1	Changes in deep-water epibenthic megafaunal assemblages in relation to seabed slope
2	on the Nigerian margin
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5	Daniel O. B. Jones <sup>1*</sup> , Charles O. Mrabure <sup>2</sup> and Andrew R. Gates <sup>1</sup>
6	
7	* corresponding author
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9	<sup>1</sup> SERPENT Project, National Oceanography Centre, European Way, Southampton,
10	SO14 3ZH, UK.
11	
12	<sup>2</sup> Total E & P Nigeria Ltd., 37 Kofo Abayomi Street, Victoria Island, Lagos, Nigeria
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15	Abstract
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17	Local-scale habitat heterogeneity associated with changes in slope is a ubiquitous
18	feature of bathyal continental margins. The response of deep-sea species to high
19	habitat heterogeneity is poorly known and slope can be used as a proxy for many
20	important ecological variables, such as current flow, sedimentation and substratum
21	type. This study determines how slope angle effects megafaunal species density and
22	diversity at the Usan field, offshore Nigeria, between 740-760 m depth. This deep-
23	water area is increasingly exploited for hydrocarbons, yet lacking in baseline
24	biological information. Replicated remotely operated vehicle video transect surveys
25	were carried out using industry infrastructure (through the SERPENT Project) at a
26	representative range of slopes (1°, 3°, 11° and 29°). Twenty-four species of benthic
27	megafaunal invertebrate were found, numerically dominated by the echinoid
28	Phormosoma placenta, and nine species of fish were observed. Megafaunal
29	invertebrate deposit feeder density decreased significantly with increasing slope
30	(density range 0.503 - 0.081 individuals m <sup>-2</sup> ). Densities of megafaunal suspension
31	feeders were very low except at the highest slope site (mean density $0.17 \text{ m}^{-2}$ ). Overall
32	species richness was greater on steeper slopes, although the richness of deposit
33	feeders was not affected. Reduced labile organic matter in sediments on steeper slopes

- 34 likely reduced deposit feeder densities, but increased current flow at higher slopes
- 35 allowed both increased richness and density of suspension feeders.
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37 Keywords:

- 38 Deep-sea; slope; habitat heterogeneity; landscape; Phormosoma placenta; Gulf of
- 39 Guinea, Nigeria; video
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- 41
- 42 1. Introduction
- 43 44

45 Habitat heterogeneity at local scales is a ubiquitous feature of bathyal continental 46 margins and exerts a fundamental influence on the diversity and structure of benthic 47 communities (Levin and Dayton, 2009). The 'habitat heterogeneity hypothesis' is one 48 of the cornerstones of ecology (e.g. MacArthur and Wilson, 1967; Simpson, 1949). It 49 assumes that structurally complex habitats may provide more niches and diverse ways 50 of exploiting the environmental resources and thus increase species diversity (Tews et 51 al., 2004). Characterisation of the seabed in terms of terrain parameters, such as slope 52 and aspect, is a potentially valuable tool for classifying the structural complexity of 53 the seabed and delineating regions of seabed that may support particular fauna and 54 therefore provide a distinct habitat (Wilson et al., 2007). These techniques, broadly 55 classified as habitat mapping, have been successful in shallow waters (Lundblad et 56 al., 2006) and are receiving increased attention in deep water (Jones and Brewer, 57 2012; Wilson et al., 2007). Slope is a commonly used and important parameter in 58 these analyses, but direct assessment of changes in deep-water assemblages in relation 59 to seabed slope have not (to the author's knowledge) been assessed quantitatively. 60 Flat areas tend to exhibit different seabed facies and support communities that are 61 different to those inhabiting steeply sloping areas (Lundblad et al., 2006; Schlacher et 62 al., 2007). Slope, as with depth, is not a direct environmental factor that controls 63 biological communities; rather it is a useful proxy for correlated but unmeasured 64 factors. In the case of slope, these environmental factors range across the scale of 65 investigation but include a complex of environmental variables, primarily habitat 66 heterogeneity (Schaff et al., 1992), local current flow (Noble and Mullineaux, 1989; 67 Palardy and Witman, 2011), sedimentation, substratum type, food availability and 68 colonisation (Butman, 1990; Butman et al., 1994; Gage, 1997). Shallow water

69 research suggests that these factors may lead to increases in density (Sebens, 1985), 70 higher diversity (Palardy and Witman, 2011), elevated growth rates and changes in species distribution (Leichter and Witman, 1997) as well as functional changes in 71 72 benthic communities, for example favouring suspension feeding over deposit feeding 73 organisms (Jumars and Nowell, 1984). It is important to assess the role of slope-74 related factors in driving the structure of deep-water benthic faunal assemblages 75 (Levin et al., 2001; Tyler, 1995), particularly in areas of high habitat complexity such 76 as canyons (De Leo et al., 2010; Rowe et al., 1982) and seamounts (Clark et al., 2010; 77 Genin et al., 1986).

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79 By combining spatially-accurate quantitative seabed imaging techniques with high-80 resolution acoustic data on submarine topography and sediment properties fine-scale 81 patterns in megafaunal biology can be linked to the broad-scale patterns in habitat 82 type and subsea landscape (Baguley et al., 2006; Wilson et al., 2007). Given the sheer size and difficulties of accessing the deep-sea environment, accurate extrapolation of 83 84 fine-scale observations using information on the important physical controls for 85 faunal distribution is vital to describe the important broad-scale patterns in benthic 86 biology (Sanchez et al., 2008; Williams et al., 2010). Characterising variation in 87 biodiversity and biological assemblage structure in response to specific terrain 88 parameters, such as slope, are important to help understand the underlying 89 mechanisms behind these patterns.

90

91 The megabenthic communities of the offshore waters of the Gulf of Guinea region are 92 poorly known (Le Loeuff, 1993; Le Loeuff and von Cosel, 1998; Lebrato and Jones, 93 2009) compared to the Canary region to the north (Duineveld et al., 1993a; Duineveld 94 et al., 1993b; Henriques et al., 2002; Jones and Brewer, 2012; Keller and Pasternak, 95 2002; Merrett and Domanski, 1985; Merrett and Marshall, 1981). The Benguela 96 region to the south of the Gulf of Guinea has also received some attention, 97 particularly on the shelf and slope (Roeleveld et al., 1992; Roy et al., 2007; Uriz, 98 1988) as well as the abyss (Kroncke and Turkay, 2003; Levin and Gooday, 2001; 99 Saiz-Salinas, 2007; Thandar, 1999; Vinogradova et al., 1990; Zibrowius and Gili, 100 1990). The Gulf of Guinea region is heavily exploited for natural resources, primarily hydrocarbon resources (Zabanbark, 2002) and fish (FAO Fishery Committee for the 101 102 Eastern Central Atlantic, 1991) and is subject to high levels of anthropogenic impact 103 from pollution, mostly of terrestrial origin (Scheren *et al.*, 2002). The limited 104 environmental information is mostly collected by oil and gas companies and not 105 available in the scientific literature. However, it is crucial that baseline quantitative 106 environmental information is available for successful management of this increasingly 107 exploited ecosystem.

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109 This paper provides the first description of deep-water megabenthic assemblages at a 110 deep-water Nigerian site. Quantitative remotely operated vehicle (ROV) imaging allowed the effect of various seabed slopes on assemblage density, composition and 111 112 diversity to be assessed. Specifically, the following hypotheses were tested: 1) 113 increases in environmental heterogeneity associated with slope will increase 114 megafaunal species richness, 2) increased slope will favour suspension feeding 115 lifestyles at the expense of deposit feeding and hence change the relative abundances 116 of these functional groups and 3) the combination of these effects will lead to 117 differences in megafaunal assemblage structure between habitats with biologically 118 significant variation in environmental conditions resulting from changes in slope.

119

120 2. Methods

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124 This study focuses on a knoll at the west of the Usan field (within Nigerian offshore

125 oil prospecting license block 222), an area that will be developed for hydrocarbon

126 production, located on the Nigerian continental margin in the Gulf of Guinea.

127 Environmental baseline surveys (Total E. & P. Nigeria Ltd., 2007) in the main Usan

128 field (centred  $03^{\circ}$  31.800' N 007° 26.400' E; slope < 1°) collected boxcore samples of

seabed sediments. The silty clay sediment was comprised of silt ( $50.1 \pm 1.1$  %; mean

 $\pm$  standard deviation), clay (34.9  $\pm$  2.0%) and sand (15.1  $\pm$  1.0%), with an organic

131 content of  $5.8 \pm 1.3\%$  and a total organic carbon content of  $3.3 \pm 0.7\%$ . Sediments

- 132 contained abundant material or terrigenous origin (e.g. seeds, pollen, plant material).
- 133 Sediments supported a macrofaunal (>0.5 mm) assemblage with low density  $(69 \pm 53)$
- 134 individuals m<sup>-2</sup>) and diversity (57 species). The macrofauna was dominated by

135 molluscs (38.6%), arthropods (22.0%), protozoans (17.6%) and annelids (16.3%),

136 with the remainder comprised of echinoderms, cnidarians, poriferans and

- 137 hemichordates (Total E. & P. Nigeria Ltd., 2007).
- 138

139 Shipboard measurements of seawater collected near the seabed at Usan had dissolved 140 oxygen levels (mean = 5.84 mg L<sup>-1</sup> [54.7 % saturation], range = 4.41 [40.8 %] to 9.54 141 [88.4 %] mg L<sup>-1</sup>) that were not hypoxic (dissolved oxygen is < 0.7 mg L<sup>-1</sup> (Helly and 142 Levin, 2004)). *In situ* physical measurements of the near-bottom seawater (at 750 ± 143 10 m water depth) made during this study showed low temperatures (5.32 ± 0.07 °C; 144 mean ± standard deviation) and consistent salinities (34.76 ± 0.01).

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146 2.2. Survey Design

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Four habitats, each with different slope conditions (1°, 3°, 11° and 29°), were 148 149 assessed at Usan (Figure 1) through the SERPENT project (Jones, 2009). The slope 150 conditions were labelled 1°, 3°, 11° and 29°. For each slope condition an area was 151 chosen within the Usan field using detailed bathymetric charts (Figure 1) that fulfilled 152 the following criteria: constant slope, a depth between 740 and 760 m and a distance 153 of at least 1 km from any drilling activity to ensure no effect of disturbance (Jones et 154 al., 2006). Three replicate remotely operated vehicle (ROV) video transects (200 m 155 length) were performed at random locations within each slope condition (S1). Survey 156 start points and survey direction were identified using a random number generator in 157 Microsoft Excel. Transects were examined by plotting them onto the chart. Any transect that extended beyond the boundaries of identified areas was discarded and 158 159 another generated using the same method. In the area of highest slope headings were 160 constrained between 350 to 360°, 0 to 10° and 170 to 190° to maintain the depth range of transects but otherwise were random. Each transect was regarded as a 161 162 sampling unit (i.e. all organism densities were totalled for each 200 m transect) and 163 used in all subsequent analysis. Three replicate transects at four slope levels resulted 164 in a total of 12 sampling stations. All positional information was recorded in 165 Universal Transverse Mercator (UTM) zone 32 N based on the World Geodetic 166 Society 1984 datum.

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Data were collected using an Oceaneering Millennium ROV equipped with a colour video camera (Insite Pacific Pegasus), sonar and ultra-short baseline navigation transponder. Cameras were mounted on a pan and tilt unit at the front of the ROV, which enabled oblique video to be taken. Before each transect the video was zoomed out to maximum extent and the camera was set to its most vertical angle (30° below the horizontal).

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177 In every transect the ROV was run in a straight line, on a set bearing and at a constant speed ( $\sim 0.3 \text{ m s}^{-1}$ ). Vehicle altitude (distance from seabed to the base of the ROV) 178 was kept constant at 1 m, which equated to a 143 cm camera altitude (from the centre 179 of the camera). Transect width (mean of 1.72 m; max variation  $\pm$  0.2 m) was 180 181 calculated from the camera acceptance angles (Jones et al., 2006). The camera 182 horizontal acceptance angle (at full wide angle) was 55° and the vertical acceptance 183 angle 43°. Transect width was also verified by passing over objects of known size on the seafloor. The optical resolution of the cameras permitted all organisms larger than 184 185 50 mm to be reliably distinguished. In addition to the transects, previous ROV suction sampling at the nearby Akpo site (03°06.950' N 006°49.120' E) and a separate survey 186 187 to carry out detailed video inspection of selected individual organisms aided species 188 identifications (Jones et al., 2006).

189

190 2.4. Data Analysis

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192 Video transects were replayed at half speed and all visible organisms were counted along the entire transect. Fauna were identified to the highest taxonomic resolution 193 194 possible, in some cases to species. Megafauna that could not be identified to species 195 were grouped into morphologically distinguishable entities (referred to as taxa). It 196 should be emphasised that this approach may underestimate the number of species 197 present. Colonial organisms were counted as single individuals. Infaunal species were 198 counted if enough of their body was visible for identification. Benthic fish were 199 recorded but not included in subsequent quantitative analysis. Megafauna were 200 classified into deposit and suspension feeders based on the literature descriptions.

202 Four indices of diversity were calculated in order to assess both the dominance and 203 species richness aspects of diversity (Magurran 2003, Gotelli & Colwell 2001). These 204 were the number of taxa in a transect or group of transects (a simple measure of 205 species richness, S), rarefied species richness (estimated species in 60 individuals:  $ES_{(60)}$ , evenness (as measured by Pielou's evenness, J') and Shannon-Wiener species 206 diversity (H' base log e). Numbers of organisms were expressed as abundances 207 (numbers in each sampling unit) or as densities (no. m<sup>-2</sup>). Diversity indices were 208 calculated using PRIMER v6 (Clarke and Warwick, 2001). 209

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211 The structure of the megabenthic assemblage was visualised using multidimensional 212 scaling plots and sites compared using multivariate community analysis, specifically 213 PERMANOVA (Anderson, 2001) and Multivariate dispersion analysis (Anderson, 214 2006). Multivariate community analysis was based on abundances of all taxa, a fourth 215 root transformation being applied to buffer the influence of dominant taxa (Field et 216 al., 1982). Similarities were calculated using Bray-Curtis coefficients (Bray and 217 Curtis, 1957). The similarity values were subjected to both classification (hierarchical 218 group-average clustering) and ordination (non-metric multi-dimensional scaling) 219 using the software PRIMER v6 (Clarke and Warwick, 2001). PERMANOVA and 220 Multivariate dispersion analysis were carried out using the 'adonis' and 'betadisper' 221 functions respectively using the R package Vegan (Oksanen et al., 2011) in the R 222 programming environment (R Development Core Team, 2010).

223

224 Data were analysed following one-way analysis of variance (ANOVA) design with 225 slope as a fixed factor. This was implemented in the univariate case by generalised 226 linear models and in the multivariate case by one-way PERMANOVA (Anderson, 227 2001). Independent generalised linear models were developed (Dobson and Barnett, 228 2008) to examine whether the taxon richness (number of species in a transect) and 229 density (as numbers of epibenthic invertebrates in a transect) of both suspension and 230 deposit feeding taxa could be explained by the seabed slope (treated as categorical 231 data). The residual deviance for the count data (48.44 for suspension feeders; 97.26 232 for deposit feeders) was greater than the residual degrees of freedom (8 for both),

indicating overdispersion. The models were, therefore, fitted with quasi-Poisson
errors. The models for species richness, evenness and species diversity were fitted
with Gaussian errors. Models were fitted using the R function glm and the ANOVA
function of the R package CAR (companion to applied regression) (Fox and
Weisberg, 2011).

238

239 3. Results

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The seabed was almost entirely composed of fine grain-sized sediment, no individual grains were visible in video (Figure 2), even when fully zoomed in. This suggested that the largest possible grain size is fine sand (< 0.5 mm) but observations of resuspension indicate that it was finer and classified as mud (< 125  $\mu$ m). Hard substrata, composed of igneous rock (identified from samples), were visible within the areas of highest slope. These hard surfaces provided habitat for numerous specialist taxa (Table 1).

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249 3.1. Nature of the benthic community

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251 A total of 24 invertebrate megafaunal taxa representing six phyla were observed in the 252 video transects at Usan (Table 1, Figure 3). In terms of species richness and numerical 253 abundance, echinoderms were most abundant: 64% of the fauna were accounted for 254 by 10 echinoderm taxa. Although eight cnidarian taxa were identified, they only 255 represented 4% of the total density. Crustaceans were numerically important (31% 256 total fauna) but only represented by three taxa, one of which, a galatheid, accounted 257 for the vast majority of individuals. Of lesser importance were the poriferans, 258 molluscs and enteropneusts, all with only one observed taxon and together 259 representing <1% of the total invertebrates.

260

The regular echinothurid echinoid *Phormosoma placenta* was overwhelmingly dominant, particularly in the flatter areas, accounting for 53% of the invertebrates observed. Two other taxa were numerically important; the galatheid *Munidopsis* sp. represented 29% of megafauna and an unidentified large ophiuroid represented 8%.

265 The other taxa observed were each of low abundance (<3% of megafauna).

266

Although not considered further in this analysis, fish were observed at all the sites (S2). A small, dark-coloured, midwater gonostomatid fish was most abundant representing over 47 % of the fish fauna. The true benthic lophilform *Dibranchus* sp. was common and represented 27% of the fish. The other major taxon was the nettastomatid anguilliform *Nettastoma melanura* which accounted for 18% of the fish. The demersal grenadier *Corphaenoides* sp. and the morid gadiform *Laenomena laureysi*? each represented around 2% of the fish seen in the video records.

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275 3.2. Variation between habitats

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277 3.2.1. Faunal density

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279 Faunal density varied between habitats of different slope, with lower total densities in 280 areas of higher slope. When total density is split into more ecologically meaningful 281 units, based on feeding mode, the trends were even clearer (Figure 4). Deposit feeders 282 made up the majority of the megafauna observed; however, when analysed alone, 283 densities of deposit feeders decreased significantly with greater slopes (L-ratio = 284 16.09, d.f. = 3, p < 0.01). Densities of suspension feeders were very low except at the habitat with the highest slope; even here the density was less than half  $(0.17 \text{ m}^{-2})$  that 285 of deposit feeders (0.40  $\text{m}^{-2}$ ). The increase in deposit feeder densities with slope was 286 significant (L-ratio = 35.41, d.f. = 3, p < 0.001). 287

288

Of the major taxa, *Phormosoma placenta* had higher densities at low slope sites (Lratio = 16.11, d.f. = 3, p < 0.01) as did the galatheid *Munidopsis* sp. (L-ratio = 88.37, d.f. = 3, p < 0.001). Ophiuroid densities were significantly higher in areas of higher slope (L-ratio = 24.88, d.f. = 3, p < 0.001).

295

The slope-related differences in the habitat had a significant effect on the number of taxa found (Figure 4; L-ratio = 9.33, d.f. = 3, p < 0.05). Pairwise tests, however, only revealed significant differences between the habitat with the lowest slope (1°) and the highest slope (29°). The principal difference in the diversity of sites was caused by the presence of obligate epilithic species at the sites where hard substratum was exposed (see Table 1). In soft sediment areas there was some variation in species complement resulting from the apparently random presence or absence of rare species.

303

Species diversity indices (Figure 4) revealed that the species richness element of diversity ( $ES_{(60)}$ : L-ratio = 35.42, d.f. = 3, p < 0.001) and the evenness component (J' L-ratio = 66.14, d.f. = 3, p < 0.001) showed a highly significant positive linear relationship with increasing slope of the habitat. As would be expected, a combined index showed similar patterns with slope (H'; Figure 4). There were significant pairwise differences between all the habitats (p < 0.05) in  $ES_{(60)}$ , J' and H' except between 1 and 3 degrees in the combined Shannon-Wiener Index (p = ns).

311

Overall, multivariate community similarity was significantly different between 312 313 habitats of different slope (PERMANOVA: F = 9.55, d.f. = 1, 11, p < 0.001; Figure 314 5). Multivariate community similarity was high (> 60 %) and between-site dispersion low between the habitats with lower slope (1, 3 and 11°), whereas the habitat with 315 316 highest slope (29°) was more heterogeneous (Multivariate dispersion analysis: average distance to centroid 0.08 for 1°; 0.14 for 3°; 0.16 for 11° and 0.30 for 29°; 317 differences not significant in permutation test F = 2.57, d.f. = 3, p = 0.13). Two sites 318 319 (29B and 29C) formed a distinct group (with 77.5 % similarity) and one site (29A) 320 was intermediate between the highest and lower slope sites, having slightly higher 321 affinity with the lower slope sites (similarity 58.8%) than with the higher slope sites 322 (similarity 35.9%).

323

<sup>324 4.</sup> Discussion

326 4.1. General characteristics of the megafaunal assemblages at Usan

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328 Knowledge of West African megafauna is limited, making it difficult to compare the 329 assemblage of invertebrates found at Usan to those at other locations. However, the 330 common species are known and the limited evidence suggests they are abundant off 331 West Africa. *Phormosoma placenta* is a common, often dominant, species on both 332 sides of the Atlantic (Jones and Brewer, 2012; Laguarda-Figueras et al., 2005; 333 Sanchez et al., 2008). The large ophuiroid observed at Usan is very similar to a 334 species found off Mauritania (Jones and Brewer, 2012), Ivory Coast and Angola 335 (author's observation; http://archive.serpentproject.com/151/), while a galatheid 336 resembling the species observed in the present study has been seen in images obtained 337 off Ivory Coast (Lebrato and Jones, 2009). It should be noted that even generic 338 differences are not always evident in photographs. The rarer species quantified in our 339 survey, and those seen outside the transect areas, likely include new species. 340 However, standard-resolution video and image data in general is not usually sufficient 341 to resolve the details necessary for species-level taxonomy.

342

343 The density of the megafaunal assemblages at Usan seems to be low, but in the range 344 of those found elsewhere (Figure 6). The nearest comparative area of an equivalent 345 depth is located off Ivory Coast where megafaunal densities were over three times 346 greater than at our study site. However, the Ivory Coast site experienced significantly 347 enhanced nutrient inputs from pyrosome carcasses (Lebrato and Jones, 2009). Few 348 other data are available for the East Atlantic at equivalent depths. Studies conducted 349 in deeper areas reported a range of megafaunal densities (Figure 6) that are usually 350 lower than at the Usan site. Of the deeper studies, only those areas that underlie 351 highly productive surface waters, for example the eutrophic EUMELI site off 352 Mauritania (Galeron et al., 2000), have yielded comparable values.

353

4.2. Environmental consequences of differences in slope

356 Although four species were present at habitats with all the levels of slope investigated, 357 ten of the taxa found at the habitat with the highest slope (29) were confined to this 358 habitat. This habitat had exposed hard substrata and hence a unique community of 359 epilithic megafauna. Most of the epilithic organisms were not found on soft sediments 360 and many, for example gorgonians, had obvious morphological adaptations for 361 attachment to rocky surfaces. Hard substratum communities are probably rare on the 362 West African margin, having only been described off Mauritania where they are 363 associated with corals (Colman et al., 2005). Other studies of hard substrata near west 364 Africa are on the mid-Atlantic ridge at non-vent (Gebruk et al., 2010) and vent sites 365 (Van Dover et al., 2002), volcanic island groups, for example the Azores, or seamounts (Hall-Spencer et al., 2007). The megafaunal assemblage found on hard 366 367 substrata at Usan superficially resembles that found at non-vent hard substrata in the 368 northern tropical Atlantic (in terms of faunal groups present) and from hard substrata 369 in the deep sea in general (e.g. Lundsten et al., 2009). However, without accurate 370 species-level identifications either here or in the megafaunal literature generally, 371 further patterns are difficult to assess.

372

373 The large reductions in density of deposit-feeding species at Usan with increases in 374 habitat slope likely reflect reductions in food availability for deposit feeders. These 375 patterns are unlikely to be related to the presence of hard substratum at the steepest 376 slope site. Slope is expected to affect food availability in two ways. Firstly, increased 377 slope will probably increase the lateral advection of food particles, reducing their 378 residence times and hence availability for deposit feeders. Decreases in food 379 availability have been repeatedly shown to be responsible for reductions in benthic 380 megafaunal density in the deep sea (Galeron et al., 2000; Thurston et al., 1994). 381 Secondly, increases in slope are likely to enhance local current speeds and hence lead 382 to reductions in the deposition and increases in the winnowing of fine, nutritious 383 particles (Levin et al., 2001). Increased flow favours suspension feeding organisms 384 over deposit feeders (Jumars and Nowell, 1984) and actively enhances recruitment, 385 driving increased species density and elevated species richness (Palardy and Witman, 386 2011).

388 Increases in slope were associated with increased overall megafaunal diversity at 389 Usan despite a decrease in the number of deposit-feeding species in the habitat with 390 the steepest slopes. There was slightly (although not significantly) higher diversity, 391 and consistent changes in multivariate community composition, in areas of steep slope 392 without hard substrata. In the absence of other factors, reduced food availability tends 393 to lead to reductions in diversity (Levin et al., 2001; Waide et al., 1999), as 394 demonstrated in the tropical Atlantic deep sea (Cosson et al., 1997). Organic particles 395 are more likely to settle and become available for deposit feeders at low slopes 396 (Fontanier et al., 2008). At high slopes resuspension and current-mediated lateral 397 transport of organic material is likely (Schlacher et al., 2007; Van Gaever et al., 398 2009), which may favour a suspension-feeding lifestyle. The trends in diversity 399 observed at Usan are consistent with predictions based on food availability. Higher 400 levels of organic matter may result in reduced diversity in areas, such as oxygen 401 minimum zones, where the organic inputs are already high and associated with 402 oxygen depletion (Levin et al., 2001), which was not observed at Usan. In addition to 403 influencing the organic content of the sediment, increases in slope are also likely to 404 affect the substratum heterogeneity. Sediment grain size will likely become coarser 405 and potentially more heterogeneous, increasing niche availability and hence deposit 406 feeder diversity (Etter and Grassle, 1992; Wheatcroft, 1992), although this was not 407 clear in the present study. In the Usan area the sites with steep slope have exposed 408 hard substrata, presumably reflecting limited deposition. This increase in habitat 409 heterogeneity is responsible for the clear and significant increases in diversity and 410 changes in multivariate community composition at the steepest slope sites. An entirely 411 different range of animals is able to colonise hard substrata and hence, where this is 412 present, diversity is likely to increase (Levin et al., 2001).

413

Deep-sea biology requires more detailed local-scale studies of faunal distribution in relation to habitat features. Physical gradients in biological assemblages associated with slope occur in many ecosystems (de Castilho *et al.*, 2006); this study extends these findings to the African deep sea. Considerable areas of the ocean margins have features of high topography and sloping seabed, such as described here. This study has shown clear reductions in deposit-feeding megafaunal density with slope. If this trend is typical for deep-water habitats, it should be incorporated into model estimates of megafaunal densities, particularly on areas of elevated topography. Diversity of
benthic megafauna does not display such a clear relationship with slope and is likely
to be controlled by site-specific responses to changes in habitat heterogeneity and
food availability.

425

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Table 1: Invertebrate megafauna densities (no m<sup>-2</sup>) at each transect at Usan. Slope is represented by numbers:  $1 = 1^{\circ}$ ,  $3 = 3^{\circ}$ ,  $11 = 11^{\circ}$ ,  $29 = 29^{\circ}$ . Site letters represent replicates. Asterisked taxa (\*) are known to prefer hard substrata and double asterisked (\*\*) only live on hard substrata (obligately epilithic). Deposit feeders (D) and suspension feeders (S) are labelled. See figure 3 for images of the taxa.

	1A	1B	1C	3A	3B	3C	11A	11B	11C	29A	29B	29C
Hexactinellid**S	0	0	0	0	0	0	0	0	0	0	0.006	0.003
White												
antipatharian**S	0	0	0	0	0	0	0	0	0	0	0.047	0.061
Whip-like												
S	0	0	0	0	0.003	0	0.003	0.003	0.003	0	0	0
Whip-like												
cnidarian (rock,				0	0	0		0	0		0.000	0.000
White)** S	0	0	0	0	0	0	0	0	0	0	0.009	0.009
S	0	0	0	0	0	0	0	0	0	0	0.006	0
Gorgonian fan**												
S	0	0	0	0	0	0	0	0	0	0	0	0.009
Actinoscypha	0.002	0	0.002	0	0	0	0	0	0	0	0	0
Sp. Edwardsiid	0.003	0	0.003	0	0	0	0	0	0	0	0	0
anemone	0	0	0.003	0	0	0	0	0	0	0	0	0
Striped												
anemone	0	0	0.003	0.003	0	0.003	0	0	0	0	0	0
placenta D	0.343	0.395	0.302	0.140	0.233	0.352	0.201	0.009	0.012	0.090	0.023	0.081
Red echinoid	0	0	0	0	0	0	0	0	0	0.003	0.012	0.003
White echinoid	0	0	0	0	0	0	0	0	0	0.003	0	0.003
White Diadema												
sp.	0	0	0	0	0	0	0	0	0	0.003	0.009	0.006
Ophiuroid D	0.012	0	0	0.009	0.020	0.032	0.064	0.070	0.038	0.029	0.052	0.026
Brisingid* S	0	0	0	0	0	0	0	0	0	0	0.012	0.003
Holothurian D	0.012	0.003	0	0	0.003	0.003	0	0	0	0	0	0
Mesothuria sp.	_				_					_	_	
D	0	0.009	0.006	0.003	0	0	0.003	0.003	0.017	0	0	0
sp D	0	0	0	0.006	0	0	0	0	0.003	0.003	0	0
Commatulid* S	0	0	0	0.000	0	0	0	0	0.000	0.000	0.003	0
Colothoid	0.000	0.070	0.047	0 1 4 2	0 1 9 0	0.116	0 1 9 0	0 172	0 172	0.022	0.000	0.015
Galatrielu	0.090	0.070	0.047	0.142	0.160	0.110	0.160	0.172	0.172	0.032	0.000	0.015
Prawn	0.003	0.009	0.015	0.003	0.009	0.003	0.006	0.003	0.009	0.012	0	0
sp.?	0	0	0	0.003	0.003	0	0.003	0.003	0	0	0	0
Opisthobranch												
D	0	0	0	0.003	0	0	0	0	0	0	0	0
Enteropneust D	0	0	0	0	0	0	0	0	0.003	0	0	0
TOTAL	0.462	0.485	0.378	0.311	0.451	0.509	0.459	0.262	0.256	0.174	0.183	0.218
Total												
suspension	0	0	0	0	0.003	0	0.003	0.003	0.003	0	0.081	0.084
feeders												
feeders	0.456	0.477	0.355	0.302	0.436	0.503	0.448	0.253	0.244	0.154	0.081	0.122

- 1 FIGURES
- 2

Figure 1: Bathymetry of Usan area showing the position of ROV seabed transects. The lower
panel shows the higher slope transects in more detail. The inset on the upper panel shows the
position of the study site in the Gulf of Guinea. The length of the 200 m long ROV transects
is drawn to scale.

7

8 Figure 2: Example pictures of seabed habitats with different slope found at Usan: A)  $1^{\circ}$  slope,

9 B) 3° slope, C) 11° slope, D) soft-substratum area with 29° slope and E) hard-substratum area

10 with  $29^{\circ}$  slope. Images A to D are images from the video transects cropped to show an

11 approximately 0.5 m wide section of seafloor at the base of the image. Image E is a more 12 oblique image cropped to show an approximately 1 m wide section of seafloor at the base of

13 the image.

14

15 Figure 3: Examples of megafauna found at Usan: A) Hexactinellid (~150 mm diameter), B)

16 White antipatharian (~200 mm height), C) Whip-like cnidarian (found on rocks, ~ 400 mm

- 17 height), D) Whip-like cnidarian (found on sand, ~ 300 mm height), E) Gorgonian fan (~ 1m
- 18 height), F) *Pennatulea* sp. (~300 mm height), G) Brisingid (arm length ~ 250mm), H)
- 19 Phormosoma placenta (50-120 mm diameter), I) Red echinoid (70 mm diameter), J) White

20 Diadema sp. (~150 mm diameter including spines), K) Ophiuroid (~150 mm arm-tip to arm-

21 tip), L) Galatheid (up to 100 mm length), M) \*Prawn (~100 mm total length), N)

22 \*Cerianthid? (~ 100 mm total diameter), O & P) \*Lithodid crab (c.f. Paralomis cristulata,

23 ~100 carapace width). \* species observed but not seen in quantitative transects.

24

25 Figure 4: Response of megafaunal assemblage indices to slope at the Usan area, offshore

26 Nigeria. A) Megafaunal invertebrate deposit (hollow circles) and suspension feeding (filled

27 circles) assemblage total densities. B) Number of megafaunal invertebrate deposit (hollow

circles) and suspension feeding (filled circles) taxa. C) Evenness of entire invertebrate

29 megafaunal assemblage measured by Pielou's evenness (J'). D) Shannon-Wiener (log base e)

30 combined diversity index (H') of entire invertebrate megafaunal assemblage. Errors bars on

31 all graphs represent standard deviations and points represent the mean density of 3 replicates.

For A) and B) the megafauna was separated into deposit (hollow circles) and suspension (filled circles) feeders.

34

35 Figure 5: Multidimensional Scaling Ordination of Bray-Curtis similarities (4th root

36 transformed) of Usan Invertebrate megafaunal assemblage. Overlaid are similarity levels

37 from hierarchical cluster analysis (as percentages). Slope is represented by numbers equating

- 38 to the slope in degrees. Letters represent replicates.
- 39

40 Figure 6: Megafaunal density (no hectare<sup>-1</sup>) compared with depth for the Eastern Atlantic.

41 East Atlantic literature data from Bay of Biscay south to Angola (Feldt *et al.*, 1989; Galeron

42 et al., 2000; Kroncke and Turkay, 2003; Lebrato and Jones, 2009; Sibuet et al., 1989; Sibuet

43 *et al.*, 1984; Sibuet and Segonzac, 1985; Thurston *et al.*, 1994; Tietjen, 1992)











