


 Cite this: *RSC Adv.*, 2024, 14, 8303

Effective extraction and determination of 24 quinolones in water and egg samples using a novel magnetic covalent organic framework combined with UPLC-MS/MS†

 Na Li,^a Mengnan Liang,^a Hao Zhang,^a Zhongxia Hua,^{bc} Ling Ma,^{*bc} Yanyu Qi ^{*a} and Ke Wang ^{*abc}

The excessive use of quinolones (QNs) has seriously threatened human health. In this study, a novel functionalized magnetic covalent organic framework Fe₃O₄@SiO₂@Ah-COF was fabricated with biphenyl-3,3',5,5'-tetracarbaldehyde and hydrazine hydrate (85%) as monomers and was used as a magnetic solid-phase extraction (MSPE) adsorbent for the determination of 24 QNs in water and egg samples through ultra-performance liquid chromatography-tandem mass spectrometry (UPLC-MS/MS). The extraction parameters of MSPE were optimized, including pH, adsorbent dosage, adsorption time, and eluent type. An effective and rapid detection method was then established, which showed good linearity ($R^2 \geq 0.9990$), low limits of detection (0.003–0.036 $\mu\text{g L}^{-1}$) and low limits of quantitation (0.008–0.110 $\mu\text{g L}^{-1}$) for QNs. The good recoveries of 24 QNs in water and egg samples were in the range of 70.3–106.1% and 70.4–119.7%, respectively, with relative standard deviations lower than 10% ($n = 5$). As a result, Fe₃O₄@SiO₂@Ah-COF is a promising magnetic adsorbent, and the established method was successfully applied for the determination of 24 QNs in water and egg samples.

 Received 10th January 2024
 Accepted 23rd February 2024

DOI: 10.1039/d4ra00247d

rsc.li/rsc-advances

1. Introduction

Quinolones (QNs) are an important and successful group of synthetic antibiotics with a 4-quinolone structure, which have been developed to the fourth generation.¹ The fourth generation of quinolones have better medicinal properties and stronger antibacterial ability; thus, they are widely used in human health care and animal disease treatment.² Their mechanism of action is that the presence of an F atom and piperazinyl group prevents the synthesis of bacterial DNA.³ On the one hand, as the consumption of QNs increases, their residues in foods of animal origin can have a variety of toxic side effects on the consumers through the migration of the food chain, seriously threatening human health.⁴ On the other hand, due to the low solubility, bioavailability and biodegradation rates,⁵ QNs are difficult to be fully absorbed by humans and animals. Most of them are discharged into the environment in their original form or as active metabolites through feces and urine, resulting in

water and soil pollution. In recent years, some reports have indicated that QNs exist at high levels in aquatic environments, interfering with microbial driven nitrogen cycling processes and becoming an important factor affecting water eutrophication and greenhouse gas emissions.⁶ Given the significant residual hazards of QNs, many countries and organizations (including the United States, the European Union and China) have set corresponding maximum residue limits (MRLs) for QNs. However, MRLs for QNs in drinking water have not been specified in GB 5749-2022, and a unified standard detection method for QNs in water quality has not yet been established. Therefore, it is of great significance to develop sensitive, reliable and high-throughput analytical methods to detect the concentration of QNs in food and water.

To date, a variety of detection methods have been applied for the analysis of QNs, such as immunoassays,⁷ fluorescence-based methods,⁸ capillary electrophoresis (CE),⁹ high performance liquid chromatography (HPLC),¹⁰ and liquid chromatography-tandem mass spectrometry (LC-MS/MS).^{11,12} Among them, ultra-high performance liquid chromatography-tandem mass spectrometry (UPLC-MS/MS)¹² has been gradually developed owing to its advantages of high sensitivity, specificity and simultaneous qualitative and quantitative analysis.

Nevertheless, the complicated matrix and extremely low concentration of QNs in food and water samples can influence

^aCollege of Chemistry and Materials Science, Hebei Normal University, Shijiazhuang 050023, China. E-mail: wkecdc@163.com; hbsdqyy@hebtu.edu.cn

^bShijiazhuang Center for Disease Control and Prevention, Shijiazhuang 050011, China. E-mail: mamalin001@163.com

^cShijiazhuang Technology Innovation Center for Chemical Poison Detection and Risk Early Warning, Shijiazhuang 050011, China

† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d4ra00247d>



detection results; therefore, an effective sample pretreatment is indispensable. Common sample pretreatment methods include liquid–liquid extraction (LLE),¹³ dispersive liquid–liquid micro-extraction (DLLME),¹⁴ solid phase extraction (SPE),^{15–17} dispersive solid-phase extraction (DSPE)^{18,19} and magnetic solid phase extraction (MSPE).^{20–22} Different from conventional sample pretreatment methods, MSPE is a dispersed solid-phase extraction technique that uses magnetic materials as adsorbents to achieve the separation and enrichment of analytes by an external magnetic field. It avoids the disadvantages of cumbersome operation and high consumption of organic solvents and has been applied in the pretreatment of environmental, biological and food samples.^{23,24}

The adsorbent is clearly the most important influential factor on MSPE. Commonly used adsorbents include nanotubes, graphene,^{25,26} molecularly imprinted polymers,²⁷ and metal–organic frameworks.²⁸ Although the above adsorbents have their characteristics, the defects of large amounts of materials, relatively lower adsorption capacity and instability limited their extensive applications as MSPE adsorbents. Relatively speaking, covalent organic frameworks (COFs) are a new type of porous polymer, which have a unique topological structure and functionalized modification characteristics. COFs have high-affinity interactions with the target compounds, high specific surface area and pore volume, and have been used in adsorption, separation, catalysis, sensing and other fields.^{29–32} Notably, COFs have shown great potential as adsorbents for the extraction and removal of various substances such as pharmaceutical contaminants,³³ glycopeptides,³⁴ and environmental contaminants.³⁵ At present, according to the types of covalent bonds formed, COFs are mainly divided into boron-based COFs, imine-based COFs, and triazine-based COFs.³⁶ Among them, the stability of boron-based COFs in water and moisture is poor. Furthermore, the preparation conditions of triazine-based COFs are harsh, the synthesis is difficult, and the crystallinity of the products is poor. Only imine-linked COFs have simple synthesis conditions, diverse synthesis methods and good chemical stability, and can be stably present in common organic solvents, water, and even acid and alkali solutions.³⁷ However, it is difficult to precipitate COFs with low density from the complex matrix, and the adsorption efficiency is low.³⁸ Fascinatingly, the combination of COFs and Fe₃O₄ perfectly makes up for this shortcoming well, making magnetic covalent organic frameworks (MCOFs) that not only have rich functional groups but also the merits of easy separation, improving the enrichment capacity and adsorption capacity of the target analyte.^{39,40}

To the best of our knowledge, there are few reports on the application of MCOF materials in the field of adsorbing QNs. In the reports on the application of the MSPE method to determine QNs,^{23,41–44} there are few reports on methods that can simultaneously detect QNs in different matrices, and the number of QNs is no more than 14. Therefore, developing a unified method capable of simultaneously detecting large amounts of QNs residues in different matrices remains a challenge. Moreover, the reusability and narrow spectrum adsorption of magnetic materials remains to be explored and improved in further development.

Herein, we first designed and fabricated a novel MCOFs (Fe₃O₄@SiO₂@Ah-COF) based on the properties of QNs *via* a simple reaction of biphenyl-3,3',5,5'-tetracarbaldehyde and hydrazine hydrate (85%). Considering the biphenyl skeleton and unsaturated bonds of Fe₃O₄@SiO₂@Ah-COF, and that QNs contain the carboxyl groups and amino group on the piperazine ring side chain,⁴⁵ the great adsorption effect was obtained through the synergistic effects of p–π interaction, π–π interaction and hydrogen bonding. Then, Fe₃O₄@SiO₂@Ah-COF was characterized and the optimal parameters affecting the extraction efficiency were investigated. Finally, the developed MSPE-UPLC-MS/MS method was applied for the determination of 24 QNs in water and egg samples.

2. Experimental

2.1 Reagents and materials

Ferric chloride hexahydrate (FeCl₃·6H₂O), ethylene glycol (EG), sodium acetate trihydrate (CH₃COONa·3H₂O), 1,4-dioxane, sodium chloride (NaCl) and anhydrous ethanol were purchased from Tianjin Yongda Chemical Reagent Co., Ltd (Tianjin, China). Biphenyl-3,3',5,5'-tetracarbaldehyde was purchased from Shanghai Macklin Biochemical Co., Ltd (Shanghai, China). Hydrazine hydrate (85%) was purchased from National Pharmaceutical Group Chemical Reagent Co., Ltd (Shanghai, China). Tetraethoxysilane (TEOS) was obtained from Aladdin Chemistry Co., Ltd (Shanghai, China). Polyethylene glycol (PEG-4000) was supplied by Xiya Chemical Technology Co., Ltd (Shandong, China). 25% (w/w) ammonia solution and 37% (w/w) hydrochloric acid were acquired from Tianjin Kernel Chemical Reagent Co., Ltd (Tianjin, China). 99.9% (w/w) formic acid was acquired from Dikma (USA). Acetonitrile was obtained from Merck (Darmstadt, Germany), and methanol was purchased from Thermo Fisher Co., Ltd (USA). Methanol and acetonitrile were of HPLC grade, and other reagents were of analytical grade. Ultra-pure water was prepared using a Milli-pore Milli-Q Gradient Water Purification System (USA).

Lomefloxacin (LOM) was purchased from Dr Ehrenstorfer, GmbH (Augsburg, Germany), and all other reagents were purchased from Tianjin Alta Technology Co., Ltd. The purity of Tosufloxacin (TOS) is 70.6%, and those of other standards are all above 95.0%. The properties of QNs are listed in Table S1.†

2.2 Instrumental and analytical conditions

The AB SCIEX Exion-TRIPLE QUAD 5500 ultra-high performance liquid chromatography-tandem mass spectrometry with electrospray ionization source (ESI) was used for the detection.

A Waters Acquity UPLC BEH C₁₈ column (2.1 mm × 100 mm, 1.7 μm) was used for the separation of QNs at the constant column temperature of 40 °C. The mobile phases were composed of acetonitrile (mobile phase B) and 0.1% formic acid aqueous solution (mobile phase A). The gradient elution procedure was as follows: (1) 0–10 min, 13% B (v/v); (2) 10–10.1 min, 13–18% B (v/v); (3) 10.1–16 min, 18% B (v/v); (4) 16–16.1 min, 18–85% B (v/v); (5) 16.1–18 min, 85% B (v/v); (6) 18–18.1 min, 85–13% B (v/v); (7) 18.1–20 min, 13% B



(v/v). The flow rate and injection volume were 0.2 mL min^{-1} and $3 \mu\text{L}$, respectively. Chromatograms of 24 QNs are shown in Fig. S3.†

Electrospray ionization (ESI) in positive ion mode was used, and data were acquired in the multiple reaction monitoring (MRM) mode. The flow rate of the drying gas was 10 L min^{-1} at the temperature of $550 \text{ }^\circ\text{C}$. The capillary voltage was 5500 V . The MRM parameters comprising the precursor ion, production ion, retention time, collision energy (CE), declustering potential (DP) of 24 QNs are optimized, and the detailed information is listed in Table S2.†

Fourier transform infrared spectroscopy (FT-IR) (Thermo Nicolet Co., Ltd, USA), Brunauer–Emmett–Teller (BET) (Quantachrome Instruments, USA), D8A X-ray diffractometer (XRD) (Bruker, Germany), SQUID XL-7 vibrating sample magnetometer (Quantum Design, USA), scanning electron microscopy (SEM) with a Regulus 8100 instrument (Hitachi, Japan), transmission electron microscopy (TEM) with a FEI Tecnai G2 F20 (FEI, USA) and energy-dispersive X-ray spectroscopy (EDS) (EDAX, USA) were used for the characterization of the prepared materials.

2.3 Synthesis of $\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{Ah-COF}$

The magnetic Fe_3O_4 nanoparticles were synthesized by the solvent thermal method.⁴⁶ Typically, $1.62 \text{ g FeCl}_3 \cdot 6\text{H}_2\text{O}$ was dissolved in 60 mL of ethylene glycol (EG) under vigorous stirring at room temperature. Then, 7.2 g of $\text{CH}_3\text{COONa} \cdot 3\text{H}_2\text{O}$ and 2.0 g of PEG-4000 were added to the solution with stirring for 30 min . Subsequently, the obtained homogeneous yellow solution was transferred into a Teflon-lined autoclave and heated to $200 \text{ }^\circ\text{C}$ for 8 h . The resultant nanoparticles were washed with ethanol three times, and then dried at $60 \text{ }^\circ\text{C}$ for 4 h in a vacuum oven.

$\text{Fe}_3\text{O}_4@\text{SiO}_2$ nanoparticles were synthesized by the sol-gel method.^{47,48} The Fe_3O_4 nanoparticles (200 mg) were dispersed in a mixed solution of 160 mL of anhydrous ethanol and 40 mL of ultrapure water. The mixture was sonicated for 30 min to ensure that it was evenly dispersed. After that, 3 mL of 25% (w/w) ammonia was added to the mixture and sonicated for 5 min . Finally, 2 mL of tetraethoxysilane (TEOS) was slowly added dropwise under magnetic stirring. The mixture was allowed to react for 24 h at $30 \text{ }^\circ\text{C}$ to get $\text{Fe}_3\text{O}_4@\text{SiO}_2$, washed three times with anhydrous ethanol and ultrapure water, and dried under vacuum at $60 \text{ }^\circ\text{C}$ for 3 h .

$\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{Ah-COF}$ was prepared based on the modified literature method.⁴⁹ A total quantity of 150 mg of $\text{Fe}_3\text{O}_4@\text{SiO}_2$ was added to a 100 mL round-bottom flask, along with 35 mL of 1,4-dioxane solvent. The mixture was ultrasonicated for about 10 min , and then placed into an oil bath agitator. Next, biphenyl-3,3',5,5'-tetracarbaldehyde (80 mg , 0.30 mmol) was dissolved in 5 mL of dimethyl sulfoxide solution, and dispersed evenly by ultrasonication. Then, hydrazine hydrate (85% , 0.80 mmol) and pure glacial acetic acid solution (1.10 mL) were separately added. The mixture was stirred for 1 h at room temperature, and then heated at $70 \text{ }^\circ\text{C}$ for 24 h . After cooling to room temperature, the blue-gray solid was isolated with

a magnet, and washed thoroughly with 1,4-dioxane solvent and anhydrous ethanol alternately three times to remove unreacted chemicals. The final product was dried under vacuum at $60 \text{ }^\circ\text{C}$ for 2.5 h to get $\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{Ah-COF}$.

2.4 Sample collection and pretreatment

Twenty water samples were collected from relevant rivers and filtered by $0.45 \mu\text{m}$ filter membrane. The pH of the water sample was adjusted to 4.0 with hydrochloric acid and stored in a refrigerator at $4 \text{ }^\circ\text{C}$ for the MSPE procedure.

Fifteen egg samples were obtained from the market. Then, 0.50 g of the homogenized sample was transferred to a 50 mL polypropylene tube and diluted with 10 mL ultra-pure water. The mixture was then centrifuged at $10\,000 \text{ rpm}$ for 10 min . All supernatants were collected, and the pH of sample solution was adjusted to 4.0 by adding hydrochloric acid for MSPE.

2.5 Procedure of MSPE

The MSPE program is shown in Scheme 1. Firstly, 10 mL of the sample solution and 10 mg of $\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{Ah-COF}$ were added to a 50 mL centrifuge tube. Sample adsorption was performed by using the vortex method for 20 min . After that, the magnetic material was collected under the action of the external magnetic field, and the supernatant was discarded. QNs were eluted from $\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{Ah-COF}$ using 8 mL of 3% (v/v) acidified methanol under vibration for 8 min . The eluent was separated with a magnet and then dried at $40 \text{ }^\circ\text{C}$ under N_2 . Finally, the residue was redissolved using 1 mL of mixed solution of acetonitrile and 0.1% formic acid solution ($2:8$, v/v), filtered through a $0.20 \mu\text{m}$ membrane, and introduced into UPLC-MS/MS for analysis. In addition, the $\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{Ah-COF}$ was washed three times with 3% acidified methanol and recycled after vacuum drying at $60 \text{ }^\circ\text{C}$.

3. Result and discussion

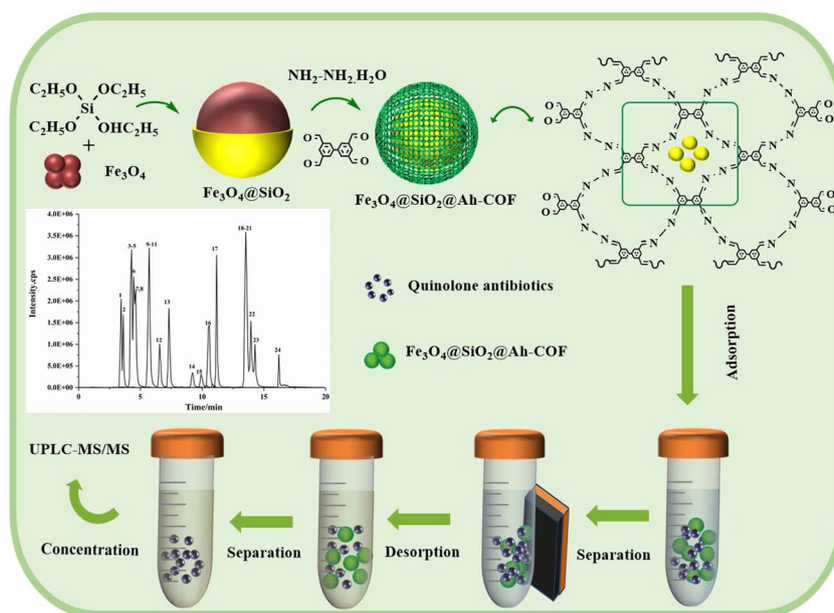
3.1 Characterization of the $\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{Ah-COF}$

The SEM and TEM images show the structural characteristics of the Fe_3O_4 nanoparticles and $\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{Ah-COF}$ nanoparticles. As shown in Fig. 1A and B, it can be easily found that Fe_3O_4 had a smooth and well-dispersed surface, while $\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{Ah-COF}$ had a rough surface. The typical Fe_3O_4 nanoparticle nuclei and the shell structure of the gray COF layer can be further clearly seen in Fig. 1C and D (TEM images).

Fig. 2 shows the energy-dispersive X-ray spectrum (EDS) of $\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{Ah-COF}$. As shown in the figure, the five elements (C, N, O, Si, and Fe) are distributed on $\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{Ah-COF}$, with contents of $59.17 \text{ wt}\%$, $6.86 \text{ wt}\%$, $17.66 \text{ wt}\%$, $2.97 \text{ wt}\%$ and $13.34 \text{ wt}\%$, respectively. These obvious differences all indicated the successful synthesis of the COF layer on the surface of Fe_3O_4 .

To further prove the successful synthesis of $\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{Ah-COF}$, the crystal morphology and characteristic spectra of the Fe_3O_4 , $\text{Fe}_3\text{O}_4@\text{SiO}_2$ and $\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{Ah-COF}$ samples were determined through XRD and FT-IR.





Scheme 1 The synthetic route for the $\text{Fe}_3\text{O}_4@SiO_2@Ah-COF$ and the procedure of MSPE.

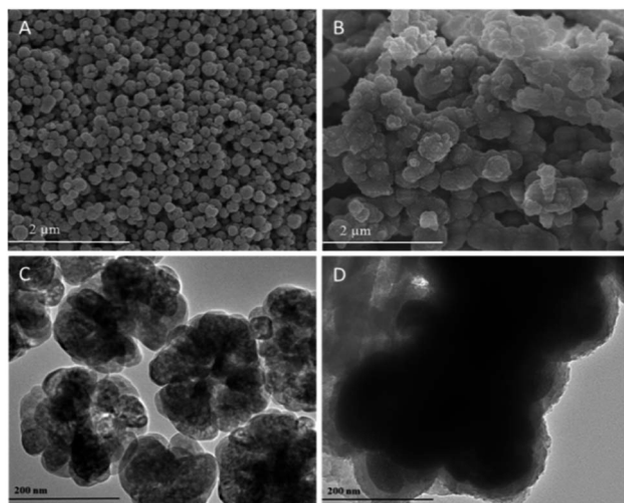


Fig. 1 The SEM of (A) Fe_3O_4 and (B) $\text{Fe}_3\text{O}_4@SiO_2@Ah-COF$; the TEM of (C) Fe_3O_4 and (D) $\text{Fe}_3\text{O}_4@SiO_2@Ah-COF$.

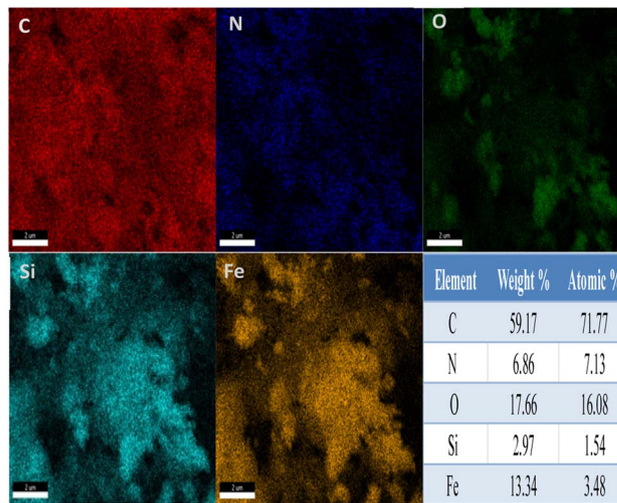


Fig. 2 EDS elemental mapping of $\text{Fe}_3\text{O}_4@SiO_2@Ah-COF$.

As seen in Fig. 3A, the XRD diagram of Fe_3O_4 showed six characteristic diffraction peaks at $2\theta = 30.1^\circ, 35.48^\circ, 43.15^\circ, 53.44^\circ, 57.01^\circ,$ and 62.61° , which corresponded to the (220), (311), (400), (422), (511) and (440) planes, respectively, and were in good agreement with the standard magnetite XRD patterns (JCPDS No. 75-1610).⁵⁰ The XRD results validated that the crystalline phase of Fe_3O_4 was not destroyed in all samples during the preparation of $\text{Fe}_3\text{O}_4@SiO_2$ and $\text{Fe}_3\text{O}_4@SiO_2@Ah-COF$.

The synthesis process of $\text{Fe}_3\text{O}_4@SiO_2@Ah-COF$ was evaluated by FT-IR. As depicted in Fig. 3B, the typical band at 587 cm^{-1} was assigned to the Fe–O–Fe vibration. The broad bands of –OH at 3441 cm^{-1} and 1631 cm^{-1} might be assigned to the stretching and bending vibrations of water molecules on the

surface of Fe_3O_4 , respectively. The peaks around 1091 cm^{-1} could be related to the Si–O bending vibration. This verified that the silica shells were successfully encapsulated onto the surface of Fe_3O_4 . Compared with the spectra of Fe_3O_4 and $\text{Fe}_3\text{O}_4@SiO_2$, the almost identical characteristic absorption peaks described above were also observed in the spectrum of $\text{Fe}_3\text{O}_4@SiO_2@Ah-COF$. Furthermore, the emerging stretching features at 1438 cm^{-1} and 1623 cm^{-1} were caused by the C=N stretching mode, demonstrating that the COFs successfully covered the surface of $\text{Fe}_3\text{O}_4@SiO_2$ via Schiff-base condensation reaction. The FT-IR results confirmed the successful fabrication of $\text{Fe}_3\text{O}_4@SiO_2@Ah-COF$.



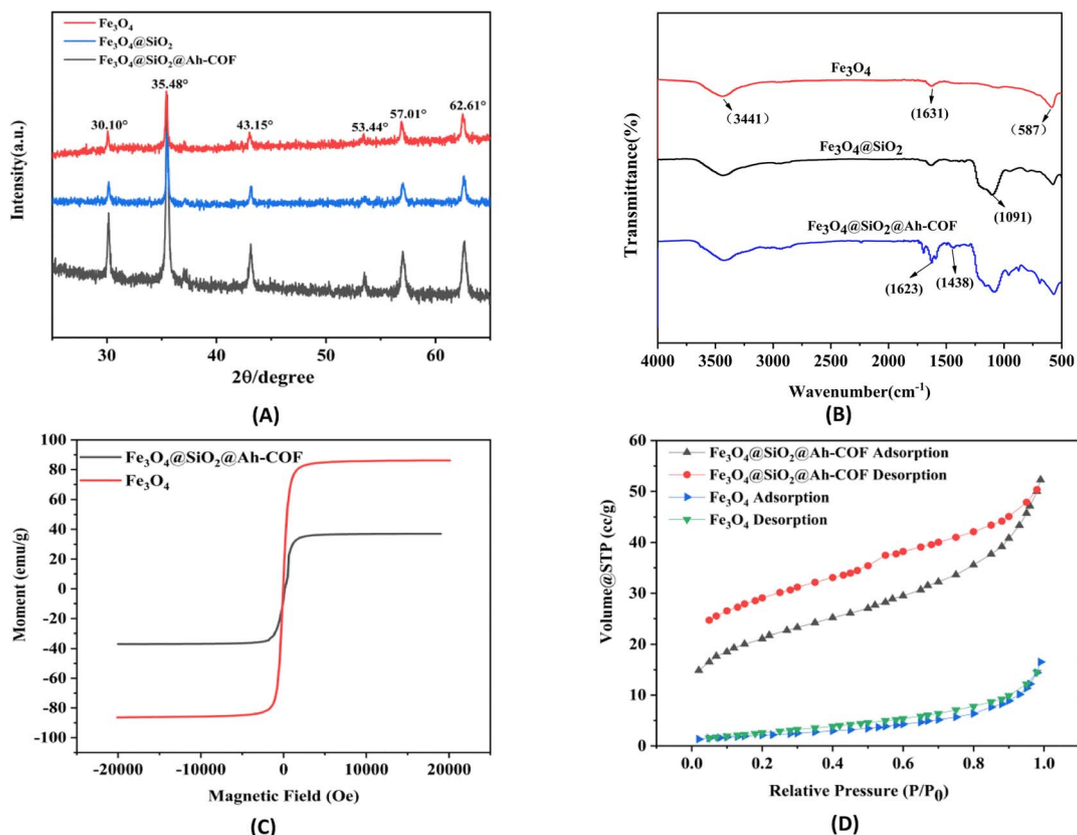


Fig. 3 (A) XRD analysis of Fe₃O₄, Fe₃O₄@SiO₂ and Fe₃O₄@SiO₂@Ah-COF. (B) FT-IR spectra of Fe₃O₄, Fe₃O₄@SiO₂ and Fe₃O₄@SiO₂@Ah-COF. (C) VSM magnetization curves of Fe₃O₄ and Fe₃O₄@SiO₂@Ah-COF. (D) N₂ adsorption-desorption isotherms of Fe₃O₄ and Fe₃O₄@SiO₂@Ah-COF.

Vibration magnetometer (VSM) was used to measure the magnetic values of Fe₃O₄ and Fe₃O₄@SiO₂@Ah-COF, and the results are shown in Fig. 3C. The saturation magnetization value of the bare Fe₃O₄ and Fe₃O₄@SiO₂@Ah-COF was 79.12 and 37.02 emu g⁻¹, respectively. Compared with Fe₃O₄, the saturated magnetization of Fe₃O₄@SiO₂@Ah-COF was somewhat reduced, which was attributed to the formation of a core-shell structure. Despite the decreased magnetization, the magnetic strength of Fe₃O₄@SiO₂@Ah-COF was still adequate for magnetic separation. Fe₃O₄@SiO₂@Ah-COF was well dispersed in water, and Fe₃O₄@SiO₂@Ah-COF can be collected in 60 s in an external magnetic field, indicating that the obtained Fe₃O₄@SiO₂@Ah-COF exhibited excellent magnetism ability and could be used for the rapid magnetic separation.

To evaluate the porosity and surface area, N₂ adsorption-desorption analysis at 77 K was carried out. As displayed in Fig. 3D, there is a well-defined type IV isotherm, indicating the presence of a mesoporous structure, and the average pore size of the magnetic material is 4.51 nm. After calculation, the BET surface area of Fe₃O₄@SiO₂@Ah-COF was 71.82 m² g⁻¹, which was higher than that of Fe₃O₄ with a surface area of 7.89 m² g⁻¹. This suggests that Fe₃O₄@SiO₂@Ah-COF can provide abundant active sites, increase the opportunity for magnetic materials to come into contact with QNs and improve the adsorption efficiency. These results confirmed the successful preparation of Fe₃O₄@SiO₂@Ah-COF.

3.2 Condition of MSPE

To obtain a high extraction recovery, parameters that influence the extraction performance (including the extraction dosage, sample solution pH, NaCl concentration, extraction time, elution solvent type, volume and elution time) were optimized in detail.

3.2.1 Effect of the adsorbent dosage. The dosage of Fe₃O₄@SiO₂@Ah-COF is a key factor that affects the extraction efficiency in the MSPE process. Various dosages of Fe₃O₄@SiO₂@Ah-COF were investigated within the range of 8–14 mg. As exhibited in Fig. 4, the experimental results showed that the best recoveries of the 24 QNs were achieved when the dosage of Fe₃O₄@SiO₂@Ah-COF was increased to 10 mg. No significant change occurred with the increase of the adsorbent dosage. The possible reason is that the adsorbent specific surface area and availability of more adsorption sites increase with the increase in adsorbent dosage, and then the adsorbent dosage reaches saturation. Thus, 10 mg was utilized for the following experiments.

3.2.2 Effect of the sample solution pH. The pH of a solution affects the existing form of QNs and the surface charge of the magnetic material,⁵¹ which have a significant influence on the extraction efficiencies. The pK_a value of QNs in aqueous solution is 5.66–8.56, and QNs exists as cations under acidic conditions. In this study, the pH of the sample solution in the



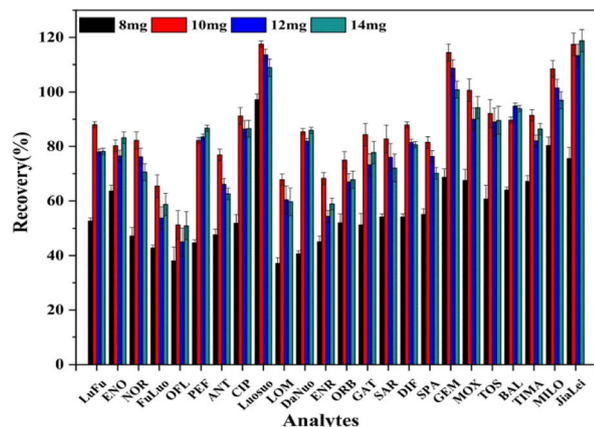


Fig. 4 Effect of the adsorbent dosage.

range of 2.0–8.0 was investigated by adding an appropriate amount of hydrochloric acid or sodium hydroxide. As shown in Fig. S1A,[†] the extraction efficiencies of all QNs increased slightly with a pH of 2.0–4.0. From pH 4.0 to 5.0, the extraction efficiencies remained nearly constant, and the recoveries of QNs decreased at a pH of 5.0–8.0. The higher extraction efficiency might be due to the electrostatic interaction between the negative charge of the oxygen-containing group on the surface of the magnetic material and the positive charge carried by QNs,⁵² as well as the mutual π - π interaction between the QNs and COFs. Therefore, a sample pH of 4.0 was used in further experiments.

3.2.3 Effect of the salt concentration. The solubility of analytes in aqueous solution is influenced by the ionic strength of the solution, which plays an important role in the distribution of analytes in the organic phase and the aqueous phase.⁵³ The effect of the ionic strength was systematically investigated by adding a certain amount of NaCl (0–20%, w/v) to the samples. The results in Fig. S1B[†] show that the recoveries of some analytes (LOM, ANT and TOS) clearly decreased with the increase of the salt concentration, and the overall recoveries of 24 QNs achieved satisfactory results without adding salt. Therefore, no salt was added in the following experiment.

3.2.4 Effect of the extraction time. In the process of MSPE, the extraction time affects the distribution equilibrium of QNs in the adsorbents and in the sample solution, so it also plays a key role in the extraction efficiency. As displayed in Fig. 5A, the vortex time spanned from 5 to 25 min. From 5 min to 20 min, the overall recovery of QNs showed an upward trend. As the vortex time was further extended to 25 min, there was no significant change in the adsorption efficiency. Herein, the optimal adsorption time was 20 min.

3.2.5 Investigation of the desorption parameters. In order to obtain a good recovery of the target compound, it is essential to select a suitable elution solvent type, volume and elution time.

The eluent is an influential factor to obtain the best elution efficiency. It has been reported that acidity is conducive to the elution of QNs,⁵⁴ so we first investigated the effects of methanol, acetonitrile, 0.5% (v/v) acidified methanol and 0.5% (v/v)

acidified acetonitrile on the recovery. Results showed that the elution efficiency of methanol is dramatically higher than that of acetonitrile. The highest recovery rate of methanol was 75.2%, while that of acetonitrile was only 15.8%. Compared with acidified acetonitrile, acidified methanol led to a higher elution efficiency. The possible reason can be explained as follows: methanol has a polarity that is more similar to that of the QNs, so it has a relatively high dissolution capacity for analytes. The concentration of formic acid in methanol was further investigated. As shown in Fig. 5B, the elution efficiencies of 24 QNs continuously increased with the increase of the formic acid concentration in methanol, and then declined at a formic acid concentration of 5% (v/v). Therefore, 3% (v/v) formic acid in methanol was selected for elution.

Fig. 5C shows that QNs can achieve good recoveries when the eluent volume increased to 8 mL. When the eluent was fixed at 8 mL, the effect of varied elution time was further studied from 3 to 11 min. As shown in Fig. 5D, the recoveries increased gradually with the increasing vortex time from 3 to 8 min, and then slowly decreased over 8 min. The results indicated that reversible adsorption reaches equilibrium after 8 min.⁵⁵ Therefore, 8 min of elution time was selected in this study.

3.3 Adsorption mechanism

$\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{Ah-COF}$ contains an imine structure, abundant carbonyl groups and a large number of benzene ring structures. Among them, the lone pair electrons on the N atom and the benzene ring structure are prone to form p- π and π - π interactions with the piperazine ring of QNs; and the carbonyl groups contained in it form hydrogen bonds with the carbonyl or carboxyl groups of QNs, which makes the magnetic materials have good adsorption effects.

In addition, the large specific surface area of the magnetic materials can provide more adsorption sites for the target analytes, improving the adsorption capacity. Moreover, the presence of mesoporous structures is beneficial for reducing the mass transfer resistance, which allows the target substance to quickly pass through the material.

3.4 Reusability of the adsorbent

Reusability is a significant parameter to evaluate the tolerance performance of a certain adsorbent. After the analytes were desorbed from the magnetic material, the magnetic material was washed and dried, and then the adsorption-desorption cycle was tested under optimal condition with spiked blank samples. As shown in Fig. S2,[†] the results proved that $\text{Fe}_3\text{O}_4@\text{SiO}_2@\text{Ah-COF}$ could be reused 5 times for the recoveries of 24 QNs without an obvious decrease, which greatly reduces the experimental cost.

3.5 Matrix effect

Matrix effect (ME) refers to matrix interferences with the target analytes, causing the mass spectrometry signal to be suppressed or enhanced during analysis and detection.⁵⁶ In this method, the ME of 24 QNs in water and egg samples was evaluated by the ratio of slopes between the matrix-matched calibration curves



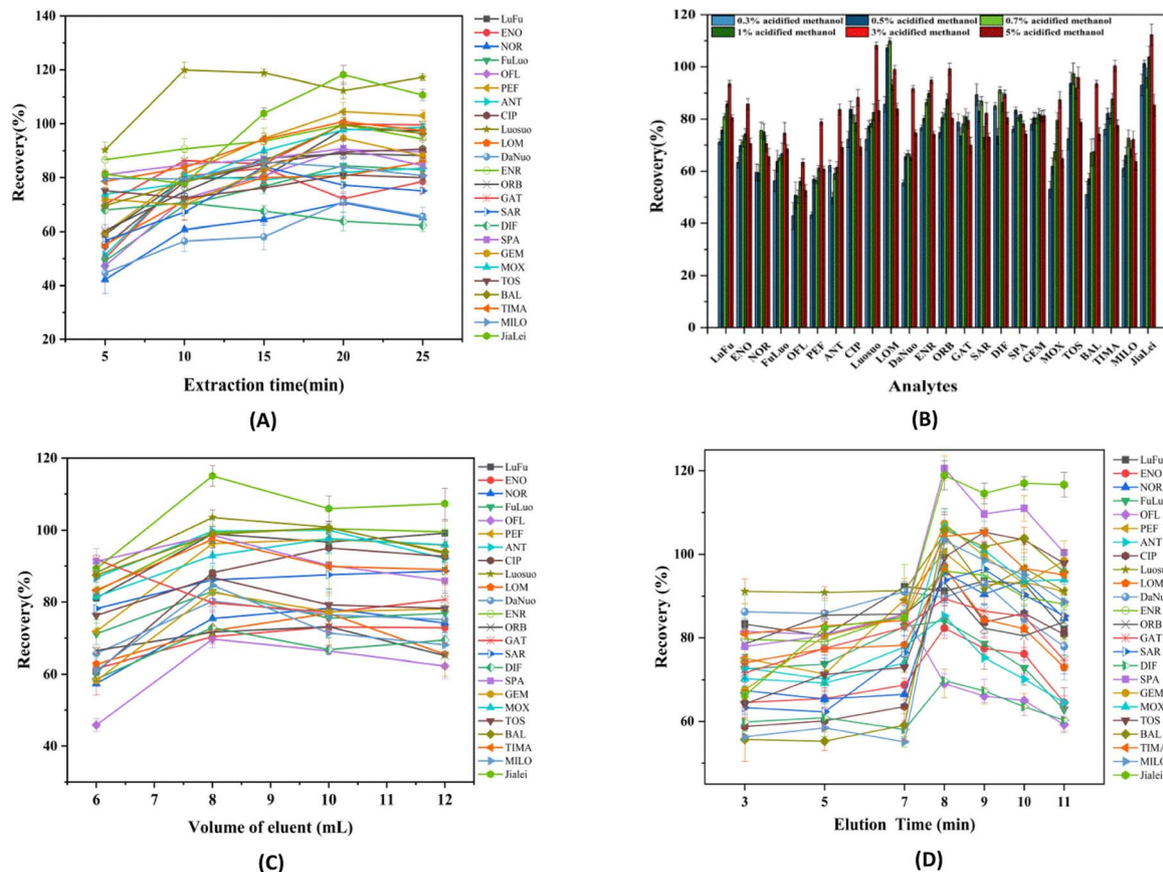


Fig. 5 Effect of the extraction time (A), elution solvent type (B), the volume of eluent (C) and elution time (D) of 24 QNs.

Table 1 Analytical performance of the proposed MSPE-UPLC-MS/MS method for the determination of QNs in water and egg samples

Analytes	Linear range (ng mL ⁻¹)	Calibration curve	R ²	LOD		LOQ	
				Water (µg L ⁻¹)	Egg (µg kg ⁻¹)	Water (µg L ⁻¹)	Egg (µg kg ⁻¹)
LuFu	0.2–200	Y = 24 230.5X + 2717.2	0.9993	0.011	0.272	0.037	0.908
ENO	0.2–200	Y = 45 259X + 4032.3	0.9996	0.007	0.083	0.023	0.276
NOR	0.2–200	Y = 26 159.1X + 3118.5	0.9999	0.004	0.188	0.014	0.625
FuLuo	0.2–200	Y = 161 000X + 10735.5	0.9999	0.009	0.083	0.030	0.277
OFL	0.2–200	Y = 177 862X + 62971.9	0.9999	0.011	0.034	0.037	0.112
PEF	0.2–200	Y = 126 828X + 1784.1	0.9998	0.003	0.026	0.010	0.087
ANT	0.2–200	Y = 153 922X + 62455.4	0.9999	0.007	0.009	0.022	0.029
CIP	0.2–200	Y = 34 507.3X + 5276.8	0.9998	0.024	0.152	0.074	0.507
LOM	0.2–200	Y = 78 931.8X + 65585.7	0.9992	0.010	0.072	0.030	0.238
Luosuo	0.2–200	Y = 149 193X + 10463.8	0.9996	0.021	0.088	0.064	0.293
DaNuo	0.2–200	Y = 218 201X + 24850.1	0.9991	0.007	0.023	0.022	0.076
ENR	0.2–200	Y = 152 947X + 11765.5	0.9999	0.013	0.025	0.038	0.082
ORB	0.2–200	Y = 206 573X + 50924.4	0.9999	0.020	0.022	0.059	0.073
GAT	0.2–200	Y = 43 867.2X + 18974.3	0.9997	0.010	0.065	0.031	0.217
SAR	0.2–200	Y = 44 213.8X + 5397.6	0.9998	0.013	0.067	0.039	0.224
DIF	0.2–200	Y = 132 196X + 10242.1	0.9997	0.009	0.027	0.029	0.091
SPA	0.2–200	Y = 218 233X + 149197	0.9996	0.006	0.017	0.018	0.058
MOX	0.2–200	Y = 32 873.5X + 6457.6	0.9997	0.011	0.146	0.033	0.486
TOS	0.2–200	Y = 124 395X + 48280	0.9998	0.015	0.042	0.047	0.071
GEM	0.2–200	Y = 118 134X + 16785.2	0.9999	0.010	0.138	0.031	0.461
BAL	0.2–200	Y = 151 779X + 3071.7	0.9997	0.003	0.034	0.008	0.115
TIMA	0.2–200	Y = 78 438.7X + 3269.3	0.9997	0.035	0.033	0.106	0.108
MILO	0.2–200	Y = 327 264X + 25903.5	0.9997	0.021	0.125	0.064	0.418
JiaLei	0.2–200	Y = 40 601.6X + 79814.7	0.9992	0.013	0.143	0.041	0.478



Table 2 Comparison with other methods reported in the literature for the determination of QNs^a

Method	Adsorbent	Sample (numbers of QNs)	Extraction time (min)	Linearity	Recovery (%)	LOD	Ref.
MSPE-HPLC-MS/MS	Fe ₃ O ₄ @COF(TpBD)@Au-MPS	Pork, chicken and bovine (6)	30	0.005–0.500 mg L ⁻¹	94.23–98.6	0.002–0.004 mg kg ⁻¹	42
MSPE-HPLC-DAD	Fe ₃ O ₄ @COFs	Milk, pork and human plasma (6)	60	2.5–1500 ng g ⁻¹	78.7–103.5	0.25–0.5 ng g ⁻¹	43
MSPE-LC-FLD	Fe ₃ O ₄ @PLS	Milk (3)	20	0.4–600 µg kg ⁻¹	96.5–118.6	0.41–0.45 µg L ⁻¹	58
RAM-MIPs-DSPE-HPLC-UV	RAM-MIPs	Milk and river water (5)	20 and 15	5–200 µg L ⁻¹	80.7–103.5	1.02–3.15 µg L ⁻¹	59
SPE-LC-MS/MS	Oasis HLB SPE cartridges	Aquaculture wastewater (17)	>25	0.5–100 µg L ⁻¹	85.1–105.9	0.93–2.87 µg L ⁻¹	60
MSPE-RPLC-MS/MS	PEI-functionalized Fe ₃ O ₄ /ATP	Chicken muscle (3)	20	0.2–100 µg kg ⁻¹	47.8–118.7	0.08–0.3 ng L ⁻¹	41
MSPE-UPLC-MS/MS	Fe ₃ O ₄ @SiO ₂ @Ah-COF	Water and egg samples (24)	20	0.2–200 µg kg ⁻¹	83.9–98.7	0.02–0.08 µg L ⁻¹	This work
					70.3–106.1	0.003–0.036 µg L ⁻¹	
					70.4–119.7	0.009–0.272 µg kg ⁻¹	

^a DAD: diode-array detector, FLD: fluorescence detection, RAM-MIPs: restricted access media-MIPs, RPLC: reversed-phase liquid chromatography.

and the solvent calibration curves. According to the calculation results (Table S3[†]), the ME values of QNs in water and egg samples were 85.4–113.7% and 80.3–116.4%, respectively, suggesting that the QNs had no obvious matrix effect.⁵⁷ Therefore, we chose to use the external standard method using a solvent standard curve for quantitative analysis of the experiment.

3.6 Method validation

The analytical performance of the developed MSPE-UPLC-MS/MS method under the optimal conditions is listed in Table 1. The linear range of 24 QNs was 0.2–200 µg kg⁻¹, with *R*² ranging from 0.9990 to 0.9999. For water samples, the limits of detection (LOD) [signal-to-noise ratio (S/N) of 3] was 0.003–0.036 µg L⁻¹ and the limits of quantitation (LOQ) (S/N of 10) was 0.008–0.110 µg L⁻¹. For egg samples, the LOD was 0.009–0.272 µg kg⁻¹ and the LOQ was 0.029–0.908 µg kg⁻¹. The results proved that the method had satisfactory linearity and high sensitivity.

As shown in Table S4,[†] with three different spiked levels, the 24 QNs obtained good recoveries in both water and egg samples, which were respectively 70.3–106.1% and 70.4–119.7% with RSD below 10%, indicating the reliability of the method.

3.7 Analysis of real samples

The method was further applied to the detection of the QNs residues in 15 water and 20 egg samples. As a result, SAR was detected in 4 egg samples and the levels varied from 4.07 to 7.54 µg kg⁻¹.

3.8 Comparison with other methods

This method was compared with other reported methods using different adsorbents for the detection of QNs residues.

As seen in Table 2, compared to traditional SPE and DSPE, MSPE simplified the extraction procedure and our method can directly extract the target QNs from real samples without protein precipitation treatment,^{59,60} which decreased the use of organic solvents and shortened the sample preparation time. It is worth noting that previous reports have mostly focused on the detection of milk and meat samples with less detection in egg samples.^{41–43,58,59} In reported methods, most of them involved the detection of 3–6 QNs. Only one method for 17 QNs detected in aquaculture wastewater was established, and the testing sample matrix was single. In comparison, this study was able to simultaneously determine 24 QNs in water and eggs with low LOD and less extraction time, which showed high throughput and high sensitivity.

4. Conclusion

This study reported the design and synthesis of Fe₃O₄@-SiO₂@Ah-COF for the first time, which possessed remarkable properties such as good magnetism, large surface area and abundance of carbonyl groups, leading to excellent adsorption efficiency for 24 QNs. Then, a new MSPE-UPLC-MS/MS method was established for the determination of 24 QNs in water and egg samples, showing a simple extraction process, low LOD and LOQ, good accuracy and wide linear range. Real samples that

possibly contained residues of QNs were analysed using the developed method, and SAR was found in four egg samples. In summary, the proposed method is suitable for the simultaneous and high-throughput determination of trace QNs in egg and water samples.

Author contributions

Na Li: conceptualization, formal analysis, methodology, writing – original draft. Mengnan Liang: data curation, formal analysis. Hao Zhang: methodology, validation. Zhongxia Hua: methodology, supervision. Ling Ma: conceptualization, supervision, funding acquisition, formal analysis. Yanyu Qi: conceptualization, supervision. Ke Wang: conceptualization, funding acquisition, project administration, resources, writing – review & editing. All authors have read and agreed to the published version of the manuscript.

Conflicts of interest

The authors declare no competing financial interest.

Acknowledgements

This work was supported by the National Natural Science Foundation of China (81903322), and the S & T Program of Hebei (223777116D).

References

- S. Guan, H. Wu, L. Yang, Z. Wang and J. Wu, *J. Sep. Sci.*, 2020, **43**, 3775–3784.
- X. Lei, X. Xu, L. Liu, H. Kuang, L. Xu and C. Xu, *J. Dairy Sci.*, 2022, **105**, 1999–2010.
- C. Fengxian and H. Reti, *Ecotoxicol. Environ. Saf.*, 2017, **136**, 111–118.
- R. Wang, K. Xu and W. Shi, *Arch. Pharm.*, 2019, **352**, 1900045.
- X. Liu, Y. Liu, J.-R. Xu, K.-J. Ren and X.-Z. Meng, *Environ. Pollut.*, 2016, **219**, 916–923.
- H. Zhang, X. Wang and Y. Li, *Environ. Chem.*, 2022, **41**, 1168–1181.
- M. Luo, K. Xing, Z. Guo, D. Guo, W. Lai and J. Peng, *J. Dairy Sci.*, 2020, **103**, 7791–7800.
- X. Xu, L. Feng, J. Li, P. Yuan, J. Feng, L. Wei and X. Cheng, *Chin. Chem. Lett.*, 2019, **30**, 549–552.
- A. Rusu, G. Hancu, G. Volgyi, G. Toth, B. Noszal and A. Gyeresi, *J. Chromatogr. Sci.*, 2013, **52**, 919–925.
- I. Varenina, N. Bilandžić, Đ. B. Luburić, B. S. Kolanović, I. Varga, M. Sedak and M. Đokić, *Food Control*, 2023, **148**, 109676.
- H. Wang, X. Zhao, J. Xu, Y. Shang, H. Wang, P. Wang, X. He and J. Tan, *J. Chromatogr. A*, 2021, **1651**, 462286.
- S. Bang Ye, Y. Huang and D.-Y. Lin, *Food Chem.*, 2022, **373**, 131466.
- H. Wang, M. Gao, M. Wang, R. Zhang, W. Wang, R. A. Dahlgren and X. Wang, *J. Chromatogr. B*, 2015, **985**, 62–70.
- L. Hou, Y. Ji, J. Zhao and L. Zhao, *Microchem. J.*, 2022, **179**, 107664.
- Y. Zhu, P. He, H. Hu, M. Qi, T. Li, X. Zhang, Y. Guo, W. Wu, Q. Lan, C. Yang and H. Jin, *J. Chromatogr. B*, 2022, **1208**, 123390.
- Z. Lu, F. Deng, R. He, L. Tan, X. Luo, X. Pan and Z. Yang, *Microchem. J.*, 2019, **151**, 104213.
- W. Huang, P. Wang, P. Jiang, X. Dong and S. Lin, *J. Chromatogr. A*, 2018, **1573**, 59–65.
- H. Yu, Z. Wang, R. Wu, X. Chen and T. W. D. Chan, *J. Chromatogr. A*, 2019, **1601**, 27–34.
- X. Di, X. Wang, Y. Liu and X. Guo, *J. Sep. Sci.*, 2019, **42**, 2187–2298.
- H.-z. Tang, Y.-h. Wang, S. Li, J. Wu, J.-w. Li, H.-y. Zhou and Z.-x. Gao, *Anal. Bioanal. Chem.*, 2019, **411**, 7039–7049.
- A. Wen, G. Li, D. Wu, Y. Yu, Y. Yang, N. Hu, H. Wang, J. Chen and Y. Wu, *J. Chromatogr. A*, 2020, **1612**, 460651.
- L. Lian, X. Zhang, J. Hao, J. Lv, X. Wang, B. Zhu and D. Lou, *J. Chromatogr. A*, 2018, **1579**, 1–8.
- H. Yu, Y. Jia, R. Wu, X. Chen and T. W. D. Chan, *Anal. Bioanal. Chem.*, 2019, **411**, 2817–2826.
- Y. Du, X. Yan, Y. Chen, Y. Wu, Q. Qiu, Y. Li and D. Wu, *J. Chromatogr. A*, 2022, **1675**, 463184.
- L. Fu, H. Zhou, E. Miao, S. Lu, S. Jing, Y. Hu, L. Wei, J. Zhan and M. Wu, *Food Chem.*, 2019, **289**, 701–707.
- X. He, G. N. Wang, K. Yang, H. Z. Liu, X. J. Wu and J. P. Wang, *Food Chem.*, 2017, **221**, 1226–1231.
- M. Hernández-Mesa, C. Cruces-Blanco and A. M. García-Campaña, *Talanta*, 2017, **163**, 111–120.
- H. Duo, X. Lu, X. Nie, L. Wang, S. Wang, X. Liang and Y. Guo, *J. Chromatogr. A*, 2020, **1626**, 461328.
- S. Wang, H. Niu, D. Cao and Y. Cai, *Talanta*, 2019, **194**, 522–527.
- H. Fan, A. Mundstock, A. Feldhoff, A. Knebel, J. Gu, H. Meng and J. Caro, *J. Am. Chem. Soc.*, 2018, **140**, 10094–10098.
- W. Huang, J. Byun, I. Rörich, C. Ramanan, P. W. M. Blom, H. Lu, D. Wang, L. Caire da Silva, R. Li, L. Wang, K. Landfester and K. A. I. Zhang, *Angew. Chem., Int. Ed.*, 2018, **57**, 8316–8320.
- D. Zhang, Y. Wang, W. Geng and H. Liu, *Sens. Actuators, B*, 2019, **285**, 546–552.
- M. Ghasemi, M. Khedri, M. Didandeh, M. Taheri, E. Ghasemy, R. Maleki, H. k. Shon and A. Razmjou, *Ind. Eng. Chem. Res.*, 2022, **61**, 8809–8820.
- A. Irfan, T. Wang, A. Wang, X. Jing, L. Yang and G. Zhu, *Anal. Chim. Acta*, 2022, **1209**, 339876.
- W. Tan, L. Zhu, L. Tian, H. Zhang, R. Peng, K. Chen, S. Zhao and F. Ye, *J. Chromatogr. A*, 2022, **1675**, 463188.
- H. Liu, Y. Zhang, D. Zhang, F. Zheng, M. Huang, J. Sun, X. Sun, H. Li, J. Wang and B. Sun, *Microchim. Acta*, 2019, **186**, 182–191.
- H. Yuan, Z. LU, Y. Li, C. Zhang and G. Li, *Chin. J. Chromatogr.*, 2022, **40**, 109–122.
- Y. Li, C.-X. Yang and X.-P. Yan, *Chem. Commun.*, 2017, **53**, 2511–2514.
- J. Xin, X. Wang, N. Li, L. Liu, Y. Lian, M. Wang and R.-S. Zhao, *Food Chem.*, 2020, **330**, 127255.



- 40 N. Li, D. Wu, N. Hu, G. Fan, X. Li, J. Sun, X. Chen, Y. Suo, G. Li and Y. Wu, *J. Agric. Food Chem.*, 2018, **66**, 3572–3580.
- 41 X. Li, Y. Chen, S. Chen, C. Hou, R. Xuan, Y. Gao, S. Ren, L. Yao, T. Wang, L. Zhang and Y. Zhang, *Anal. Bioanal. Chem.*, 2021, **413**, 3529–3540.
- 42 L. Zhang and W. Jiang, *Food Sci. Technol.*, 2022, **44**, 269–277.
- 43 M. Wang, M. Gao, K. Zhang, L. Wang, W. Wang, Q. Fu, Z. Xia and D. Gao, *Microchim. Acta*, 2019, **186**, 827–850.
- 44 A. O. Melekhin, V. V. Tolmacheva, N. O. Goncharov, V. V. Apyari, S. G. Dmitrienko, E. G. Shubina and A. I. Grudev, *Food Chem.*, 2022, **387**, 132866.
- 45 Y. Li, X. Nie, M. Yang, W. Guo, F. Chen and F. Zhang, *Food Sci.*, 2022, **43**, 295–305.
- 46 Y. Li, H. Zhang, Y. Chen, L. Huang, Z. Lin and Z. Cai, *ACS Appl. Mater. Interfaces*, 2019, **11**, 22492–22500.
- 47 D. Li, M. He, B. Chen and B. Hu, *J. Chromatogr. A*, 2019, **1601**, 1–8.
- 48 Y. Chen and Z. Chen, *Talanta*, 2017, **165**, 188–193.
- 49 W. Zhang, C. Lan, H. Zhang, Y. Zhang, W. Zhang, W. Zhao, C. Johnson, K. Hu, F. Xie and S. Zhang, *J. Agric. Food Chem.*, 2019, **67**, 3733–3743.
- 50 L. Chen, M. Zhang, F. Fu, J. Li and Z. Lin, *J. Chromatogr. A*, 2018, **1567**, 136–146.
- 51 Y. Liu, H. Wu, X. Du, Y. Feng, Y. Zhao and Y. Chang, *J. Anal. Sci.*, 2018, **34**, 196–200.
- 52 M. K. Abraham, A. S. Madanan, S. Varghese, A. I. Shkhair, G. Indongo, G. Rajeevan, N. S. Vijila and S. George, *Analyst*, 2023, **149**, 231–243.
- 53 H.-L. Jiang, Q.-B. Fu, M.-L. Wang, J.-M. Lin and R.-S. Zhao, *Food Chem.*, 2021, **345**, 128841.
- 54 C. Vakh, M. Alaboud, S. Lebedinets, D. Korolev, V. Postnov, L. Moskvina, O. Osmolovskaya and A. Bulatov, *Anal. Chim. Acta*, 2018, **1001**, 59–69.
- 55 H.-B. Zheng, J.-Z. Mo, Y. Zhang, Q. Gao, J. Ding, Q.-W. Yu and Y.-Q. Feng, *J. Chromatogr. A*, 2014, **1329**, 17–23.
- 56 N. Jian, M. Zhao, S. Liang, J. Cao, C. Wang, Q. Xu and J. Li, *J. Agric. Food Chem.*, 2019, **67**, 6892–6901.
- 57 J. Li, X. Xu, W. Guo, Y. Zhang, X. Feng and F. Zhang, *Food Chem.*, 2022, **387**, 132821.
- 58 Z. Yu, B. Liang, Z. Zhao, S. Lin, H. He and S.-X. Liang, *J. Food Compos. Anal.*, 2023, **121**, 105352.
- 59 J. Li, Y. Zhou, Z. Sun, T. Cai, X. Wang, S. Zhao, H. Liu and B. Gong, *J. Chromatogr. A*, 2020, **1626**, 461364.
- 60 Z. Ding, Y. Li and X. Wang, *Chin. J. Environ. Eng.*, 2022, **16**, 674–683.

