indicate net methanogenesis in long-term incubations with an average rate of 2 µM day⁻¹ (Fig. S2 and Table S5), similarly to the fresh incubation experiments (Bar-Or et al., 2017). This is even with the overall development of increasing Δ¹³CDOC values resulting from potential methane turnover (Fig. 2 and 3). A likely explanation for this signal is an interplay between methane production and oxidation, with the latter triggered by reverse methanogenesis, which is demonstrated among the orders of methanogens and ANMEs (Hallam et al., 2004; Timmers et al., 2017). Of these, Methanotrichs (closely related to the order Methanosarcinales) has high potential to perform reverse methanogenesis here and in other environmental settings (Valenzuela et al., 2017; 2019; Elul et al., 2021). In our sediments, Methanosarcinales were also found to increase in abundance towards the methanic zone (Bar-Or et al., 2015). Reverse methanogenesis is used in trace methane oxidation by pure cultures of various species of the Methanosarcina and the Methanobacterium genera (Zehnder and Brock, 1979; Moran et al., 2005, 2007; Luo et al., 2017; Lai et al. 2018).

Due to the overall production of methane and the lack of intensive stimulation of AOM by any electron acceptor, the high increase in Δ¹³CDOC could also theoretically result from carbon back flux during methanogenesis, which is feasible in environments close to thermodynamic equilibrium (Gropp et al., 2021). To determine whether back flux is feasible in the incubations, we assessed how much of methane is oxidized and converted to DIC using mass balance calculations. To reach the observed ¹³C-enrichment in our experiments, 3-8 % of the ¹³C-methane had to be channeled into DIC through Eq. 1 and 2, which is much higher than the previously reported methanogenesis back flux values (0.3-0.001 %, Zehnder and Brock, 1979; Moran et al., 2005). Back flux reactions have been studied before only in ANME-enrichment cultures and by modeling approaches in marine environments without indications of net methanogenesis (Holler et al., 2011; Yoshinaga et al., 2014; Chuang et al., 2019; Meister et al., 2019; Wegener et al., 2021). During net AOM conditions, however, this process was recently attributed to intracellular reversibility of enzymes involved in the reaction chain under substrate limitation without invoking methane-DIC equilibration (Wegener et al., 2021). Indeed, low methanogenesis rates in the environment may result in enhanced back flux, compared to the active methanogenic cultures (Hoehler et al., 1994; Holler et al., 2011). Yet, based on
the above, it is unlikely that back flux alone will account for the methane-DIC conversion in the Lake Kinneret sediments. Also, we observed no or very little $^{13}$C-enrichment in the DIC pool following similar incubations with marine sediments, which showed net methanogenesis and contained similar abundance of methane-metabolizing archaea to that of Lake Kinneret sediments based on the detection of \textit{mcrA} with qPCR (Sela-Adler et al., 2015; Amiel, 2018; Vigderovich et al., 2019; Yorshensky, 2019) (Table S6). Therefore, under natural conditions, methanogens alone are unlikely to produce our observed considerable amounts of DIC just by back flux.

4.4 The progression of methane oxidation over time

The geochemical and microbial profiles and fresh sediment incubations show evidence for Fe-AOM in the methanic zone of Lake Kinneret, which removes about 10-15% of the produced methane (Adler et al., 2011; Sivan et al., 2011). Anaerobic archaea appear to carry out methane turnover in these reduced sediments by reverse methanogenesis, but methanotrophic Methylococcales are also involved in methane oxidation. This fits other studies, which show more evidence pointing to the existence of aerobic bacterial activity in the deep anoxic hypolimnion of lakes and in the shallow sediments (Beck et al., 2013; Oswald et al., 2016; Martinez-Cruz et al., 2017; Cabrol et al., 2020). These bacteria live alongside strict methanogenic anaerobes and iron reducers, probably in a complex interaction, which increases the iron reduction in a cryptic cycle that should be further explored.

The presence of aerobes and anaerobes together in nature, even 20 meters below the thermocline and oxycline, means that small amounts of oxygen could be trapped in nano-niches or even in mineral layers (Wang et al., 2018), even if they are not detected by sensitive sensors. This oxygen portion may not be removed by purging the freshly collected sediments at the beginning of our experiments but is rather slowly used by the methanotrophs for their survival. However, after several incubation stages, and intensive purging and prolonged time, only archaea remained active and were involved in methane turnover. It appears that methanotrophic bacteria cannot survive the long-term slurry incubations and thus iron reduction and methane oxidation are decoupled.

To conclude, trace levels of oxygen may fuel aerobic methane oxidizers in a cryptic cycle between oxygen and iron in the natural lake methanic sediments, and they are responsible for part of the methane oxidation and maybe the iron reduction. The rest of the methane is oxidized to DIC by methanogens or ANME-1. The DIC production from methane turnover in the long-term experiments is performed only by methanogens or ANME-1. It seems less likely that this is by back flux alone, but rather by active metabolic AOM by reverse methanogenesis and an external electron acceptor. Sulfate, nitrate, nitrite, and manganese are unlikely. Humic substances are the most likely electron acceptors used with or without the natural iron oxides.

Competing interests. The authors declare that they have no conflict of interest.
Acknowledgements

We would like to thank B. Sulimani and O. Tzabari from the Yigal Allon Kinneret Limnological Laboratory for their onboard technical assistance. We thank all of O. Sivan’s lab members for their help during sampling, and especially to N. Lotem for the help with the mass balance calculations and discussions and to E. Eliani-Russak for her technical assistance. Many thanks to K. Hachmann from M. Elvert’s lab for his help during lipid analysis and to J. Gropp for insightful discussions on the back flux. This work was supported by the ERC consolidator grant (818450) and the Israel Science Foundation (857-2016) of O. Sivan. Funding for M. Elvert was provided by the Deutsche Forschungsgemeinschaft (DFG) (49926684) and EXC 2077 (390741601). Funding for M. Rubin-Blum was provided by the Israel Science Foundation (913/19), the U.S.-Israel Binational Science Foundation (2019055) and Ministry of Science and Technology (1126), and H. Vigderovich was supported by the student fellowship of the Israeli water authority.

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