Microstructure Characterization of Oceanic Polyethylene Debris

To cite this version:
Laura Rowenczyk, Alexandre Dazzi, Ariane Deniset-Besseau, Victoria Beltran, Dominique Goudounèche, et al.. Microstructure Characterization of Oceanic Polyethylene Debris. Environmental Science and Technology, American Chemical Society, 2020, 54 (7), pp.4102-4109. 10.1021/acs.est.9b07061. hal-02990067

HAL Id: hal-02990067
https://hal.archives-ouvertes.fr/hal-02990067
Submitted on 3 Dec 2020

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Microstructure characterization of oceanic polyethylene debris

Laura Rowenczyk\textsuperscript{1}, Alexandre Dazzi\textsuperscript{2}, Ariane Deniset-Besseau\textsuperscript{2}, Victoria Beltran\textsuperscript{3}, Dominique Goudounèche\textsuperscript{4}, Pascal Wong-Wah-Chung\textsuperscript{5}, Olivier Boyron\textsuperscript{6}, Matthieu George\textsuperscript{7}, Pascale Fabre\textsuperscript{7}, Clément Roux\textsuperscript{1}, Anne Françoise Mingotaud\textsuperscript{1} and Alexandra ter Halle\textsuperscript{1*}

1. Laboratoire des IMRCP, Université de Toulouse, CNRS UMR 5623, Université Paul Sabatier, 118 route de Narbonne 31062 Toulouse Cedex 9, France
2. Laboratoire de Chimie Physique (LCP), CNRS UMR 8000, Univ. of Paris-Sud, Université Paris-Saclay, Orsay, France
3. IPANEMA, CNRS, Ministère de la Culture, UVSQ, USR3461, Université Paris-Saclay, F-91192 Gif-sur-Yvette, France
4. CMEAB, IFRBMT, Université de Toulouse, 133 route de Narbonne, Toulouse, France
5. Aix Marseille Univ, CNRS, LCE, Marseille, France
6. Université de Lyon, CPE Lyon, CNRS, UMR 5265, Laboratoire de Chimie Catalyse Polymères et Procédés (C2P2), Villeurbanne, France
7. Laboratoire Charles Coulomb (L2C), Univ Montpellier, CNRS, Montpellier, France.

*Corresponding author

KEYWORDS: polymer, nanoplastic, microplastic photodegradation, plastic weathering

ABSTRACT

Plastic pollution has become a worldwide concern. It was demonstrated that plastic breaks down to nanoscale particles in the environment, forming so-called nanoplastics. It is important to understand their ecological impact, but their structure is not elucidated. In this original work, we characterize the microstructure of oceanic polyethylene debris and compare them to the non-weathered objects. Cross-sections are analysed by several emergent mapping techniques. We highlight deep modifications of the debris within a layer a few hundred microns thick. The most
intense modifications are macromolecule oxidation and a considerable decrease in the molecular weight. The adsorption of organic pollutants and trace metals is also confined to this outer layer. Fragmentation of the oxidized layer of the plastic debris is the most likely source of nanoplastics. Consequently nanoplastic chemical nature differ greatly from plastics.

**Introduction**

The scientific community has defined plastic pollution as a major worldwide concern. Indeed, since the introduction of plastic in the 1950s, 6 300 million tons of plastic waste has been generated, a very large proportion of which has accumulated in landfills or the natural environment\(^1\). In addition to large microplastics (MPs, 1–5 mm), smaller plastic particles at the micrometre (µPs)\(^2-4\) and nanometre (nanoplastics, NPs) scales\(^5,7\) in the environment have been highlighted by recent studies. Because µPs and NPs exhibit very specific physico-chemical properties and reactivities, the evaluation of their potential toxicological impact require specific investigations\(^8\). Primary MPs and µPs are defined as particles purposely manufactured at this scale, such as beads, fibres and pellets\(^9\). Secondary MPs and µPs result from the breaking down or erosion of larger objects. These mechanisms already occur early in the life of the object\(^10\). Primary NPs is still very anecdotic because there is no manufacturing at large scale, NPs mainly result from the degradation of macroscopic plastic objects\(^6\). Under laboratory conditions, the erosion of MPs presenting an advanced stage of weathering favours the formation of NPs\(^11\). A major source of plastic pollution is the mismanagement of municipal waste\(^12\), and fragmentation of these materials into micro- and nanosized particles is certainly a major degradation path\(^13\).

Plastic degradation involves hydrolysis, mechanical abrasion, thermal degradation, biodegradation and/or photodegradation\(^10,14,15\). Commonly used plastics are mostly prone to photodegradation\(^16\). Plastic photodegradation leads to structural modification of the polymer
backbone, such as oxidation with the formation of carbonyl groups\textsuperscript{17}, chain scission, radical recombination and crosslinking\textsuperscript{18}. Polymer photodegradation also leads to morphological alterations; the macromolecules can reorganize, and the crystallinity of the plastic often increases\textsuperscript{19}. These physical transformations impact the mechanical properties of the material and favour breakdown and embrittlement\textsuperscript{20}. Cracking, surface erosion and abrasion lead to the formation of μPs and NPs. Changes in the bulk properties of plastic in the ocean have been addressed\textsuperscript{21-23}, but the surface microstructure of plastic debris has not yet been thoroughly investigated.

The aim of this work is to provide new insights into modifications of the polymer microstructure upon weathering in the ocean and to elucidate the molecular structure of weathered plastics. Two oceanic plastic debris in polyethylene (PE) were selected because PE is the most commonly detected polymer in oceans\textsuperscript{29,35}. The methodology of this study could be extended to investigate the microstructure of other weathered plastic such as polystyrene or polypropylene. The physico-chemical properties of the debris were compared to those of the original boxes, which allowed us to directly evaluate the impact of oceanic weathering. Cutting-edge technologies were deployed to precisely describe cross-sections and transverse sections of the material: micro-Fourier transform infrared (µFTIR) spectroscopy, infrared spectroscopy coupled to atomic force microscopy (nano-AFMIR) to measure the material oxidation, fluorescence microscopy to monitor the adsorption of organic compounds and scanning/transmission electron microscopies coupled with energy dispersive X-ray spectroscopy (EDS-SEM/TEM) to monitor the adsorption of inorganic compounds. As μPs and NPs are generated from weathered macroscopic plastic debris, this study provides essential information for the comprehension of the physico-chemical nature of μPs and NPs.

**Materials and methods**
Sample collection

Plastic debris was collected by boat in the North Atlantic sub-tropical gyre in June 2015 during the French 7th Continent Expedition sea campaign. Two of eight plastic debris samples of PE were selected here for the microstructure investigation. We selected a box of coffee and a box of cocoa powder and purchased new boxes for comparison (Figure SI 1).

Surface and bulk characterization

Characterizations by ATR-FTIR and measurements by DSC and SEC results were published in Halle et al. The PAH contents are presented in Bouhroum et al., and the metal levels are reported in Prunier et al.

Fourier transform infrared (FTIR) spectroscopies

The μ-FTIR mappings were obtained in transmission mode using a Hyperion 3000 (BRUKER) equipped with a focal plan array (FPA) detector. Background and sample spectra were acquired using 256 scans at a spectral resolution of 4 cm⁻¹. Ten-micrometre-thick cross-sections were sealed in KBr pellets prior to observations. OPUS software allowed the CI maps to be drawn with the integration of the carbonyl absorption bands in the 1780-1660 cm⁻¹ region and the methylene absorption band in the 1490 to 1420 cm⁻¹ region. Spectra with high spectral and spatial resolution were obtained by AFMIR using a spectroscope. The spectra were generated with Analysis Studio software without any correction.

AFM characterization

Small-scale surface topographies were acquired by AFM using a Nanoscope V (Bruker) in contact mode.
Cross-section preparations

Cross-sections of the plastic samples were cut using a microtome with a diamond blade. Different thicknesses were obtained depending on the characterization technique. Cross-sections of 10 µm were used for the μ-FTIR and light and fluorescence microscopies. EDS-TEM required a thinner cut, and cross-sections of 150 nm were made specifically for this analysis. For AFMIR, the surface of the section was basically flattened using the microtome, and the measurements were performed directly on the plastic section. For EDS-MEB, the section was obtained by cryofracture.

Light and fluorescence microscopic observations

Light and fluorescence microscopies were performed using a digital slide imager NanoZoomer 2.0R.S (HAMAMATSU) equipped with an LX2000 200 W Ultrahigh-pressure mercury lamp. Observations were made at different excitation/emission wavelengths (λ<sub>em</sub>/λ<sub>ex</sub>): 359/461 nm, 495/519 nm, 552/578 nm 554/566 and 649/666 nm. The images were corrected with NDP view software to obtain a gain of 1.8 for the fluorescence observation.

EDS-SEM and EDS-TEM

TEM and scanning transmission electron microscopy (STEM) studies were performed using a JEOL cold-FEG JEM-ARM200F operated at 200 kV equipped with a probe Cs corrector reaching a spatial resolution of 0.078 nm. EDX spectra were recorded on a JEOL CENTURIO SDD detector.

Results and discussion

Mesoplastics were collected in the North Atlantic sub-tropical gyre during the 7<sup>th</sup> Continent Expedition. Two debris were easily identified as cocoa and coffee powder packaging items
named $M_1$ and $M_2$, respectively (Figure SI1). They were compared to two non-weathered original items (named $I_1$ and $I_2$, respectively). Most observations obtained for one pair of items (new and weathered) followed the same trend as those obtained for the other pair. Therefore, for clarity, we discuss the results for $M_2$ and $I_2$ in the text, and $M_1$ and $I_1$ data are presented in the Supplementary Information files.

**Characterization of plastic bulks and surfaces**

Table 1 summarizes the bulk and surface characteristics. For bulk characterization, we observed no significant alteration of the melting points and crystallinity between the mesoplastics and the original items. Polymer chain length is described by the number average molar mass ($M_n$) and the weight average molar mass ($M_w$), which can be measured by size exclusion chromatography (SEC). The $M_n$ and $M_w$ values were both significantly smaller for $M_2$ compared to $I_2$: the $M_n$ value was reduced by a factor 2.

![AFM characterization of the new coffee box ($I_2$) and the corresponding weathered object ($M_2$). The root mean square roughness values for the original box and the weathered object were 323 nm and 623 nm, respectively.](image)

For surface characterization, the mesoplastics presented a significantly higher carbonyl index than the non-weathered item. Characterization of the plastic surface by atomic force microscopy
(AFM) shows that there is a significant increase in the root mean square (RMS) roughness by at least a factor 2 (on 40x40 µm² areas, Figure 1 and Figure SI2). This roughening is induced by the appearance of micro-sized peaks at a scale that is typical of semi-crystalline arrangement²⁶. We also observe that the thickness of the mesoplastic is lower than that of the original box (reduction of 15 to 20%, Figure 2 and 4). A reduction in material thickness during production to safe material resources may explain the difference in thickness but it can also be envisaged that this thickness reduction is due to erosion; as it is observed upon polymer photodegradation under controlled conditions²⁷.

The total level of polycyclic aromatic hydrocarbons (ΣPAHs) in the plastic samples were below the detection limit for original items (Table 1), whereas mesoplastics contained measurable levels of ΣPAHs in the range of ng/g. This finding is consistent with literature data²⁸. The level in M₂ was seven times higher than that in M₁. The important variations between ΣPAH levels in plastic debris sampled in the same location have previously been noted²⁹.

For metal concentrations, the focus was made on four metals found in noticeable amounts: Fe, Pb, Ti and Cr. A detailed analysis of metal concentrations in the sample collected during this campaign is given elsewhere²⁵. The presence of metals in the original items is because some metals are incorporated during the manufacturing process (such as Ti and Fe) to improve plastic properties. All metal concentrations were systematically higher in the mesoplastics than in the original items because of sorption phenomena. The most explicit results were observed for Pb and Cr, as their quantities were 55 000 and 1 100 times higher in M₁ than in I₁, respectively. Large variability from one sample to another was also observed³⁰, and this variability is difficult to rationalize.

Table 1: Characterizations of bulk material by a) attenuated total reflectance FTIR, b) calorimetry
(DSC), and c) SEC; d) total level of polycyclic aromatic hydrocarbons measured by HPLC-FLD (sum of eight PAH levels); and e) metal concentrations determined by ICP-MS

<table>
<thead>
<tr>
<th>MATERIAL PARAMETERS</th>
<th>I₁</th>
<th>M₁</th>
<th>I₂</th>
<th>M₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbonyl Indexᵃ</td>
<td>0.1</td>
<td>0.7</td>
<td>0</td>
<td>0.3</td>
</tr>
<tr>
<td>Melting pointᵇ</td>
<td>141.8</td>
<td>135.7</td>
<td>142</td>
<td>142.7</td>
</tr>
<tr>
<td>Endset (°C)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Crystallinityᵇ (%).</td>
<td>41</td>
<td>39</td>
<td>43</td>
<td>42</td>
</tr>
<tr>
<td>Mₙᶜ</td>
<td>21.0</td>
<td>10.9</td>
<td>12.3</td>
<td>7.1</td>
</tr>
<tr>
<td>Mₘᶜ</td>
<td>85.5</td>
<td>80.8</td>
<td>94.0</td>
<td>67.5</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PAH/METAL CONCENTRATIONS</th>
<th>ΣPAHsᵈ</th>
<th>Fe (µg g⁻¹)</th>
<th>Pb (µg g⁻¹)</th>
<th>Ti (µg g⁻¹)</th>
<th>Cr (µg g⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(ng g⁻¹)</td>
<td>ND</td>
<td>1.1</td>
<td>0.05</td>
<td>42.5</td>
<td>0.8</td>
</tr>
<tr>
<td></td>
<td>2.2</td>
<td>19.2</td>
<td>2760.45</td>
<td>78.5</td>
<td>866.2</td>
</tr>
<tr>
<td></td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td></td>
<td>15.3</td>
<td>58.1*</td>
<td>0.02*</td>
<td>8.8*</td>
<td>3.0*</td>
</tr>
</tbody>
</table>

*Average of two values.

To summarize, surface characterization were more sensitive than bulk characterization to highlights some structural modifications of the polymer upon weathering. While calorimetry does not allow us to measure differences in the melting point or the percentage of crystallinity, the molar masses (Mₙ and Mₘ) was more sensitive parameters. Both metals and organic pollutants can be sorbed by plastic debris in large amounts.

Oxidation profile

For the microstructure investigation, cross-sections were obtained using different methods to
produce samples with thicknesses adapted to each characterization technique (Figure SI 3). μ-
FTIR mapping of the cross-sections allowed us to obtain a cartography of the carbonyl index
(Figure 2). The coffee box sample I₂ exhibited a low and uniform carbonyl index across the
whole cross-section (upper mapping in Figure 2). I₂ is not oxidized on its surfaces or at its
centre. In contrast, cross-section mapping of M₂ highlights a highly oxidized external face, with
a deep oxidation gradient from the edge towards the centre. Oxidation was observed at depths
of up to 500-600 µm within the material; one can note that the oxidation layer is heterogeneous
at the micrometre scale. The internal face appeared much less oxidized. The rate of oxygen
diffusion into the material presumably evolves with the material aging. With aging the surface
becomes more porous and could facilitate oxygen transport and further deep oxidation of the
debris.

AFMIR is a recently developed cutting-edge technique that combines the high spatial resolution
of an AFM and the chemical characterization offered by IR spectroscopy³¹. AFMIR presents a
high spatial resolution of ten nanometres. Spectra were acquired between 900 or 1350 cm⁻¹ and
1900 cm⁻¹. Punctual measurements (30 x 30 nm) were performed at three different locations
along the cross-section (external and internal faces and centre).

The AFMIR spectra of M₂ (Figure 2 A and SI4) indicated that the oxidation band was more
intense and broader than that of I₂ and was shifted to 1712 cm⁻¹. The broadening of the oxidation
band is explained by the presence of various oxidation products. As frequently reported in the
literature, photooxidation of PE generates ketones (peak at 1720 cm⁻¹) in the initial step. In
addition, secondary processes lead to the formation of carboxylic acids (1713 cm⁻¹), esters
(1735 cm⁻¹) and lactones (1780 cm⁻¹)³². The absorption band at 1641 cm⁻¹ is attributed to the
formation of double bonds during PE photodegradation. Both the oxidation and double bound
bands were detected in the three scanned regions of the macrodebris. Thus, even within the
material, the products of photodegradation are detected. This finding is in agreement with the
μFTIR mapping that indicated a deep oxidation gradient from the edge towards the centre of
the material section. Those local IR absorption spectra also gives an information about the
crystallinity of the sample. The relative intensity of the 1473-1464 cm⁻¹ peak may vary from
point to another. This might be an orientation effect generated during item production.
Furthermore, M₂ shows a shoulder at 1438 cm⁻¹ and several weak absorption bands at 1370-
1354 and 1303 cm⁻¹. Generally, in pure crystalline product, 1354 and 1303 cm⁻¹ (also called
amorphous bands³³) weak bands are absent³⁴ and the wagging mode of CH₂ at 1370 cm⁻¹ is
more intense. Figure 2 B. shows the IR response of crystalline hentriacontane for whom the
intensity the wagging mode at 1370 cm⁻¹ is largely higher than the two other bands (1354 and
1303 cm⁻¹). When comparing the AFMIR spectra of M₂ internal and external face we cannot
establish if one face is more crystalline than another.
Figure 2: On the top: profile of oxidation obtained by µ-FTIR with a representation of the carbonyl index mapping of the new coffee box I₂ (I) and the mesoplastic M₂ (II). At the bottom: A) local AFMIR spectra at the external face of the original coffee box (black) and the mesoplastic (red) B) local AFMIR spectra of hentriacontane (black), internal face of the mesoplastic (blue) and external face (red). The internal face appears less oxidized than the external face.

**Calorimetric characterization of the transverse section**

For the microstructure investigation, the materials were sectioned transversally with a thickness
of 100 µm for DSC analysis (Supplementary Table 1). Bulk DSC measurements did not highlight differences between the original items and the mesoplastics. While I₂ edges were only slightly more crystalline than the core material (1 to 2% ±1%). The heterogeneity of I₂ may be a result of the manufacturing process (during injection moulding) or may be because the box was exposed to light and already underwent slight photodegradation during storage. On the contrary M₂ external face was significantly different from the core material, while the internal face presented fewer differences (Figure 3). The degree of crystallinity of the external face increased by 6% compared to that of the core material, and the melting point decreased by 3°C. The increase in crystallinity of plastics debris is explained by two parameters²¹. First explanation is that amorphous regions are more easily degraded. The second is that polymer degradation increases the mobility of the macromolecules, which can form new crystals. Due to degradation, the molar mass of the chains is decreased, and therefore, the lamellae of the folded chain are shorter. The shorter macromolecules are more mobile and this allow them to crystallize³⁵. Consequently weathered polyethylene are more crystalline than pristine. The resultant crystals obtained with a shorter folding length are smaller than the original crystals; therefore, they exhibit a lower melting point (the lower the folding length is, the lower the corresponding melting point)³⁶-³⁸. The observed lower melting point of the resulting materiel indicates that the new crystals are of a lesser quality compare to the pristine ones. This can be linked either to the lower molecular weight of the polymer chains involved or to a very slow crystallization process. Since it is not possible to evaluate the aging time of the samples, we cannot discriminate between these two posibilities. We also noticed a significant narrowing of the melting curves for the external faces compared to the rest of the material (Figure SI 5 and 6).

SEC of the transverse section
The transverse sections were analysed by SEC (details in Supplementary Table 1). Generally speaking, polymer photodegradation has a greater effect on the longer macromolecular chains, and $M_n$ is more sensitive than $M_w$ to polymer chain scission. $M_w$ data is presented here and $M_n$ in supplementary material (Table SI 1). $I_2$ presented an inhomogeneity in the molar mass, but the variations in $M_2$ were much greater. Again, the external face of the mesoplastics presented the most intense modifications (Figure 3) with a $M_w$ value decreased by 80%. $M_2$ external face presents a molecular weight of only 20 000 g/mol.

Figure 3: On the left, percentage of crystallinity and on the right weight average molar mass ($M_w$). Data measured for the transverse sections (internal and external with a thickness of 100 µm) and the core material (thickness of 1 mm).

**Fluorescence and electronic microscopy of cross-sections**

When excited, $I_2$ emitted an intense fluorescence, particularly at an excitation wavelength of 554 nm (Figure 4). The initial fluorescence of $I_2$ is attributed to additives, probably dyes of the quinacridone family. The fluorescence emitted by $M_2$ at this excitation wavelength was much less intense, which may be because a result of the additive either migrating out of the materials or being degraded. Although typical results for plastic additive desorption under normal
conditions of use are generally known\textsuperscript{42, 43}, the desorption of additives under weathering conditions has been less described.

At 359 nm, although I\textsubscript{1} and I\textsubscript{2} were not fluorescent M\textsubscript{1} and M\textsubscript{2} (Figure 4 and SI 7) exhibited a fluorescence gradient from the external face towards the centre in the range of 200 to 400 µm. This fluorescence indicates the presence of new aromatic compounds in the material and could be attributed to PAHs that present characteristic fluorescence\textsuperscript{44-46} with an excitation wavelength between 255 and 365 nm. This signal could also be attributed to marine dissolved organic matter\textsuperscript{47} or the products of degradation of additives within the plastic. It is important to note that these fluorescent chemicals are sorbed within the first hundred microns of the materials and could easily leach, for example, in the case of ingestion, where the pH conditions could favour transfer to the organisms.
Figure 4: Microscopic observations of the new item (I₂) and the mesoplastic (M₂). On the left is a bright field image, and on the right are fluorescence images.

The atomic composition of the cross-sections was characterized by EDS-TEM. Numerous Ca and Ti ion-based particles were found in all four materials (Figure SI 8 to 10). Particles containing calcium were between 5 and 10 µm, and Ti ion-based particles were larger (200 to 500 nm). Inorganic compounds are included in the plastic formulation as additives to improve some of the material properties like CaCO₃ particles that are used as plastic fillers. These particles were randomly distributed in the carbon matrix and were also found in the mesoplastics, so there is no evidence for their leaching upon weathering. Some deep cracks
were observed, filled with sea salt crystals (Figure 5c and SI 11) Numerous nanoparticles were detected on M₂ external face (Figure 5a). The nanoparticles contained high amounts of Fe. Iron was systematically present with Mn, Ni and Zn (Figure SI 12 and 13). These observations confirm the occurrence of a recently described process\textsuperscript{25}: the formation of Fe or Mn precipitated minerals on plastic debris surfaces in the ocean. It was also demonstrated that polymer oxidation promotes metal sorption\textsuperscript{25}. It is noticeable to mention that high amounts of Pb was detected in bulk M₁ and EDS-TEM revealed the presence of elongated Pb particles with a size between 100 and 200 nm (Figure SI 14 and 15).

Figure 5: EDS-SEM observations of M₂. a: view of the whole transverse section, b: magnification of a part of the external face, c: magnification of deep cracks filled with sea salt
crystals. The multilayer structure of the polymer is elucidated by EDX-SEM (Figure SI 16) and is composed of ethylene vinyl alcohol copolymer.

As a conclusion, the microstructure investigation of ocean-weathered mesoplastics compared to original items has highlighted strong morphological, structural and molecular modifications of the outer layer of the material. We also could observe that the outer side was more altered certainly because more exposed to UV. Whereas bulk characterization did not allow to detect deep modifications. The microstructure investigation showed that the outer layers of the debris were remarkably more crystalline than the rest of the material. This outer layer consisted also of very shorten and highly oxidized polyethylene macromolecules. All these modifications are expected to eventually affect the mechanical properties of the polymer and lead to embrittlement\textsuperscript{48-50}. Embrittlement, fragmentation and delamination of the affected surface layer are most likely linked with the non-homogeneous erosion of the surface layer, leading to a strong increase in the surface roughness. The spatial distributions of oxidation and roughening are on the same scale as the polymer semi-crystalline microstructure (micrometre). This process mainly involves fragmentation into µPs and NPs, which explains the significant thickness decrease of the mesoplastics. The generation of NPs from microplastics was observed by Gigault et al.\textsuperscript{11}. This study also demonstrated the strong sorption of organic and inorganic chemicals, mostly localized on the outer layer. The field of NP investigation is new, and there is no description of the molecular structure of nanoplastics. What are NPs are made of? How are the macromolecules organized in NPs? As µPs and NPs result from fragmentation of the outer layer of macrodebris, their structure is certainly similar to that of the outer layer. The present study provides insight into the structure of µPs and NPs. We conclude that µPs and NPs are certainly composed of macromolecules that differ greatly from those composing plastics; these macromolecules are highly oxidized and significantly shorter. We also conclude that NPs
interact with species like trace metals or organic compounds originating from sorption on plastic debris surfaces. Overall, NP structural investigations must be conducted, or it will not be possible to understand their fate in oceans or their potential impact on ecosystem and human health.

ASSOCIATED CONTENT

Supporting Information Additional experimental details, including a photo of the macrodebris and the GPS coordinates were they have been collected. The characterization of M₁ and I₁ by AFM, AFMIR, calorimetry, GPC and fluorescence microscopy. Additional SEM, EDX-SEM and EDX-TEM data are also presented in the Supplementary Information in Figure SI 1 to 15 and Table SI 1.

Funding Sources

This project is supported by the Total Corporate Foundation Foundation and The French National Reaserch Program for Environmental and Occupational Health of Anses (EST/2017/1/219).

ACKNOWLEDGEMENT

We thank the 7th Continent Expedition association, as well as the staff and crew, for the sea sampling campaign.

REFERENCES


22. Gewert, B.; Plassmann, M. M.; MacLeod, M. Pathways for degradation of plastic


