Approaching the dDiversity and density relationships between dilemma of the 1 2 lebensspuren<u>and</u>-tracemaking organismser tandem: a study case from abyssal 3 **Northwest Pacific** Olmo Miguez-Salas^{1,2*}, Angelika Brandt^{1,3}, Henry Knauber¹ and Torben Riehl¹ 4 5 ¹Department of Marine Zoology, Senckenberg Research Institute, 60325 Frankfurt, 6 7 Germany. 8 9 ²Departamento de Estratigrafía y Paleontología, Universidad de Granada, Av. 10 Fuentenueva 18002, Granada, Spain 11 ³Department of Biological Sciences, Institute of Ecology, Evolution and Diversity, 12 Johann Wolfgang Goethe University Frankfurt, Max-von-Laue-Str. 13, Frankfurt, 60438, 13 14 Germany 15

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17 Abstract

18 In the deep -sea, the-interactions between benthic fauna and seafloor sediment. 19 substrate mainly primarily occurs through bioturbational processes which that can be preserved as traces (i.e., lebensspuren). Lebensspuren are common features of deep-20 21 seafloor landscapes and are usually-more abundant than the organism that produce them 22 (i.e., tracemakers), rendering them promising proxies to inferfor inferring biodiversity. 23 The density and diversity relationships between lebensspuren and benthic fauna are to the present dayremain unclear and contradicting correlations hypotheses have been proposed 24 suggesting (i.e., negative, positive, or even null correlations). To approach test these 25 26 variable -correlations hypotheses, in this study-lebensspuren and-, tracemakers (specific 27 epibenthic fauna that produce these traces), degrading fauna (benthic fauna that can erase 28 lebensspuren), and <u>benthic</u> fauna in general were characterized taxonomically at eight deep-sea stations in the Kuril Kamchatka Trench area; together with two novel categories: 29 30 tracemakers (specific epibenthic fauna that produce these traces) and degrading fauna (benthic fauna that can erase lebensspuren). No general correlation (over-all study area) 31

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could be observed between diversities of lebensspuren, tracemakers, degrading fauna and 32 33 fauna. However, a diversity correlation was observed at between specific stations, showing both negative and positive correlations depending on: 1) the number of unknown 34 35 tracemakers (especially significant for dwelling lebensspuren); and 2) the lebensspuren with multiple origins; and 3) tracemakers that can produce different lebensspuren. 36 Lebensspuren and total faunal density were not correlated. However, lebensspuren 37 density was either positively or negatively correlated with tracemaker densities, 38 depending on the lebensspuren morphotypes. A positive correlation was observed for 39 resting lebensspuren (e.g., ophiuroid impressions, Actinaria circular impressions), while 40 41 negative correlations were observed for locomotion-feeding lebensspuren (e.g., echinoid trails). In conclusion, lebensspuren diversity may be a good proxy for tracemaker 42 43 biodiversity when the relationship lebensspuren-tracemaker tandem can be reliable 44 characterized.; and l_Lebensspuren-density correlations vary depending on the specific lebensspuren residence time, tracemaker density and associated behaviour (rate of 45 46 movement). Overall, we suggest that lebensspuren density and diversity correlations 47 should be done with tracemaker rather than with general benthic fauna. , but oOn a global scale abiotic (e.g., hydrodynamics, substrate consistency) and other biotic factors (e.g., 48 49 microbial degradation) may also play an important role.

50 Introduction

Neoichnology studies the interactions between animals and substrates (i-e.g., bioturbation processes) in modern environments_-as well as the biogenic sedimentary resulting from these interactions well as their final products, the so-called lebensspuren (German for "life traces"; e.g., faecal casts, trails, mounds, burrows) (Ewing and Davis, 1967; Gage and Tyler, 1991). In the marine realm, -lebensspuren analyses are usually images based (e.g., Bell et al., 2013; Miguez-Salas et al., 2023). -Lebensspuren are highly-precise

portraits of the diverse linkages between environmental conditions and the animal 57 58 responses to them. Thus, neoichnological analyses provide offers a useful tool set for 59 deducingto infer environmental factors not only inin both contemporary and past 60 environments but also deliver evidences to past environments through comparisons 61 between lebensspuren and trace fossils (Buatois and Mángano, 2011). However, neoichnology as a field is not yet as developed as paleoichnology (i.e., trace fossil 62 research), and most quantitative studies are restricted to shallow marine environments 63 and tank experiments (e.g., shoreface, foreshore, marginal marine settings) (La Croix et 64 al., 2022 and references therein). Even though the abyssal zone (i.e., 3500-6500 m deep) 65 66 represents the largest marine ecosystem and covers approx. 75% of the seafloor (Ramirez-Llodra et al., 2010; Watling et al., 2013), neoichnological analyses are scarce and limited 67 by the , mainly due to the cost of observation and sampling procedures (e.g., Heezen and 68 69 Hollister, 1971; Przesławski et al., 2012; Bell et al., 2013; Miguez-Salas et al., 2022). As suchThus, neoichnological analyses are emerging e-as a promising tool for to 70 71 enhancefurthering our understanding of deep-sea environments and faunal-sediment 72 interactions.

73 Quantitative marine ecological research comprises two main components -74 dDiversity and density analyses are two main components of quantitative marine ecological research-(Halpern and Warner, 2002). Deep-sea neoichnological studies have 75 76 addressed diversity and density characterizations by considering all identified lebensspuren as morphotypes as "species" (Przesławski et al., 2012; Bell et al., 2013). 77 78 However, tracemaker (i.e., the benthic organisms that produce the observed lebensspuren) 79 diversity and density have been approached from a generalist perspective as megafauna, 80 epifauna, or lebensspuren-forming epifauna (Young et al., 1985; Dundas and Przeslawski, 2009; Przesławski et al., 2012; Bell et al., 2013). 81

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82 Early deep-sea neoichnological studies suggested suggested a correlation between that 83 lebensspuren diversity is proportional toand faunal diversity (Kitchell et al., 1978, Young et al., 1985). However, more recent studies have shownshow -no significant correlation 84 85 between epifaunal and lebensspuren richness (Przeslawski et al., 2012) and that lebensspuren diversity was not similar to that of epifaunal lebensspuren-forming diversity 86 (Bell et al., 2013). Bell et al. (2013) stated that "improvements in imaging technology 87 allow more refined classification of lebensspuren and species organisms, which may 88 affect the strength of the correlation between faunal and lebensspuren diversity, compared 89 with the more direct proportionality of faunal and lebensspuren diversity demonstrated in 90 91 earlier studies". Thus, in deep-sea research, diversity comparisons based on more precise taxonomic tracemaker identification and differentiation are a pending task, promising a 92 93 deeper understanding of the dependencies between fauna and lebensspuren variability.

In the case of lebensspuren density, earlyEarly studies revealed found an inverse 94 relationship with between lebensspuren and faunal density (Kitchell et al., 1978, Young 95 96 et al., 1985; Gerino et al., 1995). These studies suggested that this relationship is related to the fact that lebensspuren formed in low biomass regions have the capacity to persist 97 for a long time (high residence time), ultimately leading to a steady increase of the 98 99 lebensspuren density through accumulation. Nevertheless, recent data seemed to conflict 100 with this initial assumption. Przesławski et al. (2012) observed no discernible correlation 101 between that lebensspuren and epifaunal abundance-. Contrastingly, do not have any 102 relationship; and, contrastingly, Bell et al. (2013) found-reported a strong positive 103 relationship-correlation between lebensspuren and faunal densities (see Fig. 10 in Bell et 104 al., 2013). The results from These newer results Bell et al. (2013) suggest show that 105 megafaunal activity may not be the onlymight not be the only factor influencing the 106 preservation or destruction of 1-significant factor for lebensspuren destruction or

107 preservation. Small scale biotic factors (e.g., microbial degradation), as well as abiotic 108 factors (e.g., hydrodynamic regimes, sedimentations rates, sediment composition) may potentially_limit_ lebensspuren residence time and density at different across different 109 110 spatial scales in the deep -sea (Wheatcroft et al., 1989; Smith et al., 2005; Miguez-Salas 111 et al., 2020). In summary, earlier investigations and their contradicting results strong variability in the few previous studies and conflicting conclusions drawn from these 112 highlight that neoichnology and its fundamental concepts are still in their infancy-early 113 stages and needwarrant further exploration investigation. 114

115 Despite the presence of manyprevalence of lebensspuren on the deep seafloor (Heezen and Hollister, 1971), only a very-few organisms are recognized in the process of forming 116 117 these features. Thus, understanding the density-diversity relationship between lebensspuren and benthic megafauna may help decipher variability of the former 118 indirectly (i.e., without having seen the organisms). The research presented here aims to 119 compare diversityies indices and densities of lebensspuren, specific tracemakers (specific 120 121 organism that produce them), degrading fauna (benthic fauna that can erase lebensspuren), and megabenthic fauna from the Northwest Pacific Abyssal Plain, near the 122 in the direct vicinity of the Kuril Kamchatka Trench (KKT) (Fig. 1). By conducting a 123 124 detailed classification of both lebensspuren and tracemakers, this research wants to go 125 one step further with the main objective to test how previous diversity and density 126 relationships hypotheses about the relationship vary from previous results where between 127 the variability of lebensspuren was compared with and fauna diversity in a much coarser 128 taxonomic resolution. - Thise geographic region has been is well studied studied 129 extensively, with research dating back from eleven -as it has a long research history that 130 began with eleven expeditions onboard aboard theof R/V Vityaz (Russian expeditions; 131 1949, 1953 and 1966) to more recent research efforts aboard the R/V Sonne. - and was

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132	(further extended during recent campaigns with R/V Sonne (German Russian
133	expeditions; KuramBio I (2012) and KuramBio II (2016)). All of these These expeditions
134	have resulted significantly contributed to establishing the most comprehensive taxonomic
135	foundation for fauna in this region in one of the best taxonomic baseline of the fauna (e.g.,
136	Zenkevitch et al., 1955, Zenkevitch, 1963; Belyaev, 1983; Brandt and Malyutina, 2015;
137	Brandt et al., 2020; Saeedi and Brandt, 2020). among others).
138	Material and methods
139	Study sites, data acquisition, and video analysis
140	The joint German-Russian expedition KuramBio 1 (Kurile Kamchatka Biodiversity
141	Studies) aboard on board of the RV Sonne (cruise SO223) explored to the KKTKuril-
142	Kamchatka Trench and its adjacent abyssal plain from took place between JJuly 21 st st to
143	and September $\theta_{\lambda}^{\text{th}}$ = 2012 (Brandt and Malyutina, 2012). During the KuramBio 1
144	expedition, 13 Ocean Floor Observation System (OFOS) deployments were conducted
145	(Table 1). The intent of these deployments was to) to use the OFOS to study eleven deep-
146	sea stations spanning a range of depths (4,868-5,768m), located between 34°-48°N and
147	147°-157°E (Fig. 1)-with video cameras. Stations 1, 2, and 5-11 were located in the
148	abyssal plains adjacent to the KKT while stations -3 and 4 were located at the upper slope
149	of the KKT, and stations 1, 2, and 5-11 in the adjacent abyssal plains (Fig. 1) (Table 1).
150	The depths of the stations ranged from 4,868 m to 5,768 m.
151	The OFOS was lowered into the water at the C <u>Conductivity, Temperature, and</u>
152	Depth TD-position For the first 300 meters, the OFOS was lowered ating was
153	conducted with 0.5 m/s. Then, , and then the speed was increased to 0.8 m/s while the
154	ship was kept in position. The speed of the OFOS was reduced to 0.5 m/s once it was 500

- 155 <u>m above the ground, and further reduced to 0.3 m/s once it was 200 m above the ground.</u>
- 156 <u>Once the bottom was visible</u>, At 500 meters above ground, the speed was reduced to 0.5

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Con formato: Inglés (Estados Unidos) Con formato: Sangría: Primera línea: 1,25 cm 157 m/s, and further reduced to 0.3 m/s at 200 meters above ground. the winch lowering the 158 OFOS was stopped and the ship resumed steaming at 0.5 knots in a direction chosenAs soon as visual contact with the bottom was established, the winch was stopped. The ship 159 160 started moving with 0.5 knots above ground in the appropriate direction, which was chosen depending on the water current and wind situation. Then, the winch operator 161 manually kept the OFOS at an appropriate distance from the seafloor to observe the 162 seafloor benthos. Two laser pointers having with a distance of 1010 cm between each 163 other werethem were used as a scale. The first four deployments were aborted due to 164 technical problems, affecting stations 1-3 (Table 1). Thus, limited video footage was 165 166 obtained. Moreover, station 7 haslacked no HD video and was therefore not considered for (i.e., this station is not considered for the current analysis). All technical workaspects, 167 including pre-deployment preparation and post-deployment maintenance (i.e., before and 168 169 caretaking after (including video download) were managedthe deployment was conducted by the scientific-technical service ("WTD", Wissenschaftlich-Technischer 170 171 Dienst, Jörg Leptien, Reederei).

172 At each station, Sstill images were extracted from the OFOS videos were extracted 173 at a frequency of onfrom the OFOS videos at a rate of one frame per five seconds 174 (Miguez-Salas and Riehl, 2023a). These still images were then subsequently further subsampled to delete frames that were out of focus to minimize frame overlap. Out of focus 175 176 frames were defined as frames showing the OFOS moving up and down due to ocean 177 swell. - as the rolling of the ship in the ocean swell resulted in an up and down movement 178 of the OFOS and to reduce overlap between frames. Then, 50Fifty frames per station (400 still images in total), covering a seafloor area of 878 m² (109 m² per station approx.), 179 180 were randomly selected frames per station were studied (400 still images in total)and uploaded to the BIIGLE 2.0 software for later annotation and measurements 181

183	approx.). These still images were uploaded to the BHGLE 2.0 software for later
184	annotation and measurements (Langenkämper et al., 2017)SpecificSpecific frames were
185	treated with Fiji software (Schindelin et al., 2012) to enhance the visibility (CLAHE tool)
186	of certain lebensspuren features (Miguez-Salas et al., 2019).
187	Lebensspuren classification and tracemaker identification
188	Lebensspuren morphotypes were categorized in terms of inferred tracemaker behaviour
189	during the construction their formation, their morphology, and the taxonomic origin of the
190	tracemaker-staxonomic origin. The behavioural classification was adapted from
191	Seilacher's (1954) categories for marine lebensspuren: i. Resting (imprints of stationary
192	animals); ii. Locomotion-feeding (sediment displaced by the movement of deposit feeders
193	and surface sediment disturbances formed as organisms are foraging); iii. Wasting (e.g.,
194	faecal casts, pellets); and iv. Dwelling (e.g., mounds and burrows). The morphological
195	classification followed previous morphological names (e.g., Ewing and Davis, 1967;
196	Young et al., 1985; Dundas and Przesławski, 2009; Przesławski et al., 2012; Althaus et
197	al., 2015), where such exist. Morphological features measured included in the
198	classification were length, width, and diameter. Lebensspuren with unclear morphology
199	and origin (e.g., degraded faecal casts, trails with diffuse outlines) were not considered in
200	thisexcluded from analyses-study. Additionallylso, lebensspuren and fauna smaller than
201	1 cm (macrofauna and smaller) were also excluded from analyses as the resolution of the
202	still images is below high-definition (<1280x720 pixels). lebensspuren and fauna smaller
203	than 1 cm (macrofauna and smaller) have not been considered in this study. Hence, this
204	study focus <u>es only ses</u> on megafauna (i.e., fauna > 1 cm), which is implied whenever
205	throughout this study when fauna is mentioned in this study from hereon, this point

(Langenkämper et al., 2017)., covering a seafloor area of 878 m² (109 m² per station

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forward.

207	This study aimed for the The species rank, which is fundmanteal, gold standard
208	the basic, gold-standard taxonomic level to which ecological studies generally aspire
209	was aimed for in the discrimination of taxa. However, names of described species could
210	not be attached to these taxa in the majority of our image-based observations for reasons
211	of limitation in the image quality and the general difficulty to observe diagnostic
212	characters in <i>in-situ</i> photographs. SoAs such, oOpen nomenclature has been used instead
213	for megafauna taxonomic identificationfollowing the recommendations for image
214	based identifications proposed laid out by Horton et al. (2021). All differentiated
215	morphotypes are henceforward referred to as "species" for simplicity. Then, Ffauna has
216	been grouped into different categories for comparisons with the diversity and density of
217	lebensspuren: 1) tracemakers (fauna that has been clearly recognized as maker of a trace);
218	2) degrading fauna (fauna that can affect lebensspuren density negatively by eroding the
219	seafloor); and 3) benthic fauna (all fauna identified in the still images).

220 Statistical analysis

For statistical analysis, all identified lebensspuren and fauna morphotype were treated as "species". Diversity indices (Shannon–Wiener H' (loge) and Simpson's D) and evenness (J') were calculated for the four groups: lebensspuren, tracemaker fauna, degrading fauna, and fauna. As the data from all groups show non-parametric distribution throughout for all stations, diversity variability among stations was tested using Wilcoxon signed-rank test (considering all groups and all indices). Then, the The Spearman rank correlation was used to test the relationships between the diversity indices of all groups.

For density correlations (Spearman rank correlation), since <u>As</u> the number of frames was
 the same (i.e., same observation area), the <u>analysesdensity correlations analyses</u> were
 conducted considering theperformed by separately considering the total density per
 station of all groups individually for each group at every station. Additionally,

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232 lebensspuren and tracemakers densities were further subdivided into three groups:into 233 wasting, resting and locomotion-feeding. (Ddwelling was not considered because as the tracemakers of most dwelling lebensspuren are unknown). 234

Multivariate analyses was conducted tTo investigate potential differences within-235 the four groups (lebensspuren, tracemakers, degrading fauna, and total benthic fauna 236 237 groups.) between stations, multivariate analysis was conducted. First, a square root transformation was carried out to give less weight to the more abundant species taxa of 238 organisms and lebensspuren morphotypes. Then, differences in the composition of the 239 240 four groups between stations were assessed using hierarchical cluster analysis and displayed as non-metric multidimensional scaling plots (n-MDS). Both plots were 241 constructed using the Bray-Curtis similarity index. All statistical procedures were 242 conducted using PAST v. 4.12 (Hammer et al., 2001). 243

Results 244

245 A total of 9,426 lebensspuren were identified and classified from 400 still images, corresponding to 23 morphotypes associated with dwelling, wasting, resting, and 246 247 locomotion-feeding behaviours (Fig. 2; Table 2) (for raw dataset report at each station consult: Miguez-Salas and Riehl, 2023b). The fauna comprised a total of 4,009 individual 248 animals that were classified into 93 different species(terminal)-taxa, of which 66 were 249 classified as degrading fauna and 43 as tracemakers (with 790 and 676 individuals 250 respectively) (Table 3;- Miguez-Salas and Riehl, 2023bSupplementary file 1). Linking 251 252 dwelling lebensspuren with tracemakers was mostly impossible proved to be challenging, 253 with the exception except for of exception of rare and ambiguous cases where vermiform organisms, most likely polychaetes, partially emerged from paired burrows (Fig. 2P). 254 255 Tracemaker identification was possible in the majority of most of the cases for wasting lebensspuren., however, Jit is common that different for different tracemakers to produce 256 10

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257 the same lebensspuren morphotypes and it is also common for one tracemaker (taxon) to 258 produce that several lebensspuren morphotypes of lebensspuren are produced by one 259 tracemaker species taxon (see Table 2). However, in the case of cf. Elpidia - the most 260 abundant tracemaker of station 4 (see Miguez-Salas and Riehl, 2023bSupplementary file 261 1) — the complete characterization of its associated rounded faecal cast (smaller than 1 cm) was impossible due to image resolution limitations. Tracemaker identification of 262 locomotion-feeding lebensspuren was mostly possible except for mounded trails which 263 have been produced by endobenthic organisms. However, as for wasting lebensspuren, 264 also in this case different tracemakers can be responsible for similar trails (see Table 2). 265 266 Tracemaker identification of resting lebensspuren has been possible in most cases. wWas possible for most of the cases. 267

The Wilcoxon signed-rank test revealed that for all groups the median diversity* 268 was significantly different between stations for all groups and was, being lower at lowest 269 at stations 9 and 11 (Fig. 3). Moreover, faunal diversity showed a standard deviation three 270 271 orders smaller than the values reported for lebensspuren, tracemakers, and degrading fauna; showing that faunal diversity was more consistent among sites that the other 272 diversity indices. Lebensspuren diversity indices (Shannon-Wiener, Simpson's and 273 274 Evenness) of the over-all KKT area (considering all the eight stations together) showed 275 no correlation with comparable diversity indices from the other three groups the other three groups (tracemakers, degrading fauna, and benthic fauna). The only strong diversity 276 correlation resulting from the Spearman rank analysis was between tracemakers and 277 degrading fauna (R^2 >0.88, p<0.01). 278

The density correlation matrix revealed no significant correlation between the fauna and the other groups (see Fig. 4). The degrading fauna showed a positive correlation with tracemaker and <u>waste_wasting</u>-tracemakers densities. Also, tracemakers and <u>waste</u> Con formato: Sangría: Primera línea: 1,25 cm

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wasting tracemakers densities are positively correlated (Fig. 4). <u>AIn case of the</u>
lebensspuren data, a positive density correlation was obtained between lebensspuren and
wasting lebensspuren as well as resting lebensspuren and resting tracemakers. <u>while aA</u>
negative correlation was observed for locomotion-feeding lebensspuren and their
tracemakers (Fig. 4).

287 Inter station similarity of ILebensspuren assemblages composition wereas generally similar among stations high (Fig. 5 A), ranging from 75-82% similarity in the cluster 288 analysis. The n-MDS showed that lebensspuren assemblages from stations 5, 6, 8, and 10 289 290 are different from the trench (stations 3 and 4) and the southern stations (stations 9 and 11) (Fig. 6 A). The southern stations were less diverse, similar (82% similarity; Fig. 5A) 291 292 and dominated by rounded faecal casts produced by Scotoplanes spp. The trench stations were characterised by diverse and slightly less similar assemblages (75% of similarity) 293 dominated by dwelling lebensspuren (e.g., paired, lined or cluster burrows), knotted 294 faecal casts (Peniagone spp.), ophiuroid impressions (Ophiuroidea), circular impressions 295 296 (Actinaria) and thick-M-ridged trails (Asteoroidea and Echinocrepsis spp.). Stations 5, 6, 8, and 10 showed diverse lebensspuren assemblages dominated by smooth (cf. 297 298 Benthodytes, Psychropotidae) and coiled --curly faecal casts (Psychropotidae), rosette-299 shaped traces and thick-flat trails (Asteroridea, cf. Benthodytes, Psychropotidae) (Fig. 6 A). 300

The hierarchical cluster diagram for tracemakers, degrading fauna and fauna showed less similarity between stations than it was the case for lebensspuren, especially for tracemakers and degrading fauna (values ranging from 20–55% similarity in the cluster analysis) (Fig. 5 B–D). However, the trench stations (Stations 3 and 4) and the southern stations (Stations 9 and 11) seemed to have similar compositions respectively. The low inter-station similarity of tracemakers, degrading fauna and fauna assemblages Con formato: Sangría: Primera línea: 1,25 cm

was also reflected in the n-MDS plots where the spacing between stations wasconsiderably higher than in the lebensspuren plot (Fig. 6 B–D).

309 Discussion

310 The obtained results from the KKT area reveal that the relationship between lebensspuren, 311 tracemakers, and fauna is may be more complicated than initially previously hypothesized (Kitchell et al., 1978; Young et al., 1985) and may follow the complex puzzle exposed in 312 recent research (Przeslawski et al., 2012)-. On the one hand, aWhile a general null 313 diversity correlation has been observed between lebensspuren, tracemakers and fauna, --314 On the other hand, density correlations seem to be morphospecific (e.g., depending on 315 the lebensspuren-associated behaviour). But to what extent do the obtained results 316 317 contradict or corroborate previous results and what are the limitations when addressing the diversity and density of lebensspuren? 318

319 Fauna, tracemakers and lebensspuren diversity: a complex relationship

Previous comparisons between lebensspuren and faunal diversity have given rise to 320 321 different contrasting hypotheses. Though pPioneering research showed positive 322 correlations (e.g., Kitchell et al., 1978, Young et al., 1985), -- Later on, several-studies showed no correlation at all (e.g., Tilot, 1995; Turnewitsch et al., 2000; Przeslawski et 323 324 al., 2012). All-Tthese studies have in commonshare a common approach in which that the 325 diversity comparisons was were addressed conducted from a general broad perspective, especially for tracemaker organisms. Comparisons were done eitherconducted using 326 327 considering megafaunal species taxa (Young et al., 1985), epifaunal species taxa (Przeslawski et al., 2012) or certain taxonomic groups of organisms (e.g., fish, 328 holothurians, crinoids; Kitchell et al., 1978). Only Bell et al. (2013) approached the 329 330 comparison between lebensspuren and fauna in greater detailby-considering groups of 331 lebensspuren-forming epifauna, and using indices to quantify lebensspuren diversity 332 (e.g., Simpson, Shannon-Wiener). Their analyses , discoveringrevealed that "Lebensspuren diversity was generally high and not similar to that of lebensspuren-333 forming faunal diversity" (Bell et al. 2013). However, the links between specific 334 335 tracemakers and their lebensspuren and the subsequent tracemaker diversity indexes are 336 missing in Bell et al.'s (2013) study. In this study we have tried to attempt to close this knowledge gap by comparing conducting a comprehensive comparision comparison of the 337 lebensspuren diversity. We examine <u>with</u> not only the faunal diversity but also 338 encompass the tracemaker and degrading fauna (i.e., fauna that may alter the lebensspuren 339 340 assemblage by erosion/degradation). Our results show that lebensspuren diversity (Simpson, Shannon-Wiener, and Evenness) is not related to fauna, tracemaker or 341 342 degrading fauna diversity. This findingThese findings seem_s-to corroborate the latest 343 results of a non-existent correlation (Przeslawski et al., 2012; Bell et al., 2013). However, can, but can this lack of correlation be expected in all deep-seae settings? 344

345 Before answering this question, This question cannot be answered without considering the limitations of quantifying deep-sea lebensspuren diversity, which is riddled with 346 347 problems. These problems include -should be considered. There are several problems 348 when it comes to quantifying lebensspuren diversity (e.g., image resolution, camera 349 systems, unknown lebensspuren, unknown tracemakers, observation scale, trace degradation, but the most important is linked to their genesis. In other words, the same 350 351 lebensspuren morphotypes (or indistinguishable lebensspuren) can be produced by 352 different tracemakers and one tracemaker can produce different lebensspuren (see Table 353 2). For example, in case of thisthis study, several different holothurians (e.g., cf. 354 Pseudostichopus, Psychropotes, Synallactidae, Benthodytes) could have produced the 355 smooth faecal casts. Contrastingly, -could have been produced by different holothurians

356 (e.g., ef. Pseudostichopus, Psychropotes, Synallactidae, Benthodytes) and Psychropotes 357 can be linked to the production of coiled-curly and smooth faecal casts as well as thick flat trails (Fig. 2G). Thus, when comparing their diversity, the basis that each it should be 358 359 taken into account that lebensspuren morphotypes may not be related to one specific 360 species-taxon and vice versa, should be considered. However, the factwhile that in our study general lebensspuren diversity in the present study did not correlate with tracemaker 361 diversity, this does not mean that this will be the case in all deep sea settings that the same 362 applies to all deep-sea settings. It is possible that a more precise characterization of if 363 thethe tracemaker-lebensspuren relationship tandem can be characterised more precisely 364 365 or if tracemakers produce just one specific lebensspuren morphotypes may reveal a different corelation correlation in other settings. -366

367 In our study area, We observed different correlations between tracemakers and + lebensspuren could be observed when comparing the diversity among specific 368 stations. For example, when restricting the comparison to of the southern stations (stations 369 370 9 and 11), a correlation was observed between) using Simpson and Shannon-Wiener indexes of showed a correlation between tracemakers and lebensspuren diversities (Fig. 371 372 3). This was due to the fact This can be attributed to the traces dominating the that the 373 assemblage is dominated by traces for which we have been able to identify 374 thesuccessfully identified tracemakers (e.g., rounded faecal casts of Scotoplanes). On the 375 contrary, when focussing on the trench comparison of the trench stations (stations-3 and 4 376 using Simpson and Shannon-Wiener indexes revealed), a negative correlation could be observed between Simpson and Shannon Wiener indexes of tracemaker and lebensspuren 377 378 diversities (Fig. 3). This could can be attributed to the relatively large gap in our data 379 regarding the origin of most traces of the lebensspuren assemblage. -(Sstations 3 and 4 380 have a high abundance of dwelling lebensspuren (see Table 3) $_{2^{\frac{1}{7}}}$ single burrows, mounds,

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cluster burrows for which tracemakers are unknown) and dominant tracemakers (*Elpidia*)
 whose traces cannot be correctly quantified due to image resolution limitations (small
 rounded faecal casts).
 <u>-Also, the existence of unknown lebensspuren and tracemakers will</u>
 <u>contribute to the</u>contributes to the correlation variability.

The enhancement of image resolution and the increase of deep-seafloor area 385 386 covered by still image surveys may allow thee improvement of lebensspuren classification and their tracemaker identification. There is a lot of room for ample room 387 for improvement, especially with regard to locomotion and feeding lebensspuren. High 388 389 definition still images will allow researchers to characterize, for example, small morphological features of trails (e.g., podia marks from asteroids, echinoid spine 390 391 impressions), allowing for a much more detailed classification than what-could be 392 achieved within for this study. The use of artificial intelligence, which seems to be a verya seemingly promising tool in the assistance of benthic fauna recognition in imagery 393 analyses, appear to haves a restricted applicability in the characterization of lebensspuren. 394 395 This is because lebensspuren -since they are constructed with sediment which usually have that has the same texture as the seafloor (i.e., background colour). In the case of 396 dwelling lebensspuren diversity, comparison is significantly more complicated. because 397 398 <u>*t*</u><u>T</u>race morphology is largely hidden below the seafloor surface, reducing the possibility</u> 399 ability to differentiate between different-various burrow morphologies. Additionally, 400 while tracemakers are mostly unknownlargely unidentified due to a their predominantly 401 endobenthic lifestyle (e.g., Brandt et al., 2023). Furthermore Finally, it is worth noting 402 that burrows and other dwelling lebensspuren also could potentiallymay have multiple 403 potential origins (e.g., a paired burrow can be produced by multiple species of polychaetes 404 or bivalves).

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405 The fact that the same lebensspuren morphotypes can be produced by different 406 tracemakers and one tracemakers can produce different lebensspuren will affect the 407 establishment of a positive or negative diversity correlation. Also, the existence of 408 unknown tracemakers will contribute to the correlation variability. However, as tThe 409 obtained resultsOur results show that in specific stations where the, when the assemblage is dominated by traces with identifiable tracemakers, lebensspuren analysis emerges as 410 appears to be a- promising tool to-for predicting tracemaker diversity. Although these 411 results are promising, it is evident that much more research is needed, especially-Despite 412 of these optimistic results, it is fair to say that much more research is needed — with high 413 414 definition surveys (e.g., videos, images) —to close existent existing knowledge gaps in the relationship between lebensspuren-and tracemaker-tandem. Moreover, we emphasize 415 416 that when using lebensspuren as a proxy for biodiversity, the diversity correlation should 417 be made between lebensspuren and tracemakers, rather than with overall benthic fauna as no correlation has been observed in case of comparison with the latter. 418

419 Tracemaker and lebensspuren density: morphospecific relationship

Similar to previous research, tThe density comparisons between lebensspuren, degrading 420 fauna and total fauna revealed no correlation, similar to previous research (Przeslawski 421 422 et al., 2012). However, when the density comparisons comparing between lebensspuren 423 and tracemakers revealed a positive and negative correlation can be observed (Fig. 4). The density of locomotion-feeding lebensspuren is inversely correlated with their 424 tracemaker density while resting lebensspuren are positively correlated with their 425 tracemakers densities. These group-specific correlations conflict with previous research 426 427 that showed generally positive (e.g., Bell et al., 2013) or generally negative density correlations (e.g., Kitchell et al., 1978; Young et al., 1985). The difference with these 428 previous studies may be due to the fact that because their density comparisons considered 429

the total fauna instead of separate functional groups (see Fig. 10 in Bell et al., 2013), notconsidering their specific impact on the sediment.

Trace residence time is the period during which a trace is recognizable on the sea 432 floor before it is destroyed (Wheatcroft et al., 1989). It is commonly accepted that 433 Liebensspuren density values reflect the balance between lebensspuren formation and 434 435 lebensspuren destruction/degradation either by biotic (e.g., microbial degradation, degrading fauna, epifaunal rate of movement) or abiotic factors (e.g., hydrodynamics, 436 burial) (Wheatcroft et al., 1989). However, not all lebensspuren have the same residence 437 438 time. T. Thus, traces not actively maintained by animals are usually ephemeral features with lifespansshort-lived, lasting only of-days to weeks (e.g., faecal casts can be degraded 439 440 within 1-2 weeks; Smith et al., 2005). In contrast, while locomotion-feeding and resting lebensspuren have higher longer residence times as they are impressions on the seafloor 441 442 (see Fig. 8 in Wheatcroft et al., 1989 or Fig. 5 in Miguez-Salas et al., 2020). Very little is known about the residence time of dwelling lebensspuren, Ssome tracemakers live 443 444 inside inhabit them for their whole life throughout their entire life, while others change several-residence multiple times and their burrows get passively filled (Gage and Tyler, 445 1991). Thus, -a wide range of residence times may be expected. However, in any of the 446 447 cases since irrespective of scenario, the sedimentation rate is usually low in the deep-sea anda, dwelling lebensspuren should have higher residence time than wasting 448 lebensspuren, and similar or higher than locomotion-feeding and resting lebensspuren. 449

In case of this study<u>The</u> the density of locomotion-feeding lebensspuren (e.g.,
thick M-<u>ridged</u> trails), on the one hand, was inversely correlated with tracemaker density.
This could be attributed eto-for two reasons: 1) a high residence time of these
lebensspuren even if while the respective tracemakers may no longer be in the study area;
and 2) these lebensspuren represent a foraging behaviour in which the tracemakers tend

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to continuously search the seabed for food, often over a wide area (i.e., high rate of 455 456 movement). Thus, a large quantity of lebensspuren may be produced by a single individual tracemaker in continuous movement. The density of resting lebensspuren (e.g., 457 circular impressions, asteroid impressions), on the other hand, was in this study directly 458 459 correlated with tracemaker density. This is not surprising because even though while these lebensspuren have a high residence time, their tracemakers (e.g., asteroids, actiniarians) 460 have low rates of movement (Durden et al., 2015; 2019). In such cases, a high density of 461 resting lebensspuren should always be linked to a high density of their tracemakers. 462

463 The density correlation between wasting lebensspuren and their tracemakers showed a slightly positive but not significant correlation (Fig. 4). Maybe this is due to 464 465 the This may be attributable to the fact that in some cases we were not able to we were unable to quantify the exact number of faecal casts for some morphotypes for some cases. 466 For example, in station 4, the lebensspuren of the dominant tracemakers (*Elpidia*; more 467 than 150 specimens werespecimens identified) were innot-correctly quantified due to 468 469 image resolution limitations (small rounded faecal casts). Thus, presumably a positive density correlation between wasting lebensspuren and their tracemakers should be 470 expected. However, this assumption may be disturbed influenced by the behaviour of 471 their tracemakers , as their behaviours since their feeding activity can be expected to 472 depend on grain size, availability and quality of the nutrients among other environmental 473 factors (e.g., Jumars and Wheatcroft 1989; Ginger et al., 2001). 474

The observed variability in the lebensspuren density correlations show a complex scenario even without considering biotic and abiotic factors that cannot be characterized through still images. For <u>exampleinstance</u>, it has been demonstrated that meiofauna and microfauna have the ability to smoothen and eventually fully erase surficial biogenic structures through small scale, grain-by-grain jostling of particles (e.g., Cullen, 1973). 480 These "small" biotic processes are impossible to quantify through images, however, it has 481 to be kept in mind that these will have affected also the likely influenced lebensspuren density that we quantified for this study. Moreover, while previous studies assumed that 482 483 abiotic lebensspuren degradation rates are constant over the lebensspuren residence time 484 period (Bell et al., 2013), but-recent studies show that this may not be always true since 485 hourly spontaneous events (e.g., benthic storms) may erase the full lebensspuren assemblage (Miguez-Salas et al., 2020). The effects of abiotic factors (e.g., bottom 486 487 currents, substrate consistency) -on the density of the studied assemblages as well as those of some biotic factors (e.g., microbial degradation which cannot be characterized in a still 488 489 image) are out of the scope of this research but should be considered in future studies and 490 need to be kept in mind when interpreting seafloor images.

491

492 Conclusions

The neoichnological analysis of the KKT area reveals a general null diversity correlation
between lebensspuren, tracemakers, and fauna while density correlations vary depending
on the lebensspuren morphotypes. The further conclusions of this study are:

496	1. The fact that the The ability of various tracemakers to produce the same
497	lebensspuren morphotypes, and for a single tracemaker to produce various
498	lebensspuren morphotypes, will impact the - can be produced by different
499	tracemakers and one tracemakers can produce different lebensspuren will
500	affect the establishment of a positive or negative diversity correlation.
501	2. The existence of unknown tracemakers will contribute to the diversity
502	correlation variability. However, lebensspuren diversity may be a good proxy

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	for tracemaker biodiversity when the lebensspuren-tracemaker relationship	
504	tandem-can be reliablye characterized.	
505	3. Lebensspuren density can be positively or negatively correlated with	
506	tracemaker densities depending on the specific lebensspuren residence time	
507	and tracemaker behaviour (e.g., locomotion, resting).	
508	4. Lebensspuren-density correlations may be controlling on a wider spatial	
509	global scale by abiotic (e.g., hydrodynamics, grain size, organic matter,	
510	substrate consistency) and biotic factors (e.g., microbial degradation).	
511	Above all, we suggest that lebensspuren density and diversity correlations should	
512	be done with tracemaker rather than with general benthic fauna.	
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	-	
514	Special thanks to the German Federal Ministry of Education and Research (BMBF) for	
514 515	Special thanks to the German Federal Ministry of Education and Research (BMBF) for funding this project (PTJ, Grant 03G0223A to A. Brandt). We also thank the crew of $R.V$.	
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515 516	funding this project (PTJ, Grant 03G0223A to A. Brandt). We also thank the crew of <i>R.V.</i> Sonne and . We thank M. Malyutina, the Russian coordinator of the Russian expedition	
515 516 517	funding this project (PTJ, Grant 03G0223A to A. Brandt). We also thank the crew of <i>R.V.</i> Sonne and . We thank <u>M. Malyutina</u> , the <u>Russian</u> coordinator of the <u>Russian</u> expedition <u>M. Malyutina</u> . The research of O. Miguez-Salas was funded by a Humboldt Postdoctoral	
515 516 517 518	funding this project (PTJ, Grant 03G0223A to A. Brandt). We also thank the crew of <i>R.V.</i> Sonne and . We thank <u>M. Malyutina</u> , the <u>Russian</u> coordinator of the <u>Russian</u> expedition <u>M. Malyutina</u> . The research of O. Miguez-Salas was funded by a Humboldt Postdoctoral Fellowship from the Humboldt Foundation. This is contribution # <u>187</u> of the Senckenberg	Con formato
515 516 517 518 519	funding this project (PTJ, Grant 03G0223A to A. Brandt). We also thank the crew of <i>R.V.</i> <i>Sonne</i> and . We thank M. Malyutina, the Russian coordinator of the Russian expedition M. Malyutina. The research of O. Miguez-Salas was funded by a Humboldt Postdoctoral Fellowship from the Humboldt Foundation. This is contribution #187 of the Senckenberg Ocean Species Alliance (SOSA). We extend our gratitude to thanksDrs. Autun Purser and	Con formato
515 516 517 518 519 520	funding this project (PTJ, Grant 03G0223A to A. Brandt). We also thank the crew of <i>R.V.</i> <i>Sonne</i> and We thank M. Malyutina, the Russian coordinator of the Russian expedition M. Malyutina. The research of O. Miguez-Salas was funded by a Humboldt Postdoctoral Fellowship from the Humboldt Foundation. This is contribution #187 of the Senckenberg Ocean Species Alliance (SOSA). We extend our gratitude to thanksDrs. Autun Purser and Rachel Przeslawski for their constructive comments, which significantly contributed to	Con formato

524 Author's contributions

O.M.S., T.R., performed the data acquisition and treatment. O.M.S., T.R., and A.B., wrote
and designed the main manuscript text. O.M.S., H.K., prepared all figures and, tables, and
supplementary material. All authors reviewed and edited the manuscript at multiple stages
and approved it for submission.

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530 Availability of materials and data

531	All data generated or analysed during this study are included in this published article. The	
532	raw data used for this study is in	
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534	and density data obtained from KuramBio 2012 expedition still images (50 still images	Con formato: Sangría: Primera línea: 1,25 cm
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536	Supplementary Information file.	
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538	Competing interests	
539	The authors declare no competing interests	
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638	deep-sea stations) [Data set]. Zenodo. https://doi.org/10.5281/zenodo.10057636.		17,77 cm + 19,39 cm + 21 cm + 22,62 cm + 24,23 cm
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640			Con formato: Fuente de párrafo predeter., Fuente: (Predeterminada) +Cuerpo (Calibri), 11 pto, Color de fuente: Negro, Inglés (Estados Unidos)
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Con formato: Alemán (Alemania)

Con formato: Inglés (Reino Unido)

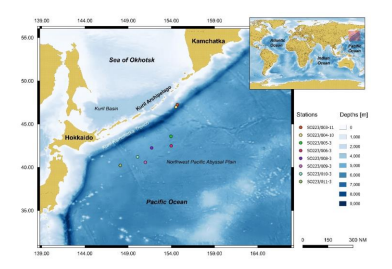
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678 Figure captions

- 679 Fig. 1 Map of the study area (Kuril-Kamchatka Trench area) and the location of the
- 680 analyzed deep-sea stations.



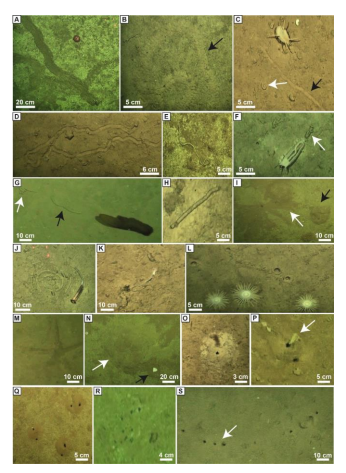
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682 Fig. 2 Examples of lebensspuren morphotypes observed and quantified in this study. A) Thick-M-ridged trail produced by Asteroidea fam. indet. gen. indet. sp.1Asteroidea. fam. 683 gen. sp. 1; B) Mounded trail (unknown tracemaker); C) Thin fFlat trail (balck arrow; 684 685 unknown tracemaker) and rounded faecal casts (white arrow) produced by Scotoplanes 686 sp.-1; D) Thick-M-ridged trail produced by Echinoidea fam. indet. gen. indet. sp.5 Echinoidea. fam. gen. sp. 5; E) Wavy faecal cast produced by Peniagone sp.1 to 687 Peniagone sp. 3; F) Knotted faecal cast produced by Peniagone sp.1 to Peniagone sp.-3; 688 G) Coiled-curly (white arrow) and smooth (black arrow) faecal cast produced by 689 690 Psychropotes morphospecies sp.2; H) Smooth (black arrow) faecal cast produced by 691 various tracemakers (see Table 2); I) Rosette-shape trace (white arrow) produced by an echiuran worm and mound shape nearby (black arrow); J) Spirals faecal cast produced by 692 Enteropneusta fam. indet. gen. indet. sp.1 Enteropneusta gen. sp. 1; K) Switchbacks faecal 693 694 cast produce by Torquaratoridae. gen. sp.-1; L) Circular impression produce by Actiniaria fam. indet. gen. indet. sp.1-Actiniaria. fam. gen. sp. 1; M) Asteroid impression produced 695 696 by an Asteoroidea (Asteroidea fam. indet. gen. indet. spp 3, 4, 7, 8, 9Asteroidea. fam. gen. sp. 3, 4, 7, 8, 9); N) Mound (white arrow) with a semi-buried asteroidean nearby 697 (black arrow); O) Single burrow located in the apex of a cone-shaped mound; P) Paired 698

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- 699 burrow with an unidentified organism coming out; Q) Three paired burrows; R) Cluster
- 700 burrows; S) Lined burrows (black arrow).



701

- 702 Fig. 3 Comparison of median diversity indices (Simpson's, Shannon-Wiener and
- 703 Evenness) of lebensspuren, tracemakers, degrading fauna and fauna at each station. Each
- roal lebensspuren morphotypes was considered a different species for calculations.

Comentado [TR1]: We have to make sure that "species" is not used for two different things. Instead, I would avoid it wherever posible and use different, more specific terms.

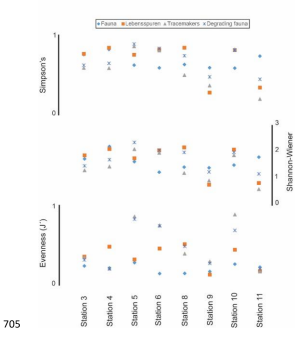
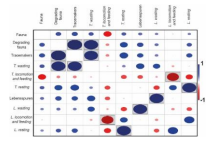


Fig. 4 Density correlation matrix for lebensspuren, tracemakers, degrading fauna and fauna. Lebensspuren and tracemakers densities were subdivide<u>d</u> into wasting, resting and locomotion-feeding (dwelling was not considered since the tracemakers of most dwelling lebensspuren are unknown). Boxed dots indicate correlations where p < 0.05.



711 Fig. 5 Hierarchical cluster diagram (constructed with Bray-Curtis similarity matrix) of

713 at each station.

the abundances of lebensspuren (A), tracemakers (B), degrading fauna (C) and fauna (D)

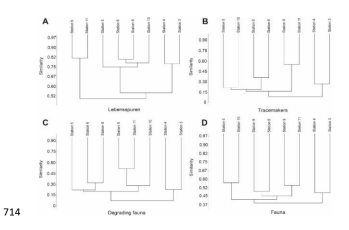
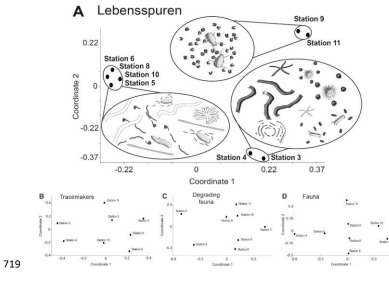


Fig. 6 Multivariate similarity represented with a non-metric multidimensional scaling (n-

MDS) plots of lebensspuren (A), tracemakers (B), degrading fauna (C) and fauna (D) at
each station. Note that the only plot that stations are together is for lebensspuren

718 abundance.



721 Table captions

Table 1. Station data of the OFOS deployments during KuramBio (2012). "Start" and
"End" coordinates refer to the time between bottom view and beginning of heaving
(survey duration). Notes: The first four deployments were aborted due to technical
problems.

Station	Start Date	Start	End	Depth (m)	Survey duration /min	Notes	
01-03	28.07.2012	44°0.03' N	44°0.01' N	5315-	7	Not enough good frames	
01.00	20.07.2012	157°18.52' E	157°18.50' E	5312		No video	
01-09	30.07.2012	- 46°14.04' N	- 46°14.04' N	-	- 4		
02-03	01.08.2012	46°14.04°N 155°33.05 E	46°14.04' N 155°33.05' E	4868- 4868	4	Not enough good frames	
03-03	04.08.2012	-	-	-	-	No video	
03-11	06.08.2012	47°14.31' N	47°13.80' N	4990-	75		
		154°42.35' E	154°43.16' E	5073			
04-10	08.08.2012	46°58.00' N	46°58.48' N	5768-	152		
		154°32.48' E	154°31.44' E	5591			
05-3	09.08.2012	43°35.03' N	43°34.64' N	5377-	125		
		153°57.95' E	153°58.60' E	5374			
06-3	13.08.2012	42°28.97' N	42°28.18' N	5298-	81		
		153°59.91' E	153°59.90' E	5308			
07-3	16.08.2012	43°2.23' N	43°1.81' N	5222-	71	Video with not	
		152°59.16' E	152°59.70' E	5221		enough definition	
08-3	19.08.2012	42°14.61' N	42°14.42' N	5125-	61		
		151°43.50' E	151°42.91' E	5125			
09-3	22.08.2012	40°34.99' N	40°34.47' N	5404-	62		
		151°0.03' E	151°0.38' E	5398			
10-3	25.08.2012	41°12.01' N	41°12.19' N	5249-	62		
		150°5.70' E	150°6.40' E	5248			
11-3	28.08.2012	40°12.93' N	40°12.92' N	5348-	61		
		148°6.04' E	148°5.41' E	5344			

726

727 Table 2. Lebensspuren and associated tracemakers identified in the present study. Note

that several lebensspuren can be produced by different tracemakers.

	Behaviour	Morphology	Description	Tracemaker taxonomy	Notes
1	Dwelling	Mounds	Large, smooth-sided cone structures. The diameter of the mounds ranged between 5 to 20 cm.	<u>Unknown</u> Unknown	Probably crustaceans
		Single burrows	Single entry holes within the flat sediment surface. Occasionally, a smooth, cone- shaped mound with a burrow entry hole at the apex. The diameters were varied, as	<u>Unknown</u> Unknown	

	Paired burrows Cluster burrows Lined	large as 2 cm, but usually between 0.5 to 1 cm. Two burrow entry holes that are closely spaced. The spacing between burrows was between 2 and 4 cm. Three or more burrow entry holes that are closely and randomly spaced. The spacing between burrows was between 2 and 10 cm. Three or more burrow entry holes that are	<u>Bivalves</u> and polychaetesBivalves and polychaetes <u>UnknownUnknown</u> UnknownUnknown	Probably crustaceans Probably crustaceans
	burrows Crater cones	aligned following a rectilinear or slightly sinuous pattern. Large central mounds surrounded by	<u>Unknown</u> Unknown	
	Crater	distinctive clusters of round, shallow impressions. Depression holes related to the collapse of horizontal burrows	<u>Actiniaria fam. indet.</u> gen. indet. sp.3Actiniaria fam.	Probably also other actiniarians
Wasting	Rounded faecal cast	Neat, short spirals of thick faecal matter	gen. sp. 3 Elpidia gen. inc. sp.1, Scotoplanes sp.1, Scotoplanes sp.2ef. Elpidia. sp. Scotoplanes sp. Scotoplanes sp. Scotoplanes sp. Scotoplanes sp.	Due to image resolution, <i>Elpidia</i> rounded faecal casts (which are commonly <1cm in size) have only been recognized on a few occasions (when it was in focus)
	Smooth faecal cast	Smooth thick faecal matter with a straight or slightly sinuous shape.	Pseudostichopus gen.inc., Psychropotes sp.1, Psychropotes sp.2, Synallactidae fam, inc. gen. indet. sp.1, Benthodytes sp.1ef. Pseudostichopus sp. Psychropotes morphospecies 1, Psychropotes morphospecies 2, Synallactidae morphospecies 1 (Amon et al. 2017), Benthodytes sp. 1	Smooth faecal cast from <i>Benthodytes</i> sp. 1 may present compressed appearance.Smooth faecal-east from <i>Benthodytes</i> sp. 1 may present compressed appearance.
	Mounded faecal cast Coiled faecal	Discrete piles of faecal matter which are not associated with burrow entry holes. Thick faecal strings appearing compressed	<u>Unknown</u> <u>Psychropotes</u> sp.1;	Due to image resolution was diffice to differentiate between war
	cast <u>Coiled</u> curly faecal cast	and curled <u>that may have with one</u> straight coil at the end. May be present along thick trail lines.	Psychropotes sp.2; Benthodytes sp.1 Psychropotes morphospecies morphospecies 1, Psychropotes morphospecies morphospecies 2, Benthodytes sp.1 1	to differentiate between pure curl control fuence. o pro
	Knotted faecal cast	Tightly loop faecal trails, often with a characteristic loop-hook at the end.	Peniagone sp.1 to Peniagone sp.3Peniagone sp.1 to Peniagone sp.3	The bigger morphotypes of this faecal cast belong to Benthodytes sp. 4 The bigger morphotypes of this faecal cast belong to Benthodytes sp. 1
	Wavy faecal cast	Tiny (less than 0.5 cm in thickness) meandering faecal remains with variable length and often in fragmented form.	<u>Peniagone</u> sp.1 to <u>Peniagone</u> sp.3Peniagone sp.1 to Peniagone sp. 3	¹ Possibly formed by uncoiling of knotted faecal cast
	Switchbacks faecal cast	Switchback or meandering feature often beginning or ending in a spiral. The acorn worm is often observed making the feature.	<u>Torquaratoridae gen.</u> indet. s <u>p.1</u> <i>Torquaratoridae</i> .	
	Spirals faecal cast	Faecal spirals with both clockwise and anti- clockwise paths. The acorn is often observed making the feature.	en. sp. 1 Enteropneusta fam. indet. gen. indet. sp. 1; Enteropneusta fam. indet. gen. indet. sp. 2Enteropneusta gen. sp. 4. Enteropneusta gen. sp. 2	
Locomotion and Feeding	Rosette-shape	Small burrow entry hole with thick, radial spokes from the central burrow. Partially completed rosettes are commonly observed. Spokes vary in thickness and	<u>Unknown</u> Unknown	This trace is usually related with echiuran worms but none has been observed in this study

	<u>M-ridged</u> <u>trails Thick M-trails</u>	length. Mounds are often found in close proximity to the rosette. Complex concave crawling structures, ranging in width from 3 to 15 cm. Both sides of the trail have small sediment ridges (forming a M- <u>ridged shape</u> trail) due to the movement of the tracemaker through the seafloor. The trails are straight and most commonly sinuous; occasionally observed with the echinoids forming the track.	Asteroidea fam. indet, gen. indet. sp.1; Asteroidea fam. indet, gen. indet. sp.2; Echinocrepis sp.1; Echinoidea fam. indet, gen. indet, sp.5Asteroidea. fam. gen. sp. 1, Asteroidea. fam. gen. sp. 4; Echinocrepis. sp. 4; Echinoidea. fam. gen.			Con formato: Inglés (Estados Unidos)
	<u>Flat trails</u> Thick flat trails	Smooth concave trails of varying length with occasional small sediment puncture marks. Thickness ranges from 2 to 10 cm. Trails may form linear, meandering, or discontinuous paths.	sp.5 <u>Benthodytes</u> sp.1, <u>Psychropotidae</u> , <u>Asteroidea fam. indet</u> , gen. indet, sp.3; Echinoidea fam. indet, gen. indet, sp.2; Echinoidea fam. indet, gen. indet, sp.7ef- Benthodytes sp. 1, <i>Psychropotidae</i> , Asteroidea, fam. gen. sp. 3; Echinoidea, fam. gen, sp. 2; Echinoidea, fam. gen, sp. 2; Echinoidea, fam. gen, sp. 7			
	<u>Variable</u> <u>t</u> Thin trails	Smooth, concave trails of varying length, up to 2 cm thick. Trails may form linear, meandering or completely random paths	fam. gen. sp. 7 Gastropoda Gastropoda fam. indet. gen. indet. sp.1-6; Echinoidea fam. indet.	Due to image resolution v to describe any ornamenta these trails. That is why the been grouped within	ation with ney have a	<u>in</u> all
			gen. indet. sp.6 <mark>Gastropoda. fam.</mark>	category.		Con formato: Alemán (Alemania)
			gen. sp. 1 to Gastropoda. fam. gen. sp. 6; Echinoidea. fam. gen. sp. 6			Con formato: Alemán (Alemania)
	Mounded trails	Smooth, with occasional ploughed features, convex trails of varying length	<u>Unknown</u> Unknown			Con formato: Alemán (Alemania)
	uans	and 3-10 cm thick. Trails may form linear, meandering or completely random paths. Craters appear sometimes in the middle of the trail.				
Resting	Asteroid impressions	Asteroid star-shaped depressions with different dimensions. Diameter ranges from 1 to 15 cm.	Asteroidea fam. indet. gen. indet. spp 3, 4, 7, 8, 9_Asteroidea. fam. gen. sp. 3, 4, 7, 8, 9			
	Ophiuroid impressions	Ophiuroid star-shaped depressions	Ophiuroidea fam. indet. gen. indet. spp 1- <u>3Ophiuroidea.</u> fam. gen. sp. 1 to Ophiuroidea. fam. gen. sp. 3			
	Circular impressions	Circular depressions with a depth of less than 4cm	Actiniaria fam, indet, gen, indet, sp.1, Actiniaria fam, indet, gen, indet, sp.3, Actiniaria fam, indet, gen, indet, Sp.7Actiniaria, fam, gen, sp. 1, Actiniaria, fam, gen, sp. 3,			

729

730 Table 3. Total number of lebensspuren, tracemakers, degrading fauna and fauna

731 identified through the 8 deep-sea stations at the Kuril Kamchatka area.

A	Tracemakers										Leber	nsspuren	Con formato: Inglés (Estados Unidos)
N=5	0 (frames	Fauna	Degrading	Total	Wasting	Locomotion	Resting	Dwelling	Total	Wasting	Locomotion	Resting	Dwelling
per	station)		fauna			and feeding					and feeding		
Sta	ation 3	560	95	91	7	1	81	Х	1207	63	84	361	699
Sta	ation 4	609	271	250	174	7	70	Х	991	257	30	195	509
Sta	ation 5	157	27	20	11	10	7	Х	974	557	18	37	361
Sta	ation 6	750	25	19	9	5	9	Х	569	257	36	32	240
Sta	ation 8	522	52	36	3	6	27	Х	321	77	32	32	178
Sta	ation 9	723	119	108	86	6	17	Х	2448	2069	25	60	292
Sta	ntion 10	181	32	13	5	8	4	Х	687	278	46	27	328
Sta	ation 11	507	169	139	130	2	5	Х	2229	1803	50	13	363