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# 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

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#### Abstract:

The Nile Deep-Sea Fan (NDSF) is located on the passive continental margin off Egypt and is characterized by the occurrence of active fluid seepage such as brine lakes, pockmarks and mud volcanoes. This study characterizes the structure of faunal assemblages of such active seepage systems of the NDSF. Benthic communities associated with reduced, sulphidic microhabitats such as sediments and carbonate crusts were sampled by remotely operated vehicles during two cruises in 2006 (BIONIL) and 2007 (MEDECO). Environmental conditions and biological factors including family-level faunal composition, density and diversity were measured at local and regional scales. Significant differences were detected at different spatial scales: (1) the fauna of reduced habitats differed substantially in activity, diversity and biomass from the non-seep environment at similar water depth, (2) cold seep microhabitats showed differences in community structure and composition related to substratum type as well as to the intensity and location of fluid emissions.

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Keywords: Nile Deep-Sea Fan; cold seeps; benthic macrofauna; alpha diversity; environmental conditions; chemosynthetic ecosystem; beta diversity.

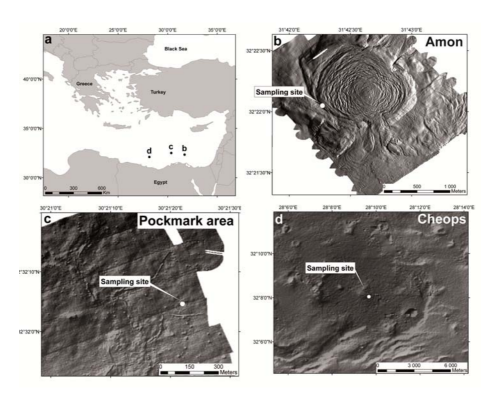
## 1. Introduction

Since their discovery on the Florida Escarpment in the Gulf of Mexico (Paull et al. 1984), cold seeps have been reported along convergent plate boundaries on active as well as passive continental margins, where over-pressure in the sediments controls the emission of fluids enriched in methane or other hydrocarbons to the seafloor and hydrosphere (Milkov 2000; Dimitrov 2002). Depending on their geophysical settings, these cold seeps are associated with distinct geological features such as gas hydrates, pockmarks, brine lakes and mud volcanoes (MVs) (Milkov 2000; Hovland et al. 2002; Judd and Hovland 2007; Foucher et al. 2009). To date, hundreds of seep sites supporting chemosynthesis-based communities have been encountered throughout the world's oceans (Sibuet and Olu 1998; Levin 2005; Campbell 2006; Baker et al. 2010).

The Eastern Mediterranean Sea is one of the world's MV and pockmark hotspots (Kopf 2002; Foucher et al. 2009). Most of the MVs are associated with the active Mediterranean Ridge that stretches over more than 1500 km (Cita et al. 1981) and the thickly sedimented Nile Deep-Sea Fan (NDSF) along the Mesozoic-rifted continental margin of northern Egypt (Bellaiche et al. 2001; Mascle et al. 2001; Loncke et al. 2004; Mascle et al. 2006). The present study focuses on sites of the NDSF that have been explored since 2003 in the frame of the European Science Foundation (ESF) MEDIFLUX program (2003-2007) and the European 6<sup>th</sup> Framework Program project HERMES (2005-2009).

The NDSF is divided into four morpho-structural provinces (Figure 1a): (1) the Levantine province, a domain of salt-related folding; (2) the Eastern province, a domain of intense salt-related tectonics; (3) the Central province with pockmark fields and gas chimneys, distinguished by active sedimentary instability and fluid-related processes; and (4) the Western province with mud cones, characterized by active turbiditic processes, salt tectonics and fluid venting (Mascle et al. 2006; Huguen et al. 2009). The accumulation of organic-rich sediments, probably since the early

Cenozoic, led to the formation of hydrocarbons in deep reservoirs that were partially sealed by the deposition of thick evaporites during the Messinian (Hsu et al. 1977). A mixture of hydrocarbons, water and mud is expelled through faults that may extrude salt deposits, which reach the seafloor (Loncke et al. 2004). Fluid seepage is recorded in geological features such as mud cones, caldera-like depressions, gas chimneys, brine pools and pockmarks (Gontharet et al. 2007; Bayon et al. 2009a). Chemosynthesis-based communities have been observed in association with these different features (Huguen et al. 2005; Zitter et al. 2005; Dupré et al. 2007).



**Fig. 1** (a) General map of the Eastern Mediterranean Sea with the localisation of the sampling sites in the three morpho-structural provinces: (b) the Amon mud volcano (MV) located in the Eastern province; (c) the Pockmark area located in the Central province and, (d) the Cheops MV in the Western province. Sampling sites are indicated on each MVs

Faunal assemblages associated with Mediterranean cold seeps from the NDSF are still relatively unknown. Only those from a few sites along the Mediterranean Ridge have been described, and symbiont-bearing species such as the siboglinid polychaete *Lamellibrachia anaximandri*, the lucinid *Lucinoma kazani* and the mytilid *Idas modiolaefomis* were identified (Salas and Woodside 2002; Olu-Le Roy et al. 2004; Werne et al. 2004; Duperron et al. 2008; Southward et al. in press). The distribution of fauna on the Amsterdam MV has been hypothesized to be linked to the

amount of methane escaping from the mud flows, which decreases from the summit to the periphery (Olu-Le Roy et al. 2004). Likewise, biological zonation has also been observed on the Barbados prism (Olu et al. 1997) and the Håkon Mosby MV (Niemann et al. 2006; Jerosch et al. 2007). In both cases, the summit of the MV was covered by fresh gassy mud flows and devoid of visible epifauna. Communities dominated by sulphide oxidizing bacterial mats were observed close to the summit. followed by symbiont-bearing fauna and heterotrophic fauna towards the periphery (Olu et al. 1997; Zitter et al. 2003; Niemann et al. 2006; Jerosch et al. 2007). Fresh mud flows are often characterized by very high rates of upward flux of reduced sulphidic fluids, excluding animals (de Beer et al. 2006). Older mud flows, located at the periphery, transport lower concentrations of methane and sustain sulphide production, fueling chemosynthetic populations of siboglinid tubeworms or bivalves like Acharax. The latter migrate to deeper sediment layers to reach for reduced chemical compounds. This environment also constitutes a favorable habitat for heterotrophic fauna, which develop in response to a local increase in microbial production (Levin and Mendoza 2007).

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The present study was the first opportunity to characterize the structure of the faunal assemblages in the NDSF seeps on two types of geological features: MVs and pockmarks. Prior to 2006, no ecological study had been performed on NDSF seep sites, with the exception of a few observations and samples taken during the NAUTINIL cruise in 2003 (Dupré et al. 2007; Bayon et al. 2009b; Huguen et al. 2009). The macro- and meiofaunal benthic communities associated with different microhabitats found in three provinces of the NDSF were sampled by remotely operated vehicles (ROVs) during two cruises: BIONIL in 2006 and MEDECO in 2007. The microhabitats were characterized with regard to their environmental conditions and faunal communities, mostly to family level. We then compared faunal composition, density and diversity at local and regional scales to test the hypotheses that 1) reduced habitats support higher biomasses but lower diversity of meio- and macrofauna compared to surrounding oxygenated habitats, 2) carbonate crusts and reduced sediments bear different faunal assemblages, 3) faunal composition is related to microhabitat type rather than to the larger geological setting, 4) beta diversity between the different microhabitats is related to differences in fluid flow and substratum type.

124 2. Materials and methods

- Faunal sampling and habitat characterizations were done during BIONIL M70/2b
- aboard the German R/V Meteor with the ROV Quest4000 (MARUM, University
- Bremen) in November 2006 and during leg 2 of MEDECO aboard the French R/V
- 128 Pourquoi Pas? with the ROV Victor6000 (Ifremer) in November 2007.

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- 2.1. Study sites
- The cold seep sites from three of the four provinces of the NDSF were investigated:
- 1) the Amon MV in the Eastern province; 2) a carbonate cemented area close to a
- pockmark field associated with large debris-flows in the Central province; and 3) the
- 134 Cheops MV in the Western province hosting large brine pools (Figure 1). A single
- reference sample located 15 km away from the centre of the Amon MV and outside
- the influence of fluid emissions, was sampled as a reference (hereafter marked "Ref")
- for oxygenated deep-sea sediments not associated with fluid flow.

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- 2.1.1. The Western province the Amon MV
- The Amon MV (32°22'05"N 31°42'27"E, Figure 1b) is circular, approximately 3 km
- in diameter, and 90 m high. It lies close to the limit of the Messinian platform, at 1150
- m water depth. The summit is covered with mud blocks and clasts and shows a
- disturbed, chaotically structured surface suggesting recent impacts of mud extrusion
- and gas expansion. Temperatures at the centre reach 45°C at 10 m below the
- seafloor, confirming high upward fluid flow at this MV (Dupré et al. 2007; Dupré et al.
- 146 2008). The periphery of the Amon MV is characterized by highly bioturbated
- hemipelagic sediments. At its south-western rise, a lateral flow of reduced muds was
- identified ('sulphur-band') surrounded by carbonate crusts that were both sampled in
- 2006 during BIONIL. The carbonate crusts were rather thick, we could not observe
- blackish muds or siboglinid colonies associated with them.

- 2.1.2. The Central province the Pockmark area
- 153 The Central province hosts large carbonate-cemented areas associated with reduced
- debris-flows, and numerous small pockmarks located between 1700 and 2100 m
- depth (Figure 1c). The pockmarks form circular depressions of a few meters in
- diameter and about 1 m deep, and are associated with the presence of authigenic

carbonate crusts and reduced sediments (Loncke et al. 2004). During BIONIL, benthic communities were sampled from one reduced blackish sediments site at midslope (site 2A, 32°32'00"N - 30°21'10"E, 1700 m) as well as from their surrounding carbonate cements. These were directly associated with blackish muds, and living siboglinid colonies were observed between the cracks of the flat, thin carbonates.

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- 2.1.3. The Eastern province the Cheops MV
- The Cheops MV is located within the Menes caldera at 3000 m depth (32°08'05"N -164 28°09'67"E, Figure 1d), above the Messinian platform. This caldera is a circular 50 m deep depression 8 km in diameter. As previously observed in 2003 during the 166 NAUTINIL cruise, a mixture of brine and mud was flowing from Cheops during the sampling operations in 2007 (MEDECO). The mud is expelled from deep layers as 168 169 indicated by high temperature anomalies (Huguen et al. 2009). This MV is also characterized by numerous brine pools, covered by bright white matter, identified as 170 171 microbial sulphur deposits (Dupré et al. 2007; Omoregie et al. 2008). Brine pools constitute direct evidence of fluid emissions, and the migration of fluids enriched in 172 salt induces high sediment instability that may influence faunal composition and 173 distribution on this MV. Faunal sampling and environmental characterizations were 174 carried out during MEDECO. Sparse carbonate crusts were observed but not 175 sampled. 176

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- 2.2. Description of the microhabitats 178
  - At all cold seep sites visited, ROV surveys indicated a mosaic of visually distinguishable microhabitats, characterized by the presence of visible fauna or microbial mats. Two types of reduced habitats were sampled: 1) reduced blackish sediments (Red) covered with whitish bacterial mats or small tubeworms at the surface (Figures 2a, c, e) and 2) carbonate crusts (CC) that were dark-colored at Amon, or "crumbly" and of whitish to light grey color at the Pockmark area Figures 2b, d).

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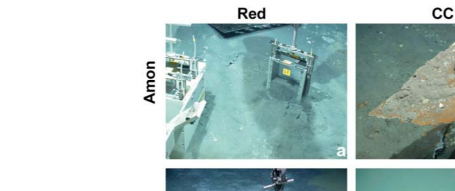
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Chemical characterization of the microhabitats was performed above the reduced sediments of Cheops MV during MEDECO. Water samples were taken for chemical analyses using the PEPITO water sampler above the organisms as close as possible to the seafloor using the Victor6000 manipulator arm. PEPITO collects water in 200

ml titanium bottles (Sarradin et al. 2009). Chemical measurements and sediment samples were taken during BIONIL as close as possible above the seafloor or in soft sediments with bottles (750 ml) and push cores (inner diameter 7.5 cm) at 2.5 to 50 m from the sampled microhabitats. Further sampling details are given in Table 1.



Pockmark





**Fig. 2** Representative photographs of the microhabitats sampled in the three study sites. On the Amon MV: (a) reduced sediments and (b) carbonate crusts; in the Pockmark area: (c) reduced sediments and (d) carbonate crusts; on the Cheops MV: (e) reduced sediments. Red: reduced sediments CC: carbonate crusts. All microhabitats were sampled during the BIONIL (2006) and MEDECO (2007) cruises. Photos a, b, c, d: MARUM, University Bremen, *QUEST4000* and photo e: Ifremer, *Victor6000* 

Epi- and endofauna were sampled with blade corers on a surface of 200 cm² (Menot et al. 2010) on all soft sediment sites (Figures 2a, c, d, g) and pieces of carbonate crusts were sampled using the claw of the ROV *Victor6000* (Figures 2b, e). A reference site 15 km away from the active centre of Amon has been sampled with a single multicorer on which 3 tubes of a surface of 74 cm² were sampled (pseudoreplicates). The number of samples for each microhabitat is reported in Table 1.

Quantitative sampling of the crusts was difficult because they had to be broken off in pieces with the submersible manipulator. No crusts were obtained from Cheops.

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2.3. Physico-chemical analyses

The 200 ml water samples were used to determine methane concentrations using the technique of headspace sampling gas chromatography with a thermal-conductivity detector (TCD) and a flame-ionisation detector (error of 4%; see method in Sarradin and Caprais 1996).

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Sediments from push corers were split horizontally at 1 cm intervals on board and porewater was extracted by centrifugation of the different sediment layers. After filtration and dilution, sulphate and chloride concentrations were measured by nonsuppressed anion exchange chromatography (Water IC-Pak anion exchange column, waters 430 conductivity detector). Total dissolved sulphide concentrations were determined with the diamine complexation method by colorimetric method (Cline 1969). Intact cores, along with supernatant water, were used for pH, oxygen and total dissolved sulphide concentration measurements. Small-scale porewater concentration profiles for O<sub>2</sub>, H<sub>2</sub>S, and pH were performed on push core samples from soft sediments using microelectrodes as described in de Beer et al. (2006). Microelectrodes with a tip diameter of ca. 20 µm were lowered into the sediment with a step resolution of 200 µm to monitor the concentration profiles within the upper sediment layer. Total oxygen consumption was measured in situ using a ROVoperated benthic chamber module (Treude et al. 2009). The chamber encloses an area of ca. 285 cm<sup>2</sup> together with approximately 10-15 cm of overlying bottom water. The change in O<sub>2</sub> concentration over time in the enclosed water volume was continuously monitored by two Clark-type mini-electrodes mounted in the chamber lid. This measurement integrates all relevant transport and consumption processes (diffusion, advection and fauna mediated transport as well as fauna respiration).

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2.4. Faunal sorting and identification

The faunal samples were processed as described in Ritt et al. (2010). Sediments from blade corers were photographed and split horizontally (0-1, 1-3, 3-5, 5-10, >10 cm) immediately after recovery. The range of the last slice depended on core lengths that varied from 10 to 20 cm (Table1). Core slices were passed through a sieve

column (2 mm, 1 mm, 500  $\mu$ m, 250  $\mu$ m) and the retained sediment was preserved in 10% buffered formalin. In the laboratory, all sediments up to 10 cm below the seafloor were rinsed and invertebrates were sorted under a dissecting microscope and identified to the lowest taxonomic level possible (here mostly family level). In this study, we considered macrofauna *sensu stricto* (>250  $\mu$ m, Hessler and Jumars 1974), and any meiofaunal taxa such as Nematoda, Copepoda, Ostracoda and Acarina were analysed separately. As a consequence, meiofaunal samples contain only the largest fraction retained by a 250  $\mu$ m mesh, and do not include the 32  $\mu$ m or 62  $\mu$ m size limit usual for meiofauna (Hessler and Jumars 1974; Thistle 2003; Van Gaever et al. 2006).

The CC samples were preserved individually in 10% buffered formalin on board after recovery. In the laboratory, the carbonates crusts were treated individually by retrieving the organisms embedded or attached on them without breaking the carbonate crusts. Then they were washed over a 250 µm mesh and the organisms retained were processed as those on the soft sediments. The surface of the sampled carbonate crusts were estimated using the IPLab Spectrum® image analysis software. Quantitative 2-D surface analyses were performed on video images of the upper face of the crusts, and we did it thrice to reduce error resulting from on-screen tracing (Sarrazin and Juniper 1999). Total surface area was used to calculate arearelated indicators, such as density and biomass by counting organisms at both faces and in the holes but divided the total of individuals only by the surface area. However, because it does not take topography into account, this method may underestimate total surface area and in turn overestimate density and biomass. The crusts from the Pockmark area had a more complex topography with numerous holes in comparison with the flat carbonates from Amon, hence, the two-side method may have added to the uncertainty with areal estimates in this case.

## 2.5. Vertical distributions

Vertical distributions within each reduced sediment microhabitat and the reference site were studied in the depth layers 0-1, 1-3, 3-5 and 5-10. Densities were calculated by summing up the sediment layers up to 10 cm below the seafloor. This was repeated for relative abundance and biomass. Biomass was assessed by measuring the mean preserved wet weight (pww) for each microhabitat. To do so, individuals of

271 all major macrofaunal taxa (bivalves, polychaetes, gastropods and crustaceans) were 272 pooled, pat-dried on absorbent paper and weighed on a microbalance with an error 273 of 0.1 mg.

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- 2.6. Diversity measurements
- Within-microhabitat ( $\alpha$ -diversity) and between-microhabitat ( $\beta$ -diversity) diversity (Whittaker 1960; Gray 2000) were estimated for all microhabitats.

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279 2.6.1. Alpha-diversity

The  $\alpha$ -diversity analyses were performed at the family level, which was reached for 280 most of the taxa except Zoantharia, Scyphozoa, Terebellida, Isopoda, Leptostraca, 281 Nematoda and Acarina. Undetermined families, individuals and larvae were removed 282 from the analysis because of the probability that they belong to a family already 283 listed. This may have resulted in an underestimation of the taxonomic richness. 284 Sample-based rarefaction curves (sensu Gotelli and Colwell 2001) were calculated 285 on macrofaunal datasets for each of the three study sites (Sanders 1968). Individual-286 based rarefaction curves were also computed on macrofaunal datasets for each 287 microhabitat type, regardless of the province of origin. These curves plot expected 288 taxonomic richness against sampling effort and allow comparisons impossible with 289 observed richness (S). All rarefaction curves use the expected number of individuals 290 as the X-axis (Sanders 1968, corrected and modified by Hurlbert 1971; Gotelli and 291 Colwell 2001; Gauthier et al. 2010). Observed within-microhabitat taxonomic diversity 292 293 was evaluated using common diversity indices as well as more robust intrinsicdiversity-based ordering methods (Liu et al. 2007; Gauthier et al. 2010). Commonly 294 used to define the  $\alpha$  diversity, the taxonomic richness (S), the number of taxonomic 295 groups observed in each microhabitat, the Shannon's entropy ( $H_e$ ; Shannon 1948) 296 and the Gini-Simpson diversity index ( $D_{GS}$ ; Gini 1912; Simpson 1949) were 297 calculated. They are presented along with their numbers equivalents, allowing 298 299 straightforward comparisons between communities (Hill 1973; Patil and Taillie 1982; Jost 2006; Jost 2007). Community evenness was also determined using Pielou's 300 index of evenness (J'; Pielou 1969). 301

The right tail-sum method (RTS) is a diversity ordering method, which is more robust and stringent than other methods and allows graphical comparisons of communities (Patil and Taillie 1982; Tothmérész 1998; Liu et al. 2007). Communities are ordered in decreasing diversity from the top most curves to the lowest ones. No clear conclusions can be drawn when curves intersect (Liu et al. 2007).

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microhabitat variation.

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# 2.6.2. Beta-diversity

Despite the modest number of samples, multivariate analyses were conducted to better illustrate the similarities and differences among faunal samples. The lowest available taxonomic level was used. Principal component analysis (PCA) and Ward's hierarchical clustering were used to indirectly evaluate the influence of habitat conditions on community structure variation within and between the different microhabitats and sites, but excluding the undetermined taxa as previously explained for the alpha-diversity. A Procrustean randomization test (Jackson 1995) was performed to compare the PCA results of the macrofaunal and meiofaunal datasets. Abundance data were first Hellinger-transformed to conserve Hellinger, rather than Euclidian, distances in PCA (Legendre and Gallagher 2001). Hellinger distances were also used for Ward's hierarchical clustering. This distance gives low weight to rare taxa in the analyses. In marine ecology in general, and even more so in the deep-sea, rare species are not well sampled, and their sporadic appearance in samples is mosly attributable to sampling error. In the PCA, the equilibrium contribution circle was computed to identify the taxa having the most impact on the position of the samples in the ordination (Legendre and Legendre 1998). The Jaccard's similarity coefficient  $(S_{iacc})$  was used to quantify similarity in terms of

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All analyses were performed in the R environment (R, Development Core Team, 2009). Rarefaction curves, diversity indices and diversity profiles were computed both with the Biodiversity R package (Kindt and Coe 2005) and functions in Gauthier et al. (2010). Multivariate analysis was carried out using the vegan package (Oksanen et al. 2008).

shared taxa among replicates within each microhabitat (Jaccard 1901); giving equal

weight to all taxa. Mean Jaccard similarity was computed to evaluate within-

- 3. Results
- 3.1. Physico-chemical characterization of microhabitats
- The length of the cores varied from 10 to 20 cm depending on the nature of the
- 339 substratum (Table 1). All samples came from surface cold seep habitats at in situ
- bottom water temperature of 13.5°C, as at the reference site. The Red cores from the
- Pockmark area contained black sediments with a strong hydrogen sulphide odor. At
- Amon, a horizon of 8-10 cm of black sediments overlaid grayish to beige hemipelagic
- sediments. On Cheops, a black layer of only 1-2 cm was found on top of beige
- 344 hemipelagic sediments. The cores of the reference site contained only beige
- 345 sediments.
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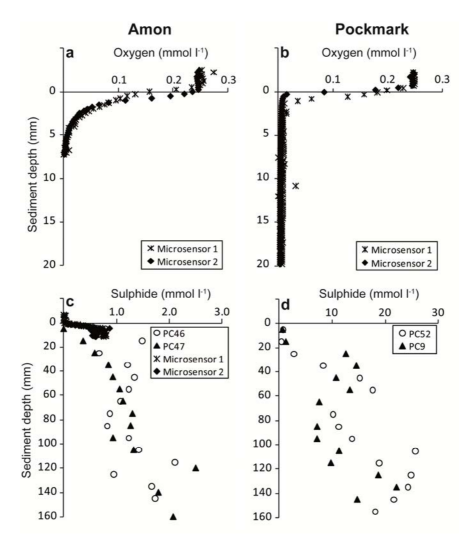
- 3.2. Chemical characterization at a larger scale around microhabitats
- 3.2.1. Bottom water measurements
- Overall, the pH was slightly lower in reduced sediments than at the reference site
- 350 (Table 2). According to microprofiler measurements, pH value reached 7.9 at the
- interface of reduced sediments on Amon while in the Pockmark area, it was around
- 352 8.1 (Table 2).
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- Oxygen bottom water concentrations were similar at all sites, including the reference
- site (200-230 µmol l<sup>-1</sup>) with the exception of the reduced sediments at Amon, where a
- temporary depletion was observed, reaching values below 200 µmol l<sup>-1</sup>. No free
- 357 sulphide was detected in the bottom waters at any of the sites, but sulphide was
- found within the porewaters at two reduced sediment sites (Table 2). Low
- concentrations of methane were measured in the bottom waters of Cheops and the
- Pockmark area while they ranged between 3.6 and 9.3 µmol l<sup>-1</sup> in the overlying water
- of Cheops samples (Table 2).
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- 363 Sulphate values of 31 mmol I<sup>-1</sup> and chloride values of 529 mmol I<sup>-1</sup> were measured
- 364 above and throughout the core from the reference site. Sulphate and chloride
- measurements above the reduced sediments on Amon showed values varying from
- 366 30 to 40 mmol l<sup>-1</sup> and 404 to 580 mmol l<sup>-1</sup> respectively (Table 2). Here, spatial
- heterogeneity between samples was high, despite the proximity of the cores (few dm,
- e.g. PC 46 and PC 47). In microbial mats from the Pockmark area, sulphate

concentrations varied from 29 to 30 mmol  $I^{-1}$  while chloride values were high, ranging from 606 to 630 mmol  $I^{-1}$  (Table 2).

3.2.2. Porewaters, oxygen consumption and sulfate reduction rates in sediments Oxygen penetration depth measured by microsensors was >4 cm at the reference site (Table 2). The porewater samples from the reference site did not contain sulphide or methane. Accordingly, sulphate reduction rate at the reference site was not measurable, and oxygen consumption was <1 mmol m<sup>-2</sup> d<sup>-1</sup> (Table 2).

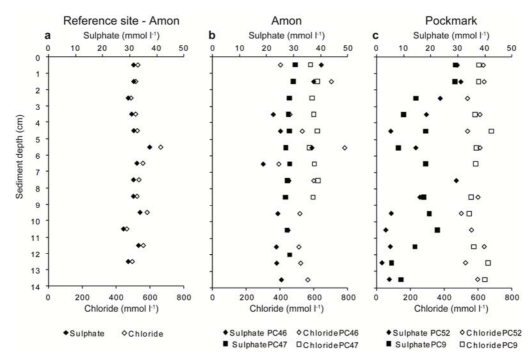
In contrast, oxygen microsensor profiles in reduced sediments of Amon, Cheops and the Pockmark area showed complete oxygen consumption within the first 1-2 millimeters of seafloor (Figures 3a, b). Total dissolved sulphide profiles in reduced sediments of Amon showed a maximum concentration of 2.5 mmol  $\Gamma^1$  at 12 cm depth and were <0.8 mmol  $\Gamma^1$  in the top 5 cm (Table 2, Figure 3c). The profile suggested that sulphide was oxidized completely within the first millimeters of sediment. In the Pockmark area, sulphide concentrations at 12.5 cm depth reached 25 mmol  $\Gamma^1$  (Figure 3d), but sulphide was also completely consumed within the surface sediments (Table 2). No porewater data were available for Cheops due to limitation in dive time.

At the reference site off Amon, sulphate and chloride concentrations were homogeneous along the whole length of the core with a mean of 32±2.4 mmol I<sup>-1</sup> for sulphate and 530±49.5 mmol I<sup>-1</sup> for chloride (Figure 4a). On Amon reduced sediments, sulphate and chloride profiles were homogeneous along the length of the core reaching values similar to the reference sample with the exception of a high sulphate concentration at the top of the core (Figure 4b). In the Pockmark area, the consumption of sulphate was visible throughout the 16 cm of the core length, whereas lower variation was observed in chloride (Figure 4c).



**Fig. 3** (a, b) Dissolved oxygen and (c, d) sulphide profiles measured by microsensors and in pore water extracted from push corers (PC) taken in reduced sediments on the (a, c) Amon MV and in (b, d) bacterial mats covering reduced sediments in the Pockmark area. The measurements were performed during the BIONIL cruise (2006)

Integrated sulphate reduction rates were negligible at the reference site, and low at the reduced sediment of Amon. Intermediate rates were measured for Cheops and high rates associated with the Pockmark area (Table 2). Likewise, total benthic oxygen fluxes were very low at the reference site, and 1-2 orders of magnitude higher at all reduced sediment sites, with the highest rates associated with the black sediments of the Pockmark area (Table 2).



**Fig. 4** Sulphate and chloride profiles measured in porewater extracted from sediment cores sampled (a) at the reference site off Amon MV, (b) in reduced sediments on the Amon MV and, (c) in bacterial mats covering reduced sediments in the Pockmark area. The measurements were performed during the BIONIL cruise (2006)

3.3. Macro- and meiofaunal community description

3.3.1. Composition, abundance, density and  $\alpha$ -diversity patterns

Mean macrofaunal densities varied from 650 to 2,100 individuals m<sup>-2</sup> at Amon (Table 3), 1,950-3,500 individuals m<sup>-2</sup> in the Pockmark area (Table 43) and from 3,200 to 5,250 individuals m<sup>-2</sup> at Cheops (Table 4). This represents approximately 3.5%, 55%, and 78% of the total fauna sampled at each site (Tables 5, 6). In comparison, the fully oxic reference site had from 541 to 1,081 macrofaunal individuals m<sup>-2</sup> (Table 3), which represented 67% of the total fauna. On the carbonate crust microhabitat (CC), densities varied from 1,852 to 7,353 individuals m<sup>-2</sup> at Amon (Table 3) and from 1,852 to 7,353 individuals m<sup>-2</sup> in the Pockmark area (Table 4) representing respectively 100% (except CC2 where it only represent 20%) and 63% of the fauna sampled (Tables 3, 4).

Despite the large sieve mesh size used (250  $\mu$ m), many specimens corresponding to meiofaunal groups (copepods, ostracods, nematodes, mites) were found in our samples (Tables 5, 6). Mean densities of meiofauna >250  $\mu$ m varied from 31,800 to

75,550 individuals m<sup>-2</sup> at Amon (Table 5), 1,450 to 3,950 individuals m<sup>-2</sup> in the Pockmark area (Table 6) and reached 600 to 2,850 individuals m<sup>-2</sup> at Cheops (Table 6). At Amon, meiofaunal mean density was higher in reduced sediments compared to carbonate crusts (Table 5). Overall, nematodes dominated the meiofauna >250μm samples, varying from 95 to 100% of total abundances on the reduced sediments from the Amon and Cheops (Tables 5, 6). However, in the reduced sediments of the Pockmark area, nematodes represented only 36% of the meiofauna (Table 6). Here, harpacticoid copepods reached up to 37.5% of the meiofaunal abundance, followed by ostracods (~16%). Surprisingly, besides four nematodes, no meiofauna was sampled from the CC microhabitat of Amon, compared to a relatively high number of nematodes found at the CC of the Pockmark area (Tables 5, 6).

Relative faunal abundances varied between microhabitats and between replicate samples of the same microhabitat, especially on carbonate crusts. Overall, the steady increase of the individual-based rarefaction curves suggests that the sampling effort was not sufficient (Figure 5a). Only the curves of the reduced microhabitats of Cheops and the Pockmark area showed that we attained a relatively good estimation of their taxonomic richness. These curves also showed that the taxonomic richness between the different reduced sediment microhabitats was highest on Amon, followed by the Pockmark area and finally Cheops. The opposite trend was observed for the CC microhabitats, where the diversity was higher in the Pockmark area (Figure 5a). Since the ranking of the curves would probably remain the same with additional macrofaunal samples, we can conclude with confidence that the total family richness (S) on Amon was highest on reduced sediments, intermediate on the Reference site and lowest on CC, while the CC microhabitat in the Pockmark area was richer than the reduced sediments (Figure 5a; Tables 7, 8).

In terms of evenness (Pielou's index J', Table 8), the reference sample harbored the most even distribution followed by CC from the Pockmark area and reduced sediments from Amon (also shown in Figure 5b). Finally, Red from Cheops was the poorest microhabitat sampled in the NDSF. It was amongst the least even of all microhabitats along with CC from Amon and Red from Pockmark (Tables 7, 8). At Amon, polychaetes were the dominant taxa in the reduced habitats and reference samples, constituting respectively ~82% and 71% of total faunal abundance. A total

of 8 polychaete families or orders (since the Terebellida were not identified at the family level) were represented in the reduced sediments (Tables 3, 7). With ~11.5% of the total macrofaunal abundance, bivalves were the second dominant taxon in the reduced sediments of Amon, whereas chidarians (~18%) ranked second in the reference sample. Gastropods and crustaceans were present in low abundance (<6%) in reduced sediments while on the reference samples, low abundances were for the crustaceans and sipunculians. On CC from Amon, cnidarians represented 72% of the total abundance, distantly followed by polychaetes (~17%; Table 8). Some gastropods and sipunculians were also present in low abundance (<6%; Table 3a). In the Pockmark area, reduced sediments were dominated by 8 polychaete taxa, reaching ~84% of the total macrofaunal abundance, followed by bivalves and gastropods (Tables 4, 7). Contrary to Amon, the fauna from CC in the Pockmark area was more evenly distributed with gastropods and polychaetes representing respectively ~51% and 40% of the total abundance (Table 4). Bivalves and crustaceans were also present, representing less than 8% of the total macrofaunal abundance (Table 4). The rarefaction curves and the Pielou's index confirm the higher evenness of the CC from the Pockmark area compared to CC from Amon (Figure 5a; Tables 7, 8), the highest evenness being observed on the reference site (Table 8). Finally, on Cheops, polychaetes largely dominated reduced sediments, with a mean of 96% of the faunal abundance, followed by low proportions of crustaceans and gastropods (Tables 4, 7).

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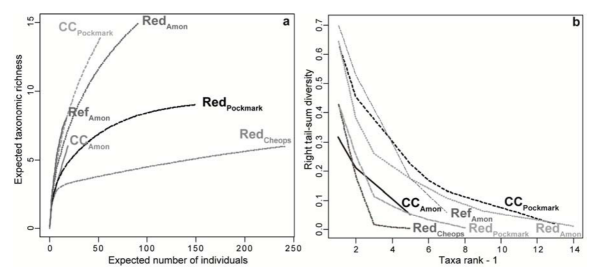
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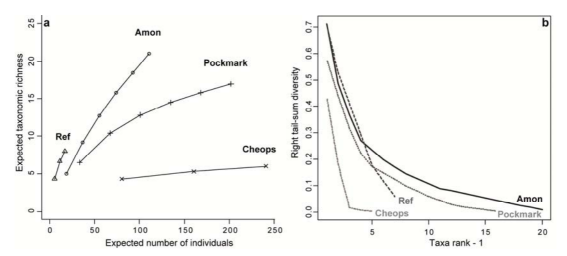
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**Fig. 5** (a) Rarefaction curves (b) and right tail-sum intrinsic diversity profiles for the pooled macrofaunal abundance data at the family level and at each microhabitat from the Amon MV (3 microhabitats including the reference sample), the Pockmark area (2 microhabitats) and the Cheops MV (1 microhabitat)

The RTS performed with the same dataset is difficult to interpret due to the crossing of the curves (Figure 5b). Using this analysis, Amon reduced sediments were the richest and most even microhabitat among the three reduced sediments. In contrast, the CC from the Pockmark area was more diverse than that from Amon (Figure 5b).

Finally, we pooled the data obtained for the different microhabitats to determine the diversity of each site, including the reference samples from Amon. For Amon and the Pockmark area, the sample-based rarefaction curves did not level-off, suggesting that sampling was insufficient to accurately estimate taxonomic richness (Figure 6a). At Cheops, despite the low sample number (n=3), the curve leveled-off, suggesting that its macrofaunal diversity was well described (Figure 6a). Our data indicate that diversity was higher at microhabitats sampled on Amon, followed by the Pockmark area, the reference samples from Amon and lastly, by Cheops (Figure 6a). Overall, the distribution of macrofauna was relatively even on Amon, since the most abundant taxon only reached ~30% for both active and reference sites, whereas the most abundant taxon represented ~45% and ~55% on the Pockmark area and Cheops respectively (Figure 6b).



**Fig. 6** (a) Rarefaction curves and (b) right tail-sum intrinsic diversity profiles for the pooled macrofaunal abundance data at the family level, according to the three study sites after pooling active microhabitats. Amon MV, n=6, Pockmark area n=3, and Cheops MV, n=3. The reference sample from Amon is reported separately, n=3 (Ref)

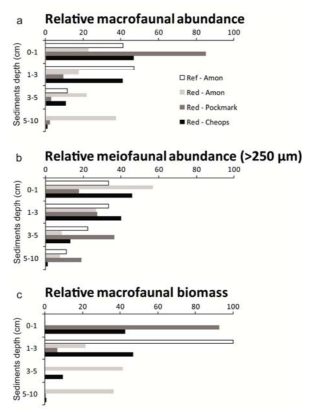
# 3.3.2. Symbiont-bearing fauna versus heterotrophic fauna

The symbiont-bearing fauna (for types of symbioses see Olu-Le Roy et al. 2004) represented 17 to 36% of the total faunal abundance in the reduced sediments from Amon, while the CC and the reference sample did not harbour any (Tables 7, 8). The symbiont-bearing fauna was represented by Frenulata polychaetes as well as by four bivalve species (*Lucinoma kazani*, *Idas modiolaeformis*, *Thyasira striata* and *Isorropodon perplex*; Table 3). The Pockmark area had between 10 and 16% of symbiont-bearing fauna both in the sediment and CC microhabitats, respectively (Tables 5, 7, 8). This fauna was represented by two known bivalve species (*L. kazani* and *I. perplexum*) and one unknown Lucinidae in the sediments. Only one bivalve species was present in the CC microhabitat (*I. modiolaeformis*; Table 4). No symbiont-bearing fauna was sampled at Cheops (Tables 7, 8) or at the reference site.

## 3.3.3. Vertical distributions within the sediments

The distribution within the sediment layers down to 10 cm below the seafloor of the macrofaunal relative abundances differed between the three reduced sediment microhabitats (Figure 7a). At all reduced sediment sites, oxygen did not penetrate deeper than a few mm, and sulfide concentration increased with sediment depth below 2 cm to 1-2 mmol I<sup>-1</sup> at Amon and to 10-20 mmol I<sup>-1</sup> in sediments of the Pockmark area. Hence, in the Pockmark area, the macrofauna was concentrated at the uppermost layer [0-1 cm] with about 85% of the total abundance (Figure 7a). However, some capitellid polychaetes and undetermined bivalves - that may be vesicomyids and lucinids - were found in the [5-10 cm] layers. In the Cheops area, the macrofauna was almost evenly distributed between the [0-1 cm] and [1-3 cm] layers and only few spionid polychaetes were found in the [5-10 cm] layer. The profile obtained for the reduced sediments of Amon was different, with a more homogeneous distribution within the 10 cm layer and a relatively high abundance at [5-10 cm] (Figure 7a) especially of frenulate, capitellid and dorvilleid polychaetes. At the reference site, most macrofauna was concentrated in almost equal proportions in the two first layers [0-1 cm] and [1-3 cm]. There, the distribution showed a clear decrease with depth, as no organism was found at the [5-10 cm] layer (Figure 7a). The meiofauna >250 µm also decreased with depth at the reference site, with up to 60% of the total meiofaunal abundance found in the two first layers (Figure 7b). The

same trend was observed in reduced sediments from Amon with this time almost 60% concentrated in the first layer. This pattern was different at the Pockmark area, where the meiofaunal abundance was highest in the [3-5 cm] layer followed by the [1-3 cm] and [5-10 cm] layers (Figure 7b). However, the few organisms found in the [5-10 cm] layer consisted of nematodes at Amon, and of harpacticoid copepods at the Pockmark area.



**Fig. 7** Vertical distribution of (a) macrofaunal and (b) meiofaunal (>250  $\mu$ m) relative abundances and (c) macrofaunal relative biomass with depth after summing the three replicates (n=3) for each sediment layer of the reduced sediment microhabitats from Amon MV, Pockmark area and Cheops MV. The reference site from Amon is also included

In general, all soft sediments showed a low macrofaunal biomass, except the reduced sediments of Amon where the mean biomass reached 0.12±0.1 kg pww m<sup>-2</sup> (Table 7). Here, the vertical distribution showed an increase with depth from 1 to 10 cm with about 40% of the total macrofaunal biomass remaining in the [5-10 cm] layer (Figure 7c) due to the presence of few lucinid and thyasirid bivalves. In the reduced sediments of the Pockmark area, most of the macrofaunal biomass (93%) was observed within the top layer [0-1 cm] and mainly consisted of dorvilleid polychaetes. Reduced sediments at Cheops hosted a similar biomass in the top two layers [0-1]

and [1-3 cm], with 43 and 47% of the total biomass respectively mainly represented by hesionid, spionid, terebellid polychates and Leptostraca crustaceans. Next, a sharp decrease was observed with depth (Figure 7c). At the reference site, mean biomass was very low (Table 8), and integrally located within the [1-3 cm] layer (Figure 7c) due to the presence of one sipunculian and few spionid polychaetes.

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# 3.3.4. Beta diversity patterns

The first two components of the principal component analysis (PCA) on macrofaunal data accounted for 41.7% of the variance in macrofaunal distribution (Figure 8a). Three clusters appear in reduced space: the carbonate crust samples; the reduced sediment samples except those from Cheops; and the Amon reference site and Cheops reduced sediments samples. Thus the variability between microhabitats was higher than that within the microhabitats, but lower than that among the geostructures (Figure 8). Polychaete taxa (Spionidae, Terebellida, Dorvilleidae, Capitellidae) and gastropods (Orbitestellidae) had the greatest impact on the variation in community structure. Focusing on these taxa, they appeared to be indicative of different groups, representing different types of microhabitats rather than different sampling locations. In the first group, Spionidae and Terebellida highly contributed to the positioning of the reduced sediments from Cheops and reference samples from Amon. In the second group, Orbitestellidae highly contributed to the positioning of both carbonate crust microhabitats (Pockmark and Amon). Finally Capitellidae and Dorvilleidae presented high contributions in reduced sediments from Amon and Pockmark area (Figure 8a). These three groups were also distinguished on the Ward's cluster (Figure 8b). According to the datasets, the pooling of both carbonate crust microhabitats seemed to be due to their low number of individuals (n=18 and 53) rather than to the presence of shared taxa (Tables 3a, b). When regarding the similarity level defined by the dotted line, the Ward's cluster also showed that the reduced sediments from Cheops had higher similarities with the carbonate crust microhabitats than with the other reduced sediment "soft" microhabitats (Figure 8b).

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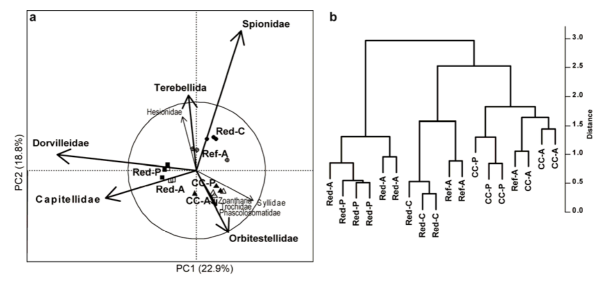
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A PCA with Hellinger-transformed meiofaunal >250  $\mu$ m data only (not shown) exhibited the same general distance patterns as the one with macrofauna *sensu stricto* (Procruste test stat=0.53, p=0.013, 1000 permutations). However, meiofaunal

sampling was very incomplete, especially on hard substratum, and these results might reflect this paucity of observations.





**Fig. 8** (a) Principal Component Analysis (PCA, scaling type 1) of Hellinger-transformed macrofaunal abundances on the Amon MV (A), Pockmark area (P) and the Cheops MV (C). The first two axes represent 41.7% of the total variance in macrofaunal abundance. The circle of equilibrium indicates the significant contribution of five taxa (radius=0.65). Vectors shorter than 0.34 were removed. (b) Ward's hierarchical clustering was performed with Hellinger-transformed macrofaunal abundances for each microhabitat types. Red: reduced sediments, CC: carbonate crusts, and Ref: reference sample

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## 4. Discussion

This study investigated the differences in faunal diversity in relation to different habitats and their biogeochemical conditions at three cold seep sites of the NDSF. Due to inevitably limited submersible time, sampling effort was relatively low given the high heterogeneity of cold seeps sites. Nevertheless, our results provide the first insights on seep faunal composition and diversity and their relationships with environmental conditions in the NDSF area.

# 4.1. Difference between the reference site and cold seep habitats at Amon MV

The reference site did not show any evidence of seepage whether direct (i.e. no detection of methane and sulphide fluxes) or indirect (i.e. absence of symbiont-bearing fauna). The fauna found at the reference site was different from that of the reduced habitats and carbonate crusts from Amon. Densities of macro- and meiofauna were two-fold lower than in reduced sediments, and total benthic oxygen

consumption was an order of magnitude lower. Accordingly, biomass was 1-4 orders of magnitude lower than at all other active sites from Amon. The reference site showed a relatively high evenness, but a lower diversity than the reduced sediment sites of Amon, therefore refuting our initial hypothesis. The reduced sediment microhabitats sampled within the present study support an overall higher faunal diversity compared to close-by oxygenated habitats. This pattern has already been observed on Eel River at about 500 m depth where diversity on clam beds was similar or higher than on non-seep sites (Levin et al. 2003; Levin et al. 2010). However, this is in contradiction with other studies from 770 to 3200 m depth were diversity appears to be lower at active seep sites (Sahling et al. 2002; Levin et al. 2010; Menot et al. 2010)

Comparing among sites, the relationship between seepage activity and diversity could best explained by a hump-shaped curve. We suggest that at the relatively low sulphide concentrations encountered at Amon the reduced sediments could be considered as an "ecotone" enhancing diversity by presenting a high variability of chemical environment and ecological niches, favoring the establishment of symbiont-bearing fauna but not limiting heterotrophic fauna by toxic concentrations of sulphide (Cordes et al. 2010). In contrast, the Pockmark area or similarily active seeps such as Hydrate ridge (Sahling et al. 2002) appear to show declined diversity, despite the high abundances/biomasses observed, potentially selecting for specialists able to withstand high sulfide concentrations and fluxes.

## 4.2. Reduced sediment microhabitats

The reduced sediment microhabitats were influenced by fluid emissions as attested by biogeochemical conditions. Sulphide concentrations at 12 cm below the seafloor in the Pockmark area were 10 times higher than what was observed on Amon for the same sediment depth. This flux was associated with a higher oxygen uptake and faster consumption at the water-sediment interface, suggesting higher microbial activity within reduced sediments of the Pockmark area compared to all other sites. According to a previous study (Girnth et al. 2010), some evidences of brine overflowing the sediments were recorded at Amon reduced sediments. However, the homogeneity of the chloride profiles at both sites does not support the presence of upward seepage of saline fluids as observed in some areas of Cheops.

Biogeochemical measurements indicate a ranking of activity with the Pockmark reduced sediments as the most sulphidic habitat, followed by Amon and Cheops.

Accordingly, the reduced sediment microhabitats at Amon and the Pockmark area were colonized by symbiont-bearing species, including siboglinid polychaetes and bivalves (Vesicomyidae, Lucinidae, Mytilidae and Thyasiridae) as well as by dorvilleid and capitellid polychaetes. The presence of these two families provides direct evidence for fluid emissions as they are usually associated with areas rich in organic matter and with sulphidic environments (Rouse and Fauchald 1997). Dorvilleid polychaetes also dominate reduced sediment microhabitats from other seep sites at 1100 m depth in the Marmara Sea (Ritt et al. 2010), at 2020 m depth in the Mediterranean Sea (Amsterdam MV, Ritt et al., in prep.) and at 500 m depth at the Eel River off California (Levin et al. 2003). Large nematodes were abundant in the reduced sediments of Amon, as previously observed on other mud volcanoes at 1220 (Van Gaever et al. 2006) and 5000 m depth (Olu et al. 1997).

The fauna in the reduced sediment microhabitat from Amon hosted the highest abundance of symbiont-bearing species, and its macrofaunal taxonomic richness was two times higher than in the Pockmark area. The bivalve species observed in the Pockmark area either harbored sulphide-oxidizing endosymbionts such as *Lucinoma kazani* and *Isorropodon perplexum* (Salas and Woodside 2002; Olu-Le Roy et al. 2004) or different types of symbionts as in the undetermined species *Idas sp.* Med, (Duperron et al. 2008).

On Cheops, the reduced sediments sampled were located about 250 m away from a brine lake which features extreme physico-chemical conditions. Methane concentrations at the surface of the lake varied from 2.4 to 3.7 mmol  $\Gamma^1$  with temperatures of 20-40°C and salinities 210 to 244 g  $\Gamma^1$  (Mastalerz, Harmegnies, pers. com.). Sediment cores revealed a dark layer covering a layer of brown hemipelagic sediments, suggesting the occurrence of a relatively recent sulphidic mud flow. Sulfate reduction rates and oxygen consumption rates were considerably lower than at the other sites, indicating a low sulphide flux, as confirmed by pore water measurements. Accordingly, despite the detection of elevated methane concentrations in the bottom waters of Cheops above the reduced sediments, no

symbiont-bearing fauna was observed. This microhabitat was dominated by spionid (~2200 individuals m<sup>-2</sup>) or terebellid (~660 individuals m<sup>-2</sup>) polychaetes, commonly found in sandy or muddy environments, from intertidal habitats to abyssal depths (Rouse and Pleijel 2001). Dense Terebellida beds, especially the Ampharetidae family have been observed at about 1100 m depth on the New Zealand margin (Sommer et al. 2009) and in the Marmara Sea (Ritt et al. 2010), as well as at 770 m at Hydrate Ridge off Oregon (Levin et al. 2010) where methane concentrations in the bottom water reach up to 2 mmol l<sup>-1</sup> and 0.7 μmol l<sup>-1</sup>, respectively for the first two first. At Cheops, methane concentrations varied between 4 and 8 μmol l<sup>-1</sup>. This suggests that terebellids are able to inhabit a wide range of environmental conditions. Further chemical measurements are needed to have a better understanding of the intriguing processes occurring within reduced sediments on Cheops.

## 4.3. Carbonate crust microhabitats

As expected, the difference in substratum between the soft sediment and carbonate crust microhabitats appears to influence the composition and distribution of the seep fauna in the NDSF. Nevertheless, the clustering of the CC microhabitats from Amon and the Pockmark area appears to be due to the low number of sampled individuals rather than the presence of common taxa. Sampling efficiency on hard substrata does not appear to be ideal as suggested by the high heterogeneity between replicates and the low Jaccard's similarity coefficients obtained. However these results may reflect the real heterogeneity of this habitat in terms of faunal composition and spatial distribution. Visual observations showed that the carbonate crusts sampled in the Pockmark area were associated with reduced sediments whereas those from Amon did not appear to be located in the vicinity of an active area. In addition, the crusts showed different facies and colour that may reflect different fluid intensities and stages of evolution (Bayon et al. 2009a). For example, the carbonates from the south-western part of Amon are thick, dark and cemented, and can reach 1 m of thickness. They may have been formed in the distant past, during a period of intense fluid emissions (Dupré et al. 2007). The absence of symbiont-bearing fauna and the dark color of crusts — due to their exposure to oxygen-rich bottom water and their iron and manganese oxide cover — corroborate the hypothesis that these carbonates may be relatively old. Fluid emissions may be very low on this part of Amon, explaining the absence of symbiont-bearing fauna. The

low faunal densities and biomass further support the fact that this environment is not favorable, neither to symbiont-bearing fauna, nor to heterotrophic fauna.

The light-colored and crumbly carbonates from the Pockmark area differed morphologically from the "inactive" crusts found on Amon as they seem to be relatively young. The precipitation of authigenic carbonates is controlled by high alkalinity of pore waters (Aloisi et al. 2002; Michaelis et al. 2002) due to the activity of consortia of archaea and bacteria involved in the anaerobic oxidation of methane (AOM) coupled to sulphate reduction (Boetius et al. 2000). Another by-product of AOM is sulphide which may support sulphide-oxidizing symbionts of chemosynthetic fauna such as the siboglinid polychaetes found beneath the crusts. In addition, the methane concentrations were high, reaching >5  $\mu$ mol  $\Gamma^1$ . Siboglinids may contribute to bio-irrigation and favor advection and diffusion processes that play a significant role in carbonate precipitation (Bayon et al. 2009a).

# 4.4. Comparison between Amon, the Pockmark area and Cheops

The reduced sediment microhabitats appeared to be distributed along a gradient related to seepage intensity and thus, to methane, oxygen and sulphide fluxes; with the Pockmark area as the most intense seepage site. The fauna from the reduced sediment microhabitats of the Pockmark area and Amon were rather similar and highly influenced by the presence of symbiotic organisms as well as dorvilleid and capitellid polychaetes, usually associated with organically enriched, reduced environments (Rouse and Pleijel 2001). Both communities showed relatively similar oxygen consumption rates. The taxonomic diversity was higher on Amon followed by the Pockmark area and Cheops. However, the reduced sediments, associated with the debris-flow of the Pockmark area, were more sulphidic and probably also more stable than the fresh Amon mud flow. The lower diversity and biomass observed in the reduced sediments of the Pockmark could be due to the high level of sulphide flux limiting the survival of most benthic species.

At the time of sampling, the mud flow at Cheops was probably recent as only a very thin sulphidic horizon was observed on top of hemipelagic sediments. These conditions may explain the lower diversity and the lack of symbiont-bearing fauna on the sampling site located 250 m away from the central brine lake. According to video

observations, instability seems to decrease from the summit toward the periphery as suggested by the presence of carbonate crusts with symbiont-bearing mytilids at about 700 m away from the lake.

4.5. Comparison at a larger scale

In terms of seepage activity, Amon is comparable to the Håkon Mosby MV (1200 m depth) from the Norwegian margin and the Amsterdam MV (2020 m depth) located on the Mediterranean Ridge. These MVs discharge mud, fluids and gases from their summits, which undergo episodic mud eruptions (Zitter et al. 2005; Dupré et al. 2007; Feseker et al. 2008). This activity induces instability and chemical gradients that influence the distribution of fauna, which is concentrically distributed around a chaotic summit (Zitter et al. 2005; Jerosch et al. 2007). The reduced site sampled at Amon was located on the base of the mud volcano, and influenced by lateral brine and mud flows (Girnth et al. 2011). Interestingly, the dimensions of the reduced sediment site was rather small (ca. 250 m) and the next larger reduced sediment patches were >2 km away.

The carbonate crusts and reduced sediments of the Pockmark area from the NDSF showed similarities with a more recently discovered site, the giant REGAB pockmark in the Gulf of Guinea at 3160 m depth. There, carbonate crusts are colonized by dense mussel beds and siboglinid assemblages (Olu-Le Roy et al. 2007). Pockmark and carbonate crust areas have also been observed at 740 m depth on the Storegga slide and the Nyegga area located on the north-west of the Norway margin which have undergone slide events. There, benthic fauna is represented by Siboglinidae, Crinoidae, Pycnogonidae and microbial mats (Hovland et al. 2005; Hovland and Svensen 2006; Paull et al. 2008). However, these features are taller than those observed in the central province of the NDSF as they reach 190 m long and 40 m wide, while pockmarks do not exceed a few meters in diameter at the NDSF study site.

The Cheops MV share similarities with the Napoli MV located on the Mediterranean Ridge at 1900 m depth, as they both harbor brine seepages (Charlou et al. 2003). However, faunal density and taxonomic richness on Cheops are lower than those observed on Napoli (Ritt et al., in prep). Faunal composition and distribution may be

strongly linked to the activity of the MV. Mud flows may initiate higher sedimentary and chemical instabilities on Cheops, limiting the colonization of benthic species. Although considered as an extreme habitat, the vicinity of brine lakes can be colonized by dense colonies of Porifera as observed on Napoli (Ritt, pers obs) or by dense mussel communities, such as those found on the shoreline of brine pools in the Gulf of Mexico at 650 m depth (Macdonald et al. 1990; Smith et al. 2000). The absence of dense assemblages of symbiont-bearing fauna close to the brines on Cheops was thus unexpected and may be explained by the effect of recent disturbances.

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In conclusion, our comparative investigation of active cold seeps on the NDSF suggested that seepage activity substantially enhanced benthic activity, biomass and diversity compared to the surrounding oxygenated and oligotrophic deep-sea environments. The biomass and biogeochemical activity of seep associated communities is 1-2 orders of magnitude higher than the surrounding and diversity was increased by a factor of 1.5. Reduced sediment microhabitats exhibited highest biomasses and diversity in comparison with surrounding oxygenated areas (reference site). As expected, reduced sediments and carbonate crusts were characterized by distinct faunal composition and faunal similarities were observed within each microhabitat type regardless of the site and geological structure (i.e. mud volcano, pockmark). However, our heterogeneous results on carbonate crusts require more investigation and emphasize the importance of developing a specific sampling tool dedicated to hard substratum in the deep sea. No simple relationship was detected between chemical conditions, sediment instability, fluid intensity and community structure although there seems to be a gradient related to seepage intensity between the different sites (Pockmark>Amon>Cheops). investigations with an appropriate sampling strategy, especially regarding the meiofauna, may help in highlighting the links between faunal distribution and environmental conditions in the NDSF. Most likely, temporal dynamics in such active geological systems may be very important in structuring community diversity. It could be studied in the future by deploying autonomous video cameras and sensors as well as by regularly monitoring a same site. Time-series data would help determine the response of the fauna to instabilities and disturbances induced by the occurrence of mud flows or brine seepage, and in which ways local diversity is related to temporal fluctuations and stability of habitats.

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**Table 1**. Location, depth, and tools used to perform physico-chemical and faunal sampling at each microhabitat from the different sampling sites of the Nile Deep-Sea Fan explored during the BIONIL (2006) and MEDECO (2007) cruises. Sampling effort, length of each sediment cores and estimated surface of each piece of carbonate crusts are also reported. Further details of samples are archived in http://www.pangaea.de/PHP/CruiseReports.php?b=HERMES

Microhabitat types	Latitude (°N)	Longitude (°E)	Depth (m)	Physico-chemical measurements	Faunal sampling
Reference site [ca. 15	5 km away fr	om Amon MV	(BIONIL, 2	2006)]	
Reference site (Ref)	32°21.42'	31°32.50'	1000	3 tubes (multicorer) M70/2b_785	3 tubes (74 cm <sup>2</sup> each) Ref1 (20 cm) Ref2 (20 cm) Ref3 (20 cm)
Eastern Province – A	mon MV (BIC	ONIL, 2006)			
Reduced sediments (Red)	32°22.05'	31°42.27′	1154	Microsensor , chamber and porewater samples PC15, 46, 47 M70/2b_765 (D115) M70/2b_790 (D121)	3 blade cores (200 cm² each) Red1 (BCROV-2, 10 cm) Red2 (BCROV-3, 10 cm) Red3 (BCROV-8, 10 cm)
Carbonate crusts (CC)	32°22.05'	31°42.27'	1153	None	3 pieces of crust CC1 (SFS-11, 84 cm²) CC2 (SFS-111, 103 cm²) CC3 (SFS-2, 139 cm²)
Central Province – Po	ockmark area	a, site 2A (BIO	NIL 2006)		
Reduced sediments (Red)	32°32.01′	30°21.13′	1697	Microsensor, chamber and porewater samples PC9, 52 M70/2b _784 (D120) M70/2b _841 (D127)	3 blade cores (200 cm² each) Red1 (BCROV-3, 20 cm) Red2 (BCROV-7, 17 cm) Red3 (BCROV-8, 20 cm)
Carbonate crusts (CC)	32°32.00′	30°21.18'	1696	None	3 pieces of crust CC1 (SFS-5, 48 cm²) CC2 (SFS-7, 81 cm²) CC3 (SFS-8, 34 cm²)
Western Province – C	Cheops MV (I	MEDECO, 200	7)		
Reduced sediments (Red)	32°08.05'	28°09.67'	3007	3x2 water samples: CH₄ MEDECO2_D343-PEPITO A-1 and PEPITO A-2, A-2, B-1, B- 2, C-1 and C-2	3 blade cores (200 cm² each) Red1 (BL-2, 20 cm) Red2 (BL-4, 19 cm) Red3 (BL-6, 17 cm)

**Table 2.** Physico-chemical characterization of the bottom water environment at the three study areas. Only the methane data from the Cheops MV were obtained directly above the organisms in this study. Amon data were provided by Girnth et al. (2011), Pockmark data by Grünke et al. (in press).  $\Sigma S = Total$  dissolved sulphides ( $H_2S+HS^2+S^2$ ), SR = Sulphate reduction rate. The reference site was ca. 15 km away from the active centre of the Amon MV.

	рН	[O <sub>2</sub> ] Bottom water (µmol l <sup>-1</sup> )	[O <sub>2</sub> ] Penetration depth (cm)	[O <sub>2</sub> ] Total benthic consumption (mmol m <sup>-2</sup> d <sup>-1</sup> )	[SO <sub>4</sub> <sup>2-</sup> ] Porewater (mmol l <sup>-1</sup> )	[Cl <sup>-</sup> ] Porewater (mmol l <sup>-1</sup> )	[CH₄] Bottom water (μmol l <sup>⁻1</sup> )	[ΣS] Porewater (mmol Γ <sup>1</sup> )	[ΣS] Porewater top 5 cm/peak conc. (mmol l <sup>-1</sup> )	SR Sediments (mmol m <sup>-2</sup> d <sup>-1</sup> )
Amon MV Reference site (Ref)	8.20 <sup>a</sup>	230 ª	>4 <sup>a</sup>	< 1 a	31.4 ª	529 ª	0.0 <sup>a</sup>	0 a		< 0.2 ª
Reduced sediments (Red)	7.88 <sup>b</sup>	150-200 <sup>b</sup>	0.0-0.25 <sup>b</sup>	10-46 °	30.7-40.3 <sup>d</sup>	404-580 <sup>d</sup>	0.0 <sup>d</sup>	< 0.7 b	0.8/2.5 <sup>d</sup>	$0.5 \pm 0.2 \text{ (n=3)}^{d}$
Pockmark area Reduced sediments (Red)	8.11 <sup>b</sup>	230 <sup>e</sup>	0.1-0.2 <sup>b</sup>	156-174°	29.1-29.8 <sup>d</sup>	606-630 <sup>d</sup>	0.2-0.3 <sup>e</sup>	0.2 <sup>b</sup> 0.7 <sup>d</sup>	0.8/25 <sup>d</sup>	22-41 <sup>d</sup>
Cheops MV Reduced sediments (Red)	-	220 <sup>e</sup>	-	111-130 °	-	-	3.60-9.34 <sup>f</sup>	-	-	5-14 <sup>d</sup>

Devices used to acquire the data: (a) multicorer, (b) microprofiler, (c) benthic chamber, (d) PCs, (e) KIPS bottle, (f) PEPITO, (-) no available data

**Table 3**. AMON - Macrofaunal (>250  $\mu$ m) densities (individuals m<sup>-2</sup>) per replicate and relative abundance (%) of each taxa in the microhabitats studied on the Amon mud volcano: reduced sediments (Red, n=3), carbonate crusts (CC, n=3) and the reference site (Ref, n=3). Total densities and relative abundances from each taxonomic group are highlighted in bold. Und. = undetermined individuals. (\*) Taxonomic level used for alpha-diversity analyses, here mostly family level. All sampling was performed during the BIONIL cruise (2006).

Taxonomic groups	F	Carbonate crusts				Reference samples						
	Red1	Red2	Red3	%	CC1	CC2	CC3	%	Ref1	Ref2	Ref3	%
Cnidaria (Total)	0	0	0	0	0	0	935	72.21	0	270	135	17.65
Anthozoa – Zoantharia*	0	0	0	0	0	0	935	72.21	0	0	0	0
Medusozoa – Scyphozoa*	0	0	0	0	0	0	0	0	0	270	135	17.65
Polychaeta (Total)	1900	400	1650	82.29	119	0	144	16.67	541	405	676	70.59
Capitellidae*	1100	100	50	26.04	0	0	0	0	0	0	0	0
Dorvilleidae*	300	50	1250	33.33	0	0	0	0	135	135	0	11.76
Hesionidae*	0	0	50	1.04	0	0	0	0	0	0	0	0
Paraonidae*	0	0	0	0	0	0	0	0	135	0	0	5.88
Pholoidae*	0	0	0	0	0	0	144	11.11	0	0	0	0
Phyllodocidae*	100	0	0	2.08	0	0	0	0	0	0	0	0
Spionidae*	100	0	50	3.13	0	0	0	0	135	135	405	29.41
Siboglinidae, Frenulata*	300	50	200	11.46	0	0	0	0	0	0	0	0
Syllidae*	0	0	0	0	119	0	0	5.56	0	0	270	11.76
Terebellida*	0	150	50	4.17	0	0	0	0	135	135	0	11.76
Larvae	0	50	0	1.04	0	0	0	0	0	0	0	0
Bivalvia (Total)	100	150	300	11.46	0	0	0	0	0	0	0	0
Lucinidae*												
Lucinoma kazani	0	150	50	4.17	0	0	0	0	0	0	0	0
Mytilidae*												
Idas modiolaeformis	50	0	0	1.04	0	0	0	0	0	0	0	0
Thyasiridae*												
Thyasira striata	0	0	100	2.08	0	0	0	0	0	0	0	0
Vesicomyidae*												
Isorropodon perplexum	0	0	150	3.13	0	0	0	0	0	0	0	0
Sareptidae*												
Und. Sareptidae	50	0	0	1.04	0	0	0	0	0	0	0	0
Gastropoda (Total)	0	50	0	1.04	0	0	72	5.56	0	0	0	0
Calliotropidae*												
Putzeysia wiseri	0	0	0	0	0	0	72	5.56	0	0	0	0
Und. Gastropoda*	0	50	0	1.04	0	0	0	0	0	0	0	0
Sipuncula (Total)	0	0	0	0	0	97	0	5.56	0	0	135	5.88
Phascolosomatidae*	U	U	U	U	U	31	U	3.30		U	133	3.00
	0	0	0	0	0	97	0	5.56	0	0	0	0
Phascolosoma aff. granulatum Golfingiidae*	0	U	U	0	U	91	U	5.56	: "	U	U	U
-	0	0	0	0	_	0	0	0	0	0	125	F 00
Nephasoma minutum	0		0	0	0			0	-	0	135	5.88
Crustacea (Total)	50	50	150	5.21	0	0	0	0	0	0	135	5.88
Decapoda-Thalassinidae*	0	0	50	1.04	0	0	0	0	0	0	0	0
Cumacea- Nannastacidae*									•			
Cumella pygmaea	0	0	50	1.04	0	0	0	0	0	0	0	0
Isopoda*	0	0	0	0	0	0	0	0	0	0	135	5.88
Leptostraca*	50	0	0	1.04	0	0	0	0	0	0	0	0

Total densitiy (ind. m <sup>-2</sup> )	2050	650	2100	119	97	1151	541	676	1081
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58									

Larvae

2.09

0 0

**Table 4.** POCKMARK area and CHEOPS - Macrofaunal (>250  $\mu$ m) densities (individuals m<sup>-2</sup>) per replicate and relative abundance (%) of each taxa in the microhabitats studied in the Pockmark area: reduced sediments (Red, n=3), carbonate crusts (CC, n=3) and on the Cheops MV: reduced sediments (Red, n=3). Total densities and relative abundances from each taxonomic group are highlighted in bold. Und. = undetermined individuals. (\*) Taxonomic level used for alpha-diversity analyses, here mostly family level. All sampling was performed during the BIONIL (2006) and MEDECO (2007) cruises.

Taxonomic groups				Pockma	ırk area				Cheops MV			
		Reduced			201		nate cru			Reduced		
	Red1	Red2	Red3	%	CC1	CC2	CC3	%	Red1	Red2	Red3	%
Polychaeta (Total)	2550	2650	1800	83.83	1042	123	4412	39.62	3800	5100	3000	95.58
Capitellidae*	450	650	250	16.16	208	0	0	1.89	0	0	0	0
Cirratulidae*	0	0	0	0	208	0	0	1.89	0	0	0	0
Dorvilleidae*	1350	1500	1350	50.30	208	0	0	1.89	0	0	50	0.40
Glyceridae*	50	0	0	0.60	0	0	0	0	0	0	0	0
Hesionidae*	500	350	200	12.57	417	0	588	7.55	950	1050	1050	24.50
Sabellidae*	0	0	0	0	0	0	2941	18.87	0	0	0	0
Spionidae*	100	100	0	2.40	0	123	0	1.89	2750	2750	1300	54.62
Syllidae*	0	0	0	0	0	0	294	1.89	0	0	0	0
Terebellida*	100	50	0	1.80	0	0	588	3.77	100	1300	600	16.06
Bivalvia (Total)	350	500	50	10.78	0	0	1176	7.55	0	0	0	0
Lucinidae*												
Lucinoma kazani	0	100	0	1.20	0	0	0	0	0	0	0	0
Und. Lucinidae	0	100	0	1.20	0	0	0	0	0	0	0	0
Mytilidae*												
Idas modiolaeformis	0	0	0	0	0	0	1176	7.55	0	0	0	0
Vesicomyidae*												
Isorropodon perplexum	100	150	0	2.99	0	0	0	0	0	0	0	0
Und. Bivalvia	150	150	50	5.39	0	0	0	0	0	0	0	0
Gastropoda (Total)	0	350	100	5.39	1667	1728	1471	50.94	0	50	50	0.80
Calliotropidae*												
Putzeysia wiseri	0	0	0	0	208	123	294	5.66	0	0	0	0
Trochidae*	0	0	0	0	208	123	588	7.54	0	0	0	0
Skeneidae*												
Akritogyra conspicua	0	0	0	0	0	123	0	1.9	0	0	0	0
Und. Skeneidae	0	100	0	1.20	0	0	0	0	0	0	0	0
Orbitestellidae*												
Lurifax vitreus	0	0	0	0	1250	1358	588	35.85	0	0	0	0
Und. Gastropoda	0	250	100	4.19	0	0	0	0	0	50	50	0.80
Crustacea (Total)	0	0	0	0	0	0	294	1.89	200	100	150	3.61
Amphipoda-Gammaridae					,	•						
Sebidae*												
Seba sp.	0	0	0	0	0	0	294	1.89	0	0	0	0
Und. Gammaridae	0	0	0	0	0	0	0	0	50	0	0	0.40
Leptostraca		v		J		J	Ū	3		v	v	5.10
Nebaliidae*	0	0	0	0	0	0	0	0	0	0	50	0.40
Und. Leptostraca	0	0	0	0	0	0	0	0	150	100	100	2.81
•			-			-	•	-			-	
Total density (ind. m <sup>-2</sup> )	2900	3500	1950		2708	1852	7353		4000	5250	3200	

**Table 5**. AMON - Meiofaunal (>250  $\mu$ m) densities (individuals m<sup>-2</sup>) per replicate and relative abundance (%) of each taxa in the reduced sediment microhabitat (Red, n=3) studied on the Amon and in the reference samples (Ref, n=3). Total densities and relative abundances from each taxonomic group are highlighted in bold. Und. = undetermined individuals. (\*) Taxonomic level used for alpha-diversity analyses, here mostly family level. All sampling was performed during the BIONIL cruise (2006).

Taxonomic groups		Reduced	sedimen			Carbona	ate crus		:	Refer	ence site	
	Red1	Red2	Red3	%	CC1	CC2	CC3	%	Ref1	Ref2	Ref3	%
Nematoda (Total)	74650	29900	28300	95.15	0	388	0	100	405	135	405	77.78
Crustacea (Total)	900	1900	4000	4.85	0	0	0	0	270	0	0	22.22
Copepoda-Harpacticoida												
Miraciidae*									-			
Bulbamphiascus imus	500	1050	200	1.24	0	0	0	0	0	0	0	0
Bulbamphiascus sp.1	0	0	0	0	0	0	0	0	0	0	0	0
Thyphlamphiascus confusus	0	0	550	0.39	0	0	0	0	0	0	0	0
Ameiridae*												
Amphiascus sp.2	0	0	0	0	0	0	0	0	0	0	0	0
Haifameira archibenthoica	0	400	2400	2.00	0	0	0	0	0	0	0	0
Haifameira sp.1	0	0	200	0.14					0	0	0	0
Agestidae*									!			
Eurycletodes sp.	0	0	0	0	0	0	0	0	270	0	0	22.22
Tisbidae*									:			
Tisbella sp.	0	0	0	0	0	0	0	0	0	0	0	0
Und. Harpacticoida	0	0	50	0.07	0	0	0	0	0	0	0	0
Copepoda-Cyclopoida												
Oncaeidae*									į			
Oncaea sp.	0	0	50	0.04	0	0	0	0	0	0	0	0
Copepoda-Calanoida*									-			
Calanoida sp.1	0	150	450	0.43	0	0	0	0	0	0	0	0
Calanoida sp.	0	300	0	0.21	0	0	0	0	0	0	0	0
Und. Calanoida	300	0	0	0.21	0	0	0	0	0	0	0	0
Ostracoda									-			
Polycopidae*									-			
Polycope sp.3M	50	0	0	0.04	0	0	0	0	0	0	0	0
Pontocyprididae*												
Propontocypris sp.2M	0	0	1	0.04	0	0	0	0	0	0	0	0
Propontocypris cf. levis	50	0	0	0.04	0	0	0	0	0	0	0	0
% Meiofauna / Total fauna	97.36	98.30	93.71		0	80.00	0		55.56	16.67	27.27	
Total density (ind. m <sup>-2</sup> )	75500	31800	32300		0	388	0		676	135	405	

**Table 6.** POCKMARK area and CHEOPS – Meiofaunal (>250 μm) densities (individuals m<sup>-2</sup>) per replicate and relative abundance (%) of each taxa in the microhabitats studied on the Pockmark area: reduced sediments (Red, n=3), carbonate crusts (CC, n=3) and on the Cheops mud volcano (MV): reduced sediments (Red, n=3). Total densities and relative abundances from each taxonomic group are given in bold. Und. = undetermined individuals. (\*) Taxonomic level used for alpha-diversity analyses, here mostly family level. All sampling was performed during the BIONIL (2006) and MEDECO (2007) cruises.

Taxonomic groups		Pockmark area							Cheops MV					
	F	Reduced	sedimer	nts		Carbon	ate crust		F	Reduced	sedimen	ts		
	Red1	Red2	Red3	%	CC1	CC2	CC3	%	Red1	Red2	Red3	%		
Nematoda (Total)	1100	500	1000	36.11	5417	247	2941	97.44	750	2850	600	100		
Crustacea (Total)	2850	1300	450	63.89	0	123	0	2.56	0	0	0	0		
Copepoda-Harpacticoida														
Miraciidae*														
Bulbamphiascus imus	1450	850	350	36.81	0	0	0	0	0	0	0	0		
Ameiridae*														
Amphiascus sp.2	0	0	0	0	0	123	0	2.56	0	0	0	0		
Tisbidae*														
Tisbella sp.	50	0	0	0.69	0	0	0	0	0	0	0	0		
Copepoda-Cyclopoida														
Cyclopina sp.	0	250	0	3.47										
Copepoda-Calaonida*														
Calanoida sp.1	100	0	0	1.39	0	0	0	0	0	0	0	0		
Calaonida sp.2	50	0	0	0.69	0	0	0	0	0	0	0	0		
Calaonida sp.3	50	0	0	0.69	0	0	0	0	0	0	0	0		
Calanoida sp.	250	0	0	3.47	0	0	0	0	0	0	0	0		
Ostracoda														
Pontocyprididae*														
Propontocypris sp.1M	50	0	0	0.69	0	0	0	0	0	0	0	0		
Propontocypris cf.levis	700	200	100	13.89	0	0	0	0	0	0	0	0		
Propontocypris cf. setosa	50	0	0	0.69	0	0	0	0	0	0	0	0		
Argilloecia sp.	50	0	0	0.69	0	0	0	0	0	0	0	0		
Chelicerata														
Und. Acarina*	50	0	0	0.69	0	0	0	0	0	0	0	0		
% Meiofauna / Total fauna	57.66	33.96	42.65		66.67	16.67	28.57		15.79	35.19	15.79			
Total density (ind. m <sup>-2</sup> )	3950	1800	1450		5417	370	2941		750	2850	600			

**Table 7**. RED - Biological descriptors of the reduced sediment microhabitats sampled on the Amon MV, the Pockmark area and the Cheops MV in the Nile Deep-Sea Fan. The highest values are highlighted in bold. The meiofaunal data includes only the large meiofauna >250  $\mu$ m. The number equivalent of Shannon and Simpson indices are given in italics

Biological descriptors	Amon	Pockmark area	Cheops		
Macrofauna -dominant	Polychaetes reaching 82.3% of total abundance (Dorvilleidae, Capitellidae)	Polychaetes reaching 83.8% of total abundance (Dorvilleidae)	Polychaetes, reaching 95.6% of total abundance (Spionidae, Hesionidae)		
Macrofauna -others	Bivalves, crustaceans, gastropods	Bivalves, gastropods	Crustaceans, gastropods		
Mean macrofaunal densities (individuals m <sup>-2</sup> )	1 600 ± 912	2 783 ± 782	4 150 ± 1 033		
Mean Jaccard's similarity	0.32	0.47	0.77		
Symbiont-bearing fauna	17.1 – 36.4%	0 - 10%	0%		
Total macrofaunal biomass (kg preserved wet weight m²)	1.2*10 <sup>-1</sup> ± 1*10 <sup>-1</sup>	$1.2*10^{-3} \pm 1*10^{-3}$	$6.4*10^{-3} \pm 3*10^{-3}$		
Macrofaunal diversity indices					
Total richness (S)	15	9	6		
Shannon (H <sub>e</sub> ') <i>Exp (H<sub>e</sub>')</i>	1.93 6.89	1.35 3.86	1.05 2.86		
Gini-Simpson (D <sub>GS</sub> ) (1 / 1-D <sub>GS</sub> )	0.78 4.62	0.63 2.68	0.59 2.44		
Evenness (J')	0.71	0.61	0.59		
Meiofauna (>250µm) -dominant	Nematodes with 95.2% of the total abundance	Copepods with 63.9% of the total abundance	100% nematodes		
Meiofauna (>250µm) -others	Copepods, ostracods (6%)	Acarina (1.3%)	-		
Mean meiofaunal densities (individuals m <sup>-2</sup> )	46 733 ± 25 004	2 400 ± 1 354	1 400 ± 1258		
Mean Jaccard's similarity	0.31	0.41	0.67		
Meiofaunal diversity indices					
Total richness (S)*	7	7	1		
Nematoda richmess	1	1	1		
Copepoda richness	4	3	0		
Ostracoda richness	2	2	0		
Acarina richness	0	1	0		
Shannon (H <sub>e</sub> ') <i>Exp (H<sub>e</sub>')</i>	0.24 1.27	1.39 <i>4.01</i>	0		
Gini-Simpson (D <sub>GS</sub> ) (1 / 1-D <sub>GS</sub> )	0.09 1.10	0.70 3.37	-		
Evenness (J')	0.12	0.71	-		

**Table 8**. CC and Ref - Biological descriptors of the carbonate crust microhabitats sampled on the Amon MV,in the Pockmark area and at the reference site (Amon MV) in the Nile Deep-Sea Fan. The highest values are highlighted in bold. The meiofauna data includes only the large meiofauna >250  $\mu$ m. The number equivalent of Shannon and Simpson indices are given in italics

Biological descriptors	Carbo	onate crusts	Reference site		
	Amon	Pockmark area	Amon		
Macrofauna -dominant	Cnidarians (Zoantharia) reaching 72.2% of total abundance	Gastropoda ( <i>Lurifax vitreus</i> ) and polychaetes (Sabellidae) in various proportions, reaching 50.9, 39.6% of total abundance	Polychaetes, reaching a mean of 70.6% of total abundance (Spionidae)		
Macrofauna -others	Gastropods, sipunculians	Bivalves, crustaceans	Cnidarians, sipunculians, crustaceans		
Mean macrofaunal densities (individuals m <sup>-2</sup> )	456 ± 602	3 971 ± 2 960	766 ± 281		
Mean Jaccard's similarity	0.0	0.31	0.34		
Symbiont-bearing fauna	0%	0 -16%	0%		
Total macrofaunal biomass (kg preserved wet weight m <sup>-2</sup> )	1.8*10 <sup>-3</sup> ± 5*10 <sup>-2</sup>	3*10 <sup>-2</sup> ± 5*10 <sup>-2</sup>	3.2*10 <sup>-5</sup> ± 6*10 <sup>-5</sup>		
Macrofaunal diversity indexes					
Total richness (S)	5	14	8		
Shannon (H <sub>e</sub> ') <i>Exp (H<sub>e</sub>')</i>	0.96 2.61	2.08 8.00	1.92 6.82		
Gini-Simpson (D <sub>GS</sub> = 1- $\lambda$ ) (1 / $\lambda$ )	0.46 1.84	0.81 5.31	0.83 5.90		
Evenness (J')	0.60	0.79	0.92		
Meiofauna (>250μm) -dominant	Only 4 individuals (nematodes)	Nematodes with 97.4% of the total abundance	Only 9 individuals (nematodes copepods)		
Meiofauna (>250µm)-others	-	Copepods (11%)	-		
Mean meiofaunal densities (individuals m <sup>-2</sup> )	129 ± 224	2 909 ± 2 523	-		
Mean Jaccard's similarity	-	0.67	-		
Meiofaunal diversity indexes					
Total richness (S)*	1	2	2		
Nematoda richness	1	1	1		
Copepoda richness	0	1	1		
Ostracoda richness	0	0	0		
Shannon (H <sub>e</sub> ') Exp (H <sub>e</sub> ')	-	0.12 1.13	-		
Gini-Simpson (D <sub>GS</sub> = 1- $\lambda$ ) (1/ $\lambda$ )	-	0.05 1.05	-		

Evenness (J') - 0.17 - 90 91 92 93 94