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Endocrine disrupting chemicals in water and recent advances on their detection using electrochemical biosensors

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The ever increasing anthropogenic activities have been producing an undesired group of substances called endocrine disrupting chemicals (EDCs), which can pose a serious threat to the health of human beings and wildlife once they are released into natural water environments. Herein, recent advances on the use of electrochemical biosensors for the determination of EDCs especially in water are thoroughly summarized. Initially, different categories of EDCs with important guidelines are briefly introduced, followed by a simple discussion of non-electrochemical detection methods. Electrochemical techniques including theoretical background and reported studies are then evaluated as the highlight of the review. Finally, current issues, technical bottlenecks, and prospects of this field are critically discussed. This review is composed of following subsections: (i) systematical classification of common EDCs with their toxicities towards human beings and animals, (ii) water contamination events, safety guidelines, and legislations, (iii) non-electrochemical methods for detection of EDCs, (iv) electrochemical monitoring systems, (v) receptors for biorecognition, (vi) detection of EDCs using electrochemical biosensors, and (vii) recent progress, issues, and further development. This review, with a strong interdisciplinary nature, across nanotechnology, biology, material science, and electrochemistry, can offer comprehensive academic assistance for future studies on the detection of EDCs using electrochemical biosensors.

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1. Introduction

In recent years, endocrine disrupting compounds (EDCs) have aroused public concern due to their adverse effects on the endocrine systems of some aquatic and terrestrial animals. Rapid industrialization such as food packaging,



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plastic manufacture, pharmaceuticals, and pesticides are mainly responsible for the increased level of EDCs in the environment and wastewater. Endocrine disrupting compounds (EDCs) often mimic endogenous hormones based on their structural similarity or interfere with normal activities of hormonal receptors by altering their behaviors. Even a trace concentration in EDC-contaminated water can induce acute or chronic health impacts on human beings or animals. As a big family, bisphenols (BPs), polychlorinated biphenyls (PCBs), phthalate esters, alkylphenols, and natural and synthetic estrogens, are the commonly reported categories of EDCs, and each category has many subtypes.¹ Moreover, a growing number of artificial and naturally occurring compounds are considered as suspected EDCs because of equivalent harmful consequences; therefore effective control and management is essential. Several critical

reviews in relation to the classifications and toxicity mechanisms of EDCs have been published during the last few years. From a clinical perspective, Diamanti-Kandarakis *et al.*² elaborated an overview on health risks after EDC exposure aiming to provide readers with an in-depth understanding of endocrine disorders, organ failures, and carcinogenesis.

EDC detection results acquired from various chromatography methods are widely regarded as gold standards by worldwide authorities. Omar *et al.*³ published a review paper concerning the determination of EDCs primarily achieved by liquid chromatography mass spectrometry-mass spectrometry (LC-MS-MS), gas chromatography-mass spectrometry (GC-MS) and other bioanalytical approaches. Nevertheless, these methods always have some drawbacks such as the use of costly equipment, complicated operation,



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and long analyzing time. Furthermore, specimens ready for testing usually need to be sent to assigned laboratories, which are, to some extent, not feasible for large scale and point-of-use applications. Hence, it has become imperative to develop low-cost, simple, and sensitive monitoring platforms with miniaturized devices for rapid determination of EDCs in water and wastewater.

To date, a variety of sensing technologies have been applied for EDC detection. Scognamiglio *et al.*⁴ conducted a review on commonly used monitoring tools for EDCs, including *in vitro* bioassays and chromatographic analysis, with biosensing techniques discussed at greater length. Similarly, Rodriguez-Mozaz *et al.*⁵ reviewed over 100 publications regarding the detection of EDCs using biosensors, where the monitoring of endocrine disorders or signals captured after the receptor–analyte conjugation were described in more detail. Compared with the conventional methods, Jaffrezic-Renault *et al.*⁶ paid more attention to the detection of EDCs by electrochemical techniques. However, advances summarized in their review were mostly reported within the years 2016–2019. As an extended version of this, Azzouz *et al.*⁷ referenced more literature associated with EDC determination *via* electrochemical strategies on nanomaterial-based platforms. Moreover, Sofen *et al.*⁸ provided a review article related to the screening of endocrine disrupting pollutants using electrochemical sensors. Despite the above good perspectives, there is really a need to expand the scale of the evaluation by analyzing more references to support each type of EDC.

This review begins with the discussion of EDCs especially for common compounds of each type, along with their sources and toxicities. Suspected EDCs are also mentioned to help readers have an entire picture. Historical water pollution and regulation issues are stated reflecting the urgency in quality control of water sources. As a methodological comparison, EDCs detected using non-electrochemical methods are firstly discussed. Analyses of electrochemical biosensors include several subsections, starting from electrochemical techniques, commonly used electrodes, surface functional layers, biorecognition elements to electrochemical biosensors for EDCs. Finally, conclusive statements regarding electrochemical biosensors for EDCs, and current challenges as well as future development are also presented.

2. Endocrine disrupting chemicals

EDCs can be roughly divided into naturally occurring products and artificial compounds, where the latter cover the larger proportion ranging from industrial additives to daily-consumed utensils, aliments, and medications. EDC contaminated effluents eventually reach natural water environments *via* different routes, increasing the health risk. Toxicity mechanisms and corresponding adverse effects after exposure are depicted in Table 1. Unlike other water pollutants, EDCs consist of multitudinous types and

subvarieties. Commonly found EDCs include bisphenols, polychlorinated biphenyls, phthalate esters, alkylphenols, and natural and artificial estrogens. Other less studied and suspected EDCs are also included in this review. Common chemical structures regarding each type of common EDC are presented in Fig. 1.

2.1 Bisphenols

Bisphenols (BPs) are the most prevalent EDCs that are always composed of two phenolic bodies linked together with a bridging component. The central connection is an essential part to differentiate from other variants. To date, BPs have approximately 20 variants. Bisphenol A (2,2-bis(4-hydroxyphenyl) propane, BPA) is the most notorious BP over other analogs and even in the entire EDC family. BPA can bind to the pregnane X receptor and different estrogen receptors (ERs) such as ER α , ER β , and estrogen related receptor γ (ERR γ), resulting in a variety of endocrine disrupting effects like immune system damage, neurological toxicities, metabolic disorders, and reproductive dysfunctions in the human body.⁹ Furthermore, BPA also exhibits toxic behavior towards wild animals particularly for the aquatic life when the concentration rises to 1–10 $\mu\text{g mL}^{-1}$.¹⁰ Due to the serious consequences, other bisphenol analogs have replaced the use of BPA. Bisphenol S (4,4-sulfonyldiphenol, BPS) is one of the ideal BPA substitutions and is extensively used in many applications. Unfortunately, recent studies reveal that BPS can induce heart problems by influencing ER β (ref. 11) and has strong toxicity to antiandrogenic activities.¹² As another alternative of BPA, bisphenol AP (4,4'-(1-phenylethylidene) bisphenol, BPAP) is also evidenced as a toxic chemical. Xiao *et al.*¹³ demonstrated that exposure to BPAP in trace amounts might lead to anti-estrogenic effects and blood glucose level disorders. Bisphenol AF (2,2-bis(4-hydroxyphenyl) hexafluoropropane) belongs to a halogenated BPA, where hydrogen atoms on the two methyl radicals are fully replaced by fluorine atoms. Unlike BPA, BPAF can activate ER α while presenting antagonistic behavior towards ER β , without the need for ERR γ . In terms of toxicity, several research groups have confirmed that BPAF can negatively impact heart and liver tissues in zebrafish.^{14,15} BPA and BPS are commonly used industrial chemicals in the manufacture of epoxy resins and polycarbonate products reaching their ubiquity in many hard plastic containers.^{8,11} BPAP and BPAF are essential components for polymer processing and can be widely found in rubber products, thermals papers, cosmetics, and electronic parts.^{11,14} There are some special BPs, where the bridging part remains the same as BPA, but has halogenated replacement (*i.e.*, chlorinated, or brominated reaction) on both phenolic rings, including tetrabromobisphenol A (TBBPA), tetrabromobisphenol S (TBBPS), and tetrachlorobisphenol A (TCBPA). Bioassays showed that these halogenated BPA derivatives are capable of interfering not only with ERs,¹⁶ but also glucocorticoid receptors (GRs), androgen receptors (ARs),¹⁷ peroxisome



Table 1 Common EDCs with their toxicities and adverse effects

EDCs	Toxicology	Adverse health effects
Bisphenols	Acting on estrogen, androgen, and thyroid receptors	Cancers, diabetes, heart diseases, and reproductive dysfunction
Polychlorinated biphenyls	Interfering with cerebral glands, thyroid receptors, and reproductive systems	Carcinogenic, mutagenic, teratogenic effects and immunotoxicity
Phthalate esters	Altering cerebral, peripheral, and reproductive hormone secretion balance	Diabetes, hepatotoxicity, precociousness, low birth weight, abortion, and respiratory issues
Alkylphenols	Abnormally affecting ERK signaling pathways	Tumorigenicity, immune and metabolic system disturbance
Natural and synthetic estrogens	Mimicking estrogenic hormones due to high structural similarity	Irregular reproductive activities, carcinogenicity, cardiovascular risks, headache, and other physiological pains

proliferator-activated receptor γ (PPAR γ), and thyroid receptors (TRs).^{18,19} However, adverse health effects on human beings need to be further investigated. Very recent studies on the carcinogenic nature of TCBPA and genotoxicity of TBBPA and TBBPS were elaborated by Lei *et al.*²⁰ and Barańska *et al.*,²¹ respectively. TBBPA, TBBPS and TCBPA are known as common brominated flame retardants, and they

are also often used in plastic paints, synthetic textiles, and electrical devices.^{22,23}

2.2 Polychlorinated biphenyls

Polychlorinated biphenyls (PCBs), another class of man-made EDCs, have 209 congeners, and all of them follow a general formula $C_{12}H_{10-x}Cl_x$. Every single PCB is identified by a specific BZ congener number according to the position and number of chlorine atoms on a biphenyl body and is usually named PCB x . Among them, tetrachlorobiphenyls such as 2,3',5,5'-tetrachlorobiphenyl (PCB72) and 3,3',4,4'-tetrachlorobiphenyl (PCB77) are frequently reported. Dioxin-like PCBs can bind to aryl hydrocarbon receptor (AhR) and are considered as the most hazardous and anti-estrogen chemicals, consisting of PCB77 and other 11 congeners.²⁴ Moreover, non-dioxin-like PCBs like PCB28, PCB101, and PCB180 are also frequently reported. Non-dioxin-like PCBs prefer to interfere with ERs and ARs, leading to various adverse effects.²⁵ Since many congeners have similar structures, PCBs mostly exist in a mixture. Aroclor, one of the commercially available products, contains over 100 variants of PCBs. Aroclor 1254, a $C_{12}H_5Cl_5$ -dominant PCB mixture, consists of four different congeners which are identified by the four to seven substitutions of chlorine atoms on the biphenyl.²⁶ PCBs are widely used in the manufacture of electrical parts, or industrial reagents such as lubricants, thermal transfer fluids, and coolants.²⁷ The persistence in the natural environment makes PCBs (especially heavier congeners) prone to accumulation in organisms through the food chain. Their strong carcinogenic nature has been verified by the International Agency for Research on Cancer (IARC).²⁸

2.3 Phthalate esters

Phthalate esters are also known as phthalates or phthalic acid esters (PAEs). They are often used as plasticizers to reduce the brittleness of raw polyvinyl chloride (PVC) products, but also can be found in many daily necessities and baby toys.²⁹ Phthalate esters usually have 26 derivatives, with molecular weights (MW) varying from 194 to 531 g mol⁻¹. Common phthalate esters incorporate dimethyl phthalate (DMP), dibutyl phthalate (DBP), and di(2-ethylhexyl)phthalate (DEHP). DEHP can be also named as

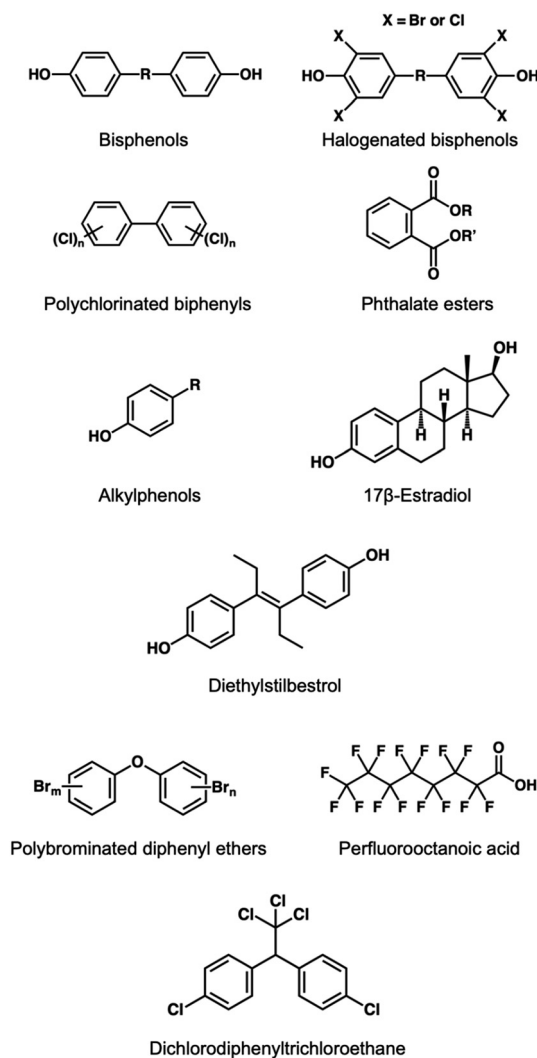


Fig. 1 General chemical structures regarding each type of common EDC.



diocetyl phthalate (DOP), which is different from di(*n*-octyl) phthalate (DNOP), although they have equivalent MW. Phthalate esters with short carbon chains (*e.g.*, DMP) are more volatile than the longer ones, thus increasing the risk of respiratory symptoms such as asthma.³⁰ DEHP is a typical long-chain phthalate ester that tends to exist in water or food rather than air due to the heavier MW. Careless exposure to phthalate esters results in imbalance on hormonal levels and therefore has toxic effects on the brain, liver, and reproductive organs, which is highly detrimental to pregnant women, infants, and adolescents.³⁰

2.4 Alkylphenols

Alkylphenols, a class of organic chemicals obtained from the alkylation of phenols, can also induce endocrine disruption in the event of exposure. Alkylphenols contain 9 subtypes that are defined according to the number of carbon atoms on the alkyl chain. Like the isomerism in phthalate esters, each of alkylphenol subtype has its own isomers. Alkylphenols have higher stability and can remain in natural environments for a long period. Octylphenols (OPs) and nonylphenols (NPs) are frequently found in water sources. Priac *et al.*³¹ not only described these two subtypes, but also emphasized their polyethoxylated precursors namely alkylphenol ethoxylates (APEOs). APEOs are not very stable and are easily subjected to a metabolic process to produce alkylphenols.³⁰ The toxicity of alkylphenols is derived from the disruption of signaling pathways associated with extracellular signal-regulated kinases (ERKs), and thus abnormally interferes with membrane estrogen receptors (mERs). ERK signaling pathways are closely related to many cellular activities such as division, differentiation, apoptosis, and carcinogenesis.³² Long chain alkylphenols are often applied as non-ionic surfactants, which are the main ingredient in detergents or cleaners. Moreover, they are also widely used in the production of oily chemicals, rubber industries, paper manufacture, or used as an alternative of flame-retardant materials.³¹

2.5 Natural and synthetic estrogens

Organisms can naturally produce a variety of estrogens in their bodies, including estrone (E1), (17 β -) estradiol (E2), estriol (E3), and estetrol (E4). These estrogens can be recognized by nuclear estrogen receptors (nERs) and mERs.^{31,33} Estrogens released at a reasonable level can maintain the normal working of reproductive and cardiovascular systems and develop cerebral and osseous tissues.³¹ The unusual release of these sex hormones can also cause endocrine disrupting consequences. Diethylstilbestrol (DES) belongs to synthetic estrogen, of which usage can be traced back to 1940. DES is initially treated as a medical reagent for pregnant women to prevent miscarriage or to solve other pregnancy-related problems. Due to the highly similar structure, DES can pretend to be an endogenic

estrogen interacting with receptors in the body, leading to endocrine disruption.

2.6 Less studied and suspected EDCs

Polybrominated diphenyl ethers (PBDEs) are another class of artificial organic halides. Like PCBs, PBDEs are also a big family that have 209 congeners. They are identified depending on the number of bromine atoms in the molecule. Common PBDEs include tetra-, penta-, octa-subtypes, and decaBDEs. As another widespread flame-resistant material, PBDEs are often added in the manufacture of polymeric foams (e.g., polyurethane foams), plastics, electronics, and textiles.³⁴ PBDEs are proven EDCs that have negative effects on estrogenic and thyroid receptors, resulting in a chaotic level of corresponding hormones.³⁵ Perfluorooctanoic acid (PFOA) belongs to a side product in the manufacture of fluoropolymer-related products, and widely used as an industrial surfactant. PFOA exists in soil, sediment, and is more likely to be in water due to the carboxylic group. It is regarded as an EDC that shows the strongest resistance against the degradation of the natural environment.³⁶ Some health reports indicate that PFOA is closely related to thyroid diseases and carcinogenesis.³⁷ However, more theoretical data is still required to interpret its in-depth toxicity mechanism. Dichlorodiphenyltrichloroethane (DDT) is one of the most famous organochlorides that has been extensively used as a pesticide and insecticide. It exhibits strong hydrophobicity and therefore prefers to seep into soil, sediment, and accumulate in lipids. Like PFOA, DDT is supposed to have carcinogenicity but needs to be confirmed by more studies.³⁸

Organic compounds with similar chemical structures or hormone-disordering effects to EDCs but have not been proven yet are classified as suspected EDCs. Many problems, such as bioaccumulation in organisms, toxicities, environmental contamination records, and official guidelines, still lack conclusive answers to date. Suspected EDCs include polychlorinated dibenzo derivatives, polycyclic aromatic hydrocarbons (PAHs), mycotoxins, pharmaceuticals, personal care products, and broad-ranging agricultural chemicals. Polychlorinated dibenzo derivatives consist of polychlorinated dibenzo dioxins (PCDDs) and polychlorinated dibenzo furans (PCDFs), which are unwanted impurities in the production of chlorinated products (*e.g.*, PCBs).³⁹ PAHs refer to the hydrocarbons with two and more benzene rings and are normally produced from the incomplete combustion in cooking (*e.g.*, use of coal and oil), smoking (*e.g.*, tobacco), and driving (*e.g.*, petrol).⁴⁰ Mycotoxins are a group of xenoestrogens produced by fungi and are sometimes also called mycoestrogens. Zearalenone (ZEN), a typical type of mycotoxin, is generated by the estrogenic metabolism of *Fusarium* and *Gibberella* species in many cereal crops.⁴¹ 17 α -Ethinylestradiol (EE2) is an artificial compound with a similar chemical structure to E2, which is used as a contraceptive medication coupled with progestins. EE2 can

induce various side effects on human beings. It has also been demonstrated that EE2 presents serious neurotoxicity towards fishes (e.g., zebrafish larvae) when it is released into natural aquatic areas.⁴² Oxybenzone is created naturally in many flowering plants and is often applied as the main constituent of sunscreen creams (i.e., one of the common personal care products).⁴³ However, skin allergies and other adverse effects caused by oxybenzone have been increasingly reported in recent years,^{44,45} making this sunscreen active ingredient more controversial. Moreover, suspected EDCs also incorporate many agricultural chemicals such as pesticides (e.g., endosulfan and kepone), herbicides (e.g., atrazine), and fungicides (e.g., vinclozolin). The endocrine disrupting effect regarding these chemicals requires more studies.

2.7 Contamination events, official warnings, and safety guidelines

The strong reliance on polycarbonate plastics and epoxy products worldwide leads to the extremely high use of BPA. China accounted for one fifth of the world's total consumption of BPA in 2013. Lin *et al.*⁴⁶ selected Yuyao, Taizhou, and Wenzhou City in Zhejiang Province, the main areas for plastic production and sale in China, to explore the influence of plastic factories on surface soils, nearby rivers and aquatic organisms. The study revealed that the level of BPA from the river in Yuyao City was 0.24–5.68 ng mL⁻¹, which was higher than the maximum quantified BPA concentrations in Asia of 317 pg mL⁻¹, North America, and Europe of 99 and 14 pg mL⁻¹, respectively.⁴⁷ On the other hand, the wide existence of BPA in baby products, such as milk bottles and infant formula cans, was demonstrated by the United States Food and Drug Administration (USFDA).⁴⁸ Many organizations or countries have implemented legislations on prohibition of BPA. Canada was the first country to ban BPA in the production of baby bottles in 2010, followed by the European Union (EU), China, and Malaysia issued similar regulations to increase the stringency upon BPA.⁴⁹ As an alternative of BPA, BPS and BPF have been widely used in many countries to make 'BPA-free' products. Yamazaki *et al.*⁵⁰ investigated the level of BPS in water sources from China, Japan, South Korea, and India, of which the highest concentration was found in the Adyar River of 7.2 ng mL⁻¹, Chennai, India. In addition, the amount of BPF in water samples was also provided by their group, indicating higher concentrations in China, Japan, and South Korea, especially the Tamagawa River of 2.85 ng mL⁻¹, Japan, to which great attention should be paid.⁴⁹ Liu *et al.*⁵¹ noted that the water specimens from the Taihu Lake in Jiangsu Province, China, not only had common BPs (i.e., BPA, BPF, and BPS), but also had BPB, BPZ, BPAF, and BPAP with comparable contents, among which an apparently high concentration of BPAF was determined in a range of 0.11–0.14 ng mL⁻¹, with a median of 0.11 ng mL⁻¹. Liu *et al.*⁵² confirmed the detection results of surface water samples

given by Yamazaki *et al.* and Liu *et al.*, and comprehensively evaluated the BPs in sewage samples from wastewater treatment plants (WWTPs), showing that BPA, BPS, and BPF were the main pollutants in China and the United States, while BPF in India was below the detection limit. Interestingly, BPFA was only detected in samples collected from China.⁵¹

Earlier electrical devices contain high levels of PCBs, where some of them are still being used at present. Natural environments will suffer from great pressure on decomposition if PCB-containing electrical devices are improperly recycled or abandoned after they reach their lifetimes. PCBs not only exist in water environment, air, and soil, but also are highly prone to be accumulated in fatty tissues due to the strong lipophilic property.⁵³ Bench *et al.* found that PCBs had been accumulated up to 10 million times in rainbow trout in Lake Ontario, which resulted in the closure of local fisheries.⁵³ Additionally, the impact of PCBs on seawater was proposed in the report as the pollution initially stored in phytoplankton could become more serious in advanced organisms through the food chain, and ultimately threaten human health.⁵³ Owing to increasing worldwide pollution concern, many countries have started taking actions. On PCBs, Canada passed a law in 1977 to ban the import, manufacture, and sale of PCBs. After that, the US, UK, Australia, and EU successively banned the use of PCBs in 1979, 1981, 1986, and 1987, respectively. To strengthen the quality management of drinking water, the United States Environmental Protection Agency (USEPA) set the highest acceptable concentration for PCBs of 500 pg mL⁻¹.⁵⁴ In 2001, governments around the world signed the Stockholm Convention aiming to eliminate PCBs by 2028. As one of the signatories to the convention, China was committed to proper inventory and harmless disposal of PCBs.⁵⁵

Like BPs, phthalate esters as plasticizers are also detected in a range of plastic products. The weak behavioral and cognitive abilities of young children can be derived from more exposure to phthalates than adults, and therefore it is significant to control the phthalate content in baby products. The EU has banned six phthalates, including butyl benzyl phthalate (BBP), di-isodecyl phthalate (DIDP), di-isononyl phthalate (DINP), DBP, DEHP, and DNOP.⁵⁶ An investigation into the health risk of phthalates was conducted by Abtahi *et al.*,⁵⁶ showing that DMP and DEHP were the primary contaminants found in different water samples from Tehran, Iran, along with other phthalates detected in relatively low levels, involving BBP, DBP, diethyl phthalate (DEP), and DNOP. The estimation on the burden of disease after drinking water contaminated by DEHP was also performed by the group based on the theory of disability-adjusted life year (DALY).⁵⁶ Paying equivalent emphasis on the health of children, the Minnesota Department of Health (MDH) listed BBP, DBP, and DEHP as Toxic Free Kids Act Priority Chemicals in 2011, with corresponding safety guidance of drinking water for BBP (100 000 pg mL⁻¹) and DBP (20 000 pg



mL⁻¹) issued in 2012 as well as DEHP (7000 pg mL⁻¹) in 2013.⁵⁷ The guideline value of DEHP was 1000 pg mL⁻¹ higher than the maximum contaminant level (MCL) provided by the FDA (6000 pg mL⁻¹) but was 1000 pg mL⁻¹ lower than the concentration set by the World Health Organization (WHO) and EU (8000 pg mL⁻¹).⁵⁸ The pollution of BBP, DBP, DEP, DEHP, DMP, and DNOP from other countries (*i.e.*, Canada, Spain, South Africa, China, and India) was reviewed by Ghosh *et al.*,⁵⁹ which showed that the content of nominated phthalates in water samples was not higher than 7 ng mL⁻¹.

A total amount of 500 000 tons of APEOs are annually produced worldwide to meet the demand for non-ionic surfactants and other applications.⁶⁰ Alkylphenols are the product of APEOs after biodegradation. Municipal wastewaters and industrial effluents (*e.g.*, factories for paper and textile production) are usually the main sources of environmental contamination.⁶² Uğuz *et al.*⁶² published a comprehensive review on the pollution of alkylphenols and their health threat to living organisms. In their work, they found that nonylphenol ethoxylates (NPEs) and NPs widely existed in water and sediment, and pointed out that the concentration of NPs in rivers from the United States, United Kingdom, Canada was much higher (*i.e.*, ranging from µg L⁻¹ to mg L⁻¹) than the level for inland surface waters of 300 pg mL⁻¹ from the Directive 2008/105/EC.⁶³ Although the current concentration of NPs in various natural waters is not high enough to cause death, the bioaccumulation in living creatures can still reach a lethal level as time passes, which should be paid more attention.⁶² Moreover, the maximum permissible standard of 7000 pg mL⁻¹ for 4-NP in ocean water was confirmed by the USPEA to ensure the normal growth of most marine animals.⁶⁴ Lei *et al.*⁶⁵ investigated several estrogens (*i.e.*, E1, E2, E3, EE2, and DES) in water samples from the Beijing–Tianjin–Hebei region, of which the total concentration in wastewater was 0.468 ± 0.0272 ng mL⁻¹ and was 0.219 ± 0.023 ng mL⁻¹ in river water. Natural or synthetic estrogens have negative effects on surface waters, particularly for fish species.⁶⁴ Similarly, Wedekind *et al.*⁶⁶ focused the analysis on the change of the sex ratio of fish caused by the presence of estrogens and endeavored to find the reason for the strong adaptation in estrogen-contaminated water in some fish populations. However, more efforts on this field were suggested to be taken in these two papers due to the current less-supporting evidence. As a major pollutant of estrogen, E2 was listed in the Water Framework Directive (WFD) of the EU in 2013, along with a safety guideline of 0.4 pg mL⁻¹.⁶⁷ It has been confirmed that PBDEs can accumulate in milk products (*e.g.*, human breast milk⁶⁸) and fatty tissues. They also mainly exist in surface water, soil, and sediment. Surface waters containing PBDEs, of which the concentrations range from a few pg L⁻¹ to hundreds of ng L⁻¹, have been registered in countries from all over the world, including Switzerland (river and lake water), Canada (lake water), the United States (lake water and estuary), Paris (river water), and the UK (river water).³⁴ PBDEs

were banned in the US and EU in 2004.^{69,70} Until recently, there is no specific maximum acceptable level for PBDEs in drinking water. PFOA is widely found in surface waters,⁷¹ coastal waters,⁷² and seawaters.⁷³ The potential adverse effects of PFOA on human beings and natural environments, especially for infants and aquatic lives, respectively, have been of significant concern, although relevant toxic events are unknown.³⁶ Aiming to reduce the extensive exposure, PFOA was discontinuously produced and gradually phased out in the United States in 2015. Canada set the maximum acceptable concentration (MAC) for PFOA in drinking water of 200 pg mL⁻¹.⁷⁴ The health advisory for PFOA in drinking water in China was firstly claimed by Zhang *et al.*,⁷⁵ as 85 pg mL⁻¹. Likewise, DDT has been banned in many countries due to its strong toxicity. However, it is still being used in some developing countries or regions to fight against malaria. Ochoa-Rivero *et al.*⁷⁶ reported that DDT with a total amount of 2804 ng mL⁻¹ was found in agricultural areas in Mexico. The value was extremely higher than those detected in water sources from other areas in Mexico or other countries where DDT was used.⁷⁶ With the consideration upon the continuous consumption of DDT, the WHO issued a guideline value of 1000 pg mL⁻¹ in drinking water.⁷⁷

2.8 Non-electrochemical methods for detection of EDCs

Traditional determination of EDCs was normally performed using bioanalytical methods such as cell assays (*i.e.*, recombinant yeast incubation, cell proliferation, and *in vitro* cell bioluminescence tests) and enzyme-based assays like luciferase induction.⁷⁸ However, these bioassays are always complicated in terms of manipulation and require lengthy analyzing duration. Recently, more advanced analytical techniques, including enzyme-linked immunosorbent assay (ELISA), surface enhanced Raman spectroscopy (SERS), and biofluorescence, have been developed to improve test efficiency and reduce batch-to-batch variation. ELISA is a bioassay where the antibody–antigen interaction is usually achieved on 96-well microplates. Biorecognition results are obtained after the introduction of signal-indicated substrate solutions. ELISA is easy to follow, and the analysis can be conducted in different specimens. Commercial ELISA kits have been available for the quantitative determination of EDCs. This method retains its predominance and is sometimes considered as a golden standard for comparing the results obtained from other approaches, although the method often suffers from a main drawback of short data-collection time.⁷⁹ EDCs measured using SERS are reported in some publications. For example, Tu *et al.*⁵⁸ developed aptamer-functionalized magnetic particles to provide binding sites for the competitive DEHP that were attached to the surface of SERS silica particles and the target DEHP. The magnetic beads were then aggregated by a magnet to yield the supernatant of silica particles containing Raman receptor molecules. Raman spectra with characteristic peaks of DEHP in different intensities were clearly observed after different



concentrations of DEHP were introduced.⁵⁸ However, this analysis can be only achieved in laboratories using a bulky analytical instrument and is not suitable for on-site tests.

Another commonly reported method for the detection of EDCs is biofluorescence. It combines biorecognition elements with fluorescent substances such as dyes, labels, or quantum dots, indicating the analyte concentration according to the change in fluorescence intensity. A non-equilibrium rapid replacement aptamer (NERRA) assay was conducted by Kim *et al.*⁸⁰ based on a portable device, where the phthalate esters were detected in the solution containing free PoPo3 dye attached aptamers. Results of biofluorescence were highly comparable to those of chromatographic methods, but the analyzing time was dramatically reduced to 30 seconds. The experiment focused only on detecting the mixture of phthalate esters rather than a single chemical.⁸⁰ Aptamer-based biosensors with fluorescence characteristics account for a large percentage of the biofluorescence technology. Lim *et al.*⁸¹ reported an aptasensor connected with two types of quantum dots (QDs) to detect DEHP. QDs 565 were immobilized on magnetic beads while QDs 655 dissociated from the DNA probes after DEHP binding to the aptamer. Apart from phthalate esters, fluorescent aptasensors are also widely used to detect other EDCs like BPs⁸² and PCBs.^{83,84} Instead of fluorescent aptamers, Tang *et al.*⁸⁵ fabricated an antibody-based biosensor based on optical fibers for the monitoring of eight phthalate esters. The antibodies coupled to fluorescent tags reacted with the analyte to create inhibition signals. In comparison to Kim *et al.*,⁸⁰ their immunosensors featured the sensitive detection of a single phthalate ester (*i.e.*, DMP), demonstrating the reliability of the protocol.⁸⁴

In recent years, chromatographic or chromatographic-spectrometric combined analyses used for the detection of EDCs, such as gas chromatography coupled to mass spectroscopy (GC-MS),⁸⁶ high-performance liquid chromatography (HPLC),⁸⁷ and liquid chromatography coupled to mass spectrometry (LC-MS),⁸⁸ have been acknowledged by more and more environmental agencies or health organizations due to their excellent sensitivity and accuracy. These methodologies enable the identification of a series of EDC variants at a single measurement, and are therefore of great significance in multi-analyte monitoring. Nevertheless, limitations regarding these techniques need to be taken into consideration at the same time, for instance, expensive apparatus, scheduled maintenance, and demand for experienced technicians. Moreover, pre-concentration, extraction (micro-extraction), and derivatization of specimens are other inconveniences when using these chromatographic methods, which often make the experiment unsuitable for *in situ* applications.⁸⁹ In fact, the complexity of EDCs worsens the limitations on the use of bioassays, chromatographical analyses, and other non-electrochemical approaches, pushing the advent of a highly efficient and miniaturized setup working under a simple detection mechanism but capable of providing competitive results. The developed prototype tends

to present more friendliness toward real samples in a point-of-care application. Compared with the conventional detection methods, biosensors in combination with electrochemical techniques have been increasingly used as a delicate sensing systems to determine EDCs in water sources relying on the higher selective and sensitive nature. Electrochemical biosensors equipped with different biorecognition components will be elucidated at the latter section of the review.

3. Electrochemical biosensing of EDCs in water

Biosensing is a process that achieves the detection of the target analyte using a device where the surface is covered with biorecognition components. Electrochemical biosensing allows biochemical activities to occur at an electrode-solution interface, followed by a signal transduction to yield readable electrical responses that are presented as original curves *via* a data processing system (Fig. 2). This technique integrates superior sensitivity with advantages of shorter analyzing duration, inexpensiveness, and miniaturization feasibility, which has been known as a promising method for the determination of small molecules. Commonly used electrochemical biosensors can be divided into immunosensors, aptasensors, and enzymatic sensors. Each class of biosensors has its own unique principle for detection. Immunosensors are a class of bioelectrodes where the surface is functionalized with antibodies, which can selectively recognize antigens *via* the antibody-antigen interaction. The affinity offered by aptasensors is derived from the chain-like structure that has strong locking effect upon the analyte. Electrodes decorated with a bioactive

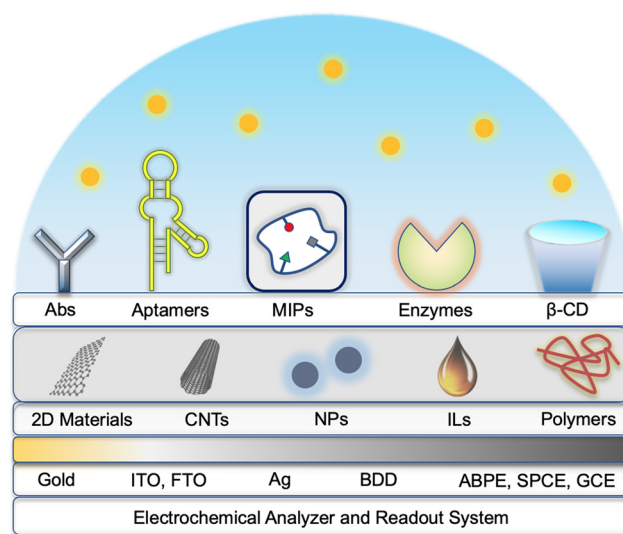


Fig. 2 Schematic diagram of electrochemical biosensors used for detection of EDCs in water (among them, images of 2D materials, CNTs, and ILs were reprinted with the permission from ref. 193–195, respectively).



enzyme exhibit electrocatalysis towards the target molecules. Some natural (*i.e.*, cyclodextrins) or artificial (*i.e.*, molecularly imprinted polymers) recognition elements have attracted tremendous interest in the past decade because of their biomimetic properties. The surface of molecularly imprinted polymer (MIP)-based sensors can provide cavities with an identical shape of the analyte, therefore showing enhanced specificity in the sensing process, while cyclodextrin-related sensors capture the target *via* their adsorptive chambers. The electrochemical biosensing system consisting of various electrochemical techniques, electrodes and surface functional supports will be firstly introduced in this section, followed by different bioreceptors, especially the binding schemes of antibodies in immunosensors will be discussed in more detail.

3.1 Electrochemical monitoring system

An electrochemical monitoring system is normally constructed by connecting a biofunctionalized electrode with an electrochemical analyzer. The analyte-containing solution is then allowed to be in contact with the specific sensing area of the transducer, triggering the conversion from biological events to electrical responses as amperometric and impedimetric forms (Fig. 3). All these measurements take

place after application of a potential on the working electrode to produce the corresponding current. Voltammetry is a method to analyze current changes over an assigned potential scanning under linear (*e.g.*, linear sweep voltammetry), periodical (*e.g.*, cyclic voltammetry), and pulsed (*e.g.*, differential pulse voltammetry and square wave voltammetry) behaviors. Linear sweep voltammetry (LSV) is the most fundamental electrochemical technique with a simple working principle in which the potential linearly changes with time.⁹⁰ An advanced version of the technique called second-order derivative linear sweep voltammetry (SDLSV) produces electrochemical results often with lower background noise and larger peak current due to the more complicated mathematical analysis.⁹¹ Cyclic voltammetry (CV) is regarded as a derivative of LSV. The linear potential in CV scans starting from a higher level to the switching point (*i.e.*, lower level), followed by a reversing of the potential from the switching point back to the initial level.⁹⁰ The workflow can be performed in multi-cycles under a fixed scan rate. In most cases, the resulting current varies as the potential scans forming a closed voltammogram that contains an oxidation and a reduction peak. CV is one of the most common electrochemical methodologies utilized not only in the sensing process, but also for the assessment of electrode modification from untreated up to a fully functionalized

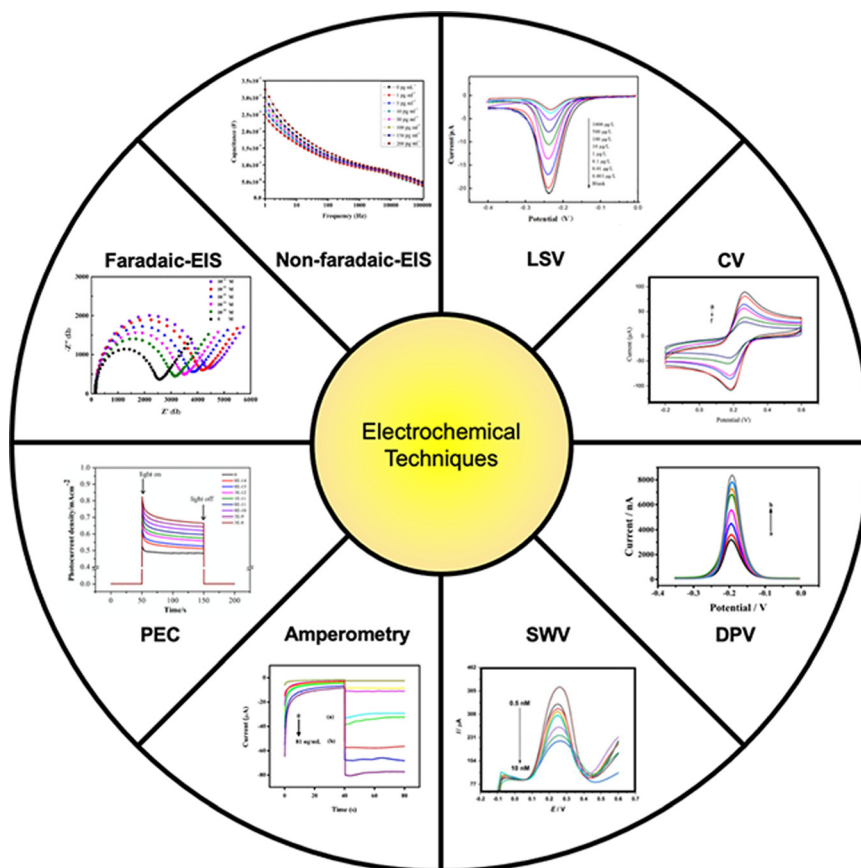


Fig. 3 Electrochemical techniques used for the detection of EDCs (reprinted with the permission from ref. 141 for LSV, ref. 164 for CV, ref. 153 for DPV, ref. 137 for SWV, ref. 129 for amperometry, ref. 161 for PEC, ref. 150 for faradaic-EIS, and ref. 136 for non-faradaic-EIS).



situation.⁹² Differential pulsed voltammetry (DPV) is another commonly reported electrochemical technique. Unlike LSV and CV, DPV embeds voltage pulses on a linearly changed potential, creating two momentary currents right before and after the pulsed voltage changes. The final current difference instead of an absolute value *versus* an assigned range of potential contributes to a characteristic DPV voltammogram.^{93,94} Like DPV, voltage pulses in square wave voltammetry (SWV) are superimposed with a staircase-shaped potential, and the current difference is measured based on the two current values recorded right before and after the pulsed voltage changes.⁹⁵ Amperometry is another frequently used electrochemical technique. It is different from voltammetry as a constant potential is usually applied on the working electrode to generate a transient current in relation to the oxidation or reduction reaction of redox couples. Amperometric measurement can diminish the undesired noises for the interferants or impurities in a tested solution by adjusting the impressed voltage, therefore possessing high sensitivity towards the detection of interest.^{96,97} Photoelectrochemistry (PEC) is an emerging technique that has been used for biosensing. Following the three-electrode set-up, a PEC system contains a working electrode modified with photosensitive semiconductor materials, coupled with a reference and counter electrode. In this method, a lamp with a medium wavelength is always required as an irradiation source. After a specific voltage is applied to the working electrode, a photocurrent is generated and varies with time. The change of photocurrent density within a light-on duration corresponds to the concentration of the analyte. PEC can produce responsive curves with a distinctive difference even in ultralow concentrations, which has been demonstrated in many studies.^{162,163,166,167,193}

Impedimetric measurement applies a certain potential to the working electrode to generate a current and calculates the resulting resistance or impedance as a final readout. Resistance can be expressed as the slope of the *I*-*V* curve according to Ohm's law. The equivalent circuit merely contains a resistor that serves as the only element subjected to a fixed potential. However, the sample environment in actual electrochemical biosensing usually includes capacitive behaviors and Warburg diffusion (Z_w), thus it is much more complicated than the model of simple resistance.⁹⁸ Electrochemical impedance spectroscopy (EIS) is a typical representative of impedimetric methods that has received a great amount of attention in a recent couple of years due to simple operation, short response time, high sensitivity, and device portability.⁹⁹ In theory, an alternating potential (*e.g.*, sinusoidal potential) is introduced into the system producing a current in the same waveform but with a phase difference.¹⁰⁰ The potential is then divided by the current to yield the impedance under each certain frequency.⁹⁸ Impedance is a characteristic EIS parameter presented in a complex expression and can be drawn in the Nyquist plot, where the real part normally reflects the charge transfer resistance against redox couples moving from the bulky

solution toward the surface of the electrode, and the imaginary part refers to the total capacitance generated at the electrode-solution interface.^{98,101,102} In other words, resistance is a special case of an impedance that lacks the imaginary part. Therefore, analyzing impedance or its parts is vital for the illustration of chemical or biological events (*e.g.*, target molecules binding to the biorecognition element) occurring at the specific sensing area.

3.1.1 Electrodes for electrochemical analysis. Electrodes are the fundamental of signal transduction in electrochemical biosensors. The three-electrode system is widely employed for electrochemical detection, consisting of a working electrode, reference electrode, and counter electrode. The working electrode refers to a sensing platform with functional supports and bioreceptors, which is ready for capturing target molecules. Unlike the two-electrode system (*i.e.*, only consisting of a working electrode and counter electrode), the reference electrode in the three-electrode system has a constant potential and negligible current, that can define the working potential after a certain voltage applied, sharing a part of the work of the counter electrode.¹⁰³ Ag/AgCl is the commonly used material for reference electrodes. The counter electrode, or sometimes called auxiliary electrode, forms a closed path for current with the working electrode. Counter electrodes are often made of platinum. Electrodes fitted in electrochemical biosensors have the following characteristics: 1) good conductivity, 2) high durability, 3) large surface area, and 4) long-term stability. Conductivity is very crucial as it improves the sensitivity and accuracy of signal transduction. Durability increases the electrode tolerance to any damage (*e.g.*, bending, scraping, and dropping). The surface area relates to the amount of bioreceptor loading that determines the specificity level. Stability ensures the electrode to be available under various conditions (*i.e.*, temperature, pressure, and pH), which is beneficial to storage or transportation. Glassy carbon electrodes (GCEs) are the most used electrodes in electrochemical biosensors due to their low cost, good conductivity, and stability. Solutions containing biorecognition elements, or antigens, or functional compounds, are often drop-casted on the surface of polished GCEs. It is ready to use after a certain length of incubation.¹⁰⁴ Carbon rod electrodes have a large surface area because of the special morphology, allowing the functionalization of bioreceptors with a high density.¹⁰⁵ Other highly conductive electrodes like carbon paste electrodes (CPE),^{160,180} carbon electrodes (CE),¹⁷¹ and silver electrodes¹³⁷ are also reported, but with fewer cases. Electrodes made of boron-doped diamond (BDD) possess advantages of high chemical stability and low background noise.^{151,164,179} Acetylene black paste electrodes (ABPE) mixed with paraffin are used as a sensing platform that integrates excellent electrical conductivity with high durability.¹⁵⁷ A unique carbon electrode can be directly created by laser-induced carbonization on a polyimide (PI) substrate.¹⁰⁶ The method for the electrode production does not require



additional carbon materials and the surface patterns of the electrode can be drawn as the design of interest. Gold electrodes are another class of common electrodes with outstanding conductivity and stability due to the inherent properties of gold. Gold surfaces prefer to form gold-thiol complexes, and therefore are capable of binding cysteamine, thiolated self-assembled monolayers (SAMs), or biorecognition components. As a noble metal, it is worth exploring the methods on how to effectively use gold in the fabrication of electrodes. For example, Yu *et al.*¹⁴² prepared a film electrode *via* sputtering gold onto a polyethylene terephthalate (PET) substrate. The loading of the gold material was well controlled using this technique. Moreover, printing methods (*e.g.*, screen printing) are advantageous for large volume production and depositing electrode materials (*e.g.*, gold¹⁰⁷ and carbon^{108,109,153,176,182}) on a wide range of cheap substrates. Some transparent materials used in solar cells such as fluorine-doped tin oxide (FTO)^{162,163} and indium tin oxide (ITO)^{166,167} are also applied to make the electrode of electrochemical biosensors. Field-effect transistors (FET) are mostly used in semiconductors, where Si/SiO₂ is often used as an underlayer substrate.¹¹⁰ Titanium or other composites (*e.g.*, HfO₂/F₁₆CuPc¹¹¹) are then coated on top of the substrate serving as a dielectric layer, with the gold drain-source electrode finally created on the dielectric layer through a lithography technique. Recent advances show that FET systems can be used in electrochemical biosensors for detection of EDCs in water, but the reports are very limited.

3.1.2 Functional supporting on electrode surfaces.

Biological events occurring on a bare electrode (*i.e.*, untreated) often lead to undesired sensing results because of the limited conductivity, small specific area, low stability, and less biocompatibility to biorecognition elements. Hence, the introduction of a functional support on electrode surfaces can rectify these issues, which is regarded as another highly crucial step in the fabrication of electrochemical biosensors. Carbon materials such as graphene (Gr),¹²⁹ graphene nanoribbons (GNRs),¹⁷² nitrogen-doped graphene nanoribbons (NGNRs),¹⁶⁹ graphene nanoplatelets (GNPs),¹⁸⁷ carbon nanotubes (CNTs),^{130,139,141,159,164,165,168,183} and carbon nanofibers (CNFs)¹³⁸ are usually deposited on the surface of the electrode as a conductive underlayer to facilitate electron transfer, increase the surface area, and improve the mechanical strength. Alternatively, the surface area can be enlarged by adding porous nanostructures like cobalt phosphide decorated porous carbon microspheres (Co_xP/NC)¹⁴⁹ and nanoporous gold leaf (NPGL).¹⁸¹ Porous three-dimensional graphene (3D-GN) can evenly load functional substances (*e.g.*, Cu and Fe₃O₄) to prevent them from aggregation owing to their surface energy.¹⁷⁴ Derikvandi *et al.*¹⁴⁰ used polyethyleneimine (PEI) to offer a similar microenvironment for the immobilization of platinum nanoparticles (PtNPs). Another purpose of increasing the surface area is to provide sufficient space for the functionalization of biorecognition elements and hence to enhance selectivity and specificity. Metal nanoparticles such

as gold nanoparticles (AuNPs),^{159,161,164,165,169,185,188} silver nanoparticles (AgNPs),¹³² PtNPs,¹⁴⁰ nickel nanoparticles (NiNPs),¹⁶⁰ bimetallic hybrids like gold/palladium nanorods (Au@Pd NRs),¹³⁶ and gold/platinum (AuPt),¹³³ or metal/metal oxide composites, including amino-functionalized magnetite and gold nanoparticles (H₂N-Fe₃O₄/Au NPs),¹⁴⁴ CoFe₂O₄,¹³⁶ functional copper magnetic nanoparticles (CuFe₂O₄-Pr-SH),¹⁴¹ and Ag-Cu₂O,¹⁴⁸ not only promote electron transportation, but also have strong electrochemical activity and enzyme-like catalytic or electrocatalytic ability, which belong to another class of common functional supports. Additionally, biochar nanoparticles (BCNPs) can also exhibit excellent electrocatalytic properties as well as a large surface area and have been well-developed in electrochemical biosensing.^{177,178} Quantum dots are a type of nanomaterial with a size less than 100 nanometers. Graphene quantum dots (GQDs)¹⁵⁶ and palladium quantum dots (PdQDs)¹⁶³ have been used to modify the electrode due to their good water solubility and biocompatibility. Apart from graphene, other two-dimensional (2D) materials have been discovered in recent years. Some studies have confirmed that the use of MoS₂ in combination with graphene can effectively improve the sensitivity of the detection platform.^{112,152} As a novel planar material, MXene has been applied to support electrodes because of its large specific surface area, high electron mobility, and hydrophilic nature.^{113,173}

Electrode-supporting substances that contain functional ends are also frequently involved to present binding sites for biorecognition components. Instead of pristine graphene, graphene oxide (GO) is often employed to bind to other electrode additions such as ionic liquids (ILs),¹⁵³ 3-aminopropyltriethoxysilane (APTES),¹⁵⁸ and metal nanoparticles,¹⁸⁴ relying on its abundant oxygen-containing functional groups. However, the limited conductivity of GO usually leads to poor sensitivity in biosensing. Facing this challenge, the GO after being loaded with functional compounds tends to be chemically^{133,136,147,148,150,175,186} or electrochemically^{157,160,171,185,187} converted into reduced graphene oxide (rGO) to maximally recreate the original conductivity of graphene. Multi-walled carbon nanotubes (MWCNTs) with carboxylic groups can provide reaction sites for antigens,¹³⁰ deoxyribonucleic acid (DNA) sequences,^{140,144} enzymes,¹⁷⁹ and cyclodextrins.¹⁸² SAMs are sometimes introduced onto metal electrodes, where the functional terminals are able to couple with receptors, for instance, cysteamine grown on a gold surface to support MIP,¹⁶⁹ mercaptoacetic acid fixed on a gold surface for antibody immobilization,¹³⁵ and mercaptoundecanoic acid (11-MUA) grafted at a silver surface for the surface anchoring of an antibody.¹³⁷ Serving as a specific organic linker for carboxylic groups, ethyl (dimethylamino propyl) carbodiimide (EDC)¹⁴⁰ or ethyl (dimethylamino propyl) carbodiimide/*N*-hydroxysuccinimide (EDC/NHS)^{135,137,144,178} is often included in an experiment to achieve activation and protection from carboxylic radicals. However, based on the conclusion of Zhang *et al.*,¹¹⁴ Liang *et al.*¹³⁴ utilized graphene oxide



nanoribbons/multi-walled carbon nanotubes (GONRs/MWCNTs) hybrids to implant antigens without using linkage chemicals. Some polymers containing amino groups such as poly (1,5-diaminonaphthalene) (P(1,5-DAN))¹²⁹ and polyaniline (PANI),¹³² can bind to antibodies through a direct and glutaraldehyde (GA)-linked interaction, respectively. AgNPs are capable of fixing thiol-terminated aptamers *via* the Ag–S bond,¹³² while AuNPs have the ability to fix antigens,¹³⁰ thiolated aptamers^{150,152,153} or complementary chains¹⁴⁹ *via* the Au–S bond. Moreover, Beiranvand *et al.*¹³⁹ functionalized Fe₃O₄/AuNPs with amino groups to react with GA at first, leaving free aldehyde groups that were used to combine with non-thiolated aptamers. Redox probes can elevate electrochemical activity and amplify sensing signals. Methylene blue (MB)^{142,143,145,153,154,181,185} is the most familiar redox probe applied to electrochemical biosensing, along with others like toluidine blue (TB),¹³⁰ ferrocene (Fc),^{146,150} thionine (Thi),¹⁵² and biotin,¹⁴⁸ are also reported. Cetyltrimethylammonium bromide (CTAB) is a cationic surfactant, can either adsorb nanoparticles by virtue of electrostatic effects,¹⁷³ or preserve the activity of biological enzymes in the event of substrate reactions.¹⁷⁹ The deposition of chitosan^{134,175} or Nafion^{168,176,177} on a biofunctionalized surface enables the electrode durability to be upgraded. ILs have the ability to form films as well and exhibit intense adsorption effects on the analyte of interest.^{153,170} Blocking agents such as bovine serum albumin (BSA)^{130,131,133,135,136,147} and 6-mercapto-1-hexanol (MCH),^{141–143,148,149,152–155} are always used as a final functional layer on the surface of immunosensors and aptasensors, respectively. These chemicals can cover unbound active sites to protect target molecules from any non-specific interactions, thus enhancing the sensitivity.

3.2 Receptors for biorecognition of estrogens

Biorecognition elements are the core constituents that contribute to electrochemical biosensors. Receptors that are commonly used to electrochemically detect EDCs in water, including antibodies (Abs), aptamers, MIPs, enzymes, and cyclodextrins, are often reported.

3.2.1 Antibody-based receptors. Antibodies belong to one of the most utilized biorecognition components and can be further classified into polyclonal and monoclonal subtypes. Electrodes functionalized with antibodies are usually called immunosensors. Polyclonal antibodies (pAbs) are the proteins generated after an exotic immunogen stimulation towards the immune system in living organisms (*e.g.*, rabbit¹²⁹ or chicken¹⁰⁷). pAbs can conjugate with multiple epitopes on a single antigen, which can easily trigger a cross-reaction in complex sample environments that contain more than one analyte.¹¹⁵ The production of monoclonal antibodies (mAbs) normally undergoes a half *in vivo* (*i.e.*, immunization activated after antigen injection) and a half *in vitro* (*i.e.*, B cell-myeloma hybridoma culture) process.¹¹⁶ mAbs are mostly obtained from mice,^{117,129,135} which are

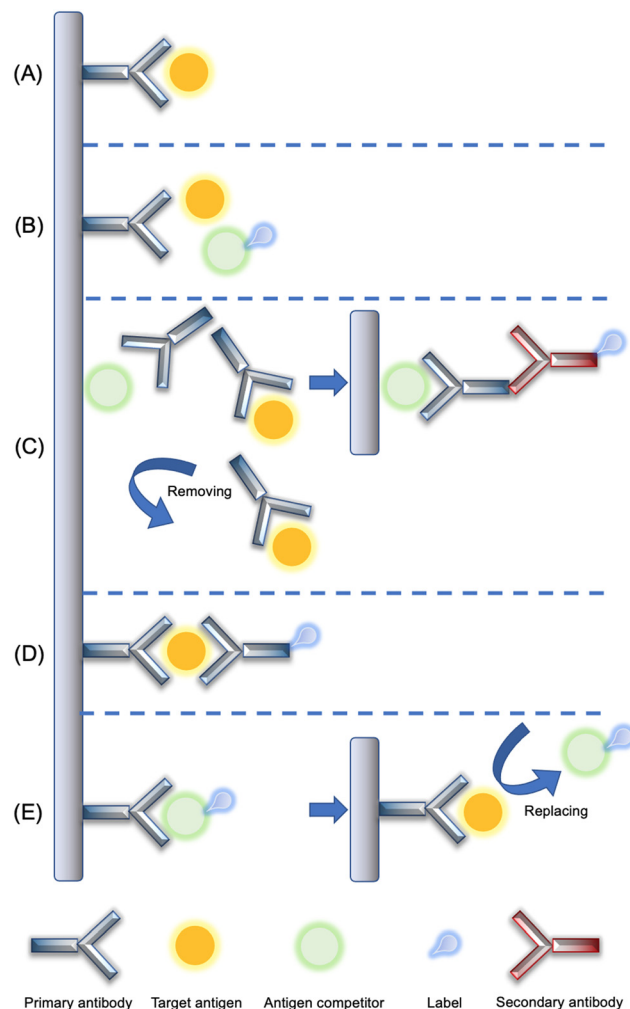


Fig. 4 Different binding schemes between antibodies and antigens in electrochemical immunosensing: (A) direct, (B) direct competition, (C) indirect competition, (D) sandwich, and (E) displacement.

different from pAbs as they interact only with a single epitope on an antigen. mAbs have the advantages of batch-to-batch reproducibility and negligible cross-reactivity but they are time-consuming to prepare and more expensive.¹¹⁸ Recent advances present that pAbs are more prevalent than mAbs used as biological receptors in electrochemical immunosensors because they can offer comparable sensitive results with a lower cost. The same as ELISA, the antibody–antigen conjugation in electrochemical immunosensing can also be divided into direct-, directly competitive-, indirectly competitive-, and sandwich schemes (Fig. 4). The direct scheme is the simplest method to detect the analyte. However, the non-specific binding of target molecules often occurs during a biorecognition process to interfere with the normal binding signals. Direct competition is another common method, where the experiment often introduces an antigen with different concentrations and a labeled competitor at a fixed concentration to simultaneously bind to the antibody. The directly competitive method, which reflects the analyte concentration by quantifying the amount of



antibody-labeled competitor conjugation, avoids the interference of non-specific binding, and thus improves sensitivity and accuracy. Most of the EDCs belong to small molecules as their molecular weights are below 900 Dalton. It is hard to form a stable dual antibody binding (*i.e.*, sandwich scheme) when the target molecules are small molecules. This is the reason for fewer reports on the sandwich-type detection method. However, the determination of small molecules can be realized by indirect competition. An indirectly competitive method usually contains two stages: 1) the introduced analyte, and the competitor immobilized on the electrode surface competitively bind to the primary antibody (Ab_1); 2) the remaining Ab_1 -competitor continuously bind to the labeled secondary antibody (Ab_2) after the removal of Ab_1 -analyte composites. It is noted that indirect competition can also cause cross-reaction due to the use of two types of antibodies, in which extra concern should be taken especially in the detection of multi-targets.¹¹⁹ Lu *et al.*¹²⁰ reported an immunosensing conducted *via* a displacement method. In their work, the competitive antigen with a butyric acid-horseradish peroxidase (HRP) was firstly recognized by the antibody, followed by a replacement of the competitor after the analyte was introduced. A lower LOD for BPA was obtained using the proposed method in comparison to the conventional direct competition.¹²⁰

3.2.2 Aptamer-based receptors. Aptamers are single-stranded DNA (ss DNA) or ribonucleic acid (RNA) sequences that are selected from the ssDNA or RNA pool by virtue of the systematic evolution of ligands by exponential enrichment (SELEX).¹²¹ Aptamers can effectively bind to target molecules due to the structural change of the chains, which are also a type of common biorecognition element. Electrodes loaded with aptamers are often called aptasensors. Some aptamers or complementary DNA (cDNA) strands possess functional ends that can be immobilized on the surface of the electrode through different bonds. Aptamer-related detections are achieved following three methods. The first one is that aptamers directly capture the target molecules causing the increase of steric hindrance against the electron transfer from the bulky solution towards the electrode surface. The second one firstly allows a cDNA fixed on the electrode surface, followed by a helical binding with the aptamer. After involving the analyte, the aptamer is not dissociated with the cDNA, yielding a larger steric hindrance. These two methods provide decreased responses with the increase of analyte concentration. The only difference of the third method from the second one is the aptamer dissociated from cDNA after forming the aptamer-analyte conjugation. Steric resistance is reduced in this method and the electrochemical responses are positively correlated with the target concentration.

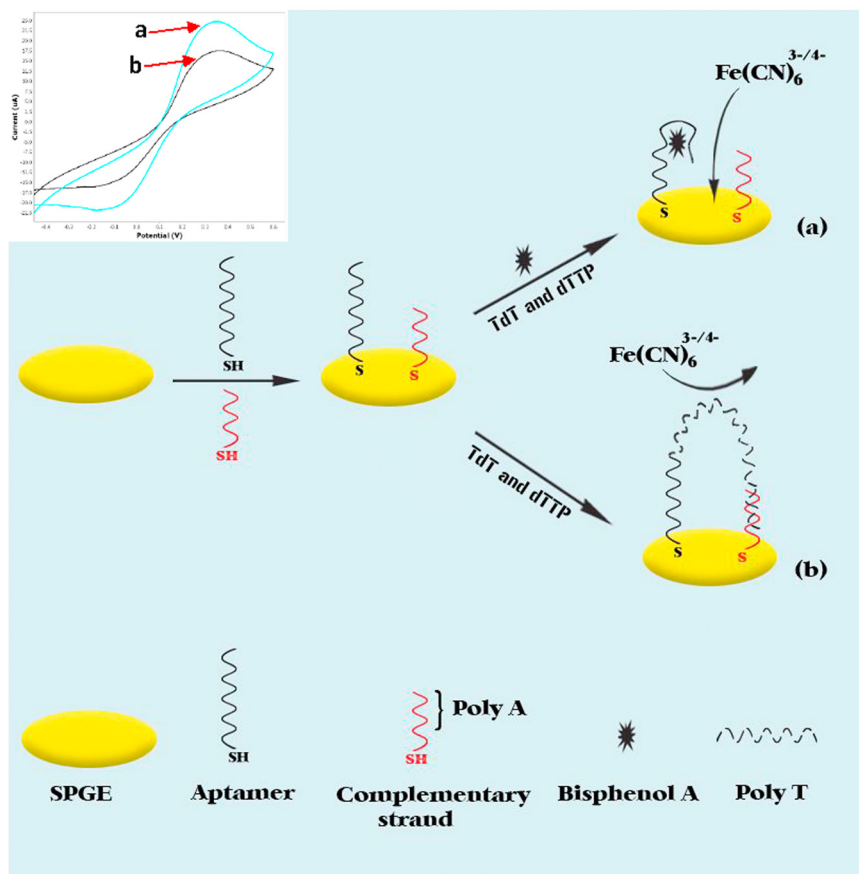


Fig. 5 Schematic illustration of the electrochemical aptasensor for BPA detection based on nontarget-induced extension of aptamer length triggered by TdT (reprinted with the permission from ref. 121).



Abnous *et al.*¹²² constructed a special aptasensor for monitoring of BPA (see Fig. 5). The aptamer and a short complementary sequence initially attached to the electrode surface, and subsequently bridged through poly T that was catalyzed by terminal deoxynucleotidyl transferase (TdT), forming a physical barrier against the redox couple (*i.e.*, Fe(CN)₆^{3-/4-}). Chemical events happened at the gold surface since the aptamer length extension reaction was not triggered after the analyte was involved.¹²²

3.2.3 Molecularly imprinted polymer-based receptors.

MIPs are a promising alternative of biorecognition elements that can exhibit strong specificity like antibodies, owing to the use of biocompatible polymers and organic functional reagents. In the past decade, more and more research groups focus their attention on studying of MIPs. A molecular imprinting process usually requires a specific monomer, cross-linker, and template molecule. The polymeric matrix is then built after a few cycles of polymerization activated by an initiator. Tailored cavities are subsequently obtained by the removal of template molecules. According to the cited papers in this review, functional monomers including pyrrole,^{123,156,159,160,171} 1*H*-pyrrole-3-carboxylic acid (py-3-COOH),¹⁷³ methacrylic acid (MAA),^{124,165,166} chitosan,¹⁵⁷ *o*-phenylenediamine (*o*-PD),¹⁶⁷ acrylamide,¹¹¹ 3,4-ethylenedioxythiophene (EDOT),¹⁷² and 4-aminothiophenol (4-ATP or *p*-ATP)¹⁶⁴ are often used. Some research groups prefer to use a comonomer to create the MIP, including MAA/4-ATP,¹⁶¹ 3-aminopropyltriethoxysilane (APTES)/phenyltrimethoxysilane (PTMS),¹⁶⁸ 2-aminothiophenol (2-ATP)/4-ATP¹⁶⁹ and *o*-phenylenediamine/*o*-toluidine.¹⁷⁰ Ethylene glycol dimethacrylate (EGDMA) and azodiisobutyronitrile (AIBN) are common chemicals for a cross-linker and initiator, respectively.^{158,161,165,166} There is an exception reported in Venkatesh *et al.*,¹¹¹ where the cross-linking and activation were conducted in the presence of *N,N*-methylene-

bisacrylamide and tetramethylethylenediamine (TEMED), respectively. Zhang *et al.*¹⁶⁸ applied tetraethoxysilane (TEOS) as a special linkage reagent (see Fig. 6). Chemicals such as methanol/acetic acid,^{156,158,159,161,165,168–170,172} ethanol/acetic acid,^{160,171} hydrochloric acid,^{157,164} and sodium hydrate mixed with ethanol¹⁷³ are used to remove template molecules after the imprinted matrix was generated. For some photoelectrochemical MIPs, the template molecules are normally removed by water with high temperature annealing,¹⁵⁶ ethanol,¹⁶³ deionized water/ethanol,¹⁶⁶ and sodium hydrate.¹⁶⁷

3.2.4 Enzymatic and cyclodextrin-based receptors.

Enzymes can also play the role of biorecognition element in electrochemical biosensors. A biological enzyme is a protein that shows strong selectivity and specificity towards target molecules. Enzymes can dramatically reduce the activation energy of chemical events to accelerate the reaction process, presenting excellent electrocatalytic nature. Common enzymes used for electrochemical detection of EDCs in water are tyrosinase (Tyr) and laccase (Lac). Tyr is a copper-containing enzyme mainly obtained from plant or animal tissues (*e.g.*, mushroom). The primary function of Tyr is to catalyze the oxidation of tyrosine and control the production of melanin and other pigments. BPA determined using Tyr-based biosensors is normally subjected to a primary oxidation to form catechol, followed by a secondary oxidation of catechol to form *o*-quinone. The reduction of *o*-quinone forms an enzymatic reaction loop to regenerate catechol, yielding reductive currents that indicate the concentrations of BPA.^{125,177–179} Lac is different from Tyr as it belongs to a multi-copper-containing oxidase family. Lac can be obtained from various species of fungi, plants, insects, and bacteria.¹²⁶ The oxidation in Lac-induced reactions usually transfers only one electron, while molecular oxygen or a redox probe often serves as an electron acceptor contributing to the reduction.^{176,181} Cyclodextrins are a class of barrel-like

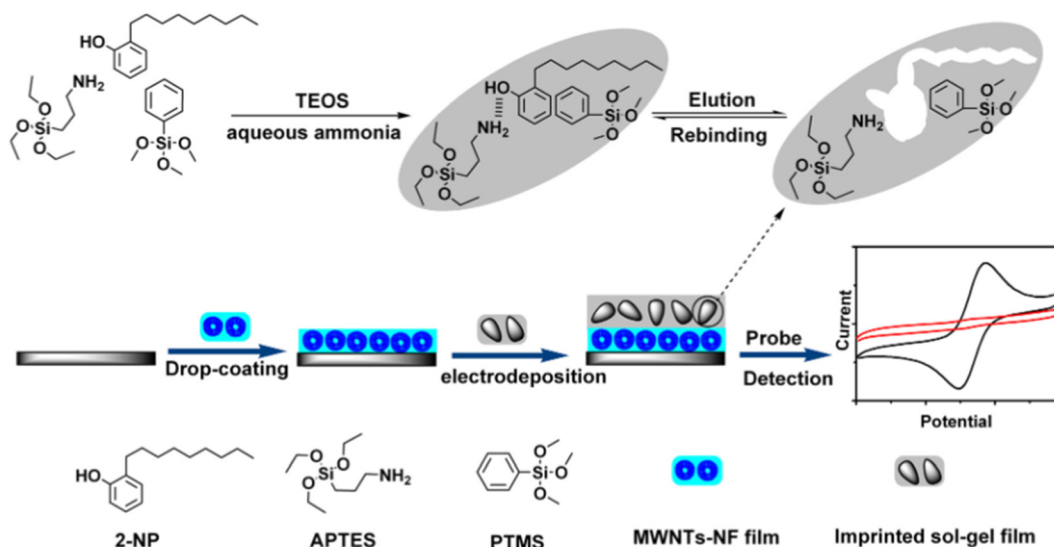


Fig. 6 Schematic representation of the preparation procedure of MIP/sol-gel/MWNT-NF/GCE (reprinted with the permission from ref. 167).



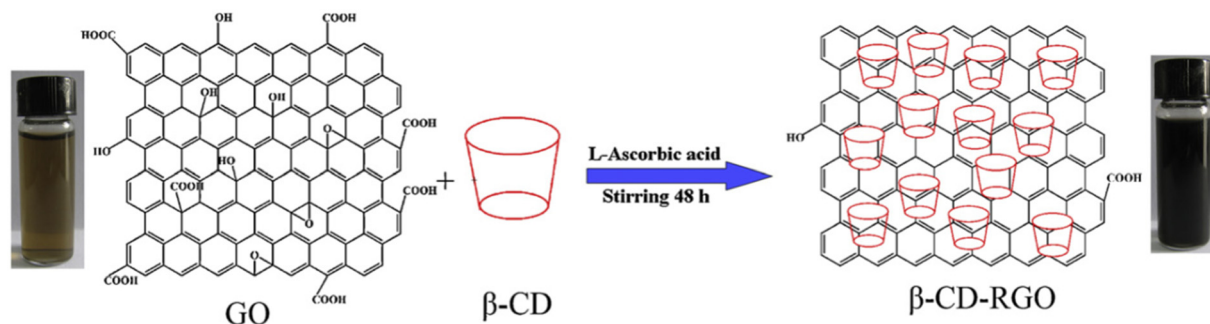


Fig. 7 Illustration of the preparation procedure of β -CD-RGO (reprinted with the permission from ref. 126).

Table 2 Electrochemical immunosensors for detection of EDCs in water

EDC	Recognition biocomponent	Detection scheme	Electrode	Electrode functional support from bottom to top	Electrochemical technique	Dynamic range (ng mL ⁻¹)	LOD (pg mL ⁻¹)	Real water sample analysis	Ref.
BPA	mAb, pAb	Sandwich	GCE	Gr/P(1,5DAN), α_1 BPA, BPA antigen, α_2 BPA, Gr-AgNPs	SWV	0.228–1.824	136.8	Drinking water	129
TBBPA	Ab	Indirect competitive	GCE	MWCNTs/Au-TB, competitor; antigen, BSA, Ab ₁ , analyte, Ab ₂ with label; MnO ₂ /Pd	Amperometry	0–81	170	Pure water, tap water, pond water, and lake water	130
TBBPA	Ab	Indirect competitive	GCE	MWCNTs, competitor; antigen, BSA, Ab ₁ , analyte, Ab ₂ with label; AuPd-NH ₂ -fMnO ₂	Amperometry	0–243	100	Pure water, tap water, pond water, and lake water	131
PCB 28	pAb	Direct	GCE	AgNPs, PANI, GA	SWV	0.2–1.2	63	Tap water	132
DMP	Ab	Direct competitive	GCE	AuPt-GS, anti-DMP, BSA, competitor; PtPb(tag)-BSA-DMP	DPV	1–1000	330	Tap water and lake water	133
DBP	Ab	Indirect competitive	GCE	MWCNTs@GONRs, chitosan, competitor; antigen, Ab ₁ , analyte, Ab ₂ with label; AuNPs	EIS	0.005–0.5	7000	Pure water, tap water, pond water, and river water	134
E2	mAb	Direct competitive	Gold	Mercaptoacetic acid, EDC/NHS, BSA, competitor; Cu ₂ S-BSA-E2	SWV	0.025–7.5	7.5	Water sample	135
E2	Ab	Direct	GCE	CoFe ₂ O ₄ /rGO, Au@Pd NRs, BSA	Amperometry	0.01–18	3.3	River water	136
E2	mAb	Direct	Silver	11-MUA, EDC/NHS	EIS	0.005–0.2	1	Water sample	137

products naturally formed through the circular cross-linking of oligosaccharides. The resulting homopolymer always with a hexagonal, heptagonal, or octagonal structure depends on the number of glucose monomers incorporated, corresponding to α -, β -, and γ -cyclodextrin, respectively (see Fig. 7).¹²⁷ Cyclodextrins have a general property of outer-hydrophilicity and inner-hydrophobicity that present adsorption onto target molecules to establish stable ‘host-guest’ inclusion complexes.¹²⁸ Among them, β -cyclodextrin (β -CD) is the one that has been widely used in electrochemical biosensors for detection of EDCs in water.

3.3 Detection of EDCs using electrochemical biosensors

In this section, detection results of EDCs in water using various electrochemical biosensors are summarized in several

tables depending on different biorecognition elements applied: Table 2 for electrochemical immunosensors, Table 3 for electrochemical aptasensors, Table 4 for MIP-based electrochemical sensors, and Table 5 for enzymatic and β -CD-based electrochemical sensors.

3.3.1 Electrochemical immunosensors. Antibody-antigen conjugation was often used in electrochemical immunosensors for detection of EDCs in water (Table 2). Nguyen *et al.*¹²⁹ fabricated an electrochemical immunosensor following a sandwich determination protocol. After the antigen conjugated with the monoclonal anti-BPA antibody (α BPA) on the electrode surface, polyclonal anti-BPA antibody (α_2 BPA) coated graphene-silver nanoparticles (Gr-AgNPs) were introduced to accomplish the sandwich-type biosensing. Under the SWV analysis, the immunosensor had a limit of detection (LOD) of 136.8 pg mL⁻¹. However, the detection



Table 3 Electrochemical aptasensors for detection of EDCs in water

EDC	Recognition biocomponent	Electrode	Electrode functional support from bottom to top	Electrochemical technique	Dynamic range (ng mL ⁻¹)	LOD (pg mL ⁻¹)	Real water sample analysis	Ref.
BPA	Thiolated aptamer	GCE	CNFs/AgNPs	SWV	0.0228–2.28	88.92	Real water sample	138
BPA	Aptamer	GCE	MWCNTs/Fe ₃ O ₄ /AuNPs-NH ₂ /GA	DPV	0.228–136.8	68.4	Environmental water sample	139
BPA	Aptamer	GCE	PEI/PtNPs/MWCNTs-COOH/EDC/ssDNA2	DPV	0.228–91.2	47.88	Environmental water sample	140
BPA	Thiolated aptamer	GCE	MWCNTs/Au/CuFe ₂ O ₄ -Pr-SH, MCH	DPV	0.0114–2.052	5.746	Mineral water	141
BPA	Thiolated aptamer	Au/PET	MB, MCH	LSV	0.001–100	0.4	Tap water	142
BPA	Thiolated aptamer	Gold	cDNA, MB, MCH	CV	0.0001–1	0.284	Drinking water	143
BPA	Aptamer	GCE	MWCNTs-COOH/EDC/NHS, NH ₂ -Fe ₃ O ₄ NPs/AuNPs, cDNA, SWCNTs	DPV	2.28×10^{-11} – 2.28×10^{-6}	1.824×10^{-8}	Lake water	144
PCB 72	Thiolated aptamer	Gold	MB@CaCO ₃ , PAH, EDTA	DPV	10^{-8} – 0.01	1	Tap water and lake water	145
PCB 77	Thiolated aptamer	Gold	Fc	DPV	0.2–200	10	Tap water	146
PCB 77	Aptamer	Gold	rGO/NiHCF NPs, BSA	DPV	0.001–0.1	0.22	Tap water and lake water	147
PCB 77	Aptamer	Gold	Ag–Cu ₂ O/rGO, thiolated HPDNa, MCH, biotin, SA-MB@CeO ₂ /AuPt	DPV	0.0001–1	0.069	Tap water	148
PCB 77	Aptamer	Gold	Co _x P/NC, AuNPs, thiolated cDNA, MCH, AuNRs/MB	DPV	0.00001–100	0.059	Tap water	149
PCB 77	Thiolated aptamer	Gold	RGO-AuNPs, Fc	DPV	10^{-6} – 10	0.0001	Tap water	150
PCB 77	Thiolated aptamer	BDD	T-Au-NPs	EIS	2.92×10^{-7} – 2.92×10^{-3}	9.344×10^{-5}	Lake water	151
PCB 77	Thiolated aptamer	GCE	MoS ₂ -rGO/Thi/AuNPs, MCH	DPV	3×10^{-7} – 0.1	8×10^{-5}	River water and tap water	152
DBP	Thiolated aptamer	SPCE	IL/GO, AuNPs, MCH, MB	DPV	0.00014–7	0.042	River water	153
DEHP	Aptamer	Gold	Thiolated CP, DNA junction, MCH, MB	DPV	0.1–5000	40	Tap water	154
DEHP	Aptamer	Gold	MCH	EIS	0.0076–2000	0.103	Tap water and lake water	155

range lay only within 0.228–1.824 ng mL⁻¹.¹³⁰ Yakubu *et al.*¹³⁰ created an indirect competitive immunosensor where the antigen was fixed on the transducer surface serving as the competitor to TBBPA. The competition was induced when the primary antibody (Ab₁) and analyte molecules were involved. After washing off the TBBPA-bound Ab₁, the remaining Ab₁-antigen combination was successively recognized by the secondary antibody (Ab₂) coupled to the MnO₂/Pd label. Amperometric results presented a linear range of 0–81 ng mL⁻¹, and a LOD of 170 pg mL⁻¹. The test for stability was performed by measuring current responses of the biosensor every three days, and a recovery of 86.85% was obtained from the one stored at 4 °C for two weeks. The reproducibility assay was conducted on five electrodes and showed a small relative standard deviation (RSD) of 3.17%.¹³⁰ From the same research group, the structure of the label was improved as AuPd decorated MnO₂ nanoflowers (AuPd-NH₂-fMnO₂), followed by the identical biosensing method to reduce the LOD to 100 pg mL⁻¹.¹³¹ Zhang *et al.*¹³⁶ provided a direct competitive sensing platform in which the analyte (*i.e.*, E2)

and Cu₂S-BSA labeled E2 competitively bound to the monoclonal antibody on the gold surface. The labeled E2-antibody composite was eventually measured reflecting the analyte concentration. A linear range of 0.025–7.5 ng mL⁻¹ was achieved and the minimum detection for E2 was 7.5 pg mL⁻¹.¹³⁶ In contrast to the competition scheme, two papers focused on the electrochemical detection of E2 using direct antibody–antigen interaction. Zhang *et al.*¹³⁶ prepared a label-free electrochemical immunosensor supported by the underlayer of spinel oxide-loaded rGO and non-biological enzyme (*i.e.*, Au@Pd NRs). A wider dynamic range of 0.01–18 ng mL⁻¹ and a lower LOD of 3.3 pg mL⁻¹ were acquired using amperometric measurement.¹³⁶ Based on this, Singh *et al.*¹³⁷ further improved the LOD to 1 pg mL⁻¹, using non-faradaic EIS.

Apart from BPs and estrogens, antibody-based electrochemical biosensors were also frequently used for monitoring of PCBs and phthalate esters in water. Nevertheless, there are few comparative studies on a single EDC. Khesuoe *et al.*¹³² reported a direct immunosensor for



Table 4 Electrochemical MIP sensors for detection of EDCs in water

EDC	Biorecognition component	Electrode	Electrode functional support from bottom to top	Electrochemical technique	Dynamic range (ng mL ⁻¹)	LOD (pg mL ⁻¹)	Real water sample analysis	Ref.
BPA	MIP	GCE	GQDs	DPV	22.8–11 400	9120	Seawater and bottled water	156
BPA	MIP	ABPE	rGO	Second-order derivative LSV	1.824–4560	1368	Drinking water and beverages	157
BPA	MIP	GCE	GO/APTES	DPV	1.368–4560	684	Mineral water	158
TBBPA	MIP	GCE	MWCNTs-AuNPs	DPV	0.272–544	130.56	Seawater and tap water	159
TBBPA	MIP	GCE	rGO/NiNPs	DPV	0.272–5440	70.72	Tap water, rainwater, and lake water	160
TBBPS	MIP	CPE	AuNPs	DPV	0.0566–5.66	16.414	Tap water, drinking water, and lake water	161
PCB 101	MIP	FTO	TiO ₂ NRs	Photoelectrochemistry	2.6×10^{-5} –9.78	0.00326	River water	162
PCB 101	MIP	FTO	Pd QDs@TiO ₂ NRs	Photoelectrochemistry	3.26×10^{-5} –0.163	0.0163	Lake water	163
DBP	MIP	BDD	MWCNTs/AuNPs	DPV	2.78–2780	917.4	Lake water and tap water	164
DBP	MIP	GCE	MWCNTs/AuNPs	CV	0.1–10 000	5.09	Tap water	165
DOP	MIP	ITO	Cu ₃ (BTC) ₂ @Cu ₂ O	Photoelectrochemistry	0.00975–39	3.5685	Bottled water	166
DOP	MIP	ITO	Bi ₂ S ₃	Photoelectrochemistry	0.000195–0.0273	0.039	Plastic bottled water, and rainwater	167
2-NP	MIP	GCE	MWCNTs-Nafion/sol-gel	DPV	44–79 200	13 200	Tap water and river water	168
4-NP	MIP	Gold	TiO ₂ -NH ₂ /AuNPs/Cysteamine	Amperometry	209–105 600	70 400	Tap water	169
4-NP	MIP	GCE	NGNRs-IL	LSV	8.8–1320	1760	Lake water, river water, and tap water	170
4-NP	MIP	CE	GP	DPV	0.01–10	3.5	Rainwater and lake water	171
4-tert-OP	MIP	GCE	GNRs	LSV	4.12–1648	206	River water and bottled water	172
DES	MIP	GCE	Ti ₃ C ₂ T _x /Cu ₂ O NPs/CNT-CTAB	DPV	2.68–18 760	1608	Lake water	173

Table 5 Electrochemical enzymatic and cyclodextrin sensors for detection of EDCs in water

EDC	Biorecognition component	Electrode	Electrode functional support from bottom to top	Electrochemical technique	Dynamic range (ng mL ⁻¹)	LOD (pg mL ⁻¹)	Real water sample analysis	Ref.
BPA	Lac	GCE	(3D-GN)/Cu/Fe ₃ O ₄	DPV	1641.6–4104	387 600	Tap water	174
BPA	Lac	GCE	rGO-Fe ₃ O ₄ NPs/Chit95	Amperometry	6–228	4104	Bottled water	175
BPA	Tyr	SPCE	Au leaf-like microstructures, Nafion	CV	114–11 400	17 556	Tap water, and mineral water	176
BPA	Tyr	GCE	BCNPs, Nafion	Amperometry	4.56–2280	725.04	Ground water	177
BPA	ML-Tyr	GCE	Mag-BCNPs-COOH, EDC/NHS	Amperometry	2.28–230.28	633.84	Environmental water	178
BPA	Tyr	BDD	MWCNTs-diazonium	CV	0.00228–22.8	2.28	River water	179
4-NP	Lac	CPE	CTAB	Amperometry	1100–66 000	200	Water sample	180
BPA	β-CD-SH	Gold	NPGL, MB	SWV	68.4–22 800	13 680	Tap water	181
BPA	β-CD	SPCE	MWCNTs	LSV	28.5–6840	3137.28	Lake water and tap water	182
BPS	β-CD	GCE	MWCNTs	SWV	125.135–15016.2	12513.5	Drinking water and tap water	183
BPA	PCD	GCE	GO, AuAgPt	DPV	BPA: 1.368–912	BPA: 456	River water	184
BPS					BPS: 2–2000	BPS: 667.5		
PCB 77	β-CD-SH	GCE	AuNPs/GR, MB	DPV	14.6–2920	8176	Canal water and lake water	185
DEHP	β-CD	GCE	Graphene	EIS	780–7020	46 800	Wastewater	186
DEHP	β-CD	GCE	DAD/rGO	EIS	78–468	3900	River water	187
4-NP	β-CD	GCE	GNP/sol-gel, AuNPs	CV	2.2–220	660	Lake water	188



PCB 28. However, cross-reactivity among the analyte, PCB 180, and benzyl chloride (BnCl) was noted because of their similar structures, leaving poor specificity. A direct competitive assay in between the target DMP and the PtPb tagged DMP (*i.e.*, competitor) was conducted by Liu *et al.*¹³³ on the surface of a GCE. The DPV result was compared with those obtained from chromatographic and SERS methods, evidencing a lower LOD of 330 pg mL⁻¹. Liang *et al.*¹³⁴ fabricated an impedimetric immunosensor for the determination of DBP relying on the mechanism of indirect competition. AuNPs enhanced Ab₂ was added into the testing system after the removal of DBP-Ab₁ complexes to amplify the electrochemical signals. A detection limit of 7000 pg mL⁻¹ was achieved, which was ten times lower than that yielded by the conventional ELISA approach.¹³⁴ A new class of biosensors has emerged which do not need separate labeling steps for their preparation and can provide rapid and *in situ* sensing in fields. In this regard an ultrasensitive immunosensor was fabricated by electropolymerization over a graphene screen printed electrode (SPE). The PANI/Gr-SPE-devices displayed linear responses to low molecular weight, poorly water soluble, estrogenic compounds, in EIS assays, from 0.0975 ng L⁻¹ to 200 ng L⁻¹ in water samples. This method was considered better compared with previous reports due to low detection limits of 0.043 pg L⁻¹ for E1, 0.19 ng L⁻¹ for E2 and 0.070 pg L⁻¹ for EE2.³¹ Another label free impedimetric biosensor was developed by immobilizing both estrogen receptor- α and bovine serum albumin on gold electrode surfaces for detecting 17 β -estradiol. The binding of the estrogen contaminant on the surface of the label free sensor increased the charge transfer resistance which was proven by EIS analysis. The present biosensor gives a linear response ($r^2 = 0.992$) for 17 β -estradiol concentration from 1.0×10^{-13} to 1.0×10^{-9} M with a detection limit of 1.0×10^{-13} M (S/N = 3).³² In a similar study, ER- α was coupled onto the gold electrode through its 6-His tag and NTA-copper complex to prepare a label free biosensor for estrogen. After interaction of estradiol with ER- α , the biosensor presents a well-defined peak at +500 mV due to estradiol oxidation (E17 peak). These rapid and innovative sensors were used to detect estradiol from hospital effluents.⁹⁴

3.3.2 Electrochemical aptasensors. Aptamer-based electrochemical biosensors have been widely reported in the past decade. Electrochemical aptasensors used for detection of EDCs in water are summarized in Table 3. Tsekeli *et al.*¹³⁸ developed an electrochemical aptasensor for BPA. The electron transfer of the redox couple near the electrode surface was monitored using SWV. Weakened electrical signals were observed when the BPA concentration was increased due to the enlarged steric hindrance caused by the aptamer-antigen complex. The biosensor reached a LOD of 88.92 pg mL⁻¹ and a good selectivity was verified by mixing BPA with other six interferents.¹³⁸ Similar studies were also performed by Beiranvand *et al.*¹³⁹ and Derikvandi *et al.*,¹⁴⁰ where the biosensors exhibited a slightly broader detection range and improved LODs. MCH was sometimes introduced

to the electrode surface after the functionalization of aptamers to avoid any non-specific adsorption. This principle was applied in the aptasensor provided by Baghayeri *et al.*,¹⁴¹ of which a well-improved LOD of 5.746 pg mL⁻¹ was achieved. Thiolated aptamers were also directly immobilized on gold electrodes. In the presence of MCH, these gold-based aptasensors presented better performances like wider dynamic ranges (*e.g.*, four or five orders of magnitude) and LODs below 1 pg mL⁻¹.^{142,143} For ultrasensitive biosensing, Zhao *et al.*¹⁴⁴ reported an aptasensor in which the electrochemical signal was significantly amplified as single-walled carbon nanotubes (SWCNTs) attached to the cDNA in lieu of the aptamer after BPA was introduced. Under the DPV measurement, the aptasensor offered a linear response from 2.28×10^{-11} to 2.28×10^{-6} ng mL⁻¹, with an ultralow LOD of 1.824×10^{-8} pg mL⁻¹.¹⁴⁴

Other than BPs, aptamer-based electrochemical biosensors used for the detection of PCBs (*e.g.*, PCB 77) in a variety of water samples have also been reported. Utilizing a PCB 77 specific DNA sequence as the bioreceptor, Wu *et al.*¹⁴⁶ developed an aptasensor with a resulting LOD of 10 pg mL⁻¹. The proposed biosensor successfully detected PCB 77 in real tap water and displayed good selectivity to the analyte even in the presence of other PCB congeners involving PCB 81, PCB 126, PCB 169, and PCB 189. Moreover, the electrochemical biosensing only took 10 seconds, which was much faster than traditional GC methods of about 10 minutes.¹⁴⁶ A lower LOD (*i.e.*, 1 pg mL⁻¹) was achieved by Fan *et al.*,¹⁴⁷ because the nickel hexacyanoferrate nanoparticles (NiHCF NPs) exerted a synergistic effect with BSA on enhancing the aptasensor sensitivity. Some aptamers were immobilized on the electrode surface *via* their complementary parts (*e.g.*, hairpin DNA¹⁴⁸ and cDNA¹⁴⁹), while the other ends of the chains were connected by a signal generator namely streptavidin (SA)-MB@CeO₂/AuPt (ref. 148) and gold nanorods (AuNRs)@MB,¹⁴⁹ respectively. Such signal generator-aptamer hybrids were prone to dissociate from the electrode surface and bind to the target molecules, leading to weakened signals. This process can significantly enlarge the range of detection and reduce the LOD but needs to prepare many functional components resulting in tedious experimental procedures. Another sensing mechanism was based on the spatial change of aptamers to shorten the distance between the signal probe (always attaching to the end of aptamers) and the electrode surface, thus improving the sensitivity. This method involved fewer electrode supports but exhibited even better electrochemical performance.¹⁴⁹ Yuan *et al.*¹⁵¹ achieved an ultrasensitive determination of PCB 77 on the twice sputtered and twice annealed gold nanoparticle (T-Au-NPs) covered BBD electrode. Under faradaic EIS, visible differences can be observed in the Nyquist plot ranging from 2.92×10^{-7} to 2.92×10^{-3} ng mL⁻¹ with a small standard deviation in each concentration, accompanied by an ultralow LOD of 9.344×10^{-5} pg mL⁻¹.¹⁵¹ Another aptamer-induced biosensing in virtue of thionine was performed by Mohammadi *et al.*¹⁵² with similar



ultrasensitive outcomes. The selectivity test was carried out by introducing structural analogs of PCB 77 as well as heavy metal ions and the ratio of DPV responses after introducing interferants was highly close to that of the control group (*i.e.*, PCB 77 only).¹⁵² Han *et al.*¹⁴⁵ constructed a special dendritic aptasensor used for the detection of PCB 72 in water. The aptamers were firstly allowed to grow on the skeleton of CaCO₃ particles loaded with MB. Ethylenediaminetetraacetic acid (EDTA) was then used to dissolve the CaCO₃ framework, leaving aptamer-based microcapsules. MB was finally released from the capsules after introducing the analyte. This biodevice possessed a LOD of 1 pg mL⁻¹ and a broad dynamic range with a six-order-of-magnitude span (*i.e.*, 10⁻⁸–10⁻² ng mL⁻¹).¹⁴⁵

The detection of phthalate esters in water samples was sometimes reported using aptamer-based electrochemical biosensors, but mainly focused on DEHP. Chen *et al.*¹⁵⁴ fabricated an aptasensor for DEHP based on the gold surface. The target-specific aptamers initially combined with the capture probe (CP). Subsequently, the analyte bound to the aptamers and formed a free composite, while the CP continuously combined with a DNA junction comprising of three hairpin DNA and a quarter sequence. The biosensor had a linear range of 0.1–5000 ng mL⁻¹ and a LOD of 40 pg mL⁻¹.¹⁵⁴ A simple biosensor that only contained aptamers and a blocking agent (*i.e.*, MCH) was made by Lu *et al.*,¹⁵⁵ where the use of EIS resulted in a wider range of detection from 0.0076–2000 ng mL⁻¹ and a lower LOD of 0.103 pg mL⁻¹, further verifying the high reliability of impedimetric methods. The prepared biosensor also had an outstanding selectivity towards DEHP in a mixture of other nine phthalate esters.¹⁵⁵ Contrary to the sensing principle of Wu *et al.*,¹⁵⁰ decreased responses were observed by Gurudatt *et al.*,¹⁵³ since the target molecules bound to the aptamers creating a larger distance between MB and the electrode surface. This biosensor realized an ultrasensitive determination of DBP in river water from South Korea.

3.3.3 Electrochemical MIP sensors. MIPs are another type of customarily used recognition element for the electrochemical detection EDCs in water (Table 4). Tan *et al.*¹⁵⁶ created a MIP sensor for the monitoring of BPA and gained a detection range of 22.8–11 400 ng mL⁻¹ with a not very low LOD of 9120 pg mL⁻¹. Real water analysis was implemented by spiking BPA in seawater and bottled water with recoveries comparable to those of HPLC. In addition, the influence of MIP thickness on performance of the proposed sensor was also discussed in their group. Increasing the thickness enlarges the surface area of the polymeric matrix, leading to adequate room for cavity production. Nevertheless, excessive thickness will result in the difficulty on the removal of template molecules. Meanwhile, cavities created in the deep inner area make the analyte unable to bind effectively because of the high mass-transfer resistance. Therefore, the thickness of MIP should be in a reasonable range *via* controlling the cycle of polymerization.¹⁵⁶ Deng *et al.*¹⁵⁷ prepared a MIP sensor based

on the ABPE electrode. Compared with Tan *et al.*,¹⁵⁶ the sensor had a competitive dynamic range and a significantly reduced LOD of 1368 pg mL⁻¹, under the measurement of SDLSV. Dadkhah *et al.*¹⁵⁸ further lowered the LOD to 684 pg mL⁻¹. The high specificity to BPA was derived from the amine groups of APTES attached to the surface of GO. Also, DPV results provided by the MIP sensor were found to be approximately five times larger than those of the non-imprinted polymer (NIP) sensor under the same conditions, demonstrating the usefulness of MIPs.¹⁵⁸ Wu *et al.*¹⁵⁹ and Chen *et al.*¹⁶⁰ developed MIP sensors for the screening of TBBPA in water samples. Using the same electrochemical technique (*i.e.*, DPV), a wider dynamic range as well as a lower detection limit was obtained by the latter group, probably owing to the modification of amine-terminated benzenediazonium (NBD) upon the GCE. Sarpong *et al.*¹⁶¹ successfully detected TBBPS in water samples using a dual-monomer polymerized MIP modified on CPE. However, the detection span was only from 0.0566 to 5.66 ng mL⁻¹ (two orders of magnitude), and there is only one paper focusing on the detection of TBBPS in water using electrochemical MIP sensors, leaving the demand for more parallel research.

Huang *et al.*¹⁶⁹ applied amine-terminated TiO₂ (TiO₂-NH₂) nanoparticles to cover the gold electrode, where the resulting surface was used to absorb AuNPs. A 4-nonylphenol (4-NP) imprinted polymer was then built on top of cysteamine self-assembled nanoparticles under polymerization with thioglycolic acid involved as a stabilizer. However, the detection results were not as low as expected. This is probably due to the use of TiO₂ nanoparticles that are likely to display semiconductive properties but are not very conductive as carbon materials. Like Sarpong *et al.*,¹⁶¹ Pan *et al.*¹⁷⁰ also prepared a MIP sensor for 4-NP based on two monomers. A highly conductive functional composite (*i.e.*, NGNRs-IL) was placed in between the MIP and GCE, serving as an excellent path for electron transfer. Compared with Huang *et al.*,¹⁶⁹ the LOD was decreased by an order of magnitude, 1760 pg mL⁻¹. Chen *et al.*¹⁷¹ reported a simple MIP sensor for the sensitive detection of 4-NP based on a home-made CE. The amine-terminated graphene oxide was electrochemically reduced and covalently bound to the electrode acting as a conductive underlayer. DPV results exhibited a dynamic range of 0.01–10 ng mL⁻¹ with a remarkably lowered minimal detection level of 3.5 pg mL⁻¹.¹⁷¹ As an isomer of 4-NP, the monitoring of 2-nonylphenol (2-NP) in environmental water samples was manifested by Zhang *et al.*¹⁶⁸ on a sol-gel MIP sensor, reaching a wide range of detection (*i.e.*, 44–79 200 ng mL⁻¹). However, the detection limit was still over ten thousand pg mL⁻¹. Pan *et al.*¹⁷² constructed two MIP-based sensing devices for 4-*tert*-OP, one of which used EDOT as a monomer, the other used AuNPs-captured EDOT. The experimental results revealed that the MIP incorporating gold nanoparticles presented a higher sensitivity, wider detection window, and lower LOD than the one without gold nanoparticles.¹⁷²



Additionally, there are also a few reports on the detection of PCBs, phthalate esters, and estrogens in water making use of electrochemical MIP sensors. Shi *et al.*¹⁶² developed a special MIP sensor, where the TiO₂ sol was initially deposited on the FTO substrate *via* spin-coating and subjected to a hydrothermal process with template molecules (*i.e.*, PCB 101) to generate cavity-embedded titanium dioxide nanorods (TiO₂ NRs). In combination with the photoelectrochemical technique, the sensor was able to determine PCB 101 with an ultralow LOD of 0.00326 pg mL⁻¹. Another similar work was also performed by the same group, in which the TiO₂ NRs were further decorated with palladium quantum dots (Pd QDs) and yielded a comparable LOD.¹⁶³ Based on these two studies, they demonstrated that photoelectrochemistry could enable a very broad detection. In the experiment performed by Wu *et al.*,¹⁶⁴ DBP was determined by the imprinted cavities on a BDD electrode ranging from 2.78–2780 ng mL⁻¹. The analyte was interfered with other structural analogs such as BPA, DMP, DEP, and DOP, to evaluate the selectivity. The peak current of DPV (RSD <3.5%) verified that the presence of interferants had a negligible effect on the electrochemical sensing of DBP.¹⁶⁴ Wang *et al.*¹⁶⁵ constructed a MIP sensing platform based on silica nanospheres. Equipped with the same functional supports as Wu *et al.*,¹⁶⁴ this sensor exhibited an extremely wide detection range of 0.1–10 000 ng mL⁻¹ and a lower LOD of 5.09 pg mL⁻¹, which was because of the large surface area provided by nanospheres. The low efficiency of charge transfer upon silica was also claimed by the research group. The issue can be solved by adding more functional substances with good electrical conductivity on the electrode surface.¹⁶⁵ MIP sensors developed on the ITO substrate were used to detect DOP in water in combination with photoelectrochemistry. A cuprous oxide–copper organic framework heterostructure (Cu₃(BTC)₂@Cu₂O) and bismuth sulphide (Bi₂S₃) were applied by Gao *et al.*¹⁶⁶ and Li *et al.*,¹⁶⁷ respectively, to improve the photoelectrochemical responses. It is observed that the range of detection and LOD are mostly at ultralow concentration, suggesting again that photoelectrochemistry can achieve ultrasensitive determination for EDCs in water. Xia *et al.*¹⁷³ introduced titanium carbide MXene composites (Ti₃C₂T_x), a type of novel 2D-material, to prepare an MIP sensor for the detection of DES in water. Cuprous oxide nanoparticles (Cu₂O NPs) were adsorbed by the CTAB-CNT composite through the electrostatic effect and generated another characteristic peak in the DPV voltammogram, showing a proportional relationship with the peak of the analyte.¹⁷³

3.3.4 Electrochemical enzymatic and cyclodextrin sensors.

In Table 5, recent progress shows that enzyme-based biosensors are mostly used to detect BPs (*e.g.*, BPA) in water. Lou *et al.*¹⁷⁴ prepared a laccase-modified biosensor for BPA, where DPV was used to analyze the sensing results after different concentrations of BPA were introduced, offering a linear range of 1641.6–4104 ng mL⁻¹ and a LOD of 387 600 pg mL⁻¹. Furthermore, they also found that the pH of phosphate buffer saline (PBS) solution of 4.0 enables laccase to have the

highest electrocatalytic activity.¹⁷⁴ A similar enzymatic sensing platform was also reported by Fernandes *et al.*,¹⁷⁵ displaying a wider detection range of 6–228 ng mL⁻¹ and a two-order-of-magnitude reduced LOD of 4104 pg mL⁻¹. The obvious improvement in detection performance was attributed to the high activity of laccase preserved by the good biocompatibility and unique spatial structure of chitosan.¹⁸⁹ Moreover, laccase-based electrochemical biosensors can be also used for the determination of 4-NP in water. Liu *et al.*¹⁸⁰ evaluated the influence of temperature (*i.e.*, 5–40 °C) and pH (*i.e.*, 4.0–7.0) on laccase. The results revealed that the enzyme showed the best catalytic ability at room temperature of 25 °C and pH of 5.5. The detection threshold of 4-NP was 200 pg mL⁻¹ obtained under optimized conditions.¹⁸⁰ In fact, electrochemical biosensors functionalized with tyrosinase as the biorecognition center are more common than the ones functionalized with laccase. For the detection of BPA, Inroga *et al.*¹⁷⁶ developed an enzymatic biosensor where tyrosinase was deposited on the leaf-like gold microstructures. The results yielded by the biosensing device had a wider detection range of 114–11 400 ng mL⁻¹ in comparison to those obtained by the abovementioned biosensors equipped with laccase. However, limited progress was made on LOD with a value of 17 556 pg mL⁻¹.¹⁷⁵ Liu *et al.*¹⁷⁷ realized a LOD to an extent as low as 725.04 pg mL⁻¹ on the BCNP coated electrode using the amperometric technique. He *et al.*¹⁷⁸ invented a multi-layered enzymatic bioelectrode through the precipitation and cross-linking of tyrosinase, and acquired similar lower results (*i.e.*, LOD of 633.84 pg mL⁻¹). After the optimization, they learned that tyrosinase provided maximum electrochemical responses at 25 °C and pH of 7.0.¹⁷⁸ Zehani *et al.*¹⁷⁹ published an ultrasensitive tyrosinase-based biosensor for BPA, with a LOD of 2.28 pg mL⁻¹ (see Fig. 8). The detection in the ultralow range (*i.e.*, 0.00228–22.8 ng mL⁻¹) could be interpreted as the small capacitive background current generated on the diazonium-modified BDD electrode, evidencing high applicability for trace level monitoring.¹⁹⁰

Cyclodextrin-based (*i.e.*, β-CD) electrochemical sensors were also used to screen BPs,^{181–183} PCBs,¹⁸⁵ phthalate esters,^{186,187} and alkylphenols¹⁸⁸ in water sources through the adsorption of hydrophobic chambers. Ye *et al.*¹⁸⁴ applied a GCE modified with gold–silver–platinum ternary alloy nanocrystals (AuAgPt) to simultaneously recognize BPA and BPS using polymerized β-CD. The sensor provided LODs below 1000 pg mL⁻¹ for these two BPs, along with satisfactory findings in practical applications.¹⁸⁴ Su *et al.*¹⁸⁸ elaborated a sensor for 4-NP, where β-CDs were mixed with a sol–gel resulting in a stable and bioactive composite, which reached a LOD of 660 pg mL⁻¹. However, there are few parallel studies on the detection of a specific EDC. Also, most of the LODs reported in this class of sensors are not low enough (*i.e.*, thousands to tens of thousands pg mL⁻¹) compared with those obtained from the biosensors with antibodies, aptamers, and MIPs. These problems indicate that cyclodextrin-based sensors require more investigation to



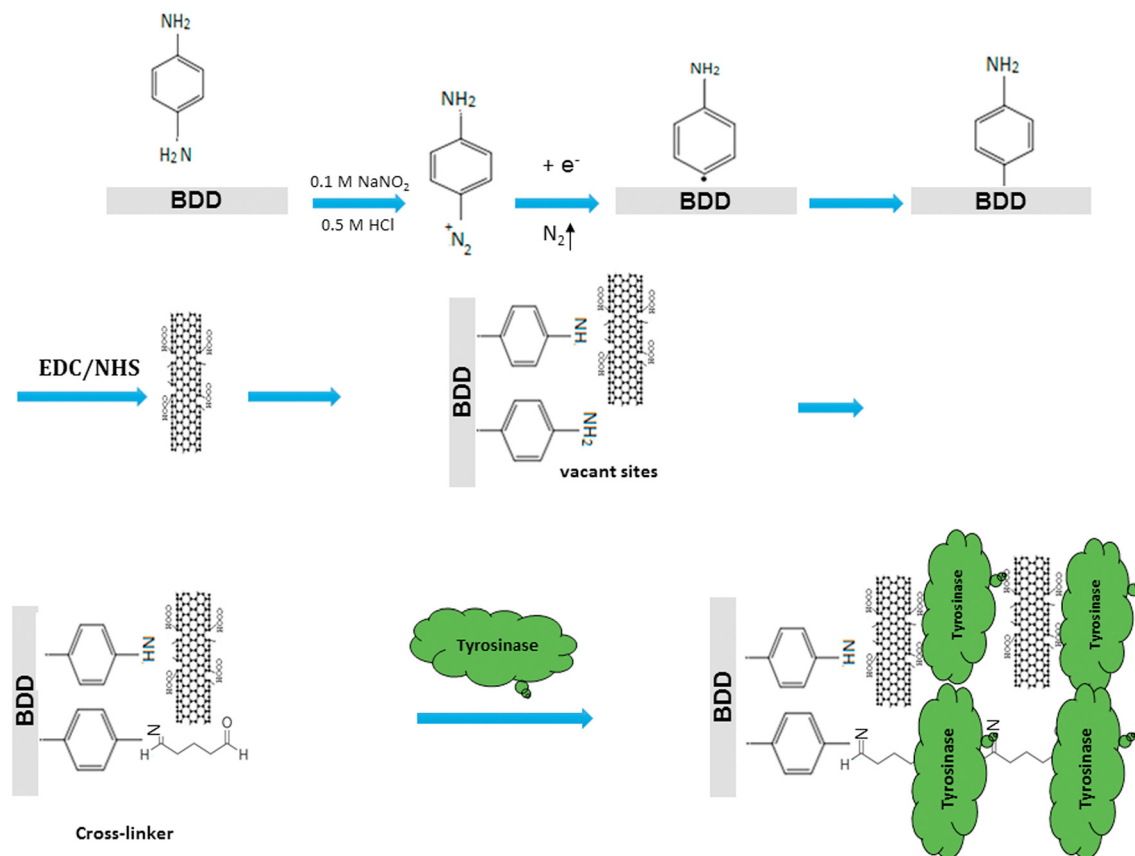


Fig. 8 Schematic illustration of the functionalizing procedure and enzyme attachment on the boron-doped diamond (BDD) surface (reprinted with the permission from ref. 178).

improve the electrochemical sensing performance. The stability of biological components is a major issue of biobased sensors. These challenges can be addressed using new class of nanozymes which will not only increase the shelf life but will also reduce the cost of sensing devices.

4. Current status and further development

Electrochemical biosensing of EDCs in water has been remarkably improved during the past decade. Research teams worldwide have made great efforts from the application of various biorecognition elements, electrode materials, and functional composites, to electrochemical techniques, to improve the ultra-sensitivity, high selectivity, long-term stability, and commercial availability of electrochemical biosensors.

Electrochemical immunosensors (*i.e.*, antibody-based biosensors) are often used to determine EDCs. Competitive immunosensing provides higher sensitivity and accuracy compared with direct antibody–antigen binding protocols. Commonly employed labels like enzymes and metallic nanoparticles are usually modified on the antigen competitor in direct competitive biosensing or on the antibody in the indirect competitive one, reflecting different analyte

concentrations. However, like sandwich-type biosensing, indirect competition requires more reagents and time-consuming preparation. Other widespread biodevices for EDCs are electrochemical aptasensors (*i.e.*, aptamer-based biosensors). Specific ssDNA or RNA sequences are simply immobilized on an electrode surface as usual or *via* binding to cDNA chains to form double-helical structures. Steric hindrance upon redox couples increases after more aptamer–analyte complexes are formed. Recently, a novel CNT induced ‘signal on’ aptasensor has been reported.¹⁴⁴ The aptamers dissociated from their cDNA chains when the analyte was introduced. Free ssDNA sequences preferred to encircle the target molecules while the complementary parts were subsequently connected with CNTs through π – π stacking, leading to amplified signals. In our opinion, this sensing method allows the analyte to be detected in a wider dynamic range coupled with a very low LOD, which is suitable for the ultrasensitive detection of EDCs in water. Electrodes functionalized with enzymes are also applied in electrochemical biosensing *via* electrocatalytic features. Apart from tyrosinase and laccase, Messaoud *et al.*¹⁹¹ fabricated a xanthine oxidase (XOD) based biosensor for the monitoring of BPA, with a LOD of 228 pg mL⁻¹ obtained using the enzyme inhibition principle. Other recognition elements like MIPs and β -CDs that exhibit strong affinity towards EDCs



have received tremendous attention. Tailored cavities are created after the polymerization of different monomers with template molecules, equipping MIP sensors with excellent reusability and stability. In addition, MIPs can be integrated with aptamers to produce super sensitive biosensors. Ensafi *et al.*¹²³ reported an aptasensor where the *p*-63 sequences were embedded in the BPA imprinted polymeric matrix. The resulting biosensor reached a broad detection range from 1.14×10^{-7} to 1.14×10^{-3} ng mL⁻¹ and an ultralow LOD of 1.824×10^{-5} pg mL⁻¹. Liang *et al.*¹⁹² provided an electrochemical sensor decorated with reduced graphene oxide (RGO), PtPd bimetallic nanoparticles (PtPd NPs), and cationic pillar [5]arene (CP5) used as the receptor for BPA. Like β -cyclodextrin, CP5 has a barrel-shaped structure with a pentagonal opening, presenting similar 'host-guest' adsorption towards target analytes. A five orders of magnitude linear range (*i.e.*, $2.28\text{--}2.28 \times 10^5$ ng mL⁻¹) and a detection limit of 752.4 pg mL⁻¹ were obtained after the DPV measurement. This device was finally utilized to successfully detect BPA in real water samples with good recoveries of 97.75–104.70%.¹⁸⁶

Electrodes are the essential part in electrochemical biosensors. Carbon (*e.g.*, carbon rod, glassy carbon, screen-printed carbon, and carbon paste) and gold (*e.g.*, Au substrate, screen-printed Au, and sputtered Au) are still the main materials for electrode fabrication. The limited electrochemical performance of bare electrodes can be significantly improved through various surface modifications. A wide range of functional compounds or composites are usually deposited on electrode surfaces *via* covalent binding, non-covalent adsorption, van der Waals force, electrostatic effects, and hydrogen bonding, to prepare micro-bioenvironments catering for different bioreceptors. In general, functional supports for electrodes have the following capacities: 1) increasing the surface area to accommodate more active binding sites, 2) improving the electrode conductivity and electrocatalytic activity, and 3) enhancing film forming and electrode stability. Moreover, the miniaturization of electrochemical biosensing systems (*i.e.*, electrode in combination with a data analyzer) is the direction for future development. Low cost, simple preparation, and portability are key factors in the commercialization of point-of-use electrochemical biosensors.

So far, a variety of electrochemical techniques have been used for biosensing of EDCs, including current varied with voltage (*e.g.*, LSV, DPV, and SWV), current varied with time (*e.g.*, amperometry and PEC), and impedance varied with frequency (*e.g.*, EIS). Some research groups believe that non-faradaic EIS (*i.e.*, capacitance as the typical response) is more promising than faradaic EIS (*i.e.*, charge transfer resistance as the typical response) because of the higher sensitivity and simpler preparation (*e.g.*, without the use of a redox couple), although faradaic EIS is still an active domain of investigation.¹³⁷ Cheng *et al.*¹⁰⁶ performed a capacitive aptasensing based on the ac electroosmotic (ACEO) effect.

Under an alternating electric field, the ACEO flows enriched target molecules relying on the charge transfer in the electrical double layer (EDL) at the electrode–solution interface. Hence, their work integrated the preconcentration of the analyte with determination, which simplified the operation procedures. Interfacial capacitance was measured at a certain frequency and transient ac voltage, yielding an ultralow LOD of 1.329×10^{-5} pg mL⁻¹ and a linear range of $2.28 \times 10^{-8}\text{--}2.28 \times 10^{-4}$ ng mL⁻¹.¹⁰⁶ Additionally, ultrasensitive detection of EDCs in water using a PEC technique has been demonstrated by many groups. An issue that is of concern is that some materials with a large band gap may not be appropriate for the fabrication of PEC electrodes as light with higher energy (*i.e.*, light with short wavelength) is often required to generate the photocurrent.¹⁹³

In conclusion, the recent development of electrochemical biosensing of EDCs in water sources has been reviewed. The sensing capabilities regarding the five types of biosensors (*i.e.*, immunosensors, aptasensors, MIP sensors, enzymatic sensors, and β -CD sensors) follow an order of aptasensors > immunosensors $\approx$ MIP sensors > enzymatic sensors $\approx$ β -CD sensors. Meanwhile, the recent advances on this field also expose some problems: 1) for immunosensing, only limited chemical variants in BPs, PCBs, phthalate esters, and estrogens are reported in the listed papers. More parallel research should be performed to increase the reliability of the result comparison. Examples for aptasensing enables comparable analyses on BPs (*i.e.*, BPA) and PCBs (*i.e.*, PCB 77). However, this group of biosensing exhibits the lowest detection diversity. In contrast, the detection diversity is improved in the cases of MIP and β -CD based electrochemical sensing, including BPS, TBBPS, PCB 101, DOP, DES, and alkylphenols (*e.g.*, 2-NP, 4-NP, and 4-*tert*-OP). However, the same issue as less comparable studies on each type of EDC should be noted. EDC determination achieved by enzymatic biosensors has only a few papers and is mostly limited to the detection of BPA. Lu *et al.*¹²⁵ used a copper-organic framework (CuMOF) modified enzymatic sensor to detect five BPs (*i.e.*, BPA, BPB, BPE, BPF, and BPZ), expanding the range of EDC detection, to some extent. Nonetheless, the LODs needed to be further improved. Moreover, some studies claimed that the specificity of the prepared electrochemical biosensors was not good enough to differentiate the target compound from its structural analogs, especially for PCBs¹³² and alkylphenols,¹⁰⁸ from which the detection result only reflected a total concentration of the mixture. Furthermore, other EDCs, like PBDEs, PFOA, DDT as well as a wider range of suspected EDCs have no report on water sample detection using electrochemical biosensors. There is still a long way to discover, study, and evaluate EDCs and their electrochemical screening methods. It is envisaged that more profound understanding, more mature detection techniques, and more standardized management of EDCs will offer stronger protection for human beings and natural environments.

Development of smart sensors for real-time monitoring of EDCs in environmental samples remains challenging due to



the complex nature of wastewater samples. Moreover, the increasing demand for electrochemical sensors will influence the cost of electrode materials. Combined efforts are needed from scientists of various fields such as analytical chemistry, bioelectrochemistry, and environmental engineers for designing reliable and cost-effective sensors for field applications. Nanozyme based sensors will open new gateways for compact and portable devices.

Abbreviations

2-ATP	2-Aminothiophenol
2D	Two-dimensional
2-NP	2-Nonylphenol
3D-GN	Three-dimensional graphene
4-ATP	4-Aminothiophenol
4-NP	4-Nonylphenol
4- <i>tert</i> -OP	4- <i>tert</i> -Octylphenol
11-MUA	Mercaptoundecanoic acid
Ab ₁	Primary antibody
Ab ₂	Secondary antibody
ABPE	Acetylene black paste electrodes
Abs	Antibodies
ACEO	AC electroosmotic
AIBN	Azodiisobutyronitrile
AgNPs	Silver nanoparticles
AhR	Aryl hydrocarbon receptor
APEOs	Alkylphenol ethoxylates
APTES	3-Aminopropyltriethoxysilane
ARs	Androgen receptors
AuAgPt	Gold–silver–platinum ternary alloy nanocrystals
AuNPs	Gold nanoparticles
AuNRs	Gold nanorod
AuPd–NH ₂ –fMnO ₂	AuPd decorated MnO ₂ nanoflowers
Au@Pd NRs	Gold/palladium nanorods
AuPt	Gold/platinum
BBP	Butyl benzyl phthalate
BCNPs	Biochar nanoparticles
BDD	Boron doped diamond
Bi ₂ S ₃	Bismuth sulphide
BnCl	Benzyl chloride
BPs	Bisphenols
BPA	Bisphenol A
BPAF	Bisphenol AF
BPAP	Bisphenol AP
BPB	Bisphenol B
BPE	Bisphenol E
BPF	Bisphenol F
BPS	Bisphenol S
BPZ	Bisphenol Z
BSA	Bovine serum albumin
cDNA	Complementary DNA
CE	Carbon electrode
Chit	Chitosan
CNFs	Carbon nanofibers

CNTs	Carbon nanotubes
Co _x P/NC	Cobalt phosphide nanoparticles embedded in three-dimensional carbon microspheres
CP	Capture probe
CP5	Cationic pillar [5]arene
CPE	Carbon paste electrode
CTAB	Cetyltrimethylammonium bromide
Cu ₃ (BTC) ₂ @Cu ₂ O	Cuprous oxide–copper organic framework heterostructure
CuFe ₂ O ₄ -Pr-SH	Thiol-functionalized copper magnetic nanoparticles
CuMOF	Copper-organic framework
Cu ₂ O NPs	Cuprous oxide nanoparticles
CV	Cyclic voltammetry
DAD1	10-Diaminodecane
DALY	Disability-adjusted life year
DBP	Dibutyl phthalate
DDT	Dichlorodiphenyltrichloroethane
DEHP	Di(2-ethylhexyl)phthalate
DEP	Diethyl phthalate
DES	Diethylstilbestrol
DIDP	Di-isodecyl phthalate
DINP	Di-isononyl phthalate
DMP	Dimethyl phthalates
DNA	Deoxyribonucleic acid
DNOP	Di(<i>n</i> -octyl)phthalate
DPV	Differential pulsed voltammetry
DOP	Diocetyl phthalate
E1	Estrone
E2	(17β-) estradiol
E3	Estriol
E4	Estetrol
EDC	Ethyl(dimethylamino propyl)carbodiimide
EDC/NHS	Ethyl(dimethylamino propyl)carbodiimide/ <i>N</i> -hydroxysuccinimide
EDCs	Endocrine disrupting chemicals
EDL	Electrical double layer
EDOT	3,4-Ethylenedioxythiophene
EDTA	Ethylenediaminetetraacetic acid
EE2	17α-Ethinylestradiol
EGDMA	Ethylene glycol dimethacrylate
EGFET	Extended gate field effect transistor
EIS	Electrochemical impedance spectroscopy
ELISA	Enzyme-linked immunosorbent assay
ERKs	Extracellular signal-regulated kinases
ERRγ	Estrogen related receptor gamma
ERs	Estrogen receptors
EU	European Union
Fc	Ferrocene
FET	Field-effect transistors
FTO	Fluorine-doped tin oxide
GA	Glutaraldehyde
GCE	Glassy carbon electrode
GC/MS	Gas chromatography coupled to mass spectrometry



GNP	Graphene nanoplatelets	pAbs	Polyclonal antibodies
GNRs	Graphene nanoribbons	PAEs	Phthalic acid esters
GO	Graphene oxide	PAH	Poly(allylamine hydrochloride)
GONRs/MWCNTs	Graphene oxide nanoribbons/multi-walled carbon nanotubes	PAHs	Polycyclic aromatic hydrocarbons
GQDs	Graphene quantum dots	PANI	Polyaniline
GR or Gr	Graphene	PBDEs	Polybrominated diphenyl ethers
Gr-AgNPs	graphene-silver nanoparticles	PBS	Phosphate buffer saline
GRs	Glucocorticoid receptors	PCBs	Polychlorinated biphenyls
H ₂ N-Fe ₃ O ₄ /Au NPs	Amino-functionalized magnetite and gold nanoparticles	PCD	β-Cyclodextrin polymer
HDSW	Heavily doped silicon wafer	PCDDs	Polychlorinated dibenzo dioxins
HPDNA	Hairpin DNA	PCDFs	Polychlorinated dibenzo furans
HPLC	High-performance liquid chromatography	PdQDs	Palladium quantum dots
HRP	Horseradish peroxidase	PEC	Photoelectrochemistry
IARC	International Agency for Research on Cancer	PEI	Polyethyleneimine
ILs	Ionic liquids	PET	Polyethylene terephthalate
ITO	Indium tin oxide	PFOA	Perfluorooctanoic acid
Lac	Laccase	PI	Polyimide
LC-MS	Liquid chromatography coupled to mass spectroscopy	PPAR _γ	Peroxisome proliferator-activated receptor γ
LOD	Limit of detection	PTMS	Phenyltrimethoxysilane
LOQ	Limit of quantification	PtNPs	Platinum nanoparticles
LSV	Linear sweep voltammetry	PtPd NPs	PtPd bimetallic nanoparticles
MAA	Methacrylic acid	PVC	Poly(vinyl chloride)
mAbs	Monoclonal antibodies	py-3-COOH	1 <i>H</i> -Pyrrole-3-carboxylic acid
MAC	Maximum acceptable concentration	QDs	Quantum dots
Mag	Magnetic	rGO or RGO	Reduced graphene oxide
MB	Methylene blue	RNA	Ribonucleic acid
MCH	6-Mercapto-1-hexanol	RSD	Relative standard deviation
MCL	Maximum contaminant level	SA	Streptavidin
MDH	Minnesota Department of Health	SAMs	Self-assembled monolayers
mERs	Membrane estrogen receptors	SDLSV	Second-order derivative linear sweep voltammetry
MIPs	Molecular imprinted polymers	SELEX	Systematic evolution of ligands by exponential enrichment
ML	Multi-layered	SERS	Surface enhanced Raman spectroscopy
MOF	Metal-organic framework	SPCE	Screen printed carbon electrode
MS	Mass spectroscopy	ssDNA	Single-stranded DNA
MW	Molecular weight	SWCNT	Single-walled carbon nanotube
MWCNTs	Multi-walled carbon nanotubes	SWV	Square wave voltammetry
NBD	Amine-terminated benzenediazonium	T-Au-NPs	Twice sputtered and twice annealed gold nanoparticles
NERRA	Non-equilibrium rapid replacement aptamer	TB	Toluidine blue
nERs	Nuclear estrogen receptors	TBBPA	Tetrabromobisphenol A
NGNRs	Nitrogen-doped graphene nanoribbons	TBBPS	Tetrabromobisphenol S
NiHCF NPs	Nickel hexacyanoferrate nanoparticles	TCBPA	Tetrachlorobisphenol A
NiNPs	Nickel nanoparticles	TdT	Terminal deoxynucleotidyl transferase
NIP	Non-imprinted polymer	TEMED	Tetramethylethylenediamine
NPEs	Nonylphenol ethoxylates	TEOS	Tetraethoxysilane
NPGL	Nanoporous gold leaf	Thi	Thionine
NPs	Nanoparticles or nonylphenols	Ti ₃ C ₂ T _x	Titanium carbide MXene composites
<i>o</i> -PD	<i>o</i> -Phenylenediamine	TiO ₂ -NH ₂	Amine-terminated TiO ₂
OPs	Octylphenols	TiO ₂ NRs	Titanium dioxide nanorods
P(1,5-DAN)	Poly(1,5-diaminonaphthalene)	TRs	Thyroid receptors
		Tyr	Tyrosinase
		USEPA	United States Environmental Protection Agency



USFDA	United States Food and Drug Administration
WFD	Water Framework Directive
WHO	World Health Organization
WWTPs	Wastewater treatment plants
XOD	Xanthine oxidase
ZEN	Zearalenone
Z _w	Warburg diffusion
αBPA	Monoclonal anti-BPA antibody
α ₂ BPA	Polyclonal anti-BPA antibody
β-CD	β-Cyclodextrin
β-CD-SH	Thiolated β-cyclodextrin

Conflicts of interest

There are no conflicts to declare.

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