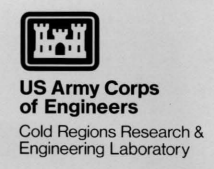
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Comparison of extraction techniques and solvents for explosive residues in soil

Thomas F. Jenkins and Daniel C. Leggett

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etics and recovery. Two solvents, a	acetonitrile and methan	nol, and four extraction tech-
niques, Soxhlet, ultrasonic bath, me	chanical shaker and ho	mogenizer-sonicator were com-
pared. The results were complex in	that some interaction	s among analyte, method and
solvent were found. Acetonitrile w	as round to be clearly	superior to methanol for RDX and

HMX. Soxhlet and ultrasonic bath generally recovered more than homogenizer or shaker, although a complicating factor is that all techniques were not necessarily at equilibrium. In terms of sample throughput, the ultrasonic bath and shaker are preferred over Soxhlet

PREFACE

This report was prepared by Thomas F. Jenkins and Daniel C. Leggett, Research Chemists, Earth Sciences Branch, Research Division, U.S. Army Cold Regions Research and Engineering Laboratory. Funding for this research was provided by the U.S. Army Toxic and Hazardous Materials Agency, Aberdeen Proving Ground, Maryland, under order no. Al-5-R0001-XX-Al-48.

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COMPARISON OF EXTRACTION TECHNIQUES AND SOLVENTS FOR EXPLOSIVE RESIDUES IN SOILS

by

Thomas F. Jenkins and Daniel C. Leggett

INTRODUCTION

The Army is interested in analyzing soils for explosive contaminants both to aid in decontaminating facilities and to stop pollution. Army laboratories or contractors hired to conduct surveys have not been supplied with a uniform protocol for extracting these contaminants from soils. As a result, various extraction techniques and solvents have been used, often without sufficient validation of their use for particular contaminants in particular soils. Thus the quantitative and in some cases the qualitative accuracy of the results of soil contaminant surveys is variable and not always known. The purpose of this work was to identify or develop a single methodology for analyzing explosive contaminants in soils that was as accurate and as rapid as possible.

Since reverse-phase HPLC (High Performance Liquid Chromatography) would most likely be the analytical method of choice for mixtures of explosive contaminants (Jenkins et al. 1984), we concentrated our efforts on those solvents that are compatible with this analytical technique. Desirable properties of such solvents are miscibility with water and low UV absorption at 254 nm. These solvents include methanol and other alcohols, acetonitrile and tetrahydrofuran. When GC (Gas Chromatography) or other type of analytical finish is used, other solvents may be more appropriate.

Goerlitz and Law (1975) extracted TNT and RDX from soils with acetone-hexane prior to GC analysis. The method involved a 12-hour equilibration period with acetone alone, followed by addition of hexane and two additional extractions with no waiting period. Three extractions were required for quantitative (93-99%) recovery because the first extraction yielded only 55%. However, it is hard to tell how much of this was attributable to incomplete solvent recovery and how much to incomplete extraction.

As part of a previous preliminary study in our laboratory, acetone, methanol, acetonitrile and tetrahydrofuran were compared for extracting TNT, TDX and HMX from a soil and a sediment (Cragin et al. 1985). A single batch extraction was used, as little additional recovery was obtained in a second extraction. The solvent-to-soil ratios were higher than those of Goerlitz and Law, however, and the solvent was not diluted with hexane prior to its separation from the soil. Samples were placed in an ultrasonic bath for 5 minutes followed by 1 hour of shaking. Extraction recovery varied with type of sample and the concentration of explosive present. Acetone interfered with quantification of HMX by HPLC. Recoveries from the other three solvents were within 20%.

Both Goerlitz and Law (1975) and Cragin et al. (1985) used unfortified, naturally contaminated soils and recoveries were based on the amounts obtained by exhaustive extractions. Miller et al. (1983) tested single batch extractions with acetone, acetonitrile and methylene chloride/methanol (95/5) on a sediment that was fortified with eight different explosive compounds. The best overall solvent was methylene chloride/methanol. Recoveries of TNT and RDX ranged from 87-107%, but tetryl was poorly recovered. It thus appears that any of several solvents or solvent mixtures could be used to extract explosive compounds. Since we are mainly interested in using reverse-phase HPLC analysis, we chose to evaluate methanol and acetonitrile since these solvents can be analyzed directly or after dilution with water. Acetone is good but causes some interference with early peaks. Tetrahydrofuran is also good and is compatible with HPLC, but appeared to offer no advantage over acetonitrile or methanol so was not evaluated further.

Unfortunately, the state of the art is not such that we can reliably match extraction solvents with particular contaminants to obtain the best solvent-contaminant pairing, and selections have been largely empirical in the past. In a review of the literature from 1960 to 1981, Dao et al. (1983) found methanol and acetone to be the most frequently used single solvents for soil pesticide analyses. Nonpolar solvents such as hexane are seldom used, presumably because they don't work as well, even for nonpolar residues. A reason for this apparent contradiction is given by Freeman and Cheung (1981), who suggest that failure to properly match the solubility parameter of the extracting solvent with that of the soil organic matter (the presumed site of sorption) will result in low recovery regardless of

equilibration time. This apparent contradiction of equilibrium thermodynamics is explained as failure of the solvent to "swell" or "wet" internal polymer surfaces, resulting in failure of the system to come to equilibrium within the practical time constraints imposed by the need to complete the analysis. On the other hand, we cannot ignore the solubility of contaminants in the extracting solvent either, particularly when they are present in concentrations that would exceed the ability of the solvent to dissolve them. Thus, solubility in extracting solvents during extraction and subsequent work-up will be considered carefully here.

In view of the foregoing discussion, careful attention will also be paid to extraction kinetics, as the shortest practicable equilibration time is desirable in a method. The usual ways of decreasing contact times are shaking, homogenization and sonication. Drying of sediments and soils prior to extraction was also considered in an earlier study (Cragin et al. 1985). This point deserves some comment here. It was found that TNT, RDX and HMX were extracted as efficiently from wet soils as from freezedried soils using methanol as the extracting solvent. It appears that this is generally the case when polar solvents are used. Use of water-imiscible solvents, however, generally leads to lower recoveries from wet sediments regardless of the extraction technique used.

Haddock et al. (1983) extracted anthracene better from dry sediments than from wet sediments 8 days after spiking, using either benzene or benzene-acetonitrile (13:7). Peterson and Freeman (1982) showed that using Soxhlet extraction for dried sediments gave higher recoveries of added phthalate esters than those that they obtained from wet sediments using methylene chloride as the extracting solvent. This same study also indicated that ultrasonication with this solvent gave recoveries that were as high as or higher than did Soxhlet (presumably when both were done on dried sediment) and required only 1 hour for completion as opposed to 48 hours for Soxhlet. This latter finding is similar to that of Wegman and Hofster (1982), who studied extraction of a group of organochlorine pesticides added to sediment for varying lengths of time. They found that shaking, either with petroleum ether or with 1:1 acetone-petroleum ether, gave recoveries as high as those that they obtained from Soxhlet extraction using these solvents. They also found that acetone-petroleum ether was superior to petroleum ether and that it extracted as efficiently from wet as from dry sediment. Acetone, however, appeared to be better than the

mixed solvent for extracting contaminants from wet sediment. These conclusions are not supported by rigorous statistical evaluation.

These studies are mentioned because they suggest that extended Soxhlet extraction may not necessarily give the highest recovery, contrary to popular expectation. One reason for this could be irregular contact with the sample caused by channeling or the nonuniform nature of the sample itself. These types of problems would naturally tend to be reduced by vigorous shaking, homogenizing or sonicating. On the other hand, Sporstol et al. (1983) claim that Soxhlet extraction is generally superior to ultrasonication for natural contaminants present in sediments. However, their study suffers from a serious flaw in that Soxhlet extraction of dried sediment is compared to ultrasonic extraction of wet sediment using dichloromethane as extractant. As pointed out above, this is not a fair comparison, and extraction techniques should be evaluated using similar sample preparation. The addition of 5% methanol to the extractant significantly improved recoveries from the wet-ultrasonicated sediment, another indication that moisture might have been the reason for low recovery.

In addition to their being the best choice of solvents for analysis by reverse-phase HPLC, methanol and acetonitrile are completely miscible with water so that they can be used for extracting either wet or dried samples. For this study, dried samples will be used since it was shown that freezedrying or air-drying produce small or insignificant losses of TNT, RDX and HMX (Cragin et al. 1985). The small losses are compensated for by the ease of working with dry material, which can be sampled more precisely. Finally, we chose to work with "naturally contaminated" sediments rather than with spiked sediment because of the additional ambiguity that the latter introduces (Peterson and Freeman 1982, Sporstol et al. 1983, Haddock et al. 1983, Cragin et al. 1985).

The objective of this study was to determine which of several alternative extraction procedures gave the best recoveries of HMX, RDX and TNT for contaminated soils. Methanol and acetonitrile were examined as extraction solvents because preliminary tests indicated good results for explosive residues and because they are suitable for use with the existing HPLC analytical procedure. The extraction methods evaluated were: continuous Soxhlet extraction, batch extraction by equilibration with solvent on a wrist-action shaker, batch extraction using a soil-plant homogenizer and batch extraction by equilibration with solvent in a bath sonifier.

EXPERIMENTAL

Analytical instrumentation

All analytical determinations for HMX, RDX, TNB, TNT and DNT were obtained using reverse-phase HPLC (Jenkins et al. 1984). The instrumental setup utilized a Perkin Elmer Series 3 pump with a Rheodyne 7125 loop injector and a Spectra-Physics SP8300 fixed 254-nm UV detector. Peak areas were quantified using an HP 3390A Integrator. Analyses were obtained on a Supelco 25 cm x 4.6 mm LC-8 column (5 μ m) using a mobile phase of methanol/acetonitrile/water, 38:12:50 (v/v/v). Samples were injected by over-filling either a 20- μ L or a 100- μ L sample loop. Analyses were obtained in duplicate and mean values reported.

Chemicals

All analytical standards for HMX, RDX, TNB, TNT and DNT were prepared from Standard Analytical Reference Materials (SARMs) obtained from the .

U.S. Army Toxic and Hazardous Materials Agency, Aberdeen Proving Ground,

Maryland. Samples were dried to constant weight in a vacuum desiccator over dry calcium chloride in the dark.

The methanol and acetonitrile used to prepare the mobile phase for HPLC determinations and as extraction solvents were either Baker HPLC grade or Mallinckrodt Chrom AR HPLC grade solvents. Water used in the preparation of the HPLC mobile phase and as sample diluent was purified using a Milli-Q Type I Reagent Grade Water System (Millipore Corporation). Methanol, acetonitrile and water were combined in the proper proportions and vacuum-filtered through Whatman GF-F glass microfibre filters to remove particulate matter and degas the solvent.

Soils

The soil samples used for extraction experiments were collected at the Iowa Army Ammunition Plant (AAP) on 25 and 26 July 1983. Soil no. 6 was obtained from an old ordnance-burning area that had not been used since 1981. Soil no. 2 was obtained from the surface of an old disposal lagoon. Each soil was air-dried to constant weight, ground with a mortar and pestle, and passed through a no. 30 mesh sieve. Both soils were placed in individual Nalgene bottles and mixed thoroughly on a roller mill to obtain as homogeneous a sample as possible to allow comparison of various extraction techniques.

Kinetic studies

Kinetic studies were conducted on four different procedures for two different soils. These were solvent extraction methods using a wrist-action shaker, a bath sonifier, a soil-plant homogenizer and a Soxhlet extractor. In each case methanol and acetonitrile solutions containing a DNT internal standard were individually tested at a soil-to-solvent ratio of 1 g/25 mL. Tests indicated that DNT did not sorb to soil at measurable levels in the presence of the extracting solvent.

Soil-plant homogenizer

For soil no. 6, two 1-g samples of soil were placed in 45-mL Pyrex centrifuge tubes; 25 mL of a methanol solution containing a DNT internal standard was added to one and 25 mL of an acetonitrile solution containing a DNT internal standard was added to the other. Each sample was ground in a soil-plant homogenizer (PT 10/35 Brinkmann Homogenizer with PTA 20S Generator) at a setting of 4 for periods of 1, 2, 4, 8 and 16 minutes (Fig. 1). After each period, the sample was centrifuged at 3500 rpm until the solution cleared (5-15 min), a 2-mL portion of clear supernatant was

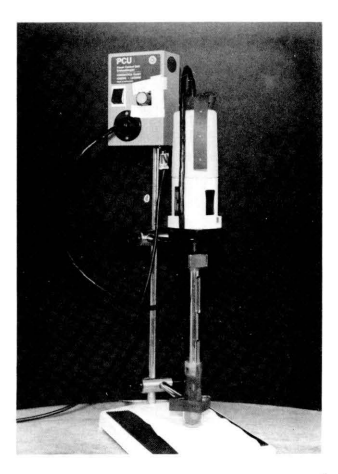


Figure 1. Brinkmann soil-plant homogenizer.

removed and the tube was replaced on the homogenizer. Each 2-mL aliquot was mixed with 2 mL of water and filtered through a 0.45-µm Gelman Acrodisc CR disposable filter assembly using a 10-cm³ B-D disposable syringe. The first 1-mL portion of the filtrate was discarded and the remainder saved for HPLC analysis. There was significant solvent evaporation during homogenizing and centrifuging, but use of the internal standard corrected for this factor. Soil no. 2 was treated in an identical manner, except grinding times were 5, 10, 20 and 60 minutes.

Bath sonifier

For this procedure, 2-g samples of soil no. 6 were placed in two 2.5-cm x 20-cm screw cap test tubes equipped with a Teflon-lined cap. We added 50 mL of either methanol or acetonitrile containing a DNT internal standard. The soil in each tube was dispersed with a vortex mixer (Vanlab Model K-550-G) for 1 minute and placed in a sonic bath (Cole-Parmer Model 8845-60) for periods of 1, 2, 4, 8, 16, 32, 64 and 240 minutes (Fig. 2). The samples were redispersed with the vortex mixer between periods. After each period, the tubes were centrifuged at 3500 rpm for 5 minutes, and a 5-mL portion of the supernatant was removed with a volumetric pipette, mixed with 5 mL of water and filtered as described above. Soil no. 2 was treated as described above, except equilibration times studied were 15, 30, 60, 120, and 420 minutes.

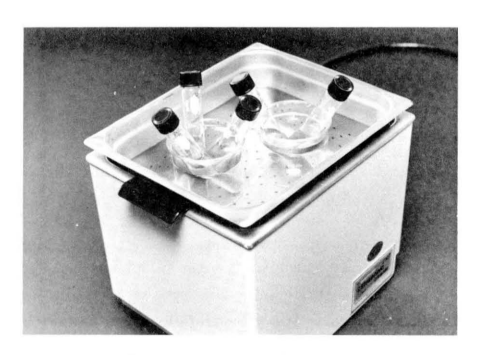


Figure 2. Vanlab model K-550 bath sonifier.

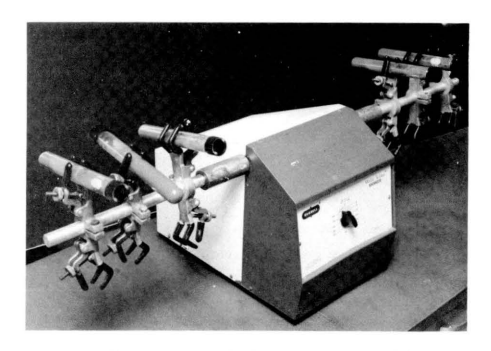


Figure 3. Burrell model 75 wrist-action shaker.

Wrist-action shaker

Two 2-g samples of soil no. 6 were placed in 2.5-cm x 20-cm screw cap test tubes and 50 mL of either methanol or acetonitrile containing a DNT internal standard was added. The soil was dispersed with a vortex mixer for 1 minute and placed on a wrist-action shaker (Burrell Model 75) at maximum speed for periods of 10, 30, 60, 90, 150, 240, 480 and 1440 minutes (Fig. 3). After each period, 5-mL samples were removed and processed as described for the bath sonifier experiment and the soil redispersed with the vortex mixer prior to beginning the next time interval. Soil no. 2 was treated in a similar fashion, except samples were taken for analysis at 30, 60, 120, 240, 1440 and 2880 minutes.

Soxhlet extractor

Two 16-g samples of soil no. 6 were placed in Soxhlet extraction thimbles (Whatman, cellulose) and extracted on Soxhlet extractors with 400 mL of either methanol or acetonitrile containing a DNT internal (Fig. 4). After 1, 2, 4, 25 and 37 hours, 5-mL samples were withdrawn with glass volumetric pipettes, mixed with 5 mL of water in 20-mL scintillation vials and filtered as described earlier. Samples were refrigerated immediately after collection and retained for HPLC analysis. The cycle time on the Soxhlet extractors was about 15 minutes. Soil no. 2 was treated identically, except samples were withdrawn at 1, 2, 4, 24 and 48 hours.

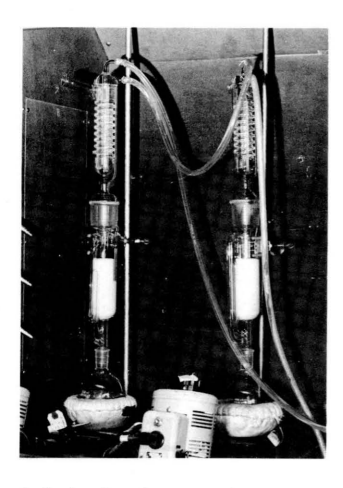


Figure 4. Soxhlet extractor.

Analysis of variance experiment

Following the completion and assessment of the kinetic experiments, the four extraction procedures were compared. This was done by extracting six replicates of soil no. 6 by each approach using the amount of time appropriate for each method. For the wrist-action shaker and the Soxhlet extractor, we used 24-hour extraction periods. For the sonic bath and the soil-plant homogenizer, the periods were 4 hours and 10 minutes respectively. This experiment was conducted twice, once with acetonitrile and once with methanol, each containing a DNT internal standard. The extraction and analytical procedures used were identical to those described in the kinetic experiments. The soil was weighed out for these two experiments at one time in random order. All analyses for a given extraction solvent were conducted on a single day in duplicate, in random sequence. Mean values were compared using a two-way analysis of variance.

Subsequently, a similar study was conducted using all four methods and soil no. 2. The extraction times for each method were 24 hours for the Soxhlet, wrist-action shaker and bath sonifier, and 30 minutes for the soil-plant homogenizer.

RESULTS AND DISCUSSION

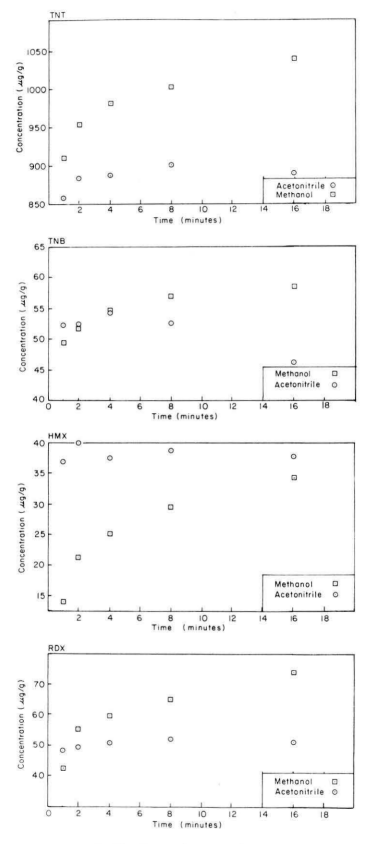
Soil no. 6

Kinetic study

Our initial work was concentrated on soil no. 6 from the Iowa AAP. This soil was collected from an ordnance-burning area and was chosen because preliminary analysis indicated that it contained rather high levels of TNT and measurable amounts of HMX, RDX and TNB (a TNT degradation product). Initially, kinetic studies were conducted on each extraction method in an attempt to determine the length of time required to achieve equilibrium in the case of the three batch methods, or complete extraction in the case of the Soxhlet procedure. These results are presented in Appendix A, Table Al for the soil-plant homogenizer, sonic bath, wrist-action shaker and Soxhlet extractor respectively. These data are plotted versus time in Figure 5. Example chromatograms for acetonitrile and methanol extracts are presented in Figure 6 for aliquots from the Soxhlet extractor versus time.

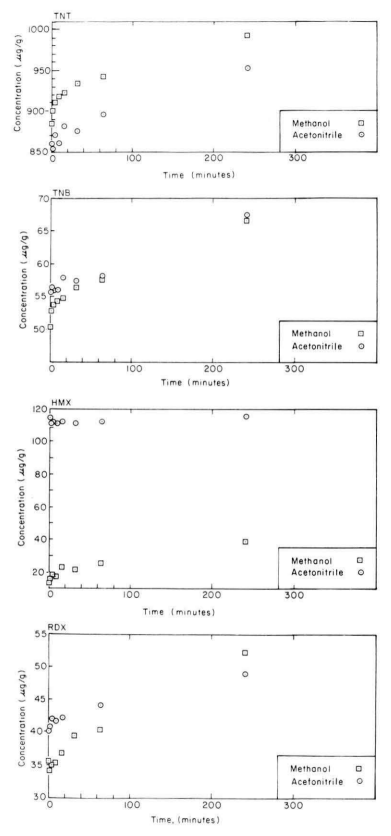
For the soil-plant homogenizer (Fig. 5a), short periods (1-16 minutes) were studied since this device grinds one sample at a time. Thus, from a practical point of view, it must reach equilibrium quickly or it is of no value, since in most USATHAMA applications a large number of samples will need to be analyzed. For the acetonitrile extracts, maximum values appear to be reached at 8 minutes for all four analytes. For methanol, there was a much slower rise and the solution was apparently not at equilibrium with the soil after 16 minutes. Care should be used when comparing the maximum values achieved in the two solvents since it is very difficult to obtain totally homogeneous subsamples of soil, and the differences in maximum values may be attributable to subsample inhomogeneity.

Since the ultrasonic bath method can process a large number of samples at a time, the use of fairly long time periods is practical. For that reason, we investigated times up to 4 hours (240 minutes)(Fig. 5b). For all four analytes, the highest values reached in both solvents were for the 4-hour samples. Since no longer periods were studied, there is no guarantee that maximum values have been obtained, although one reputed advantage of the ultrasonic bath method is its speed in reaching equilibrium. Again it is risky to compare the maximum values obtained for the two solvents on different subsamples; however, values for HMX in acetonitrile were nearly



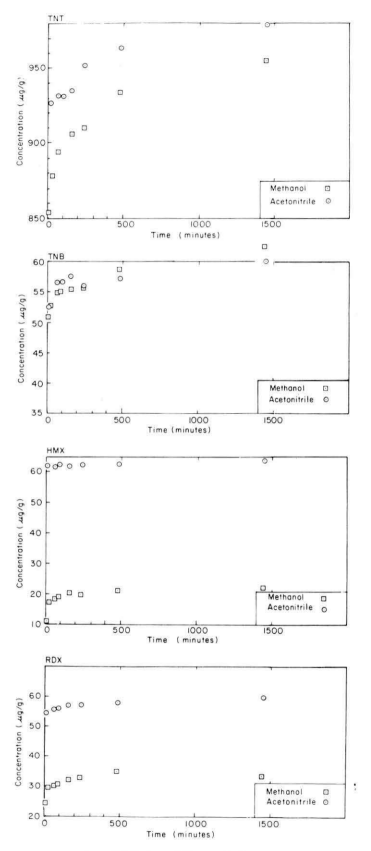
a. Soil-plant homogenizer.

Figure 5. Plots of analyte concentrations for the kinetic study of Iowa AAP soil no. 6.

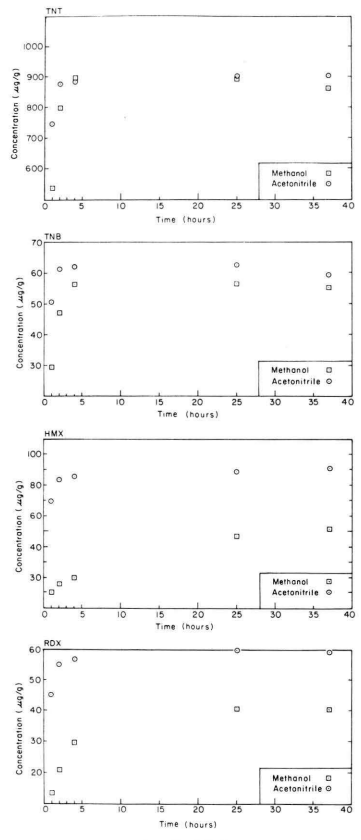


b. Sonic bath.

Figure 5 (cont'd). Plots of analyte concentrations for the kinetic study of Iowa AAP soil no. 6.



c. Wrist-action shaker.
Figure 5 (cont'd).



d. Soxhlet.

Figure 5 (cont'd). Plots of analyte concentrations for the kinetic study of Iowa AAP soil no. 6.

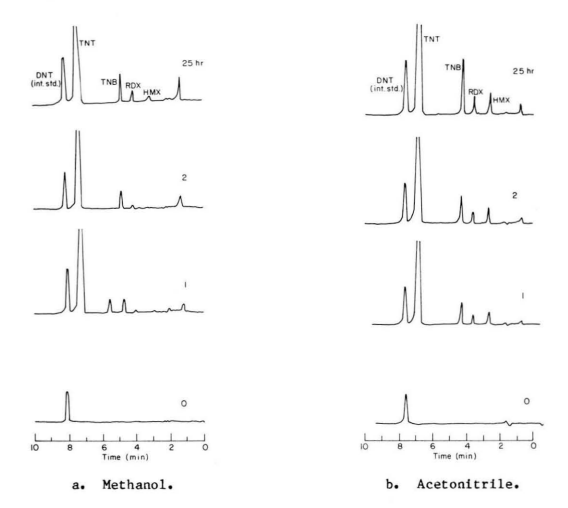


Figure 6. Example of chromatogram obtained for the kinetic study of Iowa AAP soil no. 6 with the Soxhlet extractor.

three times those found for the methanol extract. The rate at which maximum values were achieved for HMX was also greater for acetonitrile.

The values obtained for the wrist-action shaker are shown in Figure 5c. Since up to 12 samples can be shaken simultaneously on the Burrell shaker with no one in attendance, use of long equilibrium periods is practical, if necessary. We tested times up to 24 hours (1440 minutes). For RDX and HMX, it appears that fairly constant values were obtained at 4 hours. For TNT and TNB, the highest values for both solvents were clearly found at 24 hours. Because of the scale of the figure, the 24-hour values for TNT appear to be much higher than the 4-hour values; however, the increase amounts to less than 2%. For TNB the increase amounts to only about 5%.

As was observed for the sonic bath, the values obtained for HMX were much lower in the methanol extract than in acetonitrile. RDX values were

also markedly less in the methanol extract, but this trend was not observed for either TNB or TNT.

Figure 5d presents the values found for the kinetic study with the Soxhlet extractor. This device differs from the other three in that the analytes are removed from interaction with the solid soil as they are extracted, rather than relying on a favorable equilibrium condition. In addition, the extracting solvent is heated well above room temperature and the extracted components are maintained at this elevated temperature for long periods. Thus the method does not depend on a favorable equilibrium, and complete separation of the analyte from the soil is possible. For TNT, TNB and RDX, maximum values seemed to be reached at 25 hours. For HMX, slightly higher values were obtained at 37 hours, but this increase only amounted to about 2% for acetonitrile and 8% for methanol. As previously observed, maximum values for HMX were much higher in the acetonitrile extracts than in methanol. RDX followed a similar pattern, while TNT and TNB were about the same in both solvents. The maximum values obtained by the Soxhlet method are in about the same range as those found by the other methods, indicating that solvent-soil partitioning is strongly in favor of solution.

The Soxhlet method was included in the study as a means of comparing the other methods. The Soxhlet requires a separate, relatively expensive setup for each sample and would be very expensive to use for analyzing large numbers of samples.

Replicate study

To allow comparison of the four techniques, six replicate subsamples of soil no. 6 were processed by each method. The subsamples were weighed randomly and analyzed to allow comparisons of means and variances by standard analysis of variance techniques. The equilibrium times used were 10 minutes for the soil-plant homogenizer, 4 hours for the sonic bath and 24 hours for the wrist-action shaker. Soxhlet samples were also processed for 24 hours.

The individual values obtained for each method are presented in Table A2. The means and standard deviations for each analyte by each method are presented in Table 1 for acetonitrile and methanol.

Looking at the relative standard deviation (RSD), for both acetonitrile and methanol, it is clear that the analytes fall into two groups.

Table 1. Means and standard deviations for replicate study on Iowa AAP soil no. 6.

Analyte	Method	Mean* (μg/g)	Standard deviation (µg/g)	RSD (%)
	Acetoni	trile Ext	ract	
TNT	Shaker Sonic bath Homogenizer Soxhlet	880.7 a 882.9 a 848.9 b 881.2 a	9.7 13.0	1.44 1.09 1.53 0.86
TNB	Shaker Sonic bath Homogenizer Soxhlet	55.1 b 55.6 b 51.4 c 62.1 a	0.6 0.8	1.88 1.04 1.51 1.83
RDX	Shaker Sonic bath Homogenizer Soxhlet	54.2 a 54.5 a 64.4 a 65.1 a	13.3 55.8	17.9 24.3 86.7 16.3
нмх	Shaker Sonic bath Homogenizer Soxhlet	82.0 a 56.4 a 64.9 a 84.1 a	14.4 26.8	38.9 25.5 41.2 13.6
	Methan	ol Extrac	t	
TNT	Shaker Sonic bath Homogenizer Soxhlet	895.4 a 840.4 b 870.2 a 891.4 a	25.0 38.7	1.60 2.97 4.44 0.57
TNB	Shaker Sonic bath Homogenizer Soxhlet	56.0 b 52.5 c 53.4 c 58.3 a	1.5 1.1	1.37 2.90 1.97 1.08
RDX	Shaker Sonic bath Homogenizer Soxhlet	36.9 a 39.9 a 30.9 a 48.0 a	7.8 16.6	31.9 19.6 53.7 10.9
НМХ	Shaker Sonic bath Homogenizer Soxhlet	22.3 a 33.4 a 28.4 a 58.8 a	10.6 18.0	24.2 31.6 63.5 9.8

^{*}Values indicated with the same letter are not significantly different at the 95% confidence level.

The RSD for TNT and TNB is generally in the range of 1-3%, indicating very good analytical precision, as well as good soil homogenization prior to subsampling. The RSD for RDX and HMX, on the other hand, is quite large, generally at or above 20% and for the homogenizer sometimes over 50%. The mean values for HMX and RDX are about the same as that for TNB, and hence this difference in RSD is apparently not a result of differences in the mean concentration. It thus appears that RDX and HMX are distributed differently, and less homogeneously, than TNT and TNB in this soil.

Because of the much larger sample size for the Soxhlet procedure (16 g vs either 1 or 2 g), RSD values for the Soxhlet procedure are lower than those for the other methods for HMX and RDX in both solvents. In addition, RSD values for the homogenizer, where 1-g subsamples were used, were generally higher than for the shaker and sonic bath where 2-g subsamples were used, as we expected (Grant and Pelton 1976).

Because of the very good precision and low random error obtained for TNT, analysis of variance (Table 2) indicates a significant difference for the methods at the 95% confidence level. For TNT, the Soxhlet and wrist-action shaker give very similar results with both solvents. The soil-plant homogenizer gives significantly lower values with acetonitrile while the sonic bath is low for methanol. This results in a significant interaction between method and solvent at the 95% confidence level, even though the difference between solvents is not significant overall.

For TNB, excellent precision also enables sensitive comparison of the various extraction procedures and solvents (Table 1); method, solvent and interaction are all significant at the 95% confidence level (Table 2). With both solvents, the Soxhlet procedure is clearly superior to the other three methods with respect to the amount extracted, the largest difference being observed with acetonitrile. The reason for this superiority of the Soxhlet for TNB but not for TNT is unclear, particularly because of the similarity in structure for the two substances. As was observed for TNT, extraction using the sonic bath is clearly better with acetonitrile than methanol while the opposite is true for the soil-plant homogenizer.

The sensitivity of the analysis of variance for RDX and HMX is limited by the large standard deviations for both analytes. This large random error makes it impossible to observe the small differences in the various methods observed for TNT and TNB. Thus in neither case was method found to be significant at the 95% confidence level.

Table 2. Results of two-way analysis of variance for Iowa AAP soil no. 6.

	SS	df	Mean square	F
		TNT		
Method	8511.22	3	2837.07	8.02*
Solvent	11.02	1	11.02	0.03
Interaction	7732.83	3	2577.61	7.29*
Error	14146.9	40	353.67	
LSD = 29.2				
		TNB		
Method	404.13	3	134.7	139.11*
Solvent	11.43	1	11.43	11.8*
Interaction	75.58	3	25.19	26.02*
Error	38.73	40	0.9683	
LSD = 1.53				
		RDX		
Method	846.13	3	282.04	0.59
Solvent	4503.3	1	4503.3	9.39*
Interaction	348.2	3	116.0	0.24
Error	19177.9	40	479.45	
		нмх		
Method	5329.2	3	1776.4	5.53
Solvent	15668.2	1	15668.5	48.77*
Interaction	2539.2	3	846.4	2.63
Error	12850.2	40	321.3	

^{*} Values are significant at the 95% confidence level. Critical values are F(3,40) = 2.84 and F(1,40) = 4.08.

Solvent, on the other hand, was found to be significant for TNB, RDX and HMX even with the large random error variance of the latter two explosives. Use of acetonitrile clearly resulted in higher extraction efficiency for both substances. Whether this is a thermodynamic difference attributable to solubility, or to an unfavorable partition coefficient or a kinetic one, is unclear. The solubilities of RDX and HMX are higher in acetonitrile than in methanol (unpublished data, D.C. Leggett); therefore partitioning is also expected to be more favorable into acetonitrile. In

either case, from a practical point of view, acetonitrile is clearly superior for RDX and HMX extraction with this soil.

Soil no. 2

Kinetic Study

Following the work with soil no. 6, it was of interest to us to determine if similar behavior would be observed with a different soil, particularly one with much lower TNT concentrations. Preliminary analysis indicated that the Iowa AAP soil no. 2 had much lower, but measurable, concentrations of TNT and traces of TNB, RDX and HMX. This soil was from an old disposal lagoon and hence the mode of contamination should have been somewhat different from that of soil no. 6, which came from an ordnance burning area.

A kinetic study similar to that conducted with soil no. 6 was done first. All four extraction methods were tested with both acetonitrile and methanol. Because of the results with soil no. 6, equilibration times were modified from those used earlier. In particular, the longest time increments for the Soxhlet and wrist-action shaker were extended to 48 hours. The results of this kinetic study are presented in Table A3 and plotted in Figure 7 for TNT. Only trace levels were obtained for TNB, RDX and HMX, and since the peaks were not consistently integrated, no attempt was made to quantify these substances. Examples of chromatograms obtained for extracts from the wrist-action shaker are presented in Figure 8.

TNT levels were indeed much lower for soil no. 2 than for soil no. 6. The kinetics of desorption was also slower for all four methods. Thus, it is likely that TNT in soil no. 2 was associated with the soil in a somewhat different manner than it was in soil no. 6. Since soil no. 2 is from the surface of a disposal lagoon, it is likely that TNT associated with this soil was initially present in solution and became associated through adsorption as water evaporated from the surface or percolated through. Soil no. 6 was from an ordnance burning area, and it is likely that TNT became associated with this soil as discrete crystals, which may have dissolved and reassociated through rainfall and transpiration.

One explanation that may account for the slower desorption kinetics observed for soil no. 2 is that the low level of TNT present in this soil is preferentially adsorbed to high energy binding sites. Since the number of these types of sites is limited, the large amount of TNT associated with soil no. 6 probably exceeds what can be adsorbed in this fashion. The bulk

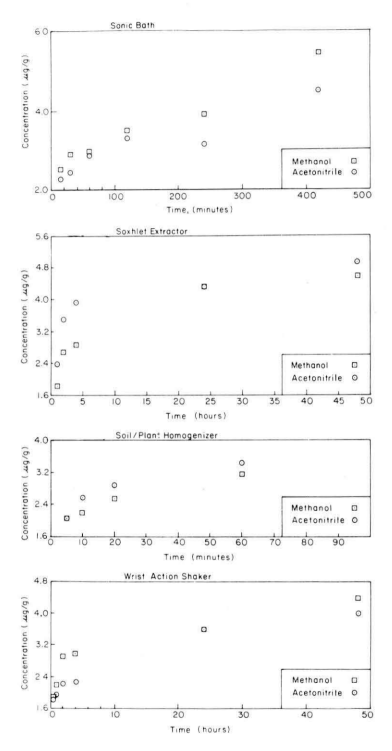


Figure 7. Plots of TNT values for the kinetic study of Iowa AAP soil no. 2.

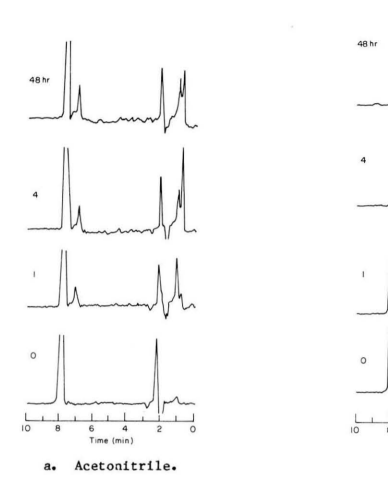


Figure 8. Example of chromatogram obtained for the kinetic study of Iowa AAP soil no. 2 with the wrist-action shaker.

Time (min)

Methanol.

of the TNT may therefore be less tightly bound. The activation energy for desorption of this material would be lower, resulting in faster desorption kinetics.

Looking at the kinetic results (Fig. 7), it appears that in none of the four methods was equilibrium definitely established during the experiment for either solvent. The Soxhlet and wrist-action shaker appear to be closer to a terminal value for the times studied than do the soil-plant homogenizer or the sonic bath, which were both studied for shorter periods. Little difference was observed between solvents. Clearly, use of the soil-plant homogenizer for extended periods is impractical since it is a one-sample-at-a-time technique and, in addition, results in a large degree of solvent evaporation since the tubes are required to be open to the atmosphere during grinding. Use of the sonic bath for longer periods is possible, however. The wrist-action shaker and Soxhlet appear to have

come fairly close to final values after 24 hours and, from a practical point of view, this is about the maximum time acceptable for soil equilibration-extraction. Thus for the next phase, the replicate study, we decided to compare the four techniques using a 24-hour period for the Soxhlet, the wrist-action shaker and the sonic bath and 30 minutes for the soil-plant homogenizer.

Replicate study

As with soil no. 6, six replicate subsamples of soil no. 2 were extracted by each of the same four techniques. Methanol and acetonitrile were both tested at the time periods described above. The individual results for TNT and TNB are presented in Table A4. Mean values and standard deviations are presented in Table 3 and the results of a two-way analysis of variance in Table 4.

Table 3. Means and standard deviations for replicate study on Iowa AAP soil no. 2.

Mean	Mean (μg/g)	Standard deviation (µg/g)	RSD (%)	Extract
rieati	(18/8)	(PB/B/	(%)	EXCIACE
		TNT		
Shaker	2.46 c	0.124	5.04	Acetonitrile
Sonic bath	3.54 ь	0.223	6.30	
Homogenizer	2.10 d	0.108	5.14	
Soxhlet	4.35 a	0.327	7.51	
Shaker	2.76 b	0.252	9.12	Methanol
Sonic bath	3.91 a	0.194	4.97	
Homogenizer	2.23 c	0.079	3.55	
Soxhlet	3.68 a	0.106	2.89	
		TNB		
Shaker	0.366 a	0.047	12.8	Acetonitrile
Sonic bath	0.449 a	0.105	23.4	
Homogenizer	0.348 a	0.127	36.5	
Soxhlet	0.332 a	0.051	15.4	
Shaker	0.311 a,b	0.087	28.0	Methanol
Sonic bath	0.444 a	0.118	26.6	
Homogenizer	0.265 ъ	0.078	29.4	
Soxhlet	0.273 b	0.059	21.6	

^{*}Values indicated with the same letter are not significantly different at the 95% confidence level.

Table 4. Results of two-way analysis of variance for Iowa AAP soil no. 2.

			Mean	
	SS	df	square	F
		TNT		
Method	27.93	3	9.31	244.8*
Solvent	0.0096	3 1	0.0096	0.5
Interaction	2.029	3	0.6764	17.78
Error	1.521	40	0.0380	
LSD = 0.302				
		TNB		
Method	0.1635	3 1	0.0545	6.96
Solvent	0.0305		0.0306	3.90
Interaction	0.0095	3	0.0032	0.40
Error	0.3132	40	0.0078	
LSD = 0.137				

*Values are significant at the 95% confidence level. Critical values are F(3,40) = 2.84 and F(1,40) = 4.08.

Experimental precision for TNT averaged about 6% (RSD), indicating very good soil homogenization prior to subsampling. This excellent precision allowed a powerful comparison among methods using analysis of variance techniques, which indicated a significant difference among methods but no consistent difference between solvents. For acetonitrile, the Soxhlet was significantly better than the other three procedures; the sonic bath was second. For methanol, the sonic bath and the Soxhlet were not significantly different but were significantly higher than the shaker or homogenizer. There was a significant interaction between method and solvent, indicating that some methods worked better in one solvent while another method worked better in the other one.

For TNB, the relative standard deviation was somewhat larger than for TNT, averaging over 20%. This is a result of the very low levels of TNB present in this soil (about 0.4 $\mu g/g$), which approached the detection limit, estimated at about 0.1 $\mu g/g$. RSD values typically increase as levels approach the detection limit.

Analysis of variance for the TNB results indicated a significant difference existed among methods but not for solvent type or method-solvent interaction. This significant difference in methods was only apparent in methanol, where the sonic bath resulted in significantly higher results than the Soxhlet or homogenizer. The shaker was not significantly different from any of the other three methods at the 95% confidence level.

In addition to lower recovery for the Soxhlet procedure for TNB, it had a second disadvantage. This was the much higher extraction of other substances that tended to interfere with HPLC analysis. The major interferant appeared to be the aminodinitrotoluenes, which elute between TNT and DNT (the internal standard).

CONCLUSIONS AND RECOMMENDATIONS

The results of this study are complicated and no simple explanation adequately describes all of the findings. Clearly methanol and acetonitrile were equally good for extraction of TNT. For RDX and HMX, acetonitrile was superior both kinetically and thermodynamically. For TNB, acetonitrile appears to be better only at higher concentrations. Overall it appears that acetonitrile is the solvent of choice.

Of the four methods tested, the ultrasonic bath and Soxhlet typically result in higher analyte recovery. For the low-level soil (soil no. 2), samples were equilibrated in the ultrasonic bath for 24 hours and generally it resulted in analyte concentrations equivalent to the Soxhlet and better than the homogenizer or shaker. For the high level sample (soil no. 6), the ultrasonic bath was tested with only a 4-hour equilibration period. Even so, the recovery with acetonitrile was excellent for TNT and only slightly less than the Soxhlet for TNB. It probably would have proved equivalent to Soxhlet overall if a 24-hour equilibration had been used.

The sonic bath is also a practical choice since it can be used to process a number of samples simultaneously. It can also be used in an unattended fashion and is relatively inexpensive. Typically, these devices run about \$800. The gentle heating that occurs with the sonic bath (\approx 38°C) also seems desirable to increase the rate at which equilibrium is attained. The sonic bath does not suffer from the problem experienced with the Soxhlet device of co-extracting significant amounts of interfering substances that have a low solvent-soil partition coefficient.

The wrist-action shaker is desirable because of its simplicity, ability to process a large number of samples in an unattended manner, and low cost. While it appeared to work rather well for TNT for the high concentration soil, it provided very poor recovery for TNT in the low level sample. For TNB it was generally equivalent to the sonic bath.

The plant-soil homogenizer suffers from the restriction of processing one sample at a time. This limits its practical equilibration time as does significant solvent evaporation. Equilibrium is not established quickly enough and the homogenizer extracts were generally much lower in concentration than any of the other procedures. Because of these findings we do not feel this device deserves further consideration.

The Soxhlet extractor has long been the accepted method for extracting trace organics from soil matrices. It generally resulted in excellent recovery of TNT and TNB compared to the other techniques in this study. However, it requires a separate, fairly expensive setup for each sample and as mentioned earlier would be very expensive to use if large numbers of sample analyses were needed. For organics with very poor solvent-soil partition coefficients, the Soxhlet is particularly desirable. Explosive residues generally have very favorable partition coefficients and hence this advantage is not particularly useful. The disadvantage of extracting interferences with low partition coefficients was observed here.

Overall, we feel that the ultrasonic bath has the greatest potential for use in extracting explosive residues from soil and possibly sediment. Additional work with more soils, and with sediments, concentrating on the kinetics of extraction, would be useful before a final recommendation is possible. It would be very desirable to understand the relationship between the rate and extent of analyte extraction and its mode of adsorption-interaction with the soil.

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Time (min)	нмх	RDX	TNB	TNT	Extract
	S	oil-Plant	homogenize	er	
1	36.8	48.2	52.2	857	Acetonitrile
2	39.9	49.4	52.3	883	
4	37.4	51.0	54.3	887	
16	37.7	51.3	46.3	891	
1	14.1	42.3	49.4	910	Methanol
2	21.1	55.3	51.7	953	
4	25.1	59.8	54.1	981	
8	29.4	65.2	56.9	1002	
16	34.0	74.1	58.6	1041	
		Sonic	Bath		
1	114	40.1	55.8	860	Acetonitrile
2	111	40.8	56.8	854	
4	112	41.9	56.8	871	
8	111	41.7	55.9	861	
16	112	42.2	57.9	882	
32	111	42.7	57.4	876	
64	112	44.1	58.2	896	
240	115	48.9	67.4	952	
ī	13.9	35.6	50.3	885	Methanol
2	16.1	34.2	52.8	901	
4	18.0	35.0	53.6	912	
8	17.6	35.3	54.3	919	
16	23.2	36.9	54.9	923	
32	21.9	39.5	56.4	934	
64	25.7	40.4	57.6	943	
240	38.9	52.2	66.5	992	
		Wrist-act	ion shaker	•	
10	62.0	54.3	52.5	926	Acetonitrile
60	61.7	55.6	56.6	931	
90	62.5	56.0	56.7	931	
150	62.0	56.7	57.8	935	
240	62.3	57.0	56.0	952	
480	62.5	58.0	57.3	963	
1440	63.7	59.8	60.1	979	

Table Al (cont'd.)

Time (hr)	HMX	RDX	TNB	TNT	Extract
10	10.9	24.3	50.8	854	Methanol
30	17.1	29.6	53.0	878	
60	18.5	30.1	54.9	894	
90	18.8	30.7	55.1	894	
150	20.6	32.3	55.5	906	
240	19.8	32.8	56.0	910	
480	21.2	34.9	58.7	933	
1440	22.2	33.4	62.4	955	
		Soxh	let		
1	69.5	45.5	50.9	746	Acetonitrile
1 2 4	83.5	55.4	61.2	877	
4	85.7	57.1	62.0	882	
25	88.2	60.1	62.5	902	
37	90.0	59.2	59.1	903	
1	10.3	13.3	29.5	537	Methano1
1 2	16.1	20.9	47.3	799	
4	20.3	30.0	56.4	899	
25	47.0	40.8	56.4	890	
37	51.1	40.4	55.1	860	

Table A2. Replicate study of Iowa AAP soil no. 6 ($\mu g/g$).

Wrist-action shaker	Sonic bath	Soil-plant homogenizer	Soxhlet extractor	Extract
	*			
		TNT		
900.7	882.2	868.5	890.7	Acetonitrile
884.1	880.8	846.4	874.4	
867.6	896.7	859.5	874.9	
872.0	891.6	832.6	876.8	
871.5	872.2	844.8	890.5	
888.4	873.9	841.3	879.6	
895.5	811.9	857.1	892.2	Methanol
882.7	828.8	863.0	893.7	
895.2	854.2	878.5	894.5	
913.9	815.1	849.4	896.9	
876.8	867.9	942.4	887.9	
908.5	864.7	831.0	882.9	

Table A2 (Cont'd.)

Wrist-action shaker	Sonic bath	Soil-plant homogenizer	Soxhlet extractor	Extract
		TNB		
55.7	56.1	50.7	63.6	Acetonitrile
55.4	55.8	51.5	61.8	
53.8	55.3	52.1	61.6	
54.2	55.9	50.3	63.1	
54.7	55.7	51.2	62.0	
56.6	54.5	52.3	60.4	
56.5	50.5	53.0	59.3	Methanol
55.4	52.4	52.0	58.6	
56.3	52.7	53.6	58.3	
57.0	51.1	53.9	57.8	
55.0	54.4	55.1	57.9	
55.5	53.9	52.9	57.6	
		RDX		
47.4	47.4	45.1	66.8	Acetonitrile
49.0	69.1	44.3	68.6	
58.6	48.4	37.5	83.0	
46.0	42.3	33.4	62.7	
52.5	46.1	177.8	53.1	
71.7	73.4	48.4	56.4	
			74.7	
40.7	31.8	33.3	48.1	Methanol
30.1	38.6	31.3	57.2	
30.6	37.7	56.9	44.7	
33.5	54.9	34.7	48.2	
27.5	39.4	30.9	41.5	
59.1	36.8	28.4	48.3	
		HMX		
		при		
71.9	74.8	92.7	85.5	Acetonitrile
131.0	55.3	104.9	68.3	***************************************
88.8	73.2	50.4	72.5	
52.1	41.9	53.2	87.1	
101.6	46.5	44.6	96.2	
46.4	46.4	43.8	94.8	
19.7	39.4	25.9	54.0	Methano1
21.4	39.4	19.3	67.6	
14.4	48.3	19.3	64.3	
22.8	28.0	27.1	57.2	
24.8	24.3	63.9	54.3 :	
30.6	21.1	14.6	55.3	

Table A3. Kinetic study of TNT extraction, Iowa AAP soil no. 2 ($\mu g/g$).

Time (min)	Time (min) Acetonitrile		
	Soil-plant homogenizer		
5	2.05	2.03	
10	2.57	2.12	
20	2.87	2.53	
60	3.42	3.16	
	Sonic bath		
15	2.25	2.51	
30	2.43	2.91	
60	2.87	2.99	
120	3.30	3.50	
240	3.15	3.89	
420	4.54	5.43	
	Wrist-action shaker		
30	1.72	1.87	
60	1.94	2.28	
120	2.22	2.92	
240	2.25	2.98	
1440	3.62	3.60	
2880	4.01	4.42	
	Soxhlet		
1	2.39	1.82	
1 2	3.49	2.68	
4	3.93	2.86	
24	4.31	4.31	
48	4.85	4.55	

Table A4. Replicate study of Iowa AAP soil no. 2 ($\mu g/g$).

Wrist-action shaker	Sonic bath	Soil-plant homogenizer	Soxhlet	Extract
	DOMEC DUCK	nomogenizer	DOMITEC	
		TNT		
		in the		
2.50	3.75	2.07	3.76	Acetonitrile
2.44	3.29	2.07	4.36	
2.36	3.32	2.02	4.30	
2.47	3.69	2.03	4.39	
2.68	3.41	2.31	4.72	
2.33	3.77	2.11	4.55	
2.86	3.80	2.15	3.66	Methanol
2.92	3.76	2.29	3.63	
2.44	3.78	2.21	3.69	
2.67	4.28	2.26	3.64	
2.54	3.92	2.33	3.88	
3.11	3.91	2.13	3.57	
		TNB		
0.417	0.331	0.321	0.380	Acetonitrile
0.285	0.556	0.294	0.263	
0.369	0.432	0.271	0.345	
0.401	0.596	0.343	0.322	
0.380	0.378	0.598	0.290	
0.342	0.403	0.260	0.392	
2204,047	200		22.000	
0.231	0.330	0.238	0.270	Methanol
0.325	0.350	0.226	0.301	
0.244	0.379	0.410	0.337	
0.250	0.541	0.201	0.246	
0.456	0.628	0.294	0.312	
0.360	0.436	0.221	0.172	

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