1	Fully Automatic In-Syringe Magnetic Stirring-Assisted Dispersive Liquid-Liquid
2	Microextraction hyphenated to High Temperature Torch Integrated Sample
3	Introduction System-Inductively Coupled Plasma Spectrometer with Direct
4	Injection of the Organic Phase
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21 Abstract

22 A proof of concept study involving the on-line coupling of automatic dispersive liquid-liquid 23 microextraction (DLLME) to ICP OES with direct introduction and analysis of the organic 24 extract is herein reported for the first time. The flow-based analyzer features a Lab-In-Syringe (LIS) setup with an integrated stirring system, a Meinhard[®] nebulizer in combination with a 25 26 heated single-pass spray chamber, and a rotary injection valve, used as on-line interface 27 between the microextraction system and the detection instrument. Air segmented flow was used for delivery of a microliter fraction of the non-water miscible extraction solvent, 12 µL of 28 29 xylene, to the nebulizer. All sample preparative steps including magnetic stirring assisted DLLME were carried out inside the syringe void volume as a size-adaptable yet sealed mixing 30 31 and extraction chamber. Determination of trace level concentrations of cadmium, copper, lead, 32 model analytes has been demonstrated by and silver as microextraction as diethyldithiophosphate (DDTP) complexes. The automatic LIS-DLLME method features 33 34 quantitative metal extraction, even in troublesome sample matrices, such as seawater, salt, and 35 fruit juices, with relative recoveries within the range of 94-103%, 93-100% and 92-99%, 36 respectively. Furthermore, no statistically significant differences at the 0.05 significance level 37 were found between concentration values experimentally obtained and the certified values of 38 two serum standard reference materials.

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47 Inductively coupled plasma (ICP)-based techniques are deemed the most universal atomic 48 spectrometric techniques for metal assays as they enable detection of practically all metals and 49 metalloids of the periodic table with excellent sensitivity, reproducibility and sample throughput. Besides, continuous improvements of instrumentation and software make ICP-50 based techniques user-friendly for routine analysis. However, limitations of instrumental 51 robustness and background interferences in the analysis of high salt content solutions or samples 52 53 with elevated organic load might jeopardize the reliability of the analytical method. In fact, the occurrence of this kind of matrices might deteriorate the nebulization efficiency, plasma 54 55 electron density, and even lead to plasma torch shutdown. The sensitivity of ICP OES and ICP-MS based methods does not in some instances suffice for the detection of elements at trace 56 57 level concentrations, as might be the case in environmental surveillance studies or health risk/exposure assessment. Several approaches have been developed to overcome or minimize 58 these drawbacks, including sorbent-based analyte preconcentration,¹⁻³ the addition of oxygen to 59 avoid carbon deposition, or the elimination of the sample matrix by electrothermal sample 60 61 vaporization prior to sample injection into the plasma.^{4,5}

62 With regard to sample handling strategies, liquid-liquid extraction (LLE) of hydrophobic metal or oxyanion complexes has proven to be a powerful pre-concentration and clean-up approach 63 for trace metal analysis by graphite furnace (GFAAS) and flame atomic adsorption 64 65 spectrometry.^{6,7} In contrast, measurements by ICP-based techniques require generally in-line desolvation, solvent emulsification, or solvent dilution to yield steady nebulization conditions.^{4,5} 66 Few papers report on LLE with back-extraction of the target species into an aqueous phase as a 67 front end to ICP detection.⁸⁻¹¹ This approach combines the advantages of LLE including salt 68 69 removal and avoiding typical problems of on-line SPE (backpressure, filter blockage, etc.) along with eluate compatibility with the detector. However, both the operational time and, if 70 automated, the instrumental complexity and effort, e.g. to yield reproducible solvent 71 introduction and reliable phase separation, refrained this LLE mode from further 72 development.1,12,13 73

74 As an alternative to matrix elimination, the use of a high efficiency micronebulizer in 75 combination with a heated spray chamber, termed high temperature torch integrated sample 76 introduction system (h-TISIS), has been reported for reliable ICP- assays of complex samples.^{14,15} With the injection of a mere few microliters of sample, matrix effects have showed 77 78 to become insignificant as the temperature of the spray chamber is set at 350°C for fuels and 79 diverse acid digested environmental samples.^{14,15} Moreover, direct analysis of hydrocarbon samples has also proven to be feasible.¹⁴ Readers are referred to a series of reviews describing 80 instrumental aspects and successful applications of this approach for metal/metalloid 81 determination in organic matrices.^{4,5} 82

83 This work was sparked by the consideration that such versatile sample introduction system could be hyphenated to automatic liquid-liquid microextraction for expedient analysis of 84 organic extracts. In this context, the Lab-In-Syringe (LIS) concept^{16,17} has gained considerable 85 attention as a sample handling tool for straightforward and versatile batch-wise automation of 86 87 liquid-phase based approaches. Taken as a sequel of the second generation of flow analysis, also called sequential injection analysis,^{18,19} LIS is featured by carrying out the entire procedure in 88 the void volume of the barrel of a gas-tight automated syringe pump operating as an enclosed 89 90 mixing chamber. Of special impact is the integration of a magnetic stirring bar into the syringe for homogenous sample/reagent mixture and solvent dispersion.^{20,21} 91

92 While there has been significant work harnessing flow-based approaches (mostly flow injection and sequential injection) for automated liquid-liquid extraction of metal species, 6,7,22-25 with 93 potential implementation in microfluidic devices,^{24,26,27} prior to on-line atomic spectrometric 94 detection, reviewed elsewhere,^{3,28,30} just few papers report on employing LIS, whose versatility 95 96 has not been fully explored yet. LIS for metal assays has been merely coupled to atomic 97 absorption spectrometric measurements, namely, mercury microextraction and cold vapor atomic absorption spectroscopy (AAS)^{31,32} and more recently to non-dispersive liquid phase 98 extraction of silver followed by GFAAS,³³ yet studies concerning on-line dispersive liquid-99

liquid microextraction (DLLME) as a front-end microextraction approach to multi-elementalICP OES/MS are still missing.

In this paper, in-syringe DLLME is explored for the first time as a "front-end" versatile 102 103 microextraction platform for ICP-based detection. Diethyldithiophosphate (DDTP) is used as a 104 selective chelating reagent on the basis of its ability of complexing metal species at the usual acidic pH values for sample conservation³⁴ as opposed to its carbamate counterparts, i.e. no 105 106 additional buffering of sample is needed, which, in turn, make the analytical method 107 straightforward (with no need of pH optimization) and less prone to blank contamination. As a 108 consequence of the high stability constants of the DDTP chelates, even in strong acidic 109 conditions, back-extraction methods with increasing of the acidity and/or the addition of 110 competing metal species are proven inappropriate for quantitative recovery of DDTP complexed metals.^{35,36} To tackle this issue, we have exploited h-TISIS as a viable interface for the direct 111 112 injection of the metal containing organic extracts into the ICP system. With this interface, 113 organic matrices are permitted whereby analyte dilution in the back-extraction solution in 114 conventional liquid-phase microextraction approaches of trace metals is circumvented. Cadmium, copper, lead, and silver were chosen as model analytes and analyzed in varied 115 116 environmental and food matrices.

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118 Material and methods

119 *Chemicals and samples*

Ultrapure water was supplied by a three-step ion-exchange system Milli-Q, fed by reverse
osmosis, Elix 3, both from Millipore (El Paso, TX, USA). Isopropanol and xylene (Panreac
Química S.A., Barcelona, Spain) were employed for the cleaning of the syringe barrel and flow
system prior to each extraction and as extraction solvent, respectively. Diethyldithiophosphate
ammonium salt (DDTP, 95 %) was obtained from Sigma Aldrich (Saint Quentin Fallavier,
France) and used as a chelating reagent, prepared in aqueous medium. 65% HNO₃ (Suprapur®,
Merck KGaA, Darmstadt, Germany) was used to prepare washing solutions and acidify the

127 standards and samples. An ICP multielement standard solution (Merck IV, Merck KGaA, 128 Darmstadt, Germany) containing 1000 mg element per litre was used to prepare the standards 129 by serial dilutions. Stock and standard solutions were prepared in 2 % (v/v) HNO₃. Organic multielement standards were prepared by dissolving a certified material (Conostan[®] S-21, 130 131 Conoco Specialty Products, Inc., Ponca City, Oklahoma, USA) in xylene. In order to evaluate 132 the reliability of the automatic system for handling complex matrices, a variety of real samples 133 were analyzed: seawater, salt, salt without sodium, grape juice and apple juice. Salt and juice 134 samples were bought in a local supermarket. Coastal seawater was collected in Alicante using 135 pre-cleaned polyethylene flasks. The sample was taken at an approximately 50 cm depth and stored at 4°C in the laboratory. Salt samples were prepared by dissolving 3.5 g of salt in 10 mL 136 137 of Milli-Q water. All samples were filtered using 0.45 µm nylon syringe filters (Filter-Lab[®], Filtros Anoia, Barcelona, Spain). Two certified lyophilized control serum samples (ClinChek® 138 139 Controls, Recipe[®], Munich, Germany) were used as quality control (QC) materials for 140 evaluation of the trueness of the analytical method. Serum samples were reconstituted in 3.0 mL 141 of ultrapure water with gentle mixing until complete dissolution of the lyophilised material.

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143 Flow setup for automated DLLME

144 The system configuration for lab-in-syringe dispersive liquid-liquid microextraction (LIS-DLLME)-ICP OES assays is illustrated in Fig. 1 and a close up is presented in Fig S1. In all 145 146 experiments, a MicroSIA device from FIAlab Instruments Inc. (Seattle, WA) was used to 147 assemble the flow manifold. It integrates a 30 mm Stroke OEM low pressure Syringe Pump (SP, 148 Cavro XCalibur) and an 8 port selection valve (SV, Vici Valvo) furnished with a PTFE rotor. The MicroSIA system contains two auxiliary supply ports of 5 and 24 V herein utilized for 149 150 stirring activation and ICP triggering. The SP is furnished with a rotary head valve (HV) with 151 three selectable ports (IN, OUT, and TOP) for tubing connections. A 5 mL-glass syringe (30 152 mm lift, 1.45 mm id, Tecan) was used for performing all solution handling including the 153 DLLME procedure inside. A commercial PTFE covered magnetic stirring bar of 14 mm size 154 (4.5 mm diameter) was placed in the syringe barrel. To diminish the resulting dead volume at 155 syringe emptying, the stirrer was flattened by sand papering to 3.5 mm height and made to 156 length in order to fit snugly into the syringe. The stirrer was forced to spin at approximately 800 157 rpm by generating a rotating magnetic field outside the syringe (see Fig. 1 and Fig. S1). To this 158 end, a pile of seven neodymium magnets (each 3 mm x 5 mm Ø) was hot-glued on top of a 159 commercial cooling ventilator (12 VDC supply) serving as a cost-effective brushless motor 160 (wings and protection removed). The motor was connected to the syringe piston bar so that the magnets were leveled with the stirring bar inside the syringe at any time. The motor was 161 162 powered by the 5 V supply port of the MicroSIA and activated (generating a rotating magnetic field) by software control. By careful adjustment of this arrangement, stirring velocities 163 164 exceeding 800 rpm were proven applicable

Lateral ports 2-6 of the SV (see Fig. 1) were connected to 2 % (v/v) HNO₃ (2), isopropanol (3) and 15 % (v/v) HNO₃ (8) for syringe chamber cleaning; extraction solvent (4), sample (5), and complexing reagent (6). Using a very short tube of PEEK piercing a wider silicone tube for drainage, port 1 allowed both syringe content discharge to waste during cleaning but also aspiration of air (see Fig. 1). Air inside the syringe enabled vortex formation by stirring, thus promoting solvent dispersion.

Port IN on the syringe HV was connected to the central port of the SV via a 15 cm long holding
coil (HC, PTFE tube, 1.0 mm i.d.). Port OUT was used to empty the syringe to waste without
passing the HC. The TOP position was connected via a 20 cm transfer line (0.5 mm i.d.) to a
low pressure (PEEK stator and rotor) six-port injection valve (IV) from Vici-Valco (Schenkon,
Switzerland), used as interface between the LIS-based microextraction system and the ICP
OES. A PEEK capillary of 8 cm (0.25 mm i.d.) was used as injection loop, the total injection
volume including the valve rotor channel was estimated as 12 μL.

Instrumental control of the extraction system was done via USB using the open-source software
Cocosoft, version 4.3 (FI-TRACE, University of the Balearic Islands).³⁷ The software is written
in Python programming language and enables the use of variables, loops, routines, and

181 conditionals, and communication via serial interface. Triggering of ICP OES activation and data

182 registration was done by relay contact using the 24 V supply port of the MicroSIA instrument.

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184 *ICP OES measurements*

An Optima 4300 DV Perkin-Elmer ICP OES spectrometer (Uberlingen, Germany) was used as
detection instrument and the emission intensity signals were axially taken. The system was
equipped with a 40.68 MHz free-running generator and a polychromator with an echelle grating.
Table 1 summarizes the operational instrumental conditions.

A glass concentric nebulizer (TR-50-C3, Meinhard[®], Golden, CA) was fitted to a 12 cm³ glass
single pass spray chamber (h-TISIS).³⁸ The h-TISIS was jacketed with a copper coil connected
to a power supply so as to heating the chamber at will. Hereto, the coil temperature was
programmed by means of a thermocouple attached to its surface (Desin Instruments, Barcelona,
Spain).¹⁴

195 Spann).

The solutions were delivered to the nebulizer by a peristaltic pump (Gilson Minipuls3 Model
M312, Villiers-le-Bel, France) and a 0.19-mm i.d. PVC-based material with plasticizer (Tygon[®]
R-3607, Ismatec, S.A.) tubing was employed.

197 An air-segmented flow injection methodology was selected to deliver sample volumes at the 5-198 $15 \,\mu$ L level to the instrument. Air was continuously aspirated by means of a peristaltic pump. At 199 a given time and precisely controlled by software, a sample plug was driven to the nebulizer 200 using a carrier stream of air to avoid sample dispersion. Images of the injection of the analyte-201 containing organic phase into the ICP torch are compiled in Fig S2. With this system, oxygen 202 was not needed to minimize background interferences in troublesome samples because of two 203 facts: (i) the injected sample volume was a mere of a few microliters; and, (ii) the oxygen in the 204 air stream continuously aspirated could boost the total carbon combustion. Therefore, negligible 205 soot deposits were found throughout the present work.

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The analytical workflows are given as supplementary materials (Tables S1 and S2). The DLLME protocol was started by cleaning the syringe with (1) isopropanol to remove any residues of the extraction solvent from the previous extraction, (2) 15% (v/v) HNO₃ and two times with 2% (v/v) HNO₃ to keep the syringe free from metal traces, and (3) with the corresponding sample solution, that is, 2%(v/v) HNO₃ for blank measurements or the sample solution itself from position 5 of the SV.

215 The in-syringe DLLME protocol is performed as follows: 250 µL of air (to promote vortex formation with the consequent solvent dispersion), 270 µL of xylene, 3600 µL of sample, a 20 216 µL air plug (to avoid contact between sample and chelating reagent in the HC), 250 µL of 217 218 reagent solution, and a final volume of 180 μ L air to empty the overall HC content into the syringe barrel were sequentially aspirated. Immediately before the aspiration of the extraction 219 220 solvent, stirring at 800 rpm was activated. After an extraction time of 120 s, the stirring was 221 deactivated for phase separation for 30 s, which allowed the xylene droplets to float and to 222 coalesce. Eight repeated activations of the stirrer for a minimum time (< 1 s, not achieving the 223 final stirring rate) were done to remove any xylene residues, which were stuck on the stirring 224 bar.

In the final step, the organic phase was pushed at 80 μ L s⁻¹ towards the injection valve first to clean the transfer line and push out any residues from the previous injection to waste. Then, aliquots of the solvent (12 μ L) were injected repeatedly into ICP OES by IV activation into the air flow carrying the injected volume to the h-TISIS at a delivery flow rate of 50 μ L min⁻¹. Every organic extract was injected three times for assessing the repeatability of the ICP readouts. Finally, the aqueous syringe content was emptied to waste with the HV in position OUT.

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235 Results and Discussion

236 Investigation of the h-TISIS-ICP OES operational conditions

237 Parameters related to the nebulization and ICP OES measurements including the injection 238 volume of the organic phase, the nebulizer gas flow rate and the spray chamber temperature 239 were evaluated. For injection volumes of xylene larger > 12 μ L, the plasma was unstable and 240 tended to shut down. The nebulizer gas flow rate was also optimized. The evaluated values were 241 in the range of 0.15-0.40 L min⁻¹. It was verified that the optimum nebulizer gas flow rate in 242 terms of sensitivity was 0.26 L min⁻¹. Higher flow rates might not ensure the quantitative 243 evaporation of the solvent in the aerosol phase within the spray chamber because of the short 244 residence times but lower flow rates might lead to excessively big aerosol droplets.

245 The effect of the evaporation chamber temperature on the analytical performance was also 246 investigated. ICP OES signal intensities for Ag, Cd, Cu and Pb were thus recorded at h-TISIS temperatures ranging from 150 to 400 °C. The h-TISIS spray chamber working at temperatures 247 248 > 300°C provided 8, 7 and 12 fold-peak height improvements with respect to those at room 249 temperature for Ag, Cd, Cu and Pb, respectively (see Fig. 2). This was due to the enhancement 250 of the aerosol solvent evaporation inside the chamber and, hence, of the analyte mass delivered 251 to the plasma. The working temperature was set to 350°C because, under these circumstances, non-spectral interferences by the solvent itself were practically neglegible.^{14,15} 252

253 The signal obtained for organic standards with h-TISIS working at the optimum experimental 254 conditions was compared with a conventional introduction system (*i.e.*, cyclonic spray chamber 255 operating at room temperature). The nebulizer gas flow rate employed for the conventional 256 system was 0.4 L min⁻¹. Table 2 shows that h-TISIS readouts were up to 13 fold improved as 257 compared to those of the cyclonic spray chamber. Limits of detection (LODs) were determined 258 according to the $3s_b$ criterion, where s_b was the standard deviation of ten consecutive blank 259 measurements. As expected from the sensitivity data, the highest LODs (Table 2) were obtained 260 for the conventional sample introduction system. It is however important to note that the

- 261 discrepancies observed across the trends in LODs and the analytical readouts are attributed to
- the dependence of the spray chamber design upon the standard deviation of the background.
- 263

264 System configuration and evaluation of the analytical protocol

265 Our experimental setup features significant advances as compared to previous works in the field 266 of LIS.^{20,21} For example, the induction of solvent dispersion by stirring bar rotation did not 267 require any additional "driving device" to generate a rotating magnetic field as reported previously.^{20,21} As the syringe pump was placed here in common up-right orientation, the 268 269 magnetic stirring bar had to move with the piston so that the motor was fixed to the piston bar to 270 assure steady leveling of both motor and stirrer. To reach the required rotation rate of 800 rpm 271 for solvent dispersion, the stirring bar had to turn smoothly inside the syringe. A 15×4 mm 272 stirring bar was thus sandpapered to a 14 mm length (syringe inner diameter was 14.5 mm). 273 Smaller stirring bars (e.g. $10 \text{ mm} \times 2 \text{ mm}$), potentially offering a lower dead volume, were not 274 able to keep up with the required rotation rate but dangle inside the syringe. Due to the inertia of 275 the liquid, the stirring bar is slowed down at the onset of stirring. Thus, a purpose-made control circuit was used for a slow turn-on of the inducing motor.²⁰ The motor then reached its final 276 speed after approximately 5 s, which enabled synchronized rotation of the stirring bar. 277

278 Regarding the analytical protocol for in-syringe DLLME, the following two operational 279 sequences for in-line sequential aspiration of solutions to the syringe were tested: 1: Air, 280 extraction solvent, sample, air, DDTP reagent and air; and, 2: Air, sample, air, DDTP reagent, 281 extraction solvent and air. The segmentation between the sample and the DDTP reagent was 282 done to prevent complex formation already inside the holding coil and the potential sorption of 283 the chelate onto the hydrophobic walls of the flow manifold, which would in turn jeopardize the 284 precision and the analyte recovery and lead to carry-over effects. Air was further found to favor 285 vortex formation with the consequent dispersion of the extraction solvent into tiny droplets. It 286 was demonstrated that the first aspiration sequence was superior in terms of peak height (1.4-1.5 287 times higher signal) and thus was kept further on. Because the extraction solvent was the first

solution introduced into the syringe, smaller droplets were formed, thus enhancing the surfacearea with the subsequent improvement of the extraction efficiency.

290 One disadvantage of the LIS-based extraction system herein proposed is the potential cross-over 291 contamination because of the syringe void volume caused by the stirring bar along with the 292 possibility of sorption of organic phase droplets onto the PTFE bar. Generally, the rinsing of the 293 syringe after extraction is done in three steps; a first cleaning step with isopropanol, to remove 294 organic solvent remnants; a second step with a concentration of nitric acid ranging from 2-15% 295 (v/v) to remove metal leftovers and, finally, with the sample, in order to rinse the system with 296 the sample matrix itself. However, the hydrophobic analyte complexes can further be retained in 297 the tubing and injection valve, potentially leading to carry-over effects. To evaluate the 298 effectiveness of several cleaning protocols (see Table S3), the concentrations of metals in three consecutive blank samples analyzed after a standard of 100 µg L⁻¹ of Ag, Cd, Cu, and Pb were 299 300 determined. Figure S3 shows the percentage of the Ag blank signals in consecutive injections with respect to that obtained at the 100 µg L⁻¹ level. The rinsing protocol capitalizing upon 15% 301 302 (v/v) HNO₃ provided the best performance because signals for the first extraction of the blank 303 corresponded to only 5% of the signal obtained for the 100 µg L⁻¹ standard. Similar results were 304 found for Cd, Cu and Pb. In the remainder of washing protocols using 2-10% (v/v) HNO₃, the 305 first blank signal amounted to as much as ca 20-95% of the initial Ag signal.

306

307 Selection of physical and chemical parameters

308 Volume of the extraction solvent, DDTP concentration and extraction time

309 The volume of the extraction solvent in the automatic LIS procedure is particularly important 310 inasmuch as large volumes facilitate quantitative extraction efficiency while microvolumes 311 (usually a few microliters) are preferable with respect to the improvement of preconcentration 312 factors. Evaluation of the volume of xylene as extraction solvent was performed by comparison 313 of the analytical readouts obtained for volumes in the range of 220 to 320 μ L at the 100 μ g L⁻¹ 314 level. Larger solvent volumes were considered unacceptable for analyte enrichment while 315 smaller volumes of solvent were unlikely to be applicable herein as the system's reliability is 316 based on the premise that the solvent droplets coalesce to one phase so that introduction of 317 droplets of the aqueous phase into the h-TISIS-ICP OES is circumvented. The ICP OES signals 318 were normalized with respect to the maximum peak height (obtained with 270 µL). Figure S4 indicates that the normalized readouts increased with the volume of extraction solvent up to 270 319 320 μ L, with repeatabilities in all instances better than 3%. Similar trends were found for peak area; 321 hence, the analytical signal was taken as peak height throughout. Note that similar behavior was 322 found for all the elements, therefore, Ag and Cd were selected as model analytes for further 323 studies.

324 In DLLME, the higher the interfacial area between immiscible phases is the shorter the 325 extraction time for attaining comparable extraction efficiencies. For a fixed stirring rate (viz., 326 800 rpm), the effect of the stirring time was evaluated. The minimum extraction time to achieve 327 pseudo-equilibrium conditions was estimated at the onset of the curvature of the regression line 328 of the peak height against extraction time for which the analytical readouts approach to steady-329 state conditions. The pseudo-equilibrium conditions were reached at 60-65 s for all the elements 330 under the experimental conditions indicated above. Moreover, it was observed that almost 100% 331 (in absolute mass) of the analytes were extracted in the organic phase for stirring times of 100-332 120 s. For stirring times >100 s the influence of the extraction time was virtually negligible as 333 the peak height remained practically unaltered. However, the intra-day precision improved with 334 the extraction time, reaching RSD values lower than 5% at 120 s. An extraction time of 120 s 335 was therefore chosen for the remaining work. The concentration of the extraction agent was also 336 evaluated. Figure S5 indicates that peak heights increased with DDTP concentration up to 50 337 mmol L⁻¹, which was selected for the remainder of the experiments.

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339 *Effect of the acid and counter ion on the extraction procedure*

340 The effect of the acid nature and counter ions on the extraction efficiency of target metals was 341 evaluated. Hence, a cohort of six standards was prepared with the same metal concentration but 342 with increasing concentrations of strong acids (HCl or HNO₃) to evaluate the potential salting-343 out effects and metal complexation. The matrix composition was: 0.21, 0.51 or 1.03 mol L⁻¹ in HNO₃ or HCl. According to previous researchers,²¹ the effect of the two counter anions as 344 345 interfering species for DDTP extraction was not statistically significant (Fig. S6). With respect 346 to the acidity of the sample matrix, a loss of signal intensity was observed at the concentration level of 1.03 mol L⁻¹ regardless of the acid nature. For nitric acid, 6% and 12 % signal losses 347 348 were observed for Ag and Cu, respectively. On the other hand, a 7% loss of peak height was 349 observed in both cases for 1.03 mol L⁻¹ HCl.

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351 Analytical method performance

352 Under the selected experimental conditions, a linear correlation of peak height against analyte 353 concentration in aqueous medium subjected to automatic DLLME was observed. The 354 calibration was performed using six concentration levels in aqueous phase from 0.4 up to $11 \mu g$ 355 L^{-1} with an injection volume of 12 µL of organic phase. Coefficients of determination (R^2) 356 higher than 0.9991 were obtained for five inter-day calibration curves. As a benchmark of inter-357 day precision, relative standard deviations were 5, 7, 4, and 8 % for the slopes of the calibration 358 curves of Ag, Cd, Cu, and Pb, respectively. Moreover, no outlying measurements (> three times 359 the standard error of the slope) were found. LODs were calculated according to the $3s_b$ criterion 360 (n=10), and in all instances were lower than 0.1 μ g L⁻¹. LOQs were 0.16, 0.14, 0.14 and 0.21 μ g L⁻¹ for Ag, Cd, Cu, and Pb, respectively. Repeatability values for six consecutive analysis of a 361 2.0 µg L⁻¹ aqueous standard were 3.1, 4.0, 2.8 and 3.9 % for Ag, Cd, Cu and Pb, respectively. 362 363 An alternative calibration method was also tested. In this case, organic standards (12 μ L) were

364 introduced directly to the ICP OES following the air-segmented injection methodology 365 described above. Organic standards were prepared using xylene as a diluent of the certified

reference material Conostan[®] S-21. Coefficients of determination (R²) higher than 0.9993 were 366 367 obtained for five calibration curves within the concentration range spanning from 5-170 µg/L on 368 5 subsequent days. The inter-day precision in terms of sensitivity was similar to that of the 369 procedure with aqueous standards followed by DLLME. Notwithstanding the deterioration in 370 sensitivity (see Table 3) as the organic standards in this second external calibration method are 371 not subjected to preconcentration, LOQs were not proportionally increased because of the 372 deterioration of the blank repeatability values for the LIS-DLLE method. Repeatability values for six consecutive analysis of a 25 μ g L⁻¹ organic standard were were 2.1, 3.4, 2.7 and 4.2 % 373 374 for Ag, Cd, Cu and Pb, respectively.

The preconcentration factor was obtained as the ratio of the slope of the straight line regression following the automatic LIS extraction procedure to that obtained by direct injection of organic standards into h-TISIS-ICP OES. Table 3 compiles the sensitivities of both calibration curves. The nominal pre-concentration factor was estimated from the ratio of the sample volume (3.60 mL) to that of the organic solvent (270 μ L), that is, 13.3. Table 3 shows that the experimentally obtained pre-concentration factors were similar to the nominal value, thus signalling that the extraction efficiency for all the metals was close to 100%.

The entire automatic LIS procedure, including mixing of the sample and reagents, extraction, phase separation, measurement and system cleaning, lasted ca. 375 s, which gives rise to a sample throughput of 9 h⁻¹. The cleaning protocol using 1.2 mL of isopropanol lasted 15 s. Shortening of the rinsing time could most likely be effected by replacing the rotary valve by a low-dead volume stainless steel stator and rotor so as to minimize carry-over effects.

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388 Analysis of real samples

With the aim of validating the extraction methodology, five real samples including seawater, salt, salt without sodium, grape juice and apple juice were analyzed by LIS-DLLME. To this end, a given aliquot was spiked with 2.0 μ g L⁻¹ of a multi-elemental solution in the aqueous phase. Consequently, the analytical concentration in the organic phase after the preconcentration step was around 25 µg L⁻¹. Note that the non-spiked samples were also analyzed. Original metal
concentrations are summarized in Table S4.

Table 4 (right) lists the relative recoveries for Ag, Cd, Cu and Pb, which were close to 100% in 395 all the cases. It can therefore be concluded that additive or multiplicative matrix effects for any 396 of the tested samples, even for typically not applicable samples of high salt content, were 397 398 insignificant. Recovery values were also calculated using a calibration curve obtained by direct 399 injection of the organic standards into the ICP (see Table 4 left). In this case, the concentration of the organic standards was divided by the preconcentration factor and used as X-axis data with 400 401 the ICP OES readouts as Y-axis for direct analysis of the spike recoveries in the aqueous phase. 402 Experimental results compiled in Table 4 demonstrated that both external calibration methods 403 provide comparable metal recoveries for all the samples with troublesome matrices. It is 404 important to point out that there is no need to subject the aqueous standards to the DLLME 405 procedure to get reliable results as the target metals regardless of the matrix composition were 406 quantitatively extracted in the organic phase.

407 For further QC/QA assessment, two serum reference materials, differentiated by the level of 408 metal concentration, were analyzed by LIS-DLLME. For further QC/QA assessment, two serum 409 certified reference materials (CRM), differentiated by the level of metal concentration, were analyzed by LIS-DLLME. Statistical assessment of experimental data for the CRMs was 410 done by comparison of the difference between the certified and the measured values 411 against the associated expanded uncertainty (U_{Δ}) because the number of accepted sets of 412 data is not provided in the CRM report. The absolute difference (Δ_m) between the mean 413 measured value (c_m) and the mean certified value (c_{CRM}) is calculated according to 414 equation 1. The combined uncertainty (u_{Λ}) was calculated, based on equation 2, from 415 the uncertainty of the certified value (u_{CRM}) and the standard deviation (s_m) of the 416 experimental data. The expanded uncertainty U_{Δ} for a confidence level of 417 approximately 95 % is obtained by multiplying the combined uncertainty (u_{Δ}) by a 418

419 coverage factor (k) equal to 2 (Equation 3). To evaluate the method performance, Δ_m 420 was compared against U_{Δ} . Because Δ_m is in all cases $\langle U_{\Delta}$, no statistically significant 421 differences were found at the 95% level between the values obtained experimentally and 422 the certified concentrations for any of the target elements (see Table 5 and Table S5).

423

424
$$\Delta_m = |c_m - c_{CRM}|$$
 Equation 1

425
$$u_{\Delta} = \sqrt{s_m^2 + u_{CRM}^2}$$
 Equation 2

426
$$U_{\Delta} = k u_{\Delta}$$
 Equation 3

427

428

429 Conclusions

430 In this work, a novel approach capitalizing on a portable flow setup has been proposed for the 431 first time for the coupling of automatic in-syringe magnetic stirring-assisted dispersive liquid-432 liquid microextraction to ICP spectrometry for direct analysis of metal laden organic extracts 433 using an h-TISIS-based total sample consumption system. With this miniaturized sample 434 introduction system, negligible matrix effects were observed in the analysis of carbon-435 containing matrixes. Because of the high stability constants of DDTP-metal chelates, back-436 extraction to aqueous phase for conventional ICP measurements in the aqueous phase is proven 437 unfeasible. Using a univariate optimization strategy suitable experimental conditions were found for DLLME-h-TISIS-ICP OES detection of trace level concentrations of target elements 438 439 in troublesome samples with enrichment factors of ca. 13. Limits of detection found for two 440 distinct calibration procedures were: 0.05, 0.04, 0.04 and 0.06 µg L⁻¹ for Ag, Cd, Cu and Pb (extraction procedure) and 0.07, 0.09, 0.06 and 0.10 µg L⁻¹ for Ag, Cd, Cu and Pb (direct 441 442 injection of standards) respectively, allowing its successful application to the analysis of 443 certified serum materials and spiked environmental samples and beverages. Efficiencies of extraction were close to 100 % with repeatabilities usually down to 8%. Therefore, external calibration can be streamlined by direct injection of organic standards into the h-TISIS-ICP detector system with no need to subject them to the extraction procedure. Further work is underway to expand the scope of the hyphenated LIS-DLLME-h-TISIS-ICP system for detection of bioaccessible metals, metalloids and organometallic compounds in complex foodstuff and soil extracts.

450

Supplementary Information. Additional experimental data and information includes (i) Images of the flow setup and plasma characteristics, (ii) Readouts of cleaning procedures and operational steps, (iii) Effect of volume of organic phase on the analytical readouts, (iv) Effect of chelating reagent concentration on the analytical readouts, (v) Effect of acid type and concentration on the analytical readouts, (vi) Detailed analytical procedure and cleansing protocol, (vii) Concentration of targeted species in the real samples and (viii) Statistical analysis of experimental data for CRM.

458

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Table 1. Operating conditions of the ICP OES furnished with h-TISIS for injection of organic samples

Variable	Value
Injected sample volume [µL]	12
Nebulizer gas flow, $Q_g [L min^{-1}]$	0.26
Outer gas flow [L min ⁻¹]	15
Intermediate gas flow [L min ⁻¹]	1.0
Rf power [kW]	1.35
Integration time [ms]	25
Sampling time [s]	1
Plasma viewing mode]	Axial
Temperature spray chamber [°C]	350
	Ag 328.068
Elements and Wavelengths [nm]	Cd 228.802
Elements and Wavelengths [nm]	Cu 324.752
	Pb 220.353

	ł	n-TISIS	Þ	Conven	tional s	system ^Φ	Peak height ^(h-TISIS) /	LOD ^(Conventional) /		
	Peak	RSD	LOD	Peak	RSD	LOD	Peak height ^(Conventional)	LOD ^(hTISIS)		
	height	(%)	$(\mu g L^{-1})$	height	(%)	$(\mu g L^{-1})$	reak neight	LOD		
Ag	6.1×10 ⁵	2.4	0.6	5.0×10 ⁴	0×10 ⁴ 11.2		12	4		
Cd	1.4×10 ⁴	7.2	0.4	1.3×10 ³	9.5	3.6	11	10		
Cu	8.1×10 ⁵	2.7	0.5	6.1×10 ⁴	1.6	1.9	13	4		
Pb	1.4×10 ⁴	4.6	0.4	1.4×10 ³	10.3	2.1	10	5		

Table 2. Peak height and LODs obtained for the h-TISIS compared against those obtained for the conventional system.*

* Metal concentration: 100 μ g L⁻¹ in xylene. Injected volume: 12 μ L. Qg (h-TISIS): 0.26 L min⁻¹, Qg (Conventional system): 0.40 L min⁻¹. Φ 10 replicates.

477 Table 3. Slopes of the calibration curves by the automatic LIS-DLLME procedure and the

478 direct injection of organic standards along with the experimental pre-concentration

479 factors

standards - LIS-DLLME procedure (L μg ⁻¹)	standards - Direct injection (L μg ⁻¹)	factor
1.1×10 ⁵	8.1×10 ³	13.6
1.7×10^{3}	0.13×10^{3}	13.1
7.9×10 ⁴	5.9×10 ³	13.4
1.9×10 ³	0.14×10^{3}	13.5
	procedure (L μg⁻¹) 1.1×10 ⁵ 1.7×10 ³ 7.9×10 ⁴	procedure (L μg ⁻¹) injection (L μg ⁻¹) 1.1×10 ⁵ 8.1×10 ³ 1.7×10 ³ 0.13×10 ³ 7.9×10 ⁴ 5.9×10 ³

480

			Standar	rds: Di	rect inje	ection*	*	Standards: Extraction procedure [#]								
Samples	Ag		Cd		Cu		Pb		Ag		Cd		Cu		Pb	
	Mean	RSD (%)	Mean	RSD (%)	Mean	RSD (%)	Mean	RSD (%)	Mean	RSD (%)	Mean	RSD (%)	Mean	RSD (%)	Mean	RSD (%)
Seawater	94	1.4	96	1.1	103	0.5	95	0.6	95	1.4	97	1.1	103	0.5	96	0.6
Salt A	98	1.1	99	0.6	95	0.2	94	0.3	99	1.1	100	0.6	97	0.2	95	0.3
Salt B (Without Na)	96	1.2	98	1.1	96	1.1	93	2.0	97	1.2	100	1.1	97	1.1	94	2.0
Apple juice	98	0.9	95	1.1	97	1.2	94	1.0	99	0.9	96	1.0	98	1.2	96	1.0
Grape juice	97	0.3	92	2.0	97	1.1	97	0.7	97	0.3	93	2.0	98	1.1	98	0.7

Table 4. Relative recoveries (%) for complex samples using the LIS-DLME-h-TISIS-ICP OES system

* The standards were prepared in xylene and directly injected in triplicate into the h-TISIS-ICP OES without the use of the extraction procedure.

[#] The standards were prepared in Ultrapure water, then analyte extraction was performed into xylene (in triplicate) and, finally, a small volume of each extract (in triplicate) was injected into the h-TISIS-ICP OES

-			Serum -	Level I	Serum - Level II ^Φ							
	I	Ag	Cd		Cu		Ag		Cd		Cu	
	Mean s		Mean	S	Mean	S	Mean	S	Mean	S	Mean	S
	$(\mu g L^{-1})$	(µg L ⁻¹)	(µg L ⁻¹)	$(\mu g L^{-1})$	(µg L ⁻¹)	(µg L ⁻¹)	(µg L ⁻¹)	(µg L ⁻¹)	(µg L ⁻¹)	$(\mu g L^{-1})$	(µg L ⁻¹)	$(\mu g L^{-1})$
Extraction procedure*	9.29¥	0.09	2.2 [¥]	0.01	0.775 [¥]	0.002	47.3 ^Φ	0.2	4.62 ^{Φ}	0.01	1.23 Φ	0.01
Direct injection [#]	9.49	0.09	2.2	0.02	0.781	0.003	47.5	0.2	4.63	0.01	1.22	0.02
Certified value*	9.85	2.00	2.28	0.47	0.801	0.122	48.0	9.8	4.54	0.93	1.34	0.20

Table 5. Concentrations for the reconstituted certified serum samples as obtained by the automatic LIS-DLLME procedure

*The standards were prepared in Ultrapure water, and analyte extraction was performed into xylene (in triplicate). A small volume of the extract (in triplicate) was injected into the h-TISIS-ICP OES.

[¥] The calibration was performed using seven concentration levels of aqueous standards ranging from 0.3 up to 11 μ g L⁻¹.

[•] The calibration was performed using eight concentration levels of aqueous standards ranging from 1 up to 15 μ g L⁻¹. For Ag determination, the sample was 1:4 diluted with Ultrapure water.

[#] The standards were prepared in xylene and directly injected in triplicate into the h-TISIS-ICP OES without applying the extraction procedure. The calibration was performed using ten concentration levels of organic standards ranging from 0.5 up to $170 \ \mu g \ L^{-1}$.

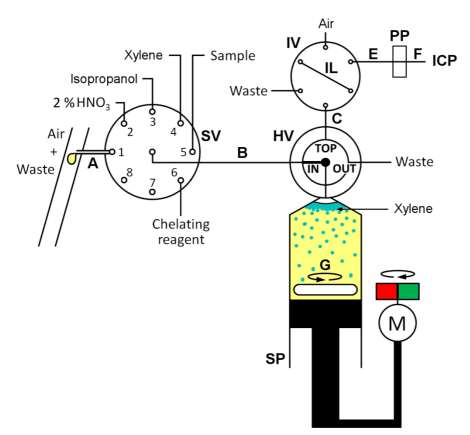
* The standard deviation was estimated as the combined standard uncertainty with a coverage factor of 1.96 at the 95% confidence level.

Figure captions

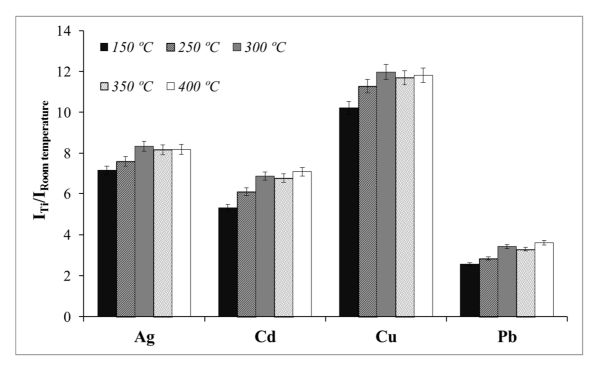
Figure 1. Outline of the automatic and miniaturized LIS-DLLME system. HV – Head valve (of syringe, positions IN, OUT, and TOP), IV – Injection valve, IL – Injection loop, 8 cm, 0.25 mm i.d., M – DC motor, PP – Peristaltic pump, SP – Syringe pump, SV – Selection valve. Tube dimensions: A – 5 cm, 0.8 mm i.d., B – 15 cm, 1.0 mm i.d., C – Transfer line 20 cm, 0.5 mm i.d., E – 20 cm, 0.25 mm i.d. (PEEK), F – red-orange peristaltic/elastic tube, 40 cm, 0.16 mm i.d., G – Magnetic stirring bar.

Figure 2. Normalized peak height with respect of that obtained at room temperature for different analytes and h-TISIS temperatures. Metal concentration: 100 μ g L⁻¹. Injected volume: 12 μ L xylene. Q_g: 0.26 L min⁻¹.









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