Microplastic Distribution at Different Sediment Depths in an Urban Estuary

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As plastic production increases, so do the threats from plastic pollution. Microplastics (defined as plastics <5 mm) are a subset of marine debris about which we know less than we do of larger debris items, though they are potentially ubiquitous in the marine environment. To quantify the distribution and change in microplastic densities through time, we sampled sediment cores from an estuary in Tasmania, Australia. We hypothesized that the type, distribution and abundance of microplastics observed would be associated with increasing plastic production, coastal population growth, and proximity to urban water outflows and local hydrodynamics. Sediments ranging from the year 1744 to 2004 were sub-sampled from each core. We observed microplastics in every sample, with greater plastic frequencies found in the upper (more recent) sediments. This time trend of microplastic accumulation matched that of global plastic production and coastal population growth. We observed that fibers were the most abundant type of microplastic in our samples. These fibers were present in sediments that settled prior to the presence of plastics in the environment. We propose a simple statistical model to estimate the level of contamination in our samples. We suggest that the current trend in the literature suggesting very high loads of fibers, particularly in remote locations such as the deep seafloor, may be largely due to contamination.

Keywords: core samples, estuary, microplastics, plastic fibers, sediment

INTRODUCTION

Microplastics (plastics <5 mm, Arthur et al., 2008) have been observed throughout bottom sediments of marine and freshwater environments, including sediments in rivers (Casta-eda et al., 2014), estuaries (Thompson et al., 2004; Sruthy and Ramasamy, 2017), lagoons (Vianello et al., 2013), lakes (Corcoran et al., 2015), seas (Zobkov and Esiukova, 2017), and deep sea trenches (Van Cauwenberghe et al., 2013; Woodall et al., 2014; Fischer et al., 2015). Microplastics are commonly found in the environment in three forms; fragments which form from mechanical and biological fragmentation of larger plastic items (ter Halle et al., 2016), microbeads which are manufactured as abrasives in cosmetics and air-blasting (Fendall and Sewell, 2009; Mason et al., 2016), and microfibers from sources such as synthetic fabrics and ropes (Browne et al., 2011). Studies have shown multiple damaging effects of microplastics in the environment, including adsorption of toxic organic contaminants (Endo et al., 2005; Teuten et al., 2007; Rochman et al., 2013), ingestion by animals with implications for human consumption (Van Cauwenberghie and Janssen, 2014; Rochman et al., 2015) and changing the heat transfer and water movement of sediment (Carson et al., 2011).
Microplastics have been observed in benthic environments since the late 1970s (Gregory, 1977; Shiber, 1979) and sediments are suggested to be a long-term sink for microplastics (Moré-Ferguson et al., 2010; Cózar et al., 2014). Benthos is an important feeding environment for many marine species (Anderson and Lovvorn, 2008; MacDonald et al., 2012; Gittman and Keller, 2013) and a recent study has shown marine benthic species ingest microplastics (Courtene-Jones et al., 2017). Knowledge of the harmful effects of microplastics on benthic marine species and communities is growing (Green, 2016; Galloway et al., 2017), however, little is known of past microplastic accumulation in benthic environments.

Several studies have examined microplastics in sediments (Claessens et al., 2011; Corcoran et al., 2015; Klein et al., 2015). For example, Claessens et al. (2011) quantified the number of microplastics in 16 year old sediments collected at the intertidal and high water mark. However, there have been no evaluations of microplastic contamination in deep sediments with known age cores, allowing analysis of deposition rates and concentrations. To address this knowledge gap, we compared sediment cores taken from an urbanized estuary and asked the following questions: Do microplastic densities change in time (with known age of sediment cores)? If microplastics are present, do we detect different densities as we move further from areas of higher human population densities?

**MATERIALS AND METHODS**

**Sampling Location**

Core samples were taken from Elwick Bay and Dogshear Point in the Derwent Estuary, Tasmania, Australia (Figure 1). The area is classed as urban/light industrial and the shoreline is partially reclaimed land. Surrounding land use includes residential areas, a horse race course and stables, low lying recreational areas, a reclaimed land. Surrounding land use includes residential areas, a horse race course and stables, low lying recreational areas, a reclaimed land.

Cores were collected in November 2004 as outlined in Townsend and Seen (2012). Core A was taken from the middle region of Elwick Bay around 3 km upstream of a zinc refinery (Townsend and Seen, 2012). Core B was taken off Dogshear Point, 3 km north-east of core A. Each core was sliced into 2 cm sections and stored in sealed containers. Core samples were stored refrigerated upright until analysis. A total of seven, 2 cm section samples from each core were processed and analyzed. Samples were systematically selected at different depth intervals of the core, with a maximum sample depth of 104 cm. This allowed us to analyse the samples in a temporal fashion, as deeper samples are from older sediments.

**Aging Sediment Core Samples**

A duplicate of core A was aged using lead isotopes (Townsend and Seen, 2012). Using the sedimentation rates and ages calculated in Townsend and Seen (2012) the age of sediment sections analyzed in this study were inferred. Sediments between 0 and 10 cm took 21 years to settle at a rate of 0.48 ± 0.05 cm/yr⁻¹ with sediments at 10 cm aged at 1983 ± 2. Sediments between 10 and 30 cm settled at a rate of 0.69 cm/yr⁻¹ with sediments at 30 cm aged at 1954 ± 5. Sediments between 30 and 50 cm took 46 years to settle at a rate of 0.43 ± 0.07 cm/yr⁻¹ with sediments at 50 cm aged at 1908 (Townsend and Seen, 2012).

**Laboatory Analysis**

Sample processing was adapted from Reeves et al. (2016). In brief, samples were processed using a stepwise approach include sieving, organic material digestion, density separation, centrifuging, and filtration to separate microplastics from the bulk sediment. All laboratory work was conducted under a vacuum hood and exposed samples and equipment were covered with foil to prevent contamination from airborne microplastics. Natural fiber clothing and laboratory coats were worn throughout the analysis to reduce microplastic contamination from synthetic clothing.

**Sievning**

Each sample was placed in a beaker with deionized water and agitated with a metal spatula to disassociate large clumps of sediment. The contents of the beaker were then poured through a sieve stack. Sieves were stacked sequentially according to mesh size, with the largest mesh size at the top. Each sieve was rinsed with deionised water and left to dry.

**Organic Material Digestion**

Each dried sample was placed in a beaker with 20 ml of 30% hydrogen peroxide, 20 ml of 0.05 M iron (II) solution and a magnetic stir bar. The sediment solution was left at room temperature for 5 min, then was placed on a heating magnetic stirrer and heated to 75°C for 30 min. If organic material was visible after 30 min, 20 ml of hydrogen peroxide was added every 15 min and stirring/heating continued until all visible organic material was digested.

**Density Separation, Centrifuging, and Filtration**

Each digested sample was placed in a 50 ml centrifuge tube. Large sediment samples were split evenly between two tubes. Sodium iodide (NaI) solution (density 1.6–1.8 g.ml) was added to each tube until 30 ml of NaI was overlaying the sediment sample. Each tube was capped, shaken manually for 20 s and then placed in a benchtop centrifuge for 5 min at 3,500 revolutions per minute. Tubes were removed gently to minimize sediment re-suspension. The top 10 ml of supernatant NaI solution was poured off into a glass Büchner vacuum apparatus fitted with a 1.2 μm polycarbonate membrane filter. The remaining sediment in the tube was topped up with NaI, so 30 ml was overlaying the sediment. The sediment then underwent the previously described treatment of manual shaking, centrifuging, and filtering of supernatant, twice. A
total of three supernatants, per sample, were poured into the vacuum apparatus and filtered. To ensure all possible separated microplastics were poured on the filter paper, on the last round of treatment (i.e., the third round), the total supernatant NaI solution was poured into the vacuum apparatus. The Büchner funnel and 1.2 µm filter were then rinsed with deionised water to capture any microplastics that may have adhered to the glass during filtration. All filtered samples were stored individually in sealed petri-dishes until microscope analysis.

**Microscope Analysis**

Filtered samples were analyzed for microplastic content using a stereomicroscope (magnification × 40). Each sample was divided into seven sorting sections using a fine-point metal probe. Samples were observed for 15 min using a constant magnification setting. Following the “Guidelines for Microscope Inspection” (Masura et al., 2015) an object was identified as plastic if it held shape or stretched when rubbed/pressed with a metal probe. Organic material would break a part under the above treatment. Positively identified microplastics were sorted into four categories according to their shape and texture (Table 1).

**Deposition Trend**

We tested whether the rate of micro fiber deposition observed in the cores correlated with plastic production using linear regression. We used the time trend in global plastic production values (PlasticsEurope, 2016) as a proxy for the relative time trend in production in the Derwent estuary. We used an exponential mode l to fit to the data available on global production to estimate the proportional change on an annual basis since plastic went into commercial production in the mid 1900s.

We adjusted our trend estimate for contamination by assuming the contamination rate was constant with respect to depth of the core slice. Based on this assumption the intercept term of the linear regression of microfiber concentration on plastic production change is an estimate of the contamination rate in the samples. The concentration of fibers in sediment slices from before 1950 serves as a second estimate of the contamination rate, as these sediments were deposited prior to the availability of plastic in the environment. The slope of the relationship between microfiber concentration and plastic production gives an estimate of the proportional increase in plastic in sediments, per unit of increase in production, which can be interpreted as the leakage rate from production.

We only used micro fiber values in the analysis as micro fibers made up nearly 90% of all microplastics in the cores.

### RESULTS

Microplastics were observed in every sediment sample of both cores (N = 14). A total of 211 microplastics were observed in core A (N = 7), and 252 microplastics were observed in core B (N = 7). The mean weight of dry sediment samples was 63.36 g for core A and 52.36 g for core B with a mean of 2.43 plastic fragments per gram of sediment for core A and 4.2 plastic fragments per gram of sediment for core B. Microplastics mainly occurred as fibers (87% of total items observed), followed by sheet (9%), fragment (3%), and beads (1%) (Figure 2). A control sample, i.e., conducting the laboratory method without a sediment sample, presented a maximum of eight microplastic fibers (range = 0–8).

### Size Classes of Microplastics Detected

In both cores, the frequency of microplastics increased with a decrease in size class. In core A, 137 microplastics were observed in size class 63 µm, 49 in size class 100 µm, 24 in size class 1 mm, and 1 in size class 4 mm (Figure 3). In core B, 132 microplastics were observed in size class 63 µm, 93 in size class 100 µm, 27 in size class 1 mm (Figure 3). In core A an unusually high number of microplastics were observed in sample 7.

### TABLE 1 | List of features used to identify different microplastic categories.

<table>
<thead>
<tr>
<th>Microplastic category</th>
<th>Shape</th>
<th>Texture</th>
<th>Color</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fiber</td>
<td>String-like with irregular bends</td>
<td>Soft, malleable</td>
<td>Any</td>
</tr>
<tr>
<td>Sheet</td>
<td>Thin, flat</td>
<td>Soft, malleable</td>
<td>Often clear, black, or translucent blue</td>
</tr>
<tr>
<td>Fragment</td>
<td>Thin, flat</td>
<td>Hard, rigid</td>
<td>Often black or red</td>
</tr>
<tr>
<td>Bead</td>
<td>Spherical, smooth</td>
<td>Hard</td>
<td>Often black or brown</td>
</tr>
</tbody>
</table>

![FIGURE 2 | The percentage of microplastic types observed in core samples A and B.](image)
Depth
In each core, we found more microplastics in the upper layer of sediments (more recent) than in the deeper (older) sediments. In core A, 38.9% of microplastics were observed in sample 1 and sample 2 depth classes, whereas only 26.1% of all microplastics were observed in the deeper layers (samples 6 and 7). In core B, we found 68.7% of microplastics in the upper, more recent layers (sample 1 and sample 2) whereas only 11.5% of microplastics were observed in the deeper layers (samples 6 and 7). Smaller microplastics were more common in shallower samples (i.e., there were more microplastics in the smallest size class (<63 µm–100 µm) in sample 1 than in sample 7 (Figure 3). In core A, 20.4% of all 63 µm microplastics were observed in sample 1 whereas only 11.7% in sample 7. In core B, 45.5% of all 63 µm microplastics were observed in sample 1 whereas only 3.0% in sample 7.

Deposition Trend
The number of micro fibers was higher in the shallower (younger) sediment samples. The rate of micro fiber deposition in the sediment correlated strongly with the expected change in production in the study region, based on the annual global plastic production (Figure 4; Table 2). The predicted accumulation model was a better model than the null (AIC: Predicted = 101, Null = 112). As global plastic production increased, the number of micro fibers deposited in sediments significantly increased ($p = <0.05$).

DISCUSSION
We observed more microplastics in the upper sediments. In core B there were six times more microplastics present in the top 15 cm (1976 and younger) than in the bottom 22 cm (1799 and older). In core A there were one and a half times the number of microplastics in the top 10 cm (1983 and younger) than in the bottom 22 cm (1799 and older). This observation is likely due to the exposure of upper sediments to a higher proportion of microplastics settling out from the water column. The production of plastics has increased from 100 million tons per year in 1993 to 322 million tons per year in 2015 (PlasticsEurope, 2008, 2016). The environments for our sediment cores differed; core B was taken 150 m off shore from a relatively high energy position in a deep channel, whereas core A was taken 1,000 m off shore in a large shallow bay with low energy and more rapid sedimentation (Figure 1). Despite the normal low energy dynamics of site core A, it is occasionally subject to the passage of large floods. This scour the embayment with fresh water and greatly changes the potential sources of microplastic contamination for short periods of time. These differences in site conditions could explain the higher concentration of plastics at site B. The site is not subject to large scouring events, thus any deposition is likely...
to remain in place. In addition, the site is much closer to sewage and stormwater outfall drains, which we found to be sources of plastic pollution in previous work (Willis et al., 2017). This result is similar to those found in other studies, which found higher numbers of plastics at sites closer to storm drain outfalls (Duckett and Repaci, 2015; Horton et al., 2017).

Most microplastics observed were fibers (87%). This phenomenon has been observed in other sediment analyses (MONAS, 2014; Woodall et al., 2015; Zobkov and Esiukova, 2017). Browne (2015) suggests microplastic fibers found in marine habitats may be derived from sewage as consequence of washing clothes. Furthermore, up to 1,770 microplastics have been reported to leave a waste water treatment plant in effluent water per hour (Magnusson and Norén, 2014). As many outfalls enter the Derwent Estuary, they are a probable source for the large quantity of microfibers observed in the samples. Outfalls as a source of micro fibers in the marine environment also indicates the dispersion from source to point of deposition is relatively local.

The strong correlation between the observed micro fibers and the predicted change in plastic production (Figure 4) indicates there is a clear temporal trend in micro fiber deposition in sediments (Table 2). This suggests that as plastic production increases, microplastic pollution is increasing proportionately. This is likely due to an increase in plastic leakage from the waste stream, as has been reported elsewhere (Thompson et al., 2009). The increase in coastal populations has been observed to increase the amount of pollution entering waterways (Jambeck et al., 2015; Lebreton et al., 2017). For example, 40% of Tasmania’s population lives around the margins of the Derwent Estuary, with the population doubling between 1950 and 2015 (Carver, 1954; Coughanowr et al., 2015). Hence, older deeper sediments were exposed to an environment with markedly lower population and plastic production rate and less opportunity for plastic contamination than the younger, shallower sediments. This pattern was also observed on Belgian beaches where plastic pollution in sediments had tripled over 20 years (Claessens et al., 2011).

Our results demonstrate micro fibers are present in marine sediments that settled pre-plastic production. Microplastics have been recorded in marine sediments since the 1970s (Gregory, 1977; Shiber, 1979). We expected to only observe microplastics in sediments shallower than 30 cm, as the sediments were aged to be younger than 1954 ± 5 years (Townsend and Seen, 2012). However, we observed micro fibers down to 104 cm. Sediment accumulation rates in Elwick Bay range from 0.4 to 0.7 cm per year (Townsend and Seen, 2012). Hence, sediment at 104 cm would have settled between 149 and 260 years ago (i.e., in 1855 to 1774). Observing micro fibers down to 104 cm does not indicate that microplastics have been settling in sediment for 260 years. Rather it raises concerns that the sediment cores were exposed to micro fiber contamination either during collection or analysis. We calculated the mean frequency of fibers in our samples pre-1950s (i.e., plastic production) to estimate the fraction of fibers from both field and laboratory contamination, which yielded an estimate of up to 17.2 fibers (∼60%) per sediment sample due to contamination, which is in line with the estimate of 21 fibers per sample, based on the intercept term in our linear regression (Figure 4). As the exact date that plastic production commenced in the Derwent Estuary is not known the values from the predicted model will give a better estimation of past microplastic levels in the sediment of the Derwent Estuary.

The fine structure of microplastics could enable them to move deeper into the older sediment via mixing due to bioturbation or water flows e.g., storm/flood events and direct transport by animals. This downward transport should affect all microplastic types, however, we only found micro fibers in the older sediment. The high number of fibers observed in the laboratory blanks indicates that the preventative contamination measures employed during laboratory analysis were ineffective. As we were unable to conduct our microscope analysis under a fume hood, airborne fibers could be one point of sample contamination. It is also possible that contamination occurred during the field collection and slicing of the cores. The cores were not originally collected for microplastic analysis. Hence, methods to prevent microplastic contamination were not a component of the field sampling program. We suspect contamination from researcher’s clothing and equipment (i.e., synthetic fibers from rope fragmentation; Thompson et al., 2004) or from airborne microplastics (Dris et al., 2015) as the most likely contamination sources. This points to a major issue with the common opportunistic use of sediment samples to look for microplastics in deep sea sediments and other places. The opportunistic nature of sampling questions whether adequate procedures to prevent microplastic contamination were undertaken. Samples in this study and others should be interpreted with contamination in mind as the results showed 20 micro fibers per 50 g of sediment can be due to contamination alone.

Considerably fewer sheet, fragment, and bead microplastics were observed than fibers (13% for these three categories combined). These non-fiber plastics were all observed in the upper sediment layers which suggests there was no contamination issue for these materials. The lower counts of non-fiber microplastics could indicate these microplastics are not a common contaminant in estuaries. However, observer bias is also a likely explanation. Microfibers may be more conspicuous under a microscope due to their unique irregular bent filament shape and commonly settling on top of the other filtered particles (i.e., sediment granules, undigested biological matter, diatom shells). Sheet, fragment, and bead microplastics may be harder to detect as their shapes were more similar to those of undigested plant material and sediment granules. Hence, they could be underrepresented in counts from sediment cores.

<table>
<thead>
<tr>
<th>Model</th>
<th>Terms</th>
<th>Estimate</th>
<th>P-value</th>
<th>AIC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Predicted</td>
<td>Number of fibers</td>
<td>21.2944</td>
<td>0.01329</td>
<td>101.0587</td>
</tr>
<tr>
<td>Accumulation</td>
<td>Plastic production estimate</td>
<td>1.2084</td>
<td>0.00142</td>
<td></td>
</tr>
<tr>
<td>Null</td>
<td>Number of fibers</td>
<td>36.45</td>
<td>0.00578</td>
<td>112.1359</td>
</tr>
</tbody>
</table>
We found size class 63\(\mu\)m had the highest count of microplastics in every sample but one. Eighty-five percent of sediments in Elwick Bay are <63\(\mu\)m (Koehnken and Eriksen, 2004), reflecting a highly organically enriched depositional area. As both sediments and microplastics are a similar size, it can be inferred that the same forces act on sediment accumulations and microplastics accumulation (Vianello et al., 2013). The high frequency of 63\(\mu\)m microplastics in sediments is quite different to the size distribution observed from net trawls of the ocean surface. In net trawls, small microplastics (i.e., 100 to 63\(\mu\)m) are not observed even though they are likely present on the surface (Law et al., 2010; Eriksen et al., 2013). This discrepancy in microplastic sizes may be largely due to the difference in sample analysis (e.g., visual versus microscope identification) (van Sebille et al., 2015) as it would be unlikely or impossible to observe microplastics between 100 and 63\(\mu\)m solely scanning with the naked eye.

**CONCLUDING REMARKS**

Our study shows that sediments are a useful record of past and present plastic leakage from the waste stream into the marine environment. This is not unlike samples from other parts of the marine ecosystems, including the water column (van Sebille et al., 2015), seabirds (Wilcox et al., 2015), and turtles (Schuyler et al., 2014). Microplastics were present in all samples, even in sediments dated from pre-plastic production. Based on our estimates, current microplastic concentrations in sediment are 115 microplastics, and are increasing at an accelerated rate of 1.208 microplastics per year. Generally speaking, however, the frequency of plastics corresponded with the increase in plastic production and coastal populations. The presence of microplastics in the older sediments indicates there was possible contamination during sampling or laboratory analysis, which is an element that must be carefully considered when estimating microplastics loads and their presumed ubiquity in the marine environment.

**AUTHOR CONTRIBUTIONS**

BDH and CW: designed the study; KAW: completed laboratory work and wrote original draft, with contributions by other authors; BDH and RE: provided technical advice and support for laboratory work; RE: collected the samples; CW: completed data manipulation, statistical analyses, and provided technical expertise on manuscript.

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