Application Note

GC Monitoring of Chlorinated Pesticides in Hazardous Waste Sites

SPB-608 capillary columns were evaluated to demonstrate their suitability for monitoring organochlorine pesticides in hazardous waste samples under US EPA Contract Laboratory Program, March 1990 Statement of Work requirements. All components in the CLP-specified pesticides test mix were baseline resolved, exceeding CLP performance requirements. The combined decomposition of endrin and 4,4'-DDT was 8.86%, easily meeting the inertness requirement that the combined decomposition of these compounds be less than 30%. Linearity was well within CLP specifications; The mean relative standard deviation of the calibration factors was 9.49% and absolute retention times of all the analytes were within the windows established in accordance with the method, establishing linearity well within CLP specifications. (ChromFax: 394003)

Key Words

- chlorinated pesticides
 hazardous waste
- polychlorinated biphenyls (PCBs)

The Contract Laboratory Program (CLP) March 1990 Statement of Work is the most recent gas chromatography procedure published by the US EPA for the monitoring of chlorinated pesticides in samples from abandoned waste sites. This method specifies stringent inertness, resolution, and linearity requirements for the wide-bore (0.53mm ID) capillary GC column.

SPB[™]-608 capillary columns, which are manufactured and tested specifically for monitoring organochlorine pesticides and PCBs in wastewater according to US EPA Method 608, are also suitable for the CLP method and for the proposed Method 8081 for current waste sites. We evaluated the SPB-608 column using the series of test mixtures specified in the CLP method to measure column resolution and inertness, and system linearity.

Column performance exceeded the CLP method requirement that resolution between consecutive peaks be greater than 60% of the height of the shorter peak. The seven target pesticides in the resolution check mix were baseline resolved, including the pairs of analytes cited in the method as having the poorest resolution on the DB-608 column — γ -chlordane/endosulfan I, 4,4'-DDE/dieldrin, and methoxychlor/endrin ketone (Figure A).

The SPB-608 column easily meets the inertness criteria established in the CLP pesticides method. Decomposition of either endrin or 4,4'-DDT must not exceed 20%, and their combined decomposition must be less than 30%. Endrin decomposition was 8.7%, all in the form of endrin ketone; no measurable amounts of endrin aldehyde were evident. Decomposition of

Figure A. Baseline Resolution of **Target Compounds**



4,4'-DDT was measured at 0.16%, in the form of 4,4'-DDE. No measurable amounts of 4,4'-DDD were evident. The combined decomposition of 4,4'-DDT and endrin on this column was 8.86%.

The linearity of the instrument, including the column, is determined by calculating percent relative standard deviation (%RSD) of the calibration factors from a three-point calibration curve for each pesticide and surrogate in individual solutions. The target compounds were analyzed in two standard mixes, A and B, to avoid coelutions. Absolute retention times were determined by calculating the mean retention time from both mixes. Linearity was well within CLP specifications (Table 1).

Absolute retention times of all the target analytes and surrogates in the performance evaluation mixture were within the windows established in the calibration runs for the three concentration ranges (Table 1). The target analytes were baseline resolved (Figure B), well exceeding the CLP requirement that the resolution between adjacent peaks of the individual standard mixes be greater than 90%.



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The performance of the SPB-608 column easily meets US EPA requirements for stability, inertness, and linearity as specified in the US EPA CLP Pesticide Method, March 1990 Statement of Work.

Table 1. SPB-608 Column Easily Meets CLP **Retention Time and Linearity Requirements**

Compound	Mean RT (n=3)	Retenti Win From	on Time dow [□] To	Calibratior Factor▲ %RSD
Tetrachloro-m-xylene [▼]	7.34	7.29	7.39	8.78
α-BHC	9.43	9.38	9.48	9.90
γ-BHC (Lindane)	10.63	10.58	10.68	6.32
β-BHC	10.92	10.87	10.97	3.28
Heptachlor	11.84	11.79	11.89	2.22
δ-BHC	12.04	11.99	12.09	17.16
Aldrin	12.88	12.83	12.93	14.61
Heptachlor epoxide	14.06	13.99	14.13	9.80
γ-Chlordane	15.17	15.10	15.24	5.84
α-Chlordane	15.65	15.58	15.72	7.17
Endosulfan I	15.69	15.62	15.76	3.33
4,4'-DDE	16.40	16.33	16.47	15.66
Dieldrin	16.61	16.54	16.68	7.13
Endrin	17.65	17.58	17.72	5.00
4,4'-DDD	18.05	17.98	18.12	4.47
Endosulfan II	18.22	18.05	18.29	27.86
4,4'-DDT	18.98	18.91	19.05	6.84
Endrin aldehyde	19.15	19.08	19.22	11.93
Endosulfan sulfate	19.62	19.55	19.69	10.65
Methoxychlor	21.41	21.34	21.48	6.62
Endrin ketone	21.73	21.66	21.80	13.62
Decachlorobiphenvl [▼]	26.76	26.66	26.86	10.52

^D Retention time windows are ±0.05 minutes for all compounds that elute before heptachlor epoxide; ±0.07 minutes for heptachlor epoxide and other compounds that elute before decachlrobiphenyl; ±0.10 minutes for decachlorobiphenyl.

▲ %RSD for up to two target compounds, not including surrogates, may be >20%, but must be ≤30.0%. %RSD for remainder of target compounds must be ≤20.0%. %RSD for surrogates must be ≤30.0%.

Surrogate retention times are measured from standard mix A analysis.

Ordering Information:

SPB-608 Fused Silica Capillary Columns

30m x 0.53mm ID, 0.5µm film	25312
15m x 0.53mm ID, 0.5µm film	25310-L

Reference

US EPA Contract Laboratory Program Statement of Work for Organics Analysis, Multi-Media, Multi-Concentration, Document Number 0LM01.0, Including Revisions 0LM01.1 (December 1990) and 0LM01.1.1 (February 1991).

Fused silica columns manufactured under HP US Pat. No. 4.293.415.

Figure B. Standard Mixes, Low-Range Concentration



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Note 3

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