Application Note No. 033

G GL Sciences

Determination of Pesticides in Aqueous Samples by On-Line Coupling Solid-Phase Extraction to Gas Chromatography with At-Column

Concentrating Interface

Ryoichi Sasano*, Takayuki Hamada, Masahiro Furuno

GL Sciences, 237-2 Sayamagahara, Iruma, Saitama, 358-0032 Japan

Key Words: On-line SPE-GC; Large volume injection; Determination of Pesticides in water

Summary

The on-line automatic Solid Phase Extraction and the capillary GC/MS were combined for the

determination of pesticides and herbicides in the aqueous samples. The at-column concentrating large

volume injection method which the authors exploited was employed as the interface between SPE and

GC/MS. In the at-column concentrating large volume injection method, the sample solvent was

evaporated in the empty liner and the target compounds were retained at the inlet of the analytical GC

capillary column. Employing this method, heat labile target compounds were not decomposed.

Automatic on-line operation was possible, since the large volume of the liquid sample could be

transferred to the capillary GC/MS system.

1 Introduction

Recently, water pollution caused by pesticides and herbicides have become one of the serious

environmental pollution. Therefore, the government regulations of water pollution due to pesticides

and herbicides have become strict for protecting human health. In the conventional methods for the

determination of pesticides and herbicides contained in the tap water and the river or lake water, solid

phase extraction sample preparation prior to GC/MS analysis has been employed. However, the

manual sample preparation methods are likely to cause human errors and require operators technical

skills.

In this study, we connected the ProspektTM (Spark Holland); automatic on-line solid phase extraction

instrument and GC/MS equipped with OPTICTM 2-200 PTV (ATAS Netherland) for at-column

concentrating large volume injection system. We employed the at-column concentrating large volume

injection method as the interface between SPE and GC/MS. Using this method, we could

GL Sciences B.V.

😘 GL Sciences

connect the automatic SPE and GC/MS without causing the degradation of target compounds. We have

examined how to determine 30 kinds of pesticide and herbicide compounds in the aqueous samples,

using this system.

2 Experimental

2.1 Chemicals and reagents

All the standard compounds of pesticides and herbicides were from GL Sciences. Acetone and

methanol of pesticide residue analysis grade were purchased from Kishida Chemical Co.. The stock

solutions of the standards were prepared by diluting 500 mg of each standard compounds in 50 ml

acetone. The solution was diluted into 0.1 mg/L using acetone. The standard solution was spiked into

the purified water as the concentration of 1 g/L. 30 kinds of pesticides and herbicides we used here were

those which are to be determined using GC, specified by Japanese national government reguration.

2.2 SPE instrument

The automatic on-line SPE instrument; Prospekt (Spark Holland) was connected with the autosampler;

MIDASTM (Spark Holland). The 2 mL stainless steel sample loop was installed to the sampling valve of

the MIDAS. The solid phase extraction cartridge employed for the Prospekt was PLRP-S, 2.0 mm i.d.~

10 mm lenghth, 15 - 25 m particle diameter. The HPLC pump PU610 (GL Sciences) was employed for

eluting the target compounds from the SPE cartridge. The outlet of SPE valve and the GC injector are

connected using a 0.25 mm i.d.~ 0.5 m length inactivated capillary tube. The schematic flow diagram of

this automated SPE system is shown in Fig. 1.

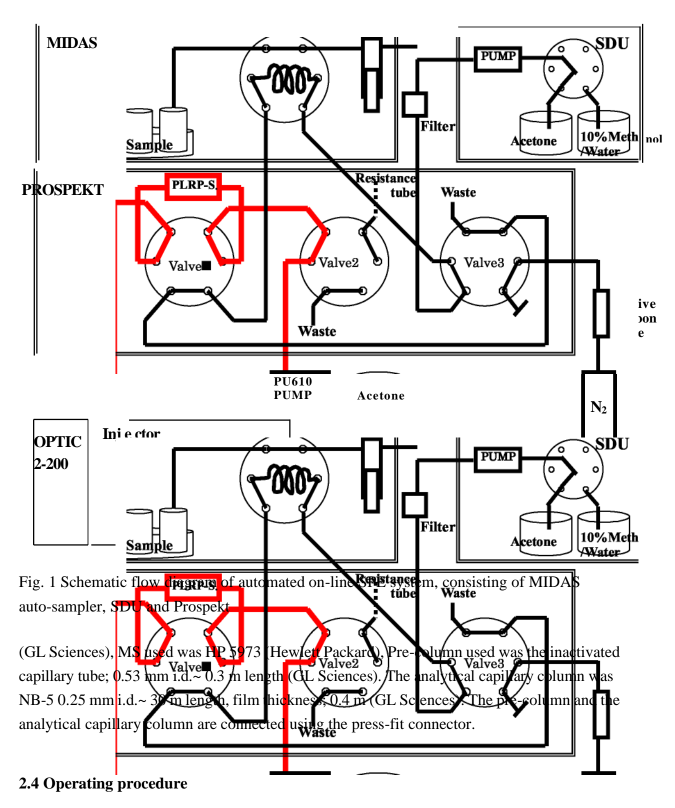
2.3 GC instrument

GC instrument used was HP6890 (Hewlett Packard) equipped with OPTIC 2-200 PTV injector (ATAS

Netherland). The liner used was the liner for at-column concentrating large volume injection

GL Sciences B.V.





2 mL sample solution was injected to the SPE cartridge employing MIDAS. The SPE cartridge was pre-conditioned using acetone and then (10 % methanol / 90 % water) mixture prior to the sample



loading. After loading the sample and washing the cartridge, the SPE cartridge was N₂ gas purged for eliminating the remaining water in the cartridge. Then, the target compounds were eluted from the SPE cartridge, using approximately 30 L acetone at the flow rate of 100 L/min. The eluted compounds were injected to the capillary GC/MS system automatically. The GC/MS system was operated using SIM mode. The operating conditions of GC/MS are shown in Table 1.

Table 1 Operating conditions of GC/MS and Opric 2-200

Oven Temp	73~Ë1 min) - 15~/min - 210~ - 4~/min - 250~ - 20~/min - 280~(6min)
Interface Temp 2	280 ~ MS
Source Temp 230	
Temp 150 ~	
Method	SIM
Injector	Optic 2-200 (At-column concentrating Liner)
3	arge Volume Method
Vent Time	AUTO
Initial Tamp	62.~
Initial Temp	02
Ramp rate	1 ~/s 280 ~
Final Temp	280 ~
Split open Time	0 min
Purge pressure	25 kPa
Transfer Time	1 min
Initial pressure	50 kPa
TT' 1	200 kPa
Final pressure	

3 Results and discussion

3.1 Interface

In this study, we employed the at-column concentrating large volume injection method. The scheme of this method is described as follows.

The temperature of the injector was set at lower than the boiling point of the sample solvent and the column oven temperature was set at higher than the boiling point of the sample solvent. The sample solution is injected into the liner shown in Fig. 2. The liquid solvent stayed in the liner, since the



carrier gas pressure and the solvent vapor presser are equilibrated. The evaporated solvent was exhausted through the solvent vent. The target compounds were concentrated at the inlet of the capillary analytical column. Then, the target compounds were separated on the analytical column by elevating column oven temperature. This injection method was employed for connecting SPE to GC/MS as an automatic on-line operation system. In this system, the eluate from the SPE cartridge was injected to GC/MS capillary system automatically as an on-line system. During evaporation of the sample solvent, the temperature of the injector was set at 60 °C and the column oven temperature was set at 73 °C, since the boiling point of the acetone was 63 °C (at 25 kPa).

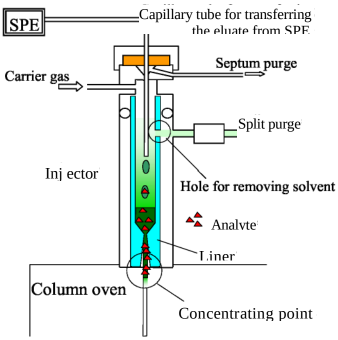


Fig. 2 Scheme of on-line interface employed for the at-column concentrating method

3.2 The recovery and reproducibility

Employing this system, 7 times running by injecting of 1pg/L pesticides and herbicides sample solution were repeated. The sample used were prepared by spiking the target compounds to the purified water at the concentration of 1 g/L. The recoveries and R.S.D. (%) was calculated from the obtained chromatograms. The results were shown in the table 2. The SIM chromatogram also shown in Fig. 3. The recoveries of most compounds were larger than 80 %. R.S.D. were



approximately within 5%. The peak shapes in the chromatogram were satisfactory.

As the results of this study, we found that on-line automatic SPE and the capillary GC/MS equipped with the at-column concentrating large volume injection apparatus could be connected with good results for the determination of pesticides and herbicides in water samples.

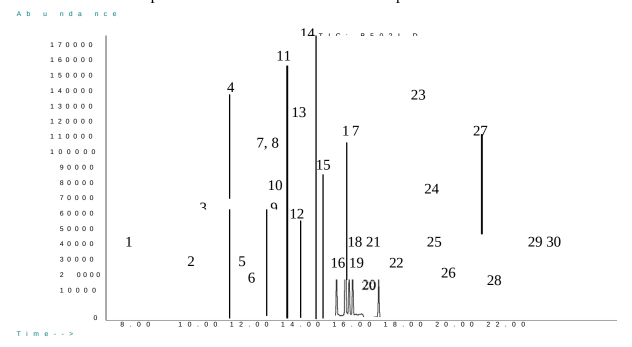


Fig. 3 GC/MS chromatogram of 1 g/L pesticides and herbicides spiked to the purified water, obtained employing this on-line system.

Table 2 Recoveries and RSD(%) obtained using this on-line system. The sample was 1 g/L pesticides and herbicides spiked to the purified water.

# Compound	(REC (%)	# Compound	R.S.D. (%) <i>REC</i> (%)	
1 Dichlorvos	9.	87.2	16 Fenitrothion	3.82	96.5
2 Etridiazole	6.	96.8	17 Thiobencarb	2.95	<i>102.0</i>
3 Chloroneb	5.	92.7	18 Cyanazine	6.65	93.6
4 Fenobucarb	3.	93.5	19 Chlorpyrifos	2.12	<i>78.3</i>
~		^	-		
11 Phenanthrene-d10	4.05	84.1	26 Isoxathion	3.50	101.4
12 Chlorothalonil	1.94	99.5	27 Mepronil	4.61	100.5
13 Iprobenfos	2.81	106.2	28 Chlornitrofen	4.42	<i>82. 9</i>
14 Terbucarb	3.33	101.9	29 Pyridaphenthion	5.67	91.8
15 Tolclofos-methyl	2.51	96.2	30 EPN	4.56	86.5