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The Role of Internal Standards and their Interaction with Soils Impact Accuracy of Volatile Organics Determinations

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Abstract: Both US Environmental Protection Agency (EPA) SW-846 Methods 8260C/5035 and 8261A include mixing soil with water and addition of internal standards prior to analyses but the equilibration of internal standards with the soil is not required. With increasing total organic carbon (TOC) and no effort to equilibrate internal standards with the matrix, results are less likely to be accurate. Adding internal standards to soils prior to diluting the sample with water gives more accurate determinations but less reliable quality control (QC). Extending times for equilibration of internal standards improves accuracy but is conducive to analyte degradation not normally observed during analyses. Soil-matrix effects on a given analyte can be greatly understated using a single internal standard as described in Method 8260C while the use of multiple internal standards as described in Method 8261A is more accurate. Method 8261A's reporting error when spiking soils before adding water provides confidence intervals with accuracy near the experimental rule (75.2, 95.7, and 99.5%) with the exception of two analytes that require overnight equilibration.

Introduction

The level of uncertainty in environmental analyses is of concern for those who use analytical data to make environmental decisions [1, 2]. Soil is a complex medium that can influence the behavior of volatile organic analytes (VOAs) as general matrix effects (such as organic phase uptake), biological activity and even varying kinetics (fast and slow) adding to the difficulty in assessing the viability of VOA determinations [3, 4]. Soil has been difficult to prepare as a reference material for VOAs. Researchers have used vapor addition of analytes to dry soils for developing performance soils [5]. It has also been found that volatile analytes react with the reference soil matrix after their addition [6]. Just how the soil matrix might impact the accuracy of volatile organic determinations is poorly understood.

The difficulties encountered in the preparation of soil reference materials also describe challenges for analysts in preparing soils for analyses and may differ depending on method [7]. Internal standards are commonly used in gas chromatography/mass spectrometry (GC/MS) analyses and they are typically added to sample extracts prior to injections to normalize variations between injections. For determination of VOAs, however, internal standards are added to the sample prior to a concentration step (such as purge-and-trap concentration) exposing the internal standards to matrix effects. US Environmental Protection Agency's (EPA's) method for determining VOAs (SW-846 Method 8260C) assumes the chemical properties of analytes and their internal standards are identical and any relative change in

response of an internal standard would be the same for analytes [8]. This assumption will introduce inaccurate results when the behavior of an analyte and its internal standard are sufficiently different during the concentration from a matrix.

The equilibration of internal standards with soils is not required in EPA's methods for soil analyses. However, when analytes are at equilibrium with a soil and internal standards are not, the internal standards may be more effectively recovered than analytes. Using recovery of the internal standard for the recovery of an analyte ignores that portion of analyte bound with the matrix resulting in an understated analyte concentration. The impact of allowing equilibration time and different approaches to adding internal standards are evaluated in this work.

Method 5032 describes vacuum distillation as the means to separate volatile compounds from the soil [9] and is used as an optional concentration step for Method 8260C. Vacuum distillation had been found to be more efficient in extracting volatile analytes from soil than headspace, ambient purge-and-trap, and heated purge-and-trap techniques [10]. Therefore observations in this work as to how compounds are affected by the soil matrix are also relevant to headspace and purgeand-trap analyses and may address problems observed using non-vapor phase (methanol) extractions [7].

Method 8261A incorporates vacuum distillation and differs from Method 8260C/5032 in that Method 8261A incorporates a battery of internal standards to interpret an analyte's response and while Method 8260C specifies multiple internal standards are to be used, each analyte is related only to its assigned internal standard. Where the variation in response of a single internal

standard translates to the same variation in its associated analytes, the battery used in Method 8261A interprets the analyte response as a function of its boiling point and relative volatility [11]. It would be expected that parsing matrix effects by chemical properties would generate more accurate results. This study processes raw experimental data by the two internal standard approaches and compares the relative accuracy of their determinations.

Method 8261A generates an error term with each analyte determination and these have been shown to be accurate in describing the analytical error for a variety of water matrices [12]. The reporting error generated as described in Method 8261A was therefore a primary tool in evaluating this study's experimental results [9]. While this study assesses the reporting errors as an analytical error it also evaluates which quality control parameters need be implemented to improve accuracy. Multiple replicate analyses provided the data for calculating the frequency that confidence intervals included the amount of analyte added to the samples. Because each analyte data set was small, a less restrictive Chebyshev's inequality was used in place of the empirical rule to identify analytes where reporting errors where inconsistent with experimental errors [13]. The analytes were also evaluated for stability in the soil samples. The matrices studied include three soils that had been used in performance studies [6], acid-washed sand, and Greenwich Bay (MA) sediment.

Experimental

GC/MS: The vacuum distiller is interfaced to a GC/MS so that the vacuum distillate is transferred directly to the GC/MS for analysis after a distillation. In this study, the GC/MS was a

Thermo DSQ mass spectrometer and Trace GC. The GC capillary column was a 30 m x 0.25 mm i.d., 1.5 μm film VOCOL (Supelco, Bellefonte, PA). The GC operating conditions were 2.5 min at -20 C, 40 C/min ramp to 60 C, 5 C/min ramp to 120 C and held at 120 C for 1 min, 20 C/min ramp to 220 C and held for 12 min resulting in a GC run time of 34 min. The injection was split 60:1 with a constant flow rate of 1.4 ml/min. The mass spectrometer scanned between 35 and 300 amu at 1 scan/sec.

Vacuum Distiller: A Cincinnati Analytical Instruments Model VDC1012 vacuum distiller (Indianapolis, IN) performed the distillations in the study. Samples were vacuum distilled for 7.5 min with a 2.5 min transfer to the GC/MS through a transfer line held at 200 C.

Quantitation: Calibration was performed as described in Method 8261A. The internal standards are listed in Table 1. Vinyl chloride- d_3 was added as a surrogate for gases [12]. The higher boiling point compounds, 1,2,3-trichlorobenzene-*d3*, 3,5-di-tert-butyl_toluene, 3,5 dibromotoluene, azulene, a,a-dichloro-o-xylene were added as part of the internal standard/surrogate suite to more fully describe the boiling point effects at the upper range of the method. The software used to perform calibration and quantitation was SMCReporter 4.2 available from EPA's web pages [14].

The surrogates used to monitor method performance are presented in Table 2. These analytes were monitored as representative of three classes of compounds: volatile class was for compounds with boiling point less than 159 ºC, non-purgeable class as the volatile class but with relative volatility greater than 100, and the semi-volatile class representing compounds that boil

at or above 159 ºC. A sub-class of *gases* was added to the volatile class and related to vinyl chloride-*d3*.

The calibration range of analytes (listed in Supplemental Information Table S3) in this study was nominally 5 to 500 ng per analyte. A review of calibration ranges was conducted to ensure the range was linear for each analyte. The lower points of some analytes were not used in generating their calibration curves when interferences at the lower point were observed. The lowest standard mass in each analyte calibration curve was used as the limit of quantitation (LOQ). Any analyte response that fell below the LOQ was considered as potentially less accurate and segregated from results that fell within the calibration range. The amount of analyte added to each sample is related to the calibration range and was at three levels, LOQ, mid-level (1/10 upper calibration amount) and high (1/2 upper calibration amount).

Samples were analyzed in the same manner as was previously conducted for studying the accuracy of reporting errors for water analyses [12]. That is a multi-point calibration was run prior to analyses, and then a check standard, blank, unspiked soil, and 5, 10, and 15 g aliquots of soil spiked at the three levels (high, mid-point, and LOQ) were analyzed daily and replicated six times. This structure of analyses did not allow for interpretation until all analyses of a matrix were completed. However, when there were additional tests to clarify results, these were conducted using a two-point calibration (before samples and after samples) and samples were spiked with the calibration amounts. The additional tests used the high spike amount (1/2 upper point in Table 3 calibration range) and only 5 g of sample to minimize impact of background analyte concentrations.

Samples: Samples were prepared for analyses in the vacuum distiller sample vessel. First the necessary amount of soil was added to the vessels and the vessels closed by connecting to the vacuum distiller. Addition of a 5-uL methanol solution containing the internals standards and surrogates to the dry samples was next. This was done by touching the glass surface under the sample with the syringe before releasing the solution. The analytes were then added to the sample in the same manner. Finally, water was added to soil. The mixing of the soil, spikes, and water varied by sample type and discussed later.

Five different soil samples were evaluated. Three sample amounts (5, 10, and 15 g) of each soil were analyzed. In addition, three different levels of analyte concentration were analyzed at each sample size. The amounts of the spikes were 0.5 and 0.2 of the upper limit of the calibration range and at the limit of quantitation (0.02 of upper limit).

Acid-washed sand, three soils (NV, GA, OR, previously used to create performance sample [6] and a sediment (SD) were used for samples.

Results and Discussion

The results are addressed in three sections and follow the sequence the study was performed. The first section discusses evaluating the Method 8261A analyses of soil and identifying any QC requirements that need to be implemented to improve accuracy. The second section addresses additional analyses that were necessary to clarify analyte behavior or other observations. The last section compares experimental results using Method 8261A (multiple internal standards per

analyte) and Method 8260C (single internal standard per analyte) quantitation procedures in order to evaluate the impact of their differing internal standard roles.

Method 8261 Accuracy

Method 8261A includes reporting an analytical error with each analyte result. The study began with evaluating if the reporting errors were consistent with random errors of measurements. For random error measurements, the frequency that one, two, and three confidence intervals include a true value follows the experimental rule (68, 95 and 99.7%) and systematic errors would cause the average result to be biased. To evaluate consistency of Method 8261A errors with being random, the errors were analyzed by analyte, matrix type and amount, and concentration. Initially only the high and mid-level spikes were investigated and those samples spike at the limit of quantitation (LOQ) were evaluated separately. This resulted in evaluating 12 results when combining the high and mid-level results or only 6 when looking at a single spike level. Therefore, due to the small number of results, the frequency that the reporting-error confidence interval includes the true value was compared to Chebyshev's rule [13]. Analytes that did not meet Chebyshev's minimum frequency values for both two and three standard deviation in a matrix were considered outliers.

On occasion, analytes would pass Chebyshev's rule criteria but their average concentration for a matrix was significantly different from the true value (systematic error). In this study, if an average result varied more than 25% from the true value, it was considered biased.

Method 8261A reporting errors were assessed after eliminating those results compromised by background or calibration defects. Analyte results were not used if that day's continuing calibration check difference exceeded 40%. Analytes were not used for a matrix when its background content in the matrix exceeded 15% of the mid-level spike amount for 15 g aliquots. Based upon a previous investigation of water samples, a minimum relative confidence interval of 6% was applied for all analyte reporting errors [12]. Determinations that the Method 8261 reports generating software, SMCReporter [14], qualified as above calibration or below calibration range were not included in evaluations.

It was an initial assumption that if an analyte was an outlier, there would be an observable cause that would also impact one of the QC parameters. Data were reviewed by matrix so that any severe effects related to a matrix would be detected. All of the determinations of an analyte for a matrix that failed Chebyshev's rule were examined to identify those analyses where the true value fell outside the three standard deviation confidence interval. The next step was to determine if these results were related to a matrix effect on a class (volatile, non-purgeable, or semivolatile compounds), or a matrix effect on subset of a class, concentration, or sample size.

Outliers that were identified were not uniquely linked to a quality control measurement. Typically the outlier condition could only be eliminated by requiring a stringent QC range that also disqualified the majority of analyses regardless of sample size or level of spike. It became evident that the mixing of soil and spikes, equilibration times, and analyte degradation were the major factors causing the observed outliers and these effects went undetected by surrogate or internal standard behavior.

Ensuring internal standards, surrogates, or matrix spikes are in equilibrium with a soil matrix is not prescribed in Methods 8260C or 8261A. In this study however, attempts were made to add internal standards, surrogates, and analytes to soil prior to adding water. Initially it was thought that the spiking of soil (before adding water) allowed sorption by the matrix to be complete and the vigorous boiling that takes place during vacuum distillation would distribute the spikes throughout the sample. However, distributing the spikes throughout the sample and uptake of the analytes by the soil both proved problematic. It was evident that there needed to be assurance internal standards, surrogates, and analytes, had been distributed throughout the soil. When spikes were not distributed throughout the soil, results were inconsistent and analytes and their labeled analogs appeared to behave differently.

One way to detect poorly distributed spikes was through a comparison of recoveries of labeled internal standards and their natural analogs for each analysis. It was expected that if the recoveries of an analyte and its analog were grossly different it would be due to insufficient mixing of the soil after spiking. The relative responses (sample response per mass unit/standard response per mass unit) would be identical if they were well distributed throughout the sample. The compound pairs that were used for identifying insufficient distribution of spikes throughout a sample were vinyl chloride (for gases), bromobenzene (VOAs), 1,2-dichlorobenzene and 1,2,3 trichlorobenzene (semi-VOAs) and their deuterium analogs allowing maximum variation limits of 75,50, 25, and 25% respectively. When a maximum variation limit was exceeded for an analysis, those results for analytes in the respective class of compounds were not used in assessing accuracy.

The procedure for distributing the spikes throughout a sample became more rigorous with each matrix examined. The frequency of exceeding maximum variation limits decreased as mixing (after spiking soil before adding water) went from none (sand), manual shaking (NV), to mechanical mixing (GA and OR). There is not a QC measure in either Method 8260C or 8261A that indicates the adding and mixing of internal standards is complete. One way to assess mixing would be to add a compound such as 1,3,5-trichlorobenzene as a surrogate and its labeled analog as an additional internal standard. Monitoring the recovery of both to ensure equivalence would identify when mixing was inadequate as long as the mixes were being added to samples separately and at different locations. However, in following studies, the use of a mechanical stirrer to mix the samples after being attached to the vacuum distiller demonstrated the maximum variation limits used in this study were easily met without losing the gas compounds.

The relative response of internal standards were monitored (the response of an internal standard in a sample divided by its response in the day's continuing calibration standard). A subset of Method 8261 internal standards was monitored for consistency as performed in Superfund's Contract Laboratory Program (CLP) methods [15]. The CLP limits on acceptable variation in the internal standards (as %) are presented in Table 1. If an internal standard relative response (high or low) corresponds with the occurrences of outlier analytes, it would be assumed to affect its class of compounds. The only relative responses of internal standards that were found to be related to outlier data was when a sample vessel seal leaked during a distillation impacting the non-purgeable class and gases.

In general, the relative response of the internal standards closely related to the total organic content in the sample (Figure 1). For low-organic content soil samples, relative responses are greater than those for the day's continuing calibration standard. This is likely due to the vigorous boiling of water mixed with soil during vacuum distillation. As organic content increases, the internal standards responses decrease with the more lipophilic compounds decreasing the most.

The surrogate compounds, their recommended recovery ranges from Method 8261A and the range of experimental recoveries are listed in Table 2. The only surrogate that was linked to an outlier analyte was vinyl chloride- d_3 . The surrogate ethyl acetate- ^{13}C was observed to degrade quickly in some soils and this tendency was previously reported [11]. The sample that was not properly sealed for analysis (NV soil) had several surrogate recoveries outside their normal range (Table 2).

Three new compounds were evaluated for use as semivolatile surrogates. Recovery of azulene was nil for most analyses other than sand when applied directly to dry soil. The recovery of the other compounds (3,5-di-tert-butyl_toluene and a,a-dichloro-o-xylene) were at times much greater than 100% yet their extreme recoveries could not be linked to an outlier.

Two analytes were found to be outliers in one of the matrices but were not outliers when samples were analyzed a day after spiking. Hexachlorobutadiene was bound tightly when spiked directly on sand with a little over 50% recovered if the analysis was within 3 hr of spiking. With an overnight equilibration time (or spiked after water added to sand) the compound recovery was

 $102 \pm 8\%$ (Table S4 in Supplemental Information). Dibromomethane behaved similarly with the OR soil matrix (Table S7 in Supplemental Information).

There was difficulty mixing the GA samples with the initial practice of mixing the samples external from the vacuum distiller resulted in losing the most volatile compounds. This loss was remedied with the improved mixing procedure (mechanical stirring of sample while container attached to vacuum distiller) and the early results for the gases were not included in accuracy determinations (Table S6 in Supplemental Information). With the exception of mentioned compounds, the results were near experimental error expectations when spiking the soils before adding water (Table 3) suggesting the reporting error for most analytes is random. Therefore the confidence intervals reported with Method 8261A results are a good measure of the analytical error for spiked soils and sediments even when there are efforts to equilibrate spikes with the sample.

The same conditions were applied to the analyses of the LOQ-spiked samples. An additional criterion was that a result was not used when the analyte was also found to be in blanks at $> 1/2$ the concentration of the LOQ spikes. The frequency these confidence intervals included the true values did not match theoretical prediction as closely as the mid-concentration spikes. This is likely due less accurate integrations or background contributions for measuring responses at the lowest calibration point. It is interesting to note that by raising the minimum one standard deviation confidence intervals to 15% as was done in the previous study [12], the confidence intervals included the true values near empirical rule frequencies (Table 3).

Additional Tests to Clarify Observations

Verification of degradation, evaluation of spiking soils after addition of water and evaluation of impacts for extending equilibration times were performed at the high spike level (1/2 upper point of calibration range Table 3) with 5 g sample amounts. Two to six replicates were analyzed and calibration was performed at the same amount as the sample spiking. Using the high spike and a smaller sample size minimized impact from background.

Degradation of analytes was determined by analyzing samples spiked with analytes, surrogates, and internal standards and held at room temperature for extended times. Many analytes were found to degrade in a matrix over a 120-hour period and these analytes were not used in the evaluation for a given matrix (Supplemental Information Table S3).

There was also an assessment of how standard practice of adding spikes with or after adding water to soil matrix impacted results. These samples were analyzed the same day. The recovery of the non-polar internal standards was generally greater than when the internal standards were added to the soils before adding water (Table 1). There was not a lot of difference in the range of surrogate recoveries for when adding compounds to dry soil and adding them after the water was added (Table 2). The analyte results by matrix (Supplemental Information Tables S4-8) did not indicate that the "wet spike" analyses were less accurate. Of course, accurate analyte results may only occur when analytes are present in a soil sample through wet spiking, and not already

present in the soil. Understating of analyte concentrations should be lessened if the internal standards and surrogates were at equilibrium with the sample matrix.

Overnight equilibration (averaging near 20 hrs) of the soil/water/spike slurry yielded results similar to when spikes were added to soils before adding water, but some degradation of analytes became evident. Extending the equilibration to 120 hours demonstrated even more sorption of the internal standards indicating the equilibration of spikes and matrices could be a slow process for the higher boiling compounds like 1,2-dichlorobenzene (Figure 2). In general, the sorption of volatile compounds by soil through a quick process and a slow process is similar to fast and slow sorption rates observed in the environment [4, 16]. Relying on only a quick equilibration (*e.g.*, analyses on day of spiking) for internal standards understates the concentrations of analytes that reach equilibrium through the slower sorption processes (and at equilibrium with the matrix).

Figure 2 illustrates the sorption of the internal standard, 1,2-dichlorobenzene-*d4*, in the different matrices from the wet spike with increasing equilibration times. For the OR matrix, the response of 1,2-dichlorobenzene-*d4* drops 66% over the first 3 hr, drops another 6% overnight, and another 19% to the 120 hr endpoint where its response is 9%. If the 120-hr endpoint reflects how analytes would be taken up by soil through exposure at the site where the soil was collected, then there is a potential to dramatically overstate the recovery of analytes and therefore understate their true concentration in a soil. For instance, if 1,2-dichlorobenzene was present in the OR soil from a site exposure, same-day spiking with internal standards and analysis would only reflect 27% of the true concentration. Figure 2 shows only 9% of the compound would be released during analysis (at equilibrium) but the spiking of internal standards on the day of

analysis indicate 34% of the compound was recovered $(9/34 = 27%)$. Overnight equilibration improves the determination (33% of true concentration) but spiking soil before adding water was a better approach (47% of true concentration from Supplemental Information Table S9).

The range of relative responses (recovery) for internal standards found experimentally (Table 1) has application to other concentration methods that are less efficient [10]. That is, the more effective the interaction of internal standards with soils (with elevated organic content), the more likely an analysis will fail the lower limits of internal standard relative response. If Method 8260C with purge and trap concentration (Method 5035) were equivalent to vacuum distillation concentration, then adding internal standards to the NV soil prior to dilution with water would cause the analyses to consistently fail criteria for Superfund's Contract Laboratory Program [15]. However, by adding internal standards to soil after adding the water and then analyzing quickly to minimize equilibration time an analysis is more likely to pass criteria. This produces a dilemma where enforcing a QC parameter encourages a practice of under reporting analyte concentrations. To address this issue, it is recommended that the time between adding internal standards to soil and analysis is reported along with the whether the internal standards were added to soil after or prior to adding water.

The sediment matrix was not included as a soil in summarized data. Even though surrogate recoveries for sediment samples consistently fell within the soil surrogate ranges recommended in Method 8261A for soils, there were still numerous outlier analytes (Table 3) as well as biased results (Supplemental Information Table S3). Taking a smaller amount of sediment (1.0-1.7 g) for analyses improved results (Table 3) and surrogate recoveries met the surrogate recovery

ranges found for water in a previous study [12]. Selecting a smaller sample size also generated less biased analytes (3 biased analytes) than for 5 g samples (12 bias analytes). The analyses of a smaller sample size yielded reporting errors that were much more consistent with random measurement errors than 5-15 g soil samples (Table 3). If taking a smaller sample size does not jeopardize meeting sensitivity requirements or does not result in a nonrepresentative sample, analyzing a smaller sample size so that surrogates meet recovery limits found previously for water would be preferred.

As the level of TOC increases, so does the potential for understating results when internal standards and the sample are not equilibrated. Because both fast and slow sorption of internal standards appear to be closely related to TOC, the knowledge of organic content in soil can be useful in interpreting data (high organic content and high recovery of internal standards likely indicate spikes are not equilibrated with matrix and therefore results likely understated).

Method 8260C Compared to Method 8261A

Method 8260C with a vacuum distillation concentration step (Method 5032) differs from Method 8261A (method specifically incorporates vacuum distillation as a concentration option) in the role of internal standards to quantify an analyte. The battery of internal standards used in Method 8261 is used to parse matrix effects into relationships relating to boiling point and relative volatility and the analyte is quantified as a function of its boiling point and relative volatility. Method 8260C uses a single internal standard for quantitation of an analyte and

assumes each analyte behaves the same as its assigned internal standard. If an analyte recovery from a matrix does not behave as the internal standard, there can be a significant bias in determinations.

The same analyses (six samples of each study matrix) used to document optimized spiking and mixing conditions were used to compare the methods. The raw analyte responses of theses analyses (originally quantified using Method 8261A) were re-quantified using Method 8260C. The analytical conditions would not ensure equilibration of spikes with matrices but would be more representative of current analytical laboratory spiking practices. It would be expected that the more thorough the spike equilibration, the more Method 8261A results would be superior to those of Method 8260C.

The occurrence of analyte bias was the means to compare the methods. The power of using the battery of internal standards per analyte in Method 8261A over the single internal standard per analyte used by Method 8260C is demonstrated by the number of analytes that had average results that differed by more than 25% from true values (Supplemental Information Table S3). Method 8261A had six analytes differing by more than 25% from true values (7% of analytes) while Method 8260C had 20 analytes (24% of analytes). The average recovery of all Method 8261A results was nearer ideal at $100 \pm 20\%$ compared to $112 \pm 37\%$ for Method 8260C. Also the range of recoveries was generally greater for the method 8260C surrogates compared to Method 8261A (Table 2). The Method 8261A internal standard corrections were superior to the Method 8260C internal standard use resulting in a 70% decrease of the occurrence of results differing more than 25% from true values.

More specific information is available as a supplement to this article. Contained in the supplement are each analyte's results by matrix, average recovery and frequency that each analyte's confidence interval included the true value, and identification of those analytes impacted by continuing calibration limits and minimum confidence intervals.

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Notice

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Table 1. Experimental Relative Responses of Internal Standards (by Matrix and spike technique)

 $\frac{a}{b}$ Range of recoveries expected for internal standards from reference 15.

^b Summary for soils not including sediment.

 \textdegree Internal standards added to soil before diluting with water. Includes sample sizes 5, 10 and 15g.

^d Internal standards added after diluting soil with water. Samples were 5g

^e This internal standard not monitored in CLP protocols.

^f Values in parenthesis were for sample with low response due to bad seal and indicate an unacceptable recovery

Table 2. Surrogate Range of Acceptable Recoveries Found Experimentally by Matrix

^a Ranges posted with Method 8261A for soil [9]

^b Surrogate windows narrowed to value in parenthesis to eliminate Chebyshev outliers

^c Results in brackets were those for a sample that was not well sealed during distillation

^d These surrogates and their ranges apply to the semivolatile analytes. Narrower limits would apply if they were used for just their analogs.

Table 3. Summary of Accuracy of Reporting Error Confidence Intervals for Determinations by Matrix

^a The sum of all experimental determinations in the study by matrix after removing analytes with more than 5% of medium spike present in matrix before spikes.

^b Number of determinations that meet criteria.

 \degree The % frequency that a result and confidence interval include the known value at 1, 2, and 3 standard deviations.

^d The average recovery of all analytes that meet criteria and one standard deviation.

e Compounds not included with results.

Figure 1. Relative Response of Internal Standards vs Total Organic Content.

Figure 2. Relative Response of 1,2-Dichlorobenzene-*d4* vs. Equilibration Time