### RESEARCH

# The dilemma of underestimating freshwater biodiversity: morphological and molecular approaches

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#### Abstract

**Background** Anthropogenic impacts on freshwater habitats are causing a recent biodiversity decline far greater than that documented for most terrestrial ecosystems. However, knowledge and description of freshwater biodiversity is still limited, especially targeting all size classes to uncover the distribution of biodiversity between different trophic levels. We assessed the biodiversity of the Lower Rhine and associated water bodies in the river's flood plain including the river's main channel, oxbows and gravel-pit lakes, spanning from the level of protists up to the level of larger invertebrate predators and herbivores organized in size classes (nano-, micro, meio- and macrofauna). Morphological diversity was determined by morphotypes, while the molecular diversity (amplicon sequencing variants, ASVs) was assessed through eDNA samples with metabarcoding targeting the V9 region of the 18S rDNA.

**Results** Considering all four investigated size classes, the percentage of shared taxa between both approaches eDNA (ASVs with 80–100% sequence similarity to reference sequences) and morphology (morphotypes), was always below 15% ( $5.4 \pm 3.9\%$ ). Even with a more stringent filtering of ASVs (98-100% similarity), the overlap of taxa could only reach up to 43% ( $18.3 \pm 12\%$ ). We observed low taxonomic resolution of reference sequences from freshwater organisms in public databases for all size classes, especially for nano-, micro-, and meiofauna, furthermore lacking metainformation if species occur in freshwater, marine or terrestrial ecosystems.

**Conclusions** In our study, we provide a combination of morphotype detection and metabarcoding that particularly reveals the diversity in the smaller size classes and furthermore highlights the lack of genetic resources in reference databases for this diversity. Especially for protists (nano- and microfauna), a combination of molecular and morphological approaches is needed to gain the highest possible community resolution. The assessment of freshwater biodiversity needs to account for its sub-structuring in different ecological size classes and across compartments in order to reveal the ecological dimension of diversity and its distribution.

Keywords Metabarcoding, Morphotypes, Species richness, V9 region

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#### Background

As an international stream, the River Rhine is the largest federal waterway in Germany and of corresponding socio-economic importance. Here, clusters of rocks, so called ripraps, can often be found as a result of river straightening [1]. Moreover, the demand of earth-bound resources, of which gravel is particularly available in the grounds of flood plains and thus mined, has furthermore resulted in the creation of artificial lake systems along the river's course. Such anthropogenically changes in the cultural landscape of Central Europe within the past decades' and our intensive utilization of freshwater habitats caused degradation, pollution, water extraction, foodweb disturbance and invasive species introduction into freshwater habitats. This is causing a recent biodiversity decline far greater than in most terrestrial ecosystems [2, 3]. Flowing waters and the associated limnic ecosystems of floodplains are particularly impacted by the biodiversity decline and other aspects of global change [4]. Despite the imminent threat to freshwater ecosystems and their socio-ecological importance, knowledge on freshwater biodiversity is still limited. However, knowledge of the species diversity and richness is essential for generating a better understanding of freshwater ecological processes which in turn provide crucial ecosystem services fundamental for human livelihoods and wellbeing [2].

Many freshwater biodiversity studies have been conducted on one or a few specific taxonomic groups [1, 5, 6]. Certain taxonomic groups within e.g. the macrozoobenthos as well as phytoplankton, have been integrated into biodiversity monitoring programs, where they are routinely surveyed to evaluate factors such as the overall ecological status of freshwater ecosystems [7-9]. However, investigations that integrate across all size classes are scarce [10, 11], although numerous studies in different frameworks and habitats have already demonstrated the complex interactions between organisms within the different trophic levels, as well as between size classes [10, 12]. While top-predators like fish are often known to regulate the population of smaller size classes, protists are shown as the essential group for the transfer of energy by connecting lower and higher trophic levels, consequently affecting the whole food web [10, 12].

Given the diversity of species within the different size classes in freshwater systems, biodiversity assessments at habitat-scale and for the magnitude of different freshwater ecosystems is unrealistic to be fulfilled in projects of reasonable size and duration when classical morphology-based identification is considered exclusively. While enabling quantitative assessments of a community, morphological methods are time-consuming because sampling techniques vary in suitability depending on the different taxa and size classes [13-15]. Furthermore, correct taxon determination based on morphological characteristics requires a notable amount of knowledge. It is necessary to include at least one expert for each size class to sufficiently cover the community's diversity. However, biodiversity monitoring is already shifting from a traditional morphological to a genetic based approach for the registration of species. Environmental DNA (eDNA) metabarcoding approaches have been identified amongst the most promising approaches to overcome previous limitations of solely morphological approaches, by their ability to detect an almost complete proportion of the species diversity in a given environmental sample [16-20]. Not only do they have the potential to significantly decrease time, costs and required knowledge but eDNA sequencing rather detects too many than too few taxa inhabiting an area [13–15, 18–21]. However, the inability to differentiate between life stages, intact and living organisms, ingested, or extraneous tissue, to calculate abundances and biomass, or to exclude taxa through unfitted barcodes are remaining obstacles with metabarcoding [16, 22-24]. Overall, it is particularly relevant to rely on standardized procedures for biodiversity monitoring, which are as less error prone as possible, time and cost efficient and independent from exclusive expert knowledge. Thus, a combination of multiple techniques is needed, especially when targeting all size classes at once.

The aim of this study was to investigate species diversity from protists up to larger invertebrates organized in size classes (nano-, micro, meio- and macrofauna) from a long-term ecological research site located in Germany at the Lower Rhine in North Rhine-Westphalia including two riprap river Rhine sections, two oxbows as well as two gravel-pit lakes in the floodplain. The aim of the LTER-D project REES is to investigate the eco-evolutionary dynamics along a trophic cascade, integrating species representatives with regards to molecular evolution as well as size, structure and distribution patterns of populations over time and space. Within this study the biodiversity of the targeted REES habitats was initially assessed to obtain a status quo of the water bodies and to create a biodiversity inventory that allows for the choice of representative species along trophic levels for the future development of the long-term study. Diversity was assessed through morphotype richness and metabarcoding, as well as accounting for sediment composition. Furthermore, this study gives insight into how much time, expertise and what kind of mismatch can be expected by targeting biodiversity from a morphological and molecular perspective when people of different expertise and career level are contributing to the assessment.

#### **Material and Methods**

#### Sampling site

The study area lies within the LTER-D REES site [25] at the Lower Rhine in North Rhine-Westphalia (district Rees) in Germany, which includes the original flood plain area of the River Rhine (partly separated by dikes) including several gravel pit lakes, Rhine oxbows and abandoned meanders, as well as the main river. Littoral benthic samples of four standing water bodies within this area (oxbow Bienen, oxbow Grietherort, gravel-pit lake Reeser Meer Süd) and thick epilithic biofilm communities of two ripraps in the River Rhine (Grietherort and Cologne) were collected between 8th of July and 1st of September 2021 (Table 1, Fig. 1).

The two oxbow sites have different connections to the main river. While oxbow Bienen (site A) is an old oxbow only connected to the River Rhine at extreme flood events, oxbow Grietherort (site B) is regularly connected to the main river. Both areas are under nature conservancy, whereby oxbow Bienen (site A) is particularly distinguished as a bird sanctuary. Both sites' shores showed occasional trees and grasses, accumulations of water lilies (*Nymphaea*), common reed (*Phragmites*) as well as nearby stinging nettles (especially site B). Additionally, site A's vegetation contained further patches of reedmace (*Typha*), mint (*Mentha*), as well as duckweed (*Lemna*) in and around the oxbows shore. The two riprap sites were dominated by basalt boulders ranging from approximately ten to 45 cm in diameter. The riprap in Cologne (site F) and the riprap in Grietherort (site E) were located at the impact slope of the Rhine. Besides thick biofilm layers overgrowing each submerged rock, no notable macrophytes were observed below water level.

Regarding the two groundwater-fed gravel-pit lakes, Reeser Meer Norderweiterung (site C) and Reeser Meer Süd (site D), the habitats' shores contained more soft sediment than big boulder, while site D showed several rock accumulations above and below the water level. Both lakes are still being used for gravel extraction. With several kilometers distance to the River Rhine and separation by the main dike, both sites' structural features are unaffected by flooding events; however, the connection via groundwater changes in correspondence with a changing water level of the River Rhine, too.

Both sites shared aquatic patches of pondweed (*Potamogeton*) as well as stoneworts (*Chara*). Both, together with the waterweed *Elodea* were especially dominant at site D's shore region. While several patches of *Phragmites* grew near site D's shore only few individuals were observed at the site of gravel-pit lake C. A unique characteristic of site C is the presence of a dense population of the great crested newt (*Triturus cristatus*) and the presence of only one fish species, the sunbleak (*Leucaspius delineatus*), which was first observed in 2019. Before that, the Reeser Meer Norderweiterung had been fish free.

Habitat	Site name (Abbr.)	Site	Sampling date	Coordinates (Lat./Lon.)
Oxbow	Bienener Altrhein (BAR), Lower Rhine region, Germany	A	2021-07-08	51°48′21.9″N / 6°21′44.9″E
Oxbow	Grietherorter Altrhein (GAR), Lower Rhine region, Germany	В	2021-08-05	51°47′29.6″N / 6°19′51.6″E
Gravel-pit lake	Reeser Meer Norderweiterung (RMNE), Lower Rhine region, Germany	С	2021-07-19	51°45′41.9″N / 6°26′15.7″E
Gravel-pit lake	Reeser Meer Süd (RMS), Lower Rhine region, Germany	D	2021-07-21	51°45′17.8″N / 6°27′32.9″E
River Rhine	Riprap Grietherort (RRG), Lower Rhine region, Germany	E	2021-08-26	51°47′25.3″N / 6°19′44.9″E
River Rhine	Riprap Cologne (RRC), Lower Rhine region, Germany	F	2021-09-01	50°54′24.0″N / 6°58′43.4″E

 Table 1
 List of the six sampling sites including sampling dates and coordinates

(See figure on next page.)

Fig. 1 Map and pictures of the six sampling sites (A-F) as part of the LTER-site REES. This site is located at the Lower Rhine in North Rhine-Westphalia (dashed red line, except for site F). The map inlet shows the location of the sampling site in Germany (light blue) within Europe. Samples were taken between June and September 2021. Sampling sites include the oxbows (green) Bienen (A) and Grietherort (B), the gravel-pit lakes (yellow) Reeser Meer Norderweiterung (C) and Reeser Meer Süd (D) and the River Rhine main channel at (blue) Grietherort (E) and Cologne (F). Map was created with QGIS [26] by Tobias Nickel



Fig. 1 (See legend on previous page.)

#### Sampling procedure

At each sampling site, nanofauna (2–20  $\mu$ m), microfauna (20–200  $\mu$ m), meiofauna (200  $\mu$ m-2 mm), and macrofauna (<2 mm) [27] samples were taken in addition to eDNA samples for metabarcoding analysis.

Macrofauna of the lentic water bodies was sampled in triplicates using a 25×25 cm benthos net (mesh size 500  $\mu$ m) covering 5 m<sup>2</sup> (oxbow site A) or 0.53 m<sup>2</sup> (oxbow site B, gravel-pit lakes site C and D) per replicate. The net was pulled across the bottom, so that the organisms of the overlaying water and top 2 cm of sediment ended up in the net. The sampling area was marked up with lines and disturbance of the area prior to the sampling was avoided. For the other three size classes, sampling corer (diameter 4.4 cm) were used to collect 3.7 cm deep sediment cores together with 7 cm (3.7 cm for site A, oxbow Bienen due to the low water level) of overlaying water. Ten cores per triplicate (final volume 1000 ml) were filtered through a 500 µm net to remove large particles, leaves and macrofauna. The mixture of one replicate was evenly transferred into four 250 ml plastic beakers (subsamples). For metabarcoding studies, one subsample (250 ml) was filtered through cellulose nitrate filters with a pore size of 0.45 µm (Sartorius Stedim Biotech, Sartorius AG, Göttingen, Germany). Filters were immediately fixated with the salt solution DESS [28] in 50 ml tubes. The other three subsamples (250 ml each) were individually filtered through a meiofaunal sieve (pore size 44 µm), while collecting the filtered water for diversity analyses of protists belonging to the nano- and microfauna. The retained meiofauna on the sieves was suspended in filtered water and transferred into 50 ml tubes. One subsample was fixated with formaldehyde for qualitative analysis, one was fixed in DESS for molecular analyses, and the third was left unfixed for quantitative analyses in the laboratory.

In case of the riprap samples, the benthic community of the submerged rocks was brushed off and sucked in by a pond vacuum cleaner (PONDOVAC 4, OASE GmbH, Hörstel, Germany). To sample eDNA as well as size-classes smaller than macrofauna, biofilms were collected along a transect of defined length with a total surface area of  $0.017 \text{ m}^2$ . The area to be sampled was marked up with the help of calibrated ropes laying on the stones. The collected suspension (~1 L) was processed identically as the previously described sediment samples. For the macrofauna, larger amounts of biofilm were sampled to increase the possibility to collect the entire benthic community. Organisms were retained by a 500 µm net placed over the vacuum's exit pipe and the volume of run-through water was determined. Sampling continued until approximately 30 L to 40 L were collected.

#### Grain size measurement

Two sediment samples (cores) of the oxbows and gravelpit lakes were taken to investigate sediment grain size composition of the upper 3.7 cm. Organic matter like leaves or branches were discarded and the wet weight of the sediment was measured. Each sample was dried overnight in a compartment dryer at 60 °C (Memmert GmbH+Co. KG, Schwabach, Germany) and the total dry weight was measured. Different sieves (63 µm, 125 µm, 250 µm, 0.5 mm, 1 mm, 2 mm) were used to determine the relative contribution of each size fraction to the total dry weight. By subtracting the weight of these size fractions from the total dry weight, the amount of fine sediment (<63 µm) was calculated and the sediment fractions associated to the sediment types gravel (>2 mm), very coarse (2 mm - 1 mm), coarse (1 mm - 0.5 mm), medium (500  $\mu$ m – 250  $\mu$ m), fine (250  $\mu$ m – 125  $\mu$ m) and very fine sized sand (125  $\mu$ m – 63  $\mu$ m), as well as silt together with clay (<63  $\mu$ m) [29].

#### Morphological identification

Morphological identification of each site's zoobenthos community was conducted using both light microscopes (Zeiss Axio Lab.A1, Carl Zeiss, Germany) and binoculars (Leica LED2500 stand (Leica Microsystems GmbH, Germany) with stereomicroscope Zeiss Stemi 2000-C attachment (Carl Zeiss AG, Oberkochen, Germany) and attached camera systems (Sony® camera HDR-XR200VE and HDR-XR160E, Sony<sup>®</sup> Group Corporation, Tokyo, Japan). Specific identification literatures were used for nanofauna [30-32], microfauna [30, 33-39], meiofauna [40-43] and macrofauna [44-53]. The diversity was assessed based on morphotype-richness using determinations of the lowest possible taxon based on literature. If multiple morphotypes were related to one shared taxon, variants were distinguished and numbered to compensate for misidentification.

#### Metabarcoding

DESS preserved sediment samples were vortexed for two minutes and centrifuged (4000×g for 20 min at 4 °C, Megafuge 2.0 R, Heraeus Instruments). Environmental DNA was then extracted from 1 g sediment of each replicate sample (a total of 3 g per site) using the DNeasy PowerLyzer PowerSoil DNA isolation kit (Qiagen, Hilden, Germany) according to the manufacturer's protocol. Prior to the application of the kit, sediment samples were washed with three washing solutions to improve the success of DNA amplification by PCR [54]. Total DNA was quantified using Quantus<sup>TM</sup> Fluorometer (Promega, Wisconsin, USA). PCR amplifications of the hypervariable V9 region of the 18S rDNA gene was performed with 12.5 µl of the 2X VWR Red Taq DNA polymerase Master

#### Results

While investigating different size classes and taxonomic groups within this study, we looked at different levels of molecular diversity, including the molecular diversity based on ASVs with 80–100% sequence similarity to reference sequences ( $ASV_{80-100\%}$ ) and at the diversity on a predicted genus level based on ASVs with 98–100% sequence similarity to reference sequences followed by a taxonomic clustering based on genus level ( $ASV_{98-100\%}$ ). These  $ASVs_{98-100\%}$  were used for a direct comparison between the morphological diversity based on the morphological approach (morphotypes) and molecular diversity on a predicted genus level based on metabarcoding.

#### Samples and sediments

In total, 12 samples were taken from the six different sampling sites including four size classes. Three replicates were sampled per size class. All samples could successfully be investigated according to the full spectrum of analysis of this study, except for metabarcoding of the oxbow Grietherort (site B) due to PCR failure. Alongside with the biological samples, sediments of each sampling site were analyzed to better define the habitat characteristics on the micro scale via grain size categories.

Four sites (oxbows and gravel-pit lakes) were characterized by fine sediment, oxbow site A and gravel-pit lake site D by gravelly muddy sand, the gravel-pit site C by gravel sand and the oxbow site B as muddy sandy gravel (Additional file 1, Fig. S1).

#### Total number of morphotypes and ASVs

This study investigates biodiversity from three different perspectives: morphological diversity (determined by morphotypes), molecular diversity (determined by ASVs with 80–100% sequence similarity to reference sequences) and the diversity on a predicted genus level (determined by ASVs with 98–100% sequence similarity to reference sequences).

Overall, zoobenthos richness summarized across all size classes and stations showed that out of the total of 191 morphotypes (determined to a lower taxon than the supergroup level) the majority belonged to macrofauna (103 morphotypes) and the lowest number of morphotypes was observed within meiofauna (22 morphotypes). For the protist communities, 24 nanofauna and 42 microfauna morphotypes were detected (Additional file 2, Table S1).

Strict bioinformatic quality control led to a final freshwater eukaryotic dataset of 9,946,586 million reads that clustered into 2,168 amplicon sequence variants (ASVs) containing only ASVs with 80–100% similarity to reference sequences (Additional file 1, Table S2 for more details). Of these ASVs the majority belonged to meiofauna (877 ASVs), followed by macrofauna (434 ASVs). For the protist communities, 425 nanofauna and 432 microfauna ASVs were detected (Additional file 1, Table S3).

Richness obtained by metabarcoding was one order of magnitude higher when directly compared with that recovered by morphological methods (2,168 ASVs versus 191 morphotypes). Metabarcoding always recovered a higher richness than morphotype detection within each of the four size classes (nanofauna: ~ 20 fold, microfauna: ~ tenfold, meiofauna: ~ 40 fold, macrofauna: ~ fourfold). When considering only ASVs with 98–100% sequence similarity followed by a taxonomic clustering on a predicted genus level assignment, the difference between ASV richness and morphotype richness was smaller (nanofauna: ~ fourfold, microfauna: ~ twofold, meiofauna: ~ threefold, macrofauna: ~ 1.3 fold).

#### Habitat specific zoobenthos richness

The NMDS analysis based on morphotype richness showed that all three different habitats clustered separately with a slight overlap between oxbows and gravelpit lakes (Fig. 2A). The mean richness considering all size classes within each of the three habitat types was almost the same in the gravel-pit lakes and the River Rhine  $(42.3 \pm 11.5 \text{ ind. and } 40 \pm 4.05 \text{ ind., respectively})$ , while the oxbows showed the lowest mean morphotype richness with 32.5 ± 8.6 individuals (Fig. 2A Violin plot). Overall, zoobenthos richness with regard to morphotypes across all size classes was not significantly different between habitats (Kruskal–Wallis chi-squared = 3.03, df = 2, p-value = 0.22).

The zoobenthic community of the River Rhine was different from all other habitats with 64 unique morphotypes, explaining 34.4% of the community richness (Fig. 2B). Oxbows inhabited the lowest number of 69 morphotypes, whereas 93 and 86 morphotypes were found in the gravel-pit lakes and the River Rhine, respectively. Only a small proportion (10 morphotypes, 5.4%) was shared between all habitat types, whereas gravel-pit lakes and oxbows shared 30 morphotypes (16.1%) as indicated in the NMDS plot (Fig. 2A).

Molecular zoobenthic richness (ASVs<sub>80-100%</sub>) was the highest within the gravel-pit lakes including 904 unique ASVs explaining 41.7% of the community richness



**Fig. 2** Richness across habitat types. **A** Non-metric multidimensional scaling (NMDS) analysis of the Jaccard distance matrix computed from morphotypes (n = 191 morphotypes) at the six sites (n = 3 replicates), associated to three different habitat types (Oxbow, Gravel-pit lakes, River Rhine). The top inset displays the distribution of morphotype richness for the three water body communities. The black dots and bars within violin plots represent means and SDs (replicates per habitat type n = 6). **B-C** Venn diagrams showing the number of unique and shared (**B**) morphotypes and (**C**) ASVs<sub>80-100%</sub> between the three different habitat types including all four size classes (nano-, micro-, meio- and macrofauna)

(Fig. 2C). Oxbows inhabited the lowest number of 788 ASVs<sub>80-100%</sub>, whereas 1,570 and 912 ASVs<sub>80-100%</sub> were found in the gravel-pit lakes and the River Rhine, respectively. A higher proportion was shared (404 ASVs<sub>80-100%</sub>, 18.6%) between all habitat types (Fig. 2C), when compared to morphotype richness (Fig. 2B). We discovered the overall highest ASVs<sub>80-100%</sub> richness as well as the highest unique ASVs<sub>80-100%</sub> richness within each investigated size class (40.6% macrofauna, 45.6% meiofauna, 40.7% microfauna, 35.8% nanofauna) in the gravel-pit lakes (Additional file 1, Fig. S2).

## Taxonomic comparison of richness assessed via ASVs and morphotypes

Hierarchical clustering of  $ASVs_{80-100\%}$  and morphotype richness showed that each habitat formed a separate cluster within the nano-, micro- and macrofauna (Figs. 3A, C, 4A and 5A). For the meiofauna, however, the clustering

differed in branching of the gravel-pit lakes, while the cluster for the River Rhine sites was consistent (Fig. 4A). Overall, nano- and microfauna richness were less similar between sampling sites (Jaccard distances between 0.56–0.93) when compared to the meio- and macrofauna (Jaccard distances between 0.39–0.63) (Figs. 3A, C, 4A and 5A).

A direct comparison of ASVs<sub>80-100%</sub> against morphotypes based on Class level revealed that the morphotype coverage of supergroups was incomplete (Figs. 3A, C, 4A and 5A). After filtering for ASVs with a 98–100% similarity to reference sequences (Additional file 1, Fig. S3 and Fig. S4), only 20.5% of the ASVs<sub>98-100%</sub> were left for the nanofauna (87 ASVs<sub>98-100%</sub>; 432,502 reads; 43 genera, 51 species; Fig. 3B), 29.4% of the ASVs<sub>98-100%</sub> for the microfauna (127 ASVs<sub>98-100%</sub>; 523,950 reads; 79 genera, 89 species; Fig. 3C), 14.8% of the ASVs<sub>98-100%</sub> for the meiofauna (130 ASVs<sub>98-100%</sub>; 1,421,912 reads; 67 genera, 73 species; Fig. 4B) and 30.0% of the for the macrofauna community (130 ASVs<sub>98-100%</sub>; 1,841,620 reads; 78 genera, 85 species, Fig. 5B). The percentage of taxa detected with both approaches ranged from 3.8-44.3%, when only ASVs<sub>98-100%</sub> with 98–100% sequence similarity were considered. When considering all ASVs<sub>80-100%</sub> from 80–100% sequence similarity, the percentage of shared taxa was always below 10% with one exception of 14.9% (Additional file 3, Table S4).

#### Nanofauna

When considering all ASVs with a sequence similarity of 80–100% for the nanofauna community,  $ASVs_{80-100\%}$ could be assigned to 32 orders. However, only morphotypes belonging to five out of these 32 classes belonging to five supergroups and Incertae sedis could be identified, including morphotypes belonging to the Filosa Sarcomonadea (Rhizaria), Cryptophyceae (Cryptista), Kinetoplastea and Euglenida ("Excavata"), Ancyromonadida (Incertae sedis) (Fig. 3A). Except for the Ancyromonadida, we also detected a high ASV<sub>80-100%</sub> richness for these division levels, where morphotypes were found. The highest morphotype richness could be detected for Kinetoplastea and Euglenida, while especially samples from the oxbow revealed a high euglenid ASV<sub>80-100%</sub> richness (Fig. 3A). Out of the 24 nanofauna morphotypes, only eight could be determined down to genus level, including kinetoplastids (Neobodo, Rhynchomonas), cercomonads (cf. Cercomonas), euglenids (Petalomonas, Entosiphon, Peranema), cryptophyceans (Goniomonas), ancyromonadids (Ancyromonas = Nutomonas) (Additional file 2, Table S1). When considering ASVs with a sequence similarity of 98-100% with a taxonomic clustering on predicted genus level, only three genera out of the 35 most abundant  $\mathrm{ASVs}_{98\text{-}100\%}$  could also be detected morphologically, however, with great discrepancies when looking at site level matches (e.g. Rhynchomonas and cf. Cercomonas) (Fig. 3B). The most abundant nanofauna genus, the cercozoan *Rhogostoma*, based on read abundance, could not be detected morphologically in the samples (Fig. 3B). The second most abundant genus, the kinetoplastid *Neobodo*, was detected in all habitats and sites by metabarcoding, but morphotype detection revealed this genus to be present only in the River Rhine and one gravel-pit lake (site D).

#### Microfauna

When considering all ASVs with a sequence similarity of 80-100% for the microfauna community, ASVs<sub>80-100%</sub> could be assigned to 32 classes belonging to five supergroups. However, only morphotypes belonging to ten out of these 32 classes could be identified, including morphotypes belonging to the Ciliophora (Litostomatea, Nassophorea, Oligohymenophorea, Phyllopharyngea, Prostomatea 1, Spirotrichea), Dinoflagellata (Dinophyceae) and Amoebozoa (Fig. 3C). Ciliophora had the highest  $\mathrm{ASV}_{\mathrm{80\text{-}100\%}}$  richness at all sites, with the majority of morphotypes assigned to this group (37 out of 42 total morphotypes). Within the Ciliophora except for the Nassophorea and Prostomatea 1, we also detected a high ASV<sub>80-100%</sub> richness for these division levels, where morphotypes were found. The highest morphotype richness within the microfauna community could be detected for the Spirotrichea, while also the highest ASV<sub>80-100%</sub> richness was observed for this group (Fig. 3A). Only four amoebozoan morphotypes were found, although the number of  $\mathrm{ASV}_{\mathrm{80\text{-}100\%}}$  and richness was much higher. While we recovered ASV<sub>80-100%</sub> for several divisions within the rhizarians, we were not able to detect them morphologically. Out of the 42 microfauna morphotypes, 21 could be determined down to genus/species level, including species belonging to the Spirotrichea, Phyllopharyngea, and Oligohymenophorea (Additional file 2, Table S1). When considering ASVs with a sequence similarity of 98-100% with a taxonomic clustering on

(See figure on next page.)

**Fig. 3** Distribution and community composition of freshwater nanofauna (**A**-**B**) and microfauna (**C**-**D**). **A** and **C** Dendrogram cluster showing the similarity (Jaccard index) of size class communities of the five sediment samples in regard to species richness based on incidence-based data (presence/absence) using UPGMA clustering. Number of freshwater ASV<sub>80-100%</sub> (grey bars) and number of morphotypes (pink bars) related to the major taxonomic protist groups are shown. Taxonomic groups correspond to class level in the PR<sup>2</sup> database classification. **B** and **D** Heatmap of nano- and microfauna ASV<sub>598-100%</sub> with a 98–100% sequence similarity and taxonomic clustering on predicted genus level. Shown are the first 35 most abundant genera (out of 43 nanofauna genera and 79 microfauna genera) with class and genus level. In several cases reference sequences were not assigned to genus level, thus, a higher taxonomic level is shown. Numbers within the heatmap correspond to the number of ASVs assigned to this genus per site. Taxonomic groups correspond to class and genus levels in the PR<sup>2</sup> database classification. The sequential color code corresponds to the relative abundance of reads with a sequence similarity of 98–100% assigned to the respective genus to either the nano- or microfauna size class. Pink asterisks indicate, if genera could also be detected morphologically. Nanofauna protist drawings are adapted from literature [31, 68]. Microfauna protist silhouettes are from PhyloPic [69] contributed by Guillaume Dera, 2023 (CC0 1.0 Universal Public Domain Dedication) and Yan Wong, 2013 (Attribution 3.0 Unported, https://creativecommons.org/licenses/by/3.0/)



**Fig. 4** Distributional patterns and community composition of freshwater meiofauna. **A** Dendrogram cluster showing the similarity (Jaccard index) of heterotrophic protist communities of the five sediment samples in regard to species richness based on incidence-based data (presence/ absence) using UPGMA clustering. Number of freshwater ASVs<sub>80-100%</sub> (grey bars) and number of morphotypes (pink bars) related to the major taxonomic groups are shown. **B** Heatmap of meiofauna ASVs with a 98–100% sequence similarity and taxonomic clustering on predicted genus level. Shown are the first 35 most abundant predicted genera (out of 67 genera) with class and genus level. In several cases reference sequences were not assigned to genus/species level, thus, the deepest taxonomic level such as the order is shown. Numbers within the heatmap correspond to the number of ASVs<sub>98-100%</sub> assigned to these genera for each of the five sites. Taxonomic genus rank corresponds to genus levels in the PR<sup>2</sup> database classification. The sequential color code corresponds to the relative abundance of reads assigned to the genus and is relative to all reads assigned to the meiofauna size classes with a sequence similarity of 98–100%. Pink asterisks indicate, if genera could also be detected morphologically. Organism silhouettes are from PhyloPic [69]. Contributed by Mathilde Cordellier, 2020 (Arachnida), Birgit Lang, 2015 (Collembola), Maxime Dahirel, 2018 (Ostracoda), Michelle Site, 2014 (Nematoda) under License Attribution 3.0 Unported (https://creativecommons.org/licen ses/by/3.0/). Contributed by Siel Wellens, 2019 (Copepoda), T. Michael Keesey, 2013 (Branchiopoda), Scott Hartmann, 2013 (Gastrotricha) and Levi Simons, 2023 (Cestoda and Rotifera) under license CC0 1.0 Universal Public Domain Dedication. Contributed by Julie Blommaert, 2020 (Rotatoria) under license Attribution-ShareAlike 3.0 Unported. Contributed by B. Duygu Özpolat, 2016 (Oligochaeta) under license Attribution-NonCommercial-ShareAlike 3.0 Unported.

predicted genus level, only two genera out of the 35 most abundant  $ASVs_{98-100\%}$  could also be detected morphologically, namely the genera *Trithigmostoma* and *Euplotes* (Fig. 3D).

#### Meiofauna

When considering all ASVs with a sequence similarity of 80-100% for the meiofauna community, ASVs<sub>80-100\%</sub> could be assigned to 49 orders. However,

only morphotypes belonging to 12 out of these 49 orders could be identified, including morphotypes belonging to the Oligochaeta (Annelida), Diplocostraca (Branchiopoda), Cyclopoida and Harpacticoida (Copepoda), Chaetonotida (Gastrotricha), unidentified nematodes (Nematoda), unidentified Eurotatoria, Ploima and Philodinida (Rotifera), Parachela (Tardigrada) (Fig. 4A). The highest ASVs<sub>80-100%</sub> richness was found within the Chaetonotida (Gastrotricha) and Monhysterida (Nematoda). While a few morphotypes could be detected for the Chaetonotida (Gastrotricha), no morphotypes could be detected for the Monhysterida (Nematoda). Out of the 22 meiofauna morphotypes, only eight could be determined down to genus/species level, including Eurotatoria (Brachionus, Euchlanis, Philodina, Synchaeta), Diplocostraca (Alona, Eurycercus), Ostracoda (Eucypris virens) and Gastrotricha (*Chaetonotus*) (Additional file 2, Table S1). When considering ASVs with a sequence similarity of 98–100% with a taxonomic clustering on predicted genus level, only one out of the 35 most abundant ASVs<sub>98-100%</sub> could be detected morphologically, namely Chaetonotus. (Fig. 4B).

#### Macrofauna

When considering all ASVs with a sequence similarity of 80–100% for the macrofauna community,  $ASVs_{80-100\%}$  could be assigned to 61 orders. However, only morphotypes belonging to 28 out of these 61 orders could be identified, including morphotypes belonging to Annelida (Hirudinea, Rhynchobdellida), Arachnida, Branchiopoda, Malocostraca, Insecta, Gastropoda and Bivalvia (Fig. 5A). The highest  $ASVs_{80-100\%}$  richness was found within the Rhabdocoela and Catenulida (Platyhelminthes), undetermined Annelida and Diptera. Except for the Platyhelminthes Rhabdocoela and Catenulida, where no morphotype could be detected, a few morphotypes were found for the other orders with a high  $ASVs_{80-100\%}$ 

(See figure on next page.)

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richness. For Amphipoda, Odonata and Trichoptera the morphotype richness was greater at some sites compared to the ASVs<sub>80-100%</sub> richness. Out of the 103 macrofauna morphotypes, 51 could be determined down to genus/ species level, including Gammaridae (e.g. *Dikerogammarus* sp.), Isopoda (e.g. *Asellus aquaticus*), Mysidae (e.g. *Limnomysis benedeni*), Odonata (*Libellula depressa*), Hemiptera (*Notonecta* sp.), Gastropoda (e.g. *Anisus vortex, Bithynia* sp., *Lymnaea stagnalis*) (Additional file 2, Table S1). When considering ASVs with a sequence similarity of 98–100% with a taxonomic clustering on predicted genus level, only three out of the 35 most abundant ASVs<sub>98-100%</sub> could be detected morphologically, namely *Asellus aquaticus, Libellula depressa and Limnomysis benedeni* (Fig. 5B).

#### Discussion

#### Comparison of methods

The performance and outcomes of biodiversity surveys have been examined within several studies with a particular focus on the comparative effectiveness of morphology and DNA-based approaches of specific taxonomic groups highlighting discrepancies that may arise in the application of these methods [70-76]. In addition, single primer pairs or combinations of primer pairs sequencing the same or different regions have been used within metabarcoding studies targeting different taxa [77-81]. The choice of a primer pair is dependent on the research question, the targeted taxonomic groups and the sequencing technique (e.g. short vs. long reads). For protists the 18S rRNA is used as a universal marker to assess protist biodiversity targeting mainly the hypervariable V4 or V9 regions (~420 bp and ~130 bp, respectively) in metabarcoding studies [55, 78, 82, 83]. For several protist taxa the V9 region can be used to detect down to the species level [84], while for others this region was only sufficient to distinguish down to the genus level [85]. A

Fig. 5 Distributional patterns and community composition of freshwater macrofauna. A Dendrogram cluster showing the similarity (Jaccard index) of macroinvertebrate communities of the five sediment samples in regard to species richness based on incidence-based data (presence/ absence) using UPGMA clustering. Number of freshwater ASVs<sub>80-100%</sub> (grey bars) and number of morphotypes (pink bars) related to the major taxonomic macroinvertebrate groups are shown. **B** Heatmap of macrofauna ASVs with a 98–100% sequence similarity and taxonomic clustering on predicted genus level. Shown are the first 35 most abundant genera (out of 115 genera) with class and genus level. In several cases reference sequences were not assigned to genus/species level, thus, the deepest taxonomic level such as the order is shown. Numbers within the heatmap correspond to the number of ASVs<sub>98-100%</sub> assigned to these genera for each of the five sites. Taxonomic genus rank corresponds to genus levels in the PR2 database classification. The sequential color code corresponds to the relative abundance of reads assigned to the genus and is relative to all reads assigned to the macrofauna size classes. Pink asterisks behind the taxonomic names indicate, if morphotypes down to genus level could also be identified. Pink asterisks indicate, if genera could also be detected morphologically. Organism silhouettes are from PhyloPic [69]. Contributed by Nathan Jay Baker, 2022 (Branchiopoda), Kamil S. Jaron, 2022 (Insecta), Tauana Cunha, 2021 (Gastropoda) and Guillaume Dera, 2023 (Rhabditophora, Trematoda, Porifera) under license CCO 1.0 Universal Public Domain Dedication. Contributed by Denis Lafage, 2020 (Malacostraca), Katie Collins, 2020 (Bivalvia) and Mathilde Cordellier, 2020 (Arachnida) under License Attribution 3.0 Unported (https://creativecommons.org/licenses/by-nc-sa/3.0/)



Fig. 5 (See legend on previous page.)

reliable taxonomic assignment of nematode species is based on the 18S rRNA together with the highly conserved mitochondrial region cytochrome oxidase COI [86]. The hypervariable V9 region has also been used to investigate the overall metazoan community composition (e.g. [87]). While for a variety of metazoan species the COI is used as a reliable marker gene for identification, several difficulties challenging distinct species identification using COI as the marker gene were addressed by the barcoding community in the past [88–90], such as the "barcoding gap" [91]. While targeting four size classes, we used the 18S V9 region as marker region to cover a broad spectrum of organisms of all size classes.

In several cases the precision and taxonomic resolution of biodiversity surveys had been increased by eDNA surveys [73, 92, 93]. When considering four different size classes, the percentage of shared taxa between both approaches eDNA (ASVs<sub>80-100%</sub>) vs. morphology (morphotypes) was always below 15%  $(5.4 \pm 3.9\%)$  within our study. A study targeting phytoplankton, zooplankton and macroinvertebrates also showed a low number of taxa shared between both approaches with 7–9% [11]. When we considered only ASVs with a sequence similarity of 98–100%, the number of shared taxa reached up to 43%  $(18.3 \pm 12\%)$  (see Additional file 3, Table S4). Different taxa were most likely recorded under the same morphotype, likely underestimating the true diversity of each site. In addition, DNA has the ability to persist for different time spans in the environment even after organisms have disappeared. In lentic water bodies this can also lead to the downstream transport of eDNA molecules beyond the natural distribution range of the living species. Extracellular DNA can be preserved from hours to days in the water column and from decades to centuries in the sediment [94]. This DNA persistence can lead to false positive results that indicate the presence of organisms that are no longer part of the ecosystem being studied. The River Rhine spans several hundred kilometers of length and thus provides the highest potential for persisting eDNA molecules of upstream occurring species that might then be detected in the lower reaches, such as the sampling sites of our study. However, the molecular diversity of the Rhine samples did not outcompete the other water bodies in neither of the size classes. The overestimation of the diversity of the River Rhine on the level of ASVs due to a high influx of persistent eDNA molecules from upstream is thus rather unlikely.

Overall, many major taxonomic groups in all size classes could not be recovered with the morphological approach and on the other hand, many  $ASVs_{98-100\%}$  of all size classes have not been found within our morphological detection (see Figs. 3B, 4B and 5B). But there were also examples of matches of both applied methods,

genus-clustered ASVs<sub>98-100%</sub> and morphotypes, that were observed within all size classes, e.g. for the macrofauna the water louse Asellus, the mysid Limnomysis as well as the dragonfly Libellula, and for the meiofauna the gastrotrich Chaetonotus. Especially for protists (nano- and microfauna), a combination of molecular and morphological approaches is needed to gain the highest possible community resolution. Within the microfauna, for example, the phyllopharyngean cyrtophorid ciliate genus Trithigmostoma, a common freshwater ciliate, was detected within both ripraps by both approaches. High read abundances of the cercozoan species Rhogostoma minus were detected by metabarcoding within the two riprap sites (E and F), but was also detected in the other two gravel-pit lakes as well as the oxbow. This delicate thecofilosean, however, could not be identified with the morphological approach during the sampling period, but was identified in the Rhine in earlier studies (H. Arndt, unpubl.). Another example is the ciliate genus Sten*tor,* which is common in the River Rhine at Cologne [5] and was highly abundant with regards to reads from the metabarcoding approach, but could not be detected by our morphological approach at the sampling date.

One advantage of metabarcoding is that it is less reliant on taxonomic expertise [16, 71] including many taxa currently not identifiable by expert taxonomists [95–97]. However, metabarcoding can only be as precise as its database [18, 22, 24]. Not only must a database include enough correctly assigned taxa to potentially find new invasive species, the aspect that many public databases are error-infected makes the availability of well-curated databases an important prerequisite [24]. Several morphotypes identified to species level in this study have been missing in the database. Within the microfauna, one example is the detected ciliate genus Aspidisca for which sequences could be found in the used reference database (e.g. A. steini, A. aculeata, A. magna), but not for the common species identified by the morphological approach (A. cicada, A. turrita, A. lynceus).

For the macrofauna, we found *Dikerogammarus* morphotypes only to be present in the samples from the Ripraps in the River Rhine, which are seen as an emerging ecosystem with new substrate-specific combinations and can be dominated by non-native species such as *Dikerogammarus villosus/haemobaphes* [1, 98, 99]. However, metabarcoding only indicated the presence of Gammaridae in the other two habitats (oxbows and gravel-pit lakes), namely *Gammarus pulex* (ASV with 100% sequence similarity) and *Gammarus tigrinus* (ASV with 100% sequence similarity). While no V9 sequence for *Dikerogammarus* was present in the used reference database, reblasting these two *Gammarus* ASVs (read abundance of 394 and 55) at GenBank showed a 100%

similarity to many species belonging to the family of Gammaridae, but only a 94.46% similarity to *Dikerogammarus* species.

#### Habitat specific communities and biodiversity inventory

Irrespective of the size class, we observed habitat specific zoobenthos communities in each of the three investigated habitat types, with few taxa or taxonomic groups overlapping. The gravel-pit lakes showed the highest overall molecular diversity (ASV<sub>80-100%</sub>) of zoobenthos, while the overall morphological diversity was similar for the gravel-pit lakes and the River Rhine. We found the highest unique morphotype richness in the ripraps of the River Rhine with a distinct community composition when compared to the other habitats. While meiofauna morphotypes, especially nematodes, were underrepresented due to the needed taxonomic expertise to identify genera or species morphologically, we recovered a high genetic diversity of nematodes belonging to the order Monhysterida and Chromadorea as well as Chaetonotida with our metabarcoding approach (ASV<sub>80-100%</sub>) within the gravel-pit lakes. Only for the macrofauna, the morphotype richness was highest in the gravel-pit lakes, while for the other size classes (i.e. the nano- and microfauna) the highest unique morphotype richness was documented from the River Rhine.

Differences in sediment composition and macrophytes at each site might strengthen the differences in community richness between those two gravel-pit lakes. Moreover, only the sunbleak *Leucaspius delineatus* was present that at the Reeser Meer Norderweiterung (site C), while the Reeser Meer Süd (site D) is inhabited by many fish species including the European perch (Perca fluviatilis), the common roach (Rutilus rutilus) and the northern pike (*Esox lucius*). While patches of macrophytes in the shore area of lakes are generally known to provide distinct ecological niches and food sources benefitting the zoobenthos diversity [100-102], the high occurrence of macrophytes (Potamogeton, Chara, Elodea), especially at gravel-pit lake site C's (Reeser Meer Norderweiterung) shoreline could be one explanation for the higher ASV<sub>80-100%</sub> within the gravel-pit lakes when compared to the River Rhine.

With this study, we tried to obtain a status quo of the water bodies that distinguishes biodiversity across the ecological size classes. With this, we display the distribution of diversity across ecological dimensions with nano- and meiofauna explaining a major proportion of the diversity that can be best assessed via metabarcoding. This ecological partitioning of diversity, however, is in stark contrast to the availability of reference information in sequence databases and highlights the need of genetic assessments of freshwater biodiversity to expand reference libraries. The biodiversity inventory of the here presented different water bodies, furthermore builds the necessary prerequisite to identify representative species along trophic cascades to aid future studies that investigate functional aspects of the freshwater communities.

#### Perspectives for biodiversity monitoring

Biodiversity assessments based on metabarcoding might not replace the traditional morphological approach until sequence availability in databases has been extended dramatically, but is suggested to be used as a complementary tool for biodiversity monitoring [72, 73, 103, 104]. The combination of morphotype detection and metabarcoding was particularly relevant for the biodiversity assessment of bodonids present in all three investigated habitats (genera *Neobodo, Rhynchomonas, Dimastigella*). This order of kinetoplastids is known to harbor a very high degree of genetic diversity as compared to whole orders of higher eukaryotes suggesting large numbers of cryptic individual species within bodonids [105].

While traditional morphological approaches as well as metabarcoding both have their advantages and limitations, metagenomics might promise a new approach for biodiversity assessment to overcome the limitations of metabarcoding such as primer dependency and PCR amplification biases. The present and further acquisition of reference genomes will help discover the genetic mechanisms underlying organisms' responses to their natural environment and with this support conservation efforts [106]. Reference genomes from a wide range of species are required to map genomic variability and ultimately contribute to the conservation of genetic diversity. Various international initiatives aim to generate reference genomes representing global diversity [107]. With high quality reference genomes, genomic diversity of individuals from the same species can be cost-efficiently unraveled by resequencing and aligning against it [108].

#### Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s12862-024-02261-y.

Additional file 1. Additional file 2. Additional file 3.

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#### Authors' contributions

H.A., A.M.W., A.S. contributed to conception and design of the study. D.S., A.K.K., P.B., A.F., A.S., H.A. were involved in the sampling. P.B., A.Scherwaß, A.F. and H.A. conducted the morphological determination of nano- and microfauna organisms. A.K.K. processed the meiofauna samples and D.S. conducted the morphological determination of the macrofauna samples. A.S. conducted the lab analyses of the sediment samples (next-generation-sequencing). Data management was conducted by D.S., and A.S. performed the bioinformatic analyses of the data. D.S. and A.S. wrote the first draft of the manuscript. All authors contributed to manuscript revision, read, and approved the final manuscript.

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#### Availability of data and materials

The data analyzed in this study are deposited at the Sequence Read Archive SRA (SRP481383) with SRA Accessions SRR27420739-SRR27420743, BioProject ID PRJNA1061115, BioSamples SAMN39255222-SAMN39255226. Moreover, data are provided within the manuscript and supplementary information files. Preliminary results have been presented at the ECOP-ISOP conference in Vienna, Austria, in 2023 [109].

#### Declarations

Ethics approval and consent to participate

Not applicable.

#### **Consent for publication**

Not applicable.

#### Competing interests

The authors declare no competing interests.

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