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Current perspectives, challenges, and future directions in the electrochemical detection of microplastics

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Microplastics (5 μm) are a developing threat that contaminate every environmental compartment. The detection of these contaminants is undoubtedly an important topic of study because of their high potential to cause harm to ecosystems. For many years, scientists have been assiduously striving to surmount the obstacle of detection restrictions and minimize the likelihood of receiving results that are either false positives or false negatives. This study covers the current state of electrochemical sensing technology as well as its application as a low-cost analytical platform for the detection and characterization of novel contaminants. Examples of detection mechanisms, electrode modification procedures, device configuration, and performance are given to show how successful these approaches are for monitoring microplastics in the environment. Additionally included are the recent developments in nanoimpact techniques. Compared to electrochemical methods for microplastic remediation, the use of electrochemical sensors for microplastic detection has received very little attention. With an overview of microplastic electrochemical sensors, this review emphasizes the promise of existing electrochemical remediation platforms toward sensor design and development. In order to enhance the monitoring of these substances, a critical assessment of the requirements for future research, challenges associated with detection, and opportunities is provided. In addition to—or instead of—the now-in-use laboratory-based analytical equipment, these technologies can be utilized to support extensive research and manage issues pertaining to microplastics in the environment and other matrices.

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a novel stationary phase in chromatographic separation, and solid-phase extraction.

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1. Microplastics: sources and occurrence

Owing to numerous transport phenomena like wind and ocean currents, microplastics (MPs) can be found in coastal areas and aquatic ecosystems worldwide in different size fractions. Household sewage discharge, polymeric particles from cosmetic and cleaning goods, feedstocks used to make plastic products, and plastic pellets or powders used for air blasting are the primary sources of these contaminants.¹ The secondary source of MPs is the progressive fragmentation of larger plastic items under atmospheric conditions, such as mechanical deterioration and UV-light exposure, which helps to introduce significant amounts of MPs into the environment.² This increases the consumption of plastic waste by a wide range of creatures and the emergence of new environmental threats.³

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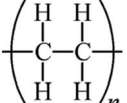
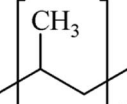
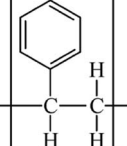
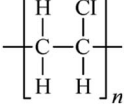
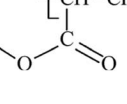
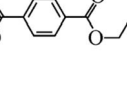
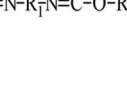
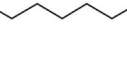
Another significant source of MP discharge is wastewater treatment facilities.^{4,5} Large plastic particles are effectively removed during wastewater treatment, but MPs frequently evade the units and enter the water.⁶ It is noteworthy that many water treatment facilities are situated close to the ocean and seawater, creating a high source of MP release. For instance, in Mainland China, about 1873 wastewater plants (56%), out of 3340, with $78 \times 10^6 \text{ m}^3 \text{ day}^{-1}$ of treatment capacity are located in coastal regions where their effluents can be directly or indirectly discharged into aquatic ecosystems.⁷ Various studies have addressed the fate, incidence, detection, and removal of MPs in water treatment plants.^{8,9}

Microbeads (<1 μm) used in cleaning products, cosmetics, and laundry soaps are additional sources of MPs. Polyethylene (PE), polypropylene (PP), polystyrene (PS), polyethylene terephthalate (PET), polyvinyl chloride (PVC), and polyesters are the major components of (Table 1) these particles, which are eventually carried into the ocean (Fig. 1).

2. Microplastics and human health

The ingestion or inhalation of MPs into the body of a person has the potential of negative health impacts (Fig. 2).¹⁰ A comparison to particulate air pollution can be made: small particles (2.5

Table 1 Major MPs found in the marine environment

Type of polymer	Density (g cm^{-3})	Biodegradation	Recycle ID code	Molecular structure
High density polyethylene (HDPE)	0.917–0.965	Not reported	02 PE-HD	
Low density polyethylene (LDPE) Linear low density polyethylene (LLDPE)		Bacteria, fungi, waxworms, mealworms	04 PE-LD	
Polypropylene (PP)	0.90–0.91	Not reported	05 PP	
Polystyrene (PS)	1.04–1.1	Bacteria, mealworms	06 PS	
Polyvinylchloride (PVC)	1.16–1.58	Fungi	03 PVC	
Polymethylacrylate (PMA)	1.17–1.2	Cyanobacteria	07 O	
Polyethylene terephthalate (PET)	1.37–1.45	Bacteria	01 PET	
Polyurethane (PU)	1.20	Fungi	07 O	
Polyesters (nylon, acrylic, etc.)	1.24–2.3	Bacteria	07 O	



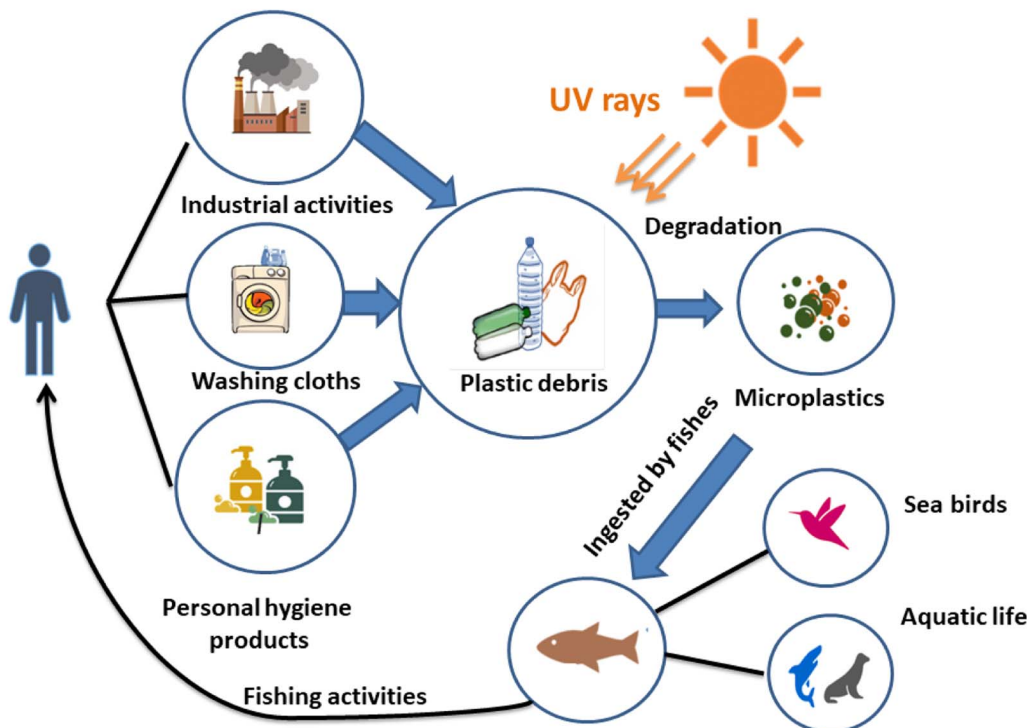


Fig. 1 MPs pathways.

mm), such as those found in diesel exhaust, have been associated with an increased risk of dying from lung cancer, cardiovascular disease, and inflammatory illnesses because they can cross the cell membranes and cause oxidative stress and inflammation.¹¹ This correlation offers strong motivation to learn more about the potential dangers of MP particles.

The reported concentrations of MPs in tap and bottled water range from 0 to 10^4 particles per liter, with smaller MPs often having higher particle counts.¹² According to the initial air observations of larger, mostly fibrous MPs, the deposition rates in central London, for instance, ranged from 575 to 1008 MPs per square meter per day,¹³ indicating that plastic particles are a major component of fine dust. Increased exposure from contaminated indoor air, ingestion of household dust or dust that settles on food,¹⁴ and direct exposure to food particles discharged from plastic bottles or containers (such as PP baby-feeding bottles¹⁵), are of particular concern.

Larger MPs are probably eliminated through feces or mucociliary clearance into the gut after depositing in the respiratory system or lungs.^{12,13} The fraction of smaller-sized particles (10 μm), which are probably toxic, is typically not included in existing assessments due to methodological limitations and measurement preference toward larger particles, which probably underestimate human external exposure.^{16,17} Remarkably, the assessments of internal exposure to plastic particles in biological fluids and tissues are still in their early stages.

Reducing the ambiguity in the human risk assessment of MPs requires a deeper comprehension of their capacity to pass through the epithelial barriers of the gastrointestinal system,

skin, and airways. A limited amount of *in vitro* and *in vivo* data indicates that only a tiny portion of administered MPs can pass through the epithelial barriers of the lungs and intestines.

These MPs have distinct uptake characteristics and generally increase in absorption efficiency as the particle size decreases.¹⁸ Because of the potential for accumulation in tissues and organs as well as lifetime exposure, this modest percentage of particle absorption is not insignificant. MPs smaller than 10 μm have been found to move from the gut cavity to the lymph and circulatory systems, inducing systemic exposure and accumulation in tissues such as the brain, kidneys, and liver.¹⁷ These findings have also been observed in rodents and aquatic species. Even though the tiniest particles (less than 0.1 μm) have the potential to penetrate all the organs and even the placenta, brain, and cell membranes,^{17,19,20} there are still significant gaps in our understanding of absorption, distribution, metabolism, and excretion (ADME). It is also unknown if MPs have dose-dependent impacts on humans.

MPs could cause physical, chemical, and microbial toxicity, as well as long-term effects, if they touch the epithelial lining of the intestines or lungs or if they are consumed. Numerous studies on human cell cultures in the lab and on rodents in the wild suggest that breathing or consuming MPs may have a wide range of biological effects, such as physical (particle) toxicity, which can lead to oxidative stress, cytokine secretion, cellular damage, immune and inflammatory responses, DNA damage, and neurotoxic and metabolic effects.¹⁷ In these studies, only a small number of pure, commercially available particle types are used. These particles are not the same as those found in the environment, and the effects found are often caused by high

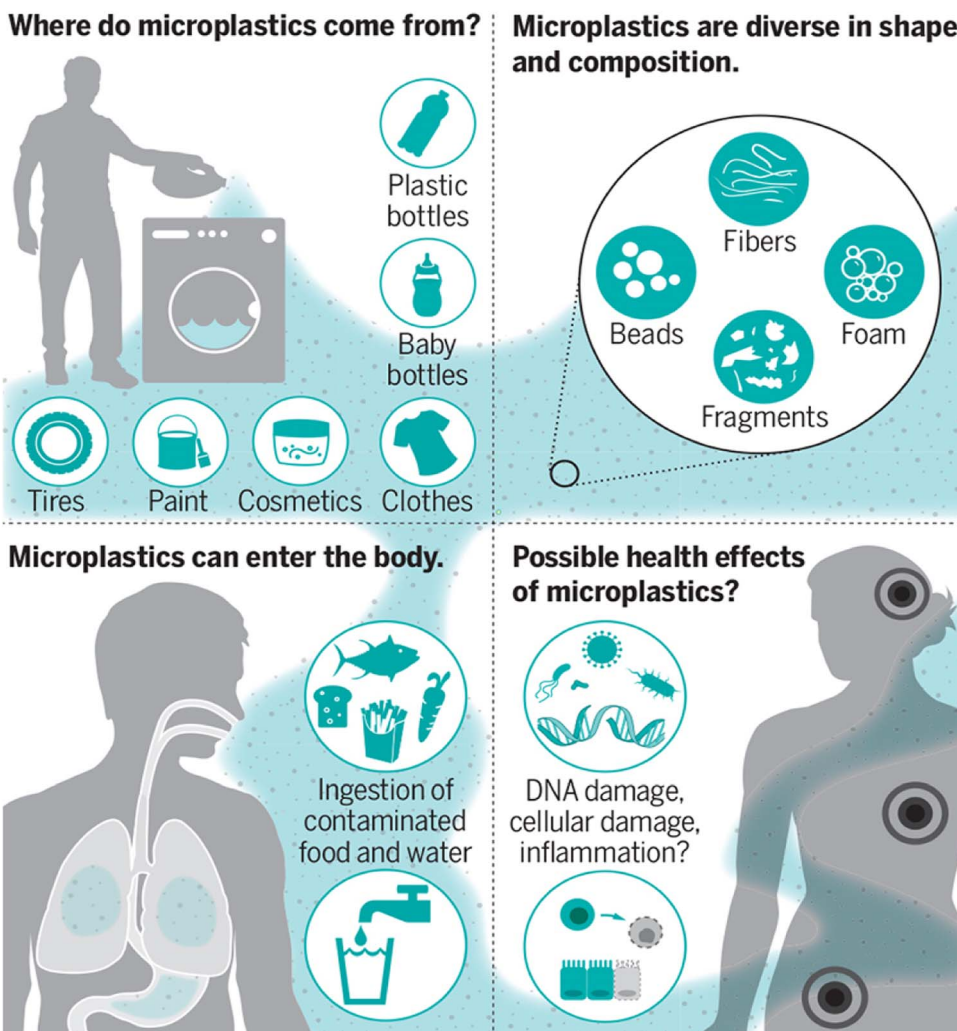


Fig. 2 MPs and their effect on human health.¹⁰

doses of MPs. Moreover, it is not always possible to rule out the chemical contamination of these test particles. Epidemiological studies have documented lung damage, including inflammation, fibrosis, and allergies, among employees in the plastic and textile sectors who are exposed to high levels of plastic fiber dust.¹⁸ These findings are consistent with the effects seen in research on ambient particle exposure.

Every day, humans are exposed to a variety of natural and synthetic particles; among them, particulate air pollution is acknowledged as one of the primary environmental risk factors for diseases worldwide. It is imperative to assess the possible contribution of MPs in overall ambient particle exposure. In comparison to other ambient particles, MPs may have unique particle properties and a wider and more diverse toxicity profile because of their persistence, wide size range, and complex structure. Important health concerns related to MPs remain largely unexplored to this day. These include internal exposure; ADME processes (including the effect of the eco- or bio-corona); interaction with the immune system; and the possible effects of nanosized plastics on the placenta, fetus, and brain. These

problems are essential to innovation, evidence-based policy-making, and risk management strategies. They are starting to be resolved by innovative interdisciplinary research programs like Microplastics & Health in the Netherlands and the European Union Horizon 2020 research programme. To address this potential health risk, multidisciplinary research projects including experts from the medical, environmental, and polymer domains are required. Although comprehensive risk assessment is still way off, it is important to fill in the major research gaps today to enable prompt decision-making about health policy and mitigation techniques.

3. Analytical techniques for the detection of microplastics

3.1. Visual inspection methods

Selecting and classifying MPs as well as sighting the color and size of the examined objects under a microscope or with the naked eye can be undertaken using visual inspection approaches such as the direct visual method, optical



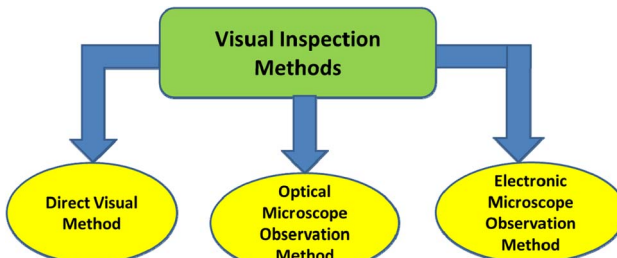


Fig. 3 Classification of visual inspection methods.

microscope observation, and electronic microscope observation²¹ (Fig. 3). The accuracy of these methods is poor, and they are time consuming.²² MPs are tiny plastic particles with a diameter of less than 1 mm that are easy to overlook or underestimate. Furthermore, the results showed that the error rate is inversely related to the particle size when employing this technique.^{23,24} Consequently, more accurate and efficient technologies for MP identification must be developed. The advantages and limitations of such methods are summarized in Table 2.

3.2. Techniques for thermal analysis

Fig. 4 shows the categorization of thermal analysis.

3.2.1. Gas chromatography/mass spectrometry (GC/MS) for pyrolysis. Combining gas chromatography-mass spectrometry (GC-MS) and pyrolysis is known as pyrolysis-gas chromatography-mass spectrometry (Pyr-GC-MS). In this method, materials are pyrolyzed in the absence of oxygen, converting the polymer into volatile small molecules. To determine the products of combustion or pyrolysis, these molecules are subsequently introduced to the GC-MS apparatus.²⁵ In pyrolysis, individual MP particles are added to a reaction tube, and the products of pyrolysis produced by different polymers are usually identified. The gaseous products of the reaction are then cold-injected, collected by the apparatus, and directed to a mass-spectrometer-connected GC column. The resulting pyrolysis product spectrum is compared with the spectrum database of commonly used plastic types to complete the MP detection process. In addition to providing information on potentially dangerous organic plastic additives (OPAs), Pyr-GC-MS can accurately identify various polymer types.²⁶ Pyr-GC-MS can also be used to precisely evaluate the materials' chemical attributes, as well as to examine their

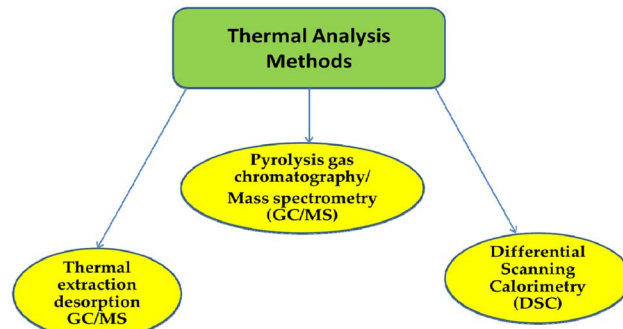


Fig. 4 Classification of thermal analysis methods.

structural features and chemical composition, by searching for the high-molecular-weight polymer pyrolysis byproducts.²⁷ This might provide a thorough sample description.

3.2.2. GC/MS for thermal extraction and desorption.

Thermal extraction desorption-gas chromatography-mass spectrometry (TED-GC-MS) is an improved thermal analytical method that combines thermogravimetric analytical solid-phase extraction (TGA-SPE) and thermal desorption-gas chromatography-mass spectrometry (TDS-GC-MS).²⁸ The substance is pyrolyzed in TGA at temperatures as high as 1000 °C before being adsorbed on solid-phase reagents for extraction. A thermal desorption device is used to desorb the products of deterioration when the temperature is elevated. After that, it is possible to remove the sample from a chromatographic column and conduct MS analysis. This method shortens the analysis time for large sample sizes while increasing the sample weight to 100 mg. Furthermore, this method prevents the issue of high-molecular-weight pyrolysis products blocking the reaction tube²⁹ and does not require the preparation of the material.³⁰

3.2.3. Calorimetry with differential scanning. Through differential scanning calorimetry (DSC), substances can be identified by measuring the melting temperature of a plastic sample heated to various temperatures.³¹ Data on the samples of MPs' gaseous products—qualitative as well as quantitative—can be obtained using this technique. To investigate the thermal properties of polymer materials, DSC is widely used. It is as simple and affordable as high-level research methodology. All the plastic products have different DSC characteristics; hence, reference materials are needed to distinguish between the different kinds of polymer. In addition, large particles create environmental disruption because of their larger mass-to-surface-area ratio compared to small particles.³²

Table 2 Advantages and disadvantages of visual inspection methods

Method	Methodology	Particle size	Advantages	Limitations
Counting microscopy	Direct particle counting is done	It can identify particles with sizes in the micrometer (μm) range	Samples containing a considerable number of large MPs are easily identified, providing a fast and reasonably priced overview of MP abundance	Since the nature of the materials cannot be ascertained, a combination of identification techniques is required

Table 3 Comparison of MP detection techniques using thermal analysis

Method	Advantages	Disadvantages	Sample mass	Detection limit	Ref.
Pyr-GC-MS	<ul style="list-style-type: none"> • Various polymer kinds can be detected • Accurate findings • High sensitivity • No sample preparation 	<ul style="list-style-type: none"> • Lengthy process • Sample deterioration • High reaction temperatures 	0.5 mg	0.007 mg g ⁻¹	27 and 33
TED-GC-MS	<ul style="list-style-type: none"> • Large sample size • No sample preprocessing • No blocking reaction cube • High sample mass • Reliable outcomes • Extensively used methodology • Affordable and uncomplicated analysis 	<ul style="list-style-type: none"> • High reaction temperatures • Sample deterioration. Excessive processing times 	100 mg	—	29, 34 and 35
DSC	<ul style="list-style-type: none"> • Prolonged processing durations • Sample degradation • Clear substrate influence • Sample preparation 		3–15 mg	—	31 and 32

The production variables that affect the transition temperature are impurities, additives, and a part of the polymerization chains. Because MP particle size has a significant impact on the DSC values, samples need to be pretreated before testing. This is yet another DSC disadvantage. Furthermore, when the melting point of the polymer mixture is the same or when the sensitivity of the DSC sensors is low, some polymers have limited thermal conductivity. Its limitations hinder its rapid application in MP detection. It was found that thermogravimetric analysis-differential scanning calorimetry (TGA-DSC) could accurately separate PE and PP in environmental samples. The phase-change signals of other types of MP cannot be recognized because of the significant overlap in the phase-change signals. The larger sample size causes the peak area to increase but the resolution to decrease, leading to inaccurate measurement results.

Table 3 provides an overview of the benefits and drawbacks, as well as some significant characteristics of thermal analysis techniques. Table 4 provides an overview of the applications of thermal analytical techniques in MPs.

3.3. Spectral analysis technique

The information obtained from the spectrum analytical method is more accurate than that obtained *via* visual recognition.⁴⁵ At present, spectroscopic methods can identify MPs and confirm their composition. The spectral signal needs to be analyzed to reflect the different characteristic peaks that each type of MP creates. Fourier-transform infrared (FTIR) spectroscopy and Raman spectroscopy have been used to identify the polymer types of MP particles with a minimum particle size of 10 and 1 μm , respectively.

3.3.1. Raman spectroscopy. The inelastic scattering of light is the foundation of the vibrational method called Raman spectroscopy.⁴⁶ As the scattering spectrum has a different frequency than the incident light, it can be used to determine the molecular structure of the material by studying or considering the rotation and vibration of the molecules. High sensitivity, high precision, high specificity, and high spatial resolution (<1 μm) are provided by Raman fingerprint

spectroscopy.⁴⁷ The advantages of this method also include the absence of requirements for sample preparation, staining, minimum sample thickness, or sample damage. To determine the polymer compositions of the materials, they are currently being swiftly treated. Due to the intrinsic resonance fluorescence phenomena, organic compounds, inorganic chemicals, and colored additives can easily cause the fluorescence background to interfere with Raman scattering.^{48,49} Furthermore, the Raman signal is usually faint, which makes accurate identification difficult to establish.

Due to its excellent signal-to-noise ratio and lack of fluorescence interference, the nonlinear Raman spectroscopy approach is becoming increasingly popular in the field of MP detection.^{50,51} Surface-enhanced Raman spectroscopy, on the other hand, is gradually expanding its ability to detect MPs because of its unique chemical specificity and high sensitivity. Future quick *in situ* MP detection in water may be possible owing to the Raman spectroscopy approach, which has the potential to solve the challenge of target identification in liquids.

3.3.2. FTIR spectroscopy. The FTIR spectra of a material are a complementary spectrum of the Raman spectra by looking at the chemical bonds in the infrared spectrum. Variations in the chemical bond configurations lead to distinct peak patterns and diagrams. It is, thus, possible to achieve the detection target by comparing the identified component materials of the particles with the standard library.⁵² Even though FTIR has a spatial resolution of as little as 20 μm ,⁵³ the drawback of time-consuming processing should be taken into account for the time being. The sample must be dried since FTIR has complex spectra when detecting wet materials, is easily disturbed by water, and has poor horizontal resolution. Microscopy and FTIR are combined by FTIR for the characterization of MPs, whose detection limit is smaller than 10 μm .⁵⁴ This approach has the potential to be used for environmental MP identification due to its benefits, including small sample sizes, high-throughput screening, and environmental friendliness.⁵⁵ This method usually produces a weak signal for tiny MPs,⁵⁶ which may result in several false positives or false negatives.⁵⁷ There are other



Table 4 Overview of the thermal analysis application in MPs^a

Method	Source of the sample	Type of sample	Limit of detection (LOD)	Sample abundance	Ref.
Pressurized liquid extraction and Pyr-GC-MS	Soil and sediments	PE, PP	NR	PE ($3.3 \pm 0.3 \text{ mg g}^{-1}$) and PP ($0.08 \pm 0.02 \text{ mg g}^{-1}$)	36
Pyr-GC-MS	Soils and sediments of freshwater	PE (fractionation ratio 15 : 2), PE (fractionation ratio 17 : 2), PE (fractionation ratio 18 : 2), PP, PS (pyrolysis product Sty), and PS (pyrolysis product aMeSty)	PE (fractionation ratio 15 : 2, $4800 \mu\text{g L}^{-1}$), PE (fractionation ratio 17 : 2, $2500 \mu\text{g L}^{-1}$), PE (fractionation ratio 18 : 2, $11300 \mu\text{g L}^{-1}$), PP ($43200 \mu\text{g L}^{-1}$), PS (pyrolysis product Sty, $500 \mu\text{g L}^{-1}$) and PS (pyrolysis product aMeSty, $1600 \mu\text{g L}^{-1}$)	NR	37
Pyr-GC-MS	Stomachs of marine fishes	PVC, PET, nylon, silica gel, and epoxy resin	NR	NR	38
Pyr-GC-MS	Surface water and wastewater	PS and PE	PS ($30 \mu\text{g L}^{-1}$) and PE ($1000 \mu\text{g L}^{-1}$)	NR	39
Pyr-GC-MS and Nile red dye	Lagoon sludge	NR	NR	Fresh sludge ($40.5 \pm 11.9 \times 10^3$ particles per kg) and dehydrated sludge ($36 \pm 9.7 \times 10^3$ particles per kg)	40
Pyr-GC-MS and FTIR TED-GC-MS	Surface water Freshwater	PVC, PP, and PE PE, PS, PET, and PP	NR PE ($20.0 \mu\text{g mg}^{-1}$), PP ($5.7 \mu\text{g mg}^{-1}$), PS ($2.2 \mu\text{g mg}^{-1}$) and PET ($18.0 \mu\text{g mg}^{-1}$)	0–110 000 particles per km ² NR	38 41
TGA, DSC	Wastewater	PE, PP, PET, PA, PES, PVC, and PU	NR	NR	42
DSC ATR-FTIR and DSC	Wastewater Dutch beaches	PE, PP, PA, and PET NR	NR NR	NR	43 44

^a NR: not reported.

requirements for the thickness and characteristics of MPs in this procedure. In Table 5, the alkalization of spectroscopic analytical techniques to examine is presented.

3.4. Other analytical methods

With its low detection limit and good sensitivity, high-performance liquid chromatography (HPLC) is an often-used terminal detection method. It is also more appropriate for finding large, polar, and thermally unstable MPs.^{58,59} Scanning electron microscopy-energy-dispersive spectroscopy (SEM-EDS) is an analytical technique that combines scanning electron microscopy (SEM) and energy-dispersive spectroscopy (EDS). Microstructures can be detected using scanning electron microscopy (SEM), which is a technique that falls between transmission electron microscopy and optical microscopy. Depending on the composition of the surface material, SEM can be directly used for microscopic imaging. The ability to capture MP surface features in clear, high-magnification pictures facilitates the ability to distinguish MPs from other organic-matter particles.⁶⁰ Since elemental analysis can only provide an image of the material's surface morphology, it is critical to integrate the surface properties of plastic particles with elemental analysis in order to more accurately identify MPs. Nowadays, the elemental composition data of MPs are

commonly examined using EDS to characterize their surface structure and discover more about their chemical composition. On the basis of the surface emission radiation of MPs, diffraction and reflection can be used to indicate the elemental composition of polymers, and SEM-EDS can be used to look into the surface morphology of MPs with a sample capacity of 5–10 mg.⁶¹ Despite the fact that this approach is currently often employed to characterize nanoplastics, it has the drawback of providing no chemical information. Table 6 provides an overview of the advantages and disadvantages of using HPLC and SEM techniques for MP identification.

3.5. Parameters used in sampling of microplastics

Following sample preparation, several techniques can be used to separate, identify, and quantify MPs from environmental samples. It is feasible to compile the frequently reported techniques in the scientific literature into grouped sampling approaches, including analytical techniques already used and established for other analytes, even though a universally accepted protocol for sampling and quantifying MPs in the environment is not available. Table 7 lists a few parameters for sampling MPs in various matrices, along with the corresponding analytical methods for finding and measuring MP particles.



Table 5 Merits and drawbacks of analytical methods using spectroscopy

Method	Methodology	Particle size by different analytical methods	Merits	Drawbacks
FTIR	<ul style="list-style-type: none"> The excitable vibrations produced by a substance rely on its molecular structure and composition The IR spectra of plastic polymers have very distinctive band structures 	<ul style="list-style-type: none"> Particles >500 μm in size can be analyzed using ATR-FTIR Particles <20 μm in size can be analyzed using microscope coupled FTIR 	<ul style="list-style-type: none"> Non-invasive methods Reliable, efficient, and dependable The newly developed automatic FTIR imaging techniques, such as FPA, which allow thousands of spectra to be quickly captured inside an area with a single measurement, greatly reduces the analysis time 	<ul style="list-style-type: none"> Samples must be IR reactive, due to insufficient absorbance Samples <20 μm may not yield spectra that may be understood. It is difficult to evaluate non-transparent particles with this technology To operate and process the data from the expensive, highly specialized instruments, competent employees are required The detection is affected by the ambient matrix (biofilm formation on polymer, for example), which complicates the interpretation of the results The sample needs to be treated to remove the IR-active water The fluorescence resulting from inorganic, organic, and biological impurities significantly hinder MP identification
Raman spectroscopy	<ul style="list-style-type: none"> The interaction between the irradiated laser light and the molecules and atoms of the sample causes the back-scattered light to have a different frequency than that of the irradiating laser The Raman shift can be used to create Raman spectra that are unique to a given molecule 	<ul style="list-style-type: none"> The microscopy-coupled Raman spectroscopy (RS) technique is suitable for particles larger than 1 μm As far as particles go, this is the only practical method for sizes between 1 and 20 μm 	<ul style="list-style-type: none"> The study of minute particles, ranging from 1 to 20 μm, with good spatial resolution and comparatively low water sensitivity is made possible by microscopy-linked RS In addition to being able to quickly execute chemical mapping and analyze opaque and dark particles, the RS approach enables automatic and speedy data collection and processing 	<ul style="list-style-type: none"> The sample needs to be cleaned before analysis can begin Three important Raman collection parameters are wavelength, laser intensity, and photo-bleaching. Automated mapping using micro-RS is still being developed RS's analysis takes a long time

4. Electrochemical detection of microplastics: a solution for low-cost monitoring

4.1. Overview

For the purpose of finding MPs in intricate matrices, extremely sensitive spectroscopic techniques are utilized. When it comes to simultaneously determining the MP concentration, these procedures are fairly adaptable. Additionally, these methods have femtomolar-range detection limits that are extremely low. Nevertheless, the production of several samples requiring challenging analytical methods and the need for highly skilled individuals to operate the sophisticated equipment are the drawbacks of these costly spectroscopic approaches.

Additionally, these methods must be combined with other chromatographic methods in order to complete sample specification, and they are only appropriate for quantitative analysis. This may increase the chance that the sample will alter during handling and storage. Furthermore, these optical techniques necessitate high-precision and high-power operations, making them unsuitable for in-field applications once more. They include expensive and sophisticated equipment with lasers, photodetectors, and so on. Thus, research is still being done to create quick, inexpensive, easy, and dependable methods that are appropriate for *in situ* and timely measurements of MPs.

Conversely, electrochemical methods are more affordable, approachable, dependable, and appropriate for field use. These electrochemical methods enable straightforward processes and



Table 6 Advantages and limitations of using HPLC and SEM methods for MP analysis

Method	Methodology	Particle size	Advantages	Limitations
SEM	Sample pictures are produced when the sample is exposed to an electron beam, which quantifies the secondary ions in the sample	For particles as small as a micron in diameter, the analysis is feasible	A high-resolution image of the samples can be produced	A high vacuum is required to cover the samples, and there is no precise identification data available
HPLC	Samples are dissolved in certain solvents. To evaluate the various molar mass distributions, size exclusion chromatography is employed, and HPLC analysis is performed to quantify the findings	A sample size of several milligrams is required for the chemical extraction	Certain polymers showed higher recoveries	Its applications are limited to environmental samples because of restrictions on polymer types and the difficulties of establishing physical properties such as size. Only a small number of samples can be assessed per run. This process can only be used to analyze specific polymers, such as PS and PET

are ideal for creating tiny circuits as transportable devices for *in situ* contamination sample monitoring. In addition, these methods require less time for analysis than conventional spectroscopic methods, which enable real-time water sample monitoring. To increase their performance in MP detection, these electrochemical approaches need to be improved in terms of design. Compared to other spectroscopic and optical techniques, these techniques have lower sensitivity and limits of detection (LODs). Different biosensing electrodes are used with different electrochemical techniques to change the electrode material, thereby increasing the sensitivity and limitations of detection. This study looks at the different electrochemical methods used to find MPs in water samples, as well as the newest progress in making different interface materials that can be used to change the electrodes that are used in these methods.

Low-cost sensors and electrochemical techniques are interesting analytical tools for MP field detection.^{79,80} Chemical sensors are generally designed by combining an electrochemical transducer with embedded identification elements and a sensing substance.^{81,82} A chemical or physical change happens when a target molecule interacts with the sensor surface because of the analyte's covalent or non-covalent binding. The target can be either qualitatively or quantitatively detected, owing to the translation of this binding into a readable signal that can be electrochemically tested. Subsequently, while fabricating electrochemical sensors, care must be taken to select and understand the materials that will be exploited to build the sensor as well as the chemical or biological processes that take place at the contact.⁸³ Recognition elements that are designed and selected at the electrode surface are chosen on the basis of their affinity and compatibility with the target analyte.

The physical characteristics of the sensing material, such as light absorbance, electrical conductivity, permittivity, and work function, are altered during the identification process between these components and the analyte. These alterations are

converted into signals that can be detected, such as changes in color, voltage, current, resistance, or impedance.⁸⁴ The work function indicates the amount of thermodynamic effort required to extract one electron from the material's Fermi level, whereas the permittivity indicates the material's ease of electromagnetic radiation transmission.

Electrochemical sensors for MP detection need transducers assembled within a carefully designed sensing interface; ideally, these sensors can be fabricated into a portable unit. These sensors require sensor materials with specific binding sites and recognition abilities to selectively identify the target analyte.^{85,86} Attaining the requisite sensitivity and selectivity requires the capacity to design specialized recognition sites. A typical architecture of the sensor design for MP and source of exposure detection is shown in Fig. 5. A successful MP-monitoring sensor should operate on par with or better than conventional chromatographic methods. These sensors ought to be lightweight, affordable, and simple to operate. They should deliver accurate, real-time data with little sample preparation.

4.2. Electrochemical identification of MPs

Distinctions between MPs and other particles in an electronic field in terms of their electronic characteristics provide the basis for electrochemistry-driven MP identification. By qualitatively and quantitatively assessing the electrochemical responses (such as current and impedance) of mixed particles, it is possible to achieve simultaneous material identification and sizing. Table 8 provides a summary of the properties of the three sample systems for the electrochemical sensing of MPs, which are addressed in this section.

4.2.1. Resistive pulse sensor. By observing the transient current changes brought about by the passage of an analyte *via* a restricted sensing region, resistive pulse sensors (RPS) are able to distinguish between the numerous analytes of MPs on the basis of their size, shape, charge, and concentration.^{90,91} Reports say that the flow RPS made using an additive fabrication



Table 7 Parameters affecting the sampling of MPs in various environmental matrices as well as the corresponding analytical methods utilized to characterize them^a

Matrix	Sampling		Equipment	Extraction	Analytical technique	Ref.
	Time elapsed (quantity collected)	Sampling				
Water	2 h (between 2.3 and 310 million L day ⁻¹) 30 min	Stacked Tyler sieves (0.355 mm and 0.125 mm stainless steel mesh) Manta trawl (rectangular opening 16 cm high by 61 cm wide, 3 m long, 333 µm mesh)	—	—	Stereo microscope	63
	15–20 min (total of 2 L)	Neuston nets (0.52 × 0.36 m) of 333 µm mesh	—	Py-GC-MS	Micro-FTIR spectroscopy Py-GC-MS	64
	<10 L 30 min	Telescopic sampling pole Plankton net (153 µm)	—	32 µm steel-wire sieve and saturated NaCl solution	Stereo microscope and ATR-FTIR	65
Water, biological tissue	5 min (water, flow rates between 0.11 and 5.04 m s ⁻¹), 30 min (biological tissue)	80 µm mesh conical net/seine nets, gillnets, conventional tackle, and minnow traps	—	10% NaClO, HNO ₃ : NaClO (1 : 10 v/v)	Stereo microscope	66
Water, sediment	—	1, 2, 4 and 10 mm mesh size 0.25 × 0.25 m ² , 5 mm sieve	—	Top layer of sediment (3–6 cm) Directly from the sediment to a depth of 2 cm	TED-GC-MS ATR-FTIR	67
Sediment	—	Metal spoon	—	Surface layer (depth of 20 m)	Stereo microscope, SEM, Py-GC-MS	68
	—	Stainless steel shovel and 20 × 20 cm wooden frame	—	Munich plastic sediment separator. Centrifugation tubes with sieves (750 mm mesh)	Stereo microscope, micro-FTIR spectroscopy, SEMEDS, ICP-MS	69
	4–6 L	Sediment cores (diameter of 10 cm to a depth of 5 cm)	Concentrated ZnCl ₂ solution (1.7–1.8 kg L ⁻¹)	Concentrated ZnCl ₂ solution (1.7–1.8 kg L ⁻¹)	Micro-Raman spectroscopy	70
	—	Stainless steel scoop (10 cm depth) in order to fill a 1 L glass Kilner jar	Saturated NaCl solution and 30% H ₂ O ₂	Saturated NaCl solution and 30% H ₂ O ₂	Raman spectroscopy	71
	3 kg	Ekman dredge	69–71% HNO ₃	—	Stereo microscope and ATR-FTIR	66
Biological tissue	Whenever 3 clams of 40–45 mm were retained	0.5 × 0.5 m ²	—	—	Stereo microscope	72
	Until approximately 50 mussels were collected	Tweezers	30% H ₂ O ₂	—	Stereo microscope, micro-FTIR spectroscopy, SEMEDS	73
	Overnight/4 weeks	Gillnets (mesh of 50 mm) per cages	—	—	Stereo microscope, ATR-FTIR	74
	—	Baka 44/60, 40/60 and GOC 73 trawl gears	Dried samples, NaOH 1 M	—	Inverted microscope and stereo microscope	75
	—	90 mm GF/A 1.6 mm glass fibre filters and magnetic hot plate stirrer	—	Digestion solution (KOH 10% solution, 60 °C, 24 h)	Py-GC-MS, Raman spectroscopy	76
	—	Gillnet, demersal trawl	—	SDS, protease, chitinase and H ₂ O ₂ treatment; vacuum dried samples, petroleum ether (60/80)	Py-GC-MS	77
	—	—	—	Dried fish and excised organs or eviscerated flesh	Micro-Raman spectroscopy, FESEM-EDX	78

^a ATR-FTIR: attenuated total reflectance-Fourier-transform infrared spectroscopy; FESEM-EDX: field-emission scanning electron microscopy with energy-dispersive X-ray spectroscopy; FTIR: Fourier-transform infrared spectroscopy; ICP-MS: inductively coupled plasma-mass spectrometry; Py-GC-MS: pyrolysis-gas chromatography-mass spectrometry; SEM: scanning electron microscopy; EDX: energy-dispersive X-ray spectroscopy; TED-GC-MS: thermal extraction desorption-gas chromatography-mass spectrometry.

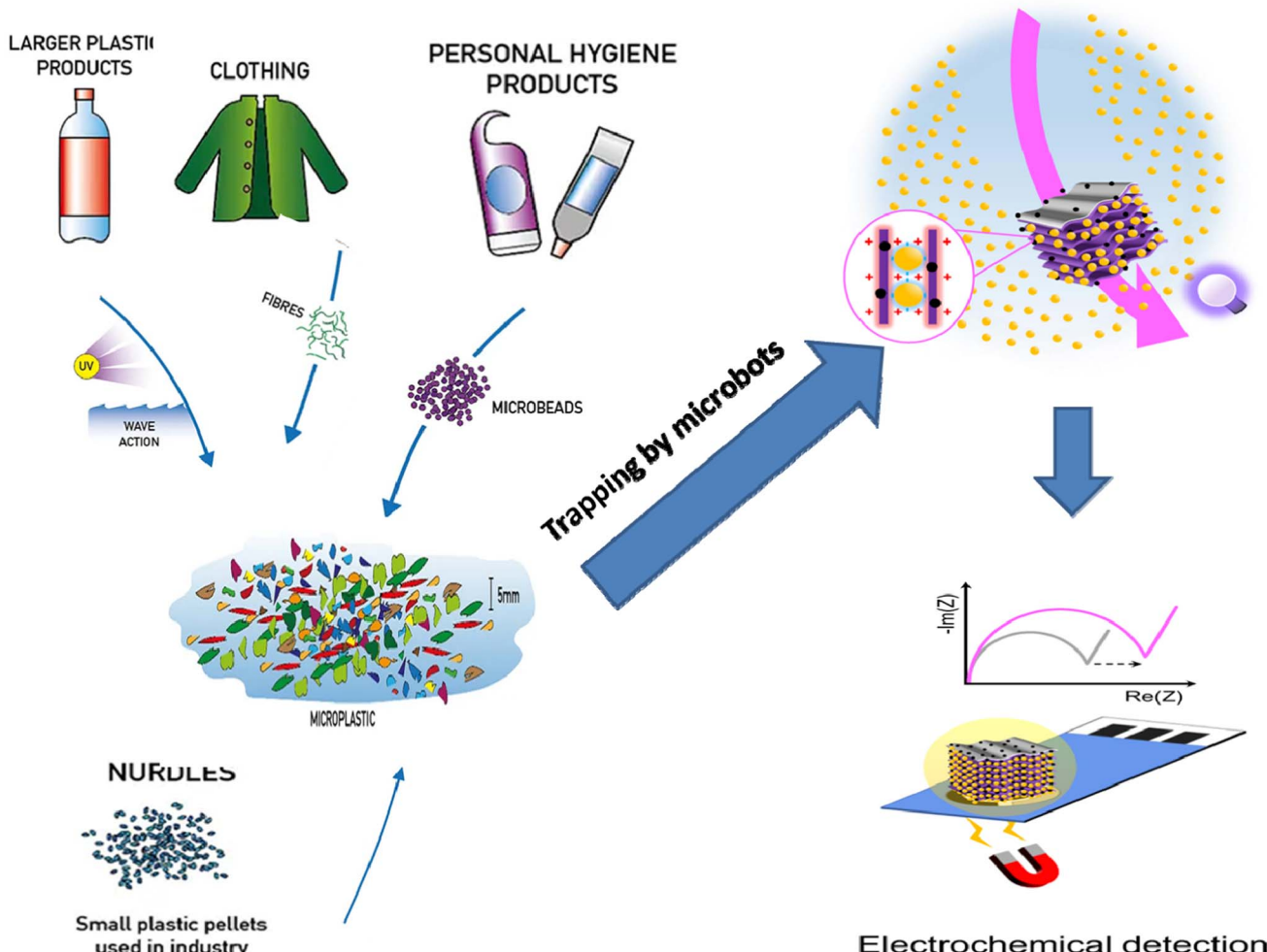


Fig. 5 Sources of MPs and illustration of sensor design for their electrochemical detection.

method can measure several different particles at very high speeds (1 mL of liquid per min) and salt concentrations ranging from 2.5×10^{-4} to 0.1 M.⁸⁷ The particles can be of any size between 2 and 30 μm . When calibrant, algae, and MPs pass through the sensing zone, their current responses significantly

vary. An average size of 21.9 μm and a concentration of 6.52×10^3 particles per mL were found in the tea sample after MPs were detected using the RPS. It should be observed that the signal is clearly affected by the ionic strength of the electrolyte (KCl), with lower ionic strengths being associated with

Table 8 Overview of some of the electrochemical techniques used to detect MPs

Method	Plastic type	Operational conditions	Advantages	Disadvantages	Ref.
Resistive pulse sensor	Poly styrene (PS)	<ul style="list-style-type: none"> Concentration range: up to 6.52×10^7 particles per L Sizing range: 2–30 μm Flow rate: up to 1 mL min⁻¹ 	<ul style="list-style-type: none"> High concentration sensing range Easy to use Suitable for most plastics Simple to carry out 	<ul style="list-style-type: none"> Restricted size range 	87
Impedance spectroscopy	Poly ethylene (PE)	<ul style="list-style-type: none"> The concentration range is 6.52×10^7 particles per L The size range is 300–1000 μm The flow rate is up to about 100 mL min⁻¹ 	<ul style="list-style-type: none"> Quite accurate Works with the majority of plastics Wide range of sensors 	<ul style="list-style-type: none"> Sophisticated sensor manufacturing procedure Sophisticated data processing Comparatively poor MP recovery rate 	88
Single micro-particle electrode	Poly ethylene (PE)	<ul style="list-style-type: none"> Concentration range: up to 1.4×10^{11} particles per L Sizing range: 1–10 μm 	<ul style="list-style-type: none"> High size precision (<2 μm) A wide concentration sensing range 	<ul style="list-style-type: none"> Restricted size range Intricate operation 	89



conductive pulses and reversed pulse directions. Furthermore, the porosity, conductivity, and shape of the MPs would affect the signal's characteristics.

4.2.2. Impedance spectroscopy. In impedance-spectroscopy-based sensing devices, electrodes measure how the impedance changes as small particles move through a medium. The impedance data from this measurement can be used to look into the particles' properties.⁸⁸ The impedance shift at higher frequencies can reflect both size and internal electrical properties of the particles, whereas at lower frequencies, it is directly proportional to the volume or size of the particles. Therefore, in order to distinguish between the effects of particle size and type, experiments are simultaneously conducted at both low and high frequencies.^{92,93} There is a way to tell the difference between PE MPs (212–1000 μm) and interferences of the same size (for example, organisms and seeds) in tap water (with an average flow rate of $103 \pm 8 \text{ mL min}^{-1}$) by looking at how the impedance changes at 1.1 MHz and 10 kHz. It is evident that the location of the particles and the corresponding impedance change allow for the differentiation of PE MPs, seeds, and living creatures in tap water.⁸⁸ Even though misclassification happens 1% of the time, a microcontroller can

sort the MPs using this straightforward logic. Further, a straight-line connection is found between the MPs' diameter and the cube root of the real impedance change at 10 kHz. This allows the size measurements and MP identification to happen at the same time in water.

4.2.3. Effects of a single microparticle electrode. Scientists have come up with the idea of single microparticle-electrode impacts (SMEI), which is when an electrode hits a biomolecule or MP/NP and changes the current response by making it stronger or weaker.⁹⁴ When electrically insulating PE MPs strike a carbon electrode in an SMEI system, the result is transient current responses or current spikes. It is interesting to note that the dissolved oxygen content of each impacting PE microbead, which is proportional to the particle's volume, is connected to the produced current spikes. As a result, the form and frequency of the current spikes provide useful information about the quantity and size of the PE MPs.⁸⁹ The size results match the SEM measurements of particles smaller than 2 μm , with 0.74 g L^{-1} of PE MPs in a 20 mM NaCl electrolyte. The SMEI system has a higher standard deviation for the coarse-sized fraction, which could potentially be attributed to the *in situ*-generated clustered MPs. The number of observed impact events during

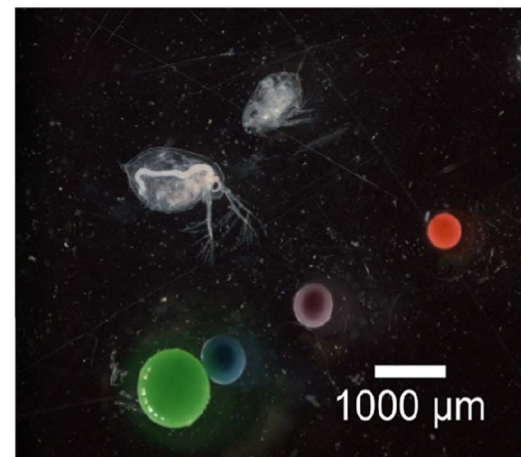
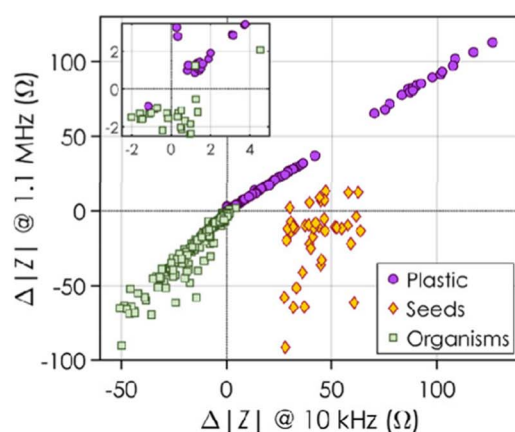
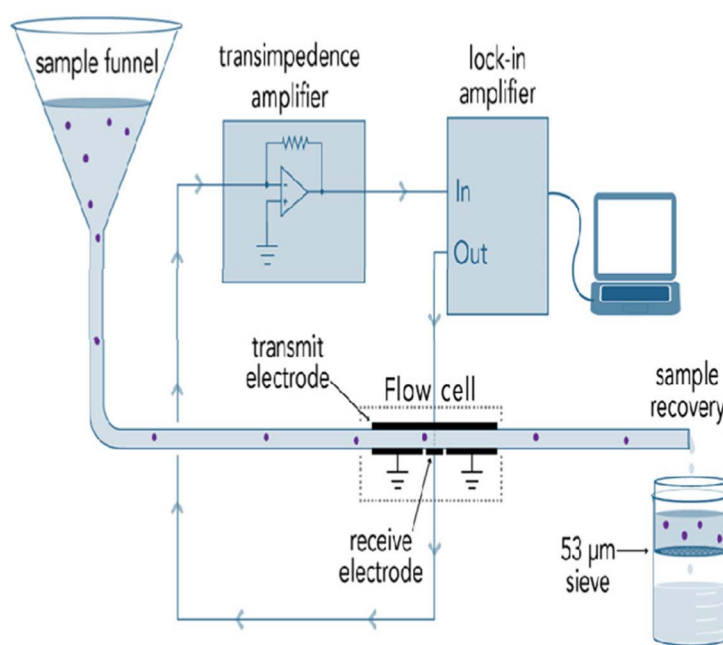


Fig. 6 Diagram showing how MPs are measured for impedance while going through a flow cell. Variation in impedance according to the type of particle: plastic, seeds, or organisms.⁸⁸



the particle-impact process can also be used to precisely measure the concentration of PE MPs in a large range, from 3.6×10^9 to 1.4×10^{11} particles per L. All things considered, the SMEI technique appears to have promise for effectively identifying the quantitative size and quantity of insulative MPs in water. The RPS, impedance spectroscopy, and SMEI work in different ways to make it possible to very accurately sense the electrochemical state of MPs.

5. Electrochemical sensors for MP determination

Although electrochemical sensors have been extensively used to detect environmental contaminants, including leachates from MPs,⁹⁵ their applications for MP detection are fairly rare and have only recently come to light. Because of the low cost, low response time, easy operation, and mobility of electrochemical devices, the development of electrochemical sensors for MPs is especially interesting. In contrast to other traditional techniques, electrochemical approaches can be readily expanded to the in-field assessment of many sample kinds, as they do not require any preliminary MP isolation or purification. Amperometry, voltammetry, and label-free electrochemical impedance spectroscopy (EIS) have been used to achieve MP monitoring and sensing up to this point.

Using an EIS-based sensor in combination with flow cytometry, PE MPs were detected.⁸⁸ The detector consisted of a flow cell for EIS detection with Au-plated circuit boards that held all the Au-plated electrodes and a flow cytometer for particle detection (Fig. 6). The idea behind this approach was that the real fraction of impedance should change according to the particle volume at low frequencies. Therefore, at low frequencies, the change in impedance is proportional to the MP particle volume and is caused by the MP particles flowing over the electrodes (Fig. 6). The plastic and biological particles may be differentiated with this method at any observed frequency. It is interesting to note that although MPs caused a positive change in impedance, biological particles caused a negative change, making this platform suitable for MP detection in complex media.

The sensitivity of the sensor and its capacity to detect and distinguish between different MP sizes were assessed in real time by testing the clean water flow against water that had been spiked with known MPs. The impedance would alter, producing peaks and enabling the measurement of particle size if MPs were present in the fluid under test. Using the linear relationship between the particle diameter and the cubic root of the actual impedance change, the impedance change and particle size were related. As a result, this sensor could measure and size both PE MPs (212–1000 μm) and MP beads (containing

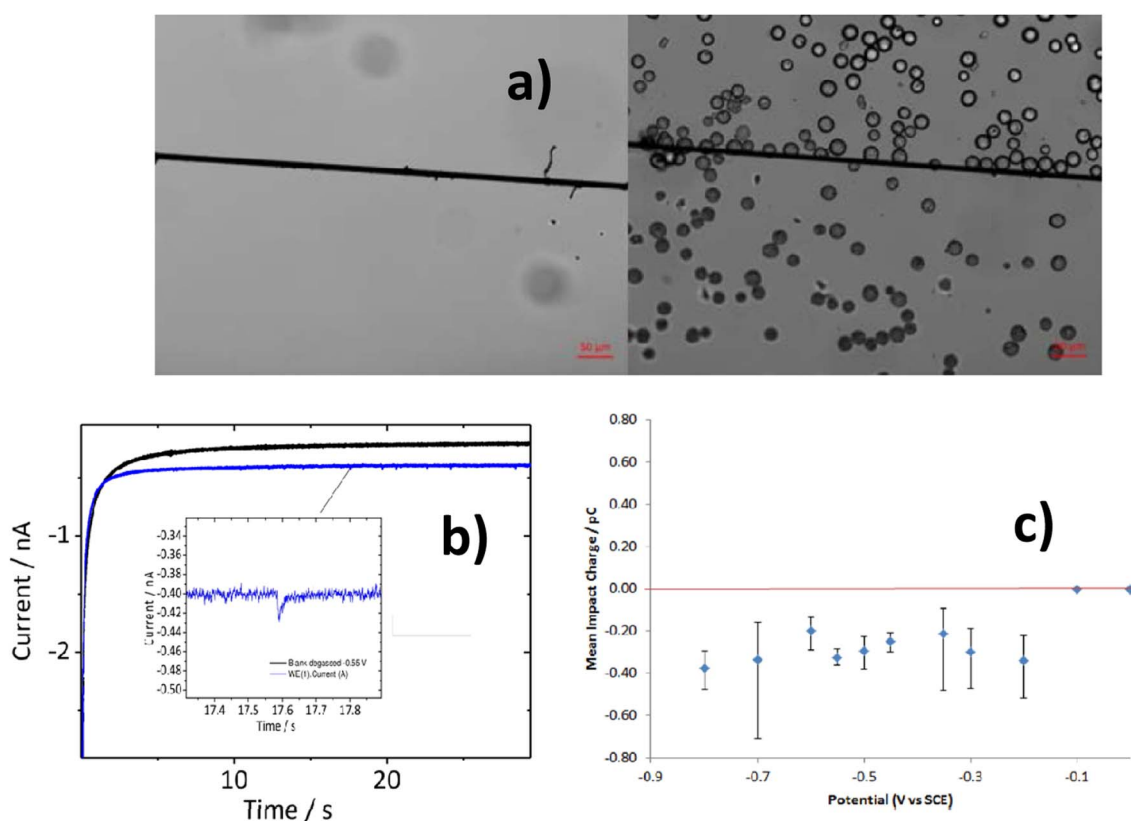


Fig. 7 SEM images for the carbon fiber electrode with and without MPs (a). An enhanced view of a transient current–time signal resulting from a collision event between MPs and electrode can be seen in the inset of the chronoamperogram (b), in which the blue line represents the PE MPs and the black line represents the MP-free state. Histogram showing the distribution of integrated charges extracted from spikes resulting from chronoamperometric measurements (c).



biological components in the 210–1200 μm size range). According to Collson and Michel,⁸⁸ the sensor had a recovery rate of over 90% for MPs in the 100–300 μm size range and a false-positive rate of 1% when it came to identifying biological materials as MPs. Although MPs were effectively detected using EIS—a label-free approach—the next generation of sensors should focus on detecting MPs in the field as well as differentiating between various types of MP particles besides PE.

Additionally, spherical PE MPs (1–22 μm) have been detected using particle impact electrochemistry.⁸⁹ A popular technique for studying particles suspended in solution is the particle-electrode impact method. Measurements using chronoamperometry revealed a fast current response due to particle interaction with the carbon fiber microwire electrode. The undivided three-electrode setup used in the electrochemical analysis setup was maintained at a particular voltage in order to observe the desired reaction.

A particle-electrode collision caused a signal change that was monitored. Fig. 7 shows the transient current response, or spike, that was produced when the particles collided with the electrode and was examined to identify the MP. The current spike in the chronoamperogram was caused by a collision between PE MP particles and the working electrode. This happened because the amount of oxygen in PE MP particles declines. With an R^2 value of 0.96, an excellent connection was found using this method between the MP particle concentrations and the frequency of spikes.⁸⁹ Comparing this detection method to other approaches, the MP measurement was shown to be more accurate and consistent overall. Although the technique was used to find electrically insulating MPs, conductive particles might also be found.

The serial faradaic ion concentration polarization technique is an alternate method for MP detection.⁹⁰ Based on their electrophoretic mobilities, the MPs are sorted by the faradaic ion concentration polarization. Their electrophoretic mobilities have an impact on the interaction between the MPs and the electric-field gradients. Elevated electrophoretic mobility particles concentrate at a low-electric-field site. On the other hand, lower electrophoretic mobilities are concentrated in greater electric fields. MPs were sorted into distinct chambers using a trifurcated trunk, with the bipolar electrodes positioned close to the trunk. The ion depletion zones and electric-field gradients are positioned across the opening of the split channel by the BPE. As a result, the MPs are sorted, with the higher-electrophoretic-mobility particles going into the lower channel and the lower ones flowing into the upper channel. The flow of PS MPs in a trifurcated microchannel was managed by the faradaic ion concentration polarization. MPs may be continuously focused on, sorted, and separated using this strategy. Because MPs interact with electric-field gradients and can be seen with optical and fluorescence microscopy, they could be put into groups based on how fast they move through an electric field.

MPs were sorted according to their size and electrophoretic mobilities in the trifurcated microchannels, which were oriented toward the MPs' trajectory by the electric-field gradients as they approached the cathodic end of the bipolar

electrodes (Fig. 8). In addition to isolating or pre-concentrating different MPs on demand without the need for membrane-based separation, this strategy makes the technology even simpler by allowing MPs in water systems to be continuously monitored and in real time. Crucially, the ability of the faradaic ion concentration polarization to interface with different methods would provide sensor developers and designers with more creative freedom; yet, MP detection has not taken advantage of this potential.

More recently, the multiuse RPS served as the foundation for the development of an inexpensive, high-throughput technique.⁸⁷ This microfluidic sensor, which was constructed on a silver wire, used an analyte's (MP) translocation *via* a tight constriction to detect changes in current. The gadget was utilized to track MPs in tea bags and identify algae to show the value of this technique for counting and tracking MPs in the presence of biological particles, which is still a difficult task.

Images of the base unit, lid, sensing zone, and electrodes are displayed in Fig. 9a–d. The lids can have various designs. The first is a flat surface that resembles a Gen 1 acetate film. In the second design, a ridge is positioned to extend into the channel and protrudes 1 mm from the surface. The third, shown in Fig. 9e, features several ridges that can alter the channel's and sensing region's shape. The gadget features printed screw threads to attach pumps because it was made to be integrated into a flow system. The integration of the electrodes' screw threads is a second change from Gen 1, in addition to the lid design (Fig. 1d). The device can quickly screen liquid volumes of 1 mL min^{-1} for the presence of algae and MPs.

There have also been reports of other dual-mode sensor instances, in addition to impedimetric and amperometric sensors for MPs. A sensor for exoelectrogenic-biofilm-based PE MP detection has been created, employing voltammetry and impedance.⁹⁷ The use of electroactive bacterium film in microbial electrochemical systems has shown a lot of promise as an energy-efficient method for MP detection in wastewater. The biofilms were subjected to MPs to find out how they responded as well as to examine the electrochemical characteristics, shape, EPS, and microbiological organization of the biofilms. A carbon fiber brush as the anode served as the working electrode in this three-electrode arrangement, which also included titanium woven wire mesh serving as the cathode counter electrode and Ag/AgCl as the reference electrode. PE-MP binding to the biofilm was the reason of the elevated internal resistance.⁹⁷ Using Nyquist plots, EIS was used to quantify the impedance of the MFCs and MECs both before and after binding. The presence of PE-MP boosted the cells' resilience. On the basis of the equivalent circuit model, the charge transfer resistance (R_{ct}) was responsible for most of the resistance. Furthermore, a notable rise in R_{ct} may be caused by an increase in the number of dead cells induced by the toxic effects of PE-MPs. When PE-MPs were present, the current density in the microbial fuel cells remained constant. On the other hand, as MP concentrations grew in the microbial electrolysis cell, the current signal decreased; this tendency persisted for more than 42 days (Fig. 10). The decrease in signal was caused by an increase in internal resistance as a result of PE-MP binding to the biofilm.⁹⁷ Therefore, the



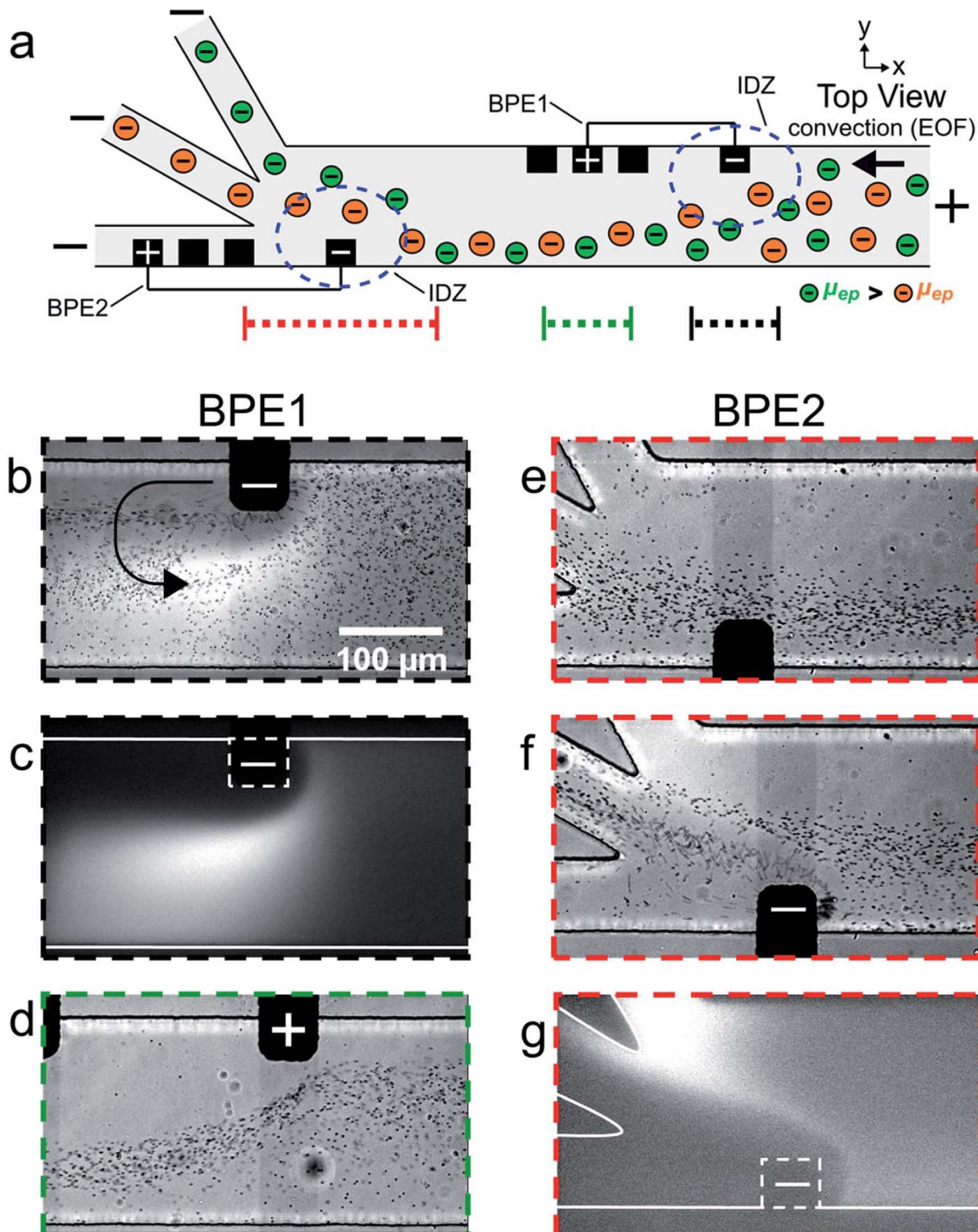


Fig. 8 (a) Schematic of the microfluidic configuration used for serial faradaic ion concentration polarization experiments. For frames (b–e), only BPE1 was active. For frames (f) and (g), both BPE1 and BPE2 were active. (b–g) Series of optical and fluorescence micrographs showing the location of mP1 and BODIPY2_ during serial faradaic ion concentration polarization. With reference to the three dotted lines at the bottom of (a), the micrographs were captured along the portion of the channel length indicated by (b and c) the dotted black line; (d) the dotted green line; and (e–g) the dotted red line. The curved black arrow in (b) indicates the location and rotation direction (counterclockwise) of the vortex downstream of the cathodic pole of BPE1.⁹⁶

microbial electrolysis cells would need to be used to measure the concentration of the MP present to sense PE-MPs using exoelectrogenic biofilms.

Furthermore, to expand its application into a real-world environment, a microbial electrolysis cell could be utilized in the future to differentiate between MP types and sizes. The same



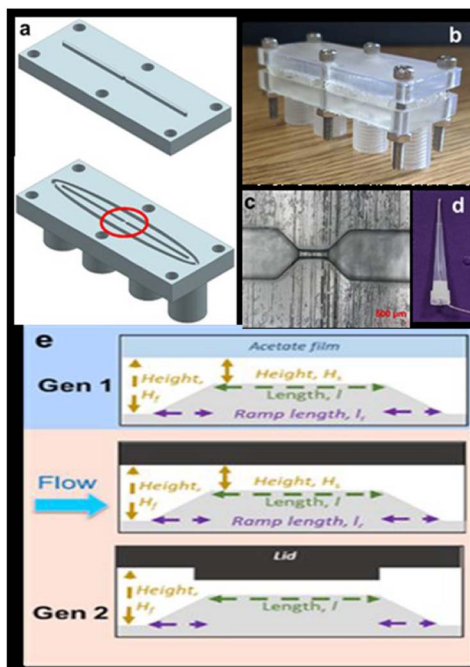


Fig. 9 (a) Design of the device, (b) image of the assembled device, (c) an optical microscope image of the sensing zones circled in part (a), (d) silver wire embedded within a pipette tip, and (e) schematic of the sensing zone when using an acetate film, a flat lid, and a ridge lid.⁸⁷

technique might be applied to MP remediation in microbial electrochemical wastewater treatment systems. Although MP cleanup has used electrochemical techniques, MP sensors are still not well understood. In order to effectively monitor MPs, further research and development of sensors is required, despite the existence of successful electrochemical sensors.

There are numerous MP cleanup procedures that give excellent chances for creating new testing instruments. By applying anodic oxidation or reduction of MPs, for instance, MP remediation can be indirectly employed for sensing applications such as MP coagulation detection, MP byproduct sensing, MP interactions with biofilms or other biologicals, and so on. MP identification and differentiation; standardized methods for MP isolation, characterization, and environmental monitoring and tracking; and MP detection in a range of sample types (soil, soil runoffs, various watersheds and pools, wastewaters and industrial effluents, agricultural watersheds, and atmosphere) are the current challenges facing the advancement of electrochemical sensors for MPs. The electrochemical gadgets may also offer a way to continuously track MPs' movements and destiny in their surroundings and in real time. The breakdown of MPs into smaller NPs is a crucial component of MP chemistry; thus, it is also necessary to identify and track plastic particles downstream from the MPs. Since MPs readily interact with and absorb other biomolecules, understanding their basic interactions through electrochemical techniques may initiate new possibilities for MP cleanup and monitoring.

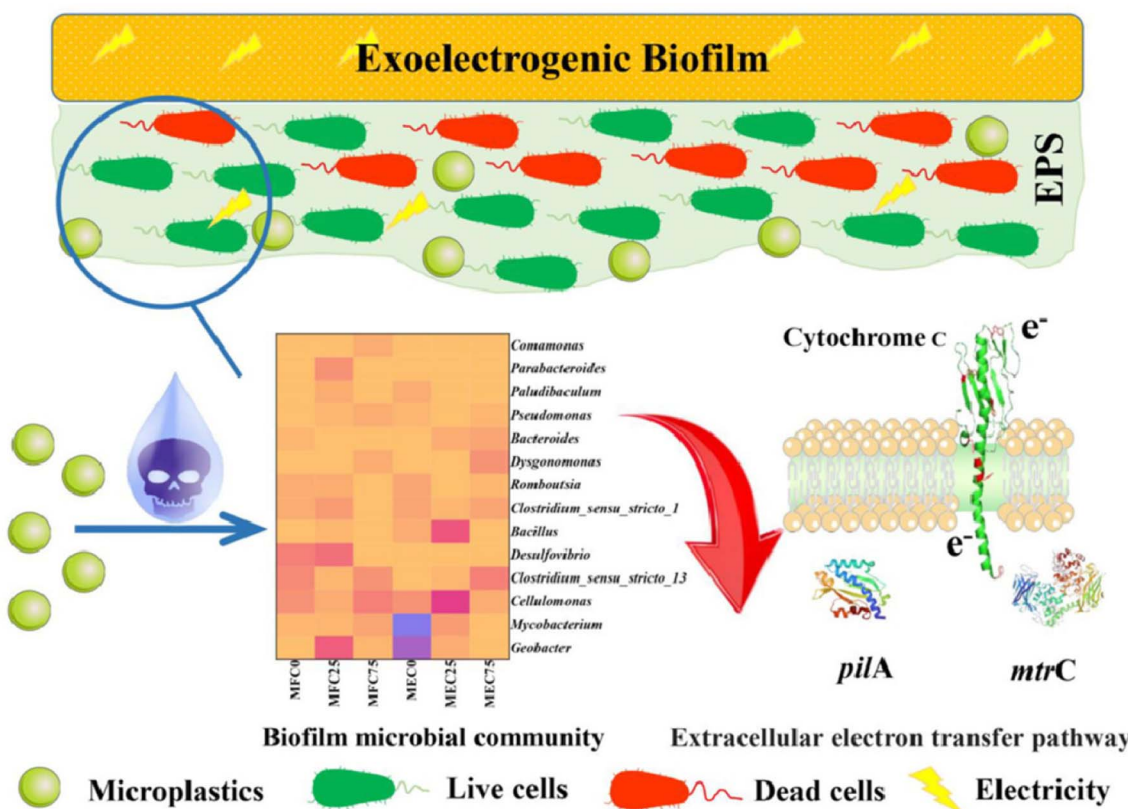


Fig. 10 Schematic of exoelectrogenic biofilm on an electrode surface and its interactions with MPs.⁹⁷



6. Important elements in MPs' electrochemical identification

Electrochemical systems are susceptible to the following factors: applied current density or potential, analyte property, electrode activity, and electrolyte property. The influence of experimental conditions should be examined in order to direct future research for the electrochemical identification and removal of MPs in water matrixes. The properties of the MPs, the applied current density or potential, and the electrolyte are thoroughly covered in this section.

6.1. Plastics' property

The size, shape, density, and type of plastics are significant characteristics that influence their electrochemical behaviors. Because PVC has a far lower electrical resistance than PE, it is more susceptible to electro-precipitation and electro-flootation. Furthermore, electro-flootation makes it simpler to remove MPs with densities lower than that of water (like PE) from water.⁹⁸ The ideal removal parameters for different-sized MPs vary due to the unique electrical characteristics of polymers of varying sizes.⁹⁹ The electrocoagulation of small MPs (20–60 μm) requires a current density of 2.88 mA cm^{-2} and a pH value of 4, whereas large MPs (350–1500 μm) require a current density of 2.88 mA cm^{-2} and a pH value of 7, possibly due to the larger bubbles generated at neutral pH.⁹⁹ Furthermore, the shape of MPs and NPs affects the effectiveness of electrochemical removal. Fiber MPs (CA and PP) have a greater electrocoagulation removal efficiency than granular MPs (PE and PMMA).¹⁰⁰ As of right now, only plastic beads have been investigated for electrochemical sensing; other MP forms have received less attention. It is important to look into the electrochemical identification of MPs with various shapes (*e.g.*, sheet, fiber, and film) in light of the shape-dependent electrical characteristics of plastics (Shimizu *et al.*, 2017).⁸⁹

6.2. Density of current

The applied voltage or current density can directly affect the reaction efficiency. Higher current densities (voltage intensities) generally encourage the elimination of MPs through processes like electrocoagulation. In the electrocoagulation/electro-flootation process, the higher current density helps flocs form (like Al(OH)_3) and speeds up the release of the coagulating species from the Al anode. The cathode also releases more hydrogen bubbles when the current density is high. These bubbles help get rid of MPs and provide electro-flootation by sticking to agglomerates.⁹⁹ In the insulator-based di-electrophoresis process, on the other hand, a high voltage intensity would make particles stick together, which could make the device less effective at separating things by blocking its tiny gaps.¹⁰¹ In addition to electrochemical performance, one should consider the high energy consumption brought about by the high current density.

6.3. Electrolyte

Many studies have been conducted on the effects of pH, carrying interfering compounds, and electrolyte concentration on electrochemical studies involving MPs. Better conductivity from

a larger electrolyte concentration benefits the electrochemical process, too.¹⁰² The electrochemical signal in the RPS-based sensing technique is clearly influenced by the ionic strength of the electrolyte (KCl), with conductive pulses and reversed pulse directions being observed at lower ionic strengths. An increase in electrolyte concentration during the electrocoagulation process might increase the MPs.¹⁰⁰ Additionally, Na_2SO_4 -electrolyte-derived reactive species (SO_4^{2-}) ($\text{SO}_4^{2-} \cdot$) electro-oxidize PS MPs during the electrooxidation – H_2O_2 process, and a concentration increase of Na_2SO_4 from 0.007 to 0.03 M results in noticeably better electro-degradation kinetics. Nevertheless, an additional concentration increase to 0.06 M has no beneficial effect. This could be because there is not enough H_2O_2 generated *in situ* to react with $\text{S}_2\text{O}_8^{2-}$ to form SO_4^{2-} species.¹⁰³

The pH of the electrolyte controls the characteristics of surface charge and the chemical structure, as well as the forms of many substances such as coagulants, analytes, and electrode materials. A near-neutral solution is advised to help create big flocs in the electrocoagulation of MPs.^{98–100,102} Comparable outcomes are shown when bisphenol A (BPA) is electrochemically detected. The acid–base dissociation of BPA is influenced by the pH of the electrolyte, which leads to a pH-dependent oxidation current and potential. When the pH goes up, the $-\text{COOH}$ groups on multiwalled carbon nanotubes (MWCNTs) become more deprotonated. This makes them act like acidic anions and would reject anionic BPA species if they were used as a BPA sensor in an alkaline solution (MWCNTs- β CD). Accordingly, BPA may be detected at a pH level of 7.4, which is almost neutral.¹⁰⁴

Metal ions (such as Pb, Ni, Cu, and Cr) can interact with naturally aged MPs by surface complexation and electrostatic adsorption. It is interesting to note that the presence of MPs and metal ions together promotes the removal of metal ions without altering the MPs' ability to electro-coagulate.¹⁰⁵ An increase in Cl^- ion concentration has no appreciable impact on the removal performance of PE MPs during electrocoagulation.¹⁰² As of right now, research on the function of dissolved organic materials in the electrochemical removal of MPs is lacking. Usually, metal ions, anions, and organics are added to the electrolyte to test how selective the electrodes are for finding contaminants that come from MPs. Some small molecules (like uric acid and ascorbic acid) and inorganics (like KBr, NaNO_2 , KCl, and NaNO_3) do not have much of an effect. However, organics (like tetrabromobisphenol A (TBBPA)) that share a chemical structure with the target pollutants (like BPA) may make identification harder.¹⁰⁶

7. Fabrication of electrochemical sensing devices

The construction of electrochemical sensing devices requires the thoughtful design and deliberate selection of electrode materials and device components that are appropriate for the intended use. For the device to consistently function in its intended environment, all of its parts must be mechanically strong and chemically stable. The next stage is to determine the



best technique for immobilizing the active material on the surface of the working electrodes after selecting the electrode materials. The approach for altering the electrode and the types of materials utilized can influence the electrode's thickness, adhesion, mass loading, distribution, and homogeneity on the surface. A range of electrodes, including metal foam, carbon-based conductive paper, glassy carbon, and ITO/FTO glass, are accessible and can be utilized as platforms for electrochemical devices involved in MP monitoring. An overview of the many techniques for depositing materials at electrode surfaces is given in this section. These techniques address issues of reproducibility, compatibility, and large-scale processing—all of which are crucial elements in the creation of sensing devices.

7.1. Electrophoretic deposition

A tried-and-true technique for directly growing thin films on electrode surfaces is electrophoretic deposition (EPD), which is renowned for its capacity to yield homogeneous coatings with adjustable thickness. Applying a DC electric field to a target substrate containing a suspension of the substance to be deposited is how EPD is carried out (Fig. 11). The electric field can create intricate 2D and 3D patterns on electrode surfaces by forcing the charged particles in suspension in an orderly manner toward the oppositely charged electrode.

Biological materials can also be deposited using this technique. Particle size, solvent type, electrophoretic mobility, and electric-field intensity can be changed to create films with diverse properties, such as thickness and homogeneity.¹⁰⁷ By providing 0.2–1.6 V, this technique has been used to deposit gold nanoparticles of various sizes onto ITO glass electrodes. Under ideal circumstances, nanoparticles of metal oxides and sulfides, including copper,^{108,109} iron,¹¹⁰ cobalt,¹¹¹ zinc,¹¹² and titanium,¹¹³ have been deposited. Graphene oxide,^{114–118} graphene quantum dots,¹¹⁹ carbon nanotubes,¹²⁰ and their composites were among the carbon-based materials that were deposited using applied voltages ranging from 2 to 100 V. These materials were chosen on the basis of their charge and mobility. Graphene composites were deposited on silicon wafers using higher voltages (300 V).^{121,122}

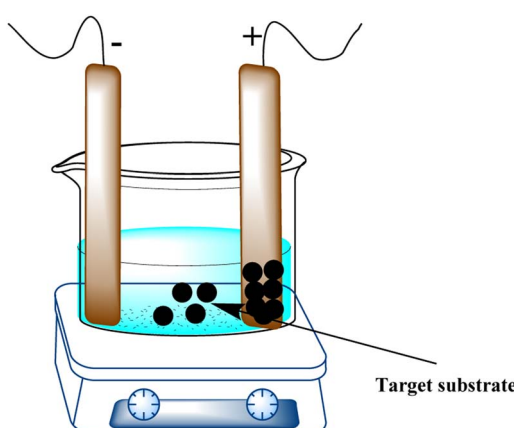


Fig. 11 Schematic of the EPD process.

Additionally, metal-organic frameworks (MOFs) were deposited using EPD to produce homogeneous films with the required shape and thickness. Because MOF particles have defect sites from missing linkers, they are partially charged. When a strong enough electric field is applied, these charges push the MOFs in the direction of the electrode that is oppositely charged. By using EPD to spread MOFs in toluene for different lengths of time, thin films of NU-1000, HKUST-1, UiO-66, and Al-MIL-53 have been micro-patterned. These films stick well to the glass electrode (FTO) and maintain their crystal structure.¹²³ To detect explosives and Cr³⁺ ions, an Ln-BTC MOF thin film was made by dispersing MOF in a CH₂Cl₂ solution.¹²⁴ Electrically conductive MOFs have recently attracted a lot of attention for thin-film manufacturing in order to be deposited as electrode materials for supercapacitors and batteries.^{125,126} To give an example, putting 0.5 V between the electrodes for 10 min made 2D MOF nickel-2,3,6,7,10,11 hexaaminotriphenylene (Ni₃(HITP)₂) nanosheets stick to nickel foam.¹²⁷

7.2. Layer-by-layer assembly

Another straightforward and reliable technique for creating multilayer films on solid substrates is layer-by-layer assembly (LBL). The electrode is alternately exposed to positively and negatively charged species in this “bottom-up” method. It is possible to create electrodes that are precisely suited by adjusting the distribution, thickness, and composition of the layers.¹²⁸ LBL does not need a conductive substrate, in contrast to EPD. Furthermore, the film can be deposited on substrates with various topologies, which makes it easier for 3D microstructured films to form.¹²⁹ Roll-to-roll,¹³⁰ spraying,¹³¹ atomization,¹³² magnetic assembly,¹³³ electrocoupling,¹³⁴ creaming,¹³⁵ and other processes are examples of LBL technology. The needs of the application and the substrate's ability to hold materials in place determine the choice of the substrate. This technique offers a broadly applicable way to embed active materials on electrode surfaces for target analyte binding or conversion. It is suitable for sequestering various sorts of materials in between the layers, such as metal nanoparticles. Surface-supported MOF thin films (SURMOFs) were made using LBL by first coating the support with a self-assembled monolayer (SAM) and then immersing it several times in solutions containing organic ligands and metal precursors, cleaning it after each immersion (Fig. 12a).^{136,137}

This process encourages the controlled development of highly aligned and uniform MOF sheets for gas sensing¹³⁸ and electrochemical catalysis.¹³⁹ Additionally, LBL enables the deposition of MNPs on solid substrates, giving good control over the density and distribution of NPs.¹⁴⁰ Electrostatic interaction was used to deposit Au NPs on graphene, which makes it easier to capture the injected electrons from the analyte (Fig. 12b).¹⁴¹ LBL assembly can be utilized to build substrates with tailored surface-enhanced Raman scattering (SERS), regulated Au NP size, and layer thickness for use in sensing applications.¹⁴² Shorter fabrication times can be achieved by automating the assembly process with this method. A thorough analysis released in 2016 covered various LBL assembly methods in great depth.¹⁴³

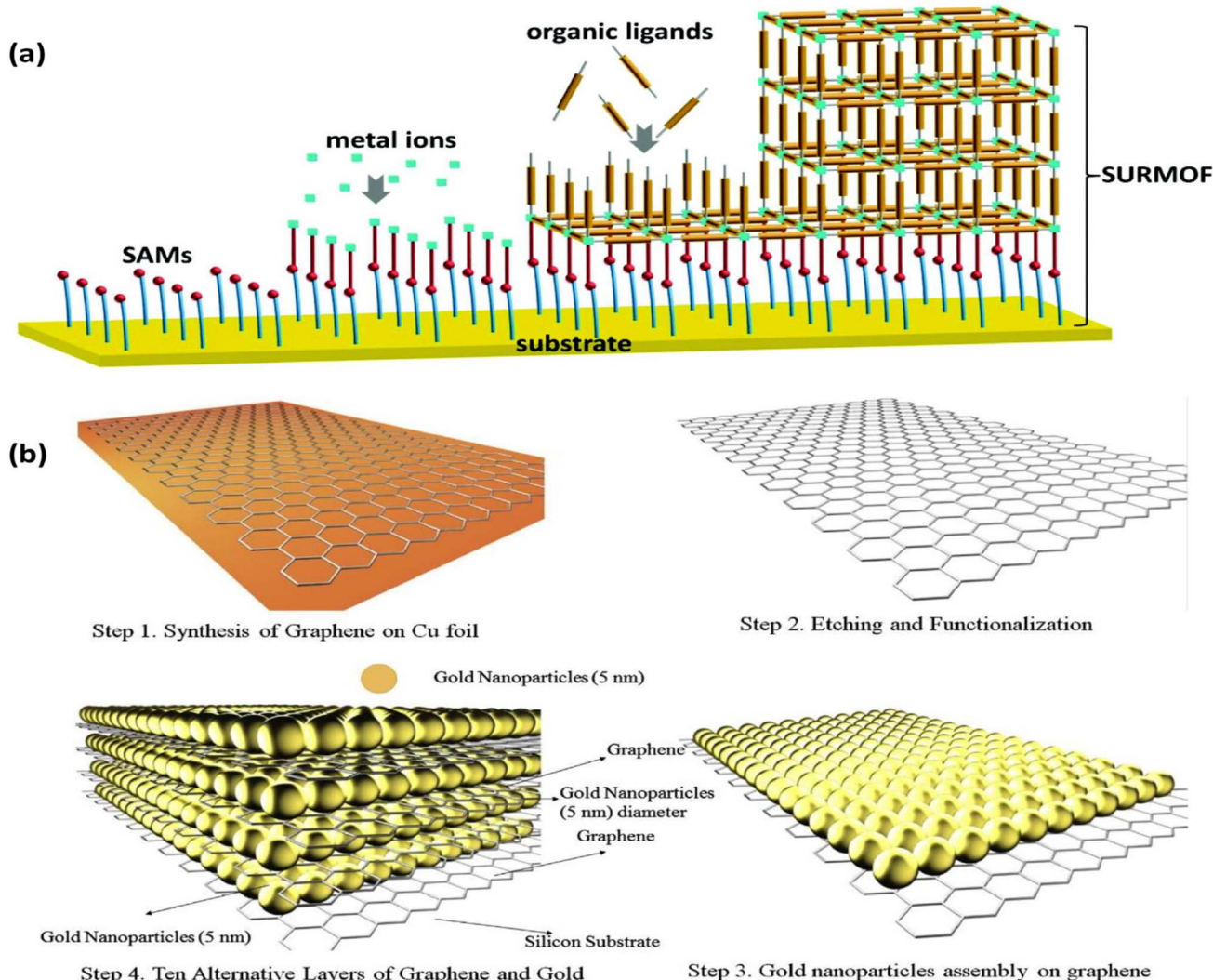


Fig. 12 (a) Illustration of SURMOF film formation via an LBL assembly.¹³⁹ (b) Schematic of the LBL assembly of Au NPs/graphene electrode. Reproduced with permission from ref. 141.

7.3. Direct 3D and inkjet printing

Large-scale production and quick processing times were made possible by the development of automated procedures, which were spurred by the rapidly growing interest in modern manufacturing. With the use of 3D printing technology, electrodes with a variety of shapes may be consistently produced, lowering the overall production costs and facilitating manufacturability.¹⁴⁴ The main obstacle in the development of 3D printable electrodes, electronics, and structures is the formulation of inks and materials that are printable. Metal oxides, carbon-based materials, ceramics, polymers, biological or biomimetic materials, and polymers are some examples of these materials. The direct printing of electrode materials enables high structural design freedom and hastens the compositional optimization of materials for sensing applications. There are several ways to print electrode materials, including binder jetting, fused deposition modeling (FDM), stereolithography (SLA), and direct ink writing (DIW), as shown in Fig. 13A–D.¹⁴⁵

The most popular 3D printing method, DIW, uses droplet-based and continuous filament techniques to create structures with a sub-50 μm resolution. This method depends on regulating the ink's viscosity, rheological characteristics, and shear-thinning behavior to promote smooth extrusion, enable printability, and preserve the printed shape. The difficulty with electrochemical devices is creating conductive inks and processes that can generate huge-volume shapes with high resolution.¹⁴⁶ SLA prints high-resolution structures by using photocurable polymers, which solidify during printing and allow for simultaneous laser scanning. Usually, this process entails printing non-conductive polymers onto conductive substrates and then covering the printed structures with a conductive layer.¹⁴⁷ Using carbon filaments that have been heated and extruded, FDM is a popular printing method for creating electrode materials. Nevertheless, commercially available filaments with weak conductivity, such as graphene or PLA, may need to be treated after printing.¹⁴⁸ Binder jetting works by putting binder droplets on a bed of powder, which is hardened



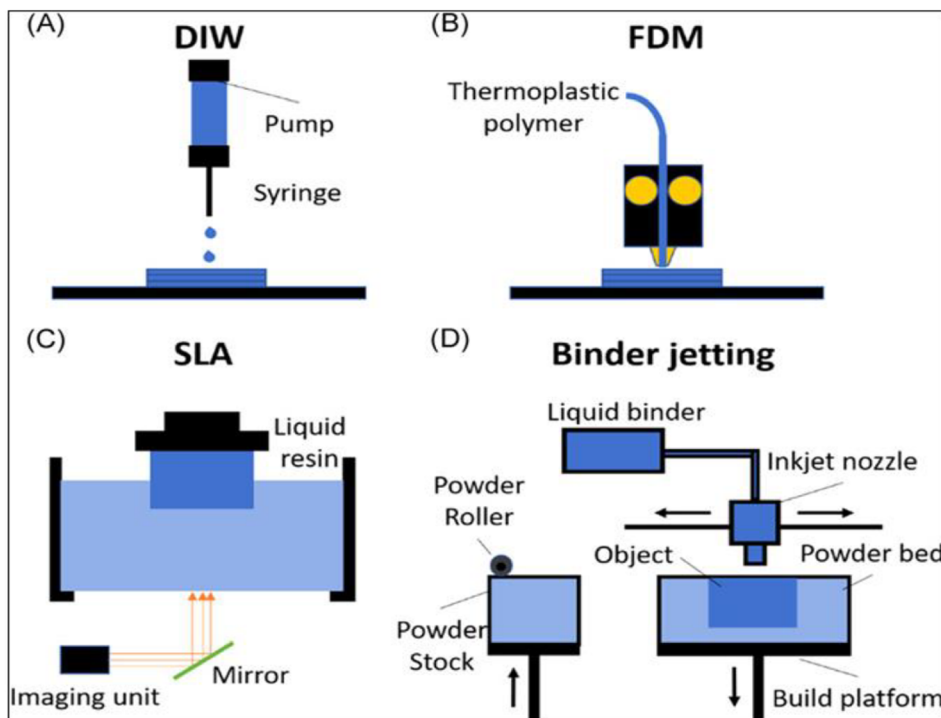


Fig. 13 Top image shows a schematic of the common printing techniques: (A) direct ink writing (DIW), (B) fused deposition modeling (FDM), (C) stereolithography (SLA), and (D) binder jetting.¹⁴⁵

to create 3D objects. Large structures with intricate geometries may be created using this method, which makes it suitable for use in industrial settings.

Electrodes can be printed by using conductive materials for the powder bed.¹⁴⁹ The quick output, repeatability of the electrode geometry, and ease of production have made printing sensor materials appealing. For example, cellulose paper and plastic substrates were inkjet printed with chemiresistor films and CNT designs.¹⁵⁰ At room temperature, the printed electrodes responded to NO_2 and Cl_2 gases in a reversible and selective manner. Charge-transfer p-type doping of CNTs serves as the foundation for the sensing process. As evident in Fig. 14a,b,¹⁵¹ nozzle-jet technology was used to print Ag-rGO composites onto a flexible substrate. These were then used as sensors to directly detect phosphate ions in eutrophic water. The printed sensor showed a low LOD of $0.2 \mu\text{M}$ and a linear range of $0.005\text{--}6 \text{ mM}$.

Screen printing is a simple, roll-to-roll method for creating inexpensive electrochemical sensors. It involves depositing layers after layers *via* open mesh screens that are specifically made to create a thick film. The conductive tracks can be built using a range of commercially available conductive pastes. The last layer necessitates the creation of a printable formulation and is usually tailored to incorporate the sensing material with or without an embedded enzyme. It is hard to put biocomponents like enzymes in the printed layer for biological sensors because they might not work during the serigraphy process or the curing step. The printed electrode can be interfaced with portable wireless potentiostats as a further step in the development of portable electrochemical sensing platforms. Additionally, methods for mass-producing

these sensors should be designed. Gravure-printed electrodes made of biocompatible inks with electrode morphologies that offer mechanical stability and produce 150 m of flexible rolls are a recent example of the roll-to-roll technique (Fig. 15).¹⁵² The low-cost electrochemical detection of heavy metals and tiny compounds has proven to be applicable. We used tyrosinase carbon enzymatic black ink to show that the technology can be used to print enzyme-based parts for making several electrochemical biosensors.¹⁵³ Printing offers benefits like low production costs, low repeatability, and manufacturability when it comes to creating sensors.

8. Future challenges

Even though MPs can be electrochemically detected in a fast, precise, selective, and reliable manner, there are still a few areas that require improvement in order to enhance electrochemical identification methods.

- Only plastic microbeads with known sizes and concentrations have been investigated for MP sensing; other MP shapes have received less attention. Determining the electrochemical reactions of various MPs and NPs in the sensing process is vital, given the size- and shape-dependent electrical characteristics of plastics.^{87,89}
- Because lab-made waters are used in current research on the electrochemical sensing of MPs, the effect of impurities found in real water, like metal ions, organisms, organic contaminants, and suspended particles, is not taken into account. For instance, metal ions would cause MPs to aggregate, which would have an impact on the sizing procedure. In light of

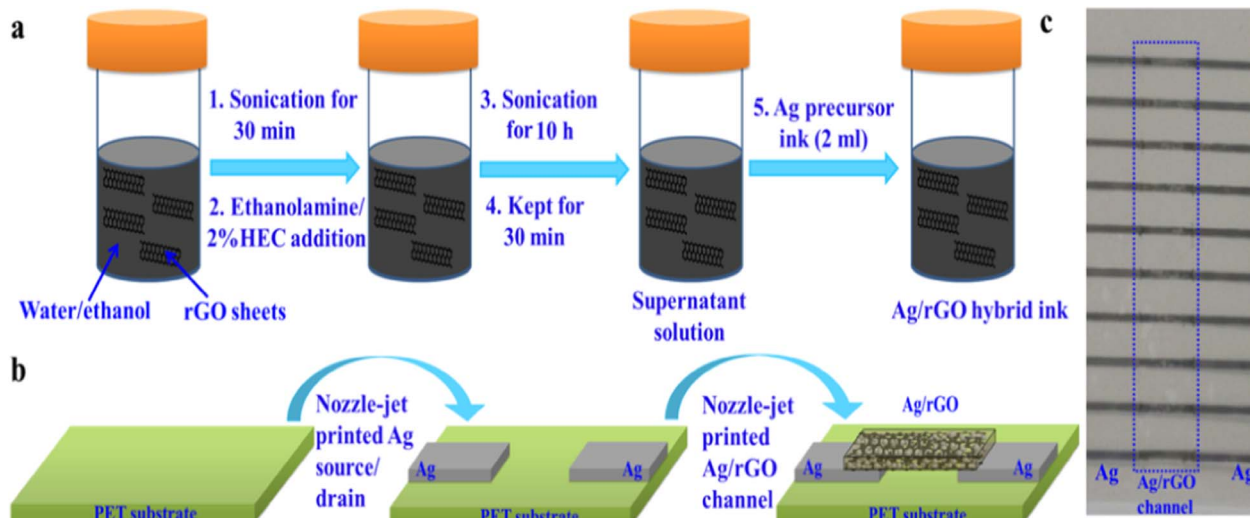


Fig. 14 (a) Schematic of Ag/rGO hybrid ink formulation, (b) nozzle-jet-printed Ag/rGO-based FET sensor, and (c) optical image of printed Ag/rGO-based FET sensors on a PET substrate.¹⁵¹

this, more research is advised to determine the function of various elements in MPs' electrochemical sensing process.

- The electrochemical device and MP identification efficiency are closely related. Small sensing zones typically improve the sensing accuracy; however, when large MPs or aggregates are present, obstructions may happen. It would be interesting to create sensing devices with numerous channels in order to

address this problem. Advanced additive manufacturing methods can also expedite the device design process.

- The majority of sensing materials require time-consuming, complex, and expensive processing in order to identify contaminants originating from MPs. Therefore, creating high-performing electrodes with simple procedures is crucial to develop electrochemical sensors that are commercially viable.

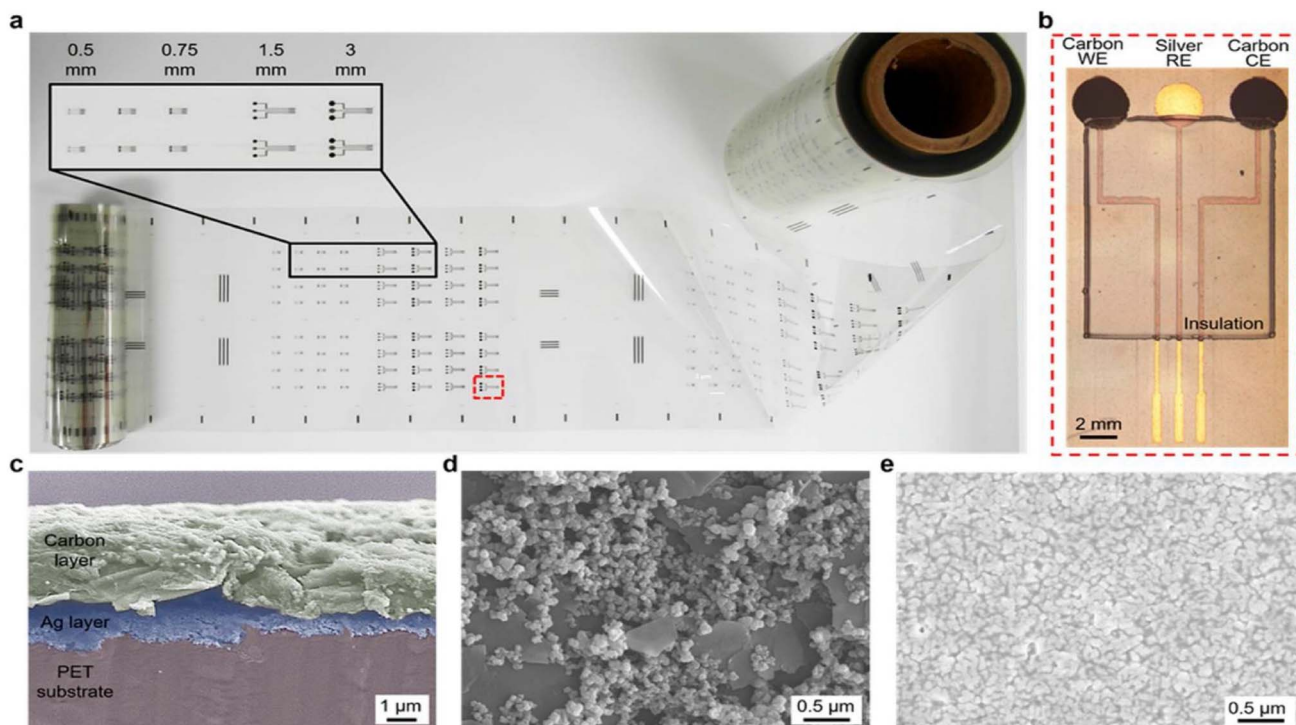


Fig. 15 Example of roll-to-roll gravure-printed electrodes on flexible PET substrates (a) with optical images showing an array of 3 mm-diameter electrodes consisting of carbon (working electrode), silver (reference electrode) and carbon (counter electrodes), and insulation layer (b) with SEM images showing the cross section of hierarchical carbon ink over silver ink (c), carbon electrode surfaces, and silver electrode (d) with nanostructured ink components (e).^{152,154}



9. Future outlooks

Benefits from electrochemical approaches include great sensitivity, cheap operating costs, and ease of use. Numerous disposable, single-use electrodes and self-powered devices are readily accessible and offer great platforms for straightforward, low-cost, and onsite contamination monitoring in decentralized laboratories. Developments in advanced material development, microfabrication, and additive manufacturing techniques, such as 3D printing and inkjet printing, have resulted in improved selectivity, stability, and shelf life of developed sensors, as well as increased detection capabilities. Nevertheless, despite the substantial advancements made, a number of enhancements are still required to transfer this technology from the lab to the real world. First off, the majority of research is still conducted using standard solutions that ignore the sample composition and matrix effect. It is still necessary to ascertain issues like interferences, the existence of naturally occurring organic compounds, and electrode passivation. Second, it must be proven that the necessary selectivity and sensitivity can be attained and that the procedure may be carried out *in situ* with or without a minimal sample pretreatment step. Even though the use of 2D, 3D, and layered materials, as well as advanced manufacturing techniques, has resulted in significant advancements in improving the detection sensitivity and portability, many electrochemical sensors are still in the “proof-of-concept stage,” and their functionality in the field and in real environmental scenarios needs to be established. Last, in order to show competitive advantages, electrochemical sensors must be validated.

Several possible paths might be imagined to enhance the potential of electrochemical approaches for MP field detection:

(1) Rationally designed 2D and 3D nanomaterials combined with biological molecules as sensing elements to increase the sensitivity and selectivity.

(2) Large-scale, scalable manufacturing techniques like 2D and 3D printing to fabricate electrodes and improve reproducibility.

(3) High-throughput multi array platforms for simultaneous MP detection, possibly in conjunction with self-reference electrodes to minimize interferences and improve selectivity.

(4) Integration of electrochemical measurements with artificial intelligence and pattern recognition.

(5) Developments in nanoimpact electrochemistry to enable ultrasensitive detection. Ultimately, a sample unit and six electrochemical sensors might be interfaced to produce a field-portable device that combines detection, recognition, and separation functions into one unit. In conclusion, electrochemical instruments may prove to be inexpensive, user-friendly platforms for tracking these newly discovered pollutants in the environment.

10. Conclusions

MPs are a part of everyday life everywhere on Earth, from the Arctic to the ocean floor. There has been a great deal of research done on the dangers that MPs present to the environment and

to people. This paper describes the present status of electrochemical approaches for MP detection. To show the different domains in which electrochemistry can be used as an alternative technique for MP field detection, examples of applications have been given. The detection of MPs and their concentrations in samples could be sped up with affordable, field-proven electrochemical techniques. These methods make it possible for extensive research to determine MP concentrations and to track and control the dangers associated with these pollutants' existence in the environment. They can also be employed as an adjunct to existing laboratory apparatus. The advantages of electrochemical methods include high sensitivity, low operating costs, and simplicity of usage. There are several self-powered instruments and disposable, one-time electrodes available on the market that provide excellent platforms for efficient, affordable, and onsite contamination detection in decentralized labs. The development of new materials, microfabrication, and additive manufacturing methods—like 3D printing and inkjet printing—has improved the selectivity, stability, and shelf life of the sensors, as well as their capacity for detection. Even with all of these advancements, a few more improvements are needed to take this technology from the lab to the real world. The advent of electrochemical technologies has facilitated the creation of instruments for cleaning, sensors for monitoring MPs, and basic knowledge about MPs. Thus, electrochemical methods are useful instruments that have the potential to revolutionize our comprehension of MPs in the environment and streamline the identification and elimination of MPs.

Author contributions

All authors listed have made a substantial, direct, and intellectual contribution to the work and approved it for publication.

Conflicts of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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