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Determination of trace antibiotics in water and milk via preconcentration and cleanup using activated carbons

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1	Determination of trace antibiotics in water and milk via
2	preconcentration and cleanup using activated carbon
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24	Abbreviations: ABX, antibiotics; SMM, sulfamonomethoxine sodium; OTC, oxytetracycline; CEF, ceftiofur hydrochloride; MAR,
25	marbofloxacin; MRLs, maximum residue limits; HPLC, high pressure liquid chromatography; SPE, solid phase extraction; RE,
26	recovery efficiency; ACs, activated carbons; CPAC, coconut powdered AC; S_{BET} , Brunauer-Emmett-Teller surface area; V_{Meso} ,
27	mesopore volume; V _{Micro} , micropore volume; PGAC, peat granular AC; WPAC, wood powder AC; MeCN, acetonitrile; TFA,
28	trifluoroacetic acid; MeOH, methanol; EtOH, ethanol; NH4OH, ammonia solution; EDTA, ethylenediaminetetraacetic acid
29	disodium salt hydrate; t-BuOH, t-butanol; n-PrOH, n-propanol; MW, molecular weight; MV, molar volume; LogKow, octanol-
30 24	water partition coefficient; S_{Water} , solubility of ABX in water; S_{EtOH} , saturated mole fraction solubility of ABX in EtOH; pK_a ,
32	ussociation constant; NHA, number of H-bond acceptors; NHD, number of H-bond donors; Kefs., references; <i>Bp</i> , boiling point; c dielectric constant; EDA, $\pi_{-\pi}$ electron donor-acceptor; TC, tetracycline; LODs, limit of detections; LOOs, limit of quantitations;
33	LRs, linear ranges: RSD, relative standard deviation: FA, formic acid: DCM, methylene chloride: MSPD, magnetic solid phase
34	dispersion; MSPE, magnetic solid phase extraction.
35	1

36 Abstract

37 CPAC-SPE-HPLC (coconut powdered activated carbon -SPE- HPLC) has been developed 38 for the determination of antibiotic (ABX), sulfamonomethoxine sodium (SMM), 39 oxytetracycline (OTC), ceftiofur hydrochloride (CEF) and marbofloxacin (MAR), in water and 40 milk. Over 99.0% SMM and OTC were recovered from 20 mL of 0.5 µg/mL ABX solution using 10 mg-CPAC for adsorption and 2 mL of 30% NH₄OH/EtOH (1/19 v/v) for elution. 41 42 Similarly, over 99.0% CEF and MAR were recovered using 15 mg-CPAC and 2 mL of 30% 43 NH₄OH/n-PrOH (1/19 v/v). Moreover, the REs of various ABX from 5-80 mL of 0.02-2.00 44 µg/mL medicated milk containing 10 mM EDTA are ordered as follows: OTC (99.3%), SMM 45 (99.1%) > CEF (68.9%) > MAR (61.4%). No interference towards HPLC analysis were 46 observed with elution using 2 mL of 30% NH₄OH/EtOH (1/19 v/v). Furthermore, much lower 47 limit of detections (0.02 µg/mL) than the maximum residual limits from European Commission 48 $(0.075-0.100 \ \mu g/mL)$ were obtained. 49 50 Keywords: Antibiotic residues in milk; Preconcentration; Cleanup; Activated carbon; Solid phase 51 extraction. 52

53

54 **1.** Introduction

55 Antibiotics (ABX) sulfamonomethoxine sodium (SMM), oxytetracycline (OTC), ceftiofur hydrochloride (CEF) and marbofloxacin (MAR), etc. are currently frequently 56 57 used in European dairy farms, and their residues in milk are cause for serious concern in Europe. The European Commission has adopted maximum residue limits (MRLs) of 58 SMM (100 μ g/L), OTC (100 μ g/L), CEF (100 μ g/L), and MAR (75 μ g/L) in milk to 59 60 guarantee the safety of dairy products and the health of human beings [1]. Thus, the 61 development of QuEChERS (quick, easy, cheap, effective, rugged and safe) methods for determining the levels of trace ABX in milk is important for dairy farmers, milk 62 63 processors, regulatory authorities and researchers [2,3].

64 The methods for determining trace ABX in milk include chromatographic methods, microbiological approaches, immunochemical techniques and biosensors, etc. [2–5], 65 66 with high pressure liquid chromatography (HPLC) seeing extensive use. As milk 67 contains complex ingredients, such as protein, lactose, fat and inorganic ions, the 68 determination of ABX can be disturbed by milk ingredients during HPLC analysis [6]. 69 This means that sample pretreatment is the most critical procedure as it can eliminate 70 interference from milk ingredients (cleanup) and enrich ABX in magnitude (preconcentration) in order to achieve accurate ABX determination [2,7,8]. Several 71 72 pretreatment methods, including solvent extraction, solid phase extraction (SPE) and 73 centrifugal ultrafiltration, etc., have been developed [2,9,10]. SPE is an updated pretreatment method that is based on low-pressure liquid chromatography and liquid-74 solid phase equilibrium, and has been widely used to separate target analytes from 75

76 samples, preconcentrate ABX and clean samples for HPLC analyses [2,11]. Compared to solvent extraction, SPE is more convenient and efficient. However, deproteinization 77 78 and the follow-up of centrifugation, sample loading, evaporation, reconstitution and 79 filtration are still necessary [10]. If the deproteinization procedure of milk is skipped, 80 high throughput of milk in the SPE column can cause blockage, an unstable flowrate 81 and inconsistent adsorption times. In addition, either MeOH or H₂O are usually used to 82 clean up SPE columns after sample loading to further eliminate the interference of 83 impurities [5,6,11].

84 Adsorption efficiency is directly associated with the recovery efficiency (RE) of ABX when using SPE. Various adsorbents, such as molecularly imprinted polymers, 85 86 metal-organic frameworks and carbon materials, etc. have been developed for milk 87 pretreatment [2,8,12]. Of these, activated carbons (ACs) besides the low cost, high adsorption capacity and recyclability, it shows the widest availability. ACs are ideal 88 89 adsorbents as they also possess distinctive physicochemical properties, such as rich 90 mesopores, high specific surface areas and a variety of functional groups [13,14]. The 91 ABX in milk can be adsorbed onto ACs via physical interactions, such as electrostatic 92 interactions, π - π bonds, H-bonds, etc., and the loaded ABX on the ACs can then be eluted by different eluents [15]. Interestingly, mesoporous ACs belong to the group of 93 "restricted access materials". The narrow pore diameter of ACs only allows low 94 95 molecular weight ABX to enter, while high molecular weight proteins or fats cannot 96 enter mesopores [16]. The procedure of chemical deproteinization can therefore 97 probably be skipped when ACs are used for the recovery of ABX from milk.

98 Furthermore, chelates that are formed between ABX, metal ions (Ca²⁺ and Mg²⁺)
99 and protein in milk can diminish the adsorption of ABX [2,5,6]. Chelating agents,
100 including EDTA, oxalic acid and citric acid, have been added to milk to promote the
101 recovery of ABX [2,5,6,9].

102 This study aims to develop an AC-SPE-HPLC approach for the determination of 103 trace SMM, OTC, CEF and MAR in milk. Based on the physicochemical properties of 104 ABX and eluents, such as acidity and alkalinity, dissociation, polarity, H-bonds, etc., the adsorption and elution conditions of the SPE for the recovery of the model ABX 105 106 from water were optimized. Furthermore, the REs, using different ACs, of ABX at various spiked concentrations in milk were evaluated in the absence and presence of 107 108 EDTA. Finally, the proposed AC-SPE-HPLC method was verified and compared with 109 methods from other published works.

110

0 2. Materials and Methods

111 **2.1.** Materials and chemicals

112 Pasteurized commercial skimmed milk was purchased from a local branch of a 113 major supermarket (Turin, Italy) and stored at room temperature. Preliminary analyses 114 demonstrated that the purchased milk was ABX-free. The main ingredients in 100 mL of milk were carbohydrate (4.8 g), sugar (4.8 g), protein (3.4 g), saturated fat (0.5 g), 115 116 fatty acid (0.3 g) and salt (0.13 g). The medicated milk samples were prepared daily by 117 adding the stock ABX solutions into the milk. The initial pH of raw, 10 mg/L CEF-, 10 mg/L SMM-, 10 mg/L MAR-, and 10 mg/L OTC-spiked milk samples were determined 118 119 to be 6.46, 6.56, 6.50, 6.56, and 6.58, respectively.

120	Coconut powdered AC (CPAC) was proven to act as a mesoporous adsorbent in
121	our previous studies [13,15], and was used as the core adsorbent for the pretreatment
122	of milk via SPE in this study. The Brunauer-Emmett-Teller (BET) surface area (S_{BET}),
123	mesopore volume (V_{Meso}), micropore volume (V_{Micro}) and average pore size of CPAC
124	were measured to be 1952 m ² /g, 1.57 cm ³ /g, 1.76 cm ³ /g and 3.95 nm, respectively.
125	CPAC was purchased from ACEF S.P.A. Piacenza (Italy). Peat granular AC (PGAC)
126	and wood powder AC (WPAC) were provided by Merck-Sigma-Aldrich, Milan (Italy).
127	The textural properties of PGAC, WPAC and CPAC have been presented in the
128	literature [13].
129	SMM (400 mg/mL, Daimeton 40, IZO Srl), OTC (92.7 mg/mL, Oxtra MV 10,
130	Huvepharma), CEF (50 mg/mL, Ceva Santé Animale), and MAR (100 mg/mL,
131	Vetoquinol) were used as the model ABX in this work. The initial pH of 10 mg/L CEF-,
132	10 mg/L SMM-, 10 mg/L MAR-, and 10 mg/L OTC aqueous solution were determined
133	to be 5.51, 6.67, 5.63, and 6.67, respectively. Their chemical structures, major
134	physicochemical properties and toxicological parameters are shown in Table S1 and in
135	Scheme 1 (in Section 3.1.3), respectively. The information of solvents and water used
136	was listed in the supplementary material (Section S1.0).
137	2.2. Apparatus and instrument
138	The analysis of water and milk samples was performed using a range of apparatus,
139	including a high-speed refrigerated centrifuge (Allegra® 64R, Beckman Coulter, US),
140	a UV-Vis spectrometer (Cary 60 UV-vis Spectrophotometer, USA), a HPLC system

141 (Waters Corp., Milford, MA, USA) coupled with a diode array detector (UV/DAD,

142 Waters Corp., Milford, MA, USA), an automatic sampler (Waters Corp., Milford, MA, USA) and a binary HPLC pump (Waters Corp., Milford, MA, USA). An Xterra RP 18 143 144 separation column (5 µm, 150 mm×4.6 mm; Phenomenex, Torrance, CA, USA) was used with gradient elution and UV-DAD acquisition. 145

146 The maximum absorption wavelengths for the UV-Vis analysis of SMM, OTC, 147 CEF and MAR in water were found to be 273, 267, 288 and 293 nm, respectively. The 148 HPLC-DAD analysis of the model ABX was performed according to the methods 149 described in Table S2. Briefly, the Xterra RP 18 separation column was pre-equilibrated 150 for 21 min with 0.1% TFA in H₂O (Phase A) and 0.1% TFA in MeCN (Phase B). A needle installed onto an automatic sampler, used for injections, was auto-washed with 151 152 MeOH/H₂O (1/1 v/v) before each sample injection. The injection volume was set as 20 153 µL. The samples were loaded and analyzed using HPLC-DAD at room temperature, and the flowrate of the mobile phase was set at 0.8 mL/min. The total running time for 154 the analysis of the model ABX ranged from 26 to 32 min. After the analysis of a batch 155 156 of samples, the separation column was washed with H₂O (phase A) and MeCN (phase B) for 50 min. The identification of ABX was performed by comparing the UV 157 158 absorption spectra and retention times of the milk samples and standard solutions.

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2.3. Preconcentration and cleanup of milk samples

The pretreatment procedures, namely, the preconcentration and cleanup of trace 160 ABX in milk for HPLC analysis, are shown in Fig. 1. 161

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164 It is necessary to note that different amounts of EDTA had previously been added 165 to the 5-80 mL milk samples so that 10 mM EDTA was contained in each. These were 166 then shaken for 5 min to ensure that the EDTA adequately interacted with the milk 167 matrix and that the milk flowed smoothly in the SPE column [2,5,6,9].

The follow-up procedures were conducted as follows. Firstly, either 10 mg powder AC or 50 mg granular AC were placed into the barrel of a glass syringe (i.d. 1.5 cm), and glass wool was fitted above and below the AC layer to prevent the AC washing out. Secondly, the 5-80 mL medicated-milk samples were passed through the AC column continuously. As a result, trace ABX were absorbed onto the AC and the white milk flowed out from the AC column. The AC column was then flushed, using 1 mL of distilled water, to remove the small amount of milk that adhered to the AC, and the small amount of water in the AC-column was extruded out. Afterwards, 2 mL of eluent was passed through the ABX-loaded AC column to elute the adsorbed ABX and the elutant was collected in a glass vial. Furthermore, 2 mL of the elutant were concentrated, via mild N₂ purge, to 0.5 mL at room temperature. Some insoluble matter was observed and was separated by centrifugation at 26000 rpm for 1 min. Finally, the supernatant was collected and stored at 4 °C for HPLC analysis.

181 **2.4.**

2.4. Evaluation of recovery efficiency

The medicated milk samples containing 0.02, 0.10, 0.50 and 1.00 μ g/mL ABX were used for the evaluation of the *REs*. The method for preconcentration and cleanup using AC-SPE is described in Section 2.3. *RE* depended on adsorption/desorption efficiency and was calculated in accordance with Eq. (1):

186
$$RE(\%) = \frac{C_d V_d}{C_0 V_0} \times 100\%$$
 (1)

187 where, C_d (µg/mL) is the determined concentration of ABX in the concentrated elutant, 188 V_d (mL) is the volume of the concentrated elutant, C_0 (µg/mL) is the spiked 189 concentration of ABX in the milk sample, and V_0 (mL) is the volume of the milk sample 190 that passed through the AC-SPE column.

All experiments were repeated at least twice and errors are shown as the difference
between the highest measured values and the average value in parallel experiments.
When errors are not visible in the figures, they are smaller than the symbols
representing the average values.

195 **3. Results and discussion**

196 3.1. Optimization of AC-SPE for ABX recovery from water

The feasibility of using ACs as an adsorbent in SPE is the primary concern here. 197 Although ACs are undoubtedly superior absorbents, the desorption of organic 198 199 adsorbates from them is generally considered to be a considerable challenge [17,18]. It was therefore necessary to demonstrate the high RE of ABX using ACs in water. CPAC 200 201 has been proven to provide superior adsorption/desorption performance in our previous studies [13,15]. In the preliminary study, using water samples instead of milk samples 202 made the whole AC-SPE operation easier, more economical and more environmentally 203 friendly. The effects of CPAC amount, the flowrate of the water sample, the type, 204 205 volume and flowrate of the eluent and the use of various ABX were evaluated in the 206 search of optimal conditions on the REs.

207 **3.1.1.** Optimization of adsorption conditions

Both the amount of adsorbent and the flowrate of the water sample through the SPE column are critical factors that influence the *RE* [8,19]. Firstly, the *REs* of the model ABX were evaluated with various doses of CPAC (5-15 mg). 20 mL of 0.5 μ g/mL aqueous solutions of ABX were passed through the CPAC-columns at a flowrate of 1.00 mL/min (Adsorption) and 2 mL of 30% NH₄OH/EtOH (1/19 v/v) were passed through the CPAC-columns at a flowrate of 0.07 mL/min (Elution) at room temperature, and the results are presented in Fig. 2a.

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Fig. 2. Effect of CPAC amount on *REs* of ABX in water. (a) *REs* using various amount of CPAC and 30%
NH₄OH/EtOH (1/19 v/v) at room temperature (Adsorption conditions: 20 mL of 0.5 µg/mL ABX were passed
through CPAC columns at 1.00 mL/min. Elution conditions: 2 mL of 30% NH₄OH/EtOH (1/19 v/v) were passed
through the ABX-loaded columns at 0.07 mL/min); (b) *REs* of MAR and CEF using various amount of CPAC and
30% NH₄OH/n-PrOH (1/19 v/v) at room temperature (Adsorption conditions: 20 mL of 0.5 µg/mL ABX were passed
through CPAC columns at 1.00 mL/min. Elution conditions: 2 mL of 30% NH₄OH/n-PrOH (1/19 v/v) were passed
through CPAC columns at 1.00 mL/min. Elution conditions: 2 mL of 30% NH₄OH/n-PrOH (1/19 v/v) were passed
through CPAC columns at 1.00 mL/min. Elution conditions: 2 mL of 30% NH₄OH/n-PrOH (1/19 v/v) were passed
through CPAC columns at 0.07 mL/min).

224	As shown in Fig. 2a, the REs of SMM and OTC are much higher than those of
225	CEF and MAR. The reason will be discussed in Section 3.1.3. In addition, increasing
226	CPAC amount initially increases the REs, the peak REs, 99.4% (SMM), 99.2% (OTC),

62.1% (CEF) and 60.4% (MAR), are achieved with 10 mg of CPAC, and then the *REs*decrease slowly with increasing the amount of CPAC. Generally, the increased amount
of adsorbent provided more adsorption sites, which ensured that ABX was efficiently
adsorbed. However, it is difficult to elute ABX at the higher adsorbent amount (> 10
mg of CPAC). The effects of eluent type and amount are discussed in Section 3.1.2.

232 To further increase the *REs* of CEF and MAR, the effect of CPAC amount on the REs of CEF and MAR with elution using 30% NH₄OH/n-PrOH (1/19 v/v) was further 233 234 investigated. 20 mL of 0.5 µg/mL aqueous solutions of CEF and MAR were passed 235 through the 5-15 mg CPAC-columns at a flowrate of 1.00 mL/min at room temperature, and 2 mL of 30% NH₄OH/n-PrOH (1/19 v/v) were used for the elution. The results are 236 237 shown in Fig. 2b. Obviously, the REs of CEF and MAR increase with increasing of 238 CPAC amount and the highest REs reach 102.9% for CEF and 99.1% for MAR with 15 mg of CPAC. As compared with elution using 30% NH₄OH/EtOH (1/19 v/v), the REs 239 of CEF and MAR increase by 66% and 81.2% using 30% NH₄OH/n-PrOH (1/19 v/v) 240 241 as eluent, respectively. Importantly, higher amount of the adsorbent (15 mg of CPAC) 242 are required to fully catch CEF and MAR from water.

In addition, the adsorption efficiency of ABX is also dependent on the flowrate of
the water sample. For example, 20 mL of 0.5 μg/mL MAR aqueous solutions were
passed through the 10 mg CPAC-columns at various water-sample flowrates (0.30-2.00
mL/min) at room temperature, and elution with 30% NH₄OH/EtOH (1/19 v/v) or 30%
NH₄OH/n-PrOH (1/19 v/v), and the results are shown in Fig. 3.

■ 30% NH₄OH/n-PrOH (1/19 v/v) ■ 30% NH₄OH/EtOH (1/19 v/v)



Fig. 3. Effect of water-sample flowrates on the *REs* of MAR at room temperature (Adsorption conditions: 20 mL of
0.5 μg/mL MAR were passed through 10 mg-CPAC columns. Elution conditions: 2 mL of eluents were passed
through the MAR-loaded columns at 0.07 mL/min).

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As shown in Fig. 3, using 10 mg of CPAC for adsorption and 2 mL of 30% NH4OH/EtOH (1/19 v/v) for elution, no obvious effects of water-sample flowrates between 0.30-1.00 mL/min were observed, but the *REs* of MAR drops from 60.4% to 51.9% as the flowrate increases from 1.00 to 2.00 mL/min. Therefore, the higher sample flowrate results in an insufficient adsorption time and lower *REs* [19]. The similar effects of water-sample flowrates were observed with 10 mg of CPAC for adsorption and 2 mL of 30% NH4OH/n-PrOH (1/19 v/v) for elution.

The effect of water-sample flowrates on *REs* of MAR with a higher CPAC amount was further studied. With 15 mg of CPAC for adsorption and 2 mL of 30% NH₄OH/n-PrOH (1/19 v/v) for elution, as expected, the *REs* reached 99.1%, 98.1%, 97.7% and 98.2% at the sample flowrates of 0.30, 0.70, 1.00 and 2.00 mL/min, respectively. It indicates that no obvious effects of water-sample flowrate on the *REs* of MAR were observed, 15 mg of CPAC is sufficient for catching trace MAR and 30% NH₄OH/n265 PrOH (1/19 v/v) is an ideal eluent.

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3.1.2. Optimization of elution conditions

The elution conditions, e.g. the type, volume and flowrate of the eluent, determine the selectivity and efficiency of elution [5,8,20]. Acidity and alkalinity, polarity (hydrophobicity, dielectric constant and Log*K*ow value) and the H-bond interaction (Hdonor/acceptor count) of eluents are the critical factors for the elution of organics from ACs [13]. The major physicochemical properties of the eluents are listed in Table 1.

272	Table 1. Major physicochemica	l properties of eluents	(Data were adopted from	Pubchem and Drugbank databases
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Eluent	MW	Вр	LogKow	<i>pK</i> _a	NHA	NHD	3
	(g/mol)	(• <i>C</i>)					
H ₂ O	18	100.0	-	15.7	1	1	80.4
HOAc	60	117.9	-0.17	4.8	2	1	6.2
NH4OH	35	38.0	-	9.2	1	2	2.5
MeCN	41	81.6	-0.34	8.8	1	0	38.8
MeOH	32	64.7	-0.77	13.1	1	1	32.7
EtOH	46	78.2	-0.31	15.9	1	1	24.6
n-PrOH	60	97.2	0.25	16.1	1	1	20.3
t-BuOH	74	84.2	0.35	19.2	1	1	10.9
n-BuOH	74	117.8	0.88	16.1	1	1	17.5

273 Note: MW, molecular weight; Bp, boiling point; $LogK_{OW}$, octanol-water partition coefficient; pK_a , dissociation 274 constant; NHA, number of H-bond acceptors; NHD, number of H-bond donors; ε , dielectric constant.

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Referring to our previous work and according to pre-experiments, 30% NH₄OH/EtOH (1/19 v/v) was proved to be an ideal eluent for the desorption of organics from CPAC. Based on Fig. 2a, the superior *REs* of SMM and OTC (> 99%) were achieved with elution using 30% NH₄OH/EtOH (1/19 v/v), but the *REs* of CEF and MAR were around 60%. In contrast, more than 99% of CEF and MAR were recovered with 15 mg of CPAC for adsorption and 2 mL of 30% NH₄OH/n-PrOH (1/19 v/v) for elution (Fig. 2b). Therefore, CEF and MAR were selected as the model ABX to evaluate

- the role of various eluents on their *REs* [8,20]. Typically, 20 mL of 0.5 µg/mL CEF or
- 284 MAR aqueous solutions were passed through the 10 mg CPAC-columns at a water-
- sample flowrate of 1.00 mL/min at room temperature. The *REs* of CEF and MAR using
- various eluents are shown in Fig. 4.

287



Fig. 4. *REs* of CEF and MAR in water using CPAC columns and various eluents at room temperature (Adsorption
 conditions: 20 mL of 0.5 μg/mL CEF or MAR were passed through 10 mg-CPAC columns at 1.00 mL/min. Elution
 conditions: 2 mL of various eluents were passed through the ABX-loaded columns at 0.07 mL/min).

As shown in Fig. 4, remarkable different *REs* of CEF and MAR were observed with elution using various eluents. CEF is a hydrophobic acidic compound with a lower S_{water} (100 mg/L), a higher Log K_{OW} value (1.2), and a lower p K_a value (2.83). In contrast, MAR is a relatively hydrophilic acidic compound with higher S_{water} (2600 mg/L), a lower Log K_{OW} value (-0.53), and a higher p K_a value (5.38), and MAR can be dissolved in EtOH, DMSO, DMF, etc.[21].

297 Considering the different physicochemical properties of CEF and MAR and the

like-dissolves-like rule, MeOH, EtOH, n-PrOH, t-BuOH or n-BuOH with various
polarities were used to elute CEF or MAR from CPAC, respectively (Table 1). The
results show that the *REs* of CEF were 57.1%, 44.3% and 28.8% with elution using nPrOH, n-BuOH and t-BuOH, respectively, while the *REs* of MAR obtained with elution
using MeOH, EtOH and n-PrOH reached 15.2%, 29.8% and 22.0%, respectively. In
addition, the very polar aprotic or protic solvents, water and MeCN are ineffective for
eluting both of CEF and MAR.

305 The acidic and alkali eluents, HOAc and 30% NH₄OH, were evaluated next. HOAc 306 is known as a common carboxylic acid with high polarity and potential for H-bonding with the -COOH, -F or -NH₂ groups that exist in CEF or MAR molecules. NH₄OH is 307 a common alkali reagent with high polarity and potential for H-bonding, which favors 308 309 the desorption of acidic CEF and MAR. The results show that the REs obtained only reached 13.2% (CEF) and 12.7% (MAR) with elution using HOAc, as well as 25.1% 310 (CEF) and 23.8% (MAR) with elution using 30% NH₄OH, respectively. The 311 312 effectiveness of 30% NH₄OH is higher than that of HOAc due to the dissociation of -COOH in the CEF and MAR molecules under alkali conditions [22]. 313

To further improve the *REs* of CEF and MAR, acidic and alkali solvents were mixed with alcohols, such as MeOH, EtOH, n-PrOH, t-BuOH or n-BuOH to form cosolvents, so that the acidity and alkalinity, polarity and H-bond interaction of the eluents can synergistically enhance the desorption. As expected, the *REs* of CEF obtained by adding 30% NH4OH to n-PrOH, t-BuOH, and n-BuOH were significantly increased to 75.8%, 46.0% and 54.9%, respectively. Meanwhile, the *REs* of MAR obtained by 320 adding 30% NH₄OH to MeOH, EtOH and n-PrOH were extremely increased to 52.0%, 60.4% and 71%, respectively. The more effective elution of CEF and MAR by adding 321 322 30% NH₄OH to n-PrOH than other alcohols could be attributed the appropriate polarity of the 30% NH₄OH/n-PrOH [8,23,24]. In contrast, the *REs* of MAR obtained using 323 324 HOAc/MeOH (1/19 v/v) and HOAc/EtOH (1/19 v/v) were 20.5% and 24.5%, 325 respectively, due to the variation of the H-bond interactions and the -COOH dissociation of MAR [8,23,24]. In summary, Lewis bases solvents as eluents exhibit 326 higher desorption efficiency than neutral and Lewis acid solvents [23,24]. The acidity 327 328 or alkalinity, the polarity and the H-bond interactions of an eluent jointly determined the desorption efficiency and the co-solvent 30% NH₄OH/n-PrOH (1/19 v/v) was an 329 330 optimal eluent.

To further improve the desorption and to minimize the consumption of eluents, the 331 effect of eluent volume on the REs of CEF and MAR was studied. In the preliminary 332 study, 20 mL of 0.5 µg/mL MAR were passed through 10 mg-CPAC columns at 1.00 333 mL/min, and then 1-10 mL of 30% NH₄OH/EtOH (1/19 v/v) were passed through the 334 MAR-loaded columns at 0.07 mL/min. As a result, the REs of MAR reached 36.6%, 335 60.4%, 60.75, 61.0%, 61.8 and 64.3% with elution using 1, 2, 4, 6, 8 and 10 mL of the 336 eluent, respectively. The REs of MAR increased remarkably as the eluent volume 337 increased from 1 to 2 mL, but only a slight increase in REs occurred after 2 mL. 338

To further understand the effect of eluent volume on the *REs*, 30% NH₄OH/n-PrOH (1/19 v/v) was used as eluent. 20 mL of 0.5 μ g/mL MAR or CEF were passed through 15 mg-CPAC columns at 1.00 mL/min, and then 1-4 mL of the eluent were passed



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344

20

0

b

0.07

0.10

345 Fig. 5. Optimizing the elution processes of CEF and MAR from CPAC at room temperature. (a) Effect of eluent 346 volumes on the REs-of CEF and MAR (Adsorption conditions: 20 mL of 0.5 µg/mL ABX were passed through 15 347 mg-CPAC columns at 1.00 mL/min. Elution conditions: 1-4 mL of 30% NH4OH/n-PrOH (1/19 v/v) were passed 348 through the ABX-loaded columns at 0.07 mL/min); (b) Effect of eluent flowrates on the REs of MAR (Adsorption 349 conditions: 20 mL of 0.5 µg/mL MAR were passed through CPAC columns at 1.00 mL/min. Elution conditions: 2 350 mL of 30% NH₄OH/n-PrOH (1/19 v/v) were passed through the MAR-loaded columns at various flowrates).

0.15

Flowrate of eluent (mL/min)

0.40

0.50

10 mg CPAC

351	As shown in Fig. 5a, the REs of MAR and CEF increased from 56.6% to 99.1%,
352	and from 76.7% to 102.9% as the volume of 30% NH ₄ OH/n-PrOH (1/19 v/v) increased
353	from 1 to 2 mL, respectively, but no changes in the <i>REs</i> were observed after 2 mL. It is
354	similar to the effects of the 30% NH4OH/EtOH (1/19 v/v) volume on the REs of MAR

and previous study's result [25]. The results reveal that 15 mg CPAC is sufficient for
full adsorption of CEF and MAR from water, and 2 mL of 30% NH₄OH/n-PrOH (1/19
v/v) is a better eluent for eluting CEF and MAR from CPAC.

Furthermore, the desorption efficiency of ABX is also dependent on the flowrate 358 359 of eluents [26]. The REs of MAR from loaded CPAC columns were evaluated at various flowrates of 2 mL of 30% NH₄OH/n-PrOH (1/19 v/v) at room temperature, and the 360 results are shown in Fig. 5b. Using 10 or 15 mg of CPAC for adsorption and 2 mL of 361 30% NH₄OH/n-PrOH (1/19 v/v) for elution, no obvious effects of eluent flowrates 362 363 between 0.07-0.15 mL/min were observed, but REs of MAR significantly drops as the flowrate increases from 0.15 to 0.50 mL/min (Fig. 5b). Obviously, a higher eluent 364 flowrate results in an insufficient desorption time and lower *REs* [26]. Consequently, 365 366 0.15 mL/min was proven to be the optimal flowrate for eluting MAR from CPAC.

367 3.1.3. Recovery mechanism of ABX from water

The *REs* of ABX are different despite the same analysis method being used as they 368 369 all have different chemicals structures and physicochemical properties [2,5]. As shown 370 in Fig. 2a, the *REs* of SMM and OTC are much higher than those of CEF and MAR, suggesting that 10 mg-CPAC as an absorbent and 2 mL of 30% NH₄OH/EtOH (1/19 371 v/v) as an eluent are highly efficient for the recovery of trace SMM and OTC. In contrast, 372 the REs of CEF and MAR reached 102.9% and 99.1%, respectively, as 15 mg CPAC 373 and 2 mL of 30% NH₄OH/n-PrOH (1/19 v/v) were used (Fig. 2b). 374 375 Generally, the S_{BET} , V_{Meso} and V_{Meso} dominate the adsorption performance of ACs

376 [17,27,28]. As mentioned in Section 2.1, CPAC possesses a very large S_{BET} and rich

377 micro- and mesopores with a reasonable fraction of micro- and mesopores (1.12) [13]. Therefore, CPAC is a superior adsorbent toward ABX in water and milk [15]. 378 Adsorption of ABX onto ACs is a complex process as it is closely associated with the 379 physicochemical property of ACs, the molecule structure, property and geometric 380 barrier of ABX, and the adsorption conditions, etc. The critical physicochemical 381 382 properties of ABX, such as electrostatic interaction and Lewis acid-base interaction (pK_a) , hydrophobicity (S_{water} and $LogK_{OW}$), electronic coupling (the π - π electron donor-383 acceptor (EDA) interactions and H-bonding), pore filling (molar volume and molecular 384 385 size), etc., influence the adsorption affinity [27–31].

386 The major physicochemical properties, chemical structures and the molecular sizes387 of model ABX are listed in Table S1 and Scheme 1.



392 Scheme 1 Chemical structures and molecular sizes of the model ABX (The molecular sizes were calculated from the

Chem3D Program).

394 As shown in Scheme 1, SMM contains a sulfonamide group, which connects with an aniline ring and an aromatic heterocyclic group. The unionized form of SMM has 395 the smallest molecule size among the four model ABX, while CPAC has a high V_{Meso} 396 and S_{BET}, which favor the adsorption of low-sized sulfonamide ABX due to the pore-397 398 filling effect [27,31,32]. In addition, the strong electron-withdrawing sulfonamide 399 group causes the decrease of the π -electron density in aromatic rings and the 400 heterocyclic group, and hence act as π -electron acceptors. In contrast, the π -electronrich regions on the graphene surface of CPAC serve as π -electron donors. Such π - π 401 402 EDA interactions enhance the adsorption of SMM on CPAC [27,29]. Furthermore, the rich O-containing groups (4.04±0.98%) of CPAC favor the adsorption of relatively 403 hydrophilic SMM [29,30]. Finally, electrostatic interactions and H-bonding may play 404 405 additional roles [30,33].

OTC belongs to tetracycline ABX, possessing one tetracycline (TC) ring and 406 multiple hydroxyl (-OH), N,N, dimethyl (-N(CH₃)₂) and amide (-C(O)NH₂) groups on 407 408 the TC ring. The molecular size of OTC is the second smaller among the four model ABX. Similar to the above, the higher adsorption affinity of OTC onto CPAC may be 409 410 mainly attributed to the high V_{Meso} , S_{BET} of CPAC, and the pore-filling effect [18,34]. In addition, the electrostatic interaction and the π - π interactions between OTC and 411 CPAC should be involved in the adsorption of OTC [35]. The multiple -OH, -N(CH₃)₂ 412 and -C(O)NH₂ groups in the OTC molecule are electron donors and induce strong 413 conjugation with the π -electrons on aromatic rings and interact electrostatically with 414 the positively charged regions on CPAC. The H-bonding between -OH group and O-415

416 containing functional groups on CPAC is also important due to the appropriate contents
417 of O-containing groups on CPAC. Also, O-containing groups facilitate the adsorption
418 of relatively hydrophilic OTC [27,36].

Compared with the higher adsorption affinity of SMM and OTC on CPAC, the 419 420 adsorption efficiencies of CEF and MAR onto CPAC are relatively lower (Fig. 2). CEF 421 is the hydrochloride salt of ceftiofur, containing a β -lactam core structure (a cephem) coupling with a furan-carboxylic-thioester, an amide, an iminomethoxy, and a 2-amino-422 4-thiazoyl group. CEF has the largest molecular size and the highest hydrophobicity 423 424 (LogKow: 1.2) among the four model ABX. The large molecular size of CEF causes a significant size-exclusion effect or geometric barrier, reducing the affinity with CPAC 425 426 [27,31,32,34,37]. Meanwhile, its high hydrophobicity is unfavorable to the complex 427 interactions of CEF with the rich O-containing groups on the surface of CPAC. Finally, acidic groups (-COOH) in hydrophobic compounds negatively influence the adsorption 428 on ACs due to the formation of large and dense water clusters [28,38]. 429

430 MAR is one of fluoroquinolones, containing a bicyclic core structure related to the substance 4-quinolone, a fluorine atom in the chemical structure, -COOH, 3,5-431 dimethylpiperazine, carbonyl, etc. CEF has the second larger molecular size and the 432 second higher hydrophobicity ($Log K_{OW}$: -0.5) among the four model ABX. As 433 mentioned above, both larger molecular size and higher hydrophobicity are unfavorable 434 to the adsorption of MAR on CPAC. It is similar to CEF, -COOH group in the MAR 435 may reduce its affinity with CPAC [28,38]. Additionally, F-containing organics may 436 exhibit a weaker affinity with ACs [39]. 437

438	The effects of eluents on the elution of CEF and MAR from CPAC have been
439	discussed in detail in Section 3.1.2. It can be concluded that the acidity or alkalinity,
440	polarity and H-bond interactions of eluent jointly dominate the elution performance of
441	ABX from CPAC and the co-solvent 30% NH ₄ OH/n-PrOH (1/19 v/v) is an appropriate
442	eluent to elute CEF and MAR from CPAC [8,23,24]. In contrast, 30% NH ₄ OH/EtOH
443	(1/19 v/v) is appropriate to effectively elute SMM and OTC. The prominent difference
444	between these two eluents is the difference in polarities of EtOH and n-PrOH. The
445	ionized SMM at the presence of NH4OH facilitates its elution by the polar eluent of 30%
446	$NH_4OH/EtOH (1/19 v/v)$ based on the like-dissolves-like rule [29]. Similarly, OTC with
447	the strongest polarity among the four model ABX can be effectively eluted by 30%
448	NH ₄ OH/EtOH (1/19 v/v). Meanwhile, the strong H-bond interactions of 30%
449	$NH_4OH/EtOH$ enhance the elution of SMM and OTC.

450 3.2. Optimization of AC-SPE for ABX recovery from milk

451 The major challenge encountered in the analysis of ABX in milk is separating the impurities, e.g. fat, protein, etc., and increasing the concentration of trace ABX [2,5]. 452 453 Using the optimization of AC-SPE for ABX recovery from water as a base, the technique was further optimized to validate the feasibility of recovering ABX from milk, 454 and this included determining the REs of ABX using various adsorbents at different 455 456 concentrations, determining the limit of detections (LODs), the limit of quantitations (LOQs), the linear ranges (LRs) and errors. Finally, the AC-SPE process was compared 457 to other methods reported for the analysis of model ABX in milk. 458

23

459 3.2.1. Recovery of ABX using various ACs at different concentrations

460	To find an appropriate adsorbent, the REs of ABX were obtained at various
461	concentrations in different milk volumes using 50 mg PGAC, 10 mg WPAC or CPAC.
462	2 mL of 30% NH ₄ OH/EtOH (1/19 v/v) were used to elute ABX from AC columns. As
463	a result, 0.5 mL HPLC samples were prepared from 10-80 mL of 0.02-1.00 $\mu\text{g/mL}$
464	medicated milk using AC-SPE; the analytes in milk were concentrated 20-160 fold.
465	It is difficult for large-volume milk to pass through the powder AC-SPE columns
466	due to the higher viscosity. Therefore, EDTA had previously been added to the milk
467	samples (containing 10 mM EDTA) to ensure the smooth flow of the large-volume milk
468	samples through the CPAC columns [2,5], which are designated as CPAC'. The results
469	of <i>REs</i> are summarized in Table 2.

ABX	$C_{\theta} (\mu g/mL)$	$V_s(mL)$	RE (%) (mean, n=3)					
			PGAC	WPAC	CPAC	CPAC'		
SMM	0.02	80	-	-	-	99.48±1.28		
	0.10	40	4.89±0.06	47.82±0.17	98.18±1.03	99.53±0.51		
	0.50	20	4.69±0.01	45.04±2.55	94.89±1.11	98.89±0.90		
	1.00	10	4.24±0.00	40.57±2.26	94.79±0.68	98.59±0.75		
OTC	0.02	80	-	-	-	99.67±0.42		
	0.10	40	5.81±0.17	49.25±0.54	81.33±3.52	99.33±0.17		
	0.50	20	4.49±0.03	45.70±0.75	79.78±1.7	99.30±0.06		
	1.00	10	4.48±0.11	42.10±0.34	78.47±2.36	99.07±1.34		
CEF	0.02	80	-	-	-	69.13±1.40		
	0.10	40	1.16±0.02	11.81 ± 0.30	38.41±1.26	69.01±0.16		
	0.50	20	1.02 ± 0.03	8.11±0.29	36.17±0.33	68.62±0.32		
	1.00	10	0.89 ± 0.00	8.30±0.02	38.36±3.33	68.79±0.32		
MAR	0.02	80	-	-	-	61.93±1.05		
	0.10	40	1.00 ± 0.01	22.80±0.69	41.83±2.54	61.89±1.19		
	0.50	20	0.90 ± 0.09	21.47±0.19	38.65 ± 0.02	60.98±0.37		
	1.00	10	0.90±0.10	19.25±0.81	37.08±1.20	60.81±0.19		

470 Table 2. *REs* of ABX at various concentrations from milk using different ACs.

471 Adsorption conditions: 10-80 mL of medicated milk were passed through 10 mg-WPAC, CPAC and 50 mg-PGAC

472 columns at 1.00 mL/min at room temperature. Elution conditions: 2 mL of 30% NH4OH/EtOH (1/19 v/v) were

473 passed through the ABX-loaded AC columns at 0.07 mL/min at room temperature.

474 Note: *C*₀, initial concentration of ABX; *V*_s, volume of the sample; *RE*, recovery efficiency.

475

494

476 As shown in Table 2, the RE ranges of various ABX using CPAC, WPAC and PGAC are 37.6%-96.0%, 9.4%-45.6% and 0.93%-4.93%, respectively. The REs 477 positively correlate with the S_{BET} of the ACs; the increasing S_{BET} provides more 478 adsorption sites. The mean REs of ABX with CPAC at different concentrations in milk 479 are ordered as follows: SMM (96.0%) > OTC (79.9%) > MAR (39.2%) > CEF (37.6%). 480 The *REs* obtained from the milk samples are somewhat lower than those from the water 481 samples, and this may be the result of the possible chelation of ABX with milk 482 components. As for the differences in the REs of various ABX in milk, the order of REs 483 is similar to that obtained in water samples and reasons are discussed in Section 3.1.3. 484 Surprisingly, adding EDTA to the milk samples not only made the milk flow more 485 486 smoothly, but also significantly improved the REs of the model ABX. As listed on 487 CPAC' in Table 2, the mean REs with EDTA of SMM, OTC, CEF and MAR at different concentrations are 99.1%, 99.3%, 68.9% and 61.4%, respectively. These are 3.3%, 488 24.4%, 56.7%, and 83.0% higher, respectively, than those obtained in the absence of 489 EDTA. This remarkable improvement in REs may be attributed to the effective 490 separation of ABX/proteins/metal ion (e.g. Ca²⁺) chelates by EDTA [9]. Thus, higher 491 492 *REs* can be obtained by CPAC-SPE for the analysis of ABX in milk. Furthermore, 15 mg of CPAC and 30% NH₄OH/n-PrOH (1/19 v/v) have been 493

495 water samples. Aiming to increase the *REs* of CEF and MAR, 15 mg of CPAC and 30%

proven to be effective adsorbent dose and eluent for the recovery of CEF and MAR in

NH4OH/n-PrOH (1/19 v/v) were used as adsorbent and eluent. Unfortunately, higher *REs* cannot be achieved after unremitting efforts due to the interference of milk
components. Hence it is necessary to promote the recovery of CEF and MAR in milk
by using AC-SPE method in the future.

500 Besides the preconcentration of ABX from milk, the separation or removal 501 efficiency of major components in milk is another concern. Fig. 6 shows the appearance 502 of the samples during SPE process.



503 504

Fig. 6 Appearance of samples collected during the pretreatment procedure for HPLC analysis.

508 Fig. S1 illustrates a HPLC comparison of samples from blank milk, samples that

As shown in Fig. 6, most of the white milk components that remained in the AC columns were washed out using 1 mL of distilled water, and very few residual white components were further removed from the elutant via centrifugation.

509 were recovered from medicated milk by using 10 mg of CPAC for adsorption and 2 mL of 30% NH₄OH/EtOH (1/19 v/v) for elution, as well as standard ABX aqueous 510 solutions. As shown in Fig. S1, only one peak appears for the standard CEF aqueous 511 solution and this is at 12.6 min, while no peak was observed around 12.6 min for the 512 513 sample from blank milk under the same HPLC conditions. This indicates that milk 514 components did not interference with the CEF analysis. In contrast, the peak for CEF appears at around 12.6 min in the sample recovered from the CEF-spiked milk using 515 CPAC-SPE. In addition, two tiny peaks, at around 7.8 min and 11.8 min, are found, and 516 517 these probably arise from the impurities in the CEF stock solution. Similarly, the milk components did not interference in the OTC analysis, although three tiny peaks are 518 found in the sample recovered from the OTC-spiked milk. Fortunately, the peaks do not 519 520 disturb the OTC analysis. The HPLC analyses of MAR and SMM are better still as no problematic peaks are observed in the samples recovered from the MAR- and SMM-521 spiked milk. Moreover, EDTA did not appear in any HPLCs, which is consistent with a 522 previous study [7]. Overall, the interference of milk components with the HPLC 523 analysis of ABX by using CPAC-SPE can be neglected. 524

525 3.2.2. Validation of the CPAC-SPE-HPLC method

The European Commission Decision (2002/657/EC) on the performance of analytical methods and the interpretation of results was published in 2002 [40]. The CPAC-SPE-HPLC method for detecting trace ABX in milk was validated, with reference to the above Decision, by investigating LOD, LOQ, LR/R^2 (correlation coefficient), reproducibility and intraday/interday precision. Based on the MRLs of

531	SMM, OTC, CEF and MAR regulated by European Commission [1], LOD and LOQ
532	value for the model ABX were set as 0.020 $\mu g/mL.$ Therefore, 80 mL of 0.020 $\mu g/mL$
533	medicated milk containing 10 mM EDTA were preconcentrated and cleaned up via
534	adsorption with 10 mg-CPAC columns, elution with 2 mL of 30% NH ₄ OH/EtOH (1/19
535	v/v)), and purging with N_2 gas to prepare 0.5 mL of HPLC samples. Hence, the trace
536	ABX was concentrated 160-fold in the HPLC samples, and the REs of SMM, OTC,
537	CEF and MAR reached 99.48%, 99.67%, 69.13% and 61.93%, respectively.
538	In this range of 0.02-2.00 $\mu g/mL$ of ABX, the linear correlation of ABX
539	concentrations in medicated milk and HPLC samples was investigated using CPAC-
540	SPE. The high R^2 values (0.993-0.998) for the four model ABX indicate that the linear
541	correlation between ABX concentrations in medicated milk and HPLC samples is
542	superior. The reproducibility of CPAC-SPE-HPLC was investigated in six repeated
543	tests. Reproducibility is expressed as the relative standard deviation (RSD, %) and
544	shown in italics in parentheses in Table 3.

545 Table 3. Mean *REs* (%) (n = 6) of ABX in milk using CPAC-SPE-HPLC, and reproducibility (RSD, %).

$C_{ABX}(\mu g/mL)$	SMM	ОТС	CEF	MAR
0.02	99.70 (6.3)	99.93 (6.5)	69.69 (4.0)	61.69 (2.4)
0.10	99.61 (2.4)	99.61 (6.4)	69.28 (3.2)	61.72 (2.0)
0.50	99.48 (4.0)	99.76 (3.1)	68.84 (2.2)	60.58 (2.2)
1.00	99.66 (5.1)	99.23 (3.2)	68.16 (3.7)	61.02 (1.2)

546 Note: *C*_{ABX}, initial concentration of ABX in milk.

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548 As listed in Table 3, the RSDs are in the range of 2.4%-6.3%, 3.1%-6.5%, 2.2%-
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549 4.0%, and 1.2%-2.4% for SMM, OTC, CEF and MAR, respectively, demonstrating that

550 CPAC-SPE-HPLC has good determination reproducibility.

551 Furthermore, the precision of this method was investigated in terms of intraday and interday assays. The intraday/interday precision of CPAC-SPE-HPLC was evaluated 552 553 using 0.1 µg/mL and 1.0 µg/mL medicated milk, which was consecutively tested five times in the same day (n = 5) and also over three different days (n = 15). For 0.1 μ g/mL 554 555 ABX in milk, the intraday RSD values obtained ranged from 1.2% to 6.4%, whereas 556 the interday RSD values were in the range of 5.5–8.3%. For 1.0 µg/mL ABX in milk, the intraday RSD values obtained ranged from 0.5% to 5.1%, while the interday RSDs 557 558 were slightly higher, ranging from 4.5% to 7.8%.

559 3.2.3. Comparison with previous analytical methods

The developed CPAC-SPE-HPLC method was compared with previous SPE methods for the determination of trace SMM, OTC, CEF and MAR in milk. The sample-preparation procedure, amount of adsorbent, the volume of eluent, analytical instrument, *REs*, LOD and LOQ of each method are summarized in Table S3 and discussed in Section S2.0.

As seen in Table S3, the higher RE of trace SMM and OTC in milk was achieved 565 566 in this work over the previous SPE methods. The LOD and LOQ values for OTC (0.02 μ g/mL) are comparable with the previous data, but the previous methods possessed 567 lower LOD and LOQ for SMM. More importantly, the LOD and LOQ of SMM and 568 OTC (0.02 μ g/mL) obtained in this work were lower than their MRLs (0.10 μ g/mL). In 569 comparison, the REs of CEF and MAR obtained in this work are lower than those 570 reported in previous studies. Due to using higher amounts of adsorbents (200-500 mg), 571 572 the previous LOD or/and LOQ of CEF and MAR are significantly lower. In contrast, 573 the LOD and LOQ of CEF and MAR (0.02 μ g/mL) obtained with 10-15 mg CPAC in 574 this work are still lower than the MRL (0.10 or 0.075 μ g/mL).

575 Overall, the simplified sample-preparation procedure for the analysis of trace ABX 576 in milk not only favors increasing *REs*, but also reduces the consumption of solvents 577 and other materials, time and energy. AC-SPE-HPLC possesses inherent advantages, 578 including the wide availability of powder AC sources, low consumption and costs, 579 convenient operation, reliability and superior reproducibility, etc. Besides, the AC-580 SPE-HPLC method can be widely applied and easily verified and any commercial SPE 581 cartridges are not required.

582 4. Conclusions

A simple, accurate, reliable, and miniaturized CPAC-SPE-HPLC method has been 583 successfully developed for the effective preconcentration, cleanup and determination 584 585 of SMM, OTC, CEF and MAR in milk. Superior REs (>99%) of the model ABX in water were obtained in preliminary study. Subsequently, the conditions were further 586 improved to obtain higher *REs* of the four model ABX in milk. 10 mg of CPAC and 2 587 588 mL of 30% NH₄OH/EtOH (1/19 v/v) were scanned as the appropriate adsorbent and eluent. The mean REs with EDTA of SMM, OTC, CEF and MAR at different 589 concentrations are 99.1%, 99.3%, 68.9% and 61.4%, respectively. The AC-SPE-HPLC 590 method exhibits a good inter/intraday precision. To further improve AC-SPE-HPLC 591 592 analysis of ABX in milk and expand the range of applications (water and food), in future, more species of ACs, such as ACs derived from different precursors, modified 593 ACs, magnetic ACs and carbon composites, should be applied and evaluated as 594

adsorbents for SPE pretreatment.

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619 **Declaration of interest**

620 The authors declare that they have no known competing financial interests or621 personal relationships that could have appeared to influence the work reported in this622 paper.

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624 **References**

- 625 [1] Commission, E. (2010). Commission Regulation (EU) No 37/2010 of 22
 626 December 2009 on pharmacologically active substances and their classification
 627 regarding maximum residue limits in foodstuffs of animal origin. *OffJ Eur Union*,
- 628 *15*, 1–72.
- [2] Khatibi, S.A., Hamidi, S., & Siahi-Shadbad, M.R. (2021). Current trends in sample preparation by solid-phase extraction techniques for the determination of antibiotic residues in foodstuffs: a review. *Critical Reviews in Food Science and Nutrition, 61,* 3361-3382. https://doi.org/10.1080/10408398.2020.1798349.
- Kantiani, L., Farré, M., & Barceló, D. (2009). Analytical methodologies for the
 detection of β-lactam antibiotics in milk and feed samples. *TrAC Trends in Analytical Chemistry*, 28, 729–744. https://doi.org/10.1016/j.trac.2009.04.005.
- 636 [4] Greño, M., Castro-Puyana, M., Garcıa, M.Á., & Marina, M.L. (2018). Analysis
 637 of antibiotics by CE and CEC and their use as chiral selectors: An update.
 638 *Electrophoresis*, 39, 235–259. https://doi.org/10.1002/elps.201700306.
- [5] Rossi, R., Saluti, G., Moretti, S., Diamanti, I., Giusepponi, D., & Galarini, R.
 (2018). Multiclass methods for the analysis of antibiotic residues in milk by liquid
 chromatography coupled to mass spectrometry: a review. *Food Additives & Contaminants PartA*, 35, 241–257.
 https://doi.org/10.1080/19440049.2017.1393107.
- Koesukwiwat, U., Jayanta, S., & Leepipatpiboon, N. (2007). Solid-phase
 extraction for multiresidue determination of sulfonamides, tetracyclines, and
 pyrimethamine in Bovine's milk. *Journal of Chromatography A*, *1149*, 102–111.
 https://doi.org/10.1016/j.chroma.2007.02.075.
- 648 [7] Narola, B., Singh, A., Mitra, M., Santhakumar, P., & Chandrashekhar, T. (2011). A validated reverse phase HPLC method for the determination of disodium EDTA 649 650 in meropenem drug substance with UV-detection using precolumn derivatization 651 technique. Analytical chemistry Insights. 6. ACI-S5953. https://doi.org/10.4137%2FACI.S5953. 652
- [8] Xu, J.-J., An, M., Yang, R., Tan, Z., Hao, J., Cao, J., Peng, L.-Q., & Cao, W.
 (2016). Determination of tetracycline antibiotic residues in honey and milk by
 miniaturized solid phase extraction using chitosan-modified graphitized
 multiwalled carbon nanotubes. *Journal of agricultural and Food chemistry*, 64,
 2647–2654. https://doi.org/10.1021/acs.jafc.6b00748.
- Kishida, K. (2011). Simplified extraction of tetracycline antibiotics from milk
 using a centrifugal ultrafiltration device. *Food chemistry*, *126*, 687–690.
 https://doi.org/10.1016/j.foodchem.2010.11.021.
- [10] Horton, R., Randall, L., Bailey-Horne, V., Heinrich, K., Sharman, M., Brunton, 661 662 L., La Ragione, R., & Jones, J. (2015). Degradation of cefquinome in spiked milk 663 as a model for bioremediation of dairy farm waste milk containing cephalosporin residues. Journal applied microbiology, 118, 901-910. 664 ofhttps://doi.org/10.1111/jam.12765. 665

- [11] Lv, Y.-K., Wang, L.-M., Yang, L., Zhao, C.-X., & Sun, H.-W. (2012). Synthesis 666 and application of molecularly imprinted poly (methacrylic acid)-silica hybrid 667 composite material for selective solid-phase extraction and high-performance 668 liquid chromatography determination of oxytetracycline residues in milk. Journal 669 670 of Chromatography Α, 1227. 48-53. 671 https://doi.org/10.1016/j.chroma.2011.12.108.
- 672 [12] Bitas, D., & Samanidou, V. (2018). Molecularly imprinted polymers as extracting
 673 media for the chromatographic determination of antibiotics in milk. *Molecules*,
 674 23, 316. https://doi.org/10.3390/molecules23020316.
- [13] Ge, X., Wu, Z., M., Manzoli, Wu, Z., & Cravotto, G. (2020). Feasibility and the mechanism of desorption of phenolic compounds from activated carbons. *Industrial & Engineering Chemistry Research*, 59, 12223–12231.
 https://doi.org/10.1021/acs.iecr.0c01402.
- [14] Wu, Z., Liu, P., Wu, Z., & Cravotto, G. (2021). In situ Modification of Activated
 Carbons by Oleic Acid under Microwave Heating to Improve Adsorptive
 Removal of Naphthalene in Aqueous Solutions. *Processes*, 9, 391.
 https://doi.org/10.3390/pr9020391.
- 683 [15] Ge, X., Wu, Z., Manzoli, M., Bonelli, B., Mantegna, S., Kunz, W., & Cravotto,
 684 G. (2021). Adsorptive decontamination of antibiotic-spiked water and milk using
 685 commercial and modified activated carbons. *Journal of Environmental Chemical*686 *Engineering*, 9, 105544. https://doi.org/10.1016/j.jece.2021.105544.
- [16] Dmitrienko, S.G., Kochuk, E.V., Tolmacheva, V.V., Apyari, V.A., & Zolotov, Y.A. 687 (2015). Determination of the total content of some sulfonamides in milk using 688 689 solid-phase extraction coupled with off-line derivatization and 690 spectrophotometric detection. Food 188. 51-56. chemistry, 691 https://doi.org/10.1016/j.foodchem.2015.04.123.
- [17] Li, L., Quinlivan, P.A., & Knappe, D.R. (2002). Effects of activated carbon surface
 chemistry and pore structure on the adsorption of organic contaminants from
 aqueous solution. *Carbon, 40,* 2085–2100. https://doi.org/10.1016/S00086223(02)00069-6.
- [18] Hubetska, T., Kobylinska, N., & Garcıa, J.R. (2020). Efficient adsorption of
 pharmaceutical drugs from aqueous solution using a mesoporous activated carbon. *Adsorption, 26, 251–266. https://doi.org/10.1007/s10450-019-00143-0.*
- [19] Bruno, F., Curini, R., Corcia, A.D., Nazzari, M., Pallagrosi, M. (2002). An original approach to determining traces of tetracycline antibiotics in milk and eggs by solid-phase extraction and liquid chromatography/mass spectrometry. *Rapid Communications Mass Spectrometry*, 16, 1365–1376. https://doi.org/10.1002/rcm.724.
- [20] Chemat, F., Vian, M.A., & Cravotto, G. (2012). Green extraction of natural products: concept and principles. *International journal of molecular sciences*, *13*, 8615–8627. https://doi.org/10.3390/ijms13078615.
- [21] Cayam chemical company. (2018). Product information. Retrieved from https://www.caymanchem.com/pdfs/24174.pdf. Accessed 2021, December 21.
- 709 [22] Gibbons, P.M. (2014). TOPICS IN MEDICINE AND SURGERY. Journal of

- 710 Exotic Pet Medicine, 23, 21-38. http://dx.doi.org/10.1053/j.jepm.2013.11.007.
- [23] Zamora, F., Sabio, E., Román, S., González-Garcia, C.M., & Ledesma, B. (2010). 711 Modelling the adsorption of p-Nitrophenol by the Boyd method in conjunction 712 with the finite element method. Adsorption Science & Technology, 28, 671–687. 713 714 https://doi.org/10.1260%2F0263-6174.28.8-9.671.
- 715 [24] Tamon, H., Saito, T., Kishimura, M., Okazaki, M., & Toei, R. (1990). Solvent regeneration of spent activated carbon in wastewater treatment. Journal of 716 chemical engineering of Japan, 23, 426–432. https://doi.org/10.1252/jcej.23.426. 717
- [25] Seyhan Bozkurt, S., Erdogan, D., Antep, M., Tuzmen, N., & Merdivan, M. (2016). 718 719 Use of ionic liquid based chitosan as sorbent for preconcentration of fluoroquinolones in milk, egg, fish, bovine, and chicken meat samples by solid 720 721 phase extraction prior to HPLC determination. Journal of Liquid Chromatography 722 Å Related Technologies, 39, 21–29. 723
 - https://doi.org/10.1080/10826076.2015.1116010.
- [26] El-Shahat, M., Burham, N., & Azeem, S.A. (2010). Flow injection analysis-solid 724 phase extraction (FIA-SPE) method for preconcentration and determination of 725 726 trace amounts of penicillins using methylene blue grafted polyurethane foam. 727 Journal ofhazardous materials. 177. 1054-1060. 728 https://doi.org/10.1016/j.jhazmat.2010.01.027.
- 729 [27] Ji, L., Liu, F., Xu, Z., Zheng, S., & Zhu, D. (2010). Adsorption of pharmaceutical antibiotics on template-synthesized ordered micro-and mesoporous carbons. 730 731 Environmental science & technology, 44, 3116-3122. https://doi.org/10.1021/es903716s. 732
- [28] Min, P. (2020). Biochar adsorption of antibiotics and its implications to 733 remediation of contaminated soil. Water, Air, & Soil Pollution, 23, 11-15. 734 735 https://doi.org/10.1007/s11270-020-04551-9.
- 736 [29] Ji, L., Chen, W., Zheng, S., Xu, Z., & Zhu, D. (2009). Adsorption of sulfonamide antibiotics to multiwalled carbon nanotubes. Langmuir, 25, 11608-11613. 737 738 https://doi.org/10.1021/la9015838.
- [30] Ahmed, M.B., Zhou, J.L., Ngo, H.H., & Guo, W. (2015). Adsorptive removal of 739 740 antibiotics from water and wastewater: progress and challenges. Science of the Total Environment, 532, 112–126. https://doi.org/10.1016/j.scitotenv.2015.05.130. 741
- [31] Li, C., Zhu, X., He, H., Fang, Y., Dong, H., Lü, J., Li, J., & Li, Y. (2019). 742 743 Adsorption of two antibiotics on biochar prepared in air-containing atmosphere: 744 influence of biochar porosity and molecular size of antibiotics. Journal of 745 Molecular Liquids, 274, 353–361. https://doi.org/10.1016/j.molliq.2018.10.142.
- 746 [32] Ji, L., Wan, Y., Zheng, S., & Zhu, D. (2011). Adsorption of tetracycline and 747 sulfamethoxazole on crop residue-derived ashes: implication for the relative importance of black carbon to soil sorption. Environmental Science & Technology, 748 749 45, 5580–5586. https://doi.org/10.1021/es200483b.
- 750 [33] Xiang, Y., Xu, Z., Wei, Y., Zhou, Y., Yang, X., Yang, Y., Yang, J., Zhang, J., Luo, 751 L., & Zhou, Z. (2019). Carbon-based materials as adsorbent for antibiotics 752 mechanisms and influencing factors. Journal of environmental removal: management, 237, 128–138. https://doi.org/10.1016/j.jenvman.2019.02.068. 753

- [34] Li, M., Zhao, Z., Wu, X., Zhou, W., & Zhu, L. (2017). Impact of mineral components in cow manure biochars on the adsorption and competitive adsorption of oxytetracycline and carbaryl. *RSC advances*, 7, 2127–2136. https://doi.org/10.1039/C6RA26534K.
- [35] Gao, M., Zhang, Y., Gong, X., Song, Z., & Guo, Z. (2018). Removal mechanism
 of di-n-butyl phthalate and oxytetracycline from aqueous solutions by nanomanganese dioxide modified biochar. *Environmental Science and Pollution Research*, 25, 7796–7807. https://doi.org/10.1007/s11356-017-1089-5.
- [36] Ma, X., & Agarwal, S. (2016). Adsorption of emerging ionizable contaminants on
 carbon nanotubes: advancements and challenges. *Molecules*, 21, 628.
 https://doi.org/10.3390/molecules21050628.
- [37] Huang, S.-C., Chung, T.-W., & Wu, H.-T. (2021). Effects of Molecular Properties
 on Adsorption of Six-Carbon VOCs by Activated Carbon in a Fixed Adsorber. *ACS omega*, 6, 5825–5835. https://doi.org/10.1021/acsomega.0c06260.
- [38] Franz, M., Arafat, H.A., & Pinto, N.G. (2000). Effect of chemical surface
 heterogeneity on the adsorption mechanism of dissolved aromatics on activated
 carbon. *Carbon*, *38*, 1807–1819. https://doi.org/10.1016/S0008-6223(00)00012-9.
- [39] Mezzari, I.A. (2006). Predicting the adsorption capacity of activated carbon for
 organic contaminants from fundamental adsorbent and adsorbate properties.
 (Master thesis) North Carolina State University, Raleigh.
- [40] Commission, E. (2002). Commission Decision 2002/657/EC of 12 August 2002
 implementing Council Directive 96/23/EC concerning the performance of
 analytical methods and the interpretation of results. *Official Journal of European Communities*, 50, 8–36.

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Determination of trace antibiotics in water and milk via

preconcentration and cleanup using activated carbon

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Abbreviations: ABX, antibiotics; SMM, sulfamonomethoxine sodium; OTC, oxytetracycline; CEF, ceftiofur hydrochloride; MAR, marbofloxacin; MRLs, maximum residue limits; HPLC, high pressure liquid chromatography; SPE, solid phase extraction; RE, recovery efficiency; ACs, activated carbons; CPAC, coconut powdered AC; S_{BET}, Brunauer-Emmett-Teller surface area; V_{Meso}, mesopore volume; V_{Micro}, micropore volume; PGAC, peat granular AC; WPAC, wood powder AC; MeCN, acetonitrile; TFA, trifluoroacetic acid; MeOH, methanol; EtOH, ethanol; NH₄OH, ammonia solution; EDTA, ethylenediaminetetraacetic acid disodium salt hydrate; t-BuOH, t-butanol; n-PrOH, n-propanol; MW, molecular weight; MV, molar volume; Log*K*_{OW}, octanolwater partition coefficient; S_{Water}, solubility of ABX in water; S_{EtOH}, saturated mole fraction solubility of ABX in EtOH; p*K*_a, dissociation constant; NHA, number of H-bond acceptors; NHD, number of H-bond donors; Refs., references; *Bp*, boiling point; ε , dielectric constant; EDA, π - π electron donor-acceptor; TC, tetracycline; LODs, limit of detections; LOQs, limit of quantitations; LRs, linear ranges; RSD, relative standard deviation; FA, formic acid; DCM, methylene chloride; MSPD, magnetic solid phase dispersion; MSPE, magnetic solid phase extraction.

S1.0 The information of solvents and water

Acetonitrile (MeCN, \geq 99.9%), trifluoroacetic acid (TFA, \geq 99%) and methanol (MeOH, 99.9%) were obtained from Sigma-Aldrich (France). Ethanol (EtOH, \geq 99.8%) was provided by Sigma-Aldrich (UK). Acetic acid (HOAc, 100%) was purchased from VWR International (European Commission). The ammonia solution (30% NH4OH) was provided by CARLO ERBA Reagents S.r.l. (Italy). Ethylenediaminetetraacetic acid disodium salt hydrate (EDTA, >99%), n-butanol (n-BuOH, >98%), t-butanol (t-BuOH, 99%), and n-propanol (n-PrOH, 99.8%) were purchased from Alfa Aesar (Thermo Fisher Scientific, Germany).

Milli-Q water was obtained from a Milli-Q Reference A+System (Merck Millipore, Darmstadt, Germany) and used for the preparation of HPLC mobile phases. Deionized water (conductivity $\leq 2 \ \mu$ S/cm) was used to prepare standard solutions of ABX, which were refrigerated at 4 °C.

S2.0 Comparison with previous analytical methods

All of the samples prepared using SPE have undergone analysis with HPLC, LC-MS or -MS/MS, or UPLC-MS in the past. The present protocol exhibited higher RE than the previous SPE methods for the analysis of trace SMM in milk [45,46], as can be seen in Table S3. The previous methods possessed lower LOD and LOQ since high amounts of adsorbents (80-500 mg) were used. In this work, only 10 mg of CPAC was used as the adsorbent and the LOD and LOQ values of SMM (0.02 µg/mL) are also lower than the MRL (0.10 µg/mL). In addition, toxic solvents, such as DCM, MeCN and MeOH, and chemical deproteinization were not required in this study. Similarly, we achieved a higher *RE* for OTC here than the two previous SPE methods [11,47]. The LOD and LOQ values for OTC (0.02 μ g/mL) are comparable with the previous data and are also lower than the MRL (0.10 µg/mL), although less CPAC (10 mg) was used as an adsorbent here, as 50 mg of adsorbent was used in the previous method. In addition, chemical deproteinization and redissolution were not required in this study. In terms of toxicity, the use of EtOH as an eluent in this study is better than the MeOH that was used in other studies.

The *RE* of CEF (68.9%) that was obtained using a 10 mg-CPAC column in this work is comparable with that (69.0%) obtained using a 500 mg-Carbograph 4 cartridge-SPE in a previous study [48]. However, the previously reported LOQ is significantly lower than the one in that study since very different amounts of adsorbents were used. Compared with the *RE* (97.0%) obtained in another study using 200 mg-Oasis HLB-MSPD, the *RE* of CEF in this work is significantly lower, but the LOD and LOQ of

CEF (0.02 µg/mL) are comparable with the previous data, and are also lower than the MRL (0.10 µg/mL), although less CPAC (10 mg) was used as the adsorbent [49]. A further benefit is that toxic solvents, such as DCM, MeCN and MeOH, and the redissolution of analytes are not required in this study. The *RE* of MAR (61.4%) that was obtained using a 10 mg-CPAC column in this work is considerably lower than those (82.5%-106.5%) obtained in three previous studies. Moreover, the previous methods had lower LOD and LOQ, but 0.020 µg/mL of LOD or LOQ in this work is still sufficient to meet the requirements for the MRL of MAR (0.075 µg/mL). However, a remarkably lower amount of adsorbent (10 mg of CPAC) was used in this study than the 200-500 mg of adsorbents used in the other studies [25,50,51]. Furthermore, the toxic solvent MeOH and chemical deproteinization were not required in this study.





Fig. S1. Comparison of HPLCs from various samples. (a) sample from blank milk, (b) sample recovered from medicated milk by using 10 mg of CPAC for adsorption and 2 mL of 30% NH₄OH/EtOH (1/19 v/v) for elution, (c) standard ABX aqueous solutions (50 µg/mL MAR, 100 µg/mL CEF, OTC or SMM).

Table S1

Table S1. Major physicochemical properties and toxicological parameters of the model ABX (Part of the data was adopted from the Pubchem and Drugbank databases).

ABX	CAS	MW	MV	LogKow	Swater	Setoh	<i>pK</i> _a	NHA	NHD	MRLs	LD50	Refs.
		(g/mol)	(cm³/mol)		(mg/L)	(10 ³)				(µg/L)	(mg/kg)	
SMM	38006-	302	213.3	-0.8	10000	-	6.33	8	2	100	5620	[15,41]
	08-5										Rat oral	
OTC	2058-	460	270.3	-0.9	300	^a 12.0	3.27	11	8	100	>2000	[15,42,43]
	46-0										Colinus	
											virginianus	
											oral	
CEF	103980-	560	290.9	1.2	100	23.3	2.83	12	4	100	-	[15]
	44-5											
MAR	115550-	362	226.4	-0.5	2600	116.6	5.38	8	1	75	>0.002	[15,44]
	35-1										Mouse	
											oral	

 $^{\rm a}$ The unit is mg/L; $^{\rm b}$ the unit is mg/L/4 hr.

Note: *MW*, molecular weight; *MV*, molar volume; *LogKow*, Octanol-water partition coefficient; *Swater*, solubility in water; *SEtOH*, saturated mole fraction solubility of ABX in EtOH, calculated according to the literature [44]; *pKa*, dissociation constant; *NHA*, number of H-bond acceptors; *NHD*, number of H-bond donors; *MRLs*, maximum residue limits; *LD*₅₀, amount of a material, given all at once, which causes the death of 50% (one half) of a group of test animals; *Refs.*, references.

Table S2

Table S2. The HPLC conditions for the determination of the model ABX.	
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ABX	Wavelength (nm)	Retention time (min)	Running time (min)	Mobile phase			
				Phase A (%)	Phase B (%)		
				0.1%TFA in H ₂ O	0.1%TFA in MeCN		
SMM	273	25.483	0	99	1		
			25	0	100		
			32	0	100		
OTC	267	13.313	0	98	2		
			25	0	100		
			26	0	100		
CEF	265	12.510	0	98	2		
			5	70	30		
			20	0	100		
			26	0	100		
MAR	298	14.018	0	98	2		
			25	0	100		
			26	0	100		

Table S3

Table S3. Comparison of the proposed CPAC-SPE-HPLC method with previous SPE methods for the determination of the model ABX in milk.

ABX	Sample-preparation procedures	Eluent	Analytical	RE	LOD	LOQ	Refs.
			instrument	(%)	(µg/mL)	(µg/mL)	
SMM	Agitation, 500 mg-carbograph 4 cartridge-	1.5 mL MeOH-	LC-ES	97.3	^a 0.001	^a 0.003	[45]
	SPE, redissolution, filtration.	6 mL DCM/ MeOH (80/20	-MS				
		v/v) acidified with 10 mM					
		TFA continuously					
	Deproteinization, 80 mg-magnetic	1.0 mL MeCN containing	HPLC	86.8	0.00008	0.00025	[46]
	graphene oxide nanocomposite -MSPE,	5 % NH4OH (v/v)					
	evaporation, redissolution.						
	10 mg-CPAC-MSPE, concentration by N_2 ,	2 mL 30% NH ₄ OH/EtOH	HPLC	^b 99.1	0.02	0.02	This
	centrifugation.	(1/19 v/v)					study
OTC	Deproteinization, desolvation,	2 mL MeOH/ HOAc (60/40	LC-MS/MS	^b 99.2	0.014	-	[11]
	redissolution, filtration, 50 mg-molecularly	v/v)					
	imprinted poly (methacrylic acid) -silica						
	hybrid composite -SPE.						
	Deproteinization, C ₁₈ -SPE, constant	0.75 mL MeOH	UPLC-	^b 98.8	-	-	[47]
	volume.		MS/MS				
	10 mg CPAC-SPE, concentration by N_2 ,	2 mL 30% NH ₄ OH/EtOH	HPLC	^b 99.3	0.02	0.02	This
	centrifugation.	(1/19 v/v)					study
CEF	500 mg-carbograph 4 cartridge-SPE,	6 mL DCM/ MeOH (80/20	HPLC	69.0	-	0.001	[48]
	desolvation by N_2 , redissolution, filtration.	v/v) acidified with 50 mM					
		FA					
	200 mg-Oasis HLB- MSPD, evaporation to	1 mL MeOH –2 mL MeCN	HPLC	97.0	^a 0.0118	^a 0.0357	[49]
	dryness, redissolution.	continuously					
	10 mg-CPAC-SPE, concentration by N_2 ,	2 mL 30% NH ₄ OH/EtOH	HPLC	^b 68.9	0.02	0.02	This
	centrifugation.	(1/19 v/v)					study
MAR	Centrifugation, PLRP-cartridges-SPE.	$9~mLH_2O/MeOH$ ($80/20$	HPLC	106.5	0.0001	-	[50]
		v/v) plus 0.5% FA (1.5 mL)					
	Deproteinization, filtration, redissolution,	10 mL MeOH containing	LC-MS	^b 100.5	0.00009	0.00031	[51]
	centrifugation, 500 mg-Oasis HLB	1.5% (w/v) HOAc					
	cartridges-SPE, evaporation to dryness,						
	redissolution.						
	Deproteinization, 200 mg-ionic liquid	5 mL 20% NH3 (v/v MeOH)	LC-MS/MS	82.5	0.00423	-	[25]
	based chitosan-SPE, desolvation by N2,						
	constant volume.						
	10 mg-CPAC-SPE, concentration by N2,	2 mL 30% NH ₄ OH/EtOH	HPLC	^b 61.4	0.02	0.02	This
	centrifugation.	(1/19 v/v)					study

^a The unit: ppm; ^b with EDTA in medicated milk.

Note: *RE*, recovery efficiency; *LODs*, limit of detections; *LOQs*, limit of quantitations; *Refs*, references; *FA*, Formic acid; *DCM*, Methylene chloride; *MSPD*, Magnetic solid phase dispersion; *MSPE*, Magnetic solid phase extraction.

References

- [11] Lv, Y.-K., Wang, L.-M., Yang, L., Zhao, C.-X., & Sun, H.-W. (2012). Synthesis and application of molecularly imprinted poly (methacrylic acid)–silica hybrid composite material for selective solid-phase extraction and high-performance liquid chromatography determination of oxytetracycline residues in milk. *Journal* of Chromatography A, 1227, 48–53. https://doi.org/10.1016/j.chroma.2011.12.108.
- [15] Ge, X., Wu, Z., Manzoli, M., Bonelli, B., Mantegna, S., Kunz, W., & Cravotto, G. (2021). Adsorptive decontamination of antibiotic-spiked water and milk using commercial and modified activated carbons. *Journal of Environmental Chemical Engineering*, 9, 105544. https://doi.org/10.1016/j.jece.2021.105544.
- [25] Seyhan Bozkurt, S., Erdogan, D., Antep, M., Tuzmen, N., & Merdivan, M. (2016). Use of ionic liquid based chitosan as sorbent for preconcentration of fluoroquinolones in milk, egg, fish, bovine, and chicken meat samples by solid phase extraction prior to HPLC determination. *Journal of Liquid Chromatography* & *Related Technologies*, 39, 21–29. https://doi.org/10.1080/10826076.2015.1116010.
- [41] Fu, L., Huang, T., Wang, S., Wang, X., Su, L., Li, C., & Zhao, Y. (2017). Toxicity of 13 different antibiotics towards freshwater green algae Pseudokirchneriella subcapitata and their modes of action. *Chemosphere*, 168, 217–222. https://doi.org/10.1016/j.chemosphere.2016.10.043.
- [42] O'Neil, M.J., Smith, A., Heckelman, P.E., & Budavari, S. (2001). The merck index-An encyclopedia of chemicals, drugs, and biologicals. whitehouse station. NJ: Merck and Co, *Inc.*, 767, 4342.
- [43] Yalkowsky, S., & He, Y. (2003). An extensive compilation of aqueous solubility data for organic compounds extracted from the AQUASOL database. Handbook Aqueous Solubility Data (p 377). 377.
- [44] Wu, Z., Li, W., Yu, P., Fan, X., Sun, H., Zhao, H., & Zhang, Y. (2019). Measurement and Correlation of Solubility of Marbofloxacin in 12 Pure Solvents from 283.15 to 328.15 K. *Journal of Chemical & Engineering Data*, 64, 5275– 5281. https://doi.org/10.1021/acs.jced.9b00490.
- [45] Cavaliere, C., Curini, R., Di Corcia, A., Nazzari, M., & Samperi, R. (2003). A simple and sensitive liquid chromatography- mass spectrometry confirmatory method for analyzing sulfonamide antibacterials in milk and egg. *Journal of* agricultural Food chemistry, 51, 558–566. https://doi.org/10.1021/jf020834w.
- [46] Wang, Y., Liu, L., Xiao, C., Chen, L., Yang, P., Liu, Q., Wang, J., & Liu, X. (2016). Rapid determination of trace sulfonamides in milk by graphene oxide-based magnetic solid phase extraction coupled with HPLC–MS/MS. *Food Analytical Methods*, 9, 2521–2530. https://doi.org/10.1007/s12161-016-0433-6.
- [47] Mei-Ratliff, Y. (2012). Determination of the antibiotic oxytetracycline in commercial milk by solid-phase extraction: a high-performance liquid chromatography (HPLC) experiment for quantitative instrumental analysis. *Journal of Chemical Education*, 89, 656–659. https://doi.org/10.1021/ed900065y.
- [48] Bruno, F., Curini, R., Corcia, A.D., Nazzari, M., & Samperi, R. (2001). Solid-

phase extraction followed by liquid chromatography- mass spectrometry for trace determination of β -lactam antibiotics in bovine milk. *Journal of agricultural and Food chemistry*, 49, 3463–3470. https://doi.org/10.1021/jf010046r.

- [49] Karageorgou, E.G., Samanidou, V.F., & Papadoyannis, I.N. (2012). Ultrasoundassisted matrix solid phase dispersive extraction for the simultaneous analysis of β-lactams (four penicillins and eight cephalosporins) in milk by high performance liquid chromatography with photodiode array detection. *Journal of Separation Science*, 35, 2599–2607. https://doi.org/10.1002/jssc.201200514.
- [50] Kantiani, L., Farré, M., & Barceló, D. (2011). Rapid residue analysis of fluoroquinolones in raw bovine milk by online solid phase extraction followed by liquid chromatography coupled to tandem mass spectrometry. *Journal of Chromatography* A, 1218, 9019–9027. https://doi.org/10.1016/j.chroma.2011.09.079.
- [51] Herrera-Herrera, A.V., Hernández-Borges, J., Rodr*i*guez-Delgado, M.A., Herrero, M., & Cifuentes, A. (2011). Determination of quinolone residues in infant and young children powdered milk combining solid-phase extraction and ultra-performance liquid chromatography–tandem mass spectrometry. *Journal of Chromatography* A, 1218, 7608–7614. https://doi.org/10.1016/j.chroma.2011.05.066.

Graphical abstract



Simple preconcentration and cleanup of trace antibiotics in milk using activated carbon (AC)based solid phase extraction for HPLC analysis.