

1 **Liquid chromatography with diode array detection and**
2 **multivariate curve resolution for the selective and sensitive**
3 **quantification of estrogens in natural waters**

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5 Rocío L. Pérez, Graciela M. Escandar*

6 *Instituto de Química Rosario (CONICET-UNR), Facultad de Ciencias*

7 *Bioquímicas y Farmacéuticas, Universidad Nacional de Rosario, Suipacha 531*

8 *(2000) Rosario, Argentina. E-mail: escandar@iquir-conicet.gov.ar*

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24 **Abstract**

25 Following the green analytical chemistry principles, an efficient strategy involving second-
26 order data provided by liquid chromatography (LC) with diode array detection (DAD) was
27 applied for the simultaneous determination of estriol, 17 β -estradiol, 17 α -ethinylestradiol and
28 estrone in natural water samples. After a simple pre-concentration step, LC-DAD matrix data
29 were rapidly obtained (in less than 5 min) with a chromatographic system operating
30 isocratically. Applying a second-order calibration algorithm based on multivariate curve
31 resolution with alternating least-squares (MCR-ALS), successful resolution was achieved in
32 the presence of sample constituents that strongly coelute with the analytes. The flexibility of
33 this multivariate model allowed the quantification of the four estrogens in tap, mineral,
34 underground and river water samples. Limits of detection in the range between 3 and 13 ng
35 L⁻¹, and relative prediction errors from 2 to 11% were achieved.

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43 **Keywords**

44 Liquid chromatography

45 Diode array detection

46 Multivariate curve resolution

47 Estrogens

48 Natural waters

49 **1. Introduction**

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51 Estrogens are steroidal hormones which play an important role in human physiology,
52 including, among others, reproductive female functions, modulation of tissues growth and
53 bone integrity [1]. The three major naturally occurring estrogens 17β -estradiol (E2), estriol
54 (E3), estrone (E1), and the synthetic estrogen 17α -ethynylestradiol (EE2), widely used in
55 contraceptive pills, are the main contributors to the total estrogenicity in waterways [2]. In
56 fact, active estrogen forms are constantly excreted into the aquatic environment and may
57 cause serious health effects in animals and humans, especially in regards to reproduction
58 [3,4]. Since estrogens are the most potent endocrine disrupting compounds (EDCs) which, in
59 turn, are defined as chemicals that may negatively interfere with the endocrine system of
60 humans and wildlife [5], it is not surprising that continuous efforts are devoted to find
61 sensitive and selective methods for their quantification in natural samples.

62 Complete overviews on the development of the analysis of steroidal hormones in
63 environmental matrices can be found in the literature [2,6–9]. As indicated in the latter works,
64 liquid chromatography (LC) and gas chromatography (GC) followed by detection with mass
65 spectrometry (MS) or tandem MS are the most employed analytical tools to determine
66 estrogens and other EDCs in many different water sources. However, this instrumental is
67 sophisticated and usually requires important capital investment and personnel training. In
68 addition, because of the complexity of certain environmental matrices, a great effort must be
69 devoted to sample preparation, with the additional risk of loss of analytes during extensive
70 extraction and clean up steps [2].

71 In such situations, multivariate data analysis can be used for improving the selectivity of
72 data collected in less expensive equipment by mathematical means. Specifically, multi-way
73 calibration based on higher-order data (e.g. second-order LC-diode array detection or LC-

74 DAD data) allows the prediction of analyte concentrations in samples containing potential
75 interferences. This useful property, named the “second-order advantage” [10,11], avoids the
76 requirement of interference removal, with the concomitant saving of experimental work and
77 analysis time. Further, toxic organic solvents frequently used for clean up procedures are
78 prevented.

79 As part of a program devoted to the development of high performance methods within the
80 framework of green chemistry principles [12,13], the use of isocratic LC-DAD data coupled
81 to second-order multivariate calibration, was proposed as a useful approach for rapid and
82 selective detection of estrogens. The LC-DAD matrix data were obtained in short times and
83 using minimal solvent volumes. In a first phase, determinations were carried out in solutions
84 containing the studied estrogens and additional compounds selected as potential interferences.
85 In a second step, the proposed methodology was applied to real samples.

86 Two issues had to be taken into account when choosing the appropriate algorithm to
87 process the present data: 1) component profiles in the elution time mode usually change in
88 shape and/or position from sample to sample, and 2) the absorption spectra of the studied
89 analytes are very similar. These problems were overcome applying the so-called extended
90 multivariate curve resolution-alternating least-squares (MCR-ALS) algorithm [14], using
91 specific strategies which will be discussed below. It is important to remark that this algorithm
92 has been proposed for handling different types of chromatographic challenges [15,16] and, in
93 the present report, it was successfully used for improving both the sensitivity and selectivity
94 of the applied chromatographic method.

95

96 **2. Experimental**

97

98 *2.1. Instrumentation*

100 Chromatographic runs were performed on an HP 1200 liquid chromatograph (Agilent
101 Technologies, Waldbronn, Germany) consisting of a quaternary pump, a manual injector
102 fitted with a 50 μ L loop and a diode array UV–visible detector set at a wavelength range from
103 200 to 330 nm. Three C18 chromatographic columns provided by Agilent Technologies
104 (Santa Clara, CA, USA) were checked: Zorbax Eclipse XDB (4.6 \times 150 mm, 5 μ m particle
105 size), Poroshell 120 EC (4.6 \times 100 mm, 2.7 μ m particle size), and Poroshell 120 EC (4.6 \times 50
106 mm, 2.7 μ m particle size). The data were collected using the software HP ChemStation for
107 LC Rev. HP 1990–1997.

108

109 *2.2. Reagents and solutions*

110

111 All reagents were of high-purity grade and used as received. Estriol, 17 β -estradiol, 17 α -
112 ethynylestradiol, estrone, naproxen (NX), drospirenone (DRSP), norethisterone acetate
113 (NETA), androstenedione (AED), and diazepam (DZM) were purchased from Sigma-Aldrich
114 (Milwaukee, WI, USA). Methanol and acetonitrile were obtained from Merck (Darmstadt,
115 Germany). Water was purified using a MilliQ system (Millipore, Bedford, USA).

116 Methanol stock solutions of estrogens and potential interferents were prepared and stored
117 in dark flasks at 4 $^{\circ}$ C. A set of five calibration solutions by duplicate (10 samples) containing
118 E3, E2, EE2 and E1, each equally spaced in the range 0–110 ng mL⁻¹, were prepared by
119 measuring appropriate aliquots of standard solutions, placing them in 2.00 mL volumetric
120 flasks, evaporating the solvent with a nitrogen stream, and completing to the mark with the
121 solvent mixture used as mobile phase. A test set of additional 19 samples, containing the four
122 analytes and also NX, DRSP, NETA, AED, and DZM, were similarly prepared. The

123 concentrations of each potential interferent ranged between 70–340 ng mL⁻¹, and were
124 randomly selected.

125

126 *2.3. Real samples*

127

128 Because the evaluated water samples (tap, mineral, underground and river waters) did not
129 contain the studied estrogens at levels higher than the attained detection limits, a recovery
130 study was carried out by spiking them with standard solutions of E3, E2, EE2 and E1,
131 obtaining concentration levels in the range 10–100 ng L⁻¹. These water samples were
132 prepared in duplicate and, with the exception of river water, they underwent no previous
133 treatment. River water was collected from Paraná River (Rosario, Argentina) in a 4 L amber
134 glass bottle rinsed with methanol and Milli-Q water, stored at 4 °C immediately after
135 sampling, and analyzed as soon as possible (within 48 h after collection) in order to avoid
136 addition of chemical preservatives. River samples were filtered twice prior to injection: first
137 through a paper filter and then through a cellulose acetate 0.2 µm pore size filter.

138 The solid-phase extraction (SPE) procedure was carried out using SPE disks
139 Empore Octadecyl C18 (Supelco, Bellefonte, PA, USA). The membrane was conditioned
140 with 1 mL of methanol and then the extraction of 250 mL of the sample was carried out in
141 approximately 12 min per sample. This flow rate is in the optimum range for maximum
142 breakthrough volume (10–30 mL min⁻¹) [17]. The retained estrogens were eluted with
143 methanol, and this solvent was evaporated with a nitrogen stream. Then, the solutions were
144 reconstituted with 0.200 mL of mobile phase and subjected to the same chromatographic
145 analysis as the test samples. In this way, the preconcentration factor was 2500.

146

147 *2.4. LC-DAD procedure*

148

149 The data matrices were collected from 0 to 4.5 min each 1.8 s in the elution time axis, at
150 wavelengths from 200 to 330 nm each 1 nm. The slit width was 1 nm. The LC-DAD matrices
151 of size 149×131 (time and spectral data points respectively) were saved in ASCII format, and
152 transferred to a PC for subsequent manipulation. The mobile phase used for all
153 chromatographic runs was a 50:50 (v/v) mixture of water and acetonitrile, delivered at a flow
154 rate of 1.0 mL min⁻¹ with a chromatographic system operating under isocratic mode.

155

156 *2.5. Software*

157

158 The data were handled using the MATLAB computer environment [18]. The calculations
159 involving MCR-ALS have been made using *mvc2_gui*, a MATLAB graphical interface
160 toolbox which is a new version of that already reported in the literature [19], freely available
161 at www.iquir-conicet.gov.ar/descargas/mvc2.rar

162

163 **3. Theory**

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165 The MCR-ALS algorithm has been discussed in detail [14], and thus only a brief
166 description is presented here. In this algorithm, an augmented data matrix is created from the
167 test data matrices and the calibration data matrices. These individual matrices are of size $J \times K$,
168 where J is the number of elution times (number of rows of each data matrix) and K the
169 number of emission wavelengths (number of columns of each data matrix). Augmentation can
170 be performed either column-wise or row-wise, depending on the type of experiment being
171 analyzed [20]. In the presently studied case, the augmentation was implemented column-wise,
172 i.e. in the elution time direction, because in this way the chemical rank of the augmented
173 matrix is better preserved.

174 In the column-wise augmentation mode, the bilinear decomposition of the augmented
175 matrix is performed according to the expression:

$$176 \quad \mathbf{D} = \mathbf{C} \mathbf{S}^T + \mathbf{E} \quad (1)$$

177 where the columns of \mathbf{D} contain the chromatograms measured at J times for $(I_{\text{cal}} + 1)$ different
178 samples at K wavelengths, the columns of \mathbf{C} contain the augmented elution time profiles of
179 the intervening species, the columns of \mathbf{S} their related spectra, and \mathbf{E} is a matrix of residuals
180 not fitted by the model. The sizes of these matrices are \mathbf{D} , $J(I_{\text{cal}} + 1) \times K$, \mathbf{C} , $J(I_{\text{cal}} + 1) \times N$, \mathbf{S} ,
181 $K \times N$, \mathbf{E} , $J(I_{\text{cal}} + 1) \times K$ (N is the number of responsive components). As can be observed, \mathbf{D}
182 contains data for the I_{cal} calibration samples and for a given test sample. Decomposition of \mathbf{D}
183 is achieved by iterative least-squares minimization of the residuals contained in \mathbf{E} , under
184 suitable constraining conditions such as non-negativity, unimodality, correspondence,
185 selectivity, trilinearity, closure, etc. [20].

187 MCR-ALS requires initialization with parameters as close as possible to the final results.
188 For example, the species spectra can be supplied, as obtained from either pure analyte
189 standards or estimated from the analysis of the so-called 'purest' spectra [21–23], applying a
190 multivariate algorithm which extracts pure component spectra from a series of spectra of
191 mixtures of varying composition [21]. Another option is to provide estimated elution time
192 profiles, as obtained from procedures such as evolving factor analysis (EFA) [24]. Specific
193 constraints and initialization applied in the present case will be explained below.

194 After MCR-ALS decomposition of \mathbf{D} , concentration information contained in the elution
195 profiles (\mathbf{C} matrix) can be used for quantitative predictions, by first defining the analyte score
196 as the area under the profile for the i th sample:

$$197 \quad a(i,n) = \sum_{j=1+(i-1)J}^{iJ} C(j,n) \quad (2)$$

198 where $a(i,n)$ is the score for the analyte n in the sample i , and $C(j,n)$ is the element of the
199 analyte profile in the augmented mode. The analyte scores in the calibration samples are

200 employed to build a pseudo-univariate calibration graph against the nominal analyte
201 concentrations, predicting the concentration in the test samples by interpolation of the test
202 sample score.

203

204 **4. Results and discussion**

205

206 *4.1. Selection of optimal experimental conditions*

207

208 In order to achieve the resolution of the studied estrogens in the shortest possible time and
209 using the least amount of organic solvent, chromatographic conditions were optimized.
210 According to previous experience related to the chromatographic determination of estrogens
211 [25,26], mobile phases containing different ratios of acetonitrile and water were tested, and a
212 mobile phase constituted by acetonitrile-water in a 50:50 ratio provided the best resolved
213 peaks. Thus the latter mobile phase was used in all runs.

214 For the three C18 chromatographic columns of 50, 100 and 150 mm length packed with
215 particles of 2.7, 2.7 and 5 μm average diameter, respectively, different loop volumes (5, 20,
216 50 and 100 μL) and flow rates in the range 0.8–1.5 mL min^{-1} were probed. It was
217 corroborated that the 100 mm column packed with 2.7 μm particles, a 50 μL loop sample and
218 a flow rate of 1 mL min^{-1} produced better signals. The pH values of the sample solutions
219 were approximately neutral (the pH was not adjusted).

220 A model system of the four analytes prepared in a mobile phase solution was tested using
221 the working conditions summarized in Table 1. Estrogens peaks were resolved in less than 4.5
222 min using an isocratic regime, the elution order being E3, E2, EE2 and E1.

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225

Table 1

Instrumental and chemical parameters.

	Values/reagents
Mobile phase	Acetonitrile/water (50:50, v/v)
Column	Poroshell 120 EC (4.6×100 mm, 2.7 μm particle size)
Volumetric flow-rate (mL min ⁻¹)	1.0
Temperature	Room-temperature
Injection volume (μL)	50
Time range (min)	From 0 to 4.5
Wavelength range (nm)	From 200 to 330
Calibration range (ng mL ⁻¹)	From 0 to 110

226

227 *4.2. Multivariate calibration results*

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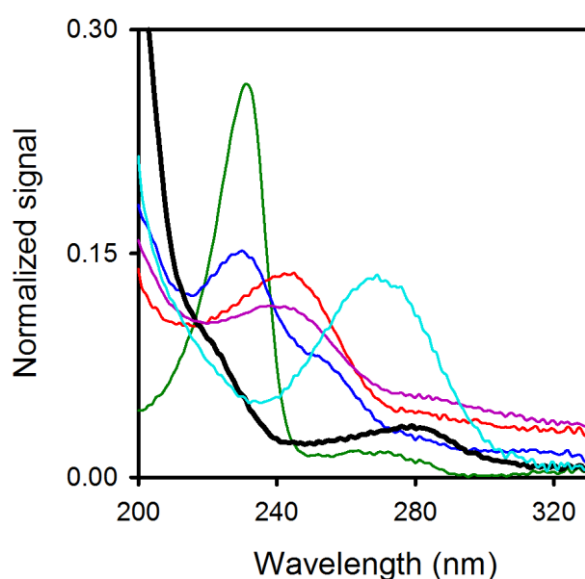
229 In real samples, the simultaneous presence of additional matrix constituents, which overlap
230 both in the spectral and time modes with the analytes, precludes the estrogens quantification
231 through classical zeroth-order calibration. In this latter case, it is highly convenient to use
232 second-order calibration with suitable algorithms for the quantitation of the analytes, because
233 of the need of achieving the second-order advantage [10,11]. However, an additional
234 limitation inherent to chromatographic second-order data is the lack of repeatability in the
235 elution time profiles between successive runs, which prevents the use of algorithms requiring
236 that the data show the trilinearity property [27]. In this regard, MCR-ALS was selected for
237 data processing because this algorithm achieves the second-order advantage and has the
238 additional benefit of not requiring that a given component shows the same chromatographic
239 profile in each experimental run [28].

240 In a first stage, samples containing the studied analytes and potential interferences were
241 processed. Real samples containing their own constituents were then studied.

242

243 *4.2.1. Synthetic samples*

244 With the purpose of mimic a real situation, test samples were prepared containing the
245 estrogens solutions and also foreign compounds which could be concomitantly present in
246 natural waters. It was verified that emerging pollutants such as naproxen (an anti-
247 inflammatory drug), drospirenone and noresthisterone acetate (two progestins),
248 androstenedione (a sex hormone precursor), and diazepam (a psychiatric drug) coelute with
249 the analytes under the established working conditions, and also strongly overlap their spectral
250 signals (Fig. 1). Therefore, these compounds were selected as potential interferents.

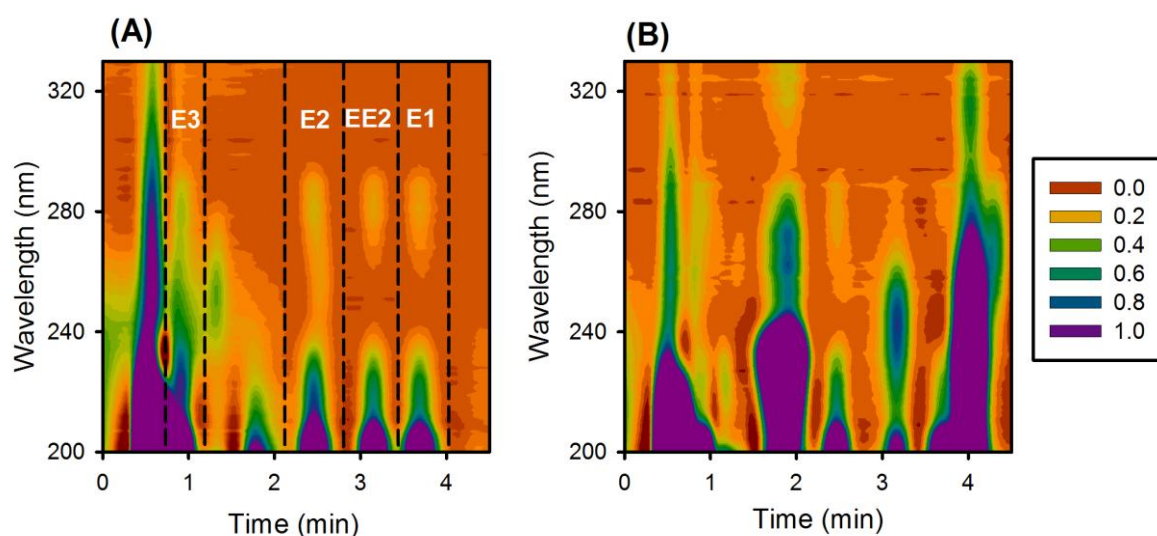


251
252 **Fig. 1** Normalized absorption spectra in acetonitrile-water (50:50 v/v) for the assayed
253 estrogens (thick black line), NX (green line), DZM (blue line), DRSP (cyan line), NETA
254 (violet line), and AED (red line).
255

256 Figure 2 shows the contour plots for typical LC-DAD matrices recorded for a calibration
257 and for a test sample, where the high complexity of the analytical problem under study can be
258 appreciated.

259 Because the four studied estrogens have very similar UV spectral profiles (see Fig. 1), it
260 was not possible to perform MCR-ALS analysis with matrix augmentation in the temporal
261 direction (i.e., column-wise) when working with the full chromatogram (e.g. involving the
262 complete elution time range). This would lead to almost zero spectral selectivity, making it

263 difficult the decomposition of the augmented matrix. An alternative in this case is to augment
 264 the matrices in the spectral direction (i.e., row-wise) [29]. However, due to elution time shifts
 265 and band shape changes among calibration and test samples, matrix augmentation in the
 266 spectral direction is also inconvenient. Hence, MCR-ALS was applied by column-wise
 267 augmentation of chromatographic data matrices, but dividing the elution time axis in four
 268 time regions, each one including a single analyte (see Table 2 and [Fig. 2A](#)).



269
 270 **Fig 2.** Two-dimensional contour plots of LC-DAD matrices for samples only containing the
 271 studied estrogens (A), and in the presence of NX, DZM, DRSP, NETA and ADE as
 272 interferences (B). Dashed lines in (A) delimit the selected chromatographic/spectral regions
 273 used for data processing of each analyte, as indicated. Concentrations are as follows (all in ng
 274 mL⁻¹): (A) E3, 102; E2, 110; EE2, 109; E1, 101. (B) E3, 66; E2, 83; EE2, 38; E1, 96; NX,
 275 252; DZM, 308; DRSP, 78; NETA, 60 and AED, 70.
 276

277 **Table 2**
 278 Selected chromatographic/spectral ranges used for MCR-ALS data processing.

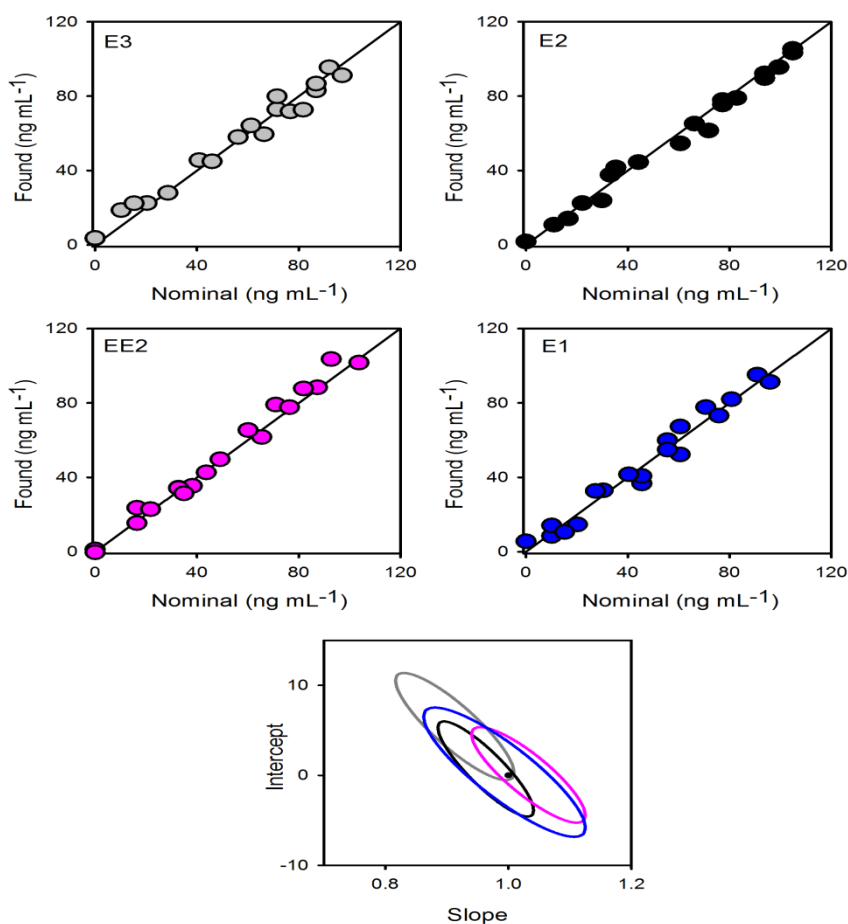
Analyte	Time (min)	Wavelength (nm)
E3	0.82 – 1.19	200 – 330
E2	2.10 – 2.86	200 – 330
EE2	2.86 – 3.40	200 – 330
E1	3.40 – 4.07	200 – 330

281 In this latter case, the chemical rank would be equal to the mathematical pseudorank,
 282 because the component spectra do not change from sample to sample. Thus, data processing
 283 comprised the building of augmented column-wise **D** matrices containing, for each time

284 region and in the whole wavelength range, data for each of the analyzed samples and for the
285 calibration samples.

286 Before starting MCR-ALS resolution, the estimation of the number of spectrally active
287 components in each **D** data matrix was made from the plot of singular values as a function of
288 a trial number of components, locating a number for which the plot stabilizes. The latter
289 number is initially employed for MCR-ALS analysis, and is afterwards refined (increased or
290 decreased) until an appropriate solution is found, with a reasonable least-squares fit and
291 physically recognizable profiles. For a given number of responsive components, their spectra
292 were then estimated from the analysis of the so-called purest variables [21]. The profiles
293 provided by the latter analysis were suitable to perform the resolution and, therefore, it was
294 not necessary to include reference spectra for the analyte as initial estimates for MCR-ALS.

295 In order to drive the iterative procedure to chemically interpretable solutions, non-
296 negativity constraints in both modes, correspondence restriction, and unimodality constraint
297 in the temporal mode were applied. The selected MCR convergence criterion was 0.1%
298 (relative change in fit for successive iterations) and the maximum number of iterations was
299 2500. However, convergence was achieved after less than 10 iterations in most of the
300 evaluated samples.



301
 302 **Fig. 3** Plots of E3 (gray), E2 (black), EE2 (pink), and E1 (blue) predicted concentrations as a
 303 function of the nominal values in test samples (as indicated), and elliptical joint regions (at
 304 95% confidence level) for the slopes and intercepts of the regressions for the corresponding
 305 predictions. The black dot in the elliptical plots marks the theoretical (intercept = 0, slope = 1)
 306 point.
 307

308 Figure 3 shows the prediction results corresponding to the application of MCR-ALS to a
 309 set of 19 test samples. As can be observed, the predictions for the four estrogens are in good
 310 agreement with the corresponding nominal values. If the elliptical joint confidence region
 311 (EJCR) [30] is analyzed for the slope and intercept of the above plot, we conclude that ellipse
 312 includes the theoretically expected values of (1,0), indicating the accuracy of the used
 313 methodology. The good recoveries obtained after the application of MCR-ALS suggest that
 314 interacting background effects, which could be present in chromatographic analysis of
 315 complex matrices [31], are not significant and, therefore, the use of external calibration was
 316 an adequate option. The statistical results are completed with the values shown in Table 3.

Table 3

Statistical results for the studied estrogens in samples with NX, DZM, DRSP, NETA and ADE as potential interferents (test set) and in spiked water samples using LC-DAD matrices and MCR-ALS.

	E3	E2	EE2	E1
<i>Synthetic test set</i>				
SEL	0.93	0.77	0.90	0.72
γ (ng ⁻¹ mL)	4.3	4.7	5.0	4.8
LOD (ng mL ⁻¹)	14	10	10	8
LOQ (ng mL ⁻¹)	42	30	30	24
RMSEP (ng mL ⁻¹)	5.1	4.1	4.4	5.1
REP (%)	10	7.4	8.0	10
<i>Tap water^a</i>				
SEL	0.56	0.85	0.83	0.81
γ (ng ⁻¹ mL)	1000	1800	3300	3400
LOD (ng mL ⁻¹)	0.012	0.012	0.010	0.006
LOQ (ng mL ⁻¹)	0.036	0.036	0.030	0.018
RMSEP (ng mL ⁻¹)	0.005	0.003	0.005	0.001
REP (%)	11	7	11	2
<i>Mineral water^a</i>				
SEL	0.90	0.67	0.86	1.0
γ (ng mL ⁻¹)	4400	4700	4600	4100
LOD (ng mL ⁻¹)	0.013	0.008	0.008	0.008
LOQ (ng mL ⁻¹)	0.039	0.024	0.024	0.024
RMSEP (ng mL ⁻¹)	0.003	0.002	0.003	0.001
REP (%)	8	5	6	3
<i>Underground water^a</i>				
SEL	0.47	0.53	0.88	0.57
γ (ng mL ⁻¹)	1300	3400	4400	4800
LOD (ng mL ⁻¹)	0.003	0.007	0.006	0.007
LOQ (ng mL ⁻¹)	0.009	0.021	0.018	0.021
RMSEP (ng mL ⁻¹)	0.001	0.002	0.003	0.001
REP (%)	2	5	8	3
<i>River water^a</i>				
SEL	0.60	0.83	0.68	0.54
γ (ng mL ⁻¹)	3700	4000	3600	5600
LOD (ng mL ⁻¹)	0.010	0.008	0.011	0.010
LOQ (ng mL ⁻¹)	0.030	0.024	0.033	0.030
RMSEP (ng mL ⁻¹)	0.002	0.001	0.002	0.002
REP (%)	5	3	6	6

^a The results refer to water samples before SPE. For comparison with the test samples, values for water samples are given in ng mL⁻¹.

SEL, selectivity calculated according to ref. 32.

γ , analytical sensitivity

LOD, limit of detection calculated according to ref. 29.

LOQ, limit of quantification calculated as LOD \times 3.

RMSEP, root-mean-square error of prediction.

REP, relative error of prediction.

318 The relative errors of prediction (all below 15 %) indicate good precision. The obtained
319 values of both limits of detection (LODs) and quantification (LOQs), in the order of parts-per-
320 billion, demonstrate the positive effect of second-order data in the sensitivity of the method
321 [29]. However, considering the estrogen levels which can be found in water samples in parts-
322 per-trillion (see below), it is evident that a pre-concentration step is required for the
323 subsequently evaluated real systems.

324

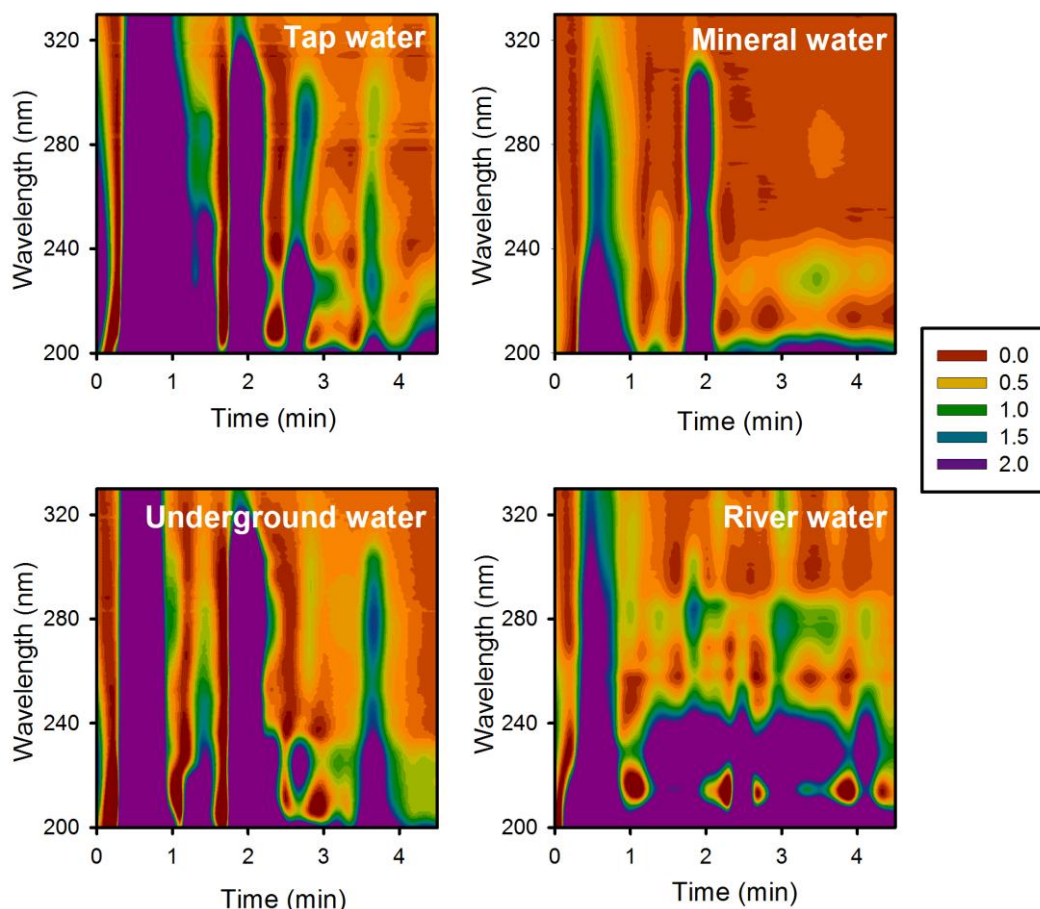
325 4.2.2. *Real water samples*

326 With the purpose of testing the present method in real samples and demonstrating its
327 ability of overcoming the interference from background constituents, waters from different
328 origins were analysed.

329 In water bodies, estrogens are detected in a wide range of concentrations, generally in the
330 order of parts-per-trillion levels [7]. Therefore, the sensitivity of the method was increased
331 through a pre-concentration step employing C18 membrane-SPE. It is necessary to point out
332 that the selection of C18 membranes is based on our excellent experience with this solid-
333 support as extractor of low-polarity compounds, such as the studied analytes [28,33].
334 Additionally, these membranes are easily and rapidly conditioned, decreasing the laboratorist
335 effort. Because the selectivity between the analytes and interferences is provided by the
336 chemometric tool, the complete physical separation of target analytes from the matrix
337 constituents is not required, as in traditional extraction techniques.

338 Figure 4 shows contour plots of LC-DAD matrices corresponding to different real water
339 samples after the SPE procedure. As expected, the C18 membrane also retains other matrix
340 constituents which can interfere, as in the present case, co-eluting with the estrogens and
341 overlapping their absorption spectra. For applying classical zeroth-order calibration, these
342 interferences should be completely removed before the quantification is performed. However,

343 this fact does not represent a problem when using an appropriate second-order calibration
344 approach.



345
346 **Fig. 4.** Two-dimensional contour plots of LC-DAD matrices for different water samples, after
347 SPE, spiked with the studied analytes. Original concentrations are as follows (all in ng L^{-1}):
348 E3, 53; E2, 51; EE2, 50; E1, 53 (tap water); E3, 42; E2, 35; EE2, 49; E1, 41 (mineral water);
349 E3, 53; E2, 42; EE2, 80; E1, 74 (underground water); E3, 46; E2, 40; EE2, 41; E1, 40 (river
350 water).

351
352
353 The chromatographic and spectral regions processed for each analyte were the same as
354 those used for test samples. Table 4 shows the results of the recovery study performed by
355 spiking water samples with appropriate amounts of estrogens, in duplicate, at three different
356 concentration levels. The average recovery of the four estrogens in each type of water at the
357 three different fortification levels was tested for significance by using the Student *t*-test: the
358 null hypothesis corresponds to the recovery of 100% [30]. The *t* values obtained for $n - 1$
359 degrees of freedom (where n is the number of evaluated levels) at a 95% of significance

360 compare favorably with the corresponding tabulated value [$t_{crit}(0.05,2) = 4.30$], suggesting
 361 that the proposed method is appropriate for the determination of the studied compounds.

Table 4
 Recovery study for the studied estrogens in spiked water samples using MCR-ALS^a

	E3				E2				EE2				E1			
	Taken	Found ^b	R	t^c	Taken	Found ^b	R	t^c	Taken	Found ^b	R	t^c	Taken	Found ^b	R	t^c
TW ^d	105	102(2)	97		103	107(1)	104		100	114(2)	114		105	99(1)	94	
	53	54(7)	102		51	49(3)	96		50	46(5)	92		52	51(3)	98	
	25	27(1)	108	0.69	22	25(4)	114	0.96	26	24(4)	92	0.14	24	23(2)	96	3.46
MW ^e	18	22(1)	122		18	18(2)	100		18	15(2)	83		16	15(2)	94	
	13	13(2)	100		12	13(1)	108		13	13(1)	100		10	9(5)	94	
	42	43(7)	102	1.15	35	34(4)	97	0.58	49	51(6)	104	0.63	40	41(6)	103	0.99
UW ^f	53	47(7)	100		41	43(4)	104		80	81(3)	101		74	73(4)	99	
	33	33(5)	100		20	19(6)	95		35	33(3)	94		16	15(2)	94	
	14	15(2)	107	0.19	12	14(2)	117	0.79	15	19(1)	126	0.76	11	12(1)	109	0.22
RW ^g	21	23(1)	110		16	17(4)	106		22	22(3)	100		20	17(1)	85	
	46	48(3)	104		40	44(7)	110		41	41(1)	100		40	40(3)	100	
	84	84(5)	100	1.73	80	82(3)	103	2.60	82	87(5)	106	1.15	79	81(2)	103	0.69

^a Concentrations are given in ng L⁻¹ and recoveries (R) in percentage.

^b Means of duplicates. Standard deviation between parentheses.

^c Calculated student t for the average recovery. The critical t value for $n - 1$ degrees of freedom at a 95% significance level is $t_{crit}(0.05,2) = 4.30$ (see text).

^d TW, tap water from Rosario city (Santa Fe, Argentina).

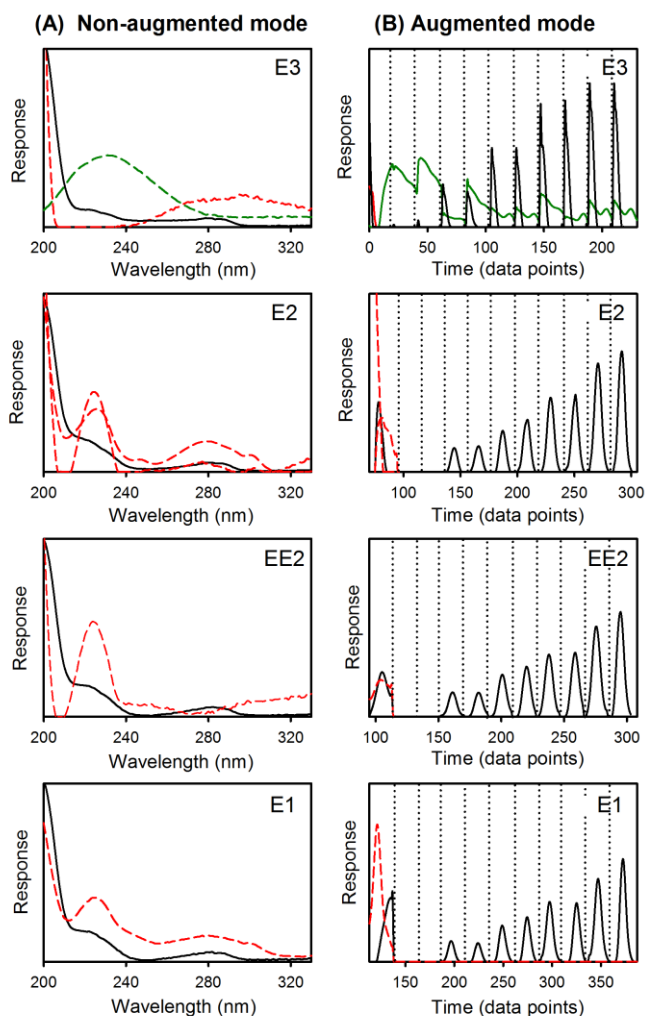
^e MW, mineral water from Villavicencio hills (Mendoza, Argentina)

^f UW, underground water from Funes City (Santa Fe, Argentina).

^g RW, river water from Paraná river (Argentina).

362
 363 The outstanding results obtained after MCR-ALS was applied to the data suggest that the
 364 method can overcome the problem of the presence of unexpected interferents from the
 365 background of the real samples. As an example, Fig. 5 shows the profiles retrieved by MCR-
 366 ALS in both spectral and temporal modes for a real matrix (underground water in this
 367 example) added with the analytes. From top to bottom, each pair of plots in Fig. 5 shows the
 368 retrieved spectral and augmented time profiles for the studied analytes (these augmented time
 369 profiles contain successive sub-profiles for the unknown and calibration samples). For each
 370 analyte, the specific elution time region was the same as for the test samples, and repeats itself

371 in each sub-profile of the augmented time profile. It can be concluded that, although
 372 interferences are present in all regions, the spectra are correctly distinguished, and the
 373 chromatographic bands are recognized as belonging to the corresponding estrogen and
 374 background (present in water and calibration samples) or interferences (only present in the
 375 real sample).



376

377 **Fig. 5** Profiles retrieved by MCR-ALS when processing an underground water sample spiked
 378 with the four estrogens. (A) Spectral profiles in the selected E3, E2, EE2 and E1
 379 chromatographic regions, as indicated. (B) The corresponding time profiles (the dotted
 380 vertical lines separate, from left to right, the studied sample and the successive calibration
 381 samples). In all plots, the solid black line indicates estrogen, and dashed green and red lines
 382 indicate background and interferences, respectively.
 383

384 Table 3 displays the statistical results corresponding to the estrogen determination in real
 385 water matrices. These results indicate that neither the selectivity nor the relative error of

386 prediction is significantly affected by the fact that real matrices are being studied. Besides, the
 387 analytical sensitivity and limits of detection and quantification reflect the benefits of the
 388 preconcentration, and the possibility of determining the studied analytes at part-per-trillion
 389 levels. In comparison with the performances of selected chromatographic methods reported in
 390 the last years for the determination of estrogens in water samples (Table 5), limits of detection
 391 from 0.03–10, 0.01–36, and 1– 1100 ng L⁻¹ have been found using CG-mass spectrometry,
 392 LC- mass spectrometry, and LC-spectroscopic methods, respectively. In the present case, low
 393 limit of detections are achieved (LODs = 3–13 ng L⁻¹) applying a non-sophisticated method
 394 such as LC-DAD, with a relatively small use of organic solvents and without the need of
 395 derivatization as most CG procedures [7] and some LC methods [39].

Table 5

Analytical performance of selected chromatographic methods reported in the last years for estrogens in natural waters.

Sample pretreatment	Method	Estrogen	Other	LOD ^a	Sample	Ref
<i>Gas chromatography</i>						
SPE(C18)	LVI-GC-MS	E1, E2, EE2		0.041, E1; 0.046, E2; 0.031, EE2	SW, WW	34
HF-MMLLE-deriv	GC-MS	E1, E2, EE2		3, E1; 1.6, E2; 10, EE2	TW, SWW	35
SPE(Oasis HLB)-deriv	GC-MS/MS	E1, 17 α - and 17 β -E2, E3, EE2, MES	andr	0.7–3	SW, treated effluents	36
<i>Liquid chromatography-mass spectrometry</i>						
SPE(Strata)	LC-MS/MS	E1, 17 α -E2, 17 β -E2, E3, EE2	andr, prog	0.02, E1, 0.01, 17 β -E2; 0.03, 17 α -E2, 0.03, E3; 0.2, EE2	SW, UW	37
Deriv-SPE(Oasis HLB)	LC-MS/MS	E1, E2, E3	ECF	0.038, E1; 0.13, E2; 0.11, E3	RW	38
SPE(Oasis HLB + Florisil)-deriv	LC-MS/MS	E1, 17 α -E2, 17 β -E2, E3, EE2		0.07, E1; 0.084, 17 α -E2; 0.078, 17 β -E2; 0.28, E3; 0.067, EE2	RW	39
On line SPE (Hypersil Gold C18)	LC-LC-MS/MS	E1, E2, E3, EE2	prog	10, E1; 3, E2; 50, E3; 7, EE2	SW, SWW	40
SPE(C18 disk)	UPLC-MS/MS	E1, E2, E3, EE2	prog	0.5, E1; 0.6, E2; 1.0, E3; 1.2, EE2	TW, RW, LW, WW	41
SPE(Oasis HLB + Florisil)	LC-MS/MS	E1, 17 α - and 17 β -E2, E3, EE2	ECF	0.4-3 ^b	RW, WWTP, WWTPPE	26
SPE(Oasis HLB + silica or Florisil)	LC-MS/MS	E1, 17 α -E2, 17 β -E2	andr, prog, cort	0.008-0.5	RW, DSW, effluents	42

Table 5. Continued

Sample pretreatment	Method	Estrogen	Other	LOD ^a	Sample	Ref
MM-SPE-MPS	LC-MS/MS	E1, E2, E3, EE2	BPA	1-36	TW, sea W, SWW	43
On line SPE(NG1)	LC-MS/MS	E1, E2, E3, EE2, andr DES		0.5-2	RW, WWTP, WWTPPE	44
US + SPE	LC-MS/MS	E1, E2, E3		0.07-60 ^c	Sea W	45
<i>Liquid chromatography-spectroscopy</i>						
On line SPE(CF)	LC-UV	E1, E2, E3, DES		16.2, E1; 78.1, E2; 5.6, E3; 0.98 DES	RW, LW, well W	46
SPE(MIP)	LC-UV	E1		5.7	Well W, LW	47
SPE(ENNFMM)	LC-UV	E1, E2, EE2		170, E1; 50, E2; 80, EE2	Natural waters	48
DLLME	LC-UV	E1, E2		100, E2; 200, E1	SPW, TW, RW	49
DLLME	LC-DAD-FD	E1, E2, E3, EE2, DES		80-500	RW, sea W, WW	50
On line SPME	LC-FD	E3, EE2	Prog, BPA	5-30	Natural waters	51
CP-CPE	LC-UV	E1, E2, EE2, DES, DHS		200-700	RW, LW	52
Coated SBSE	LC-UV	E1, E2, EE2, DES, DIS	BPA, OP	290, E1; 280, E2; 350, EE2; 260, DES; 180, DIS	RW, LW, FW	53
SPE(Strata)	LC-UV-FD	E1, E2, EE2	PPs	10-1100 ^b	RW, WWTP, WWTPPE	54
DLLME	LC-FD	E2, EE2		2.0, E2; 6.5, EE2	TW, SW, WW	55
SPE(C18 disk)	LC-DAD-MCR-ALS	E1, E2, E3, EE2		6-10, E1; 7-12, E2; 3-13, E3; 6-11, EE2	TW, MW, UW, RW	This work

^a For comparison, concentration units were unified to ng L⁻¹. ^b Limits of quantification. ^c Levels of measured concentrations. Abbreviations: andr, androgens; BPA, bisphenol A; CF, cigarette filter; cort, corticosteroids; CP-CPE, co-precipitation assisted cloud point extraction; DAD, diode array detector; deriv, derivatization; DES, diethylstilbestrol; DHS, dihydrostilbestrol; DIS, Dienestrol; DSW, discharging sites water; DLLME, dispersive liquid-liquid microextraction; ECF, estrogen conjugated forms; ENNFMM, electrospun nylon6 nanofibrous membrane; FD, fluorescence detector; FW, fishpond water; GC, gas chromatography; HF-MMLLE, hollow-fiber microporous membrana liquid-liquid extraction; LC, liquid chromatography; LVI, large volume injection; LOD, limit of detection; LOQ, limit of quantification; LW, lake water; MES, mestranol; MIP, molecularly imprinted polymer; MM-SPE-MPS, magnetic-mediated solid-phase extraction micro-particle sorbent; MS, mass spectrometry; MS/MS, tandem mass spectrometry; MW, mineral water; OP, octylphenol, PPs, pharmaceutical products; prog, progestagens; RW, river water; SBSE, Stir bar sorptive extraction; SPE, solid-phase extraction; SPW, spring water; SW, surface water; SWW, sewage water; TW, tap water; UPLC, ultra performance liquid chromatography; US, ultrasonication; UW, underground water; W, water; WW, wastewater; WWTP, wastewater treatment plant influent; WWTPPE, wastewater treatment plant effluent.

398 **5. Conclusions**

399

400 Liquid chromatography-diode array detection associated to multivariate curve resolution-
401 alternating least-squares (MCR-ALS) has demonstrated to be a powerful tool for the
402 determination of estradiol, estriol, estrone and ethynylestradiol in water samples.

403 Additional properties should be remarked beyond the outstanding sensitivity and
404 selectivity achieved in the present work, which led to limits of detection between 3 and 13 ng
405 L⁻¹, and excellent precision (prediction errors were equal to or below 11%). The use of an
406 appropriate chemometric tool makes it unnecessary to apply extraction and clean up steps for
407 the removal of coeluting compounds, avoiding the use of toxic organic solvents (essential for
408 environmental safety), and saving experimental time and operator effort.

409 The good quality of the obtained results suggests that the proposed method is appropriate
410 for the rapid analysis of the studied endocrine disrupting agents in natural waters, favorably
411 competing with sophisticated methods usually employed in this type of determinations, which
412 require expensive equipment and derivatization and sample extraction/clean up procedures.

413

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415

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419

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