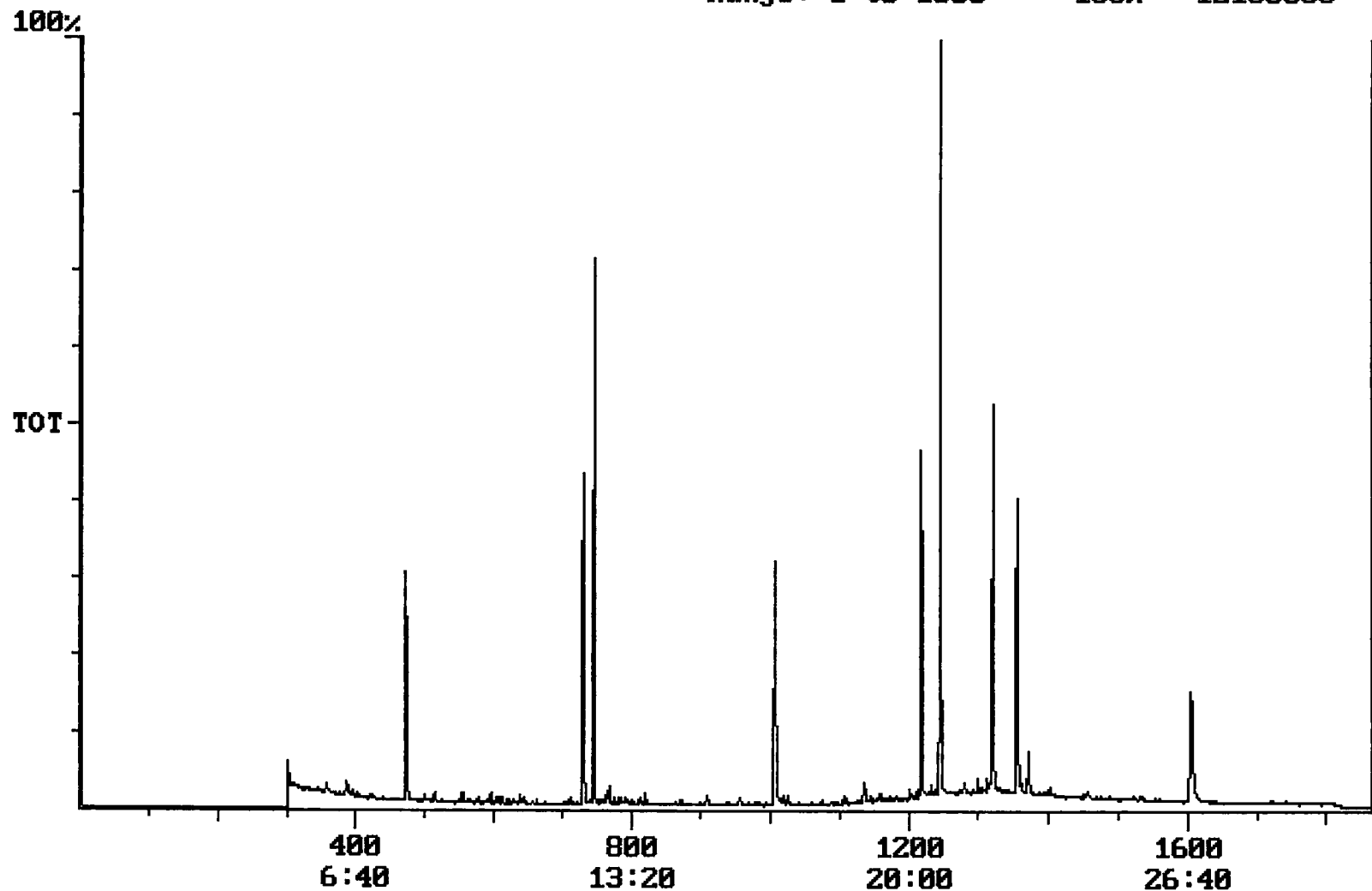


User: Bohlk, Ted
 Westchester Dept. of Labs & Research
 Date: 01/08/2002 Time: 13:07:47

Sample: AD27623
 Analysis: \$525 Organic Chemicals - 525.2
 Units: ug
 Started: 11/23/01 13:03 Ended: 12/8/01 13:03 Analyst: TB
 Result source: manual entry
 Page: 1

	Component Name	Viol	Result	Qualifier	MDL
1	Hexachlorocyclopentadiene		< MDL		0.10
2	Hexachlorobenzene		< MDL		0.10
3	Simazine		< MDL		0.40
4	Atrazine		< MDL		0.20
5	Alachlor		< MDL		0.20
6	bis(2-Ethylhexyl)adipate		< MDL		0.60
7	bis(2-Ethylhexyl)phthalate		< MDL		0.60
8	Benzo(a)pyrene		< MDL		0.20
9	EPTC		< MDL		1.0
10	2,6-Dinitrotoluene		< MDL		2.0
11	2,4-Dinitrotoluene		< MDL		2.0
12	Molinate		< MDL		0.90
13	Terbacil		< MDL		2.0
14	Acetochlor		< MDL		2.0
15	Metribuzin		< MDL		0.15
16	Metolachlor		< MDL		0.75
17	Butachlor		< MDL		0.40
18	4,4-DDE		< MDL		0.80
19	Surr_1,3-Dimethyl-2-nitrobenzen		6.0		1.0
20	Surr_Triphenyl phosphate		5.8		1.0
21	Surr_Perylene-d12		4.6		1.0

Chromatogram Plot File: E:\27623BB Date: Dec-08-1991 20:51:41
Comment: BOHLK; METHOD 525; INST 204; 12/08/01; SAMPLE 27623
Scan No: 1 Retention Time: 0:01 RIC: 0 Mass Range: 0 - 0
Plotted: 1 to 1860 Range: 1 to 1860 100% = 12153353



Integration Report

Quanfile: 27623BB

Quan Entries: 17

Comment: BOHLK; METHOD 525; INST 204; 12/08/01; SAMPLE 27623

Sorted via: Entry Number ↑

No	Name of Compound	R Time	Scan#	Mode	Peak Area	Pk Height
1	acenaphthene d-10(IS)	12:11	731	Auto	2,388,133	1,319,676
2	phenanthrene d-10 (IS)	16:46	1006	Auto	3,590,870	1,329,566
3	chrysene d-12(IS)	22:34	1354	Auto	3,048,709	1,461,997
4	p-terphenyl d-14(IS)	20:45	1245	Auto	5,798,951	4,293,498
5	acenaphthene d-10(SURR)	12:11	731	Auto	2,388,133	1,319,676
6	phenanthrene d-10(SURR)	16:46	1006	Auto	3,590,870	1,329,566
7	chrysene d-12 (SURR)	22:34	1354	Auto	3,048,709	1,461,997
8	1,3-dimethyl-2-nitrobenzene	7:52	472	Auto	1,342,810	712,695
9	triphenyl phosphate (S)	21:59	1319	Auto	1,705,666	908,796
10	perylene d-12 (SURR)	26:45	1605	Auto	2,059,904	468,133
11	hexachlorobenzene	15:11	911	Auto	3,760	1,914
12	bis(2-ethylhexyl)adipate	21:52	1312	Auto	26,463	9,593
13	bis(2-ethylhexyl)phthalate	22:51	1371	Auto	436,451	201,811
14	benzo(a)pyrene	26:32	1592	Auto	855	855
15	2,6-dinitrotoluene	11:51	711	Auto	15,348	8,877
16	butachlor	20:18	1218	Auto	19,880	10,162
17	4,4'-dichlorodiphenyl ether	20:40	1240	Auto	3,857	2,072

reviewed IS/ENR's

Quantitation Report Quanfile: 27623BB Quan Entries: 17
 Comment: BOHLK; METHOD 525; INST 204; 12/08/01; SAMPLE 27623
 Sorted via: Entry Number ↑ (S) = Standard

Cal	Name of Compound	S	Scan#	R Time	Me	Calc Amt(A)	Units
1	acenaphthene d-10(IS)	I	731	12:11	BV	5.000	UG/L
2	phenanthrene d-10 (IS)	I	1006	16:46	BV	5.000	UG/L
3	chrysene d-12(IS)	I	1354	22:34	BV	5.000	UG/L
4	p-terphenyl d-14(IS)	I	1245	20:45	BB	5.000	UG/L
5	acenaphthene d-10(SURR	A	731	12:11	BV	2.268	UG/L
6	phenanthrene d-10(SURR	A	1006	16:46	BV	2.664	UG/L
7	chrysene d-12 (SURR)	A	1354	22:34	BV	2.764	UG/L
8	1,3-dimethyl-2-nitrobe	A	472	7:52	BV	6.026	UG/L
9	triphenyl phosphate (S	A	1319	21:59	BB	5.774	UG/L
10	perylene d-12 (SURR)	A	1605	26:45	BB	4.638	UG/L
12	hexachlorobenzene	A	911	15:11	BB	0.016	UG/L
16	bis(2-ethylhexyl)adipa	A	1312	21:52	MM	0.059	UG/L
17	bis(2-ethylhexyl)phtha	A	1371	22:51	VB	0.493	UG/L
18	benzo(a)pyrene	A	1592	26:32	BB	0.002	UG/L
20	2,6-dinitrotoluene	A	711	11:51	BB	0.109	UG/L
27	butachlor	A	1218	20:18	BB	0.061	UG/L
28	4,4'-dde	A	1240	20:40	BB	0.013	UG/L

. Comments for Sample AD27623 - Analysis \$525

Westchester Dept. of Labs & Research

User: Vinci, David L

Date: 01-22-2002 Time: 14:12:48

The recoveries for 2 of the 18 compounds in the LCS Standard were below their acceptance ranges. The recoveries for 3 of the 18 compounds in the Spiked Sample were outside of their acceptance ranges. The breakdown for Endrin in the Breakdown Standard was 61%. The injection port is kept at 280 degrees Celcius which may be contributing to high Endrin Breakdown. An investigation into modifying injection port parameters will be made.