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1. Introduction

Fluoroquinolones (FQs) are a group of synthetic antibiotics, which are derived from quinolone nalidixic acid by introduction of a fluorine atom at position 6 and a piperazine moiety at position 7¹ (ESI Table S1[†]). Since the first introduction of FQs for human use in the 1970s² and therapeutic treatment of livestock in the mid-1990s,³ they have become the most commonly prescribed antibiotics because of their broad activity spectrum towards mycoplasma, Gram-positive and Gramnegative bacteria.⁴ However, due to the extensive use of FQs, they enter environmental water *via* human excretion into wastewater or dispersion of manure onto agricultural soils, which could probably cause allergic reactions in humans and the emergence of food-borne bacteria.¹ The development of robust and sensitive methods to monitor FQ residues in environmental water is necessary.

Different techniques for FQs determination in aquatic environment have been developed such as high performance liquid

Highly-sensitive detection of eight typical fluoroquinolone antibiotics by capillary electrophoresis-mass spectroscopy coupled with immunoaffinity extraction[†]

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Fluoroquinolones (FQs) are a class of synthetic antibiotics with a broad active spectrum towards mycoplasma and bacteria (both Gram positive and negative). The contamination of environmental water samples with FQs has been a concern due to the extensive use of them in the veterinary and pharmaceutical industries. Reliable and simple analytical methods are needed for the monitoring of these FQs. An off-line procedure of immunoaffinity extraction (IAE) followed by capillary electrophoresis-mass spectroscopy (CE-MS) has been developed for simultaneous determination of eight typical FQs in environment water. A broad-specific polyclonal antibody was home-prepared and immobilized on a Sepharose 4B stationary phase for sample affinity extraction and the enrichment factor of our columns was about 100. CE-MS was subsequently applied to FQs separation and quantification. The immunoaffinity column was evaluated in the parameters of selectivity, binding capacity, elution protocol, and recovery. Under the optimized conditions, the developed method was successfully applied to analyze FQs spiked with 2.5 nM to 30 nM of environmental water from a local lake, providing satisfactory recoveries from 72% to 112%, and LODs and LOQs ranging from 1.2 nM to 5.0 nM and 4.0 nM to 17 nM, respectively.

chromatography with fluorescence detection (HPLC-FD),5-8 immunoassay,^{9,10} HPLC with ultraviolet (UV)¹¹ or mass spectroscopy (MS)^{12,13} detection, capillary electrophoresis (CE) both with UV14,15 or FD,16 and flow injection method with chemiluminescence detection.17 In our previous work,18 we detected five typical FQs in low-fat milk by field-enhanced sample injection-based CE in bubble cell capillary. As an online sample stacking strategy to overcome the low loading capacity of CE, field-enhanced sample injection was performed at the boundary of sample matrix with low-conductivity and the background electrolyte (BGE) zone with high-conductivity. That's to say such a developed method is not suitable to detect FQs in high-salt contained matrix like environmental water. Among those, immunoassay and methods using optical detectors did not provide structure confirmatory analysis, while the instrumentbased methods combining separation techniques and MS identification require an effective sample cleanup procedure to remove the complex matrix interference and extract the analytes.

The reported cleanup methods for FQs include solid phase extraction (SPE),^{6,11,12,15} pressurized liquid extraction (PLE),^{19,20} matrix solid-phase dispersion (MSPD),^{21,22} microwave-assisted extraction,^{23,24} microdialysis.²⁵ The most common used SPE was reverse-phase SPE based upon the hydrophobic interaction between the analytes and the sorbents, which was lack of

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selectivity, leading to the co-extraction of analytes and matrix interference, especially when interested analytes are at trace levels and interference at higher concentration. A special SPE, named immunoaffinity extraction (IAE) based on the highly specific recognition between antigen and antibody, allows for a high degree of molecular selectivity.26 IAE was generally realized by loading a large volume of the sample into the immunoaffinity column and afterwards washing the unbounded or nonspecifically adsorbed interferences, eluting the specifically captured compounds for the subsequent qualitative or quantitative analysis. This strategy has been applied in environmental monitoring,27,28 pharmaceutical and biomedical analysis,²⁹⁻³¹ and food analysis.^{32,33} As a specific extraction technique, IAE has been successfully used to online enrich four FQs in milk³⁴ prior to HPLC-FD, providing a satisfactory enrichment with good selectivity.

In this work, a generic antibody for a group of immunological similar FQs was prepared for immunoaffinity column formation to simultaneously extract those FQs analogs selectively. By combining IAE and CE-MS, detecting 8 FQs simultaneously was achieved. Samples were specifically purified and concentrated by IAE. The eluent from immunoaffinity column was concentrated and subsequently analyzed by CE-MS. We applied this method to determine FQs spiked water sample from a local lake. The LODs ranged from 1.2 nM to 5.0 nM, which matched the result of the National standard method, HPLC-FD, with the advantages at simplicity and structural specificity.

2. Experimental section

2.1 Reagents and solutions

Bovine serum albumin (BSA), ovalbumin (OVA), Tween-20, incomplete Freund's adjuvant, and complete Freund's adjuvant were purchased from Sigma (St. Louis, MO, USA). N-Hydroxysuccinimide (NHS) and 1-(3-dimethylaminopropyl)-3ethyl carbodiimide (EDC) were from Alfa Aesar (Johnson Matthey, USA). 3,3',5,5'-Tetramethylbenzidine (TMB), and horseradish peroxidase labeled goat anti-rabbit IgG (HRP-SecAb) were obtained from Balb Corp. (Beijing, China). Defatted milk was purchased from the local supermarket. Ciprofloxacin (CIP) and enrofloxacin (ENR) were from TCI (Tokyo, Japan), and pefloxacin (PEF), ofloxacin (OFL), gatifloxacin (GAT), fleroxacin (FLE), lomefloxacin (LOM) and enoxacin (ENO) from National Institutes for Food and Drug Control (Beijing, China). Methanol was of HPLC grade and obtained from Avantor (Avantor Performance Materials, Inc. USA). All other chemicals, including formic acid (FA), acetic acid (AA), ethylenediamine (EDA), hydrogen peroxide (H₂O₂), quinine (QUIN), and ethylenediamine (EDA) were of analytical grade or better and obtained from Beijing Chemical Reagent Co. (Beijing, China). Unless specified, all buffers were prepared in high purity water from a Milli-Q purification system (Millipore, Bedford, MA, USA).

In enzyme-linked immunosorbent assay (ELISA), 0.01 M phosphate buffered saline (PBS, pH 7.4) was prepared. 0.05 M Na_2CO_3 -NaHCO₃ (CBS, pH 9.6) was used as coating buffer. Blocking solution was made by dissolving 5 g of milk in 100 mL

of PBS, and washing buffer (PBST) consisted of 250 μ L of Tween-20 and 500 mL of PBS. The substrate solution was prepared before use by mixing 0.1 M phosphate buffer (PB, pH 6.0), TMB (6 mg mL⁻¹ in DMSO) and H₂O₂ (30%) at a volume ratio of 1000 : 20 : 2.

Stock solutions of the above mentioned FQs and QUIN were prepared in 20 mM NaOH or 1 mM HCl at a concentration of 10 mM and stored at -20 °C. The working solutions were prepared by diluting the stock solutions with assay buffer.

2.2 Apparatus

CE was performed using a PA 800 plus System (Beckman Coulter Inc., USA) equipped with a UV detector. MS analyses were realized by a 6320 Ion Trap (Agilent Technologies Inc., USA) with a commercial ESI source. The coupling of CE to MS was carried out using a G1608A Agilent CE-MS sprayer kit (Agilent Technologies Inc., USA), which is arranged orthogonally to MS inlet capillary and constituted of a triple-tube design accommodating one tube for separation capillary, one for sheath liquid delivery and another one to provide the nebulizing gas. All UV absorbance was measured on a HITACHI U-4100 spectrophotometer (Tokyo, Japan), and all polystyrene 96-well microtiter plates (Costar, USA) were read on Multiskan FC Microplate Photometer (Thermo Fisher Scientific, Waltham, MA, USA).

2.3 Preparation and characterization of broad-specific antibody against FQ analogs

2.3.1 Synthesis of immunogen and coating antigen. CIP was selected because of its common molecular structure. It was coupled with BSA (CIP-BSA) to immunize the rabbits, with OVA (CIP-OVA) using as coating antigen in ELISA. The conjugation procedure was based on the previously reported NHS ester method³⁵ with slightly modifications. In particular, CIP (90 mg) was dissolved in DMF (30 mL), followed by the addition of EDC (120 mg) and NHS (120 mg) to obtain the active NHS solution. The mixture was stirred at room temperature overnight. The carrier proteins (140 mg BSA, and 90 mg OVA) were dissolved in 30 mL carbonate buffer (0.1 M, pH 8.0), respectively. The above active NHS mixture was added dropwise to the carrier protein solutions with continuous stirring and further stirred at room temperature for 4 h. After being dialyzed against PBS for 12 h and deionized water for 24 h at 4 °C to remove the unbound hapten and residual solvent, the conjugate was lyophilized to white powder and stored at -20 °C before use. The ratio of CIP and protein in immunogen was estimated through the molecular weights of BSA and CIP-BSA determined by MALDI-TOF-MS.

2.3.2 Production of polyclonal antibody. New Zealand rabbits were purchased from Xinglong experimental animal farm (Beijing). Three healthy New Zealand rabbits were immunized with the immunogen, CIP-BSA. The immunization procedure involved five subcutaneous injections at different body parts on rabbits.³⁶ Anti-serum was obtained seven days after the last injection and purified based on a modified method of octanoic acid-saturated ammonium sulfate.³⁷ All the

experiments conform to the US National Institutes of Health "Guide for Care and Use of Laboratory animals" and were approved by the Institutional Animal Care and Use Committee of Peking University. The concentration of IgG was 16 mg mL⁻¹, which was estimated by its absorbance at 280 nm. The purified pAb was stored at -20 °C.

2.3.3 Titer level and affinity constant of the purified antibody. The optimization of coating concentration of CIP-OVA and the determination of titer level of purified pAb were performed by ELISA based on a chessboard method.³⁷ First of all, a 96-well microplate was coated with 100 μ L of coating buffer containing CIP-OVA at different concentrations overnight at 4 °C. Afterwards, the microplate was blocked with 200 μ L blocking solution (5% milk) at 37 °C for 2 h. Next, 100 μ L of tenfold serially diluted purified pAbs were added to each well and incubated for 1 h at 37 °C. The followed is the addition of 100 μ L HRP-SecAb (1 : 5000 diluted) for 1 h at 37 °C, then with 100 μ L substrate solution added to react for 15 min in dark. The enzyme reaction was stopped by 50 μ L of 2 M H₂SO₄ before the absorbance of each well read at 450 nm.

The affinity constant was estimated in the process of optimizing coating concentration using the equation presented by Beatty.³⁸

2.3.4 Cross-reactivity of polyclonal antibody with CIP analogs. The specificity of the obtained anti-CIP pAb was investigated by testing its cross-reactivities with the other seven FQs and QUIN by competitive ELISA. The microplate was coated with 100 μ L of CIP-OVA at an optimized concentration overnight at 4 °C, and blocked by defatted milk as described above. 50 μ L competitive compounds were then added at various concentrations together with 50 μ L of the 10⁴-fold diluted purified pAb. Other steps were the same as described in Section 2.3.3.

2.4 Preparation of immunoaffinity column

The immunoaffinity column was prepared according to the method reported before.²⁹ 0.3 g CNBr-activated Sepharose 4B was swelled and washed with 1 mM HCl, and then dispersed in 5 mL coupling buffer (0.1 M NaHCO₃, 0.5 M NaCl, pH 8.3) containing 18 mg pAb. The coupling of pAb to Sepharose 4B was performed in a rotated vessel for 2.5 h at room temperature. After the slurry was transferred into an injector (5 cm \times 0.7 cm id), the excess antibody was removed from the column by eluting with the coupling buffer. The unreacted groups in the gel were blocked with 0.1 M Tris–HCl buffer (pH 8.0) at room temperature for 2 h. The column was then washed for three times by repeating the cycles of 5 mL of 0.1 M NaAc–HAc buffer (containing 0.5 M NaCl, pH 4.0) followed by 5 mL of 0.1 M Tris–HCl buffer (containing 0.5 M NaCl, pH 8.0). 0.01 M PBS was finally applied for the column equilibration and preservation.

2.5 Procedure of immunoaffinity solid-phase extraction

10 mL sample was loaded on the immunoaffinity column, which has already been conditioned with 0.01 M PBS. The column was then washed with 3 mL PBS to remove the nonspecifically captured analytes and subsequently washed by 3 mL water to reduce the negative effect of salt on CE-MS. After

eluting the immunoaffinity column with 5 mL methanol, the collected effluent was concentrated to about 1 mL by high purity nitrogen and transferred to centrifugal vacuum concentrator (Eppendorf, Germany) for further concentration and drying. The extracted analyte was re-dissolved in deionized water (100 μ L) before the following CE-MS analysis.

2.6 CZE separation coupled with ESI-MS detection

Fused-silica capillaries (50/375 μ m id/od, 40/50 cm effective/ total length) (Ruifeng Inc., Hebei, China) were used for FQs separation. Capillary zone electrophoresis conditions were set as follows: applied voltage, 25 kV; sample injection, 0.5 psi for 5 s; capillary temperature, 25 °C. All data were collected at 280 nm. Before the first use, the capillaries were conditioned by sequentially rinsing by 20 psi with 1 M NaOH for 30 min, deionized water for 15 min and running buffer for 5 min. Between consecutive analyses, the capillary was only rinsed with running buffer for 4 min. The running buffer was 50 mM EDA-FA (pH 8.0).

When MS was hyphenated with CE *via* a coaxial sheath liquid interface for post-column detection, the length of the fusedsilica capillary was extended to 65 cm. The sheath liquid was composed of methanol/water (50 : 50 v/v) containing 0.1% (v) FA, and introduced at a flow rate of 5 μ L min⁻¹ by a syringe pump. An electrospray potential of -4.0 kV was applied at the inlet of the MS (positive mode) while the CE-MS interface was grounded. A nebulizing gas pressure of 5.0 psi was provided to assist the spraying. Drying gas temperature and flow were set at 270 °C and 2.0 L min⁻¹, respectively.

Unless specified, all solutions for CE-MS were filtered through 0.22 μm membrane and degassed by ultrasonication before use.

2.7 Real sample analysis

Fresh water was collected from Weiming Lake at Peking University (Beijing, China), and used to dilute the standard FQs solution to defined concentrations. FQs spiked samples were immediately filtered through 0.22 μ m cellulose nitrate membrane filters (Jinteng, China) before extraction in order to prevent the influence of the matrix from fresh water on immunoaffinity column. The blank water sample and the spiked real samples (both in 10 mL) were performed in the same way by the developed IAC and CE-MS off-line coupling procedure.

3. Results and discussion

3.1 Synthesis of artificial antigen

Antigen designing plays the most crucial role in antibody preparation, especially in the case of producing a broad-specific antibody. Herein, our aim lies in preparing an antibody that could immunologically recognize eight FQs and developing an immunoaffinity solid-phase extraction system for simultaneous enrichment of these FQs. Thus, the hapten, which was proposed to conjugate with a carrier protein to generate an immunogen for antibody selection, should retain the most of common structural features in FQ analogs. One or more FQs has been reported to be used as a hapten to produce a generic antibody,¹ and CIP might be the desirable hapten considering its availability and reliability. The immunogen, CIP-BSA, was prepared through CIP conjugated to BSA with the NHS method and the coating antigen was prepared through CIP coupled with OVA in the same way. The conjugation ratio of CIP and BSA was estimated to be 11.7 *via* the molecule weights of BSA (66.4 kDa) and its conjugation with CIP (70.3 kDa).

3.2 Antibody characterization

The purified polyclonal antibody was assessed by ELISA in the respect of titer level, affinity constant and cross-reaction.

The initial concentrations (C_0) of the antibody was 16 mg mL⁻¹ by UV measurement. The optimal concentration for coating was estimated by coating the microtiter plates with 100 µL CIP-OVA of different concentrations (2 µg mL⁻¹, 4 µg mL⁻¹, 5 µg mL⁻¹, 6 µg mL⁻¹, 8 µg mL⁻¹ and 10 µg mL⁻¹). According to the shape of sigmoid curves in Fig. 1(a), 8 µg mL⁻¹ was chosen as optimal concentration for CIP-OVA coating, and the titer of this purified pAb was 1 : 10^{4.8} (pAb dilution corresponding to 50% of the maximum absorbance).

Affinity constant (K_{aff}) of antibody represents the combination stability of immunocomplex. Herein, K_{aff} of pAb with

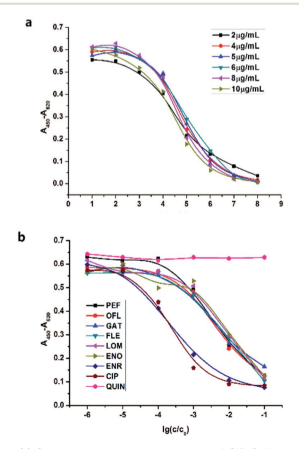


Fig. 1 (a) Coating concentration optimization of CIP-OVA and titer level determination of antibody by ELISA. (b) Cross-reactivities of pAb with eight FQ analogs by competitive ELISA (QUIN as control test).

CIP-OVA can be calculated as 1.53×10^7 L mol⁻¹ in the optimization procedure of the coating concentration, which was high enough for immunoaffinity enrichment.

The broad-specificity of antibody was determined by a screening competitive ELISA (Fig. 1(b)) and calculated according to 50% displacement method.³⁹ The microtiter plates were coated with 100 µL CIP-OVA (8 µg mL⁻¹). The purified pAb was diluted by 10⁴-fold before addition (50 µL per microwell). The initial concentrations (C_0) of the analytes (including QUIN for control test) were 0.1 mM. The crossreactivity of this purified pAb with CIP was defined as 100%, and it was found to be 10% with PEF, 10% with OFL, 10% with GAT, 10% with FLE, 8% with LOM, 10% with NOR, 8% with ENO, and 100% with ENR, indicating its broad-specificity to all these eight FQs.

3.3 Evaluation of the immunoaffinity column

The total amount of pAb immobilized in 0.3 g of CNBr-Sepharose 4B was estimated to be 17.7 mg (59 mg g^{-1}) by measuring the concentrations of pAb before and after the coupling reaction. The final gel volume of the affinity column (dead volume) was about 2.0 mL. In order to investigate retention behavior of CIP in the immunoaffinity column, breakthrough curve of CIP (0.4 µg mL⁻¹ in PBS) was compared with that of QUIN (0.4 μ g mL⁻¹ in PBS). As shown in Fig. 2(a), QUIN was detected in the first effluent fraction after discarding the dead volume of 2.5 mL. In contrast, CIP was strongly retained in the column, and the breakthrough of CIP was not observed until 5.5 mL, which demonstrated the specific retention of CIP in the immunoaffinity column. Binding capacity, another critical parameter in immunoaffinity column, is defined as the maximal amount of antigen captured by immobilized antibodies, and overloading the capacity would lower the accuracy of quantitative analysis. The theoretical binding capacity was calculated to be 20 µg according to the amount of immobilized antibody and its two antigen binding sites without consideration of the antibody inactivation. Otherwise, the experimental value of binding capacity was obtained as 1.03 μ g by overloading the column with CIP (20 μ g, 1 mL) and measuring the uncaptured (washed by PBS) or captured (eluted by methanol) target. It was much lower than the theoretical one, mainly resulted from the steric hindrance between analyte and antibody, as well as the random orientation and inactivation of the antibody.

3.4 Elute conditions

The antigen–antibody complex, formed *via* electrostatic forces, hydrogen bonds and van der Waals forces, was generally disrupted by low-pH solution or water–organic modifier mixture to dissociate the analytes without adversely affecting the immunological activity of antibodies. Here, high percentage of an organic solvent was chosen to dissociate captured small molecules efficiently considering the following two aspects. First, a large volume of low-pH solution was required for complete dissociation²⁹ which was not suitable for off-line procedure. In addition, the volatility of organic solvent would

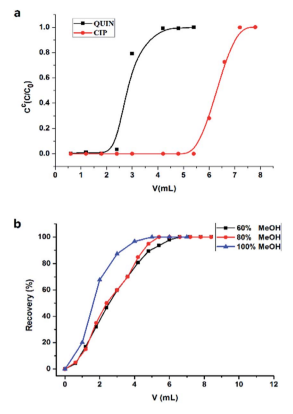


Fig. 2 (a) Breakthrough curves of CIP and QUIN obtained in anti-CIP pAb immobilized column. Sample concentration was 0.4 μ g mL⁻¹ both for CIP and QUIN. (b) Elution profiles of CIP with different percent of methanol.

make the off-line pre-concentration easier for the subsequent CE-MS analysis. The actual methanol concentration needed relied on the affinity constant between the analytes and the immobilized antibody. Thus, different percentages of methanol in water (60%, 80%, 100%) were investigated to make a compromise between complete elution of the analytes and minimal damage to the antibody. After the immuno-capture of CIP (20 μ g mL⁻¹), the elution fraction was collected in one Eppendorf tube after another and detected by UV respectively. The recovery of target compound increased with the solvent volume for eluting (Fig. 2(b)). All captured compound was completely eluted under three conditions, demonstrating a good target release performance. Nevertheless, the total volume of water-methanol mixture required for thorough dissociation decreased along with the increasing of methanol percentage. Moreover, the immobilized antibody was not observed to be damaged even when 100% methanol was applied for elution owing to the short exposing time of the antibody to organic solvent. Decrease of enrichment efficiency of our columns was not observed after about 50 times use. For the process efficiency and simplicity, 100% methanol was chosen as the optimum elution solvent. The elution volume was increased to be 5 mL rather than 4 mL to ensure a complete dissociation of the immuno-complexes of analytes and immobilized antibody.

3.5 CZE separation with UV or MS detection for quantitative analysis

Basic BGE is generally reported for FQs separation.15,40 Considering its compatibility with ESI-MS, the BGE should be constituted of volatile salts such as ammonium carbonate.41 However, FQs cannot be separated completely by using ammonium carbonate. Recently, BGE based on EDA with pH 8.0 adjusted by FA18 was proposed as volatile and compatible with MS system for FQs separation. FQs were detected in positive mode in CE-MS and quantitative ions for MS detection were listed in Table S1.[†] As indicated in Fig. 3(a), the considered eight FQs can be basely separated under optimized conditions within 15 min both for UV and MS detection. Theoretically, CE-MS was more time-consuming than CE-UV due to its longer separation channel and lowered electric field intensity, which cannot be found in the electropherograms. This can be explained by the syphonic effect between the two ends of CE separation capillary when there was no outlet vial assembled in CE equipped with MS detector.

The mixtures of eight FQs at different concentrations were prepared and analyzed both by CE-UV and CE-MS to investigate the linearity of the proposed procedure. The dependence of FQs peak heights (obtained in CE-UV) and peak areas (obtained in CE-MS) on their concentrations were fitted linearly, respectively,

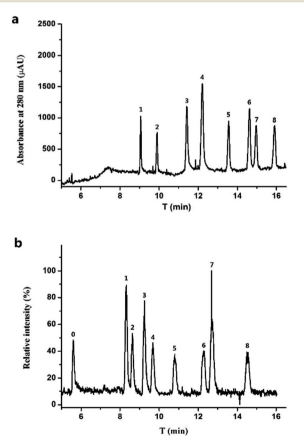


Fig. 3 Electropherograms with UV detection (a) and MS detection (b) of eight FQs (20 μ M for each) and QUIN (5 μ M). Peaks are labelled as follows: QUIN as 0; GAT as 1; LOM as 2; ENO as 3; CIP as 4; OFL as 5; ENR as 6; FLE as 7; PEF as 8.

FQs	CE-MS	CE-UV				
	Calibration curves	Calibration range (µM)	LOD/LOQ (µM)	Calibration curves	Calibration range (µM)	LOD/LOQ (µM)
GAT	$y = 4.89 \times 10^{6} x + 7.96 \times 10^{4}, R^{2} = 0.998$	0.16-20	0.050/0.16	$y = 24.7x + 39.2, R^2 = 0.998$	1.9-60	0.60/1.9
LOM	$y = 2.45 \times 10^{6} x + 1.11 \times 10^{5}, R^{2} = 0.998$	0.25-20	0.075/0.25	$y = 26.4x + 10.1, R^2 = 0.999$	5.6-60	1.7/5.6
ENO	$y = 4.95 \times 10^{6} x - 1.12 \times 10^{5}, R^{2} = 0.996$	0.16-20	0.050/0.16	$y = 14.6x + 6.2, R^2 = 0.999$	11-60	3.4/11
CIP	$y = 2.77 \times 10^6 x + 4.96 \times 10^5, R^2 = 0.997$	0.25-20	0.075/0.25	$y = 22.6x - 12.3, R^2 = 0.999$	10-60	3.0/10
OFL	$y = 2.33 \times 10^{6} x + 9.61 \times 10^{4}, R^{2} = 0.998$	0.50-20	0.15/0.50	$y = 26.5x + 14.7, R^2 = 0.999$	5.0-60	1.5/5.0
FLE	$y = 2.77 \times 10^{6} x + 3.55 \times 10^{5}, R^{2} = 0.999$	0.33-20	0.10/0.33	$y = 26.8x + 17.3, R^2 = 0.999$	4.3-60	1.3/4.3
ENR	$y = 5.16 \times 10^{6} x + 2.04 \times 10^{5}, R^{2} = 0.999$	0.33-20	0.10/0.33	$y = 30.0x + 9.2, R^2 = 0.999$	5.0-60	1.5/5.0
PEF	$y = 3.22 \times 10^{6} x + 1.23 \times 10^{5}, R^{2} = 0.999$	0.33-20	0.10/0.33	$y = 21.4x + 39.2, R^2 = 0.998$	6.6-60	2.0/6.6

and presented in Table 1. The LODs (3 S/N) and LOQs (10 S/N) were calculated and summarized in Table 1, which demonstrated that the LODs obtained in CE-MS were 10 to 40 times lower for different FQs than those in CE-UV. Thus, combining CE-MS of such an improved sensitivity with immunoaffinity

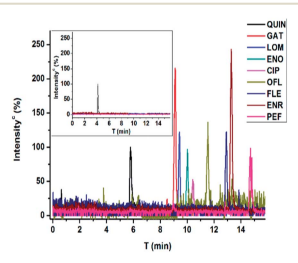


Fig. 4 Electropherograms with MS detection (extracted ion chromatography) for FQs spiked lake water (30 nM for FQs and 5 μ M for QUIN). The inset is the blank water sample for control.

extraction for enrichment provided a potential way to monitor the trace FQs in environmental water.

3.6 Environmental water analysis

Organics in environmental samples are generally in trace level and IAE provides an attractive approach to enrich structurally closed analytes in a small volume by off-line being coupled with a chromatographic step for separation. Herein, an evaluated immunoaffinity column was utilized for FQs extraction from spiked lake water and the developed CE-MS was applied for the subsequent qualitative and quantitative analysis. The environmental water samples were simultaneously spiked with eight FQs at different concentrations (30 nM, 15 nM, 5 nM and 2.5 nM, respectively) (Fig. 4). As indicated in Table 2, the presented procedure provided excellent recovery values from 72% to 112% and satisfying RSDs between 1.4% and 17%. Considering the recovery and the difference of the volume of sample (10 mL) loaded in immunoaffinity column and the volume of redissolved sample (100 µL) for CE-MS analysis, the enrichment factor of our columns was about 100 times. The LODs ranging from 1.2 nM for GAT to 5.0 nM for FLE, were comparable or much better than those of reported reverse-phase SPE followed by HPLC-FD,42 HPLC-MS,43,44 immunoassay10 and CE-based method.14,45 It's far below the permissible MRL established by European Union. Although the immunoaffinity column was

FQs	Spiked environmental water samples $(n = 3)$								
	30 nM		15 nM		5 nM		2.5 nM		
	Recovery (%)	RSD (%)	Recovery (%)	RSD (%)	Recovery (%)	RSD (%)	Recovery (%)	RSD (%)	LOD/LOQ (nM)
GAT	102	7.1	98.2	5.1	88.7	9.4	88	13.6	1.2/4.0
LOM	111	1.4	101	7.4	108	12.1	72	16.7	2.5/8.3
ENO	99.6	2.9	103	7.9	96.7	3.2	<lod< td=""><td>ND</td><td>3.0/10</td></lod<>	ND	3.0/10
CIP	103	2.4	108	13.2	72.0	8.3	<lod< td=""><td>ND</td><td>3.0/10</td></lod<>	ND	3.0/10
OFL	107	12	99	5.9	74	15	<lod< td=""><td>ND</td><td>3.0/10</td></lod<>	ND	3.0/10
FLE	112	9.7	75	2.1	<lod< td=""><td>ND</td><td><lod< td=""><td>ND</td><td>5.0/17</td></lod<></td></lod<>	ND	<lod< td=""><td>ND</td><td>5.0/17</td></lod<>	ND	5.0/17
ENR	102	2.7	96.9	2.4	95.3	11.6	107	9.4	1.2/4.0
PEF	103	6.1	103	5.7	88.7	4.7	<lod< td=""><td>ND</td><td>2.5/8.3</td></lod<>	ND	2.5/8.3

Table 2 FQs spiked lake water analyzed by off-line coupling of IAE and CE-MS

more expensive, it could significantly improve the specificity of extraction step and be used for many times (maybe more than 50 times).

4. Conclusions

The presented off-line coupling of IAE with CE-MS was successfully utilized for FQs determination in fresh water collected from a local lake, demonstrated to be stable, sensitive and reliable. A generic polyclonal antibody was prepared and developed to construct an immunoaffinity column for simultaneous extraction and concentration of FQs with similar structure in environmental water. For further gualitative identification and quantitative determination, the hyphenation of CE-MS was applied for high-resolution separation and highsensitivity detection of eight FQs in a single analysis. The satisfactory recoveries of all FQs between 72% and 112% shows the usefulness the broad-specificity of the antibody, and the obtained LODs are in range of 1.2 nM to 5.0 nM, comparable with that of the widely recommended HPLC-FD. Moreover, the proposed analytical technique can be easily extended to the analysis of FQs in soil or food sample at trace level by adding another sample cleanup step, such as extraction, filtration or centrifugation prior to IAE.

Conflicts of interest

There are no conflicts to declare.

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