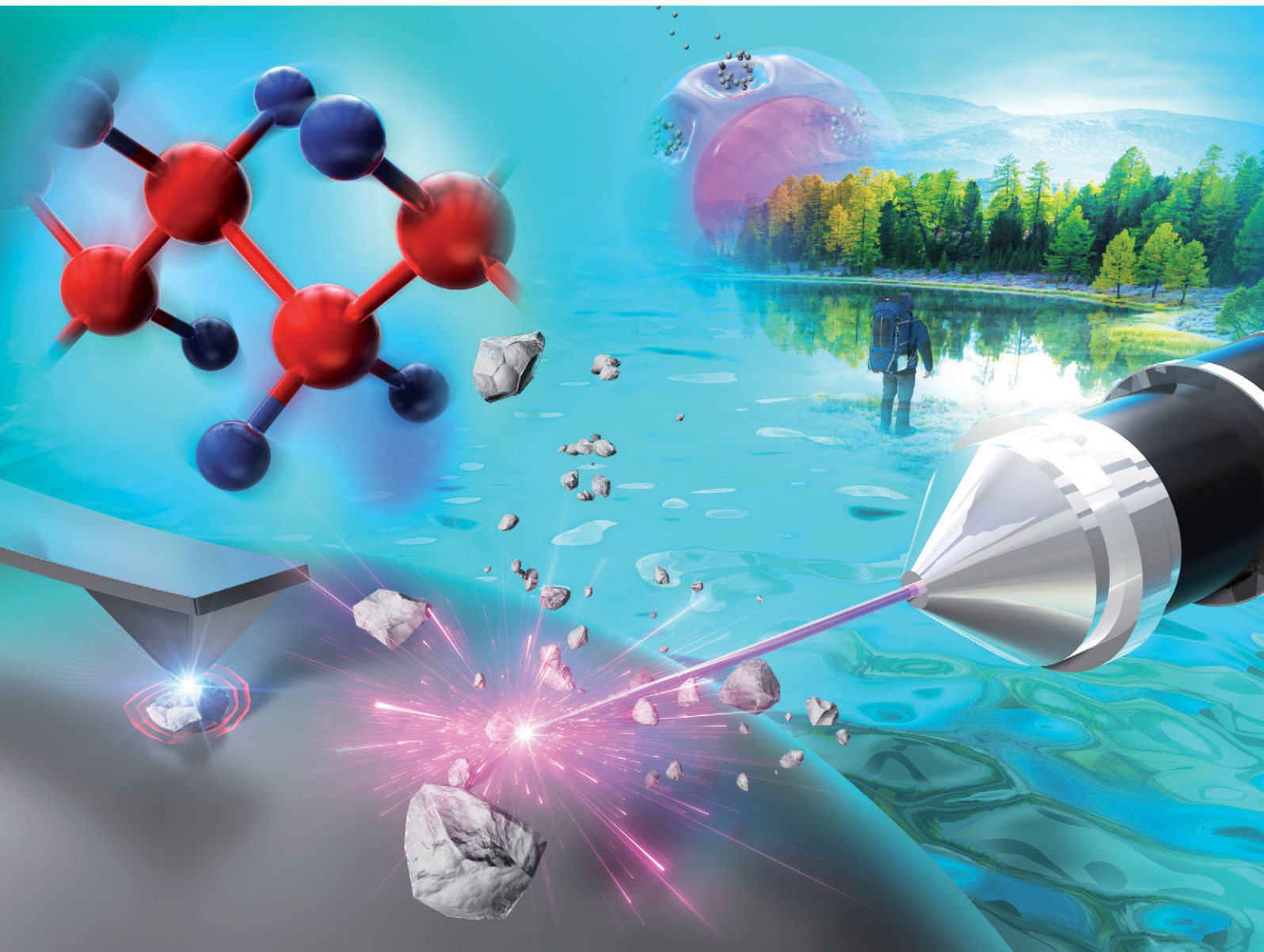


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PAPER

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Local infrared spectral measurement system for the inspection of independent nano-plastic particles in water-based solutions†

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This study proposes a new method of bubble accumulation that enables the capture of individual nanoparticles diluted in water-based solvent and the evaluation of the shape and local infrared spectra of each independent nanoparticle. We have demonstrated this system using nanoparticles of defined size generated by nano-second laser ablation. Following a process of concentration of microbubbles, we have been able to analyze the material properties of individual nanoparticles by AFM-IR. The AFM images and IR spectra results indicate the presence of independent nanoparticles, and the IR spectra showed that the particle size is considered to decrease as the oxidation reaction progresses. This system approach for the concentration and analysis of nanoparticles can particularly contribute to bio adaptation research, since the identification of the physical properties of nanoparticles can provide a better understanding of the environmental/biological effects and relationships, the mechanism of nanoparticle aggregation and the interatomic forces between particles.

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Environmental significance

Currently, micro-plastics floating in the ocean are misidentified as phytoplankton or seaweed and bioaccumulated through the food chain, potentially causing chronic health effects on apex predators. The biological effects and nanophysical properties of nano-plastics, which are much smaller in size than micro-plastics, are not well understood. It is also difficult to measure and analyze the morphology and physical properties of nano-plastics collected from nature. For low molecular weight polyethylene with nano-sized particles whose size is defined by laser ablation, this study reports on the analytical processing design to provide the accumulation of nano-plastics diluted in water-based solvents and evaluation of their individual IR spectral properties. Identifying the physical properties of nanoparticles could be one means of solving environmental problems through a better understanding of environmental and nanotoxicology and relationships, aggregation mechanisms, and interatomic forces between particles.

1 Introduction

Currently, methods for the identification of micro-plastics (MPs, 1 μm –5 mm in size)^{1,2} floating in the ocean can

misidentify these as phytoplankton or seaweed. When ingested by sea-life, such MPs can bioaccumulate through the food chain, with the potential for chronic health effects on apex predators.^{1–29} Whilst a great deal of research around the world has studied the interaction of MPs with biological systems, their toxicity, the formation mechanisms, and an understanding of the biological effects and nano-physical properties of nano-plastics (NPs, 1 nm–1 μm in size),^{2,14–16} which are much smaller in size than MPs, are not fully understood. Further research on NPs needs to be conducted as soon as possible. However, it is difficult to collect NPs dispersed in nature, and to measure and analyze their morphology and material properties. Therefore, in order to advance our understanding of biological effects in NPs, it is necessary not only to artificially create model samples of dispersed NPs,^{3–6} but also to clarify the physical and chemical properties of individual artificially created NPs.

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i) Preparation of nanoparticles: many studies have used polystyrene nano-plastics (PS-NPs, monodisperse, smooth and spherical in morphology) for applied studies of interactions with biological systems and toxicity.^{4,17,18} However, because of their different properties from other polymers, and because they differ significantly from the rough surfaces and irregular shapes of their natural counterparts, these PS-NPs lack relevance to the environment and cannot represent realistic NPs in the environment.^{4,17,18} In addition, the presence of solvent residues, surfactants, and impurities make it inappropriate to use NPs produced by liquid-phase synthesis or pulverization for applied studies of interactions with biological systems and toxicity.³ In contrast, it is becoming possible to use laser ablation to produce model samples of NPs that are artificially dispersed in nature, free of solvent residues, surfactants, and impurities. Magri (2018) *et al.* performed nano-second laser ablation (irradiation wavelength 248 nm, pulse duration 20 ns, repetition frequency 20 Hz, Coherent-CompexPro 110) on a commercial PET film (Goodfellow Cambridge Ltd) in Milli-Q water under an irradiation fluence of 4.5 J cm⁻² and with an ablation area of 4 cm². The average particle size of the generated PET-NPs was 26.7 ± 14.2 nm.⁶ It is also becoming clear that the conditions under which laser ablation is performed can control the particle size.³ Kanehara (2023) *et al.* performed nano-second laser ablation (irradiation wavelength 266 nm, pulse duration 12–15 ns, repetition frequency 10 Hz, Lotis Tii LS-2134D) on a reagent LDPE film (Sigma-Aldrich Co. LLC) in ultra-pure water under an irradiation fluence of 8.0 mJ cm⁻² for 0.5–1.5 h on a 0.023 cm² area. As a result, it was found that the longer the laser irradiation time, the smaller the particle size of NPs in the generated LDPE.³ Typical concentrations of micro/nano plastics in the environment range from a few dozen of particles m⁻³³⁰ to more than 1000 particles m⁻³,^{31,32} but the concentration of the model nanoparticles produced is only 10⁸ particles per ml,³ which is not sufficient for the particle concentration needed to measure biotoxicity and bio immunity.^{33,34}

ii) Collection: concentration by centrifugation methods is not applicable due to the possibility of condensation, transformation, and non-dispersibility of the fine particles. In addition to centrifugation methods, other NP collection methods include membrane filtration, field flow fractionation, cloud point extraction, pressurized fluid extraction, alkali-assisted thermal hydrolysis,¹⁸ and emulsions,¹⁹ but none of these methods are suitable for NPs extraction because of the difficulty of separating and identifying NPs from sand and organic contaminants²⁰ (see the ESI† for the advantages and disadvantages of each method for collecting nanoscale plastic particles (Table S1†)). Although the dilute state is not suitable for measurement, the dilute state makes it possible to evaluate a single particle. It is expected that Marangoni convection created by bubbles could be used to properly analyze fine particles.^{35–37} The study has reported that marine nano-samples are actually concentrated and accumulated at the microbubble interface by inducing Marangoni convection through the generation of microbubbles by focused laser heating at the interface between the substrate and the sample solution.³⁸

iii) Physical and chemical analysis: for the evaluation of NPs, a standard physical property evaluation such as shape observation by electron microscopy, average particle size evaluation by DLS, chemical characterization by Raman spectroscopy, and acquisition of infrared spectra of aggregated nanoparticles by FTIR are mainly studied.^{19–21} However, these physical and chemical analysis methods are limited in their ability to characterize the presence of such particles in liquid samples^{22–26} (see the ESI† for the advantages and disadvantages of each method for visual and spectral analyzing nanoscale plastic particles (Table S2 and S3†)). Ecological studies using micro- and nanoparticles of polystyrene (PS), which are easy to fabricate with uniform particle size and shape, have been conducted worldwide,^{26–29} and PET-NPs, a crystalline plastic other than PS, has only been found to pass through intestinal epithelial cells.⁶ Thus, studies on the aggregate and standard physical properties of particles and their biological effects have been initiated, but the physical properties of individual NPs themselves remain unexplored. Although IR spectroscopy has been widely used to evaluate the physical properties of polymers, it is difficult to evaluate the chemical properties of nanoparticles due to the limitation of spatial resolution, which only provides an average IR spectrum for many particles with a spatial resolution of several micrometers or more. Recently, however, AFM-IR has been developed as a nano IR spectroscopy method by Dazzi *et al.*^{39–43} AFM-IR is a technique in which light from a tunable infrared laser is irradiated onto a localized region of a sample and the resulting amplitude change in photothermal expansion of the sample is measured using the cantilever of an atomic force microscope to obtain IR absorption spectra and absorption images of individual independent particles with a spatial resolution on the 50–100 nm scale.^{39–45} Although the concentration of NPs produced by laser ablation is highly dilute and needs to be concentrated, the physical properties of individual NPs can be potentially evaluated by allowing independent particles to accumulate without aggregation on the prism used in AFM-IR.

As yet, the processes involved in this study have been developed independently and have not been optimised as a complete method. This study proposes and demonstrates a new method to evaluate NPs to measure the shape and local infrared properties of concentrated nanoparticles by adapting bubble accumulation, as shown in Fig. 1. Our method realizes the capture of individual nanoparticles diluted in solvent and will be available for the study of biological effects. Once the physical properties of the nanoparticles themselves are clarified, a better understanding of the environmental and biological effects and relationships, the mechanism of NP aggregation, and the interatomic forces between particles can be promoted based on the correct nanoparticle properties.

2 Experimental

In order to develop a new method that can evaluate the local infrared spectral properties of nanoparticles by adapting bubble collection, three processes need to be optimised.



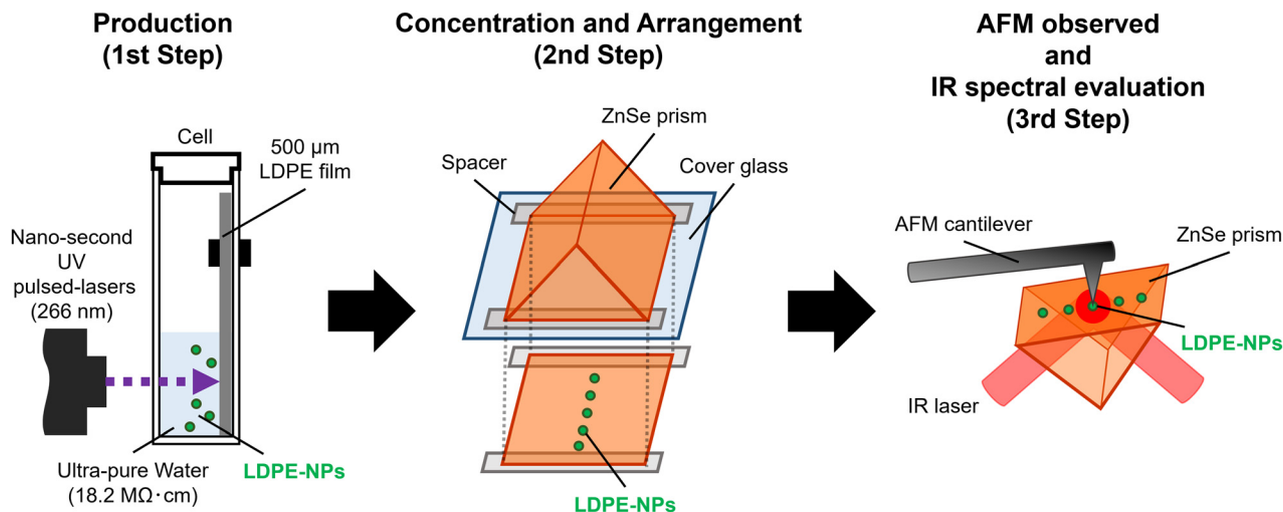


Fig. 1 Schematic of our system designed to evaluate the local infrared spectral properties of nanoparticles by adapting bubble accumulation to capture individual nanoparticles diluted in solvent.

Firstly, appropriate nanoparticles need to be produced by nano-second laser ablation. The conditions we have used are shown in Table 1. A double pulsed Q-switched Nd:YAG laser (Irradiation wavelength 266 nm, Pulse duration of 12–15 ns, Repetition rate of 10 Hz, Lotis Tii LS-2134D) is used. It consists of two independent Nd:YAG lasers pumped by a single flash lamp. A 500 μm thick LDPE film sample (Aldrich LDPE (melt index 25 g/10 min (190 $^{\circ}\text{C}$ /2.16 kg)), Sigma-Aldrich Co. LLC) is fixed inside a quartz glass cell filled with ultrapure water (18.2 $\text{M}\Omega\text{ cm}$, Simplicity UV). There, a laser

beam with a UV wavelength of 266 nm is irradiated for 1 h on the LDPE film sample, and NPs are produced.

In the second process, the NPs produced by nano-second laser ablation in the above are concentrated to analyze the local chemical properties of one particle using microbubbles and placed one particle at a time on a zinc selenide (ZnSe) substrate.⁴⁶ Normally, ITO substrates are applied. The laser-induced microbubbles on the Au film (10 nm) or ITO substrate were formed with 1064 nm irradiation because the substrates absorb the wavelength of 1064 nm and provide photothermal effect.^{35–37} However, in this study, ZnSe prisms were chosen to avoid the influence of infrared absorption in the AFM-IR measurements. For ZnSe, absorption of light occurs at less than 600 nm. In addition, thermal conductivity of ZnSe (18 $\text{Wm}^{-1}\text{ K}^{-1}$) is similar to that of the Au film (38.5 $\text{Wm}^{-1}\text{ K}^{-1}$) and ITO (50 $\text{Wm}^{-1}\text{ K}^{-1}$). Therefore, irradiation with a 450 nm laser ($\sim 45\text{ mW}$) leads to the formation of the bubble.

Fig. 2 shows a model diagram of the enrichment method using microbubbles. Two pieces of extended Parafilm (thickness: $\sim 40\ \mu\text{m}$) as spacers placed on a cover glass, and a ZnSe prism (Nihon Thermal Consulting. Co) was placed on the spacers with its surface to be measured by AFM-IR facing inward. Then, the chamber constructed with the prism was placed on the stage of an inverted optical microscope (IX 71, Olympus), equipped with an oil-immersion objective lens ($\times 100/1.30\ \text{NA}$, Olympus). The solution containing LDPE-NPs by nano-second laser ablation was introduced into the chamber by capillary action. A continuous wave (CW) light beam (450 nm) from a diode laser (Civil Laser) was focused on the ZnSe prism surface through the objective lens of the microscope. The laser intensity at 450 nm was monitored before passing the beam through the objective lens. The focus point of the laser beam was controlled by x - y stage movement of the microscope. Microscopy images were recorded on a CMOS camera (WRAYCAM-VEX120, WRAYMER) mounted on the microscope.

Table 1 The Nano-second laser ablation conditions for producing suitable nanoparticles

Material	Aldrich LDPE (melt index 25 g/10 min (190 $^{\circ}\text{C}$ /2.16 kg))	
Shape of the material	500 μm film	
Laser	A double pulsed Q-switched Nd:YAG laser (Lotis Tii LS-2134D)	
Specifications of laser	Pulse duration	12–15 ns
	Repetition rate	10 Hz
	Irradiation wavelength	266 nm
	Output energy	0.91 mJ
	Shooting area	0.023 cm^2
	Energy density	40 mJ cm^{-2}
	Irradiation time	1.0 h
Cell	Type	Full-sided clear
	Optical path length	10 mm
	Capacity	3.5 ml
	Size	12.5 \times 12.5 \times 56 mm
	Material	Quartz glass
	Model number	SC10104
	Transmittance at 266 nm	90%
Fluence on sample surface	36 mJ cm^{-2}	
Dispersing solvent	Water	Ultra-pure water
	Electrical resistivity	18.2 $\text{M}\Omega\text{ cm}$



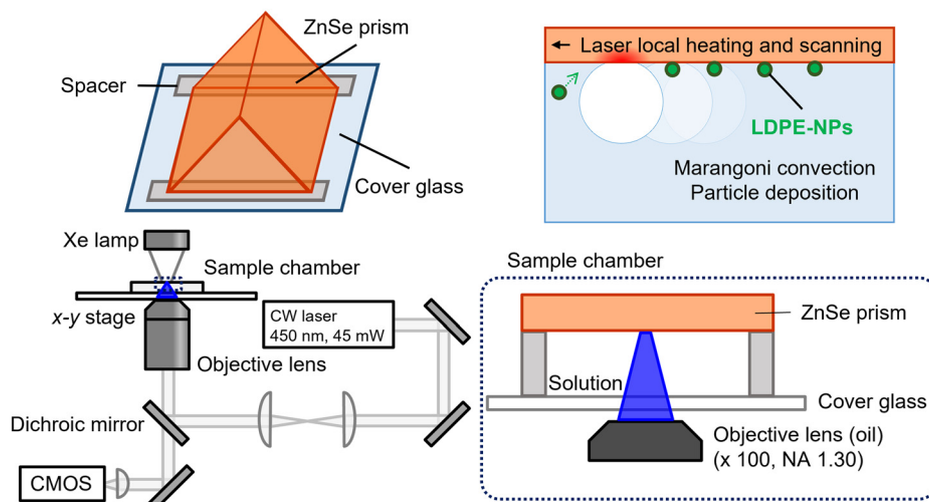


Fig. 2 Schematic of the collection method of LDPE-NPs using microbubbles. The diameter of microbubbles obtained was 5–10 μm .

Finally, nanoIR (Anasys Instruments)⁴⁷ AFM-IR is used to obtain the infrared spectra of the individual nanoparticles. The nanoIR system uses a pulsed, wavelength-variable IR source to analyze the individual pumped molecular vibrations within the LDPE-NPs. As in conventional ATR spectroscopy, the IR beam is irradiated onto the LDPE-NPs by total internal reflection, and rapid thermal expansion occurs as the LDPE-NPs absorb the energy. The cantilever vibrations induced by the thermal expansion are damped by a characteristic ring down, and the amplitude and frequency of the vibrations are extracted by Fourier transform.⁴⁷ Next, the local IR absorption spectrum of LDPE-NPs is obtained by measuring the amplitude of the cantilever vibration while sweeping the wavelength of the irradiating IR laser.

3 Results and discussion

Fig. 3(a) shows the particle size distribution of LDPE-NPs generated from nano-second laser ablation against the

dynamic light scattering (DLS) intensity for 1.0 h (3.6×10^4 pulses) laser exposure time. Fig. 3(b) shows the graph of the autocorrelation function. The autocorrelated polystyrene latex particles (100.5 ± 2.6 nm, monodispersed) at a concentration of 10 mg ml^{-1} are shown as a black line as a reference in the DLS measurement.

The particle size distribution against the scattering intensity in Fig. 3(a) shows that the particle size of LDPE-NPs generated by nano-second laser ablation is distributed in the range of 50–500 nm, and that LDPE-NPs with a diameter of 200 nm are particularly abundant. However, the graph of the autocorrelation function in Fig. 3(b) shows that the value of the autocorrelation function is smaller than that of the polystyrene latex particles in the reference, suggesting that the number of generated LDPE-NPs particles is still rather small. This strongly indicates the need for microbubble accumulation in this study.

Fig. 4(a) shows an optical image of the bubble generation by 450 nm laser irradiation on the ZnSe prism surface. The

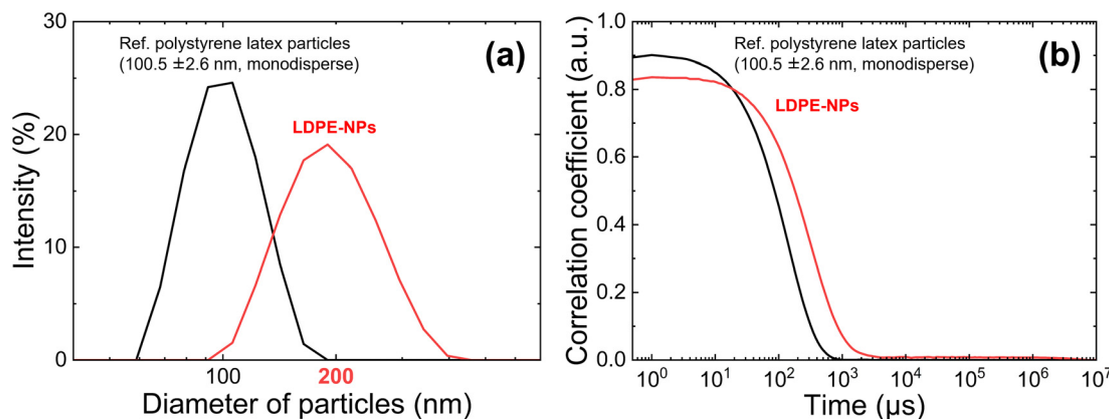


Fig. 3 Particle size evaluation of LDPE-NPs by DLS measurement. Polystyrene latex particles (100.5 ± 2.6 nm, monodisperse) at a concentration of 10 mg mL^{-1} are shown as a black line as a reference. (a) Particle size distribution by scattering intensity versus laser irradiation time of 1.0 h (3.6×10^4 pulses) and (b) graph of the autocorrelation function.



ZnSe prism was irradiated with a 450 nm laser to induce a photothermal effect, which generated microbubbles in the solvent with micro- and nano-plastic particles, and the micro- and nano-plastic particles were collected by Marangoni convection. After the laser irradiation was stopped and the bubbles disappeared, the substrate surface was observed, and no holes or other damage was found on the substrate. For reference, an aqueous solution in which polyethylene particles 1–4 μm in diameter (Ref. PE-MPs, CPMS-0.96 1–4 μm –0.2 g, Cospheric) were dispersed and introduced into the chamber to generate the bubbles, and the Ref. PE-MPs gathered around the bubbles and accumulated on the surface of the ZnSe substrate (Fig. 4(b)). These demonstrations on the ZnSe substrate show that the previously reported bubble enrichment and collection of micro- and nanoparticles is also possible on the ZnSe substrate and can be deployed in a combined AFM-IR measurement system. Unfortunately, nano-sized plastic particles cannot be observed under an optical microscope, but they are assumed to have accumulated, and samples were then prepared for AFM-IR spectral measurements.

Fig. 5 shows the IR spectra of the LDPE-NPs prepared on the ZnSe prism, along with AFM images of the micro-particles that were present in the observation area. The IR spectrum of Ref. PE-MPs used as a reference is also shown. PE-MPs were focused on a ZnSe prism, and the IR spectra were measured from nanoIR.

The AFM image of the nanoparticles and the IR spectra (Fig. 5) confirm that the LDPE particles are not only independently spherical and nano-sized (Fig. 5(A)), but also that some LDPE particles are distorted or agglomerated in shape or exist in aggregates, as seen in Fig. 5(B) and (C). (see the ESI† for the height profiles of (A) LDPE-NPs of about 300 nm diameter, (B) LDPE-MPs.1, and (C) LDPE-MPs.2 (Fig. S1†)). The IR spectra show signals originating from C=O stretching (1735 cm^{-1}), C–O stretching (1240 cm^{-1} , 1165 cm^{-1}), C=C stretching (1600 cm^{-1} , 1580 cm^{-1}), and O–H stretching (3400 cm^{-1}), confirming the progress of the oxidation reaction.⁴⁶ In particular, CH₂ asymmetric C–H stretching (2918 cm^{-1}), CH₂ symmetric C–H stretching (2848 cm^{-1}), CH₃ asymmetric stretching (2960 cm^{-1}), and CH₃ symmetric stretching (2870 cm^{-1}) can be also found following

the oxidation reaction. This is thought to be due to the formation of the terminal methyl group.^{48–51} In addition, the methyl group shows a peak at a higher wavenumber than the methylene group in the IR spectra of polypropylene with both methyl and methylene groups.^{48–51} Furthermore, in LDPE-NPs with a diameter of about 300 nm (Fig. 5(A)), the peak intensity of the C=C bond increased compared to that of the C=O bond, and a lower wavenumber shift was observed compared to the C=C bond of mono-olefins, suggesting the formation of relatively long conjugated systems. It is thought that the photo-oxidation first occurs on the surface of micro- and nano-plastics (MNPs) and produces highly reactive organic radicals and ROS that are involved in the radical reactions.⁵² Typically, along with the generation of alcohols, ketones, olefins, and aldehydes, carboxylic acids and esters, vinyl groups and O-containing functional groups, such as carbonyl, carboxyl, and hydroxyl groups can be formed during the photo-oxidation of MNPs.⁵² In addition, new peaks in the carbonyl ($1500\text{--}1800\text{ cm}^{-1}$) and hydroxyl ($3000\text{--}3800\text{ cm}^{-1}$) regions were observed in UV-irradiated polypropylene-microplastics (PP-MPs) in ultrapure water, indicating the formation of oxygen related functional groups on the surface of PP-MPs during the aging process from ATR-FTIR analysis.⁵³ Please refer to the ESI† for an example of a possible decomposition reaction scheme (Fig. S2†). Note that for Ref. PE-MPs, standard PE absorption spectra are obtained. This suggests that the oxidation was promoted by nano-second laser ablation at ultraviolet wavelengths or in ultrapure water, and not by laser-induced oxidation at the time of accumulation.^{54–60} From the above, we confirmed the relation between the size and state of the nanoparticles. One of the most possible explanations is that greater oxidation produces smaller particles.

4 Conclusions

This study proposes and demonstrates a new method to evaluate the shape and local infrared properties of nanoparticles by adapting bubble accumulation, which achieves the capture of individual nanoparticles diluted in solvent. Three steps are completed: first, nano-second laser ablation is utilized to fabricate nanoparticles to fabricate a

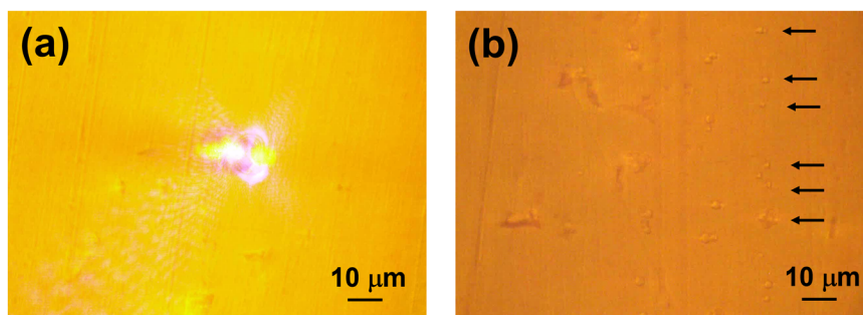


Fig. 4 Optical images of the interface between the ZnSe prism surfaces and solution under a microscope. (a) Generation of a microbubble by 450 nm laser irradiation and (b) ref. PE-MPs (1–4 μm) deposited by the bubble. Black arrows indicate some of the particle locations.



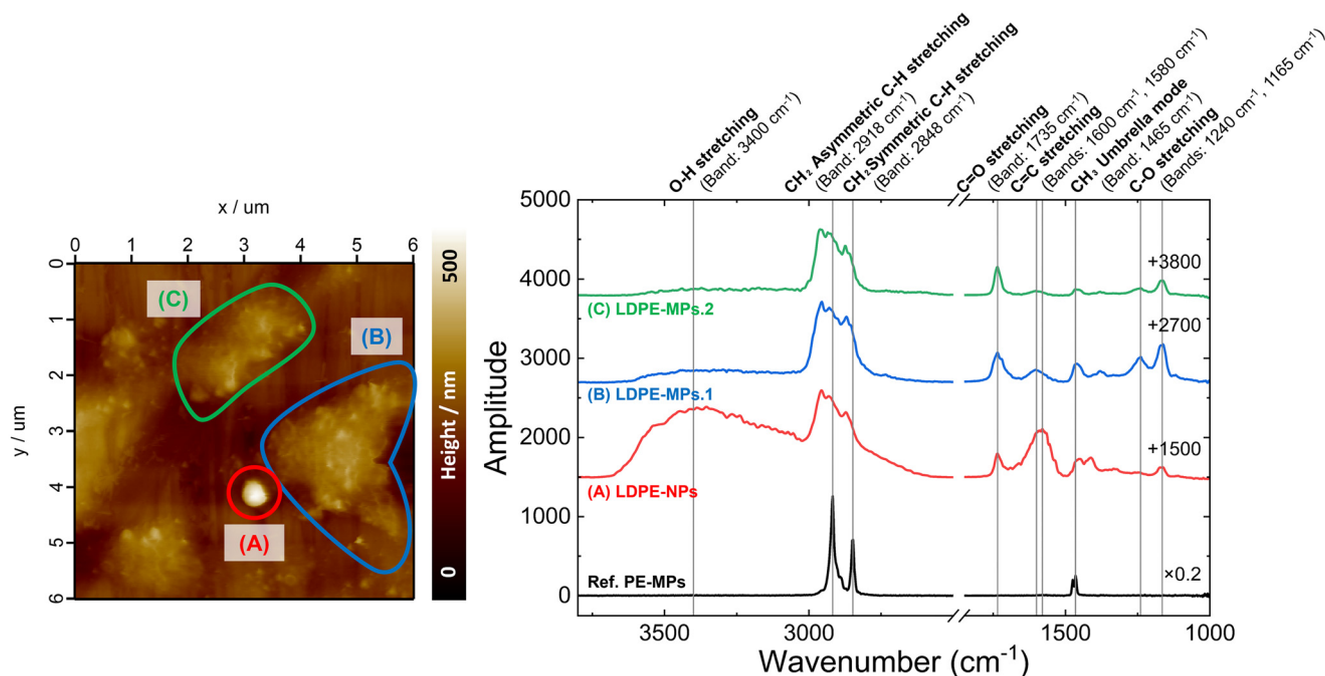


Fig. 5 AFM images and IR spectra of nanoparticles. It was found that the particle size is considered to decrease as the oxidation reaction progresses. (A) LDPE-NPs of about 300 nm diameter, (B) LDPE-MPs.1 and (C) LDPE-MPs.2.

defined model sample that is artificially dispersed in nature without solvent residues, surfactants, or impurities. The specific conditions were as follows: nano-second laser ablation (irradiation wavelength of 266 nm, pulse duration of 12–15 ns, repetition rate of 10 Hz, an irradiation fluence of 36 mJ cm^{-2} , and an ablation area of 0.023 cm^2) for a 500 μm thick LDPE film sample was performed; second, LDPE-NPs were concentrated and prepared to analyze the local chemical properties of single particles. Using Marangoni convection of microbubbles, LDPE-NPs were placed one particle at a time on the ZnSe substrate surface. Thirdly, AFM-IR of nanoIR (Anasys Instruments)⁴⁷ was used to analyze the chemical properties of nanoparticles as well as shape observations for individual nanoparticles. AFM observations confirmed the presence of independent nanoparticles. For these, infrared resonance peaks attributed to C=O stretching, C–O stretching, C=C stretching, O–H stretching, *etc.* were observed, confirming the progress of the oxidation reaction. For nanoparticles with a diameter of about 300 nm, the intensity of the C=C bond increased compared to the C=O bond, and a red shift was observed compared to the C=C bond of mono-olefins, suggesting the formation of relatively long conjugated systems.

Based on these results, whilst the formation mechanism, understanding of biological effects, and nano-physical properties in NPs dispersed in natural solutions have not been clarified, this study significantly advances the processes of NP generation, preparation, and analysis. This system design approach, as proposed in Fig. 1, can contribute to NP research as a system for rapid generation, concentration, and analysis. The clarification of the physical properties of the

nanoparticles will help to better understand their environmental and biological effects and relationships, the mechanism of NP aggregation, and the interatomic forces between particles.

Data availability

The data supporting this article have been included as part of the ESI.†

Author contributions

Ikuna Kanehara: investigation (lead); visualization (equal); formal analysis (equal); writing – original draft (lead); writing – review and editing (equal). Tatsuhiro Nagasaka: resources (lead); formal analysis (equal); writing – review and editing (equal). Hirofumi Seki: resources (supporting); formal analysis (supporting); writing – review and editing (equal). Sho Fujii: visualization (equal); formal analysis (equal); writing – original draft (supporting); writing – review and editing (equal). Tsuyoshi Kimura: supervision (equal); writing – review and editing (equal). Masaya Yamamoto: supervision (lead); funding acquisition (lead); writing – review and editing (equal). Tadao Tanabe: supervision (equal); writing – original draft (supporting); writing – review and editing (equal).

Conflicts of interest

The authors have no conflicts to disclose.



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