



Microplastics in sediments: A review of techniques, occurrence and effects



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ABSTRACT

Microplastics are omnipresent in the marine environment and sediments are hypothesized to be major sinks of these plastics. Here, over 100 articles spanning the last 50 year are reviewed with following objectives: (i) to evaluate current microplastic extraction techniques, (ii) to discuss the occurrence and worldwide distribution of microplastics in sediments, and (iii) to make a comprehensive assessment of the possible adverse effects of this type of pollution to marine organisms. Based on this review we propose future research needs and conclude that there is a clear need for a standardized techniques, unified reporting units and more realistic effect assessments.

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1. Introduction

Plastic has changed the way we live. It possesses a unique set of properties making it extremely popular for use in everyday life: it can be used at a wide range of temperatures, has low thermal conductivity, a high strength-to-weight ratio, is bio-inert, durable and above all it is cheap (Andrady, 2011; Andrady and Neal, 2009). This has led to the use of plastic in a myriad of applications, ranging from household and personal goods, clothing and packaging to construction materials. As a result, the global plastic production has grown exponentially ever since its mass production started in the 1950s, with 288 million tonnes produced worldwide in 2012 (PlasticsEurope, 2013). Even though the societal benefits of plastic are undeniable (Andrady and Neal, 2009), there are some serious environmental concerns associated with the material. While a part of the plastic waste is properly managed (through combustion or recycling), it has been estimated that millions of tonnes of plastic waste (4.8–12.7 million tonnes in 2010) end up the marine environment (Jambeck et al., 2015).

Plastics are present in the environment in a wide variety of sizes, ranging from metres to micrometers (Barnes et al., 2009). The smallest form of plastic litter is called microplastic. These are present in the environment as ‘microplastics by design’, so-called primary microplastics, or arise from the degradation of larger plastic litter. While the former are typically resin pellets and microbeads associated with industrial spillages (EPA, 1992) and the use of cosmetics (Fendall and Sewell, 2009; Zitko and Hanlon, 1991), the latter (secondary microplastics) are formed through the action of degrading forces such as UV radiation and physical abrasion (Barnes et al., 2009; Cole et al., 2011). Another important source comes from synthetic clothing: a single synthetic garment can release up to 1900 fibres per washing cycle (Browne et al., 2011).

At present, there is no universally accepted definition regarding the size of microplastics. When first described in 2004, the term microplastic was adopted to refer to microscopic plastic debris in the 20 µm region (Thompson et al., 2004). A motion to broaden the definition to all fragments smaller than 5 mm was made in 2009 (Arthur et al., 2009). While the value of 5 mm is more commonly accepted, the 1 mm upper size limit is a more intuitive one as ‘micro’ refers to the micrometer range. As a result, this more strict

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definition is also often used in scientific literature (e.g. Browne et al., 2011; Claessens et al., 2011; Van Cauwenberghe et al., 2013; Vianello et al., 2013; Dekiff et al., 2014).

Microplastics have been reported in the water column and marine sediments worldwide (Claessens et al., 2011; Law et al., 2010; Moore et al., 2001; Thompson et al., 2004). While the first reports on microplastics in surface waters already date back to the early 1970s (Carpenter et al., 1972; Carpenter and Smith, 1972), it took another 5 years until the first records of plastic pellets on beaches were made (Gregory, 1977, 1983; Shiber, 1979) and another thirty years until the first microplastics (<1 mm) in sediments were reported (Thompson et al., 2004). Sediments are suggested to be a long-term sink for microplastics (Cózar et al., 2014; Law et al., 2010; Morét-Ferguson et al., 2010). Logically, plastics with a density that exceeds that of seawater ($>1.02 \text{ g cm}^{-3}$) will sink and accumulate in the sediment, while low-density particles tend to float on the sea surface or in the water column. However, through density-modification even low-density plastics can reach the seafloor. Biomass accumulation due to biofouling can lead to an increase in density resulting in the sinking of the microplastic (Andrady, 2011; Reisser et al., 2013; Zettler et al., 2013). Using nitrogen as a proxy, Morét-Ferguson et al. (2010) concluded that the reported change in microplastic density is due to attached biomass. Indeed, analysis of polyethylene bags submerged in seawater for 3 weeks showed a significant increase in biofilm formation over time, accompanied by corresponding changes in physicochemical properties of the plastic, such as a decrease in buoyancy (Lobelle and Cunliffe, 2011). These studies suggest that biofouling can contribute towards the settling and eventual burial in sediments of previously buoyant plastic. Biomass accumulation on plastic may even partly explain the recent finding that the global plastic load in the open-ocean surface is estimated to be two orders of magnitude lower than expected from estimates of plastic releases in the marine environment (Cózar et al., 2014).

The main objective of this review is to assess the state of the science in the exposure and effects assessment of microplastics in the marine environment, more specifically in marine sediments. This was achieved by analysing available literature to: (1) provide an in-depth evaluation of the current and commonly used techniques for extracting microplastics from sediments, (2) discuss the occurrence and distribution of microplastics in marine sediments worldwide and (3) make a comprehensive assessment of the known effects to benthic and sediment-associated wildlife.

2. Review of available literature

We conducted an extensive literature review using the ISI Web of Knowledge and Google Scholar databases. Based on the search parameters detailed below, a total of 122 original publications were retrieved, dating back to 1977. The majority of publications (90%) were published from 2004 onwards, with 75% of all literature published in the last five years (Fig. 1). Next to peer-reviewed papers, conference proceedings, posters and dissertations were also included in this review.

From these publications, all necessary information regarding (i) the extraction technique, (ii) microplastic abundance and distribution and in the case of effect assessments (iii) exposure concentration and observed effects was extracted and processed.

In the ISI Web of Knowledge, a literature search using the keywords 'plastic pellet or microplastic' in combination with 'sediment or beach' generated a list of 139 peer-reviewed papers. These date back to 1982 and cover the period until the beginning of 2015. From these publications a list of 32 papers on occurrence and distribution, 9 reviews and 5 papers presenting and discussing extraction

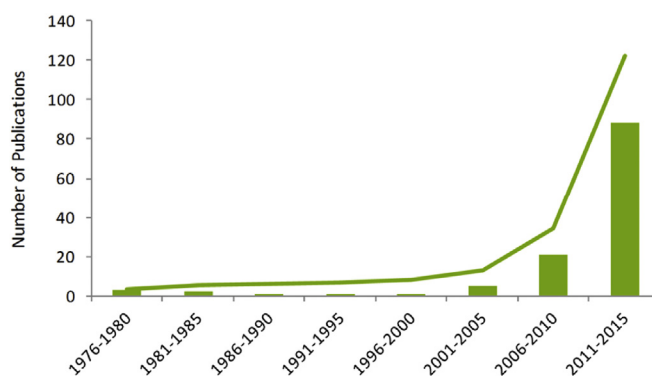


Fig. 1. Evolution in the publication of 'microplastic in sediment' literature. The bars represent the number of publications published in the corresponding 5-year period, while the curve represents the cumulative distribution of the published literature since 1975.

techniques was compiled. An additional search on the Google Scholar search engine, using the same keywords, yielded and additional 19 publications, posters and dissertations on occurrence and distribution of microplastics in sediments (1977–2015).

Using the ISI Web of Knowledge database, the queries 'microplastic, organism, ingestion' and 'microplastic contaminant or microorganism' resulted in two publication lists of 18 original publications each. These publications go back to 1994 and cover the period until the beginning of 2015. Still, this collection of publications revealed not all the relevant information on the direct and indirect effects of microplastics to epibenthic species. The Google Scholar search engine revealed additional hits for these queries, including conference posters, conference proceedings and dissertations. From these lists, a final relevant literature list of 57 publications, posters and dissertations was composed.

3. Sampling and extraction techniques

Due to the rapid development of microplastic research, there is a lack of consistency in sampling and extraction techniques used to quantify microplastics in sediments. As a result of the large variety in techniques applied, comparison of reported microplastic concentrations between studies is often impossible or requires additional calculations based on assumptions (e.g. sediment densities). The majority of these method inconsistencies can be related to (i) differences in the lower and upper size limit implemented, (ii) the sensitivity of the applied extraction technique and (iii) differences in sampling technique leading to a wide variety of reporting units.

The lack of an unequivocal size-based definition of microplastic has resulted in the reporting of several different size fractions in literature, all using the same term: microplastics. In practice, this means that the results of a substantial body of microplastic literature cannot be compared directly. As microplastics include particles up to 5 mm (Arthur et al., 2009) and both extraction and identification becomes more challenging with decreasing dimensions, authors often opt to only include plastics larger than 1 mm (e.g. Baztan et al., 2014; Jayasiri et al., 2013; McDermid and McMullen, 2004) or even >2 mm (e.g. Heo et al., 2013; Ivar do Sul et al., 2009; Turner and Holmes, 2011). Even among those studies that do include the smallest of microplastics (down to $1.6 \mu\text{m}$) different upper size limits are applied: either 1 mm (Browne et al., 2011, 2010; Claessens et al., 2011; Van Cauwenberghe et al., 2013a) or 5 mm (Martins and Sobral, 2011; Mathalon and Hill, 2014; Ng and Obbard, 2006; Reddy et al., 2006). As both different lower and

upper size limits are used throughout microplastic literature, a vast amount of data on microplastic occurrence and distribution worldwide is lost. Yet, this inconsistent use of the term ‘microplastic’ can be easily addressed by introducing a more comprehensive classification to differentiate between small microplastics (SMPs: < 1 mm) and large microplastics (LMPs: 1–5 mm) (Fig. 2) as proposed by European MSFD technical subgroup on Marine Litter (Galgani et al., 2013). Another earlier study suggests the following: micro- (<0.5 mm) and mesolitter (0.5–10 mm) (Gregory and Andrady, 2003). While the discussion often focuses on the upper size limit of microplastics, it can be argued that the adoption of a lower size limit is equally important. To date, the lower size limit used in microplastic assessment studies is highly dependent on the sensitivity of the sampling and extractions techniques applied. Often, the technical constraints associated with the extraction of small microplastics (SMPs) result in the omission of this lower size limit. However, not including the sub-1 mm fraction can result in reporting highly underestimated concentrations. Indeed, it has been demonstrated repeatedly that these small microplastics represent 35–90% of all microplastics present in the marine environment (Browne et al., 2010; Eriksen et al., 2013; McDermid and McMullen, 2004; Song et al., 2014; Zhao et al., 2014).

A wide range of sampling techniques is used for monitoring microplastics in sediments (reviewed in Hidalgo-Ruz et al., 2012 and Rocha-Santos and Duarte, 2015). As a result, the reported abundances are often expressed in different units. While a simple conversion can sometimes be made to compare among studies, often comparison is impossible or requires assumptions that lead to biased results. The choice of sampling strategy and sampling approach (reviewed by Hidalgo-Ruz et al., 2012) will eventually determine the unit in which observed abundances will be reported. Those studies sampling an area (using quadrants) will often report abundances per unit of surface (m^2 ; e.g. Ivar do Sul et al., 2009; Lee et al., 2013; Martins and Sobral, 2011). If areal bulk samples up to a specific depth are taken the reporting unit is m^3 (e.g. Ballent et al., 2012; Turra et al., 2014). Conversion between these type of abundances is possible, if sufficient information is available on sampling depth. Yet, for 20% of the studies this is not the case as reported sampling depths can range from 0 to 50 cm. Other widely used reporting units are volume (mL to L; e.g. McDermid and McMullen, 2004; Norén, 2007; Thompson et al., 2004) or weight (g to kg; e.g. Claessens et al., 2011; Ng and Obbard, 2006; Reddy et al., 2006). Conversion between these two types of units is not straight forward as detailed information on the density of the sediment is required. As this is never (as far as we could establish) reported in microplastic studies, assumptions have to be made, as Claessens et al. (2011) did for the conversion of microplastic abundances in sediment. Additionally, within studies reporting weight, a distinction can be made among those reporting wet (sediment) weight and those reporting dry weight. This adds to the constraints of converting from weight to volume units, or vice versa. Sediment samples from different locations or even different zones on one beach (e.g. high littoral vs. sub littoral zone) have different water content. Therefore, a (limited) number of authors choose to express microplastic concentrations as dry weight eliminate this variable

(Claessens et al., 2011; Dekiff et al., 2014; Ng and Obbard, 2006; Nor and Obbard, 2014; Van Cauwenberghe et al., 2013a; Vianello et al., 2013).

After sampling, either from beach sediments or the seabed, different approaches can be used to separate the microplastic fragments from the sandy or muddy matrix. The most common approach is to extract plastic particles from the sediment using a density separation, based on the differences in density between plastic and sediment particles. Typically, this is achieved by agitating the sediment sample in concentrated sodium chloride (NaCl) solution, as described by Thompson et al. (2004). However, as the density of the NaCl solution is only 1.2 g cm^{-3} , only low-density plastics will float to the surface and can hence be extracted. Different authors have addressed this issue by using different salt solutions to obtain higher densities. Liebezeit et al. (2012) and Imhof et al. (2013) extracted microplastics from sediments using zinc chloride (ZnCl_2 , $1.5\text{--}1.7 \text{ g cm}^{-3}$), while others (Dekiff et al., 2014; Van Cauwenberghe et al., 2013a; Van Cauwenberghe et al., 2013b) used a sodium iodide (NaI, $1.6\text{--}1.8 \text{ g cm}^{-3}$) solution. These modifications of the commonly used method of Thompson et al. (2004) result in an increased extraction efficiency for high-density microplastics such as polyvinylchloride (PVC, density $1.14\text{--}1.56 \text{ g cm}^{-3}$) or polyethylene terephthalate (PET, density $1.32\text{--}1.41 \text{ g cm}^{-3}$). As these high-density plastics make up over 17% of the global plastic demand (PlasticsEurope, 2013), not including these types of microplastic can result in a considerable underestimation of microplastic abundances in sediments. Especially as these high-density plastics are the first to settle and incorporate into marine sediments (density of seawater is 1.02 g cm^{-3}).

As sampling, extraction and detection methods and techniques are being developed worldwide (Claessens et al., 2013; Fries et al., 2013; Harrison et al., 2012; Imhof et al., 2012; Nuelle et al., 2014) it is clear that in order to completely understand the distribution of microplastics in the marine environment, a harmonisation and standardisation of techniques and protocols is urgently needed to enhance microplastic research and monitoring.

4. Occurrence of microplastics in sediments

The first reports of microplastics associated with sediments date back to the late 1970s. These early observations comprised industrial resin pellets (2–5 mm) on beaches in New Zealand, Canada, Bermuda, Lebanon and Spain (Gregory, 1977, 1978, 1983; Shiber, 1979, 1982), demonstrating -already back then-their worldwide distribution. Even in these first reports, pellet concentrations regularly exceeded 1000 pellets per metre of beach, with extreme abundances reported from 20,000 to 100,000 pellets m^{-1} (Gregory, 1978). Large ports and local plastic industry were considered major sources, while for Bermuda -which lacks such local sources-the influence of oceanic circulation patterns (located in the west of the North Atlantic Gyre) explain the high concentrations (Gregory, 1983). Large numbers of beached industrial pellets in association with labelled, intact bags detected on beaches in the United Arabian Emirates and Oman confirmed the importance of local contamination sources (Khordagui and Abu-Hilal, 1994). Ever since these

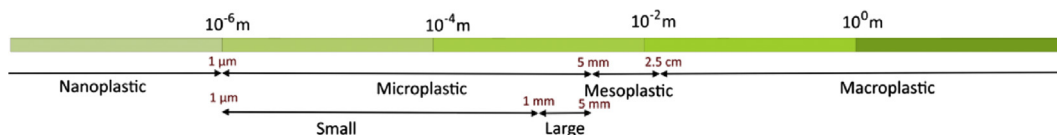


Fig. 2. Size matters. Suggestion of plastic debris nomenclature based on size, as proposed by the European MSFD technical subgroup on Marine Litter (MSFD GES Technical Subgroup on Marine Litter, 2013). The overall term “microplastic” is composed of small microplastics (SMPs, smaller than 1 mm) and large microplastics (LMPs, 1–5 mm), to differentiate between two commonly used definitions of microplastics.

first studies, pellet contamination of beaches worldwide has been reported (Table 1). For the majority of these studies the main focus was not to assess the occurrence and abundance of these pellets, but rather to evaluate the contaminant load present on these pellets. Indeed, their size, long environmental persistence and worldwide distribution, make them especially suitable for chemical analysis (Mato et al., 2001). Many hydrophobic compounds (including polychlorinated biphenyls (PCBs), polycyclic aromatic hydrocarbons (PAHs), dichlorodiphenyltrichloroethane (DDT) and degradation products) have been detected on pellets collected from marine environments. Concentrations of PCBs on polypropylene pellets collected in Japan were up to 10^6 times that of the surrounding seawater (Mato et al., 2001). Recently, Fotopoulou and Karapanagioti (2012) demonstrated that surface alterations in pellets, resulting from environmental erosion, can explain the increased affinity for contaminants of pellets (Endo et al., 2005). While virgin pellets have smooth and uniform surfaces, eroded pellets exhibited an uneven surface with an increased surface area and polarity, affecting the efficiency of sorption (Fotopoulou and Karapanagioti, 2012).

While the occurrence of industrial resin pellets in marine environments were already described in the 1970s, it took another 30 years before the first reports on other types of microplastics were published. By analysing subtidal, estuarine and sandy sediments from 18 locations across the UK, Thompson et al. (2004) were the first to demonstrate the presence of μm -sized (<1 mm) microplastics in marine sediments. Soon, reports from Singapore (Ng and

Obbard, 2006), India (Reddy et al., 2006) and Sweden (Norén, 2007) illustrated the widespread distribution of these small microplastics. Currently, small and large microplastics are detected in sediments worldwide: especially beaches, subtidal and offshore sediments have been examined (Table 2, Fig. 3). Recently, even deep oceanic sediments have been shown to contain microplastics: up to 2000 particles per m^2 are detected in sediments at a depth of 5000 m (Fisher et al., 2015; Van Cauwenberghe et al., 2013b). It has also been demonstrated that the level of plastic pollution is increasing: sediment core analysis revealed that over the last 20 years microplastic deposition on Belgian beaches tripled (Claessens et al., 2011).

Due to their easy accessibility, sandy beaches have been the main focus of studies assessing microplastic abundance (over 80% of reviewed abundance studies). The zone sampled, however, differs among studies: while some studies cover entire beach transects (perpendicular to the shoreline), others studied specific littoral zones. As was already remarked by Hidalgo-Ruz et al. (2012), this lack in uniformity between studies explains why the distribution of microplastics on beaches is still little understood, and that there is a need to systematically examine potential accumulation zones of microplastics. In a recent attempt to elucidate the distribution of microplastics across the different beach zones, Heo et al. (2013) analysed the entire cross section (from back-to foreshore) of an impacted South Korean beach. Their results indicated that, unlike macroplastics, which accumulated at the high tide line, microplastics (2–10 mm) were most abundant in the

Table 1
Available literature on pollution of marine sediments by industrial resin pellets. Origin and main focus of the research (i.e. assessing occurrence and abundance, assessing contaminant load or investigating surface characteristics) is provided.

Continent	Location	Main focus	Reference
Africa	Canary Islands	Contaminant load	Heskett et al., 2011
	Saint Helena	Contaminant load	Heskett et al., 2011
	South Africa	Contaminant load	Ryan et al., 2012
America	Barbados	Contaminant load	Heskett et al., 2011
	Bermuda	Occurrence	Gregory, 1983
	Brazil	Occurrence	Costa et al., 2010
		Occurrence	Turra et al., 2014
		Contaminant load	Fisner et al., 2013a
		Contaminant load	Fisner et al., 2013b
	California	Contaminant load	Rios et al., 2007
		Contaminant load	Van et al., 2012
	Canada	Occurrence	Gregory, 1983
	Hawaii	Occurrence	McDermid and McMullen, 2004
		Contaminant load	Rios et al., 2007
		Contaminant load	Heskett et al., 2011
Asia	Cocos Islands	Contaminant load	Heskett et al., 2011
	Hong Kong	Contaminant load	Zurcher, 2009
	Japan	Contaminant load	Mato et al., 2001
		Contaminant load	Endo et al., 2005
		Characteristics	Kuriyama et al., 2002
	Jordan	Occurrence	Abu-Hilal and Al-Najjar, 2009
	Lebanon	Occurrence	Shiber, 1979
	Malaysia	Occurrence	Ismail et al., 2009
	Oman	Occurrence	Khordagui and Abu-Hilal, 1994
	United Arabian Emirates	Occurrence	Khordagui and Abu-Hilal, 1994
	New Zealand	Occurrence	Gregory, 1977
		Occurrence + Contaminant load	Gregory, 1978
Europe	Belgium	Occurrence	Van Cauwenberghe et al., 2013a
	Greece	Contaminant load	Karapanagioti and Klontza, 2008
		Contaminant load	Karapanagioti et al., 2011
	Malta	Occurrence + Characteristics	Turner and Holmes, 2011
	Portugal	Contaminant load	Frias et al., 2010
		Occurrence + Contaminant load	Antunes et al., 2013
		Contaminant load	Mizukawa et al., 2013
	Spain	Occurrence	Shiber, 1982
		Occurrence	Shiber, 1987
	United Kingdom	Contaminant load	Ashton et al., 2010
		Contaminant load	Holmes et al., 2012

Table 2

Abundance of microplastics in sediments worldwide. Location and location specification (i.e. 'sediment type') are provided, as well as the microplastic size range (particle size) applied during the assessment.

Continent	Location	Location specification	Particle size	Measured abundance	Reference
Africa	Canary Islands	Beach	1 mm–5 mm	<1 – >100 g/L	Baztan et al., 2014
	Hawaii	Beach	1 mm–15 mm	541–18,559 items/260 L	McDermid and McMullen, 2004
America	US	Florida subtidal	250 µm–4 mm	116–215 items/L	Graham and Thompson, 2009
	Maine subtidal			105 items/L	
	Brazil	Beach	2 mm–5 mm	60 items/m ²	Ivar do Sul et al., 2009
	Brazil	Beach	0.5 mm–1 mm	200 items/0.01 m ²	Costa et al., 2010
			1 mm–20 mm	100 items/0.01 m ²	
	Hawaii	Beach	250 µm–10 mm	0.12%–3.3% plastic by weight	Carson et al., 2011
	Brazil	Tidal plain	1 mm–10 cm	6.36–15.89 items/m ²	Costa et al., 2011
	Chile	Beach	1 mm–4.75 mm	<1–805 items/m ²	Hidalgo-Ruz and Thiel, 2013
	Québec	River sediment	400 µm–2.16 mm	52–13,832 beads/m ²	Castañeda et al., 2014
	Nova Scotia	Beach	0.8 µm–5 mm	20–80 fibres/10 g	Mathalon and Hill, 2014
Asia	Singapore	Beach	1.6 µm–5 mm	0–4 items/250 g dry	Ng and Obbard, 2006
	India	Ship-breaking yard	1.6 µm–5 mm	81.4 mg/kg	Reddy et al., 2006
	South Korea	High tide line	2 mm–10 mm	913 items/m ²	Heo et al., 2013
	India	Beach	1 mm–5 mm	10–180 items/m ²	Jayasiri et al., 2013
	South Korea	Beach dry season	1 mm–5 mm	8205 items/m ²	Lee et al., 2013
		Beach rainy season		27,606 items/m ²	
	Singapore	Mangrove	1.6 µm–5 mm	36.8 items/kg dry	Nor and Obbard, 2014
	NW Pacific	Deep sea trench	300 µm–5 mm	60–2020 items/m ²	Fisher et al., 2015
	South Korea	Beach	50 µm–5 mm	56–285,673 items/m ²	Kim et al., 2015
	UK	Beach	1.6 µm–5 mm	0.4 fibres/50 mL	Thompson et al., 2004
Europe		Estuary		2.4 fibres/50 mL	
		Subtidal		5.6 fibres/50 mL	
	Sweden	Subtidal	2 µm–5 mm	2–332 items/100 mL	Norén, 2007
	UK	Beach	1.6 µm–1 mm	<1–8 items/50 mL	Browne et al., 2010
	UK	North Sea beach	38 µm–1 mm	0.2–0.8 fibres/50 mL	Browne et al., 2011
		English Ch. beach		0.4–1 fibres/50 mL	
	Belgium	Harbour	38 µm–1 mm	166.7 items/kg dry	Claessens et al., 2011
		Continental Shelf		97.2 items/kg dry	
		Beach		92.8 items/kg dry	
	Portugal	Beach	1.2 µm–5 mm	133.3 items/m ²	Martins and Sobral, 2011
	Germany	Urban beach	1 mm–15 mm	5000–7000 items/m ³	Ballent et al., 2012
		Rural beach		150–700 items/m ³	
	Germany	Tidal flat	1.2 µm–5 mm	0–621 items/10 g	Liebezeit and Dubaish, 2012
	Italy	Sub-alpine lake	9 µm–5 mm	1108 items/m ²	Imhof et al., 2013
	Greece	Beach	1 mm–2 mm	57–602 items/m ²	Kaberi et al., 2013
			2 mm–4 mm	10–575 items/m ²	
	Belgium	High tide line	38 µm–1 mm	9.2 items/kg dry	Van Cauwenberghe et al., 2013
		Low tide line		17.7 items/kg dry	
	Italy	Subtidal	0.7 µm–1 mm	672–2175 items/kg dry	Vianello et al., 2013
	Germany	Beach	<1 mm	1.3–2.3 items/kg dry	Dekiff et al., 2014
	Slovenia	Beach	0.25–5 mm	177.8 items/kg dry	Laglbauer et al., 2014
Worldwide		Infralittoral		170.4 items/kg dry	
		Deep sea	5 µm–1 mm	0.5 items/cm ²	Van Cauwenberghe et al., 2013

upper intertidal zone, closer to the backshore. These results indicate that the mechanisms influencing macroplastic distribution on beaches, like wind and currents (Carson et al., 2013a; Thornton and Jackson, 1998), affect microplastic distribution in a different way. As a result, choosing the appropriate site or zone for microplastic assessment on beaches may not be as straight forward as previously thought, yet presents a critical factor in the assessment of microplastic pollution in coastal regions (Kim et al., 2015).

Differences in macro-versus microplastic distribution on beaches was also demonstrated by Browne et al. (2010) in the Tamar estuary (UK). In this study, plastic density and beach orientation (up- or downwind) best explained the observed macroplastic distribution, indicating the influence of wind created currents in the distribution of large floating debris. It was hypothesized that, due to their small sizes, microplastics in the water column will behave in the same way as sediment particles. Yet, no clear relationship was observed between microplastic (<1 mm) abundance and the proportion of clay in the sediment (Browne et al., 2010). It was therefore argued that other processes such as aggregation with organic material might play a more important role in the movement of microplastics. Indeed, Long et al. (2015) demonstrated in a

laboratory study that different algae species (*Chaetoceros neogracile* and *Rhodomonas salina*) incorporate and concentrate microplastics, substantially increasing microplastic sinking rates. Moreover, Strand et al. (2013) demonstrated that there is a strong relationship between microplastic abundance and both organic (%TOC) and fine fraction (<63 µm) content in sediments, supporting the hypothesis that microplastics will accumulate in depositional areas. In the Lagoon of Venice, Vianello et al. (2013) detected the lowest microplastic concentrations in the outer Lagoon, where water currents are higher (>1 m s⁻¹). Consequently, the highest concentrations were encountered in the inner Lagoon which is characterised by lower hydrodynamics and a higher fine particle (<63 µm) fraction in the sediment. Aggregation with organic matter (i.e. marine snow) was also considered the main route of transport for microplastics to deep-sea sediments (Van Cauwenberghe et al., 2013b).

Microplastics are categorised in different classes, based on their overall appearance using simple features such as shape, colour, etc. Several categories are used throughout literature, depending on the criteria applied by the authors. Types that re-occur frequently are: pellets, fragments, granules, fibres, films and Styrofoam. Due to

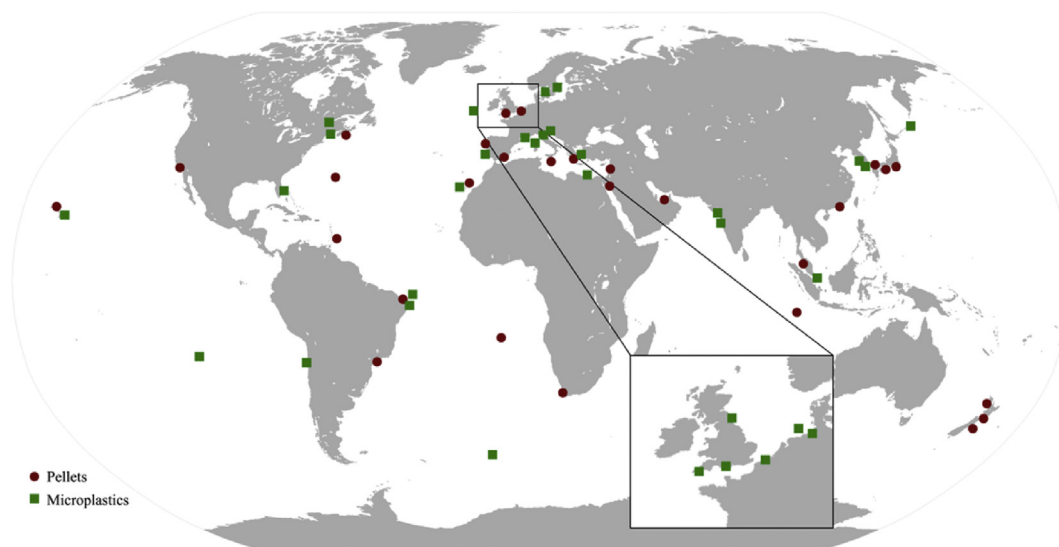


Fig. 3. Geographical distribution of studies reporting industrial resin pellets and other microplastic types in sediments. Black circles indicate studies that reported on the abundance or presence of industrial resin pellets, black squares indicate studies that focus on other microplastic types (i.e. fragments, microbeads and fibres).

their distinctive shape microplastic fibres are easily recognised in environmental samples. As a result, some studies primarily focus on fibres rather than particles (Browne et al., 2011, 2010; Fisher et al., 2015; Mathalon and Hill, 2014; Thompson et al., 2004). It was demonstrated by Browne et al. (2011) that such fibres are indicative of a sewage origin: an increased microfibre load (>250%) was detected in sewage-sludge disposal sites compared to reference sites. As the majority of microplastic fibres were either polyester or acrylic, synthetic garments were considered important sources of microplastics. Browne et al. (2011) investigated the contribution of the use of domestic washing machines and concluded that washing synthetic garments contribute considerable numbers of microplastics to marine environments: up to 1900 fibres can be released into the environment from washing a single piece of clothing.

Microplastics appear to be more abundant in densely populated areas. In a study analysing sediments from 18 locations representing 6 continents, Browne et al. (2011) demonstrated a positive relationship between microplastic and human population density. Indeed, microplastics are detected in large numbers in highly populated areas, such as at locations in the North Sea (Claessens et al., 2011; Liebezeit and Dubaish, 2012; Norén, 2007; Thompson et al., 2004; Van Cauwenberghe et al., 2013a) and the Mediterranean Sea (Kaberi et al., 2013; Klostermann, 2012; Vianello et al., 2013), as well as in Asia (Ismail et al., 2009; Ng and Obbard, 2006; Nor and Obbard, 2014; Reddy et al., 2006) and the highly populated coast of Brazil (Costa et al., 2010; Ivar do Sul et al., 2009; Turra et al., 2014). On heavily polluted beaches, microplastics (0.25–10 mm) can make up 3.3% of the sediment by weight, as opposed to 0.12% plastic by weight on control beaches (Carson et al., 2011). On these Hawaiian beaches, plastics ranging in size from 0.25 to 4 mm were most abundant (55.5%), yet proportions of microplastics (1–4.75 mm) of up to 90% have been reported as well (McDermid and McMullen, 2004). The link between microplastic pollution in sediments and human activities has also been demonstrated by Claessens et al. (2011), who detected particularly high concentrations of microplastic granules in the sediments of coastal harbours. However, as not all types of microplastics could be linked to sources in the harbours, the importance of rivers as potential sources of microplastics to the marine environment was stressed. Recently, this was confirmed by Vianello et al. (2013), who

detected the highest microplastic concentrations in those areas influenced most by freshwater inputs. Recently, the importance of rivers as sources of microplastics to the marine environment was demonstrated by Castañeda et al. (2014), who detected high concentrations of microbeads in the sediment of the Saint Lawrence river, Canada. These microbeads were suggested to originate from municipal and industrial sewage effluents.

5. Uptake and effects in marine organisms

As microplastic abundances in the environment increase, organisms inhabiting marine systems are more likely to encounter these particles. Numerous factors such as size, density, shape, charge, colour, aggregation and abundance of the plastic particles affect their potential bioavailability to a wide range of aquatic organisms (Kach and Ward, 2008; Wright et al., 2013a). The opportunity for encountering or uptake of microplastics by marine organisms is mainly attributed to two key properties of the particles: the size and density. For example, particles with a density higher than that of seawater will become available to benthic suspension and deposit feeders (as they sink to the sea floor). As the size fraction of these microplastics is similar (or even smaller) to the grain sizes of sediments, microplastics can be ingested not only by lower trophic-level organisms which capture almost anything of the appropriate size class, but also by other sediment-dwelling organisms (Moore, 2008; Wright et al., 2013a). Consequently, plastic particles may accumulate within these organisms upon ingestion, potentially resulting in direct effects caused by physical injury in the intestinal tract or even translocation to other tissues or organs. Sediment-dwelling organisms are sensitive indicator species for many kinds of naturally and anthropogenically induced disturbances, and are used worldwide as bio-indicators of ecosystem health (OSPAR, 2010; Thain et al., 2008; Van Hoey et al., 2010). Given that this paper deals with the contamination of sediments by microplastics, only species such as echinoderms, polychaetes, crustaceans, bivalves and demersal fish are considered to review the uptake of microplastics and potentially associated effects.

Uptake of microplastics by marine biota has both been investigated in organisms living in natural conditions (Table 3), as well as in laboratory trials (Table 4). Mussels, such as the blue mussel

Table 3

Microplastic ingestion in the natural environment. Origin of the species investigated is provided (BE: Belgium, NL: the Netherlands, FR: France, UK: United Kingdom, NPSG: North Pacific Subtropical Gyre, CA: Canada, DE: Germany), as well as the particle sizes targeted by the study and extraction protocol (if provided by the authors, otherwise 'unclear').

Biota	Origin	Assay	Microplastic load	Particle size	Reference
Polychaete	<i>Arenicola marina</i>	BE, NL, FR Whole organism HNO ₃ digestion Faecal analysis	1.2 ± 2.8 MP/g ww 0.3 ± 0.6 MP/g ww	>5 µm	Van Cauwenberghe et al., 2015
Crustacea	<i>Nephrops norvegicus</i>	UK Gut analysis	83% contained MP	<5 mm	Murray and Cowie, 2011
	<i>Crangon crangon</i>	BE Whole organism HNO ₃ :HClO ₄ (4:1 v:v) digestion	0.64 ± 0.53 MP/g ww	>20 µm	Devriese et al., in press
	<i>Lepas</i> spp.	NPSG Gut analysis	33.5% contained MP	>0.5 mm	Goldstein and Goodwin, 2013
Bivalve	<i>Mytilus edulis</i>	BE Whole organism HNO ₃ digestion	0.36 ± 0.07 MP/g ww	>5 µm	Van Cauwenberghe and Janssen, 2014
		BE, FR, NL Whole organism HNO ₃ digestion Faecal analysis	0.2 ± 0.3 MP/g ww 0.1 ± 0.2 MP/g ww	>5 µm	Van Cauwenberghe et al., 2015
		NL Whole organism HNO ₃ :HClO ₄ (4:1 v:v) digestion	3.5 fibres/10 g ww	>20 µm	De Witte et al., 2014
		BE groyne 2.6 fibres/10 g ww			
		BE quay 5.1 fibres/10 g ww			
		CA Whole organism H ₂ O ₂ digestion	34–178 MP/ind	>0.8 µm	Mathalon and Hill, 2014
	<i>Crassostrea gigas</i>	FR Whole organism HNO ₃ digestion	0.47 ± 0.16 MP/g ww	>5 µm	Van Cauwenberghe and Janssen, 2014
Fish	Demersal fish	UK Gut analysis	1.90 ± 0.10 MP/ind	unclear	Lusher et al., 2013
		DE Gut analysis	3.4% contained MP	unclear	Rummel, 2014
	<i>Gobio gobio</i>	FR Gut analysis	12% contained MP	unclear	Sanchez et al., 2014

Table 4

Direct effects of microplastic exposure to aquatic (benthic) organisms, demonstrated under controlled laboratory conditions. Details on the exposure conditions, i.e. exposure route, particle type and size (if provided by the authors) and exposure concentration, are provided. (UPVC: unplasticised polyvinylchloride, PS: polystyrene, HDPE: high-density polyethylene).

Biota		Exposure route	Particle type	Exposure concentration	Assay	Effect	Reference
Polychaete	<i>Arenicola marina</i>	Spiked sediment	125–150 µm UPVC	5% by weight	Energy budget	Decreases in energy budget and feeding	Wright et al., 2013b
		Spiked sediment	20–2000 µm	1.5 g MP/L	Feeding activity	Ingestion	Thompson et al., 2004
		Spiked sediment	10 µm PS	50 MP/mL	Faecal analysis	No significant effect	Van Cauwenberghe et al., 2015
			30 µm PS		Energy budget		
Crustacea	<i>Mysis</i> spp.	Spiked seawater	10 µm PS	10 MP/mL	Ingestion	Ingestion No accumulation Trophic transfer	Setälä et al., 2014
				1000 MP/mL			
				2000 MP/mL			
	<i>Carcinus maenas</i>	Pre-exposed zooplankton		10,000 MP/mL			
		Spiked seawater	8–10 µm PS	9.4 × 10 ⁵ MP/L	Tissue analysis	Retention	Watts et al., 2014
				4.0 × 10 ⁴ MP/L	Faecal analysis		
	<i>Semibalanus balanoides</i>	Spiked mussels	0.5 µm PS	4.0 × 10 ³ MP/g	Tissue analysis	Translocation	Farrell and Nelson, 2013
		Pre-exposed mussels			Trophic transfer		
Spiked seawater		20–2000 µm		1 g/L	Gut analysis	Ingestion	Thompson et al., 2004
Bivalve	<i>Nephrops norvegicus</i>	Spiked fish	5 mm PP	10 fibres/cm ³	Stomach analysis	Retention	Murray and Cowie, 2011
					Accumulation		
	<i>Mytilus edulis</i>	Spiked seawater	3.0 µm PS	15,000 MP/400 mL	Gut and hemolymph analysis	Translocation to circulatory system	Browne et al., 2008
			9.6 µm PS				
		Spiked seawater	10 µm PS	50 MP/mL	Energy budget	No significant effect	Van Cauwenberghe et al., 2015
			30 µm PS				
		Spiked seawater	90 µm PS	10 MP/mL			
			>0–80 µm HDPE	2.5 g/L	Histological and histochemical assays	Accumulation in lysosomal system and digestive cells	von Moos et al., 2012
					Inflammatory response		
					Pseudofaeces production		
<i>Crassostrea virginica</i>	Spiked seawater	30 nm PS	0.1 g/L	Feeding activity	Reduced feeding	Wegner et al., 2012	
			0.2 g/L				
			0.3 g/L				
	Spiked seawater	0.5 µm	12,000 MP/mL	Ingestion rate	(Aggregate) Ingestion	Kach and Ward, 2008	
		1 µm					
	Spiked seawater	100 nm	13,000 MP/mL	Ingestion rate	Ingestion	Ward and Kach, 2009	
Echinoderm	<i>Holothuria</i>	Spiked seawater	100 nm	13,000 MP/mL	Ingestion rate	Ingestion	Ward and Kach, 2009
	<i>Paracentrotus lividus</i>	Spiked sediment	0.25–15 mm PVC	10 g	Ingestion rate	Selective ingestion	Graham and Thompson, 2009
		0.25–1.5 mm Nylon	65 g				
		Spiked seawater	50 nm PS	2 g			
				3 µg/mL	Embryotoxicity Gene expression	Developmental defects	Della Torre et al., 2014
				25 µg/mL			

Mytilus edulis is often selected as model species as they inhabit a wide geographic range, are sedentary, and filter large volumes of water. Four recent studies confirmed the contamination of field-collected *M. edulis* with microplastics (Table 3). These studies demonstrated that mussels collected in Europe contained on average 0.2–0.5 microplastics/g wet weight (ww) (De Witte et al., 2014; Van Cauwenberghe and Janssen, 2014; Van Cauwenberghe et al., 2015), while mussels sampled in Canada revealed a much higher microplastic load (34–178 microplastics/mussel) (Mathalon and Hill, 2014). Decapod crustaceans, such as Norway lobster (*Nephrops norvegicus*) and brown shrimp (*Crangon crangon*), are opportunistic feeders and have been shown to consume plastic present in the natural environment (Table 3). A high prevalence of plastic contamination in *Nephrops* (83% of investigated individuals) was observed in the Clyde Sea area (Murray and Cowie, 2011). These *Nephrops* ingested plastic strands (attributed to fishing waste), and some individuals were contaminated with tightly tangled balls of synthetic monofilaments. Devriese et al. (in press) noticed that plastic contamination in wild *C. crangon* mainly consisted of microscopic synthetic fibres at concentrations of 0.64 ± 0.53 microplastics/g ww, while only few other types of microplastics were detected in this species. In Gooseneck barnacles (*Lepas anatifera* and *L. pacifica*) originating from the North Pacific Subtropical Gyre (NPSG), 33.5% of individuals had ingested plastic (Goldstein and Goodwin, 2013). The observed plastic contamination in this filter feeder consisted of 99% degraded fragments and 1% of monofilament. Controlled lab studies with crustaceans were based on two types of exposure routes: exposure through the surrounding matrix or exposure through contaminated/spiked food items (Table 4). Using seawater spiked with microplastics, lab exposures with barnacles (*Semibalanus balanoides*) and *Carcinus maenas* revealed uptake for both crustaceans (Thompson et al., 2004; Watts et al., 2014). In *C. maenas*, uptake of these microspheres was established through inspiration across the gills. Ingestion due to dietary exposure was established in trials with three different organisms, *C. maenas*, *Nephrops norvegicus* and mysid shrimp (Murray and Cowie, 2011; Farrell and Nelson, 2013; Setälä et al., 2014; Watts et al., 2014). *N. norvegicus* fed with plastic seeded fish revealed the presence of the spiked filaments in the lobsters' stomachs 24 h following ingestion (Murray and Cowie, 2011). Both Farrell and Nelson (2013) and Watts et al. (2014) confirmed natural trophic transfer of microplastics (0.5 μm and 8–10 μm , respectively) from mussels (*M. edulis*) to crab (*C. maenas*) using pre-exposed mussels. Crabs retained these particles for up to 14 days after ingestion (Watts et al., 2014). Trophic transfer of microplastics from zooplankton to the crustacean *Mysis relicta* was demonstrated by Setälä et al. (2014) in a laboratory setting using zooplankton pre-exposed to 10 μm spheres (Table 4). After three hours, examination of *M. relicta* showed a 100% prevalence of microplastics in the animals' intestine. However, exposure of another mysid species (*Mysis mixta*) to contaminated prey did not result in microplastic transfer (Setälä et al., 2014). Levels of microplastics in five different demersal fish species from the English Channel were evaluated by Lusher et al. (2013). Overall, 35% of fish were contaminated with plastic, representing an average environmental microplastic load of 1.90 ± 0.10 particles per individual (Table 3). The ingested plastic consisted primarily of fibres, with the most common size class being 1–2 mm. Microplastics ingestion by wild gudgeons (*Gobio gobio*) from French rivers was also demonstrated (Sanchez et al., 2014).

The blue mussel *Mytilus edulis* is by far the species used most to study microplastics effect studies. Given that *M. edulis* living in natural habitat takes up microplastics, a number of lab trials have been performed to assess the potential adverse effects of microplastics uptake (Table 4). These, often, short-term effect assays are

typically conducted with a single type and/or size of plastic at particle concentrations much higher than realistic environmental levels. Wegner et al. (2012) demonstrated the increased production of pseudofaeces and reduced filter activity after exposure to 30 nm nanopolystyrene (0.1, 0.2 and 0.3 g/L), which according to the authors may lead to increases in the energy expenditure and reduce the organism's food uptake at long term exposure. However, no significant reduction in feeding activity or decrease in energy budget were demonstrated by Browne et al. (2008) and Van Cauwenberghe et al. (2015). Von Moos and co-workers (2012) did observed adverse effects, such as a strong inflammatory response, induced by the uptake of small plastic particles after only 3 h of exposure (Table 4). Short-term exposure experiments with small polystyrene (PS) spheres (3.0 μm and 9.6 μm ; 1.5×10^4 particles/400 mL) and HDPE spheres (>80 μm ; 2.5 g/L) revealed their translocation (especially of smaller microspheres) from the digestive tract to the circulatory system and digestive cells of *M. edulis* (Browne et al., 2008; Von Moos et al., 2012). Translocation of microplastics after ingestion was also demonstrated for the crab *Carcinus maenas* (Farrell and Nelson, 2013). Using pre-exposed mussels, this study demonstrated the translocation of small microplastics to the hemolymph of the crabs after indirect exposure, i.e. exposure through contaminated prey. In a similar setup, however, Watts et al. (2014) did find any indications of translocation to the hemolymph in exposed crabs. An important sediment-associated marine organisms that has been the subject of several microplastic effect assessments is the lugworm *Arenicola marina* (Table 4). In a short-term exposure (14 days) experiment, lugworms were exposed to sediment spiked with 10 μm , 30 μm and 90 μm PS spheres (total concentration of 100 particles.g⁻¹ sediment). While these short-term exposure did not demonstrate a significant effect on the energy metabolism (Van Cauwenberghe et al., 2015), mid-term trials (28 days) revealed clear severe effects (Wright et al., 2013b). After 28 days of exposure to 5% by weight unplasticised PVC (mean diameter 130 μm), a significant decrease in body weight and a significant reduction of the feeding activity was observed, which was ultimately reflected by a depletion of up to 50% of the energy reserves (Wright et al., 2013b).

Regrettably, due to the lack of consistency in the assays used and technical challenges (e.g. difficulties in dissecting invertebrates), environmental levels of microplastics in invertebrate organisms are difficult to interpret. As a result, intra- and interspecies evaluation is very difficult. The most common discrepancies can be related to the organ or tissues examined, the extraction protocol (e.g. digestion of tissues), the risk of airborne contamination (Woodall et al., 2015), the particle size range assessed, the reporting unit and the identification of plastics (Song et al., 2015). For example, hot acid digestion using HNO₃ (69%) was proposed by Claessens et al. (2013), while an adaptation using a 4:1 (v:v) mixture of nitric acid (65% HNO₃) and perchloric acid (68% HClO₄) was used by Devriese et al. (in press). Furthermore, Mathalon and Hill (2014) used an oxidizing agent (30% H₂O₂) to remove animal tissue. Besides the digestion assay, the particle retention of the used filters to filtrate the digest outlines the observed particle size range. For this reason Mathalon and Hill (2014) assessed microplastics >0.8 μm , Van Cauwenberghe and Janssen (2014) >5 μm , while De Witte et al. (2014) evaluated microplastics >20 μm .

The published microplastics effect assessments are typically conducted with only one type or size of plastic (mostly microspheres) at particle concentrations much higher than the environmental levels. Strikingly, all the lab trials are based on short- to mid-term (hours to 28 days) exposure to unrealistically high concentrations. These papers revealed a range of effects exhibited ingestion by a number of species, e.g. decrease of energy reserves, inhibition or reduction of feeding/filtering activity, translocation

to the circulatory system, inflammatory response and developmental defects. A few papers observed trophic transfer of microplastics and suggest an impact on the food web. Although more research is needed to determine whether plastics of any dimensions can be transferred through the food chain, translocation effects do suggest that particle size really matters. For evaluating the environmental risk of microplastics knowledge is required on the environmental levels and types of plastic, the translocation size limit and the relevant biological endpoints. Additionally, long-term exposures under controlled conditions with environmentally relevant microplastics concentrations and types are needed to allow a realistic assessment of potential microplastic-associated risks.

5.1. Indirect effects

Due to their specific characteristics, microplastics not only pose a direct threat to (marine) organisms, but they are also believed to have indirect effects on organisms. We define indirect effects as an effect caused when microplastics act as a vector for either chemicals (i.e. chemical threat) or bacteria (i.e. bacterial threat).

The chemical threat of microplastic is complex and works at different levels. Plastic polymers, owing to their large size, are considered to be biochemically inert. However, as polymerization reactions are rarely complete, residual monomers can still be found in the polymer matrix. Residual monomer content of a plastic can vary from 0.0001% to 4% (Araújo et al., 2002). These monomers can leach out of the polymeric material and, as some of these are considered toxic (including carcinogenic and mutagenic effects), they can pose a threat to the environment. This effect can be estimated based on the monomer hazard ranking as described by Lithner et al. (2011). Most hazardous polymers belong to the families of polyurethanes, polyvinyl chloride and styrene, amongst others (Lithner, 2011). Additional toxic effects of microplastics can also be caused by the wide array of plastic additives added during plastic manufacturing. Examples are the initiators, catalysts and solvents, all of which are added to obtain specific features of the final polymer. But also antimicrobial agents, such as Triclosan, plasticisers, flame retardants (PBDEs), pigments and fillers are used in the compounding of plastic. All these non-polymeric components are of low molecular weight and therefore able to migrate or diffuse from the plastic polymer, potentially causing (adverse) effects (Crompton, 2007).

This migration behaviour is similar for chemical contaminants (POPs) adsorbed on microplastics. It is known that a plethora of persistent organic pollutants (POPs) can sorb from the environment (i.e. seawater and sediment) on/in the plastic matrix of (micro) plastics. The presence of such POPs on marine plastics (especially industrial resin pellets) has been demonstrated for a wide variety of chemicals and for different geographic areas (e.g. Mato et al., 2001; Endo et al., 2005; Hirai et al., 2011; Bakir et al., 2014) (see Table 1 for additional studies on contaminant assessment on industrial resin pellets). These contaminants have a greater affinity for the plastic matrix than the surrounding seawater leading to an accumulation onto the plastic particle. This accumulation was found to be up to one million times higher in some cases (Hirai et al., 2011). Polymer type plays an important role in this contamination accumulation: under identical sorption conditions, polychlorinated biphenyls (PCBs) and polycyclic aromatic hydrocarbons (PAHs) are consistently found in a higher concentration on high-density polyethylene (HDPE), low-density polyethylene (LDPE) and polypropylene (PP), compared to polyethylene terephthalate (PET) and polyvinyl chloride (PVC), while phenanthrene sorbs more to PE than PP or PVC (Bakir et al., 2012; Rochman et al., 2013; Bakir et al., 2014). As a result, possible effects of both the polymer and

associated contaminants have to be considered when assessing the potential risks of microplastics.

Although frequently suggested, the evidence to support this chemical threat is rather limited. So far controlled lab exposures have been performed with the lugworm (*Arenicola marina*), the model organism for deposit feeders. Exposure of *A. marina* to PCB-loaded microplastics (at a dose of 7.4% microplastics by dry weight) showed an effect on feeding activity, resulting in weight loss (Besseling et al., 2013). Browne et al. (2013) demonstrated a decreased phagocytic activity by over 60% in *A. marina* exposed to sand with 5% microplastic (PVC, 230 µm) presorbed with nonylphenol. However, no such effect was reported for phenanthrene. While nonylphenol and phenanthrene desorbed from the PVC particles and transferred to the animals' tissue, the lugworms accumulated >250% more of these contaminants when exposed to contaminated sand (Browne et al., 2013).

The bioaccumulation of persistent organic pollutants (POPs) has been theoretically investigated by Gouin et al. (2011) and Koelmans et al. (2013) using a modelling approach. Both studies suggested that microplastics are only of minor importance as vectors of POPs to organisms. Koelmans et al. (2013) even predicted a decrease in contaminant body burden due to a cleaning mechanism of strong sorbent plastics, counteracting biomagnification. In a similar modelling exercise, Koelmans et al. (2014) investigated the leaching of plastic associated chemicals, i.e. additives, to marine organisms. The rationale behind this modelling approach is the fact that for additives plastic ingestion by marine organisms may be more relevant than for diffusely spread POPs as the microplastics act as a source of the additives (Koelmans et al., 2014). The results showed that ingestion of microplastics can be considered a substantial pathway for additive exposure. It is clear that further research on this topic is essential to fully understand the impact of sorbed and plastic-associated contaminants on marine organisms, and by extension the entire marine and human food web. So far, studies to assess the transfer of (environmentally relevant concentrations of) chemicals, both pollutants and additives, have not been performed on resident organisms, clearly indicating that this is an area that needs further research.

The bacterial threat of marine litter and by extension microplastics arises from the fact that they represent new habitats in the marine environment and, as such, can serve as a substrate for (micro)biological interactions. Microplastics have a hydrophobic surface that stimulates rapid biofilm formation (Zettler, 2013). So far, conventional microbial identification methodologies require a bacterial cultivation step using a growth medium, has hampered the full characterization of the microbial biofilm due to the 'great plate count anomaly' (Staley and Konopka, 1985). This term has been used to describe the fact that the majority (99–99.9%) of cells within an environmental sample are not recoverable in pure culture using classical microbial plating. However, the recent breakthrough of 'Next Generation Sequencing' technology allows for the full characterization of complex microbial samples without a cultivation step. This was nicely demonstrated in the pioneering work of Zettler et al. (2013) and will contribute significantly to biofilm characterization.

Due to their persistence in nature, (micro)plastics exhibit a longer half life than other marine substrates, creating an interesting habitat for microorganisms. For fouling, microbial biofilm formation is the initial step (Dobretsov, 2010), while in the consecutive steps so-called epiplastic organism like diatoms, ciliates and a wide variety of other organisms will attach on the formed biofilm (Reisser et al., 2014a). The formation of these biofilms on microplastics is of concern as they might contain human or animal pathogens that could potentially endanger animal and human health, and impact economic activities. Additionally, the nutritional

value of these biofilms might encourage grazing and ingestion of the covered microplastics (Reisser et al., 2014b). Associated impacts hence include food web effects.

In 2011, Harrison et al. published a call for research, calling upon the scientific community to investigate the colonisation, taxonomic composition and functional potential of microplastic-associated biofilms, in order to understand ecological implications and to develop management measures to safeguard marine life. As a response, research started focussing on characterizing the microbial assemblages of floating microplastic particles (Carson et al., 2013b; Reisser et al., 2014a; Zettler et al., 2013). Carson et al. (2013) investigated the biofilms of 100 particles (1–10 mm) collected in the Northeast Pacific, and determined that plastic type appeared to influence bacteria abundance. Zettler et al. (2013) discovered that microplastic-associated communities differ significantly from those in the surrounding seawater. For example, several hydrocarbon degraders were detected on the plastic but not in the seawater, indicating microorganisms may possibly play a role in plastic degradation (Zettler et al., 2013). Colonisation and biofilm characterisation of microplastics in marine sediments has been far less investigated. Harrison et al. (2014) used a coastal sediment microcosm and demonstrated that bacteria in marine sediments rapidly colonise low density polyethylene (LDPE) microplastic fragments (5 mm). As was the case for seawater (Zettler et al., 2013), the bacterial communities detected on the plastic were significantly different from those in the surrounding sediment. Interestingly, the dominant taxa (*Acrobacter* and *Colwellia* spp.) on microplastics from sediments were not found to be present on floating microplastics, indicating that distinct biofilms are likely to occur between different marine habitat types (Harrison et al., 2014).

6. Conclusions and outlook

Important advances have been made with respect to our understanding of the occurrence and impacts of microplastics in marine environment. However, as this research field is rapidly evolving, especially in the last 10 years as is reflected in the exponential growth of peer-reviewed publications, several issues can be identified regarding the nomenclature and classification of microplastics and applied methodologies and techniques. The current lack in standardisation and harmonisation greatly hampers the inter-study comparison and data transfer, not only for reported abundances but also for (measured) effects and impacts. We therefore recommend the implementation of an unequivocal size-based definition for microplastics, based on both upper and lower size limits, and a uniform nomenclature. Also practical issues concerning the assessment of occurrence and effects should be addressed and standardised. Today, a plethora of sampling, extraction and identification techniques are in use. An important point of interest is the harmonisation of extraction techniques. While the majority of extraction techniques are based on the same principle, i.e. density separation, a wide assortment of variations on this principle exist. Some are more efficient in extracting different types of microplastics (i.e. differences in density) than others, but in some cases this comes at an extra cost. It is clear that a standard extraction technique, especially for monitoring purposes, should be adopted by the research and regulatory community. In general, depending on the research question addressed, sampling strategies will differ. Yet, by reporting the complete set of sampling details (i.e. sampling depth, sediment weight or volume, but also sediment density, water content, etc.) differences between sampling techniques can be bypassed, and inter-study comparison facilitated. As such, this proposed harmonisation will assist future, uniform microplastic abundance assessments, and allow science-based geographical comparison and time trend assessments.

A general conclusion regarding the assessment of potential (adverse) effects following microplastic uptake in marine organisms concerns the experimental set-up of such experiments. In general, experimental microplastic concentrations are several orders of magnitude higher than current environmental concentrations, and all lab trial exposure periods are short-to mid term. While such approaches are often performed using 'proof of principle' experiments, and deemed necessary to assess the importance of this type of pollution, testing at high, environmentally unrealistic, concentrations does not provide any information on the current adverse effects on or risks to marine ecosystems. Future effect assessments of microplastics should therefore focus on mimicking more 'natural' exposure conditions. More specifically, there is a need for more long-term exposure assessments of environmentally relevant concentrations of naturally occurring assemblages of microplastics (i.e. different sizes, shapes and types).

The chemical threat of microplastics has been studied elaborately in the past years, raising some concerns. While adverse biological effects have been measured in the lab, some studies suggest (small) microplastics play only a minor part in the total body load of such environmental contaminants in marine organisms. While there is a growing body of literature regarding pollutants on microplastics, additives, or plastic-associated chemicals, are far less studied. Yet, due to the lower concentrations of these additives in the environment, transfer of these chemicals from microplastics to organisms might be more relevant than the sorbed chemicals. However, it is clear that further research on this topic is essential to fully understand the impact of sorbed and plastic-associated contaminants on marine organisms, and by extension the entire marine, and human, food web.

A far less researched potential threat of microplastics, is the presence and transfer of (potentially harmful) marine microorganisms associated with these plastics. To date, only limited literature is available on microplastic biofilm characterisation. We need to understand the colonisation dynamics and taxonomic composition (more specifically the presence and transport of pathogens and other harmful species) to properly assess the ecological implications, as these organisms could result in ecological and economical consequences to the marine food webs and human health.

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