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An electrochemiluminescence device for visualized detection of lead in practical samples†

Huaanzi Hu, D‡a Cheng Wang, D‡b Chen Qian,*c Chengqi Li,d Yulin Li,d Jianbin Pan*e and Xinye Ni*a

Lead (Pb²⁺) pollution poses a significant threat to human health due to its potential accumulation through the food chain. In response to this challenge, an array of electrochemiluminescence (ECL) devices has been developed for the accurate and visualized detection of trace Pb²⁺, achieving an ultra-low limit of detection (LOD) of 9.8 pg L⁻¹. The device utilizes a Pb²⁺-specific aptamer DNA chain, modified on gold nanoparticles (AuNPs), to create an efficient ECL probe. The integration of this ECL probe into an indium tin oxide (ITO) substrate results in a Pb²⁺ specific array device. With the assistance of an up-response ECL imaging system, this setup enables the accurate and visualized determination of trace Pb²⁺, not only in standard solution containing interference ions, but also in practical samples of *Lycium ruthenicum* Murr., *Glycyrrhiza uralensis* and lake water. This work advances the visual detection of Pb²⁺ using ECL-based technology, demonstrating significant potential for enhancing food safety.

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1 Introduction

Lead (Pb) contamination poses a significant threat to both the environment and human health.^{1,2} The contamination is primarily attributed to anthropogenic activities, such as mining, smelting, and the use of lead in various industrial products, including batteries and paints.³ This toxic metal, although naturally occurring in trace amounts in the Earth's crust, has experienced a substantial increase in global production due to the high demand for automobile and mobile phone batteries.⁴ The presence of Pb ions (Pb²⁺) in soil, water, and air has increased significantly, leading to widespread food contamination.⁵⁻⁷ The hazards of Pb²⁺ pollution are multifaceted, affecting both ecosystems and human health. In the environment, Pb²⁺ disrupts the natural balance by contaminating soil and water sources and biomagnifying through food

chains, which poses a significant risk to flora and fauna.^{8,9} For humans, Pb²⁺ contamination in foods can result in severe toxicity, posing significant health risks to the population. Children and developing fetuses are particularly vulnerable, as Pb²⁺ exposure can lead to impaired cognitive development and intellectual decline.¹⁰ It also has detrimental effects on the cardiovascular, renal, and immune systems.¹¹ Long-term exposure can lead to more severe health problems, including neurological disorders and cancer.¹² Therefore, there is an urgent need to develop a device that can afford convenient and sensitive detection of Pb²⁺.^{13,14}

As a quintessential heavy metal, lead (Pb2+) detection often relies on traditional methods such as Atomic Absorption Spectroscopy (AAS) and its advanced form, Electrothermal Atomic Absorption Spectroscopy (ET-AAS), which is known for their sensitivity and accuracy but require skilled technicians and can be time-consuming. Inductively Coupled Plasma Mass Spectrometry (ICP-MS) offers high sensitivity and a wide dynamic range, yet it is associated with higher costs and necessitates complex sample preparation. 15,16 In contrast, emerging nanomaterial-based sensors provide a promising alternative. These sensors leverage the high surface reactivity, large surface area, strong adsorption capacity, and high catalytic efficiency of nanomaterials to significantly enhance sensitivity and selectivity.17 In recent years, the introduction of nanotechnology for the detection of Pb2+ is gaining increasing attentions. 18,19 Among these approaches, electrochemiluminescence (ECL) detection stands out due to its low background interference, rapid response, high precision, and portability, which led to the widespread adoption of the ECL method for the precise quantification of lead ions.20-23 However, most of the current studies

[&]quot;Department of Radiotherapy, the Affiliated Changzhou No. 2 People's Hospital of Nanjing Medical University, Changzhou Medical Center, Nanjing Medical University, Changzhou 213003, P. R. China. E-mail: nxy@njmu.edu.cn

^bSchool of Pharmacy, Changzhou University, No. 21 Middle Gehu Road, Changzhou, 213164, P. R. China

Department of Orthopedics, Wujin Hospital Affiliated with Jiangsu University, Changzhou 213164, P. R. China. E-mail: qianchen88@126.com

^dState Key Laboratory of Radiation Medicine and Protection, School for Radiological and Interdisciplinary Sciences (RAD-X), Soochow University, Suzhou 215123, P. R. China

[&]quot;State Key Laboratory of Analytical Chemistry for Life Science, School of Chemistry and Chemical Engineering, Nanjing University, Nanjing 210023, P. R. China. E-mail: jbpan@nju.edu.cn

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 $[\]ddagger$ These authors contributed equally to this work.

fall short in providing a visualized method for the rapid and convenient assessment of Pb²⁺ levels in samples.

In response to this challenge, we have crafted a novel electrochemiluminescence (ECL) array system designed for the swift and clear identification of trace levels of Pb²⁺. This system features a remarkably low detection limit and superior selectivity, making it an advanced tool for identifying Pb2+ with high precision and reliability. It is particularly suitable for monitoring Pb2+ in practical samples. Specifically, we have functionalized gold nanoparticles (AuNPs) with a specific sequence of aptamer DNA, creating an efficient Pb²⁺-sensitive ECL probe designated as AuNPs@dsDNA. The -NH2 groups within this construct serve as a co-reactant in the ECL reaction. The AuNPs, acting as the core material of the probe, facilitate electron transfer during the ECL process due to their excellent electrical conductivity. Additionally, the Au-S bonds guarantee that the DNA sequences attached to the AuNPs' surface remain stable in aqueous environments. This design ensures that the ECL signal from the probe-modified electrodes is robust and reliable. The probe can be integrated into an array of indium tin oxide (ITO) electrode tunnels, thereby forming a high-performance ECL detection platform. Concurrently, tris(2,2'-bipyridyl)ruthenium(II) dichloride (Ru(bpy)₃Cl₂), when dissolved in a phosphate-buffered saline (PBS) solution, acts as an ECL luminophore. In practical applications, Pb²⁺ from the sample selectively displaces the NH2-ssDNA chains, thereby diminishing diminish the ECL signal from Ru(bpy)₃Cl₂. This interaction results in an exceptionally low limit of detection (LOD) for Pb²⁺ of 9.8 pg L^{-1} . To the best of our knowledge, this LOD represents one of the most sensitive detection limits reported for Pb²⁺ detection methodologies, as detailed in Table S1.† Moreover, the device is equipped with an ECL imaging system that enables the rapid and visual monitoring of Pb2+ at trace levels. In conclusion, the development of this ECL detector marks a significant advancement in the field of trace Pb²⁺ detection. It offers an efficient and precise method for the rapid assessment of Pb²⁺ concentrations, which is crucial for advancing related scientific research and holds substantial importance for ensuring food safety.

2 Experimental section

2.1 Preparation of array device for Pb2+ detection

A solution of AuNPs (10 μ L) at a concentration of 25 nM was mixed with a tris(carboxyethyl)phosphine (TCEP) solution (20 μ L) at 10 mM, along with a PBS solution (70 μ L) at 0.01 M, within a 10 mL centrifuge tube. This mixture was thoroughly homogenized. Subsequently, 100 μ L of a 12.5 μ M chain A solution was introduced into the tube. Upon the addition of 1800 μ L of n-butanol, the aqueous phase was carefully separated following a gentle centrifugation and adjusted to a final volume of 200 μ L, yielding a concentration of 6.25 μ M for the AuNPs@ssDNA solution. This resultant mixture was then applied to the array of ITO electrodes, with an aliquot of 10 μ L dispensed into each tunnel. Once the electrodes were air-dried, chain B solution at a concentration of 25 μ M was applied to the ITO electrodes, again using 10 μ L per tunnel. After the ITO

electrodes were fully dried, the assembly was completed, yielding an array of Pb²⁺ detection devices featuring the modified AuNPs@dsDNA.²⁴

2.2 Pb2+ detection and selectivity

A series of Pb2+ solutions with varying concentrations were applied to the array-based Pb²⁺ detection devices. After a 30 minute incubation period, the devices were rinsed with deionized water to remove any unbound species. For the preparation of the ECL reagent, 1 mL volume of a 10 mM Ru(bpy)₃Cl₂ solution was diluted to a final volume of 10 mL using a 0.1 M PBS solution (pH 7.4) within the measurement cell. The photomultiplier tube (PMT) was set to an operating voltage of 400 V, and the scanning rate for the potential was maintained at 100 mV s⁻¹. To assess the selectivity of the detection system, a range of potential interfering ions were tested,25 including potassium (K⁺), calcium (Ca²⁺), sodium (Na⁺), magnesium (Mg²⁺), strontium (Sr²⁺), mercury (Hg²⁺), iron (Fe³⁺), cadmium (Cd²⁺), cesium (Cs⁺), silver (Ag⁺), and copper (Cu²⁺). Each of these ions was present at a concentration of 100 $\mu g L^{-1}$, while the Pb $^{2+}$ concentration was set at 100 ng L $^{-1}$. The procedure for these selectivity tests was conducted in accordance with the established protocol for Pb2+ detection.

3 Results and discussion

3.1 Characterization on structure and ECL activity

The AuNPs, with an average size of 5 nm, were commercially available and further functionalized to specifically detect Pb²⁺ by conjugating them with a DNA aptamer sequence (referred to as chain A), following established protocols.^{26,27} The addition of TCEP facilitated the attachment of chain A to the AuNPs' surface. Successful functionalization was evidenced by a significant decrease in the zeta potential of the AuNPs@ssDNA compared to that of the pristine AuNPs, as depicted in Fig. 1A. Furthermore, the zeta potential of the AuNPs@dsDNA was notably lower than that of the AuNPs@ssDNA, confirming the formation of the duplex structure.

Cyclic voltammetry (CV) was employed to substantiate these modifications. The CV responses of glassy carbon electrodes modified with AuNPs, AuNPs@ssDNA, AuNPs@dsDNA were recorded in a PBS solution containing 1 mM Ru(bpy)₃Cl₂ (Fig. 1B). The oxidation peak for Ru(bpy)₃Cl₂ was observed at approximately +1.30 V and was significantly enhanced upon AuNPs modification, indicating improved electrode conductivity due to the AuNPs. This peak was further intensified after the addition of chain A, resulting in AuNPs@ssDNA. The negative zeta potential of ssDNA (Fig. 1A) likely promoted electrostatic interactions with Ru(bpy)₃Cl₂, leading to the observed increase of current (Fig. 1B). Supporting evidence was provided by the CV data for ethylenediamine (Fig. S1†), which showed an oxidation peak at +1.35 V, akin to that of Ru(bpy)3Cl2. The comparison between AuNPs@dsDNA

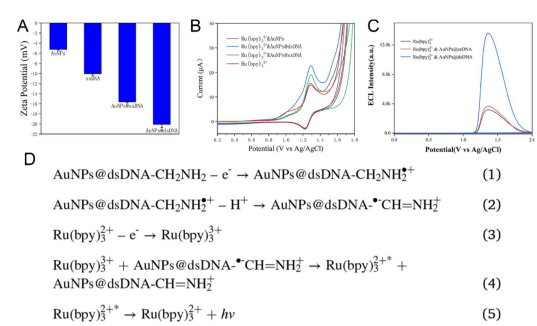


Fig. 1 (A) The zeta potential for AuNPs, chain A, AuNPs@ssDNA (AuNPs modified with chain A) and AuNPs@dsDNA (AuNPs@ssDNA further modified with chain B); (B) the CV data of GCE modified with Ru(bpy) $_3$ Cl $_2$ solution (blank), AuNPs, AuNPs@ssDNA and AuNPs@dsDNA. (C) The ECL data for ITO electrodes modified with Ru(bpy) $_3$ Cl $_2$ solution (blank), AuNPs, AuNPs@ssDNA and AuNPs@dsDNA. The CV and ECL measurements were conducted in 0.1 M PBS (pH = 7.4) solution with 1 mM Ru(bpy) $_3$ ²⁺. The PMT was set at 400 V with scan rate of 100 mV s⁻¹. (D) The proposed ECL mechanism for the device.

and AuNPs@ssDNA in Fig. S2 and S3† further corroborated the role of -NH₂ groups in enhancing the oxidation peak of AuNPs@dsDNA, and the increased DNA content and Ru(bpy)₃-Cl₂ intercalation into dsDNA, respectively.

The ECL emission peak of Ru(bpy)₃Cl₂ was detected at around +1.40 V, with an onset potential of +1.08 V (Fig. 1C). This signal was notably enhanced upon the modification with AuNPs@ssDNA (Fig. 1C), potentially due to the increased current observed in Fig. 1B. A marked enhancement in the ECL signal was observed upon the addition of NH₂-ssDNA (chain B) to AuNPs@ssDNA, highlighting the reactivity of ¬NH₂ groups (Fig. 1C). The ECL peak at +1.40 V mirrored the oxidation peak of Ru(bpy)₃Cl₂, suggesting that the ECL signal is a result of the oxidation of Ru(bpy)₃Cl₂. The proposed ECL mechanism is illustrated in Fig. 1D.

3.2 Detection mechanism of Pb²⁺

ITO electrodes offer distinct benefits over conventional GCEs, including ease of fabrication, integration potential, and superior light transmission capabilities for ECL imaging assessments. These attributes make ITO electrodes a promising platform for the development of visual detection devices for Pb²⁺. To facilitate rapid and visual Pb²⁺ monitoring, an array of Pb²⁺ detection devices was fabricated by immobilizing AuNPs@dsDNA within the arrayed tunnel structures of the ITO electrode (as depicted in Fig. 2A). The ECL responses from these devices, upon exposure to Pb²⁺ solutions of varying concentrations, were recorded in a 0.1 M PBS solution containing 1 mM Ru(bpy) $_3$ Cl $_2$ (Fig. 2B). A progressive decrease in ECL signals was observed as Pb²⁺ concentrations rose from 0 to 10 μ g L⁻¹. The ECL intensity exhibited a strong linear relationship with the

logarithmic concentration of Pb²⁺, yielding a low limit of detection (LOD) for Pb²⁺ at 9.8 pg L⁻¹ (Fig. 2C). The observed reduction in ECL signal is attributed to the displacement of NH₂-ssDNA by Pb²⁺. To elucidate this mechanism, the CV of the Pb²⁺ detection devices was measured before and after exposure to Pb²⁺ solutions in a 0.1 M PBS solution containing 1 mM Ru(bpy)₃Cl₂ (Fig. 2D). A noticeable decrease in the oxidation peak current of the AuNPs@dsDNA-modified GCEs was observed post-Pb²⁺ treatment, aligning with the current levels of AuNPs@ssDNA (as indicated by the blue line in Fig. 2D). This finding provides corroborative evidence for the proposed displacement mechanism.

To assess the selectivity of the probe, a panel of common interfering ions were tested at concentrations 1000-fold higher than that of Pb^{2+} (Fig. 2E). The selection encompassed both ubiquitous environmental cations and representative heavy metal ions, given Pb^{2+} is classified as a heavy metal. The Pb^{2+} detection devices demonstrated exceptional selectivity for Pb^{2+} , underscoring their reliability and potential for practical Pb^{2+} detection applications.

Moreover, to prove its storage stability, we have conducted a comparative experiment on two batches of device, one is freshly prepared device while another is old device stored for one month under room temperature. As shown in Fig. S4,† we can see that both batches of device maintain similar ECL response to the same sample, suggesting the high storage stability of the device.

3.3 Visualized detection of Pb²⁺

Additionally, the incorporation of ECL imaging technology has been employed to facilitate the rapid and visual monitoring of

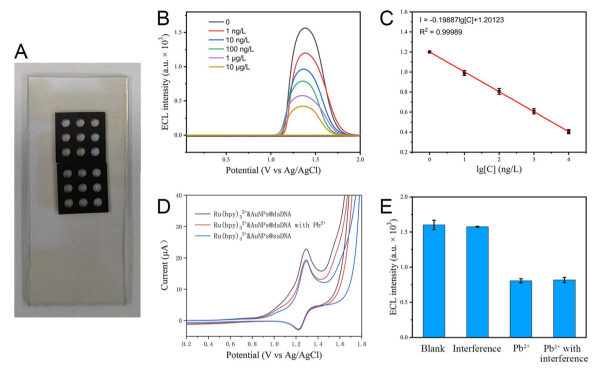


Fig. 2 (A) A visual representation of the ITO electrodes modified with AuNPs@dsDNA, which serve as part of an array device designed for the visual detection of Pb2+; (B) the ECL signals of array devices treated by various standard Pb2+ concentrations. (C) The calibration curve between the ECL intensity and the concentration (expressed as logarithm value) of Pb2+; (D) the CV of GCE modified with AuNPs@ssDNA and AuNPs@dsDNA treated with or without 100 ng L^{-1} of Pb²⁺; (\bar{E}) assessment of resistance to interference of the Pb²⁺ detection array device. The CV and ECL measurements were conducted in 0.1 M PBS (pH = 7.4) solution with 1 mM Ru(bpy)₃²⁺. The PMT was set at 400 V with scan rate of 100 mV s⁻¹. For the interference assay, the Pb²⁺ concentration was 100 ng L⁻¹ while the interference ions were fixed at an equal concentration of 100 μ g L⁻¹.

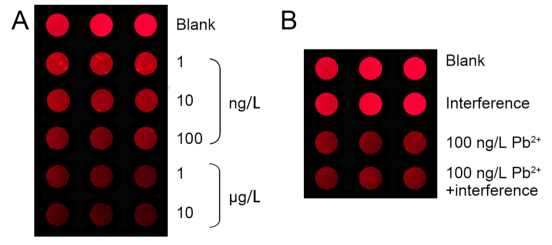


Fig. 3 The visualization of Pb²⁺ by the device. (A) The device response upon treatment with Pb²⁺ solutions across varying concentrations, illustrating dose-dependent ECL intensity changes; (B) the selectivity assessment in the presence of different designated interfering ions, confirming minimal cross-reactivity. All scans were conducted at a rate of 400 mV s⁻¹.

Pb²⁺ in practical samples, as depicted in Fig. 3. The array of Pb²⁺ detection devices exhibit vivid red ECL emissions, which are indicative of the luminescence generated by Ru(bpy)₃Cl₂. A discernible reduction in ECL intensity is observed as the Pb2+ concentration escalates from 1 ng L^{-1} to 10 μ g L^{-1} , paralleling

the trend previously illustrated in Fig. 2B and C. Concurrently, the findings in Fig. 3B corroborate the high selectivity of this ECL-based device for Pb2+ detection. These outcomes underscore the device's efficacy and potential in providing precise and visual Pb²⁺ monitoring in food samples.

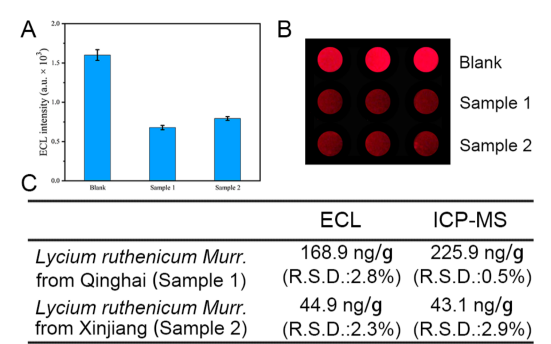


Fig. 4 The detection performance of the device on practical food samples. (A) The ECL signal of the device in response to the different practical samples; (B) ECL images of the device in response different practical samples; (C) the summary of Pb²⁺ concentration in different practical samples determined by our device and ICP-MS.

3.4 The detection performance of the device on practical samples

Lycium ruthenicum Murr., recognized for its rich content of anthocyanins, which provides exceptional antioxidant capabilities, is crucial for developing functional foods and beverages that promote health. Its use in traditional medicine and ongoing research into its phytochemicals further enhance its role in creating innovative, health-beneficial food products.²⁸ Consequently, the development of a swift, quantitative, and visual analytical method for detecting Pb²⁺ contamination in *Lycium ruthenicum* Murr. is of paramount importance to ensure its safety for food and medicinal application. To verify the practicability of the Pb²⁺ detection device, we applied it to the quantitative analysis of Pb²⁺ in *Lycium ruthenicum* Murr. from

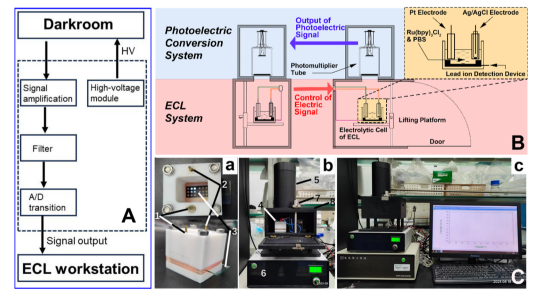


Fig. 5 (A) The schematic structure of the controller in ECL detection system. (B) The schematic structure of the newly designed ECL detector. (C) The practical picture and corresponding illustration of the ECL detector (a) the measuring pool for ITO electrode based Pb²⁺ detection devices: (1) counter electrode (Pt), (2) reference electrode (Ag/AgCl), (3) working electrode (ITO electrode based Pb²⁺ detection device); (b) newly designed ECL detector: (4) measuring pool, (5) PMT, (6) controller, (7) lock, (8) darkroom; (c) the practical application of this instrument in ECL detection using ITO electrodes.

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Qinghai (sample 1) and Xinjiang (sample 2) respectively. It was found that the ECL signal was significantly quenched in both groups of samples (Fig. 4A and B). Quantitative analysis of Ph²⁺ However, the preparation of the ECL detection system.

found that the ECL signal was significantly quenched in both groups of samples (Fig. 4A and B). Quantitative analysis of Pb²⁺ content in Lycium ruthenicum Murr. showed that the contents of sample 1 and sample 2 were 168.9 ng g⁻¹ and 44.9 ng g⁻¹, respectively. In comparison with the ICP-MS method (Fig. 4C), the concentration of Pb2+ in sample 1 and sample 2 were 225.9 ng g⁻¹ and 43.1 ng g⁻¹, respectively, which is consistent with the results measured by our DNA-based biosensor. To further confirm the detection capability of our device, we also introduced two more practical samples, Glycyrrhiza uralensis from Inner Mongolia and lake water from Dushu lake, as representative food and environmental samples. As shown in Fig. S5,† our device can give comparable results to ICP-MS at higher Pb2+ condition while remain sensitive under lower Pb2+ condition where ICP-MS is unapplicable. These results demonstrate that our device can accurately detect trace amounts of Pb2+ in real samples. Therefore, it is of special significance that this device can effectively identify potential trace Pb2+ contamination in practical objects at an early stage, thereby enhancing human health.

In alignment with the practical application requirements for Pb²⁺ detection, we have engineered a corresponding ECL detection apparatus, which encompasses a control module (as shown in Fig. 5A) and an up-response ECL detection component (illustrated in Fig. 5B). The physical images of the system are displayed in Fig. 5C, with photographs (a) and (b) showcasing the device's distinct parts. Integrated with an electrochemical workstation, this ECL detection setup is capable of delivering consistent CV and ECL signals when used with ITO electrodes, as depicted in picture (c) of Fig. 5C. This newly crafted system has been effectively utilized for Pb²⁺ detection in the current study. The array-based Pb²⁺ detection device, which emits ECL signals in an upward direction, is particularly compatible with the up-response ECL detector, offering a distinct advantage over GCE setups.

4 Conclusions

In conclusion, this study presents a groundbreaking array ECL device for the ultra-sensitive and selective detection of Pb2+ in practical samples, addressing a critical need in food safety. By leveraging the unique properties of AuNPs modified with a Pb²⁺specific aptamer DNA chain, the device achieves an unprecedented low LOD of 9.8 pg L⁻¹, significantly outperforming existing technologies. The innovative integration of this ECL probe into an ITO electrode array allows for rapid and visualized detection, with high selectivity and short measurement times, making it suitable for on-site monitoring of trace Pb²⁺. The successful application of the device in multiple practical samples, demonstrates its practicality and potential for early identification of Pb²⁺ contamination, thereby enhancing food safety. Moreover, the newly designed ECL detection system, with its controller and up-response ECL detector, ensures stable and efficient operation, showcasing the device's readiness for field applications. This work not only advances the field of trace heavy metal ECL detection but also provides a vital tool for

However, the preparation of the ECL detection system involves multiple steps, including the functionalization of AuNPs with aptamer DNA and the assembly of the ITO electrode array. This complexity may pose challenges in scaling up. Therefore, in our future work, we will focus on simplifying the preparation procedure to improve its reproducibility and reliability for potential widespread use or commercial applications. Moreover, the ECL detection system requires a custom-designed apparatus, including an electrochemical workstation and a specialized ECL detector. Therefore, our future work focus also includes miniaturizing and marketization of this detection equipment to improve its accessibility for laboratories or field applications.

Data availability

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

Conflicts of interest

There are no conflicts to declare.

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