From macro to micro: dataset on plastic contamination along and across a sandy tide-less coast (the Curonian Spit, the Baltic Sea)

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\textbf{Article history:}
Received 19 April 2020
Accepted 22 April 2020
Available online 30 April 2020

\textbf{Keywords:}
Beach litter
Microplastics
National park
Beach zones
Curonian Spit

\textbf{A B S T R A C T}

The contamination by macrolitter (>25 mm), mesolitter (5-25 mm), large microlitter (2-5 mm), large and small microplastics (L-MPs (2-5 mm) and S-MPs (0.5-2 mm), accordingly) in the surface beach sand at 6 locations along the 100-km-long marine coast of the Curonian Spit UNESCO National Park and the neighboring city beaches is quantified. In total, 55 samples obtained during 1-2 May 2018 are analyzed. Primary data is provided, along with exhaustive information on sampling dates and coordinates, sampling methods, extracting procedures, control measures, detection techniques, and \textit{μ}-Raman spectroscopy verification. The number of items per m\textsuperscript{2} and items per kg dry weight (for MPs) is determined separately for fibres, films, and fragments. Distributions by size and plastic type are presented. Standard protocols, a modified NOAA method, and \textit{μ}-Raman spectroscopy were applied to obtain the data, thus they can be used for comparative analyses.
**Table 1**

Dates, sampling sites locations and general characteristics.

| No | Location | Coordinates| Number of sections in stripe (5 m) | Number of beach zones |
|---|---|---|---|---|
| 1 | Lithuania | 55.67671 20.4 | 8 | 4 |
| 2 | Lithuania | 55.03030 21.03051 | 9 | 4 |
| 3 | Russia | 55.23906 35.18 | 8 | 4 |
| 4 | Russia | 55.37723 35.8 | 9 | 4 |
| 5 | Lithuania | 55.73098 30.2 | - | 4 |

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**Specifications Table**

**ESubject**

Unitary Science, Ecology

**ESpecific subject area**

Marine plastic Contamination, Environment

**Table of data**

Graph Figure

**How data were acquired**

Sand Rake method [1]; a square sampling frame (18 cm x 18 cm) and stainless steel spatula [2,3]; NOAA extraction (Zn concentration was measured using flame atomic absorption spectrometry).

**Data format**

Text file

**Description of data collection**

Macro-, meso- and microlitter, large and small microplastics extraction according to the modified Sand Rake method [1]. Spatula spans a surface area of 0.085 m² (25 mm), large and small microplastics (5-25 mm), large microlitter (2-5 mm), mesolitter (5-25 mm) and macroplastics (>25 mm) were collected from samples collected during 1-2 May 2018. The study site to this article (Fig. 1), geographic reference, and general characteristics of sampling locations and sample characteristics are presented in Table 1. The sampling scheme at every location is presented in (Fig. 2). The data of Sand Rake method [1] for macro- and meso- and large microlitter are presented in all commonly used units: number of items in a sample, number of items per square meter (items per m²), and number of items per m of the coast.

**Value of the Data**

Macro-, meso- and microlitter, large and small microplastics (MPs) contamination in surface beach sands of the Curonian Spit UNESCO National Park and the neighboring city beaches is documented. Sampling was specially designed to grasp quasi-instant “natural” plastic contamination patterns in a large area with minor anthropogenic influence.

The idea is to develop a science-based cost-effective method for monitoring of beach plastic contamination.

Data allow for comparisons of plastic contamination along and across the National Park area.

Data can be used for comparative analysis of plastic contamination in sandy beach sediments of other sandy coasts.

**1. Data**

The dataset contains information about macrolitter (>25 mm), mesolitter (5-25 mm), large microlitter (2-5 mm), large and small microplastics (L-MPs (2-5 mm) and S-MPs (0.5-2 mm) accordingly) concentration in 55 sandy beach sediments samples collected at 6 locations along the 100-km-long marine coast of the Curonian Spit UNESCO National Park (located in-between the cities of Klaipeda (Lithuania) and Zelenogradsk (Russia)) and the neighboring cities during 1-2 May 2018. The study site to this article (Fig. 1), geographic reference, and general characteristics of sampling locations and sample characteristics are presented in (Table 1). The sampling scheme at every location is presented in (Fig. 2). The data of Sand Rake method [1] for macro-, meso- and large microlitter monitoring are presented in all commonly used units: number of items in a sample, number of items per square meter (items per m²), and number of items per m of the coast.
Fig. 1. The study area in the southeastern Baltic Sea. Sampling locations are indicated by white circles, the closest villages (all located at the lagoon site) – by black circles.
Fig. 2. The sampling scheme, repeated at every location: raking for litter objects > 2 mm and sampling nearby for MPs. The raking area at different locations varied between 10 and 35 m² (see Table 1, Appendix 1). The zones of the beach and the scheme of sampling for MPs are shown: (I) the beach face, (II) the first (current) wrack line, (III) the middle part of the winter berm, and (IV) the strongest winter-storm wrack line; two replicates ca. 5 m apart were taken in every beach zone.
length (Table 2). The data of the square sampling frame method for MPs monitoring for two size classes (S-MPs (0.5-2 mm) and L-MPs (2-5 mm)) from 4 beach zones are presented in the number of items in a sample, the number of items per square meter (items per m²), and the number of items per kg dry weight (items per kg DW) (Table 3). The laboratory analysis procedures are presented in (Fig. 3). The photos of twelve selected MPs specimens extracted from the sediments are presented in (Fig. 4). The polymer types identified with Raman spectroscopy are presented in (Table 4), and the types of polymers in three groups (shapes) of MPs (in percent) are presented in (Table 5).

The dataset containing a detailed information about macro-, meso- and microlitter and large and small MPs contamination for each station in MS Excel format is provided in Supplementary Material (Appendix 1). The data on identification of S-MPs (0.5-2 mm) by μ-Raman spectroscopy are presented in Appendix 2. The polymer types, types of synthetic dyes, images of MPs, the hit ratio between the specimen spectra and reference spectra, which were identified by μ-Raman spectroscopy, are presented in Appendix 3.

2. Experimental Design, Materials, and Methods

2.1. Sediment sampling

The samples were collected at 6 locations along the 100-km-long marine coast of the Curonian Spit UNESCO National Park (located in-between the cities of Klaipeda (Lithuania) and Zenlenogradsk (Russia)) and the neighboring city beaches in the southeastern Baltic Sea during 1-2 May 2018 (Fig. 1). The sand samples for analysis of L-MPs (2-5 mm) and S-MPs (0.5-2 mm) content were collected at 6 locations along the coast (4 beach zones, in 2 replicates each), while the abundance of macrolitter (>25 mm), mesolitter (5-25 mm), and microlitter (2-5 mm) was quantified only at 5 of them, due to weather conditions. Two sampling methods were simultaneously applied: the Sand Rake method for litter larger than 2 mm [1], and the sampling frame method for MPs (see [2,3]) for MPs (0.5-5 mm). Throughout the text, we keep the exact meaning of the
Fig. 3. Analysis procedures: the modified NOAA method.
Fig. 4. Examples of MPs particles found in this study.
Table 3
Data of the square (18 cm × 18 cm) sampling frame method for total MPs, and separately for two size classes (S-MPs (0.5-2 mm) and L-MPs (2-5 mm)) from 4 beach zones (in 2 replicates): (i) the number of items in a sample (items), (ii) the number of items per square meter (items per m²), and (iii) the number of items per kg dry weight (items per kg DW).

| Location               | S-MPs | L-MPs | S-MPs | L-MPs |
|------------------------|-------|-------|-------|-------|
| Lesnoe                 | 3/1   | 3/1   | 3/2   | 3/2   |
| Zelenogradsk           | 3/1   | 3/1   | 3/2   | 3/2   |
| Klaipeda               | 3/1   | 3/1   | 3/2   | 3/2   |
| Smiltyn                | 3/1   | 3/1   | 3/2   | 3/2   |

terms for anthropogenic debris items: macro-, meso-, and microlitter include all anthropogenic items (glass, paper, ceramics, plastic, etc), while macro-, meso-, and microplastic is solely plastic.

Anthropogenic (both plastic and non-plastic) litter in the surface 3-5 cm of the beach sediments was quantified directly on-site by the modified Sand Rake method [1]. Following this method, debris was collected from the entire width of the beach (from 25 to 65 m) between
the waterline (current wrack line in Fig. 2) and the vegetation line / cliff using a metallic rake with the mesh size of 2 mm (see photo on the right-hand side of Fig. 2). The exact location of the sampling sections at the coastline was chosen randomly since wide and flattened beaches under investigation did not show evident topographic peculiarities or large litter patches. Raking was impossible at St. 6 (Zelenogradsk): sands became wet due to heavy rain. The total raked area amounts to 135 m². All the collected litter was further divided by fractions and analyzed in the laboratory.

The sand samples for analysis on MPs (0.5–5 mm) content were collected from four zones across the beach, with two replicates (about 5 m apart) in each zone (Fig. 2): the beach face, the current wrack line, the middle of the winter berm, and the wrack line left after the past storm. The sand sediments were collected from the upper 2-cm layer using a wooden square sampling frame (18 cm × 18 cm) and a clean stainless steel spatula. In total, 50 samples were collected by this method, making an integral sampled area of 1.625 m². All the sand samples were packed into new polyethylene bags with a string lock, and transported into the laboratory for further analysis.

Table 4
Polymer type and types of synthetic dyes identified using μ-Raman spectroscopy.

| Polymer type                  | Acronym | %    | Types of Synthetic Dyes (SD):                              |
|-------------------------------|---------|------|-----------------------------------------------------------|
| 1 Polyethylene                | PE      | 30.0 | Hostasol-Green G-K                                       |
| 2 Polypropylene               | PP      | 17.1 | Motoperm Blue                                             |
| 3 Polystyrene                 | PS      | 11.4 | Pigment red                                               |
| 4 Strong background fluorescence | fluorescence | 10.0 | Van Duke Brown                                            |
| 5 Low density polyethylene    | LDPE    | 8.6  | Amido Black 10B                                           |
| 6 Synthetic dyes              | SD      | 4.3  | Cobalt phthalocyanine                                     |
| 7 Cellulose/Cellulose acetate | CE/CA   | 2.9  | Astra Blue Base                                           |
| 8 Polyethylene terephthalate/Polyester | PET/PES | 2.9  |                                                            |
| 9 Plastic wax                 | Plastic wax | 2.9  |                                                            |
| 10 Polyvinyl chloride acetate | PVCA    | 2.9  |                                                            |
| 11 Nylon 6                    | Nylon   | 1.4  |                                                            |
| 12 Polymethylphenylsiloxane   | PMPS    | 1.4  |                                                            |
| 13 Polyvinyl acetate          | PVA     | 1.4  |                                                            |
| 14 Polyvinyl Butiral          | PVB     | 1.4  |                                                            |
| 15 Polivinylidene chloride    | PVDC    | 1.4  |                                                            |

Table 5
Types of polymers in three groups (shapes) of microplastics (in percent).

|                  | Percentage from items in each individual group (shape), % | Percentage of total number of: |
|------------------|----------------------------------------------------------|--------------------------------|
|                  | Fragments | Films | Fibres | Fragments | Films | Fibres |
| PE               | 31.4      | 54.5  | 16.7   | PE        | 15.7  | 8.6    |
| PP               | 22.9      | 0.0   | 16.7   | PP        | 11.4  | 0.0    |
| PS               | 8.6       | 0.0   | 20.8   | PS        | 4.3   | 0.0    |
| fluorescence     | 2.9       | 18.2  | 16.7   | fluorescence | 1.4  | 2.9    |
| LDPE             | 17.1      | 0.0   | 0.0    | LDPE      | 8.6   | 0.0    |
| SD               | 5.7       | 9.1   | 0.0    | SD        | 2.9   | 1.4    |
| CE/CA            | 2.9       | 0.0   | 4.2    | CE/CA     | 1.4   | 0.0    |
| PET/PES          | 0.0       | 0.0   | 8.3    | PET/PES   | 0.0   | 0.0    |
| Plastic wax      | 2.9       | 9.1   | 0.0    | Plastic wax | 1.4  | 1.4    |
| PVCA             | 0.0       | 0.0   | 8.3    | PVCA      | 0.0   | 2.5    |
| Nylon            | 2.9       | 0.0   | 0.0    | Nylon     | 1.4   | 0.0    |
| PMPS             | 0.0       | 9.1   | 0.0    | PMPS      | 0.0   | 1.4    |
| PVA              | 0.0       | 0.0   | 4.2    | PVA       | 0.0   | 0.0    |
| PVB              | 2.9       | 0.0   | 0.0    | PVB       | 1.4   | 0.0    |
| PVDC             | 0.0       | 0.0   | 4.2    | PVDC      | 0.0   | 1.4    |
| SUM, %           | 100       | 100   | 100    | SUM, %    | 50.0  | 15.7   |

(18.2 cm × 18.2 cm) and a clean stainless steel spatula. In total, 50 samples were collected by this method, making an integral sampled area of 1.625 m². All the sand samples were packed into new polyethylene bags with a string lock, and transported into the laboratory for further analysis.
2.2. Methods

2.2.1. Sample Preparation

Microplastics were extracted from the beach sand samples using the method employed in [4] with modifications [3,5]. Initial steps included drying, weighing and sieving the samples through the cascade of four sieves (mesh sizes of 5, 2, 1, and 0.5 mm). Visually detected MPs (as well as organic debris, amber, glass, paraffin, etc.) were removed directly from the sieves, while the residue remaining between the sieves 2 and 0.5 mm was treated using the modified NOAA method for the extraction of MPs from a sediment sample (see [2,3,5,6]), developed on the base of the NOAA recommendations [4]. It includes (I) density separation in the solution of ZnCl$_2$ (density 1.6 g mL$^{-1}$), filtering (174 μm), wet peroxide oxidation (H$_2$O$_2$ (30%) at 75 °C), calcite fraction removal by HCl solution; (II) once again - filtering (174 μm), density separation (1.6 g mL$^{-1}$), filtering (174 μm), (III) examination under a stereomicroscope (Micromed MC2 Zoom Digital) with the magnification from 10 $\times$ to 40 $\times$ directly on the surface of the filter according to [7], and (IV) MPs identification with a Raman spectrometer (Fig. 3). The extracted microparticles were classified into three generic groups: fragments, films, and fibers according to [8].

2.2.2. Analytical techniques

Larger particles were picked up, and “plastics” were identified visually, with the aid of a UV-lamp, mechanical stretching, and testing by hot needle, according to the recommendations for the microscopic determination [7]. The extracted small microparticles were optically analyzed and photographed using a stereomicroscope (Micromed MC2 Zoom Digital) with magnification from $\times$ 10 to $\times$ 40, and a UV-lamp was used when required (similar to the process described in [3]). The single operator performed all the detection and analysis procedures to exclude inter-operator variability. Raman spectroscopy was used to verify the result and attain the composition of plastic-like particles [9]. A Raman Centaur U (LTD “NanoScanTechnology”, Russia) spectrometer was used to obtain plastic spectra [10,11].

2.2.3. Contamination and quality controls

All instruments used during the extraction process were washed with distilled water and dried before the analysis. Along with usual caution to prevent the external contamination of the samples (cotton clothes, glass/metal containers, metal laboratory equipment, glass tableware), quality control measures were applied whenever possible: control white paper sheets were disposed in working space during all the time of sample handling to estimate possible contamination from laboratory air. Fifty blank samples were run to assess the level of background contamination. The numbers of fibers in controls was not statistically significant compared with MPs concentration found in samples.

Artificial reference particles (ARPs) were added to each sample prior to the extraction procedure as an additional measure to control the extraction efficiency. A detailed description of this effective method of extraction control is provided [3,6,10,11].

2.2.4. Verification by μ-Raman spectroscopy

In order to maximize the verification efficiency, the procedure of preliminary analysis and particle sorting was applied. The items for verification were selected not randomly, but as representatives for larger groups of particles, similar by their visual appearance (shapes, colours), mechanical quality (rigid, soft, elastic, foamed, etc.), and behaviour during the hot-needle test. In total, out of 5102 items (0.5–2 mm) found in sand samples, 85 items (about 2%) were selected for verification by Raman spectrometry. From them, for example, only 2 items of polystyrene foam fragments were selected out of 714 similar items, 22 coloured fibers out of 1048 similar ones, 6 out of 39 coloured films, etc. (Appendix 2).

The analysis procedure followed [10]. The polymer type and types of synthetic dyes identified using μ-Raman spectroscopy are presented in Table 4 and Table 5. In other cases, the core polymer type of some specimens was impossible to identify because of the strong signal induced by strong background fluorescence, by synthetic dyes (SD) or chemical compounds remaining on
the surface of a particle. Still, the fact of the presence of SD was considered as confirmation of the synthetic origin of a particle. So, all such specimens were accounted for as MPs (see photos in Fig. 4).

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgments

The field campaign was designed and supported within the ERA.Net RUS Plus S&T project 429 (RFBR, 18-55-76001). Laboratory facilities are maintained within the framework of the state assignment No. 0149-2019-0013. Raman spectrometry analyses were performed with the support of RSF (19-17-00041). We are thankful to Arūnas Balčiūnas for instructions on field sampling.

Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.dib.2020.105635.

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