First Evidence of Retrospective Findings of Microplastics in Harbour Porpoises (*Phocoena phocoena*) From German Waters

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Microplastic ingestion by lower trophic level organisms is well known, whereas information on microplastic ingestion, egestion and accumulation by top predators such as cetaceans is still lacking. This study investigates microplastics in intestinal samples from harbour porpoises (*Phocoena phocoena*) found along the coastline of Schleswig-Holstein (Germany) between 2014 and 2018. Out of 30 individuals found along the North Sea (NS) and the Baltic Sea (BS) coast, 28 specimens contained microplastic. This study found a relationship between the nutritional status of cetaceans and the amount of found microplastics. Harbour porpoises with a good or moderate nutritional status contained a higher number of microplastics, when compared with specimens in a poor nutritional status. In addition, when individuals died accidently due to suspected bycatch in gillnets, where a feeding event is highly assumed or a pharyngeal entrapment happened, the microplastic burden was higher. In total, 401 microplastics (≥100 µm), including 202 fibres and 199 fragments were found. Intestines of the specimens of the BS contained more microplastics than the ones from the NS. Differences in the share of fibres could be revealed: for BS fibres constituted 51.44% and for NS, fibres constituted 47.97%. The polymers polyester, polyethylene, polypropylene, polyamide, acrylic (with nitrile component) and an acrylic/alkyd paint chip (with styrene and kaolin components) were identified. This is the first study investigating the occurrence of microplastics in harbour porpoises from German waters and will, thus, provide valuable information on the actual burden of microplastics in cetaceans from the North and Baltic Seas.

Keywords: microplastic burden, FTIR, marine mammals, cetacean, North Sea, Baltic Sea, nutritional status, health

INTRODUCTION

The ubiquitous presence of marine litter, and especially the occurrence of small particles called microplastics (<5 mm) (Arthur et al., 2009) is already confirmed in different marine habitats and organisms (Fossi et al., 2014; Lusher et al., 2015b; Pereira et al., 2020). A trophic transfer of microplastic particles between species of different trophic levels can be assumed as this has previously been determined by other studies (Farrell and Nelson, 2013; Setälä et al., 2014;
Porpoises from the NS and BS are collected in the course of a health monitoring (Siebert et al., 2001, 2020; Lehnert et al., 2005). This monitoring is established since 1990 at the Institute for Terrestrial and Aquatic Wildlife Research (ITAW), which regularly conducts necropsies of harbour porpoises using a standardised protocol (Siebert et al., 2001, 2020). Since 2014, intestinal samples of marine mammals, including harbour porpoises, were exclusively collected for microplastic analysis. Based on the necropsies, the age, sex, health status and the location in which each individual was found is assessed and recorded. Thus, this information is available for the investigated intestinal samples from harbour porpoises found between 2014 and 2018.

The following criteria were applied for choosing the most suitable samples: (i) the gastrointestinal tract (GIT) had to be intact, (ii) faeces were present, and (iii) the individual was already weaned. 30 individuals were chosen for analysis: 14 individuals from the NS and 16 individuals from the BS (Figure 1).

The intestinal samples were stored in pre-cleaned glass jars at −20°C until further processing. Then, each defrosted and opened intestinal sample was placed into a double-layered washing sachet made of nylon cloths. The inner bag of the washing sachet has a mesh size of 300 µm, and the outer bag has a mesh size of 100 µm. Both cloths, including the sample, were sewn together with the help of a conventional sewing machine, resulting in a so-called washing sachet. These washing sachets were washed in a conventional washing machine at 60°C without spinning cycle. For the removal of biogenic matter, an enzyme based detergent and a conventional detergent were added for facilitating the washing procedure. Subsequently, a density separation, a vacuum filtration onto cellulose filters (Rotilabo®, Typ 11A, Ø 55 mm, retention 12–15 µm) and fluorescence microscopy enabled by Nile Red (diluted with chloroform) staining were conducted for microplastic isolation and identification. Subsequently, all potential microplastics found on the cellulose filters were photographed, counted, and measured in size. All steps of sample processing were conducted in a closed acrylic box to avoid airborne contamination. The whole implementation of sample handling and processing is described in detail in Philipp et al. (2020).

For polymer identification, selected microplastic particles were manually collected. In addition, a disinfectant step was conducted to exclude a passing on of bacterial or parasitical zoonosis. For this purpose, the cellulose filters containing the stained particles were sprayed with ethanol (70%). After evaporation, the particles showed the same fluorescence qualities as before. Thus, the potential microplastics were selected and manually collected with tweezers or needle pins and placed into a droplet of ethanol (70%) onto an aluminium oxide membrane filter (Anodisc, Ø 47 mm, 0.2 µm pore size, Whatman, Freiburg, Germany). The filter was kept still until the droplet was evaporated and the particles had attached to the filter. Since the transfer of particles was done manually, a loss of particles needs to be taken into account.

The polymer composition of 77 potential microplastics (incl. fragments and fibres) from intestinal sample were analysed by using a µFTIR spectroscope (Hyperion 2000, Bruker, Ettlingen, Germany). All measurements were conducted in transmission.
mode with 32 co-added scans (sometimes 100 scans for very thin fibres) and a spectral resolution of 4 cm\(^{-1}\) in a wavenumber range of 4,000–1,250 cm\(^{-1}\) as aluminium oxide membrane filters are infrared inactive between 3,800 and 1,250 cm\(^{-1}\). For background measurements, the blank aluminium oxide membrane filter was used. For thick particles, for which transmission mode was not suitable, the measurements were conducted in attenuated total reflectance (µATR) mode. Those µATR measurements were conducted between 4,000 and 600 cm\(^{-1}\).

Procedural blanks of the used detergents and materials, e.g., nylon sachets (n = 3), and the working environment (n = 10) were taken into account for avoiding an overestimation caused by secondary contamination. The analysed blank filters of the working environment accompanied the samples from time of collection until the staining procedure was finished.

On average, one fibre and seven particles were found in those procedural blanks and were finally subtracted from the microplastic counts in each parallel sample. Four of those potential microplastics could be collected, manually placed on the aluminium oxide membrane filter and were considered for µFTIR analysis. Moreover, the polymer composition of different equipment materials like the nitrile gloves and shavings of the used acrylic box were additionally determined by FTIR in ATR mode (Vertex 70; Bruker, Ettlingen, Germany) or by µFTIR to avoid an overestimation. For the Vertex measurements, ATR measurements were performed in a wavenumber range of 4,000–370 cm\(^{-1}\) with 8 co-added scans and a spectral resolution of 4 cm\(^{-1}\).

The quantity of found microplastics in comparable groups is given in mean ± standard deviation (M ± SD) to enable a comparison between findings. Moreover, the results were statistically analysed by determining the Cohen’s d and applying a paired t-test using the package “pwr” in the R software Version 4.0.2 (Champely et al., 2020; R Core Team, 2020). Thus, results were described as significant if \(p < 0.05\). In addition, the Figures 2, 5–7 were visualised using the package “ggplot2” (Wickham, 2016).

**RESULTS**

**Quantity and Size**

In total, 30 intestinal samples were available for analyses. An amount of 611 potential microplastics (incl. fragments and fibres, > 100 \(\mu\)m) were found. A secondary contamination of one fibre and seven fragments were considered and subtracted from each sample. Thus, 401 microplastics were finally determined. This amount of microplastics was found in 28 intestinal samples,
FIGURE 2: Width-length distribution of all suspected microplastic particles in intestine samples of 30 harbour porpoise in the size range of 100 µm up to 5,000 µm (n = 611). Secondary contamination was not considered.

in the remaining an absence of microplastics was noticed. When categorising into particle type, 202 fibres and 199 fragments were found. Hence, only two intestines were free from microplastics. Most of the found fibres had a length between 100 and 2,000 µm (Figure 2).

Four additional fibres longer than 5,000 µm, thus defined as mesoplastics (Gregory and Andrady, 2003), were found. Three of them occurred in a sample of an adult male harbour porpoise found in 2017 (lengths: 8,450, 6,964, and 8,029 µm). The fourth fibre (7,365 µm in length) was found in a juvenile male stranded in 2014. Both carcasses were found in the BS. Based on the size, those four fibres were excluded from the results.

FTIR Results

Out of all 611 potential microplastics found in the 30 intestinal samples from the harbour porpoises, originally 94 particles (16%) were selected for polymer identification by µFTIR. Those fibres and fragments were manually collected and placed onto Anodisc membrane filters. Subsequently, 77 particles (12%, n_{fibres} = 28, n_{fragments} = 49) found in the intestinal samples were finally analysed by µFTIR. The remaining 17 microplastics (n_{fibres} = 7, n_{fragments} = 10) were either lost during the sample transport in closed petri dishes to the analysing site or could not be measured due to their small size. Polyester (PEST) was the most frequently found polymer in those investigated intestinal samples (n_{PEST} = 30), followed by polyethylene (PE, n = 17) and polyamide (PA, n = 12) (Figure 3). Furthermore, two polypropylene (PP) particles, one paint chip (acrylic/alkyd with kaolin and styrene) (see Figure in the Supplementary Material), one none further identified polyolefin and one cellulose acetate fibre (which is a semi-synthetic cellulose) were determined. Three acrylic particles, including two with nitrile component, were additionally found. A visualisation of the found polymers (n = 67) is given in Figure 4.

Moreover, the polymer composition of two fragments (one fragment found in an intestine and one from a procedural blank) could not be identified. However, both showed strong similarities and were excluded from the analysis. Only four potential microplastic particles (n_{fibres} = 2, n_{fragments} = 2) were found on all procedural blank filters, and were additionally analysed by µFTIR. One fibre from the blanks was lost and one fragment could not be clearly spectroscopically identified.
However, the other fibre was identified as PEST and the second fragment was determined as varnish with kaolin, styrene and calcium carbonate. Furthermore, two fragments from the intestinal samples had spectra which were highly similar to the varnish which was found in a procedural blank. Hence, those particles were excluded from the analysis. In five cases of potential microplastic particles, biogenic matter was identified and a sixth particle was clearly different from plastic. In addition,
one particle could not be identified due to its small size, since it was broken during the manual collection and placement on the aluminium oxide filter.

**Differences Between Seas**
Comparing the samples based on their origin (NS or BS), there was a significantly higher amount of microplastics in individuals from the BS if compared to the NS (n\textsubscript{BS} = 278; M ± SD = 18.27 ± 14.54; n\textsubscript{NS} = 123; M ± SD = 8.2 ± 7.89; p-value = 0.03). Furthermore, the highest number of 48 microplastic particles was found in an adult female from the BS. When comparing the share of fibres in both seas, significant differences could be revealed (BS: 51.44%; NS: 47.97%; p-value = 0.02). The share of fragments, however, was similar across locations (BS: 48.56%; NS: 52.03%; p-value = 0.1).

**Differences Per Year**
The annual mean values for each sea revealed a higher number of microplastic particles in harbour porpoises from the BS. Furthermore, the range of microplastics found in individuals from the NS was mostly between zero and up to 10 particles per individual in 2015, 2016, and 2018. Only in 2014 and 2017, more than 20 particles were found in the intestinal samples from the NS (2014: 29 and 2017: 21). However, in the BS samples more than 30 particles were found in 2015, 2017, and 2018. The years 2015 and 2018 were the ones with the highest number of findings per individual (44 particles in 2015 and 48 particles in 2018). In two cases from the NS, microplastics were not present (2014 and 2016). In comparison, in the BS microplastic was found in all samples. All this information is presented in Figure 5. However, no significant differences could be determined between the two sample sites (NS and BS; p-value = 0.21), mainly because of the low power of the statistical analysis, which resulted from the low sample size within each year and sea. Following the power analysis, a sample size of at least 12 individuals per year for each sea would be necessary for a reliable trend interpretation (power 80%, p-value = 0.05).

**Differences in Age and Sex**
This study investigates intestinal samples of 13 female and 17 male harbour porpoises (Figure 6). The microplastic burden in females is slightly higher (M ± SD = 13.38 ± 15.41), when compared to the amount of microplastics in males (M ± SD = 13.35 ± 10.56). Certainly, no significant difference in microplastic load could be revealed between sexes (p-value = 0.99). Moreover, no significant differences between adult harbour porpoises (n = 21; M ± SD = 13.82 ± 13.25) and juvenile ones (n = 9; M ± SD = 12.18 ± 12.31) were confirmed (p-value = 0.82), although the microplastic amount in adult ones seemed higher. The two unburdened samples from the NS originated from a juvenile male and an adult female. In both age classes the highest amount of microplastics was found in females (adult: 48 particles; juvenile: 44 particles; see Figure 6).

**Health Status**
The evaluation of the whole GIT revealed an absence of parasite specimens in all investigated intestines. A mild enteritis was found for seven individuals. In detail, in one of those harbour porpoises a mild diffuse eosinophilic enteritis was determined. Furthermore, in two cases (juveniles) a mild diffuse eosinophilic enteritis and a focal mural one were determined. A most likely parasitic etiology was observed in those two harbour porpoises. A fourth individual was affected by a diffuse mild lymphocytic-plasmacellular and eosinophilic infiltration of the lamina propria in combination with a moderate hyperplasia of the Peyer’s plaques. Three further porpoises showed evidence of gastritis (mild and high grade) and enteritis. Whereof two individuals suffered from a mild non-suppurative enteritis, and the third one was also affected by a diffuse moderate lymphocytic-plasmacellular and eosinophilic infiltration of the lamina propria. In total, parasite infestations of e.g. *Pholeter gastrophilus* and *Anisakis simplex* in the multi-chambered stomach was confirmed in 12 harbour porpoises. Harbour porpoises investigated in this study, which were either accidentally bycaught (Siebert et al., 2020) or affected by a pharyngeal entrapment (Gross et al., 2020), showed a microplastic burden of 19.8 particles (SD = 12.77; n = 11) per individual. Compared to the remaining ones (n = 19), where no accidental death could be diagnosed (incl. the three pregnant females), a lower mean value of 10 (SD = 11.59) microplastics per porpoise was identified. Furthermore, if the individual was in a good (n = 9) or moderate (n = 14) nutritional status, the mean number of particles was significantly higher (Mean\textsubscript{good} = 14.11; Mean\textsubscript{moderate} = 16.07) in contrast to a bad nutritional status (n = 7; Mean\textsubscript{bad} = 7) (Figure 7). This is also confirmed by the statistical analysis (p-value = 0.04).
DISCUSSION

This study is the first to evaluate the microplastic burden in marine mammals inhabiting German waters focusing on particles smaller than 1 mm, in marine mammals inhabiting German waters. Furthermore, intestinal samples of harbour porpoises originating from the BS were investigated in microplastic research for the very first time. In total, 93% of all investigated samples from the NS and the BS show a burden of microplastic particles. Minor differences in the range of detected fibres, and no differences in the quantity of found fragments were revealed for both seas. Based on the loss of only 3% of hard parts during sample processing, and the considered secondary pollution, revealed by the preceding publication, the results are reliable and not overrated (Philipp et al., 2020).

Evaluation of the Method and Results

The Nile Red staining is a well-established method in microplastic pre-identification to preselect particles for further investigation (Erni-Cassola et al., 2017). Since some polymer types are melted or dissolved when stained with Nile Red diluted in chloroform (Tamminga et al., 2017), a loss of, e.g., polystyrene or cellulose acetate particles is highly likely. Hence, melted particles were excluded from pre-selection and further polymer identification based on their deficit in quality and the fragile consistency. Furthermore, based on the manual transferring of single particles onto the Anodisc filter, a loss of promising polymer particles has to be taken into account. For future analyses it is advisable to use the Anodisc filter straight away for filtering the samples, instead of transferring it manually after the filtration process.

Based on the fact that many potential microplastics (91%, 73 out of 80, incl. 77 analysed particles found in intestines and 3 analysed particles found on procedural blanks) could be identified as microplastics (incl. n_{intestine} = 77 and n_{proceduralsamples} = 3) by µFTIR analysis, it is highly likely that most particles counted in our study were microplastic. Taken this evaluation and the validation of Philipp et al. (2020) into account (90%), a reliable number of 361 microplastics out of the 401 suspected particles is assumed. Thus, the results show the applicability of the protocol introduced by Philipp et al. (2020) for intestinal samples of cetaceans and determine the actual burden in a reliable way.

Comparison With Other Studies

The determined percentage of 93% (28 out of 30 examined intestines are burdened) coincides with the results of Lusher et al. (2018) investigating carcasses of cetaceans from Irish waters (98%), and Nelms et al. (2019) analysing marine mammals found along the coastline of Great Britain (100%). If focussing on harbour porpoises, the study by Lusher et al. (2018) determined
a microplastic presence in only 6.25% of all investigated cetacean carcasses. One explanation for these differences in microplastic occurrences could be the chosen time period of the review by Lusher et al. (2018). It was conducted between 1990 and 2015. A second explanation could be the used mesh size of at least 300 μm, resulting in the loss of smaller particles, which are included in the study presented herein. Furthermore, no detailed information on the stranding site is given, which would be useful for comparison purpose, since differences in microplastic loads around Ireland (Irish Sea, Celtic Sea and the western coastline facing the open North Atlantic) were determined when the microplastic occurrence was compared at different prawn fishing grounds in 2016 (Hara et al., 2020). Furthermore, the study by Lusher et al. (2018), confirmed the microplastic burden in 21 individuals covering six different cetacean species summing up to 528 investigated GITs. In addition, microplastic (≥300 μm) was only found in Odontoceti species. The study of van Franeker et al. (2018) conducted on harbour porpoises stranded at the Dutch coast, revealed the presence of marine litter items [incl. macroplastic and microplastics (≥1 mm)] in 7% of all investigated stomachs. Van Franeker and colleagues are aware of the fact that due to the mesh size range of used sieves, particles smaller than 1 mm were lost and not considered during their study (van Franeker et al., 2018). As the here presented study confirmed that the main part of found microplastics are smaller than 1 mm (85%), the results of van Franeker et al. (2018) are not comparable with our study. In addition, the size limits of 100 μm and 5 mm, which are based on the used mesh sizes of the washing sachets (Philipp et al., 2020) and the definition of microplastics (Arthur et al., 2009), overlap with the size of zooplankton species, which a variety of invertebrates feed on (Devriese et al., 2015; Fischer, 2019). Thus, investigations in predatory fish or marine mammal species should also focus on these small-sized microplastic particles.

Reference studies of microplastics in marine mammals, especially from the BS, are scarce. Thus, studies investigating fish, sediment and water samples are considered for further discussion. The microplastic burden in different fish species is higher in the BS (11–22%) (Lenz et al., 2016; Beer et al., 2018) when compared to the southern NS (5.4%) (Foekema et al., 2013). In contrast, the microplastic concentrations in surface waters and sediment samples show higher concentration in the southern NS (Karlsson et al., 2017; Lorenz et al., 2019), compared to findings of the BS (Graca et al., 2017; Tammenga et al., 2018). Whereas, a model on the global fibre distribution in surface waters estimated a higher accumulation in the BS (∼1,760 ± 4,500 m⁻³), compared to the North Atlantic region (∼1,800 ± 1,720 m⁻³) (Lima et al., 2021). Nevertheless, an ubiquitous distribution of
FIGURE 7 | Quantity of suspected microplastics (n = 401) in combination with the nutritional status and further information about the carcasses (bycaught: the harbour porpoise was (suspected to be) bycaught by a fishing boat or in a gillnet; flatfish: the harbour porpoise was affected by a pharyngeal entrapment; na: no extraordinary finding could be revealed; pregnant: pregnancy in female was noticed). The nutritional status is coded as follows: (1): bad, (2): moderate, and (3): good.

Polymer Findings
In this study, the most frequently found polymer was PEST. Based on the fact that six fragments and 24 fibres were found in the intestinal samples and only one PEST fibre was identified in one of the procedural blanks, PEST microplastics were still taken into account and were not excluded from this study. Additionally, the procedural blanks show a low amount of fibres (Øfibres = 1 and Øfragments = 7 per procedural blank) and were already subtracted from the results presented here. To control for microplastic contamination, only cotton gloves and lab coats were worn while processing the samples (Philipp et al., 2020). Other studies investigating GIT samples of marine mammals from the North Atlantic found PEST particles, even though high protective measurements were used (Lusher et al., 2015a; Nelms et al., 2019). In addition, a high amount of synthetic fibres like PEST fibres were determined in the Northeast Atlantic region (Thompson, 2004; Lusher et al., 2014), and in inhabiting fish species (Lusher et al., 2013; McGoran et al., 2017; Pereira et al., 2020). The twelve found polyamide particles were not excluded from the analysis, since fibres of the used nylon cloth (PA) are obviously identifiable due to their unique fibre pattern (Philipp et al., 2020), and are clearly different from the PA particles found in the intestinal samples. Thus, those fibres were immediately excluded while pre-selecting, counting and collecting particles for microplastic burden of individuals of both seas should be evaluated. Thus, a continuation of the herein presented approach is advisable and is intended.

synthetic particles along the German BS coast is assumed (Stolte et al., 2015). Furthermore, the fact that marine mammals of the BS might be more exposed to marine litter than in the NS was already confirmed (Unger et al., 2017). Thus, a higher risk of microplastic burden in the BS could be hypothesised. The results obtained in this study underline the findings of marine litter and support the following hypothesis: Investigated harbour porpoises of the BS show a higher number of microplastic particles (incl. fibres and fragments), in contrast to individuals from the NS in each year. In particular, two females from the BS show a high amount of microplastics (44 and 48 particles). For avoiding an overassessment in future research, higher sample sizes per sea are highly recommended.

It will be worth to strive for a reference study investigating the area of the Baltic Proper, since this area is assumed to accumulate pollutants from the whole BS (Stolte et al., 2015), and the here occurring harbour porpoise subspecies is critically endangered (Carlén et al., 2018). Nevertheless, the time span of the available sample collection and the quantity of samples is still too low for identifying a trend in both seas. After statistical assessment with paired $t$-test and Cohen’s d, the samples size has to be increased if reliable comparisons in microplastic burden of individuals of both seas should be evaluated. Thus, a continuation of the herein presented approach is advisable and is intended.
analyses. Furthermore, not only PA fibres but also PA fragments were found. Previously, high numbers of PA microplastics were identified in GITs of marine mammals (Nelms et al., 2019) and in fish from the North-East Atlantic region (Lusher et al., 2013; McEnerney et al., 2017).

Both, the NS and the BS are enclosed by highly populated countries (Halpern et al., 2008, 2015). The system of waste management is relatively sophisticated in European countries, but there is still a high amount of produced waste, thus the likelihood of litter input into the marine environment is increasing (Andreassi Bassi et al., 2017). Another source of microplastic particles, and in particular of fibres which originate from washed clothes are waste water treatment plants (WWTP) (Browne et al., 2011; Bretas Alvim et al., 2020). Moreover, it is well known that WWTP are not able to hold back microplastic fibres and particles (Dubaish and Liebezeit, 2013; Mani et al., 2016). Thus, it could be assumed that the found PA and PEST microplastics originated from waste water, which is transported by rivers and ends up in the marine environment (Browne et al., 2011; Dubaish and Liebezeit, 2013; Mani et al., 2016).

Besides land-based sources, which are responsible for 80% of marine litter inputs (Jambeck et al., 2015), sea-based sources like lost fishing gear or waste generated on ships or offshore installations need to be taken into account (Galgani et al., 2013). Furthermore, both seas are highly impacted by anthropogenic exploitations, such as fishing activities (Catchpole et al., 2005; Halpern et al., 2008, 2015). Thus, the fishing industry is a potential source of found microplastics such as PA and PEST, as well as PE and PP (Pruter, 1987; Deshpande et al., 2020).

Globally, the demand for PE is high (PlasticsEurope, 2020). Thus, it is not surprising that a high amount of PE microplastics was identified in this study. This result is comparable to findings in surface waters of the BS (Gewert et al., 2017). Studies investigating microplastics in marine mammals or fish from the North Atlantic found lower amounts of PE and PP (Lusher et al., 2013; Nelms et al., 2019). Due to the relatively low proportion of microplastics investigated with µFTIR spectroscopy, no final conclusions can be drawn on differences in polymer occurrences in the NS and the BS.

One of the microplastics found was identified as an acrylic or alkyd chip with evidence of kaolin and styrene. Such polymers and additives are commonly used for car (Yang et al., 2012) and ship paints (Lee et al., 2021). Hence, the microplastic chip that was found in the intestine of one harbour porpoise originating from the NS could have derived from abraded ship paint. It is well known that weathered paint from ship and boat surfaces is regularly released into the ocean (Song et al., 2014; Lacerda et al., 2019). Hence, paint fragments can be ingested by marine mammals such as cetaceans as also suggested by Lusher et al. (2018). Unfortunately, this pre-mentioned study did not provide information on paint polymer types. Furthermore, paint chips have already been found in Australian sea turtles (Caron et al., 2018) or in pelagic fish (Ory et al., 2017). Thus, an unintentional prey intake of paint chips by, e.g., fish while preying on zooplankton is assumed (Ory et al., 2017), and the transport through the food web could be considered.

Trophic Accumulation via the Food Web?
The study of van Franeker et al. (2018) pointed out, that the microplastic presence in harbour porpoises most likely occurred due to an unintentional intake while feeding on burdened fish specimens close to the sediment. Besides pelagic species like herring (Clupea harengus), gadoids (e.g., Merlangius merlangus) and sprat (Sprattus sprattus), different benthic sandeel species (e.g., Ammodytes marinus) and gobies (Gobiidae) form the main prey of harbour porpoises (Leopold, 2015). The linkage between burdened prey species and top predators was previously confirmed for grey seals preying on gadoids (Hernandez-Milian et al., 2019). Moreover, three of the here analysed harbour porpoises asphyxiated by pleuronectiforms (e.g., Solea solea) and two further individuals were bycaught in gillnets for benthic and demersal fish species. Those harbour porpoises showed a high mean microplastic amount per individual (M ± SD = 17.7 ± 12.42). Thus, a relationship between preying on benthic fish and the presence of microplastics in predator species is highly assumed. Furthermore, the bycaught or asphyxiated porpoises were in a better nutritional status in comparison to the remaining ones. However, a good nutritional status is not synonymous with a good health of a harbour porpoise (Siebert et al., 2020). Based on the results presented herein, it can be assumed that a harbour porpoise, which is not physically hampered in feeding or hunting, will accumulate more microplastics in the GIT than a diseased one. The results of this study are supported by the assumptions of Leopold (2015) and Rummel et al. (2016) who claimed that 5,000 individual gobs per day are needed for maintaining a harbour porpoise’s good physical condition, and the mean burden of 0.03 particles in each benthic fish may result in a daily intake of almost 150 microplastic particles by one adult harbour porpoise. Kastelein et al. (1997) detected a passage time of 143–196 min in captive harbour porpoises. If the assumption is that a full digestion takes ~3 h, a harbour porpoise will approximately defecate eight times a day. Thus, the potential accumulation of 150 microplastics has to be divided by eight, resulting in 18.75 particles per defecation. Certainly, this is just a rough estimate and many factors influence egestion and ingestion rates, but this calculation coincides with the findings of microplastics in samples from the BS (M ± SD = 18.27 ± 14.54).

Impact on the Health Status
Across all 30 individuals no intralesional parasites could be identified in the intestines. However, seven individuals showed an enteritis, whereof two are attributed to parasites and another one accompanied by a hyperplasia of Peyer’s plaques. These accumulations of follicles are known to adsorb different kinds of particles (biogenic and synthetic), and to transfer it into the lymph system when the particle is adequately small in size (Ensign et al., 2012). In total, only two juvenile harbour porpoises originating from the BS showed changes in the Peyer’s plaques, where a low number of suspected microplastics was found (n = 17 and n = 15), in comparison to the other 28 specimens. However, the stomachs were affected by parasites in 12 cases, whereof 11 suffered from a gastritis. Two harbour porpoises from the BS showed changes in the stomach tissue, including gastritis,
Atlantic population. Thus, a higher microplastic burden in the western Baltic population, if compared to the North-East. A higher risk of exposure to microplastics was revealed for faecal samples of alive individuals are the most feasible approach based by field conditions. Furthermore, ethical concerns arise as especially in free-ranging marine mammals, are complicated microplastic investigations and experiments in mammals, and intake and egestion rate, further research is needed. Nevertheless, confirm the egestion of microplastic particles. For evaluating the health monitoring, and (3) the findings in the rectum and faeces of focussing on only one part of the GIT: (1) avoidance of secondary contamination in smaller samples is easier, (2) the remaining carcass and GIT can be entirely evaluated for a health monitoring, and (3) the findings in the rectum and faeces confirm the egestion of microplastic particles. For evaluating the intake and egestion rate, further research is needed. Nevertheless, microplastic investigations and experiments in mammals, and especially in free-ranging marine mammals, are complicated based by field conditions. Furthermore, ethical concerns arise as indicated by Nelms et al. (2018). Thus, samples of carcasses and faecal samples of alive individuals are the most feasible approach to assess the microplastic burden in marine mammals. This is the first study investigating harbour porpoises from different subpopulations for microplastics, and revealed differences in microplastic presence in the NS and the BS. A higher risk of exposure to microplastics was revealed for the western Baltic population, if compared to the North-East Atlantic population. Thus, a higher microplastic burden in the BS is assumed. Furthermore, evidence for the continuous accumulation of microplastics via the food web was given, but could not significantly be confirmed in adult individuals, compared to juvenile ones. Additionally, there is no significant difference in the quantity of synthetic particles in male or female harbour porpoises. To gain further knowledge on differences in sex or age, the quantity of samples has to be increased in future research.

An important relationship between a good or moderate nutritional status and the occurrence of microplastics is demonstrated in this study. Moreover, the egestion and thus, a discharge of microplastic particles out of the organism could be confirmed. No relationship between parasites, tissue damage and microplastic presence could be identified. Therefore, a histological investigation of cell damage or tissue damage localisation with the help of biomarkers would be advisable in future research. Further investigations are needed for evaluating the rate of accumulation and burden in harbour porpoises in the different seas. Indeed, this study outlines first evidence in retrospective microplastic burden. Nevertheless, a higher sample size, as well as a larger temporal coverage is needed to reliably estimate trends in the microplastic burden in harbour porpoises. Furthermore, this study supports the need for a comprehensive marine litter monitoring in predatory species to gain knowledge on accumulation processes and health assessment in apex species of the marine food web.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author/s.

ETHICS STATEMENT

Ethical review and approval was not required for the animal study because the investigations on marine mammals for scientific and conservation purpose were conducted in accordance with national and international regulations. During this study samples of dead found specimens were analysed and thus no invasive methods were used.

AUTHOR CONTRIBUTIONS

CP, BU, and US conceptualized the study and acquired funding. CP conducted the laboratory analysis of processing the samples and isolate microplastics, assisted by BU. CP did the statistical analysis. JHEK provided the µFTIR and FTIR for polymer type analysis. SME conducted the polymer identification by µFTIR and FTIR. CP and SME generated the figures. The manuscript was prepared by CP, and contributed editing with perspectives and arguments was done by BU, SME, JHEK, and US. All authors approved the submitted version.

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REFERENCES

Andersen, L. W. (2003). Harbour porpoises (Phocoena phocoena) in the north atlantic: distribution and genetic population structure. NAMMCO Sci. Publ. 5, 11–29.

Andreasi Bassi, S., Christensen, T. H., and Damgaard, A. (2017). Environmental performance of household waste management in europe – An example of 7 countries. Waste Manag. 69, 545–557. doi: 10.1016/j.wasman.2017.07.042

Arthur, C., Baker, J., and Bamford, H. (2009). Proceedings of the international research workshop on the occurrence, effects, and fate of microplastic marine debris. Sept 9-11 2008.

Beer, S., Garm, A., Huwer, B., Dierking, J., and Nielsen, T. G. (2018). No increase in marine microplastic concentration over the last three decades – a case study from the baltic sea. Sci. Total Environ. 621, 1272–1279. doi: 10.1016/j.scitotenv.2017.10.011

Bretas Alvim, C., Mendoza-Roca, J. A., and Bes-Piá, A. (2020). Wastewater treatment plant as microplastics release source – quantification and identification techniques. J. Environ. Manage. 255:109739. doi: 10.1016/j.jenvman.2019.109739

Browne, M. A., Crump, P., Niven, S. J., Teuten, E., Tonkin, A., Galloway, T., et al. (2011). Accumulation of microplastic on shorelines worldwide: sources and sinks. Environ. Sci. Technol. 45, 9175–9179. doi: 10.1021/es201811s

Carlen, L., Thomas, L., Carlström, J., Aminudin, M., Teilmann, J., Tregenza, N., et al. (2018). Basin-scale distribution of harbour porpoises in the baltic sea provides basis for effective conservation actions. Biol. Conserv. 226, 42–53. doi: 10.1016/j.biocon.2018.06.031

Caron, A. G. M., Thomas, C. R., Berry, K. L. E., Mott, C. A., Ariel, E., and Brodie, J. E. (2018). Ingestion of microplastic debris by green sea turtles (Chelonia mydas) in the great barrier reef: validation of a sequential extraction protocol. Mar. Pollut. Bull. 127, 743–751. doi: 10.1016/j.marpolbul.2017.12.062

Carr, K. E., Smyth, S. H., McCullough, M. T., Morris, J. F., and Moyes, S. M. (2012). Morphological aspects of interactions between microparticles and mammalian cells: intestinal uptake and onward movement. Prog. Histochem. Cytochem. 46, 185–252. doi: 10.1016/j.phpro.2011.11.001

Catchpole, T. L., Frid, C. L. I., and Gray, T. S. (2005). Discards in north sea fisheries: causes, consequences and solutions. Mar. Policy 29, 421–430. doi: 10.1016/j.marpol.2004.07.001

Champely, S., Ekstrom, C., Dalgaard, P., Gill, J., Weibelzahl, S., Anandkumar, A., et al. (2020). Package “pwr.”. Available Online at: https://github.com/heliosdrm/pwr.

Deshpande, P. C., Philip, G., Brattebo, H., and Fet, A. M. (2020). Using material flow analysis (MFA) to generate the evidence on plastic waste management from commercial fishing gears in norway. Resour. Conserv. Recycl. X 5:100024. doi: 10.1016/j.rcrx.2019.100024

Dreviése, L. L., van der Meulen, M. D., Maes, T., Bekart, K., Paul-Pont, I., Frère, L., et al. (2015). Microplastic contamination in brown shrimp (crangon crangon, linnaeus 1758) from coastal waters of the southern north sea and channel area. Mar. Pollut. Bull. 98, 179–187. doi: 10.1016/j.marpolbul.2015.06.051

Dubash, F., and Liebezeit, G. (2013). Suspended microplastics and black carbon particles in the jade system, southern north sea. Water Air Soil Pollut. 224:1352. doi: 10.1007/s11270-012-1352-9

Ehrens, L. M., Cone, R., and Hanes, J. (2012). Oral drug delivery with polymeric nanoparticles: the gastrointestinal mucus barriers. Adv. Drug Deliv. Rev. 64, 557–570. doi: 10.1016/j.addr.2011.12.009

Erni-Cassola, G., Gibson, M. L., Thompson, R. C., and Christie-Oleza, J. A. (2017). Lost, but Found with nile red: a novel method for detecting and quantifying small microplastics (1 mm to 20 µm) in environmental samples. Environ. Sci. Technol. 51, 13641–13648. doi: 10.1021/acs.est.7b04512

Farrell, P., and Nelson, K. (2013). Trophic level transfer of microplastic: mytilus edulis (L.) to carcinus maenas (L.). Environ. Pollut. 177, 1–3. doi: 10.1016/j.envpol.2013.01.046

Fischer, E. (2019). Distribution of microplastics in marine species of the Wadden Sea along the coastline of Schleswig-Holstein, Germany. Hamburg, Germany: Hamburg University.

Foekema, E. M., De Gruijter, C., Mergia, M. T., van Franeker, J. A., Mark, A. J., and Koolmans, A. A. (2013). Plastic in north sea fish. Environ. Sci. Technol. 47, 8818–8824. doi: 10.1021/es400931b

Fossi, M. C., Coppola, D., Baini, M., Giannetti, M., Guerranti, C., Marsili, L., et al. (2014). Large filter feeding marine organisms as indicators of microplastic in the pelagic environment: the case studies of the mediterranean basking shark (cetorhinus maximus) and fin whale (balaenoptera physalus). Mar. Environ. Res. 100, 17–24. doi: 10.1016/j.marenvres.2014.02.002

Frid, C., Hammer, C., Law, R., Loeng, H., Pawlak, J. F., Reid, P. C., et al. (2003). Environmental Status of the European Seas. Available online at: https://archimer.ifremer.fr/doc/00040/15135/12473.pdf.

Galgani, F., Hanke, G., Werner, S., and De Vrees, L. (2013). Marine litter within the european marine strategy framework directive. ICES J. Mar. Sci. 70, 1055–1064. doi: 10.1093/icesjms/fsi122

García-Cegarra, A. M., Jung, J.-L., Orrego, R., Padilha, J., de, A., Malm, O., et al. (2021). Persistence, bioaccumulation and vertical transfer of pollutants in long-finned pilot whales stranded in chilean patagonia. Sci. Total Environ. 770:145259. doi: 10.1016/j.scitotenv.2021.145259

Gaskin, D. E. (1984). The harbour porpoise (Phocoena phocoena) regional populations, status, and information on direct and indirect catches. Rep. Int. Whal. Comm. 24:18.

Gewert, B., Ogonowski, M., Barth, A., and MacLeod, M. (2017). Abundance and composition of near surface microplastics and plastic debris in the stockholm archipelago, baltic sea. Mar. Pollut. Bull. 120, 292–302. doi: 10.1016/j.marpolbul.2017.04.062

Graca, B., Szewc, K., Zakrzewska, D., Dołęga, A., and Szczerbowska-Boruchowska, M. (2017). Sources and fate of microplastics in marine and beach sediments of the southern baltic sea—a preliminary study. Environ. Sci. Pollut. Res. 24, 7650–7661. doi: 10.1007/s11356-017-8419-5

SUPPLEMENTARY MATERIAL

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Roch, S., Friedrich, C., and Brinker, A. (2020). Uptake routes of microplastics in fishes: practical and theoretical approaches to test existing theories. *Sci. Rep.* 10:3896. doi: 10.1038/s41598-020-60630-1

Rummel, C. D., Löder, M. G. J., Fricke, N. F., Lang, T., Griebeler, E.-M., Janke, M., et al. (2016). Plastic ingestion by pelagic and demersal fish from the north sea and baltic sea. *Mar. Pollut. Bull.* 102, 134–141. doi: 10.1016/j.marpolbul.2015.11.043

Setälä, O., Fleming-Lehtinen, V., and Lehtiniemi, M. (2014). Ingestion and transfer of microplastics in the planktonic food web. *Environ. Pollut.* 185, 77–83. doi: 10.1016/j.envpol.2013.10.013

Siebert, U., Pawliczka, I., Benke, H., von Vietinghoff, V., Wolf, P., Pilats, V., et al. (2020). Health assessment of harbour porpoises (PHOCOENA PHOCOENA) from baltic area of denmark, germany, poland and latvia. *Environ. Int.* 143:105904. doi: 10.1016/j.envint.2020.105904

Unger, B., Herr, H., Benke, H., Böhmert, M., Burkhardt-Holm, P., Döhne, M., et al. (2017). Marine debris in harbour porpoises and seals from german waters. *Mar. Environ. Res.* 130, 77–84. doi: 10.1016/j.marenvres.2017.07.009

van Franeker, J. A., Bravo Rebolledo, E. L., Hesse, E., IJsseldijk, L. L., Kühn, S., Leopold, M., et al. (2018). Plastic ingestion by harbour porpoises *Phocoena phocoena* in the netherlands: establishing a standardised method. *Ambio* 47, 387–397. doi: 10.1007/s13280-017-1002-y

Wickham, H. (2016). *ggplot2: elegant graphics for data analysis*. New York City: Springer.

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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