Automation in multiresidue analysis of pesticides using on-line solid-phase extraction and liquid chromatography

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Automatic devices which couple on-line the sample pretreatment by solid-phase extraction and the liquid chromatographic separation in one analytical run are nowadays commercially available. This represents a fast, modern and reliable approach for monitoring pesticides over a wide range of polarity.

Solid-phase extraction simplifies the sample pretreatment step, but there is still some evaporation and transfer of the extract. The on-line SPE-LC-UV diode array detection (UV-DAD) system is being used routinely in many laboratories and an increasing number of applications is currently described using on-line SPE-LC coupled to various interfaces and MS detection [1]. Important advantages in coupling on-line the pretreatment and the LC separation and detection are a decrease of the risk of contamination of the sample and sample extract, the removal of analyte losses by evaporation and finally the transfer and the analysis of the totality of the extracted species. In contrast to off-line SPE, where only an aliquot of the extract is injected into the chromatograph, the analysis of the complete sample allows the sample volume to be dramatically reduced to less than 5 – 10 mL when SPE is coupled to GC or to 100 – 150 mL when it is coupled to LC. It was surprising to see that GC has been the preferred method of environmental chemists for a long time, but that on-line systems using LC became the first robust on-line techniques. This is explained by the good compatibility of the LC aqueous mobile phases with the SPE of the samples. On-line coupling of SPE with GC is more delicate because of the inherent incompatibility between the aqueous part of the SPE step and the dry part of the GC system. Although not reported here, much work has been done in this area and automated devices have recently become available.

On-line set-up and precolumn design

The on-line set-up coupling SPE to LC is particularly easy to perform in any laboratory, and has been described extensively in general reviews [1-9]. The apparatus is represented in figure 1 [10]. The trace-enrichment is carried out on a small precolumn, generally made of stainless steel in order to be pressure-resistant, which is placed at the sample-loop position of a six-port liquid switching valve. A solvent delivery unit (SDU) provides the solvents necessary to purge, wash, and activate the precolumn. After conditioning, sample application, and eventual clean-up via the pump of the SPE part, the precolumn is placed in front of an analytical column by switching the valve into the “inject” position. The trapped compounds are then eluted directly from the precolumn into the analytical column by a suitable mobile phase allowing the chromatographic separation of the extracted analytes. Quantitative results of good accuracy can be expected as there is no sample manipulation between the preconcentration and the analysis. The LC system is often run in the reversed-phase mode, using C18 analytical columns because the mobile phase is a partly aqueous solvent mixture. Therefore, residual water in the precolumn after the preconcentration of aqueous samples does not have to be removed before the desorption. The addition of a second switching valve allows both direct injection onto the analytical column and preconcentration via the precolumn. Automation is very easy and several devices are now commercially available (such as the Prospekt from Spark Holland, and OSP-2 from Merck). In these systems, a new disposable precolumn is used for each run and the exchange is automatic. The whole sequence can be programmed and can be performed on a sample whilst the on-line analysis of a previous sample occurs [10-18]. This full automation has been used for on-site monitoring of pesticides in surface waters as part of an early warning alarm system [8,13,17-20]. These studies have contributed greatly in demonstrating that on-line SPE-LC is a robust and reliable technique that can be applied routinely in the field.

The quality of the coupling can easily be controlled by comparing chromatograms obtained by direct injection with those obtained by on-line preconcentration via the precolumn. The dimensions of the precolumn should be adapted to those of the analytical column and are typically 2 – 15 mm long and 1 – 4.6 mm I.D. for a classical 15 – 25 cm long analytical column. The size of the precolumn is an important parameter because the profile of the concentrated species transferred from the precolumn to the analytical column should be as narrow as possible at the beginning of the separation in order to avoid band-broadening. Although it was first recommended to pack precolumns with 5 – 10 µm packings [5], the trend now is to use 15 – 40 µm packings in order to have a high sampling rate during the loading of the sample and to prevent clogging with, for example, a surface-water sample. Despite the granulometry, one is recommended to use LC-grade- and pressure-resistant sorbents. Prepacked precolumns with different sorbents are now available from various manufacturers. It is also easy to pack a precolumn in the laboratory, thus allowing the potential of using new sorbents.
Backflush-desorption of the precolumn should give the least amount of extra band-broadening but has also the drawback of creating problems of clogging of the analytical column when real samples are used. In the forward-desorption mode, the precolumn has the additional role of acting as a guard column and thus preserving the life-time of the analytical column.

Selection of the sorbent in the precolumn

The compatibility between the sorbent and the precolumn is important. The most efficient system is ideally obtained for a precolumn and an analytical column of the same nature [5]. But, a serious limitation of SPE-LC systems is that they use small precolumns which therefore contain a small amount of sorbent. In contrast to off-line SPE where there is the possibility of increasing the breakthrough volumes, $V_b$, by increasing the amount of sorbent in the cartridge, in on-line SPE, when compounds are poorly retained the only solution is to select a more retentive sorbent. For an average LOD of 5 ng, if one wants determination in water at a concentration of 1 µg/L, a sample volume of 5 mL will be sufficient. In real water samples, the presence of humic substances and other contaminants will often require the handling of a 10-fold higher volume. When determination at levels of 50 ng/L are required, the sample volume should be increased to at least 100 – 150 mL.

Table I compares the recoveries obtained when handling 200 mL samples through commercial precolumns of 10 mm × 2 mm I.D. packed with $C_8$, $C_{18}$ and an apolar styrene divinylbenzene (SDB) copolymer PLRP-S [21]. The limitation of using $C_{18}$ silica in conventional on-line precolumns for the preconcentration of the more polar pesticides is demonstrated by low recovery data. The classification of pesticides by increasing hydrophobicity is a great help in rapidly estimating whether $C_{18}$ will be the appropriate sorbent, depending on the trace-level required [1,2,22].

As expected the SDB polymer provides the highest recoveries. This polymer is often selected for multiresidue analysis containing pesticides over a wide range of polarity. Figure 2 shows the chromatograms corresponding to the one-line analysis of 150 mL of a drinking water sample originating from ground water, non-spiked in figure 2a and spiked with 0.1 µg/L of a mixture containing 21 pesticides with a wide range of polarities [10]. Although breakthrough has occurred for de-ethylatrazine, there was no problem in identifying de-ethylatrazine at a concentration of 0.09 ± 0.01 µg/L in such waters. De-isopropylatrazine is part of polar solutes which become difficult to determine at the 0.1 µg/L level, owing to its too early breakthrough. Another advantage of copolymer sorbents over $C_{18}$ silica is that they can be used in the pH range 1–13.

When pesticides over a wide range of polarity are analysed on-line losses can occur for the more polar ones, as results of breakthrough, but losses can also occur for the more apolar one as results of adsorption in connection and sample bottles, as in off-line methods. One example was reported in the screening of organophosphorus pesticides [19]. This class of compounds is often analysed by GC but contains some thermolabile compounds. With the handling of 100 mL samples and using a PLR-S precolumn, excellent recoveries were obtained for the medium polarity compounds and lower recoveries were observed for the more polar compounds such as monocrotophos (15%) and vamidothion (70%) due to breakthrough. But, recoveries around

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Table I. Percent recovery of pesticides using cartridges packed with different phase material (sample volume 200 mL, precolumn size: 10 × 2 mm I.D.). Adapted from reference [21].

<table>
<thead>
<tr>
<th>Compound</th>
<th>$C_8$</th>
<th>$C_{18}$</th>
<th>PLRP-S</th>
</tr>
</thead>
<tbody>
<tr>
<td>De-isopropylatrazine</td>
<td>15</td>
<td>30</td>
<td>55</td>
</tr>
<tr>
<td>Desethylatrazine</td>
<td>20</td>
<td>32</td>
<td>60</td>
</tr>
<tr>
<td>Metoxuron</td>
<td>20</td>
<td>56</td>
<td>95</td>
</tr>
<tr>
<td>Simazine</td>
<td>25</td>
<td>60</td>
<td>97</td>
</tr>
<tr>
<td>Monuron</td>
<td>10</td>
<td>35</td>
<td>98</td>
</tr>
<tr>
<td>Chlortoluron</td>
<td>45</td>
<td>80</td>
<td>97</td>
</tr>
<tr>
<td>Atrazine</td>
<td>75</td>
<td>85</td>
<td>97</td>
</tr>
<tr>
<td>Isoproturon</td>
<td>75</td>
<td>81</td>
<td>92</td>
</tr>
<tr>
<td>Diuron</td>
<td>50</td>
<td>75</td>
<td>95</td>
</tr>
<tr>
<td>Metobromuron</td>
<td>35</td>
<td>75</td>
<td>97</td>
</tr>
<tr>
<td>Terbutylazine</td>
<td>70</td>
<td>90</td>
<td>97</td>
</tr>
<tr>
<td>Linuron</td>
<td>80</td>
<td>89</td>
<td>95</td>
</tr>
<tr>
<td>Metolachlor</td>
<td>80</td>
<td>89</td>
<td>95</td>
</tr>
<tr>
<td>Neburon</td>
<td>85</td>
<td>90</td>
<td>97</td>
</tr>
</tbody>
</table>

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Figure 1. On-line set-up. 1: LC switching valve, 2: precolumn, 3: switching valve of the solvent delivery unit, 4: preconcentration pump, 5: LC pump, 6: analytical column, 7: detector.
80% were obtained for the more apolar carbophenothion and bromophos ethyl due to adsorption problems in the inner walls of the preconcentration system. This underlines the inherent problem in analysing compounds too large and a range of polarity because adsorption can be minimised only by adding a surfactant or an organic solvent which has the effect of increasing the losses of the more polar compounds.

New SDB with high specific surface areas have been shown to strongly retain polar analytes [23,24]. The problem is that the on-line coupling with a C$_{18}$ analytical column is difficult for on-line analysis of very polar analytes because the difference in retention by the two sorbents is too large.olar com pounds can only be separated on C$_{18}$ columns whereas this did not appear using a C$_{18}$ analytical column. This is due to the carbon inherent problem in analysing compounds over too large an injection volume or too long an injection time at the higher flow rates. Porous graphitic carbon is of interest because of its suitability to trap very polar analytes [26]. But some band broadening occurs when coupling a PGC precolumn to a C$_{18}$ column for the same reasons as those occurring when coupling a SDB precolumn. This is not due to the carbon because on-line systems using a PGC precolumn and an analytical PGC column have been described [25,27-29].

Ion-exchangers should be appropriate sorbents for selectively trapping ionic pesticides. Cation-exchangers have been used for the on-line preconcentration of aniline derivatives [30]. However, it was shown that when a sample is percolated through cation-exchanger sorbents and one is looking for less than 1 µg/L of an organic cation, the much higher amounts of inorganic cations which are present in natural waters rapidly overload the cation-exchanger capacity. A chemical clean-up pretreatment consisting of oxalate precipitation of calcium ions, then EDTA complexation of metal ions, has been described for removing most of the inorganic cations before preconcentration [30]. Nevertheless, over-loading still occurs rapidly, as demonstrated by the preconcentration study on aminotriazole [31]. When this analyte was dissolved in deionized LC-grade waters, the breakthrough was measured as 150 mL, using a 1 cm $\times$ 0.2 cm I.D. precolumn packed with a polymer-based cation-exchanger. It was below 30 mL with drinking water samples, after the chemical pretreatment described above. In other work, a low volume -10 mL- was percolated through an ion-pair-exchanger sorbent for the determination of aniline and chloridazon, without chemical pretreatment for the removal of anionic inorganic compounds [32].

**Precolumns in series**

The on-line configuration usually includes a single precolumn. However there are many examples where more selectivity can be obtained by using two precolumns. For compounds that are ionizable in the pH range 2 – 10, direct percolation of samples through ion-exchanger sorbents can be avoided by using a two-step preconcentration. It is based on the fact that solutes are retained on a SDB sorbent when in their neutral form but not in their ionic form. This approach was applied to chlorotriazines and their hydroxylated derivatives which have ionization constants around 2 and 5 respectively [33]. The water sample, adjusted to pH 7, was percolated through a single SDB (PRP-1) precolumn. Then, this precolumn was coupled to a second one packed with the cation-exchanger precolumns and a small volume of deionized water containing 25% acetonitrile, adjusted to using mobile phases which contain a high proportion of water but which are unable to desorb the polar analytes trapped on the SDB precolumn [24]. More retentive analytical columns should be used to solve this problem. Recent commercially available polymeric columns were shown to be efficient, but their specific area is low and consequently, for very polar compounds, retention factors are very similar to those obtained with C$_{18}$ silica columns. The possibility of coupling precolumn packed with high surface area SDB with an analytical porous graphic analytical columns allowed the determination of clopyralid, oxamyl, dicamba, monocrotophos and other very polar pesticides at the 0.1 µg/L level from 100 mL samples [24].
Pesticide analysis

Anion-exchangers have also been combined with PRP–1 for a two-step preconcentration. Applications to phenol and phenoxyacid herbicides have been presented [34-36].

Reproducibility and robustness of the method

One advantage of automation in on-line preconcentration is that more reproducible results can be expected, provided the precolumns are packed with the same amount of sorbent and have the same efficiency. The overall reproducibility of the method includes both the reproducibility of the preconcentration and of the LC system. The repeatability of peak-areas and heights obtained by direct-loop injections into the analytical column has been studied, using an acetonitrile gradient for the analytical separation. The relative standard deviation (RSD) was between 3 and 7%, and 3 and 5% for measurements of peak areas and peak heights, respectively [14]. In the same study, the reproducibility between cartridges was measured by preconcentrating 50 mL of LC-grade water spiked with 0.5 µg/L of pesticides, using a Prospekt system with a new precolumn packed with the PLRP-S copolymer in each run. The RSD was around 10% (n = 5) for measurements of both peak areas and peak heights. RSD values below 10% have been also confirmed in other studies, thus indicating that the precolumns were packed under reproducible conditions. The flow-rate applied for the preconcentration varied from 1 to 5 mL/min and the same average 10% RSD was observed [18].

In the framework of the Rhine Basin programme, an automated LC monitoring system (SAMOS-LC, or System for Automated Monitoring of Organic compounds in Surface waters) has been studied extensively. The procedure includes the loading of 100 – 150 mL of surface water onto PLRP-S precolumns of a Prospekt device at 5 mL/min. The on-line analysis is carried out using a C18 analytical column with an acetonitrile gradient at pH 3. The data are automatically evaluated, with the production of a report for compounds present at, or above, a certain concentration level between 1 and 3 µg/L [18]. The reproducibility of the retention times with a set of 25 – 30 pesticides was excellent with a RSD value of 0.2 – 1.5% (n = 20). At an analyte concentration of 1 µg/L, the RSD of peak areas was in the range 1 – 15%, with a new precolumn in each run. The highest RSD were observed only for analytes eluting between 12 and 25 min and were explained partly by matrix interferences and partly by breakthrough of the more basic compounds on the PLRP-S cartridges [13]. The SAMOS system was made to act as an early warning system for use in the field. The robustness of the system was studied in two laboratories during 5- and 7 month periods. No major problem was encountered for over 1000 analyses, apart from the exchange of a deuterium lamp and clogging of the preconcentration system with non-filtered waters.

Validation

The most appropriate means for testing the accuracy of results given by an analytical procedure including a sample pretreatment should use natural matrix reference materials which are similar to environmental ground- or surface-waters. These materials are now under study but are not yet available [37]. The actual quality control of analyses can only be performed through interlaboratory calibrations. Lacorte et al. [14,38,39] were the first to validate an automated on-line solid-phase extraction system for the first time by participating in the Aquacheck interlaboratory comparison study organized by the WRC (Medmenham, UK) where more conventional sample preparation methods and gas few. Several means of quantitation can be used. With on-line systems, it is not advisable to carry out quantitative analysis by comparison with direct injections. First, the volume of many injection loops is specified to an average accuracy of 20% and calibration of a loop is a rather delicate and time-consuming operation. This does not have to be considered with off-line procedures because the same loop is used for both analysis of unknown extracts and construction of calibration curves. Secondly, slight but imperceptible band-broadening may occur.

For the above reasons, any quantitation method (calibration curves, standard addition, etc.) should be performed using the whole procedure, i.e. with the same experimental conditions (same types of precolumns, sample volume, analytical column, and on-line gradient elution) as selected for the analysis of unknown water samples. Therefore, it is not necessary to know the recovery of the extraction process for each analyte. When possible, it is better to handle a sample-volume lower than the lowest breakthrough volume for more reproducible results. However, when multiresidue analyses are carried out, the sample volume is selected in order to detect most of the compounds at the required level. With a sample volume of 150 mL and using the Prospekt cartridges packed with PLRP-S, the recoveries of de-isopropyl- and de-ethyl-atrazine are not 100% because breakthrough has occurred on PLRPS, but it is possible to detect these compounds with reproducible results.

The calibration can be made by spiking LC-grade-, drinking-, surface- or other real water samples. In practice, especially in multiresidue analysis, it is easier to construct calibration curves once for all the analytes and to use them for any kind of water. It has been shown that calibration curves constructed with spiked LC-grade water samples and from spiked drinking water samples were similar [14]. Good linearity and correlation coefficients were obtained in the range 0.1 – 1.5 µg/L. Calibration curves have also been constructed with surface waters and good linearity was obtained in the range 0.1 - 7 µg/L. In surface water, the calibration curves depend on the interferences which show up in the real chromatogram and on the possibility of identifying the unknown peaks and their purity. In recent years, much attention has been given to identification and peak purity in diode-array software.
chromatographic determination were being used. The overall RSD between values obtained by the authors and the average value obtained by fourteen or fifteen other laboratories varied between 1.6 and 36% for atrazine and organophosphorus pesticides in finished drinking waters at levels ranging from 0.02 to 0.2 µg/L.

Application to various samples: Limits of determination

The suitability of the SPE-LC-DAD system for multiresidue purposes is provided in figure 3 where metamitron was present in the groundwater at a concentration level of 0.12 µg/L and was just detected at 220 nm since this analyte was coeluted with an interfering hump [10]. However, the same chromatogram drawn at 306 nm illustrates the low detection limit that can be reached for this analyte and the confirmation that can be provided by the DAD when the UV spectrum presents some characteristics in the scan range. It was possible to use a full scale of 1 mV, ten times lower than that at 220 nm, because a very low background in the baseline is detected at 306 nm. Unfortunately, this property does not apply to all the pesticides.

In general, the limit of determination depends primarily on the detection mode and the properties (i.e. spectral properties) of the analytes, but also on the type and matrix of waters that are analysed. There is always a great difference (up to a factor of 10) between LODs obtained in LC-grade water and those obtained in ground- or drinking-waters. LODs in contaminated surface waters are also higher (about 5 times, depending on the organic carbon content) than those observed in drinking waters.

A survey of the results for many drinking- or groundwater indicates that the limit of determination of 0.1 µg/L is easily obtained for most pesticides using UV detection and with the handling of 150 – 300 mL of sample. The LOD is often lower, around 10 – 30 ng/L, which allows...
Pesticide analysis

quantitative analyses at the 0.1 µg/L level. Only the more polar compounds and/or degradation products such as de-isopropylatrazine are difficult to determine at levels below 0.1 µg/L. In surface waters, LOD of 0.1 µg/L can also be obtained, as can be estimated from figure 4 by the on-line analysis of a river Marne sample spiked with 0.5 µg/L of the same pesticides as the drinking water sample in figure 2 [2]. The chromatograms are very similar, the experimental conditions being the same apart from the attenuation range of the UV detector which is four times higher in figure 4. However, for many pesticides, the LODs are far below the 0.5 µg/L range, as can be seen from the height of some peaks at 220 nm.

For the more polar range of compounds, the on-line system PGC/PGC could provide accurate determination of the more polar analytes. In a long-term survey of a ground-water source, monitoring using the PLRP-S precolumn and C18 analytical column on-line system indicated constant and rather high amounts of atrazine and de-ethylatrazine, with average concentrations of 0.5 and 0.6 µg/L respectively (see Fig. 3). Because of the bad detection obtained with this system for the second metabolite de-isopropylatrazine, a PGC precolumn-PGC analytical column coupling was used. Figure 5 shows the advantage of such a system, since de-isopropylatrazine (DIA) is eluted after de-ethylatrazine (DEA) and can easily be delayed to 40 min in the chromatogram, after the interfering compounds [10]. The breakthrough volume of DIA on PGC is above 100 mL so that detection limits using 100 mL samples are in the low-0.1 µg/L range in LC-grade water, as shown in figures 5a,b. In the non-spiked ground water (Figs. 5c,d), DEA was confirmed at a concentration of 0.6 µg/L and the concentration of DIA was 0.05 ± 0.01 µg/L.

Further developments

Since many recent pesticides and/or degradation products cannot be analysed by GC, the only possibility of multiresidue analysis including compounds over a wide range of polarity is given by LC. On-line systems using LC can be fully automated and works as an early-warning or on-site monitoring system. They can also be used as a powerful routine tool in the laboratory. An automated post-column reaction can also be added to the on-line SPE-LC-UV DAD system, thus broadening the range of analyte analysed in one run. N-methyl carbamates and glyphosate can be thus determined.

The numerous applications of SPE-LC on-line systems have shown this method to be well suited for multiresidue analysis. Its wide acceptance has contributed greatly to the development of SPE-LC-MS systems, along with its identification potential.

References

Pesticide analysis