Niche partitioning between planktivorous fish in the pelagic Baltic Sea assessed by DNA metabarcoding, qPCR and microscopy

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Marine communities undergo rapid changes related to human-induced ecosystem pressures. The Baltic Sea pelagic food web has experienced several regime shifts during the past century, resulting in a system where competition between the dominant planktivorous mesopredatory clupeid fish species herring (Clupea harengus) and sprat (Sprattus sprattus) and the rapidly increasing stickleback (Gasterosteus aculeatus) population is assumed to be high. Here, we investigate diet overlap between these three planktivorous fishes in the Baltic Sea, utilizing DNA metabarcoding on the 18S rRNA gene and the COI gene, targeted qPCR, and microscopy. Our results show niche differentiation between clupeids and stickleback, and highlight that rotifers play an important role in this pattern, as a resource that is not being used by the clupeids nor by other zooplankton in spring. We further show that all the diet assessment methods used in this study are consistent, but also that DNA metabarcoding describes the plankton-fish link at the highest taxonomic resolution. This study suggests that rotifers and other understudied soft-bodied prey may have an important function in the pelagic food web and that the growing population of pelagic stickleback may be supported by the open feeding niche offered by the rotifers.

Marine ecosystems around the globe experience rapid regime shifts induced by human activity, altering the community composition and ecosystem functions1. While some species are pushed to the edge of extinction, new opportunities open for those that can adapt to the new conditions and fill open niches that the regime shift brings with it. To monitor ecosystem resilience and understand why some populations decline while others thrive, a detailed understanding of the food web composition and structure is required1. Despite this, the basic trophic interactions remain poorly resolved in most marine food web studies. Planktivorous fish are often clustered in the same feeding niche, even though they and their prey (e.g., zooplankton) span a broad diversity of sizes, taxa, and ecological functions3-5. By contributing substantially to the pelagic biomass, small pelagic fish occupy a key function connecting lower and upper trophic levels3-6. Thus, variation in zooplankton availability directly affects the growth and survival of these mesopredators, and ultimately the success of fish recruitment, while the top-down mechanisms induced by small pelagic fish on zooplankton indirectly control phytoplankton7.

The Baltic Sea is strongly influenced by human stressors8, and consequently has undergone a sequence of ecosystem-wide regime shifts during the last century9. A dramatic decline of the main pelagic predator, the piscivorous Atlantic cod (Gadus morhua) around the year 1990 created the pelagic ecosystem regime that we, to a large extent, still experience today9,10, dominated by the clupeid mesopredators European sprat (Sprattus sprattus, hereafter ‘sprat’) and Atlantic herring (Clupea harengus, hereafter ‘herring’). Density-dependent competition for food resources has resulted in reduced stomach fullness and body mass of the two clupeids in the Baltic Sea proper11. However, a large partition of the pelagic zooplankton production remains unutilized by the clupeids, leaving an open niche that could potentially be occupied12. In parallel to the increase in clupeid abundance, research highlighted an exponential increase in another pelagic mesopredator, the three-spined

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zooplankton, being able to detect a broad diversity of prey items for zooplankton in the pelagic ecosystem\textsuperscript{20,22}. A fish gut content. DNA metabarcoding has proven to be a successful tool investigating diverse groups of diets of clupeids and stickleback compete for the same resources. bodied plankton, such as rotifers and ctenophores, are important prey items of planktivorous fishes and whether and quantitative PCR (qPCR) to reveal prey items that are difficult to assess microscopically. Here we ask if soft- Baltic Sea, sprat, herring, and stickleback. We utilized traditional microscopic analysis, DNA metabarcoding mesopredators in the open pelagic water. in the Baltic Sea\textsuperscript{12}. At the same time, soft and gelatinous zooplankton (i.e., rotifers and ctenophores) can at times being the main food for planktivorous fish, leading to a high diet overlap between sprat, herring, and stickleback residuial of undigested body parts, often hard exoskeletons. As a result, copepods and cladocerans are described 18S rRNA (hereafter ‘stickleback’) over the last two decades\textsuperscript{13,14} that caused a trophic cascade in coastal spawning areas, down to the algal level\textsuperscript{15,16}. Although the dramatic increase of stickleback was triggered by alterations in the dominance of predators and prey in coastal areas\textsuperscript{17}, the open pelagic habitat, where stickleback are assumed to complete a substantial part of their life cycle\textsuperscript{13,17}, also needs to offer enough resources to support the growing population. Most studies of fish stomach content are based on microscopic observations, where identification relies on residuals of undigested body parts, often hard exoskeletons. As a result, copepods and cladocerans are described being the main food for planktivorous fish, leading to a high diet overlap between sprat, herring, and stickleback in the Baltic Sea\textsuperscript{15}. At the same time, soft and gelatinous zooplankton (i.e., rotifers and ctenophores) can at times dominate the zooplankton biomass in the Baltic Sea\textsuperscript{18–20}. However, the digestion rate for soft and gelatinous prey is faster compared to crustaceans\textsuperscript{21}, and potentially contribute to an overestimation of reported empty fish gut content. DNA metabarcoding has proven to be a successful tool investigating diverse groups of diets of zooplankton, being able to detect a broad diversity of prey items for zooplankton in the pelagic ecosystem\textsuperscript{20,22}. A recent study using DNA metabarcoding conducted in coastal spawning areas was able to detect a much broader diversity of stickleback prey preference than previously described, indicating flexibility and adaptivity of the stickleback\textsuperscript{23}. However, the same method has not yet been used to study diet overlap between the three pelagic mesopredators in the open pelagic water. In this study, we assess diet overlap between the three most abundant planktivorous mesopredators in the Baltic Sea, sprat, herring, and stickleback. We utilized traditional microscopic analysis, DNA metabarcoding and quantitative PCR (qPCR) to reveal prey items that are difficult to assess microscopically. Here we ask if soft-bodied plankton, such as rotifers and ctenophores, are important prey items of planktivorous fishes and whether clupeids and stickleback compete for the same resources.

### Results

The metabarcoding of adult fish gut contents produced 3.3 and 2.9 million 18S rRNA (18S) and COI gene sequences, respectively, that passed quality control, of which 0.5 and 0.1 million of the sequences were identified as fish (belonging to family Teleostei; 18S) and host (COI), respectively, and were excluded in all downstream analyses (Table 1). Sprat larval stages produced 0.6 million 18S and 0.9 million COI reads, of which 20,264 and 0.9 million reads were kept after filtration, respectively (Table 1). The relative abundances of 18S and COI reads revealed overall consistent prey composition for all fish life stages, however, the taxonomic resolution differed among the barcodes (Fig. 1 and Supplementary Fig. S1). More specifically, for adult fish, 18S had a higher resolution of copepod prey, revealing the consumption of the genera Pseudocalanus, Acartia, Centropages and Temora, whereas COI was limited to the detection of Pseudocalanus and Acartia (Fig. 1). Additionally, 18S amplified Mertensia prey that were not amplified by COI. Contrarily, COI was better able to identify rotifers at the species level revealing the consumption of Syncphaeta baltica and this barcode also detected cladocerans (e.g., Podon) and annelids, including Bylgides and Marenzelleria, at the genus level. These taxonomic differences resulted in an overall higher alpha diversity of 18S than COI (Supplementary Fig. S2).

The sequences of adult fish gut contents were dominated by several copepod species, reaching almost three quarters of their diet (73% for 18S and 74% for COI), followed by cladocerans (11%, 18S) and rotifers (11% for 18S and 14% for COI) (Fig. 2a). Together, these prey taxa were responsible for at least 70% of the Bray–Curtis dissimilarity between all adult fish species (Fig. 2b). The copepod Pseudocalanus occupied a higher proportion of sequence reads in the gut content of herring (18S: 57%; COI: 60%) and sprat (18S: 63%; COI: 79%) compared to stickleback (18S: 1.5%; COI: 8%; $P < 0.001$, Supplementary Table S1). Based on the 18S, the copepods Temora, Centropages, and Acartia were abundant in the sequence reads of all fish species. However, the relative

| Gene region | Life stage | Fish species | Before filtration | After filtration | After rarefaction |
|-------------|------------|--------------|------------------|------------------|------------------|
| 18S         | Adult      | Herring      | 10 209 1.1       | 10 196 0.9       | 155              |
|             |            | Sprat        | 14 301 1.8       | 14 255 1.6       | 183              |
|             |            | Stickleback  | 7 120 0.4        | 7 116 0.3        | 99               |
|             |            | Total        | 31 451 3.3       | 31 401 2.8       | 302              |
|             | Larvae     | Sprat        | 8 102 0.6        | 8 39 0.02        |                  |
| COI         | Adult      | Herring      | 10 408 1.1       | 10 404 1.1       | 378              |
|             |            | Sprat        | 13 440 1.1       | 13 439 1.1       | 404              |
|             |            | Stickleback  | 8 96 0.6         | 8 93 0.6         | 89               |
|             |            | Total        | 31 645 2.9       | 31 638 2.8       | 589              |
|             | Larvae     | Sprat        | 9 437 0.9        | 7 40 0.06        |                  |

Table 1. Summary of the sample size (n), sequence reads (million) and unique amplicons variants (ASVs) for the 18S and COI barcodes associated to all adult fish and larvae species and individual adult and larvae species at each step of the downstream data processing. The first filtration was performed to remove host sequences and the rarefaction was performed for adult life stages based on the sample with lowest read count (see “Methods”).

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abundance of Temora reads were twice as high in stickleback (27%) compared to herring (13%, \( P = 0.038 \)) and sprat (10%, \( P = 0.0064 \), Supplementary Table S1). The ctenophore Mertensia was detected in higher proportions in the sequence reads of herring (15%) and sprat (13%) compared to stickleback (3%), but the beta regression revealed no significant differences between the fish species (\( P > 0.05 \), Supplementary Table S1). The 18S showed that rotifers constituted about a third of stickleback’s gut content sequence reads (34%), while being almost absent in the gut content of herring (0.2%) and sprat (0.4%) (\( P < 0.001 \), Supplementary Table S1). More specifically, the COI revealed that the rotifer S. baltica was more associated with stickleback (39%) than with sprat and herring (ca. 1%, \( P < 0.001 \), Supplementary Table S1). Based on the Bray–Curtis distance of both gene markers reads, the diet of the two clupeids sprat and herring overlapped at more than 60% (Fig. 2c), which was confirmed by pairwise permANOVAs (Supplementary Table S2). Stickleback occupied a different niche than sprat (BCI \text{18S} = 19.9%, BCI \text{COI} = 18.08%, \( P < 0.001 \)) and herring (BCI \text{18S} = 22.74%, BCI \text{COI} = 25.66%, \( P < 0.001 \)).

Results from qPCR verified high abundances of rotifers in the gut content of stickleback (on average, a signal equivalent to \( > 2.29 \times 10^6 \) rotifers 18S gene copies per stickleback gut) (Fig. 3). This concentration was more than 40 times as high in stickleback as in herring or sprat (ANOVA of the log-transformed concentrations, \( F_{2,26} = 8.45, P = 0.0015 \)). The qPCR found no detectable concentration of rotifer 18S associated to the copepods Acartia, Temora, Centropages, Eurytemora, and Pseudocalanus, nor to the cladocerans Evadne and Bosmina (data not shown).

The microscopic analysis of gut content revealed that copepods were more abundant in sprat (on average 192 copepods per gut, representing 54% of their prey) and herring (565 copepods, 62%) compared to stickleback (105 copepods, 3%) (Fig. 4a). We also found cladocerans to be abundant in some of the clupeid’s guts (on average 120 cladocerans per gut in sprat, 34% of their prey and 347 in herring guts, 38% of their prey) but less abundant in the stickleback guts (13 cladocerans per gut, 0.4%). Stickleback guts were mainly filled with rotifer eggs and contained on average 3400 eggs per gut, while no rotifer eggs were detected in sprat nor herring (Fig. 4b,c).
Figure 2. (a) Prevalent prey (lower) of the planktivorous fish herring (*Clupea harengus*), sprat (*Sprattus sprattus*) and stickleback (*Gasterosteus aculeatus*) (upper) based on their gut-content analysis using DNA sequences in the southern Baltic Sea. The thickness of the links is proportional to the relative 18S rRNA (left) and COI (right) read abundance. (b) Nonparametric multidimensional scaling (NMDS) plot of fish prey composition based on Bray–Curtis distance of 18S rRNA (left) and COI (right) gut content sequences. The prey species are represented with the coloured squares and the prey contributing to the most difference between fish species is shown with their names. Ellipses following the t-distribution of the NMDS scores are shown for each fish species. (c) Bray–Curtis similarity Index (BCI) in percent with the 95% confidence interval in the bracket between and across (diagonal) each fish species. A low BCI indicates low niche overlap. The output of the pairwise permANOVA is shown with the asterisks (*** = P < 0.001).
Figure 3. Quantitative PCR assay of rotifers in the gut content of the planktivorous fish sprat, herring and stickleback in the southern Baltic Sea targeted by rotifer-specific 18S rRNA gene primers. The results are shown as equivalents to rotifers 18S gene copies.

Figure 4. (a) Microscopic count of gut content of the planktivorous fish. Rotifers were identified as rotifer eggs. (b) Eggs were observed in the gut content of stickleback, originating from the rotifer genus Synchaeta. (c) Female of the rotifer genus Synchaeta carrying three eggs as identified from water samples. Scales are the same for (b) and (c).
Discussion

High diet overlap is assumed to cause competition between the three dominant pelagic planktivorous mesopredators in the Baltic Sea, sprat, herring, and stickleback\(^1\)\(^2\)\(^3\)\(^4\). Despite this assumption, stickleback populations have increased dramatically over the past decades, which raises the question of whether and how resources are partitioned\(^5\). While previous studies of fish diet overlap have mainly relied on microscopic identification of gut content, we implemented a DNA metabarcoding approach targeting two different gene regions, the 18S rRNA gene (18S) and the mitochondrial cytochrome c oxidase 1 gene (COI) to reveal the taxonomic diversity of prey, and a qPCR step to quantify rotifers that are at times abundant in the Baltic Sea. Our study highlights consistency between methods, with DNA metabarcoding resolving the plankton–fish link at the highest taxonomic resolution. Our results suggest a unique niche of stickleback that may enable high population growth in the open water, despite high competition between mesopredators, although this finding needs to be confirmed at larger scale. More than half of the DNA found in herring and sprat stomach contents was assigned to *Pseudocalanus*, supporting previous observations of high diet overlap between the two clupeids\(^4\)\(^6\). On the other hand, the diet of stickleback differed substantially from the two clupeids, with rotifers appearing as main prey DNA in spring. The high rotifer biomass in the environment and lack of competition from other predators indicate that this novel niche utilization may support the drastic increase of pelagic stickleback in the Baltic Sea.

We find that copepods dominated the gut content of the two clupeids sprat and herring. *Pseudocalanus* and *Temora* occupied most of the sequence reads of the clupeid metabarcoding, two species that are often reported as preferred prey in previous studies\(^1\)\(^2\)\(^3\). Despite high contributions of these two copepods, *Pseudocalanus* was more than four times as abundant as *Temora* in clupeid gut contents. A strong preference for this copepod with marine origin can further confirm the increased competition between the clupeids, as *Pseudocalanus* has decreased due to decreased salinity\(^7\) and shares a similar vertical distribution as clupeid during daytime\(^8\). Our study using metabarcoding further reveals a large relative quantity (11%) of the ctenophore *Mertinia* in the gut samples of both clupeids. Similar, Clarke et al.\(^9\) reported an important contribution of gelatinous zooplankton to upper trophic levels in the Southern Ocean. Despite high abundances of ctenophores in the Baltic Sea and their assumed importance in marine foodwebs\(^10\), they are not reported as food for planktivorous fish. A possible explanation is the difficulty observing them microscopically, as their digestion rate is faster than crustaceans\(^11\), and no hard parts remain in the digestive system. Further, COI detected the presence of cladocerans, which was confirmed by the microscopic survey, but underrepresented with 18S that strongly amplify copepods\(^2\)\(^9\). Interestingly, more than twice annelid COI reads, including the benthic macroinvertebrates *Bylgides* and *Marenzelleria*, were associated to stickleback (15%) and herring (8%) than to sprat (4%), highlighting their ability to migrate vertically. These interactions suggest that together stickleback and herring contribute to benthic-pelagic coupling when oxygen is not restricting vertical migration in the southern Baltic Sea\(^10\).

Spat and herring share a similar feeding niche, which may explain previously observed declines in body mass and stomach fullness, and supports the theory of competition between the two species\(^1\). In contrast, stickleback revealed little diet overlap with the other mesopredators. The low relative abundances of *Pseudocalanus* (1–8%) in metabarcoding analyses indicates that the density-dependent competition may not limit the population growth of stickleback. The copepods that were shared in the diet of stickleback, sprat, and herring were *Temora*, *Acartia*, and *Centropages* have increased over the last decades, as opposed to *Pseudocalanus*\(^1\)\(^2\). Our results show that stickleback are able to feed on a broader spectrum of prey and highlight that stickleback utilizes the rotifer *Synchaeta baltica* as prey, which is an important component of the plankton community composition in the Baltic Sea\(^1\)\(^2\)\(^3\)\(^4\)\(^5\). Due to the difference of prey size, we can expect an overrepresentation of copepod to rotifer sequences compared with microscopic count data. High predation rate on *S. baltica* is supported by both the qPCR assay as well as microscopic counts, although only the eggshells were visible but not the soft-bodied rotifer. Despite the considerably lower carbon content per *S. baltica* (ca. 6 µg C ind\(^{-1}\)) compared to copepods (ca. 20 µg C ind\(^{-1}\))\(^13\), the high number of rotifers likely act as a major food source for stickleback. These results propose that stickleback, due to their opportunistic feeding behaviour\(^4\)\(^2\)\(^2\)\(^2\)\(^2\)\(^2\)\(^2\), have a distinct feeding niche from sprat and herring in the open water, as they feed on a smaller size class of zooplankton compared to the clupeids. Thus, we cannot assume the same process of competition between clupeids and stickleback.

Rotifers can at times be very abundant in the Baltic Sea, reaching densities up to 25,000 ind m\(^{-3}\), but their natural predators are poorly studied. An increasing trend in biomass of the main two rotifer genera (*Synchaeta* and *Keratella*) was observed since the 1990s\(^15\). In a recent study, we showed that rotifers might occupy a unique feeding niche, as direct grazers of dinoflagellate spring bloom, as well as in the recycling of organic matter in summer\(^16\). The low level of predation on rotifers by clupeid adults (< 1% of the reads) observed here indicate that this trophic niche may not be fully utilized. Further, qPCR did not identify predation on rotifers by other zooplankton species, including several species of copepods and cladocerans, which is supported by previous observations showing that limited DNA reads were associated to dominant zooplankton species in spring\(^1)\(^2\)\(^3\). Thus, stickleback appears to have little or no competition for rotifers as a food resource in the southern Baltic Sea in spring. This abundant and increasing resource\(^17\) may therefore sustain the expanding Baltic stickleback population during its pelagic phase.

Similar feeding patterns but different taxonomic resolutions were found between approaches for adult fish. DNA metabarcoding has the highest taxonomy resolution, targeting the full prey spectrum, which allow for an exploration of the whole prey community for fish and other organisms\(^1)\(^2\)\(^3\)\(^4\). As previously reported by Clarke et al.\(^9\) both barcodes showed consistent outcomes despite their different taxonomy resolutions. 18S identified the highest prey diversity by having an increased resolution for copepods genera revealing the consumption of *Pseudocalanus*, *Acartia*, *Centropages*, and *Temora*, while COI was limited to the identification of the genera *Pseudocalanus* and *Acartia*. Moreover, 18S identified gelatinous zooplankton prey, while COI was more efficient in identifying cladocerans, annelids and rotifers at lower taxonomy levels. qPCR, that is more sensitive than DNA
metabarcoding for identifying a single species\textsuperscript{40}, allowed us to confirm and quantify the interaction between the rotifer \textit{S. baltica} and stickleback. However, this approach does not allow for a diet assessment, but rather for a post-hoc quantification, due to the specific primers used. Microscopy is the traditional method to assess fish diet, but is limited to prey with hard remaining parts, such as exoskeleton\textsuperscript{23,41}. Our visual observations are consistent with the fish prey species identified at the same sampling location in 2020 (J. Hentati-Sundberg, pers. comm.) and both show the above-mentioned limitation. Our study confirms that combining molecular tools, including DNA metabarcoding and qPCR, with traditional microscopy observations is a robust approach to explore the full prey spectrum of planktivorous fish, and reveals a diverse prey spectrum for mesopredators in the Baltic Sea.

In this study, we highlight the importance of soft-bodied zooplankton organisms as potential prey for planktivorous fish, suggesting that these neglected prey taxa might be important players in the ecosystem. While several prey species are difficult to observe using traditional gut content analysis, molecular techniques such as metabarcoding and qPCR can be used as a suitable complement. In this study, we show that rotifers, which are often not identified with microscopic analysis of gut samples, may contribute to niche partitioning between sprat, herring, and stickleback. Although this result must be confirmed by more intensive sampling, the opportunistic feeding in the pelagic Baltic Sea may reduce competition with clupeids and support the increasing stickleback population.

**Methods**

**Sampling, DNA extraction and microscopic analysis.** Adult and larval fish were sampled during cruise ALS21 with Research Vessel ALKOR at five locations in three basins of the Southern Baltic Sea proper in April 2019 (Fig. 5a, Supplementary Table S3). Based on the database for Swedish national monitoring data\textsuperscript{42}, potential fish prey (i.e., zooplankton) abundance in the upper 60 m (43 m in the Arkona Basin) of the water column in April 2019 was dominated by copepods, including \textit{Pseudocalanus}, \textit{Acartia}, \textit{Temora} and \textit{Centropages} contributing to more than 69% of the mesozooplankton community across all basins, and the rotifer \textit{Synchaeta} that was more than twice as abundant in the Gotland Basin (27%) than in the Arkona Sea (11%, Fig. 5b). Adults of the three dominant pelagic planktivorous fish species, the three-spined stickleback (\textit{Gasterosteus aculeatus}), Atlantic herring (\textit{Clupea harengus}), and European sprat (\textit{Sprattus sprattus}), were sampled using a pelagic fishing trawl net ("Jungfischtrawl", 0.5 cm mesh size in the cod) deployed at specific depths during daytime between 47 and 73 m depth (Supplementary Table S3). Full stomachs of individual close to the average population size were immediately dissected, preserved in 80% ethanol and frozen at – 20 °C until further analysis. Planktonic larvae of sprat were sampled using a Bongo Net (300 and 500 µm, Hydrobios, Kiel, Germany) with single-oblique tows to 6 m above ground and back to the surface, identified and sorted under a stereomicroscope, preserved in 80% ethanol and frozen at − 20 °C until further analysis. Fish larvae were soaked in a 1% bleach solution to remove contamination of external DNA, rinsed multiple times in DNA-free water (Qiagen), and soaked in 180 µl ALT lysis buffer. The content of the fish guts was removed and homogenized, half of the content was mixed with 360 µl ALT lysis buffer (Qiagen, Hilden, Germany), and the other half was preserved in 80% ethanol for visual observation. DNA from all samples was extracted using QIAamp DNA Micro Kit (Qiagen) according to manufacturer’s instructions for tissue samples with an additional step of bead beating with 1 mm glass beads and an overnight incubation at 56 °C with proteinase K (Qiagen). We also included archived DNA samples from rotifer, copepod, and cladoceran sampled in the northern Baltic Sea proper in 2017 and 2018 (for details see Zamora-Terol et al.\textsuperscript{22}) to quantify rotifer to zooplankton prey contribution. The gut content of a subsample,
including six herrings, 13 sprats and six sticklebacks (Supplementary Table S3), was identified under a stereomicroscope to validate the findings of the DNA metabarcoding.

**Metabarcoding analysis.** A 400 bp long fragment of the V4 region of the 18S rRNA gene (18S) was amplified in a polymerase chain reaction (PCR) with universal primers 528F and 706R\(^\text{41}\) and a 313 bp long fragment of the mitochondrial cytochrome c oxidase I gene (COI) was PCR amplified with versatile primers mCOIintF and dgHCO2198F.\(^{42}\) The thermocycler conditions are described in Zamora-Terol et al.\(^{22}\) for the 18S. For the COI, the thermocycler was programmed as follows: denaturation (98 °C) for 1 min, followed by 30 cycles of denaturation (98 °C) for 20 s, annealing (46 °C) for 20 s, and elongation (72 °C) for 15 s. The program finished with a final elongation (72 °C) for 2 min and a cooling (4 °C). Illumina sequencing library preparation was performed according to best practices described by Hu et al.\(^{45}\) Sequence clustering was done “onboard” and sequenced on MiSeq (MSC 2.5.0.5/RTA 1.18.54) pair-end setup (2 × 300 bp, version 3, Illumina, San Diego, California) with the addition of 10% genomic PhiX. FastQ files were processed in the cutadapt software\(^{46}\) to remove primers and adapters. The next steps of quality control, including trimming, filtering, and taxonomy assignment were performed in R using the DADA2 pipeline\(^{47}\). 18S sequences were assigned to the Prokaryotic Ribosomal Reference database\(^{48}\). COI sequences were assigned to a custom-made database combining the MARES database\(^{49}\) with a 636 bp length sequence of *Synchaeta baltica* retrieved in GenBank (accession: MK905848\(^{50}\)). Library preparation, sequencing, and subsequent bioinformatic analyses were performed as described elsewhere\(^{51}\). For each step in the library preparation a negative control was included, and analysed with Qubit and gel electrophoresis after the full library preparation. No detectable bands or DNA concentrations were observed in the negative controls.

**Quantitative PCR assessment of rotifers.** To accurately assess rotifer abundance in fish and zooplankton guts, we designed a quantitative PCR (qPCR) assay by constructing a degenerate rotifer-specific primer pair to target a 94 bp long fragment of the V4 region of 18S: Rotifer18SF (AYCGGTGCTGCYTTDTATG) and Rotifer18SR (CAGGCGTAWRGCGCTGTTTTA). The primers were analysed in the Test Prime software of (arb-SILVA) and match with 59.4% of the species entries of the rotifer order Monogononta, including *Keratella* spp., the two most dominant rotifer genera in the Baltic Sea. The primers did not match any other sequence in the SILVA database\(^{52}\). The primers’ self-dimerization and heterodimerization formation was tested by “Multiple Primer Analyzer” software (Thermo Fisher) with no positive detection. The qPCR assay was performed on all fish gut samples (Supplementary Table S3) and DNA samples from copepods and cladocerans sampled previously\(^{22}\).

Each PCR reaction contained 10 µl SYBR Green I master mix (Roche, Basel, Switzerland), 1 µl Rotifer18SF and 1 µl Rotifer18SR (10 nM, Eurofins Genomics, Ebersberg, Germany), 6 µl PCR grade water (Roche) and 2 µl DNA template. qPCR was performed on LightCycler® 480 II (Roche) with an initial denaturation and polymerase activation at 95 °C for 5 min followed by 40 cycles of 95 °C for 30 s denaturation, 64 °C for 30 s annealing and 74 °C for 30 s elongation and fluorescent acquisition. All analyses were terminated by a melting curve analysis of 95 °C denaturation for 1 min followed by a gradual increase of 2.2 °C s\(^{-1}\) from 65 to 97 °C under constant acquisition. For each qPCR plate we included a standard curve to correct for primer efficiency bias, and a negative control to detect contamination. Standard curves were made from PCR products extracted and amplified DNA of *S. baltica*. Residuals of the PCR reaction was removed with GeneJET PCR purification kit (Thermo Fisher, Waltham, Massachusetts), quantified with Qubit dsDNA HS assay and fluorometer (Thermo Fisher), and subsequently diluted with a factor of 10X, starting at an absolute concentration of 10\(^8\) gene copies reaction\(^{-1}\) (2 µl). The standard curve was used for absolute sample copy number estimation in the Light Cycler 454 software (Roche), with Cq values defined as the highest value of the second derivative of the RT amplification curve. While amplification bias was measured and corrected for in each run, the average runs had an efficiency of 1.82 (91%) with an R\(^2\) of 0.993.

**Data analysis.** Data filtering and statistical analysis were facilitated by the Phylloseq R package\(^{52}\). 18S sequences belonging to the Teleostei family and COI sequences belonging to the sampled fish genus were removed. Subsampling using the function “rarefy_even_depth” from the Phylloseq R package\(^{53}\), based on the sample with the lowest reads count (8437 (18S) and 20,334 (COI)) were performed to homogenize the prey diversity among adult fish samples. This step was not performed for larval stages as the number of reads were too low in some samples. Different cut-offs between larvae and adult fish were used to keep dominant prey reads. For fish larvae, unidentified sequences and taxa occupying less than 5% of the reads as well as samples with less than 50 reads after filtration were removed. For adult fish, the most prevalent prey, determined as taxa occupying at least 0.1% of the sequences in at least 70% of the sequences in each sample group (i.e., fish species) were kept for further analyses.

For adult fish, we used the Bray–Curtis Index ((1 – Bray–Curtis distance) × 100) to measure the percentage of dietary overlap. Differences in diet composition were tested by perMANOVA using the “adonis” function from the Vegan R package\(^{55}\), and pairwise comparisons between fish species were performed using the “pairwise.adonis2” function from the pairwiseAdonis R package\(^{54}\). Prey taxa contributing to at least 70% of the difference between all fish species were assessed using the “simper” function from the Vegan R package\(^{55}\), and their relative proportions were modelled with beta regression, using the “betareg” function from the betareg R package\(^{56}\). Nonmetric multidimensional scaling plots were based on Bray–Curtis distances and scores were calculated with the “metaMDS” function in the Vegan R package\(^{53}\). Figures were made in the ggplot2 R package\(^{56}\). The most important prevalent taxa were visualized in bipartite networks made using the Circos R package\(^{57}\).

**Ethics statement.** The study protocol was approved on the 22nd of January 2019 by the Department of Ecology Environment and Plant Sciences (DEEP), Stockholm University, in collaboration with
the GEOMAR Helmholtz Centre for Ocean Research Kiel, Kiel, in compliance with the German animal protection laws “TierSchG §4” and “TierSchV §1 (2)”. The German animal protection laws do not require a permit for the mass capture and killing of fish from trawl net hauls and fish larvae from Bongo net tows. We confirm that the study was undertaken with all procedures that minimize the pain and suffering, and improve animal welfare.

Data availability
Raw DNA sequences generated during the current study were uploaded with associated metadata to the European Nucleotide Archive (ENA) under the project accession number PRJEB51972 (https://www.ebi.ac.uk/ena/browser/view/PRJEB51972). R scripts and data analyzed for this study are available in the Dryad repository, https://doi.org/10.5061/dryad.vq83bk3vk. For data request, please contact the corresponding author Kinlan M.G. Jan.

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A.N.: Conceptualization, Data curation, Formal Analysis, Investigation, Methodology, Project administration, Software, Validation, Visualization, Writing—original draft. K.J.: Conceptualization, Data curation, Formal Analysis, Software, Validation, Visualization, Writing—original draft. J.D.: Methodology, Writing—review and editing. M.W.: Conceptualization, Funding acquisition, Project administration, Resources, Supervision, Writing—review and editing.

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Competing interests
The authors declare no competing interests.

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