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# Fluorogenic hyaluronan nanogels for detection of micro- and nanoplastics in water†

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Environmental pollution from plastics is exponentially increasing due to human activities. While larger microplastics can be detected with various methods, retrieving micron-sized fragments and nanoplastics remains challenging. Yet, these smaller-sized plastics have been raising considerable toxicological concern. Here, we show that a poorly emissive hyaluronan functionalized with rhodamine B (HA-RB) adheres with high affinity to various microplastic surfaces, becoming brightly emissive. Micro- and nanoplastics (MNPs) can be successfully detected with size as small as the diffraction limit of confocal microscopy (ca. 250 nm). FLIM images show that the fluorescence lifetime of the dye moieties changes according to the plastics, making possible a discrimination of the nature of MNPs based on lifetime. HA-RB, compared to previous reports, eliminates false-positive results caused by formation of dye aggregates, resulting in a higher S/N ratio which allows the unequivocal detection of nano-sized fragments.

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#### **Environmental significance**

This manuscript reports an attempt to face a great challenge of our times, the pollution of water resources by micro- and nanoplastics. Quantitatively, this problem is simply huge, in which most of the enormous global production of plastics generates plastic wastes being dispersed in the environment. The detection of nano- and micron-sized fragments represents an unmet challenge, with these smaller-sized plastics also raising considerable toxicological concern. Our strategy relies on water-dispersed nanogels based on hyaluronic acid fluorescently functionalized. This nanomaterial is in a dark state in water and adheres to microplastics becoming brightly emissive. This interaction occurs with high affinity, such that even 5 nM HA-RB provides a clear analytical signal. This functionalized biopolymer has proven biocompatibility, and it is water soluble and largely biodegradable.

#### Introduction

Plastics are materials based on polymers of synthetic or semisynthetic origin, which - since their first preparations at the beginning of the last century - have now reached a central role in contemporary society with a huge global production of around 300 million tons each year. Despite the increasing worldwide attention to proper disposal of plastics after their use - either by recycling, converting to energy or landfill their large majority (ca. 70%) accumulates in the environment, where they can persist for a long time. The presence of plastic debris in the environment poses a safety concern to human health owing to direct or indirect effects, and it undoubtedly disturbs ecology - intended as the interrelationships of organisms with their environment and with each other - under many aspects and thus represents a threat also for human supplies and life.2 The toxicological

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impacts of microplastics and of nanoplastics (overall size below 1 µm) are also associated with their sizes, raising considerable concern due to their possible direct interaction with cells. In addition, these smallest fragments are also the most difficult to detect and to eliminate. With decreasing size, plastic particles become more bioavailable to organisms: for example, micro- and nanoplastics (MNPs) have been reported in various taxa across trophic levels, such as plankton, benthic invertebrates, shrimp, bivalves, fish, seabirds.3,4 Furthermore, MNPs can be carriers of several kinds of pollutants (heavy metals, plasticizers, etc.) or covered by biofilms, increasing even more their potentially detrimental impacts on the ecosystem.

The analysis of MNPs, including the determination of their concentration, chemical nature, size distribution and morphology, remains a goal far to be fulfilled. Several analytical methods have been developed for microparticle analysis, with Fourier transform infrared (FTIR) and Raman spectroscopy representing, at present, the gold standard for microplastic detection and analysis.5,6 The analysis of nanoplastics is even more challenging, with protocols still under development. Fluorescence methods - typically

endowed with excellent versatility and high sensitivity, low cost and handiness of use<sup>7</sup> – have found only limited application in the analysis of real samples of MNPs. Fluorescence has been used to trace dye-labelled plastic particles or to recognize plastic debris with fluorescence microscopy after staining them with suitable dyes such as Nile Red.<sup>8-13</sup> This dye has been advantageously employed to stain various microplastics in water and also used for high-throughput techniques based on single particle tracking, 14 but its tendency to aggregation causes an increase of the background signal and the occurrence of false-positive features. In addition, its affinity is relatively low towards microplastics, leading to possible desorption and thus posing additional problems. 10-12 Very recently, more complex probes based on polyhedral oligomeric silsesquioxane (POSS) derivatized with a coumarin dye were reported and underwent an emission spectral change from yellow to blue upon the interaction with plastic particles of PS, PLA and PMMA in water. 15

Inspired by previous work,16 in which we observed a strong affinity between RITC-functionalized hyaluronic acid (HA-RB) and PEGylated nanoparticles, in this contribution, we investigate the onset of similar interactions between this longchain HA functionalized with RB dyes and the surface of different MNPs dispersed in water. Such an interaction provides a useful fluorescence analytical signal to highlight the presence of MNPs in water due to two overlapping effects.

First, HA-RB accumulates on the surface of MNPs driven by a very high affinity; second, the fluorescence quantum yield of the RB moieties increases owing to a hydrophobicityinduced unquenching mechanism, as witnessed by the increase of their average fluorescence lifetime. Finally, the different fluorescence lifetime observed for HA-RB adsorbed on various MNPs and, in particular, on polystyrene MNPs, can provide an absolute parameter to discriminate different types of plastics.

## Results and discussion

#### HA-RB testing for micro- and nanoplastics

HA-RB was prepared by adapting a previous synthetic method.<sup>16</sup> Briefly, HA (200-600 kDa), was covalently functionalized with rhodamine B isothiocyanate (RITC) in DMSO and then purified with dialysis (cut-off 12 kDa), yielding a dispersion of functionalized HA-RB in water with 1 RB moiety every ca. 60 HA monomers (1.7% doping degree of HA monomer units, i.e., ~18 RB dyes per HA polymer chain). The yield of RITC binding to HA was obtained by measuring the absorbance of a known amount of HA-RB in a good solvent (ethanol) and using the molar extinction coefficient of RB in the same solvent.17 Native HA has an intrinsic tendency to form aggregates in water, largely driven by the rigidity of this polyanion;<sup>18</sup> yet, the presence of RB moieties with their hydrophobic contribution - greatly enhances this tendency and further destabilizes HA-RB polymer chains, leading to the formation of aggregated nanogels in the 100-300 nm size range.16 The role of RB dyes in promoting this

aggregation in water is proved by the distinctive absorption spectrum of aggregated RB, showing a higher shoulder at 530 nm and a broader peak at 565 nm with respect to the monomeric RB dyes (see Fig. S1†). RB moieties in the HA-RB nanogels in water suffer from heavy self-quenching, displaying an average PLOY of 0.004 and a lifetime of 1.8 ns, which results in a low fluorescence background for sensing schemes both in a cuvette and in fluorescence microscopy. On the contrary, HA-RB is well soluble in ethanol (whereas native HA is not), and RB moieties here display their typical absorption spectrum and bright emission (PLQY = 0.39).

We obtained water-dispersed MNPs from two different sources: for the first sample type, lab-grade microplastics were purchased from Sigma-Aldrich; for the second type, we fragmented intact, non-degraded commercial plastic objects via ball-milling and/or with a pestle and mortar. Then, each kind of MNP - in the form of a solid powder - was dispersed in distilled water, even though the dispersions were not fully stable since a large part of the MNP fragments was separated either by precipitation or by floating on the water surface. To these heterogeneous dispersions, a small volume of HA-RB in water was added (typically 50 µL to 1 mL of the MNP dispersion), yielding a final HA-RB concentration of ~170 nM (estimated concentration of polymer chains, using 400 kDa as  $M_{\rm W}$ ). After shaking for 30 seconds, the MNP dispersions were ready for fluorescence microscopy observations, which were performed on a drop of 50 µL of the dispersion deposited on a coverglass.

As readily observed by confocal fluorescence microscopy under 561 nm excitation, after mixing with HA-RB, MNPs become brightly luminescent in the 590 ± 30 nm range. Confocal images (Fig. 1) show that the MNPs are effectively labelled and clearly distinguishable from the background both when the fragments are very large (with a few fragments reaching up to 100 microns in diameter) and when the fragments are about 1 micron in diameter (Fig. S8†). Confocal sectioning with oil immersion, high numerical aperture objective (NA = 1.45, magnification 100×) allows the confirmation that HA-RB labelling is a surface process, and

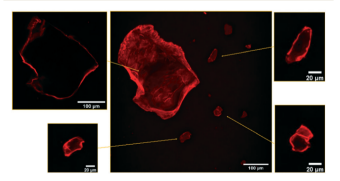


Fig. 1 Confocal fluorescence microscopy of PMMA microparticles in water, in the presence of 170 nM HA-RB. Center image: an overlap of 55 stacks spanning 66 micrometers in depth. The inset squares show selected individual Z-stacks showing the cross-sections of PMMA microparticles.

that no penetration of HA-RB in the MNPs occurs, as proven by the dark inner part of the MNPs, which in turn displays a bright fluorescent surface (Fig. 1).

Comparison of the same MNP samples treated with Nile Red (NR), a standard fluorescent dye in available fluorescence-based techniques for microplastic detection, reveals a comparable intensity onto the surface of MNPs owing to NR adsorption, but also a relevant amount of NR aggregates with different sizes, from sub-micron to several tens of micrometers, that provide "false positive" fluorescent objects, as also previously reported. 19 These NR aggregates can be distinguished from MNPs only via lifetime imaging analysis, owing to their short lifetime (Fig. S4†).

Yet, under these conditions and without any purification step, the reference sample of HA-RB in water, in the absence of MNPs, contains small (<500 nm diameter) and poorly emissive HA-RB aggregates. The presence of nanogels of HA-RB in this size range impedes the unambiguous recognition of nanoplastics of similar size, even if they do not interfere with the identification of microplastics with diameters larger than ca. 1-2 micron. To overcome this issue and allow for clear recognition also of nanoplastics with sizes smaller than 1 micron, we added a centrifugation step (5000 rpm, 10 minutes) that separates MNP fragments, while non-adsorbed HA-RB nanogels - owing to their small size, hydration and density - remain in the supernatant and are eliminated. In the absence of MNPs, with this additional purification step, we could not identify any emissive aggregate (Fig. S7†). Therefore, sub-micron emissive features in the MNP samples, when treated with this centrifugation step, can also be explicitly attributed to nanoplastics with size as small as the resolution of the microscope (ca. 250 nm).

We acquired large images of 5.5 × 5.5 mm<sup>2</sup> (multi-frame mosaic acquisitions of 10 × 10 images at a constant height from the coverglass surface, in the range 5-50 microns from the surface) of the MNP samples labelled with HA-RB. To best observe the fluorescently stained MNP samples, we immersed them in a hyaluronan hydrogel (16 mg mL<sup>-1</sup> native HA 200-600 kDa in water) rather than pure water, in order to slow down water evaporation and movement of fragments during acquisition. Native hyaluronan was selected - among other hydrogels - because it does not interfere with the photophysical properties of HA-RB and with its interaction with MNP fragments (Fig. S2†). We then analysed the confocal images with a particle tracking routine based on ImageJ, which selects particles based on an intensity threshold value taken well above the background noise (e.g., threshold = ca. 500 counts for background signal = 64 ± 12 counts, detailed description in ESI† and Fig. S7 and S8). Plotting the integrated pixel intensity versus the size (equivalent diameter) of the MNP fragment shows that, as expected, the overall fluorescence signal builds up in the fragments with increasing size (Fig. 2). The submicron part of the plots reveals that a large number of nanoplastics are brightly labelled with HA-RB and that they stand out of the background with an overall intensity which is from  $\sim 10$  to > 200 times the background noise. In a log-log

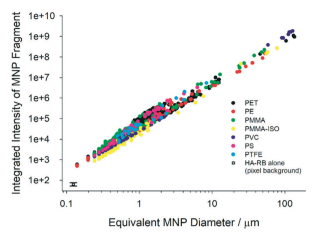


Fig. 2 Plots of integrated pixel intensity versus the size (equivalent diameter, calculated from the area of the fragments,  $d = (A/2\pi)^{0.5}$ ) of the MNP fragment.

plot, the intensity shows a clear linear trend, well fitted with a slope close to 2, as expected for a surface adsorption process. The slope, which is slightly larger than 2 for all plastic samples, suggests that microplastics feature rough surfaces with higher fractal dimension.<sup>20</sup> From the fitting, we can calculate the intensity of a nanoplastic fragment of size corresponding to a single pixel (square pixel of side 124 nm), which allows a ready comparison with the background, showing that MNPs stained with HA-RB display between 3.5 (PMMA-iso) and 7.4 times (PMMA) the average intensity of the background for 124 nm size (Table 1). We compared these results with the use of molecular dye rhodamine B (RB), which is soluble in water. RB shows good affinity towards MNP surfaces, with high luminescence from the microplastic fragments. Yet, RB is highly emissive also in water, and therefore the high signal from the solution hinders the possibility to detect the small nanoplastic fragments.

#### Fluorescence lifetime imaging of MNPs using HA-RB

The variation of the excited state lifetime of RB moieties upon interaction with MNPs was also investigated. As previously found, the lifetime of HA-RB in water is short and

Table 1 Fitting parameters of the plots shown in Fig. 2. The log-log data points have been fitted with a linear fit; the table reports the intercept (i.e., log(I) at diameter =  $10^0 \mu m$ ), slope and calculated intensity (and log(I)) at a single pixel (square of lateral size = 124 nm), which is compared with the average background intensity at a single pixel

	Intercept	Slope	Intensity $(\log I)$ at single pixel
PS	4.75	2.43	352 (2.55)
PTFE	4.67	2.25	422 (2.63)
PMMA	4.64	2.17	472 (2.67)
PMMA-iso	4.35	2.19	229 (2.36)
PVC	4.51	2.24	300 (2.48)
PE	4.54	2.12	398 (2.60)
PET	4.64	2.25	416 (2.62)
Background	_	_	$64 \pm 12 (1.81)$

multiexponential, corresponding to a broadly self-quenched state. Interestingly, the average lifetime displays relevant variations upon interaction with MNPs, pointing to a conformational change of HA-RB nanogels with a fraction of self-quenched RB moieties becoming emissive from a monomer-like state.

In particular, at low concentrations of HA-RB, the fluorescence decay of HA-RB onto MNP surfaces appears longer and, in some cases, is also more similar to a singleexponential decay with respect to the HA-RB nanogels in water (Fig. S3†). This can be explained by a hydrophobicityinduced unquenching mechanism: breaking of the RB dimers and aggregates (responsible for the initial selfquenching and for HA-RB aggregation as nanogels) occurs due to conformational changes of HA-RB upon adsorption on the MNP surface, which results in the local increase of the average lifetime, narrowing of the lifetime distribution and, consequently, in the enhancement of the fluorescence quantum yield. Yet, the conformational changes appear to be quite sensitive to the type of surface, so that an increase of the lifetime of different magnitude is observed for different types of MNPs. Therefore, it is noteworthy to underline that, under these conditions, the chemical nature of MNPs can be made evident via FLIM analysis, allowing for a straightforward distinction as shown in Fig. 3 for PS and PMMA microplastics, which can be clearly recognized even when mixed together.

The lifetime dependence on the specific plastic surface could be explained by the interplay of two different interactions: the first is the hydrophobic interaction, which causes the increase of lifetime, and which characterizes the binding of HA-RB onto the surfaces of purely hydrophobic plastics such as PTFE and PVC. The second interaction that can play a role in the affinity with MNP surfaces is the H-bond formation due to the hyaluronan functional groups of HA-RB, which may be more relevant in the binding with less hydrophobic surfaces of PMMA and PET MNPs. Compared to the other plastic polymers, PS shows a peculiar behaviour, in which electronic interactions with the styrene units may be at the basis of the short and approximately monoexponential lifetime of HA-RB.

Yet, the build-up of HA-RB layers on MNP surfaces, at higher HA-RB concentrations, changes photophysics of the RB moieties. The local intensity on the

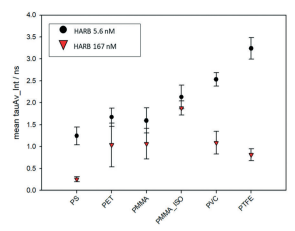


Fig. 4 Mean values with standard deviations of "\(\tau\_\text{Av\_Int"}\), i.e., the average lifetime weighted on intensity, for MNP samples incubated with 170 nM HA-RB (red triangles) and 5.6 nM HA-RB (black circles).

surface of MNPs increases, but the emission decay gradually becomes shorter and multiexponential. The additional HA-RB on the surface of MNPs indeed increases the local density of RB dyes and therefore the probability of self-quenching interactions among RB moieties is enhanced, resulting in the observed shorter and multiexponential lifetime (Fig. 4). For these reasons, a low concentration of HA-RB should be used for discriminating among the different kinds of plastics; this concentration is, however, highly sufficient to give a clear image (Fig. S3 and S6†).

The ability of HA-RB to readily stain micro- and nanoplastics even at extremely low concentration (~5 nM polymer chain and ~100 nM rhodamine B dyes, to be compared with the typical concentration used for the tests based on Nile Red, in the range 1-10 µM) can be explained by the concomitant effect of a high affinity of the HA-RB nanoprobe towards many plastic surfaces and by high PLQY due to the effective switch-ON of the luminescence of RB moieties upon adhesion of the probe onto these surfaces.

## Conclusions

In conclusion, we report here a fluorogenic material, i.e., hyaluronic acid functionalized with rhodamine B (HA-RB), with a very weak fluorescence intensity in water, but readily

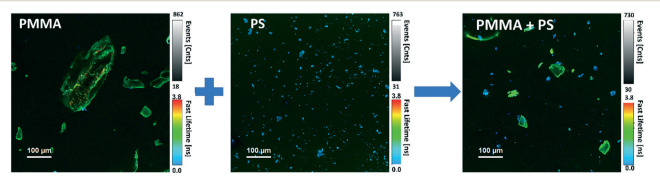


Fig. 3 FLIM microscopy images of PMMA and PS MNPs labelled with HA-RB, before and after mixing the two samples.

adsorbs onto the surface of various MNPs with a concomitant enhancement of brightness, making them fluorescent. Fluorescently stained MNPs are clearly visible with size as small as the resolution of a confocal microscope, with nanoplastic fragments of approximately 250 nm standing out of the background. This high sensitivity can be achieved due to the recovery of the fluorescence quantum yield of the RB moieties thanks to a hydrophobicity-induced unquenching mechanism together with a high affinity interaction with MNPs. These properties allow for MNP detection even at a very low concentration of the probe (~5 nM). Furthermore, the lifetime parameter reveals a different degree of hydrophobicity-induced unquenching of HA-RB on the surface of different microplastics, making possible the identification of the types of MNPs based on their fluorescence lifetime.

These results represent a feasibility study, under controlled conditions, and its applicability to real MNP debris in environmental samples remains a challenge. The intrinsic versatility of the reported fluorogenic hyaluronan, though, represents a valuable strength: in fact, various properties of fluorogenic hyaluronan nanogels can be finely tuned by changing the mass, the derivatization or even the chemistry of HA and of the fluorogenic moiety. This versatility is potentially very advantageous to design custom probes for different applications, in which sensitivity requires proper balance with selectivity towards various categories of interferents and with different experimental conditions.

## **Experimental**

#### Synthetic procedure

Synthesis of HA-RB was adapted from a previously reported procedure. 16 Briefly, HA (200-600 kDa, 30 mg) was dispersed in DMSO (8 mL) and rhodamine B isothiocyanate (RITC, 4.4 mg) was added under stirring; the reaction was left to proceed for 24 h at room temperature. The product was then diluted with water (8 mL) and unreacted RITC was eliminated via dialysis (at least 3 days at room T, regenerated cellulose, cut-off 12 kDa), to finally obtain a dispersion of functionalized HA-RB in water with 1 RB moiety every ca. 60 HA monomers (1.7% doping degree of HA monomer units, i.e., ~18 RB dyes per HA polymer chain).

#### Materials and methods

Lab-grade plastics were either purchased from Sigma Aldrich polystyrene 9003-53-6; "PET", poly (ethylene terephthalate) 25038-59-9; "PMMA\_ISO", methacrylate)-isotactic 25188-98-1; "PVC", poly(vinyl chloride, 9002-86-2; "PTFE" poly(tetrafluoroethylene) 9002-84-0; "PE", polyethylene 9002-88-4) or obtained by grinding clean and new plastic objects ("PMMA" poly(methyl methacrylate) from a ball mill-grinded disposable fluorescence cuvette). PET and PMMA show macroscopic/granular fragments and were ground with an analytical mill IKA A10 basic until a fine powder was obtained. The other microplastic samples were already in powder form and were finely ground in a mortar to further reduce mesh size and increase the fraction of nanoplastics and small microplastics.

#### Scanning electron microscopy

Scanning electron microscopy (SEM) was carried out by using a Philips 515 microscope at an accelerating voltage of 15 kV on the samples fixed with a conducting bi-adhesive tape on an aluminium stub and coated with gold. Images were analysed with ImageJ software.

#### Sample preparation for fluorescence imaging

MNP samples were weighed and incubated under mechanical agitation with 1 mL of water solution containing 170 nM HA-RB. After that, 30 µL volumes of the MNP dispersion were mixed together with a drop of 100 µL hyaluronan hydrogel (16 mg mL<sup>-1</sup>) directly on the glass slide. This solution volume assures that the hydrogel drop remains viscous during the experiments, to homogeneously disperse the sample and to delay evaporation. The same procedure has been used to incubate microplastics with Nile Red. In the second set of experiments, we added a purification procedure. Two sets of each microplastic sample were incubated in glass vials under mechanical agitation for 30 min with 1 mL of water solution of 170 nM HA-RB for one set and 5.6 nM HA-RB for the other one. After the incubation, 2 mL of water was added to each vial and then centrifuged at 5000 rpm for 10 min. This process separates MNP fragments, while non-adsorbed HA-RB nanogels remain in the supernatant and 2 mL of it is carefully eliminated with a Pasteur pipette. Then 2 mL of new water was added to each sample and centrifuged again. This washing process was repeated three times. Finally, we mixed the centrifuged MNPs with HA hydrogel for confocal and FLIM imaging.

#### Photophysical characterization

Confocal and FLIM. Confocal images were registered on a Nikon A1R microscope with an oil immersion, high NA objective (NA = 1.45, magnification 100×), an excitation laser at 561 nm and a GaAsP PMT with an emission filter at 595/50 nm. The acquisition parameters were set to 1 A.U., 8× signal average, resonant scanning mode, laser power = 1.64%, PMT GaAsP HV = 70. This setting was kept the same for all the samples analyzed. To obtain large statistic ensembles of MNPs, we acquired 10 × 10 mosaic images, with a final field of view of  $1.2 \times 1.2 \text{ mm}^2$  and a single pixel size of 124 nm.

FLIM images were acquired with a time-correlated single photon counting (TCSPC) system (PicoQuant GmbH Berlin), using a 405 nm pulsed excitation laser at a 10 MHz repetition rate, a 560 nm long-pass emission filter, a hybrid PMA detector and a PicoQuant PicoHarp 300 correlation board. FLIM images were then analysed with SymPhoTime 64, PicoQuant GmbH.

#### Image analysis

Confocal images of the different samples have been analysed with a particle tracking macro based on ImageJ commands. The macro was written to process large images with comparable conditions and to obtain relevant information on all the fragments in the area. It processes images to find particles with size greater than the pixel size (124 nm) and pixel intensity higher than a pre-set threshold. The output includes area, minimum, maximum and mean pixel intensities together with the integral of the identified area. Additional detailed information and examples on the image analysis process can be found in ESI.†

## **Author contributions**

Methodology, software and data curation: M. C. and D. G.; supervision: E. R., D. G. and N. Z.; conceptualization and supervision: E. R., D. G., N. Z. and L. P.; funding acquisition: D. G. and L. P.; writing – original draft, review & editing: all authors contributed equally.

## Conflicts of interest

There are no conflicts to declare.

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