Influence of minor hydrocarbon seepage on sulfur cycling in marine subsurface sediments

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20 Abstract.

All hydrocarbon (HC) reservoirs leak to some extent. When small quantities of HCs escape offshore reservoirs and migrate through overlying organic-poor marine sediments towards the surface, these HCs are often completely metabolized by microorganisms before reaching the sediment-water interface. However, these low and often unnoticed upwards HC fluxes still influence the geochemistry of surrounding sediments

25 and potentially stimulate metabolic activity by microbial populations in shallow subseafloor environments.

In this study, we examined how localized HC seepage affects microbial sulfate reduction in organicpoor sediments from the SW Barents Sea, focusing on three sampling areas overlying known HC deposits and two pristine seabed reference areas. The analysis of 50 gravity cores revealed significant variability in the predicted sulfate depletion depth, ranging from 3 to 12 m below the seafloor. We observed nearly linear

30 pore water sulfate and alkalinity profiles, alongside low rates of sulfate reduction (pmol \times cm³ \times d⁻¹).

Metagenomic and metatranscriptomic data indicated metabolic potential and activity towards sulfate reduction and anaerobic oxidation of methane (AOM). Expression of functional marker genes (*aprAB*, *dsrAB*, *mcrA*) revealed metabolic activities by sulfate-reducing Desulfobacterota and methanotrophic ANME-1 archaea sustained by HC traces in the sediment. Further, in concomitance to AOM processes, we found that

35 archaea amongst the class Lokiarchaeia and Thoarchaeia were expressing genes involved in sulfur reduction (*hydB*, *hydG*).

Overall, our study demonstrates that the gradient in pore water geochemistry, the rates of sulfate reduction processes, and the genetic features of microbial populations actively involved in sulfate-driven AOM processes are all affected by low and often unnoticed upwards HC fluxes. This slight HC seepage

40 resulted in sulfate depletion at shallower depths compared to what would be expected at a comparable, nonaffected site, and produced concomitant biogeochemical signatures in the shallow subsurface that enable the inference of deeply buried reservoirs.

1. Introduction

All hydrocarbon (HC) reservoirs leak to some degree (Hunt, 1995; Yergin, 2009; Heggland, 1998), with over

- 45 80 % of all seeps occurring directly above the reservoirs (Ciotoli et al. (2020). While large leakages result in conspicuous manifestations at the seafloor, (e.g. Cramm et al. (2021), minor seeps, characterized by low, primarily diffusive HC fluxes usually remain invisible at the sediment-water interface (SWI), as there are no visible signs of seepage, e.g. carbonate crussible bioherms. In cases where HC fluxes are so low that hydrocarbons are fully mineralized and do not reach the SWI, sediment geochemistry and microbiology at
- 50 minor seepage sites will be altered (Rasheed et al., 2013; Abrams, 2020; Riedinger et al., 2014; Joye et al., 2004). Even if HC fluxes were higher and hydrocarbons do indeed reach the SWI, they are often overlooked (Abrams, 2020; Joye, 2020).

HC seepage triggers both geochemical and microbiological processes, changing physicochemical properties, such as sediment porosity, mineral and pore water composition, as well as microbial community

- 55 composition and activity (Abrams, 2020; Hvoslef et al., 1996; Joye, 2020). Likewise, the presence of free gas in methane-containing pore fluids alters sediment porosity and compressibility (Jang et al., 2018), which can be detected through seismic profiling due to the resulting impedance contrast (Rovere et al., 2020). These changes may also affect the sediment's geochemical properties, such as clay texture, organic matter (OM) reactivity, and metal content (Chen et al., 2023). HC influx and subsequent HC consumption will also alter
- 60 pore water geochemistry and pH, promoting ion exchange reactions, dissolution and precipitation (Smrzka et al., 2019; Turchyn et al., 2021; Torres et al., 1996) and structural conversion of minerals (Jiang, 2012). As

a consequence, specific trace metals and metalloids (e.g. Cr, Cu, As, Se, Sb) that can be metabolized by specific microorganisms (Raab and Feldmann, 2003) tend to become enriched in the sediment pore water at seepage sites (Rasheed et al., 2013; Guseva et al., 2021). Finally, most offshore sediments are composed of

- 65 pelagic deposits, i.e. clays, silts or biogenic oozes, containing only low amounts of bioavailable OM that becomes increasingly recalcitrant with depth of burial (Middelburg, 2018). Thus, the influx of HCs in such sedimentary systems represents an additional source of electron donors, providing chemical energy for microorganisms and potentially fueling various metabolic processes. The metabolic potential to break down and assimilate HCs has been identified in a large number of microorganisms, e.g. reviewed in Joye (2020),
- 70 whose activities may induce concomitant effects in the pore water, organic and mineral fraction of the sediment (Kim et al., 2004).

The quantitatively most important anaerobic OM degradation process in the upper layers of marine sediments, where oxygen is depleted, but sulfate is still present in relatively high concentrations is sulfate reduction (Jørgensen et al., 2024; Bowles et al., 2014; Kasten and Jørgensen, 2000; Jørgensen, 1982). Due to

75 its abundance in seawater, sulfate diffuses into the sediment pore water where it is used as an electron acceptor by sulfate-reducing microorganisms composed of sulfate reducing bacteria and ANME (Widdel et al., 2010). Sulfate reduction (SR) can be divided into two main pathways, namely organoclastic and methanotrophic SR. The respective overall reactions are:

80 Organoclastic SR: $2 \text{ CH}_2\text{O} + \text{SO}_4^{2-} \rightarrow 2 \text{ HCO}_3^- + \text{H}_2\text{S}$ (Equation 1) Methanotrophic SR: $\text{CH}_4 + \text{SO}_4^{2-} \rightarrow \text{HS}^- + \text{HCO}_3^- + \text{H}_2\text{O}$ (Equation 2)

Due to the abundance of easily biodegradable OM and sulfate close to the sediment-water interface, organoclastic SR normally prevails in anaerobic OM degradation within the uppermost part of the sediment column, leading to a decrease in sulfate concentration. Methanotrophic SR becomes predominant only when biogenic or thermogenic methane diffusing from deeper sediments reaches the zone where sulfate is still available but approaches depletion (Martens and Berner, 1974). The narrow zone where both methane and sulfate are available is referred to as the sulfate-methane transition zone (SMTZ). Within the SMTZ, mainly methane, and potentially also other OM compounds (Beulig et al., 2019; Jørgensen et al.,

90 2019b), are oxidized by microorganisms while sulfate is reduced (Equation 2).

Thus, the respective methane and sulfate fluxes exert control on the depth of the SMTZ (Borowski et al., 1996, 1999; Jørgensen and Kasten, 2006), which, under steady-state conditions, is usually a reflection

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of methanogenesis rates (Henrichs and Reeburgh, 1987) or fluxes of thermogenic methane (Hu et al., 2017). However, the SMTZ is highly dynamic and responds relatively rapidly (i.e. within decades) to shifts in pore

- 95 water geochemistry (Sultan et al., 2016; Hong et al., 2016). Thus, microbially active subseafloor sediments exhibit significant variability in the depth of the SMTZ across different oceans, from 1 to 10 of meters on the continental shelf (Egger et al., 2018; Treude et al., 2020; Treude et al., 2014; Riedinger et al., 2017; Riedinger et al., 2018; Niewöhner et al., 1998). On the continental slope, the SMTZ can be down to 100 m below seafloor (mbsf) in extreme cases (Nunoura et al., 2009), but is usually not deeper than 25 mbsf.
- 100 The subseafloor biosphere hosts phylogenetically diverse and metabolically active microbial communities (Parkes et al., 2014; D'hondt et al., 2004). Their metabolic activity depends mainly on biogeochemical cycling of nitrogen, phosphate and sulfur. At HC seepage sites, microbial communities are well-defined in terms of taxonomy and highly specialized in terms of metabolic functions, and are referred to as "seep biomes" (Ruff et al., 2015). It has been known for many decades that methanogenic hydrocarbon
- 105 degradation is carried out jointly by bacteria and archaea in underground hydrocarbon reservoirs. New studies suggest that *Candidatus methanoliparum* may be able to independently degrade long-chain alkanes and produce methane (Wegener et al., 2022; Zhou et al., 2021; Zhang et al., 2024). Seep biomes show spatial heterogeneity, but are distinct from non-seep environments (Pop Ristova et al., 2015). The reported key microbial guilds include methane and oil oxidizers, sulfate reducers, and nitrogen fixers. Taxa found near
- 110 HC seeps involve members of the bacterial phyla *Desulfobacterota* and *Chloroflexota* that can respectively use simple and halogenated HCs as electron acceptors (Kleindienst et al., 2014; Zhang et al., 2023), whereas archaeal groups, like the ANME or Asgardarchaeota, consume methane or make use of HCs or sulfur compounds (Firrincieli et al., 2021; Macleod et al., 2019).
- Our study aims to determine in what ways minor HC seepage modifies the diversity and activity of anaerobic methanotrophs and sulfate-reducing bacteria in surface sediments of the SW Barents Sea. We hypothesize that minor HC seepage influences pore water geochemical properties, even in near-surface sediment, and that resident microbial communities respond quickly in terms of composition and activity to the geochemical alteration of their habitats.

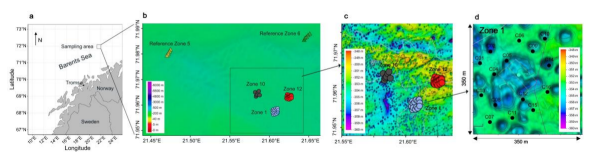
For this, we collected a total of 50 gravity cores, of which 40 cores from areas affected by light HC seepage from underlying HC reservoirs (i.e., HC-affected sites) but without any visible surface manifestations like carbonate crusts otherms, and 10 cores from seabed areas without seepage (i.e., reference sites). We combine detailed analyses of pore water sulfate and alkalinity with potential and modeled sulfate reduction rates, dissolved methane (CH₄) and carbon dioxide (CO₂) concentrations, and supplement them with metagenomic (i.e. total DNA) and metatranscriptomic (i.e. total RNA) data on taxonomic and functional

125 marker genes in order to trace metabolic activity by microbial sulfate-reducing consortia in their geochemical context.

2. Material and methods

2.1 Geological context of the Barents Sea, sampling sites

- The Barents Sea is an epicontinental shelf sea surrounded by the marginal Norwegian Sea to the southwest, 130 the Arctic Ocean to the north and the Russian island Novaya Zemlya to the east (Fig. 1a). Since its formation during the Caledonian orogeny (i.e. 490-390 Ma), the Barents Sea Basin experienced several phases of uplift and subsidence, with subsequent tilting and erosion. These geologic events resulted in the characteristic structure of the Barents Sea in the form of a succession of structural highs and basins (Gabrielsen et al., 1990; Larssen et al., 2002).
- HC exploration in the Barents Sea has a history stretching back several decades (Doré, 1995; Johansen et al., 1993). The sediments have been characterized in numerous studies, e.g. Sættem et al. (1991); Elverhøi and Solheim (1983), and are mainly composed of organic-poor silty clays (Knies and Martinez, 2009; Nickel et al., 2013) originating from eroded sedimentary and igneous rocks.
- During an expedition to the southern Loppa High region of the Barents Sea (Fig. 1a) from October 29th to November 8th 2021, we collected 50 gravity cores (up to 3 m in length) sediment at about 350 m water depth at three areas with known underlying HC reservoirs (i.e. Zone 1, Zone 10, Zone 12) and at two reference areas (i.e. Ref 5, Ref 6) without underlying HC reservoirs (Fig. 1b), labelled thereafter HC-affected sites and reference sites, respectively. The cores were subsampled for sediment and pore water on board the research vessel.



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Figure 1. Sampling locations. (a) Map of the Norwegian coast. The square marks the sampling area. **(b)** Fifty cores were retrieved from three HC-affected sites (Zone 1, Zone 10, Zone 12) and two reference sites (Ref 5, Ref 6) (bathymetric map: https://emodnet.ec.europa.eu/geoviewer/). **(c)** bathymetric map of the 3 HC-

affected sites. (d) Each site was sampled in a grid of cores, Zone 1 is enlarged here as an example. The

150 missing samples in the middle are due to the increased occurrence of dropstones in this area, preventing the deployment of a gravity corer.

2.2 Pore water and sediment sampling

- Pore water samples were extracted from all cores at a resolution of 10 cm, using rhizons (Rhizosphere Research Products) (Seeberg-Elverfeldt et al., 2005), collecting a minimum volume of ca. 5 mL over 24 h. The pore water was filtered through 0.2 μm pore size cellulose esters (MCE) membrane syringe filters (Merck MF-Millipore) and aliquoted as follows: 1.5 mL transferred into a plastic screw cap vial without further treatment for anion and cation measurements; 2 mL into a glass vial amended with 50 μL saturated HgCl₂ solution to inhibit microbial activity and sealed without headspace for alkalinity measurements (Edenborn et
- 160 al., 1985); and 1.5 mL into a plastic screw cap vial mixed with 200 μ L ZnCl₂ (20 % weight × vol⁻¹) to precipitate dissolved hydrogen sulfide as ZnS for subsequent sulfide measurements. All samples were stored at +4 °C until analysis in the home lab.

Immediately after core recovery, the lowermost 20 cm of each sediment core were cut off and subsampled for DNA and RNA extraction. The sediment was pushed out of the liner and the outer 1-2 cm of sediment rims were removed with a sterile spatula. Aliquots of the remaining sediment were transferred into

165 sediment rims were removed with a sterile spatula. Aliquots of the remaining sediment were transferred into gas-tight aluminum foil bags, flushed with nitrogen gas, heat-sealed and stored at -80 °C for later extraction of various biomolecules.

2.3 Pore water sulfate, sulfide and alkalinity

- 170 Anion concentrations were determined using an Ion Chromatography (IC) System equipped with a SykroGel A \times 300 AB-A01 column (all Sykam). The eluent contained 7.3 mg \times L⁻¹ NaSCN and 636 mg \times L⁻¹ NaCO₃. The pump rate was set to 1 mL \times min⁻¹ and the injected sample volume was 50 µL. All samples were measured in triplicates and the results were averaged. Because of their high salt concentrations, all samples had to be diluted 1:50 with MilliQ water before injection. The detection limit of the IC system is 0.5 mg \times L⁻¹. The
- 175 average standard deviation of replicate measurements was always better than 3 %.

Sulfide concentrations were measured according to the protocol of (Cline, 1969). Aliquots with absorption values greater than 1 were diluted and remeasured. Sulfide concentrations were calculated by comparing adsorption values measured for pore water samples to those of a Na₂S standard. The detection

limit of the method is 0.1 µM. All samples were measured in triplicates and the results were averaged, with

180 standard deviation ≤ 3.5 %.

Pore water alkalinity was determined using the Visocolor HE alkalinity AL 7 kit (Macherey-Nagel GmbH). As we only had a total of ca. 2 mL of pore water per sample, we scaled down the volume of sample to 500 μ L in order to perform all measurements in triplicates. For this, 10 μ L of indicator was added to 500 μ L of pore water and titrated with the Visocolor solution until we observed a change in color from blue

to orange. Alkalinity of the pore water samples was calculated based on the titration solution volume.

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Replicates differed by less than 3 % on average.

2.4 Sulfate reduction rates, modeled sulfate reduction rates

- For sulfate reduction measurements, each gravity core was subsampled in triplicate every 40-50 cm and at 190 the bottom end. Structurally intact sediment plugs (volume ca. 3.5 cm³) were recovered by inserting 5 mL glass barrels fitted with a syringe plunger into the sediment. Following the whole core injection method (Jørgensen, 1978), 15 μL of radioactive ³⁵SO₄-sulfate tracer (200 kBq) were injected into the glass barrels containing the samples retrieved on the same day (Fossing et al., 2000). The glass barrels were closed with butyl rubber stoppers, and the samples incubated at *in situ* temperature (ca. 4 °C) for 24 hours. Each sample
- 195 was then transferred into a previously weighed 50 mL centrifuge tube containing 10 mL of 20 % zinc acetate solution to stop all microbial activity and to precipitate all volatile hydrogen sulfide as zinc sulfide. Each vial was then thoroughly shaken to break down all sediment aggregates and stored frozen until analysis. Blank samples were prepared by adding the radiolabeled sulfate tracer just before transferring the sample into the zinc acetate solution. The tracer blank was prepared in the same way without sediment.
- 200 Sulfate reduction rates (SRR) were quantified in the home lab, using single-step cold chromium distillation (Kallmeyer et al., 2004). The sample vials were thawed, centrifuged and the supernatant carefully removed. A small aliquot of supernatant was kept for quantification of total radioactivity, the rest was discarded. The sediment pellet was then quantitatively transferred into a distillation flask. Due to the clayey consistence of the sediments, we had to re-freeze the vials and transfer the frozen sediment pellet from the
- 205 centrifuge tube to the distillation flask. SRR were calculated according to the following formula:

Sulfate reduction rate: $SRR = \frac{SO_4 * \varphi * a_{tris} * 1.06 * 10^6}{a_{tot} * t}$ (Equation 3)

SRR: sulfate reduction rate [pmol cm⁻³ × day⁻¹]; SO₄: sulfate concentration in the pore water [mmol × L⁻¹]; φ : porosity cm⁻³] of the sediment, set to 0.7; a_{tris}: radioactivity of total reduced inorganic sulfur [cpm];

 a_{tot} : total radioactivity used in the spiking [cpm]; t: incubation time [d]; 1.06: correction factor for isotopic 210 fractionation; 10⁶: unit conversion factor from [µmol × L⁻¹] to [pmol x L⁻¹].

Assuming steady state conditions, the net rate of sulfate production, or consumption, can be modeled based on the measured sulfate concentration profiles by the diffusion-reaction modeling software PROFILE (Berg et al., 1998). This software considers three different kinds of vertical transport (i.e. diffusion, bioturbation, irrigation) together with the flux across the SWI. The final output corresponds to the minimum number of intervals, as defined by their respective sulfate production or consumption rates, that are necessary

- to reproduce the measured pore water profiles. For the modeling calculation, we used: coefficient of sulfate diffusion in water at 4 °C $D_{Sulfate}$ = 0.56 × 10⁻⁵ [cm² × s⁻¹] (Iversen and Jørgensen (1993); porosity φ = 0.7; and sulfate seawater concentration 28 mmol × cm³. Because the sediment appeared to be free of macrobenthos, the biodiffusivity and irrigation coefficients were set to D_B = 0 cm² × s⁻¹, and α = 0,
- 220 respectively.

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2.5 Methane and carbon dioxide concentrations

For dissolved CH₄ and CO₂ concentrations, we subsampled the lowermost 20 cm of sediment from each gravity core directly after recovery. Using 5 mL cut-off syringes, we extracted ca. 4 cm³ of sediment per core.
The first and last 0.5 cm³ of sediment in the syringe were discarded while the central 3 cm³ were transferred into a 10 mL glass crimp vial containing a saturated NaCl solution. The vials were immediately sealed with thick butyl rubber stoppers, crimped, and stored upside down without headspace until analysis in the home

lab.

- Prior to measurement, we introduced 3 mL of ultrapure helium gas as headspace while withdrawing the same amount of NaCl solution from the vial. To equilibrate dissolved gases with the headspace, the content of the vials was mixed at 220 rpm on an orbital shaker for 18 h, and further vortexed to break up the remaining small clayey aggregates. We extracted ca. 350 µL of headspace from the vial using a gas-tight syringe. Prior to measurement, ca. 100 µL of the sampled gas were flushed through the injection needle, and 250 µL of gas sample were then injected into a 7890A Gas Chromatography System equipped with a flame
- ionization detector (FID), a thermal conductivity detector (TCD) and HP PLOT Q column (all Agilent). Oven temperature was set to 50°C, flow rate to 17.2 mL × min⁻¹ and pressure to 13 psi. The detectors both worked at 200°C with flow rates of 40 mL × min⁻¹ (FID) and 15 mL × min⁻¹ (TCD). The system was calibrated, using 250 μ L of analytic pure standards, injecting CO₂ concentrations of 310 ppm and 5270 ppm, and CH₄

concentrations of 10 ppm and 5170 ppm. The initial CH_4 and CO_2 concentrations were converted from ppm 240 to molar concentrations by applying the ideal gas law.

2.6 DNA/RNA extractions, sequencing libraries, functional marker genes

DNA was extracted for a total of 12 samples from 8 different cores corresponding to 4, 2 and 2 samples from HC-affected sites, i.e. Zone 1, Zone 10 and Zone 12, respectively, along with 3 and 1 samples from Ref 5 and Ref 6, respectively. Samples were subjected to DNA extraction using the DNEasy PowerMax Soil Kit (Qiagen, Germany) following manufacturer's instructions with few adaptations. For 2.5 g of starting sediment material, the final elution was performed in two steps using twice 2.5 mL of solution C6, and successively incubated and centrifuged. Eluted DNA was then concentrated by adding 510 μL of a 3M sodium acetate (pH 5.2) / glycogen (0.4 μg × μL⁻¹) solution and incubated for one hour at -20°C and then centrifuged to pellet DNA. After ethanol cleaning and air-drying, DNA was resuspended in 100 μL of 10 mM Tris buffer. Finally, DNA was cleaned and concentrated using the Zymo DNA Clean & Concentrator (Zymo, UK) following the manufacturer's instructions and eluted in a final elution volume of 35 μL.

DNA sequencing was achieved by using 32.5 µL of purified and concentrated DNA for metagenomic library preparation with the QIAseq FX DNA library UDI A/B kit (Qiagen, Germany)
irrespective of the individual sample concentrations. The genomic DNA was enzymatically fragmented for 10 min and DNA libraries were prepared with 7 PCR cycles for the PCR-less procedure, and without dedicated PCR step for the PCR-free procedure. The average library sizes were ±400 base pairs. Prepared libraries were quantified using a Qubit fluorometer (ThermoFischer, USA) and quality-checked on a Bioanalyzer (Agilent, USA). Sequencing was performed on an Illumina NextSeq2000 platform using 2 × 151
bp read length at the Luxembourg Centre for Systems Biomedicine (LCSB) Sequencing Platform, aiming at

an average of 10 Giga base pairs per sequencing library.

RNA was extracted from 2×5 g of sediment per sample, using the RNeasy PowerSoil Total RNA kit (QIAGEN) according to the manufacturer's instructions, yielding 30 µL of RNA-containing extract, and quantified on a Qubit4 fluorometer, using the RNA high sensitivity kit (ThermoFisher Scientific). RNA

265 extracts were directly subjected to library preparation in technical duplicates, using the Revelo[™] RNA-Seq High Sensitivity kit (Tecan Life Sciences) with 8 µL of RNA extracts as template. This trio kit includes first and second strand cDNA synthesis, adaptor ligation and library amplification through 10 PCR cycles. Prepared libraries were quantified using a Qubit4 fluorometer (ThermoFisher Scientific) and quality-checked on a Bioanalyzer (Agilent) to determine the molarity and overall size distribution of RNA molecule

- 270 fragments. The average library fragment size was \pm 300 bps and the yields were between 4 and 12 ng $\times \mu L^{-1}$. ¹. Metatranscriptomic libraries displaying high and low molarities were pooled into two different batches, and both sequenced at the EMBL GeneCore (https://www.comprg/) on a NextSeq 2000 with a P3 kit and an Illumina micro kit (2 \times 150 bps), respectively.
- *De novo* assembly of metagenomic reads was performed (Table S1) using a customized Snakemake 275 workflow (Mölder et al., 2021), including quality-control, binning into contigs, gene prediction and annotation as described in (Bornemann et al., 2023). Paired-end reads were interleaved, Illumina adapters and controls removed, reads quality-based trimmed, and de-interleaved into forward and reverse read files. Bacterial and archaeal scaffolds were concatenated and mapped back to raw reads to retrieve coverage, % GC and contig lengths. Genes were then predicted by extracting open reading frames (ORFs) from
- 280 metagenome-assembled scaffolds using Prodigal (Hyatt et al., 2010). Metatranscriptomic reads were qualityfiltered and trimmed using Trimmomatic (Bolger et al., 2014). Functional profiles were obtained by mapping the filtered and trimmed reads to their corresponding metagenomic databases using BWA-mem (Li and Durbin, 2009). ORFs were retrieved using Bakta (Schwengers et al., 2021) and combined with the mapping using the featureCounts tool (Liao et al., 2014).
- 285 Taxonomic identifications of all ORFs were further integrated with functional annotations by performing BLASTp searches against an aggregated database of predicted proteins (https://github.com/williar MetaProt-database), using DIAMOND protein aligner v. 0.9.24 (Buchfink et al., 2015). Cut-off values for assigning the best match to specific taxa present in the aggregated database were performed at a minimum bit score of 50, minimum amino acid similarity of 60, and an alignment length of 50 residues. We used this approach to draw conclusions about metabolic traits derived specifically from – 290

phylum to class taxonomic levels only (Vuillemin et al., 2020a; Vuillemin et al., 2020b).

Here, we focus on ORFs encoding genes involved in sulfur cycling and methane-related processes only. To trace metabolic potential in anaerobic respiration of sulfur compounds (i.e. sulfate, sulfite, thiosulfate, sulfur, polysulfide), we looked for the presence of ORF- encoding genes with similarity to

- 295 adenylylsulfate reductase (*apr*), dissimilatory sulfite reductase (*dsr*), anaerobic sulfite reductase (*asr*), thiosulfate/polysulfide reductase (*phs/psr*), and sulfhydrogenase (*hyd*) (Vuillemin et al., 2022). For methane production and anaerobic consumption, we targeted ORFs encoding proteins of the methyl coenzyme M reductase (*mcr*). To identify and characterize metabolically active taxa, we targeted the small subunit ribosomal ribonucleic acid (SSU rRNA) as a taxonomic marker and the filamenting temperature-sensitive
- 300 mutant A-Z (*FtsAZ*) for cell division.

To confirm taxonomic assignments and metabolic activity of taxa involved in sulfate reduction and AOM processes, we performed a phylogenetic analysis of the RNA polymerase sigma factor (*rpoD*) and ribosomal protein S3 (*rps3*) gene proteins, *aprAB*, *dsrAB*, *mcrA* and *hydB* that we could detect in the transcriptomes (Vuillemin et al., 2020a). All ORFs annotated to these genes were aligned against their top

two BLASTp hits in the NCBI database, using MUSCLE (Edgar, 2004). Conserved regions of the alignments were selected using Gblocks 0.91b (Castresana, 2000) with the following settings: allowing for smaller final blocks, gap positions within the final blocks and less strict flanking positions (http://phylogenetic analysis of the resulting amino acid alignments of the predicted proteins were conducted in SeaView version 5.0.5 (Gouy et al., 2010), using PhyLM maximum likelihood (Guindon et al., 2010), with
 BLOSUM62 as the evolutionary model and 100 bootstrap replicates

3. Results

3.1 Pore water geochemical profiles

implies that sulfate would be depleted at ca. 12 mbsf.

All sulfate concentrations in the uppermost profiles roughly match the sulfate concentration of the Barents
Sea bottom waters (ca. 28 mM). In all cores, sulfate concentrations decrease almost linearly with depth, linear regressions showing correlation coefficients of R² > 0.85 (Fig. 2a-2b and Supplementary Fig. S1). The slopes of the regression lines can be divided into two groups corresponding to one group encompassing reference sites Ref 5 and Ref 6, and another group including the three HC-affected sites Zone 1, Zone 10 and Zone 12. Based on a t-test using the software PAST v. 3.20 (Hammer et al., 2001), the slopes of these two groups are statistically significantly different (t = 5.574, p = 1.25E-6) from each other (Fig 2). The sulfate profiles from HC-affected sites show a steeper decrease with depth than those from reference sites. On average, sulfate concentrations decrease by 10 mmol × m⁻¹, which would correspond to complete sulfate depletion at roughly 3 mbsf at HC-affected sites. At reference sites, the average slope is 3 mmol × m⁻¹ with little variations, which

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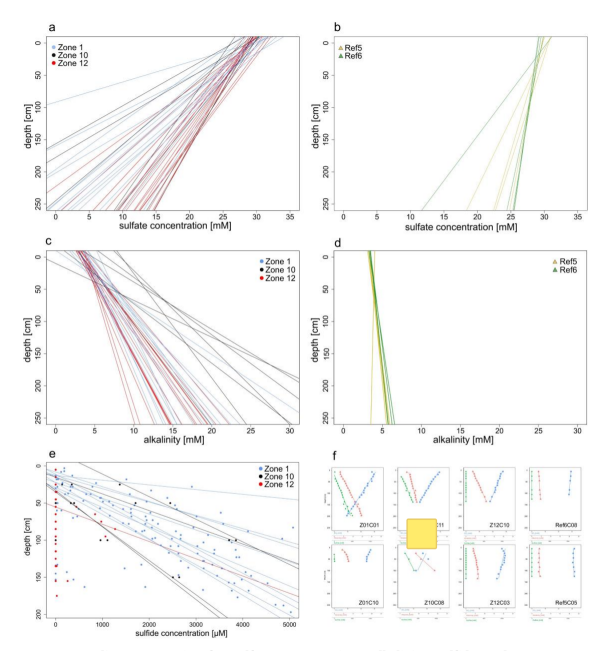


Figure 2. Pore-water linear regression for sulfate concentrations, alkalinity, sulfide, and some representative profiles.

(**a**, **b**) Regression lines for sulfate profiles (derived from measured pore water concentrations) are linear (R^2 330 > 0.85) and exhibit decreasing trends with depth. The regression lines shown in (**a**) represent HC-affected sites (Zone 1, Zone 10, Zone 12), which display a steeper slope compared to those in (**b**) reference sites (Ref 5, Ref 6). (**c**, **d**) Regression lines for pore-water alkalinity (calculated from measured pore water concentrations) are linear ($R^2 \ge 0.85$) and increase with depth. The regression lines for (**c**) HC-affected sites show a more gradual increase compared to (**d**) reference sites. (**e**) Sulfide concentrations measured across all

- 335 cores from HC-affected sites (represented by dots), with their corresponding regression lines shown where sulfide concentrations were quantifiable. (f) Subset of representative profiles selected for each sampling site. The uppermost and lowermost rows depict cores with the highest and lowest R² values, respectively, from HC-affected and reference sites.
- All alkalinity profiles display a clear linear increase with depth ($R^2 \ge 0.85$). From 5 mmol × L⁻¹ at the SWI, pore water alkalinity increases differently with depth, corresponding to 8, 5 and 6 mmol × L⁻¹ × m⁻¹ at HCaffected sites Zone 1, Zone 10, and Zone 12, and ≤ 1 mmol × L⁻¹ × m⁻¹ at references sites Ref 5 and Ref 6, respectively (Fig. 2c-2d and Supplementary Fig. S2).
- Based on the slopes of pore water profiles, the calculated alkalinity fluxes are larger at HC-affected 345 sites than at reference sites. The same is true for sulfate fluxes, but differences across HC-affected sites and reference sites are less pronounced (Supplementary Fig. S3). In comparison, variations observed within each sampling site are small.

Sulfide could hardly be detected in pore water samples collected from reference sites, most values being below 1 μ M. In contrast, sulfide was detected in most of the cores from HC-affected sites. Especially at Zone 1 and Zone 10, sulfide concentrations increase linearly with depth (Fig. 2e). The majority (ca. 70 %)

- of the cores from Zone 1 show an increase of 2.5 mM \times m⁻¹ in sulfide concentrations. One core (Z01C03) stands out, reaching sulfide concentrations of ca. 11 mM at 1 mbsf. Several cores (ca. 57 %) from Zone 10 show a binary distribution of sulfide concentrations, corresponding either to linear profiles reaching ca. 1mM sulfide at 1 mbsf, or to a steeper increase reaching ca. 3.7 mM at the same depth. At Zone 12, sulfide is
- 355 detectable in only 2 of the 17 cores, with much lesser increase in concentrations compared to Zone 1 and Zone 10.

3.2 Measured and modeled sulfate reduction rates

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SRR were detectable in only 29 out of the 50 cores (Fig. 3), with rates between 4 and 320 pmol \times cm⁻³ \times d⁻¹. In 403 of the 455 samples across the 50 cores analyzed (i.e. 89 %), SRR were below the detection limit (1 pmol \times cm⁻¹ \times d⁻¹). The occurrence of very low but detectable SRR increased with sediment depth, which mostly stands true for HC-affected sites (Fig. 3).

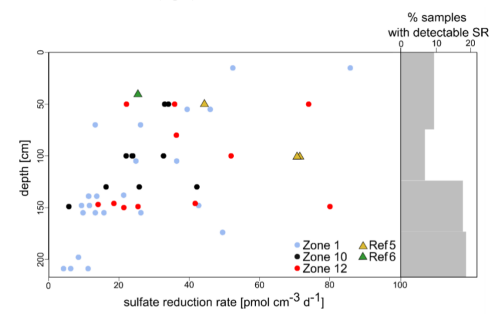


Figure 3. Sulfate reduction rates (SRR) and percentages of samples with sulfate reduction rates above
the detection limit. Low SRR became increasingly detectable with increasing sediment depth. The number of samples with measurable SRR (% grey bars) is higher in sediments from HC-affected sites (dots) compared to reference sites (triangles), *n.B.* due to a lack of samples, the two uppermost intervals were merged.

Diffusion-reaction modeling with PROFILE (Berg et al., 1998) suggests steady SRR in 47 of the 50 370 cores (Supplementary Fig. S1), implying that SRRs remain constant over the entire length of the cores. The modeled SRR profiles show SR activity (i.e. >0 pmol × cm⁻³ × d⁻¹) in 33 cores, of which 6 display rates ≤0.01 pmol × cm⁻³ × d⁻¹, and 27 with supposed SRR ≥1 pmol × cm⁻³ × d⁻¹. For three specific cores (i.e. Z01C02, Z01C09, Z12C17), PROFILE proposes two intervals with different SRRs, suggesting that SR processes may only be active in the upper 50 cmbsf and inactive in the sediment below.

375 The modeled and measured SRR are indicative of net and gross turnover rates, respectively. The gross rate reflects microbial turnover whereas the net turnover rate includes the recycling of reduced sulfur compounds (Berg et al., 1998). The modeled (i.e. net) rates for most samples are higher than the measured (i.e. gross) rates, which is rather unusual and can be explained by very low values whose absolute changes

have a drastic effect on the calculated values. The modeled and measured SRR are nevertheless within the 380 same order of magnitude, and thus comparable.

3.3 Methane and carbon dioxide concentrations

All samples contain measurable methane concentrations, ranging from 0.08 to 19.79 μ mol × L⁻¹. All samples from reference sites contain less than 5 μ mol \times L⁻¹ of methane, detection of any higher concentrations being restricted to some samples from HC-affected sites and only below 100 cmbsf (Fig. 4). CO₂ was detectable in all samples, with concentrations ranging from 29 μ mol \times L⁻¹ to 512 μ mol \times L⁻¹, but without any specific trend with depth (Fig. 4). The reason is probably that any CO_2 from whichever source enters the carbonate equilibrium and the gas is removed from the pore water, both in HC-affected sites and reference sites. The low methane and carbon dioxide concentrations prevented any measurement of their stable isotopic signatures.

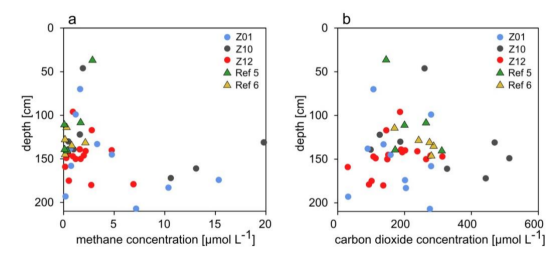


Figure 4. Concentrations of dissolved methane and carbon dioxide. (a) At HC-affected sites, methane concentrations are highest in samples originating from 100 cmblf or deeper, whereas methane concentrations at reference sites remain below 5 μ mol \times L⁻¹. (b) CO₂ concentrations across samples do not display any specific trend with sediment depth.

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3.4 Statistical correlations

To identify geochemical differences between HC-affected and reference sites, we performed a Pearson correlation analysis. Correlation coefficients calculated for each separate gravity core (Table S2) indicate that

- 400 alkalinity is strongly negatively correlated ($p \ge -0.85$) with sulfate concentrations in sediments from HCaffected sites. The same applies to quantifiable sulfate and sulfide concentrations ($p \ge -0.7$). For reference sites, alkalinity and sulfate concentrations are also strongly correlated ($p \approx 0.8$), whereas sulfide concentrations are below the detection limit and hence have been omitted for the correlation analysis.
- We established ratios of sulfate to alkalinity (HCO₃⁻) fluxes (Fig. 5). As about 90% of seawater 405 alkalinity can be contributed to bicarbonate (89.8% HCO₃⁻, 2.9 % CO₃²⁻) (Kerr et al., 2021)we set alkalinity synonymous with bicarbonate concentration. We compared the measured fluxes to those expected from organoclastic (1:2 based on Eq. 1) and methanotrophic (1:1 based on Eq. 2) SRR activities. At reference sites, the average ratio of sulfate to alkalinity fluxes equals 2:1 whereas it is 1.25:1 (5:4) at HC-affected sites, imply that either HCO₃⁻ is being removed from the pore water or that pore water sulfate is eventually being 410 replenished via sulfide oxidation.

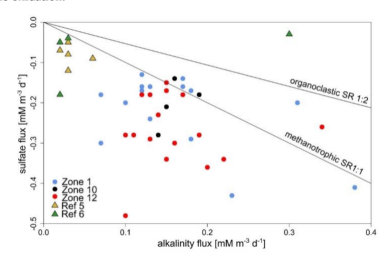


Figure 5. Relationship between sulfate and alkalinity fluxes at HC-affected sites and reference sites. The majority of samples from HC-affected sites (dots) have lower alkalinity than sulfate fluxes. This is even more pronounced at reference sites (triangles). Black lines signify theoretical ratios for methanotrophic (1:1)

415 and organoclastic (1:2) SR.

3.5 Functional marker genes specific to the SMTZ

From the 12 sequenced metagenomes, we extracted a total of 307 234 ORFs with multiple annotations that corresponded to 821 850 predicted genes. From the 5 sequenced metatranscriptomes, we extracted a total of

420 31 093 ORFs corresponding to 47 316 predicted genes (Supplementary Table S2).

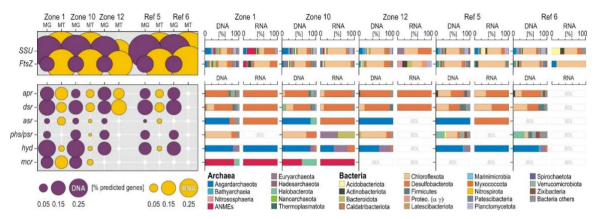


Figure 6. Open reading frames encoding functional marker genes involved in sulfate reduction, production and anaerobic consumption of methane and their taxonomic assignments at the phylum to class level. The relative abundances of certain phyla increase at HC-affected sites compared to reference

sites. Concomitant expression of *apr* and *dsr* genes by Desulfobacterota, *phs* genes by Chloroflexota, *hyd* genes by Asgardarchaeota, and *mcr* genes by ANMEs and Halobacterota are consistent with methanotrophic sulfate-reducing activities. *Abbreviations* - *apr*: adenylylsulfate reductase; *dsr*: dissimilatory sulfite reductase; *asr*: anaerobic sulfite reductase; *phs/psr*: thiosulfate and polysulfide reductase; *hyd*: sulfhydrogenase; *mcr*: methyl coenzyme M reductase; MG: Metagenomics; MT: Metatranscriptomics; BDL:

For each of the targeted marker genes, we plotted the relative percentage of total predicted genes and their corresponding taxonomic assignments at the phylum and class level (Fig. 6). Taxonomic assignments of *SSU* rRNA and *FstAZ* proteins show that metabolically active and growing microbial populations at HC-affected sites are affiliated with the phyla Chloroflexota, Desulfobacterota and Euryarchaeota, and more specifically with anaerobic methanotrophs (i.e. ANMEs). The phyla Desulfobacterota and Chloroflexota are actively engaged in processes of sulfate reduction, as shown by the expression of ORFs encoding the *aprAB* and *dsrAB* genes (Fig. 7a-7b). Taxonomic assignments of expressed ORFs encoding *mcrA* genes show an increased abundance of methanotrophic ANMEs (Fig. 7c) over

440 methanogenic Halobacterota. ORFs encoding *hydB-G* genes are expressed by taxa related to the class Lokiarchaeia and Thorarchaeia (Fig. 7d) among the phylum Asgardarchaeota, indicating metabolic capability to use reduced sulfur compounds (e.g. polysulfide, elemental sulfur). However, their overall expression levels appear limited and these Asgardarchaeota do not seem to actively divide (Fig. 6). At reference sites,

⁴³⁰ below detection limit.

Chloroflexota appear to be the only phylum capable of growing and dividing in these organic-poor sediments

445 (Fig. 6). However, concomitant expression of functional marker genes involved in sulfate reduction remains below the limit of detection. We acknowledge that the non-detection of certain genes may also result from limited sequencing depths.

Phylogenetic trees plotted for transcript ORFs assigned to the *rps3* and *rpoD* taxonomic marker genes (Supplementary Figs. S4-S5) provided concomitant evidence for metabolic activity by the phyla

450 Desulfobacterota, Chloroflexota (class Dehalococcoidia), Acidobacterota (class Aminicenentia), Asgardarchaeota (class Thorarchaeia), and Halobacterota (class ANME-1).

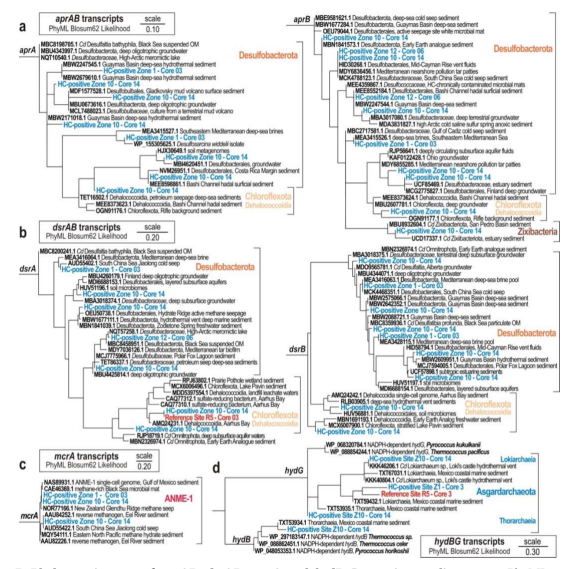


Figure 7. Phylogenetic trees of aprAB, dsrAB, mcrA and hydB-G protein-encoding genes. PhyML

maximum likelihood trees of transcript ORFs encoding: (a) the adenylylsulfate reductase (714 amino acids) subunit alpha (*aprA*) and beta (*aprB*); (b) the dissimilatory sulfite reductase (493 amino acids) subunit alpha (*dsrA*) and beta (*dsrB*); (c) the methyl coenzyme M reductase reductase (698 amino acids) subunit alpha (*mcrA*); and (d) the sulfhydrogenase, or sulfur reductase (286 amino acids), subunit beta (*hydB*) and gamma (*hydG*). Phylogenetic trees are based on 100 bootstrap replicates with BLOSUM62 as evolutive model.
Boldface types signify sequence accession numbers to the NCBI database.

4. Discussion

4.1 Pore water geochemical profiles correlate to meta-omics features

Pore water profiles reflect the spatial distribution of net turnover processes. The shape of the profile allows to distinguish intervals in which transport is purely diffusive (i.e. linear profile) from those in which there is either net production (i.e. convex profile) or consumption (i.e. concave profile) of chemical species (Schulz, 2006; Hesse and Schacht, 2011).

Based on the largely linear profiles for pore water alkalinity, sulfate and sulfide (Figs. 2-3, Figs. S1-S2), we infer the absence or near-absence of net sulfate consumption in the depth interval covered by the

- 470 cores, and a sink for sulfate as well as a source of sulfide and alkalinity at greater depth. Microbially-mediated AOM with sulfate as electron acceptor yields hydrogen sulfide, bicarbonate and water (Eq. 2), providing an explanation for both sink and source (Martens and Berner, 1974; Iversen and Jorgensen, 1985), assuming sufficient supply of methane at greater depth. The occurrence of AOM at deeper depth is further supported by the relationship between sulfate and alkalinity fluxes (Fig. 5) and negligibly small methane concentrations
- 475 (< 10 μM L⁻¹) at the bottom end of all cores from HC-affected sites and reference sites (Fig. 4), indicating almost complete consumption of methane originating from biogenic or thermogenic sources below our sampling interval.

Linear concentration profiles are only obtained in regions with no net consumption or production of the respective compound (Schulz, 2006). Despite the apparent linearity of sulfate profiles (Fig. 2), we observed very low SRR (\leq 300 pmol × cm⁻³ × d⁻¹), confirming previous reports of sporadic SRR in Barents Sea sediments with similar rates above the SMTZ (Nickel et al., 2012). In our case, sulfate reduction is mainly detectable at HC-affected sites (Fig. 3), with a greater prevalence in deeper sediments (Fig. 5). With seepage supplying electron donors, like methane and/or other HCs, from deeper reservoirs to near-surface sediment, sulfate reduction is fuelled by those HCs rather than from the buried and more recalcitrant OM. Inconspicuous

- 485 supplies of methane and/or other HCs to the overlying sediment can stimulate SRR by providing alternative electron donors. Such sporadic energy supply promotes geochemical conditions that stimulate benthic HCdegrading microbial communities. Although these communities can be physiologically and phylogenetically diverse, they are mainly composed of anaerobic guilds of methanogens, methanotrophs, sulfate reducers and fermenters (Teske, 2019).
- 490 The analysis of functional marker genes supports the notion of sporadic HC supply, providing complementary insights into the metabolic potential and activity of specific microbial populations in the

sediment. The occurrence and abundance of aprAB, dsrAB, and mcrA genes supports interpretations of geochemical data in terms of sulfate-driven AOM processes and metabolic activity (Fig. 6). Taxonomic profiling of populations expressing ORFs encoding genes involved in sulfate reduction (i.e. aprAB, dsrAB)

- reveals Desulfobacterota (family Desulfobacteraceae) and Chloroflexota (class Dehalococcoidia) as the prevalent bacterial phyla constituting the consortium of sulfate-reducing bacteria at HC-affected sites (Fig. 7, Supplementary Figs. S4-S5). The *aprAB* and *dsrAB* phylogenetic trees clearly show that none of these genes correspond to their reverse function and the ORFs annotated as *aprAB* and *dsrAB* are predicted to function in dissimilatory sulfate reduction. Moreover, the sediment being fully anoxic, any detection of sulfur
- 500 oxidation via reversible *aprAB* or reverse *dsrAB* was unexpected as such activities are mostly reported for aerobic Gammaproteobacteria (Vuillemin et al., 2022).

Further, ORFs encoding genes involved in anaerobic production and consumption of methane (i.e. mcrA) clearly show that the clade ANME-I exhibits concomitant, although low metabolic activity towards anaerobic oxidation of methane (Fig. 7c), namely they express ORFs encoding the reversible mcrA gene.

- 505 The higher expression levels of methanotrophic clade ANME-I over methanogenic Halobacterota (Fig. 6) also suggest that metabolic activity by ANMEs could be sustained by an alternative source of methane (Dong et al., 2020), although cryptic production of biogenic methane by the ANME-1 clade has also been observed in experiments replicating SMTZ conditions (Beulig et al., 2019). Because methane concentrations measured at both HC-affected sites and reference sites were similarly low (Fig. 4), it appears unlikely that AOM
- 510 processes are sustained by in situ production of biogenic methane only (Dong et al., 2020). Finally, the class Lokiarchaeia and Thorarchaeia in the phylum Asgardarchaeota expressed ORFs encoding beta and gamma subunits of hydB-G genes (Fig. 7d), and thereby appear metabolically active, using various sulfur and sulfide species, potentially produced as by-products of sulfate reduction.

In contrast, Chloroflexota (class Dehalococcoidia) was the only phylum at reference sites showing 515 sufficent metabolic activity was adequate to achieve cell division (i.e. FstAZ). In the absence of an univocal signal towards active dissimilatory sulfate reduction by Dehalococcoidia at HC-affected sites (Fig. 7b), we infer that these Chloroflexota could persist and gain energy through fermentation processes in organic-lean marine sediments instead (Vuillemin et al., 2020a). Due to this metabolic versatility, Chloroflexota were found to be active at both HC-affected and reference sites, which rank them as poor microbial indicators for

520 minor HC seepage (Iasakov et al., 2022).

Together, these microbial patterns extracted from meta-omics corroborate the geochemical data, arguing in favor of discrete AOM processes concomitant to sulfate reduction at HC-affected sites, and for

limited in situ production of biogenic methane at both HC-affected sites and reference sites. Alongside an active AOM consortium composed of Desulfobacterota and ANME-1, concomitant production of reduced

525 sulfur compounds could apparently stimulate metabolic activity by Lokiarchaeia and Thorarchaeia (Figs. 6 and 7).

4.2 Geochemical indicators for minor HC seepage

The combined use of meta-omics microbial patterns with the determination of SR and related geochemical features and fluxes in upper sediments, enables to a certain extent to distinguish between HC-affected sites

- 530 and reference sites. Sediments of the SW Barents Sea are organic-poor (TOC <0.5 %), and thus characterized by a lack of electron donors (Knies and Martinez, 2009) and low microbial activity (Nickel et al., 2012; Nickel et al., 2013) with consequent sulfate penetration to greater sediment depths. In organic-rich sediments, any signal inherent to minor HC seepage fueling SR would be difficult to detect as it would be covered by organoclastic SR processes. The organic-lean nature of Barents Sea sediments should enable the detection of</p>
- 535 discrete HC seeps, because the slightest supply of electron donors leads to a direct increase in the metabolic activity of HC-degrading microbial populations. This slight increase in metabolic activity can then be discriminated from the background signal observed at reference sites. Diffusion of HCs was thus expected to promote sulfate consumption, resulting in its depletion in the pore water at shallower sediment depths. This, combined with variable fluxes of thermogenic and biogenic methane, can account for variations in the depth
- 540 of the SMTZ.

At different locations of the SW Barents Sea, the depth of the SMTZ, or rather the sulfate depletion depth, was postulated, for instance, at 37 mbsf in the Hammerfest/Loppa High and the Tromsø Basin/Ingøydjupet area (Nickel et al., 2012), and between 3.5 and 29.2 mbsf in the Ingøydjupet area (Argentino et al., 2021a). These values are consistent with the global average depth of the SMTZ (12.8 ± 12.1 mbsf) on the continental slope (Egger et al., 2018). For comparison, at the Vestnesa Ridge which is under the influence of gas hydrates, the depth of the SMTZ fluctuates massively between seepage (ca. 1 mbsf)

and non-seepage (ca. 3 to 75 mbsf) sites (Hong et al., 2016). Similarly, Barents Sea sediments with and without methane seeps show an SMTZ located between a few cm and about 2 mbsf (Argentino et al., 2021b).

Linear extrapolation of sulfate profiles (Fig. 2) indicates complete sulfate depletion, indicative of an 550 SMTZ, at 12 and 3 mbsf for reference and HC-affected sites, respectively. We suggest that the difference in sulfate penetration depth is caused by upwards migration of HCs. The sulfate and alkalinity fluxes inherent to organoclastic (Eq. 1) and methanotrophic (Eq. 2) SR have respective theoretical ratios of 1:2 and 1:1 while mixed SR processes have intermediate values. Current ratios of sulfate to alkalinity fluxes (Fig. 5) at reference sites imply that either HCO_3^{-1} is being removed from the pore water, or that pore water sulfate is eventually

- 555 being replenished via sulfide oxidation. From previous work in the area, we know that the sediment is fully anoxice v the uppermost few cm. Phylogenetic analysis of *aprAB* and *dsrAB* genes clearly shows that none of the identified amino acid sequences correspond to their reverse function. Further, we did not identify any *dsrC* sequences, which potentially work in cellular sulfur oxidation (Ruiz-Blas et al., 2024). While dissolved inorganic carbon (DIC) can be assimilated via acetogenesis, methanogenesis, or methanotrophy
- 560 (Kellermann et al., 2012), authigenic carbonate minerals are known to precipitate from the pore water as byproducts of AOM processes (Turchyn et al., 2021), sometimes even forming carbonate hardground at the seabed. However, from the low level of detection of AOM-related transcripts, we inferred only a minor imprint on the geochemistry and therefore assume that bicarbonate ions originating from AOM processes at greater depths are diffusing upward and eventually precipitate as calcium carbonate. Up to 80 % of the sulfide
- 565 produced through SR can also be converted back into sulfate through reoxidation of reduced sulfur compounds (Jørgensen et al., 2004) and disproportionation (Pellerin et al., 2015; Jørgensen et al., 2019a). Finally, the dissolution of sulfate minerals, such as barite (BaSO₄), may also act as an additional source of sulfate (Griffith and Paytan, 2012). Altogether, this suggests either the reoxidation of sulfide, or the removal of bicarbonate from the pore water, e.g. by mineral precipitation, or both. The former is further supported by
- 570 concomitant sulfate reduction, detected by radiotracer incubations (Fig. 3) and microbial gene expression patterns (Fig. 6), the latter by the ratios of sulfate to alkalinity flux (Fig. 5). Thus, we could identify several biogeochemical features that argue for ongoing sulfate reduction (most likely AOM-related) at HC-affected sites below our sampled depth range, which allow their discrimination from reference sites. Given the sulfate concentration profiles and the small amounts of methane detected in the samples, it becomes clear that AOM
- 575 is not a quantitatively important process in the cored depth interval, producing an erratic signal in HC-affected sediments.

5. Conclusions

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Our study, combining modeled and measured SRR as well as pore water geochemistry with functional marker genes, revealed remarkable differences between samples originating from a pristine seafloor and those from areas affected by small and inconspicuous HC seepage. The linear shape of pore water profiles was indicative of purely diffusive transport under steady-state conditions and allowed estimation of the depth of sulfate depletion, which appeared markedly different between HC-affected and reference sites. At the HC-affected sites, we found higher pore water fluxes of sulfate, sulfide and alkalinity, and could also demonstrate active

- 585 microbial SR by turnover measurements and modeling. Furthermore, meta-omics provided additional evidence for enhanced metabolic activity related to microbial sulfate reduction coupled with minor AOM processes at HC-affected sites, which could be clearly discriminated from the background signal inherent to fermentative activities and minor *in situ* production of methane by the subseafloor biosphere. These observations demonstrate that minor HC seepage, which is not even visible at the seafloor, significantly
- 590 influences sedimentary biogeochemical cycles driven by microbial populations resident in organic-lean sediments.

6. Data availability

Geo ical data, ORFs sequences and their BlastP outputs are available as Supplementary Information.

7. Author contributions

595 JK and RDP organized the sampling cruise. ES and JK designed the study and sampled the cores. ES performed sulfate, alkalinity, SRR and gas analyses. AV, BK, CL and AS performed DNA and RNA analyses. ES and AV wrote the manuscript and all authors contributed to writing and revision.

8. Competing interests

The authors declare that they have no conflict of interest.

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