

1 Are seamounts refuge areas for fauna from polymetallic
2 nodule fields?

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20 **Abstract**

21 Seamounts are abundant and prominent features on the deep-sea floor and intersperse with the
22 nodule fields of the Clarion-Clipperton Fracture Zone (CCZ). There is a particular interest in
23 characterising the fauna inhabiting seamounts in the CCZ because they are the only other ecosystem
24 in the region to provide hard substrata besides the abundant nodules on the soft sediment abyssal
25 plains. It has been hypothesised that seamounts could provide refuge for organisms during deep-sea
26 mining actions or that they could play a role in the (re-)colonisation of the disturbed nodule fields.
27 This hypothesis is tested by analysing video transects in both ecosystems, assessing megafauna
28 composition and abundance.

29 Nine video transects (ROV dives) from two different license areas and one Area of Particular
30 Environmental Interest in the eastern CCZ were analysed. Four of these transects were carried out as
31 exploratory dives on four different seamounts in order to gain first insights in megafauna
32 composition. The five other dives were carried out in the neighbouring nodule fields in the same
33 areas. Variation in community composition observed among and along the video transects was high,
34 with little morphospecies overlap on intra-ecosystem transects. Despite the observation of
35 considerable faunal variations within each ecosystem, differences between seamounts and nodule
36 fields prevailed, showing significantly different species associations characterising them, thus
37 questioning their use as a possible refuge area.

38 1. Introduction

39 Seamounts are abundant and prominent features on the deep-sea floor (Wessel et al., 2010). They
40 are common in all the world's oceans, occurring in higher abundances around mid-ocean ridges,
41 island-arc convergent areas, and above upwelling mantle plumes (Kitchingman et al., 2007).
42 Seamounts are defined as isolated sub-surface topographic feature, usually of volcanic origin, of
43 significant height above the seafloor (International Seabed Authority (ISA), 2019). They are generally
44 isolated, typically cone shaped undersea mountains rising relatively steeply at least several hundred
45 meters from the deep-sea floor. Seamounts comprise a unique deep-sea environment, characterised
46 by substantially enhanced currents and a fauna that is dominated by suspension feeders, such as
47 corals (Rogers, 2018). They represent hard substrata in the otherwise soft sediment deep sea and
48 can thus be considered habitat islands (Beaulieu, 2001). Given the growing evidence that seamounts
49 differ substantially across a range of spatial scales, the concept of seamounts as a single, relatively
50 well-defined habitat type is outdated (Clark et al., 2012). Depth and substrate type are key elements
51 in determining the composition and distribution of benthic fauna on seamounts, while location is
52 likely the subsequent most important driver of faunal composition and distribution patterns (e.g.
53 Tittensor et al., 2009). Connectivity varies substantially between seamounts, resulting in the
54 presence of taxa with very localised to very wide distributions (Clark et al., 2010).

55
56 The Clarion-Clipperton Fracture Zone (CCZ), in the equatorial eastern Pacific Ocean, is most known
57 for its extensive polymetallic nodule fields that will potentially be mined in the future. In this area,
58 nodules represent the most common hard substrate on the soft-sediment abyssal plains, and many
59 organisms rely on them for survival (Vanreusel et al., 2016). Removal of hard substrate through
60 mining actions will impact all these organisms, which were estimated at about 50% of all megafaunal
61 species in the CCZ (Amon et al., 2016). Nodule fields in the CCZ are interspersed by seamounts
62 (Wedding et al., 2013), the only feature offering hard substrata besides the nodules. Based on this
63 feature/characteristic, it has been hypothesised that seamounts could provide refuge for organisms
64 during deep-sea mining activities or that seamounts could play a role in the (re-)colonisation of the
65 disturbed nodule fields. Whether or not this is true may have important implications for
66 management of the impacts of polymetallic nodule mining in the CCZ. However, knowledge on the
67 biodiversity inhabiting seamounts in this region is currently lacking.

68 The objectives of the current study were twofold: (i) Provide first insights in seamount megafauna
69 within the CCZ, (ii) Compare the benthic fauna inhabiting seamounts and nodule fields in the eastern
70 CCZ. Since this is the first time the seamounts at the eastern CCZ were visited, a separate section is
71 dedicated to describe these first insights.

72 2. Material and Methods

73 2.1. Study site and data

74 During the SO239 ECORESPONSE cruise in 2015 (Martinez Arbizu and Haeckel, 2015), four
75 seamounts were visited for the first time within two different license areas and one area of
76 particular environmental interest (APEI) within the Clarion-Clipperton Fracture zone (CCZ) (Table 1).
77 Nodule fields within the same license areas were visited and sampled as well. Video imagery and
78 faunal samples were collected by a Remotely Operated Vehicle (ROV Kiel 6000 (GEOMAR), equipped
79 with a high definition Kongsberg OE14-500 camera).

80 Seamount transects were carried out uphill, towards the summit resulting in a depth gradient along
81 the transect (Table 1). The four seamount transects were characterised by different depth ranges
82 and lengths and were, due to the vessel's positioning and the predominant South-East surface
83 currents, all carried out downstream, on the north to north-western flanks of the seamounts (Table
84 1 and Fig. 1). The names of the seamounts used here, Rüppel and Senckenberg (BGR, German
85 License area), Heip (GSR, Belgian License area) and Mann Borgese (APEI3), are the ones agreed upon
86 by the scientist during the ECORESPONSE cruise (Martinez Arbizu and Haeckel, 2015), pending
87 incorporation of these names in the GEBCO gazetteer. The seamounts differed in shape and size
88 with Senckenberg and Heip being a sea-mountain range, while Rüppel and Mann Borgese were more
89 isolated, stand-alone seamounts (Fig. 1). Nodule field dives were carried out on relatively flat
90 surfaces (maximum depth range covered during a dive or transect was 30m difference, Table 1) and
91 were referred to by the dive number and license area. The five nodule transects were all located
92 between 4000-5000m depth and the transects differed in length between dives as well (Table 1).
93 Within the same license area, distance between different transects was 16 to 60km, while distance
94 between license areas added up to several hundreds of kilometres (minimum ~700kms BGR – GSR,
95 Fig. 1).

96 Investigated areas were restricted to the eastern part of the CCZ with APEI3 being the most north-
97 and westward bound area. The optical resolution of the camera enabled reliable identification of
98 organisms larger than 3 cm (Martinez Arbizu and Haeckel, 2015). The combination of exploration
99 and opportunistic sampling restricted a systematic image collection. Target ROV travelling altitude
100 was <2m and travelling speed was~0.2m/s which, along with the camera zoom, were kept constant
101 whenever possible.

102 2.2. Video analysis and statistics

103 All videos were annotated to the lowest taxonomic level possible. The number of morphospecies,
104 defined as morphologically different organisms within the lowest taxonomic group identified, were
105 assessed. Identifications were double checked with scientists working in the same area as well as
106 taxonomic experts and comprise different taxonomic levels (e.g. Genus, Family) and organism
107 samples were used for proper identification whenever possible. Those identifications restricted to
108 higher taxon groups (Family, Class, etc.) and for which it was impossible to attribute a
109 morphospecies, were referred to as taxa and are likely to morphologically differ between transects.
110 Xenophyophores, living on the soft sediment deep-sea floor, were less prominently present at
111 seamounts than at nodule fields and were not quantified. Fish (Actinopterygii), Crustacea
112 (Nematocarcinidae, Aristeidae, Peracarida) and Polychaeta were quantified but left out of the
113 comparing statistical analysis due to their lack of representativity and possible attraction due to ROV
114 lights. The same was done for jellyfish and other doubtful identifications that could not be
115 confidently assigned to a higher taxonomic group (Table A1). A subset of the nodule field transects
116 from BGR, GSR and APEI3 was presented by Vanreusel et al. (2016), corresponding to 44% of what
117 was studied here and limited organism identification to a higher taxonomic level (Order (e.g.
118 Alcyonacea) or Class (e.g. Ophiuroidea)). In our study, the entire transects (100%) were annotated to
119 morphospecies level, allowing a detailed comparison between seamounts and nodule fields.

120 Three categories of substratum types were distinguished: (1) Predominant soft substrata (<40% hard
121 substrata), (2) mix or transition (between 40 and 60% hard substrata) and (3) predominant hard

122 substrata (>60% hard substrata), and were annotated per 10m distance units based on the video
123 footage and tested for correlations with taxonomic abundances.

124 ROV transects on the seamounts were carried out as exploratory dives. Sampling strategy both at
125 seamounts and nodule fields combined video and sampling or specimen collection. Travelling
126 altitude was easier maintained at the relatively flat nodule field transects, where an average of 93%
127 of the time was spent at altitudes <2m. Contrastingly, the uphill seamount transects were more
128 variable in ROV altitude with on average 61% of the time spent at <2m altitudes, and the remaining
129 ~39% spent at higher altitudes, which generally resulted in a higher surface covered at the
130 seamounts. Approximate surface covered (m^2) was then estimated by using the ROV altitude, time
131 spent at a predefined altitude, travelled distance, and the image widths at predefined altitudes.
132 Following altitude ranges (and image widths, following Vanreusel et al. 2016 and extrapolated
133 thereon) were taken into account: <1m (2m), 1-2 m (4m), 2-3m (6m), 3-4 m (8m), 4-5m (10m).
134 Ranges from >5m, adding up to 12% for seamount transects and 3% for nodule field transects that
135 were left out since these were the parts where the seafloor was not visualised or organisms could not
136 be quantified. Due to the explorative nature of the dives, the pan and tilt of the ROV camera were
137 not kept constant and thus represents a bias on the surface estimations. Visualisation of ancient
138 disturbance tracks were omitted as well, as these fell out of the scope of the article. Faunal densities
139 were calculated as individuals per square meter (ind./100m 2). Statistical testing was carried out in R
140 (R core team, 2018). Non-metric multidimensional scaling analysis (NMDS) was based on Bray-Curtis
141 dissimilarity and carried out with the vegan package (Oksanen et al., 2018). The Kendall's coefficient
142 of concordance (W) was calculated to identify significantly associated groups of species, based on
143 correlations and permutations (Legendre, 2005).

144 3. Results

145 About 80% of all taxa observed across the two adjacent ecosystems, could be identified to a
146 morphospecies level. At a first view, morphospecies revealed to be quite different between
147 seamounts and nodule fields (Fig. 2). While the number of faunal observations at the seamount
148 transects were within similar ranges (4.4-7.6 ind./100m 2), those at the nodule transects featured
149 both highest and lowest values (3.8-30.3 ind./100m 2) (Table 1). The lowest number of faunal
150 observations were done at the two APE13 nodule transects (ROV13 and 14) and highest at the GSR
151 nodule transect ROV08. What follows is a first description of eastern CCZ seamount megafauna
152 (section 3.1.) and a detailed comparison with the neighbouring nodule fields (section 3.2.)

153 3.1. Insights in CCZ seamount megafauna

154 The most abundant and diverse (most morphospecies) taxa at the seamount transects comprised
155 Echinodermata (Asteroidea, Crinoidea, Holothuroidea and Ophiuroidea), Anthozoa (Actiniaria,
156 Alcyonacea, Pennatulacea) and Porifera (Hexactinellida) (Table A1, Fig. 3, Fig. A1). Keeping in mind
157 the limitation of the video sampling, differences among the benthic seamount community
158 composition are described here. The transect at Mann Borgese (APE13) was characterised by high
159 densities of Antipatharia, more specifically Antipathidae (3.5 ind./100m 2), and solitary Scleractinia
160 (1.5 ind./100m 2) (Table A1, Fig. 3). Antipathidae observations were mostly grouped at the end of the
161 video transect, i.e. at the summit. Densities of both Antipatharia and Scleractinia were much lower
162 on the other seamount transects (<0.2 ind./100m 2) with Scleractinia being absent from Heip and
163 Senckenberg transects. Alcyonacea corals were observed on all seamount transects. Isididae were
164 found at Senckenberg and Heip transects, and one individual from the Chrysogorgiidae family was
165 observed at the latter as well. Varying numbers of Primnoidae were observed on all transects (Table

166 A1). High abundances of Pennatulacea were observed at Senckenberg (0.7 ind./100m²), representing
167 about 20% of sessile fauna annotations for this transect.

168 Enteropneusta were only observed on Rüppel and Senckenberg transects in the BGR area,
169 represented by two different morphospecies, namely *Yoda* morphospecies (Torquaratoridae) at
170 Rüppel and *Saxipendium* morphospecies (Harrimaniidae) at Senckenberg.

171 Highest Polychaeta densities were observed at Heip transect in the GSR area, which was mainly due
172 to high densities of free-swimming Acrocirridae (0.5 ind./100m² vs. 0.02-0.03ind./100m² in BGR area
173 Table A1). Aphroditidae polychaetes were only present at the BGR transects (0.02 ind./100m² ,
174 corresponding to 3 individuals along the transect at Rüppel and 1 individual along the transect at
175 Senckenberg) (Table A1).

176 Porifera densities were highest at the Heip transect (0.93 ind./100m²), followed by Senckenberg
177 (0.38 ind./100m²), Mann Borgese (0.36 ind./100m²) and lastly Rüppel (0.31 ind./100m²) (Table 2, Fig.
178 A1(c)). Six Porifera families were annotated featuring >7 to >10 morphospecies per transect (Table
179 2). Cladorhizidae (two individuals) were only observed on Heip transect, and one *Poliopogon* sp.
180 (Pheronematidae) was observed at Mann Borgese transect. Rossellidae gen. sp. nov. was present on
181 three seamount transects, exception being Mann Borgese.

182 Overall Echinodermata densities were highest at Senckenberg seamount (3.5 ind./100m²), adding up
183 to 51% of all image annotations for this transect. For comparison, echinoderms at Heip (1.5
184 ind./100m²) and Rüppel (1.4 ind./100m²) were responsible for 37 and 32% of all image annotations
185 for these transects, followed by Mann Borgese (0.62 ind./100m²) or 8.2% of the annotations. The
186 number of morphospecies for all echinoderm taxa (Asteroidea, Echinoidea, Holothuroidea and
187 Crinoidea) was also highest at these 2 seamounts in the BGR area (Fig. A1., Table A1). Crinoidea and
188 Holothuroidea densities were highest at Senckenberg (0.9 ind./100m² and 0.7 ind./100m²,
189 respectively). The holothuroid families of Elpidiidae and Laetmogonidae were only observed at
190 Senckenberg and Rüppel (BGR). Psychropotidae and Synallactidae were observed on all seamounts,
191 represented by different morphospecies. Deimatidae were not observed on Mann Borgese, but were
192 present on the three other seamount transects, again with different morphospecies and densities.
193 Velatid Asteroidea were only observed at Senckenberg and Rüppel (BGR), while Brisingida and
194 Paxillosida were observed on all four seamounts. Aspidodiadematid Echinoidea were absent from
195 the Heip transect and urchinid Echinoidea were absent from the Mann Borgese transect.

196 A species accumulation curve (Fig. 4a) confirmed the limitations of the restricted and exploratory
197 nature of the sampling as no asymptote was reached. The rarefaction curves (Fig. 4b) showed that
198 the transects with the most faunal observations, which corresponded here to the longer transects,
199 were more diverse. However, at smaller sample sizes curves did not cross, thus maintaining the
200 differences observed at higher sample sizes with the Senckenberg transect (ROV04) as most diverse
201 followed by Rüppel (ROV02) (both BGR). The video transect carried out at Mann Borgese (ROV15,
202 APEI3) was the least diverse.

203 A comparison of all morphospecies observed along the 4 transects was presented in a Venn diagram
204 (Fig. 5a). Each seamount transect was characterised by a highest number of unique morphospecies,

205 only observed on the transect in question and not elsewhere. Only three morphospecies were
206 present in all seamount transects, namely Ceriantharia msp. 2, a small red galatheid crab and a
207 foliose sponge. Highest number of overlapping morphospecies (n=16) was observed between Rüppel
208 and Senckenberg, both in the BGR area (Fig. 5a). Mann Borgese showed the smallest degree of
209 overlap with the other transects (Fig. 5a).

210
211 About 57% of all sessile fauna was associated with predominantly hard substrata, followed by 31%
212 on the mixed substrata. For the mobile taxa, the pattern was less pronounced with 41 and 42%
213 associated with predominantly hard and mixed hard/soft substrata respectively. The amount of
214 predominantly hard and soft substrata was negatively correlated, though not significantly. This was
215 due to the equal amounts (40-60%) of mixed hard/soft substrata. Over all seamount transects
216 pooled together, no taxa were significantly correlated with the amount of hard substrata, nor with
217 soft substrata. When looking at the individual transects, no significant correlations were found
218 between taxa and substrata for ROV02 or ROV04 or ROV09, most likely due to the equal distribution
219 of the amount of hard/soft/mix substrata. In this perspective, ROV15 stood out, as it was dominated
220 by predominantly hard substrata (56%). For this transect, Pennatulacea were significantly
221 negatively correlated with the amount of hard substrata and Zoantharia/Octocorallia were
222 significantly and positively correlated with hard substrata, as were Ophiuroidea, Asteroidea,
223 Crinoidea and Mollusca.

224
225 Due to the limited sample size, the representativity of the observed biological patterns remains to
226 be corroborated by a more elaborate sampling strategy.

227 3.2. Comparison of seamount and nodule field faunal composition and variation

228 The faunal composition and richness of the nodule transects can be consulted in Fig. 3, Fig. A1 and
229 Table A1. The only taxon showing significant difference in density between seamounts and nodule
230 fields were the Porifera (T-test assuming unequal variances, $t=-3.7$, $p<0.05$). In concordance with the
231 seamount transect, the species accumulation curve of the nodule transects did not reach an
232 asymptote either (Fig. 4c). The rarefaction curves showed that the relations among transects were
233 less straightforward for the nodule transects versus the seamount ones and did cross at smaller
234 sample sizes (<100 individuals, Fig. 4d). ROV13 and ROV14 transects (both APEI3) were the longest in
235 distance travelled (Table 1) but featured less faunal observations. At small sample sizes, the richness
236 at ROV13 and 14 was highest. ROV08 and ROV10 (both GSR) showed parallel curves with ROV08
237 being more diverse (Fig. 4d).

238 A venn diagram showing the morphospecies overlap among the nodule transects showed a total of 5
239 species re-occurring on all 5 transects (Fig. 5b). These were: Munnopsidae msp. 1 (Isopoda,
240 Crustacea), Actiniaria msp.7 (Cnidaria), Ophiuroidea msp. 6 (Echinodermata), *Holascus* sp. and
241 *Hyalonema* sp. (Hexactinellida, Porifera). There was a high number of unique morphospecies for
242 each transect, though not as high as for the seamount transects (Fig. 5). ROV13 and 14 (both APEI3)
243 showed least overlap with the other transects, which is similar to what was observed at the
244 seamounts.

245 Observations and quantifications of morphospecies confirmed the high degree of dissimilarity
246 between the two neighbouring ecosystems. Porifera, Ophiuroidea (Echinodermata), Actiniaria and
247 Alcyonacea (Cnidaria) were more abundant at nodule fields (Fig. 3). These taxonomic groups were
248 also most diverse on nodule fields (i.e. highest number of morphospecies), exception being the
249 Alcyonacea which featured more morphospecies on the seamounts (12 to 8 morphospecies for

250 seamounts and nodule fields respectively) (Fig. 3). Of all Porifera, Cladhorizidae were more diverse
251 at nodule fields than at seamounts (7 to 1 morphospecies, respectively).

252 There were only 21 morphospecies (10%) that were observed both on seamounts and nodule fields
253 (Fig. 6). While this subset of morphospecies occurred in both ecosystems, they did so in very
254 different densities, i.e. very abundant in one ecosystem and very low in abundance in the other:
255 examples are Galatheidae small red msp. (Decapoda, Crustacea), *Synallactes* white msp.
256 (Holothuroidea), Ophiuroidea msp. 5 and 6, Comatulida msp. 1 (Crinoidea), *Hyalonema* sp. and
257 *Hyalostylus* sp. (both Hexactinellida, Porifera) (Fig. 6).

258 Three Ophiuroidea morphospecies were present at both seamounts and nodule fields (Fig. 2, 3 and
259 6). Most of the Ophiuroidea observed at the CCZ seamounts that could be identified to
260 morphospecies level, were small and situated on hard substrata (morphospecies 5), while those at
261 nodule fields (including morphospecies 6) were observed on the soft sediments. Morphospecies 6
262 was only rarely observed on the seamounts (Fig. 3). Another easily recognisable morphospecies was
263 found on Porifera, coral and animal stalks and was more abundant at seamounts than at nodule
264 fields (morphospecies 4) (Fig. 3).

265 Crinoidea, Asteroidea (both Echinodermata) and Antipatharia (Cnidaria) were more abundant on the
266 seamounts (Fig. 3). This coincided with a higher diversity for Asteroidea and Antipatharia on the
267 seamounts as well. Crinoidea diversity was similar (5 to 4 morphospecies comparing seamounts to
268 nodule fields). Holothuroidea occurred in similar densities in both ecosystems (Fig. 3, though they
269 were characterised by different morphospecies (Table 2, Fig. A1(b)). Overall densities of Echinoidea
270 were comparable between seamounts and nodule fields, though for the nodule fields this was
271 mostly due to one very abundant morphospecies, namely Aspidodiadematidae msp 1, which was
272 absent at the seamounts (Table 2, Fig. A1(b)). Besides this, Echinoidea were more diverse at
273 seamounts (11 morphospecies vs. 5 at nodule fields).

274 There was no morphospecies overlap for Tunicata, Antipatharia, and Actiniaria. Alcyonacea,
275 Ceriantharia, Corallimorphidae and Crinoidea only shared 1 morphospecies between seamounts and
276 nodule fields, namely *Callozostron* cf. *bayeri*, Ceriantharia msp. 2, *Corallimorphus* msp. 2 and
277 Comatulida msp. 1 respectively (Fig. 6).

278 There were no observations of Enteropneusta, Scleractinia and Zoantharia (Cnidaria), Aphroditidae
279 (Polychaeta) or holothuroid Deimatidae at the nodule fields transects (Table A1, Fig. A1). While
280 Actinopterygii were left out of the analysis, it should be noted that fish observations were more
281 diverse at the seamounts than on the nodule fields.

282 There was quite some faunal variation observed among the video transects of, both seamounts and
283 nodule fields (see Fig. 5 and 7). The (dis)similarities were analysed by a nMDS analysis, which
284 grouped the 9 different video transects based on their taxonomic composition. Despite the large
285 intra-ecosystem variation, they pooled in two distinct groups separating the nodule fields from the
286 seamounts (Fig. 7a). Within each group, BSR and GSR transects were more similar to one another
287 both for seamounts and nodule fields, whilst APE13 transects stood out more.

288 The Kendall's coefficient of concordance (W, Legendre, 2005) corroborated the existence of two
289 significantly different species associations, whose composition corresponded to the fauna

290 characterising the nodule fields ($W=2.03$, $p<0.001$, after 999 permutations) and the seamounts
291 ($W=3.04$, $p<0.001$, after 999 permutations).

292 Depth was fitted as a vector on top of the nMDS plot (Fig. 7b) and showed that the discrepancy in
293 faunal composition between the two ecosystems also corresponded to a difference in depth, with
294 the nodule transects all being situated below the 4000m isobath and the seamount transects ranging
295 from 1650 to >3500m (Fig. 7b).

296 4. Discussion

297 4.1. Intra-ecosystem faunal variation

298 Community composition varied markedly at seamounts and nodule fields. The limited sampling ($n=9$
299 transects), at different locations and additionally, for the seamounts, different depth ranges,
300 precluded any general conclusions on quantifications of biodiversity *per se*. However, taking this into
301 account, it was also the first time seamounts were visited in the area, thus granting first insights in
302 the fauna inhabiting these seamounts and allowing a first comparison with nodule faunal
303 composition.

304 The two BGR seamount transects were most similar in faunal composition, followed by the Heip
305 seamount transect (GSR). These seamount video transects were characterised by more similar depth
306 ranges, and the two BGR transects were also geographically closest to each other. Although for
307 seamounts, distance separating them might be a less determining factor than depth since
308 (mega)faunal communities can be very different even between adjacent seamounts (Schlacher et al.,
309 2014; Boschen et al., 2015). Overall, parameters that vary with depth, such as temperature, oxygen
310 concentration, substratum type, food availability, and pressure are considered major drivers of
311 species composition on seamounts (Clark et al., 2010; McClain et al., 2010). The quantification of the
312 amount of hard and soft substrata was not distinctive enough to explain differences observed here.
313 The difference in depth could also explain the higher dissimilarity with Mann Borgese (APEI3) who
314 featured the shallowest transect and summit, which was dominated by Antipatharia. Antipatharians
315 were previously reported to be more dominant towards peaks as compared to mid-slopes at
316 corresponding depths (Genin et al., 1986). Based on their filter-feeding strategy, Porifera (except
317 carnivorous Cladorhizidae), were also thought to benefit from elevated topography (peaks) or
318 exposed substrata in analogy to corals (Genin et al., 1986; Clark et al., 2010), though no such pattern
319 was apparent here. Porifera are notoriously difficult to identify based on imagery. Although the
320 sampled individuals allowed some identifications to genus or species level (Kersken et al., 2018a and
321 b), identifications remained hard to extrapolate across the different video transects. Generally, as in
322 our study, seamount summits have been more intensively sampled (Stocks, 2009) although the little
323 work done at seamount bases and deep slopes indicated that these areas support distinct
324 assemblages (Baco, 2007).

325 Among the nodule transects a considerable amount of variation in faunal composition was observed
326 (this study, Vanreusel et al., 2016). The two APEI3 nodule transects (ROV13 and 14) stood out in
327 faunal composition, diversity and in low number of faunal observations. They were also the only two
328 transects situated below the 4500m isobaths. But rather than depth, the nodule coverage may be
329 considered an important driving factor, since the density of nodule megafauna was shown to vary
330 with nodule size and density/coverage (Stoyanova, 2012; Vanreusel et al., 2016, Simon-Llédo et al.,
331 2019). Here as well, the APEI3 transects were characterised by a high nodule coverage (~40-88%,

332 Vanreusel et al., 2016), whereas the BGR and GSR nodule transects (ROV3 and ROV 8 + 10,
333 respectively) had a nodule coverage <30% and were also more similar in faunal composition
334 (Vanreusel et al., 2016). Other factors that could be at play are the more oligotrophic surface waters
335 of the northern CCZ which could be the cause of the overall lower faunal densities at APEI3 nodule
336 fields (Vanreusel et al., 2016). Volz et al. (2018) corroborated this, with the location of the APEI3 site
337 in the proximity of the carbon-starved North Pacific gyre being characterised by a reduced POC-flux
338 quantified to being 22-46% lower than the GSR and BGR areas respectively.

339 The species accumulation curves showed that an asymptote was reached neither at seamounts, nor
340 at nodule fields. Consequently, longer transect lengths might be necessary to representatively
341 quantify and assess megafauna density and diversity (Simon-Lledó et al., 2019). In addition, for a first
342 in-depth description and assessment of seamount fauna composition, one video transect is
343 insufficient to describe the diversity and shifts in faunal assemblages of the surveyed seamounts.
344 Rather, an ampler imaging strategy should be developed, with a minimum transect length exceeding
345 1000 m (Simon-Llédo et al., 2019) and replicate transects carried out on different faces of the
346 seamount, on slopes with varying degree of exposure to currents and different substrate types.
347 Wider depth ranges should be taken into account as well. Alternatively, across slope transects,
348 following depth contours should be considered as these could provide observation replicates for a
349 given depth. Despite its limitations, this study grants first insights in the seamount inhabiting
350 megafauna of the eastern CCZ and an important first comparison with nodule fauna.

351 4.2. Faunal (dis)similarities between seamounts and nodule fields
352 In other areas, seamounts were shown to share fauna with surrounding habitats (Clark et al., 2010)
353 and could thus potentially serve as source populations for neighbouring environments (McClain et
354 al., 2009). While generally few species seemed restricted to seamounts only (Clark et al., 2010),
355 morphospecies in this study revealed to be quite different between seamounts and nodule fields
356 with little overlap between both. Despite the high degree of variation observed among all the video
357 transects, these grouped into two distinctly separate clusters, separating nodule from seamount
358 transects. The few overlapping morphospecies did occur in different densities in each ecosystem,
359 implying a different role or importance in the ecological community and its functioning.

360 Overall, nodule fields showed higher faunal densities than seamounts. Shifts in density patterns
361 between nodule fields and seamounts were more evident in a number of taxa, where the variety of
362 morphospecies and feeding strategy within each group was likely to be at play. An example of this
363 are the Echinodermata, which include Asteroidea (predators and filter feeders (Brisingida)),
364 Crinoidea (filter feeders), Echinoidea (deposit feeders), Holothuroidea (deposit feeders) and
365 Ophiuroidea (omnivores). Asteroidea were more abundant on seamounts and both Echinoidea and
366 Asteroidea were more diverse in this ecosystem as well. Ophiuroidea were most abundant on the
367 nodule fields (ratio 7 to 1 when compared to seamounts). Same ophiuroid morphospecies were
368 present at seamounts and nodule fields but in very different abundances and they were observed on
369 different substrata types, which indicates different lifestyles, feeding behaviour and corresponding
370 dietary specialisations (Persons and Gage, 1984). Previously it was already demonstrated that
371 Ophiuroidea did not show high levels of richness or endemism on seamounts (O'Hara, 2007). At
372 nodule fields, Ophiuroidea were often observed in association with xenophyophores (Amon et al.,
373 2016, this study) and a similar observation was done at east Pacific seamounts off Mexico (Levin et
374 al., 1986), though no such associations were observed on the seamounts studied here.

375 Holothuroidea densities were thought to possibly decrease when less soft sediment was available
376 since they feed mainly on the upper layers of the soft-bottom sediment (Bluhm and Gebruk, 1999).
377 No significant link was established between holothuroid densities and the amount of hard substrata
378 in this study, but their community composition varied distinctly between nodule fields and
379 seamounts with more families being observed at the latter. Additionally, at the seamounts, many
380 holothurians were observed on top of rocks, possibly reflecting different feeding strategies and
381 explaining the observations of different morphospecies. Geographical variations, different bottom
382 topography, differences in nodule coverages and sizes and/or an uneven distribution of holothurians
383 on the sea floor were thought to play a role in holothuroid community composition (Bluhm and
384 Gebruk, 1999). On the other hand, variability in deep-sea holothuroid abundance was proposed to
385 depend primarily on depth and distance from continents (see Billet, 1991 for a review).

386 Stalked organisms, such as Crinoidea (Echinodermata) and Hexactinellida (except for
387 Amphidiscophora, Porifera) rely on hard substrata for their attachment and are considered being
388 among the most vulnerable organisms when mining is concerned. Crinoidea were more abundant on
389 seamounts, possibly because hard substrata were less limiting than in the nodule fields. Porifera
390 densities (stalked and non-stalked) varied among all analysed transects, revealing no particular
391 trends in abundance. However, the species composition of deep-sea glass sponge communities from
392 seamounts and polymetallic nodule fields was distinctly different. Polymetallic nodule field
393 communities were dominated by widely-distributed genera such as *Caulophacus* and *Hyalonema*,
394 whereas seamount communities seemed to have a rather unique composition represented by
395 genera like *Saccocalyx*.

396 Corals were generally considered to be more abundant on seamounts than adjacent areas, due to
397 their ability to feed on a variety of planktonic or detritus sources suspended in the water column
398 (Rowden et al., 2010). In this study, the Alcyonacea densities were lower on the seamounts than on
399 the nodule transects. The majority of Alcyonacea morphospecies of the seamounts did not occur on
400 the nodule fields and vice versa, with exception of *Callozostron cf. bayeri* which was also present at
401 the nodule fields but in very low densities (1/8 of those observed at seamounts). The Antipatharia
402 were most abundant at the Mann Borgese seamount (APE13) compared to all other transects. The
403 depth difference of more than 3000m between this particular seamount and the nodule fields could
404 explain the abundance in Antipatharia which were shown to be more abundant at lower depths
405 (Genin et al., 1986). Additional presence of Pennatulacea at seamounts, a taxon that was virtually
406 absent from the nodule field transects and that appeared more linked to predominant soft substrata
407 at seamounts, resulted in completely distinct coral communities for each ecosystem.

408 Actiniaria were denominated the second most common group at CCZ nodule fields, after the
409 xenophyophores (Kamenskaya et al., 2015) and, in our study, were also more abundant on nodule
410 fields than on seamounts. Depending on the species and feeding strategy, the ratio hard/soft
411 substrata and their preference for either one could play a role. Since morphospecies were distinct
412 between seamounts and nodule fields, their role in the respective communities are likely to differ as
413 well. Combinations of deposit feeding and predatory behaviour in Actiniaria have been observed, as
414 well as burrowing activity, preference for attachment to hard substrata and exposure to currents
415 (Durden et al., 2015a; Lampitt and Paterson, 1987; Riemann-Zürneck, 1998), all factors that could
416 influence the differences in morphospecies observed.

417 Some taxa were only observed on the seamounts in this study, while they occurred on nodule fields
418 elsewhere, be it in low densities. For instance, Enteropneusta, which in this study were found only
419 on seamounts, were observed previously at CCZ nodule fields though observations were rather rare
420 (Tilot, 2006). They appeared more abundant at the nodule fields of the Deep Peru Basin (DISCOL
421 area), though a wide range in abundances was displayed there as well (Bluhm, 2001). The exception
422 were the Scleractinia, which were quite common on seamounts, as also reported in other studies
423 (e.g. Baco, 2007, Rowden et al., 2010), but distinctly absent at nodule fields.

424 Explanation for the discrepancies in faunal composition and the low degree of morphospecies
425 overlap between seamount and nodule fields, as observed here, can be multiple. For one, nodules
426 may not be considered a plain hard substratum, with their metal composition, microbial colonisation
427 and the nodule/sediment interface influencing the epi-and associated megafaunal composition. The
428 possibility of a specific deep-sea faunal community that tolerates or benefits from manganese
429 substrata has been previously proposed (Mullineaux, 1988). The comparison between seamounts
430 and nodule fields as two neighbouring hard-substrata ecosystems also entailed a comparison
431 between depth gradients and possible thresholds (>4000m for nodule fields and 1500>x <4000m for
432 seamounts). Related to this is the steepness of the seamount slope and its current exposure playing
433 a role in the faunal colonisation (Genin et al., 1986; Rappaport et al., 1997). Other studies showed
434 that habitat heterogeneity increased megafaunal diversity at seamounts (Raymore, 1982) and
435 elsewhere, such as abyssal plains (Lapointe and Bourget, 1999; Durden et al., 2015b, Leitner et al.,
436 2017, Simon-Llédó et al., 2019). Within this perspective the smaller-scale substratum heterogeneity
437 transcending the ratio hard/soft substrata or amount of hard substrata could play a role as well.

438 5. Conclusions

439 Based on our current knowledge; seamounts appear inadequate as refuge areas to help maintain
440 nodule biodiversity. In order to conclusively exclude seamount habitats as a refuge for nodule fauna,
441 a more comprehensive sampling should be carried out. The sampling strategy wielded in this study
442 lacked replicates, uniformity and was limited in sample size. Seamount bases should be taken into
443 consideration as well as they can be characterised by distinctly different assemblages than the
444 summits and they feature depth ranges more similar to nodule fields.

445 While their role as refuge area for nodule field fauna is currently debatable, the possible uniqueness
446 of the seamount habitat and its inhabiting fauna implies that seamounts need to be included in
447 management plans for the conservation of the biodiversity and ecosystems of the CCZ.

448 Author Contributions

449 DC, PAR, SPR, DK analysed the images. DC analysed the data. PMA, PAR, AC conceptualised and
450 carried out the sampling. All authors contributed to the redaction of the manuscript.

451 Data Availability

452 Data sets are made available through OSIS-Kiel data portal, BIIGLE and PANGAEA.

453 Competing interest

454 The authors declare that they have no conflict of interest

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Tables

575
 576 Table 1. Overview table on details of imagery transects analysed in the Clarion-Clipperton license
 577 areas. Video duration includes time spent sampling. Number of observations include undetermined
 578 organisms. Transect lengths do not include parts visualising ancient disturbance tracks or parts when
 579 the seafloor was not visualised or visible.

Station/Dive	License Area	Seamount (SM) or Nodule field (NF)	Depth (m)	Transect length	Approximate surface covered (m ²)	Faunal densities (ind. /100m ²)
SO239_29_ROV02	BGR	SM: Rüppell	3000-2500	1250m	9458.6	4.4
SO239_41_ROV03	BGR	NF	4080-4110	1590m	5309.1	19.3
SO239_54_ROV04	BGR	SM: Senckenberg	3350-2850	2500m	12288.5	6.9
SO239_131_ROV08	GSR	NF	4470-4480	710m	1602.5	30.3
SO239_135_ROV09	GSR	SM: Heip	3900-3550	1000m	6905.4	5.3
SO239_141_ROV10	GSR	NF	4455-4480	520m	1683.4	27.6
SO239_189_ROV13	APEI 3	NF	4890-4930	1790m	3580.0	3.8
SO239_200_ROV14	APEI 3	NF	4650-4670	1490m	2980.0	6.2
SO239_212_ROV15	APEI 3	SM: Mann Borgese	1850-1650	900m	4805.3	7.6

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585 Table 2. Overview of all densities (ind./100m²) observed within each video transect. Higher taxa are
586 in bold. * indicates taxa left out of the statistical analyses due to lack of representativity. Indets were
587 organisms impossible to attribute to a lower taxonomic group. ROV02=Rüppel, ROV04=Senckenberg,
588 ROV09=Heip, ROV15=Mann Borgese

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	SEAMOUNTS				NODULE FIELDS				
	ROV2 Ind./100m ²	ROV4 Ind./100m ²	ROV9 Ind./100m ²	ROV15 Ind./100m ²	ROV3 Ind./100m ²	ROV8 Ind./100m ²	ROV10 Ind./100m ²	ROV13 Ind./100m ²	ROV14 Ind./100m ²
Annelida*									
Polychaeta indet. * (No Serpulidae)	0.02	0.02		0.02	0.09		0.12	0.03	0.03
Acrocirridae	0.02	0.03	0.52		0.17	0.06	0.18	0.89	0.97
Aphroditidae	0.03	0.01			0.17	0.50	0.36		0.10
Echiura msp 1						0.06	0.18		
Polynoidae						0.06		0.03	0.07
Polynoidae msp 2									
Polynoidae white msp									
Bryozoa									
Bryozoa msp 2			0.02						
Bryozoa indet.		0.01			0.13	0.69	0.06	0.06	0.07
Cnidaria									
Anthozoa									
Ceriantharia									
Ceriantharia msp 1	0.04					0.06			
Ceriantharia msp 2		0.01				0.12	0.06		
Ceriantharia indet.			0.05						
Hexacorallia									
Actiniaria									
Actinoscyphiidae									
Actiniidae/ <i>Bolocera</i> msp.	0.13	0.02							
Actiniaria msp 15	0.01	0.04							
Actiniaria msp 4			0.02						
Actiniaria msp 5	0.01	0.02		0.02					
				0.06					

Actiniaria msp 10					0.09				0.17
Actiniaria msp 2					0.32	0.50	0.06		0.13
Actiniaria msp C					0.11	0.19	0.12		
Actiniaria msp D					0.02				
Actiniaria msp 7					0.19	0.06	0.18	0.03	0.07
Actiniaria msp 8					0.04	1.62	0.95		0.10
Actiniaria msp 9							0.06		
Actiniaria msp A							0.06		0.03
Actiniaria msp B					0.08	0.06	0.12	0.03	
Actiniaria indet.	0.02	0.03			0.47	0.62	1.37	0.06	0.07
<u>Antipatharia</u>									
Antipathidae									
<i>Antipathes</i> msp 1					1.59				
<i>Antipathes</i> msp 2					0.02				
<i>Stichopathes</i> msp 1					1.75				
Antipathidae indet.					0.10				
Schizopathidae									
<i>Abyssopathes</i> cf. <i>lyra</i>						0.15	0.25	0.18	
<i>Bathypathes</i> cf. <i>alternata</i>							0.06	0.06	
<i>Bathypates</i> cf. <i>alternata</i> msp 1									
<i>Bathypates</i> cf. <i>alternata</i> msp 2									
<i>Bathypathes</i> sp.						0.06	0.06		
<i>Bathypathes</i> msp 1									
cf. <i>Parantipathes</i> msp 1									
<i>Umbellapathes</i> aff. <i>bipinnata</i>									
<i>Umbellapathes</i> aff. <i>helioanthes</i>									
Antipatharia indet.	0.01	0.02	0.01	0.12	0.08	0.12	0.06		
<u>Corallimorpharia/Corallimorphidae</u>									
<i>Corallimorphus</i> msp 1		0.01							
<i>Corallimorphus</i> msp 2		0.09	0.01		0.08	0.12	0.06		
Corallimorpharia msp 3		0.01							

Corallimorpharia msp 4			0.01		0.02		0.06		
Corallimorpharia msp A					0.02				
Corallimorpharia msp B					0.02				
Scleractinia									
Scleractinia msp 1	0.02			1.47					
Zoantharia									
Zoantharia msp 2		0.09		0.02					
Zoantharia indet.				0.04					
Octocorallia									
Alcyonacea									
Alcyoniidae									
<i>Anthomastus</i> msp 1	0.03								
<i>Anthomastus</i> msp 2	0.00	0.03		0.02					
Coralliidae				0.02					
<i>Corallium</i> sp. nov.				0.02					
Chrysogorgiidae			0.01						
<i>Chrysogorgia</i> cf. <i>pinnata</i>									
Isididae									
<i>Bathygorgia</i> aff. <i>abyssicola</i> 1						0.06	0.06		
<i>Bathygorgia</i> aff. <i>profunda</i> 1		0.03	0.01						
<i>Bathygorgia</i> aff. <i>profunda</i> 2			0.01						
<i>Keratoisis</i> aff. <i>flexibilis</i> msp 2			0.01						
Isididae msp 1	0.02	0.01		0.11	0.02	0.04	2.50	0.71	
Isididae indet.									
Taiaroidea									
Taiaroidae msp 1								0.06	
Primnoidae									
<i>Abyssoprimnoa</i> cf. <i>gemina</i>							0.31	0.18	
<i>Callozostron</i> cf. <i>bayeri</i>	0.01	0.11				0.02			
<i>Calyptrophora</i> cf. <i>persephone</i>						0.02			
<i>Narella</i> msp 1		0.02		0.02					

Primnoidea indet.	0.08		0.02		0.81	1.50	0.48	0.06	0.07
Alcyonacea msp 1		0.03	0.21	0.50	0.04	2.93	1.25		0.03
Alcyonacea indet.					2.67				
<u>Pennatulacea</u>									
Umbellulidae									
<i>Umbellula</i> msp 1_White									0.07
<i>Umbellula</i> msp 1_orange		0.06		0.02					
<i>Umbellula</i> msp 2			0.02						
Umbellulidae indet.		0.03							
Protoptilidae					0.02				
Protoptilum msp 1			0.01		0.04				
Pennatulacea msp 2			0.01						
Pennatulacea msp 5			0.05						
Pennatulacea msp 6			0.02						
Pennatulacea msp 7			0.08						
Pennatulacea msp 8			0.02						
Pennatulacea indet	0.02		0.42		0.02		0.06		
Octocorallia msp 1					0.04				
Octocorallia msp 2									
Anthozoa indet.	0.02	0.02	0.07	0.12	0.04	0.06	0.06		
<i>Hydrozoa</i>									
<i>Brachiocerianthus</i> msp			0.02						
Hydrozoa indet.		0.02	0.01	0.04		0.06			
Crustacea*									
<i>Decapoda</i>									
Caridea	0.46		0.52	0.47	0.04	0.06		0.36	0.06
Decapoda msp 3			0.02						0.10
Decapoda msp 4	0.01								
Decapoda/Aristeidae	0.01		0.02		0.10	0.02	0.25	0.18	0.06
Decapoda msp 1								0.03	

Galatheidae									
Galatheidae small red msp	0.37	0.11	0.02	0.08	0.02				
Galatheidae small white msp	0.01	0.02							
Munidopsis spp.	0.11	0.07	0.07						0.10
Galatheidae indet.	0.02	0.03	0.02	0.02			0.19		
Parapaguridae									
Parapaguridae msp 1/ <i>Proberebei</i> sp.	0.07	0.05							
<i>Peracarida</i>									
Amphipoda			0.01		0.02	0.12		0.03	0.13
Podoceridae msp 1								0.03	
Amphipoda msp 1		0.02	0.02						
Isopoda									
Munnopsidae msp 1					0.17	0.19	0.06	0.08	0.27
Decapoda indet.		0.02	0.10						
Crustacea indet.	0.01	0.06	0.07						0.07
Echinodermata									
<i>Astroidea</i>									
<i>Brisingida</i>									
Brisingida msp 1 (6 arms - orange)		0.03	0.07		0.08				
Brisingida msp 1 (8 arms - orange)	0.02	0.08	0.04						0.03
Brisingida msp 3 (6 arms - white)		0.08	0.13	0.04	0.06	0.19	0.06		0.07
Brisingida msp 4 (9-10 arms)	0.02	0.08							
Brisingida indet.	0.04	0.02		0.04					
<i>Paxillosida</i>									
<i>Solaster</i> msp		0.01							
Paxillosida cf AST_009/AST_007		0.10	0.06						
Paxillosida msp 1	0.01	0.01		0.02					
Paxillosida msp 2a									
Paxillosida msp 2b			0.01						
Paxillosida msp 3		0.02	0.02						

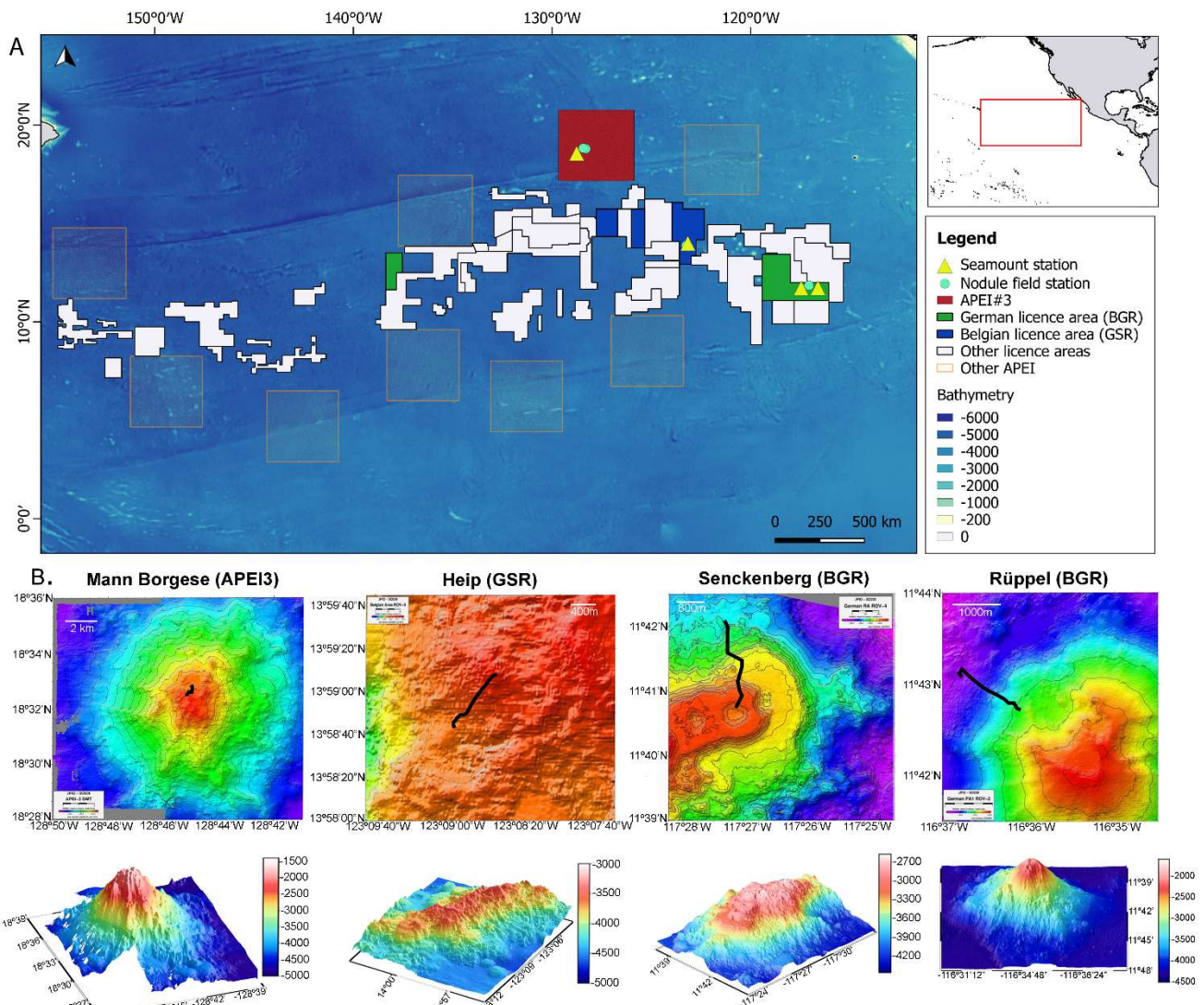
<i>Enypniastes</i> sp.									0.03
Psychropotidae									
<i>Benthodytes</i> cf. <i>incertae</i> purple msp		0.03	0.01						
<i>Benthodytes</i> cf. <i>incertae</i> red msp		0.09							
<i>Benthodytes</i> msp		0.04							
<i>Benthodytes</i> msp 1				0.02					0.03
<i>Benthodytes</i> pink msp			0.01	0.01					
<i>Benthodytes</i> purple msp									
<i>Benthodytes</i> red msp					0.02				
<i>Psychropotes</i> cf. <i>semperiana</i>								0.03	
<i>Psychropotes</i> <i>longicauda</i>							0.12		
<i>Psychropotes</i> msp 3							0.06		
<i>Psychropotes</i> <i>verrucosa</i>						0.08	0.06		
Psychropotidae msp 1_Nodules						0.02	0.06	0.06	
Psychropotidae msp 1		0.07	0.01						
Psychropotidae msp 2_Nodules							0.19		
Psychropotidae msp 2			0.01						
Psychropotidae msp 3						0.04	0.06		
Psychropotidae msp 4							0.06		
Psychropotidae red msp	0.02								
Psychropotidae indet.	0.16	0.09		0.02					
<u>Holothuriida</u>									
Mesothuriidae									
<i>Mesothuria</i> msp	0.01	0.02							
<u>Synallactida</u>									
Deimatidae									
<i>Deima</i> msp.		0.01							
Deimatidae - irregular papillae length msp		0.05	0.01						
<i>Oneirophanta</i> msp	0.01		0.02						
Deimatidae indet.		0.01	0.01						
Synallactidae									

<i>Benthothuria</i> msp					0.08					
<i>Paelopatides</i> "orange" msp	0.01	0.01								
<i>Synallactes</i> msp 1 (<i>Synallactidae</i> purple msp)	0.01	0.01								
<i>Synallactes</i> msp 2										
<i>Synallactes</i> msp 2 pink					0.04	0.25	0.06			
<i>Synallactes</i> msp 2 pink (smooth)	0.03	0.02				0.31				
<i>Synallactes</i> sandy-coloured msp	0.02									
<i>Synallactes</i> white msp	0.02				0.70	0.19	0.30			0.03
<i>Synallactidae</i> indet.	0.04									
<u>Persiculida</u>										
<i>Molpadiodemidae</i>										
<i>Molpadiodemas</i> msp		0.02								
<i>Pseudostichopodidae</i>										
<i>Pseudostichopus</i> msp						0.06				
<i>Molpadiodemas/Mesothuria</i>	0.17	0.15	0.04	0.04	0.06	0.12	0.06			0.07
<i>Holothuroidea</i> indet.					0.06	0.06	0.12			
<u>Ophiuroidea</u>										
<i>Ophiuroidea</i> msp 1					0.02	0.06	0.06			
<i>Ophiuroidea</i> msp 3						0.12				
<i>Ophiuroidea</i> msp 5	0.02	0.39	0.49							
<i>Ophiuroidea</i> msp 6		0.03	0.01		0.32	1.31	0.65			
<i>Ophiuroidea</i> msp 4	0.04	0.21			0.11					
<i>Ophiuroidea</i> msp 7		0.01								
<i>Ophiuroidea</i> indet.		0.02	0.04	0.08	5.67	6.68	7.31			0.13
<u>Enteropneusta</u>										
<i>Enteropneusta</i> msp 1 cf. <i>Yoda</i>		0.10								
<i>Enteropneusta</i> msp 2 cf. <i>Saxipendulum</i> msp.	0.07									
<u>Mollusca</u>										
<u>Gastropoda</u>										

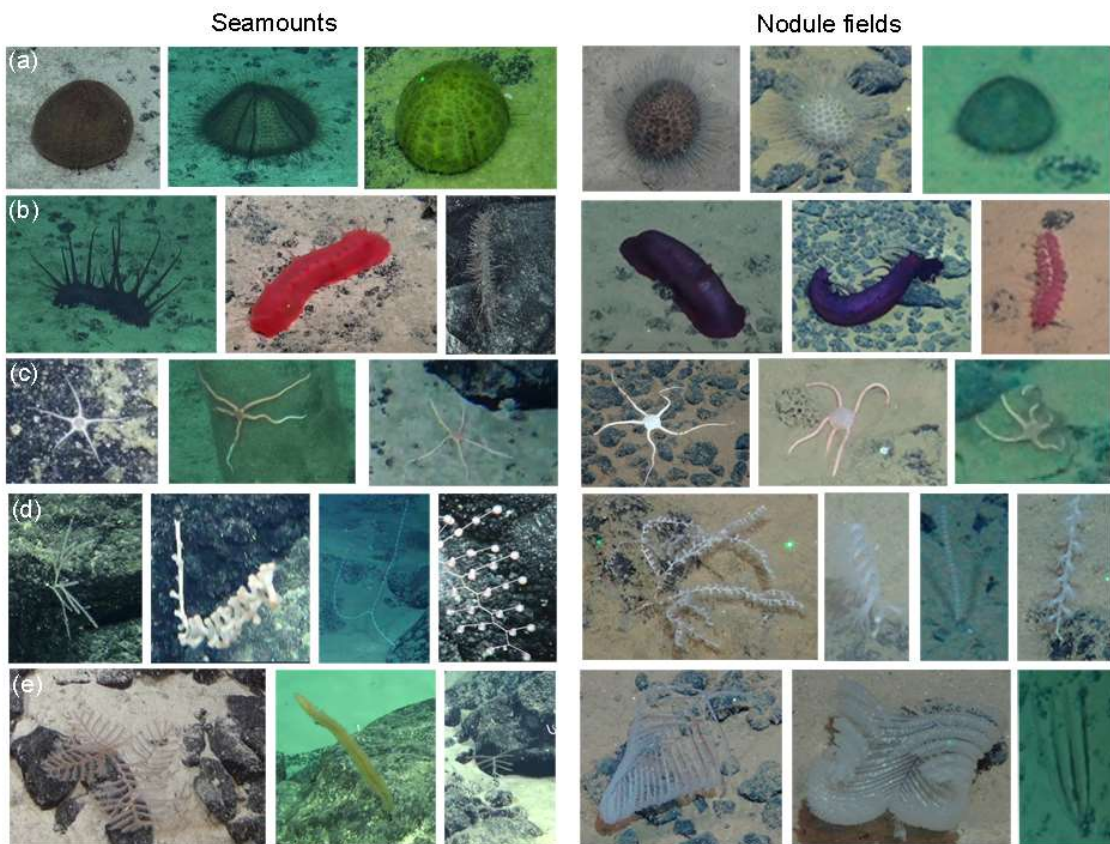
Limpet									
Gastropoda msp 1									
Polyplacophora	0.04		0.01 0.02		0.04				
Gastropoda indet.							0.06		
<i>Cephalopoda</i>									
Octopoda msp 1	0.01								
Pisces*	0.33	0.28	0.07	0.06	0.47	0.19	0.48	0.17	0.17
Porifera									
<i>Demospongiae</i>									
Cladorhizidae									
Cladorhizidae msp 1				0.02					0.07
Cladorhizidae msp 1(soft)									0.03
Cladorhizidae msp 2								0.03	0.03
Cladorhizidae msp 3									0.03
Cladorhizidae msp 4								0.06	0.13
Cladorhizidae msp 5									
Cladorhizidae msp 6									
Cladorhizidae indet								0.03	0.07
<i>Hexactellinida</i>									
Euplectellidae									
<i>Bathydorus spinosus</i>	0.01								
<i>Bolosoma</i> sp.									
<i>Corbitella discasterosa</i>	0.01				0.02				
<i>Docosaccus maculatus</i>								0.06	0.03
<i>Docosaccus nidulus</i>								0.06	
<i>Holascusspp</i>									
<i>Hyalostylus schulzei</i>								0.03	0.03
<i>Hyalostylus</i> sp.								0.03	0.03
<i>Sacocalyx pedunculatus</i>		0.02	0.15	0.02 0.06	0.19	0.12	0.06	0.03	0.03

<i>Sacocalyx</i> sp.	0.04	0.02	0.02						
Euretidae									
<i>Bathyxiphis subtilis</i>									0.03
<i>Chonelasma bispinula</i>	0.01			0.01				0.06	
<i>Chonelasma choanoides</i>									
<i>Chonelasma</i> sp.									
Hyalonematidae									
<i>Hyalonema</i> spp.		0.02	0.10	0.02	0.11	0.31	0.24	0.08	0.23
Rosselidae									
<i>Caulophacus</i> sp.		0.06	0.07	0.02	0.17	0.06	0.06		
<i>Crateromorpha</i> sp.		0.02		0.02					
Rossellidae gen. sp.	0.04	0.01	0.02						
Pheronematidae					0.02				
<i>Poliopogon</i> sp.									
Hexactellinida/foliose sponge msp	0.01	0.02	0.01	0.06					
Hexactellinida - Stalked					0.26	0.50	0.53		
Hexactinellida black msp		0.01							
Hexactellinida indet.	0.20	0.20	0.52	0.12	0.98	1.06	1.60	0.45	0.37
Pycnogonida	0.02		0.01						0.03
Tunicata									
Octacnemidae									
<i>Megalodicopia</i> msp. 1	0.02	0.01	0.01						0.03
<i>Megalodicopia</i> msp. 2									
<i>Dicopia</i> msp.	0.04								
Pyuridae									
<i>Culeolus</i> msp.									0.03
Tunicata indet.	0.02	0.01	0.01	0.02					
<i>Paleodictyon nodosum</i>								0.03	

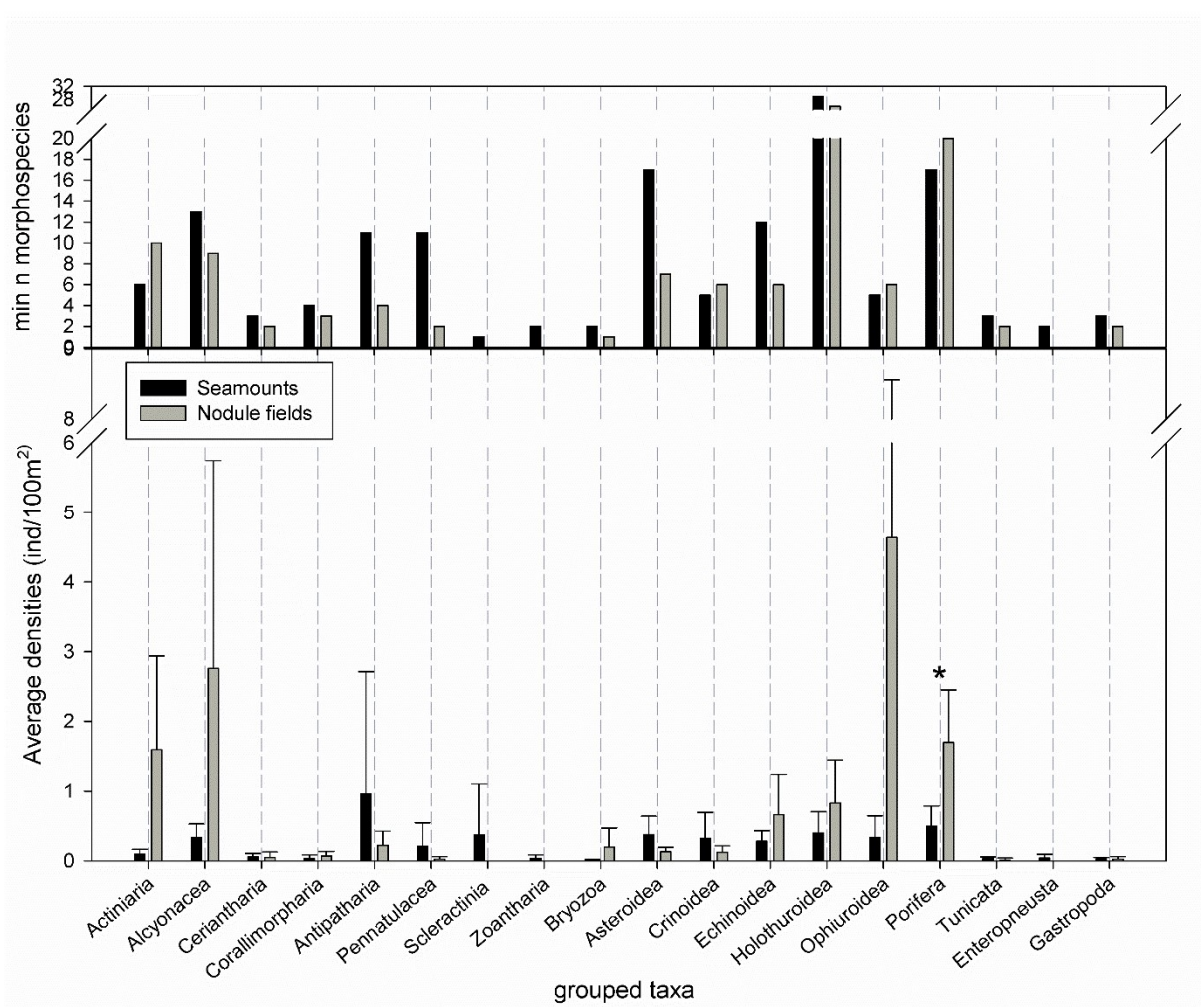
Figures



608 Fig. 1. (A). Location of the Clarion-Clipperton Fracture zone in the equatorial eastern Pacific Ocean
 609 featuring the contract areas from the International Seabed Authority (ISA) and the positions of the
 610 sampled areas (seamounts and nodule fields). Information on transect length and depth gradients
 611 can be found in Table 1. (B). Location of the seamount transects carried out towards the summit on
 612 the north –north-western flank and seamount profiles. Rüppel (BGR, ROV02) and Mann Borgese
 613 (APEI3, ROV15) are single seamounts, while Senckenberg (BGR, ROV04) and Heip (GSR, ROV09) are
 614 sea-mountain ranges.



617
 618 Fig. 2. Some examples of different morphospecies at seamounts and nodule fields in the CCZ.
 619 Selected taxa were (a) Echinoidea (from left to right, Urechinidae msp 4 (URC_019), Urechinidae msp
 620 2 (URC_013), Urechinidae msp 3 (URC_009), Urechinidae msp. A (URC_020), Urechinidae msp. B
 621 (URC_021), Urechinidae msp. C (URC_005), (b) Holothuroidea (from left to right, Psychropotidae
 622 msp 1 (HOL_088), *Benthodytes* red msp. (HOL_101), Deimatidae - irregular papillae msp. (HOL_070),
 623 *Psychropotes verrucosa* (HOL_045), Laetmogonidae (HOL_030), *Synallactes* msp 2 pink (HOL_008)(c)
 624 Ophiuroidea (from left to right, Ophiuroidea msp. 5 (OPH_003), Ophiuroidea msp. 4 (OPH_005),
 625 Ophiuroidea msp. 6 (OPH_006), Ophiuroidea msp. 6 (OPH_006), Ophiuroidea (OPH_012),
 626 Ophiuroidea msp. 4 (OPH_005)), (d) Alcyonacea (from left to right, *Callozostron* cf. *bayeri* (ALC_009),
 627 *Bathygorgia* aff. *profunda* 2 (ALC_005), *Keratoisis* aff. *flexibilis* msp 2 (ALC_029), *Chrysogorgia* cf.
 628 *pinnata*, *Abyssoprimnoa* cf. *gemina* (ALC_008), *Bathygorgia* aff. *profunda* 1, *Calyptrophora* cf.
 629 *persephone* (ALC_007), *Bathygorgia* aff. *abyssicola* 1 (ALC_003), (e) Antipatharia (*Umbellapathes* aff.
 630 *helioanthes* (ANT_018), cf. *Parantipathes* morphotype 1 (ANT_017), *Bathypates* cf. *alternata* msp 1
 631 (ANT_010), *Bathypates* cf. *alternata* (ANT_006), *Abyssopathes* cf. *lyra* (ANT_022), *Bathypates* sp.
 632 (ANT_003)). Codes refer to an ongoing collaboration in creating one species catalogue for the CCZ
 633 and align all morphospecies of different research groups. Copyright: SO239, ROV Kiel 6000, GEOMAR
 634 Helmholtz Centre for Ocean Research Kiel
 635
 636

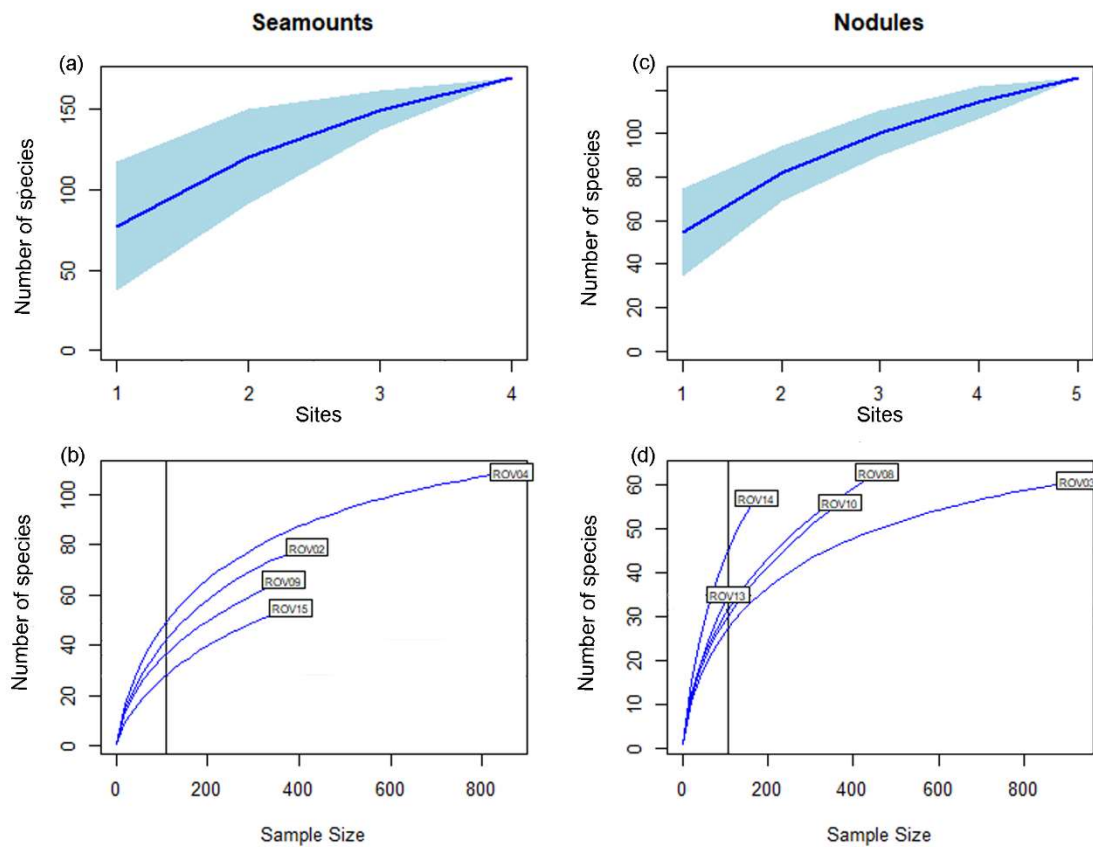


637

638 Fig. 3. Average densities at higher taxa level per ecosystem and standard deviation in the lower
 639 panel and minimum number of morphospecies per taxon and ecosystem in the upper panel. *=
 640 Significant difference in density ($t=-3.7$, $p<0.05$).

641

642

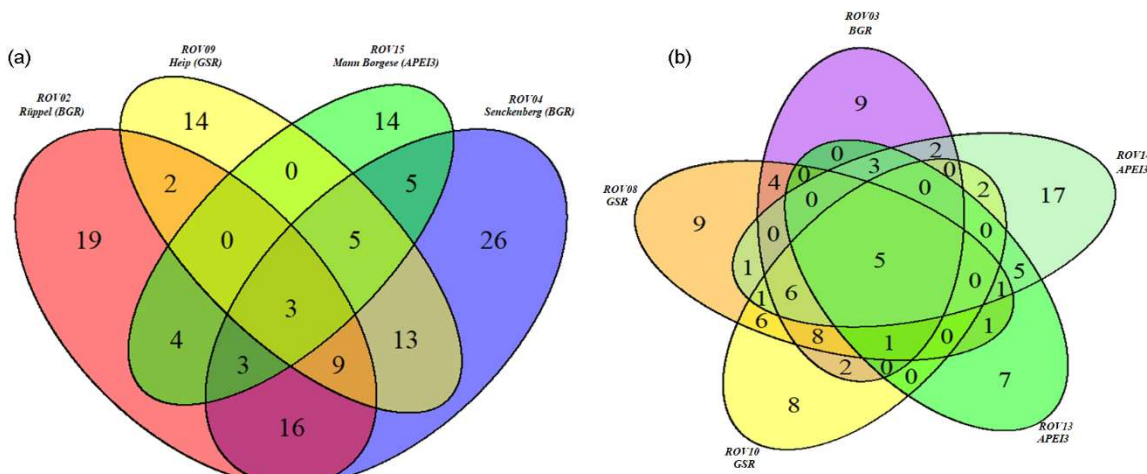


643
644 Fig. 4. Species accumulation (upper panel, a and c) and rarefaction curves (lower panel, b and d) for
645 the seamount ($n=4$) and nodule field ($n=5$) transects. Seamount dives: ROV02= Rüppel (BGR),
646 ROV04=Senckenberg (BGR), ROV09=Heip (GSR), ROV15=Mann Borgese (APEI3) in the lower left
647 panel (b). Nodule field dives: ROV03 was carried out in the BGR area, ROV08 and 10 in the GSR area
648 and ROV13 and 14 in the APEI3, presented in the lower right panel (d). Sample size is the number of
649 individuals. Vertical line in the lower panel shows sample size=100.

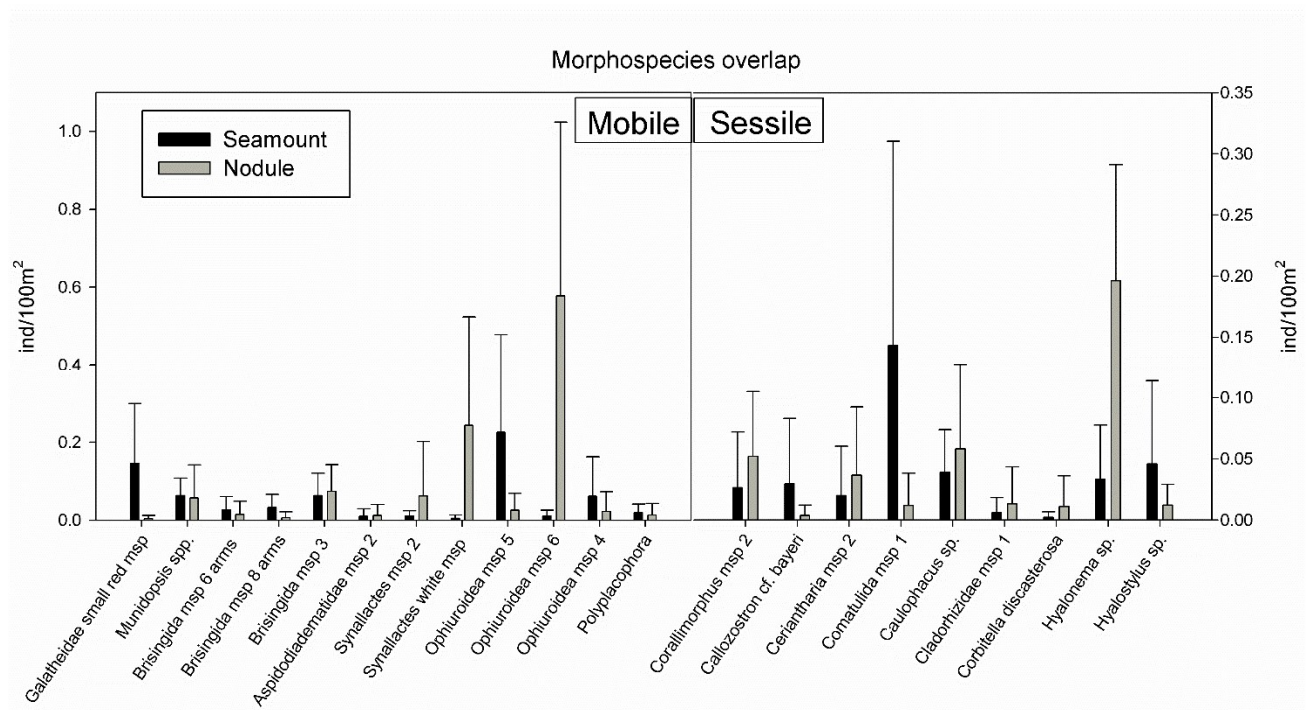
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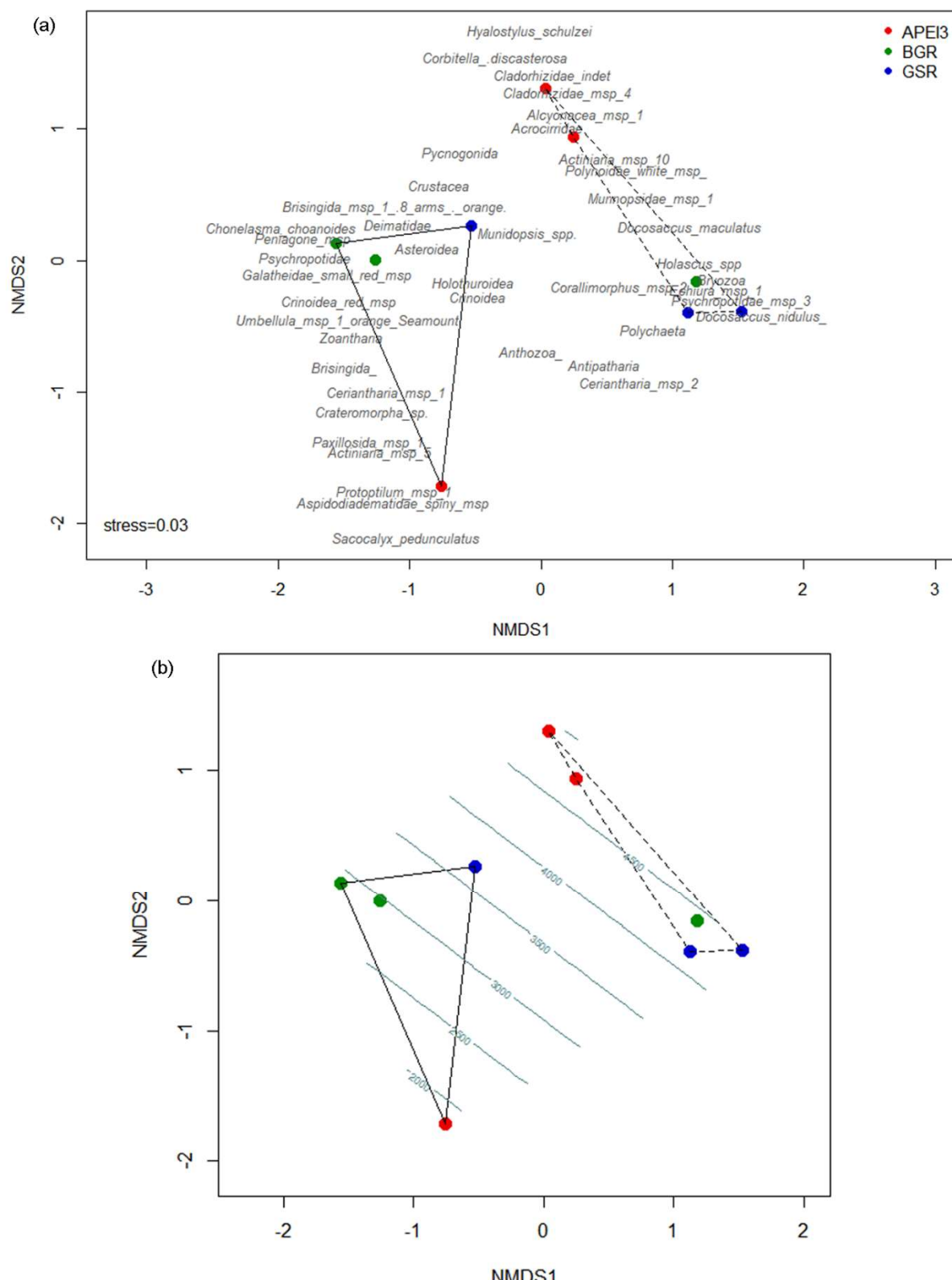
653
654 Fig. 5. A Venn diagram showing the unique and shared morphospecies among seamount video
655 transects. Values are indicative rather than absolute due to different transect lengths and
656 differences in richness. Left panel (a) features seamount transects and the right panel features the 5
657 nodule field transects. Colour codes were adapted among panels, with APEI3 nodule transects in
658 green, related to Mann Borgese seamount transect. BGR (ROV03) transect was purple in
659 correspondence to BGR seamount transects (red=Rüppel and blue=Senckenberg). GSR transects
660 (ROV08 and 09) were shades of yellow.



666

667 Fig. 6. Morphospecies present in both seamounts and nodule field transects and their average
668 density (ind./100m²) and standard deviation per ecosystem.

669



670

671 Fig. 7. nMDS-plot with faunal densities and Bray-Curtis distances. Upper panel (a) presents the
672 grouping of the video transects based on their faunal composition and lower panel (b) features the

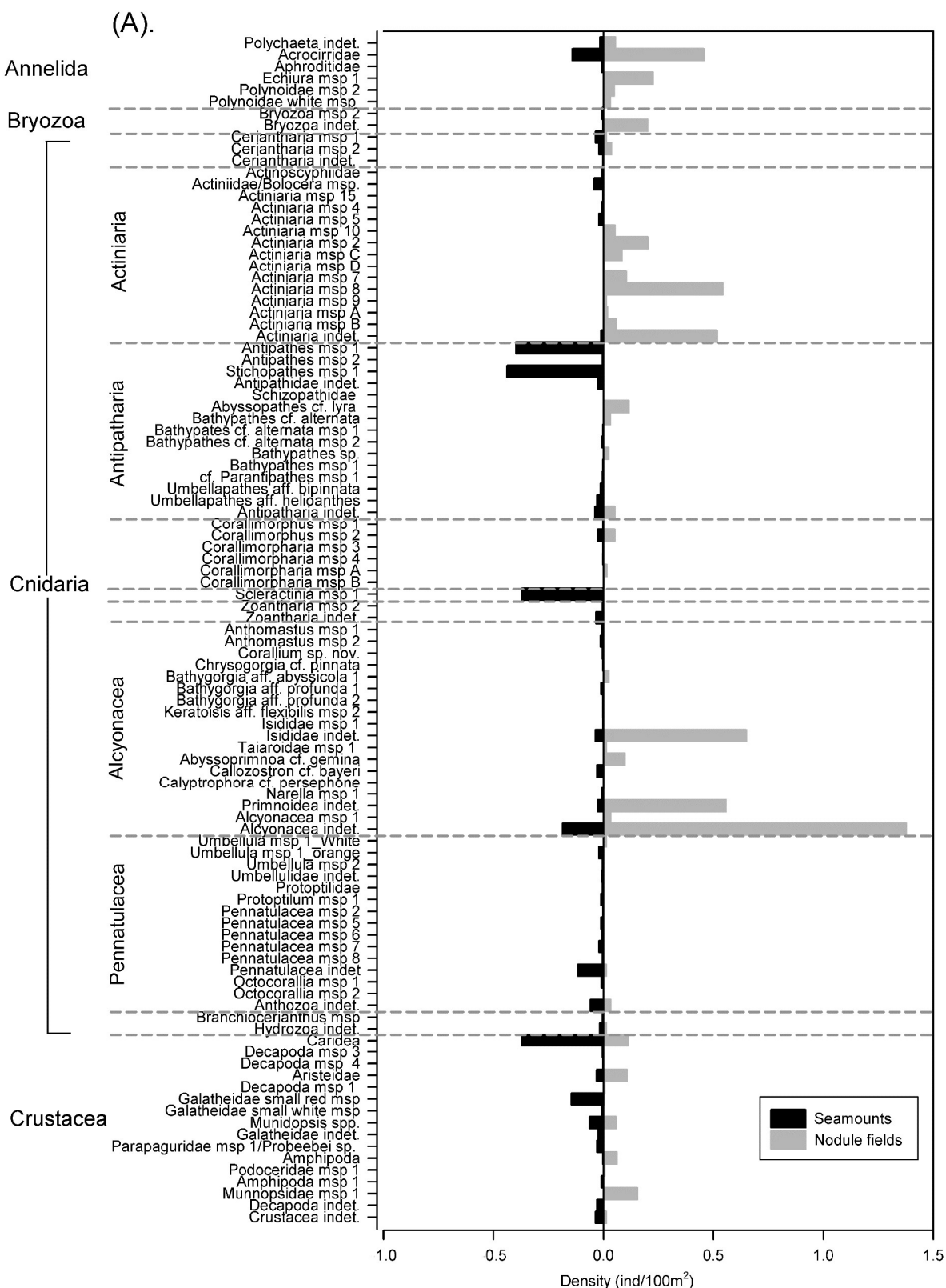
673 same plot but with depth as a vector fitting. Dotted lines group the nodule transects while the full
674 line groups the seamount transects.

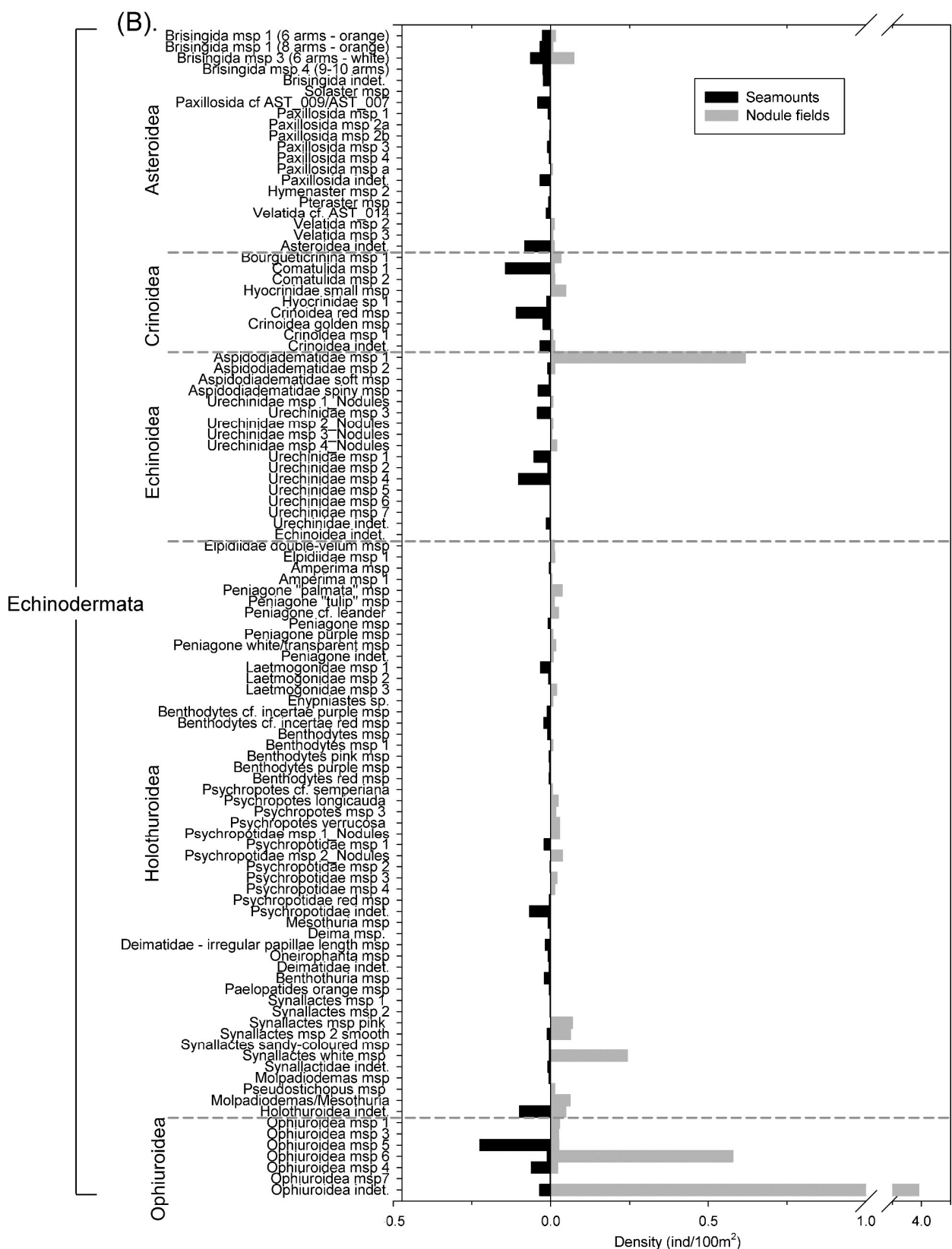
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Appendix





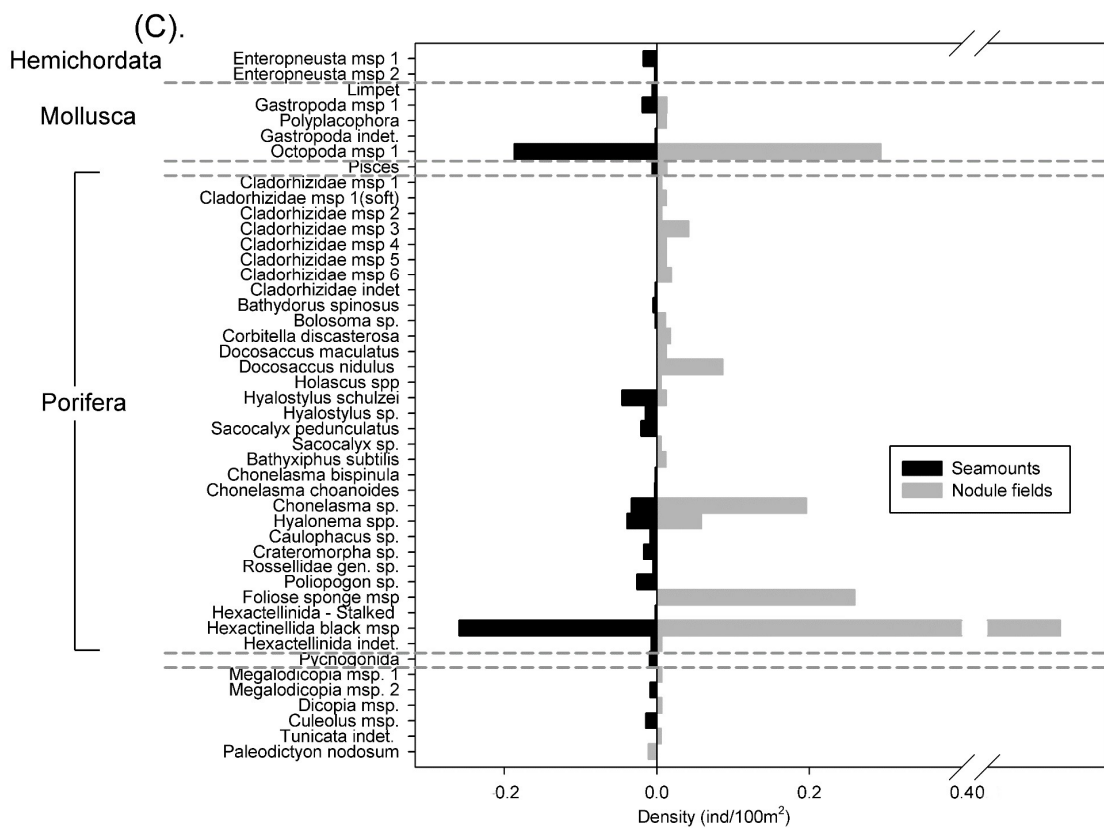


Fig. A1. Back-to-back histogram comparing average densities of morphospecies and taxa (ind/100m²) for seamount (n=4) and nodule field (n=5) video transects. (A) Annelida, Bryozoa, Cnidaria and Crustacea, (B) Echinodermata and (C) Mollusca, Porifera, Hemichordata and Chordata (Tunicata).