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You Are What You Eat, Microplastics in Porbeagle Sharks From the North East Atlantic: Method Development and Analysis in Spiral Valve Content and Tissue

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Researchers worldwide are studying the environmental distribution and impacts of manufactured or environmentally fragmented small pieces of plastics, so called microplastics (<5 mm). These microplastics eventually build up in the marine environment, threatening marine ecosystems. The magnitude, fate and effects of these microplastics across the food web are largely unknown. Here, we measured digested microplastics in a top predator and critically endangered species, the North-East Atlantic Porbeagle shark (Lamna nasus), and compared this with general health conditions. A method for quantifying microplastics in spiral valves of porbeagle sharks was developed. Microplastics were detected in all spiral valves, up to 10.4 particles per g wet weight (w.w.) content and 9.5 particles per g w.w. tissue. This equates to individual microplastics loads as high as 3850 particles per spiral valve, most likely a result of trophic transfer. No statistically significant correlations were found between the average number of plastic particles in spiral valve content and tissue and the Condition and Hepatosomatic Index of porbeagle sharks. The results of this research show that North-East Atlantic porbeagle sharks ingest and digest microplastics and that there is a potential for microplastic biomonitoring using this species. More research is needed to detect possible health effects of microplastic contamination in these apex predators.

Keywords: marine litter, microplastics, top predator, porbeagle shark, Lamna nasus, Raman spectroscopy, trophic transfer, food web accumulation

INTRODUCTION

Concerns about plastics in the environment are mounting. Plastic items of all sizes end up in marine waters by inappropriate waste disposal and human behavior, resulting in ever-increasing marine litter concentrations (Jambeck et al., 2015). The anthropogenic litter currently floating at sea, according to estimates, accounts for at least 5.25 trillion particles of plastic (Eriksen et al., 2014) and can be found in the most remote marine (Lusher et al., 2015b) and fresh water systems (Eerkes-Medrano et al., 2015).
They have a wide range of sizes, causing impacts across the entire food web. The larger sized items, such as the macroplastics (>20 mm diameter) and mesoplastics (5–20 mm), are known to cause entanglement and/or obstruction of the gut when ingested (Jacobsen et al., 2010; Foekema et al., 2013; Lusher et al., 2013, 2015a). The smaller particles, microplastics (<5 mm), are available to the smallest marine organisms, building up in food webs (Setlää et al., 2014), causing amongst other things abrasion, blockage of digestive tracts, starvation (Werner et al., 2016), and death (Maes et al., 2020). Laboratory studies have shown that plastic particles in the lower micro- and nano-sized range can cross cell membranes, causing tissue damage (Brown et al., 2008; UNEP, 2016). In addition, the easily ingestible microplastics can form a pathway in the entry of chemical contaminants. These chemicals are either leached from the plastic material itself (e.g., additives) or adsorbed to plastic from the surrounding water and thus taken up together with the plastic by the organism when ingested (Teuten et al., 2009; Koelmans et al., 2014; Yeo et al., 2017). The chemical concentrations of these persistent pollutants accumulate in biota and are often higher at the top of the food chain (Whitacre, 2010; Bendall et al., 2014; Nicolaus et al., 2016).

Microplastic ingestion has been reported in a wide range of marine organisms, such as sea cucumbers, mussels, lobsters, amphipods, lugworms, barnacles, and zooplankton (GESAMP, 2015a; UNEP, 2016). In addition, it has been reported in higher trophic levels, such as fish (e.g., herring), birds (e.g., Northern Fulmars) and marine mammals (e.g., whales) (Van Franeker et al., 2011; Foekema et al., 2013; Lusher et al., 2013, 2016; Besseling et al., 2015; Bellas et al., 2016; Steer et al., 2017; Nelms et al., 2018). It is suggested that the larger marine animals obtain the microplastics directly via ingestion (e.g., filter feeding) (Sampaio et al., 2018) or indirectly by trophic transfer via fish which consumed microplastics (Lusher et al., 2015a; Fossi et al., 2017).

Therefore, the impact of microplastics might be an additional problem for a critically endangered species, such as the North East Atlantic top-predator porbeagle shark (Bendall et al., 2014; IUCN, 2015; Nicolaus et al., 2016). Porbeagle sharks, Lamna nasus (Bonnerterre, 1788), are stout-bodied sharks with large black eyes and pointed snouts attaining a maximum length of about 355 cm (total length) (Francis et al., 2009). They are mostly found in the cold-temperate areas in the upper pelagic zone of the North Atlantic, South Atlantic and South Pacific oceans. Their diet consists of small to medium-sized pelagic and ground fishes, such as lancet fish, herring, mackerel, lance, lumpfish, flounders, hake, and cod, but feed also on squid and invertebrate (Joyce et al., 2002). Their low fecundity, late maturation age and prolonged gestation period makes them susceptible to overfishing (Bendall et al., 2012). In addition, they are long-lived species and are therefore potential targets for contaminant accumulation, such as mercury (Bendall et al., 2014; Nicolaus et al., 2016). Moreover, these sharks might be subject to direct or indirect plastic ingestion, as they reside in marine waters with reported high levels of plastic debris (Pham et al., 2013; Eriksen et al., 2014; Woodall et al., 2015b; Hartley et al., 2018), including microplastics (Lusher et al., 2014; van Sebille et al., 2015; Maes et al., 2017b), and the presence of microplastics in their prey (Foekema et al., 2013; Lusher et al., 2013, 2016; GESAMP, 2015b).

The concerns about the ecological impact of microplastics match the increasing trend in microplastic studies and publications of the last 4–5 years (Barboza and Gimenez, 2015; Barboza et al., 2018). One of the main topics of these studies is the development of analytical techniques and detection methods for the presence of microplastics that can be applied to sediment, water and biota samples (GESAMP, 2019). Differences in biota types, sample size and a range of other parameters (e.g., feeding behavior) have led to a variety of procedures for the analysis of microplastics in biota (GESAMP, 2019). Different steps to clean-up biota samples without affecting the microplastics (i.e., removal of biological tissues, sediment grains, and other non-plastic organic fractions) make harmonization difficult. A standard procedure that can be applied to different organisms is, therefore, not yet available (GESAMP, 2019). Moreover, most research has been conducted on small amounts of soft tissue and opts to exclude particles of certain sizes (Besseling et al., 2019; GESAMP, 2019). Harmonization in applied methodologies is needed to improve the comparison between studies (Karlsson, 2014), but methodologies should also be stretchable in order to obtain more knowledge of the magnitude, fate and effects of microplastic particles in marine organisms throughout the food web (GESAMP, 2019).

Macro debris ingestion and entanglement has been commonly observed in a range of sharks species (Cliff et al., 2002; Bernardini et al., 2018; Sampaio et al., 2018; Smith, 2018; Barreto et al., 2019). There are only few published studies identifying plastics and/or microplastics in sharks (Diana et al., 2014; Fossi et al., 2014, 2017; Cartes et al., 2016; Panti et al., 2016; Alomar and Deudero, 2017; Bernardini et al., 2018; Sampaio et al., 2018; Smith, 2018). Available microplastic studies mostly analyzed small types of Elasmobranchs where the entire stomach and intestines was dissected and/or digested for subsequent microscopic analysis (Alomar and Deudero, 2017; Smith, 2018). Studies dealing with larger sharks are limited to stomach content analysis and report only macro litter items (Sampaio et al., 2018) and/or microplastics within the range 5–1 mm (Bernardini et al., 2018). There are many opportunities for bias in estimating diets from the stomach contents of elasmobranchs in the field (Cortés, 1997; Bucking, 2015). Predatory fish frequently regurgitate their stomach contents (Hyslop, 1980; Cortés, 1997), potentially removing part of the plastic load. To reduce such monitoring bias and to focus on the ingested and digested microplastic fraction, we only analyzed the spiral valve content and tissue.

In this study we aim to assess the impact of microplastics on the general health condition of marine top predators, Porbeagle sharks (Lamna nasus) from the North-East Atlantic Ocean. Existing studies focus on ingestion rather than digestion and use stomach, instead of spiral valve, contents. Different existing methods were adapted and tested to extract the spiral valve content and analyze this content for microplastics. The best performing method was then applied to test the hypotheses that (i) porbeagle sharks ingest and digest microplastic and (ii) the uptake of microplastics adversely affects the general health condition (as assessed by Condition Index and Hepatosomatic Index) of Lamna nasus sharks.
MATERIALS AND METHODS

Sampling and Study Design

General Health Indications and Observations

Cefas, the Centre for Environment, Fisheries and Aquaculture Science, received 53 dead porbeagle sharks provided by commercial fishers for research purposes. These specimens were captured as seasonal by-catch in Celtic Sea gillnet fisheries with nearly all fish captured between August and October 2014 (Ellis and Bendall, 2015; Nicolaus et al., 2016). Individual specimens were frozen after being brought ashore and were subsequently transferred to Cefas (Lowestoft) for processing, where they were sexed, measured and weighed (Ellis and Bendall, 2015; Figures 1A–D). Body length was determined by the Total Length and refers to the length measured from the tip of the snout to the top of the upper lobe of the caudal fin in a flexed down position, with this measurement made in a direct line under the body (Ellis and Bendall, 2015). After examination and opening of the internal cavity, samples of the spiral valve were tightened at the top and bottom (Bernardini et al., 2018) with cotton strings before cutting, bagged separately and stored in the freezer (Figures 2A–D). The spiral valve, part of the intestinal tract, was made available for research on microplastics. Due to the large size of each porbeagle spiral valve, time and budget limitations, 13 randomly chosen frozen spiral valves were sent to The Vrije Universiteit Amsterdam for further microplastic analysis. The spiral valves were inspected, and measurements were taken while collecting the content and tissue. Background information on the digestive tract of the porbeagle shark is found in Supporting Information.

Method Development

Sample Collection

The frozen spiral valves were put in a lukewarm water bath for approximately 3 h to speed up the thawing process, with 2 additional zip lock plastic bags to prevent the risk of leaking. The spiral valve was weighed (Sartorius CP2202S) before and after excess material was cut away (part of gonads, rectal gland and blood vessel remains). Other remaining material was removed from the exterior tissue with milliQ water.

To determine the presence of microplastics, the content of each spiral valve and the inside tissue of the valve were examined. Abnormalities, such as tissue damage and the presence of parasites, were also noted before subsampling, the valve content was homogenized by 1 min hand-stirring with a metal spoon. Spiral valve tissue samples were used to estimate the amounts of plastic particles trapped inside.

Four methods to collect the spiral valve content were explored (Figures 3A–D) on 4 individual spiral valves, the best method was then used on the remaining 9 spiral valves. In the first two methods, the content material was recovered from the spiral valves by hanging the spiral valve vertically from a tripod. In the first method, the spiral valve was hung above six different sieves (1 mm, 710, 500, 355, 200, and 100 µm) to separate the collected content immediately based on size. A glass funnel was used to keep the top part of the spiral valve open and 1 L of milliQ water was flushed through the spiral valve to rinse the remaining content from the spiral valve. In the second method, the spiral valve was hung on the tripod without the funnel and sieves. The content was forced out of the spiral valve by squeezing the spiral valve from top to bottom in downwards strokes, collecting the content in a glass bowl (Figure 4). In the third method, the spiral valve was cut open to scrape out the content with a metal spoon. For the fourth method, the spiral valve was cut open in a glass bowl with 200 mL of milliQ water to collect the content through washing. The collected content of all spiral valves was stored separately in glass bottles and the weight was determined.
The bottles were covered with aluminum foil and stored in a freezer at minus 20°C. The weight of the emptied spiral valve was determined afterward. The emptied spiral valve was cut open with scissors from the rectum up to the top part of the spiral valve. Pieces of the inside tissue were collected by cutting small fractions of the top, middle (2×) and lower pleat-shaped chambers. The 1 g tissue samples were placed in separate 25 mL glass containers, covered with foil and stored in the freezer at −20°C.

Digestion and Validation

Different methodologies for the digestion of spiral valve content and tissue were tested. The procedures tested included an enzymatic (Cole et al., 2014; Karlsson, 2014), acid (De Witte et al., 2014) and alkaline (Foekema et al., 2013; Enders et al., 2017; Kühn et al., 2017; Thiele et al., 2019) digestion. In addition to these selected procedures some further acid and alkaline digestion tests, using different sample volumes and/or digestion times, were performed. An overview of all digestion tests is given in Figure 5, more methodological details can be found in Supporting Information. Different types of microplastic particles were also subjected to the different digestion methods to assess whether plastic particles could be recovered and/or were affected by the digestion procedures. After exposure to the acid and alkaline solutions, recovery and changes in color and shape of the 12 tested plastic types were evaluated under the microscope. In addition, the best performing digestion procedure was further validated, by assessing the impact of the digestion procedure on the weight of the spiked plastic material (Supporting Information). All chemicals were purchased from Sigma-Aldrich unless stated otherwise.

Following the results of method testing, the alkaline potassium hydroxide (KOH, Riedel-de Haén, 10%) method (Foekema et al., 2013; Enders et al., 2017; Kühn et al., 2017; Thiele et al., 2019) was adopted to extract microplastics from the spiral valve content and tissue. The removed content of each spiral valve was homogenized and five subsamples of 2 g were taken from each spiral valve. The subsamples were added into 25 mL glass containers and a 6 mL solution of 10% KOH was added. The samples were incubated in a fume hood at room temperature for 2–9 days. Of each spiral valve, four tissue samples of 1 g were placed into 25 mL glass containers and a 3 mL solution of 10% KOH was added. The tissue samples were incubated for 17 days. During the digestion procedures, parasites were included in the subsamples. When only one parasite was found in a spiral valve, the parasite was digested with one of the subsamples. When multiple parasites were found, they were homogenized with the contents and included in the samples. To provide some additional information on microplastics in parasites, the content of one rectal gland (1.2 g), where high amounts of parasites were observed, was incubated with a 3.6 mL solution of 10% KOH for 3 days and one sample with nine small parasites, individually picked from the spiral valves, was incubated with a 2 mL solution of 10% KOH for 2 days.

Filtration

With digestion completed, samples were vacuum filtered through a Whatman GF/F filter (pore size 0.7 μm, Ø 47 mm) to retain the microplastics (Claessens et al., 2013). The emptied glass containers were rinsed with milliQ water and vacuum filtered again. The effect of combining KOH with an additional alkaline solution (e.g., bleach or soap) to remove lipid residues was tested (Enders et al., 2017). To simplify existing procedures (Foekema et al., 2013; Lusher et al., 2016; Enders et al., 2017), after filtration, the filter was covered with 2.5 mL 20% Extraneo® (MA 01 liquid, alkaline soap) for a maximum of 30 min to decrease the amount of digest. After adding the soap, the glassware was rinsed three times to make sure all sample was filtered through. A total of 50 mL of filtered milliQ water was used for every sample. Samples of both the content and tissue were filtered separately. When a filter became clogged, multiple filters were used. The difference between filters of various tests was determined by visual and microscopic inspection. Filters were stored and frozen in a petri-dish until further analysis.

Microplastic Analysis

Quality Control

An essential aspect of this study was the monitoring and prevention of contamination and an adequate recovery of microplastics from the sample matrix. Therefore, all tests were
FIGURE 4: Overview of the different digestion methods, sample sizes, and digestion time frames. *Included boiling.
performed in a closed environment (e.g., fume hood), except for spiral valve weighing and the microscope analysis. The fume hood was cleaned regularly throughout the study with ethanol. A cotton lab coat was worn, and blue nitrile gloves were used during the sample collection, digestion and filtration procedures. Only sterile glassware was used and thoroughly rinsed three times with milliQ water prior to use. Metal spoons and scissors were acetone-washed prior to use and all equipment and samples were covered with aluminum foil or glass. In between filtrations, glassware was cleaned with milliQ water. In addition to these steps, it is important to note that all samples were processed by a single researcher. Although Quality Assurance and Quality Control (QA/QC) tools and certified reference materials are lacking for this emerging field (GESAMP, 2019), some additional precautionary steps were introduced to prevent and control contamination (Woodall et al., 2015a). To eliminate any contamination from the chemical solutions, all solutions were filtered on GF/F filters (pore size 0.7 µm, Ø 47 mm) before use. Only GF/F (pore size 0.7 µm, Ø 25 mm) filtered milliQ water was used to make up solutions or clean glassware. The fume hood was cleaned before every procedure. Potential airborne contamination for processes in the fume hood was determined by leaving 3 clean GF/F filters (pore size 0.7 µm, Ø 47 mm) in 3 open petri-dishes for 30 min (the maximum time needed to collect the content and tissue samples in one spiral valve). To determine air contamination during sample weighing, 3 GF/F filters (pore size 0.7 µm, Ø 25 mm) were exposed for 2 min (blank control samples). All filters were visually checked for microplastic contamination with a microscope after exposure. During the digestion and filtration steps, for every two spiral valves, blank samples were included for both spiral valve content and tissue analyses. The concentrations of microplastics in spiral valve content and tissue were corrected for the average value of the blank samples. The Limit of Detection (LoD) was defined as $3 \times \text{SD}$ of the blank results. The Limit of Quantification (LoQ) is defined as $3.3 \times \text{LoD}$ or $10 \times \text{SD}$ of the blank results. The LoD and LoQ are reported for the spiral valve content and spiral valve tissue separately (Supplementary Table SC), they give an indication of the level of significance of the results (LOW < LoD, MEDIUM < LoQ and HIGH > LoQ).

**Identification and Classification**

In this study, microplastics were defined as plastic items between 10 µm and 5 mm in size. Based on the anatomy of the digestive tract of the porbeagle shark and their potential to regurgitate larger items, it was expected that items larger than 5 mm were not present in the spiral valve. All filters were analyzed under a light microscope (Olympus CX31 – 4x) to evaluate the performance of the digestion and the presence of microplastics. Following recommendations of Ferreira (Ferreira, 2014), microplastics were categorized according their physical characteristics: size, shape and color. Plastic particles were categorized in size bins of approximately 100 µm (20–100, 101–200, etc.) and were measured using MicroCamLab for Microsoft Windows. Their shape was described as: fragments, pellets (spheres), fibers, plastic films, foamed plastic and granulates. A needle was used to carefully turn the particle and to help identify the shape when there was uncertainty. Results of the microscopic analysis were reported as number of microplastic particles per g w.w.

**Validation**

Raman spectroscopy (Renishaw® inVia Raman Microscope) was used to determine the polymer type (Koenig, 1999). The results of this technique were compared with existing references of (virgin) polymers and non-polymer particles. Only the distinguishable items, clearly visible under the microscope (>80 µm), were selected for the Raman analyses. We did not analyze the particles below 80 µm with Raman spectroscopy, because they proved extremely difficult to transfer from the GF/F to the edge filters for Raman spectroscopy without introducing contamination. In addition, the blue nitrile gloves and blue plastic from the filter packaging was analyzed with Raman to examine their potential role in contamination. Items were located with a magnification lens of 20×, illuminated with a red (785 nm) or green (532) laser at an intensity laser power from 0.5 up to 20%, and imaged with a Philips SP1030NC webcam.
TABLE 1 | Overview of measurements during sampling and sample collection.

| Fish no. | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
|----------|---|---|---|---|---|---|---|---|---|----|
| Sex      | M | F | M | M | F | F | M | F | M | F  |
| Maturity | C | A | B | B | A | A | A | C | A | C  |
| Total length (cm) | 218 | 170 | 144 | 113 | 139 | 119 | 183 | 194 | 221.4 | 216 |
| Total body weight (kg) | 74.2 | 43.1 | 24.1 | 11.2 | 22.6 | 15 | 48.2 | 65.8 | 84.5 | 77.6 |
| Liver weight (kg) | 6.376 | 5.382 | 2.156 | 1.35 | 1.89 | 2.158 | 2.888 | 6.852 | 10.42 | 8.15 |
| Weight full spiral valve (g) | 865.6 | 626.6 | 376.5 | 220.6 | 403.3 | 270.3 | 974.2 | 675.7 | 871.9 | 785.2 |
| Wet weight spiral valve content (g) | 370.2 | 240.7 | 128.2 | 89.1 | 167.6 | 120.2 | 424 | 254.9 | 278.7 | 278.7 |
| Wet weight spiral valve tissue (g) | 495.4 | 385.9 | 248.3 | 131.5 | 235.7 | 150.1 | 550.2 | 420.8 | 598.3 | 506.5 |
| Parasites (Y/N) | Yes** | No | Yes* | No | Yes** | Yes* | No | No | Yes** | No  |
| Tissue damage (Y/N) | No | No | No | No | No | No | Yes | No | No | No  |
| Fullness index | 49.89 | 55.85 | 53.2 | 79.55 | 74.15 | 80.14 | 91.76 | 38.74 | 32.38 | 35.91 |
| Condition index | 0.716 | 0.877 | 0.807 | 0.776 | 0.842 | 0.89 | 0.754 | 0.901 | 0.779 | 0.77 |
| Hepatosomatic index | 8.593 | 12.49 | 8.948 | 12.05 | 8.363 | 14.39 | 6.251 | 10.41 | 10.41 | 10.5 |

Maturity is defined as: A: immature, B: maturing, C: mature. All weights are in g, except for total body weight (kg). Parasites: ** multiple parasites found, *only one parasite found. Fullness Index: (weight of stomach contents * 10.000)/weight of fish). Condition Index: K = 100 × (fish body weight/total length)². Hepatosomatic Index: IH = Liver weight/Body weight × 100.

Statistical Correlation Between Ingested Microplastics and General Health Indices

Plastic ingestion was defined as the average plastic particle per g w.w. content and the average plastic particle per g w.w. tissue of spiral valve. The general health condition of the porbeagle shark is determined by the Condition Index [K = fish body weight (g)/length (cm)² × 100] (Goede and Barton, 1990). In addition, the Hepatosomatic Index [IH = liver weight (kg)/body weight (kg) × 100] (Goede and Barton, 1990), also an indication of the status of energy reserve, was used as an additional general health estimate (Goede and Barton, 1990). Higher numbers for both variables indicate a good fitness condition and general health. Correlations were controlled for the Fullness Index [IF = weight of spiral valve contents (kg)/weight of fish (kg) × 10,000], because it is hypothesized that a fuller spiral valve results in a higher amount of plastic. Simple statistics were performed in Microsoft Excel 2010. The correlation analyses were performed with IBM SPSS Statistics 21 and statistical significance was set at p < 0.05. To determine the correlation between plastic ingestion and the Condition Index of the porbeagle shark, normal distribution was tested with Shapiro-Wilk test and Spearman’s rho test was used on blank subtracted results from the microplastic analyses. To determine the correlation between plastic ingestion and the Hepatosomatic Index of the porbeagle shark, a Pearson correlation test was performed. To control the Fullness Index, only partial correlation tests were performed. By holding this third variable constant, the influence of the fullness of the spiral valve could be eliminated.

RESULTS

Sampling and Study Design

Overall, the spiral valves and their contents looked similar. The dark brown/reddish colored content was a thick sticky substance and contained no large items (>5 mm). However, some contained small gray unidentified fragments which may have been remains of bones. Parasites were found in 50% of the spiral valves (Supplementary Table SA). They were assigned, with some uncertainty, to one tapeworm species, Dinobothrium septaria and one unidentified nematode species. The tapeworms were approximately 8 cm long and the nematodes were approximately 3 cm long (Supplementary Figure SE). In addition, parts of arthropod exoskeleton and appendages of different organisms were observed during microscope inspection (Supplementary Figure SF). An overview of the collected measurements during the sampling and during the sample collection is given in Table 1. Condition Index and Hepatosomatic Index, together with other general health indicators are included.

Method Development

Sample Collection

Four spiral valves were used to test four different procedures (Nock and Caira, 1988; Dailey and Vogelbein, 1990; Curran and Caira, 1995; Henderson et al., 2002; Alarcos et al., 2006; Beveridge and Justine, 2007) for the collection of spiral valve content. This included hanging of the spiral valve and collecting the content via sieves (1) or massaging into a glass bowl (2), scraping the content out with a metal spoon (3) or washing the content out with milliQ water (4) (Figure 3). Although most of the content could be collected with the use of milliQ water, one of the main issues with the use of milliQ water is that the w.w. of the content cannot be determined. An additional problem was that some water was residing in the spiral valve. Therefore, it seemed inappropriate to flush the spiral valve with milliQ water and wash the content out of the spiral valve. Multiple problems occurred when using the sieves. Most of the content stuck to the sieves and clogged the mesh. This made it impossible to collect the content from the sieves and separate fractions based on size. To remove the content from the sieves, large volumes of hydrogen...
peroxide (30%) were used and (wet) weight of the spiral valve content could again not be measured. The hanging of the spiral valve seemed the most effective, part of the content was collected passively using gravity. In addition, actively squeezing aided to retrieve the remaining content. Cutting open and collecting content by scraping or washing the spiral valve seemed to be ineffective because of the potential issues it caused in relation to content loss or contamination. Therefore, it was decided that the remaining spiral valves would be by hung of a tripod and massaging would be applied to retrieve the content. The glass bowl was replaced with a glass bottle with a smaller opening to decrease the risk of air borne contamination. The lower part of the spiral valve was placed in the glass bottle to prevent spilling of the content. An overview is shown in Figure 4. After collecting the spiral valve content, tissue samples were taken from the empty spiral valve. Any content that could still be retrieved was added to the already collected content with a metal spoon. The blank filters were analyzed with a microscope after the exposure. No microplastics were detected. Therefore, it was assumed that airborne contamination in the fume hood was of low concern while sampling and weighing spiral valve content and tissue.

Digestion and Validation

To extract microplastics from spiral valve content and tissue samples, different digestion methods, sample sizes and time frames were explored and compared. These included enzymatic digestion, acid digestion and alkaline digestion techniques (Supplementary Figure S1.1). Their performance was determined by how well the sample could consequently be filtered through a glass fiber filter and how much residual content was observed by visual and microscopic inspection after filtration. The results are summarized below, more methodological details and pictures can be found in Supplementary Figures S1.1, S1.2. The filters using the acid digestions were covered with digest residue and could therefore cover possibly microplastics (Supplementary Figure S1.2). The enzymatic digestion was an effective method for small samples, but inappropriate for larger samples, such as spiral valve content and tissue, due to the high costs of Proteinase K. The 10% KOH solution seemed to perform best with a spiral valve content sample of 2 g and a tissue sample of 1 g, leaving a minimal digest residue on the filter (Figure 6). Some filters were stained brownish and/or contained gray fragments, most likely dietary remains. The gray fragments were not considered a problem during the identification as they could be crushed into a powder with a needle, making them distinguishable from plastic. The filters were increasingly covered when using larger sample sizes. Still, larger volumes or even better, a large series of smaller replicate samples would cover when using larger sample sizes. Still, larger volumes or even better, a large series of smaller replicate samples would be a clear improvement with longer digestion times. The results of the KOH digestion differed, however, between spiral valves. Most likely due to different diets and stages of digestion at the time of death. The composition of the remaining spiral valves was unknown, therefore, it was decided to digest the content samples of the next spiral valves slowly over a period of several days until contents were, homogenous and fluid, ready for filtration (Figure 7).

Almost all tested particles (8 types of plastic, 12 particles each) could be recovered with the KOH content (94.8%) and tissue (98%) method. Unrecovered particles were attributed to the rinsing procedure (transparent plastics stuck on the glassware wall) and not to the digestion method. In addition, the spiked PVC particles were difficult to count and to filter as a result of their small size. No apparent changes in color and shape of the spiked plastics occurred as a result of the KOH digestion (Supplementary Figure SC). Also, no particles were stuck together. In addition, the use of 10% KOH and extended digestion times (>24 h) was further validated on 100 spiked particles, by assessing recovery and weight disparities. The recovery for all plastics was 100%, except for PET (99%) due to the loss of one PET particle before weighing. The weight of the plastics increased at first due to the dried KOH residue on the particles. After a milliQ rinsing step, the average weight of the particles did not differ significantly (Supporting Information) from the weight before the digestion. Only the mean differences in weight for PA at all 3 exposures and PVC after 9 days exposure were significantly higher after the KOH digestion. The average weight difference of the spiked plastics before and after 10% KOH digestion together with the significance levels is shown in Supplementary Tables S6.1–S6.4).

Filtration

Filters that were rinsed by covering them with a layer of 2.5 mL solution of 20% Extragen appeared visually cleaner than the ones which were only rinsed with milliQ water. Therefore, the use of the alkaline soap (prefiltered) was added to the filtration procedure. The difference between simply milliQ rinsed filters and milliQ with soap rinsed filters tested on 5 and 2 g samples can be seen in Supplementary Figure SD.

Microplastic Analysis

Quality Control

The blank measurements showed contamination of mostly fibers and blue fragments but were not consistent. The average number of plastic particles in the content blanks was 6.6 ± 6.5 particles and for the tissue 1 ± 0.7. Supplementary Figure SG illustrates also the range in number of particles per blank sample.

For spiral valve content, the LoD (3 × SD of the average plastic particle in the blanks) was 19.6 and the LoQ (3.3 × LoD) was 64.8. For the spiral valve tissue, the LoD was 2.1 and the LoQ was 7.0. Supplementary Table SC shows the LoD and LoQ described for 1 g and includes blue fragments, fibers, black fragments and other plastics.

Concentration of Plastic Particles

A total of 878 plastic particles were identified by visual identification. These were identified as fragments (65.9%), fibers (32.9%), pellets (0.9%), and films (0.5%). The most abundant colors were blue (44.8%), black (23.7%), red (9.6%), and transparent (6.5%). Other colors that were present were orange, brown, green, gray, yellow, purple, white, pink, or multicolored (all < 5%). Almost all particles were below 100 μm
(fibers excluded) and the largest plastic particle identified was approximately 930 µm long (Supplementary Figure SH). An overview of the average number of blue fragments, fibers, black fragments, other and total plastic particles is given in Table 2 and Figure 8. It also includes the standard deviation (SD) per spiral valve content and tissue, before and after correction with the blank values (adjusted per type). One spiral valve (1) was not included in the analysis due to the presence of large amounts of sand particles in its content which interfered with the detection of microplastics. Instead of analyzing the entire sample, the total amount of microplastics per spiral valve was recalculated by multiplying the subsample concentrations with total weight of content and tissue and adding both together (Table 3). In the content of the rectal gland (1.2 g) three blue fibers were found, but after blank subtraction, no plastics can be reported. Also, nine parasitic nematodes were examined, but no plastic particles were discovered within. A raw data file is presented in Supporting Information.

**Validation**

After the identification of all isolated particles by microscope, Raman spectroscopy was used as an additional tool to identify...
### TABLE 2

The number of plastic particles in spiral content and tissue, expressed as average* number of plastic particles per g w.w. before and after blank correction (per type); SD, Standard Deviation; NA, Not Available; *5 content and 4 tissue replicates per spiral valve.

| Spiral valve | Blank correction | Particles | 1   | 2    | 3    | 4    | 5    | 6    | 7    | 8    | 9    | 10   | Average | Average | SD    | Average | Average | SD    | Average | Average | SD    | Average | Average | SD    | Average | Average | SD    |
|--------------|------------------|-----------|-----|------|------|------|------|------|------|------|------|------|--------|---------|-------|---------|---------|-------|---------|---------|-------|---------|---------|-------|
|              |                  |           |     |      |      |      |      |      |      |      |      |      |        |         |       |         |         |       |         |         |       |         |         |       |         |         |       |
| Content      | Before Blue      | NA        | 7.9 | 2.1  | 0.5  | 0.5  | 9.4  | 3.8  | 0.2  | 0.3  | 3.4  | 1.2  | 0.8   | 0.4    | 1.1   | 1.0    | 0.5    | 0.6   | 1.0    | 0.5    | 1.0   |
|              | Fiber Blue       | NA        | 2.1 | 1.1  | 0.5  | 0.4  | 1.3  | 0.8  | 1.3  | 0.6  | 3.7  | 3.0  | 3.0   | 2.3    | 2.4   | 1.6    | 1.9    | 0.9   | 1.4    | 0.7    | 1.4   |
|              | Black Blue       | NA        | 2.0 | 2.0  | 0.1  | 0.2  | 0.3  | 0.3  | 3.3  | 1.4  | 3.3  | 1.2  | 2.1   | 1.9    | 1.1   | 1.0    | 0.3    | 0.4   | 0.7    | 0.3    | 0.7   |
|              | Other Blue       | NA        | 1.4 | 0.7  | 0.6  | 0.9  | 1.8  | 0.8  | 0.7  | 1.1  | 3.2  | 1.5  | 0.5   | 0.9    | 0.1   | 0.2    | 0.8    | 0.8   | 0.3    | 0.3    | 0.3   |
|              | Total Blue       | NA        | 13.4| 3.6  | 1.7  | 1.0  | 12.8 | 3.9  | 5.5  | 2.6  | 13.6 | 4.1  | 6.4   | 3.8    | 4.7   | 2.9    | 3.5    | 1.9   | 3.4    | 1.1    | 3.4   |
|              | After (per type) | Blue      | NA  | 6.4  | 2.1  | 0.0  | 0.0  | 7.9  | 3.8  | 0.0  | 0.0  | 1.9  | 1.2  | 0.0   | 0.0    | 0.2   | 0.4    | 0.0    | 0.0   | 0.0    | 0.0    | 0.0   |
|              | Fiber Blue       | NA        | 0.8 | 0.8  | 0.0  | 0.0  | 0.2  | 0.4  | 0.1  | 0.2  | 2.2  | 3.0  | 1.9   | 1.8    | 1.0   | 1.5    | 0.6    | 0.5   | 0.2    | 0.4    | 0.2   |
|              | Black Blue       | NA        | 1.8 | 1.9  | 0.0  | 0.1  | 0.1  | 0.1  | 3.0  | 1.4  | 3.0  | 1.2  | 1.9   | 1.8    | 0.9   | 0.9    | 0.2    | 0.3   | 0.4    | 0.3    | 0.3   |
|              | Other Blue       | NA        | 1.4 | 0.7  | 0.6  | 0.9  | 1.8  | 0.8  | 0.7  | 1.1  | 3.2  | 1.5  | 0.5   | 0.9    | 0.1   | 0.2    | 0.8    | 0.8   | 0.3    | 0.3    | 0.3   |
|              | Total Blue       | NA        | 10.4| 3.5  | 0.6  | 0.9  | 10.0 | 4.0  | 3.8  | 2.3  | 10.3 | 4.1  | 4.3   | 3.0    | 2.2   | 2.2    | 1.6    | 1.0   | 0.9    | 0.3    | 0.3   |
| Tissue       | Before Blue      | NA        | 1.8 | 1.3  | 5.3  | 2.8  | 1.0  | 0.8  | 1.3  | 1.9  | 0.0  | 0.0  | 0.3   | 0.5    | 2.0   | 2.2    | 1.5    | 0.6   | 2.0    | 1.8    | 2.0   |
|              | Fiber Blue       | NA        | 2.8 | 1.3  | 2.0  | 2.4  | 2.5  | 1.7  | 0.8  | 1.0  | 5.5  | 4.4  | 3.3   | 1.7    | 3.5   | 0.6    | 5.3    | 2.6   | 3.0    | 2.2    | 3.0   |
|              | Black Blue       | NA        | 0.5 | 0.6  | 0.5  | 0.6  | 1.0  | 1.4  | 0.0  | 0.0  | 0.0  | 0.0  | 0.3   | 0.5    | 1.0   | 0.8    | 2.5    | 1.9   | 0.0    | 0.0    | 0.0   |
|              | Other Blue       | NA        | 0.5 | 0.6  | 0.3  | 0.5  | 0.5  | 1.0  | 0.3  | 0.5  | 2.5  | 2.1  | 0.8   | 1.0    | 1.0   | 0.0    | 1.3    | 1.3   | 1.0    | 1.4    | 1.0   |
|              | Total Blue       | NA        | 5.5 | 2.6  | 8.0  | 4.5  | 5.0  | 1.8  | 2.3  | 1.7  | 8.0  | 5.1  | 4.5   | 0.6    | 7.5   | 2.6    | 10.5   | 3.4   | 6.0    | 2.2    | 6.0   |
|              | After (per type) | Blue      | NA  | 1.8  | 1.3  | 5.3  | 2.8  | 1.0  | 0.8  | 1.3  | 1.9  | 0.0  | 0.0  | 0.3   | 0.5    | 2.0   | 2.2    | 1.5    | 0.6   | 2.0    | 1.8    | 2.0   |
|              | Fiber Blue       | NA        | 1.8 | 1.3  | 1.5  | 1.9  | 1.5  | 1.7  | 0.3  | 0.5  | 4.5  | 4.4  | 2.3   | 1.7    | 2.5   | 0.6    | 4.3    | 2.6   | 2.0    | 2.2    | 2.0   |
|              | Black Blue       | NA        | 0.5 | 0.6  | 0.5  | 0.6  | 1.0  | 1.4  | 0.0  | 0.0  | 0.0  | 0.0  | 0.3   | 0.5    | 1.0   | 0.8    | 2.5    | 1.9   | 0.0    | 0.0    | 0.0   |
|              | Other Blue       | NA        | 0.5 | 0.6  | 0.3  | 0.5  | 0.5  | 1.0  | 0.3  | 0.5  | 2.5  | 2.1  | 0.8   | 1.0    | 1.0   | 0.0    | 1.3    | 1.3   | 1.0    | 1.4    | 1.0   |
|              | Total Blue       | NA        | 4.5 | 2.6  | 7.5  | 4.0  | 4.0  | 1.8  | 1.8  | 1.7  | 7.0  | 5.1  | 3.5   | 0.6    | 6.5   | 2.6    | 9.5    | 3.4   | 5.0    | 2.2    | 5.0   |
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FIGURE 8 | Average number of plastic particles per g w.w. measured in spiral valve content and tissue after blank correction.

polymer types. There were on average 4.9 ± 2.4 particles per g w.w. content and 5.5 ± 2.7 particles per g w.w. tissue. From all filters, a total of 24 particles, at least one particle from each spiral valve, between 84 and 572 µm were examined by Raman spectroscopy for polymer identification purposes by cataloging a collection of fingerprints. These included the five main observed types of microplastics: a black fiber and transparent, black, red, and blue fragments. Pigment spectra and environmentally driven changes to surface properties, such as weathering and biofouling, hinder spectroscopic identification (Harrison et al., 2014; Beltrán-Sanahuja et al., 2020). The process of degradation, autoxidation of hydrocarbon polymers, entails the formation of novel oxygen-containing groups in the main chain through a series of primary and secondary radical reactions that involve chain scissions and cross-linking of polymer backbone, the formation of polar carbonyls (C = O) and vinyl (CH₂ = CH) groups, and finally, changes in the conformation and crystallinity of the polymer (Gómez et al., 2019).

The spectra of the items showed typical characteristics of weathered synthetic polymeric material, but none can be assigned with complete confidence. The water solvent was easily identified at about 1640 cm⁻¹ and used as an internal intensity standard. Although the spectra were not clear enough for full identification, significant signals were present in the OH/NH stretch region between 3000 and 3700 cm⁻¹. These signals were weaker relative to the C-H stretches than would typically be seen for natural carbohydrate-based polymers such as cellulose, carrageenan or chitin, or for proteinaceous material, suggesting that they are indeed synthetic. Although difficult to be certain, the transparent particles showed characteristic features of PET: the signals around 1270 and 1450 cm⁻¹, and 1710 cm⁻¹moments in the spectra of the remaining fragments suggest these might be polyethylene. In addition, four blue fragments from the samples (Supplementary Figure SI) were compared with the spectra of the blue nitrile gloves and the blue plastic from the filter packaging. The blue fragments in the samples gave a clear spectrum, comparable to the spectrum of the blue packaging. Since all blue fragments resulted in the same spectrum, it was concluded that this was not due to contamination: not the polymer spectra, but pigment spectra were obtained. Supplementary Figure SI illustrates the similarities in spectra.

Statistical Correlation Between Ingested Microplastics and General Health Indices

The correlation between the average plastic particle per g w.w. content (4.9 ± 2.4) and w.w. tissue (5.5 ± 2.7) and the general health indices of porbeagle sharks were examined by comparing the corrected average microplastic concentrations with the Condition Index (0.82 ± 0.06) and Hepatosomatic Index (10.6 ± 2.5). Spearman's rho correlation showed no statistically significant correlation between the average particle per g w.w. content and the Condition Index (r = -0.008, p = 0.983) and the Hepatosomatic Index (r = 0.165, p = 0.651). Pearson correlation tests showed no statistically significant correlation between the average plastic particle per g w.w. tissue and the Condition Index (r = 0.167, p = 0.668). When controlled for the Fullness Index with partial correlation no significant correlation between the average plastic particle per g w.w. tissue and the Condition Index was found (r = 0.151, p = 0.722). Both tests were also performed for the Hepatosomatic Index and again no statistically significant correlation was observed (Pearson: r = 0.597, p = 0.09 and Partial: r = 0.582, p = 0.130). While no statistically significant correlation was found here due to the significance level set at 0.05, the p-value
DISCUSSION AND CONCLUSION

A large part of the study focused on method development as standard techniques for microplastic analysis in large top predator sharks are missing. In a microplastic study using Blue sharks, stomachs were opened and the contents were washed through a 1 mm metal sieve with pre-filtered water (Bernardini et al., 2018). In this study, spiral valve content and tissue were used, representing digestion compared to ingestion. Studies investigating tapeworms or parasites presence in sharks, cut open the spiral valves and remove parts of the lumen tissue (Curran and Caira, 1995; Alarcos et al., 2006) or the contents via suspension, washing or shaking (Henderson et al., 2002; Randhawa and Brickle, 2011). Different methodologies to extract spiral valve content and tissue were explored and the most appropriate method, a combination of gravity and massaging, was applied on the remaining spiral valves. Several studies report microplastic ingestion based on the examination of the whole or substantial parts of the digestive tract (Foekema et al., 2013; Lusher et al., 2013; Phillips and Bonner, 2015; Bernardini et al., 2018; Sampaio et al., 2018; Smith, 2018). Most laboratory equipment was not readily available for the rather large volumes of porbeagle spiral valve content we encountered (e.g., one porbeagle spiral valve content measured 424 g w.w.). Hence, only subsamples of the spiral valve content were analyzed.

This study supports the use of a 10% KOH solution to digest spiral valve content and tissue. The (lipid) content of the spiral valves left fatty residues on the filter after digestion. To improve results, the filters were washed with alkaline soap to decrease the amount of residue. The success of this mixture aligns with previous findings which suggest that a combined alkaline digestion with KOH and NaClO are useful compounds when digesting biota and/or gastrointestinal content for microplastic analysis (Cole et al., 2014; Strand and Tairova, 2016; Enders et al., 2017). This method, using a post soap wash step, proved to be safe, cost effective, with less procedural steps, without affecting any of our spiked microplastics. Although good results were obtained with the alkaline soap, it was decided to harmonize procedures by adopting the combined KOH:NaClO digestion for future biota analysis in our laboratory.

To exclude non-plastic particles during the microscopic analysis, a step to determine the consistency of the larger particles (>100 µm) was added. Fish bone remains were easily distinguished from plastics by carefully touching the item with a needle. On applying pressure, the remains of prey broke down into a powder following alkaline digestion (Kaye et al., 2004).
One spiral valve contained a large amount of sand, making visual microplastic analysis impossible, although this was not an issue in the other spiral valves, it would be useful to introduce a density separation and simplify microplastic detection (e.g., Nile red; Maes et al., 2017a) by introducing additional steps after the digestion procedure. This shows that while certain steps of the protocol can be harmonized (e.g., KOH digestion and microplastic quantification), small adaptations might be required to make protocols species or case specific.

Although no values were reported for spiral valve 1, due to the presence of sand, plastics particles were detected in both the content and tissue of all spiral valves. Fibers, blue fragments and black fragments were most prevalent. Remarkably, all spiral valves and some of the blanks contained blue fragments. It was suspected that these fragments were a result of contamination due to the high numbers of particles with this specific combination of shape and color. Although Raman spectroscopy was applied, there is still some uncertainty in terms of the origin of these fragments. At the moment, certified reference materials are unavailable and polymer identification with spectroscopy is fraught with challenges, especially in this small size range of weathered and pigmented particles detection (Lenz et al., 2015; Araujo et al., 2018). The Raman spectroscopy picked up the spectra of the blue pigments more easily than polymer spectra and confirmed that the blue pigment in the packaging of the filters was comparable with some blue fragments in the samples. This could be due to resonance enhancement, where signals from the pigment are enhanced, but signals from the polymer are hidden. The pigment is known as copper phthalocyanine and is used in multiple applications. Several studies mentioned blue plastic fragments in environmental samples and attribute the Raman spectra to this pigment (Van Cauwenberghhe et al., 2013, 2015; Karlsson, 2014; Van Cauwenberghhe and Janssen, 2014) which makes source tracking rather difficult. It indicates that non-plastic materials, including filter packaging, should be used, wherever possible, to minimize contamination and/or when in doubt, screening for plastic contamination should take place prior to usage. Extra precautions should be sought to further lower the blank values. Only a small portion of samples were higher than the LoQ. This may be partly due to the low number of blanks used. It is therefore recommended to increase the number of procedural blanks and to repeat this over time as contamination might be day dependent.

The high numbers of microplastics in the spiral valves imply that plastic is ingested and digested by porbeagle sharks. In 4 out of 9 spiral valves, more plastic particles per g w.w. were found in tissue samples than in spiral valve content. This indicates that taking samples from solely the content might not represent the actual amount of microplastics in the spiral valve. Moreover, the results of the lipid content and tissue analysis demonstrate that the sample collection via the massaging method might not be enough to collect all content (Supporting Information). This was already observed during the tissue sample collection, small amounts of spiral valve content were stuck to the surface of the tissue. Therefore, it is uncertain whether and how many microplastics in the tissue samples were resulting from the small amounts of content in the tissue samples or if the microplastics were absorbed into the 3D structured lining of the tissue where nutrients are absorbed into the shark’s body. Eventually, this might result in adverse effects on the shark’s health if other organs and tissues are exposed (Browne et al., 2008; Wright et al., 2013; GESAMP, 2015a; Panti et al., 2016). Due to this uncertainty both spiral valve content and tissue should be analyzed and reported jointly for monitoring purposes. Parasites were present within the spiral valves but contained no plastic particles. This could be due to visual cut off point of 10 μm. The collected parasites might ingest particles much smaller than the particle size examined in this study. In addition, the parasites of only one spiral valve were examined and therefore, no conclusions about microplastic ingestion in parasites can be drawn.

To examine if the porbeagle shark’s health was adversely affected by microplastic ingestion, we looked for a correlation between the average plastic particle per g w.w. in spiral valves content/tissue and several general health indices. In this case, the Hepatosomatic Index seemed to be the most relevant health indicator in relation to microplastic ingestion, maybe because it relates more directly to toxic effects (Scacco et al., 2010). Some microplastics will be excreted together with the rest of the spiral valve content. A portion gets stuck within the tissue and could potentially exchange chemicals (Koelmans et al., 2013, 2014) during their extended stay in the digestive tract. Previous studies reported chemical concentration levels in the liver (Bendall et al., 2014) and muscles (Nicolaus et al., 2016) of porbeagles and concluded that current levels were mostly low or undetectable. To preserve these and other vulnerable marine top predator species, examining both microplastics and concentrations of toxic chemicals adsorbed or leaching from microplastics and in specimens itself would be recommended to provide more insights in the toxicity of microplastic particles and associated chemical equilibriums (Koelmans et al., 2013). Moreover, additional ecosystem variables should be considered, such as the presence of parasites and a range of environmental stressors. Although no statistically significant correlation between the average plastic particle per g w.w. tissue and Hepatosomatic Index was observed, the correlation was moderately positive. An explanation could be that liver weight increases due to pathological changes (Scacco et al., 2010) as a result of the increased residence of microplastics in the gut, causing a higher Hepatosomatic Index. All other correlations were not statistically significant (all p > 0.05). This could be due to the small sample size in the analyses (n = 9). However, it is likely that the found concentrations of microplastics in the spiral valve of the porbeagles did not cause effects measurable by general health condition indexes such as the Hepatosomatic Index. Similarly, some fish studies reported no statistical significant relation between plastic ingestion and their condition (Foekema et al., 2013; Zoeter Vanpoucke, 2015).

The presence of microplastics in porbeagle sharks is most likely the result of the contamination of their food supply (Foekema et al., 2013; Lusher et al., 2013) (indirect) and/or internal fragmentation of the larger plastics they ingested (Cliff et al., 2002) (direct). DNA analysis did not allow us to ascertain
what the porbeagle sharks diet consisted of and whether prey was the source of microplastics (Supporting Information). Earlier studies indicated potential accumulation and trophic transfer of microplastics across parts of the foodweb (Farrell and Nelson, 2013; Setälä et al., 2014; Lusher et al., 2015a). Studies looking at microplastics in fish from the North Atlantic region report around one microplastic per fish (Foekema et al., 2013; Lusher et al., 2013; GESAMP, 2015b). When recalculating the total amounts of microplastics in the spiral valves of porbeagle sharks, rather large numbers were observed (Table 3). Concentration went up as high as 6116 microplastics per individual spiral valve, suggesting that microplastics are accumulating across the food web, potentially harming critically endangered North East Atlantic porbeagle sharks. The present work emphasizes the potential for top predator application in microplastic monitoring. Porbeagle sharks are apex predators feeding on a wide range of organisms, including teleosts and cephalopods (Ellis and Shackley, 1995; Joyce et al., 2002; Francis et al., 2009) and play a key role in controlling ecosystem dynamics (Francis et al., 2009). Distances of over 1,000 nautical miles (nm) were recorded by tagged porbeagle sharks, though over 90% of the 143 sharks tagged moved less than 500 nm from their original tagging location (Campana and Gibson, 2008; Pade et al., 2009). Although one porbeagle has been recorded crossing the Atlantic (Francis et al., 2009), tagged sharks in the Celtic Sea mostly remained in that area (Pade et al., 2009). The spatial distribution of incidental porbeagle bycatch reported by the participating vessels confirm that porbeagles are widespread within the Celtic Sea (Ellis and Bendall, 2015). There may be a separate North Atlantic stock off Iceland (Matsumoto, 2006), this North-East Atlantic stock is generally considered to be distinct from those in the North-West Atlantic and Mediterranean (ICES and WGEF, 2007; FAO, 2010). As such, porbeagle sharks could be an ideal species for integrated monitoring across a wider (sub)region. In future, it would be recommended to target spiral valve content in by-caught Elasmobranches, such as the porbeagle shark, for microplastics monitoring purposes. This avoids potential bias from gastric evacuation (Eggers, 1977; Bromley, 1994; Cortés, 1997; Bucking, 2015) and might give a better link to probable impacts and pollutant loads as it relates to digestion (Bucking, 2015). To overcome temporal changes in bycatch rates of porbeagle sharks in gillnets (Bendall et al., 2012; McCully et al., 2013; Ellis and Bendall, 2015) and to support an appropriate microplastics monitoring program, the analysis should be expanded to other by-caught top predator species.

DATA AVAILABILITY STATEMENT

All datasets generated for this study are included in the article/Supplementary Material.

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ETHICS STATEMENT

Ethical review and approval was not required for the animal study because the porbeagle sharks were obtained dead, captured as seasonal by-catch in gillnet fisheries, and processed for research purposes.

AUTHOR CONTRIBUTIONS

TM: setup and organization of study and sampling protocols, spiral valves collection, preservation and transport, design microplastic analysis protocols and layout, overview student work, manuscript writing and editing, and manuscript submission. JD: sample analysis and processing, data analysis, reporting, and report writing. AV: analysis protocols, student supervision, and data interpretation. MD: sample collection, dissections, porbeagle measurements. VB: porbeagle collection and organization of dissections. MV: student support, microplastic analysis, and lab supervision. HL: student support, data interpretation, and report writing.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2020.00273/full#supplementary-material
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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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