

Journal of the Serbian Chemical Society

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J. Serb. Chem. Soc. 00(0) S1-S4 (2025)

## SUPPLEMENTARY MATERIAL TO

# Amino-starch derivates for adsorption of specific pharmaceuticals and pesticides in contaminated water: Examination in both spiked and real water samples

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EXPERIMENTAL

Materials and chemicals

The following materials and chemicals were used to prepare the starch samples: potato starch (loss on drying at 105 °C <10%, sulfured ash <0.5%, SuperLab, Serbia), melamine (2,4,6-triamino-1,3,5-triazine, 99%, Thermo Fisher), L(-)-cysteine ((R)-2-amino-3-mercaptopropionic acid,  $\geq$ 97%, Thermo Fisher), L-histidine ((S)-2-amino-3-(4-imidazolyl)propionic acid,  $\geq$ 98.5%, Carl Roth), bentonite clay (nanoclay, Sigma-Aldrich) and diatomaceous earth (SiO<sub>2</sub> 95%, Sigma-Aldrich).

Pharmaceuticals and pesticides selected for the study were the most commonly used and frequently detected in the investigated area (Table 1). High purity (> 95%) analytical standards of four chosen pharmaceuticals: erythromycin, lorazepam, diazepam, and clopidogrel were provided by national pharmaceutical companies (Hemofarm, STADA Group, Vršac, Serbia, and Zorka-Pharma, Šabac, Serbia). The analytical standards of four selected pesticides: atrazine, propazine, malathion, and tebufenozide were supplied from Riedel-de Haën (Seelze, Germany).

The stock standard solutions were prepared in methanol at the concentration of 100  $\mu$ g mL<sup>-1</sup>. The working standard solutions were prepared by mixing the appropriate amounts of the stock standard solutions and diluting them with methanol. All solutions were preserved at -4 °C. All solvents used were HPLC grade from J.T. Baker (Center Valley, US) or Sigma-Aldrich (St. Louis, US), and all reagents were of analytical grade. Deionized water was obtained by passing the distilled water through a GenPure ultrapure water system (TKA, Niederelbert, Germany).





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## TABLE S-I. Labels of samples

Modific	Biocomposite abbreviation	
	+ Melamine	= SM
	+ Cysteine	= SC
Native starch	+ Histidine	= SH
	+ Clay (bentonite)	= S-clay
	+ Diatomeuse earth	= S-d.e
Starah Malamina (SM)	+ Clay (bentonite)	= SM-clay
Starch-Melannine (SM)	+ Diatomeuse earth	= SM-d.e
Starch Custoine (SC)	+ Clay (bentonite)	= SC-clay
Starch-Cysterne (SC)	+ Diatomeuse earth	= SC-d.e
Starah Histidina (SU)	+ Clay (bentonite)	= SH–clay
Staten-ristidine (SH)	+ Diatomeuse earth	= SH-d.e

### Characterization methods

Fourier-transform infrared spectroscopy (FTIR) was performed using a Nicolet iS10 spectrometer (Thermo Scientific) in the attenuated total reflectance (ATR) mode with a single bounce 45 °F Golden Gate ATR accessory with a diamond crystal, and DTGS detector. FTIR spectra were obtained at 4 cm<sup>-1</sup> resolution with ATR correction. The FTIR spectrometer was equipped with OMNIC software and the spectra were recorded in the wavelength range from 2.5  $\mu$ m to 20  $\mu$ m (i.e., 4000-500 cm<sup>-1</sup>).

The morphology of samples was examined using a scanning electron microscope (SEM) (type of instrument – FE-SEM, TESCAN Mira3 XMU) operating at 10 kV. Before analysis, samples were coated with gold to reduce the charging effect and improve the image quality.

X-ray diffraction (XRD) was performed using an Ultima IV Rigaku diffractometer, equipped with  $CuK\alpha 1,2$  radiations, using a generator voltage (40.0 kV) and a generator current (40.0 mA). The range of 5–40°  $2\theta$  was used for all powders in a continuous scan mode with a scanning step size of 0.02° and at a scan rate of 2° min<sup>-1</sup>, using D/TeX Ultra high-speed detector. A monocrystalline silicon sample carrier for sample preparation was used.

### Adsorption performance and regeneration studies

The separation of pesticides and pharmaceuticals was conducted using a Dionex UltiMate 3000® LC system (Thermo Scientific, USA). For detection and quantification of pesticides and pharmaceuticals, LTQ XL (Thermo Scientific, USA) mass spectrometer was used with an electrospray ion source and linear ion trap mass analyzer. The gradient of the mobile phase consisting of methanol (A), water (B), and 10 % acetic acid (C) is shown in Table S-I. Selected reaction monitoring (SRM) chromatograms of investigated pharmaceuticals and pesticides are given in Fig. S-1, and LC/MS-MS quantification parameters are presented in Table S-II.



#### SUPPLEMENTARY MATERIAL



	TABLE S	-II. Gra	dient and	flow	rate of	the	mobile	phase
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Time	Flow rate	Co	ntent, <sup>6</sup>	%
(min)	$(\text{cm}^3 \text{min}^{-1})$	А	В	С
0	0.5	49	50	1
0	0.5	49	50	1
15.00	1	0	100	0
18.00	1	0	100	0
18.01	0.5	49	50	1
23.00	0.5	49	50	1

TABLE S-III. LC/MS and MS<sup>n</sup> optimized parameters for identification of the selected pharmaceuticals and pesticides

Pollutant	Retention time, min	m/z	Collision energy	m/z
		Precursor ion	a. u.*	Product ion
Erythromycin	4.72	734.1	28	576.1
Lorazepam	6.49	321.0	32	302.9
Diazepam	8.23	285.2	40	257.2
Clopidogrel	10.72	321.9	28	211.8
Atrazine	6.50	216.0	38	174.0
Propazine	7.91	230.0	36	188.0
Malathion	8.62	331.0	28	284.7
Tebufenozide	10.01	375.0	34	225.0

\* arbitrary units defined by LCQ system



Fig. S-1. SRM chromatograms of the selected pharmaceuticals and pesticides

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	Pollutant									
	Erytrom.	Atraz.	Loraz.	Propaz.	Diazep.	Malath.	Tebuf.	Chlop.		
Sample			A	dsorption e	efficiency,	%				
S	33.44	12.64	21.34	19.71	17.86	18.73	19.15	19.07		
SM	87.5	29.39	41.32	39.41	63.21	39.46	35.47	39.62		
SC	100	30.93	51.64	50.27	68.51	61.75	77.52	78.62		
SH	100	33.36	51.44	58.39	70.92	69.51	87.06	82.56		
Clay	32.12	14.53	16.78	17.89	24.66	28.97	32.85	35.46		
S-clay	59.6	22.77	26.69	34.65	34.49	51.45	46.66	49.11		
SM-clay	94	28.72	35.72	36.9	52.57	65.45	70.66	75.67		
SC-clay	100	30.05	43.67	43.75	53.19	78.58	80.54	82.57		
SH–clay	100	32.58	50.78	44.91	55.97	81.53	82.54	86.86		
d.e	35.58	14.76	22.72	22.21	25.37	27.85	24.62	26.29		
S-d.e	83	29.79	34.45	29.27	28.27	38.49	47.44	44.19		
SM-d.e	94.82	38.38	37.22	40.27	41.99	56.7	66.69	71.51		
SC-d.e	98	44.02	39.33	39.74	41.6	59.03	72.75	75.97		
SH-d.e	98.21	45.04	41.41	28.39	45.24	62.89	79.49	86.03		

TABLE S-IV. Adsorption efficiency of tested materials for the removal of selected pharmaceuticals and pesticides

'	TABLE	S-V.	Adsorpt	ion e	fficiency	of	tested	materials	for	the	removal	of	selected
1	pharmace	euticals	and pes	ticides	from rea	l wa	ter sam	ples					

		Pollutant									
		Erytrom.	Atraz.	Loraz.	Propaz.	Diazep.	Malath.	Tebuf.	Chlop.		
	Sample		А	dsorption	n efficiency	y, % (distil	led water)				
	SM	87.5	29.39	41.32	39.41	63.21	39.46	35.47	39.62		
	SC	100	30.93	51.64	50.27	68.51	61.75	77.52	78.62		
	SH	100	33.36	51.45	58.39	70.92	69.51	87.07	82.56		
			A	Adsorptio	n efficienc	y, % (surfa	ace water)				
	SM	96.94	12.39	21.83	19.45	63.18	91.28	80.19	92.53		
	SC	92.29	14.48	30.04	16.34	70.76	89.88	77.09	90.52		
	SH	69.22	32.59	29.68	32.62	32.89	76.67	83.33	82.18		
			Adsor	ption effi	ciency, %	(groundwa	ater)				
	SM	96.13	14.62	18.28	18.4	77.11	95.29	81.75	88.08		
	SC	91.26	18.39	32.87	22.05	76.06	84.92	74.99	83.99		
	SH	64.31	31.98	24.55	32.31	22.45	83.71	82.2	82.11		
			Adso	rption eff	ficiency, %	b (wastewa	ter)				
•	SM	98.7	48.05	62.19	69.99	79.57	96.6	90.86	95.15		
	SC	94.05	57.98	72.31	25.91	73.65	91.59	89.79	92.49		
	SH	53.78	38.93	42.08	28.97	44.84	85.51	89.46	88.44		