

Diversity and density relationships between lebensspuren and tracemaking organisms: a study case from abyssal northwest Pacific

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Abstract. In the deep sea, interactions between benthic fauna and seafloor sediment primarily occur through bioturbation that can be preserved as traces (i.e. lebensspuren). Lebensspuren are common features of deep-sea landscapes and are more abundant than the organisms that produce them (i.e. tracemakers), rendering lebensspuren promising proxies for inferring biodiversity. The density and diversity relationships between lebensspuren and benthic fauna remain unclear, and contradicting correlations have been proposed (i.e. negative, positive, or even null correlations). To approach these variable correlations, lebensspuren and benthic fauna were characterized taxonomically at eight deep-sea stations in the Kuril-Kamchatka Trench area, together with two novel categories: tracemakers (specific epibenthic fauna that produce these traces) and degrading fauna (benthic fauna that can erase lebensspuren). No general correlation (overall study area) was observed between diversities of lebensspuren, tracemakers, degrading fauna, and fauna. However, a diversity correlation was observed at specific stations, showing both negative and positive correlations depending on: (1) the number of unknown tracemakers (especially significant for dwelling lebensspuren); (2) the lebensspuren with multiple origins; and (3) tracemakers that can produce different lebensspuren. Lebensspuren and faunal density were not correlated. However, lebensspuren density was either positively or negatively correlated with tracemaker densities, depending on the lebensspuren morphotypes. A positive correlation was observed for resting lebensspuren (e.g. ophiuroid impressions, Actiniaria circular impressions), while negative correlations were observed for locomotion-feeding lebensspuren (e.g. echinoid trails). In conclusion, lebensspuren diversity may be a good proxy for tracemaker biodiversity when the lebensspuren-tracemaker relationship can be reliable characterized. Lebensspuren-density correlations vary depending on the specific lebensspuren residence time, tracemaker density, and associated behaviour (rate of movement). Overall, we suggest that lebensspuren density and diversity correlations should be studied with tracemakers rather than with general benthic fauna. On a global scale, abiotic (e.g. hydrodynamics, substrate consistency) and other biotic factors (e.g. microbial degradation) may also play an important role.

1 Introduction

Neoichnology studies the interactions between animals and substrates (e.g. bioturbation) in modern environments as well as the biogenic sediment resulting from these interactions, 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 image based (e.g. Bell et al., 2013; Miguez-Salas et al., 2023). Lebensspuren are portraits of the diverse linkages between environmental conditions and the animal responses to them. Thus, neoichnological analyses offer a useful tool set for deducing environmental factors both in contemporary and in past environments through comparisons 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 research), and most quantitative studies are restricted to shallow marine environments and tank experiments (e.g. shoreface, foreshore, marginal marine settings) (La Croix et al., 2022, and references therein). Even though the abyssal zone (i.e. 3500– 6500 m deep) 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 by the cost of observation and sampling procedures (e.g. Heezen and Hollister, 1971; Przeslawski et al., 2012; Bell et al., 2013; Miguez-Salas et al., 2022). As such, neoichnological analyses are emerging as a promising tool for furthering our understanding of deep-sea environments and faunal–sediment interactions.

Quantitative marine ecological research comprises two main components – diversity and density analyses (Halpern and Warner, 2002). Deep-sea neoichnological studies have addressed diversity and density characterizations by considering all identified lebensspuren as morphotypes (Przeslawski et al., 2012; Bell et al., 2013). However, tracemaker (i.e. the benthic organisms that produce lebensspuren) diversity and density have been approached from a generalist perspective as megafauna, epifauna, or lebensspuren-forming epifauna (Young et al., 1985; Dundas and Przeslawski, 2009; Przeslawski et al., 2012; Bell et al., 2013).

Early deep-sea neoichnological studies suggested a correlation between lebensspuren and faunal diversity (Kitchell et al., 1978; Young et al., 1985). However, more recent studies show no significant correlation between epifaunal and lebensspuren richness (Przeslawski et al., 2012). Bell et al. (2013) stated that "improvements in imaging technology allow [for] more refined classification of lebensspuren and organisms, which may affect the strength of the correlation between faunal and lebensspuren diversity, compared with the more direct proportionality of faunal and lebensspuren diversity demonstrated in earlier studies". Thus, in deep-sea research, diversity comparisons based on more precise taxonomic tracemaker identification and differentiation are a pending task, promising a deeper understanding of the dependencies between fauna and lebensspuren variability.

Early studies found an inverse relationship between lebensspuren and faunal density (Kitchell et al., 1978; Young 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 for a long time (high residence time), ultimately leading to a steady increase of the lebensspuren density through accumulation. Nevertheless, recent data seemed to contradict this initial assumption. Przeslawski et al. (2012) observed no discernible correlation between lebensspuren and epifaunal abundance. By contrast, Bell et al. (2013) reported a strong positive correlation between lebensspuren and faunal densities (see Fig. 10 in Bell et al., 2013). The results from Bell et al. (2013) suggest that megafaunal activity might not be the only factor influencing the preservation or destruction of lebensspuren. Small-scale biotic factors (e.g. microbial degradation), as well as abiotic factors (e.g. hydrodynamic regimes, sedimentation rates, sediment composition) potentially limit lebensspuren residence time and density across different spatial scales in the deep sea (Wheatcroft et al., 1989; Smith et al., 2005; Miguez-Salas et al., 2020). In summary, earlier investigations and their contradicting results highlight that neoichnology and its fundamental concepts are still in their early stages and warrant further investigation.

Despite the prevalence of lebensspuren on the deep seafloor (Heezen and Hollister, 1971), only a few organisms are recognized in the process of forming these features. Thus, understanding the density-diversity relationship between lebensspuren and benthic megafauna may help decipher indirectly the variability of the former (i.e. without having seen the organisms). The research presented here aims to compare diversity indices and densities of lebensspuren, tracemakers (specific organisms that produce them), degrading fauna (benthic fauna that can erase lebensspuren), and megabenthic fauna from the northwest Pacific abyssal plain, near the Kuril-Kamchatka Trench (KKT) (Fig. 1). By conducting a detailed classification of both lebensspuren and tracemakers, this research aims to go one step further, with the main objective of testing how diversity and density relationships vary from previous results where lebensspuren was compared with fauna diversity in a much coarser taxonomic resolution. This geographic region has been studied extensively, with research dating back from 11 expeditions aboard the R/V Vityaz (1949, 1953, and 1966) to more recent research efforts aboard the R/V Sonne (KuramBio I (2012) and KuramBio II (2016)). These expeditions have significantly contributed to establishing the most comprehensive taxonomic foundation for fauna in this region (e.g. Zenkevitch et al., 1955; Zenkevitch, 1963; Belyaev, 1983; Brandt and Malyutina, 2015; Brandt et al., 2020; Saeedi and Brandt, 2020).

2 Material and methods

2.1 Study sites, data acquisition, and video analysis

The joint German–Russian expedition KuramBio 1 (Kuril-Kamchatka Biodiversity Studies) aboard the RV *Sonne* (cruise SO223) explored the KKT and its adjacent abyssal plain from 21 July to 7 September 2012 (Brandt and Ma-lyutina, 2012). During the KuramBio 1 expedition, 13 Ocean Floor Observation System (OFOS) deployments were conducted (Table 1). The aim of these deployments was to use the OFOS to study 11 deep-sea stations spanning a range of depths (4868–5768 m), located between 34-48° N and 147-157° E (Fig. 1). Stations 1, 2, and 5–11 were located in the abyssal plains adjacent to the KKT, while stations 3 and 4 were located at the upper slope of the KKT (Fig. 1) (Table 1).



Figure 1. Map of the study area (Kuril-Kamchatka Trench area) and the location of the deep-sea stations analysed.

The OFOS was lowered into the water at the conductivity, temperature, and depth position. For the first 300 m, the OFOS was lowered at $0.5 \,\mathrm{m\,s^{-1}}$. Then, the speed was increased to $0.8 \,\mathrm{m \, s^{-1}}$ while the ship was kept in position. The speed of the OFOS was reduced to $0.5 \,\mathrm{m \, s^{-1}}$ once it was 500 m above the ground, and further reduced to $0.3 \,\mathrm{m\,s^{-1}}$ once it was 200 m above the ground. Once the bottom was visible, the winch lowering the OFOS was stopped and the ship resumed steaming at 0.5 kn in a direction chosen depending on the water current and wind situation. Then, the winch operator manually kept the OFOS at an appropriate distance from the seafloor to observe the seafloor benthos. Two laser pointers with 10 cm between them were used as a scale. The first four deployments were aborted due to technical problems, affecting stations 1-3 (Table 1). Thus, limited video footage was obtained. Moreover, station 7 lacked high-definition video and was therefore not considered for the current analysis. All technical aspects, including pre-deployment preparation, and post-deployment maintenance (i.e. including video download) were managed by a scientific-technical service ("WTD", Wissenschaftlich-Technischer Dienst, Jörg Leptien, Reederei).

Still images were extracted from the OFOS videos at a frequency of one frame per 5 s (Miguez-Salas and Riehl, 2023a). These still images were then sub-sampled to delete frames that were out of focus so as to minimize frame over-

lap. Out-of-focus frames were defined as frames showing the OFOS moving up and down due to ocean swell. Fifty frames per station (400 still images in total), covering a seafloor area of 878 m^2 (109 m² per station approx.), were randomly selected and uploaded to the BIIGLE 2.0 software for later annotation and measurements (Langenkämper et al., 2017). Specific frames were treated with Fiji software (Schindelin et al., 2012) to enhance the visibility (CLAHE tool) of certain lebensspuren features (Miguez-Salas et al., 2019).

2.2 Lebensspuren classification and tracemaker identification

Lebensspuren morphotypes were categorized in terms of inferred tracemaker behaviour during their formation, their morphology, and the taxonomic origin of the tracemakers. The behavioural classification was adapted from Seilacher's (1954) categories for marine lebensspuren: (1) resting (imprints of stationary animals); (2) locomotion–feeding (sediment displaced by the movement of deposit feeders and surface sediment disturbances formed as organisms are foraging); (3) wasting (e.g. faecal casts, pellets); and (4) dwelling (e.g. mounds and burrows). The morphological classification followed previous morphological names (e.g. Ewing and Davis, 1967; Young et al., 1985; Dundas and Przeslawski, 2009; Przeslawski et al., 2012; Althaus et al., 2015), where such names exist. The morphological features measured in-

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

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). Note: The first four deployments were aborted due to technical problems.

cluded in the classification were length, width, and diameter. Lebensspuren with unclear morphology and origin (e.g. degraded faecal casts, trails with diffuse outlines) were excluded from analyses. Additionally, lebensspuren and fauna smaller than 1 cm (macrofauna and smaller) were also excluded from analyses as the resolution of the still images is below high-definition ($< 1280 \times 720$ pixels). Hence, this study focuses only on megafauna (i.e. fauna > 1 cm), implied whenever fauna is mentioned in this study henceforth.

This study aimed for the species rank, which is the fundamental, gold standard taxonomic level to which ecological studies generally aspire. However, names of described species could not be attached to these taxa in the majority of our image-based observations for reasons of limitations in the image quality and the general difficulty of observing diagnostic characteristics in the in situ photographs. As such, open nomenclature has been used instead, following the recommendations for image-based identifications laid out by Horton et al. (2021). Fauna has been grouped into different categories for comparisons with the diversity and density of lebensspuren: (1) tracemakers (fauna that has been clearly recognized as makers of a trace); (2) degrading fauna (fauna that can affect lebensspuren density negatively by eroding the seafloor); and (3) benthic fauna (all fauna identified in the still images).

2.3 Statistical analysis

Diversity indices (Shannon-Wiener H' (log_e) and Simpson's D) and evenness (J') were calculated for four groups: lebensspuren, tracemaker fauna, degrading fauna, and fauna. As the data from all groups show non-parametric distribution for all stations, diversity variability among stations was tested using the Wilcoxon signed-rank test (considering all groups and all indices). The Spearman rank correlation was used to test the relationships between the diversity indices of all groups. As the number of frames was the same (i.e. same observation area), density correlation analyses were performed by separately considering the total density for each group at every station. Additionally, lebensspuren and tracemaker densities were further subdivided into three groups: wasting, resting, and locomotion-feeding. Dwelling was not considered as the tracemakers of most dwelling lebensspuren are unknown.

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

3 Results

A total of 9426 lebensspuren were identified and classified from 400 still images (see Supplement), corresponding to 23 morphotypes associated with dwelling, wasting, resting, and locomotion-feeding behaviours (Fig. 2 and Table 2) (for raw dataset report at each station, see Miguez-Salas and Riehl, 2023b). The fauna comprised a total of 4009 individual animals that were classified into 93 different taxa, of which 66 were classified as degrading fauna and 43 as tracemakers (with 790 and 676 individuals, respectively) (Table 3; Miguez-Salas and Riehl, 2023b). Linking dwelling lebensspuren with tracemakers proved to be challenging, with the exception of rare and ambiguous cases where vermiform organisms, most likely polychaetes, partially emerged from paired burrows (Fig. 2P). Tracemaker identification was possible in most of the cases for wasting lebensspuren. It is common for different tracemakers to produce the same lebensspuren morphotypes and it is also common for one tracemaker (taxon) to produce several lebensspuren morphotypes (see Table 2). However, in the case of cf. *Elpidia* – the most abundant tracemaker of station 4 (see Miguez-Salas and Riehl, 2023b) - the complete characterization of its associated rounded faecal cast (smaller than 1 cm) was impossible due to image resolution limitations. Tracemaker identification of locomotion-feeding lebensspuren was mostly possible except for mounded trails which have been produced by endobenthic organisms. However, as for wasting lebensspuren (see Supplement), also in this case different tracemakers can be responsible for similar trails (see Table 2). Tracemaker identification of resting lebensspuren was possible for most of the cases.

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

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 tracemakers and waste tracemaker densities. Also, tracemakers and waste tracemaker densities are positively correlated (Fig. 4). A positive density correlation was obtained between lebensspuren and wasting lebensspuren as well as resting lebensspuren and resting tracemakers. A negative correlation was observed for locomotion–feeding lebensspuren and their tracemakers (Fig. 4).

Lebensspuren assemblages were generally similar among stations (Fig. 5a), ranging from 75 %-82 % similarity in the cluster analysis. The n-MDS showed that lebensspuren assemblages from stations 5, 6, 8, and 10 are different from the trench (stations 3 and 4) and the southern stations (stations 9 and 11) (Fig. 6a). The southern stations were less diverse, similar (82% similarity; Fig. 5a) and dominated by rounded faecal casts produced by Scotoplanes spp. The trench stations were characterized by diverse and slightly less similar assemblages (75% of similarity) dominated by dwelling lebensspuren (e.g. paired, lined or cluster burrows), knotted faecal casts (Peniagone spp.), ophiuroid impressions (Ophiuroidea), circular impressions (Actiniaria) and M-ridged trails (Asteroidea and Echinocrepsis spp.). Stations 5, 6, 8, and 10 showed diverse lebensspuren assemblages dominated by smooth (cf. Benthodytes, Psychropotidae) and coiled-curly faecal casts (Psychropotidae), rosetteshaped traces and flat trails (Asteroidea, cf. Benthodytes, Psychropotidae) (Fig. 6a).

The hierarchical cluster diagram for tracemakers, degrading fauna, and fauna showed less similarity between stations than was the case for lebensspuren, especially for tracemakers and degrading fauna (values ranging from 20% to 55% similarity in the cluster analysis) (Fig. 5b–d). However, the trench stations (stations 3 and 4) and the southern stations (stations 9 and 11) seemed to have similar compositions. The low inter-station similarity of tracemakers, degrading fauna, and fauna assemblages was also reflected in the n-MDS plots where the spacing between stations was considerably higher than in the lebensspuren plot (Fig. 6b–d).

4 Discussion

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

Behaviour	Morphology	Description	Tracemaker taxonomy	Notes
Dwelling	Mounds	Large, smooth-sided cone structures. The diameter of the mounds ranged between 5 and 20 cm.	Unknown	Probably crustaceans
	Single hurrows	Single entry holes within the flat sediment surface. Occasion- ally, a smooth, cone-shaped mound with a burrow entry hole	Unknown	
		at the apex. The diameters were varied, as large as 2 cm, but usually between 0.5 and 1 cm.		
	Paired	Two burrow entry holes that are closely spaced. The spacing	Bivalves and polychaetes	
	burrows	between burrows was between 2 and 4 cm.		
	Cluster	Three or more burrow entry holes that are closely and ran-	Unknown	Probably crustaceans
	burrows	domly spaced. The spacing between burrows was between 2 and 10 cm.		
	Lined	Three or more burrow entry holes that are aligned following	Unknown	Probably crustaceans
	burrows Crater cones	a rectilinear or slightly sinuous pattern. Large central mounds surrounded by distinctive clusters of	Unknown	
		round, shallow impressions.		
-	Crater	Depression holes related to the collapse of horizontal burrows	Actiniaria fam. indet. gen. indet. sp. 3	Probably also other actiniarians
Runspan	faecal cast	וזיכמו, אוטור אטוומה טו חווכה ומככמו ווומונכו	<i>Exprame</i> generation sp. 1, <i>sconoptanes</i> sp. 1, <i>Scotoplanes</i> sp. 2	rounded faecal casts (which are com-
				monly < 1 cm in size) were only rec- ognized 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, Renhodvres sp 1	Smooth faecal cast from <i>Benthodytes</i> sp. 1 may present compressed appear- ance.
	Mounded	Discrete piles of faecal matter which are not associated with	Unknown	
	faecal cast Coiled-curly	burrow entry holes. Thick faecal strings annearing compressed and curled that	Psychronotes on 1. Psychronotes on 2.	Due to image resolution it was dif-
	faecal cast	may have one straight coil at the end. May be present along	Benthodytes sp. 1	ficult to differentiate between pure
		thick trail lines.		curly-segmented faecal cast and other similar morphotypes. Thus, all were included in this category.
	Knotted	Tightly looped faecal trails, often with a characteristic loop-	Peniagone sp. 1 to Peniagone sp. 3	The bigger morphotypes of this faecal
	faecal cast	hook at the end.	, , ,	cast belong to <i>Benthodytes</i> sp. 1
	Wavy faecal	Tiny (less than 0.5 cm thick) meandering faecal remains with variable lenoth and offen in fragmented form	Peniagone sp. 1 to Peniagone sp. 3	Possibly formed by uncoiling of knot- ted faecal cast
	Switchbacks	Switchback or meandering feature often beginning or ending	Torquaratoridae gen. indet. sp. 1	
	faecal cast	in a spiral. The acorn worm is often observed making the fea-		
	Spiral faecal	Faecal spirals with both clockwise and anti-clockwise paths.	Enteropneusta fam. indet. gen. indet. sp. 1;	
	cast	The acorn is often observed making the feature.	Enteropneusta fam. indet. gen. indet. sp. 2	

Behaviour	Morphology	Description	Tracemaker taxonomy	Notes
Locomotion and feeding	Rosette-shape	Small burrow entry hole with thick, radial spokes from the central burrow. Partially completed rosettes are com- monly observed. Spokes vary in thickness and length. Mounds are often found in close proximity to the rosette.	Unknown	This trace is usually related with echi- uran worms but none was observed in this study
	M-ridged trails	Complex concave crawling structures, ranging in width from 3 to 15 cm. Both sides of the trail have small sedi- ment ridges (forming a M-ridged trail) due to the move- ment 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. 5	
	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 dis- continuous paths.	<i>Benthodytes</i> sp. 1, Psychropotidae, Asteroidea fam. indet. gen. indet. sp. 3; Echinoidea fam. indet. gen. indet. sp. 2; Echinoidea fam. indet. gen. indet. sp. 7	
	Variable thin trails	Smooth, concave trails of varying length, up to 2 cm thick. Trails may form linear, meandering, or com- pletely random paths	Gastropoda Gastropoda fam. indet. gen. indet. sp. 1–6; Echinoidea fam. indet. gen. indet. sp. 6	Due to image resolution it was dif- ficult to describe any ornamentation within these trails. Therefore they have all been grouped in this size cat- egory.
	Mounded trails	Smooth, with occasional ploughed features, convex trails of varying length and 3–10 cm thick. Trails may form linear, meandering, or completely random paths. Craters sometimes appear in the middle of the trail.	Unknown	
Resting	Asteroid impressions Ophiuroid impressions	Asteroid star-shaped depressions with different dimen- sions. Diameter ranges from 1 to 15 cm. Ophiuroid star-shaped depressions	Asteroidea fam. indet. gen. indet. spp. 3, 4, 7, 8, 9 Ophiuroidea fam. indet. gen. indet. spp. 1– 3	
	Circular impressions	Circular depressions with a depth of less than 4 cm	Actiniaria fam. indet. gen. indet. sp. 1, Actiniaria fam. indet. gen. indet. sp. 3, Actiniaria fam. indet. gen. indet. sp. 7	

Table 2. Continued.



Figure 2. Examples of lebensspuren morphotypes observed and quantified in this study. (**A**) M-ridged trail produced by Asteroidea fam. indet. gen. indet. sp. 1. (**B**) Mounded trail (unknown tracemaker). (**C**) Flat trail (black arrow; unknown tracemaker) and rounded faecal casts (white arrow) produced by *Scotoplanes* sp. 1. (**D**) M-ridged trail produced by Echinoidea fam. indet. gen. indet. sp.5; (**E**) Wavy faecal cast produced by *Peniagone* sp. 1 to *Peniagone* sp. 3. (**F**) Knotted faecal cast produced by *Peniagone* sp. 1 to *Peniagone* sp. 3. (**G**) Coiled-curly (white arrow) and smooth (black arrow) faecal cast produced by *Psychropotes* sp. 2. (**H**) Smooth (black arrow) faecal cast produced by various tracemakers (see Table 2). (**I**) Rosette-shape trace (white arrow) produced by an echiuran worm and mound shape nearby (black arrow). (**J**) Spiral faecal cast produced by Actiniaria fam. indet. gen. indet. sp. 1. (**M**) Asteroid impression produced by an Asteroidea fam. indet. gen. indet. sp. 1. (**D**) Mound (white arrow) with a semi-buried asteroidean nearby (black arrow). (**O**) Single burrow located in the apex of a cone-shaped mound. (**P**) Paired burrow with an unidentified organism coming out. (**Q**) Three paired burrows. (**R**) Cluster burrows. (**S**) Lined burrows (black arrow).

Table 3. Total number of lebensspuren, tracemakers, degrading fauna, and fauna identified through the eight deep-sea stations at the Kuril Kamchatka area.

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



Figure 3. Comparison of median diversity indices (Simpson's, Shannon–Wiener, and evenness) of lebensspuren, tracemakers, degrading fauna, and fauna at each station. Each lebensspuren morphotype was considered a different species for the calculations.

4.1 Fauna, tracemakers, and lebensspuren diversity: a complex relationship

Previous comparisons between lebensspuren and faunal diversity have given rise to different contrasting hypotheses. Though pioneering research showed positive correlations (e.g. Kitchell et al., 1978; Young et al., 1985), later studies showed no correlation at all (e.g. Tilot, 1995; Turnewitsch

et al., 2000; Przesławski et al., 2012). These studies share a common approach in which the diversity comparisons were conducted from a broad perspective, especially for tracemaker organisms. Comparisons were conducted using megafaunal taxa (Young et al., 1985), epifaunal taxa (Przeslawski et al., 2012), or certain taxonomic groups of organisms (e.g. fish, holothurians, crinoids; Kitchell et al., 1978). Only Bell et al. (2013) approached the comparison between lebensspuren and fauna in greater detail by considering groups of lebensspuren-forming epifauna, and using indices to quantify lebensspuren diversity (e.g. Simpson, Shannon-Wiener). Their analyses revealed that "lebensspuren diversity was generally high and not similar to that of lebensspurenforming faunal diversity" (Bell et al., 2013). However, the links between specific tracemakers and their lebensspuren and the subsequent tracemaker diversity indexes are missing in the study by Bell et al. (2013). In the present study we attempt to close this knowledge gap by conducting a comprehensive comparison of lebensspuren diversity. We examine not only the faunal diversity but also encompass tracemaker and degrading fauna (i.e. fauna that may alter the lebensspuren assemblage by erosion/degradation). Our results show that lebensspuren diversity (Simpson, Shannon-Wiener, and evenness) is not related to fauna, tracemaker, or degrading fauna diversity. These findings seem to corroborate results of a non-existent correlation (Przeslawski et al., 2012; Bell et al., 2013). However, can this lack of correlation be expected in all deep-sea settings? This question cannot be answered without considering the limitations of quantifying deep-sea lebensspuren diversity, which is riddled with problems. These problems include image resolution, camera systems, unknown lebensspuren, observation scale, and trace degradation, but the most important is linked to their genesis. In other words, the same lebensspuren morphotype (or indistinguishable lebensspuren) can be produced by different tracemakers and one tracemaker can produce different lebensspuren (see Table 2). For example, in this study, several different holothurians (e.g. cf. Pseudostichopus, Psychropotes, Synallactidae, and Benthodytes) could have pro-



Figure 4. Density correlation matrix for lebensspuren, tracemakers, degrading fauna, and fauna. Lebensspuren and tracemaker densities were subdivided 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.



Figure 5. Hierarchical cluster diagram (constructed with Bray–Curtis similarity matrix) of the abundances of lebensspuren (a), tracemakers (b), degrading fauna (c), and fauna (d) at each station.



Figure 6. Multivariate similarity represented with 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 where stations are together is for lebensspuren abundance.

duced the smooth faecal casts. By contrast, *Psychropotes* can be linked to the production of coiled-curly and smooth faecal casts as well as flat trails (Fig. 2G). Thus, when comparing their diversity, it should be taken into account that lebensspuren morphotypes may not be related to one specific taxon, and vice versa. However, while general lebensspuren diversity in the present study did not correlate with tracemaker diversity, this does not mean that the same applies to all deep-sea settings. It is possible that a more precise characterization of the tracemaker–lebensspuren relationship, or of whether tracemakers produce specific lebensspuren morphotypes, may reveal a different correlation in other settings.

We observed different correlations between tracemakers and lebensspuren when comparing diversity among specific stations. For example, comparison of the southern stations (stations 9 and 11) using Simpson and Shannon– Wiener indexes showed a correlation between tracemakers and lebensspuren diversities (Fig. 3). This can be attributed to the traces dominating the assemblage for which we have successfully identified tracemakers (e.g. rounded faecal casts of *Scotoplanes*). On the contrary, comparison of the trench stations 3 and 4 using Simpson and Shannon–Wiener indexes revealed a negative correlation between tracemaker and lebensspuren diversities (Fig. 3). This can be attributed to the relatively large gap in our data regarding the origin of most traces of the lebensspuren assemblage. Stations 3 and 4 have a high abundance of dwelling lebensspuren (see Table 3), single burrows, mounds, 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). Also, the existence of unknown lebensspuren and tracemakers contributes to the correlation variability.

The enhancement of image resolution and the increase in deep-seafloor area covered by still image surveys may enable the improvement of lebensspuren classification and their tracemaker identification. There is ample room for improvement, especially with regard to locomotion and feeding lebensspuren. High-definition still images will allow researchers to characterize small morphological features of trails (e.g. podia marks from asteroids, echinoid spine impressions), allowing for a much more detailed classification than could be achieved in this study. The use of artificial intelligence, a seemingly promising tool in the assistance of benthic fauna recognition in imagery analyses, appears to have restricted applicability in the characterization of lebensspuren. This is because lebensspuren are constructed with sediment that has the same texture as the seafloor (i.e. background colour). In the case of dwelling lebensspuren diversity, making a comparison is significantly more complicated. Trace morphology is largely hidden below the seafloor surface, reducing the ability to differentiate between various burrow morphologies. Additionally, tracemakers are largely unidentified due to their predominantly endobenthic lifestyle (e.g. Brandt et al., 2023). Finally, it is worth noting that burrows and other dwelling lebensspuren may have multiple potential origins.

Our results show that in specific stations where the assemblage is dominated by traces with identifiable tracemakers, lebensspuren analysis appears to be a promising tool for predicting tracemaker diversity. Although these results are promising, it is evident that much more research is needed, especially with high-definition surveys (e.g. videos, images) to close existing knowledge gaps in the relationship between lebensspuren and tracemakers. Moreover, we emphasize that when using lebensspuren as a proxy for biodiversity, the diversity correlation should be made between lebensspuren and tracemakers, rather than with overall benthic fauna, as no correlation has been observed in the case of comparisons with the latter.

4.2 Tracemaker and lebensspuren density: morphospecific relationship

Similar to previous research, the density comparisons between lebensspuren, degrading fauna, and fauna revealed no correlation (Przesławski et al., 2012). However, the density comparisons between lebensspuren and tracemakers revealed a positive and negative correlation (Fig. 4). The density of locomotion-feeding lebensspuren is inversely correlated with their tracemaker density, while resting lebensspuren are positively correlated with their tracemaker density. These group-specific correlations contradict previous research 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 previous studies may be because their density comparisons considered the total fauna instead of separate functional groups (see Fig. 10 in Bell et al., 2013), not considering their specific impact on the sediment.

Trace residence time is the period during which a trace is recognizable on the seafloor before it is destroyed (Wheatcroft et al., 1989). Lebensspuren density values reflect the balance between lebensspuren formation and lebensspuren destruction or degradation either by biotic (e.g. microbial degradation, degrading fauna, epifaunal rate of movement) or abiotic factors (e.g. hydrodynamics, burial) (Wheatcroft et al., 1989). However, not all lebensspuren have the same residence time. Thus, traces not actively maintained by animals are short lived, lasting only days to weeks (e.g. faecal casts can be degraded within 1–2 weeks; Smith et al., 2005). By contrast, locomotion-feeding and resting lebensspuren have longer residence times as they are impressions on the seafloor (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. Some tracemakers inhabit them throughout their life, while others change residence multiple times and their burrows get passively filled (Gage and Tyler, 1991). Thus, a wide range of residence times may be expected. However, irrespective of scenario, the sedimentation rate is usually low in the deep sea and dwelling lebensspuren should have higher residence time than wasting lebensspuren, and similar or higher residence time than locomotion-feeding and resting lebensspuren.

The density of locomotion-feeding lebensspuren (e.g. Mridged trails) was inversely correlated with tracemaker density. This could be attributed to two reasons: (1) a high residence time of these lebensspuren even if the respective tracemakers are no longer in the study area; and (2) these lebensspuren represent a foraging behaviour in which the tracemakers tend to continuously search the seabed for food, often over a wide area (i.e. high rate of 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. circular impressions, asteroid impressions) was directly correlated with tracemaker density. This is not surprising because while these lebensspuren have a high residence time, their tracemakers (e.g. asteroids, actiniarians) have low rates of movement (Durden et al., 2015, 2019). In such cases, a high density of resting lebensspuren should always be linked to a high density of their tracemakers.

The density correlation between wasting lebensspuren and their tracemakers was slightly positive but not significant (Fig. 4). This may be attributable to the fact that we were unable to quantify the exact number of faecal casts for some morphotypes. For example, in station 4, the lebensspuren of the dominant tracemakers (*Elpidia*; more than 150 specimens) were incorrectly quantified due to image resolution limitations (small rounded faecal casts). Thus, a positive density correlation between wasting lebensspuren and their tracemakers should be expected. However, this assumption may be influenced by the behaviour of their tracemakers , as their feeding activity can be expected to depend on the grain size, availability, and quality of the nutrients among other environmental factors (e.g. Jumars and Wheatcroft, 1989; Ginger et al., 2001).

The variability observed in the lebensspuren density correlations shows a complex scenario even without considering biotic and abiotic factors that cannot be characterized through still images. For instance, 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). These "small" biotic processes are impossible to quantify through images; however, they most

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likely influenced the lebensspuren density that we quantified for this study. Moreover, while previous studies assumed that abiotic lebensspuren degradation rates are constant over the lebensspuren residence period (Bell et al., 2013), recent studies show that this may not be always true since 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 currents, substrate consistency) on the density of the assemblages studied as well as those of some biotic factors (e.g. microbial degradation which cannot be characterized in a still image) are out of the scope of this research but should be considered in future studies.

5 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. Further conclusions of this study are:

- 1. The ability of various tracemakers to produce the same lebensspuren morphotypes, and for a single tracemaker to produce various lebensspuren morphotypes, will impact the establishment of a positive or negative diversity correlation.
- 2. The existence of unknown tracemakers will contribute to the diversity correlation variability. However, lebensspuren diversity may be a good proxy for tracemaker biodiversity when the lebensspuren-tracemaker relationship can be reliably characterized.
- 3. Lebensspuren density can be positively or negatively correlated with tracemaker densities depending on the specific lebensspuren residence time and tracemaker behaviour (e.g. locomotion, resting).
- 4. Lebensspuren–density correlations may be controlled on a wider spatial scale by abiotic (e.g. hydrodynamics, grain size, organic matter, substrate consistency) and biotic factors (e.g. microbial degradation).

Above all, we suggest that lebensspuren density and diversity correlations should be studied with tracemakers rather than with general benthic fauna.

Data availability. All data generated or analysed during this study are included in this article. The raw data used for this study are available at: https://doi.org/10.5281/zenodo.10057636 (Miguez-Salas and Riehl, 2023b).

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