# Animal burrowing at cold seep ecotones boosts productivity by linking macromolecule turnover with chemosynthesis and nutrient cycling

Maxim Rubin-Blum<sup>1,2\*</sup>, Eyal Rahav<sup>4</sup>Rahav<sup>2</sup>, Guy Sisma-Ventura<sup>4</sup>Ventura<sup>2</sup>, Yana Yudkovski<sup>4</sup>Yudkovski<sup>2</sup>, Zoya Harbuzov<sup>4</sup>Harbuzov<sup>2</sup>, Or Bialik<sup>2,3</sup>, Oded Ezra<sup>2</sup>Ezra<sup>1</sup>, Anneleen Foubert<sup>4</sup>, Barak Herut<sup>1,2</sup>, Yizhaq Makovsky<sup>2</sup>Makovsky<sup>1</sup>

<sup>4</sup>Israel<sup>1</sup>Charney School of Marine Sciences (CSMS), University of Haifa, Haifa, Israel

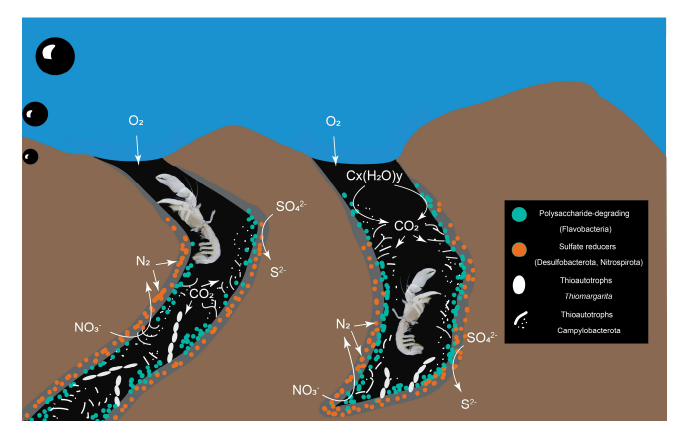
<sup>2</sup>Israel Oceanographic and Limnological Research, National Institute of Oceanography, Haifa, Israel
 <sup>2</sup>Charney School of Marine Sciences (CSMS), University of Haifa, Haifa, Israel
 <sup>3</sup>Institute of Geology and Palaeontology, University of Muenster, Münster, Germany
 <sup>4</sup>Department of Geosciences, University of Fribourg, Fribourg, Switzerland
 \*Correspondence: mrubin@ocean.org.il

# Abstract

Hydrocarbon seepage at the deep seafloor fuels flourishing chemosynthetic communities. These seeps impact the functionality of the benthic ecosystem beyond hotspots of gas emission, altering the abundance, diversity and activity of microbiota and fauna, and affecting geochemical processes. Yet, these chemosynthetic ecotones (chemotones) are far less

- 5 explored than the foci of seepage. To better understand the functionality of chemotones, we: i) mapped seabed morphology at the periphery of gas seeps in the deep Eastern Mediterranean Sea, using video analyses and synthetic aperture sonar; ii) sampled chemotone sediments and described burrowing using computerized tomography; iii) explored nutrient concentrations; iv) quantified microbial abundance, activity and N<sub>2</sub> fixation rates in selected samples and v) extracted DNA and explored microbial diversity and function using amplicon sequencing and metagenomics. Our results indicate that the
- 10

gas seepage yields gradients of burrowing intensity at the seep ecotones, especially by the ghost shrimp *Calliax lobata*. This burrowing alters nitrogen and sulfur cycling through the activity of diverse microbes. Burrow walls form a unique habitat, where macromolecules are degraded by Bacterioida, and their fermentation products fuel sulfate reduction by Desulfobacterota and Nitrospirota. These in turn support chemosynthetic Campylobacterota and giant sulfur bacteria *Thiomargarita*, which can aid *C. lobata* nutrition. These interactions may support enhanced productivity at seep ecotones.



15

#### 1. Introduction

Most organisms in the deep-sea biosphere thrive under extreme energy limitation (Orsi, 2018). In this dark, energy-limited environment, the natural discharge of fossil hydrocarbons results in accelerated biogeochemical dynamics, creating unique

- 20 geobiological habitats - cold seeps (Joye, 2020). Cold seeps form the richest and most productive ecosystems that are widespread across continental margins (Joye, 2020; Levin and Sibuet, 2012). The transition zones between these highly productive ecosystems and the often impoverished deep-sea benthic habitats are considered chemosynthetic ecotones, that is, chemotones (Ashford et al., 2021). The chemotone communities benefit from chemosynthetic production, which can deliver food supplements through the water column, and intrinsic chemosynthetic producers in the sediments, resulting in 25 intermediate biomass, richness and distinct species composition (Ashford et al., 2021). These areas of interactions and transition affect the deep-sea ecosystem services, most importantly, carbon cycling and sequestration, as well as fisheries production (Levin et al., 2016). However, our knowledge of chemotone communities is limited, and their functionality has not been studied in detail, e.g. (Amon et al., 2017; Ashford et al., 2021; Åström et al., 2022; Ritt et al., 2011).
- Chemical and temperature clines around seep edifices may determine the biogeochemical processes at chemotones (Levin 30 et al., 2016). In turn, surface water productivity and fluxes of photosynthetic carbon to the seep area may also impact the extent of seep chemotones (Ashford et al., 2021). For example, the strong pelagic-benthic coupling is limiting the chemosynthetic inputs in some Arctic seeps (Åström et al., 2022). Cold seeps are also pervasive in the warm (13.5 and 15.5 °C) deep waters of the ultraoligotrophic Southeastern Mediterranean Sea (SEMS) (Bayon et al., 2009; Coleman and Ballard, 2001; Coleman et al., 2011; Herut et al., 2022; Kormas and Meziti, 2020; Lawal et al., 2023; Olu-Le Roy et al., 2004). In 35 high contrast to the Arctic seeps, the weak benthic-pelagic coupling in the ultraoligotrophic SEMS may lead to sharp gradients between the active seep sites and the energy-limited deep benthos (Sisma-Ventura et al., 2022). The warm deepsea temperatures may alter the availability of dissolved gases and boost microbial activity in SEMS, potentially affecting

the biogeochemical gradients at seep clines (Mavropoulou et al., 2020; Rahav et al., 2019; Roether and Well, 2001).

Here we focus on cold seep chemotones at the toe region of Palmahim Disturbance, an evaporite rooted, progressive, large-40 scale (~15 x 50 km), rotational submarine slide on the SEMS continental margin, offshore Israel (Gadol et al., 2020; Garfunkel et al., 1979); (Fig. 1). These seeps are characterized by local hotspots of slow diffusive discharge, where authigenic carbonates are formed (Rubin-Blum et al., 2014a, 2014b; Weidlich et al., 2023) with dispersed bivalve beds, indicative of substantial past, and limited present activity (Beccari et al., 2020). Yet, hotspots of marked discharge of gasrich brinesbrine, forming small (<40 m wide) brine pools, were recently discovered (Herut et al., 2022). The sediments at 45 Palmahim Disturbance seep chemotones do not host the typical seep fauna but are inhabited by dense populations of a burrowing ghost shrimp, Calliax lobata, which alter the morphology of the sediment surface for tens of meters away from the seepage hotspots (Basso et al., 2020; Beccari et al., 2020; Makovsky et al., 2017). Similar burrowing is prominent in the periphery of SEMS seeps, as it was, for example, observed in the hemipelagic sediments near the Amon mud volcano at the Nile Deep Sea Fan (Blouet et al., 2021; Ritt et al., 2011). At Palmahim Disturbance, the sediments in the seep 50 periphery are characterized by elevated nutrient fluxes and oxygen consumption, as well as increased microbial activity

compared to the background stations (Sisma-Ventura et al., 2022).

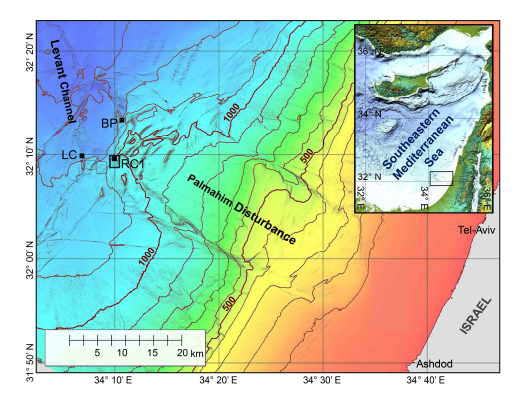


Figure 1. A bathymetric map of Palmahim Disturbance and its vicinity (color-coded and contoured with a 100 m spacing). This study is focused on seafloor gas seeps in the ridge-crest pockmark system (RC1) at the southwestern edge of the Disturbance, the proximate part of the Levant Channel (LC), and the brine pool (BP) area to the north. Black dots mark the sites of box-core sampling, and the black frames outline the ROV surveying analysis maps of Figure 2D-E.

60

Thus, microbes likely play a key role not only in hotspots of seepage but also in the seep ecotones. However, to date, most studies of seep microbiota focus on seepage hotspots. At these hotspots, seep ecosystems support diverse assemblages of microorganisms, with cosmopolitan distribution and key biogeochemical functions (Ruff et al., 2015; Teske and Carvalho, 2020). At the seepage hotspots, microbes catalyze key processes such as anaerobic oxidation of methane (AOM), sulfate reduction and chemosynthesis (Boetius and Wenzhöfer, 2013; Knittel and Boetius, 2009; Teske and Carvalho, 2020). The

marked primary productivity at seeps may boost secondary microbial productivity, via, for example, the fermentation of organic macromolecules (Zhang et al., 2023). Such microbial communities are characteristic of Palmahim Disturbance seepage hotspots (Rubin-Blum et al., 2014b). Yet, little is known about the diversity and function of microbial communities
that occupy the seep periphery, in the SEMS seeps in particular.

Here, we aimed to understand the extent and functionality of the burrowed sediments at Palmahim Disturbance seep chemotones. We thus mapped the bioturbation using analyses of video collected with remotely operated vehicles (ROVs) and high-resolution acoustic mapping with synthetic aperture sonar (SAS), mounted on an autonomous underwater vehicle (AUV). Hypothesizing that the bioturbated sediments at Palmahim Disturbance seep chemotones host unique microbial

70 communities, which may function synergistically to exploit limited fluxes of seep carbon, we investigated their diversity and function using metagenomics analyses of samples collected at the seep periphery (Sisma-Ventura et al., 2022). We investigated the activity of chemotone microbes using the leucine uptake assay and evaluated their ability to fix dinitrogen. Previous studies suggest that burrowing shrimps, in particular, ghost shrimps (Axiidea), garden and groom their burrows by incorporating organic matter into the walls through mucus excretions, creating microniches within the burrow walls

75 (Abed-Navandi et al., 2005; Astall et al., 1997; Coelho et al., 2000; Dworschak et al., 2006; Gilbertson et al., 2012; Laverock et al., 2010; Papaspyrou et al., 2005). In seeps, burrowing may enhance carbonate precipitation in areas with very low methane flux and/or diffusive seepage (Weidlich et al., 2023). We thus hypothesized that the burrowing activity of *C*. *lobata* can alter the diversity and functionality of these microbes, introducing new metabolic adaptations and handoffs. Therefore, we focused on the specific sediment layers where burrows were present.

# 80 2. Methods

#### 2.1 Exploration of the Palmahim Disturbance gas seeps

Palmahim Disturbance toe region comprises a set of hundreds of meters high and ~1 km wide fold ridges and is bounded to the west by the ~500 m wide and ~40 m deep easternmost channel of the deep sea fan of the Nile River, the Levant channel (Gvirtzman et al., 2015). We investigated three seepage domains: i) Hundreds of meters wide pockmark systems
at the crest of three ridges marking the southwest of the Palmahim Disturbance toe, with a particularly large pockmark at the crest of the eastern ridge (RC1; Figure 1); ii) Elongate pockmarks along the sides of the Levant Channel (LC; Figure 1); iii) Brine pools at the northern part of Palmahim Disturbance toe region (BP; Figure 1). We collected sediment samples and survey data during several cruises to the Palmahim Disturbance offshore Israel, comprising: i) The E/V Nautilus cruise in 2011 (Coleman et al., 2012); ii) EUROFLEETS2-SEMSEEP cruise in the framework of the EUROFLEETS2 expeditions
in 2016, onboard R/V AEGAEO (Basso et al., 2020; Makovsky et al., 2017) and iii) R/V Bat-Galim cruises in 2020-23 (Herut et al., 2022; Rubin-Blum et al., 2024; Sisma-Ventura et al., 2022). In the latter cruises, sediment collection was supplemented with an AUV-based seafloor survey using a SAS (Kraken Robotics Inc. MinSAS-120), providing a backscatter image of the seafloor with a constant ultra-HD 3x3 cm resolution across a 120 m wide swath along each side

of the AUV track. The SAS data were collected in the framework of a 70 km long reconnaissance survey that traversed thePalmahim Disturbance toe region. SAS images collected in some specific areas were manually correlated with the relevant

# ROV videos by identifying specific seafloor morphologies.

# 2.2 Burrowing quantification

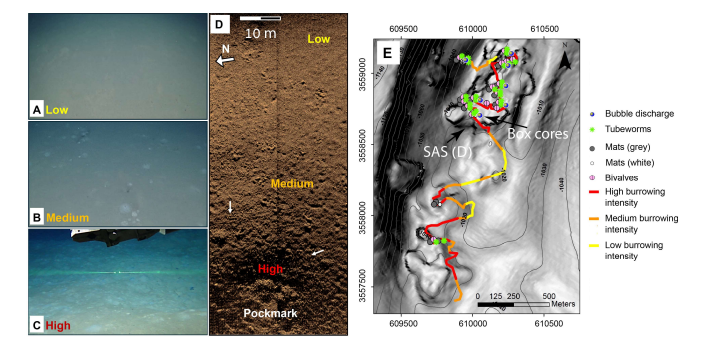
The HD video data of three consecutive dives of the EV Nautilus 2011 cruise to the three ridge-crest pockmarks in the southwestern part of Palmahim Disturbance toe (RC1, Figure 1) were systematically analyzed in an ArcGIS desktop (Esri).
The intensity of burrowing was semi-quantified by classifying to three levels the density of burrowing mounds observed on the seafloor in the field of view of the ROV camera during surveying, and by SAS (Figure 2, A-D): i) Low – where the seafloor is generally smooth with the possible presence of a few sporadic burrowing mounds; ii) Medium – when multiple burrowing mounds are continuously viewed; and iii) High – where burrowing mounds are pervasive across the majority of the field of view. Analysis mapped markers of active seepage, including bubbles discharge, bacterial mats, chemosynthetic fauna, and seafloor burrowing. The results were combined to generate a mapsmap of seepage indicator distribution (Figure 2, E).

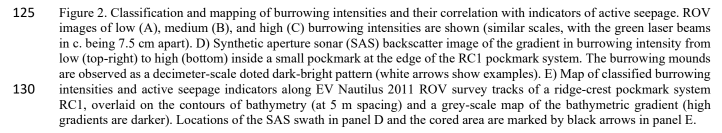
### 2.3 Sediment sample collection

Sediments were sampled using box-corers, targeted with an ultra-short baseline (USBL) positioning system in the large ridge-crest pockmark RC1 (AG16-17BC1 (2016), RC20-BC1 and RC20-BC2 (2020), all within ~20 m positioning accuracy

- of 32° 09.60' N, 34° 10.00' E, water depth circa 1035 m) (Figure 1, Table 1). Additional RC20-Flare1 core was collected within RC1, corresponding to a gas flare detected by acoustics (32° 9.76' N, 34° 10.12'). In 2016, we collected an additional core within the pockmark in the western flank of Levant Channel (LC; AG16-15BC1, 32°09.88' N, 34°06.88' E, water depth 1261 m). In 2023, we collected two box corer samples near the Palmahim Disturbance brine pool site (BP; BP22-BC1 and BP22-BC2, 32° 13.33' N, 34° 10.72' E and 32° 13.32' N, 34° 10.71' E, ~10 and 20 m away from the brine pool, respectively, water depth ~1135 m). The sediments in each of the box corers were subsampled using ~40 cm-long push corers (60 mm diameter) and sectioned onboard at 1 cm resolution. We also collected distinct samples from the same section, based on the black coloration of the burrow walls and bright coloration of the ambient sediments. The sections for DNA extractions were frozen and kept at -20 °C until further processing. Additional push cores were collected from the same box-core samples, kept intact, and stored at 4 °C for X-ray computerized tomography (CT) scanning.
- 120 Table 1: Box core samples used in this study. RC1 is a ridge-crest pockmark system, LC is Levant Channel, BP is a brine pool area (Figure 1). The following analyses were performed: CT scan (1), nutrient profiles (2), flow cytometry (3), activity measurements (4), amplicon sequencing of the 16S rRNA gene (5), metagenomics (6) and dinitrogen fixation (7).

Sample	Year	Depth (m)	Latitude	Longitude	Location	Burrowing	Analyses
AG16-17BC1	2016	1034	32°09.62′	34°10.02′	RC1	High	1,2,3,4,5,7
AG16-15BC1	2016	1242	32°09.89′	34° 06.92′	LC	Low	1,2,3,4,5,7
RC20-BC1	2020	1035	32° 09.60′	34°10.00′	RC1	High	5,6
<i>RC20-BC2</i>	2020	1035	32° 09.60′	34°10.00′	RC1	High	5,6
<i>RC20-BC3</i>	2020	1035	32° 09.60′	34°10.00′	RC1	Low	5
RC20-Flare1	2020	1035	32° 09.76′	34° 10.12′	RC1	Low	5
BP22-BC1	2022	1135	32° 13.33′	34° 10.72′	BP	Low	5
<i>BP22-BC2</i>	2022	1135	32° 13.32′	34° 10.71′	BP	High	5





#### 2.4 Computerized tomography (CT) scans

- We performed computer tomographic (CT) scanning using a Siemens Somatom Definition AS62 installed at the Institute
  of Forensic Medicine (University of Bern, Switzerland). Whole core sections were scanned at a spatial resolution of 300 µm, using energies of 120 kV and effective power of 35 mA. Image analyses and processing (combined single thresholding and watershed segmentation, labeling and quantification) were done using Avizo 9.4 (FEI Thermo Scientific). Before segmentation and labeling, images were filtered using a non-local mean filter. Bioturbation features have been segmented and quantified for each slice across the full scanned volume. Results are presented as volume % vs. depth (each 400 µm
  from top to bottom). For the overall core section, bioturbation has been described according to the Bioturbation Index (BI) (Reineck, 1963) and assessed qualitatively (taking a descriptive approach) for approximately every 10 cm interval of boxcore. This method assesses bioturbation on a scale from 0-6, where 0 indicates 0% bioturbation, 1 corresponds to 1-4 % bioturbated sediment, 2 between 5 to 30 % bioturbation, 3 between 31 to 60 % bioturbation, 4 between 61 to 90 % bioturbation, 5 between 91 to 99 % bioturbation and 6 reflects 100% bioturbation. The shape and size of the burrows were
- described according to the length and thickness/diameter of the burrowing and their orientation (Jumars et al., 2007).

#### 2.5 DNA extraction and sequencing

DNA was extracted from the sediment (~500 mg wet weight) samples with the legacy DNeasy PowerSoil kit (Qiagen, USA), or the FastDNA Spin Kit for Soil (MPBio, USA), using the FastPrep-24<sup>™</sup> Classic (MP Biomedicals, USA) beadbeating to disrupt the cells (2 cycles at 5.5 m sec<sup>-1</sup>, with a 5 min interval). The V4 region (~ 300 bp) of the 16S rRNA gene

150 was amplified from the DNA (~50 ng) using the 515Fc/806Rc primers (5'- GTGYCAGCMGCCGCGGTAA, 5'-GGACTACNVGGGTWTCTAAT) amended with either CS1/CS2 tags (2020 samples) or SP1/SP2 tags (2023 samples) (Apprill et al., 2015; Parada et al., 2016). PCR conditions were as follows: initial denaturation at 94 °C for 45 s, 30 cycles of denaturation (94 °C for 15 sec), annealing (15 cycles at 50 °C and 15 cycles at 60 °C for 20 sec) and extension (72 °C for 30 s) (Sisma-Ventura et al., 2022). Library preparation from the PCR products and sequencing of 2x250 bp Illumina 155 MiSeq reads was performed by HyLabs (Israel) in 2020 and Syntezza Bioscience (Israel) in 2021. Five metagenomic libraries were constructed and sequenced to a depth of ~70 million 2x150 Illumina reads at Hylabs (Israel) (2 libraries from the burrows: 9-10 cm section in RC20-BC1, 8-9 cm section in BOX 2; 3 control libraries from RC20-BC1: 1, 2 and 5, 25

and 29 cm below sediment surface).

#### **2.6 Bioinformatics**

160 Demultiplexed paired-end reads were processed in QIIME2 V2021.4 environment (Bolyen et al., 2019). Primers and sequences that didn't contain primers were removed with cutadapt (Martin, 2011), as implemented in QIIME2. Reads were truncated based on quality plots, checked for chimeras, merged and grouped into amplicon sequence variants (ASVs) with DADA2 (Callahan et al., 2016), as implemented in QIIME2. The amplicons were classified with scikit-learn classifier that was trained on Silva database V138 (16S rRNA, Glöckner et al., 2017). Mitochondrial and chloroplast sequences were 165 removed from the 16S rRNA amplicon dataset. Downstream analyses were performed in R V3.6.3 (R Core Team, 2024),

using packages phyloseq (McMurdie and Holmes, 2013) and ampvis2 (Andersen et al., 2018).

Metagenomic libraries were processed using ATLAS V2.3 (Kieser et al., 2020), with SPAdes V3.14 de-novo assembler (Prjibelski et al., 2020) with --meta -k 21,33,66,99 flags. We used metaBAT2 (Kang et al., 2015) and maxbin2 (Wu et al., 2015) as binners, DAS Tool (Sieber et al., 2018) as a final binner for metagenome-assebled genome (MAG) curation.

- 170 MAGs were dereplicated with dREP (Olm et al., 2017), using a 0.975 identity cutoff. We used CheckM2 to assess the completeness of MAGs (Chklovski et al., 2023), assigned taxonomy with GTDB-Tk V1.5 and GTDB R202 taxonomy (Chaumeil et al., 2020) and annotated them with the SEED and the rapid annotation of microbial genomes using Subsystems Technology (RAST) (Overbeek et al., 2014). Identification of key functions was based on the hidden Markov model (HMM) profiles within METABOLIC (Zhou et al., 2022). Metagenomic reads and MAGs were deposited to the NCBI 175 under project number PRJNA1072319.

#### 2.7 Prokaryote abundance

Total prokaryotes abundance (bacteria and archaea) was quantified from both the upper sediment (top 18-25 cm, every  $\sim 2$ cm) and above the seabed (50 m, every 5-20 m) using flow cytometry. Sediment samples (~3 gr) were collected into sterile 15 ml plastic tubes containing 6 ml of pre-filtered seawater (0.2 µm) from the same station, fixed with a flow cytometry

- 180 grade glutaraldehyde solution (1% final concentration, Sigma, G7651) and kept in 4 °C until analyses within ~2 weeks. At the lab, the chelating agent sodium pyrophosphate (final concentration 0.01 M) and the detergent Tween 20 (final concentration 0.5%) were added. The samples were then vortexed at 700 rpm, placed in an ultrasonic water bath (Symphony, VWR) for 1 min to disperse the grain-attached prokaryotic cells into the liquid seawater phase, and left overnight at 4 °C. At the lab, subsamples were stained for 10 min in the dark with 1 µl SYBR green (Applied Biosystems)
- 185 cat #S32717) per 100 µl of the sample. The samples were analyzed by an Attune® acoustic focusing flow cytometer at a flow rate of 25 µL min<sup>-1</sup>. Taxonomic discrimination was based on cell side scatter, forward scatter, and green fluorescence. Cell abundance was normalized to the sediment's dry weight.

#### 2.8 Heterotrophic activity

Samples of ~3 gr were collected across the upper sediment (18-25 cm, see subsection 2.5), re-suspended in 3.5 ml of prefiltered seawater/porewater (0.2 µm) and spiked with 500 nmol L<sup>-1</sup> (final concentration) of [4,5-<sup>3</sup>H]-leucine (Perkin Elmer USA, specific activity 160 Ci mmol<sup>-1</sup>). Porewater from 5-25 cm was first degassed from O<sub>2</sub> by bubbling into them N<sub>2</sub> for ~10 min, thus keeping them anoxic. The samples were incubated in the dark for 4-5 h under in situ temperatures. At the end of the incubation, 100 µl of 100% trichloroacetic acid solution was added to stop any microbial assimilation of leucine, followed by sonication (Symphony, VWR) for 10 minutes to remove bacterial biomass from the sediment grains (Frank et

- 195 al., 2017). The bacterial biomass extract from the liquid phase and divided into three 1 ml aliquots. For the seawater samples, the collected material was added with 100 nmol L<sup>-1</sup> of [4,5-<sup>3</sup>H]-leucine (same working solution as above, final concentration), incubated for 4-5 h in the dark, and 100 µl of 100% trichloroacetic acid solution was added to stop the incubation. The sediment's liquid extracts were then processed using micro-centrifugation (Smith et al., 1992). Disintegration per minute (DPM) from each sample was read using a Packard Tri carb 2100 liquid scintillation counter. A
- 200

0 conversion factor of 1.5 kg C mol<sup>-1</sup> with an isotope dilution factor of 2.0 was used to calculate the carbon assimilation rate (Simon and Azam, 1989). Blanks included sediments added with [4,5-<sup>3</sup>H]-leucine and immediately with trichloroacetic acid, incubated and processed under the same conditions as described above.

#### 2.9 Nitrogen fixation rates

We evaluated N<sub>2</sub> fixation rates in the top sediment (upper 3 cm) and overlying seawater ~1 m above the seabed, in push
cores AG16-15BC1 and AG16-15BC1 sampled in 2016. The sediment samples (~150 ml) were placed in 250 ml gas-tight
bottles filled with porewater from the same sediment layer (~100 ml) and <sup>15</sup>N<sub>2</sub> enriched seawater (10% of the water volume),
as previously described (Mohr et al., 2010). In short, <sup>15</sup>N<sub>2</sub> gas (99%, Cambridge Isotopes) was injected at a 1:100 ratio (*v:v*)
to filtered (0.2 µm, PALL) and degassed (MiniModule G543) seawater (FSW) or porewater (FPW) and shaken vigorously
for ~12 h to completely dissolve the <sup>15</sup>N<sub>2</sub> gas. The sample bottles were incubated in the dark under an ambient temperature
of ~15 °C for 168-172 h. After incubation, the bottles were vortexed for 30 min at 700 rpm, placed in an ultrasonic water
bath (Symphony, VWR) for an additional 10 min, left on the bench for ~1 h, and filtered using pre-combusted GF/F filters,
following 100 µm mesh pre-filtration to remove large aggregates. For the overlying seawater measurements, 4.5 L were
placed in darkened Nalgene bottles with <sup>15</sup>N<sub>2</sub> enriched seawater (10% of the water volume). The bottles were incubated for
72 h under ambient temperature and the content was filtered onto pre-combusted GF/F filters. Triplicate bottles without

<sup>15</sup>N enrichment were used for determining the natural abundance of the particulate matter at each station. The filters were kept at -20 °C in sterile petri plates until dried at 60 °C overnight and analyzed on a CE Instruments NC2500 elemental analyzer interfaced with a Thermo-Finningan Delta Plus XP isotope ratio mass spectrometer (IRMS). For isotope ratio mass spectrometry, a standard curve to determine N mass was generated for each sample run. Based on natural abundance, N mass on the filters, incubation times, and precision of the mass spectrometer, our calculated detection limit for <sup>15</sup>N uptake
 was 0.01 nmol N L<sup>-1</sup> d<sup>-1</sup>. N<sub>2</sub> fixation rates were calculated based on equations 2 and 3 after Mulholland et al., (2006) using

Equation 2 
$$N2 \ fixation = \frac{[\text{Atom}\% \ (\text{final}) - \text{Atom}\% \ (\text{t0})]}{[\text{Atom}\% \ N2 - \text{Atom}\% \ (\text{t0})]} x \frac{\text{PN}(\text{final})}{\Delta t}$$

Equation 3 
$$Atom\% = \frac{15N}{(15N + 14N)} \times 100$$

Where the Atom%<sub>(final)</sub> and Atom%<sub>(t0)</sub> represent the fractional <sup>15</sup>N enrichment of the particulate nitrogen (PN) pool after the incubation and of the ambient seawater, respectively. Atom% N<sub>2</sub> is the fractional <sup>15</sup>N enrichment of the N<sub>2</sub> source pool. PN<sub>(final)</sub> is the concentration of the particulate nitrogen at the end of the incubation, and  $\Delta t$  is the incubation length (72 h for the seawater and 168-172 h for the sediment incubations).

# 3. Results and Discussion

225

N solubility factors of Weiss (1970).

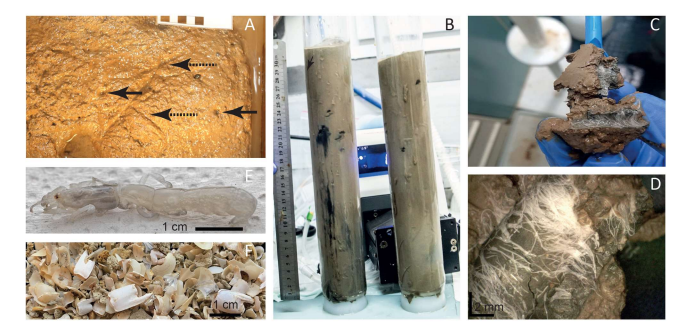
### 3.1 Burrowed sediments are concentrated around Palmahim Disturbance seepage hotspots

- While burrow mounds are rare in the generally smooth seafloor in the bathyal SEMS, these seabed features were particularly abundant at the toe region of Palmahim Disturbance. The seabed burrow extensions appeared in two main forms: i)
  decimeter-scale circular sediment mounds with mostly bright color, often similar to that of the surrounding seafloor and ii) several millimeters to decimeter-scale holes, commonly surrounded by scrape marks (Figure 2). The borrow density varied from only a few sporadic borrows (low intensity) to a complete convolution of the seafloor within an ROV field of view (high intensity; Figure 2). In the regions of intense burrowing, we observed rugged seafloor with localized depressions and mounds (bright color, often dark heaps).
- 235 The video analysis and mapping of active seepage markers along the ROV dive tracks, as well as the SAS surveying, revealed that bioturbation intensities increased towards the inner part of a pockmark (Figure 2E). The high burrowing intensity is often associated with the presence of seepage indicators, including microbial mats, bubbles, or chemosynthetic fauna. Within the large pockmark RC1, nearly the entire soft sediment seafloor was intensely bioturbated, except in the vicinity of rock outcrop. The domes of intensely burrowed sediments reached over a meter in diameter and sometimes
- 240 overly neighboring ones. Our observations suggest that burrowing is markedly enhanced by nearby seepage, whereas the burrowed areas largely exceed the vicinity of visible seepage hotspots, often beyond 100 m distances.

#### 3.2 Calliax lobata ghost shrimps inhabit near-seep large burrows

Box core sediment samples within RC1 area (AG16-17BC1, RC20-BC1 and RC20-BC1) -were intensely burrowed (Figure 3). The sediment surface within these samples appeared fluffy with decimeter-scale mounding and multiple holes and scratches (Figure 3A). We observed sporadic black coloration at various horizons, indicative of large, ~0.5 cm wide burrows (Figure 3B-D). Random sieving of leftover sediments in the respective box-corer with a 2 mm mesh revealed that macrofauna was usually scarce, except for 6 live specimens *Calliax lobata* ghost shrimp, measuring ~5 cm long and ~1 cm
wide (Figure 3E). *C. lobata* claws consistently dominated the >2 mm sieved fraction, varying in size up to ~1 cm long and ~5 mm wide (Figure 3F), following previous observations in the same area (Beccari et al., 2020). We, therefore, suggest that *C. lobata* has a key role in near-seep sediment burrowing.

CT scans of AG16-17BC1 and AG16-15BC1 cores showed considerable burrowing (Figure 4). The burrows were imaged as lower density to empty, sub-vertical to sub-horizontal, linear and helicoidal, and generally cylindrical tubes that vary in
their diameter between ~1 to ~10 mm. Larger burrows were usually surrounded by bright halos or rings of relatively dense material. Several burrows had chamber-like regions (Figure 4c). Pervasive tone variations across the CT scans suggest the impact of multiple generations of burrowing that convolute a major part of the cored sections, with more pronounced tone-contrasts imaging the most recent generation of burrowing.



- Figure 3. Burrows of the ghost shrimp *Calliax lobata* near active seepage sites within PalamhimPalmahim Disturbance in the Eastern Mediterranean Sea. A) The seafloor, as exposed in the AC16-17BC1 box-core (RC1 area, 2016), shows burrowing mounds and burrow openings, as well as (black arrows) and scratches (dashed arrows) on the sediment surface. B) Image of RC20-BC1 and RC20-BC2 cores (RC1 area, 2020), showing dark coloration of burrows, unusual for the ambient sediments. C and D) Zooming into the burrows of RC20-BC1 and RC20-BC2 cores showed the presence of large filamentous microbes. E) Live *C. lobata* specimen found in AC16-17BC1. F) A multitude of *C. lobata* claws recovered through sporadic sieving (2 mm mesh) of AC16-17BC1.
  - 11

Segmentation of the CT images captures these more pronounced contrasts, marking the current burrowing activity. Core AG16-17BC1 (47 cm recovery) was composed of homogeneous oxidized clayey sediments with a high amount of shell fragments and pteropod fragments. In this core, the burrow segmentation had an average value of 0.9 %, with a maximum

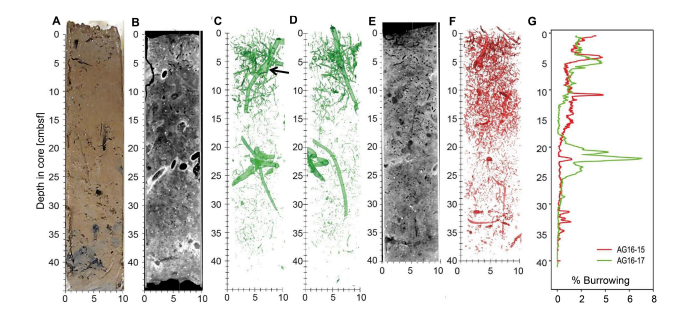
of 7.0 % of the total sediment volume. We noted the highest burrowing at 5 cmbsf (~ 4%) and between 20-22 cmbsf (~ 7%). Between 0 and 15 cmbsf, the sediments had an overall sparse burrowing (BI = 1) with a few discrete traces. Discrete traces were composed of elongated sub-vertical burrows having a length of approximately 3 cm and a diameter of ~1-5 mm, as well as helicoidal burrows similar in size. One large vertical burrow being 12 cm in length and with a diameter of 1 cm is present from the surface to 15 cmbsf. From 15 cmbsf to 25 cmbsf, the overall BI was again close to 1 with few

275 discrete traces composed of very small sub-vertical burrows being 1 cm in length and 1 mm in diameter, as well as two large-sized burrows: 15 cm in length (1 cm diameter) and 6 cm in length (4 cm thickness). From 25 to 41 cmbsf, no clear bioturbation features were quantified (BI = 0).

280

285

Core AG16-15BC1 was composed of oxidized clayey sediment with minor horizontal layering. The surface was highly bioturbated showing a wavy and slightly disturbed surface with small mounds, as well as a high amount of pteropod shells. The average volume % burrowing was 0.7 % with peak burrowing at 4 cmbsf and 10 cmbsf (up to 4%). The box-core sediments (0-40 cmbsf) present overall sparse burrowing (BI = 1). From 0-20 cmbsf, a few discrete traces (BI = 1) were composed of elongated sub-vertical burrows of 5 cm in length and 0.5 cm in diameter and horizontally elongated burrows of 6 cm in length and 0.5 cm in diameter. Also, small curved burrows were present (1 cm in length and 1-2 mm thick). From 20-35 cmbsf, few discrete traces (BI = 0-1) were composed of vertically elongated burrows of 5 cm length and 2-3 mm diameter, as well as small curved burrows of 2 cm length and 1-2 mm thickness with <u>a</u> sub-vertical orientation. The amount of bioturbation decreased towards 35 cmbsf. From 35 – 40 cmbsf, no bioturbation (BI=0) was visible.



12

Figure 4. Burrowing in AG16-15BC1 and AG16-17BC1, as detected by computerized tomography (CT) scans. A) A vertical section through a sub-core of box-core AC16-17BC1 cut and imaged immediately after sampling. B) A vertical section through the 3D CT scan of another sub-core of -AC16-17BC1 (standard medical CT display where higher relative X-ray intensities are darker and low are brighter, depicting lower and higher material densities, respectively). C and D) Two perpendicular 3D displays of segmentation of the relatively vacant (high intensity) burrows, imaged in the CT scan of the same core (in B). E) A vertical section through the 3D CT scan of a sub-core of -AC16-15BC1 (displayed as in B). F) A 3D display of AC16-15BC1 segmentation of the vacant (high intensity) burrows. G) Graphs comparing the volumetric

295 % of relatively vacant burrows in CT scanned sub-cores of AC16-17BC1 (green) and AC16-15BC1 (red), as estimated based on the respective segmentation results (in C-D and F).

Based on the burrow morphology, three distinct burrow traces/ichnofabrics were distinguished: i) large ( $\geq$ 5 mm wide) vertical to sub-horizontal burrows (*Thalassinoides*); ii) smaller (2-5 mm wide) vertical and sub-horizontal burrows forming a dense network (*Ophiomorpha*), iii) small ( $\leq$ 2 mm wide) curved and helicoidal burrows (*Gyrolithes*) (Seilacher, 2007). The presence of significant sub-horizontal segments and chambers in the *Thalassinoides* burrows suggests they were

- 300 The presence of significant sub-horizontal segments and chambers in the *Thalassinoides* burrows suggests they were created by arthropods. First, it is unlikely that sub-horizontal burrows would be created by a bivalve, which would usually produce vertical to oblique burrows (Knaust, 2015; Trueman et al., 1966). Second, arthropods, and in particular decapods, excavate chambers that allow them to turn their direction within their burrows (Coelho et al., 2000; Hembree, 2019; Stamhuis et al., 1997). In turn, the sparse spacing of the burrows also suggests a decapod rather than a wormlike animal,
- 305 which often produces tightly packed burrows (Seilacher, 2007). We suggest that *C. lobata* is the primary inhabitant of the two larger *Thalassinoides* and *Ophiomorpha* burrow morphologies, prominent in AG16-17BC1, based on the marked abundance of these animals and matches in size. It is impossible to determine at this stage the provenance of the smaller *Gyrolithes*-type burrows, possibly worms-like animals.

#### 3.3 Burrowing alters geochemical processes at seep chemotones

- 310 Our data indicates that intense near-seep burrowing can affect geochemical processes via the exchange of solutes across the sediment and the overlying water. The measurements of nutrient concentrations indicate that NO<sub>3</sub><sup>-</sup> depth profiles in cores AG16-15BC1 and AG16-17BC1 show different penetration depths and standing stocks, correlated to the presence of burrows (**Figure 5A**, we assumed negligible nitrite concentrations). In core AG16-15BC1, we observed an increasing trend of NO<sub>3</sub><sup>-</sup> concentrations from ~35 cm upwards. In comparison, core AG16-17BC1 showed significant enrichment above 20
- cm depth. Assuming steady-state conditions, the amount of NO<sub>3</sub><sup>-</sup> in the upper 35 cm of core 15 doubled that of core 17 (~260 and ~105 nmol NO<sub>3</sub><sup>-</sup> depth cm<sup>-2</sup>, respectively). Nitrification driven by ammonia-oxidizing archaea is the key process in background sediments of the abyssal SEMS (Rubin-Blum et al., 2022). However, in the seepage area, this process may be regulated by burrowing, altering the diffusive access of ammonium from the ambient sediment and supplies of oxygen via irrigation by the burrow inhabitant (Kristensen and Kostka, 2005). Thus, macrofaunal burrowing near the sediment surface may aid nitrification (Glud et al., 2009; Mayer et al., 1995). The gradual upward change in the NO<sub>3</sub><sup>-</sup>/PO<sub>4</sub><sup>3-</sup> ratios (Figure 5B) further supports the enhancement of nitrification. Nonetheless, *C. lobata* burrowing may enhance denitrification (removal of nitrate), which was suggested to be enhanced and correlated with the presence of burrowing
  - macrofauna (Nielsen et al., 2004). In core AG16-17, denitrification is likely prominent, given the rapid depletion of nitrate (**Figure 5A**). Our results are thus in line with the previous observations of infauna and  $NO_3^-$  depth distribution at 1450 m

water depth using precise microprofiling, which suggested that burrow irrigation may stimulate both the NO<sub>3</sub><sup>-</sup> production and consumption (Glud et al., 2009).

Previous observations suggest that nitrifying bacteria have a positive correlation to Fe(III) and a negative correlation to dissolved sulfide, which inhibits nitrification (Kristensen and Kostka, 2005). Experimental studies supported field data, confirming that increased amounts of Fe(III) react with sulfide, intensifying the effect of nitrification-denitrification

- 330 processes. In both cores, the depth distribution of porewater  $SO_4^-$  concentrations show values slightly lower than the ambient seawater (31 mM; Herut et al., 2022), suggesting the occurrence of sulfate reduction (**Figure 5C**). Nonetheless, the sulfate levels in core AG16-17BC1 are consistently lower than in AG16-15BC1, indicating that sulfate reduction in this core, where *C. lobata* burrows were found, was enhanced. This is in line with previous studies, showing that sedimentary infaunal burrows and irrigation impact the depth distribution of early diagenetic dissolved reactants/electron acceptors (e.g.  $O_2$ ,  $NO_3^-$
- 335 , and SO<sub>4</sub><sup>2-</sup>) and products (e.g. CO<sub>2</sub>, NH<sub>4</sub><sup>+</sup>, and H<sub>2</sub>S) in porewaters (Aller, 2001; Kristensen and Kostka, 2005; Meile and Cappellen, 2003).

Our data suggests that seep chemotones are potential hotspots of dinitrogen fixation. Although we did not measure nitrogen

fixation rates directly in the burrows, the 0.02±0.004 fmol N gr<sup>-1</sup> d<sup>-1</sup> estimates in the top 3 cm of the chemotone sediments (AG16-17BC1) were significantly higher than the 0.01±0.005 fmol N gr<sup>-1</sup> d<sup>-1</sup> background values (AG16-15BC1, ANOVA
followed by a Fisher LSD multiple comparison post hoc test, p=0.04, Supplementary Figure S1-A). Given that most putative diazotrophs were found in burrow walls, (see below), our results likely underestimate itsN2 fixation rates in the sediments. N2 fixationThese rates were also enhanced in the water column approximately 1 m above the chemoteone sediments (0.04±0.01 nmol N L<sup>-1</sup> d<sup>-1</sup> as opposed to 0.01±0.009 nmol N L<sup>-1</sup> d<sup>-1</sup> at the control site, ANOVA followed by a Fisher LSD multiple comparison post hoc test, p=0.02, Supplementary Figure S1-B), with different diazotrophic populations found at that habitat than those in the sediments (Sisma-Ventura et al., 2022). Altogether, our results suggest that nitrogen cycling at chemotones is substantially altered, either by diazotrophy or denitrification (see above), and that these processes may be boosted by burrowing.

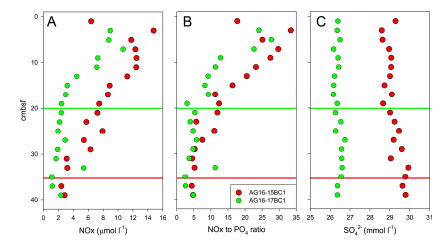
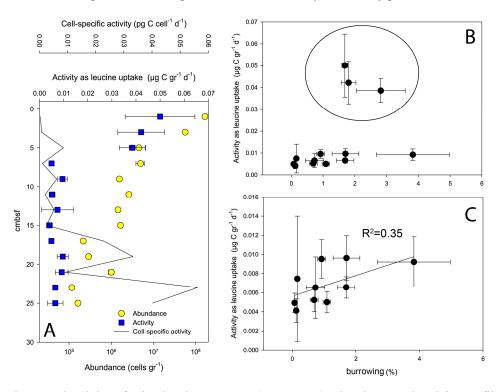


Figure 5: Pore water characteristics in cores AG16-15BC1 and AG16-17BC1: A) Nitrate+nitrite (NOx) concentration. B)
 NOx to phosphate ratio, C) Sulfate concentration. Red and green lines indicate the depths at which NOx concentrations appear to stabilize.

14

#### 3.4 Burrowing may alter microbial abundance and activity at seep chemotones

Following the previous observations of elevated microbial activity in seep chemotone sediments (Sisma-Ventura et al., 2022), our measurements in core AG16-17BC1 suggest a weak, but feasible link between burrowing and microbial activity in these sediments (Figure 6). While the highest abundance and activity values were found in the upper 5 cm, we observed a substantial variation in activity below 5 cm in this core (Figure 6A). In sediments deeper than 5 cm, we observed a weak trend linking burrow intensity and the rate of leucine uptake (Figure 6B, C). We note that activity was not estimated in burrows, but in random subsamples from the respective sections, thus may reflect only partial burrow effects.



**360** Figure 6: Abundance and activity of microbes in core AG16-17BC1: A) Abundance and activity profiles. B) Activity versus burrowing. C) Activity versus burrowing – upper 5 cm removed.

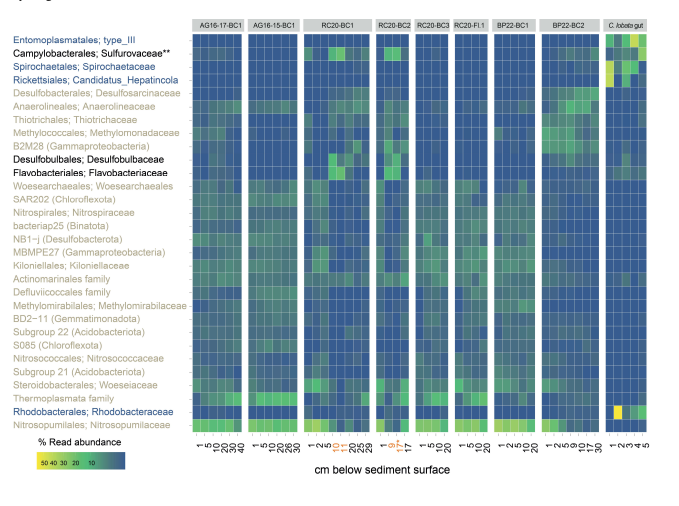
#### 3.5 Seep chemotones host unique microbial communities that are altered by burrowing

Amplicon sequencing of the 16S rRNA gene highlighted the unique microbial communities that existed at seep chemotones (Figure 7, Supplementary Figure S2). These communities markedly differed from those found in seepage hotspots in the same area (Rubin-Blum et al., 2014b), and those of background sediments in the deep SEMS (Rubin-Blum et al., 2022). We identified different levels of seepage-related perturbations in the sediments. RC20-Flare1 and BP22-BC1 samples represent the poorly disturbed, typical background sediments in the SEMS. The most altered sediments were in RC20-BC1, collected within the vicinity of authigenic carbonate in 2020 (Sisma-Ventura et al., 2022) and BP22-BC2, collected 40 meters away from the brine pools in 2022. Ammonia oxidizers such as Nitrosopumilaceae and Nitrospiraceae, as well as
other microbes prominent in background SEMS sediments, such as Thermoplasmata and Binatota lineages (Rubin-Blum et al., 2022), were depleted in these profiles, whereas the putative methane oxidizers, sulfate reducers and chemoautotrophs

were found (Figure 7, Supplementary Figure S2). In particular, the anaerobic methane oxidizers ANME-1 were found at

depths below 25 and 5 cm, in RC20-BC1 and BP22-BC2, respectively. The aerobic Methylomonadaceae methane oxidizers were prominent in the upper sections of these cores. The read abundance of Desulfosarcinae sulfate reducers increased with distance from the sediment surface, and the putative chemoautotrophs Thiotrichales were often found in these samples. This provides evidence for chemosynthetic activity in these chemotone sediments, following previous observations of enhanced microbial activity (Sisma-Ventura et al., 2022).

These results are in line with visual inspection and microscopic observations that revealed: i) black coloration of burrow walls, potentially indicative of precipitation of minerals following sulfate reduction and ii) large, white filaments typical of 380 giant sulfur bacteria inhabiting the burrow inner perimeter (Ionescu and Bizic, 2019) (Figure 3 C-D). According to amplicon sequencing, the key families that were prominent specifically in the burrow subsamples included Desulfobulbaceae, known mainly as sulfur and/or cable bacteria (Trojan et al., 2016), Sulfurovaceae autotrophic sulfur oxidizers, often found in association with marine invertebrates (Bai et al., 2021; Hui et al., 2022) and Flavobacteriaceae versatile degraders of macromolecules, especially carbohydrates (Chen et al., 2021; Gavriilidou et al., 2020). Similar key taxa appear to be associated with bioturbation in other habitats, such as the intertidal flats (Fang et al., 2023). These findings lead to the hypothesis that unique metabolic handoffs take place within the burrows, altering carbon, nitrogen and sulfur cycling.



385

375

Figure 7: Read abundance of the top 30 most prevalent prokaryote taxa at the family level, in Palmahim sediments (see Table 1) and the hindgut of the ghost shrimp *Calliax lobata* (black: enriched in burrows, blue: enriched in *C. lobata*, brown: enriched in sediments). Sediment depths where burrow microbes were collected are highlighted in orange. \* Both samples are from the 16-17 cm section, yet the first one is taken from the burrow and the second from background sediments. \*\* abundant in burrows and associated with *C. lobata* hindgut.

#### 3.6 Metabolic handoffs at burrow walls may fuel dark productivity

395 Based on two DNA samples extracted specifically from burrows in cores RC20-BC1 and RC20-BC2, we curated 58 highquality metagenome-assembled genomes spanning 15 phyla (57 bacteria, 1 archaeon, median quality 92%, median contamination 2%, Figure 8). We observed considerable genotype diversity within Bacteroidota, Desulfobacterota and Proteobacteria phyla (mostly Gammaproteobacteria in the latter). 51 MAGs were found in both samples, although the read abundance patterns often differed. In particular, we found the putative autotrophic sulfur oxidizers. Thiohalobacteraceae 400 (MAGs 20 and 113, reclassified as UBA9214 by GTDB) and Thiomargarita (MAG 115) autotrophic sulfur oxidizers in RC20-BC2, in which the giant bacteria were observed, whereas Thiohalomonadaceae (MAG-83), Sedimenticolaceae (MAGs 22 and 46), and Sulfurimonadaceae (Campylobacterota, MAGs 2 and 80) appear to be the key autotrophs) were found in RC20-BC1. Flavobacteriaceae (mainly CG1-02-35-72 lineage, MAGs 10 and 69) were abundant in both samples, but most dominant in RC20-BC1 (14.4% read abundance). Desulfobacterota comprised Desulfobacterales (MAGs 55e.g. 405 Desulfobacula) and 75), and mainly Desulfobulbales lineages BM004 and BM506, the latter associated with the dissimilatory reduction of perchlorate (Barnum et al., 2018), as well as  $Desulfocapsa_{\overline{2}}$  (4.7% read abundance in RC20-BC1), known to specialize in disproportionating inorganic sulfur compounds (Finster et al., 2013; Ward et al., 2021). BM004 MAG 88 was the dominant one in RC20 BC2 (4.1% read abundance) and Desulfobacula (Desulfobacterales) MAG 55 in RC20-BC1 (4.7% read abundance). Themodesulfovibrionia UBA6902 (Nitrospirota) MAG-11-was also found in 410 RC20-BC2 (2.2% read abundance). In RC20-BC1, Anarerolineales UBA11858 MAG 99 was prominent (MAG 99, 4.5% read abundance). Gemmatimonadetes (MAGs 9 and 89, ~5% read abundance each) were abundant in RC20-BC2. We hypothesized that these dominant lineages can perform an array of metabolic functions, using a range of substrates and electron donors/acceptors, markedly altering the functionality of ecotones compared to the unaltered bathyal sediments in the SEMS.

# 415 <u>3.6 Metabolic handoffs at burrow walls may fuel dark productivity</u>

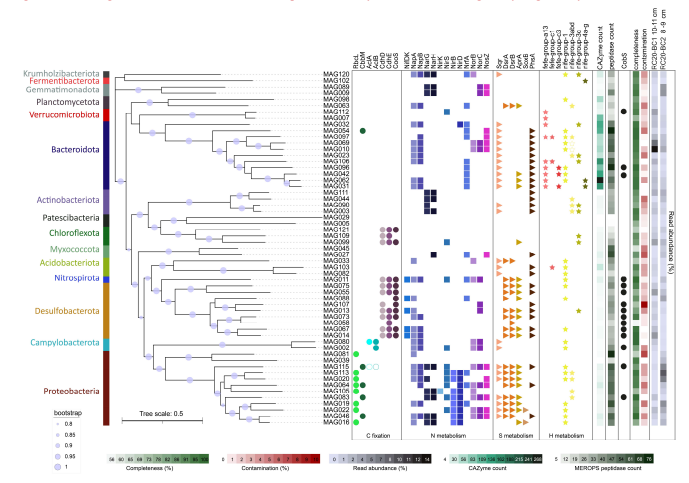
420

In both samples, the chemotrophs<u>Metagenomics suggest</u> that <u>can fix inorganic carbon comprised mostvarious</u> gammaproteobacterial lineages <u>and Campylobacterota</u>. Most of these organisms may fix carbon via the Calvin–Benson–Bassham (CBB) cycle, given the presence of *cbbLS* and/or *cbbM* genes encoding the form I and II ribulose-1,5-bisphosphate carboxylase/oxygenase (RubisCO), respectively (**Figure 8**). Sulfurimonadaceae MAGs encoded the form I ATP citrate lyase, needed for carbon fixation via the reverse tricarboxylic acid cycle (rTCA). In agreement with previous studies (Flood et al., 2016), *Thiomargarita* encoded not only the form II RubisCo but also the form II ATP citrate lyase, indicating the potential to fix carbon via more than one pathway (Rubin-Blum et al., 2019). Partial or complete Wood Ljungdahl (WL) pathway was widespread in Chloroflexota, Desulfobacterota and NitrospirotaSulfurimonadaceae (Campylobacterota) encoded the form I ATP citrate lyase, needed for carbon fixation via the reverse ficarboxylic acid cycle (rTCA).

425 Proteobacteria and Campylobacterota may fuel carbon fixation based on dissimilatory sulfur cycling, encoding proteins

needed for oxidation of reduced sulfur compounds, suggesting the potential for carbon fixation using energy from dihydrogen oxidation and sulfur reduction (Drake, 1994). Alternatively, some of these organisms may use this pathway for energy conservation during acetoclastic growth (Alves et al., 2020; Fang et al., 2022; Santos Correa et al., 2022).

Either oxidation of reduced sulfur, or sulfur/thiosulfate disproportionation may be the key drivers of chemoautotrophy in
 the burrows, especially by Proteobacteria and Campylobacterota. This is indicated by the widespread occurrence of genes encoding proteins needed for dissimilatory sulfur cycling, such as Sqr, DsrAB, AprAB, components of the Sox system, as well as the PhsA, a key protein in thiosulfate disproportionation (Anantharaman et al., 2018; Finster, 2008) (Figure 8). While oxygen is the likely electron acceptor for chemosynthesis, denitrification appears to play a substantial role in energy metabolism. This is These lineages may respire oxygen, or denitrify, as indicated by the frequent occurrence of genes needed for dissimilatory nitrogen cycling, e.g., those encoding Nap and Nar (nitrate reduction), NirK, NirS and NirBD (nitrite reduction, as well as NrfA for dissimilatory nitrite reduction to ammonia, DNRA), NorBC (nitric oxide reduction) and NosZ (nitrous oxide reduction to dinitrogen) (Figure 8). Most gammaproteobacterial MAGs encoded all or part of these genes, indicating that denitrification could be completed either by an individual lineage or synergistically.



440 Figure 8: Phylogeny, taxonomy and key functions of bacteria, based on the reconstruction of metagenome-assembled genomes (MAGs) from *Calliax lobata* burrows. FastTree phylogeny is based on the GTDB-tk marker gene set. Functions were assigned using METABOLIC Hidden Markov Model (HMM) set. Empty circles and stars indicate the functions discovered through manual annotation of functions, using SEED and NCBI BLAST.

In turn, the sulfurSulfur cycling in marine sediments can be driven by fermentation, evolving hydrogen, or other products
 of this process, such as hydrogen, acetate, butyrate and lactate (Campbell et al., 2023; Ruff et al., 2023). Similar to groundwater communities (Ruff et al., 2023), the dominant Bacteroidota are the likely key fermenters of macromolecules appear to bein the highly abundant Bacteroidotaburrows (Figure 8). These organisms are ubiquitous degraders of macromolecules such as complex glycans (Fernández-Gómez et al., 2013; Martens et al., 2008, 2009; Ndeh and Gilbert, 2018). They encode carbohydrate-active enzymes (CAZymes) that are often organized in polysaccharide utilization loci (PULs), with typical SusCD transporters (Grondin et al., 2017; Martens et al., 2009; Reeves et al., 1996). We found multiple SusCD in-copies in diverse Bacteroidota MAGs, but also discovered a multitude of encoded CAZymes, as well as

peptidases, highlighting the potential of these bacteria to use a broad scale of macromolecules. (Figure 8).

Bacteroidota may ferment-these macromolecules to produce various substrates, most importantly, hydrogen, as suggested by the presence of Fe-Fe hydrogen evolving hydrogenases in Bacteroidales (MAGs 31, 42, 62, 96 and 106-(, Figure 8).
The highly abundant Flavobacteriales MAGs 10 and 69 did not encode the Fe-Fe or the canonical type 4 H<sub>2</sub> evolving hydrogenases. Instead, they encoded hydrogenase I (*hyaABCDEF*), and, most importantly, an adjacent gene cluster encoding an aberrant putative HoxHYFU NAD-dependant hydrogenase, apparently widespread in Bacteroidota (based on multiple NCBI nr BLAST hits of HoxH protein in this clade, data not shown). These soluble, oxygen-tolerant, bidirectional hydrogenases described in *Ralstonia eutropha* H16 (Burgdorf et al., 2006) and cyanobacteria (Appel et al., 2000; Gutekunst et al., 2014), can promote hydrogen evolution in bacteria that ferment sugars (Cheng et al., 2019). We also identified *fdhAB*

genes encoding the NAD-dependent formate dehydrogenase in these Flavobacteriales MAGs, hinting at the potential coupling of formate oxidation and hydrogen evolution. Other genes involved in fermentation, including those needed for acetogenesis and acetate conversion to acetyl-CoA (*ack, pta* and *acs*), were found in all the Bacteriodota MAGs. Thus, most Bacteriodota that we discovered in burrows, including the dominant CG1-02-35-72 Flavobacteriales lineages, are
 likely the key fermenters that fuel the cycling of organic matter at the burrow walls.

Fermentation products can provide electron donors Desulfobacterota and Nitrospirota sulfate reducers, which encode the Dsr/Apr systems, highlighting the key role of sulfur cycling in these communities (**Figure 8**). Hydrogen can directly fuel chemosynthesis, for example, by Campylobacterota, which encode uptake NiFe hydrogenases and often thrive based on hydrogen in hydrothermal vents (Molari et al., 2023). Some Gammaproteobacteria, including the symbiotic Sedimenticolaceae and Thioglobaceae, also use hydrogen as an energy source in chemosynthetic habitats (Petersen et al., 2011). In line with these observations, we identified uptake NiFe hydrogenases in all the diverse Sedimenticolaceae MAGs (16, 19, 22, 46 and 155), as well as in Thiohalobacteraceae MAGs 20 and 113, hinting that hydrogen is a potential energy source for burrow autotrophs.

470

In turn, fermentation products can provide electron donors to the Desulfobacterota and Nitrospirota sulfate reducers that likely drive sulfate depletion in these sediments (Figure 5). Partial or complete Wood-Ljungdahl (WL) pathway was widespread in Desulfobacterota and Nitrospirota, as well as Chloroflexota, suggesting the potential for carbon fixation using energy from dihydrogen oxidation and sulfur reduction (Drake, 1994). Alternatively, some of these organisms may use this pathway for energy conservation during acetoclastic growth (Alves et al., 2020; Fang et al., 2022; Santos Correa et al., 2022). This is in line with the observation of hydrogenogenic fermentation and hydrogenotrophic sulfate reduction that co-occur in anoxic sandy sediments (Chen et al., 2021). These interactions between Bacteroidota and Desulfobacterota are not limited to marine sediments, but for example, facilitate dihydrogen turnover in the human gut (Gibson et al., 1993; Thomas et al., 2011; Wolf et al., 2016). The resulting sulfide fluxes may further fuel the productivity of sulfur-oxidizing bacteria.

Hydrogen can also directly fuel chemosynthesis, for example, by Campylobacterota, which encode uptake NiFe hydrogenases, In summary, our-and often thrive based on hydrogen in hydrothermal vents (Molari et al., 2023). Some Gammaproteobacteria, including the symbiotic Sedimenticolaceae and Thioglobaceae, also use hydrogen as an energy source in chemosynthetic habitats (Petersen et al., 2011). In line with these observations, we identified uptake NiFe hydrogenases in all the diverse Sedimenticolaceae MAGs (16, 19, 22, 46 and 155), as well as in Thiohalobacteraceae MAGs 20 and 113, hinting that hydrogen is a potential energy source for burrow autotrophs.

490 Our results suggest the presence of productive burrow communities, that cycle carbon, sulfur and nitrogen, potentially through the turnover of hydrogen or other fermentation products. Examples of similar communities, comprising Bacteroidota degraders of macromolecules, sulfate reducers (Desulfobacterota and others), and chemotrophs (mainly Campylobacteria and Gammaproteobacteria) are common among a wide range of biomes, including burrowed sediments (Fang et al., 2023), epibionts of invertebrates in chemosynthetic habitats (Bai et al., 2021; Xu et al., 2022) and 495 endosymbiotic communities in Idas mussels (Zvi-Kedem et al., 2023). The interactions between these organisms are not limited to the exchange of key metabolites (Zoccarato et al., 2022). One example of synergy in this system is the potential exchange of cobalamin (vitamin B12) among the community members. Our data suggest that the taxonomic diversity of cobalamin producers in the burrow walls is limited, primarily to Desulfobacterota and some thiotrophs (Thiomargarita MAG 115 and Thiohalomonadaceae MAG 43), as well as several others (Figure 8). On the other hand, the keycommon 500 Flavobacteriaceae (MAGs 10 and 69) do not produce cobalamin but may take it up, as they encode the outer membrane vitamin B12 receptor BtuB and BtuFCD components of the vitamin B12 ABC transporter, providing this cofactor to B12dependent enzymes such as methylmalonyl-CoA mutase (EC 5.4.99.2) and class II ribonucleotide reductase (EC 1.17.4.1). The resilience of these communities may depend on redundancy in the production of such shared commodities, for example, cobalamin production by diverse Desulfobacterota genotypes.

# 505 3.7 Calliax lobata hosts specific gut microflora but may ingest burrow microbes

The amplicon sequencing of DNA extracted from hindgut or midgut in several *C. lobata* specimens identified specialized microflora, but also microbes found in the burrow walls (**Figure 7, Supplementary Figure 1<u>S2</u>**). The four key lineages associated with the gut sections included Entoplasmatales, Spirochaetales, Rickettsiales and Rhodobacterales. For example, we found abundant *Candidadus* Hepatincola (Rickettsiales), a common nutrient-scavenging bacterium in the gut of crustaceans, and isopods in particular, from terrestrial and aquatic environments (Dittmer et al., 2023). We found that someSome lineages, which were prominent in burrow walls, may occur in *C. lobata* gut. For: for example, we found trace quantities of *Desulfobulbus* and *Desulfocapsa* in the gut of some *C. lobata* specimens (**Supplementary Figure S2**). Campylobacterota were abundant in the gut and the burrow walls, whereas *Sulfurospirillum* species were associated only with the gut, and *Sulfurovum* were found in both the burrows and the gut, indicating that these lineages occupy distinct

515 niches. Altogether, these findings hint that *C. lobata* may be feeding in the burrows, providing evidence for the 'gardening' hypothesis.

# 4. Conclusions

Our results indicate that SEMS seep chemotones are characterized by extensive burrowing by ghost shrimp *C. lobata.* The burrowed sediments can span tens to hundreds of meters away from the seepage hotspots, extending the seep area of influence by at least an order of magnitude. The high microbial activity in these burrowed sediments alters the biogeochemical cycles at the sediment-water interface of the oligotrophic SEMS (Sisma-Ventura et al., 2022), and our new results indicate that unique microbial communities at the burrow walls can contribute new functions needed to sustain this activity. These functions include the fermentation of complex macromolecules by Bacterioida, coupling oxidation of these fermentation products to sulfate reduction by Desulfobacterota and Nitrospirota, and chemotrophy by Campylobacterota and chemosynthetic Gammaproteobacteria, including the giant sulfur bacteria *Thiomargarita*. The enhancement of nitrogen cycling at seep chemotones may play a particularly important role in the oligotrophic SEMS. The discovery of these functions underscores the underestimated role of seep chemotones in deep-sea biogeochemistry.

The functionality of the burrow communities may depend on the gardening or grooming of burrow walls by *C. lobata*. Our observations indicate that this behavior is ubiquitous in ghost shrimps, altering benthic—pelagic nutrient exchange in both shallow and deep habitats (Laverock et al., 2010; Papaspyrou et al., 2005). While the burrow wall communities are most likely fueled by *C. lobata* secretions, producing an organic matter sink ('detrital trap') in the burrow (Papaspyrou et al., 2005), it remains to be tested if *C. lobata* burrows can reach the sulfate-methane transition zone (e.g., the deeper sections of the sampled sediments, where ANME-1 were found, Figure 3), providing substrates for chemosynthesis. It is also intriguing to investigate the evolutionary aspects of crustacean associations with microbes in chemosynthetic habitats, given the similarity between the burrow and ectosymbiont communities (Bai et al., 2021; Cambon-Bonavita et al., 2021; Xu et al., 2022).

#### Author contributions

MR-B, YM, AF, OB and BH conceived this study and acquired funding. YM and OE performed mapping and led
 ROV/AUV work. AF and OB performed CT scans. GS-V and BH analyzed the chemical parameters. ER estimated microbial abundance and productivity. ZH and YY performed molecular work. MR-B performed molecular and bioinformatics analyses. MR-B wrote the paper with the contributions of all co-authors.

#### Acknowledgments

This research used samples and data provided by the E/V Nautilus Exploration Program, expedition NA019. The authors thank all individuals who helped during the expeditions, including onboard technical and scientific personnel of E/V Nautilus, SEMSEEPS 2016 and IOLR-University of Haifa cruises, and the captains and crews of E/V Nautilus, IOLR R/V Bat-Galim and HCMR R/V Aegaeo. We thank the University of Haifa Hatter Department of Marine Technologies team, Ben Herzberg and Samuel Cohen, who operated the Yona ROV. We thank Drs. Moshe Tom and Hadas Lubinevsky (IOLR) for their assistance in sample collection.

# 550 Funding

This study is funded by the Israeli Science Foundation (ISF) grants 913/19 and 1359/23, the Israel Ministry of Energy grants 221-17-002 and 221-17-004, the Israel Ministry of Science and Technology grant 3-17403, and the Mediterranean Sea Research Center of Israel (MERCI). The EUROFLEETS2 SEMSEEP cruise was funded by the European Union FP7 Programme under grant agreement no. 312762 and farther analysis was funded by the Swiss National Science Foundation grantGrant Ref. 200021\_175587. This work was partly supported by the National Monitoring Program of Israel's

555 <u>grantGrant</u> Ref. 200021\_175587. This work was partly supported by the National Monitoring Mediterranean waters and the Charney School of Marine Sciences (CSMS), University of Haifa.

## **Declaration of competing interest**

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

# 560 Data availability statement

The raw metagenomic reads and metagenome-assembled genomes are available on NCBI as BioProject PRJNA1072319.

## 5. Bibliography

Abed-Navandi, D., Koller, H. and Dworschak, P. C.: Nutritional ecology of thalassinidean shrimps constructing burrows
 with debris chambers: The distribution and use of macronutrients and micronutrients, Mar. Biol. Res., 1(3), 202–215, doi:10.1080/17451000510019123, 2005.

Aller, R. C.: Transport and reaction in the bioirrigated zone, in The Benthic Boundary Layer: Transport Processes and Biogeochemistry, edited by B. P. Boudreau and B. B. Jorgensen, pp. 269–301, Oxford University Press., 2001.

Alves, J. I., Visser, M., Arantes, A. L., Nijsse, B., Plugge, C. M., Alves, M. M., Stams, A. J. M. and Sousa, D. Z.: Effect of sulfate on carbon monoxide conversion by a thermophilic syngas-fermenting culture dominated by a *Desulfofundulus* species, Front. Microbiol., 11(November), 588468, doi:10.3389/fmicb.2020.588468, 2020.

Amon, D. J., Gobin, J., Van Dover, C. L., Levin, L. A., Marsh, L. and Raineault, N. A.: Characterization of methane-seep communities in a deep-sea area designated for oil and natural gas exploitation off Ttrinidad and Tobago, Front. Mar. Sci., 4(October), 342, doi:10.3389/fmars.2017.00342, 2017.

575 Anantharaman, K., Hausmann, B., Jungbluth, S. P., Kantor, R. S., Lavy, A., Warren, L. A., Rappé, M. S., Pester, M., Loy, A., Thomas, B. C. and Banfield, J. F.: Expanded diversity of microbial groups that shape the dissimilatory sulfur cycle, ISME J., 12(7), 1715–1728, doi:10.1038/s41396-018-0078-0, 2018.

Andersen, K. S., Kirkegaard, R. H., Karst, S. M. and Albertsen, M.: ampvis2: an R package to analyse and visualise 16S rRNA amplicon data, bioRxiv, 299537, doi:10.1101/299537, 2018.

580 Appel, J., Phunpruch, S., Steinmüller, K. and Schulz, R.: The bidirectional hydrogenase of Synechocystis sp. PCC 6803 works as an electron valve during photosynthesis, Arch. Microbiol., 173(5–6), 333–338, doi:10.1007/s002030000139, 2000.

Apprill, A., Mcnally, S., Parsons, R. and Weber, L.: Minor revision to V4 region SSU rRNA 806R gene primer greatly increases detection of SAR11 bacterioplankton, Aquat. Microb. Ecol., 75(2), 129–137, doi:10.3354/ame01753, 2015.

585 Ashford, O. S., Guan, S., Capone, D., Rigney, K., Rowley, K., Orphan, V., Mullin, S. W., Dawson, K. S., Cortés, J., Rouse, G. W., Mendoza, G. F., Lee, R. W., Cordes, E. E. and Levin, L. A.: A chemosynthetic ecotone—"chemotone"—in the sediments surrounding deep-sea methane seeps, Limnol. Oceanogr., 66(5), 1687–1702, doi:10.1002/lno.11713, 2021.

590

600

605

Astall, C. M., Taylor, A. C. and Atkinson, R. J. A.: Behavioural and physiological implications of a burrow-dwelling lifestyle for two species of upogebiid mud-shrimp (Crustacea: Thalassinidea), Estuar. Coast. Shelf Sci., 44(2), 155–168, doi:10.1006/ecss.1996.0207, 1997.

Åström, E. K. L., Bluhm, B. A. and Rasmussen, T. L.: Chemosynthetic and photosynthetic trophic support from cold seeps in Arctic benthic communities, Front. Mar. Sci., 9(August), 1–19, doi:10.3389/fmars.2022.910558, 2022.

Bai, S., Xu, H. and Peng, X.: Microbial communities of the hydrothermal scaly-foot snails from kairei and longqi vent fields, Front. Mar. Sci., 8(October), 764000, doi:10.3389/fmars.2021.764000, 2021.

595 Barnum, T. P., Figueroa, I. A., Carlström, C. I., Lucas, L. N., Engelbrektson, A. L. and Coates, J. D.: Genome-resolved metagenomics identifies genetic mobility, metabolic interactions, and unexpected diversity in perchlorate-reducing communities, ISME J., 12(6), 1568–1581, doi:10.1038/s41396-018-0081-5, 2018.

Basso, D., Beccari, V., Almogi-Labin, A., Hyams-Kaphzan, O., Weissman, A., Makovsky, Y., Rüggeberg, A. and Spezzaferri, S.: Macro- and micro-fauna from cold seeps in the Palmahim Disturbance (Israeli off-shore), with description of *Waisiuconcha corsellii* n.sp. (Bivalvia, Vesicomyidae), Deep. Res. Part II Top. Stud. Oceanogr., 171(January), 104723, doi:10.1016/j.dsr2.2019.104723, 2020.

Bayon, G., Loncke, L., Dupré, S., Caprais, J.-C., Ducassou, E., Duperron, S., Etoubleau, J., Foucher, J.-P. J.-P., Fouquet, Y., Gontharet, S., Henderson, G. M., Huguen, C., Klaucke, I., Mascle, J., Migeon, S., Olu-Le Roy, K., Ondreas, H., Pierre, C., Sibuet, M., Stadnitskaia, A. and Woodside, J.: Multi-disciplinary investigation of fluid seepage on an unstable margin : The case of the Central Nile deep sea fan, Mar. Geol., 261(1–4), 92–104, doi:10.1016/j.margeo.2008.10.008,

2009.

Beccari, V., Basso, D., Spezzaferri, S., Rüggeberg, A., Neuman, A. and Makovsky, Y.: Preliminary video-spatial analysis of cold seep bivalve beds at the base of the continental slope of Israel (Palmahim Disturbance), Deep. Res. Part II Top. Stud. Oceanogr., 171, 104664, doi:10.1016/j.dsr2.2019.104664, 2020.

610 Blouet, J. P., Imbert, P., Ho, S., Wetzel, A. and Foubert, A.: What makes seep carbonates ignore self-sealing and grow vertically: The role of burrowing decapod crustaceans, Solid Earth, 12(10), 2439–2466, doi:10.5194/se-12-2439-2021, 2021.

Boetius, A. and Wenzhöfer, F.: Seafloor oxygen consumption fuelled by methane from cold seeps, Nat. Geosci., 6(9), 725–734, doi:10.1038/ngeo1926, 2013.

- 615 Bolyen, E., Rideout, J. R., Dillon, M. R., Bokulich, N. A., Abnet, C. C., Al-Ghalith, G. A., Alexander, H., Alm, E. J., Arumugam, M., Asnicar, F., Bai, Y., Bisanz, J. E., Bittinger, K., Brejnrod, A., Brislawn, C. J., Brown, C. T., Callahan, B. J., Caraballo-Rodríguez, A. M., Chase, J., Cope, E. K., Da Silva, R., Diener, C., Dorrestein, P. C., Douglas, G. M., Durall, D. M., Duvallet, C., Edwardson, C. F., Ernst, M., Estaki, M., Fouquier, J., Gauglitz, J. M., Gibbons, S. M., Gibson, D. L., Gonzalez, A., Gorlick, K., Guo, J., Hillmann, B., Holmes, S., Holste, H., Huttenhower, C., Huttley, G. A., Janssen, S.,
- 620 Jarmusch, A. K., Jiang, L., Kaehler, B. D., Kang, K. Bin, Keefe, C. R., Keim, P., Kelley, S. T., Knights, D., Koester, I., Kosciolek, T., Kreps, J., Langille, M. G. I., Lee, J., Ley, R., Liu, Y.-X., Loftfield, E., Lozupone, C., Maher, M., Marotz, C., Martin, B. D., McDonald, D., McIver, L. J., Melnik, A. V., Metcalf, J. L., Morgan, S. C., Morton, J. T., Naimey, A. T., Navas-Molina, J. A., Nothias, L. F., Orchanian, S. B., Pearson, T., Peoples, S. L., Petras, D., Preuss, M. L., Pruesse, E., Rasmussen, L. B., Rivers, A., Robeson, M. S., Rosenthal, P., Segata, N., Shaffer, M., Shiffer, A., Sinha, R., Song, S.
- 625 J., Spear, J. R., Swafford, A. D., Thompson, L. R., Torres, P. J., Trinh, P., Tripathi, A., Turnbaugh, P. J., Ul-Hasan, S., van der Hooft, J. J. J., Vargas, F., Vázquez-Baeza, Y., Vogtmann, E., von Hippel, M., et al.: Reproducible, interactive, scalable and extensible microbiome data science using QIIME 2, Nat. Biotechnol., 37(8), 852–857, doi:10.1038/s41587-019-0209-9, 2019.
- Burgdorf, T., Lenz, O., Buhrke, T., Van Der Linden, E., Jones, A. K., Albracht, S. P. J. and Friedrich, B.: [NiFe]hydrogenases of *Ralstonia eutropha* H16: Modular enzymes for oxygen-tolerant biological hydrogen oxidation, J. Mol. Microbiol. Biotechnol., 10(2–4), 181–196, doi:10.1159/000091564, 2006.

Cambon-Bonavita, M.-A., Aubé, J., Cueff-Gauchard, V. and Reveillaud, J.: Niche partitioning in the Rimicaris exoculata

holobiont: the case of the first symbiotic Zetaproteobacteria, Microbiome, 9(1), 87, doi:10.1186/s40168-021-01045-6, 2021.

635 Campbell, A., Gdanetz, K., Schmidt, A. W. and Schmidt, T. M.: H2 generated by fermentation in the human gut microbiome influences metabolism and competitive fitness of gut butyrate producers, Microbiome, 11(1), 133, doi:10.1186/s40168-023-01565-3, 2023.

Chaumeil, P. A., Mussig, A. J., Hugenholtz, P. and Parks, D. H.: GTDB-Tk: A toolkit to classify genomes with the genome taxonomy database, Bioinformatics, 36(6), 1925–1927, doi:10.1093/bioinformatics/btz848, 2020.

640 Chen, Y.-J. J., Leung, P. M., Wood, J. L., Bay, S. K., Hugenholtz, P., Kessler, A. J., Shelley, G., Waite, D. W., Franks, A. E., Cook, P. L. M. M. and Greening, C.: Metabolic flexibility allows bacterial habitat generalists to become dominant in a frequently disturbed ecosystem, ISME J., 15(10), 2986–3004, doi:10.1038/s41396-021-00988-w, 2021.

Cheng, J., Li, H., Zhang, J., Ding, L., Ye, Q. and Lin, R.: Enhanced dark hydrogen fermentation of Enterobacter aerogenes/HoxEFUYH with carbon cloth, Int. J. Hydrogen Energy, 44(7), 3560–3568, doi:10.1016/j.ijhydene.2018.12.080, 2019.

645

675

Chklovski, A., Parks, D. H., Woodcroft, B. J., Tyson, G. W., Chkolovski, A., Parks, D. H., Woodcroft, B. J., Tyson, G. W., Chklovski, A., Parks, D. H., Woodcroft, B. J. and Tyson, G. W.: CheckM2: a rapid, scalable and accurate tool for assessing microbial genome quality using machine learning, Nat. Methods, 20(8), 1203–1212, doi:10.1038/s41592-023-01940-w, 2023.

650 Coelho, V., Cooper, R. and de Almeida Rodrigues, S.: Burrow morphology and behavior of the mud shrimp *Upogebia omissa* (Decapoda:Thalassinidea:Upogebiidae), Mar. Ecol. Prog. Ser., 200, 229–240, doi:10.3354/meps200229, 2000.

Coleman, D. and Ballard, R.: A highly concentrated region of cold hydrocarbon seeps in the southeastern Mediterranean Sea, Geo-Marine Lett., 21(3), 162–167, doi:10.1007/s003670100079, 2001.

Coleman, D. F., Austin Jr., J. A., Ben-Avraham, Z., Makovsky, Y. and Tchernov, D.: Seafloor pockmarks, deepwater
 corals, and cold seeps along the continental margin of Israel, Oceanography, 25(1, Supplement), 40–41, 2012.

Coleman, D. F. D., Austin Jr., J. A., Ben-Avraham, Z., Ballard, R. D., Austin, J. A., Ben-Avraham, Z. and Ballard, R. D.: Exploring the continental margin of Israel: "Telepresence" at work, Eos Trans. AGU, 92(10), 81–82, doi:10.1029/2011EO100002, 2011.

Dekas, A. E., Chadwick, G. L., Bowles, M. W., Joye, S. B. and Orphan, V. J.: Spatial distribution of nitrogen fixation in methane seep sediment and the role of the ANME archaea, Environ. Microbiol., 16(10), 3012–3029, doi:10.1111/1462-2920.12247, 2014.

Dekas, A. E., Fike, D. A., Chadwick, G. L., Green-Saxena, A., Fortney, J., Connon, S. A., Dawson, K. S. and Orphan, V. J.: Widespread nitrogen fixation in sediments from diverse deep-sea sites of elevated carbon loading, Environ. Microbiol., 20(12), 4281–4296, doi:10.1111/1462-2920.14342, 2018.

665 Dittmer, J., Bredon, M., Moumen, B., Raimond, M., Grève, P. and Bouchon, D.: The terrestrial isopod symbiont *Candidatus* Hepatincola porcellionum' is a potential nutrient scavenger related to Holosporales symbionts of protists, ISME Commun., 3(1), 18, doi:10.1038/s43705-023-00224-w, 2023.

Drake, H. L.: Acetogenesis, Acetogenic Bacteria, and the Acetyl-CoA "Wood/Ljungdahl" Pathway: Past and Current Perspectives, in Acetogenesis, pp. 3–60, Springer US, Boston, MA., 1994.

670 Dworschak, P. C., Koller, H. and Abed-Navandi, D.: Burrow structure, burrowing and feeding behaviour of Corallianassa longiventris and *Pestarella tyrrhena* (Crustacea, Thalassinidea, Callianassidae), Mar. Biol., 148(6), 1369–1382, doi:10.1007/s00227-005-0161-8, 2006.

Fang, J., Jiang, W., Meng, S., He, W., Wang, G., Guo, E. and Yan, Y.: Polychaete bioturbation alters the taxonomic structure, co-occurrence network, and functional groups of bacterial communities in the intertidal flat, Microb. Ecol., 86(1), 112–126, doi:10.1007/s00248-022-02036-2, 2023.

Fang, Y., Liu, J., Yang, J., Wu, G., Hua, Z., Dong, H., Hedlund, B. P., Baker, B. J. and Jiang, H.: Compositional and metabolic responses of autotrophic microbial community to salinity in lacustrine environments, edited by R.

Mackelprang, mSystems, 7(4), e0033522, doi:10.1128/msystems.00335-22, 2022.

Fernández-Gómez, B., Richter, M., Schüler, M., Pinhassi, J., Acinas, S. G., González, J. M. and Pedrós-Alió, C.: Ecology of marine bacteroidetes: A comparative genomics approach, ISME J., 7(5), 1026–1037, doi:10.1038/ismej.2012.169, 2013.

Finster, K.: Microbiological disproportionation of inorganic sulfur compounds, J. Sulfur Chem., 29(3–4), 281–292, doi:10.1080/17415990802105770, 2008.

Finster, K. W., Kjeldsen, K. U., Kube, M., Reinhardt, R., Mussmann, M., Amann, R. and Schreiber, L.: Complete
genome sequence of desulfocapsa sulfexigens, a marine deltaproteobacterium specialized in disproportionating inorganic sulfur compounds, Stand. Genomic Sci., 8(1), 58–68, doi:10.4056/sigs.3777412, 2013.

Flood, B. E., Fliss, P., Jones, D. S., Dick, G. J., Jain, S., Kaster, A.-K., Winkel, M., Mußmann, M. and Bailey, J.: Singlecell (meta-)genomics of a dimorphic *Candidatus* Thiomargarita nelsonii reveals genomic plasticity, Front. Microbiol., 7(May), 603, doi:10.3389/fmicb.2016.00603, 2016.

690 Gadol, O., Tibor, G., ten Brink, U., Hall, J. K., Groves-Gidney, G., Bar-Am, G., Hübscher, C. and Makovsky, Y.: Semiautomated bathymetric spectral decomposition delineates the impact of mass wasting on the morphological evolution of the continental slope, offshore Israel, Basin Res., 32(5), 1166–1193, doi:10.1111/bre.12420, 2020.

Garfunkel, Z., Arad, A. and Almagor, G.: The Palmahim disturbance and its regional setting, Geol. Surv. Isr. Bull., 56, 1979.

695 Gavriilidou, A., Gutleben, J., Versluis, D., Forgiarini, F., Van Passel, M. W. J., Ingham, C. J., Smidt, H. and Sipkema, D.: Comparative genomic analysis of Flavobacteriaceae: Insights into carbohydrate metabolism, gliding motility and secondary metabolite biosynthesis, BMC Genomics, 21(1), 569, doi:10.1186/s12864-020-06971-7, 2020.

Gibson, G. R., Macfarlane, G. T. and Cummings, J. H.: Sulphate reducing bacteria and hydrogen metabolism in the human large intestine., Gut, 34(4), 437–9, doi:10.1136/gut.34.4.437, 1993.

Gilbertson, W. W., Solan, M. and Prosser, J. I.: Differential effects of microorganism-invertebrate interactions on benthic nitrogen cycling, FEMS Microbiol. Ecol., 82(1), 11–22, doi:10.1111/j.1574-6941.2012.01400.x, 2012.

Glöckner, F. O., Yilmaz, P., Quast, C., Gerken, J., Beccati, A., Ciuprina, A., Bruns, G., Yarza, P., Peplies, J., Westram, R. and Ludwig, W.: 25 years of serving the community with ribosomal RNA gene reference databases and tools, J. Biotechnol., 261(June), 169–176, doi:10.1016/j.jbiotec.2017.06.1198, 2017.

705 Glud, R. N., Thamdrup, B., Stahl, H., Wenzhoefer, F., Glud, A., Nomaki, H., Oguri, K., Revsbech, N. P. and Kitazatoe, H.: Nitrogen cycling in a deep ocean margin sediment (Sagami Bay, Japan), Limnol. Oceanogr., 54(3), 723–734, doi:10.4319/lo.2009.54.3.0723, 2009.

Grondin, J. M., Tamura, K., Déjean, G., Abbott, D. W. and Brumer, H.: Polysaccharide utilization loci: Fueling microbial communities, J. Bacteriol., 199(15), doi:10.1128/JB.00860-16, 2017.

710 Gutekunst, K., Chen, X., Schreiber, K., Kaspar, U., Makam, S. and Appel, J.: The bidirectional NiFe-hydrogenase in synechocystis sp. PCC 6803 Is reduced by flavodoxin and ferredoxin and is essential under mixotrophic, nitrate-limiting conditions, J. Biol. Chem., 289(4), 1930–1937, doi:10.1074/jbc.M113.526376, 2014.

Gvirtzman, Z., Reshef, M., Buch-Leviatan, O., Groves-Gidney, G., Karcz, Z., Makovsky, Y. and Ben-Avraham, Z.: Bathymetry of the Levant basin: interaction of salt-tectonics and surficial mass movements, Mar. Geol., 360, 25–39, doi:10.1016/j.margeo.2014.12.001, 2015.

715

720

Hembree, D. I.: Burrows and ichnofabric produced by centipedes: modern and ancient examples, Palaios, 34(10), 468–489, doi:10.2110/palo.2019.059, 2019.

Herut, B., Rubin-Blum, M., Sisma-Ventura, G., Jacobson, Y., Bialik, O. M., Ozer, T., Lawal, M. A., Giladi, A., Kanari, M., Antler, G. and Makovsky, Y.: Discovery and chemical composition of the eastmost deep-sea anoxic brine pools in the Eastern Mediterranean Sea, Front. Mar. Sci., 9, 1040681, doi:10.3389/fmars.2022.1040681, 2022.

Hui, M., Wang, A., Cheng, J. and Sha, Z.: Full-length 16S rRNA amplicon sequencing reveals the variation of epibiotic

microbiota associated with two shrimp species of Alvinocarididae: possibly co-determined by environmental heterogeneity and specific recognition of hosts, PeerJ, 10, e13758, doi:10.7717/peerj.13758, 2022.

Ionescu, D. and Bizic, M.: Giant Bacteria, in Encyclopedia of Life Sciences, p. DOI:
10.1002/9780470015902.a0020371.pub2, John Wiley & Sons, Ltd, Chichester., 2019.

730

735

760

Joye, S. B.: The geology and biogeochemistry of hydrocarbon seeps, Annu. Rev. Earth Planet. Sci., 48, 205–231, doi:10.1146/annurev-earth-063016-020052, 2020.

Jumars, P. A., Dorgan, K. M., Mayer, L. M., Boudreau, B. P. and Johnson, B. D.: Material constraints on infaunal lifestyles. May the persistent and strong forces be with you., Trace Foss. Concepts, Probl. Prospect., 442–457, doi:10.1016/B978-044452949-7/50152-2, 2007.

Kang, D. D., Froula, J., Egan, R. and Wang, Z.: MetaBAT, an efficient tool for accurately reconstructing single genomes from complex microbial communities, PeerJ, 3, e1165, doi:10.7717/peerj.1165, 2015.

Kieser, S., Brown, J., Zdobnov, E. M., Trajkovski, M. and McCue, L. A.: ATLAS: a Snakemake workflow for assembly, annotation, and genomic binning of metagenome sequence data, BMC Bioinformatics, 21(1), 257, doi:10.1186/s12859-020-03585-4, 2020.

Knaust, D.: Siphonichnidae (new ichnofamily) attributed to the burrowing activity of bivalves: Ichnotaxonomy, behaviour and palaeoenvironmental implications, Earth-Science Rev., 150, 497–519, doi:10.1016/j.earscirev.2015.07.014, 2015.

Knittel, K. and Boetius, A.: Anaerobic oxidation of methane: progress with an unknown process., Annu. Rev. Microbiol., 63(1), 311–34, doi:10.1146/annurev.micro.61.080706.093130, 2009.

Kormas, K. A. and Meziti, A.: The microbial communities of the East Mediterranean Sea mud volcanoes and pockmarks, in Marine Hydrocarbon Seeps: Microbiology and Biogeochemistry of a Global Marine Habitat, edited by A. P. Teske and V. Carvalho, pp. 143–148, Springer Nature Switzer-land., 2020.

Kristensen, E. and Kostka, J. E.: Macrofaunal burrows and irrigation in marine sediment: Microbiological and
 biogeochemical interactions, in Interactions Between Macro- and Microorganisms in Marine Sediments, edited by E.
 Kristensen, R. R. Haese, and J. E. Kostka, pp. 125–157, AGU, Washington, D.C., US, D.C., US., 2005.

Laverock, B., Smith, C. J., Tait, K., Osborn, A. M., Widdicombe, S. and Gilbert, J. A.: Bioturbating shrimp alter the structure and diversity of bacterial communities in coastal marine sediments, ISME J., 4(12), 1531–1544, doi:10.1038/ismej.2010.86, 2010.

750 Laverock, B., Gilbert, J. A., Tait, K., Osborn, A. M. and Widdicombe, S.: Bioturbation: Impact on the marine nitrogen cycle, Biochem. Soc. Trans., 39(1), 315–320, doi:10.1042/BST0390315, 2011.

Lawal, M. A., Bialik, O. M., Lazar, M., Waldmann, N. D., Foubert, A. and Makovsky, Y.: Modes of gas migration and seepage on the salt-rooted Palmahim Disturbance, southeastern Mediterranean, Mar. Pet. Geol., 153(November 2022), 106256, doi:10.1016/j.marpetgeo.2023.106256, 2023.

755 Levin, L. A. and Sibuet, M.: Understanding continental margin biodiversity: A new imperative, Ann. Rev. Mar. Sci., 4(1), 79–112, doi:10.1146/annurev-marine-120709-142714, 2012.

Levin, L. A., Baco, A. R., Bowden, D. A., Colaco, A., Cordes, E. E., Cunha, M. R., Demopoulos, A. W. J., Gobin, J., Grupe, B. M., Le, J., Metaxas, A., Netburn, A. N., Rouse, G. W., Thurber, A. R., Tunnicliffe, V., Van Dover, C. L., Vanreusel, A. and Watling, L.: Hydrothermal vents and methane seeps: rethinking the sphere of influence, Front. Mar. Sci., 3(October), 72, doi:10.3389/fmars.2016.00072, 2016.

Makovsky, Y., Rüggeberg, A., Bialik, O., Foubert, A., Almogi-Labin, A., Alter, Y., Bampas, V., Basso, D., Feenstra, E. and Fentimen, R.: South East Mediterranean SEEP Carbonate, R/V Aegaeo Cruise, EuroFLEETS2 SEMSEEP, R/V Aegaeo Cruise EuroFLEETS2 SEMSEEP., 2017.

Martens, E. C., Chiang, H. C. and Gordon, J. I.: Mucosal glycan foraging enhances fitness and transmission of a saccharolytic human gut bacterial symbiont, Cell Host Microbe, 4(5), 447–457, doi:10.1016/j.chom.2008.09.007, 2008.

Martens, E. C., Koropatkin, N. M., Smith, T. J. and Gordon, J. I.: Complex glycan catabolism by the human gut microbiota: The bacteroidetes sus-like paradigm, J. Biol. Chem., 284(37), 24673–24677, doi:10.1074/jbc.R109.022848, 2009.

Martin, M.: Cutadapt removes adapter sequences from high-throughput sequencing reads, EMBnet.journal, 17(1), 10–12, doi:10.14806/ej.17.1.200, 2011.

Mavropoulou, A. M., Vervatis, V. and Sofianos, S.: Dissolved oxygen variability in the Mediterranean Sea, J. Mar. Syst., 208(April), 103348, doi:10.1016/j.jmarsys.2020.103348, 2020.

Mayer, M., Schaffner, L. and Kemp, W.: Nitrification potentials of benthic macrofaunal tubes and burrow walls: effects of sediment NH4+ and animal irrigation behavior, Mar. Ecol. Prog. Ser., 121, 157–169, doi:10.3354/meps121157, 1995.

775 McMurdie, P. J. and Holmes, S.: phyloseq: An R Package for reproducible interactive analysis and graphics of microbiome census data, edited by M. Watson, PLoS One, 8(4), e61217, doi:10.1371/journal.pone.0061217, 2013.

Meile, C. and Cappellen, P. Van: Global estimates of enhanced solute transport in marine sediments, Limnol. Oceanogr., 48(2), 777–786, doi:10.4319/lo.2003.48.2.0777, 2003.

Mohr, W., Großkopf, T., Wallace, D. W. R. and LaRoche, J.: Methodological underestimation of oceanic nitrogen fixation rates, PLoS One, 5(9), e12583, doi:10.1371/journal.pone.0012583, 2010.

Molari, M., Hassenrueck, C., Laso-pérez, R., Wegener, G., Offre, P., Scilipoti, S. and Boetius, A.: A hydrogenotrophic *Sulfurimonas* is globally abundant in deep-sea oxygen-saturated hydrothermal plumes, Nat. Microbiol., 8(April), 651–665, doi:10.1038/s41564-023-01342-w, 2023.

Mulholland, M. R., Bernhardt, P. W., Heil, C. A., Bronk, D. A. and Neil, J. M. O.: Nitrogen fixation and release of fixed nitrogen by *Trichodesmium* spp. in the Gulf of Mexico, Limnol. Oceanogr., 51(4), 1762–1776, doi:10.4319/lo.2006.51.4.1762, 2006.

Ndeh, D. and Gilbert, H. J.: Biochemistry of complex glycan depolymerisation by the human gut microbiota, FEMS Microbiol. Rev., 42(2), 146–164, doi:10.1093/femsre/fuy002, 2018.

Nielsen, O., Gribsholt, B., Kristensen, E. and Revsbech, N.: Microscale distribution of oxygen and nitrate in sediment
 inhabited by *Nereis diversicolor*: spatial patterns and estimated reaction rates, Aquat. Microb. Ecol., 34, 23–32, doi:10.3354/ame034023, 2004.

Olm, M. R., Brown, C. T., Brooks, B. and Banfield, J. F.: DRep: A tool for fast and accurate genomic comparisons that enables improved genome recovery from metagenomes through de-replication, ISME J., 11(12), 2864–2868, doi:10.1038/ismej.2017.126, 2017.

- 795 Olu-Le Roy, K., Sibuet, M., Fiala-Médioni, A., Gofas, S., Salas, C., Mariotti, A., Foucher, J. P. J. P. J. P. J. P. and Woodside, J.: Cold seep communities in the deep eastern Mediterranean Sea: composition, symbiosis and spatial distribution on mud volcanoes, Deep Sea Res. Part I Oceanogr. Res. Pap., 51(12), 1915–1936, doi:10.1016/j.dsr.2004.07.004, 2004.
- Orphan, V., Hinrichs, K., Ussler III, W., Paull, C., Taylor, L., Sylva, S. P., Hayes, J. M. and Delong, E. F.: Comparative analysis of methane-oxidizing archaea and sulfate-reducing bacteria in anoxic marine sediments, Appl. Environ. Microbiol., 67(4), 1922–1934, doi:10.1128/AEM.67.4.1922, 2001.

Orsi, W. D.: Ecology and evolution of seafloor and subseafloor microbial communities, Nat. Rev. Microbiol., 16(11), 671–683, doi:10.1038/s41579-018-0046-8, 2018.

Overbeek, R., Olson, R., Pusch, G. D., Olsen, G. J., Davis, J. J., Disz, T., Edwards, R. A., Gerdes, S., Parrello, B., Shukla,
M., Vonstein, V., Wattam, A. R., Xia, F. and Stevens, R.: The SEED and the Rapid Annotation of microbial genomes using Subsystems Technology (RAST), Nucleic Acids Res., 42(D1), D206–D214, doi:10.1093/nar/gkt1226, 2014.

Papaspyrou, S., Gregersen, T., Cox, R. P., Thessalou-Legaki, M. and Kristensen, E.: Sediment properties and bacterial community in burrows of the ghost shrimp *Pestarella tyrrhena* (Decapoda: Thalassinidea), Aquat. Microb. Ecol., 38(2), 181–190, doi:10.3354/ame038181, 2005.

810 Parada, A. E., Needham, D. M. and Fuhrman, J. A.: Every base matters: Assessing small subunit rRNA primers for marine microbiomes with mock communities, time series and global field samples, Environ. Microbiol., 18(5), 1403– 1414, doi:10.1111/1462-2920.13023, 2016.

Petersen, J. M., Zielinski, F. U., Pape, T., Seifert, R., Moraru, C., Amann, R., Hourdez, S., Girguis, P. R., Wankel, S. D., Barbe, V., Pelletier, E., Fink, D., Borowski, C., Bach, W. and Dubilier, N.: Hydrogen is an energy source for hydrothermal vent symbioses., Nature, 476(7359), 176–80, doi:10.1038/nature10325, 2011.

Prjibelski, A., Antipov, D., Meleshko, D., Lapidus, A. and Korobeynikov, A.: Using SPAdes de novo assembler, Curr. Protoc. Bioinforma., 70(1), e102, doi:10.1002/cpbi.102, 2020.

R Core Team: R: A Language and Environment for Statistical Computing, [online] Available from: http://www.r-project.org/, 2024.

820 Rahav, E., Silverman, J., Raveh, O., Hazan, O., Rubin-Blum, M., Zeri, C., Gogou, A., Kralj, M., Pavlidou, A. and Kress, N.: The deep water of Eastern Mediterranean Sea is a hotspot for bacterial activity, Deep Sea Res. Part II Top. Stud. Oceanogr., 164, 135–143, doi:10.1016/j.dsr2.2019.03.004, 2019.

Reeves, A. R., D'Elia, J. N., Frias, J. and Salyers, A. A.: A Bacteroides thetaiotaomicron outer membrane protein that is essential for utilization of maltooligosaccharides and starch, J. Bacteriol., 178(3), 823–830, doi:10.1128/jb.178.3.823-830.1996.

Reineck, H.-E.: Sedimentgefüge im Bereich der südlichen Nordsee, Schweizerbart Science Publishers, Stuttgart,

815

825

845

Germany. [online] Available from: http://www.schweizerbart.de//publications/detail/isbn/9783510612956/Abhandlungen%5C\_d%5C\_Senckenb%5C\_Naturf %5C\_Gesell, 1963.

830 Ritt, B., Pierre, C., Gauthier, O., Wenzhöfer, F., Boetius, A. and Sarrazin, J.: Diversity and distribution of cold-seep fauna associated with different geological and environmental settings at mud volcanoes and pockmarks of the Nile Deep-Sea Fan, Mar. Biol., 158(6), 1187–1210, doi:10.1007/s00227-011-1679-6, 2011.

Roether, W. and Well, R.: Oxygen consumption in the Eastern Mediterranean, Deep. Res. Part I Oceanogr. Res. Pap., 48(6), 1535–1551, doi:10.1016/S0967-0637(00)00102-3, 2001.

- 835 Rubin-Blum, M., Shemesh, E., Goodman-Tchernov, B., Coleman, D. F., Ben-Avraham, Z. and Tchernov, D.: Cold seep biogenic carbonate crust in the Levantine basin is inhabited by burrowing *Phascolosoma* aff. *turnerae*, a sipunculan worm hosting a distinctive microbiota, Deep Sea Res. Part I Oceanogr. Res. Pap., 90(1), 17–26, doi:10.1016/j.dsr.2014.04.014, 2014a.
- Rubin-Blum, M., Antler, G., Turchyn, A. V., Tsadok, R., Goodman-Tchernov, B. N., Shemesh, E., Austin, J. A.,
  Coleman, D. F., Makovsky, Y., Sivan, O. and Tchernov, D.: Hydrocarbon-related microbial processes in the deep sediments of the Eastern Mediterranean Levantine Basin, FEMS Microbiol. Ecol., 87(3), 780–796, doi:10.1111/1574-6941.12264, 2014b.

Rubin-Blum, M., Dubilier, N. and Kleiner, M.: Genetic evidence for two carbon fixation pathways (the Calvin-Benson-Bassham Cycle and the reverse tricarboxylic acid cycle) in symbiotic and free-living bacteria, edited by S. J. Hallam, mSphere, 4(1), 00394–18, doi:10.1128/mSphere.00394-18, 2019.

Rubin-Blum, M., Sisma-Ventura, G., Yudkovski, Y., Belkin, N., Kanari, M., Herut, B. and Rahav, E.: Diversity, activity, and abundance of benthic microbes in the Southeastern Mediterranean Sea, FEMS Microbiol. Ecol., 98(2), fiac09, doi:10.1093/femsec/fiac009, 2022.

Rubin-Blum, M., Makovsky, Y., Rahav, E., Belkin, N., Antler, G., Sisma-Ventura, G. and Herut, B.: Active microbial
 communities facilitate carbon turnover in brine pools found in the deep Southeastern Mediterranean Sea, Mar. Environ.
 Res., 198(April), 106497, doi:10.1016/j.marenvres.2024.106497, 2024.

Ruff, S. E., Biddle, J. F., Teske, A. P., Knittel, K., Boetius, A. and Ramette, A.: Global dispersion and local diversification of the methane seep microbiome, Proc. Natl. Acad. Sci., 112(13), 4015–4020, doi:10.1073/pnas.1421865112, 2015.

855 Ruff, S. E., Humez, P., de Angelis, I. H., Diao, M., Nightingale, M., Cho, S., Connors, L., Kuloyo, O. O., Seltzer, A., Bowman, S., Wankel, S. D., McClain, C. N., Mayer, B. and Strous, M.: Hydrogen and dark oxygen drive microbial productivity in diverse groundwater ecosystems, Nat. Commun., 14(1), 3194, doi:10.1038/s41467-023-38523-4, 2023.

Santos Correa, S., Schultz, J., Lauersen, K. J. and Soares Rosado, A.: Natural carbon fixation and advances in synthetic engineering for redesigning and creating new fixation pathways, J. Adv. Res., 47, 75–92, doi:10.1016/j.jare.2022.07.011, 2022.

Seilacher, A.: Trace Fossil Analysis, Springer Berlin Heidelberg, Berlin, Heidelberg, Heidelberg., 2007.

860

870

875

890

Sieber, C. M. K., Probst, A. J., Sharrar, A., Thomas, B. C., Hess, M., Tringe, S. G. and Banfield, J. F.: Recovery of genomes from metagenomes via a dereplication, aggregation and scoring strategy, Nat. Microbiol., 3(7), 836–843, doi:10.1038/s41564-018-0171-1, 2018.

865 Simon, M. and Azam, F.: Protein content and protein synthesis rates of planktonic marine bacteria, Mar. Ecol. Prog. Ser., 51, 201–213, doi:10.3354/meps051201, 1989.

Sisma-Ventura, G., Bialik, O. M., Makovsky, Y., Rahav, E., Ozer, T., Kanari, M., Marmen, S., Belkin, N., Guy-Haim, T., Antler, G., Herut, B. and Rubin-Blum, M.: Cold seeps alter the near-bottom biogeochemistry in the ultraoligotrophic Southeastern Mediterranean Sea, Deep Sea Res. Part I Oceanogr. Res. Pap., 183, 103744, doi:10.1016/j.dsr.2022.103744, 2022.

Smith, D. C., Smith, D. C., Azam, F. and Azam, F.: A simple, economical method for measuring bacterial protein synthesis rates in seawater using <sup>3</sup>H-leucine, Mar. Microb. food web, 6(2), 107–114, 1992.

Stamhuis, E. J., Schreurs, C. E. and Videler, J. J.: Burrow architecture and turbative activity of the thalassinid shrimp *Callianassa subterranea* from the central North Sea, Mar. Ecol. Prog. Ser., 151(1/3), 155–163 [online] Available from: http://www.jstor.org/stable/24857699, 1997.

Teske, A. and Carvalho, V.: Marine hydrocarbon seeps: microbiology and biogeochemistry of a global marine habitat, edited by A. Teske and V. Carvalho, Springer International Publishing, Cham, Switzerland., 2020.

Thomas, F., Hehemann, J. H., Rebuffet, E., Czjzek, M. and Michel, G.: Environmental and gut Bacteroidetes: The food connection, Front. Microbiol., 2(MAY), 93, doi:10.3389/fmicb.2011.00093, 2011.

880 Trojan, D., Schreiber, L., Bjerg, J. T., Bøggild, A., Yang, T., Kjeldsen, K. U. and Schramm, A.: A taxonomic framework for cable bacteria and proposal of the candidate genera *Electrothrix* and *Electronema*, Syst. Appl. Microbiol., 39(5), 297–306, doi:10.1016/j.syapm.2016.05.006, 2016.

Trueman, E. R., Brand, A. R. and Davis, P.: The dynamics of burrowing of some common littoral bivalves, J. Exp. Biol., 44(3), 469–492, doi:10.1242/jeb.44.3.469, 1966.

885 Ward, L. M., Bertran, E. and Johnston, D. T.: Expanded genomic sampling refines current understanding of the distribution and evolution of sulfur metabolisms in the Desulfobulbales, Front. Microbiol., 12(May), 666052, doi:10.3389/fmicb.2021.666052, 2021.

Weidlich, R., Bialik, O. M., Rüggeberg, A., Grobéty, B., Vennemann, T., Neuman, A., Makovsky, Y. and Foubert, A.: Occurrence and genesis of cold-seep authigenic carbonates from the South-Eastern Mediterranean Sea, Depos. Rec., (May), 239, doi:10.1002/dep2.239, 2023.

Weiss, R. F.: The solubility of nitrogen, oxygen and argon in water and seawater, Deep. Res. Oceanogr. Abstr., 17(4), 721–735, doi:10.1016/0011-7471(70)90037-9, 1970.

Wolf, P. G., Biswas, A., Morales, S. E., Greening, C. and Gaskins, H. R.: H2 metabolism is widespread and diverse among human colonic microbes, Gut Microbes, 7(3), 235–245, doi:10.1080/19490976.2016.1182288, 2016.

895 Wu, Y.-W., Simmons, B. A. and Singer, S. W.: MaxBin 2.0: an automated binning algorithm to recover genomes from multiple metagenomic datasets, Bioinformatics, 32(4), 605–607, doi:10.1093/bioinformatics/btv638, 2015.

Xu, Z., Wang, M., Zhang, H., He, W., Cao, L., Lian, C., Zhong, Z., Wang, H., Fu, L., Zhang, X. and Li, C.: Metabolism interactions promote the overall functioning of the episymbiotic chemosynthetic community of *Shinkaia crosnieri* of cold

seeps, mSystems, 7(4), e0032022, doi:10.1128/msystems.00320-22, 2022.

900 Zhang, C., Fang, Y. X., Yin, X., Lai, H., Kuang, Z., Zhang, T., Xu, X. P., Wegener, G., Wang, J. H. and Dong, X.: The majority of microorganisms in gas hydrate-bearing subseafloor sediments ferment macromolecules, Microbiome, 11(1), 37, doi:10.1186/s40168-023-01482-5, 2023.

Zhou, Z., Tran, P. Q., Breister, A. M., Liu, Y., Kieft, K., Cowley, E. S., Karaoz, U. and Anantharaman, K.:
 METABOLIC: high-throughput profiling of microbial genomes for functional traits, metabolism, biogeochemistry, and community-scale functional networks, Microbiome, 10(1), 33, doi:10.1186/s40168-021-01213-8, 2022.

Zoccarato, L., Sher, D., Miki, T., Segrè, D. and Grossart, H. P.: A comparative whole-genome approach identifies bacterial traits for marine microbial interactions, Commun. Biol., 5(1), 276, doi:10.1038/s42003-022-03184-4, 2022.

Zvi-Kedem, T., Vintila, S., Kleiner, M., Tchernov, D. and Rubin-Blum, M.: Metabolic handoffs between multiple symbionts may benefit the deep-sea bathymodioline mussels, ISME Commun., 3(1), 48, doi:10.1038/s43705-023-00254-4, 2023.

910 4, 20

905

h	1 5	
P	12	

# 920

#### **Figure legends:**

- 925 Figure 1. A bathymetric map of Palmahim Disturbance and its vicinity (color-coded and contoured with a 100 m spacing). This study is focused on seafloor gas seeps in the ridge-crest pockmark system (RC1) at the southwestern edge of the Disturbance, the proximate part of the Levant Channel (LC), and the brine pool (BP) area to the north. Black dots mark the sites of box-core sampling, and the black frames outline the ROV surveying analysis maps of Figure 2D-E.
- Figure 2. Classification and mapping of burrowing intensities and their correlation with indicators of active seepage. ROV
  images of low (A), medium (B), and high (C) burrowing intensities are shown (similar scales, with the green laser beams in c. being 7.5 cm apart). D) Synthetic aperture sonar (SAS) backscatter image of the gradient in burrowing intensity from low (top-right) to high (bottom) inside a small pockmark at the edge of the RC1 pockmark system. The burrowing mounds are observed as a decimeter-scale doted dark-bright pattern (white arrows show examples). E) Map of classified burrowing intensities and active seepage indicators along EV Nautilus 2011 ROV survey tracks of a ridge-crest pockmark system
  RC1, overlaid on the contours of bathymetry (at 5 m spacing) and a grey-scale map of the bathymetric gradient (high
- gradients are darker). Locations of the SAS swath in panel D and the cored area are marked by black arrows in panel E.
- Figure 3. Burrows of the ghost shrimp *Calliax lobata* near active seepage sites within <u>PalamhimPalmahim</u> Disturbance in the Eastern Mediterranean Sea. A) The seafloor, as exposed in the AC16-17BC1 box-core (RC1 area, 2016), shows

burrowing mounds and burrow openings, as well as (black arrows) and scratches (dashed arrows) on the sediment surface.
 B) Image of RC20-BC1 and RC20-BC2 cores (RC1 area, 2020), showing dark coloration of burrows, unusual for the ambient sediments. C and D) Zooming into the burrows of RC20-BC1 and RC20-BC2 cores showed the presence of large filamentous microbes. E) Live *C. lobata* specimen found in AC16-17BC1. F) A multitude of *C. lobata* claws recovered through sporadic sieving (2 mm mesh) of AC16-17BC1.

Figure 4. Burrowing in AG16-15BC1 and AG16-17BC1, as detected by computerized tomography (CT) scans. A) A vertical section through a sub-core of box-core AC16-17BC1 cut and imaged immediately after sampling. B) A vertical section through the 3D CT scan of another sub-core of -AC16-17BC1 (standard medical CT display where higher relative X-ray intensities are darker and low are brighter, depicting lower and higher material densities, respectively). C and D) Two perpendicular 3D displays of segmentation of the relatively vacant (high intensity) burrows, imaged in the CT scan of the same core (in B). E) A vertical section through the 3D CT scan of a sub-core of -AC16-15BC1 (displayed as in B). F)

950 A 3D display of AC16-15BC1 segmentation of the vacant (high intensity) burrows. G) Graphs comparing the volumetric % of relatively vacant burrows in CT scanned sub-cores of AC16-17BC1 (green) and AC16-15BC1 (red), as estimated based on the respective segmentation results (in C-D and F).

Figure 5: Pore water characteristics in cores AG16-15BC1 and AG16-17BC1: A) Nitrate+nitrite (NOx) concentration. B) NOx to phosphate ratio, C) Sulfate concentration. Red and green lines indicate the depths at which NOx concentrations appear to stabilize.

Figure 6: Abundance and activity of microbes in core AG16-17BC1: A) Abundance and activity profiles. B) Activity versus burrowing. C) Activity versus burrowing – upper 5 cm removed.

Figure 7: Read abundance of the top 30 most prevalent prokaryote taxa at the family level, in Palmahim sediments (see Table 1) and the hindgut of the ghost shrimp *Calliax lobata* (black: enriched in burrows, blue: enriched in *C. lobata*, brown: enriched in sediments). Sediment depths where burrow microbes were collected are highlighted in orange. \* Both samples are from the 16-17 cm section, yet the first one is taken from the burrow and the second from background sediments. \*\* abundant in burrows and associated with *C. lobata* hindgut.

Figure 8: Phylogeny, taxonomy and key functions of bacteria, based on the reconstruction of metagenome-assembled genomes (MAGs) from *Calliax lobata* burrows. FastTree phylogeny is based on the GTDB-tk marker gene set. Functions
 were assigned using METABOLIC Hidden Markov Model (HMM) set. Empty circles and stars indicate the functions discovered through manual annotation of functions, using SEED and NCBI BLAST.

Figure 9: N<sub>2</sub> fixation rates at the overlying water (A,  $\sim$ 1 m above seabed) and in the top sediment (B, upper 3 cm) in an active SEEP site (AG16 17BC1, grey) and in an background station (AG16 15BC1). The plot shows averages and corresponding standard deviation of three replicates. Filtered seawater (FSW) or filtered porewater (FPW) served as a control.

970

955