

Cover Sheet for

ENVIRONMENTAL CHEMISTRY METHOD

Pesticide Name: Quintozene (PCNB)

MRID #: 430615-01

Matrix: Soil

Analysis: GC/ECD

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SECRET

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(S) (U) (C) (E) (R) (E) (T)

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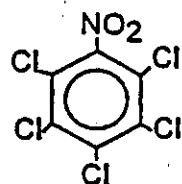
(S) (U) (C) (E) (R) (E) (T)

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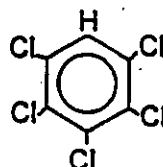
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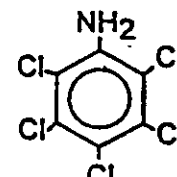
Figures



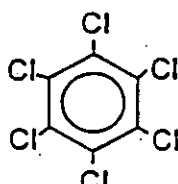
Pentachloronitrobenzene
(PCNB)



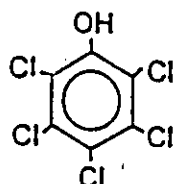
Pentachlorobenzene
(PCB)



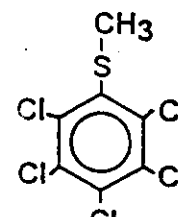
Pentachloroaniline
(PCA)



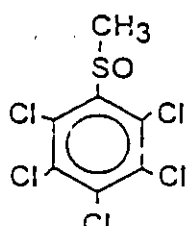
Hexachlorobenzene
(HCB)



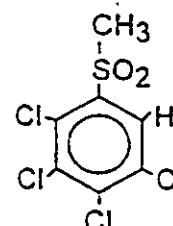
Pentachlorophenol
(PCP)



Pentachlorothioanisole
(PCTA)



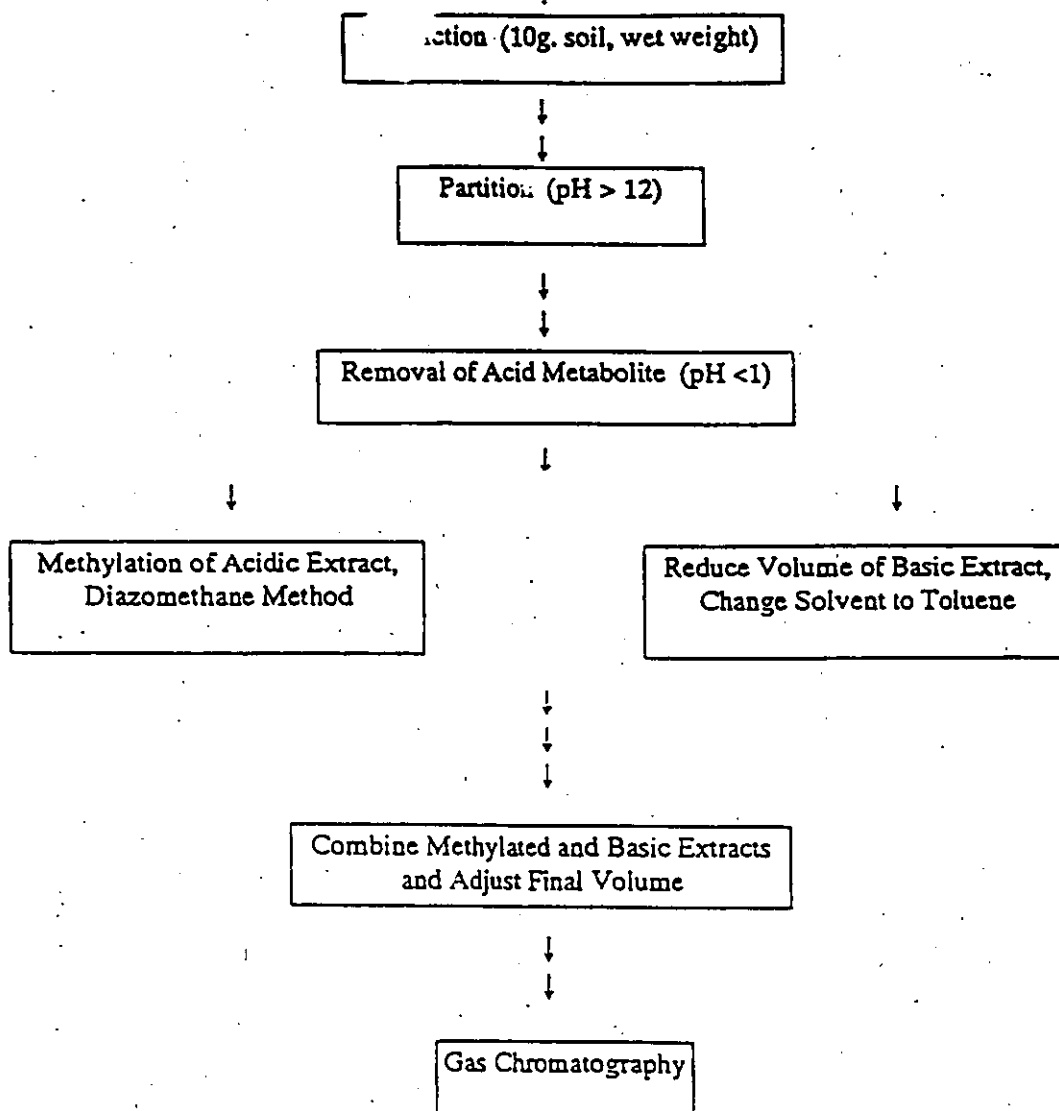
Pentachlorothioanisole sulfoxide
(PCTASO)



2,3,4,5-tetrachlorothioanisole sulfone
(TCTASOO)

Figure 1: Molecular structure of PCNB, PCB, PCA, HCB, PCP, PCTA, PCTASO, and TCTASOO

Figure 2: PCNB and Metabolites Analysis Method
Flowchart



TITLE

**Residue Analytical Method for Terraclor
and its Metabolites and Impurities**

Authors

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Performing Laboratory

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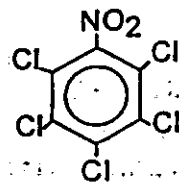
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I. Summary

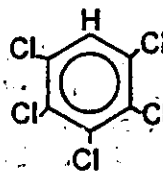
Soil samples were extracted with acetone/hexane. Water (pH > 12) was added to the acetone/hexane extract and the mixture was re-extracted in hexane. The water was then acidified (pH < 1) and once again extracted with hexane. The extract of the acidified water was methylated with diazomethane and combined with the remainder of the sample. The solvent was changed to toluene and the volume of the combined extracts was adjusted. The samples were analyzed for PCNB and metabolites and impurities using GC/ECD.

II. Introduction

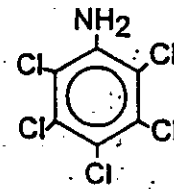
This method was developed by Centre Analytical Laboratories to determine pentachloronitrobenzene (PCNB), pentachlorobenzene (PCB), hexachlorobenzene (HCB), pentachloroaniline (PCA), pentachlorothioanisole (PCTA), pentachlorophenol (PCP), 2,3,4,5-tetrachlorothioanisole sulfone (TCTASOO) and pentachlorothioanisole sulfoxide (PCTASO). The molecular structure of these compounds is shown in Figure 1. The following references were used as a guideline in the initial development of this method: "Determination of Terrazole (5-Ethoxy-3-Trichloromethyl-1,2,4-Thiadiazole) and Terraclor (Pentachloronitrobenzene) and Allied Metabolites in Plant Tissues or Harvest Samples" and "Determination of Terraclor (Pentachloronitrobenzene) and Terrazole (5-Ethoxy-3-Trichloromethyl-1,2,4-Thiadiazole) in Soil".



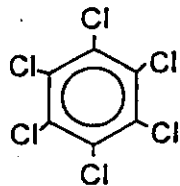
Pentachloronitrobenzene (PCNB)



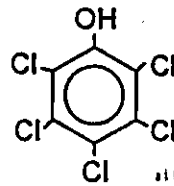
Pentachlorobenzene (PCB)



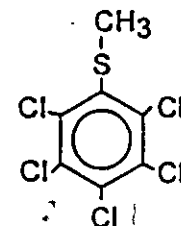
Pentachloroaniline (PCA)



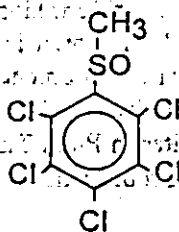
Hexachlorobenzene (HCB)



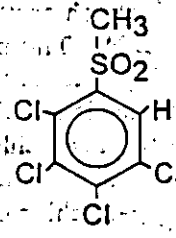
Pentachlorophenol (PCP)



Pentachloroanisol (PCTA)



Pentachloroanisol sulfoxide (PCTASO)



2,3,4,5-tetrachloroanisol sulfone (TCTASOO)

Figure 1: Molecular structure of PCNB, PCB, PCA, HCB, PCP, PCTA, PCTASO, and TCTASOO

III. Method

A. Chemicals/Supplies

Acetone, residue grade	Baker
1-Decanol	Baker
Diazomethane in diethyl ether solution	Aldrich
Diethyl ether, residue grade	Baker
Dry Ice	Penn State University
n-Hexane, residue grade	Baker
HCB Analytical Standard AC-1194-38C	Uniroyal Chemical Co.
Nitrogen	MG Gases
PCA Analytical Standard AC-1234-4	Uniroyal Chemical Co.
PCB Analytical Standard AC-1166-14	Uniroyal Chemical Co.
PCNB Analytical Standard AC-1261-133	Uniroyal Chemical Co.
PCP Analytical Standard AC-1261-84	Uniroyal Chemical Co.
PCTA Analytical Standard AC-1166-16	Uniroyal Chemical Co.
PCTASO Analytical Standard AGD-1384-005	Uniroyal Chemical Co.
Sodium Hydroxide	Baker
Sulfuric Acid	Baker
TCTASOO Analytical Standard AGD-1384-024	Uniroyal Chemical Co.
Toluene, residue grade	Baker

B. Equipment

Balance	Mettler PE 3000
Centrifuge	Damon/IEC
Centrifuge bottle, teflon 250 ml	Nalgene
Hobart Food Chopper	Hobart Mfg. Co.
Erlenmeyer Flask, 250 ml	Pyrex, Kimax
pH meter	Beckman
Rotary evaporator, Buchi Rotovap	Brinkman
Round bottom flasks, 500 ml	Pyrex, Kimax
Separatory funnel, 250 ml	Nalgene
Standard laboratory equipment: beakers, pipets, test tubes etc.	Pyrex, Kimex
TurboVap LV evaporator	Zymark

C. Instrumentation

The gas chromatograph and integrator models, column type and operating conditions were as follows:

Instrument	Hewlett Packard model 5890 series gas chromatograph
Column	Restek RTX-35, 30 m, 0.53 mm ID, 0.25 μ m df
Oven	initial temp. 100°C, initial time 2 min rate A: 5°C/min to 200°C, final time 0 min rate B: 20°C/min to 270°C, final time 5 min
Detector	Electron Capture Detector (ECD) temp. 300°C
Injector	Direct Injection temp. 270°C
Carrier Gas Flow	Hydrogen, 10 ml/min
Make-up Flow	Nitrogen, 35 ml/min
Integrator	Shimadzu C-4RA Chromatopac

D. Preparation of Standard and Spiking Solutions

Analytical standards received from the sponsor were used to prepare individual compound stock solutions from which working standard and method day spiking solutions were prepared. Stock solutions of each compound, at a concentration of 1.0 mg/ml, were made by weighing out 10 mg of the analytical standard on an analytical balance, and dissolving it in 10 ml of toluene. The amount of toluene added was corrected considering the percent purity of the standard. For example, if HCB was 99.8% pure then 10 mg would be weighed out and dissolved in 9.98 ml of toluene (10.0×0.998). The PCP stock solution was prepared using methanol.

A solution of the combined seven compounds in toluene, at a concentration of 100 $\mu\text{g/ml}$, was made by adding 2 ml of each of the individual compound stock solutions at 10 mg/ml of PCB, HCB, PCNB, PCA, PCTA, TCTASOO and PCTASO, to 6 ml of toluene, so that the final volume was 20 ml. A solution of PCP at 100 $\mu\text{g/ml}$ was made by diluting the 1.0 mg/ml stock solution of PCP ten-fold with methanol.

A Method day spiking solution of the combined seven compounds at a concentration of 10 $\mu\text{g/ml}$ was made by a ten-fold dilution of the 100 $\mu\text{g/ml}$ seven compound solution with toluene. Likewise, PCP day spiking solution at 10 $\mu\text{g/ml}$ was made by a ten-fold dilution with methanol of the 100 $\mu\text{g/ml}$ PCP solution. Fortification of the method day spike samples at a 1 μg level was accomplished by adding 100 μl of the 10 $\mu\text{g/ml}$ spiking solutions to a control sample. Fortification of the method day spike samples at a 10 μg level was done by adding 100 μl of the 100 $\mu\text{g/ml}$ spiking solutions to a control sample.

A 10 $\mu\text{g/ml}$ standard stock solution was made by adding 200 μl of each individual compound stock solutions of PCB, HCB, PCP, PCNB, PCA, PCTA, TCTASOO and PCTASO at 1 mg/ml, and bringing the final volume to 20 ml with toluene. A 1 $\mu\text{g/ml}$ standard stock solution of the combined eight compounds was prepared by diluting the 10 $\mu\text{g/ml}$ standard stock solution ten-fold with toluene. Dilutions of the 10 $\mu\text{g/ml}$ and 1 $\mu\text{g/ml}$ standard stock solutions were made to prepare working 0.100 $\mu\text{g/ml}$, 0.050 $\mu\text{g/ml}$, 0.010 $\mu\text{g/ml}$ and 0.003 $\mu\text{g/ml}$ standards.

D. Analytical Procedures

1. Sample Processing

The frozen soil core samples were received in the laboratory. The frozen cores were divided into smaller pieces with a cleaver and rubber mallet, or by other appropriate means. The stones and debris were removed. A Hobart Food Chopper was pre-chilled with dry ice and the frozen soil pieces were put inside. The soil was chopped and homogenized with dry ice. The soil was then placed in sample containers and stored in the freezer where the dry ice was allowed to sublime. The soil samples were kept under freezer conditions ($-24^{\circ}\text{C} \pm 7^{\circ}\text{C}$) until analysis.

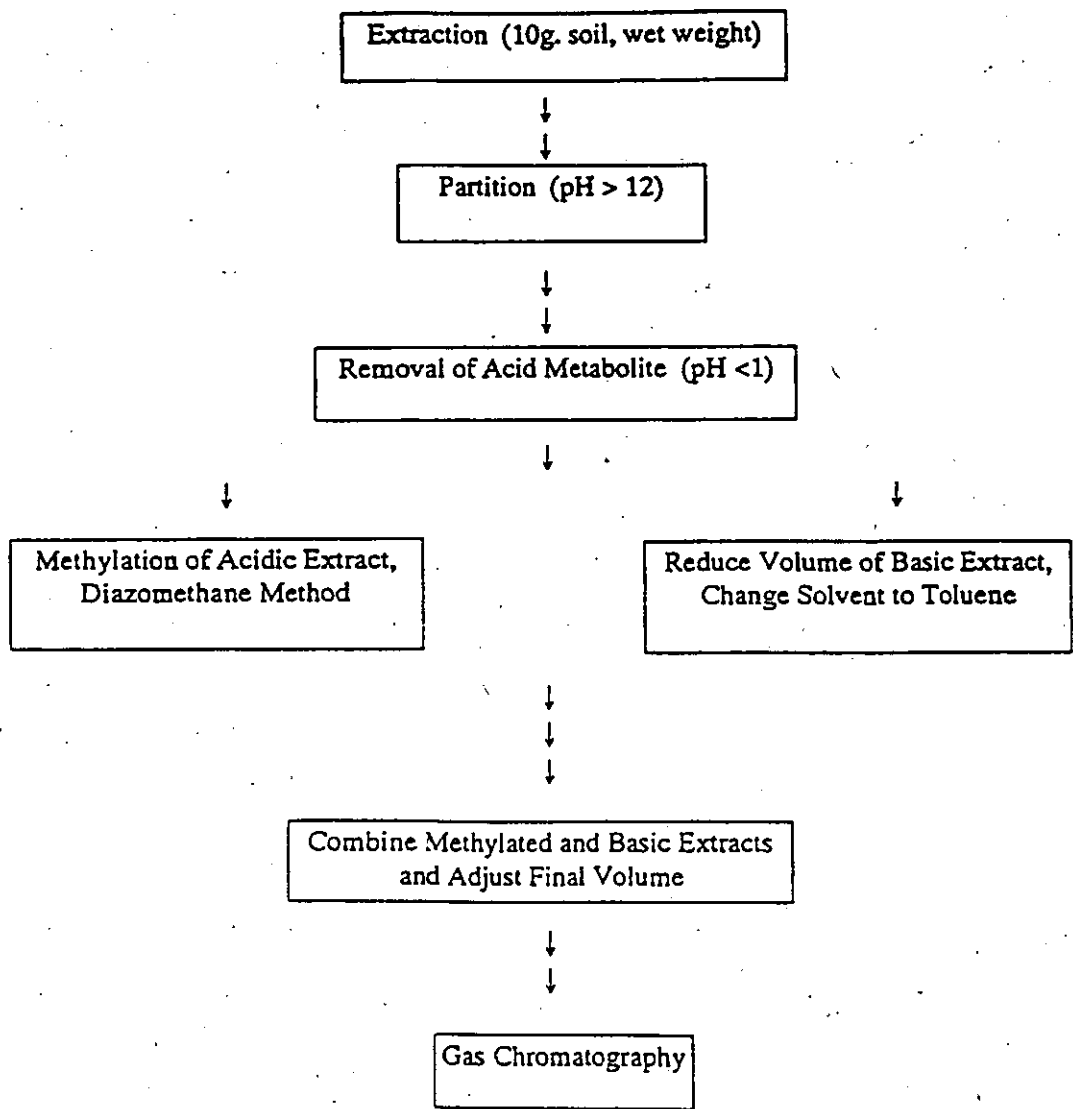
2. Analysis

A flow diagram of the analysis procedure is shown in Figure 2. Detailed explanations of each step are as follows:

Step 1 Extraction (10 g soil, wet weight)

Weigh 10 g of soil in a 250 ml teflon centrifuge bottle. Fortify the two spike samples used to determine extraction method recoveries for the set of samples. Add 100 ml 50:50 v/v acetone:hexane and shake vigorously for 2 min. Centrifuge at 1500 RPM for 5 min. Pour the supernatant into separatory funnel leaving the soil in the bottle. Add another 50 ml of 50:50 v/v acetone:hexane to soil, shake 1 min and centrifuge. Add supernatant to separatory funnel.

Figure 2: PCNB and Metabolites Analysis Method
Flowchart



Step 2 Partition

Add 50 ml distilled/deionized water (pH >12, adjusted with 25% NaOH) to separatory funnel and shake 20 sec. Drain the water/acetone layer into a 250 ml beaker and collect the hexane layer in a 250 ml Erlenmeyer flask. Re-extract the water/acetone layer with 50 ml hexane, shaking for 1 min. Drain the water/acetone layer into the 250 ml beaker and add the remaining hexane layer to the flask. The basic extract in the Erlenmeyer flask contains compounds PCB, HCB, PCNB, PCA, PCTA, TCTASOO, and PCTASO.

Step 3 Removal of Acidic Metabolite

Pour the water/acetone portion back into the separatory funnel and add 10 ml 10 N H_2SO_4 to lower the pH < 1. Add 50 ml hexane, shake vigorously for 1 min and drain the water/acetone layer into the beaker. Pour the hexane layer into a 500 ml round bottom flask. Re-extract the water/acetone with another 50 ml hexane, by shaking for 1 min. Drain the water/acetone layer into the beaker and add the remaining hexane layer to the round bottom flask. The acidic extract in the round bottom flask contains compound PCP.

Step 4 Methylation, Diazomethane Method

Add 10 drops of decanol to the acidic extract to prevent the sample from going to dryness during evaporation and reduce the volume to about 5 ml using a rotary evaporator. Transfer this portion of the sample into a methylation vial, rinsing the round bottom flask with hexane. Further reduce the volume of the sample to 0.5 ml using a TurboVap LV evaporator under nitrogen. Add 0.5 ml diazomethane, or enough to turn the sample yellow; let it stand under a hood for 10 minutes. Evaporate off the diazomethane using the TurboVap, reducing the volume again to 0.5 ml.

Step 5 Combine the Extracts and Adjust the Volume

Rinse a round bottom flask with acetone and transfer the basic extract, prepared in step 2, from the Erlenmeyer flask to the round bottom flask. Reduce the volume to about 5 ml using the rotary evaporator, then add 10 ml toluene. Pour the methylated portion of the sample into the round bottom rinsing the vial with 15 ml toluene. Reduce the volume of the combined extracts to about 5 ml with the rotary evaporator, then and bring the final volume up to 10 ml with toluene. The sample is now ready for GC analysis.

Step 6 Gas Chromatography

Inject a 1 μ l aliquot of each eight-component standard in the range of 0.003 μ g/ml to 0.100 μ g/ml into the gas chromatograph. Record the resulting peak areas, or peak heights, and plot this data versus concentration (μ g/ml) of the corresponding standard to obtain standard calibration curves. Prepare standard curves for each analysis day.

Inject a 1 μ l aliquot of the sample into the gas chromatograph. If necessary, dilute the sample with toluene so that the signal response is within the standard curve range. Record peak areas, or peak heights, and determine the concentration of each component compound relative to the standard curves generated for that day.

IV. Method of Calculation

The peak areas corresponding to the eight compounds (PCB, HCB, PCP, PCNB, PCA, PCTA, TCTASOO, and PCTASO) in the standards were obtained from the chromatograms and regressed versus the concentration of the compounds in the standards. Statistics were generated on a Swan Corporation 386/33 computer using Axum program capable of performing quadratic regression (second order polynomial regression) on the peak areas versus their corresponding concentrations to generate standard curves. The following quadratic equation was used:

$$y = b_0 + b_1 x + b_2 x^2$$

A corrected peak area value, if required, was determined using the following formula:

$$\text{Peak area in sample corrected} = \text{Peak area in sample} - \text{Peak area in control}$$

The corrected peak area of each sample was used to calculate the amount in $\mu\text{g/ml}$ of each compound found in the samples analyzed relative to the generated standard curves. The square of the correlation coefficient (R^2) was used to evaluate the fit of the curve. The $\mu\text{g/ml}$ compound found value was then multiplied by the final volume of the sample to yield the μg compound found.

$$\mu\text{g compound found} = [\mu\text{g/ml compound found}] \times [\text{final volume (ml)}]$$

The μg compound found values were converted to ppm compound found value by dividing by the sample weight. The ppm compound found value was then divided by ppm compound added to obtain the percent recovery in fortified method spikes.

If the average percent recovery for the two spiked samples of the set was below 100%, the amount of compound found in the sample was divided by the average recovery of the spikes to give the corrected value. No correction was made for average recoveries above 100%.

$$\mu\text{g compound found corrected} = \mu\text{g compound found} / \text{average spike recovery}$$

The ppm compound found in the samples was calculated using the μg compound found corrected for percent recoveries divided by sample weight.

V. Results and Discussion

An eight day Validation (CAL 004-05) was conducted by Centre Analytical Laboratories on the determination of PCNB, PCB, HCB, PCP, PCA and PCTA in soil using the method described herein. The validation consisted of five spiking levels in duplicate for each of the compounds, a total of 60 fortified samples. The spiking levels were 0.005ppm, 0.250ppm, 1.00ppm, 3.00ppm and 10.0ppm. The limit of detection (LOD) was 0.003ppm; at this level the signal to background ratio was greater than or equal to 5. The limit of quantitation (LOQ) was 0.005ppm; this level is set above the LOD so that the method was not pushed to its limits. The average recoveries for the eight day study were as follows:

<u>COMPOUND</u>	<u>8 DAY AVERAGE RECOVERY</u>	<u>STD-DEVIATION</u>
PCNB	100%	4 %
PCB	88%	2 %
HCB	97%	3 %
PCA	101%	3 %
PCTA	101%	3 %
PCP (as PCP-OMe)	90%	5 %

This data indicates the analytical method is suitable for these compounds.

A one day validation was conducted on the determination of TCTASOO and PCTASO in soil using this same analytical method. It consisted of three spiking levels in duplicate, a total of six fortified samples. The spiking levels were 0.005ppm, 0.100ppm and 1.00ppm. As in the previous validation, the limit of detection (LOD) was 0.003ppm; at this level the signal to background ratio was greater than or equal to 5. The limit of quantitation (LOQ) was 0.005ppm; this level is set above the LOD so that the method was not pushed to its limits. The average percent recoveries were as follows:

<u>COMPOUND</u>	<u>AVERAGE RECOVERY</u>	<u>STD-DEVIATION</u>
TCTASOO	102%	15 %
PCTASO	103%	6 %

This data indicates the analytical method is also suitable for these compounds.

Examples of representative chromatograms of a standard, control sample and a spiked control, in addition to a typical standard calibration curve are shown in Figures 3-6.

SAMPLE NUMBER: 1
 SAMPLE NAME: STANDARD
 MISC. DATA: 0.050 µg/ml
 DILUTION FACTOR: NA

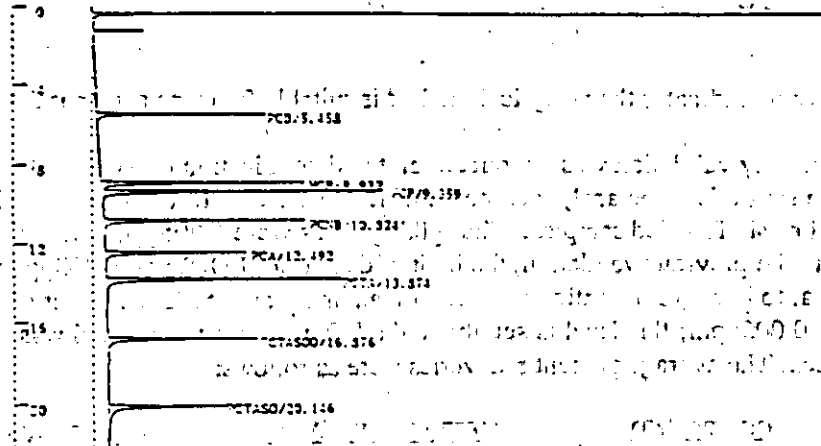
C-RIA CHROMATOGRAPHY REPORT No. 1783 : CHROMATOGRAM-1:SDIS.CDD 92/09/30 20:37:26

Analysis File : 1:R000P.

Centre Analytical Labs

Analysis of PCB, PCP, HCB, PCNB, PCA, PCTA, PCTASO & TETASO

COLUMN: RTX-35 (30m, 0.33mm ID, 1.25µm df)
 OVLN: 100 C, 5 min, 5 C/min to 200C, 20 C/min to 270, 5 min
 DETECTOR: ECD, 300 C
 INJECTOR: Direct, 270 C
 CARRIER GAS FLOW: N2, 10mL/min
 MAKE-UP: N2, 35 mL/min
 INSTRUMENT: HP 5890
 RANGE: 3, ATTENUATION: 5
 Operator: E. K. SOLGARIS



*** CALCULATION REPORT ***

CH	PKNO	TIME	AREA	HEIGHT	MR	TOND	CONC	NAME
1	1	5.458	72232	12287	1			PCB
1	2	8.932	100945	12513	2			HCB
1	3	9.359	137389	19597	3			PCP
1	4	10.824	102126	14287	4			PCNB
1	5	12.492	74921	10336	5			PCA
1	6	13.374	136461	16265	6			PCTA
1	7	16.374	80257	10559	7			PCTASO
1	8	23.154	73663	8639	8			TETASO
TOTAL			729542	105771				

Figure 3: Chromatogram of a 0.050 µg/ml Standard Solution

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SAMPLE NUMBER: 1
 SAMPLE NAME: 113
 REPT. DATE: 11/04/83 CONTROL
 DILUTION FACTOR: NA

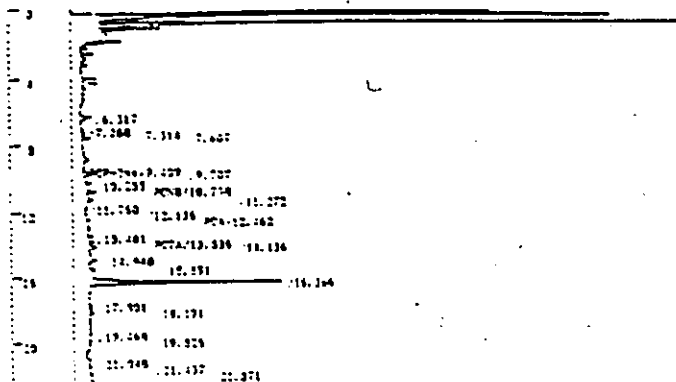
C-916 CHROMATOGRAPHY CIP-1 REPORT No. 1725 CHROMATOGRAM-2:8916.C16 12/09/80 16:49:06

Analysis File: 1:18CNDP.

Control Analytical Log

Analysis of PCB, PCP, PCB, PCB, PCA, PCFA, PCFAS & TETABD

COLUMN: RTX-12 30m, 0.53mm ID, 1.25um df
 OVEN: 100 C, 2 sec; 1 C/min to 200C, 25 C/min to 270, 2 min
 DETECTOR: ECD, 100 C
 INJECTOR: Direct, 270 C
 CARRIER GAS (FLOW): 42, 100 mL/min
 MAKE-UP: N2, 15 mL/min
 INSTRUMENT: HP 5890
 RANGE: 5, ATTENUATION: 4
 Operator: E. E. COLWARTH



*** CALCULATION REPORT ***

CH NO	TIME	AREA	HEIGHT	W. FREQ	CONC	NAME
9	6.317	2218	4120			
11	7.245	2211	270			
12	7.314	3233	237			
13	7.607	3819	370			
22	9.203	2297	213	1		PCB
23	9.727	13248	250			
24	10.258	2182	223			
27	10.739	7272	393	4		PCB
29	11.272	1463	232			
31	11.75	2296	223			
32	12.136	7253	217			
33	12.142	1275	218	1		PCA
36	13.161	1860	373			
37	13.536	2232	158	1		PCFA
38	14.136	3825	277			
39	14.740	13991	353			
40	15.151	4296	223			
42	16.166	14911	1545			
43	17.491	1827	226			
46	18.191	1338	276			
49	18.763	2105	220			
50	19.825	2123	221			
51	21.910	2117	297			
52	21.937	2149	229			
53	21.971	1287	226			
TOTAL		26967	11223	3		

Figure 4: Chromatogram of a Control Sample

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SAMPLE NUMBER: 1
 SAMPLE NAME: SPIKE
 MISC. DATA: 9110886 10 ug
 DILUTION FACTOR: 20

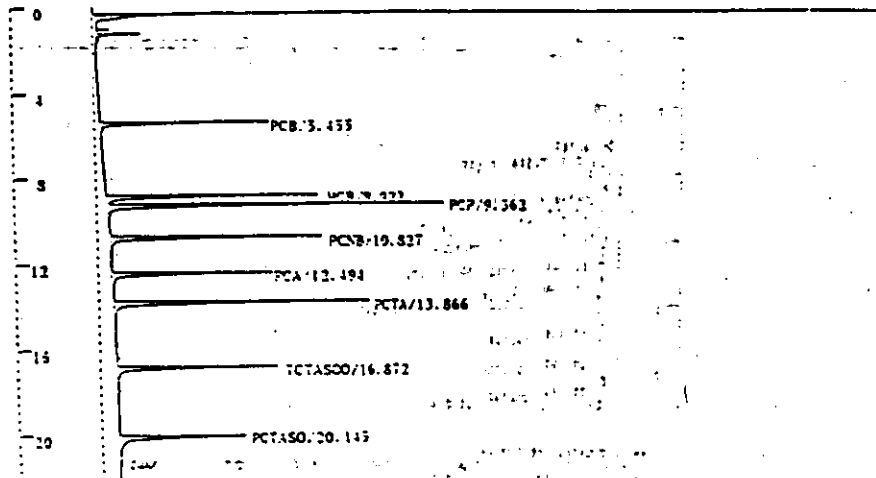
C-RIA CHROMATOPAC CH-1 REPORT No. 1784 CHROMATOGRAM-3:SD15.C23 92/09/30 22:32:58

Analysis File: 1:RCROP.

Centre Analytical Labs

Analysis of PCB, PCP, HCB, PCNB, PCA, PCTA, PCTASO & TCTASO

COLUMN: RTX-35 30m, 0.33mm ID, 0.25um df
 OVEN: 100 C, 2 min, 5 C/min to 200C, 20 C/min to 270, 5 min
 DETECTOR: ECD, 300 C
 INJECTOR: Direct, 270 C
 CARRIER GAS FLOW: H2, 10mL/min
 MAKE-UP: N2, 35 mL/min
 INSTRUME: HP 5890
 RANGE: 3, ATTENUATION: 6
 Operator: K. GOLNARIS



** CALCULATION REPORT **

CH	PKNO	TIME	AREA	WEIGHT	UK	IDNO	CONC	NAME
1	3	5.455	45016	13331	1			PCB
	13	9.352	58931	15509	2			HCB
	14	9.352	158930	26422	3			PCP
	16	10.327	99774	15468	4			PCNB
	17	12.494	82215	12672	5			PCA
	19	13.866	137898	19732	6			PCTA
	25	16.872	81100	12717	7			TCTASO
	30	20.145	73917	9936	8			PCTASO
		TOTAL	795779	127787			0	

Figure 5: Chromatogram of a Control Spiked at 10 µg and Diluted 20 fold

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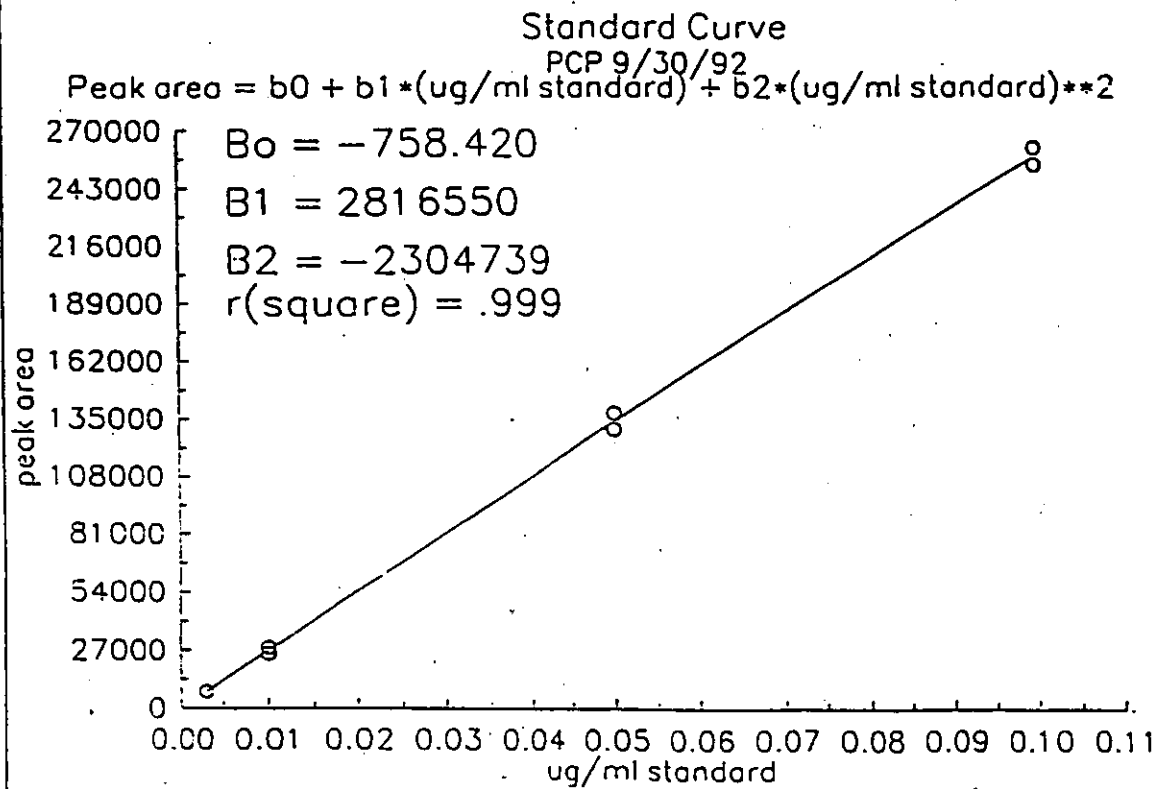


Figure 6: A Typical Calibration Curve

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VI. References

1. "Determination of Terrazole (5-Ethoxy-3-Trichloromethyl-1,2,4-Thiadiazole) and Terraclor (Pentachloronitrobenzene) and Allied Metabolites in Plant Tissues or Harvest Samples", Uniroyal Method CAM-24-73, July 3, 1973.

2. "Determination of Terraclor (Pentachloronitrobenzene) and Terrazole (5-Ethoxy-3-Trichloromethyl-1,2,4-Thiadiazole) in Soil", W.P. Griffith, April 28, 1970.

