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Screen-printed electrode based electrochemical detector coupled with ionic liquid dispersive liquid—liquid microextraction and microvolume back-extraction for determination of mercury in water samples

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1	Screen-Printed Electrode based Electrochemical Detector Coupled with		
2	Ionic Liquid Dispersive Liquid-Liquid Microextraction and Microvolume		
3	Back-Extraction for Determination of Mercury in Water Samples		
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12			
13	Abstract		
14	A novel approach is presented, whereby gold nanostructured screen-		
15	printed carbon electrodes (SPCnAuEs) are combined with in-situ ionic liquid		
16	formation dispersive liquid-liquid microextraction (in-situ IL-DLLME) and		

id d 17 microvolume back-extraction for the determination of mercury in water samples. In-situ IL-DLLME is based on a simple metathesis reaction between a water-18 19 miscible IL and a salt to form a water-immiscible IL into sample solution. Mercury complexes with ammonium pyrrolidinedithiocarbamate (APDC) are 20 21 extracted from sample solution into the water-immiscible IL formed in-situ. 22 Then, an ultrasound-assisted procedure is employed to back-extract the mercury into 10 µL of a HCl solution which is finally analyzed using SPCnAuEs. 23 Sample preparation methodology was optimized using a multivariate 24 25 optimization strategy. Under optimized conditions, a linear range between 0.5 and 10 µg L⁻¹ was obtained with a correlation coefficient of 0.997 for six 26 calibration points. The limit of detection obtained was 0.2 µg L-1. The 27

repeatability of the proposed method was evaluated at two different spiking levels (3 and 10 µg L⁻¹) and a coefficient of variation of 13% was obtained in both cases. The performance of the proposed methodology was evaluated in real-world water samples including tap water, bottled water and river water. Relative recoveries between 97 and 107% were obtained.

1. Introduction

Mercury is one of the most well-known toxic elements and even the World Health Organization places it between the first ten chemicals or group of chemicals of major public health concern [1]. Mercury exists in different forms with different properties, namely elemental or metallic (*i.e.* Hg⁰); inorganic (*i.e.* Hg²⁺); and organic (*i.e.* MeHg⁺, EtHg⁺, PhHg⁺). Several factors determine the adverse effects from mercury exposure including the chemical form of mercury, the dose, the age and health of the person exposed, and the duration and kind of exposure (e.g. inhalation, ingestion, etc.) [2]. Among the most relevant health effects we can mention damage to the gastrointestinal tract, nervous system, kidneys, respiratory failures and problems during the development of organs in unborn.

Mercury enters in the environment through both biogenic and anthropogenic vias. However, human activities such as mining, burning of fossil fuels, agriculture, paper and electrochemical industries, and household wastes, are the main responsible of the concerning increase of mercury levels in air, soil and water of certain contaminated areas. Monitoring the presence of mercury in natural and drinking waters is of great interest due to its high toxicity and bioaccumulation factor [3]. Mercury concentrations are commonly in the range

of low ng L⁻¹ in environmental waters [3] whereas the permitted level of mercury in drinking water depends on the responsible authorities of each territory. For example, the Environmental Protection Agency (EPA) set the threshold level at 2 µg L⁻¹ [4] but the European Union establishes the limit at 1 µg L⁻¹ [5].

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Numerous analytical methods using capillary electrophoresis [6, 7], gas [7, 8] and liquid [7, 9] chromatography, cold-vapor atomic absorption [7, 10] or fluorescence [7] spectrometry, inductively coupled plasma atomic emission [7] or mass spectrometry [7, 11] have been developed to determine mercury in natural [7] and drinking waters [8-11]. In addition, electrochemical techniques have also been widely employed and a proof of this are two excellent and recently published reviews about the latest advances in electrochemical, mainly voltammetric, determination of mercury [12, 13]. Electrochemistry offers sensitivity, simplicity, rapid response and inexpensive instrumentation with miniaturization and portable options. A major drawback to be considered results from the difficulty of removing mercury from electrode surface between measurements which lead to memory effect problems [12, 13]. However, tedious and time consuming cleaning steps can be avoided with the use of screen-printed electrodes (SPEs), which can be disposable after a single use due to their high cost effectiveness. Several methods based on SPEs have been reported for the determination of mercury in different water samples, including the use of bare gold SPEs [14], and modified SPEs with carbon nanomaterials [15-17], gold films [18, 19], gold nanoparticles [20, 21], nanohybrid materials [20] and chelating agents [22, 23]. As can be seen in Table 1, the vast majority of the reported works include a preconcentration step over the working electrode followed by anodic stripping voltammetry. Gold is commonly employed in working electrodes due to its high affinity for mercury which lead to an improvement in its preconcentration. In addition, mercury suffers from a process named underpotential deposition (UPD) on gold electrodes [13]. The UPD process occurs by the strong interaction between the two metals once the mercury is reduced forming an adsorbed layer. The formation of this layer makes that UPD occurs at a more positive potential than in normal conditions and, as a consequence, the selectivity of the method is generally improved. In this work, screen-printed carbon electrodes (SPCEs) modified with gold nanoparticles are employed as electrochemical transducers in the detection stage. The use of nanoparticles in electroanalysis is continuously growing due to its numerous advantages, related to the unique properties of nanoparticulate materials [24] (e.g. increase surface area, enhanced mass transport and improve selectivity, catalytic activity and signal to noise ratio).

Liquid-phase microextraction (LPME) [25] appeared in the latest nineties offering undoubted advantages as miniaturized extraction techniques, such as simplicity, easiness to handle, low sample and solvent consumptions, and an important reduction of residues generated. One of the most popular LPME technique is dispersive liquid-liquid microextraction (DLLME) [26] which has even come to dominate LPME research publications in the recent years [27]. DLLME is based on the complete dispersion of the extractant solvent into the sample, normally assisted by a disperser agent. During DLLME, there is a high contact between phases therefore the extraction is really rapid and effective. After the extraction, phases are separated normally by centrifugation and the enriched phase is analyzed. Numerous modifications of the original DLLME

procedure [26] have been reported up to now [28] including the use of new extractant solvents such as ionic liquids (ILs) [29]. Within the use of ILs, a novel methodology called in-situ IL formation dispersive liquid-liquid microextraction (in-situ IL-DLLME) [30, 31] has recently been developed. In-situ IL-DLLME is based on the formation of a water-immiscible IL using a metathesis reaction between a water-miscible IL and an ion exchange salt into sample solution. Thereby, the extractant phase is generated in-situ in form of homogeneously dispersed fine drops, the disperser agent is totally avoided and the extraction efficiency generally increases.

Different LPME techniques including single-drop microextraction [9, 32], DLLME [33–35], in-situ IL-DLLME [30] and task-specific IL ultrasound-assisted DLLME [36] have been employed for the determination and speciation of mercury in water samples. In these works, chromatographic systems [9, 34, 35], UV-Vis spectrometry [30], cold vapor [36] and electrothermal vaporization [32] atomic absorption spectrometry, and capillary electrophoresis [33] were used as analytical techniques.

The approach presented here employs an in-situ IL-DLLME followed by an ultrasound-assisted microvolume back-extraction and SPCnAuEs as electrochemical transducers for the determination of mercury in water samples. This combination exploits the advantages of including a miniaturized sample preparation step with the high sensitivity and specificity that offers the electrochemical determination of mercury using SPCnAuEs. LPME provides a high preconcentration of the analyte and a clean-up step for dirty matrixes employing low amounts of sample and chemicals. In addition, considering the low volume of sample needed for analysis with SPEs, they appear as an

alternative and perfectly compatible detection methodology after miniaturized extraction techniques, thus avoiding classical and bulky analytical instrumentation [37]. A multivariate optimization strategy has been adopted for the optimization of the sample preparation and the applicability of the method has been tested studying real-world water samples.

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2. Experimental part

2.1. Reagents and water samples

A stock standard solution of 1000 mg L⁻¹ of Hg²⁺ was prepared by dissolving Hg(OAc)₂ (≥ 99%) from Fluka (Stenhein, Germany) in ultrapure water. Working solutions were prepared by proper dilution of this stock standard. The IL 1-hexyl-3-methylimidazolium chloride ([Hmim][CI]) (98%) was purchased from Iolitec (Heilbronn, Germany). The lithium bis[(trifluoromethyl)sulfonyl]imide (LiNTf2) salt and the chelating ammonium pyrrolidinedithiocarbamate (APDC) (~ 99%) were supplied by Sigma-Aldrich (St. Louis, MO, USA). A solution of 2 mg mL⁻¹ of the chelating agent was prepared by dissolving APDC in ultrapure water. NaCl reactive grade and NaOH (≥ 97%, pellets) were from ACS Scharlau (Barcelona, Spain). Fuming HCI (37%) was supplied by Merck (Madrid, Spain). The ultrapure water employed for preparing all solutions was obtained with a Millipore Direct System Q5TM purification system from Ibérica S.A. (Madrid, Spain).

Standard Au³⁺ tetrachloro complex (1.000 \pm 0.002 g of AuCl₄⁻ in 500 mL of 1.0 M HCl) was purchased from Merck (Madrid, Spain). Solutions of AuCl₄⁻ 1 mM were prepared by suitable dilution of this standard solution in HCl 0.1 M.

Tap water was collected from the water-supplied network of the lab in the Departament of Physical and Analytical Chemistry of the University of Oviedo (Spain). Bottled water (San Benedetto mineral water, Valencia, Spain) was purchased in the supermarket. River water from Nora river was collected in Tiñana (Siero, Spain). All water samples were stored at 4 °C and were used without any further pretreatment. Initial analysis confirmed that mercury levels were undetectable in the three selected water samples.

2.2. Apparatus and electrodes

An ultrasonic bath from Elma (Singen, Germany) was used to assist the back-extraction procedure.

An Autolab PGSTAT 12 potentiostat from EcoChemie (Ultrecht, The Netherlands) controlled by Autolab GPES software version 4.8 was used for electrochemical experiments.

SPCEs (ref. DRP-110) with three electrode configuration were purchased from DropSens (Oviedo, Spain). The working electrode, with a disk-shaped of 4 mm of diameter, and the counter electrode were made of a carbon ink whereas the pseudo-reference electrode was made of silver. Specific connectors obtained from DropSens (ref. DRP-DSC) were used for the conexion of the SPCEs with the potentiostat.

2.3. In-situ IL-DLLME and microvolume back-extraction

Under optimum conditions, 20 mg of [Hmim][CI] were placed in a test tube and dissolved in 4 mL of aqueous standards or sample solutions containing the analyte and the chelating agent (40 μ L of 2 mg mL⁻¹). The ionic

exchange salt LiNTf₂ was added in an equimolar ratio (*i.e.* 28.3 mg) with the IL [Hmim][CI] and a cloudy solution was immediately formed. The mixture was manually shaken for 0.5 minutes. In order to accelerate phases separation, the tube was then introduced in an ice bath for 5 minutes. Next, the phases were separated by centrifugation for 10 minutes at 5000 rpm. The aqueous phase was removed with a glass pipette, and the formed IL-phase (*i.e.* [Hmim][NTf₂]) was withdrawn with a micropipette and deposited in an Eppendorf tube of 0.5 mL. For the back-extraction, 10 µL of HCl 4 M were added to the IL phase and the mixture was sonicated in an ultrasounds bath for 14 min at 90% of power and 37 KHz of frequency. After back-extraction, phases were separated by centrifugation for 5 min at 5000 rpm and the enriched acidic phase that remained in the upper part was analyzed.

2.4. Electrochemical analysis

Gold nanoparticles were generated over the SPCEs surface employing the procedure developed by Martínez-Paredes et al. [38] and previously optimized by D. Martín-Yerga et al. for the determination of mercury [20]. Briefly, 40 µL of a 1 mM AuCl₄- solution in HCl 0.1M were dropped onto the SPCE surface and a constant current of -100 µA was applied for 180 s. After gold nanoparticles deposition, the electrode surface was generously rinsed with ultrapure water and dried at room temperature before use. A new SPCnAuEs was prepared and employed for each experiment.

The electrochemical behavior of mercury on SPCnAuEs was previously and deeply studied [20], therefore, no further discussion will be included in the present work. After back-extraction, 5 µL of the resulting upper acidic phase

was mixed with 37 μ L of NaOH 0.5 M in order to obtain a suitable electrolytic medium. A volume of 40 μ L of this solution was deposited on the electrode surface for the voltammetric measurements. Mercury was determined by square-wave anodic stripping voltammetry (SWASV) employing previous optimized conditions [20]. Mercury was preconcentrated over SPCnAuEs by applying a constant potential of +0.3 V for 240 s. Thereafter, the potential was recorded between +0.3 V and +0.55 V at a frequency of 80 Hz, amplitude of 30 mV and step potential of 4 mV. All experiments were carried out at room temperature.

2.5. Data processing

An anodic peak corresponding to the reoxidation of mercury appears at approximately +0.42 V and the height of this peak was employed for the quantification of the analyte. The "base line correction" option provided by GPEs software was employed to get more defined peaks, specially at low concentrations, and to obtain more reliable and accurate measurements.

A two-step multivariate optimization strategy, using Plackett-Burman and central composite designs, was carried out to determine the optimum conditions of sample preparation. Minitab 15 statistical software (State College, PA, USA) was employed to construct the experimental design matrices and evaluate the results.

3. RESULTS AND DISUSSION

3.1. Optimization

3.1.1. Screening step

Plackett-Burman design is a two-level fractional factorial design that ignores interaction between variables and therefore main effects can be calculated with a reduced number of experiments leading to a saving in resources and time. The Plackett-Burman design results very useful in the first steps of a project when many variables are initially considered but finally only a few show important effects [39]. A saturated Plackett-Burman design was used to construct the matrix of experiments, including 11 variables: eight real variables and three dummy variables. The effects of dummy variables were used to evaluate the experimental error [40, 41]. The eight real experimental variables selected at two levels were: amount of [Hmim][CI], amount of chelating agent, ionic strength, pH, HCl volume, back-extraction time, power and frequency of the ultrasounds bath. Table 2 shows the experimental variables and levels considered in the Plackett-Burman design. A total of twelve experiments were randomly performed using aqueous standards of 25 µg L-1.

The data obtained were evaluated using an ANOVA test and the results were visualized with the Pareto chart shown in Figure 1. The length of each bar was proportional to the influence of the corresponding variable and the effects that exceed each reference vertical line can be considered significant with a 95% and 90% probability, respectively.

According to Figure 1, the ultrasounds frequency and HCI volume were statistically significant variables, with 95% probability, having a negative effect. The negative effect of the frequency is in agreement with the fact that at high ultrasounds frequencies, cavitation bubbles are more difficult to create as a result of the shorter duration of rarefraction cycles. Higher amplitudes (*i.e.* power) would be necessary to ensure that cohesive forces in the liquid were

overcome and maintain a certain cavitational energy [42]. For the HCl volume, the negative effect is easily explained considering that if less volume of acid is used, a higher concentration of the analyte is obtained. The ultrasounds device employed during this work only accepted two discrete values of frequency, namely 37 and 80 KHz, thus this significant variable could not be included in the following optimization step and was fixed in its lower level. As a consequence, back-extraction time and amount of [Hmim][CI], which showed important effects with 90% probability (see Fig. 1) were included in the optimization step. Back-extraction time showed a positive effect whereas for the amount of [Hmim][CI] the effect was negative. These effects revealed that the mass transfer during back-extraction is not as instantaneous as during in-situ IL-DLLME and it is enhanced if low amounts of [Hmim][CI] are used, probably because of diffusion effects and the fact that a smaller volume of immiscible IL-phase is formed during microextraction, and therefore a higher concentration of analyte is obtained.

The other four real variables considered in screening step with non-significant effects were fixed at the following levels: chelating agent, 40 μ L (2 mg mL⁻¹); ionic strength, 0% of NaCl; pH, the pH of water without any adjustment; and ultrasounds power, 90%.

3.1.2. Optimization of significant variables.

Central composite design (CCD) combines a two-level full factorial design (2^k) with 2k star points, where k is the number of variables being optimized, and one point at the center of the experimental region. In order to ensure the rotatability of the model, star points were set at $\alpha=\sqrt{k}=1.682$ whereas

the central point was repeated five times to provide an orthogonal design [39]. CCD was used to evaluate and optimize main effects, interaction effects and quadratic effects of the three considered variables. Table 3 shows the low and high levels, the central and star points of the considered variables in the optimization step. Nineteen experiments were randomly performed using aqueous standards of 25 µg L⁻¹.

The data obtained were also evaluated using an ANOVA test. The coefficients of the variables and the p-values are listed in Table 4.

Significant variables with 95% probability (*i.e.* p-value < 0.05) were HCl volume, back-extraction time and the quadratic effects of back-extraction time and amount of [Hmim][Cl], which confirms the curvature of the system and its fitting with the proposed second-grade polynomial system. The adjustment obtained expressed as r² value was 92%.

The response surfaces obtained using the CCD are shown in Figure 2. Pairs of variables were considered separately in order to easily interpret the effect of each one on the response of the system. Thus, Figure 2a shows the response surface which results of plotting HCl volume vs back-extraction time, with a retention value of 40 mg for the amount of [Hmim][Cl]; Figure 2b shows the response surface obtained as a function of HCl volume and amount of [Hmim][Cl], whilst back-extraction time is fixed at 10 min; and Figure 2c shows the surface response corresponding of the effects of back-extraction time and amount of [Hmim][Cl], with established HCl volume at 40 μ L. As expected, HCl volume has a negative effect (Fig. 2a and b) and the response of the system increases when the HCl volume decreases. For the back-extraction time, the response of the system increases with the time (Fig. 2a and c) until reaching a

maximum at 14 min. Both, 10 μ L for HCl volume and 14 min for the back-extraction time, were adopted as the optimum conditions for the proposed methodology. As can be seen in Fig. 2b and c, the effect of the amount of [Hmim][Cl] also presents a maximum over 40 mg, although the variation of the response is really slight between 40 and 20 mg. Thus, considering the sign of the effect of this variable obtained in the Plackett-Burman design, which was negative, and the importance of waste reduction, 20 mg of [Hmim][Cl] were finally chosen for the validation of the method.

In summary, the results obtained from the optimization process lead to the following experimental conditions: amount of [Hmim][CI], 20 mg; chelating agent, 40 μ L (2 mg mL⁻¹); ionic strength, 0% of NaCI; pH, the pH of water without any adjustment; HCI volume, 10 μ L; back-extraction time, 14 min; ultrasounds power, 90%; and ultrasounds frequency, 37 KHz.

3.2. Analytical figures of merit

Quality parameters of the proposed method were evaluated. Under optimized conditions, a concentration range from 0.5 to 25 μ g L⁻¹ was studied. Finally, the linear range was established between 0.5 and 10 μ g L⁻¹. The calibration curve was constructed using six concentration levels, evaluated by triplicate. The voltammograms corresponding to the blank and the aqueous standards of concentrations from 0.5 to 10 μ g L⁻¹ are shown in Figure 3. The resulting calibration curve gave a high level of linearity with a correlation coefficient (r) of 0.997 (N=6). The sensitivity of the instrumental measurements estimated by the slope of the calibration curve was (3.0 \pm 0.3) μ A μ g⁻¹ L. The repeatability of the proposed method, expressed as coefficient of variation (CV),

was evaluated by five consecutive analysis of aqueous standards at concentrations of 3 and 10 μ g L⁻¹. CV values of 13% were found in both cases. An enrichment factor of 25 was obtained for the proposed procedure, defined as the slope ratio of the calibration curves with and without preconcentration.

The limit of detection (LOD) was estimated according to the Directive 98/83/EC, on the quality of water intended for human consumption, as the concentration corresponding to a signal that is five times the standard deviation of the blank. The LOD was found to be 0.2 μg L⁻¹, which is lower than the most of the reported work up to now using SPEs (see Table 1), and stands lower than the threshold value established by both, the EPA and the European Union (*i.e.* 2 μg L⁻¹ and 1 μg L⁻¹, respectively). In addition, it is important to point out that the sensitivity and LOD of the proposed method are significantly better than those obtained in a previous work [20] (*i.e.* 0.120 μA μg⁻¹ L and 3.3 μg L⁻¹, respectively) using the same kind of SPCnAuEs under equal conditions but without sample preparation. Therefore, the great but scarcely explored advantages that offer the combination of LPME with electrochemical detection using SPEs have been demonstrated.

3.3 Real-world water samples analysis

The feasibility of the proposed method to determine mercury in real-world water samples was evaluated studying matrix effects. Three water samples (namely tap water, bottled water and river water) were employed for recovering studies. As mentioned before, previous analysis revealed that mercury levels in the samples were under the LOD of the present approach. Three replicated analysis of each water sample were carried out at two different spiking levels (1

and 7 µg L⁻¹). Relative recoveries were calculated as the ratio of the signals found in real and ultrapure water samples spiked at the same concentration level. As can be observed in Table 5, relative recoveries ranged from 97 to 108 % in the three performed water samples, whereas the CV values were between 9 and 15 %. According to these results, it can be concluded that the matrix effects were not significant for the determination of mercury in the three selected water samples.

4. Conclusions

SPCnAuEs have been successfully combined with in-situ IL-DLLME and microvolume back-extraction methodologies for the determination of mercury in water samples, reaching a limit of detection that satisfied the established legal threshold levels and proving its application in real-world water sample analysis.

Higher sensitivity and lower LOD were obtained with the proposed methodology compared to those obtained with the same electrochemical transducers but omitting the sample preparation. Therefore, the great and up to now practically unexplored benefits that offer the combination of miniaturized sample preparation techniques with the electrochemical analysis using SPEs have been demonstrated.

Although the ice-bath, centrifugation and ultrasounds limit the in-field application of the proposed methodology, authors strongly believe in a promising future for the combination of LPME with SPEs as detection methodology within the perspectives of developing inexpensive analytical methodologies with portable options for rapid and on-site measurements.

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Table 1. Comparison of different methods using SPEs for the determination of mercury in water samples.

Electrode	Lineal range	LOD	Real water samples	Comments/Analytical technique (deposition time in parentheses)	Ref.
SPGE	5-30 ng mL ⁻¹	1.1 ng mL ⁻¹	Wastewater and rain water	SWASV (60 s)	[14]
SPE/carbon black	2.5·10 ⁻⁸ -1·10 ⁻⁷ M (5-20 μg L ⁻¹)	5·10 ⁻⁹ Μ (1 μg L ⁻¹)	Drinking water	Indirect determination by amperometric measurements of thiols	[15]
SPBE/MWCNTs	0.2-40 μg L ⁻¹	0.09 µg L ⁻¹	Tap water	SWASV (180 s)	[16]
Carbon NP-based SPEs	1-10 μg L ⁻¹	-	Seawater	Heated electrodes/ SWASV (120 s)	[17]
SPE/gold film	2-16 μg L ⁻¹	1.5 µg L ⁻¹	Tap water	SWASV (120 s)	[18]
SPE/gold film	0.1-0.8 μg L ⁻¹	0.08 μg L ⁻¹	-	Preconcentration step using magnetic nanoparticles modified with thiols/ SWASV (120 s)	[18]
SPCE/gold film	0-100 μg L ⁻¹	0.9 μg L ⁻¹	-	SWASV (120 S)	[19]
SPGOnAuEs	2-50 μg L ⁻¹	1.9 μg L ⁻¹	-	SWASV (200 s)	[20]
SPCNTnAuEs	0.5-50 μg L ⁻¹	0.2 μg L ⁻¹	Tap and river waters	SWASV (200 s)	[20]
SPCnAuEs	5-100 μg L ⁻¹	3.3 µg L ⁻¹	-	SWASV (240 s)	[20]
SPCnAuEs	5-20 ng mL ⁻¹	0.8 ng mL ⁻¹	Rain and river waters, industrial wastewater	SWASV (120 s)	[21]
SPCE/Sumichelate a10R	-	12 pM (0.002 μg L ⁻¹)	Lagoon water	DPASV (300 s)	[22]
CTS-SPE	20-80 ng mL ⁻¹	2 ng mL ⁻¹	-	DPASV (30 s)	[23]

445 SPGE, screen-printed gold electrode; SWASV, square-wave anodic stripping voltammetry; 446 SPBE, screen-printed bismuth electrode; MWCNTs, multi-walled carbon nanotubes; NP, nanoparticles; SPGOnAuEs, screen-printed graphene oxide/gold nanoparticles electrodes; 447

SPCNTnAuEs, screen-printed carbon nanotubes/gold nanoparticles electrodes; DPASV, 448 449

differential-pulse anodic stripping voltammetry; CTS-SPE, chitosan-modified screen-printed

450 electrodes.

 Table 2. Experimental variables and levels of the Plackettt-Burman design.

Variable	Level		
variable	Low (-1)	High (+1)	
Amount of [Hmim][CI] (mg)	20	40	
Chelating agent (µL, 2 mg mL ⁻¹)	20	40	
Ionic strength (NaCl concentration, %, w/v)	0	10	
рН	5	10	
HCl volume (μL)	20	50	
Back-extraction time (min)	5	10	
Ultrasounds power (%)	50	90	
Ultrasounds frequency (KHz)	37	80	

Table 3. Variables and low and high levels, central and star points used in CCD design.

Variable	Level			Star points (α=1.682)	
variable	Low (-1)	Central (0)	High (+1)	-α	+α
HCl volume (μL)	22	40	58	10	70
Back-extraction time (min)	6	10	14	3	17
Amount of [Hmim][CI] (mg)	28	40	52	20	60

 Table 4. Coefficients and p-values obtained in CCD design.

Variable	Coefficient	p-value
HCl volume (C ₁)	-15,429	0*
Back-extraction time (C ₂)	8,581	0,002*
Amount of [Hmim][CI] (C ₃)	-3,668	0,101
$C_1^*C_1$	-0,572	0,869
$C_2^*C_2$	-13,322	0,003*
C ₃ *C ₃	-11,672	0,007*
$C_1^*C_2$	-2,863	0,533
C ₁ *C ₃	-3,429	0,458
$C_2^*C_3$	2,722	0,553

^(*) Significant variables with 95% probability (*i.e.* p-value < 0.05).

Table 5. Relative recoveries and CV values (in parentheses) for the analysis of mercury in real-world water samples.

Water sample	Spiking level			
water sample	1 μg L ⁻¹	7 μg L ⁻¹		
Tap water	106 (11)	108(7)		
Bottled water	98 (11)	103 (15)		
River water	97 (10)	98 (9)		

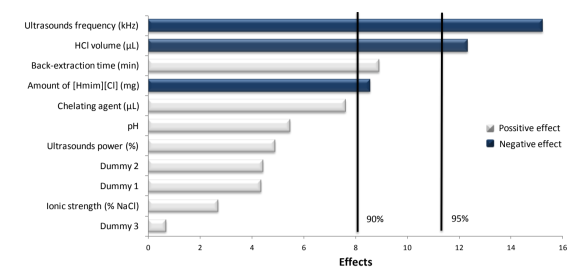


Fig. 1 Pareto chart of Plackett-Burman design.

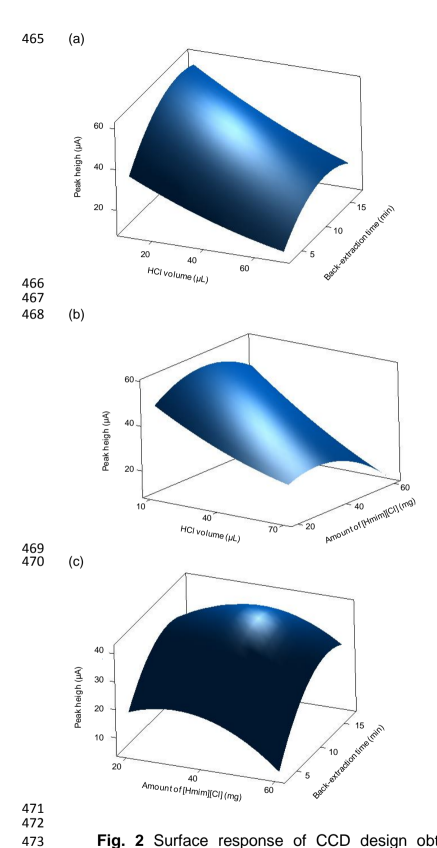


Fig. 2 Surface response of CCD design obtained by plotting: (a) HCl volume *vs.* back-extraction time (amount of [Hmim][Cl]: 40 mg); (b) HCl volume *vs.* amount of [Hmim][Cl] (back-extraction time: 10 min); (c) amount of [Hmim][Cl] *vs.* back-extraction time (HCl volume: 40 μL).

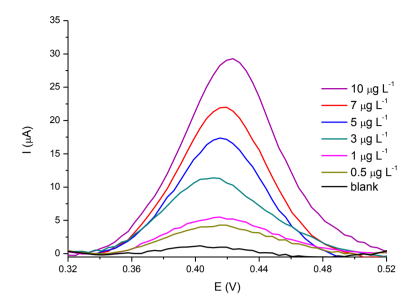


Fig. 3 Square-wave voltammograms, after baseline correction, of a blank and mercury aqueous standards of 0.5, 1, 3, 5, 7 and 10 μg L⁻¹ after in-situ IL-DLLME and back-extraction under optimum conditions.