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Interlaboratory Analytical Comparison Study to Support Deepwater Horizon Natural Resource Damage Assessment: Description and Results for Mussel Tissue QA10TIS01

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Interlaboratory Analytical Comparison Study to Support
Deepwater Horizon Natural Resource Damage
Assessment: Description and Results for Mussel Tissue
QA10TIS01

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ABSTRACT

To support natural resource damage assessment (NRDA) in response to the Deepwater Horizon (DWH) oil spill in the Gulf of Mexico, a large number of coastal sediment and tissue (i.e., oysters) samples have been collected outside of the spill zone to define baseline environmental conditions prior to being exposed to oil. Analysis of oiled sediments and oil-exposed oysters will continue for the foreseeable future. To support these efforts, NOAA will require additional analytical laboratories to perform NRDA sample analyses in addition to the commercial laboratories currently providing analyses in support of NRDA. To compare the data among these laboratories, inter-laboratory comparison studies have been initiated with the results from the third exercise, mussel tissue QA10TIS01 reported here. In this exercise, selected polycyclic aromatic hydrocarbons (PAHs), alkylated PAHs, hopanes, and steranes were determined in the exercise material, which consisted of a cryogenically homogenized mussel tissue sample, and in SRM 1974b Organics in Mussel Tissue (*Mytilus edulis*). The results from this third exercise are reported along with a summary of the analytical methods used.

INTRODUCTION

On April 20, 2010, a fatal explosion, fire, and sinking of BP's Deepwater Horizon drilling rig occurred approximately 40 miles off the Louisiana coast. The disaster resulted in the discharge of tens of thousands of barrels of oil per day from the seafloor into the Gulf of Mexico. In what has become the worst offshore oil spill in U.S. history, a wide expanse and variety of natural resources have become exposed and potentially impacted by oil and other consequences of the spill. Under the Oil Pollution Act, those responsible for an oil spill are liable for clean-up and for natural resource damages. Several federal and state agencies are conducting a natural resource damage assessment (NRDA) to determine what resources have been injured and what uses of the resources have been lost due to the spill.

To support this NRDA, the trustees and BP's representatives have been collecting and analyzing tens of thousands of environmental samples to characterize both pre-spill and post-spill environmental conditions. A broad range of sample types have been collected including oil in various forms, water, sediment, and biota. For the foreseeable future, subsequent sampling and analysis will be required. In addition, numerous other entities have collected environmental samples for hydrocarbon analysis and submitted them to different laboratories throughout the country.

In the past the National Institute of Standards and Technology (NIST) has helped benchmark and improve the quality of analytical data gathered on the marine environment by administering interlaboratory comparison exercises. To compare the data among the many laboratories analyzing samples from this spill, the National Oceanic and Atmospheric Administration (NOAA) has requested that NIST coordinate interlaboratory comparison studies with sediment, crude oil, and bivalve tissue being the three matrices of interest. These studies are performance-based with each laboratory using its current methods for analysis of similar matrices that it would use for its program customers. The target analytes for each study are selected polycyclic aromatic hydrocarbons (PAHs), alkylated PAHs, hopanes, and steranes. More than three dozen laboratory facilities were contacted by NOAA and invited to participate in the studies; for all three matrix studies a large number of laboratories agreed to receive samples and report their analytical results to NIST.

The data received from 34 laboratories for the mussel tissue QA10TIS01 are summarized in this report along with summaries of the analytical methods used by each laboratory. Numerical indices, z- and p-

scores, are used to assess and track laboratory performances for accuracy and precision, respectively, and to provide a mechanism for assessing the comparability of data produced by the participating laboratories for the target analytes.

SOURCE OF MATERIAL

A previous tissue control material (NIST/NOAA NS&T QA Program Tissue Control Material III QC90TC Prepared Summer 90) was relabeled for use as the unknown material in this exercise. The new labels read:

Intercomparison Exercise to Support DWH NRDA
Sample: QA10TIS01 Mussel Tissue
Bottle # ---
Prepared Nov 2010

The mussels used for preparation of QC90TC were collected and stored at one of the NOAA NS&T laboratories prior to being shucked and shipped to NIST. QC90TC was prepared by NIST in 1990 as a frozen bivalve tissue homogenate still containing its endogenous water. The material was used by the NS&T laboratories in an interlaboratory study in 1991 [3].

SAMPLE DISTRIBUTION

Three bottles of QA10TIS01 Mussel Tissue each containing approximately 15 g of frozen mussel tissue were distributed to each of 45 laboratories during the week of February 21, 2011. Each laboratory was requested to analyze three samples of QA10TIS01 mussel tissue and at least one or more samples of SRM 1974b Organics in Mussel Tissue (*Mytilus edulis*) [4] (not provided to the laboratories) with their laboratory's and/or program's current analytical protocols being used for the determination of the concentrations (mass fractions) of the parent polycyclic aromatic hydrocarbon (PAH) compounds, alkylated PAH compounds, hopanes, and steranes.

The instructions including the list of target analytes sent to participants are attached in Appendix A.

EVALUATION OF EXERCISE RESULTS

Establishment of the Assigned Values

Laboratory data submission: Each participating laboratory was asked to submit data from three replicate determinations of the “unknown” material QA10TIS01 and was requested to report results of concurrent analyses of NIST SRM 1974b Organics in Mussel Tissue (*Mytilus edulis*). Laboratories were requested to report these results to three significant figures and to provide brief descriptions of their cleanup and analytical procedures.

Determination of laboratory analyte means: For each laboratory, the laboratory analyte mean of the three sample results (S1, S2, and S3) was calculated for each analyte. Non-numerical data were treated

as follows: A mean "<value" was used when three "<values" were reported; NA (not analyzed/determined) was used for three reported NAs; and, if the reported results were of mixed type, e.g., S1 and S2 were numerical values and S3 was reported as "<value", the two similar "types" were used to either determine the mean or to set a non-numerical descriptor.

Determination of assigned values: The assigned values are the means of the acceptable data as defined here. For a particular analyte, the performance on the reference material, SRM 1974b, was initially deemed acceptable for the purpose of this exercise if the laboratory result was within 30 % of the upper and lower limits of the confidence interval for analytes listed in the Certificate of Analysis for SRM 974b [4]. The criterion of 30 % is the same as the one that was used for the National Oceanic and Atmospheric Administration (NOAA) Mussel Watch and National Status and Trends Quality Assurance Programs [5]. If a laboratory demonstrated acceptable performance on a particular analyte in the reference material, that laboratory's results for that analyte in the corresponding "unknown" exercise material was then used in the calculation of the analyte's exercise assigned value, unless it was deemed an outlier. For evaluation of potential outliers, statistical tests and expert analyst judgment were used after viewing both normal and log-normal plots of the data. This judgment utilized knowledge of potential outliers based on the laboratory's reported methods. In instances for which the analyte concentration was below the detection limit of most participating laboratories, no exercise assigned value was calculated. In data sets where a number of laboratories report results as "not detected" at various detection limits, there is no consensus as to what numerical value should be assigned to these results in the computation of consensus means and other values.

REPORTED RESULTS

Laboratories were assigned numerical identification codes in order of receipt of data with the exception of the two NIST laboratories which are Lab 1 and 14 in this exercise. The laboratory mean replicate data are shown in Tables 1 to 3 for QA10TIS01. Included in these tables are the exercise assigned mean values and the standard deviation of the assigned mean values along with the exercise assigned median values. Summaries of the methods used by each laboratory are in Appendix B, and notes included by a laboratory with its data are listed in Appendix C. In Appendix D, charts of the mean numerical results reported by each laboratory for each analyte are shown for the exercise material and the corresponding reference material, SRM 1974b.

Performance Scores

The exercise coordinators recognize that different environmental monitoring programs have different data quality objectives and needs. The acceptability of the results submitted by a particular laboratory will be decided by the individual program(s) for which the laboratory provides data. Typically, each program will use these exercise results in conjunction with the laboratory's performance in the analysis of certified reference materials and/or control materials, and of other quality assurance samples. These exercise results are exhibited in a number of ways in this report to facilitate their use by most environmental monitoring programs in their acceptability assessments.

IUPAC guidelines describe the use of z-scores and p-scores for assessment of accuracy and precision in intercomparison exercises such as those described in this report. These indices assess the difference between the result of the laboratory and the exercise assigned value and can be used to compare the

performance of different laboratories among the participants on different analytes and on different materials.

Accuracy Assessment (z-score)

$$z\text{-score} = (\text{bias estimate}) / (\text{performance criterion}) = (x - X) / \sigma$$

where x is the individual laboratory result, X is the "Exercise Assigned Value," and σ is the target value for standard deviation.

The choice of σ is dependent upon data quality objectives of a particular program, or as ISO 17043 suggests "a fitness for purpose goal for performance as determined by expert judgement" [6]. It can be "fixed" and arrived at by perception, prescription, or referenced to validated methodology (e.g., $\sigma = 0.025 X$; X is the exercise assigned value,), or it can be an estimate of the actual variation (e.g., the calculated sample standard deviation, s , from the exercise data). The "fixed" performance criterion is more useful in the comparison of a laboratory's performance on different materials while the use of the actual variation may be more useful within a given exercise, for example, if the determination of a particular analyte is exceptionally problematic. The measurement of analytes targeted in this study is not particularly problematic.

We have calculated and reported z-scores using the fixed performance criterion for each analyte for each laboratory using "25 % of the exercise assigned value" as the fixed target value for standard deviation for this program. The use of z-score (25 % X) is also taken from the NOAA Mussel Watch and National Status and Trends Quality Assurance Programs [5]. The z-scores calculated for these exercises can thus be interpreted as shown in the following examples:

z-score (25 % X):

- +1 \Rightarrow laboratory result is 25 % higher than the assigned value
- 2 \Rightarrow laboratory result is 50 % lower than the assigned value.

A common classification of z-scores is [7]:

$ z \leq 2$	Satisfactory
$2 < z < 3$	Questionable
$ z \geq 3$	Unsatisfactory.

This classification has been deemed acceptable within the measurement community.

Tables 4 through 6 summarize the z-scores (25 %) for each laboratory for each reported analyte in QA10TIS01.

Precision Assessment (p-score)

$$p\text{-score} = \sigma_{\text{lab}} / \sigma_{\text{target}}$$

For the calculation of p-scores for this program, the σ values used are coefficients of variation (CV calculated as relative standard deviations) with the current target σ (CV) for the three replicates being

15 %. Tables 7 through 9 summarize the relative standard deviations (RSDs) calculated from the three concentrations reported by the laboratory for each analyte quantified while Tables 10 through 12 summarize the p-scores (15%). A p-score of 1 indicates that the laboratory's CV was 15%, and a p-score of 2 indicates that the laboratory's CV was 30%.

DISCUSSION

NOAA's NRDA office solicited laboratories involved in the analysis of samples shortly after the DWH disaster for their interest in participating in this interlaboratory study for analytes of interest in bivalve tissue. The participation by the laboratories was voluntary, and samples of QA10TIS01 were provided free of charge. Laboratories were provided with information for ordering SRM 1974b (See Appendix A). Interested laboratories were requested to quantify selected PAHs, alkylated PAHs (some individual and some as groups), and biomarkers (hopanes and steranes) in three aliquots of QA10TIS01 and SRM 1974b using their laboratories' analytical protocols for these analyses. A total of 45 laboratories received samples of which 34 laboratories submitted data. The 34 participating laboratories are listed in alphabetical order in Appendix E.

Tables 1 through 3 summarize the laboratory means and exercise assigned values for the PAHs, alkylated PAHs, and hopanes and steranes, respectively. The consensus value for a given compound in QA10TIS01 was derived by combining data where corresponding values in SRM 1974b were within 30 % of the expanded uncertainty of the SRM value. In the absence of a corresponding SRM value, individual results were screened using outlier tests and included in the consensus value if values were shown not to be statistical outliers. Appendix D contains the charts of the QA10TIS01 and SRM 1974b data by analyte. In these charts, the analytes that are not included on the Certificate of Analysis for SRM 1974b are shown with no target value; however, a median value calculated from the data submitted is included in each chart. Note that laboratory 24 submitted two sets of data, 24a and 24b. The differences in the two sets of data included the extraction method, accelerated solvent extraction (ASE) for 24a versus QuEChERS for 24b, and final analysis step, gas chromatography with mass spectrometric detection (GC/MS) for 24a and liquid chromatography with fluorescence detection (LC-FLD) for 24b.

No assigned values were calculated for chrysene, triphenylene, or benzo[*j*]fluoranthene in QA10TIS01. Chrysene and triphenylene coelute when using a 5% phenyl phase in GC analysis as do several of the benzofluoranthene isomers. Although several laboratories recognized and reported coelutions for chrysene with triphenylene and among the benzofluoranthenes using this phase, several did not, thus reporting the combined value for chrysene and triphenylene as only the chrysene peak and in a similar fashion for the benzofluoranthenes.

The spread in the PAH data for QA10TIS01 among the laboratories is highest for biphenyl, acenaphthylene, anthracene, benzo[*a*]fluoranthene and dibenz[*a,h*]anthracene, analytes that are either relatively volatile (biphenyl and acenaphthylene) or at relatively low mass fractions and determined by fewer laboratories. The range in mass fractions for the PAHs in the mussel tissue QA10TIS01 is from 3 ng/g for benzo[*a*]fluoranthene and dibenz[*a,h*]anthracene to 220 ng/g for fluorene. The agreement among the data for the PAHs was better in the previous crude oil study [2] than in this study or in the previous sediment study [1] suggesting that extraction and sample concentration may be issues.

The alkylated PAH data for crude oil QA10 OIL01 [2] do not show as much spread among the laboratories as for the mussel tissue reported here or the sediment material QA10SED01 [1]. As for the previous studies, there are issues with the choice of the representative compound used for quantification of the alkylated PAHs, as well as the interpretation of the chromatograms as to what peaks should and should not be included in the summation for the alkylated groups. For an alkylated group of PAHs, the mass spectra are different based on where the PAH is alkylated (or if it is alkylated) thus resulting in different response factors for isomers in a homolog group. These issues need to be explored in more detail in future discussions.

Fewer laboratories reported data for the selected biomarkers with seven or fewer laboratories reporting quantitative results. The largest spread in the data seen for 5 α (H),14 α (H),17 α (H)-cholestane 20R while very good agreement was seen for 17 α (H),21 β (H)-hopane (an RSD of 12%). The quantification issues for the hopanes and steranes include the lack of authentic calibration standards.

A range of extraction methods was used in this study as summarized in Appendix B by laboratory: nine data sets used Soxhlet extraction (Figure 1), five used QuEChERS (Figure 2), six used sonication (Figure 3), ten used ASE (Figure 4), three used a tissumizer (Figure 5), one used base digestion (Figure 5), and one used microwave extraction (Figure 5). Figures 1 through 5 show the comparison among laboratories using the same extraction method for selected compounds along with the median for the data sets. Figure 6 compares the medians for each extraction method for the same selection of compounds. The data for all of the compounds are summarized by extraction method in Tables 13, 14, and 15 for the PAHs, alkylated PAHs, and hopanes and steranes, respectively. No concrete conclusions can be drawn from comparing the data obtained by the different extraction methods as there is a large spread in the data (see table below) from each extraction method, with the tissumizer results (from three laboratories being the least variable. This table summarizes the number of laboratories who performed each extraction method (not necessarily reporting data for each analyte), the average relative standard deviations (avg RSD in %) for each class of compounds, and the number of compounds in each class for which more than one data set was received by the indicated extraction method (for n=). No data were excluded when calculating the RSDs shown.

Extraction Method	# labs reporting	PAHs avg rsd (%)	total n=34 for n=	alkylated		hopanes/ steranes avg rsd (%)		total n=18 for n=
				PAHs avg rsd (%)	total n=40 for n=	steranes avg rsd (%)	steranes avg rsd (%)	
Soxhlet	9	50	27	50	31	35	2	
QuEChERS	5	51	13		0			0
Tissumizer	3	18	24	27	38	27	17	
Sonication	6	116	17	123	15			0
ASE	10	32	24	44	29			0
Base digestion	1		20		27			8
Microwave extraction	1		5		8			0

Caution should be used when interpreting these data as for the sonication method, one laboratory (17) reported high values for many of the analytes. These high values are probably not a result of the extraction method since their data for the corresponding analytes in SRM 1974b generally fell within the 30% criterion mentioned in the “*Determination of assigned values*” section. See Appendix D for the charts of the individual analytes.

The majority of the z-scores based on 25 % (Tables 4 through 6) are within ± 2 while the majority of the p-scores based on 15 % (Tables 10 through 12) are within ± 1 . This indicates that the laboratories are internally consistent, but there is still a fair amount of spread in the data among the laboratories.

It is important to evaluate the non-quantitative results reported by each laboratory as well. Although these results are not easily presented or numerically evaluated, they are included in Tables 1 through 3 of this report. The laboratory and its data users should closely examine these non-quantitative results. Decisions based on false negative or false positive results from a laboratory can lead to significant environmental and/or economic consequences. Some laboratories reported detection limits in the exercise material that may be too high for the data quality objectives and needs of their program(s), and these issues should be assessed as well.

Intercomparison exercises provide an important mechanism for assessing the comparability, repeatability, and trueness of data being produced by the participating laboratories. Exercise materials similar in matrix, form, and analyte concentration to typical samples routinely analyzed by the laboratories are most useful for demonstrating the level of comparability and for revealing potential measurement and method problems. Minimizing the among-laboratory biases so that the analytical variability is significantly less than the field sampling variability should be an achievable goal in environmental monitoring.

Acknowledgments

The time and effort of the analysts and management of the participating laboratories are gratefully acknowledged.

Disclaimer

Certain commercial equipment, instruments, or materials are identified in this report to specify adequately the experimental procedure. Such identification does not imply recommendation or endorsement by the National Institute of Standards and Technology, nor does it imply that the materials or equipment identified are the best available for the purpose.

References

1. Schantz, M.M. and Kucklick, J.R., Interlaboratory Analytical Comparisons Study to Support Deepwater Horizon Natural Resource Damage Assessment: Description and Results for Marine Sediment QA10SED01, NISTIR ???, Gaithersburg, MD (2011).
2. Schantz, M.M. and Kucklick, J.R., Interlaboratory Analytical Comparisons Study to Support Deepwater Horizon Natural Resource Damage Assessment: Description and Results for Crude Oil QA10OIL01, NISTIR ???, Gaithersburg, MD (2011).
3. NOAA Technical Memorandum NOS ORCA 79 Quality Assurance Project Intercomparison Exercise Results 1991-1993, August, 1995
4. Certificate of Analysis for Standard Reference Material (SRM) 974b Organics in Mussel Tissue (*Mytilus edulis*), National Institute of Standards and Technology (NIST), Gaithersburg, MD, 2003. (https://www-s.nist.gov/srmors/view_detail.cfm?srm=1974B)

5. Schantz, M.M., Parris, R.M., and Wise, S.A., NIST Intercomparison Exercise Program for Organic Contaminants in the Marine Environment: Description and Results of 2007 Organic Intercomparison Exercises, NISTIR 7501, Gaithersburg, MD (2008).
6. ISO/IEC 17043: 2010 Conformity Assessment-General Requirements for Proficiency Testing
7. IUPAC “The International Harmonized Protocol for the Proficiency Testing of (Chemical) Analytical Laboratories,” Pure & Appl. Chem., 65 (9), 2123-2144 (1993).

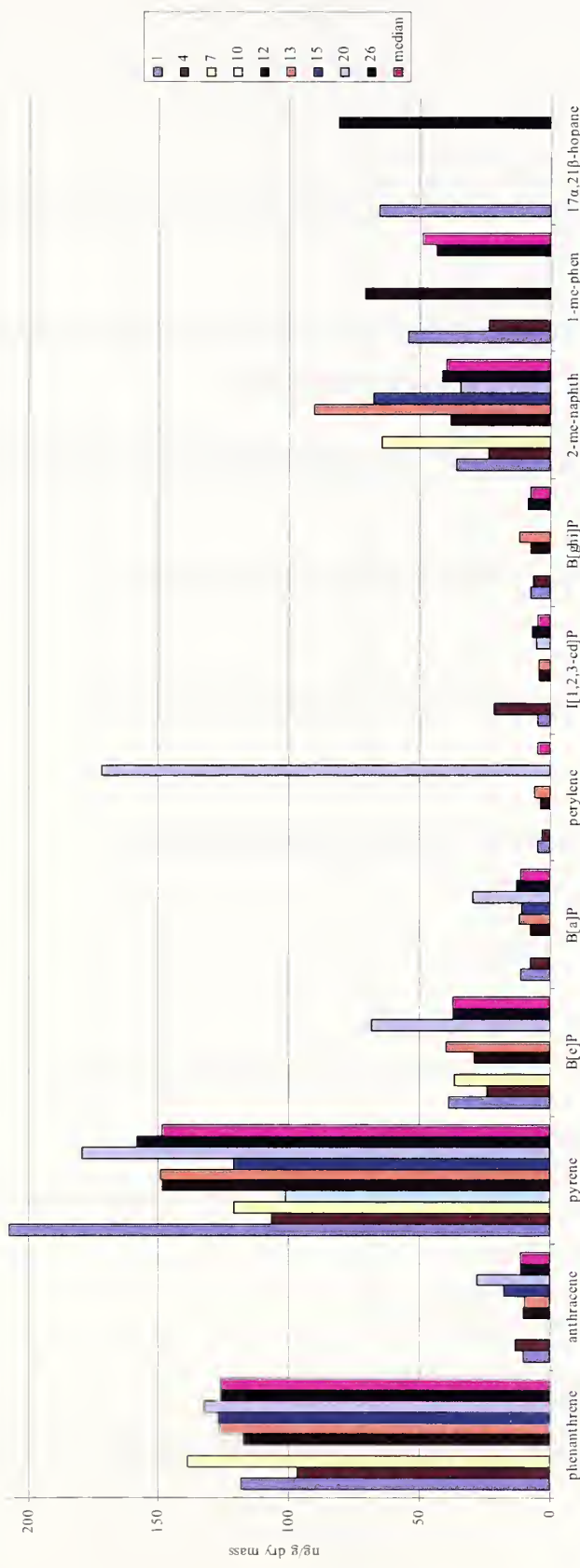


Figure 1. Data from laboratories performing Soxhlet extraction. Medians are not calculated when there are two or fewer sets of data.

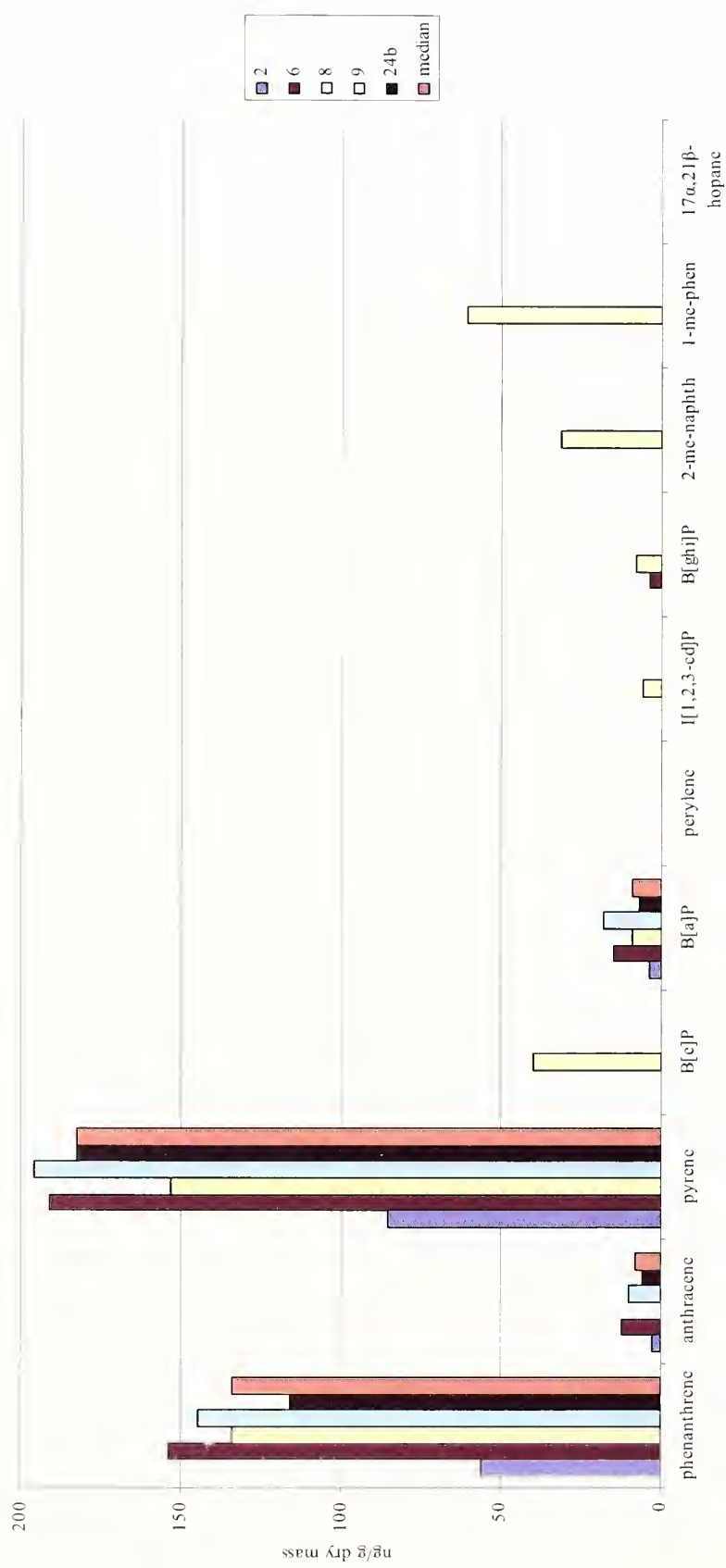


Figure 2. Data from laboratories performing QueChERS. Medians are not calculated when there are two or fewer sets of data.

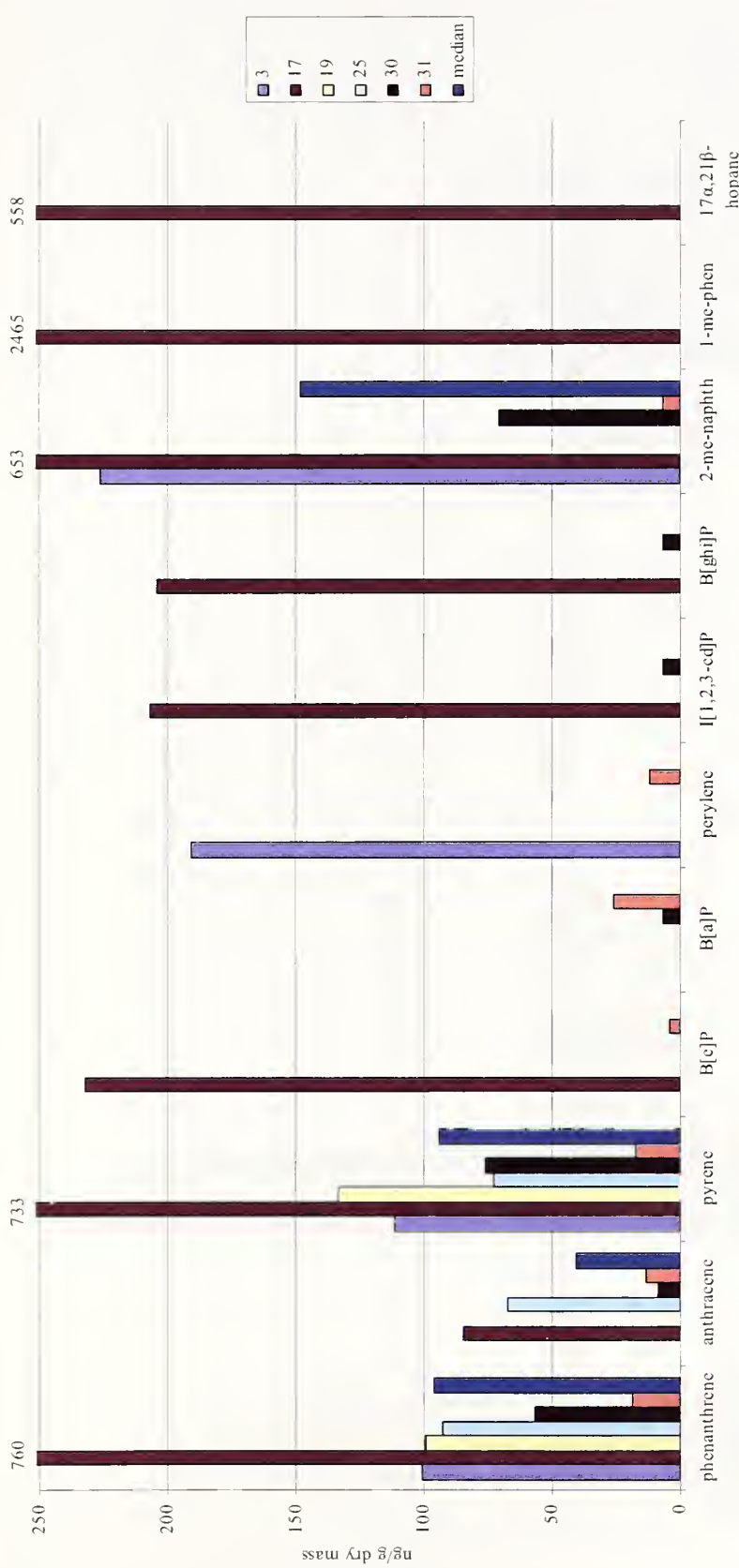


Figure 3. Data from laboratories performing sonication. Medians are not calculated when there are two or fewer sets of data.

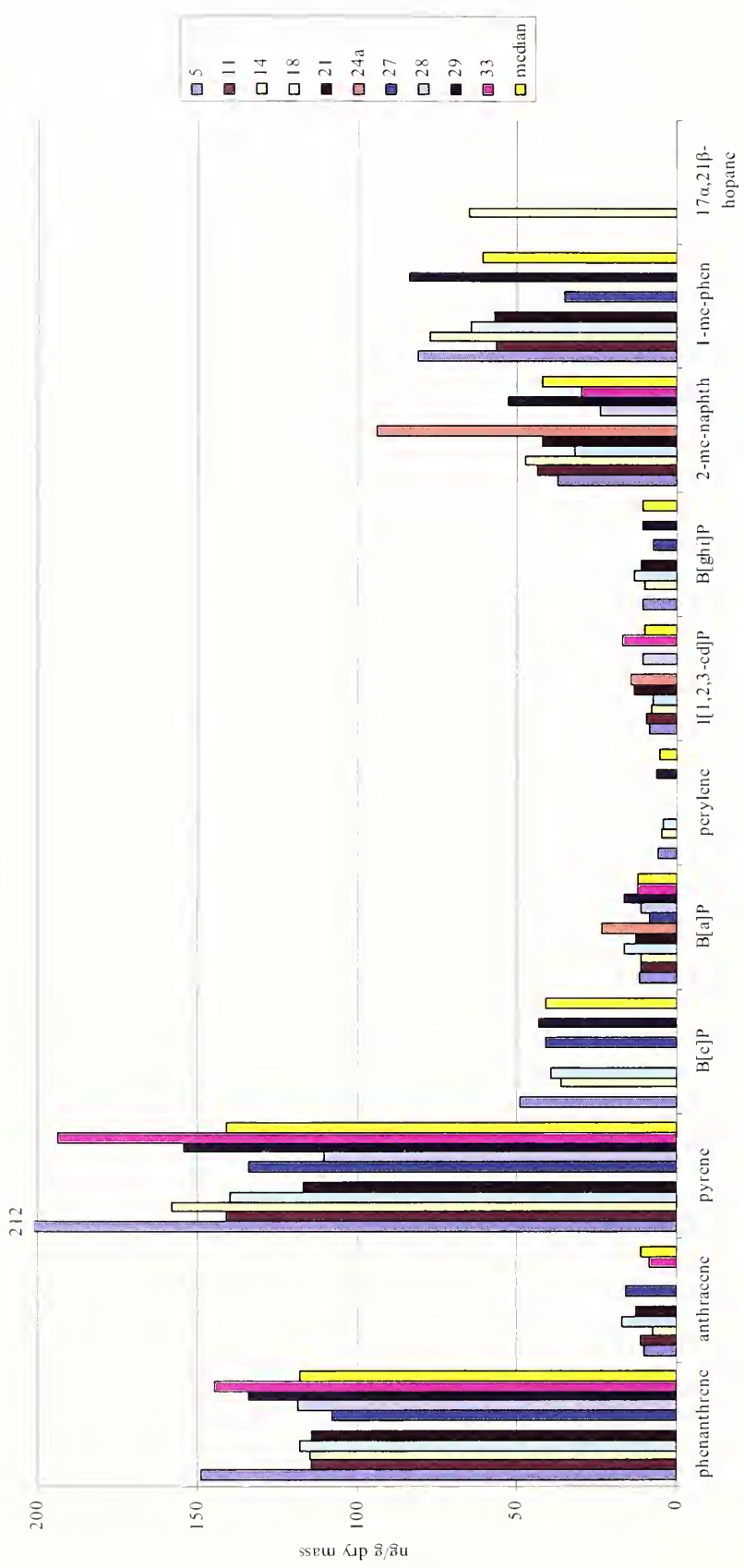


Figure 4. Data from laboratories performing ASE. Medians are not calculated when there are two or fewer sets of data.

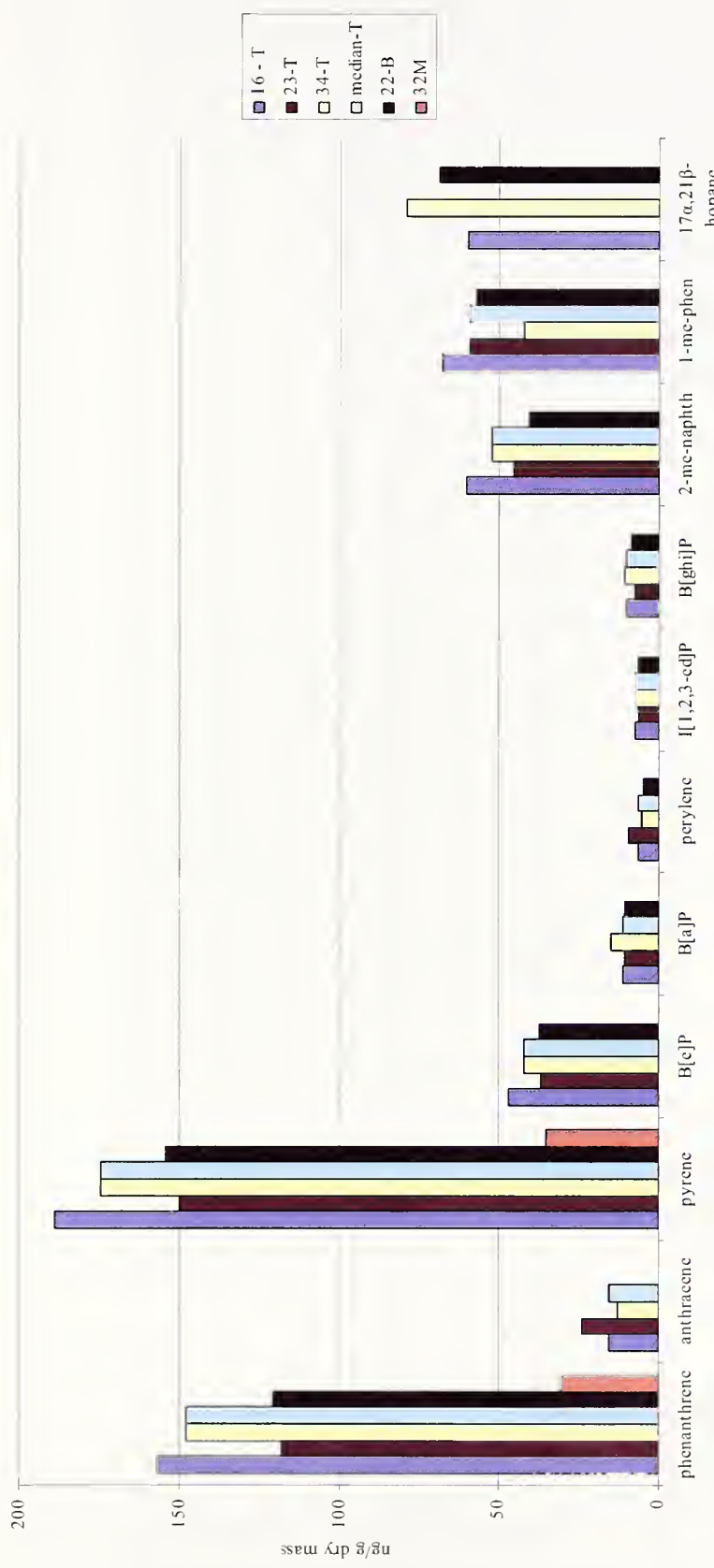


Figure 5. Data from the three laboratories using the tissumizer method (T) along with the medians for that method and from the one laboratory using base digestion (B) and one laboratory using microwave extraction (M). Medians are not calculated when there are two or fewer sets of data.

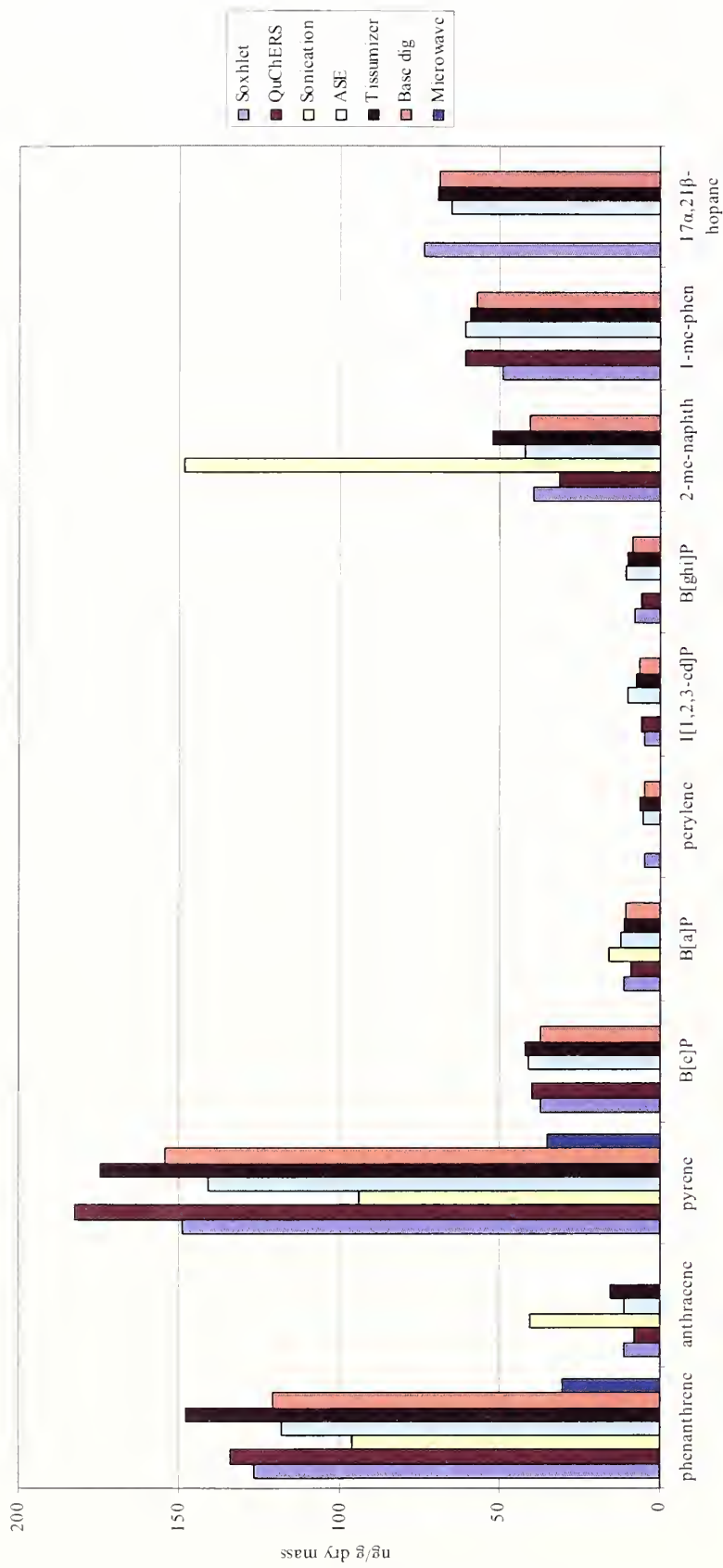


Figure 6. Data comparison by extraction method. Medians are shown for the data sets with 2 or data points that were not excluded from the assignment of the consensus means. Otherwise, the data represent one set of data. See Tables 13 to 15 for the complete summary.

Table 1. Mussel Tissue (QAL01TS01): Laboratory means of three replicates and exercise assigned values - Water and PAHs

	Laboratory No.	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19
Percent Water	85.4	87.8	86.0	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19
ng/g dry mass																				
naphthalene	43.0	37.2	<294	19.7	12.9	104	36.2	31.3	other	<5700	18.4	24.2	54.5	NA	350	34.6	625	30.2	<78	
biphenyl	12.9	13.0	16.0	Other	<14.8	NA	NA	<5700	NA	9.18	21.1	NA	18.3	13.6	567	19.1	<78			
aceanthiphene	14.0	nd	<294	11.3	9.59	0.00	<14.8	<0.2	<11	<5700	NA	7.42	<30	9.70	<10	11.7	<3.2	9.64	<78	
aceanthiphene	<10	<294	9.03	2.74	NA	<14.8	<0.2	NA	<5700	NA	4.50	<30	8.08	12.7	10.7	<3.2	6.77	<78		
fluorene	33.9	14.3	<294	24.3	26.0	43.6	19.8	25.2	17.7	<5700	19.7	14.8	33.3	30.9	25.3	24.0	20.3	<78		
phenanthrene	119	56.1	101	97.0	149	153	139	134	144	<5700	114	117	126	114	127	15.7	760	118	99.3	
anthracene	10.3	2.76	<294	13.3	10.0	12.1	<14.8	other	10.0	<5700	11.4	10.3	9.69	7.70	17.7	85	17.3	<78		
fluoranthene	246	11.5	181	140	305	247	196	241	260	183	221	269	210	209	173	28.8	1160	303	207	
Pyrene	207	85.4	112	107	212	191	121	153	196	101	141	149	149	158	121	189	733	140	133	
benzofluoranthene	NA	NA	17.3	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	<3.2	NA	<78	
benz[a]anthracene	37.3	8.87	<294	19.7	57.8	49.9	16.3	28.3	other	<5700	35.8	32.1	30.8	38.6	26.7	35.4	198	53.2	<78	
chrysene	59.1	<294	54.7	Other	94.8	93.9	49.8	107	<5700	96.2	73.0	52.9	35.3	88.3	coelute	82.5	<78			
triphenylene	NA	NA	NA	Other	NA	NA	NA	NA	NA	NA	NA	NA	NA	38.3	NA	coelute	82.5	NA		
chrysene/triphenylene	144				139											10.8	421			
benzo[b]fluoranthene	26.3	14.4	<294	20.7	38.1	29.8	21.9	27.3	30.8	<5700	44.0	25.0	35.1	31.0	Other	34.6	261	32.6	<78	
benzo[f]fluoranthene	28.6	NA	<294	13.0	Other	NA	NA	NA	NA	<5700	NA	NA	NA	16.6	Other	NA	coelute	NA	<78	
benzo[k]fluoranthene	15.4	3.54	<294	16.6	20.3	11.9	16.8	<5700	25.4	24.7	14.4	15.1	11.0	coelute	coelute	14.7	<78			
benzo[b+k]fluoranthene														0.027		31.7	194			
benzo[b+j+k]fluoranthene																				
benzo[a]fluoranthene	<10	NA	1.33	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	5.42	NA	<3.2	NA	<78	
benzo[c]fluorene	38.8	<294	24.3	48.9	NA	36.6	39.9	NA	<5700	NA	29.3	39.9	36.1	NA	46.7	222	39.4			
benzo[al]pyrene	11.4	3.91	<294	7.40	11.8	14.7	<14.8	9.03	18.4	<5700	11.5	7.55	11.7	11.0	10.7	11.0	<3.2	16.7	<78	
Dibenzene	4.96	191	2.97	5.68	NA	<14.8	NA	NA	<5700	NA	3.93	5.86	4.74	<10	6.55	<3.2	4.16	<78		
indeno[1,2,3-cd]pyrene	4.75	nd	<294	21.3	8.35	0.00	<14.8	5.83	<21	<5700	9.39	4.39	4.58	8.22	<10	7.70	207	7.65	<78	
benzo[b]phenanthrene	7.81	<294	6.67	10.8	3.83	<14.8	8.03	<7	<5700	NA	7.81	11.9	9.88	<10	9.92	204	13.1			
dibenz[b]phenanthrene	1.12	nd	<294	0.00	Other	0.00	<14.8	<0.2	<9	<5700	18.2	2.28	2.11	<1.5	<10	2.54	175	3.14	<78	
dibenz[a]anthracene						2.35														
dibenz[a,h]anthracene	NA	NA	63.3	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	14.2	<3.2	NA	<78	
cis/trans-decalin	NA	<294	21.3	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	18.2	143	NA	<78	
dibenzofuran	NA	NA	42.3	11.8	NA	NA	NA	NA	NA	NA	50.1	NA	28.3	30.0	56.2	245	NA			
retene	NA	<294	12.7	20.9	NA	<14.8	NA	NA	<5700	NA	<40	NA	NA	NA	1.30	<3.2	NA	NA	<78	
benzothiophene	<15														12.0	20.4	<3.2	11.8		
dibenzothiophene	NA	NA	11.7	Other	NA	NA	NA	NA	NA	NA	13.6	NA	NA	NA	7.41	<3.2	NA	NA		

NA=Not analyzed; DL=Detection limit; No value in space=nothing reported by lab. Bolded values were not used in the calculation of the consensus statistics

Table 1 (cont.). Mussel Tissue (Q.A101FS01): Laboratory means of three replicates and exercise assigned values - Water and PAHs
(reported as if three figures were significant)

ng/g dry mass	Laboratory No.	Consensus Values (%)																		
		Percent Water	20	21	22	23	24a	24b	25	26	27	28	29	30	31	32	33	34	mean	std dev
naphthalene	37.0	23.0	24.3	26.5	47.0	117	<conc	24.3	<12.2	15.0	31.0	14.7	5.60	<26	22.5	28.4	27.2	9.2	30.2	
phenanthrene	35.0	NA	11.8	16.5	NA	NA	<conc	16.3	13.3	NA	NA	<39.8	6.87	<26	NA	12.2	16.6	6.4	16.1	
acenaphthene	<7.4	6.67	10.4	9.06	NA	DL	<conc	11.1	7.12	NA	DL	6.69	11.7	<26	NA	10.4	9.63	2.13	9.67	
acenaphthylene	13.8	4.45	7.68	16.0	NA	NA	<conc	10.1	12.5	NA	DL	6.29	<3.0	<26	NA	7.86	8.88	3.76	8.08	
fluorene	23.5	20.6	18.6	20.8	DL	<conc	18.7	19.5	18.5	29.2	18.7	<3.0	<26	25.2	21.5	23.4	6.6	21.5		
phenanthrenec	133	11.4	121	118	DL	116	92.8	126	108	119	134	56.6	18.7	30.3	14.5	14.8	120	2.4	11.9	
anthracene	28.2	12.8	<40	24.2	DL	6.10	67.7	11.4	16.1	<6.21	DL	8.75	13.7	<26	8.29	12.7	12.6	5.7	12.1	
fluoranthene	127	23.8	200	221	DL	215	11.4	227	242	231	215	176	24.0	52.7	277	262	217	50	218	
pyrene	179	11.7	15.4	15.0	DL	182	73.0	158	134	111	154	75.8	17.3	35.0	19.4	17.5	146	3.8	149	
benzo[b]fluorene	NA	NA	NA	NA	NA	NA	<conc	37.0	<15.4	22.4	DL	<39.8	NA	NA	NA	NA	12.2	20.9	9.7	
benzo[flanthracene	63.6	33.7	27.4	28.9	DL	50.3	<conc	28.3	34.2	33.8	37.2	24.0	5.20	<26	55.9	34.8	36.6	12.1	34.0	
chrysene	88.7	58.7	other	Other	29.2	85.7	43.8	Other	cooclite	83.3	cooclite	56.0	15.3	26.0	71.2	cooclite	No Target	No Target	65.2	
triphénylene	NA	NA	other	NA	NA	NA	Other	NA	NA	cooclite	NA	NA	NA	NA	NA	NA	NA	No Target	60.4	
chrysene/triphenylene	84.4	69.6																102	103	105
benzo[b]fluoranthene	36.0	55.5	25.0	37.6	48.3	20.8	<conc	37.5	42.1	22.4	28.1	21.8	<3.0	cooclite	57.2	30.5	32.4	10.5	30.9	
benzo[b]fluoranthene	NA	NA	other	NA	NA	NA	NA	Other	cooclite	NA	NA	cooclite	NA	NA	NA	NA	No Target	No Target	22.6	
benzo[k]fluoranthene	65.8	14.3	other	12.5	DL	7.83	<conc	Other	NA	30.1	cooclite	14.2	3.20	cooclite	19.3	cooclite	15.8	6.2	14.7	
benzo[b+k]fluoranthene																	No Target	No Target		
benzo[b+k]fluoranthene	NA	NA	NA	NA	NA	NA	<conc	3.99	<2.94	NA	DL	NA	NA	<26	NA	NA	NA	No Target		
benzo[a]fluoranthene	68.6	NA	37.5	36.9	NA	NA	<conc	37.4	40.8	NA	43.1	<39.8	4.00	<26	NA	42.3	38.6	5.9	39.4	
benzo[a]pyrene	29.5	12.9	10.6	10.4	23.3	7.05	<conc	12.7	84.2	11.4	16.5	6.36	26.0	<26	12.3	15.2	12.3	5.3	11.4	
perylene	172	NA	4.57	9.40	NA	NA	<conc	<10	<7.65	NA	6.50	<39.8	12.1	<26	NA	5.22	5.38	1.64	5.68	
indeno[1,2,3- <i>cd</i>]pyrene	5.12	13.4	6.61	6.44	14.2	DL	<conc	7.07	<4.58	10.5	DL	6.36	<3.0	<26	16.8	7.24	8.05	4.00	7.45	
benzo[<i>b</i>]phenylene	<7.4	11.0	8.75	7.73	NA	DL	<conc	8.68	7.29	NA	10.5	6.36	<3.0	<26	NA	10.7	8.92	2.25	8.75	
dibenz[a,h]anthracene	<7.4	5.09	other	1.77	DL	DL	<conc	7.83	<3.28	<4.81	DL	<39.8	<3.0	<26	3.23	cooclite	3.00	2.07	2.54	
dibenz[a,h]anthracene	NA	NA	NA	NA	NA	NA	NA	91.9	33.9	NA	DL	<39.8	NA	38.3	NA	3.63	2.63	0.89	2.35	
cis/trans-decalin	<7.4	NA	NA	NA	NA	<conc	1.58	17.1	NA	DL	<39.8	<3.0	<26	NA	1.13	No Target	77.6			
dibenzofuran	NA	NA	30.7	NA	NA	NA	NA	25.4	<9.61	NA	DL	NA	NA	<26	NA	17.4	2.6	17.7		
retene	NA	NA	NA	NA	NA	NA	NA	<1.3	<5.15	NA	DL	<39.8	NA	<26	NA	<2.26	34.4	14.4	30.7	
benzothiophene	<7.4	14.0	14.6	15.8	NA	NA	<conc	16.7	17.6	NA	18.6	<39.8	<3.0	<26	NA	1.66	No Target	1.30		
dibenzothiophene	NA	NA	NA	NA	NA	NA	NA	12.7	20.4	NA	NA	NA	NA	<26	NA	18.4	15.7	3.0	15.8	
naphthalenethiophene																	14.0	4.6	13.1	

NA = Not analyzed; DL = Detection limit; No value in space=nothing reported by lab. Bolded values were not used in the calculation of the consensus statistics (Lab 31 reported values on a wet mass basis)

Table 2. Mussel Tissue (QA10UTS01): Laboratory means of three replicates and exercise assigned values - Alkylated PAHs (ng/g dry mass) (reported as if three figures were significant)

Laboratory No.	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19
1-methyl/naphthalene	27.2		<294	15.0	21.0	NA	113	50.3	NA	<5700	27.4	23.0	50.5	29.8	37.7	40.1	<6.4	19.9	<78
2-methyl/naphthalene	226	226	23.7	37.1	NA	64.4	31.5	NA	<5700	43.8	38.0	90.3	47.6	67.7	60.2	653	31.8	<78	
2,6-dimethyl/naphthalene	67.0	102	42.3	68.8	NA	NA	70.5	NA	NA	46.9	65.4	101	41.5	NA	90.2	1062	46.5	<78	
1,6,7-trimethyl/naphthalene	46.5	112	46.3	39.4	NA	NA	NA	NA	NA	50.5	74.3	NA	NA	76.8	325	41.4	<78		
1-methylphenanthrene	54.6	<294	23.7	81.1	NA	NA	61.1	NA	NA	56.4	70.9	Other	77.1	NA	67.5	2465	64.8	<78	
C1-decalins	NA	NA	390	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	831	2467	NA	NA	
C2-decalins	NA	NA	1333	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	2137	14233	NA	NA	
C3-decalins	NA	NA	1900	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	1727	21200	NA	NA	
C4-decalins	NA	NA	2533	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	2707	24533	NA	NA	
C1-naphthalenes	63.4	596	23.7	58.2	NA	NA	NA	NA	NA	<5700	NA	35.5	NA	NA	NA	64.7	862	NA	<78
C2-naphthalenes	255	378	78.3	236	NA	NA	NA	NA	NA	<5700	NA	125	NA	NA	NA	313	194	2728	NA
C3-naphthalenes	629	391	153	606	NA	NA	NA	NA	NA	207	NA	253	NA	NA	NA	419	406	4270	NA
C4-naphthalenes	498	594	170	600	NA	NA	NA	NA	NA	110	NA	210	NA	NA	NA	466	362	5307	NA
C1-benzothiophenes	NA	NA	0.00	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	16.9	<3.2	NA	NA
C2-benzothiophenes	NA	NA	0.00	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	34.6	<3.2	NA	NA
C3-benzothiophenes	NA	NA	0.00	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	55.8	<3.2	NA	NA
C4-benzothiophenes	NA	NA	0.00	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	41.6	<3.2	NA	NA
C1-fluorenes	76.8	<294	34.0	68.6	NA	NA	NA	NA	NA	<5700	NA	37.7	NA	NA	NA	75.7	72.4	856	NA
C2-fluorenes	181	<294	107	186	NA	NA	NA	NA	NA	<5700	NA	109	NA	NA	NA	188	209	2563	NA
C3-fluorenes	296	<294	473	262	NA	NA	NA	NA	NA	<5700	NA	173	NA	NA	NA	436	300	<3.2	NA
C1-phenanthrenes/anthracenes	280	<294	163	425	NA	NA	NA	NA	NA	<5700	NA	226	NA	NA	NA	415	321	<3.2	NA
C2-phenanthrenes/anthracenes	521	662	263	626	NA	NA	NA	NA	NA	102	NA	326	NA	NA	NA	689	510	2353	NA
C3-phenanthrenes/anthracenes	467	<294	323	508	NA	NA	NA	NA	NA	<5700	NA	232	NA	NA	NA	1044	361	3257	NA
C4-phenanthrenes/anthracenes	145	<294	230	209	NA	NA	NA	NA	NA	<5700	NA	113	NA	NA	NA	597	130	1380	NA
C1-dibenzothiophenes	95.9	<294	55.3	92.1	NA	NA	NA	NA	NA	<5700	NA	72.3	NA	NA	NA	64.7	99.4	385	NA
C2-dibenzothiophenes	237	<294	140	245	NA	NA	NA	NA	NA	<5700	NA	193	NA	NA	NA	192	269	946	NA
C3-dibenzothiophenes	223	<294	143	206	NA	NA	NA	NA	NA	<5700	NA	163	NA	NA	NA	155	256	1113	NA
C4-dibenzothiophenes	110	<294	87.0	103	NA	NA	NA	NA	NA	<5700	NA	68.7	NA	NA	NA	<40	110	877	NA
C1-fluoranthenes/pyrenes	145	<294	70.0	114	NA	NA	NA	NA	NA	<5700	NA	91.9	NA	NA	NA	66.7	134	717	NA
C2-fluoranthenes/pyrenes	623	166	42.7	68.3	NA	NA	NA	NA	NA	<5700	NA	55.3	NA	NA	NA	43.0	67.1	<3.2	NA
C3-fluoranthenes/pyrenes	33.4	<294	31.7	37.1	NA	NA	NA	NA	NA	<5700	NA	24.3	NA	NA	NA	<40	40.2	<3.2	NA
C4-fluoranthenes/pyrenes	<15	NA	12.6	12.2	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	<3.2	NA	NA
C1-naphthalobenzoithiophenes	NA	NA	55.7	20.4	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	12.2	NA	NA
C2-naphthalobenzoithiophenes	NA	NA	0.00	13.8	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	8.7	<3.2	NA	NA
C3-naphthalobenzoithiophenes	NA	NA	0.00	6.97	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	7.59	<3.2	NA	NA
C4-naphthalobenzoithiophenes	NA	NA	0.00	3.10	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	0.00	<3.2	NA	NA
C1-chrysenes	42.6	<294	25.7	56.9	NA	NA	NA	NA	NA	<5700	NA	28.9	NA	NA	NA	44.4	308	NA	<78
C2-chrysenes	25.7	<294	21.3	31.2	NA	NA	NA	NA	NA	<5700	NA	16.2	NA	NA	NA	32.2	<3.2	NA	<78
C3-chrysenes	<10	<294	9.93	13.5	NA	NA	NA	NA	NA	<5700	NA	6.20	NA	NA	NA	0.00	<3.2	NA	<78
C4-chrysenes	<10	<294	0.00	<1.79	NA	NA	NA	NA	NA	<40	NA	NA	NA	NA	NA	<3.2	NA	<78	

NA=Not analyzed; DL=Detection limit; No value in space=nothing reported by lab; Bolded values were not used in the calculation of the consensus statistics

Table 2 (cont). Mussel tissue (QA10T1S01): Laboratory means of three replicates and exercise assigned values - Alkylated PAHs (ng/g dry mass) (reported as if three figures were significant)

	Laboratory No.	20	21	22	23	24a	24b	25	26	27	28	29	30	31	32	33	34	mean	std dev	median
1-methylnaphthalene		37.0	25.6	24.8	30.4	58.3	NA	<conc	29.7	<24.3	38.3	31.2	16.1	7.90	<52	47.6	34.2	31.7	11.1	30.1
2-methylnaphthalene		34.4	42.3	40.8	45.1	94.0	NA	<conc	41.4	<24.3	24.0	52.5	70.5	6.90	<52	30.0	52.2	43.6	15.3	43.1
2,6-dimethylnaphthalene		106	41.6	76.2	53.7	118	NA	104	78.8	48.4	69.7	DL	27.1	13.2	<52	87.5	73.3	70.7	24.9	69.7
1,6,7-trimethylnaphthalene		168	NA	41.7	NA	NA	<conc	81.4	30.6	NA	46.0	NA	17.7	<52	NA	35.6	63.6	37.6	46.4	
1-methylphenanthrene		<7.4	57.3	57.3	59.2	DL	NA	<conc	43.7	35.1	<8.65	83.8	NA	<3.0	<52	0.00	42.3	58.5	16.4	58.3
C1-decalins		NA	NA	NA	NA	NA	NA	<conc	560	30.6	NA	DL	NA	NA	<52	NA	547	582	18.3	554
C2-decalins		NA	NA	NA	NA	NA	NA	<conc	1733	45.6	NA	DL	NA	NA	<52	NA	1353	1639	37.9	1543
C3-decalins		NA	NA	NA	NA	NA	NA	<conc	2967	310	NA	DL	NA	NA	<52	NA	1127	1930	7.66	1813
C4-decalins		NA	NA	NA	NA	NA	NA	<conc	3533	506	NA	DL	NA	NA	<52	NA	1493	2367	838	2620
C1-naphthalenes		47.2	63.4	65.5	75.4	NA	NA	<conc	48.3	<43.3	62.3	90.0	NA	13.0	<52	77.5	55.3	59.3	17.0	63.4
C2-naphthalenes		212	277	298	173	508	NA	<conc	180	102	263	229	NA	26.0	<52	321	138	232	106	229
C3-naphthalenes		314	390	615	306	497	NA	<conc	290	226	498	695	NA	27.0	54.3	43.5	235	386	165	390
C4-naphthalenes		319	240	554	326	683	NA	<conc	400	279	138	Other	NA	18.3	76.0	313	247	173	316	
C1-benzothiophenes		NA	NA	NA	NA	NA	NA	<conc	<39	21.1	NA	DL	NA	NA	<52	NA	14.4	17.5	3.4	15.7
C2-benzothiophenes		NA	NA	NA	NA	NA	NA	<conc	<39	12.4	NA	DL	NA	NA	<52	NA	19.5	22.2	11.3	15.9
C3-benzothiophenes		NA	NA	NA	NA	NA	NA	<conc	<39	20.4	NA	DL	NA	NA	<52	NA	49.4	41.8	18.9	34.9
C4-benzothiophenes		NA	NA	NA	NA	NA	NA	<conc	54.0	37.8	NA	DL	NA	NA	<52	NA	50.2	45.9	7.5	41.6
Cl-fluoranes		<7.4	84.9	140	68.6	98.7	NA	<conc	68.3	70.1	21.5	446	NA	<3.0	<52	103	51.8	71.4	29.3	72.4
C2-fluoranes		<7.4	195	312	146	193	NA	<conc	373	190	181	658	NA	<3.0	60.0	77.2	150	177	77	186
C3-fluoranes		<7.4	48.4	530	242	DL	NA	<conc	780	354	<6.89	558	NA	<3.0	67.0	277	228	328	190	287
C1-phenanthrenes/anthracenes		<7.4	312	328	258	685	NA	<conc	187	230	360	554	NA	<3.0	59.3	433	257	331	138	296
C2-phenanthrenes/anthracenes		<7.4	113	546	378	1457	NA	106	493	393	536	934	NA	<3.0	93.3	63.9	398	434	219	502
C3-phenanthrenes/anthracenes		<7.4	155	547	294	469	NA	<conc	460	307	356	837	NA	<3.0	77.9	482	240	457	224	361
C4-phenanthrenes/anthracenes		<7.4	NA	590	141	548	NA	<conc	260	159	<9.18	426	NA	<3.0	<52	237	92.5	246	170	184
Cl-dibenzothiophenes		46.2	NA	103	85.1	NA	NA	<conc	82.7	71.6	NA	188	NA	<3.0	<52	NA	71.4	86.8	35.0	83.9
C2-dibenzothiophenes		<7.4	NA	355	215	NA	NA	<conc	287	226	NA	651	NA	<3.0	<52	NA	201	211	134	226
C3-dibenzothiophenes		61.7	NA	321	190	NA	NA	<conc	303	234	NA	705	NA	<3.0	<52	NA	186	196	73	206
C4-dibenzothiophenes		55.6	NA	132	NA	NA	<conc	263	108	NA	433	NA	<3.0	NA	NA	95.4	113	57	109	
Cl-fluoranthenes/pyrenes		373	338	376	123	NA	NA	<conc	127	115	NA	112	NA	<3.0	<52	NA	107	159	107	119
C2-fluoranthenes/pyrenes		<7.4	91.6	291	59.1	NA	NA	<conc	92.7	71.7	NA	84.2	NA	<3.0	<52	NA	59.3	74.1	31.8	67.7
C3-fluoranthenes/pyrenes		57.9	NA	99.1	27.0	NA	NA	<conc	52.3	36.2	NA	40.9	NA	<3.0	<52	NA	30.2	42.5	20.3	36.7
C4-fluoranthenes/pyrenes		NA	NA	22.4	NA	NA	<conc	<39	16.9	NA	18.5	NA	<3.0	NA	NA	15.2	16.3	3.9	16.0	
Cl-naphthalenobenzothiophenes		NA	NA	NA	NA	NA	<conc	<39	19.1	NA	NA	<3.0	<52	NA	18.9	25.2	17.3	19.1		
C2-naphthalenobenzothiophenes		NA	NA	NA	NA	NA	<conc	<39	16.0	NA	NA	<3.0	<52	NA	15.9	13.6	3.4	13.8		
C3-naphthalenobenzothiophenes		NA	NA	NA	NA	NA	<conc	<39	8.19	NA	NA	<3.0	<52	NA	9.75	8.11	1.46	7.28		
C4-naphthalenobenzothiophenes		NA	NA	NA	NA	NA	<conc	<39	8.19	NA	NA	<3.0	<52	NA	<0.707	No Target	38.7	42.3	9.9	43.5
Cl-phenylenes		35.5	NA	60.9	44.7	NA	NA	<conc	43.5	41.6	NA	43.9	NA	<3.0	<52	NA	NA	23.9	7.6	24.8
C2-phenylenes		210	NA	30.6	19.5	NA	NA	<conc	43.0	22.4	NA	21.8	NA	<3.0	<52	NA	NA	26.2	9.9	24.8
C3-phenylenes		<7.4	NA	6.74	6.84	NA	NA	<conc	<39	<13.8	NA	12.3	NA	<3.0	<52	NA	<0.595	9.25	3.14	6.84
C4-phenylenes		<7.4	NA	0.540	2.93	NA	NA	<conc	<39	<13.8	NA	7.81	NA	<3.0	<52	NA	<0.595	No Target	0.540	

NA=Not analyzed; DL=Detection limit; No value in space=nothing reported by lab. Bolded values were not used in the calculation of the consensus statistics (Lab 31 reported values on a wet mass basis)

Table 3. Mussel Tissue (QA101HS01): Laboratory means of three replicates and exercise assigned values - Biomarkers (ng/g dry mass)

		Biomarkers (ng/g dry mass)																	
		(reported as if three figures were significant)																	
Laboratory No.	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19
Carbazole	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	<40	NA	NA	NA	NA	NA	NA	NA	NA
18a(H)-22,29,30-Trisnorbornane	NA	<294	NA	<5700	NA	<40	NA	NA	NA	NA	223	NA	NA						
17a(H)-22,29,30-Trisnorbornane	<25	<294	NA	<5700	NA	<40	NA	NA	NA	15.3	NA	NA	<78						
18a(H)-30-Norbornane	NA	<294	NA	<5700	NA	<40	NA	NA	NA	15.7	NA	NA	<78						
17a(H)-Dihydropane	NA	<294	NA	<5700	NA	<40	NA	NA	NA	0.00	NA	NA	<78						
17a(H)-2,3[β]H-Hopane	NA	<294	NA	<5700	NA	<40	NA	NA	NA	48.2	NA	NA	<78						
17a(H)-2,3[β]H-22R-Homohopane	65.7	<294	NA	<5700	NA	<40	NA	NA	NA	0.00	NA	NA	<78						
17a(H)-2,3[β]H-22R-Homohopane	<20	<294	NA	<5700	NA	<40	NA	NA	NA	65.3	Other	558	NA						
17a(H)-2,3[β]H-22S-Homohopane	23.1	<294	NA	<5700	NA	<40	NA	NA	NA	12.4	Other	18.4	NA						
13[α]H,7[α]H,17[α]H-Diacholestan-20S	NA	<294	NA	<5700	NA	<40	NA	NA	NA	20.8	Other	NA	<78						
5a(H),14a(H),17a(H)-Cholestan-20S	NA	<294	NA	<5700	NA	<40	NA	NA	NA	18.2	NA	NA	<78						
5a(H),14a(H),17a(H)-Cholestan-20R	38.2	<294	NA	<5700	NA	<40	NA	NA	NA	26.8	NA	NA	<78						
5a(H),14a(H),17a(H)-24-Ethylcholestan-20S	29.0	<294	NA	<5700	NA	<40	NA	NA	NA	36.8	Other	47.1	NA						
5a(H),14a(H),17a(H)-24-Ethylcholestan-20R	29.0	<294	NA	<5700	NA	<40	NA	NA	NA	16.8	Other	12.4	NA						
5a(H),14a(H),17a(H)-24-Ethylcholestan-20R	<20	<294	NA	<5700	NA	<40	NA	NA	NA	31.0	NA	NA	<78						
5a(H),14b(H),17b(H)-Cholestan-20R	<20	<294	NA	<5700	NA	<40	NA	NA	NA	20.0	Other	17.1	NA						
5a(H),14b(H),17b(H)-Cholestan-20S	NA	<294	NA	<5700	NA	<40	NA	NA	NA	18.5	NA	NA	<78						
5a(H),14b(H),17b(H)-24-Ethylcholestan-20R	<20	<294	NA	<5700	NA	<40	NA	NA	NA	8.75	Other	19.8	NA						
5a(H),14b(H),17b(H)-24-Ethylcholestan-20S	NA	<294	NA	<5700	NA	<40	NA	NA	NA	20.0	Other	NA	NA						
NA = Not analyzed; DL = Detection limit; No value in space=nothing reported by lab; Bolded values were not used in the calculation of the consensus statistics																			

Table 3 (cont.). Mussel Tissue (QA101HS01): Laboratory means of three replicates and exercise assigned values - Biomarkers (ng/g dry mass)

		Biomarkers (ng/g dry mass)																	
		(reported as if three figures were significant)																	
Laboratory No.	20	21	22	23	24	24b	25	26	27	28	29	30	31	32	33	34	mean	std dev	median
Carbazole	NA	<10	NA	NA	NA	NA	<none	<1.7	NA	NA	NA	NA	NA	NA	NA	NA	6.85	No Target	11.5
18a(H)-22,29,30-Trisnorbornane	131	NA	NA	NA	NA	NA	<none	NA	NA	NA	NA	<3.0	NA	NA	NA	NA	19.8	No Target	19.8
17a(H)-22,29,30-Trisnorbornane	79.6	NA	23.2	NA	NA	NA	<none	NA	NA	NA	NA	<3.0	NA	NA	NA	NA	20.8	No Target	22.0
17a(H)-2,3[β]H-30-Norbornane	<7.4	NA	60.2	NA	NA	NA	<none	NA	NA	NA	NA	412	NA	NA	NA	NA	51.2	No Target	51.2
18a(H)-30-Norbornane	<7.4	NA	NA	NA	NA	NA	<none	NA	NA	NA	NA	86.7	NA	NA	NA	NA	18.1	No Target	48.2
17a(H)-Dihydropane	<7.4	NA	NA	NA	NA	NA	<none	NA	NA	NA	NA	<3.0	NA	NA	NA	NA	5.38	No Target	2.74
17a(H)-2,3[β]H-Hopane	<7.4	NA	68.8	NA	NA	NA	<none	81.1	NA	NA	NA	<3.0	NA	NA	NA	NA	78.9	69.9	8.3
17a(H)-2,3[β]H-22R-Homohopane	<7.4	NA	23.8	NA	NA	NA	<none	NA	NA	NA	NA	<3.0	NA	NA	NA	NA	20.5	18.8	4.8
17a(H)-2,3[β]H-22S-Homohopane	<7.4	NA	29.4	NA	NA	NA	<none	NA	NA	NA	NA	<3.0	NA	NA	NA	NA	24.4	24.5	3.2
18a(H)-24-Ethylcholestan-20S	50.6	NA	NA	NA	NA	NA	<none	NA	NA	NA	NA	<3.0	NA	NA	NA	NA	17.7	No Target	18.2
5a(H),14a(H),17a(H)-24-Ethylcholestan-20S	<7.4	NA	NA	NA	NA	NA	<none	NA	NA	NA	NA	<3.0	NA	NA	NA	NA	29.1	No Target	28.0
5a(H),14a(H),17a(H)-24-Ethylcholestan-20R	<7.4	NA	36.4	NA	NA	NA	<none	NA	NA	NA	NA	<3.0	NA	NA	NA	NA	52.4	42.2	7.2
5a(H),14a(H),17a(H)-24-Ethylcholestan-20S	21.0	NA	NA	NA	NA	NA	<none	NA	NA	NA	NA	<3.0	NA	NA	NA	NA	11.3	15.4	4.4
5a(H),14a(H),17a(H)-24-Ethylcholestan-20R	66.2	NA	27.6	NA	NA	NA	<none	NA	NA	NA	NA	24.7	NA	NA	NA	NA	32.4	37.2	16.3
5a(H),14b(H),17b(H)-Cholestan-20R	<7.4	NA	<30	NA	NA	NA	<none	NA	NA	NA	NA	67.3	NA	NA	NA	NA	17.7	18.3	1.5
5a(H),14b(H),17b(H)-Cholestan-20S	<7.4	NA	NA	NA	NA	NA	<none	NA	NA	NA	NA	59.7	NA	NA	NA	NA	18.8	No Target	18.8
5a(H),14b(H),17b(H)-24-Ethylcholestan-20R	<7.4	NA	13.0	NA	NA	NA	<none	NA	NA	NA	NA	<3.0	NA	NA	NA	NA	23.7	16.3	6.7
5a(H),14b(H),17b(H)-24-Ethylcholestan-20S	<7.4	NA	NA	NA	NA	NA	<none	NA	NA	NA	NA	<3.0	NA	NA	NA	NA	18.1	No Target	19.1

NA = Not analyzed; DL = Detection limit; No value in space=nothing reported by lab; Bolded values were not used in the calculation of the consensus statistics (Lab 31 reported values on a wet mass basis)

Table 4. Mussel Tissue (QA1011S0): α scores (25% by laboratory)- Water and PAHs

Percent Water	Laboratory No.	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18
naphthalene		2.3	1.5	-1.1	-2.1	11.3	1.3	0.6	-1.3	-0.4	4.0	-47.4	1.1	87.9	0.4				
biphenyl		-0.7		28.8	0.0					-1.7	1.3		0.6	-0.6	1.388	0.8			
acenaphthene		1.7		0.6	-0.1					-1.0			0.8		-0.1				
acenaphthylene				0.1	-2.8					-2.0			-0.4	1.7	0.8		-1.0		
fluorene		1.8	-1.6	0.2	0.4	3.5	-0.6	0.3	-1.0	-0.6	-1.5	1.7	1.3	0.3	0.1	34.8	-0.5		
phenanthrene		-0.1	-2.1	-0.7	-0.8	0.9	1.1	0.6	0.4	0.8	-0.2	-0.1	0.2	-0.2	0.2	1.2	21.1	-0.1	
anthracene		-0.7	-3.1	0.2	-0.8	-0.2			-0.8	-0.4	-0.8	-0.9	-1.6	1.6	1.0	22.8	1.5		
fluoranthene		0.5	-1.9	-0.7	-1.4	1.6	0.5	-0.4	0.4	0.8	-0.6	0.1	-0.1	-0.2	-0.8	1.3	17.3	1.5	
pyrene		1.7	-1.7	-1.0	-1.1	1.8	1.2	-0.7	0.2	1.3	-1.2	-0.2	0.1	0.1	0.3	-0.7	1.1	16.0	
benz[b]fluorene				-0.7								-1.0						-0.2	
benz[a]anthracene		0.1	-3.0			-1.8	2.3	1.5	-2.2	-0.9	-0.1	-0.5	-0.6	0.2	-1.1	-1.1	-0.1	17.7	
ethynene																		1.8	
triphenylene																			
chrysene/triphenylene		1.6				1.4										0.2	12.4		
benz[b]fluoranthene		-0.8	-2.2		-1.4	0.7	-0.3	-1.3	-0.6	-0.2	1.4	-0.9	0.3	-0.2	0.3	0.3	28.2	0.0	
benzo[f]fluoranthene																			
benzo[k]fluoranthene		0.0	-3.1			-0.6		0.4	1.3	-0.9	0.4	2.7	2.5	-0.2	0.0	-1.1		-0.1	
benzo[b,f]fluoranthene																			
benzo[<i>g,h</i>]fluoranthene																0.5	23.3		
benzo[<i>a,h</i>]fluoranthene																			
benzo[<i>e,f</i>]fluoranthene																2.8			
benzo[e]phenylene		0.0				-1.5	1.1		-0.2	0.1		-1.0	0.1	-0.3	0.8	20.0	0.1		
benzo[f]phenylene		0.5	-2.8			1.7	0.3	0.6		-1.2	1.7	-0.4	-1.6	-0.4	-0.6	-0.7	0.6	1.2	
phenylene		-0.6			125.4	-2.0	-0.1					-1.3	0.0	-0.8	0.4			-1.2	
indeno[1,2,3-cd]phenylene		-1.6				6.6	0.2		-1.1		0.7	-1.8	-1.7	0.1	-0.2	0.89	-0.2		
benzo[en]phenylene		-0.5				-1.0	0.8	-2.3		-0.4		-0.5	1.3	0.4	0.4	87.4	1.9		
dibenz[a,b]anthracene		-2.5									20.2	-1.0	-1.2	-1.2	-0.6	229.2	0.2		
dibenz[a,b]anthracene																			
cis/trans-decalin																			
dibenzofuran																0.2	28.9		
retene																2.5	24.5		
benzothiophene																			
dibenzothiophene																			
naphthobenzothiophene																			
naphthobenzothiophene																			

Table 4 (cont). Mussel Tissue (Q_{A10}TIS0): *t* scores (25% by laboratory)- Water and PAHs
(*z*=+1 is 25% higher than the exercise assigned value; *z*=-1 is 25% lower than the exercise assigned value.)

Percent Water	Laboratory No.	19	20	21	22	23	24a	24b	25	26	27	28	29	30	31	32	33	34
naphthalene		1.4	-0.6	-0.4	-0.1	2.9	13.2	-0.4	-1.8	0.6	-1.8	-3.2	-0.7	0.2				
biphenyl		4.5	-1.2	0.0	-0.1	-0.1	-0.8						-2.3		-1.0			
acenaphthene			-1.2	0.3	-0.2		0.6	-1.0					-1.2	0.8		0.3		
acenaphthylene		2.2	-2.0	-0.5	3.2		0.5	1.6					-1.2			-0.5		
fluorene		0.0	-0.5	-0.8	-0.4		-0.8	-0.7	-0.8	1.0	-0.8		0.3	-0.3				
phenanthrene		-0.7	0.4	-0.2	0.0	-0.1	-0.1	-0.9	0.2	-0.4	0.0	0.5	-2.1	-3.4	-3.0	0.8	0.9	
anthracene		5.0	0.1	3.7		-2.1	17.5	-0.4	1.1				-1.2	0.3	-1.4	0.0		
fluoranthene		-0.2	-1.7	0.4	-0.3	0.1	0.0	-1.9	0.2	0.4	0.3	0.0	-0.8	-3.6	-3.0	1.1	0.8	
pyrene		-0.3	0.9	-0.8	0.2	0.1	1.0	-2.0	0.3	-0.3	-1.0	0.2	-1.9	-3.5	-3.0	1.3	0.8	
benzo[b]fluoranthene			3.0	-0.3	-1.0	-0.8		1.5								-1.7		
benz[a]anthracene																2.1	-0.2	
ethylene																		
triphenylene																		
ethylenetriphenylene																		
benzo[<i>b</i>]fluoranthene		0.4	2.9	-0.9	0.6	2.0	-1.4	0.6	1.2	-1.2	-0.5	-1.3			3.1	-0.2		
benzo[<i>f</i>]fluoranthene																		
benzo[<i>k</i>]fluoranthene		12.6	-0.4			-0.8		-2.0					3.6	-0.4	-3.2	0.9		
benzo[<i>b-f</i>]fluoranthene																		
benzo[<i>f+k</i>]fluoranthene																		
benzo[<i>b+f+k</i>]fluoranthene																		
benzo[d]fluoranthene																		
benzo[<i>c</i>]phenene		3.1		-0.1	-0.2			-0.1	0.2		0.5					-1.5		
benzo[<i>a</i>]phenene		5.6	0.2	-0.6	3.6	-1.7		0.1	-1.3	-0.3	1.3	-1.9	4.4		0.4	0.9		
perylene		123.8		-0.6	3.0						0.8	5.0			-0.1			
indenof[1,2,3- <i>cde</i>]pyrene		-1.5	2.6	-0.7	-0.8	3.0		-0.5	1.2		-0.8			4.4	-0.4			
benzo[<i>ghi</i>]perylene			0.9	-0.1	-0.5			-0.1	-0.7	0.7	-1.1			0.8				
dibenz[<i>a,h</i>]anthracene		2.8		-1.6				6.4						0.3				
dibenz[<i>a,h-i</i>]anthracene				-1.1											1.5			
cis/trans-decalin																		
dibenzofuran																-0.4		
retene								-0.4							-1.0			
benzodioxepine																		
naphthobenzoethiophene								-0.4	-0.3	0.0			0.2	0.5	0.7	0.7		
													-0.4	1.8		1.2		

Lab 31 reported values on a wet mass basis

Table 5. Mussel Tissue (QAT0T1S0): z scores (25% by laboratory)-Alkylated PAHs

		z=-1 is 25% higher than the exercise assigned value; z=-1 is 25% lower than the exercise assigned value.)																	
		Laboratory No.																	
		1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18
1-methylnaphthalene	-0.6		-2.1	-1.3		10.2	2.4			-0.5	-1.1	2.4	-0.2	0.8	1.1		-1.5		
2-methylnaphthalene	-0.7		16.7	-1.8	-0.6		1.9	-1.1		0.0	-0.5	4.3	0.4	2.2	1.5	55.9	-1.1		
2,6-dimethylnaphthalene	-0.2		1.8	-1.6	-0.1		0.0		-1.3	-0.3	1.7	-1.7		1.1	56.1	-1.4			
1,6,7-trimethylnaphthalene	-1.1		3.0	-1.1	-1.5				-0.8	0.7				0.8	16.4	-1.4			
1-methylphenanthrene	-0.3		-2.4	1.5		0.2		-0.1	0.8		1.3			0.6	164.6	0.4			
C1-decalins			-1.3												1.7	13.0			
C2-decalins			-0.7												1.2	30.7			
C3-decalins			-0.1												-0.4	39.9			
C4-decalins			-0.1												0.2	34.2			
C1-naphthalenes	0.3		36.2	-2.4	-0.1										0.4	54.2			
C2-naphthalenes	0.4		2.5	-2.6	0.1										1.4	-0.7			
C3-naphthalenes	2.5		0.1	-2.4	2.3										0.3	40.3			
C4-naphthalenes	-0.4		0.3	-2.8	0.3										-0.6	-1.4	34.5		
C1-benzothiophenes																-0.1			
C2-benzothiophenes																2.2			
C3-benzothiophenes															1.3				
C4-benzothiophenes															-0.4				
C1-thiophenes	0.3		-2.1	-0.2											0.2	0.1	44.0		
C2-thiophenes	0.1		-1.6	0.2											0.3	0.7	54.0		
C3-thiophenes	-0.4		1.8	-0.8											1.3	-0.3			
C1-phenanthrenes/anthracenes	-0.6		-2.0	1.1											1.0	-0.1			
C2-phenanthrenes/anthracenes	0.8		2.1	-1.6	1.8										2.4	0.7	17.7		
C3-phenanthrenes/anthracenes	0.1		-1.2	0.4											5.1	-0.8	24.5		
C4-phenanthrenes/anthracenes	-1.6		-0.3	-0.6											5.7	-1.9	18.4		
C1-dibenzothiophenes	0.4		-1.4	0.2											-1.0	0.6	13.7		
C2-dibenzothiophenes	0.5		-1.3	0.6											-0.4	1.1	13.9		
C3-dibenzothiophenes	0.5		-1.1	0.2											0.8	1.2	18.7		
C4-dibenzothiophenes	-0.1		-0.9	-0.4											-1.6	-0.1	26.9		
C1-fluoranthenes/pyrenes	-0.4		-2.2	-1.1											-1.7	-2.3	-0.6		
C2-fluoranthenes/pyrenes	-0.6		5.0	-1.7	-0.3										-1.0	-1.7	-0.4		
C3-fluoranthenes/pyrenes	-0.9		-1.0	-0.5											-1.7	-0.2			
C4-fluoranthenes/pyrenes			-0.9	-1.0															
C1-naphthobenzothiophenes			4.8	-0.8													-2.1		
C2-naphthobenzothiophenes			0.1													-1.4			
C3-naphthobenzothiophenes			-0.6													-0.3			
C4-naphthobenzothiophenes			0.0													-1.3			
C1-chrysenes			-0.1													0.9			
C2-chrysenes			0.3													-1.3			
C3-chrysenes			0.3																
C4-chrysenes																			

Table 5 (cont.). Mussel Tissue (QMA01TS01): z scores (25% by laboratory)-Alkylated PAHs

	Laboratory No.	19	20	21	22	23	24a	24b	25	26	27	28	29	30	31	32	33	34
1-methylnaphthalene		0.7	-0.8	-0.9	-0.2	3.4			-0.2		0.8	-0.1	-2.0	-3.0	2.0	0.3		
2-methylnaphthalene		-0.8	-0.1	-0.3	0.1	4.6			-0.2		-1.8	0.8	-2.5	-3.4	-1.2	0.8		
2,6-dimethylnaphthalene		2.0	-1.6	0.3	-1.0	2.7		1.9	0.5	-1.3	-0.1	2.5	-3.3	0.9	0.1			
1,6,7-trimethylnaphthalene		6.6			-1.4				1.1	-2.1		-1.1		-2.9			-1.8	
1-methylphenanthrene		-0.1	-0.1	0.0					-1.0	-1.6	1.7						-1.1	
C1-decalins									-0.2	-3.8							-0.2	
C2-decalins									0.2	-3.9							-0.7	
C3-decalins									2.1	-3.4							-1.7	
C4-decalins									1.5	-3.2							-1.7	
C1-naphthalenes		-0.8	0.3	0.4	1.1				-0.7	0.2	2.1		-3.1			1.2	-0.3	
C2-naphthalenes		-1.9	-0.3	0.8	1.2	-1.0	4.8		-0.9	-2.2	0.5	0.0	-3.6			1.5	-1.6	
C3-naphthalenes		-1.9	-0.7	0.0	2.4	-0.8	1.2		-1.0	-1.7	1.2	3.2	-3.7			0.5	-1.6	
C4-naphthalenes		-2.4	-1.7	-2.3	0.0	-1.6	1.0		-1.1	-2.0	-3.0		-3.9			-3.4	-1.7	
C1-benzothiophenes										0.8							-0.7	
C2-benzothiophenes																	-0.5	
C3-benzothiophenes																	0.7	
C4-benzothiophenes									0.7	-0.7							0.4	
C1-fluorenes		0.8	3.8	-0.2	1.5				-0.2	-0.1	-2.8	21.0				1.8	-1.1	
C2-fluorenes		-0.7	0.4	3.1	-0.7	0.4			4.5	0.3	0.1	10.9				-2.6	-2.3	-0.6
C3-fluorenes		-1.4	-3.4	2.5	-1.0				5.5	0.3		2.8				-3.2	-0.6	-1.2
C1-phenanthrenes/anthracenes		-1.7	-0.2	0.0	-0.9	4.3			-1.7	-1.2	0.4	2.7				-3.3	1.2	-0.9
C2-phenanthrenes/anthracenes		-1.2	-3.0	1.0	-0.5	9.4			-3.0	0.5	-0.4	4.6				-3.1	1.9	-0.3
C3-phenanthrenes/anthracenes		-1.7	-2.6	0.8	-1.4	0.1			0.0	-1.3	-0.9	3.3				-3.3	0.2	-1.9
C4-phenanthrenes/anthracenes		-2.1			5.6	-1.7	-3.9		0.2	-1.4	2.9					-0.2	-2.5	
C1-dibenzothiophenes			-1.9		0.8	-0.1			-0.2	-0.7	4.7						-0.7	
C2-dibenzothiophenes		-2.0			2.7	0.1			1.4	0.3	8.3					-3.0		-0.2
C3-dibenzothiophenes		-1.8	-2.7		2.5	-0.1			2.2	0.8	10.4						-0.2	
C4-dibenzothiophenes		-2.0			0.7				5.3	-0.2		11.3					-0.6	
C1-difluoranthenes/pyrenes		-1.5	5.4	4.5	5.4	-0.9			-0.8	-1.1		-1.2				-1.3		
C2-difluoranthenes/pyrenes			0.9	11.7	-0.8				1.0	-0.1	0.5					-0.8	-0.8	
C3-difluoranthenes/pyrenes			1.4		5.3	-1.5			0.9	-0.6	-0.2					-1.2		
C4-difluoranthenes/pyrenes					1.5				0.1	0.5						-0.3		
C1-naphthobenzothiophenes												-1.0					-1.0	
C2-naphthobenzothiophenes												0.7					0.7	
C3-naphthobenzothiophenes																	0.8	
C4-naphthobenzothiophenes																		
C1-chrysenes			-0.6		1.8	0.2			0.1	-0.1		0.2				-0.3		
C2-chrysenes		28.0		0.7	-1.0				2.6	-0.6		-0.7				-0.4		
C3-chrysenes					-1.1	-1.0						1.3						
C4-chrysenes																		

Lab 31 reported values on a wet mass basis

Table 6. Mussel Tissue (Q10T1S01): α scores (25% by laboratory)-Biomarkers ($\alpha=1$ is 25% higher than the exercise assigned value; $\alpha=-1$ is 25% lower than the exercise assigned value.)

Laboratory No	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18
Carbazole																		
5a(H)-22,29,30-Triisnorhopane																		
17a(H)-22,29,30-Triisnorhopane																		
17b(H),21[β(H)-30-Norhopane																		
18a(H)-30-Norneohopane																		
17a(H)-Diaphopane																		
17a(H),21[β(H)-Homopane	-0.2																	
17a(H),21[β(H)-22R-Homohopane																		
17a(H),21[β(H)-22S-Homohopane	-0.2																	
13b(H)17a(H)-Diacholestan-20S																		
5a(H),14a(H),17a(H)-Cholestan-20S																		
5a(H),14a(H),17a(H),17a(H)-Cholestan-20R	-0.4																	
5a(H),14a(H),17a(H)-24-Ethylcholestan-20S																		
5a(H),14a(H),17a(H)-24-Ethylcholestan-20R	-0.7																	
5a(H),14b(H),17a(H)-Cholestan-20S																		
5a(H),14b(H),17b(H)-C-Cholestan-20S																		
5a(H),14b(H),17b(H),17b(H)-C-Cholestan-20S																		
5a(H),14b(H),17b(H)-24-Ethylcholestan-20R																		
5a(H),14b(H),17b(H)-24-Ethylcholestan-20S																		

Table 6 (cont). Mussel Tissue (Q10T1S01): α scores (25% by laboratory)-Biomarkers ($\alpha=1$ is 25% higher than the exercise assigned value; $\alpha=-1$ is 25% lower than the exercise assigned value.)

Laboratory No	19	20	21	22	23	24a	24b	25	26	27	28	29	30	31	32	33	34
Carbazole																	
17a(H)-22,29,30-Triisnorhopane																	
17a(H)-22,29,30-Triisnorhopane																	
17a(H)-30-Norhopane																	
17a(H)-Diaphopane																	
17a(H)-21[β(H)-Homopane																	
17a(H),21[β(H)-22R-Homopane																	
17a(H),21[β(H)-22S-Homopane																	
13b(H)17a(H)-Diacholestan-20S																	
5a(H),14a(H),17a(H)-Cholestan-20S																	
5a(H),14a(H),17a(H)-24-Ethylcholestan-20R	-0.1																
5a(H),14a(H),17a(H),17a(H)-Cholestan-20S																	
5a(H),14b(H),17b(H)-C-Cholestan-20S																	
5a(H),14b(H),17b(H)-24-Ethylcholestan-20R																	
5a(H),14b(H),17b(H)-24-Ethylcholestan-20S																	
5a(H),14b(H),17b(H)-24-Ethylcholestan-20R																	
5a(H),14b(H),17b(H)-24-Ethylcholestan-20S																	

Lab 31 reported values on a wet mass basis

Table 7. Mussel Tissue (QA1011S01); Laboratory relative standard deviations of three replicates - Water and PAHs

Percent Water	Laboratory No.	%															
		1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16
naphthalene		7.2%	36.5%	21.2%	4.3%	2.9%	16.7%	16.7%	26.7%	22.4%	11.7%	5.5%	29.8%	13.4%	24.5%		
biphenyl	7.8%	65.6%	16.5%	10.2%	9.8%					7.2%	9.0%	2.4%	13.7%	14.8%	31.7%		
acenaphthene	4.2%										8.4%	2.0%	6.4%	4.1%	1.2%		
acenaphthylene		15.7%	23.3%														
fluorene	6.1%	18.6%	12.6%	1.4%	8.1%	4.2%	5.4%	9.6%	7.4%	9.8%	4.6%	4.6%	6.0%	1.5%	1.1%	2.0%	
phenanthrene	2.7%	1.3%	24.1%	15.2%	2.1%	4.7%	5.0%	3.8%	12.7%	4.4%	11.9%	3.6%	4.4%	12.2%	1.1%	1.9%	9.3%
anthracene	4.0%	78.2%	22.1%	3.2%	3.4%					3.9%	11.9%	24.0%	3.8%	3.3%	13.6%	6.3%	
fluoranthene	2.2%	9.2%	19.5%	14.3%	2.6%	3.7%	3.8%	3.4%	16.9%	3.1%	5.6%	11.4%	2.2%	16.4%	1.1%	1.2%	7.8%
pyrene	3.6%	3.6%	7.6%	14.3%	2.6%	3.1%	4.1%	3.2%	25.2%	8.5%	3.3%	12.5%	2.7%	3.2%	13.8%	1.0%	1.4%
benzo[b]fluorene				8.8%													
benz[a]anthracene	7.4%	14.9%		17.9%	3.5%	5.0%	1.1%	4.2%			12.0%	8.0%	8.8%	0.4%	12.1%	8.9%	31.7%
chrysene		7.9%		16.0%		3.3%	3.1%	6.2%	12.2%		2.7%	10.7%	5.6%	3.0%	3.6%	15.2%	
triptycene																	
benzo[b]fluoranthene	3.1%	5.0%		14.8%	3.0%	2.4%	4.6%	5.1%			4.7%	11.4%	5.8%	3.9%	3.9%	13.6%	37.4%
benzo[f]fluoranthene	7.7%																
benzo[k]fluoranthene	10.0%	5.1%			20.4%		8.3%	3.0%	11.2%	16.7%	16.3%	18.2%	7.5%	5.0%	0.0%		
benzo[4]fluoranthene						173.2%											
benzo[e]pyrene	5.4%					16.6%	3.6%	0.9%	6.3%		3.5%	19.8%	14.5%	9.0%	2.0%	12.2%	8.9%
benzo[a]pyrene						10.6%	12.0%	1.5%	4.1%		5.5%	36.3%	6.0%	6.0%	5.4%	14.3%	
perylene						6.1%		22.4%	3.6%				13.4%	12.8%	1.4%	14.4%	
indeno[1,2,3-ed]pyrene						10.6%	7.2%	1.4%		12.3%		4.2%	17.2%	35.2%		9.8%	
benzo[ghi]perylene	6.5%						23.4%	2.8%	25.5%	5.9%			10.8%	23.2%		1.5%	
dibenz[a,h]anthracene						11.6%						8.4%	11.9%	13.8%		19.7%	
cis/trans-decalin									12.9%					3.5%		0.2%	
dibenzofuran									16.5%				11.4%	3.4%		1.1%	2.5%
retene									17.1%	3.4%			10.5%	2.6%	10.0%	0.6%	22.1%
benzothiophene															17.8%		
dibenzothiophene									12.1%	4.8%				12.1%	5.5%	14.4%	1.2%
naphthobenzothiophene									4.9%					10.5%		13.1%	

Table 7 (cont.). Mussel Tissue (QAT01TS01): Laboratory relative standard deviations of three replicates - Water and Paths

Percent Water	Laboratory No.	18	19	20	21	22	23	24a	24b	25	26	27	28	29	30	31	32	33	34
		0.7%	0.1%	0.2%	0.0%	0.8%	0.6%	1.5%	1.5%	0.0%	0.0%	0.3%	0.8%	0.3%	0.1%	0.7%	0.41%	0.2%	
naphthalene	28.9%	9.8%	8.9%	6.2%	2.7%	43.6%	7.4%	14.6%	6.4%	10.2%	6.3%	5.8%	5.0%	5.8%	5.0%	5.8%	5.0%	5.0%	
benzylbenzene	6.6%	10.9%	4.2%	5.8%	5.8%	1.3%	1.3%	1.3%	1.3%	1.0%	1.0%	1.0%	1.0%	1.0%	1.0%	1.0%	1.0%	1.0%	
acenaphthene	15.1%	2.1%	5.9%	8.0%	11.6%	1.7%	1.0%	1.0%	1.0%	4.5%	4.5%	34.1%	34.1%	34.1%	34.1%	34.1%	34.1%	34.1%	
acenaphthylene	29.7%	24.7%	17.5%	11.6%	3.7%	3.2%	4.3%	9.1%	0.6%	2.0%	2.0%	2.0%	2.0%	2.0%	2.0%	2.0%	2.0%	2.0%	
fluorene	1.8%	49.1%	3.7%	3.4%	3.4%	3.2%	4.3%	7.5%	7.5%	11.0%	11.0%	17.4%	17.4%	17.4%	17.4%	17.4%	17.4%	17.4%	
phenanthrene	0.6%	19.3%	7.9%	5.6%	2.9%	7.0%	33.5%	5.2%	5.2%	5.9%	5.9%	18.8%	18.8%	18.8%	18.8%	18.8%	18.8%	18.8%	
anthracene	6.7%	69.7%	6.8%	4.1%	4.1%	4.0%	10.6%	4.9%	4.9%	5.9%	5.9%	11.2%	11.2%	11.2%	11.2%	11.2%	11.2%	11.2%	
fluoranthene	7.6%	2.8%	17.4%	2.4%	7.1%	3.3%	8.1%	47.7%	5.3%	5.4%	10.9%	2.3%	13.3%	13.3%	13.3%	13.3%	13.3%	13.3%	13.3%
pyrene	32.5%	25.1%	13.6%	5.8%	3.9%	8.6%	58.0%	6.8%	6.8%	11.6%	2.3%	71.6%	35.3%	35.3%	35.3%	35.3%	35.3%	35.3%	35.3%
benzo[a]fluorene	30.4%	33.9%	1.0%	5.7%	4.4%	7.7%	6.2%	2.0%	2.0%	9.7%	10.2%	3.8%	10.4%	10.4%	10.4%	10.4%	10.4%	10.4%	10.4%
benz[a]anthracene	7.9%	20.4%	13.8%	18.6%	18.6%	9.3%	37.1%	11.0%	11.0%	15.2%	15.2%	49.4%	49.4%	49.4%	49.4%	49.4%	49.4%	49.4%	
chrysene	7.9%	105.8%	5.9%	5.0%	2.7%	10.2%	20.7%	8.0%	8.0%	4.2%	9.7%	7.5%	15.3%	15.3%	15.3%	15.3%	15.3%	15.3%	15.3%
triphenylene	11.4%	10.5%	10.5%	10.5%	10.5%	10.5%	10.5%	10.5%	10.5%	10.5%	10.5%	10.5%	10.5%	10.5%	10.5%	10.5%	10.5%	10.5%	10.5%
benzo[b]fluoranthene	3.1%	70.0%	8.1%	4.8%	9.1%	9.1%	13.6%	13.6%	13.6%	18.9%	18.9%	3.6%	3.6%	3.6%	3.6%	3.6%	3.6%	3.6%	3.6%
benzo[k]fluoranthene	7.0%	66.2%	2.4%	2.2%	7.9%	5.9%	1.7%	1.7%	1.7%	10.2%	10.2%	10.2%	10.2%	10.2%	10.2%	10.2%	10.2%	10.2%	10.2%
benzo[e]pyrene	6.9%	82.4%	11.7%	9.9%	3.9%	6.2%	5.9%	10.6%	10.6%	7.8%	7.8%	10.2%	10.2%	10.2%	10.2%	10.2%	10.2%	10.2%	10.2%
perylene	7.4%	52.7%	3.9%	5.8%	10.2%	11.6%	40.8%	5.0%	5.0%	4.4%	4.4%	50.6%	50.6%	50.6%	50.6%	50.6%	50.6%	50.6%	50.6%
indenof[1,2- <i>c</i>]pyrene	66.6%	35.0%	10.2%	29.5%	8.1%	8.8%	3.5%	3.5%	3.5%	5.9%	5.9%	13.4%	13.4%	13.4%	13.4%	13.4%	13.4%	13.4%	13.4%
benzo[ghi]perylene	1.5%	18.4%	22.3%	28.9%	28.9%	28.9%	52.5%	#DIV/0!	#DIV/0!	21.7%	20.3%	8.7%	8.7%	8.7%	8.7%	8.7%	8.7%	8.7%	8.7%
dibenzo[a,h]anthracene										7.2%	4.7%						16.6%	7.7%	
cis/trans-decalin																			10.5%
dibenzofuran																			
retene																			
benzo[ghi]phenanthrene																			
dibenzothiophene																			
naphthobenzo[ghi]phenic																			

Table 8. Mussel Tissue (QA1011S01): Laboratory relative standard deviations of three replicates - Alkylated PAHs

	Laboratory No.	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17
1-methylnaphthalene	0.8%	11.5%	0.5%	38.7%	11.5%	8.3%	4.7%	17.0%	5.7%	17.7%	12.1%							
2-methylnaphthalene	2.4%	28.4%	13.6%	1.1%	5.7%	13.1%	4.6%	48.8%	4.9%	18.4%	13.8%							
2,6-dimethylnaphthalene	2.1%	15.6%	13.0%	0.9%	4.8%	8.0%	4.3%	9.5%	6.6%	10.2%	12.8%							
1,6,7-trimethylnaphthalene	0.7%	4.5%	27.0%	10.9%				2.7%		8.8%	4.6%							
1-anethylphenanthrene	5.5%			14.8%	2.6%			1.4%		15.3%	5.9%							
C1-decalins				16.0%														
C2-decalins				11.5%														
C3-decalins				11.5%														
C4-decalins				9.1%														
C1-naphthalenes	1.4%	11.8%	9.8%	0.9%														
C2-naphthalenes	2.9%	4.4%	13.8%	1.3%														
C3-naphthalenes	2.1%	11.0%	13.6%	0.8%														
C4-naphthalenes	3.1%	13.4%	15.6%	1.3%														
C1-benzothiophenes																		
C2-benzothiophenes																		
C3-benzothiophenes																		
C4-benzothiophenes																		
C1-fluorenes	4.1%			15.6%	0.5%													
C2-fluorenes	5.0%			14.1%	1.1%													
C3-fluorenes	9.1%			1.2%	3.3%													
C1-phenanthrenes/anthracenes	6.0%			18.7%	3.3%													
C2-phenanthrenes/anthracenes	4.8%			16.0%	1.9%													
C3-phenanthrenes/anthracenes	2.2%			17.6%	2.2%													
C4-phenanthrenes/anthracenes	7.0%			15.7%	0.6%													
C1-dihydroethiophenes				13.6%	2.0%													
C2-dihydroethiophenes				7.1%	1.7%													
C3-dihydroethiophenes				14.5%	2.8%													
C4-dihydroethiophenes				16.1%	4.1%													
C1-thiophenenes/pyrenes	2.5%			14.3%	1.5%													
C2-thiophenenes/pyrenes	5.4%			13.4%	18.8%	0.2%												
C3-thiophenenes/pyrenes	4.9%			14.9%	1.6%													
C4-thiophenenes/pyrenes					19.2%	3.1%												
C1-naphthobenzothiophenes					12.0%	3.6%												
C2-naphthobenzothiophenes																		
C3-naphthobenzothiophenes																		
C4-naphthobenzothiophenes																		
C1-chrysenes	7.6%																	
C2-chrysenes																		
C3-chrysenes																		
C4-chrysenes																		

Table 8 (cont.). Mussel Tissue (Q1101TS01): Laboratory relative standard deviations of three replicates - Alkylated PAHs

Laboratory No.	18	19	20	21	22	23	24a	24b	25	26	27	28	29	30	31	32	33	34
1-methylnaphthalene	6.6%		45.5%	5.8%	17.2%	5.9%	31.5%				4.3%	9.5%	13.8%	2.5%	6.5%	6.0%		
2-methylnaphthalene	12.8%		23.6%	9.8%	11.0%	4.5%	26.9%				10.1%	6.1%	2.8%	14.0%	17.1%	4.6%	8.3%	
2,6-dimethylnaphthalene	9.9%		12.6%	5.3%	3.5%	7.1%	39.6%				8.7%	4.4%		21.4%	75.4%	6.8%	5.5%	
1,6,7-trimethylnaphthalene	11.2%		23.1%			5.4%					6.1%							21.0%
1-methylphenanthrene	9.0%		6.7%	10.0%	2.5%						4.8%	6.6%						1.9%
C1-decalins																		3.0%
C2-decalins																		1.5%
C3-decalins																		5.4%
C4-decalins																		16.7%
C1-naphthalenes		24.8%	8.4%	13.4%	5.1%						12.6%		4.9%		2.5%			
C2-naphthalenes		4.7%	21.2%	5.8%	2.0%	33.5%					5.6%	9.6%	3.3%		0.0%	5.7%		1.7%
C3-naphthalenes		0.0%	15.4%	14.2%	13.2%	2.3%					6.0%	6.2%	3.2%		24.0%	8.0%	7.1%	
C4-naphthalenes		6.7%	8.7%	8.7%	15.4%	2.5%	19.7%				7.5%	5.3%	5.4%		23.1%	7.4%	10.6%	11.3%
C1-benzothiophenes																		1.9%
C2-benzothiophenes																		12.9%
C3-benzothiophenes																		1.5%
C4-benzothiophenes																		15.1%
C1-phenanthrenes																		8.6%
C2-phenanthrenes																		5.2%
C3-phenanthrenes																		8.6%
C4-phenanthrenes																		8.8%
C1-fluoranthrenes																		3.7%
C2-fluoranthrenes																		2.6%
C3-fluoranthrenes																		3.7%
C1-phenanthrenes/anthracenes																		4.1%
C2-phenanthrenes/anthracenes																		2.7%
C3-phenanthrenes/anthracenes																		21.1%
C4-phenanthrenes/anthracenes																		7.5%
C1-phenanthrenes/anthracenes																		2.8%
C2-phenanthrenes/anthracenes																		6.8%
C3-phenanthrenes/anthracenes																		1.9%
C4-phenanthrenes/anthracenes																		8.9%
C1-fluoranthrenes/anthracenes																		14.5%
C2-fluoranthrenes/anthracenes																		1.9%
C3-fluoranthrenes/anthracenes																		23.6%
C4-fluoranthrenes/anthracenes																		1.8%
C1-phenanthrenes/pyrenes																		2.3%
C2-phenanthrenes/pyrenes																		2.3%
C3-phenanthrenes/pyrenes																		8.5%
C4-phenanthrenes/pyrenes																		5.2%
C1-fluoranthrenes/pyrenes																		8.5%
C2-fluoranthrenes/pyrenes																		7.6%
C3-fluoranthrenes/pyrenes																		3.2%
C4-fluoranthrenes/pyrenes																		1.1%
C1-naphthalenobenzothiophenes																		1.7%
C2-naphthalenobenzothiophenes																		1.7%
C3-naphthalenobenzothiophenes																		5.7%
C4-naphthalenobenzothiophenes																		3.4%
C1-naphthalenobenzothiophenes																		5.6%
C2-naphthalenobenzothiophenes																		5.6%
C3-naphthalenobenzothiophenes																		11.4%
C4-naphthalenobenzothiophenes																		5.5%
C1-naphthalenobenzothiophenes																		6.9%
C2-naphthalenobenzothiophenes																		13.3%
C1-chrysenes																		4.1%
C2-chrysenes																		5.8%
C3-chrysenes																		
C4-chrysenes																		

Table 9. Mussel Tissue (QA1011801): Laboratory relative standard deviations of three replicates - Biomarkers

Carbozole	Laboratory No.	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17
[8a(H)-22,29,30-Trisnorhopane																	1.90%	23.1%
[17a(H)-22,29,30-Trisnorhopane																	6.8%	
[8a(H)-30-Norhopane																		
[17a(H)-Diaphopane																		
[17a(H),2,[3b(H)-Hopane																		
[17a(H),2,[3b(H)-22R-Homohopane																		
[17a(H),2,[3b(H)-Homohopane																		
[17a(H),2,[3b(H)-22S-Homohopane																		
[13b(H),1,7a(H)-Diacholestan-20S																		
[5a(H),1,4a(H),17a(H)-Cholestan-20R																		
[5a(H),1,4a(H),17a(H)-Cholestan-20S																		
[5a(H),1,4a(H),17a(H)-24-Ethylcholestan-20R																		
[5a(H),1,4a(H),17a(H)-24-Ethylcholestan-20S																		
[5a(H),1,4b(H),17b(H)-Cholestan-20R																		
[5a(H),1,4b(H),17b(H)-Cholestan-20S																		
[5a(H),1,4b(H),17b(H)-24-Ethylcholestan-20R																		
[5a(H),1,4b(H),17b(H)-24-Ethylcholestan-20S																		

Table 9 (cont). Mussel Tissue (QA1011801): Laboratory relative standard deviations of three replicates - Biomarkers

Carbozole	Laboratory No.	18	19	20	21	22	23	24a	24b	25	26	27	28	29	30	31	32	33	34
[8a(H)-22,29,30-Trisnorhopane																		2.5%	
[17a(H)-22,29,30-Trisnorhopane																			
[17a(H),2,[3b(H)-30-Norhopane																			
[17a(H)-Diaphopane																			
[17a(H),2,[3b(H)-Homohopane																			
[17a(H),2,[3b(H)-22S-Homohopane																			
[17a(H),2,[3b(H)-Homohopane																			
[17a(H),17a(H)-Cholestan-20S																			
[5a(H),14a(H),17a(H)-Cholestan-20R																			
[5a(H),14a(H),17a(H)-24-Ethylcholestan-20S																			
[5a(H),14b(H),17a(H)-Cholestan-20R																			
[5a(H),14b(H),17a(H)-Cholestan-20S																			
[5a(H),14b(H),17b(H)-Cholestan-20R																			
[5a(H),14b(H),17b(H)-Cholestan-20S																			
[5a(H),14b(H),17b(H)-24-Ethylcholestan-20R																			
[5a(H),14b(H),17b(H)-24-Ethylcholestan-20S																			

Table 10. Mussel Tissue (Q A1011S01); p-scores (15%) - Water and PAHs

Percent Water	Laboratory No.	1	2	3	4	5	6	7	8	9	10	11	n=1	12	13	14	15	16	0.04
naphthalene		0.48	2.43	1.41	0.28	0.19	1.11	1.11			1.78	1.50	0.75	0.37	1.99	0.90	1.63		
benzyl		0.52	4.37	1.10	0.68	0.65					0.88	0.60	0.16	0.92	0.99	2.12			
acenaphthene		0.28									0.56		0.43					0.80	
acenaphthylene													0.13						
fluorene													0.27						0.74
phenanthrene		0.40	1.24	0.84	0.09	0.54	0.28	0.36	0.64	0.49	0.65	0.30	0.31	0.40	1.01				1.37
anthracene		0.18	0.09	1.60	1.01	0.14	0.32	0.33	0.25	0.85	0.29	0.80	0.24	0.29	0.81	0.80	0.62		
fluoranthene		0.27	5.21	1.48	0.21	0.23	0.23	2.09		0.26	0.79	1.63	0.25	0.22	0.91	0.42			
pyrene		0.15	0.61	1.30	0.95	0.18	0.25	0.23	1.10	0.21	0.37	0.76	0.15		1.10	0.75	0.52		
benzobifluorene		0.24	0.24	0.51	0.95	0.18	0.20	0.28	0.21	1.68	0.56	0.22	0.83	0.18	0.21	0.92	0.71	0.76	
benzofluoranthene		0.49	0.99	1.19	0.23	0.34	0.07	0.28	0.80	0.80	0.53	0.59	0.02	0.80	0.59	0.59	0.59	0.59	2.12
chrysene		0.53		1.07		0.22	0.21	0.42	0.82	0.18	0.71	0.37	0.20	0.01					
triphenylene																0.24			
benzo β fluoranthene		0.21	0.33		0.99	0.20	0.16	0.31	0.34	0.80	0.31	0.76	0.39		0.26		0.91	2.49	
benzo γ fluoranthene		0.51																	
benzo δ fluoranthene		0.66	0.34	1.36		0.55	0.20	0.75	1.11	1.09	1.21	0.50	0.33						
benzo α fluoranthene				1.55															
benzo ϵ ethyrene		0.36			1.11	0.24		0.06	0.42										
benzo α pyrene		0.70	0.80		1.86	0.10	0.28	0.37	2.42	0.23	1.32	0.96	0.40	0.36	0.36	0.82	0.59		
perylene		0.40		0.29	1.50	0.24						0.90	0.85	0.10	0.95		0.96		
indenol[1,2,3-cd]pyrene		0.71			0.48	0.10		0.82		0.28	1.15	2.35							0.66
benzof[b]bicylene		0.43			1.56	0.19	1.70	0.40			0.72	1.57							1.04
dbenz[a,h]anthracene		0.77									0.56	0.79	0.92						1.32
cis/trans-decalin					0.86										0.23	0.68			
dbenzofuran						1.10					0.76	0.22				0.75	0.17		
retene					1.14	0.23					0.70		0.17	0.67	0.64	0.64	1.49		
henzoanthophene																1.19			
benzoanthorphen					0.80	0.32					0.81	0.36		0.96	0.82				
naphthobenzothiophene				0.33							0.70		0.01	0.02	0.02	0.16	0.04	0.04	

Table 10 (cont). Mussel Tissue (QAT0TTS01): p scores (15%) - Water and PAHs

Percent Water	Laboratory No.	18	19	20	21	22	23	24a	24b	25	26	27	28	29	30	31	32	33	34
		0.05	0.01	0.01	0.00	0.05	0.04	0.10	0.10	0.00	0.00	0.05	0.14	0.02	0.01	0.04	0.03	0.01	
naphthalene		1.93		0.65	0.59	0.42	0.18	2.91	0.49	0.97	0.42	0.43	0.68	0.42	0.39	0.33			
biphenyl		0.44		0.73		0.28	0.39			0.69	0.35								
acenaphthene		1.01		0.14	0.39	0.54				0.88	0.70								
acenaphthylene		1.98		1.65	1.17	0.78	0.11												
fluorene		0.12		3.27	0.24	0.22	0.21			0.29	0.61								
phenanthrene		0.65	0.04	1.29	0.53	0.38	0.20		0.47	2.24	0.35	0.50	0.73	0.13	1.53		0.46	0.83	
anthracene		0.44		4.65	0.45	0.28			0.27	0.70	0.33	0.40		0.13	1.16	1.25	0.55	0.11	
fluoranthene		0.50	0.19	1.16	0.16	0.47	0.22			0.54	3.18	0.35	0.36	0.73		0.75		0.23	
pyrene		2.16	0.29	1.67	0.90	0.39	0.26		0.57	3.87	0.20	0.46	0.77	0.16	4.77	1.00	0.60	0.13	
benz[b]fluorene		2.03		2.26	0.07	0.38					0.13			0.65				0.26	0.57
benz[a]anthracene		0.53		1.36	0.92					0.51			0.41	0.27		0.68	0.25	0.70	0.48
chrysene		0.53								1.24	0.62	2.47			0.74		1.01	3.29	0.45
triphenylene		0.76		7.65	0.39	0.34	0.18												
benz[b]fluoranthene										0.68	1.38		0.53	0.28		0.65	0.50	1.02	
benz[k]fluoranthene		0.20		4.67	0.54		0.32			0.61									
benz[e]fluoranthene																			
benz[c]fluoranthene																			
benz[c]porene																			
benz[a]porene																			
benz[a]Diprene																			
perylene																			
indenol[2,3-c]pyrene		4.44			2.33	0.68	0.77	2.72		0.33	0.73								
benzo[ghi]perylene		0.10		1.97	0.54	0.58				0.24	0.24						0.89	2.09	
dibenz[a,h]anthracene		1.23		1.49		1.93				3.50								0.58	2.14
cis/trans-decalin																			
dibenzoluran																			
retene																			
benzothiophene																			
dibenzothiophene		0.69				0.52	0.42	0.16						0.24	0.86	0.40			
naphthalobenzothiophene														0.29	0.72				

Table 11. Mussel Tissue (QAT01TS01): p scores (1S%) - Alkylated PAHs

Laboratory No.	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17
1-methylnaphthalene	0.05		0.77	0.04		2.58	0.77		0.55	0.31	1.13	0.38	1.18	0.81			
2-methylnaphthalene	0.16		1.89	0.91	0.07		0.38	0.88		0.54	0.30	3.25	0.32	1.22	0.92	0.85	
2,6-dimethylnaphthalene	0.14		1.04	0.87	0.06		0.32		0.53	0.29	0.63	0.44		0.68		1.01	
1,6,7-trimethylnaphthalene	0.06		0.30	1.80	0.73				0.59	0.31				0.95	0.64		
1-methylphenanthrene	0.37			0.99	0.17		0.18		0.09	1.02		0.39		0.57	1.67		
C1-decalins					1.07									0.72	1.24		
C2-decalins					0.76									0.86	0.64		
C3-decalins					0.61									0.88	1.87		
C4-decalins					0.66									0.75	1.39		
C1-naphthalenes	0.09		0.79	0.65	0.06						0.31			0.86	1.05		
C2-naphthalenes	0.19		0.29	0.92	0.09						0.36			0.46	0.78		
C3-naphthalenes	0.14		0.73	0.91	0.06						0.55			0.40	0.81	1.03	
C4-naphthalenes	0.21		0.89	1.04	0.09						0.56			0.32	0.83	1.15	
C1-benzothiophenes														1.13			
C2-benzothiophenes														0.69			
C3-benzothiophenes														1.43			
C4-benzothiophenes														0.66			
C1-fluorenes	0.28		1.04	0.04										0.13	0.63	0.35	
C2-fluorenes	0.34		0.95	0.07										0.53	0.76	0.73	
C3-fluorenes	0.61		0.08	0.22										0.39	1.49	0.77	
C1-phenanthrenes/anthracenes	0.40		1.25	0.22										0.73	0.51	0.78	
C2-phenanthrenes/anthracenes	0.32		1.06	1.02	0.13						5.36			0.59	0.67	0.31	
C3-phenanthrenes/anthracenes	0.14			1.17	0.15						0.66			0.91	0.76	0.53	
C4-phenanthrenes/anthracenes	0.46		1.05	0.04							0.62			0.17	0.96	1.30	
C1-dibenzothiophenes			0.90	0.13							0.68			0.16	0.87	0.58	
C2-dibenzothiophenes			0.38	0.11							0.68			0.05	0.80	1.39	
C3-dibenzothiophenes			0.97	0.19							0.65			0.24	0.71	0.47	
C4-dibenzothiophenes			1.07	0.27							0.51			0.80	0.55		
C1-fluoranthenes/pyrenes	0.17		0.95	0.10							0.63			0.59	0.71	0.47	
C2-fluoranthenes/pyrenes	0.36		0.89	1.25	0.01						0.81			0.66	0.90		
C3-fluoranthenes/pyrenes	0.33		0.99	0.11							0.59			0.71			
C4-fluoranthenes/pyrenes			1.28	0.21										0.00			
C1-naphthobenzoanthracenes			0.80	0.24										0.87			
C2-naphthobenzoanthracenes				0.17										0.69			
C3-naphthobenzoanthracenes				0.29										1.44			
C4-naphthobenzoanthracenes				0.70										0.58			
C1-chrysenes	0.51		1.05	0.21										0.67	0.89		
C2-chrysenes			1.41	0.11										1.25			
C3-chrysenes			2.14	0.40										0.76			
C4-chrysenes																	

Table 11 (cont.) - Mussel Tissue Q(A)1011S01: p scores (15%) - Alkylated PAHs

	Laboratory No	18	19	20	21	22	23	24a	24b	25	26	27	28	29	30	31	32	33	34
1-methylnaphthalene	0.44	3.04	0.39	1.15	0.39	2.10	0.63	0.29	0.79	0.92	0.17					0.43	0.40		
2-methylnaphthalene	0.35	1.57	0.65	0.73	0.30	1.79		0.67	0.67	0.40	0.18	0.93	1.14		0.31	0.55			
2,6-dimethylnaphthalene	0.66	0.84	0.36	0.23	0.48	2.64		0.53	0.58	0.29	1.43	5.03		0.45	0.37				
1,6,7-trimethylnaphthalene	0.75	1.54			0.36			0.41	0.71	0.72			1.43						
1-methylphenanthrene	0.60		0.45	0.67	0.17			0.32	0.44	0.26									
C1-decalins																			
C2-decalins																			
C3-decalins																			
C4-decalins																			
C1-naphthalenes																			
C2-naphthalenes	0.31	1.41	0.39	0.38	0.14	2.23		0.84	0.33	0.17					0.38	0.12			
C3-naphthalenes		1.03	0.95	0.88	0.15	0.85		0.40	0.41	2.17	0.36			1.60	0.53	0.47			
C4-naphthalenes	0.45	0.58	0.58	1.03	0.16	1.31		0.50	0.35	5.06	1.38	0.92	0.71	0.75	1.03	0.86			
C1-benzothiophenes															0.00		1.01		
C2-benzothiophenes															1.06				
C3-benzothiophenes															2.22				
C4-benzothiophenes															0.57				
C1-fluorenes																	0.58		
C2-fluorenes	0.52	0.67	0.72	0.23	3.32			0.31	0.57	1.43									
C3-fluorenes	0.18	1.33	0.95	0.07	1.32			0.27	0.52	1.39	0.16								
C4-fluorenes	0.35	0.22	0.51	0.16	1.29			1.28	0.46	0.33					1.41	0.50	0.18		
C1-phenanthrenes/anthracenes	0.13	0.30	0.38	0.11	1.47			4.85	0.43	0.98	0.08				0.45	1.05	0.19		
C2-phenanthrenes/anthracenes	0.29	0.19	0.95	0.25	1.11			2.74	0.28	0.50	1.36				0.59	0.96	0.13		
C3-phenanthrenes/anthracenes	0.33			1.07	0.39	2.76		0.25	0.48	0.82	0.41				1.58	0.12	0.15		
C4-phenanthrenes/anthracenes				1.08	0.44	0.20		0.26	0.63	0.28					0.56	0.35			
C1-albenzoanthophenes															0.33	0.44	0.05		
C2-albenzoanthophenes	0.36														0.94	0.49	0.18		
C3-albenzoanthophenes															0.34	0.53	0.31		
C4-albenzoanthophenes															0.15	0.75	0.15		
C1-fluoranthenes/pyrenes	0.16	2.78	0.75	1.07	0.34			0.25	0.48	0.82	0.41				0.59	0.38	0.38		
C2-fluoranthenes/pyrenes			1.02	0.55	0.28			0.26	0.63	0.28					0.54	0.35	0.20		
C3-fluoranthenes/pyrenes	n=1				0.10	0.19		0.51	0.43	0.66					0.51	0.37	0.37		
C4-fluoranthenes/pyrenes						3.04		0.54	0.54	0.54	0.47				0.75	0.49	0.76		
C1-naphthobenzoanthophenes															0.68				
C2-naphthobenzoanthophenes															0.11				
C3-naphthobenzoanthophenes															0.31	0.20	0.14		
C4-naphthobenzoanthophenes															0.35	0.44	0.27		
C1-ethryenes															0.31	0.53	0.20		
C2-ethryenes															0.51	0.50	0.14		
C3-ethryenes															0.34	0.14	0.13		
C4-ethryenes															2.35	3.58	0.13		

Table 12. Mussel tissue (QA1011S01): p scores (12%) - Biomarkers

Laboratory No.	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17
Carbazole																	
18a(H)-22,29,30-Trienothropane																	1.54
17a(H)-22,29,30-Trienothropane																	1.26
17a(H),21b(H)-30-Norhopane																	0.46
18a(H)-30-Normethopane																	0.96
17a(H)-Diaphopane																	
17a(H)-21b(H)-Horane																	0.45
17a(H),21b(H)-2R-Homotropane																	
17a(H),21b(H)-22S-Homophopane																	0.89
13b(H)-7a(H)-Diacholestane 20S																	0.57
5a(H),14a(H),7a(H)-Cholestan-20S																	0.53
5a(H),14a(H),7a(t)-Cholestan-20R																	0.77
5a(H),14a(H),7a(H)-24-Ethylcholestan-20S																	0.30
5a(H),14a(H),7a(H)-24-Ethylcholestan-20R																	0.72
5a(H),14b(H),7b(H)-7a(H)-Cholestan-20R																	0.42
5a(H),14b(H),7b(H)-7a(H)-Cholestan-20S																	0.35
5a(H),14b(H),7b(H)-24-Ethylcholestan-20S																	0.47
5a(H),14b(H),7b(H)-24-Ethylcholestan-20R																	0.47
5a(H),14b(H),7b(H)-24-Ethylcholestan-20S																	

Table 12 (cont). Mussel tissue (QA1011S01): p scores (15%) - Biomarkers

Laboratory No.	18	19	20	21	22	23	24 _a	24b	25	26	27	28	29	30	31	32	33	34
Carbazole																		
18a(H)-22,29,30-Trienothropane																		0.17
17a(H)-22,29,30-Trienothropane																		0.53
17a(H),21b(H)-30-Norhopane																		0.19
18a(H)-30-Normethopane																		0.06
17a(H)-Diaphopane																		0.29
17a(H)-Horane																		1.73
17a(H),21b(H)-10panic																		0.34
17a(H),21b(H)-22R-Homotropane																		0.44
17a(H),21b(H)-23S-Homotropane																		0.28
13b(H)-7a(H)-Diacholestane 20S																		0.47
5a(H),14a(H),7a(H)-Cholestan-20S																		0.43
5a(H),14a(H),7a(H)-Cholestan-20R																		0.33
5a(H),14a(H),7a(H)-24-Ethylcholestan-20S																		0.36
5a(H),14a(H),7a(H)-24-Ethylcholestan-20R																		0.28
5a(H),14b(H),7b(H)-7a(H)-Cholestan-20R																		0.18
5a(H),14b(H),7b(H)-7a(H)-Cholestan-20S																		0.28
5a(H),14b(H),7b(H)-24-Ethylcholestan-20S																		0.05
5a(H),14b(H),7b(H)-24-Ethylcholestan-20R																		0.21
5a(H),14b(H),7b(H)-24-Ethylcholestan-20S																		0.36

Table 13. Misted-Tissue (Q101TS01): Laboratory means of three replicates summarized by extraction method - PAHs (ng/g dry mass) (reported as if three figures were significant)

	Laboratory No.	Sobkot				QNECHERS				QuEChERS				tissumizer				
		1	4	7	10	12	13	15	20	26	median	2	6	8	9	23	34	
naphthalene	43.0	19.7	36.2	<5700	24.2	54.5	350	37.0	24.3	36.6	37.2	104	31.3	other	11.7	70.6	28.4	
biphenyl	12.9	16.0	<14.8	<5700	9.18	21.1	18.3	35.0	16.3	16.3	NA	NA	NA	NA	13.6	26.5	28.4	
acenaphthene	14.0	11.3	<14.8	<5700	7.42	<30	<10	<7.4	11.1	11.2	nd	0.00	<0.2	<11	DL	11.7	9.06	10.4
acenaphthylene	<10	9.03	<14.8	<5700	4.50	<30	12.7	13.8	10.1	10.1	NA	<0.2	NA	NA	10.7	16.0	10.7	
thiophene	33.9	24.3	19.8	<5700	14.8	33.3	25.3	18.7	23.9	14.3	43.6	25.2	17.7	DL	21.4	24.0	20.8	
phenanthrene	11.9	97.0	139	<5700	11.7	126	127	131	126	56.1	153	134	144	116	134	157	118	
anthracene	10.3	13.3	<14.8	<5700	10.3	9.69	17.7	28.2	11.4	11.4	2.76	12.1	other	10.0	6.10	8.06	15.7	
fluoranthene	246	140	196	183	269	210	173	127	227	196	11.5	247	241	260	21.5	241	221	
pyrene	207	107	121	101	149	149	121	179	158	95.4	191	153	156	182	189	150	175	
benzo[b]fluorene	NA	17.3	NA	NA	15.8	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	
benzo[a]fluorene	37.3	19.7	16.3	<5700	32.1	30.8	26.7	63.6	28.3	29.6	8.87	49.9	28.3	other	50.3	39.1	35.4	
cobaltite	54.7	93.9	<5700	73.0	52.9	88.3	88.7	80.6	59.1	94.8	49.8	107	85.7	Other	NA	NA	34.8	
coelute	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	
triphenylene	144	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	
chrysene/triphenylene	26.3	20.7	21.9	<5700	25.0	35.1	Other	36.0	37.5	26.3	14.4	29.8	27.3	30.8	20.8	27.3	30.5	
benzo[b]fluoranthene	28.6	NA	NA	NA	NA	Other	NA	Other	28.6	NA	NA	NA	NA	NA	NA	NA	34.6	
benzo[k]fluoranthene	15.4	13.0	20.3	<5700	24.7	14.4	11.0	65.8	Other	15.4	3.54	16.6	11.9	16.8	7.83	11.9	NA	NA
benzo[a]fluoranthene	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	
benzo[b,f,k]fluoranthene	<10	1.33	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	
benzo[a]fluoranthene	36.8	24.3	36.6	<5700	29.3	39.9	NA	68.6	37.4	37.4	NA	NA	NA	NA	39.9	NA	NA	
benzo[a]pyrene	11.4	7.40	<14.8	<5700	7.55	11.7	10.7	29.5	12.7	11.4	3.91	14.7	9.03	18.4	7.05	9.03	46.7	
perylene	4.96	2.97	<14.8	<5700	3.93	5.36	<10	172	<10	4.96	NA	NA	NA	NA	6.55	9.40	5.22	
indeno[1,2,3-cd]pyrene	4.75	21.3	<14.8	<5700	4.39	4.38	<10	5.12	7.07	4.93	nd	0.00	5.83	<21	DL	7.70	6.44	
benzofluoranthene	7.81	6.67	<14.8	<5700	7.81	11.9	<10	<7.4	8.68	7.81	nd	3.83	8.03	<7	DL	5.93	9.92	
dibenz[a,h]anthracene	1.12	0.00	<14.8	<5700	2.28	21.1	<10	<7.4	7.83	2.11	nd	0.00	<0.2	<9	DL	2.54	1.77	
dibenz[a,h]anthracene	NA	63.3	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	3.63	NA	
cis/trans-decalin	NA	21.3	NA	NA	13.8	19.8	NA	<7.4	17.8	NA	NA	NA	NA	NA	14.2	NA	11.3	
dibenzofuran	NA	42.3	NA	NA	50.1	30.0	NA	25.4	36.2	NA	NA	NA	NA	NA	18.2	NA	15.7	
retene	NA	0.00	<14.8	NA	<40	NA	NA	<1.3	NA	NA	NA	NA	NA	NA	56.2	NA	56.2	
benzothiophene	<15	12.7	<14.8	<5700	16.0	12.9	12.0	<7.4	16.7	NA	NA	NA	NA	NA	13.0	NA	1.66	
dibenzothiophene	NA	11.7	NA	NA	13.6	NA	NA	NA	12.7	NA	NA	NA	NA	NA	20.4	NA	1.48	
naphthalobenzoanthrone	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	7.41	NA	18.4	
															18.0	NA	12.7	

NA=Not analyzed; DL=Detection limit; No value in space=nothing reported by lab. Bolded values were not used in the calculation of the consensus means in Tables 1 through 3.

Table 13 (cont). Mussel Tissue (Q101801): Laboratory means of three replicates summarized by extraction method along with the median values for the extraction method - PAHs (mg/g dry mass) (reported as if three figures were significant)

	Laboratory No.	Sonication						ASE						base digestion					
		3	17	19	25	30	31	median	5	11	14	18	21	24a	27	28	29	33	median
naphthalene	<294	625	<78	<conic	14.7	5.50	14.7	14.7	12.9	18.4	N/A	30.2	23.0	47.0	<12.2	15.0	31.0	22.5	24.3
benzylbenzene	130	567	<78	<conic	<39.8	6.87	13.0	Other	N/A	N/A	19.1	N/A	13.3	N/A	N/A	13.3	N/A	22.8	<26
acenaphthene	<294	<3.2	<78	<conic	6.69	11.7	9.18	9.59	N/A	9.70	9.64	6.67	N/A	7.12	N/A	10.4	N/A	16.2	11.8
acenaphthylene	<294	<3.2	<78	<conic	6.29	<3.0	6.29	2.74	N/A	8.08	6.77	4.45	N/A	12.5	N/A	6.77	N/A	7.68	<26
fluorene	<294	227	<78	<conic	18.7	<3.0	18.7	26.0	19.7	30.9	20.3	20.6	DL	19.5	18.5	29.2	25.2	20.6	18.6
phenanthrene	101	760	99.3	92.8	56.6	18.7	96.1	149	11.4	11.4	11.4	11.4	DL	10.8	11.9	13.4	14.5	11.8	12.1
anthracene	<294	84.6	<78	67.7	8.75	13.7	10.0	11.4	17.3	12.8	12.8	<6.21	DL	16.1	8.29	11.4	<40	<26	
fluoranthene	181	1160	207	11.4	17.6	24.0	17.9	30.5	22.1	20.9	30.3	23.8	DL	24.2	23.1	21.5	27.7	20.0	52.7
pyrene	112	731	133	73.0	75.8	17.3	93.7	21.2	14.1	15.8	14.0	11.7	DL	13.4	11.1	15.4	19.4	14.1	15.4
benzylfluorene	N/A	<3.2	<conic	<39.8	N/A	N/A	N/A	N/A	N/A	N/A	N/A	<15.4	DL	22.4	N/A	N/A	N/A	N/A	35.0
benz[a]anthracene	<294	198	<78	<conic	24.0	52.0	24.0	57.8	35.8	36.6	53.2	33.7	DL	34.2	33.8	37.2	55.9	37.2	<26
chrysene	<294	coelute	<78	41.8	56.0	15.3	43.8	96.2	35.3	82.5	58.7	29.2	coelute	83.3	coelute	71.2	71.2	other	26.0
triphenylene	N/A	coelute	N/A	N/A	N/A	N/A	N/A	Other	N/A	38.3	82.5	N/A	N/A	N/A	coelute	N/A	60.4	other	N/A
chlorobenzene/anthracene	421	N/A	N/A	N/A	N/A	N/A	N/A	13.9	N/A	N/A	N/A	N/A	N/A	N/A	N/A	91.6	107	107	84.4
benz[a]fluoranthene	<294	261	<78	<conic	21.8	<3.0	38.1	44.0	31.0	32.6	55.5	49.3	42.1	22.4	28.1	57.2	40.1	25.0	coelute
benz[a]fluoranthene	N/A	coelute	N/A	N/A	N/A	N/A	N/A	Other	N/A	16.6	N/A	N/A	coelute	N/A	coelute	N/A	N/A	16.6	other
benz[a]fluoranthene	<294	coelute	<78	<conic	14.2	3.20	8.70	25.4	15.1	14.7	14.3	DL	30.1	coelute	N/A	19.3	17.2	coelute	coelute
benz[a]fluoranthene	194	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	14.7	32.8	32.8	24.7
benz[a]fluoranthene	N/A	<3.2	N/A	N/A	N/A	N/A	N/A	N/A	5.42	N/A	N/A	N/A	<2.94	N/A	DL	N/A	5.42	N/A	<26
benz[e]pyrene	<294	232	<78	<conic	<39.8	4.00	48.9	36.1	39.4	N/A	N/A	40.8	43.1	N/A	40.8	37.5	N/A	N/A	
benz[a]pyrene	<294	<3.2	<78	<conic	6.36	26.0	16.2	11.8	11.5	11.0	16.7	12.9	23.3	8.42	11.4	16.5	12.3	12.0	<26
perylene	191	<3.2	<78	<conic	<39.8	12.1	12.1	5.68	N/A	4.74	4.16	N/A	N/A	<7.65	N/A	6.50	N/A	5.21	<26
indenol[1,2,3-cd]pyrene	<294	207	<78	<conic	6.36	<3.0	6.36	8.35	9.39	8.22	7.65	13.4	14.2	<4.58	10.5	DL	16.8	10.0	<26
benz[b]fluoranthene	<294	204	<78	<conic	6.36	<3.0	6.36	10.8	N/A	9.88	13.1	11.0	N/A	7.29	N/A	10.5	N/A	10.6	<26
benz[a]anthracene	<294	175	<78	<conic	<39.8	<3.0	2.35	<1.5	3.14	5.09	DL	<3.28	<4.81	DL	3.23	4.16	other	<26	
dibenz[a+e]anthracene	N/A	<3.2	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	DL	N/A	1.92	N/A	
cistrans-decalin	<294	143	<78	<conic	<39.8	<3.0	N/A	N/A	N/A	N/A	N/A	N/A	N/A	53.9	DL	N/A	53.9	38.3	
dibenzofuran	N/A	245	N/A	N/A	N/A	N/A	N/A	N/A	11.8	N/A	28.3	N/A	N/A	17.1	DL	N/A	17.1	N/A	
retene	N/A	<3.2	<78	<conic	<39.8	<3.0	N/A	N/A	N/A	N/A	N/A	N/A	N/A	<9.61	DL	N/A	20.1	30.7	<26
benzothiophene	<294	<3.2	<78	<conic	<39.8	<3.0	N/A	N/A	N/A	N/A	N/A	N/A	N/A	<5.15	DL	N/A	N/A	N/A	<26
naphthalenothiophene	N/A	<3.2	N/A	N/A	N/A	N/A	N/A	Other	N/A	N/A	N/A	N/A	N/A	17.6	N/A	18.6	N/A	14.6	<26
N/A=Not analyzed; DL=Detection limit; N/A=space=n/a nothing reported by lab; Bolded values were not used in the calculation of the consensus means in Tables 1 through 3.																			

Table 14. Mussel Tissue (QAX101S01): Laboratory means of three replicates summarized by extraction method along with the median values for the extraction method - Alkylated PAHs (ng/g dry mass) (reported as if three figures were significant)

	Laboratory No	Soxhlet			QMECHERS			QMECHERS			24b			tissueinizer		
		1	4	7	10	12	13	15	20	median	16	23	34	median		
1-methylindophenole	27.2	15.0	11.3	<5700	23.0	50.5	37.7	37.6	29.7	33.6	N.A.	50.3	40.1	30.4	34.2	34.2
2-methylindophenole	36.2	23.7	64.4	<5700	38.0	90.3	67.7	34.4	41.4	39.7	N.A.	31.5	60.2	45.1	52.2	52.2
2,6-dimethylnaphthalene	67.0	42.3	N.A.	N.A.	65.4	101	N.A.	106	78.8	72.9	N.A.	N.A.	70.5	60.2	51.7	73.3
6,7-trimethylnaphthalene	46.5	46.3	N.A.	N.A.	50.5	74.3	N.A.	168	81.4	62.4	N.A.	N.A.	NA	NA	41.7	41.7
1-methylphenanthrenene	54.6	23.7	N.A.	N.A.	<7.4	43.7	49.1	N.A.	61.1	N.A.	N.A.	N.A.	61.1	67.5	59.2	42.3
C1-decalins	N.A.	390	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	NA	NA	NA	NA
C2-decalins	N.A.	1333	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	NA	NA	NA	NA
C3-decalins	N.A.	1900	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	NA	NA	NA	NA
C4-decalins	N.A.	2533	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	NA	NA	NA	NA
C1-naphthalenes	63.4	23.7	NA	<5700	35.5	NA	NA	47.2	48.3	47.2	NA	NA	NA	NA	NA	NA
C2-naphthalenes	255	78.3	NA	<5700	125	NA	31.3	21.2	180	196	NA	NA	NA	NA	NA	NA
C3-naphthalenes	629	153	NA	207	253	NA	419	314	290	290	NA	NA	NA	NA	NA	NA
C4-naphthalenes	498	170	NA	110	210	NA	466	319	400	319	NA	NA	NA	NA	NA	NA
C1-benzothiophenes	N.A.	0.00	N.A.	N.A.	N.A.	N.A.	N.A.	<39	N.A.	N.A.	N.A.	N.A.	N.A.	NA	NA	NA
C2-benzothiophenes	N.A.	0.00	N.A.	N.A.	N.A.	N.A.	N.A.	<39	N.A.	N.A.	N.A.	N.A.	NA	NA	NA	NA
C3-benzothiophenes	N.A.	0.00	N.A.	N.A.	N.A.	N.A.	N.A.	<39	N.A.	N.A.	N.A.	N.A.	NA	NA	NA	NA
C4-benzothiophenes	N.A.	76.8	34.0	<5700	37.7	NA	75.7	<7.4	68.3	68.3	NA	NA	NA	NA	NA	NA
C1-fluorenes	181	107	NA	<5700	109	NA	188	<7.4	373	181	NA	NA	NA	NA	NA	NA
C2-fluorenes	296	473	NA	<5700	173	NA	436	<7.4	780	436	NA	NA	NA	NA	NA	NA
C3-fluorenes	280	163	NA	<5700	226	NA	415	<7.4	187	226	NA	NA	NA	NA	NA	NA
C1-phenanthrenes/anthracenes	521	263	NA	102	326	NA	68.9	<7.4	493	410	NA	NA	NA	NA	NA	NA
C2-phenanthrenes/anthracenes	467	323	NA	<5700	232	NA	104.4	<7.4	460	460	NA	NA	NA	NA	NA	NA
C3-phenanthrenes/anthracenes	145	230	NA	<5700	113	NA	597	<7.4	260	230	NA	NA	NA	NA	NA	NA
C4-phenanthrenes/anthracenes	95.9	55.3	NA	<5700	72.3	NA	64.7	<46.2	82.7	68.5	NA	NA	NA	NA	NA	NA
C1-dihenzothiophenes	23.7	14.0	NA	<5700	193	NA	192	<7.4	287	193	NA	NA	NA	NA	NA	NA
C2-dihenzothiophenes	223	14.3	NA	<5700	163	NA	155	61.7	303	159	NA	NA	NA	NA	NA	NA
C3-dibenzothiophenes	110	87.0	NA	<5700	68.7	NA	<40	55.6	263	87.0	NA	NA	NA	NA	NA	NA
C4-dibenzothiophenes	145	70.0	NA	<5700	91.9	NA	66.7	373	127	109	NA	NA	NA	NA	NA	NA
C1-fluoranthenes/pyrenes	62.3	42.7	NA	<5700	55.3	NA	43.0	<7.4	92.7	55.3	NA	NA	NA	NA	NA	NA
C2-fluoranthenes/pyrenes	33.4	31.7	NA	<5700	24.3	NA	<40	57.9	52.3	33.4	NA	NA	NA	NA	NA	NA
C3-fluoranthenes/pyrenes	<1.5	12.6	NA	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	<39	12.6	NA	NA	NA	NA
C4-fluoranthenes/pyrenes	N.A.	55.7	NA	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	<39	55.7	NA	NA	NA	NA
C1-naphthobenzothiophenes	N.A.	0.00	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	NA	NA	NA	NA
C2-naphthobenzothiophenes	N.A.	0.00	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	NA	NA	NA	NA
C3-naphthobenzothiophenes	N.A.	0.00	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	N.A.	NA	NA	NA	NA
C4-naphthobenzothiophenes	N.A.	42.6	25.7	NA	<5700	28.9	NA	35.5	43.5	35.5	NA	NA	NA	NA	NA	NA
C1-chrysenes	25.7	21.3	NA	<5700	16.2	NA	210	43.0	25.7	NA	NA	NA	NA	NA	NA	NA
C2-chrysenes	<10	9.93	NA	<5700	6.20	NA	<7.4	<39	8.07	8.07	NA	NA	NA	NA	NA	NA
C3-chrysenes	<10	0.00	NA	NA	<40	NA	<7.4	<39	0.00	0.00	NA	NA	NA	NA	NA	NA
C4-chrysenes											2.93	<0.595	2.93			

NA = Not analyzed, DL = Detection limit; No value in space=nothing reported by lab. Bolded values were not used in the calculation of the consensus statistics

Table 14 (cont.). Mussel Tissue (Q1011S01): Laboratory means of three replicates summarized by extraction method along with the median values for the extraction method - Alkylated PAHs (mg/g dry mass) (reported as if three figures were significant)

	Laboratory No.	ASE										base digestion median 22	microwave median 32		
		17	19	25	30	31	median	11	14	18	21	24a	27	28	33
1-methylbenzene	<294	<6.4	<78	<cone	16.1	7.90	12.0	21.0	27.4	29.8	19.9	25.6	58.3	<24.3	38.3
2-methylbenzene	226	653	<78	<cone	70.5	6.90	70.5	37.1	41.8	47.6	31.8	42.3	94.0	<24.3	24.8
1,6-dimethylbenzene	102	1062	<78	104	27.1	13.2	102	68.8	46.9	41.5	41.6	11.8	48.4	69.7	87.5
1,6,7-trimethylbenzene	112	325	<78	<cone	N/A	17.7	112	39.4	N/A	41.4	N/A	30.6	N/A	46.0	N/A
1-methylphenanthrene	<294	2465	<78	<cone	N/A	<3.0	N/A	81.1	56.4	77.1	64.8	57.3	DL	35.1	<8.65
C1-decalins	N/A	2467	N/A	<cone	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	30.6	N/A	83.8
C2-decalins	N/A	14233	N/A	<cone	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	45.6	N/A	N/A
C3-decalins	N/A	21200	N/A	<cone	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	310	N/A	N/A
C4-decalins	N/A	24533	N/A	<cone	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	506	N/A	N/A
C1-undecanes	506	462	<78	<cone	N/A	13.0	N/A	86.2	N/A	N/A	63.4	N/A	<24.3	62.3	77.5
C2-undecanes	378	2728	123	<cone	N/A	26.0	251	236	N/A	N/A	277	508	102	263	90.0
C3-undecanes	391	4270	200	<cone	N/A	27.0	295	606	N/A	N/A	300	497	226	498	229
C4-undecanes	504	5307	227	<cone	N/A	18.3	410	600	N/A	N/A	240	683	279	138	266
C1-benzoethiophenes	N/A	<3.2	N/A	<cone	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	21.1	N/A	N/A
C2-benzoethiophenes	N/A	<3.2	N/A	<cone	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	12.4	N/A	N/A
C3-benzoethiophenes	N/A	<3.2	N/A	<cone	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	20.4	N/A	N/A
C4-benzoethiophenes	N/A	<3.2	N/A	<cone	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	37.8	N/A	N/A
C1-fluorenes	<294	856	<78	<cone	N/A	<3.0	N/A	68.6	N/A	N/A	84.9	98.7	70.1	21.5	446
C2-fluorenes	<294	2563	147	<cone	N/A	<3.0	147	186	N/A	N/A	195	19.0	181	658	77.2
C3-fluorenes	<294	<3.2	217	<cone	N/A	<3.0	217	362	N/A	N/A	48.4	DL	354	<6.89	558
C1-phenantrenes/anthracenes	<294	<3.2	190	<cone	N/A	<3.0	190	425	N/A	N/A	312	68.5	230	360	554
C2-phenantrenes/anthracenes	662	2358	307	106	N/A	<3.0	484	626	N/A	N/A	113	1457	393	934	433
C3-phenantrenes/anthracenes	<294	3257	263	<cone	N/A	<3.0	1760.0	508	N/A	N/A	155	469	307	356	636
C4-phenantrenes/anthracenes	294	1380	117	<cone	N/A	<3.0	117	209	N/A	N/A	5.38	1.59	<9.48	426	469
C1-dibenzofurans	<294	385	<78	<cone	N/A	<3.0	N/A	92.1	N/A	N/A	N/A	N/A	71.6	N/A	590
C2-dibenzofurans	<294	946	107	<cone	N/A	<3.0	107	245	N/A	N/A	N/A	N/A	226	N/A	92.1
C3-dibenzofurans	<294	1113	110	<cone	N/A	<3.0	110	206	N/A	N/A	N/A	N/A	651	N/A	245
C4-dibenzofurans	<294	877	<78	<cone	N/A	<3.0	N/A	103	N/A	N/A	N/A	N/A	234	N/A	328
C1-fluoranthenes/pyrenes	<294	717	99	<cone	N/A	<3.0	N/A	114	N/A	N/A	338	N/A	108	433	639
C2-fluoranthenes/pyrenes	166	<3.2	<78	<cone	N/A	<3.0	166	68.3	N/A	N/A	91.6	N/A	115	N/A	132
C3-fluoranthenes/pyrenes	<294	<3.2	<78	<cone	N/A	<3.0	N/A	37.1	N/A	N/A	N/A	N/A	71.7	N/A	115
C4-fluoranthenes/pyrenes	N/A	<3.2	N/A	<cone	N/A	<3.0	N/A	12.2	N/A	N/A	N/A	N/A	36.2	N/A	136
C1-phenanthrenes/biphenes	N/A	<3.2	N/A	<cone	N/A	<3.0	N/A	20.4	N/A	N/A	N/A	N/A	19.1	N/A	19.7
C2-phenanthrenes/biphenes	N/A	<3.2	N/A	<cone	N/A	<3.0	N/A	13.8	N/A	N/A	N/A	N/A	16.0	N/A	<5.2
C3-phenanthrenes/biphenes	N/A	<3.2	N/A	<cone	N/A	<3.0	N/A	6.97	N/A	N/A	N/A	N/A	<8.19	N/A	<5.2
C4-phenanthrenes/biphenes	N/A	<3.2	N/A	<cone	N/A	<3.0	N/A	31.0	N/A	N/A	N/A	N/A	<8.19	N/A	N/A
C1-alkylbenzothiophenes	<294	308	<78	<cone	N/A	<3.0	N/A	56.9	N/A	N/A	N/A	N/A	41.6	N/A	43.9
C2-alkylbenzothiophenes	<294	<3.2	<78	<cone	N/A	<3.0	N/A	31.2	N/A	N/A	N/A	N/A	22.4	N/A	22.4
C3-alkylbenzothiophenes	<294	<3.2	<78	<cone	N/A	<3.0	N/A	13.5	N/A	N/A	N/A	N/A	22.4	N/A	22.4
C4-alkylbenzothiophenes	<294	<3.2	<78	<cone	N/A	<3.0	N/A	<1.79	N/A	N/A	N/A	N/A	<13.8	N/A	12.9

N/A=Not analyzable; DL=Detection limit. No value in space=nothing reported by lab. Bolded values were not used in the calculation of the consensus statistics

Table 15. Moused Tissue (QA101180): Laboratory means of three replicates summarized by extraction method along with the median values for the extraction method - Biomarkers (ng/g dry mass) (reported as if three figures were significant)

	Sow/lot	Laboratory No.	1	4	7	10	12	13	15	20	26	median	QuEChERS	16	23	34	tissueinizer
Carbazole			NA	NA	NA	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[8]ad(I)-22,29,30-Trisnorbornane			NA	NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[17a(I),22,29,30-Trisnorbornane]	<25		NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[7a(I),21(I),30-Norbornane]	<25		NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[8a(I)-30-Normalcyclopane]			NA	NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[7a(I)-Dihydropane]			NA	NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[17a(I),21(I)-1-Epoxide]			NA	NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[7a(I),21(I)-1-Epene]	65.7		NA	NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[7a(I),21(I)-2aR-1-Limonophane]	<20		NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[7a(I),21(I)-2aS-1-Limonophane]	23.1		NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[13b(I)] ¹ [7a(I)-Dihydrostane 20S]			NA	NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4a(I),17a(I)-Cholestan-20S]			NA	NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4a(I),17a(I)-Cholestan-20R]	38.2		NA	NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4a(I),17a(I)-24-Ethylcholestan-20S]			NA	NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4a(I),17a(I)-24-Ethylcholestan-20R]	29.0		NA	NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4b(I),17b(I)-24-Ethylcholestan-20R]	<20		NA	NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4b(I),17b(I)-Cholestan-20S]			NA	NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4b(I),17b(I)-24-Ethylcholestan-20R]	<20		NA	NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4b(I),17b(I)-Cholestan-20S]			NA	NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4b(I),17b(I)-24-Ethylcholestan-20S]	<20		NA	NA	<700	<40	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[NA = Not analyzed; DL = Detection limit; No value in space=nothing reported by lab; Bolded values were not used in the calculation of the consensus statistics]																	

Table 15 (cont). Mousel Tissue (QA101180): Laboratory means of three replicates summarized by extraction method along with the median values for the extraction method - Biomarkers (ng/g dry mass) (reported as if three figures were significant)

	Sonication	Laboratory No.	3	17	19	25	30	31	median	ASE	5	11	14	18	21	24a	27	28	29	33	median
Carbazole			NA	23.1	NA	<conc	NA	NA	22.1	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[8]ad(I)-22,29,30-Trisnorbornane	<294		NA	<78	<conc	NA	<3.0	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[17a(I),22,29,30-Trisnorbornane]	<294		NA	<78	<conc	NA	<3.0	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[7a(I),21(I)-30-Norbornane]	<294		NA	<78	<conc	NA	11.2	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[7a(I)-Dihydropane]	<294		NA	<78	<conc	NA	86.7	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[7a(I)-2,0(I)-Homolopane]	<294		NA	<78	<conc	NA	<3.0	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[7a(I),2,1(I)-22S-Homolopane]	<294		NA	<78	<conc	NA	<3.0	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[13b(I)] ¹ [7a(I)-Dihydrostane 20S]	<294		NA	<78	<conc	NA	<3.0	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4a(I),17a(I)-Cholestan-20S]	<294		NA	<78	<conc	NA	<3.0	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4a(I),17a(I)-Cholestan-20R]	<294		NA	<78	<conc	NA	<3.0	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4a(I),17a(I)-24-Ethylcholestan-20S]	<294		NA	<78	<conc	NA	<3.0	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4b(I),17b(I)-24-Ethylcholestan-20R]	<294		NA	<78	<conc	NA	24.7	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4b(I),17b(I)-Cholestan-20R]	<294		NA	<78	<conc	NA	67.3	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4b(I),17b(I)-24-Ethylcholestan-20S]	<294		NA	<78	<conc	NA	<3.0	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4b(I),17b(I)-Cholestan-20S]	<294		NA	<78	<conc	NA	59.7	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[5a(I),4b(I),17b(I)-24-Ethylcholestan-20S]	<294		NA	<78	<conc	NA	<3.0	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[NA = Not analyzed; DL = Detection limit; No value in space=nothing reported by lab; Bolded values were not used in the calculation of the consensus statistics]																					

[NA = Not analyzed; DL = Detection limit; No value in space=nothing reported by lab; Bolded values were not used in the calculation of the consensus statistics]

APPENDIX A

Interlaboratory Analytical Comparison Study to Support Deepwater Horizon Natural Resource Damage Assessment

Intercomparison Exercise: Mussel Tissue QA10TIS01 Description of Materials and Instructions

Intercomparison Exercise Materials:

Each of the three jars contains approximately 15 g (wet basis) of Mussel Tissue QA10TIS01. This cryogenically homogenized fresh frozen material was prepared from mussels collected from an urban area. This material has not been enriched or spiked and still contains its endogenous water. Each 2-oz clear glass jar has a Teflon-lined screw cap and is labeled with an individual jar number as well as the above name.

In addition, three concurrent analyses of SRM 1974b Organics in Mussel Tissue (*Mytilus edulis*) are recommended. This material can be obtained from the NIST Standard Reference Materials Program (\$798/5 x 8 g to 10 g bottles). See the following link for information on ordering on-line: https://www-s.nist.gov/srmors/view_detail.cfm?srm=1974B

Storage of Materials:

The tissue material should be stored in the dark at temperatures of -20 °C or lower. If allowed to thaw or if stored for extended periods at temperatures higher than -40 °C, it will lose its powder-like form. If this happens, use the contents from the entire jar for analysis. This material has been stored at NIST at -80 °C and was shipped to you on dry ice. If only a portion of the contents of a jar is used, the jar should be tightly closed immediately after removal of a subsample to preserve the integrity of the remaining material for later analysis.

Instructions for Use:

You are to analyze Mussel Tissue QA10TIS01 and SRM 1974b using **your** laboratory's and/or program's analytical protocols, for the concentrations (mass/mass [dry-mass basis]) of the parent polycyclic aromatic hydrocarbon (PAH) compounds, alkylated PAH compounds, and biomarkers currently being determined in your laboratory. A target list of compounds are presented in Table 1; however, participants do not need to quantify all of these compounds and can add additional compounds when reporting data.

The percentage of water in the mussel tissue should be determined so that the results can be reported on a dry basis. You should have received sufficient material so that you can perform separate determinations for the water content if you do not dry your tissue samples prior to analysis.

The amount of material used for each analysis should correspond to the amount (wet basis) of marine tissue that you would typically analyze as prescribed in your protocols. It is best if the Mussel Tissue QA10TIS01 and the SRM 1974b samples are not allowed to thaw prior to the taking of samples for analysis; however, if the material has been even partially thawed, you should use the contents of the entire jar as a single sample as it is difficult to take representative samples from a jar once the material

has thawed. After removing the material for analysis from the jars, the samples should be used without delay.

You should analyze three samples of Mussel Tissue QA10TIS01 and at least one or more samples of SRM 1974b using your protocol for tissue samples. If time allows, we are asking that you analyze one sample of Mussel Tissue QA10TIS01 and one sample of SRM 1974b with one batch of laboratory samples; analyze a second sample of each material with another batch; and the third sample with yet another batch. This will allow a more realistic assessment of laboratory precision over a longer term than the assessment obtained when a laboratory places all three samples in the same extraction and cleanup batch and the resulting extracts are analyzed using the same calibration curve, etc.

Reporting of Results:

Please report one result, as if three figures were significant, for each of the analytes quantified in each of the three replicates of the Mussel Tissue QA10TIS01 and of SRM 1974b. Report results in units of ng/g **dry-mass** basis. Report the date of measurement of each sample in the requested m/d/y format. Also, report the results of your percentage water determinations of Mussel Tissue QA10TIS01.

If you know that a target or non-target compound is interfering (coeluting) with the determination of a target analyte, please identify this issue by qualifying the data and note the data qualifier used at the bottom of your table of results. Please note that any changes you make to the column or row headings **within** the tables will **not** be seen by the coordinators because only the table entries and comments at the bottom of the tables are automatically transferred to the exercise database.

We prefer that concentration values be reported for each analyte determined. If the measured concentration is below your typical reporting concentration for an analyte in a particular matrix, you can report the number and list the appropriate detection limit, quantification limit, etc. at the bottom of the data table. However, if you need to report non-numerical data please use the following conventions:

NA	"Not analyzed", "not determined"
<"value"	"Less than specified concentration", e.g., <8 ng/g
Other	"Other"; add note of explanation at end of data table, e.g., interference
DL	"Below detection limit" may be used, however, <"value" is preferable

Do not use negative numbers or parentheses to indicate "less than detection limits".

The attached file is an EXCEL file, QA10TIS01.xls. If you have any software/hardware conversion problems, please contact Michele Schantz. The data file templates also include places for you to list the surrogate/internal standards and type of calibration curve used, and to provide a brief description of the analyses. Please **do not** add spaces before entering numbers in the table cells and enter them as "numbers" not as "labels". Please **do not** insert any columns or rows **within** the table in the data file. If you wish to include additional data and/or other information or comments, you may add it to the bottom of the data table in the attached file.

Submit your results by **April 8, 2011** as an attached file via e-mail to:

michele.schantz@nist.gov

Further Information:

If you need further information, please contact Michele at the following address or phone numbers:

Michele M. Schantz
NIST
100 Bureau Drive Stop 8392
Gaithersburg, MD 20899-8392

Phone: (301)975-3106
FAX: (301)977-0685

Table 1: Preliminary List of Analytes of Interest in the Interlaboratory Analytical Comparison Study to Support Deepwater Horizon Natural Resource Damage Assessment

PAHs

Naphthalene	cis/trans-Decalin
Biphenyl	Dibenzofuran
Acenaphthene	Retene
Acenaphthylene	Benzothiophene
Fluorene	Dibenzothiophene
Phenanthrene	Naphthobenzothiophene
Anthracene	
Fluoranthene	
Pyrene	
Benzo[<i>b</i>]fluorene	
Benz[<i>a</i>]anthracene	
Chrysene	
Triphenylene	
Benzo[<i>b</i>]fluoranthene	
Benzo[<i>j</i>]fluoranthene	
Benzo[<i>k</i>]fluoranthene	
Benzo[<i>a</i>]fluoranthene	
Benzo[<i>e</i>]pyrene	
Benzo[<i>a</i>]pyrene	
Perylene	
Indeno[1,2,3- <i>cd</i>]pyrene	
Benzo[<i>ghi</i>]perylene	
Dibenz[<i>a,h</i>]anthracene	

Alkylated PAHs

1-Methylnaphthalene	C1-Naphthalenes
2-Methylnaphthalene	C2-Naphthalenes
2,6-Dimethylnaphthalene	C3-Naphthalenes
1,6,7-Trimethylnaphthalene	C4-Naphthalenes
1-Methylphenanthrene	
C1-Decalins	C1-Benzothiophenes
C2-Decalins	C2-Benzothiophenes
C3-Decalins	C3-Benzothiophenes
C4-Decalins	C4-Benzothiophenes

Table 1 (cont.)

C1-Fluorenes	C1-Naphthobenzothiophenes
C2-Fluorenes	C2-Naphthobenzothiophenes
C3-Fluorenes	C3-Naphthobenzothiophenes
	C4-Naphthobenzothiophenes
C1-Phenanthrenes/anthracenes	
C2- Phenanthrenes/anthracenes	C1-Chrysenes
C3- Phenanthrenes/anthracenes	C2-Chrysenes
C4- Phenanthrenes/anthracenes	C3-Chrysenes
C1-Dibenzothiophenes	C4-Chrysenes
C2-Dibenzothiophenes	
C3-Dibenzothiophenes	
C4-Dibenzothiophenes	
C1-Fluoranthenes/pyrenes	
C2-Fluoranthenes/pyrenes	
C3-Fluoranthenes/pyrenes	
C4-Fluoranthenes/pyrenes	

Biomarkers

Carbazole
18 α (H)-22,29,30-Trisnorhopane
17 α (H)-22,29,30-Trisnorhopane
17 α (H),21 β (H)-30-Norhopane
18 α (H)-30-Norneohopane
17 α (H)-Diahopane
17 α (H),21 β (H)-Hopane
17 α (H),21 β (H)-22R-Homohopane
17 α (H),21 β (H)-22S-Homohopane
13 β (H) 17 α (H)-Diacholestane 20S
5 α (H),14 α (H),17 α (H)-Cholestane 20S
5 α (H),14 α (H),17 α (H)-Cholestane 20R
5 α (H),14 α (H),17 α (H)-24-Ethylcholestane 20S
5 α (H),14 α (H),17 α (H)-24-Ethylcholestane 20R
5 α (H),14 β (H),17 β (H)-Cholestane 20S
5 α (H),14 β (H),17 β (H)-Cholestane 20R
5 α (H),14 β (H),17 β (H)-24-Ethylcholestane 20S
5 α (H),14 β (H),17 β (H)-24-Ethylcholestane 20R

Appendix B

Summary of Method Information Provided by the Participating Laboratories

Extraction parameters used by labs for tissue samples

Lab #	Revised	Extracted QAI(01TSU)	Extracted SRM (9-4b)	Determination % water	Extraction Method	Extraction Solvent	Extraction Time	Extraction Solvent
1	3-22-2011	S	S	Open drying at 110°C until stable mass	Sorbet	deionized water (D/W)	20 h	mixed with hydroxylamine in the extraction cell
2	3-16-2011	S	S	Loss on drying (10°C for 12 h)	QuEChERS	Acetonitrile		
3	3-28-2011	S	10	ASTM D2246	SW-846 SS-08	DCM	3 times at 5 min each	
4	3-30-2011	S	S	Modified EPA (eo 3)	EPA 3550	DCM	4h	ASF conditions: Pressure = 2000 psi (13.8 MPa), Temperature = 100°C
5	4-5-2011	S	S	Open drying at 110°C overnight	Accelerated solvent extraction (ASF)	DCM	~16 min	18.00 mL to 5g shake min with ceramic homogenizer, use ceramic media to shake (500 rpm)
6	4-6-2011	S	S	AOAC Official Method 952 (Oil Solids (Total) in Seafood)	QuEChERS	Acetonitrile	1 min	2, 3 cycles per hour - overnight
7	4-7-2011	S	S	Small amount of sample placed in dram shell vial and heated to ~105°C on a heating block for over 8 h	Sorbet		12-16 h	
8	4-8-2011	S	S	Weighted aliquot of sample before and after drying in vacuum oven at 90°C	QuEChERS	Acetonitrile	shake 1 min, rotate 10 min	
9	4-8-2011	S	S	Weighted about 0.5 g and dried at 100°C for 24 h. Repeated drying until constant weight	modified QuEChERS	Acetonitrile / ACN / Acetonitrile		
10	4-8-2011	18	0					
11	4-7-2011	2	2	Gravimetric Determination of Water by Drying and Weighing Ruiz, R. P. 2001. Current Protocols in Food Analytical Chemistry. John Wiley & Sons	ASF (sample + sodium sulfate + magnesium sulfate)	DCM	2 min via Centrifugation	2h program starts at 150°C bowl 15 min + 15 min reductions spin extraction, 3-15 min reductions. Cycles for 45 min pull off & blow down
12	4-8-2011	0	10	SW-846 Guidance	3540	DCM Acetone	10 min per sample	
13	4-8-2011	S	S	Dried an aliquot of sample in the oven at 105°C for 24 h	Sorbet	water, hexane, acetone	> 18 h	
14	4-8-2011	S	S	Dry for 24 h at 110°C	ASF	DCM	5 min, static time 5 min, cell pressure was 2000 psi (13.8 MPa) and there were three cycles	
15	4-7-2011	S	N/A	Open dry 105°C gravimetric	Modified EPA 3550	DCM	Minimum 16 h	
16	4-8-2011	S to 10	S to 10	Drying oven at 105°C overnight	SCP-S-100 - Thinnerizer extraction	DCM	2 to 2 min	Open 3 h extraction, 2 h static cleaning, 1 h concentration
17	4-8-2011	10	10	Iod 3 Modified TS	EPA 3550	DCM		
18	4-8-2011	S	S	gramine	ASL: KD Silica Cicciolini KD	DCM Hexane	1 day	

Extraction parameters used by labs for tissue samples (cont)

Lab #	Reported	% extracted	g extracted	% water	Determination	Extraction Method	Extraction Solvent	Extraction Time	Extraction Fraction
19	QA101NS01	SRM 1074b				Sonication	Acetone/DCM (50:50)	3 min in triplicate	other
20	4.8.2011	13.2 - 14.6	9.9 - 12.2	160.3	SM 2540G	SW 86.3541	DCM / Acetone	2.5 h	
21	4.8.2011	10.2	6.8						21 min total time with two cycles extraction
22	4.8.2011	8	8		Freeze drying 7 days at -20°C	ACU-350 NOAA procedure	DCM		
23	4.8.2011	18	10		Gravimetric	Rise digest	Pentane	4.5 h	
24a	4.8.2011	20	7		An aliquot (~1 g) was removed and dried to a consistent weight after drying at 40°C	Tissumizer with sodium sulfate	DCM	three times for 3 min each time	
24b	4.8.2011	21	2		Microwave moisture analyzer	NOAA Technical Memorandum NMFS-NWFSC-59	isooctane	24 h	
25	4.8.2011	8			Microwave moisture analyzer	QuICHERS-based method for LC-FLD screening	acetone/tile	1.5 h	
26	4.8.2011	10	8		2g of each sample was dried in an oven set to 105°C for 1 h.	35500B sonication extraction	1:1 DCM/Acetone	3 min with sonic disrupter hem	
27	4.8.2011	10	8		EPA 160.3	EPA 3541	DCM	312	
28	4.8.2011	9.6	8		oven drying at 105°C to constant weight	ANF	DCM	13 min per sample	1500 psi (10.4 MPa), 100°C Celsius, 2 static extraction cycles sample, reduction to 2.0 to 3.0 ml using water bath
29	4.4.2011	2	2		Drying overnight in an oven set at 90°C	NOAA Method (July 2010)	DCM	20 min	
30	4.15.2011	6	6		gravimetric 1g dried at 120°C for 24 h aliquot weighed into aluminum dish then dried at 110°C for 4 h, then weighed again and % solids calculated	ANF	DCM	~30min	ground w. sodium sulfate
31	4.22.2011	15	4			3550C	DCM		
32	4.26.2011	17	4		Dry at 105°C	SW 3540 (Microwave)	DCM/Acetone 9:10	4 (9 11.4) min	
33	4.27.2011	28	28		University of Massachusetts - Determination of Moisture and Total Solids, Gravimetric drying at 100 °C for 8 h	NOAA Technical Memorandum NMFS-NWFSC-59	ACF - DCM	20 min (2x)	Exchanged to hexane
34	5.6.2011	8	8		Gravimetric SM2540G	Tissumizer, Lab SOP - OP-003	DCM	30 min total (3 x 20 min serial extractions)	

Sample clean-up and quantification methods used by laboratories for tissue study

Lab #	Sample extract cleanup method	Method of quantification
1	concentrate extract to 0.5 mL with solvent change to hexane; pass through two aminopropyl solid phase extraction (SPE) columns in series using 20 mL of 20% methylsulfide in hexane (isane solvent mixture and volume used to condition the SPE column) concentrate fraction to 0.5 mL and transfer to autosampler vial	IS
2	QuEChERS	ES
3	Gel Permeation Column (GPC)	IS
4	Silica Gel	IS
5	Gravity flow column with silica gel and neutral alumina, followed by size-exclusion HPLC to elute fraction containing PAHs and alkyl-PAHs.	IS
6	QuEChERS packet added (6 g magnesium sulfate and 1.5 g of sodium acetate), shake vigorously 1 min, centrifuge at 3000 rpm for 10 min, filter through 0.2 μ m PTFE syringe filter	ES
7	1. SX-3 Size Exclusion Chromatography (SEC) with DCM Elution; 2. KOH-Impregnated Silica Gel 60 (SG-60; 5g) Flash Chromatography with 5% Methyl t Butyl Ether (MBE) in Hexanes Elution; 3. Phenogel (Phenomenex) SEC with DCM Elution; 4. 3% Aqueous Deactivated SG-60 (5g) Flash Chromatography with Hexanes (Impurities - Discarded) Followed by 25% MBE in Hexanes (PAHs) Elution	IS
8	4.2g 60 mesh activated Silica gel with 1cm layer sodium sulfate. Column rinsed with 3:1 hexane:MeCl2 followed by additional Hexane rinse. Sample added followed by 40mL Hexane. PAHs eluted with 3:1 Hexane: MeCl2 Solvent exchange to Iso-octane for analysis	IS
9	No post extraction sample cleanup.	ES
10	GPC	IS
11	Silica Alumina Column Chromatography (gravity-flow), Gel Permeation Chromatography (GPC) - J2 Scientific	IS
12	3640-3630	IS
13	Gel Permeation Chromatography	IS
14	Alumina solid phase extraction.	IS
15	GPC followed by a neutral alumina/silica column cleanup	IS for PAH; ES for biomarkers
16	Alumina cleanup followed by silica gel cleanup	IS
17	Silica gel cleanup performed.	IS
18		IS
19	filter through sodium sulfate, followed by GPC, followed by silica gel	IS
20	None	IS
21	GPC	IS
22	Column Chromatography, PAHs & biomarkers - 5% deactivated silica gel; PAHs-5% deactivated silica gel, 2%-deactivated alumina, biobead.	IS
23	Silica gel/Alumina columns and HPLC Phenogel	IS
24a	GPC	IS
24b	None	ES
25	None	IS
26	EPA 3630	IS
27	silica/alumina column chromatography, HPLC (Phenomenex 100A columns), reduction to 0.5 mL in water bath	IS
28	Silica/Alumina Column Chromatography: Size Exclusion High-Performance Liquid Chromatography	IS
29	Extracts were filtered through silicone-treated filter papers and sodium sulfate, GPC'd, and cleaned with alumina.	IS
30	SW 846 3630 modified - silica gel	IS
31		IS
32	Silica Gel	IS
33	Alumina/Silica Cleanup Column followed by 12 - GPC Cleanup	IS
34	pAH 3610 - alumina cleanup, followed by 3640 - GPC cleanup, Biomarker: 3610 - alumina cleanup, followed by 3640 - GPC cleanup, followed by 3630 Silica fractionation, F1 portion collected for Biomarkers	IS

Analytical methods used for PAHs

Lab #	Instrument	PAHs Phase	Dimensions	mode of injection (split/splitless/on-column)	Calibration	
					# points	Curve range*
1	GC/MS	DB-17MS	60m x 0.25mm, 0.25μm film	on-column	6	varied by compound
2	HPLC/FID	Zorbax PAH	5 cm x 4.6 mm, 1.8 um part	10 ul direct	3	2.5 ppb to 50 ppb
3	GC/MS	DB-5	60m x 0.25mm, 0.25μm film	splitless	6	0.1 ug/mL to 4.0 ug/mL
4	GC/MS	ZB-5MS	30m x 0.25mm, 0.25μm film	split/splitless	10	2 ppb to 2000ppb
5	GC/MS	DB-5	60m x 0.25mm, 0.25μm film	on-column	6	0.001 ng/μL to 0.33 ng/μL
6	LC	Zorbax Eclipse PAH Rapid Resolution HT	50 mm x 4.6 mm, 1.8 um part		3	2.5 ng/g to 50 ng/g
7	GC/MS	DB-5	30m x 0.25mm, 0.25μm film	on-column	7	10ng/mL to 4000ng/mL
8	GC/MS	Crossbond® silarylene phase; similar to polymeric C18 50% phenyl/50% dimethyl polysiloxane	30m x 0.25mm, 0.25μm film	LVI/ solvent vent	6	1ng/mL to 500 ng/mL (0.2 ng/g to 100ng/g)
9	HPLC	polymeric C18	501mm x 4.6 mm, 1.8 um part	on-column	3	2.5 mg/mL - 50 mg/mL
10	GC/MS	ZB-MS-5si	60m x 0.25mm, 0.25μm film		6	0.1 ug/mL to 4.0 ug/mL
11	GC/MS	DB-5MS	60m x 0.25mm, 0.25μm film	on-column	6	1 ng/mL to 300ng/mL
12	GC/MS	ZB-5MSi	60m x 0.25mm, 0.25μm film	splitless	6	0.008 ug/ml to 0.80 ug/ml
13	GC/MS	DB-5	30m x 0.25mm, 0.25μm film		6	5 ng/mL to 1000 ng/mL
14	GC/MS	Rxi-17 sil	60m x 0.25mm, 0.25μm film	splitless	6	0.1 ng/g to 1000 ng/g
15	GC/MS	DB-5	30m x 0.25mm, 0.25μm film	splitless	6	0.05 mg/L to 5.0 mg/L
16	GC/MS	DB-5	60m x 0.25mm, 0.25μm film	splitless	7	0.01 to 7.2
17	GC/MS	Rtx-5MS	30m x 0.32mm, 0.25μm film	split	8	4ug/g - 1000ug/g
18	GC/MS	DB-EUPAH CF	20m x 0.18mm, 0.14μm film	pulsed splitless	6	0.5pg/uL - 100pg/uL
19	GC/MS	5% phenyl/polydimethyl	60m x 0.25mm, 0.25μm film	splitless	6	0.1 ug/ml to 4 ug/ml
20	GC/MS	ZB5MSi	60m x 0.25mm, 0.25μm film		6	0.02 ug/L to 0.8 ug/L
21	QQQ	HP-5MS	30m x 0.25mm, 0.25μm film	pulsed splitless	7 or 8	1 ppb to 500 ppb
22	GC/MS	RTX-5	30m x 0.25mm, 0.25μm film	split/splitless	5	50 ng/mL to 5000ng/mL
23	GC/MS	DB5-MS	30m x 0.25mm, 0.25μm film	split/splitless	5	20 ng/mL to 1000 ng/mL
24-a	GC/MS	DB-5MS	60m x 0.25mm, 0.25μm film	on-column	6	1ng/mL to 330 ng/mL
24b	LC-FID	C18 Rapid resolution	501mm x 4.6 mm, 1.8 um part	on-column	3	2.5 μg/mL to 50 μg/mL
25	GC/MS	ZB-5MSi	60m x 0.25mm, 0.25μm film	pulsed splitless	6	0.1 ug/m to -4.0 ug/m
26	GC/MS	ZB-5MS	30m x 0.25mm, 0.25μm film	splitless	10	2.0 ng/mL to 2000 ng/mL
27	GC/MS	HP-5MS	60m x 0.25mm, 0.25μm film	split	5	(20, 100, 250, 500, 1000)
28	GC/MS	DB-5	60m x 0.25mm, 0.25μm film		6	3 ppm to 333 ppm
29	GC/MS	DB-5MS	30m x 0.25mm, 0.25μm film	PTV	6	10ng/mL to 5000ng/mL
30	GC/MS	5MS	30m x 0.25mm, 0.25μm film	split/splitless	5	0.050 ug/mL to 10ug/mL
31	GC/MS	ZB-5msi	60m x 0.25mm, 0.25μm film	splitless	6	0.1 ppm to 4.0 ppm
32	GC/MS	ZB5 ms	30m x 0.25mm, 0.5μm film	splitless	6	0.1-1.0
33	GC/MS	DB-5	60m x 0.25mm, 0.25μm film	on-column	6	1 ppb to 300 ppb
34	GC/MS-SIM	ZB-5	60m x 0.25mm, 0.25μm film	splitless	7	10 ng/mL to 20,000 ng/mL

*Units are those reported by the participating laboratory

Analytical methods used for Alkylated PAHs

Lab #	Instrument	Alkylated PAH		Dimensions	mode of injection (split/splitless on-column)	Calibration Curve		range*
		Phase	# points					
1	GC/MS	DB-17MS	60m x 0.25mm, 0.25µm film	on-column	6			varied by compound
2								
3	GC/MS	DB-5	60m x 0.25mm, 0.25µm film	splitless	6			0.1 µg/mL to 4.0 µg/mL
4	GC/MS	ZB-5MS	30m x 0.25mm, 0.25µm film	split/splitless				
5	GC/MS	DB-5	60m x 0.25mm, 0.25µm film	on-column	6			0.001ng/µL to 0.33 ng/µL
6								
7	GC/MS	DB-5	30m x 0.25mm, 0.25µm film	on-column	7			10ng/mL to 400ng/mL
8	GC/MS	Crossbond® silarylene phase; similar to 50% phenyl/50% dimethyl polysiloxane	30m x 0.25mm, 0.25µm film	LV/V solvent vent	6			1ng/mL to 500 ng/mL (0.2 ng/g to 100ng/g)
9								
10	GC/MS	ZB-MS-5Si	60m x 0.25mm, 0.25µm film		6			0.1 µg/mL to 4.0 µg/mL
11	GC/MS	DB-5MS	60m x 0.25mm, 0.25µm film	on-column	6			1 ng/mL to 300ng/mL
12	GC/MS	ZB-5MSi	60m x 0.25mm, 0.25µm film	splitless	6			0.008 µg/mL to 0.80 µg/mL
13	GC/MS	DB-5	30m x 0.25mm, 0.25µm film		6			5 ng/mL to 1000 ng/mL
14	GC/MS	Rxi-17 sil	60m x 0.25mm, 0.25µm film	splitless	6			0.1 ng/g to 1000 ng/g
15	GC/MS	DB-5	30m x 0.25mm, 0.25µm film	splitless	6			0.05 mg/L to 5.0 mg/L
16	GC/MS	DB-5	60m x 0.25mm, 0.25µm film	splitless	7			0.01 to 7.2
17	GC/MS	Rtx-2MS	30m x 0.32mm, 0.25µm film	split	8			4ug/g to 10000ug/g
18	GC/MS	DB-EUPAH-1CF	20m x 0.18mm, 0.14µm film	pulsed splitless	6			0.5pg/µL to 100pg/µL
19	GC/MS	5% Phenyl/polydimethyl	60m x 0.25mm, 0.25µm film	splitless	6			0.1 ug/mL to 4 ug/mL
20	GC/MS	ZB5MSi	60m x 0.25mm, 0.25µm film		6			0.02 ug/L to 0.8 ug/L
21	GC/MS QQQ	HP-5MS	30m x 0.25mm, 0.25µm film	pulsed splitless	8			1ppb to 500 ppb
22	GC/MS	RTX-5	30m x 0.25mm, 0.25µm film	split/splitless	1			2000 ng/mL
23	GC/MS	DB5-MS	30m x 0.25mm, 0.25µm film	split/splitless	5			20ng/mL to 1000 ng/mL
24a	GC/MS	DB-5MS	60m x 0.25mm, 0.25µm film	on-column	6			1 ng/mL to 330 ng/mL
24b								
25	GC/MS	ZB-5MSi	60m x 0.25mm, 0.25µm film	pulsed splitless	6			0.1 ug/mL to 4.0 ug/mL
26	GC/MS	ZB-5MS	30m x 0.25mm, 0.25µm film	splitless	10			2.0 ng/mL to 2000 ng/mL
27	GC/MS	HP-5MS	60m x 0.25mm, 0.25µm film	split	5			(20, 100, 250, 500, 1000) ng/mL
28	GC/MS	DB-5	60m x 0.25mm, 0.25µm film		6			3 ppm to 333 ppm
29	GC/MS	DB-5MS	30m x 0.25mm, 0.25µm film	PTV	6			10ng/mL to 500ng/mL
30	GC/MS	5MS	30m x 0.25mm, 0.25µm film	split/splitless	5			0.050 ug/mL to 100ug/mL
31	GC/MS	ZB-5MSi	60m x 0.25mm, 0.25µm film	splitless	6			0.1 ppm to 4.0 ppm
32	GC/MS	ZB5 msii	30m x 0.25mm, 0.5µm film	splitless	6			0.1-1.0
33	GC/MS	DB-5	60m x 0.25mm, 0.25µm film	on-column	6			1ppb to 300 ppb
34	GC/MS-SIM	ZB-5	60m x 0.25mm, 0.25µm film	splitless	7			10 ng/mL to 20,000 ng/mL

*Units are those reported by the participating laboratory

Analytical methods used for Biomarkers

Lab #	Instrument	Biomarkers Phase	Dimensions	mode of injection		# points	Calibration Curve	range*
				(split/splitless/on-column)	on-column			
1	GC/MS	DB-17MS	60m x 0.25mm, 0.25μm film			4		varied by compound
2		DB-5	60m x 0.25mm, 0.25μm film	splitless		6		0.1 μg/mL to 4.0 μg/mL
3								
4								
5								
6								
7								
8								
9								
10	GC/MS	ZB-MS-5si	60m x 0.25mm, 0.25μm film			6		0.1 μg/mL to 4.0 μg/mL
11								
12	GC/MS	ZB-5MSi	60m x 0.25mm, 0.25μm film	splitless		6		0.008 μg/mL to 0.80 μg/mL
13								
14	GC/MS	Rxi-17 sil	60m x 0.25mm, 0.25μm film	splitless		6		0.1 ng/g to 2000 ng/g
15	GC/MS	DB-5	30m x 0.25mm, 0.25μm film	splitless		1		SINTEF reference oil
16	GC/MS	DB-5	60m x 0.25mm, 0.25μm film	splitless		6 to 7		0.03 to 7.2 / 0.01 to 7.2
17	GC/MS	Rtx-2MS	30m x 0.32mm, 0.25μm film	split		8		4 μg/g to 10000 μg/g
18								
19	GC/MS	5% phenyl/polydimethyl	60m x 0.25mm, 0.25μm film	splitless		6		0.1 μg/mL to 4 μg/mL
20	GC/MS	ZB5MSi	60m x 0.25mm, 0.25μm film			6		0.02 μg/L to 0.8 μg/L
21								
22	GC/MS	Rtx-5	30m x 0.25mm, 0.25μm film	split/splitless		1		300 ng/mL to 11500ng/mL
23								
24a								
24b								
25	GC/MS	ZB-5MSi	60m x 0.25mm, 0.25μm film	pulsed splitless		6		0.1 μg/mL to 4.0 μg/mL
26								
27								
28								
29								
30								
31	GC/MS	ZB-5msi	60m x 0.25mm, 0.25μm film	splitless		6		0.1ppm to 4.0ppm
32								
33								
34	GC/MS-SIM	ZB-5	60m x 0.25mm, 0.25μm film	splitless		6		10 ng/mL to 10,000 ng/mL

*Units are those reported by the participating laboratory

Quantification of PAHs

		PAHs			
Lab #	[S] surrogate added prior to extraction	Used?	added prior to analysis	Used?	recovery?
1	naphthalene-d8, biphenyl-d10; acenaphthalene-d10; phenanthrene-d10; fluoranthene-d10; pyrene-d10; B[a]A-d12; B[a]P-d12; perylene-d12; B[ghi]P-d12; DB[a,h]A-d14	x			
2					
3	Nitrobenzene-d5, 2-Fluorobiphenyl, Tephenyl-d14		Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Chrysene-d10	x	no
4	surrogates only		Internal standards		
5	NPH-d8, ACE-d10, BaP-d12	x	HMB		PHN-d10 (prior to size-exclusion LC)
6					
7	Deuterated-Primary PAHs (16 total)	x	d10-2-Methylnaphthalene and d12-Benz(a)pyrene		
8	13C labeled EPA 16 PAH @ 10ng/g	x			
9					
10	Nitrobenzene-d5, 2-Fluorobiphenyl; Tephenyl-d14		Naphthalene-d8, Acenaphthalene-d10; Chrysene-d12	x	
11	Naphthalene-d8, Acenaphthalene-d10 and Benzo(a)pyrene-d12	x			yes
12	Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Chrysene-d12, Perylene-d12		Fluorene-d10, Benzo(a)pyrene-d12	x	no
13	Acenaphthene-d10, Acenaphthylene-d8, Anthracene-d10, Benz[a]Anthracene-d12, Benz[b]Fluoranthene-d12, Benz[k]Fluoranthene-d12, Benzog[h,j]Perylene-d12, Benz[a,z]Pyrene-d12, Chrysene-d12, Dibenz[a,b]Anthracene-d14, Fluoranthene-d10, Fluorene-d10, Indenol[1,2,3-cd]Pyrene-d12, Naphthalene-d8, Phenanthrene-d10, Pyrene-d10				
14	Naphthalene-d8; Acenaphthene-d10; Phenanthrene-d10; Fluoranthene-d10; Pyrene-d10; Benz[a]anthracene-d12; Benzo[a]pyrene-d12; Perylene-d12; Bibenz[a,h]anthracene-d14; Benzo[[ghi]pyrene-d12]	x			
15	naphthalene-d8, acenaphthylene-d8, phenanthrene-d10, pyrene-d10, benzo(a)pyrene-d12, benzodghi)perylene-d12		2-fluorobiphenyl, p-terphenyl-d14, chrysene-d12, dibenz[oh]anthracene-d14	x	
16	Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Benzo(a)pyrene-d12		Fluorene-d10, Chrysene-d12	x	yes
17	Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, 5 alpha-Androstanone, Chrysene-d12, Perylene-d12		Fluorene-d10, Pyrene-d10, Benzo(a)pyrene-d12	x	yes
18	Fluorene-d10, Fluoranthene-d10		Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Chrysene-d12, Perylene-d12	x	no

Quantification of PAHs (cont)

			PAHs	
Lab #	IS/surrogate added prior to extraction	Used?	added prior to analysis	corrected for recovery?
19	sur - 2-fluorobiphenyl, nitrobenzene-d5, terphenyl-d14		IS - Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Chrysene-d12, Perylene-d12	x no
20	Surrogates: Nitrobenzene-d5, 2-Fluorobiphenyl, Terphenyl-d14		Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Chrysene-d12, Perylene-d12	
21	Surrogates: Nitrobenzene-d5, 2-Fluorobiphenyl, Terphenyl-d14		Internal standards: Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Chrysene-d12, Perylene-d12	x no
22	d8-naphthalene, d10-2-methylnaphthalene, d10-biphenyl, d12-2,6-dimethylnaphthalene, d8-acenaphthylene, d10-phenanthrene, d10-fluoranthene, d12-benz[a]anthracene, d12-chrysene, d12-benz[b,k]fluoranthene, d12-indeno[1,2-e]pyrene, d12-benz[a,h]anthracene, d12-indeno[1,2-e]pyrene, d12-hexzo[gh]perylene	x	d10-acenaphthene, d10-pyrene, d12-benz[e]pyrene, used to quantify labeled surrogates only.	
23	d8-Naphthalene, d10-Acenaphthene, d10-Phenanthrene, d12-Chrysene, d12-Perylene	x	d10-Fluorene, d12-Benzo(a)pyrene	
24a	Naphthalene-d8, Acenaphthalene-d10, Benzo(a)pyrene-d12	x	Hexamethylbenzene	
24b	2-Fluorophenyl, phenol-d5, nitrobenzene-d5, 2,4,6-tribromophenol, Terphenyl-d14, 2-Fluorobiphenyl		1,4-dichlorobenzene-d4, Naphthalene-d8, Acenaphthene-d10, Phenanthrene-d10, Chrysene-d12, Perylene-d12	x yes
25				
26	Fluorene-d10, Fluoranthene-d10, Terphenyl-d14			
27	Surrogates: Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Chrysene-d12, Perylene-d12	x	Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Chrysene-d12, Perylene-d12	IS yes
28	naphthalene-d8, acenaphthene-d10; benzo[e]pyrene-d12	x	IS-Fluorene-d10, Pyrene-d10, Benzo[a]pyrene-d12	x yes
				phenanthrene-d10 (prior to size-exclusion LC)
29	d8-Naphthalene, d8-1-Methylnaphthalene, d8-Acenaphthylene, d10-Fluorene, d8-Dibenzo[othiophene, d10-Phenanthrene, d10-Anthracene, d10-Fluoranthene, d10-Pyrene, d12-Benz[a]anthracene, d12-Chrysene, d12-Benzo[b]fluoranthene, d12-Benzo[k]fluoranthene, d12-Benz[e]pyrene, d12-Benzo[a]pyrene, d12-Perylene, d12-Benzo[g,h,i]perylene	x	d14-p-terphenyl	
30	2-fluorobiphenyl, terphenyl d14		naph d8, acenaph d10, phenan d10, chry d12, pery d12	x no
31	Nitrobenzene-d5, 2-fluorobiphenyl, Terphenyl-d14	x	Naphthalene-d8, acenaphthalene-d10, phenanthrene-d10, chrysene-d12, perylene-d12	
32	d8-Naphthalene, d10-Acenaphthene, d10-Phenanthrene, d12-Chrysene, d12-Perylene		d10-Fluorene, d10-Pyrene, d12-Benzo(a)pyrene	x no
33	D8Naphthalene, D10Acenaphthene, D12 BAP	x	Hexamethylbenzene	
34	Surrogates: Naphthalene-d8, Phenanthrene-d10, Benzo[a]pyrene-d12, 5b(H)Cholane	x	IS, Acenaphthene-d10, Chrysene-d12	x yes

Quantification of Alkylated PAHs

				Alkylated PAHs			corrected for
Lab #	IS/surrogate added prior to extraction	Used?		added prior to analysis	Used?	recovery?	others?
1	naphthalene-d8, biphenyl-d10, acenaphthalene-d10, phenanthrene-d10, fluoranthene-d10, pyrene-d10, B[a]A-d12, B[ghi]P-d12, B[ghi]P-d12,DB[a,h]A-d14	x					
2							
3	Nitrobenzene-d5, 2-Fluorobiphenyl, Terphenyl-d14			Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Chrysene-d10	x	no	
4	surrogates only			Internal standards			PhN-d10 (prior to size-exclusion LC)
5	NPH-d18, ACE-d10, BaP-d12		HMB				
6		x					
7	Deuterated-Primary PAHs (16 total)	x	d10-2-Methylnaphthalene and d12-Benzo(e)pyrene				
8	13C labeled EPA 16 PAH (@10ng/g	x					
9							
10	Nitrobenzene-d5, 2-Fluorobiphenyl, Terphenyl-d14		Naphthalene-d8; Acenaphthalene-d10, Chrysene-d12, Perylene-d12	x	yes		
11	Naphthalene-d8, Acenaphthalene-d10 and Benzo[a]pyrene-d12	x					
12	Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Chrysene-d12, Perylene-d12		Fluorene-d10, Benzo(a)pyrene-d12	x	no		
13							
14	Naphthalene-d8, Phenanthrene-d10, Fluoranthene-d10	x					
15	naphthalene-d8, acenaphthylene-d8, Phenanthrene-d10, pyrene-d10, benzo(a)pyrene-d10, benzo(gh)perylene-d12		2-fluorobiphenyl, p-terphenyl-d14, chrysene-d12, dibenz(a,h)anthracene-d14	x			
16	Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Benzo(a)pyrene-d12		Fluorene-d10, Chrysene-d12	x	yes		
17	Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, 5 alpha-Androstanone, Chrysene-d12, Perylene-d12		Fluorene-d10, Pyrene-d10, Benzo(a)pyrene-d12	x	yes		
18	1-methyl Naphthalene-d10			x	no		

Quantification of Alkylated PAHs (cont)

			Alkylated PAHs			
Lab #	IS/surrogate added prior to extraction	Used?	added prior to analysis	Used?	recovery?	corrected for
19	sur - 2-fluorobiphenyl, nitrobenzene-d5, terphenyl-d14		IS - Naphthalene-d8, Acenaphthalene-d10, Chrysene-d12, Perylene-d12	x	no	
20	Surrogates: Nitrobenzene-d5, 2-Fluorobiphenyl, Terphenyl-d14		Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Chrysene-d12; Perylene-d12			
21	Surrogates: Nitrobenzene-d5, 2-Fluorobiphenyl, Terphenyl-d14		Internal standards: Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Chrysene-d12.			
22	dl0-2-methylnaphthalene, d12-2,6-dimethylnaphthalene, d8-acenaphthylene, d10-pheanthrene, d10-2-chrysene	x	dl0-acenaphthene, d10-pyrene, d12-benz[e]pyrene, used to quantify labeled surrogates only.			
23	db3-Naphthalene, dl0-Acenaphthene, dl0-Phenanthrene, d12-Chrysene, d12-Perylene	x	dl0-Fluorene, d12-Benz[a]pyrene			
24a	Naphthalene-d8, Acenaphthene-d10, Benzo[a]pyrene-d12	x	Hexamethylbenzene			
24b						
25	2-Fluorophenyl, phenol-d5, nitrobenzene-d5, 2,4,6-tribromophenoI, Terphenyl-d14, 2-Fluorobiphenyl		1,4-dichlorobenzene-d4, Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Chrysene-d12, Perylene-d12	x	yes	
26	Fluorene-d10, Fluoranthene-d10, Terphenyl-d14		sur Naphthalene-d8, Acenaphthene-d10, Phenanthrene-d10, Chrysene-d12, Perylene-d12	IS	yes	
27	SI-S-Naphthalene-d8, Acenaphthene-d10, Phenanthrene-d10, Chrysene-d12, Perylene-d12		IS-Fluorene-d10, Pyrene-d10, Benzo[a]pyrene-d12	x	yes	
28	naphthalene-d8, acenaphthene-d10, benzo[a]pyrene-d12	x	hexamethylbenzene			
29	ds-Naphthalene, db-1-Methylnaphthalene, d10-Acenaphthene, d10-Fluorene, ds-Dibenzothiophene, dl0-Anthracene, d10-Fluoranthene, dl2-Chrysene, dl2-Benzo[e]pyrene	x	dl14-p-terphenyl			
30	2-fluorobiphenyl, terphenyl d14		napth d8, acenaph dl0, phenan dl0, chry dl2, pery dl2	x	no	
31	Nitrobenzene-d5, 2-fluorobiphenyl, Terphenyl-d14	x	Naphthalene-d8, acenaphthalene-d10, phenanthrene-d10, chrysene-d12, perylene-d12			
32	ds-Naphthalene, dl0-Acenaphthene, dl0-Phenanthrene, d12-Chrysene, d12-Perylene		dl0-Fluorene, dl0-Pyrene, d12-Benzo[a]pyrene	x	no	
33	DSS-Naphthalene, Dl0Acenaphthene, D12 BAP	x				
34	Surrogates: Naphthalene-d8, Phenanthrene-d10, Benzo[a]pyrene-d12, 5b(H)Cholane	x	IS- Acenaphthene-d10, Chrysene-d12	x	yes	

Quantification of Biomarkers

Lab #	IS/surrogate added prior to extraction	Biomarkers		Used?	recovery?	corrected for
		Used?	added prior to analysis			
1	naphthalene-d8; biphenyl-d10; acenaphthalene-d10; phenanthrene-d10; fluoranthene-d10; pyrene-d10; B[a]A-d12; B[a]P-d12; perylene-d12; B[ghi]P-d12; DB[a,h]A-d14	x				
2						
3	Nitrobenzene-d5; 2-Fluorobiphenyl; Terphenyl-d14		Naphthalene-d8; Acenaphthalene-d10; Phenanthrene-d10; Chrysene-d10	x	no	
4						
5						
6						
7						
8						
9						
10	Nitrobenzene-d5; 2-Fluorobiphenyl; Terphenyl-d14		Naphthalene-d8; Acenaphthalene-d10; Chrysene-d12; Perylene-d12	x	yes	
11						
12	Naphthalene-d8; Acenaphthalene-d10; Phenanthrene-d10; Chrysene-d12; Perylene-d12		Fluorene-d10; Benzo(a)pyrene-d12	x	no	
13						
14	n-triacontane-d162	x				
15						
16	Sb(II)-Cholane		Chrysene-d12	x	yes	
17	Naphthalene-d8; Acenaphthalene-d10; Phenanthrene-d10; 5 alpha-Aandrostan, Chrysene-d12; Perylene-d12		Fluorene-d10; Pyrene-d10; Benzo(a)pyrene-d12	x	yes	
18						

Quantification of Biomarkers (cont)

Lab #	IS/surrogate added prior to extraction	Biomarkers		Used?	Used prior to analysis IS - Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Chrysene-d12, Perylene-d12	Used? recovery? x no	corrected for recovery?
		Used?	Biomarkers				
19	sur - 2-fluorobiphenyl, nitrobenzene-d5, terphenyl-d14						
20	Surrogates: Nitrobenzene-d5, 2-fluorobiphenyl; Terphenyl-d14		Naphthalene-d8, Acenaphthalene-d10; Phenanthrene-d10; Chrysene-d12; Perylene-d12				
21							
22	d10-phenanthrene, d50-tetracosane	x	d10-pyrene, used to quantify labeled surrogates only				
23							
24							
25	2-Fluorophenyl, phenol-d5, nitrobenzene-d5, 2,4,6-tribromophenol, Terphenyl-d14, 2-[Fluorobiphenyl]		1,4-dichlorobenzene-d4, Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Chrysene-d12, Perylene-d12				
26	Fluorene-d10, Fluoranthene-d10, Terphenyl-d14	sur	Naphthalene-d8, Acenaphthalene-d10, Phenanthrene-d10, Chrysene-d12, Perylene-d12	IS	IS	yes	
27							
28							
29							
30							
31	Nitrobenzene-d5, 2-fluorobiphenyl, Terphenyl-d14	x	Naphthalene-d8, acenaphthalene-d10,phenanthrene-d10, chrysene-d12, perylene-d12				
32							
33							
34	Surrogates: Naphthalene-d8, Phenanthrene-d10, Benzo[a]pyrene-d12, 5b(H)Cholane	x	IS; Acenaphthalene-d10, Chrysene-d12	x	yes		

PAHs - IS/surrogate used for quantitation

	Lab 1	Lab 2	Lab 3	Lab 4	Lab 5	Lab 6
naphthalene	naphthalene-d ₈		Naphthalene-d ₈	Naphthalene-d ₈	Naphthalene-d ₈	
biphenyl	biphenyl-d ₁₀		Naphthalene-d ₈	Saphthalene-d ₈	ACE-d ₁₀	
acenaphthene	acenaphthene-d ₁₀		Acenaphthene-d ₁₀	Acenaphthene-D ₁₀	ACE-d ₁₀	
acenaphthylene	acenaphthylene-d ₁₀		Acenaphthylene-d ₁₀	Acenaphthene-D ₁₀	ACE-d ₁₀	
fluorene	phenanthrene-d ₁₀		Acenaphthene-d ₁₀	Acenaphthene-D ₁₀	ACE-d ₁₀	
phenanthrene	phenanthrene-d ₁₀		Phenanthrene-d ₁₀	Phenanthrene-D ₁₀	ACE-d ₁₀	
anthracene	phenanthrene-d ₁₀		Phenanthrene-d ₁₀	Phenanthrene-D ₁₀	ACE-d ₁₀	
fluoranthene	fluoranthene-d ₁₀		Phenanthrene-d ₁₀	Phenanthrene-D ₁₀	BaP-d ₁₂	
pyrene	pyrene-d ₁₀		Chrysene-d ₁₂	Chrysene-D ₁₂	BaP-d ₁₂	
benzo[b]fluorene	NA		NA	Perylene-D ₁₂		
benzo[a]anthracene	Bi[_a]A-d ₁₂		Chrysene-d ₁₂	Chrysene-D ₁₂	BaP-d ₁₂	
chrysene	Bi[_a]A-d ₁₂		Chrysene-d ₁₂	Chrysene-D ₁₂	BaP-d ₁₂	
benzo[a]phenylene	Bi[_a]A-d ₁₂		NA			
benz[<i>h</i>] fluoranthene	Bi[_a]P-d ₁₂		Chrysene-d ₁₂	Perylene-D ₁₂	BaP-d ₁₂	
benzo[<i>f</i>] fluoranthene	Bi[_a]P-d ₁₂		NA			
benzo[<i>k</i>] fluoranthene	Bi[_a]P-d ₁₂		Chrysene-d ₁₂	Perylene-D ₁₂	BaP-d ₁₂	
benzo[<i>al</i>] fluoranthene	Bi[_a]P-d ₁₂		NA			
benzo[<i>el</i>] pyrene	Bi[_a]P-d ₁₂		Chrysene-d ₁₂	Perylene-D ₁₂	BaP-d ₁₂	
benzo[<i>al</i>] pyrene	Bi[_a]P-d ₁₂		Chrysene-d ₁₂	Perylene-D ₁₂	BaP-d ₁₂	
perylene	perylene-d ₁₂		Chrysene-d ₁₂	Perylene-D ₁₂	BaP-d ₁₂	
indeno[1,2,3- <i>cd</i>]pyrene	B[_{gh}]P-d ₁₂		Chrysene-d ₁₂	Perylene-D ₁₂	BaP-d ₁₂	
benzo[<i>ghi</i>]perylene	B[_{gh}]P-d ₁₂		Chrysene-d ₁₂	Perylene-D ₁₂	BaP-d ₁₂	
dibenz[<i>ah</i>]anthracene	DB[_{ah}]A-d ₁₄		Chrysene-d ₁₂	Perylene-D ₁₂	BaP-d ₁₂	
cis/trans-decalin	NA		NA	Naphthalene-d ₈		
dibenzofuran	NA		Acenaphthene-d ₁₀	Acenaphthene-D ₁₀		
retene	NA		NA	Chrysene-D ₁₂	ACE-d ₁₀	
benzothiophene	NA		NA	Naphthalene-d ₈		
dibenzothiophene	fluoranthene-d ₁₀		Acenaphthene-d ₁₀	Phenanthrene-D ₁₀	ACE-d ₁₀	
naphthalenethiophene	NA		NA	Chrysene-D ₁₂		

PAHs - IS/surrogate used for quantitation

	Lab 7	Lab 8	Lab 9	Lab 10	Lab 11	Lab 12
naphthalene	d8-Naphthalene	13C-naphthalene		Naphthalene-d8	Naphthalene-d8	Fluorene-d10
phenyl	d10-Acenaphthylene	NA		Acenaphthene-d10		Fluorene-d10
acenaphthene	d10-Acenaphthene	13C-acenaphthene		Acenaphthene-d10		Fluorene-d10
acenaphthylene	d10-Acenaphthylene	13C-acenaphthylene		Acenaphthene-d10		Fluorene-d10
fluorene	d10-Fluorene	13C-Fluorene		Acenaphthene-d10	Acenaphthene-d10	Fluorene-d10
phenanthrene	d10-Phenanthrene	13C-phenanthrene		Phenanthrene-d10	Acenaphthene-d10	Fluorene-d10
anthracene	d10-Aanthracene	13C-anthracene		Phenanthrene-d10	Aceanthracene-d10	Fluorene-d10
fluoranthene	d10-Fluoranthene	13C-Fluoranthene		Phenanthrene-d10	Aceanthracene-d10	Fluorene-d10
pyrene	d10-Pyrene	13C-pyrene		Chrysene-12	Aceanthracene-d10	Benz(a)pyrene-d12
benz[b]fluorene		NA		NA		Benz(a)pyrene-d12
benz[a]anthracene	d12-Benz[a]anthracene	13C-benz[a]anthracene		Chrysene-12	Benz[a]pyrene-d12	Benz(a)pyrene-d12
chrysene	d12-Chrysene	13C-chrysene		Chrysene-12	Benz[a]pyrene-d12	Benz(a)pyrene-d12
phenanthrene		NA		NA		Benz(a)pyrene-d12
benz[b]fluoranthene	d12-Benz[b]fluoranthene	13C-benz[b]fluoranthene		Chrysene-12	Benz[a]pyrene-d12	Benz(a)pyrene-d12
benz[a]fluoranthene		NA		NA		Benz(a)pyrene-d12
benz[a]anthracene	d12-Benz[a]anthracene	13C-benz[a]anthracene		Chrysene-12	Benz[a]pyrene-d12	Benz(a)pyrene-d12
benz[a]fluoranthene		NA		NA		Benz(a)pyrene-d12
benz[a]anthracene		NA		NA		Benz(a)pyrene-d12
benz[b]fluoranthene		NA		NA		Benz(a)pyrene-d12
benz[a]anthracene		NA		NA		Benz(a)pyrene-d12
benz[a]anthracene		NA		NA		Benz(a)pyrene-d12
benz[a]anthracene		NA		NA		Benz(a)pyrene-d12
benz[a]anthracene		NA		NA		Benz(a)pyrene-d12
terephene	d12-Terphenyl	13C-terphenyl		Chrysene-12	Benz[a]pyrene-d12	Benz(a)pyrene-d12
indeno[1,2,3-cd]pyrene	d12-Indeno[1,2,3-cd]pyrene	13C-indeno[1,2,3-cd]pyrene		Chrysene-12	Benz[a]pyrene-d12	Benz(a)pyrene-d12
benz[e]phenylene	d12-Benz[e]phenylene	13C-benz[e]phenylene		Chrysene-12	Benz[a]pyrene-d12	Benz(a)pyrene-d12
allophen[a]anthracene	d14-Allophen[a]anthracene	13C-dibenz[a]anthracene		Chrysene-12	Benz[a]pyrene-d12	Benz(a)pyrene-d12
cyclofrans-decalin		NA		NA		Fluorene-d10
dibenzofuran		NA		NA		Fluorene-d10
retene		NA		NA		
benzothiophene	d8-Naphthalene	NA		NA		Fluorene-d10
dibenzothiophene	d10-Fluorene	NA		Phenanthrene-d10		Fluorene-d10
naphthalenozethiophene		NA		NA		Benz(a)pyrene-d12

PAHs - IS/surrogate used for quantitation

	Lab 13	Lab 14	Lab 15	Lab 16	Lab 17	Lab 18
naphthalene	naphthalene-d8		2-fluorobiphenyl	Fluorene-d10 / Naphthalene-d8	Fluorene-d10 / Naphthalene-d8	Naphthalene-d8
phenyl	naphthalene-d8		2-fluorobiphenyl	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Acenaphthene-d10	Naphthalene-d8
acenaphthene	acenaphthene-d10	naphthalene-d8	2-fluorobiphenyl	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Acenaphthene-d10	Acenaphthene-d10
acenaphthylene	acenaphthylene-d18	naphthalene-d8	2-fluorobiphenyl	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Acenaphthene-d10	Naphthalene-d8
fluorene	fluorene-d10	naphthalene-d8	2-fluorobiphenyl	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Acenaphthene-d10	Acenaphthene-d10
phenanthrene	phenanthrene-d10	phenanthrene-d10	2-fluorobiphenyl	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Phenanthrene-d10	Phenanthrene-d10
anthracene	anthracene-d10	phenanthrene-d10	2-fluorobiphenyl	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Phenanthrene-d10	Phenanthrene-d10
fluoranthene	fluoranthene-d10	fluoranthene-d10	p-phenyl-d14	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	Phenanthrene-d10
pyrene	pyrene-d10	pyrene-d10	p-phenyl-d14	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	Phenanthrene-d10
benzo(b)fluorene			NA		Pyrene-d10 / Chrysene-d12	
benz(2)anthracene	benz(2)anthracene-d12	benz(2)anthracene-d12	chrysene-d12	Chrysene-d12 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	Phenanthrene-d10
chrysene	chrysene-d12	benz(2)anthracene-d12	chrysene-d12	Chrysene-d12 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	Chrysene-d12
triphenylene	benz(2)anthracene-d12	benz(2)anthracene-d12	NA	Chrysene-d12 / Benz(a)pyrene-d12	Pyrene-d10 / Chrysene-d12	Chrysene-d12
benzo(b)fluoranthene	benzo(b)fluoranthene-d12	benzo(a)pyrene-d12	chrysene-d12	Chrysene-d12 / Benz(a)pyrene-d12	Benz(a)pyrene-d12 / Perylene-d12	Perylene-d12
benzo(f)fluoranthene		benzo(a)pyrene-d12	chrysene-d12	Chrysene-d12 / Benz(a)pyrene-d12	Benz(a)pyrene-d12 / Perylene-d12	Perylene-d12
benzo(k)fluoranthene	benzo(k)fluoranthene-d12	benzo(a)pyrene-d12	chrysene-d12	Chrysene-d12 / Benz(a)pyrene-d12	Benz(a)pyrene-d12 / Perylene-d12	Perylene-d12
benzo(a)fluoranthene		benzo(a)pyrene-d12	NA	Chrysene-d12 / Benz(a)pyrene-d12	Benz(a)pyrene-d12 / Perylene-d12	Perylene-d12
benzo(a)fluoranthene	benzo(a)fluoranthene-d12	benzo(a)pyrene-d12	chrysene-d12	Chrysene-d12 / Benz(a)pyrene-d12	Benz(a)pyrene-d12 / Perylene-d12	Perylene-d12
perylene	benzo(a)pyrene-d12	perylene-d12	chrysene-d12	Chrysene-d12 / Benz(a)pyrene-d12	Benz(a)pyrene-d12 / Perylene-d12	Perylene-d12
indeno[1,2,3-cd]pyrene	indeno[1,2,3-cd]pyrene-d12	benz(g)perylene-d12	dibenz(a,h)anthracene-d14	Chrysene-d12 / Benz(a)pyrene-d12	Benz(a)pyrene-d12 / Perylene-d12	Perylene-d12
benz(g)phenylene	benz(g)phenylene-d12	perylene-d12	dibenz(a,h)anthracene-d14	Chrysene-d12 / Benz(a)pyrene-d12	Benz(a)pyrene-d12 / Perylene-d12	Perylene-d12
phenanthrene	phenanthrene-d14	dibenz(a,h)anthracene-d14	dibenz(a,h)anthracene-d14	Chrysene-d12 / Benz(a)pyrene-d12	Benz(a)pyrene-d12 / Perylene-d12	Perylene-d12
cis/trans-decalin			NA	Fluorene-d10 / Naphthalene-d8	Fluorene-d10 / Naphthalene-d8	
dibenzofuran	acenaphthylene-d8		NA	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Acenaphthene-d10	
retene		fluoranthene-d10	p-phenyl-d14	Fluorene-d10 / Chrysene-d12	Pyrene-d10 / Acenaphthene-d10	
benzohopophene			NA	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Naphthalene-d8	
dihenzothiophene	fluorene-d10	phenanthrene-d10	p-phenyl-d14	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Phenanthrene-d10	Acenaphthene-d10
naphthalenethiophene			NA	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	

PAHs - ls/surrogate used for quantitation

	Lab 19	Lab 20	Lab 21	Lab 22	Lab 23	Lab 24a	Lab 24b
naphthalene	Naphthalene-d8	Naphthalene-d8	naphthalene-d8	d8-naphthalene	d8-Naphthalene	NA	NA
phenyl	Naphthalene-d8	Acenaphthene-d10		d10[4-phenyl]	d10-Acenaphthene	NA	NA
acenaphthene	Acenaphthalene-d10	Acenaphthene-d10	acenaphthene-d10	d8-acenaphthylene	d10-Acenaphthene	NA	NA
acenaphthylene	Acenaphthalene-d10	Acenaphthene-d10	acenaphthene-d10	d8-acenaphthylene	d10-Acenaphthene	Acenaphthene-d10	Acenaphthene-d10
fluorene	Acenaphthalene-d10	Acenaphthene-d10	acenaphthene-d10	d10-phenanthrene	d10-Acenaphthene	Acenaphthene-d10	Acenaphthene-d10
phenanthrene	Phenanthrene-d10	Phenanthrene-d10	phenanthrene-d10	d10-phenanthrene	d10-Phenanthrene	Acenaphthene-d10	Acenaphthene-d10
anthracene	Phenanthrene-d10	Phenanthrene-d10	phenanthrene-d10	d10-phenanthrene	d10-Phenanthrene	Acenaphthene-d10	Acenaphthene-d10
fluoranthene	Phenanthrene-d10	Phenanthrene-d10	phenanthrene-d10	d10-fluoranthene	d10-Phenanthrene	Acenaphthene-d10	Acenaphthene-d10
pyrene	Chrysene-d12	Chrysene-d12	chrysene-d12	d10-fluoranthene	d10-Phenanthrene	Acenaphthene-d10	Acenaphthene-d10
benzo[b]fluoranthene				NA		Benzo[b]pyrene-d12	Benzo[b]pyrene-d12
benz[a]anthracene	Chrysene-d12	Chrysene-d12	chrysene-d12	d12-phenanthrene	d12-Chrysene	Benzo[a]pyrene-d12	Benzo[a]pyrene-d12
chrysene	Chrysene-d12	Chrysene-d12	chrysene-d12	d12-chrysene	d12-Chrysene	NA	NA
triphenylene			other			Benzo[b]pyrene-d12	Benzo[b]pyrene-d12
benzo[b]fluoranthene	Chrysene-d12	Chrysene-d12	perylene-d12	d12-benz[b]fluoranthene	d12-Chrysene	NA	NA
benzo[f]fluoranthene			other			Benzo[b]pyrene-d12	Benzo[b]pyrene-d12
benzo[k]fluoranthene	Chrysene-d12	Chrysene-d12	perylene-d12	d12-benz[k]fluoranthene	d12-Chrysene	NA	NA
benzo[a]fluoranthene			NA			NA	NA
benzo[e]pyrene	Chrysene-d12	Chrysene-d12		d12-benz[e]pyrene	d12-Chrysene	Benzo[e]pyrene-d12	Benzo[e]pyrene-d12
benzo[a]pyrene	Chrysene-d12	Chrysene-d12	perylene-d12	d12-benz[a]pyrene	d12-Chrysene	NA	NA
perylene	Chrysene-d12	Chrysene-d12		d12-perylene	d12-Perylene	Benzo[b]pyrene-d12	Benzo[b]pyrene-d12
indeno[1,2,3- <i>cd</i>]pyrene	Chrysene-d12	Chrysene-d12	perylene-d12	d12-indeno[1,2,3- <i>cd</i>]pyrene	d12-Chrysene	NA	NA
benzo[ghi]perylene	Chrysene-d12	Chrysene-d12	perylene-d12	d12-benz[ghi]perylene	d12-Chrysene	Benzo[ghi]perylene-d12	Benzo[ghi]perylene-d12
dibenz[<i>a,h</i>]anthracene	Chrysene-d12	Chrysene-d12	perylene-d12	d14-dibenz[<i>a,h</i>]anthracene	d12-Chrysene	NA	NA
cistram-declinin				NA		NA	NA
dibenzofuran	Acenaphthalene-d10	Acenaphthene-d10		NA		NA	NA
retene				d10-fluoranthene		NA	NA
benzothiophene				NA		NA	NA
dibenzothiophene				d10-phenanthrene		NA	NA
naphthalenethiophene				NA			

PAHs - IS/surrogate used for quantitation

	Lab 25	Lab 26	Lab 27	Lab 28	Lab 29	Lab 30
naphthalene	Naphthalene-d8	Naphthalene-d8(Fluorene-d10)	Fluorene-d10(Naphthalene-d8)	naphthalene-d8	d8-Naphthalene	naphthalene-d8
phenyl	Naphthalene-d8	Naphthalene-d8(Fluorene-d10)	Fluorene-d10(Acenaphthene-d10)	ds-1-Methylnaphthalene	naphthalene-d8	naphthalene-d8
acenaphthene	Acenaphthene-d10	Acenaphthene-d10(Fluorene-d10)	Fluorene-d10(Acenaphthene-d10)	d10Acenaphthene	acenaphthene d10	acenaphthene d10
acenaphthylene	Acealaphthene-d10	Acenaphthene-d10(Fluorene-d10)	Fluorene-d10(Acenaphthene-d10)	d8-Acenaphthylene	acenaphthene d10	acenaphthene d10
fluorene	Acenaphthene-d10	Acenaphthene-d10(Fluorene-d10)	Fluorene-d10(Acenaphthene-d10)	d10 Fluorene	acenaphthene d10	acenaphthene d10
phenanthrene	Phenanthrene-d10	Phenanthrene-d10(Fluoranthene-d10)	Pyrene-d10(Phenanthrene-d10)	acenaphthene-d10	d10-Phenanthrene	phenanthrene d10
anthracene	Phenanthrene-d10	Phenanthrene-d10(Fluoranthene-d10)	Pyrene-d10(Phenanthrene-d10)	acenaphthene-d10	d10-Anthracene	phenanthrene d10
fluoranthene	Phenanthrene-d10	Phenanthrene-d10(Fluoranthene-d10)	Pyrene-d10(Phenanthrene-d10)	acenaphthene-d10	d10-Fluoranthene	phenanthrene d10
pyrene	Chrysene-d12	Chrysene-d12(Fluoranthene-d10)	Pyrene-d10(Phenanthrene-d10)	acenaphthene-d10	d10-Pyrene	phenanthrene d10
benz[b]fluorene	Chrysene-d12	Chrysene-d12(Fluoranthene-d10)	Pyrene-d10(Phenanthrene-d10)	acenaphthene-d10	d12-Benz[b]fluoranthene	phenanthrene d10
benz[a]anthracene	Chrysene-d12	Chrysene-d12(Fluoranthene-d12)	Pyrene-d10(Chrysene-d12)	benzo[a]pyrene-d12	d12-Benz[a]anthracene	chrysene d12
chrysene	Chrysene-d12	Chrysene-d12(Fluoranthene-d12)	Pyrene-d10(Chrysene-d12)	benzo[a]pyrene-d12	d12-Chrysene	chrysene d12
trifluorophene			NA		d12-Chrysene	NA
benzo[b]fluoranthene	Chrysene-d12	Perylene-d12(Terphenyl-d14)	Benzol[ab]pyrene-d12(Chrysene-d12)	benzo[a]pyrene-d12	d12-Benzol[b]fluoranthene	perylene d12
benzo[f]fluoranthene		Perylene-d12(Terphenyl-d14)	Benzol[ab]pyrene-d12(Chrysene-d12)		d12-Benzol[f]fluoranthene	perylene d12
benzo[k]fluoranthene	Chrysene-d12	Perylene-d12(Terphenyl-d14)	benzo[a]pyrene-d12		d12-Benzol(k)fluoranthene	perylene d12
benzo[a]fluoranthene		Perylene-d12(Terphenyl-d14)	benzo[a]pyrene-d12(Chrysene-d12)		d12-Benzol(a)pyrene	NA
benzo[e]pyrene	Chrysene-d12	Perylene-d12(Terphenyl-d14)	Benzol[ab]pyrene-d12(Chrysene-d12)		d12-Benzol(e)pyrene	perylene d12
benzo[a]pyrene	Chrysene-d12	Perylene-d12(Terphenyl-d14)	Benzol[ab]pyrene-d12(Chrysene-d12)		d12-Benzol(a)pyrene	perylene d12
perylene	Chrysene-d12	Perylene-d12(Terphenyl-d14)	Benzol[ab]pyrene-d12(Perylene-d12)		d12-perylene	perylene d12
indeno[1,2,3-cd]pyrene	Chrysene-d12	Perylene-d12(Terphenyl-d14)	Benzol[ab]pyrene-d12(Chrysene-d12)	benzo[a]pyrene-d12	d12-henzo(cd)perylene	perylene d12
benzo[ghi]perylene	Chrysene-d12	Perylene-d12(Terphenyl-d14)	Benzol[ab]pyrene-d12(Chrysene-d12)		d12-henzo(ghi)perylene	perylene d12
phenanthro[1,2,3-ij]anthracene	Chrysene-d12	Perylene-d12(Terphenyl-d14)	Benzol[ab]pyrene-d12(Chrysene-d12)	benzo[a]pyrene-d12	d12-henzo(ghj)perylene	perylene d12
cis/trans-decalin		Naphthalene-d8(Fluorene-d10)	Fluorene-d10(Naphthalene-d8)		d8-Naphthalene	naphthalene d8
dibenzofuran	Acenaphthene-d10	Acenaphthene-d10(Fluorene-d10)	Fluorene-d10(Acenaphthene-d10)	d10-Acenaphthene	acenaphthene d10	acenaphthene d10
retene		Chrysene-d12(Fluoranthene-d10)	Pyrene-d10(Phenanthrene-d10)	d10-Pyrene	NA	NA
benzothiophene		Naphthalene-d8(Fluorene-d10)	Fluorene-d10(Acenaphthene-d10)	d8-Naphthalene	phenanthrene d18	phenanthrene d18
dibenzothiophene	Acenaphthene-d10	Phenanthrene-d10(Fluoranthene-d10)	Pyrene-d10(Phenanthrene-d10)	d10-Phenanthrene	acenaphthene d10	acenaphthene d10
naphthobenzothiophene		Chrysene-d12(Terphenyl-d14)	Pyrene-d10(Phenanthrene-d10)			

PAHs - IS/surrogate used for quantitation

	Lab 31	Lab 32	Lab 33	Lab 34
naphthalene		d10-Fluorene	d8-Naphthalene	Acenaphthene-d10
biphenyl		d10-Fluorene	D10 Acenaphthene	Acenaphthene-d10
acenaphthene		d10-Fluorene	D12 Benzo(a)pyrene	Acenaphthene-d10
acenaphthyrene		d10-Fluorene	Acenaphthene-d10	Acenaphthene-d10
fluorene		d10-Fluorene	Acenaphthene-d10	Acenaphthene-d10
phenanthrene		d10-Fluorene	Acenaphthene-d10	Acenaphthene-d10
anthracene		d10-Fluorene	Acenaphthene-d10	Acenaphthene-d10
fluoranthene		d10-Pyrene	Acenaphthene-d10	Acenaphthene-d10
pyrene		d10-Pyrene	Acenaphthene-d10	Acenaphthene-d10
hepta[1]fluorene			Acenaphthene-d10	Acenaphthene-d10
tert[1]anthracene		d12-Benzo(a)pyrene		Chrysene-d12
chrysene		d12-Benzo(a)pyrene		Chrysene-d12
triptycene				Chrysene-d12
hepta[1]fluoranthene				Chrysene-d12
benzo[1]fluoranthene				Chrysene-d12
benzo[4]fluoranthene		d12-Benzo(a)pyrene		Chrysene-d12
benzo[1]fluoranthene				Chrysene-d12
benzo[1]fluorene				Chrysene-d12
benzo[a]pyrene		d12-Benzo(a)pyrene		Chrysene-d12
perylene		d12-Benzo(a)pyrene		Chrysene-d12
indeno[1,2,1-cd]pyrene		d12-Benzo(a)pyrene		Chrysene-d12
benzo[ghi]perylene		d12-Benzo(a)pyrene		Chrysene-d12
dibenz[f,h]anthracene		d12-Benzo(a)pyrene		Chrysene-d12
cis/trans-decalin		d10-Fluorene		Acenaphthene-d10
dihydrofuran		d10-Fluorene		Acenaphthene-d10
retene		d10-Pyrene		Acenaphthene-d10
benzothiophene		d10-Fluorene		Acenaphthene-d10
dibenzothiophene		d10-Fluorene		Acenaphthene-d10
naphthalothiophene		d10-Pyrene		Acenaphthene-d10

Alkylated PAHs - IS/surrogate used for quantitation

	Lab 1	Lab 2	Lab 3	Lab 4	Lab 5	Lab 6
1-methylnaphthalene	naphthalene-d8		Naphthalene-d8	Naphthalene-D8	NPH-d8	
2-methylnaphthalene	naphthalene-d8		Naphthalene-d8	Naphthalene-D8	NPH-d8	
2,6-dimethylnaphthalene	naphthalene-d8		Naphthalene-d8	Naphthalene-D8	ACE-d10	
1,6,7-trimethylnaphthalene	naphthalene-d8		Acenaphthene-d10	Naphthalene-D8	ACE-d10	
1-methylphenanthrene	phenanthrene-d10		Phenanthrene-d10	Phenanthrene-D10	ACE-d10	
C1-decalins	NA		NA	Naphthalene-D8		
C2-decalins	NA		NA	Naphthalene-D8		
C3-decalins	NA		NA	Naphthalene-D8		
C4-decalins	NA		NA	Naphthalene-D8		
C1-naphthalenes	naphthalene-d8		Naphthalene-d8	Naphthalene-D8	ACE-d10	
C2-naphthalenes	naphthalene-d8		Naphthalene-d8	Naphthalene-D8	ACE-d10	
C3-naphthalenes	naphthalene-d8		Naphthalene-d8	Naphthalene-D8	ACE-d10	
C4-naphthalenes	naphthalene-d8		Naphthalene-d8	Naphthalene-D8	ACE-d10	
C1-benzothiophenes	NA		NA	Naphthalene-D8		
C2-benzothiophenes	NA		NA	Naphthalene-D8		
C3-benzothiophenes	NA		NA	Naphthalene-D8		
C4-benzothiophenes	NA		NA	Naphthalene-D8		
C1-fluorenes	phenanthrene-d10		Acenaphthene-d10	Acenaphthene-D10	ACE-d10	
C2-fluorenes	phenanthrene-d10		Acenaphthene-d10	Acenaphthene-D10	ACE-d10	
C3-fluorenes	phenanthrene-d10		Acenaphthene-d10	Acenaphthene-D10	ACE-d10	
C1-phenanthrenes/anthracenes	phenanthrene-d10		Phenanthrene-d10	Phenanthrene-D10	ACE-d10	
C2-phenanthrenes/anthracenes	phenanthrene-d10		Phenanthrene-d10	Phenanthrene-D10	ACE-d10	
C3-phenanthrenes/anthracenes	phenanthrene-d10		Phenanthrene-d10	Phenanthrene-D10	ACE-d10	
C4-phenanthrenes/anthracenes	phenanthrene-d10		Phenanthrene-d10	Phenanthrene-D10	ACE-d10	
C1-dibenzothiophenes	phenanthrene-d10		Acenaphthene-d10	Phenanthrene-D10	ACE-d10	
C2-dibenzothiophenes	phenanthrene-d10		Acenaphthene-d10	Phenanthrene-D10	ACE-d10	
C3-dibenzothiophenes	phenanthrene-d10		Acenaphthene-d10	Phenanthrene-D10	ACE-d10	
C4-dibenzothiophenes	phenanthrene-d10		Acenaphthene-d10	Phenanthrene-D10	ACE-d10	
C1-fluoranthenes/pyrenes	fluoranthene-d10		Chrysene-d12	Phenanthrene-D10	BaP-d12	
C2-fluoranthenes/pyrenes	fluoranthene-d10		Chrysene-d12	Phenanthrene-D10	BaP-d12	
C3-fluoranthenes/pyrenes	fluoranthene-d10		Chrysene-d12	Phenanthrene-D10	BaP-d12	
C4-fluoranthenes/pyrenes	fluoranthene-d10		NA	Phenanthrene-D10	BaP-d12	
C1-naphthobenzo[b]phenes	NA		NA	Chrysene-D12	ACE-d10	
C2-naphthobenzo[b]phenes	NA		NA	Chrysene-D12	ACE-d10	
C3-naphthobenzo[b]phenes	NA		NA	Chrysene-D12	ACE-d10	
C4-naphthobenzo[b]phenes	NA		NA	Chrysene-D12	ACE-d10	
C1-chrysenes	B[a]A-d12		Chrysene-d12	Chrysene-D12	BaP-d12	
C2-chrysenes	B[a]A-d12		Chrysene-d12	Chrysene-D12	BaP-d12	
C3-chrysenes	B[a]A-d12		Chrysene-d12	Chrysene-D12	BaP-d12	
C4-chrysenes	B[a]A-d12		Chrysene-d12	Chrysene-D12	BaP-d12	

Alkylated PAHs - IS/surrogate used for quantitation

	Lab 7	Lab 8	Lab 9	Lab 10	Lab 11	Lab 12
1-methylnaphthalene	d8-Naphthalene	13C Acenaphthylene		Naphthalene-d8	Naphthalene-d8	Fluorene-d10
2-methylnaphthalene	d8-Naphthalene	13C Acenaphthylene		Naphthalene-d8	Naphthalene-d8	Fluorene-d10
2,6-dimethylnaphthalene		13C Acenaphthylene		NA	Naphthalene-d8	Fluorene-d10
1,6,7-trimethylnaphthalene		NA		NA		Fluorene-d10
1-methylphenanthrene		13C Phenanthrene		NA	Acenaphthene-d10	Fluorene-d10
C1-decalins		NA		NA		
C2-decalins		NA		NA		
C3-decalins		NA		NA		
C4-decalins		NA		NA		
C1-naphthalenes		NA		Naphthalene-d8		Fluorene-d10
C2-naphthalenes		NA		Naphthalene-d8		Fluorene-d10
C3-naphthalenes		NA		Naphthalene-d8		Fluorene-d10
C4-naphthalenes		NA		Naphthalene-d8		Fluorene-d10
C1-benzothiophenes		NA		NA		
C2-benzothiophenes		NA		NA		
C3-benzothiophenes		NA		NA		
C4-benzothiophenes		NA		NA		
C1-fluorenes		NA		Acenaphthene-d10		Fluorene-d10
C2-fluorenes		NA		Acenaphthene-d10		Fluorene-d10
C3-fluorenes		NA		Acenaphthene-d10		Fluorene-d10
C1-phenanthrenes/anthracenes		NA		Phanthrene-d10		Fluorene-d10
C2-phenanthrenes/anthracenes		NA		Phanthrene-d10		Fluorene-d10
C3-phenanthrenes/anthracenes		NA		Phanthrene-d10		Fluorene-d10
C4-phenanthrenes/anthracenes		NA		Phanthrene-d10		Fluorene-d10
C1-dibenzothiophenes		NA		Phanthrene-d10		Fluorene-d10
C2-dibenzothiophenes		NA		Phanthrene-d10		Fluorene-d10
C3-dibenzothiophenes		NA		Phanthrene-d10		Fluorene-d10
C4-dibenzothiophenes		NA		Phanthrene-d10		Fluorene-d10
C1-fluoranthenes/pyrenes		NA		Chrysene-12		Benz(a)pyrene-d12
C2-fluoranthenes/pyrenes		NA		Chrysene-12		Benz(a)pyrene-d12
C3-fluoranthenes/pyrenes		NA		Chrysene-12		Benz(a)pyrene-d12
C4-fluoranthenes/pyrenes		NA		NA		
C1-naphthobenzothiophenes		NA		NA		
C2-naphthobenzothiophenes		NA		NA		
C3-naphthobenzothiophenes		NA		NA		
C4-naphthobenzothiophenes		NA		NA		
C1-chrysenes		NA		Chrysene-12		Benz(a)pyrene-d12
C2-chrysenes		NA		Chrysene-12		Benz(a)pyrene-d12
C3-chrysenes		NA		Chrysene-12		Benz(a)pyrene-d12
C4-chrysenes		NA		NA		Benz(a)pyrene-d12

Alkylated PAHs - IS/surrogate used for quantitation

	Lab 13	Lab 14	Lab 15	Lab 16	Lab 17	Lab 18
1-methylnaphthalene	naphthalene-d8	naphthalene-d8	2-fluorobiphenyl	Fluorene-d10 / Naphthalene-d8	Fluorene-d10 / Naphthalene-d8	Naphthalene-d8
2-methylnaphthalene	naphthalene-d8	naphthalene-d8	2-fluorobiphenyl	Fluorene-d10 / Naphthalene-d8	Fluorene-d10 / Naphthalene-d8	Naphthalene-d8
2,6-dimethylnaphthalene	naphthalene-d8	naphthalene-d8	NA	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Acenaphthene-d10	Naphthalene-d8
1,6,7-trimethylnaphthalene	acenaphthylene-d8		NA	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Acenaphthene-d10	Acenaphthene-d10
1-methylphenanthrene	phenanthrene-d10	phenanthrene-d10	NA	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	Phenanthrene-d10
C1-decalins			NA	Fluorene-d10 / Naphthalene-d8	Fluorene-d10 / Naphthalene-d8	
C2-decalins			NA	Fluorene-d10 / Naphthalene-d8	Fluorene-d10 / Naphthalene-d8	
C3-decalins			NA	Fluorene-d10 / Naphthalene-d8	Fluorene-d10 / Naphthalene-d8	
C4-decalins			NA	Fluorene-d10 / Naphthalene-d8	Fluorene-d10 / Naphthalene-d8	
C1-naphthalenes			NA	Fluorene-d10 / Naphthalene-d8	Fluorene-d10 / Naphthalene-d8	
C2-naphthalenes			2-fluorobiphenyl	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Naphthalene-d8	
C3-naphthalenes			2-fluorobiphenyl	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Naphthalene-d8	
C4-naphthalenes			2-fluorobiphenyl	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Naphthalene-d8	
C1-benzothiophenes			NA	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Naphthalene-d8	
C2-benzothiophenes			NA	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Naphthalene-d8	
C3-benzothiophenes			NA	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Naphthalene-d8	
C4-benzothiophenes			NA	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Naphthalene-d8	
C1-fluorenes			2-fluorobiphenyl	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Acenaphthene-d10	
C2-fluorenes			2-fluorobiphenyl	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Phenanthrene-d10	
C3-fluorenes			2-fluorobiphenyl	Fluorene-d10 / Acenaphthene-d10	Fluorene-d10 / Phenanthrene-d10	
C1-phenanthrenes/anthracenes			p-terphenyl-d14	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Phenanthrene-d10	
C2-phenanthrenes/anthracenes			p-terphenyl-d14	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Phenanthrene-d10	
C3-phenanthrenes/anthracenes			p-terphenyl-d14	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Phenanthrene-d10	
C4-phenanthrenes/anthracenes			p-terphenyl-d14	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Phenanthrene-d10	
C1-dibenzothiophenes			p-terphenyl-d14	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Phenanthrene-d10	
C2-dibenzothiophenes			p-terphenyl-d14	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Phenanthrene-d10	
C3-dibenzothiophenes			p-terphenyl-d14	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Phenanthrene-d10	
C4-dibenzothiophenes			p-terphenyl-d14	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Phenanthrene-d10	
C1-fluoranthenes/pyrenes			p-terphenyl-d14	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	
C2-fluoranthenes/pyrenes			p-terphenyl-d14	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	
C3-fluoranthenes/pyrenes			p-terphenyl-d14	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	
C4-fluoranthenes/pyrenes			NA		Pyrene-d10 / Chrysene-d12	
C1-naphthobenzothiophenes			NA	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	
C2-naphthobenzothiophenes			NA	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	
C3-naphthobenzothiophenes			NA	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	
C4-naphthobenzothiophenes			NA	Fluorene-d10 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	
C1-chrysenes			NA	Chrysene-d12 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	
C2-chrysenes			NA	Chrysene-d12 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	
C3-chrysenes			NA	Chrysene-d12 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	
C4-chrysenes			NA	Chrysene-d12 / Phenanthrene-d10	Pyrene-d10 / Chrysene-d12	

Alkylated PAHs - IS/surrogate used for quantitation

	Lab 19	Lab 20	Lab 21	Lab 22	Lab 23	Lab 24a	Lab 24h
1-methylnaphthalene	Naphthalene-d8	Naphthalene-d8	naphthalene-d8	methylnaphthalene d10-2-	d8-Naphthalene	Naphthalene-d8	
2-methylnaphthalene	Naphthalene-d8	Acenaphthene-d10	acenaphthene-d10	methylnaphthalene d12-2-	d8-Naphthalene	Naphthalene-d8	
2,6-dimethylnaphthalene	Naphthalene-d8	Acenaphthene-d10	naphthalene-d8	dimethylnaphthalene d12-2,6-	d10-Acenaphthene	Naphthalene-d8	
1,6,7-trimethylnaphthalene	Phenanthrene-d10	Acenaphthene-d10		NA	d10-Acenaphthene	NA	
1-methylphenanthrene	Phenanthrene-d10	Phenanthrene-d10	phenanthrene-d10	d10-phenanthrene	d10-Phenanthrene	Acenaphthene-d10	
C1-decalins				NA		NA	
C2-decalins				NA		NA	
C3-decalins				NA		NA	
C4-decalins				NA		NA	
C1-naphthalenes	Naphthalene-d8	Naphthalene-d8	naphthalene-d8	d10-2- methylnaphthalene		Naphthalene-d8	
C2-naphthalenes	Naphthalene-d8	Acenaphthene-d10	naphthalene-d8	d12-2,6- dimethylnaphthalene		Naphthalene-d8	
C3-naphthalenes	Naphthalene-d8	Acenaphthene-d10	naphthalene-d8	d12-2,6- dimethylnaphthalene		Acenaphthene-d10	
C4-naphthalenes	Naphthalene-d8	Phenanthrene-d10	naphthalene-d8	d12-2,6- dimethylnaphthalene		Acenaphthene-d10	
C1-benzothiophenes				NA		NA	
C2-benzothiophenes				NA		NA	
C3-benzothiophenes				NA		NA	
C4-benzothiophenes				NA		NA	
C1-fluorenes	Acenaphthalene-d10	Phenanthrene-d10	acenaphthene-d10	d10-phenanthrene		Acenaphthene-d10	
C2-fluorenes	Acenaphthalene-d10	Phenanthrene-d10	acenaphthene-d10	d10-phenanthrene		Acenaphthene-d10	
C3-fluorenes	Acenaphthalene-d10	Phenanthrene-d10	acenaphthene-d10	d10-phenanthrene		Acenaphthene-d10	
C1-phenanthrenes/anthracenes	Phenanthrene-d10	Phenanthrene-d10	phenanthrene-d10	d10-phenanthrene		Acenaphthene-d10	
C2-phenanthrenes/anthracenes	Phenanthrene-d10	Phenanthrene-d10	phenanthrene-d10	d10-fluoranthene		Acenaphthene-d10	
C3-phenanthrenes/anthracenes	Phenanthrene-d10	Chrysene-d12	phenanthrene-d10	d10-fluoranthene		Acenaphthene-d10	
C4-phenanthrenes/anthracenes	Phenanthrene-d10	Chrysene-d12		d10-fluoranthene		Acenaphthene-d10	
C1-dibenzothiophenes	Phenanthrene-d10	Phenanthrene-d10		d10-phenanthrene		NA	
C2-dibenzothiophenes	Phenanthrene-d10	Phenanthrene-d10		d10-phenanthrene		NA	
C3-dibenzothiophenes	Phenanthrene-d10	Phenanthrene-d10		d10-phenanthrene		NA	
C4-dibenzothiophenes	Phenanthrene-d10	Chrysene-d12		d10-phenanthrene		NA	
C1-fluoranthenes/pyrenes	Chrysene-d12	Chrysene-d12	chrysene-d12	d10-fluoranthene		Acenaphthene-d10	
C2-fluoranthenes/pyrenes	Chrysene-d12	Chrysene-d12	chrysene-d12	d10-fluoranthene		Acenaphthene-d10	
C3-fluoranthenes/pyrenes	Chrysene-d12	Chrysene-d12		d10-fluoranthene		Acenaphthene-d10	
C4-fluoranthenes/pyrenes				d10-fluoranthene		Acenaphthene-d10	
C1-naphthobenzothiophenes				NA		NA	
C2-naphthobenzothiophenes				NA		NA	
C3-naphthobenzothiophenes				NA		NA	
C4-naphthobenzothiophenes				NA		NA	
C1-chrysenes	Chrysene-d12	Chrysene-d12		d12-chrysene		NA	
C2-chrysenes	Chrysene-d12	Chrysene-d12		d12-chrysene		NA	
C3-chrysenes	Chrysene-d12	Chrysene-d12		d12-chrysene		NA	
C4-chrysenes	Chrysene-d12	Chrysene-d12		d12-chrysene		NA	

Alkylated PAHs - IS/surrogate used for quantitation

	Lab 25	Lab 26	Lab 27	Lab 28	Lab 29	Lab 30
1-methylnaphthalene	Naphthalene-d8	Naphthalene-d8/Fluorene-d10	Fluorene-d10/Naphthalene-d8	naphthalene-d8	d8-1-Methylnaphthalene	naphthalene d8
2-methylnaphthalene	Naphthalene-d8	Naphthalene-d8/Fluorene-d10	Fluorene-d10/Naphthalene-d8	naphthalene-d8	d8-1-Methylnaphthalene	naphthalene d8
2,6-dimethylnaphthalene	Naphthalene-d8	Naphthalene-d8/Fluorene-d10	Fluorene-d10/Acenaphthene-d10	naphthalene-d8	d8-1-Methylnaphthalene	naphthalene d8
1,6,7-trimethylnaphthalene		Naphthalene-d8/Fluorene-d10	Fluorene-d10/Acenaphthene-d10		d10-Fluorene	NA
1-methylphenanthrene	Phenanthrene-d10	Phenanthrene-d10/Fluoranthene-d10	Pyrene-d10/Phenanthrene-d10	acenaphthene-d10	d8-Dibenzothiophene	NA
C1-decalins		Naphthalene-d8/Fluorene-d10	Fluorene-d10/Naphthalene-d8		d8-1-Methylnaphthalene	NA
C2-decalins		Naphthalene-d8/Fluorene-d10	Fluorene-d10/Naphthalene-d8		d8-1-Methylnaphthalene	NA
C3-decalins		Naphthalene-d8/Fluorene-d10	Fluorene-d10/Naphthalene-d8		d8-1-Methylnaphthalene	NA
C4-decalins		Naphthalene-d8/Fluorene-d10	Fluorene-d10/Naphthalene-d8		d8-1-Methylnaphthalene	NA
C1-naphthalenes	Naphthalene-d8	Naphthalene-d8/Fluorene-d10	Fluorene-d10/Naphthalene-d8	naphthalene-d8	d8-1-Methylnaphthalene	NA
C2-naphthalenes	Naphthalene-d8	Naphthalene-d8/Fluorene-d10	Fluorene-d10/Acenaphthene-d10	naphthalene-d8	d10Acenaphthene	NA
C3-naphthalenes	Naphthalene-d8	Naphthalene-d8/Fluorene-d10	Fluorene-d10/Acenaphthene-d10	acenaphthene-d10	d10Fluorene	NA
C4-naphthalenes	Naphthalene-d8	Naphthalene-d8/Fluorene-d10	Fluorene-d10/Acenaphthene-d10	acenaphthene-d10	d10Fluorene	NA
C1-benzothiophenes		Naphthalene-d8/Fluorene-d10	Fluorene-d10/Acenaphthene-d10		d8-1-Methylnaphthalene	NA
C2-benzothiophenes		Naphthalene-d8/Fluorene-d10	Fluorene-d10/Acenaphthene-d10		d8-1-Methylnaphthalene	NA
C3-benzothiophenes		Naphthalene-d8/Fluorene-d10	Fluorene-d10/Acenaphthene-d10		d8-1-Methylnaphthalene	NA
C4-benzothiophenes		Naphthalene-d8/Fluorene-d10	Fluorene-d10/Acenaphthene-d10		d8-1-Methylnaphthalene	NA
C1-fluorenes	Acenaphthene-d10	Acenaphthene-d10/Fluorene-d10	Fluorene-d10/Acenaphthene-d10	acenaphthene-d10	d10-Fluorene	NA
C2-fluorenes	Acenaphthene-d10	Acenaphthene-d10/Fluorene-d10	Fluorene-d10/Acenaphthene-d10	acenaphthene-d10	d10-Fluorene	NA
C3-fluorenes	Acenaphthene-d10	Acenaphthene-d10/Fluorene-d10	Fluorene-d10/Acenaphthene-d10	acenaphthene-d10	d10-Fluorene	NA
C1-phenanthrenes/anthracenes	Phenanthrene-d10	Phenanthrene-d10/Fluoranthene-d10	Pyrene-d10/Phenanthrene-d10	acenaphthene-d10	d10-Anthracene	NA
C2-phenanthrenes/anthracenes	Phenanthrene-d10	Phenanthrene-d10/Fluoranthene-d10	Pyrene-d10/Phenanthrene-d10	acenaphthene-d10	d10-Anthracene	NA
C3-phenanthrenes/anthracenes	Phenanthrene-d10	Phenanthrene-d10/Fluoranthene-d10	Pyrene-d10/Phenanthrene-d10	acenaphthene-d10	d10-Anthracene	NA
C4-phenanthrenes/anthracenes	Phenanthrene-d10	Phenanthrene-d10/Fluoranthene-d10	Pyrene-d10/Phenanthrene-d10	acenaphthene-d10	d10-Anthracene	NA
C1-dibenzothiophenes	Phenanthrene-d10	Phenanthrene-d10/Fluoranthene-d10	Pyrene-d10/Phenanthrene-d10		d8-Dibenzothiophene	NA
C2-dibenzothiophenes	Phenanthrene-d10	Phenanthrene-d10/Fluoranthene-d10	Pyrene-d10/Phenanthrene-d10		d8-Dibenzothiophene	NA
C3-dibenzothiophenes	Phenanthrene-d10	Phenanthrene-d10/Fluoranthene-d10	Pyrene-d10/Phenanthrene-d10		d8-Dibenzothiophene	NA
C4-dibenzothiophenes	Phenanthrene-d10	Phenanthrene-d10/Fluoranthene-d10	Pyrene-d10/Phenanthrene-d10		d8-Dibenzothiophene	NA
C1-fluoranthenes/pyrenes	Chrysene-d12	Phenanthrene-d10/Fluoranthene-d10	Pyrene-d10/Phenanthrene-d10		d10-Fluoranthene	NA
C2-fluoranthenes/pyrenes	Chrysene-d12	Phenanthrene-d10/Fluoranthene-d10	Pyrene-d10/Phenanthrene-d10		d10-Fluoranthene	NA
C3-fluoranthenes/pyrenes	Chrysene-d12	Phenanthrene-d10/Fluoranthene-d10	Pyrene-d10/Phenanthrene-d10		d10-Fluoranthene	NA
C4-fluoranthenes/pyrenes		Phenanthrene-d10/Fluoranthene-d10	Pyrene-d10/Phenanthrene-d10		d10-Fluoranthene	NA
C1-naphthobenzothiophenes		Chrysene-d12/Terphenyl-d14	Pyrene-d10/Phenanthrene-d10			NA
C2-naphthobenzothiophenes		Chrysene-d12/Terphenyl-d14	Pyrene-d10/Phenanthrene-d10			NA
C3-naphthobenzothiophenes		Chrysene-d12/Terphenyl-d14	Pyrene-d10/Phenanthrene-d10			NA
C4-naphthobenzothiophenes		Chrysene-d12/Terphenyl-d14	Pyrene-d10/Phenanthrene-d10			NA
C1-chrysenes	Chrysene-d12	Chrysene-d12/Terphenyl-d14	Pyrene-d10/Chrysene-d12		d12-Chrysene	NA
C2-chrysenes	Chrysene-d12	Chrysene-d12/Terphenyl-d14	Pyrene-d10/Chrysene-d12		d12-Chrysene	NA
C3-chrysenes	Chrysene-d12	Chrysene-d12/Terphenyl-d14	Pyrene-d10/Chrysene-d12		d12-Chrysene	NA
C4-chrysenes	Chrysene-d12	Chrysene-d12/Terphenyl-d14	Pyrene-d10/Chrysene-d12		d12-Chrysene	NA

Alkylated PAHs - IS/surrogate used for quantitation

	Lab 31	Lab 32	Lab 33	Lab 34
1-methylnaphthalene		d10-Fluorene		Acenaphthene-d10
2-methylnaphthalene		d10-Fluorene		Acenaphthene-d10
2,6-dimethylnaphthalene		d10-Fluorene		Acenaphthene-d10
1,6,7-trimethylnaphthalene		d10-Fluorene		Acenaphthene-d10
1-methylphenanthrene		d10-Pyrene		Acenaphthene-d10
C1-decalins		d10-Fluorene		Acenaphthene-d10
C2-decalins		d10-Fluorene		Acenaphthene-d10
C3-decalins		d10-Fluorene		Acenaphthene-d10
C4-decalins		d10-Fluorene		Acenaphthene-d10
C1-naphthalenes		d10-Fluorene		Acenaphthene-d10
C2-naphthalenes		d10-Fluorene		Acenaphthene-d10
C3-naphthalenes		d10-Fluorene		Acenaphthene-d10
C4-naphthalenes		d10-Fluorene		Acenaphthene-d10
C1-benzothiophenes		d10-Fluorene		Acenaphthene-d10
C2-benzothiophenes		d10-Fluorene		Acenaphthene-d10
C3-benzothiophenes		d10-Fluorene		Acenaphthene-d10
C4-benzothiophenes				Acenaphthene-d10
C1-fluorenes		d10-Fluorene		Acenaphthene-d10
C2-fluorenes		d10-Fluorene		Acenaphthene-d10
C3-fluorenes		d10-Fluorene		Acenaphthene-d10
C1-phenanthrenes/anthracenes		d10-Pyrene		Acenaphthene-d10
C2-phenanthrenes/anthracenes		d10-Pyrene		Acenaphthene-d10
C3-phenanthrenes/anthracenes		d10-Pyrene		Acenaphthene-d10
C4-phenanthrenes/anthracenes		d10-Pyrene		Acenaphthene-d10
C1-dibenzothiophenes		d10-Fluorene		Acenaphthene-d10
C2-dibenzothiophenes		d10-Fluorene		Acenaphthene-d10
C3-dibenzothiophenes		d10-Fluorene		Acenaphthene-d10
C4-dibenzothiophenes				Acenaphthene-d10
C1-fluoranthenes/pyrenes		d10-Pyrene		Acenaphthene-d10
C2-fluoranthenes/pyrenes		d10-Pyrene		Acenaphthene-d10
C3-fluoranthenes/pyrenes		d10-Pyrene		Acenaphthene-d10
C4-fluoranthenes/pyrenes				Acenaphthene-d10
C1-naphthobenzenothiophenes		d10-Pyrene		Acenaphthene-d10
C2-naphthobenzenothiophenes		d10-Pyrene		Acenaphthene-d10
C3-naphthobenzenothiophenes		d10-Pyrene		Acenaphthene-d10
C4-naphthobenzenothiophenes				Acenaphthene-d10
C1-chrysenes				Chrysene-d12
C2-chrysenes		d12-Benzo(a)pyrene		Chrysene-d12
C3-chrysenes		d12-Benzo(a)pyrene		Chrysene-d12
C4-chrysenes		d12-Benzo(a)pyrene		Chrysene-d12

Biomarkers - IS/surrogate used for quantitation

	Lah 1	Lah 3	Lah 10	Lah 12	Lah 14	Lab 16	Pyrene-d10 / Phanthrene-d10							
Carbazole	NA	NA	NA					d10-phenanthrene	Chrysene-d12			Phenanthrene-d10 / fluoranthene-d10		Chrysene-d12
18a(1b),22,29,30-Fistuorophane	NA	Chrysene-d12	Chrysene-12			Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	NA					NA
17a(1b),22,29,30-Fistuorophane	Bi[al]A-d12	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	Chrysene-d12					NA
17a(1b),2,[4(1b)-30]-Norbornane	Bi[al]A-d12	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	Chrysene-d12					Chrysene-d12
18a(1b),30-Normethophane	NA	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	Chrysene-d12					NA
17a(1b)-Dihopane	NA	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	Chrysene-d12					Chrysene-d12
17a(1b),2,[6(1b)-1]-lopane	Bi[al]A-d12	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Ben[alpha]pyrene-androstane	Chrysene-d12					Chrysene-d12
17a(1b),2,[4(1b)-22S]-Homohopane	Bi[al]A-d12	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	Chrysene-d12					Chrysene-d12
17a(1b),2,[6(1b)-22S]-Homohopane	Bi[al]A-d12	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	Chrysene-d12					NA
13b(1b)7a(1b)-Diacholestane 20S	NA	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	Chrysene-d12					Chrysene-d12
5a(1b)14(1b)7a(1b)-Cholestane 20S	NA	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	Chrysene-d12					NA
5a(1b)14(1b)7a(1b)-Diacholestane 20R	NA	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	Chrysene-d12					Chrysene-d12
5a(1b)14(1b)7a(1b)-24-Ethylcholestane 20S	NA	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	Chrysene-d12					NA
5a(1b)14(1b)7a(1b)-24-Ethylcholestane 20R	NA	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	Chrysene-d12					Chrysene-d12
5a(1b)14(1b)17a(1b)-24-Ethylcholestane 20R	NA	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	Chrysene-d12					NA
5a(1b)14(1b)17b(1b)-Cholestane 20S	NA	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	Chrysene-d12					Chrysene-d12
5a(1b)14(1b)17b(1b)-24-Ethylcholestane 20R	NA	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	Chrysene-d12					NA
5a(1b)14b(1b)17b(1b)-24-Ethylcholestane 20S	NA	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	Chrysene-d12					Chrysene-d12
5a(1b)14b(1b)17b(1b)-24-Ethylcholestane 20R	NA	Chrysene-d12	Chrysene-12		Benz[a]anthracene-d12	Chrysene-d12 / 5b(1b)-Cholane		Chrysene-d12	Chrysene-d12					NA

PAHs - Associated % recovery / acceptability ranges

		Lab 10	Lab 16	Lab 17	Lab 19	Lab 20	Lab 21	Lab 23	Lab 25	Lab 26	Lab 27	Lab 28	Lab 30	Lab 31	Lab 33	Lab 34
naphthalene	minus 50 to plus 100	50 to 120	1% to 33%	60 - 140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA	60-130%	40-120		
phenyl	60-140	minus 50 to plus 100	40 to 120	1% to 42%	60-140			50-120	50-200% of IS value in CCV	40-120		NA				40-120
acephenethiene	60-140	minus 50 to plus 100	40 to 120	1% to 12%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120		NA			40-120
acetylenylene	60-140	minus 50 to plus 100	40 to 120	1% to 42%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120		NA			40-120
fluorene	60-140	minus 50 to plus 100	40 to 120	1% to 42%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA			40-120
phenanthrene	60-140	minus 50 to plus 100	40 to 120	4% to 54%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA			40-120
anthracene	60-140	minus 50 to plus 100	40 to 120	4% to 54%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA			40-120
fluoranthene	60-140	minus 50 to plus 100	40 to 120	24% to 68%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA			40-120
pyrene	60-140	minus 50 to plus 100	40 to 120	24% to 68%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA			40-120
benz[b]fluorene	NA	minus 50 to plus 100		24% to 68%									NA			40-120
benz[a]anthracene	60-140	minus 50 to plus 100	40 to 120	24% to 68%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA			40-120
chrysene	60-140	minus 50 to plus 100	40 to 120	24% to 68%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA			40-120
uriphenslene	NA	minus 50 to plus 100		24% to 68%									NA			40-120
benz[b]fluoranthene	60-140	minus 50 to plus 100	40 to 120	1% to 73%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA			40-120
benzofluranthenene	NA	minus 50 to plus 100		1% to 73%									NA			40-120
benzofluranthenene	60-140	minus 50 to plus 100	40 to 120	1% to 73%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA			40-120
benzofluranthenene	NA	minus 50 to plus 100		1% to 73%									NA			40-120
benzofluranthenene	60-140	minus 50 to plus 100	40 to 120	1% to 73%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA			40-120
benzofluranthenene	60-140	minus 50 to plus 100	40 to 120	1% to 73%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA			40-120
benzofluranthenene	60-140	minus 50 to plus 100	40 to 120	1% to 73%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA			40-120
perylene	60-140	minus 50 to plus 100	40 to 120	1% to 73%	60-140			50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA			40-120
indeno[1,2,3-c,d]phenanthrene	60-140	minus 50 to plus 100	40 to 120	1% to 73%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA			40-120
benzofluranthenene	60-140	minus 50 to plus 100	40 to 120	1% to 73%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA			40-120
dibenz[a,h]anthracene	60-140	minus 50 to plus 100	40 to 120	1% to 73%	60-140	60 - 140	50-200	50-120	50-200% of IS value in CCV	40-120	60-120	+/-30%	NA			40-120
cis/trans-decalin	NA	minus 50 to plus 100	40 to 120	1% to 33%									NA			40-120
dibenzofuran	60-140	minus 50 to plus 100	40 to 120	1% to 42%	60-140								NA			40-120
retene	NA	minus 50 to plus 100	40 to 120	24% to 68%									NA			40-120
benzophenone	NA	minus 50 to plus 100	40 to 120	1% to 33%									NA			40-120
dibenzophenone	60-140	minus 50 to plus 100	40 to 120	4% to 54%	60-140								NA			40-120
naphthalene-ethophenone	NA	minus 50 to plus 100	40 to 120	24% to 68%									NA			40-120

Biomarkers - Associated % recovery / acceptance ranges

	Lab 3	Lab 10	Lab 16	Lab 17	Lab 19	Lab 25	Lab 26	Lab 27	Lab 30	Lab 34
Carbazole		minus 50 to plus 100		4% to 54%		50-200% of IS value in CCV	40-120	NA	NA	50-130
18a(1)-22,29,30-Trisnorhopane	NA	minus 50 to plus 100	40 to 120		60-140	50-200% of IS value in CCV		NA	NA	50-130
17a(1)-22,29,30-Trisnorhopane	NA	minus 50 to plus 100	40 to 120		60-140	50-200% of IS value in CCV		NA	NA	50-130
17a(1),2,3[β]H-30-Norhopane	NA	minus 50 to plus 100	40 to 120		60-140	50-200% of IS value in CCV		NA	NA	50-130
18a(1)-30-Norhopane	NA	minus 50 to plus 100	40 to 120		60-140	50-200% of IS value in CCV		NA	NA	50-130
17a(1)-Dahopane	NA	minus 50 to plus 100	40 to 120		60-140	50-200% of IS value in CCV		NA	NA	50-130
17a(1),2,[β]H-1-topane	60-140	minus 50 to plus 100	40 to 120	17% to 60%	60-140	50-200% of IS value in CCV	40-120	NA	NA	50-130
17a(1),2,[β]H-22R-Homohopane	NA	minus 50 to plus 100	40 to 120		60-140	50-200% of IS value in CCV		NA	NA	50-130
17a(1),2,[β]H-22S-Homohopane	NA	minus 50 to plus 100	40 to 120		60-140	50-200% of IS value in CCV		NA	NA	50-130
13b(1),17a(1)-Diacholestane 20S	NA	minus 50 to plus 100	40 to 120		60-140	50-200% of IS value in CCV		NA	NA	50-130
5a(1),14a(1),17a(1)-Cholestane 20S	NA	minus 50 to plus 100	40 to 120		60-140	50-200% of IS value in CCV		NA	NA	50-130
5a(1),14a(1),17a(1)-Cholestane 20R	NA	minus 50 to plus 100	40 to 120		60-140	50-200% of IS value in CCV		NA	NA	50-130
5a(1),14a(1),17a(1)-24-Ethylcholestane 20S	NA	minus 50 to plus 100	40 to 120		60-140	50-200% of IS value in CCV		NA	NA	50-130
5a(1),14a(1),17a(1)-24-Ethylcholestane 20R	NA	minus 50 to plus 100	40 to 120		60-140	50-200% of IS value in CCV		NA	NA	50-130
5a(1),14b(1),17b(1)-Cholestane 20R	NA	minus 50 to plus 100	40 to 120		60-140	50-200% of IS value in CCV		NA	NA	50-130
5a(1),14b(1),17b(1)-Cholestane 20S	NA	minus 50 to plus 100	40 to 120		60-140	50-200% of IS value in CCV		NA	NA	50-130
5a(1),14b(1),17b(1),17b(1)-Cholestane 20R	NA	minus 50 to plus 100	40 to 120		60-140	50-200% of IS value in CCV		NA	NA	50-130
5a(1),14b(1),17b(1),17b(1)-Cholestane 20S	NA	minus 50 to plus 100	40 to 120		60-140	50-200% of IS value in CCV		NA	NA	50-130

PAHs -If "representative compound" used for quantitation, list the compound

	Lab 5	Lab 8	Lab 18	Lab 19	Lab 27
naphthalene				naphthalene	
biphenyl				biphenyl	NA
acenaphthene				acenaphthene	NA
acenaphthylene				acenaphthylene	NA
fluorene				fluorene	NA
phenanthrene				phenanthrene	NA
anthracene				anthracene	NA
fluoranthene				fluoranthene	NA
pyrene				pyrene	NA
benzol[b]fluorene					NA
benz[a]anthracene				benz[a]anthracene	NA
chrysene				chrysene	NA
triphenylene					chrysene
benzo[<i>b</i>]fluoranthene				benzo[<i>b</i>]fluoranthene	NA
benzo[<i>f</i>]fluoranthene					benzo[<i>k</i>]fluoranthene
benzo[<i>k</i>]fluoranthene				benzo[<i>k</i>]fluoranthene	NA
benzo[<i>a</i>]fluoranthene					benzo[<i>k</i>]fluoranthene
benzo[<i>e</i>]pyrene				benzo[<i>e</i>]pyrene	NA
benzo[<i>a</i>]pyrene				benzo[<i>a</i>]pyrene	NA
perylene				perylene	NA
indenol[1,2,3- <i>cd</i>]pyrene				indenol[1,2,3- <i>cd</i>]pyrene	NA
benzo[<i>ghi</i>]perylene				benzo[<i>ghi</i>]perylene	NA
dibenzo[<i>a,h</i>]anthracene				dibenzo[<i>a,h</i>]anthracene	NA
cis/trans-decalin					NA
dibenzofuran				dibenzofuran	NA
retene					NA
benzothiophene					NA
dibenzothiophene				dibenzothiophene	NA
naphthobenzothiophene					NA

Alkylated PAHs - If "representative compound" used for quantitation, list the compound

	Lab 1	Lab 3	Lab 4	Lab 5	Lab 8	Lab 10	Lab 12	Lab 15
1-methylanthracene	NA	NA	NA	NA	13C Acenaphthylene	NA	NA	NA
2-methylanthracene	NA	NA	NA	NA	13C Acenaphthylene	NA	NA	NA
2,6-dimethylanthracene	NA	NA	NA	NA	13C Acenaphthylene	NA	NA	NA
1,6,7-trimethylanthracene	NA	NA	NA	NA	13C Phenanthrene	NA	NA	NA
1-methylphenanthrene	NA	NA	NA	NA	13C Phenanthrene	NA	NA	NA
C1-decalins	NA	NA	cis/trans-decalin	NA	NA	NA	NA	NA
C2-decalins	NA	NA	cis/trans-decalin	NA	NA	NA	NA	NA
C3-decalins	NA	NA	cis/trans-decalin	NA	NA	NA	NA	NA
C4-decalins	NA	NA	Naphthalene	Naphthalene	Naphthalene	Naphthalene	Naphthalene	Naphthalene
C1-naphthalenes	1-ene and 2-methylnaphthalene	Naphthalene	Naphthalene	2,6-dimethylnaphthalene	2,6-dimethylnaphthalene	Naphthalene	Naphthalene	Naphthalene
C2-naphthalenes	2,6-dimethylnaphthalene	Naphthalene	Naphthalene	1,6,7-trimethylnaphthalene	1,6,7-trimethylnaphthalene	Naphthalene	Naphthalene	Naphthalene
C3-naphthalenes	2,3,5-trimethylnaphthalene	Naphthalene	Naphthalene	1,6,7,trimethylnaphthalene	1,6,7,trimethylnaphthalene	Naphthalene	Naphthalene	C3-naphthalenes
C4-naphthalenes	1-benzothiophene	Naphthalene	Naphthalene	benzothiophene	benzothiophene	NA	NA	NA
C2-benzothiophenes	benzothiophene	NA	NA	benzothiophene	benzothiophene	NA	NA	NA
C3-benzothiophenes	benzothiophene	NA	NA	benzothiophene	benzothiophene	NA	NA	NA
C4-benzothiophenes	benzothiophene	NA	NA	benzothiophene	benzothiophene	NA	NA	NA
C1-fluorenes	Fluorene	Fluorene	Fluorene	Fluorene	Fluorene	Fluorene	Fluorene	Fluorene
C2-fluorenes	Fluorene	Fluorene	Fluorene	Fluorene	Fluorene	Fluorene	Fluorene	Fluorene
C3-fluorenes	1-ene, 2-ene, 3-ene, and 9-phenanthrene plus 2-methyrene	Phenanthrene	Phenanthrene	1-methylphenanthrene	1-methylphenanthrene	Phenanthrene	Phenanthrene	Phenanthrene
C1-phenanthrenes/anthracenes	1,7-dimethyphenanthrene	Phenanthrene	Phenanthrene	1-methylphenanthrene	1-methylphenanthrene	Phenanthrene	Phenanthrene	Phenanthrene
C2-phenanthrenes/anthracenes	1,7-dimethyphenanthrene	Phenanthrene	Phenanthrene	1-methylphenanthrene	1-methylphenanthrene	Phenanthrene	Phenanthrene	Phenanthrene
C3-phenanthrenes/anthracenes	1,7-dimethyphenanthrene	Phenanthrene	Phenanthrene	1-methylphenanthrene	1-methylphenanthrene	Dibenzothiophene	Dibenzothiophene	C1-phenanthrenes/anthracenes
C1-dihenzoethiophenes	dihenzoethiophene	Dibenzothiophene	Dibenzothiophene	dibenzoethiophene	dibenzoethiophene	Dibenzothiophene	Dibenzothiophene	C1-dihenzoethiophenes
C2-dihenzoethiophenes	dihenzoethiophene	Dibenzothiophene	Dibenzothiophene	dibenzoethiophene	dibenzoethiophene	Dibenzothiophene	Dibenzothiophene	dibenzoethiophene
C3-dihenzoethiophenes	dihenzoethiophene	Dibenzothiophene	Dibenzothiophene	dibenzoethiophene	dibenzoethiophene	Dibenzothiophene	Dibenzothiophene	dibenzoethiophene
C4-dihenzoethiophenes	dihenzoethiophene	Dibenzothiophene	Dibenzothiophene	dibenzoethiophene	dibenzoethiophene	Dibenzothiophene	Dibenzothiophene	dibenzoethiophene
C1-fluoranthenes/pyrenes	1-ene and 2-methyfluoranthene plus 1-pyrene and 1-arycene	Fluoranthene	Fluoranthene	fluoranthene	fluoranthene	Pyrene	Pyrene	Pyrene
C2-fluoranthenes/pyrenes	1-ene and 2-arycene	Pyrene	Fluoranthene	fluoranthene	fluoranthene	Pyrene	Pyrene	Pyrene
C3-fluoranthenes/pyrenes	1-ene and 2-methyfluoranthene plus 1-pyrene and 1-arycene	Pyrene	Fluoranthene	fluoranthene	fluoranthene	Pyrene	Pyrene	Pyrene
C4-fluoranthenes/pyrenes	1-ene and 2-methyfluoranthene plus 1-pyrene and 1-arycene	NA	Fluoranthene	fluoranthene	fluoranthene	NA	NA	NA
C1-naphthalene/anthracenes	NA	NA	naphthalene	naphthalene	naphthalene	NA	NA	NA
C1-naphthalene/anthracenes	NA	NA	naphthalene	naphthalene	naphthalene	NA	NA	NA
C2-naphthalene/anthracenes	NA	NA	naphthalene	naphthalene	naphthalene	NA	NA	NA
C3-naphthalene/anthracenes	NA	NA	naphthalene	naphthalene	naphthalene	NA	NA	NA
C4-naphthalene/anthracenes	NA	NA	naphthalene	naphthalene	naphthalene	NA	NA	NA
C1-chrysenes	3,4 and 4,5-chrysenes	Chrysene	Chrysene	chrysene	chrysene	Chrysene	Chrysene	Chrysene
C2-chrysenes	3,4 and 6,7-chrysenes	Chrysene	Chrysene	chrysene	chrysene	Chrysene	Chrysene	Chrysene
C3-chrysenes	3,4 and 6,7,8-chrysenes	Chrysene	Chrysene	chrysene	chrysene	Chrysene	Chrysene	Chrysene
C4-chrysenes	3,4 and 6,7,8,9-chrysenes	Chrysene	Chrysene	chrysene	chrysene	Chrysene	Chrysene	Chrysene

Alkylated PAHs - If "representative compound" used for quantitation, list the compound

	Lab 16	Lab 17	Lab 19	Lab 20	Lab 21	Lab 22	Lab 26	Lab 27
1-methylnaphthalene			1-methylnaphthalene					NA
2-methylnaphthalene			2-methylnaphthalene					NA
2,6-dimethylnaphthalene			2,6-dimethylnaphthalene					NA
1,6,7-trimethylnaphthalene			1,6,7-trimethylnaphthalene					NA
1-methylphenanthrene		phenanthrene	1-methylphenanthrene					phenanthrene
C1-decalins	trans-decalin	cis/trans-decalin						cis/trans-decalin
C2-decalins	trans-decalin	trans-decalin						cis/trans-decalin
C3-decalins	trans-decalin	trans-decalin						cis/trans-decalin
C4-decalins								
C1-naphthalenes	naphthalene	naphthalene	Naphthalene	1-methylnaphthalene				naphthalene
C2-naphthalenes	naphthalene	naphthalene	Naphthalene	Naphthalene	2,6-dimethylnaphthalene	2,6- & 1,2-dimethylnaphthalene	Naphthalene	naphthalene
C3-naphthalenes	naphthalene	naphthalene	Naphthalene	1,3,5-trimethylnaphthalene	1,3,5-trimethylnaphthalene	1,3,5-trimethylnaphthalene	Naphthalene	naphthalene
C4-naphthalenes	naphthalene	naphthalene	Naphthalene	1,3,5-trimethylnaphthalene	1,3,5-trimethylnaphthalene	1,3,5-trimethylnaphthalene	Naphthalene	naphthalene
C1-phenanthrenes	benzo[b]phenanthrene	benzo[b]phenanthrene					Benzophenone	benzo[b]phenanthrene
C2-phenanthrenes	benzo[b]phenanthrene	benzo[b]phenanthrene					Benzophenone	benzo[b]phenanthrene
C3-phenanthrenes	benzo[b]phenanthrene	benzo[b]phenanthrene					Benzophenone	benzo[b]phenanthrene
C4-phenanthrenes	benzo[b]phenanthrene	benzo[b]phenanthrene					Benzophenone	benzo[b]phenanthrene
C1-fluorenes	Fluorene	fluorene	fluorene	fluorene	fluorene	2-methylfluorene	Fluorene	fluorene
C2-fluorenes	Fluorene	fluorene	fluorene	fluorene	fluorene	1,7-dimethylfluorene	Fluorene	fluorene
C3-fluorenes	Fluorene	fluorene	fluorene	fluorene	fluorene	1,7-dimethylfluorene	Fluorene	fluorene
C1-phenanthrenes/anthracenes	phenanthrene	phenanthrene	phenanthrene	phenanthrene	1-methylphenanthrene	1-methylphenanthrene	phenanthrene	phenanthrene
C2-phenanthrenes/anthracenes	phenanthrene	phenanthrene	phenanthrene	phenanthrene	1-methylphenanthrene	1,2,6-trimethylphenanthrene	phenanthrene	phenanthrene
C3-phenanthrenes/anthracenes	phenanthrene	phenanthrene	phenanthrene	phenanthrene	1-methylphenanthrene	retene	phenanthrene	phenanthrene
C4-phenanthrenes/anthracenes	phenanthrene	phenanthrene	phenanthrene	phenanthrene	1-methylphenanthrene	2,3-methyldibenzo[othiophenes]	dibenzothiophene	dibenzothiophene
C1-dibenzothiophenes	dibenzothiophene	dibenzothiophene	dibenzothiophene	dibenzothiophene	Dibenzothiophene	2,4-dimethyldibenzothiophene	Dibenzothiophene	dibenzothiophene
C2-dibenzothiophenes	dibenzothiophene	dibenzothiophene	dibenzothiophene	dibenzothiophene	Dibenzothiophene	2,4-dimethyldibenzothiophene	Dibenzothiophene	dibenzothiophene
C3-dibenzothiophenes	dibenzothiophene	dibenzothiophene	dibenzothiophene	dibenzothiophene	Dibenzothiophene	2,4-dimethyldibenzothiophene	Dibenzothiophene	dibenzothiophene
C4-dibenzothiophenes	dibenzothiophene	dibenzothiophene	dibenzothiophene	dibenzothiophene	Dibenzothiophene	2,4-dimethyldibenzothiophene	Dibenzothiophene	dibenzothiophene
C1-fluoranthenes/pyrenes	pyrene	pyrene	pyrene	pyrene	Pyrene	3-methylfluoranthene	Fluoranthene	fluoranthene
C2-fluoranthenes/pyrenes	pyrene	pyrene	pyrene	pyrene	Pyrene	3-methylfluoranthene	Fluoranthene	fluoranthene
C3-fluoranthenes/pyrenes	pyrene	pyrene	pyrene	pyrene	Pyrene	3-methylfluoranthene	Fluoranthene	fluoranthene
C4-fluoranthenes/pyrenes			pyrene	pyrene	Pyrene	3-methylfluoranthene	Fluoranthene	fluoranthene
C1-naphthalene/anthropheines	benzo[b]naphthalene/2,1-dihydrophenanthrene						Naphthalenobenzophenone	naphthalenobenzophenone
C2-naphthalene/anthropheines	benzo[b]naphthalene/2,1-dihydrophenanthrene						Naphthalenobenzophenone	naphthalenobenzophenone
C3-naphthalene/anthropheines	benzo[b]naphthalene/2,1-dihydrophenanthrene						Naphthalenobenzophenone	naphthalenobenzophenone
C4-naphthalene/anthropheines	benzo[b]naphthalene/2,1-dihydrophenanthrene						Naphthalenobenzophenone	naphthalenobenzophenone
C1-1-phenylbenzothiophenes	Chrysene	chrysene	chrysene	chrysene	Chrysene	1- & 6-dimethylchrysene	Chrysene	chrysene
C2-1-phenylbenzothiophenes	Chrysene	chrysene	chrysene	chrysene	Chrysene	5,9-dimethylchrysene	Chrysene	chrysene
C3-1-phenylbenzothiophenes	Chrysene	chrysene	chrysene	chrysene	Chrysene	5,9-dimethylchrysene	Chrysene	chrysene
C4-1-phenylbenzothiophenes	Chrysene	chrysene	chrysene	chrysene	Chrysene	5,9-dimethylchrysene	Chrysene	chrysene

Alkylated PAHs - If "representative compound" used for quantitation, list the compound

	Lab 28	Lab 29	Lab 32	Lab 33	Lab 34
1-methylanthracene					
2-methylanthracene					
2,6-dimethylanthracene					
1,6,7,8-tetramethylanthracene					
1-methylphenanthrene					
C1-decalins					
C2-decalins					
C3-decalins					
C4-decalins					
C1-naphthalenes	1-methylnaphthalene/2-methylnaphthalene	1,2-methylenaphthalene	Naphthalene	1-methyl naphthalene	naphthalene
C2-naphthalenes	2,6-dimethylnaphthalene	Sum of five DMNs	Naphthalene	2,6 dimethyl naphthalene	naphthalene
C3-naphthalenes	2,3,5-trimethylnaphthalene	1,6,7-TMN	Naphthalene	2,3,5 trimethyl naphthalene	naphthalene
C4-naphthalenes	1,2,5,6-tetramethylnaphthalene	1,6,7-TMN	Benzothiophene	1,2,5,6 tetramethyl naphthalene	naphthalene
C1-benzo[b]phenes	benzo[b]phenes	benzo[b]phenes	Benzothiophene	benzo[b]phenes	benzo[b]phenes
C2-benzo[b]phenes	benzo[b]phenes	benzo[b]phenes	Benzothiophene	benzo[b]phenes	benzo[b]phenes
C3-benzo[b]phenes	benzo[b]phenes	benzo[b]phenes	Benzothiophene	benzo[b]phenes	benzo[b]phenes
C1-fluorenes	1-methylfluorene	1-methylfluorene	Fluorene	1 methyl fluorene	fluorene
C2-fluorenes	1,7-dimethylfluorene	1,8-dimethylfluorene	Fluorene	1,7 dimethyl fluorene	fluorene
C3-fluorenes	9,9-erythrofluorene	1-methylfluorene	Fluorene	9-n-propyl fluorene	fluorene
C1-phenaanthrenes/anthocenes	1-methylphenanthrene	Sum of five methylphenanthrenes and anthracenes	Anthracene	1-methylphenanthrene	phenanthrene
C2-phenaanthrenes/anthocenes	1,3-dimethylphenanthrene	9,10-dimethylphenanthrene	Anthracene	1,3 dimethylphenanthrene	phenanthrene
C3-phenaanthrenes/anthocenes	1,2,6,8-tetramethylphenanthrene	9,10-dimethylanthracene	Anthracene	1,2,6 trimethylphenanthrene	phenanthrene
C4-phenaanthrenes/anthocenes	1,2,6,8-tetramethylphenanthrene	9,10-dimethylanthracene	Anthracene	1,2,6,9 tetramethylphenanthrene	phenanthrene
C1-dibenz[cd]ethophenes		2,1,4-methyl(dibenzo)ethophene	Dibenzothiophene		dibenzothiophene
C2-dibenz[cd]ethophenes		2,1,4-methyl(dibenzo)ethophene	Dibenzothiophene		dibenzothiophene
C3-dibenz[cd]ethophenes		2,1,4-methyl(dibenzo)ethophene	Dibenzothiophene		dibenzothiophene
C4-dibenz[cd]ethophenes		2,1,4-methyl(dibenzo)ethophene	Dibenzothiophene		dibenzothiophene
C1-fluoranthene/pyrenes		benzo(a)bifluorene	Fluoranthene		pyrene
C2-fluoranthene/pyrenes		benzo(a)bifluorene	Fluoranthene		pyrene
C3-fluoranthene/pyrenes		benzo(a)bifluorene	Fluoranthene		pyrene
C4-fluoranthene/pyrenes		benzo(a)bifluorene			pyrene
C1-naphthalene/anthophenes				Naphthalene/anthophenes	naphthalene/anthophenes
C2-naphthalene/anthophenes				Naphthalene/anthophenes	naphthalene/anthophenes
C3-naphthalene/anthophenes				Naphthalene/anthophenes	naphthalene/anthophenes
C4-naphthalene/anthophenes					naphthalene/anthophenes
C1-decynes					decyne
C2-decynes					decyne
C3-decynes					decyne
C4-decynes					decyne

Biomarkers - If "representative compound" used for quantitation, list the compound

	Lab 3	Lab 10	Lab 12	Lab 16	Lab 19	Lab 20	Lab 24
Carbazole	NA						
18a(H)-22,29,30-Trisnorhopane	17[B(H)2][M(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17b(H)2,[b(H)]-Hopane	17b(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane
17a(H)-22,29,30-Trisnorhopane	17[B(H)2][M(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17b(H)2,[b(H)]-Hopane	17b(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane
17a(H)2,[B(H)30-Norhopane	17[B(H)2][M(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17b(H)2,[b(H)]-Hopane	17b(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane
18a(H)-30-Stercolopane	17[B(H)2][M(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17b(H)2,[b(H)]-Hopane	17b(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane
17a(H)-Dahngane	17[B(H)2][M(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17b(H)2,[b(H)]-Hopane	17b(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane
17a(H)2[B(H)-Hopane	17[B(H)2][M(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17b(H)2,[b(H)]-Hopane	17b(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane
17a(H)2,[B(H)2-2R-Homohopane	17[B(H)2][M(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17b(H)2,[b(H)]-Hopane	17b(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane
17a(H)2,[B(H)2-2S-Homohopane	17[B(H)2][M(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17b(H)2,[b(H)]-Hopane	17b(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane	17a(H)2,[B(H)]-Hopane
5a(H)-Diacholestane 20S	5a-Cholestane	5(a) Cholestane	5a-Cholestane	5a-Cholestane	5a-Cholestane	5a-Cholestane	5b(H)Cholane
5a(H),14a(H),7a(H)-Cholestane 20S	5a-Cholestane	5(a) Cholestane	5a-Cholestane	5a-Cholestane	5a-Cholestane	5a-Cholestane	5b(H)Cholane
5a(H),14a(H),7a(H)-24-Ethylcholestane 20S	5a-Cholestane	5(a) Cholestane	5a-Cholestane	5a-Cholestane	5a-Cholestane	5a-Cholestane	5b(H)Cholane
5a(H),14a(H),7a(H)-24-Ethylcholestane 20R	5a-Cholestane	5(a) Cholestane	5a-Cholestane	5a-Cholestane	5a-Cholestane	5a-Cholestane	5b(H)Cholane
5a(H),14b(H),7b(H)-Cholestane 20R	5a-Cholestane	5(a) Cholestane	5a-Cholestane	5a-Cholestane	5a-Cholestane	5a-Cholestane	5b(H)Cholane
5a(H),14b(H),7b(H)-Cholestane 20S	5a-Cholestane	5(a) Cholestane	5a-Cholestane	5a-Cholestane	5a-Cholestane	5a-Cholestane	5b(H)Cholane
5a(H),14b(H),7b(H)-24-Ethylcholestane 20R	5a-Cholestane	5(a) Cholestane	5a-Cholestane	5a-Cholestane	5a-Cholestane	5a-Cholestane	5b(H)Cholane
5a(H),14b(H),7b(H)-24-Ethylcholestane 20S	5a-Cholestane	5(a) Cholestane	5a-Cholestane	5a-Cholestane	5a-Cholestane	5a-Cholestane	5b(H)Cholane

APPENDIX C

Laboratory Notes Submitted with the Data

[Notes including data for additional analytes

Lab		QA10TIS01 Sample 1 (ng/g dry mass)	QA10TIS01 Sample 2 (ng/g dry mass)	QA10TIS01 Sample 3 (ng/g dry mass)	SRM 1974b Sample 1 (ng/g dry mass)	SRM 1974b Sample 2 (ng/g dry mass)	SRM 1974b Sample 3 (ng/g dry mass)	SRM 1974b 103 B[<i>a</i>]A-dl2	IS/surrogate used
1	Chrysene + triphenylene	145	140	146	111	100	103		
5	benzo[b]naphtho[2,1-d]thiophene benzo[b]naphtho[1,2-d]thiophene benzo[b]naphtho[2,3-d]thiophene chrysene + triphenylene benzo[f-k]fluoranthene dibenz[a,h+ac]anthracene 1,6,7-trimethylnaphthalene (also known as 2,3,5-trimethylnaphthalene, CAS # 2245-58-7), partially coelutes with another C3-naphthalene, this coeluting peak was split off during peak integration	16.0 3.31 2.70 1.30 31.6 2.36	16.1 3.27 2.45 139 33.6 2.37	16.3 3.07 2.38 147 33.2 2.32	23.4 5.72 5.26 155 62.3 7.17	24.0 5.90 5.41 156 62.2 6.64	23.7 5.70 4.99 168 61.9 6.70	ACI-dl0 ACI-dl0 ACI-dl0 Bap-dl2 Bap-dl2 Bap-dl2	"representative compound" used for quantitation dibenzothiophene dibenzothiophene dibenzothiophene
6	Benzog[ghi]perylene results (wet tissue) is below MDL of 0.60 ng/g (wet tissue) for oyster tissue acenaphthene MDL (wet tissue) is 0.18 ng/g dibenz[ah]anthracene MDL (wet tissue) is 0.55 ng/g indeno[1,2,3-cd]pyrene MDL (wet tissue) is 0.07 ng/g Conservative Estimate of total PAH dry basis Sample 1 5824 ng/g, sample 2 5318 ng/g, sample 3 5744 ng/g Benzo[ghi]perylene has a consistent interfering peak, the peak is corrected for the interference in standards and samples	Batch A Sample 1	Batch B Sample 2	Batch C Sample 3	Batch A Sample 1	Batch B Sample 2	Batch C Sample 3	IS/surrogate used	
7	ALKYLATED PAH ANALYSIS	QA10TIS01 Batch A Sample 1	QA10TIS01 Batch B Sample 1	QA10TIS01 Batch C Sample 3	SRM 1974b Batch A Sample 1	SRM 1974b Batch B Sample 2	SRM 1974b Batch C Sample 3	d10-Acenaphthylene d10-Acenaphthylene d10-Fluorene d10-Phenanthrene d10-Phenanthrene	
8	Other: QA10TIS01 anthracene coeluting with an interfering peak at both quantitation and confirmation ions 178 and 177 respectively. Ion ratios out of range.								
9	note: calibration is based on wet weight basis. other = interference noted for naphthalene and benz[a]anthracene								

Lab Notes including data for additional analytes

10 Any values less than RL will be considered as estimated "n" values.

NA are compounds not calibrated for

Samples were analyzed at a dilution due to the matrix of the sample

11 Other - Blank contaminated with naphthalene

12 *Percent moisture value used for the SRM Material (89.92%) was taken from the SRM Certificate of analysis.

Other - An unknown intertent peak did not allow for quantitation

13 (NOTES: Typical quantitation limit for all analytes in samples and SRM in this matrix are <30 ng/g)

14 (NOTES: Typical quantitation limit for all analytes in samples and SRM in this matrix are <30 ng/g)

	QA10TIS01 Sample 1 (ng/g dry mass)	QA10TIS01 Sample 2 (ng/g dry mass)	QA10TIS01 Sample 3 (ng/g dry mass)	SRM 1974b Sample 1 coelution	SRM 1974b Sample 2 coelution	SRM 1974b Sample 3 coelution
abb 20R 24R-nethylcholestane	19.6	18.8	19.2	59.8	68.6	67.2
1,2-dimethylnaphthalene	13.7	15.4	14.3	11.6	11.5	12.6
3-methylphenanthrene	11.6	12.4	11.6	16.9	17.6	20.3
2-methylphenanthrene	14.3	15.8	14.0	16.9	17.6	17.6
2-naphthylanthracene	17.8	19.0	17.9	9.6	9.8	10.5
9-methylphenanthrene	76.1	79.4	79.0	13.0	11.7	12.7
3-methyl+1-methylfluoranthene	12.5	13.8	13.2	14.0	13.5	15.0
1-methylpyrene	24.8	26.4	24.2	24.1	26.5	25.7
4-methylpyrene	13.6	14.3	13.8	15.1	16.7	16.2
3-methylchrysene	3.47	3.96	3.86	4.65	4.71	5.00
6-methylchrysene	2.97	3.20	2.98	3.60	3.85	3.69
benzof[b]fluoranthene	19.7	21.3	19.9	20.7	23.2	23.4
benzo[e]phenanthrene	14.8	15.4	15.1	19.1	19.6	21.2
dibenz[a]anthracene	<1.5	<1.5	<1.5	4.64	4.58	5.32
dibenz[a]anthracene	<1.5	<1.5	<1.5	4.24	4.59	3.86
benzo[b]chrysene	<1.5	<1.5	<1.5	4.02	4.00	3.88
picene	<1.5	<1.5	<1.5	7.98	7.78	7.98

Lab | Notes including data for additional analytes

Lab	Notes including data for additional analytics																
15	Unable to obtain SRM 1974b as NIST could not ship it across the border to Canada.																
	QA10TIS01 Sample 1 (ng/g dry mass)	QA10TIS01 Sample 2 (ng/g dry mass)	QA10TIS01 Sample 3 (ng/g dry mass)	benzo[b&e]/fluoranthene 0.031 0.021	benzo[b&e]/fluoranthene 0.029	Biomarkers calculated as ratios only:	SUMMARY	Source	Source	Source	Source	Mean	Std Dev	RSD%			
<i>Tarpanes</i>							DR-27Ts/27Tm - (DR-27Ts)	1.190246359	1.228033753	1.093023256	1.284924116	1.199056871	0.080692	6.7			
DR-29(ab)30(ab) - (DR-29(ab))	0.492225719	0.481056548	0.481349995	0.48606703	0.485174823	0.0052313	1.1										
DR-31(ab)31(abR) - (DR-31(ab))	1.438697831	1.442665855	1.377923908	1.385527025	1.411203655	0.0342181	2.4										
DR-32(ab)32(abR) - (DR-32(ab))	1.510096344	1.46006357	1.613340105	1.504377921	0.0792057	5.3											
DR-33(ab)33(abR) - (DR-33(ab))	1.318269399	1.35235746	1.245795021	1.29398136	1.3025349825	0.0447926	3.4										
<i>Sterane</i>							Source	Source	Source	Source	Mean	Std Dev	RSD%				
DR-(27bbR + 27bbS)/(28bbR + 28bbS)	0.706424785	0.721822493	0.723710502	0.719873991	0.717975943	0.0078467	1.1										
DR-(28bbR + 28bbS)/(27bbR + 27bbS)	0.3663375776	0.368973186	0.3579144677	0.364834552	0.364524548	0.004726	1.3										
DR-(29bbR + 29bbS)/(27bbR + 27bbS)	0.466025548	0.451915025	0.463301472	0.4579969	0.459784736	0.0062116	1.4										
<i>Trisubstituted steranes</i>							Source	Source	Source	Source	Mean	Std Dev	RSD%				
DR-C27TA(20S)C28TA(20S) - (DR-C)	0.632985279	0.641149081	0.62770133	0.638608421	0.6108445	1.7											
DR-C27TA(20R)C28TA(20R) - (DR-C)	0.965588774	0.965805785	0.940931607	0.92509641	0.0158668	1.7											
DR-C28TA(20R)C28TA(20S) - (DR-C)	0.762307833	0.763949175	0.793219152	0.770889741	0.0149081	1.9											
<i>PAHs</i>							C3-D/C3-P - (DR-C3D/C3P)	0.199698226	0.201890811	0.200803325	0.200797521	0.0010963	0.5				
C2-D/C2-P - (DR-C2D/C2P)	0.155768311	0.160931939	0.156201044	0.157633764	0.0028645	1.8											
CRITICAL DIFFERENCE ANALYSIS																	
L981972-1							DR-27Ts	DR-29ab	DR-32ab	DR-33ab	DR-27bbR	DR-29bbR	DR-C26TA	DR-C27-TA	DR-C28TA	C3-D/C3-P	C2-D/C2-P
Difference	1.140323092	0.809296365	1.22763466	1.942802669	1.44149766	0.63555281	0.37271401	0.5156381	0.6980482	1.2392996	0.5901263	0.1394943	0.2543507				
Critical Difference	0.058733779	-0.324121542	0.183368994	-0.438424748	-0.139920834	0.0824298	-0.007616	-0.055853	-0.05944	-0.28723	0.1807634	0.0613032	-0.096717				
Conclusion	0.163756597	0.090612983	0.184718862	0.24130641	0.19286474	0.094744	0.0515665	0.0682796	0.093566	0.1533958	0.0952711	0.0238204	0.0288389				
L981972-2							Match	Match	Match	Match	No Match	No Match	No Match	No Match	No Match	No Match	
Difference	1.350643777	0.787543882	1.169922762	1.574833174	1.461805556	0.6183365	0.3655331	0.5390136	0.866129	1.4419263	0.5693548	0.1687429	0.3083067				
Critical Difference	-0.151586946	-0.865468954	-0.421280892	-0.070455253	-0.15921073	0.0996214	-0.001009	-0.227521	-0.489857	0.2015349	0.032046	-0.150673					
Conclusion	0.17897045	0.089090309	0.180678489	0.215544777	0.19358027	0.0935406	0.0511014	0.0699159	0.1053316	0.1675797	0.0938171	0.0258678	0.0320158				
L981972-3							No Match	No Match	No Match	No Match	No Match	No Match	No Match	No Match	No Match	No Match	
Difference	-0.211646838	0.791306183	1.221881838	1.673452769	1.236477987	0.6592533	0.3662597	0.502869	0.8145859	1.4906977	0.5315204	0.142206	0.276259				
Critical Difference	-0.202589967	-0.30613136	-0.18921817	-0.16904448	-0.066116838	0.0567047	-0.001735	-0.043084	-0.175977	-0.338628	0.2396973	0.0585915	-0.118625				
Conclusion	0.16874926	0.089353567	0.184315884	0.222448148	0.177755097	0.0664048	0.0511549	0.0709937	0.0916187	0.1017236	0.0709937	0.0240140	0.0303725				

Notes: including data for additional analytes

		< symbol refers to values less than our MDL.
27	The reported value of chrysene is the sum of chrysene and triphenylene. The reported value of benzo(k)fluoranthene is the sum of benzo(k)fluoranthene and benzo(j)fluoranthene	
	QA10TIS01 QA10TIS01 QA10TIS01 QA10TIS01 QA10TIS01 QA10TIS01 QA10TIS01	SRM 1974b SRM 1974b SRM 1974b SRM 1974b SRM 1974b SRM 1974b SRM 1974b
	Sample 1 Sample 2 Sample 3 Sample 1 Sample 2 Sample 3 Sample 1	(ng/g dry mass) (ng/g dry mass)
	90.9 86 98 116 102 NA NA	
	15.8 14.5 13.9 55.4 55.4 NA NA	
29	Most of our detection limits ranged ~10 to 50 ng/g dry, some were lower or as high as ~200. Other = We used a methyl PAH cocktail to calibrate and assess recoveries. Although it did not contain C4-Naphthalenes, we measured significant amounts using the standard protocol.	
	Our numbers were DL in 1974b and ~1300ng/g dry in QATISS10.	
	We also analyzed a suite of 28 Methyl PAHs; only the above were detected.	
	Figures lacking three significant figures should be assumed to have lost trailing zeros.	
	Chrysene and triphenylene quantified together as one chromatographic peak.	
	Benzo(j)fluoranthene and benzo(k)fluoranthene quantified together (although we do not assume both are present in samples) as one chromatographic peak.	
	2,6- and 2,7-methylnaphthalene were quantified together (although we do not assume both are present in samples) as one chromatographic peak.	
	Other Analytes:	
	QA10TIS01 QA10TIS01 QA10TIS01 QA10TIS01 QA10TIS01 QA10TIS01 QA10TIS01	SRM 1974b SRM 1974b SRM 1974b SRM 1974b SRM 1974b SRM 1974b SRM 1974b
	Sample 1 Sample 2 Sample 3 Sample 1 Sample 2 Sample 3 Sample 1	(ng/g dry mass) (ng/g dry mass)
	1.26 1.37 1.37 15.0 16.5 12.9 1S/surrogate	
	3-McPhenanthrene benzo[g,i]fluoranthene+cyclopenta[c,d]p.	used
	benzo[c]phenanthrene chrysene + triphenylene	d10-Anthracene
	benzo[b+j+k]fluoranthene	d10-Pyrene
32	QA10TIS01 QA10TIS01 QA10TIS01 QA10TIS01 QA10TIS01 QA10TIS01 QA10TIS01	SRM 1974b SRM 1974b SRM 1974b SRM 1974b SRM 1974b SRM 1974b SRM 1974b
	Sample 1 Sample 2 Sample 3 Sample 1 Sample 2 Sample 3 Sample 1	(ng/g dry mass) (ng/g dry mass)
	<25 <26 <25 <25 <25 <25 106	
34	1) chrysene is reported as a coelution of both chrysene and triphenylene. 2) benzo(k)fluoranthene is reported as a coelution of both benzo(l)fluoranthene and benzo(k)fluoranthene. 3) Dibenz[a,h]anthracene is reported as a coelution of both Dibenz[a,h]anthracene and Dibenz[a,c]anthracene.	
	QA10TIS01 QA10TIS01 QA10TIS01 QA10TIS01 QA10TIS01 QA10TIS01 QA10TIS01	SRM 1974b SRM 1974b SRM 1974b SRM 1974b SRM 1974b SRM 1974b SRM 1974b
	Sample 1 Sample 2 Sample 3 Sample 1 Sample 2 Sample 3 Sample 1	(ng/g dry mass) (ng/g dry mass)
	107 96.4 104 106 113 115 1S/surrogate	
	41.1 26.6 34 54 53.1 58.3 used	acceptance
	4.04 2.72 4.14 7.21 4.83 4.84	Chrysene-d12
		Chrysene-d12
		Chrysene-d12

APPENDIX D

Charts of QA10TIS01 and SRM 1974b Results by Analyte

See Tables 1 through 3 for results reported as <number>, detection limit, etc.

For QA10TIS01 plots:

Solid line: exercise assigned value

Dotted line: $z = \pm 1$, i. e., 25 % from assigned value

Dotted/dashed line: $z = \pm 2$, i. e., 50 % from assigned value

Dashed line: $z = \pm 3$, i. e., 75 % from assigned value

For SRM 1974b plots:

Solid line: material certified concentration or target value (see caption of each plot)

Dotted line: 95 % confidence interval (CI)

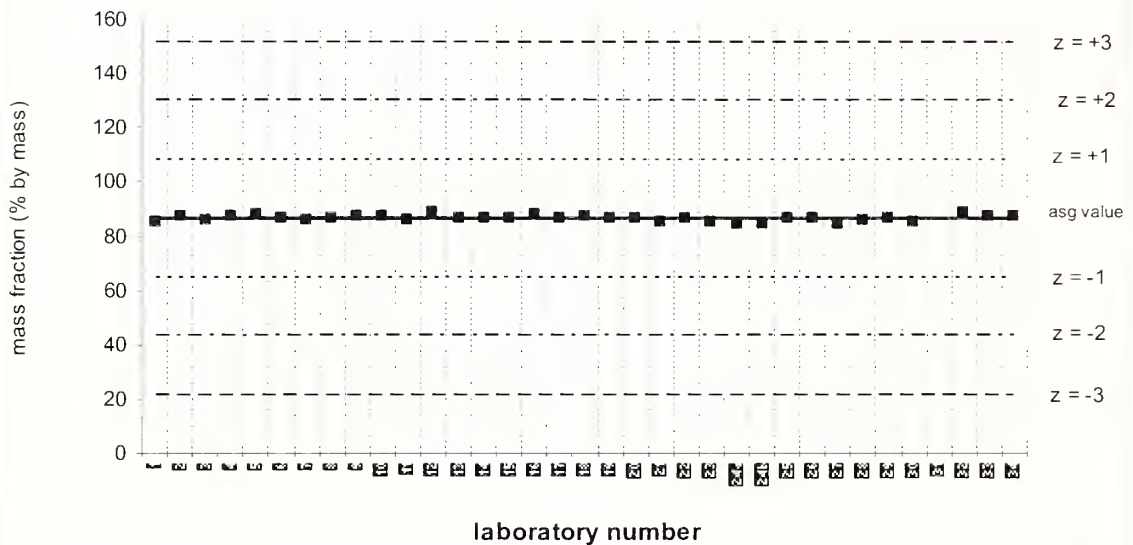
Dashed line: 30 % from 95 % confidence interval (CI)

Percent Water

QA10TIS01

Assigned value = 86.7 % by mass $s = 1.1$ % by mass 95% CI = 0.4 % by mass Median value = 86.8 % by mass

Reported Results: 34 Quantitative Results: 34



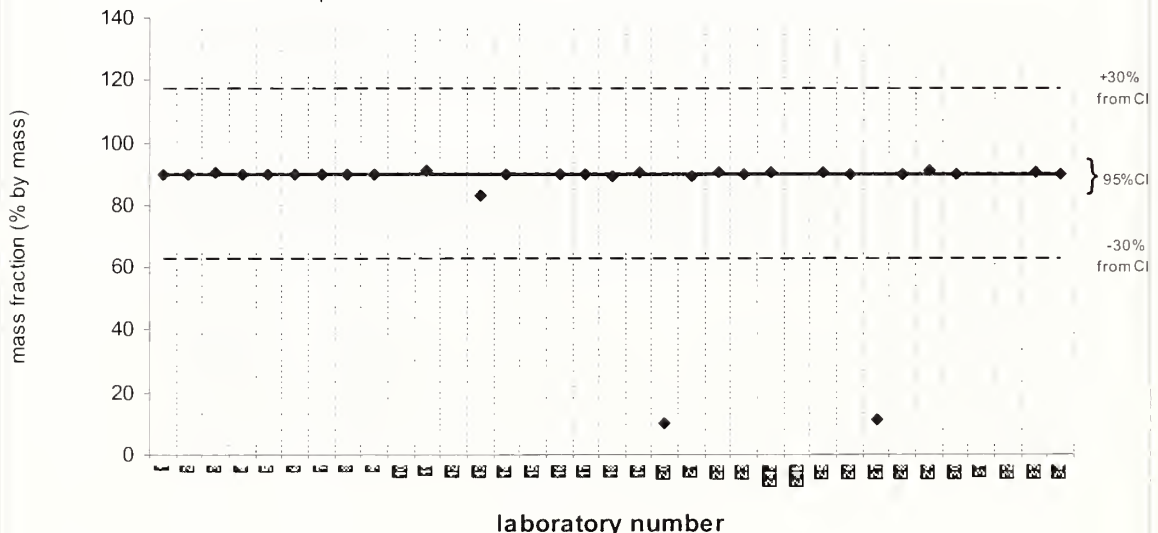
Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

Percent Water

SRM 1974b

Target Value = 89.9 % by mass ; 95% CI 0.1 % by mass: Median value = 89.9 % by mass

Reported Results: 29 Quantitative Results: 29

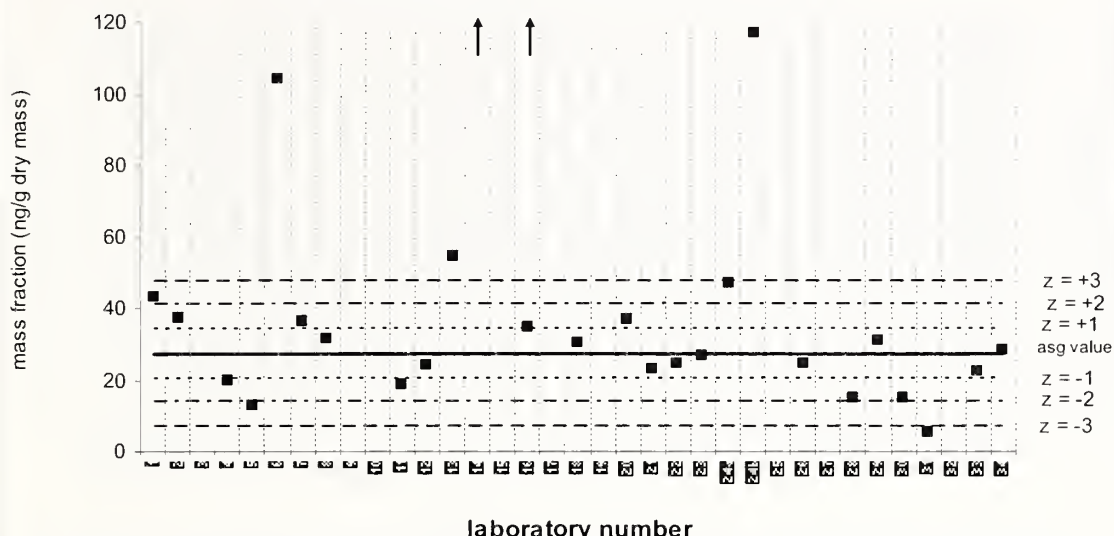


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

naphthalene**QA10TIS01**

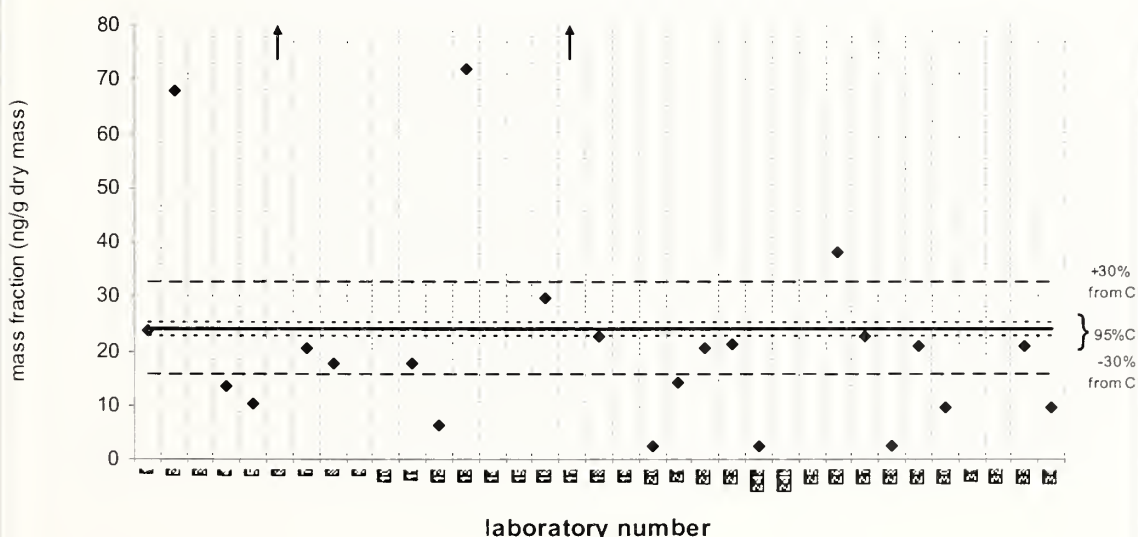
Assigned value = 27.2 ng/g dry mass s = 9.2 ng/g dry mass 95% CI = 4.1 ng/g dry mass Median value = 30.2 ng/g dry mass

Reported Results: 33 Quantitative Results: 27

**naphthalene****SRM 1974b**

Certified Value = 24.0 ng/g dry mass ; 95% CI 1.2 ng/g dry mass: Median value = 20.7 ng/g dry mass

Reported Results: 29 Quantitative Results: 25

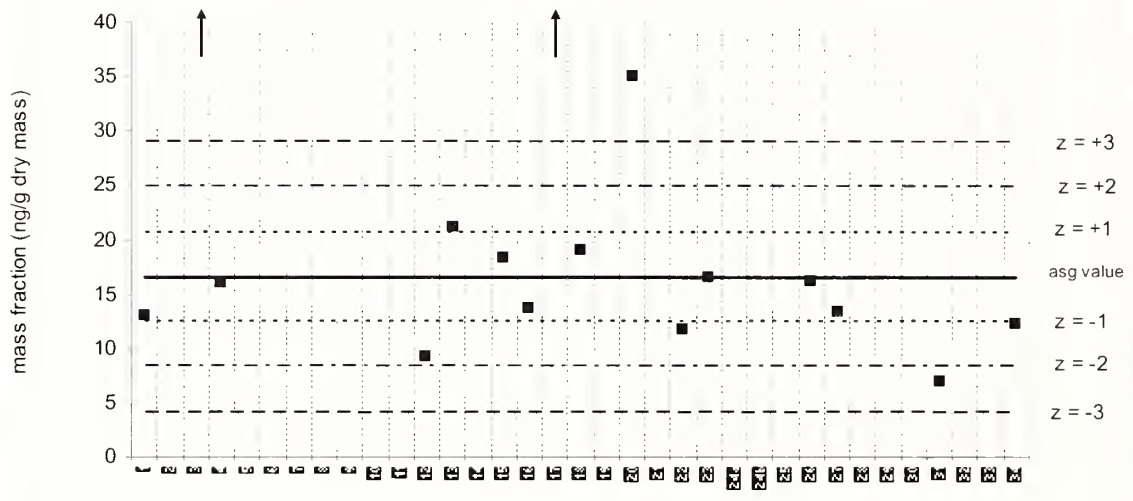


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

biphenyl**QA10TIS01**

Assigned value = 16.6 ng/g dry mass s = 6.4 ng/g dry mass 95% CI = 3.5 ng/g dry mass Median value = 16.1 ng/g dry mass

Reported Results: 22 Quantitative Results: 16

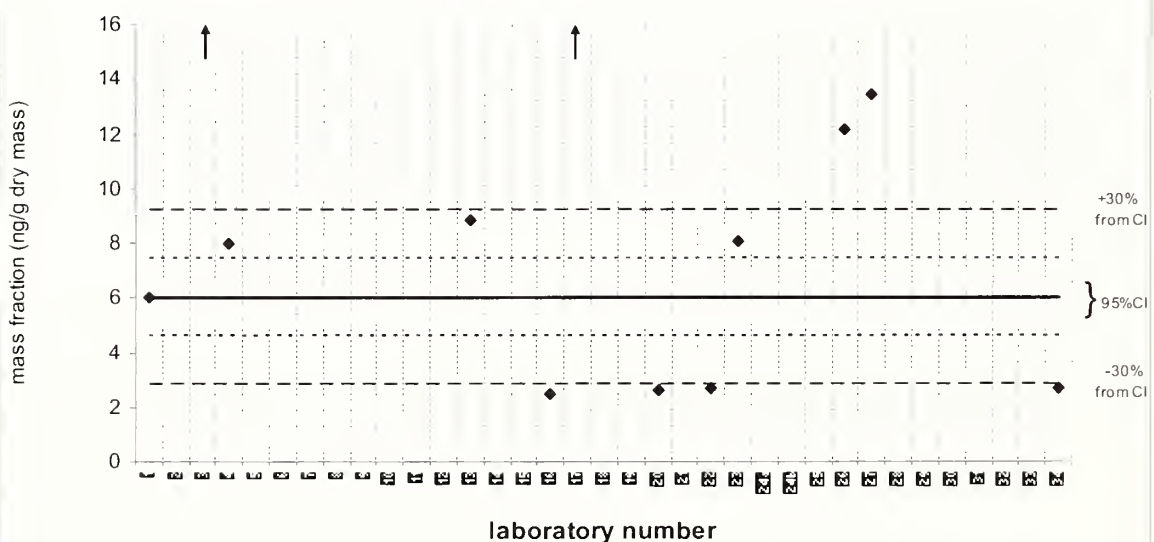


Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

biphenyl**SRM 1974b**

Reference Value = 6.0 ng/g dry mass ; 95% CI 1.4 ng/g dry mass: Median value = 8.0 ng/g dry mass

Reported Results: 18 Quantitative Results: 12



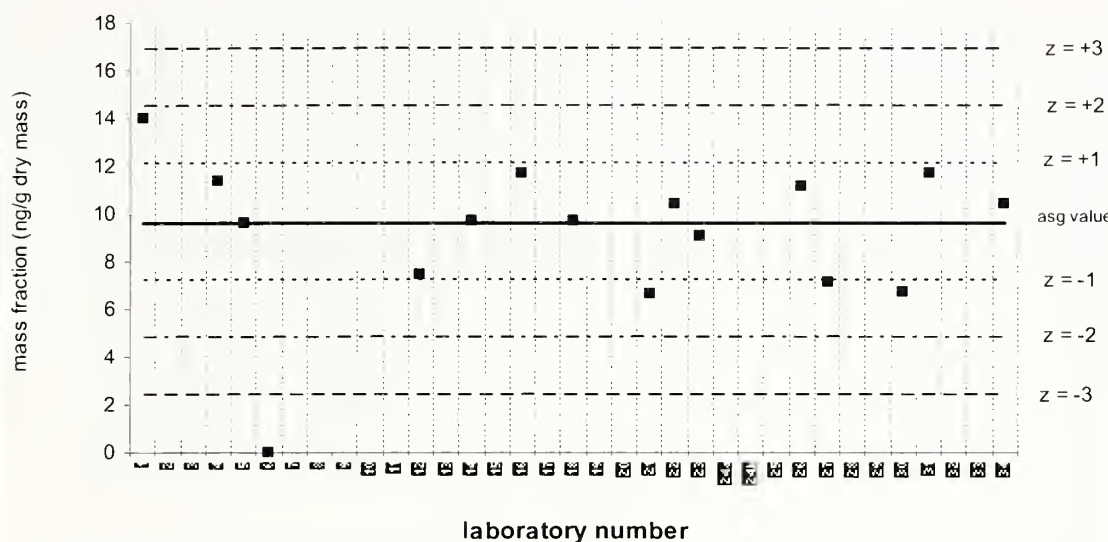
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

acenaphthene**QA10TIS01**

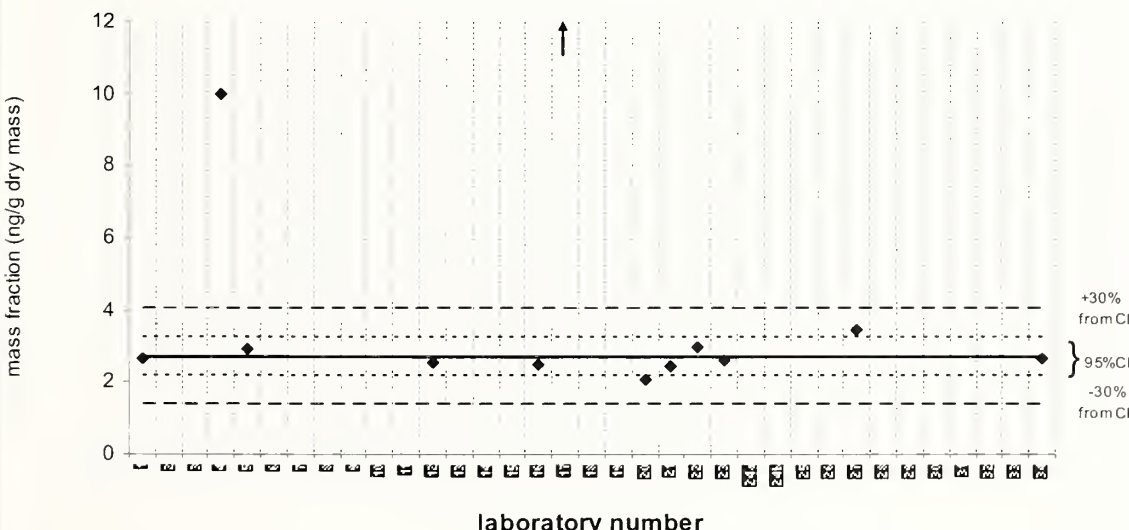
Assigned value = 9.63 ng/g dry mass s = 2.13 ng/g dry mass 95% CI = 1.11 ng/g dry mass

Median value = 9.67 ng/g dry mass

Reported Results: 30 Quantitative Results: 16

**acenaphthene****SRM 1974b**

Reference Value = 2.70 ng/g dry mass ; 95% CI 0.53 ng/g dry mass: Median value = 2.66 ng/g dry

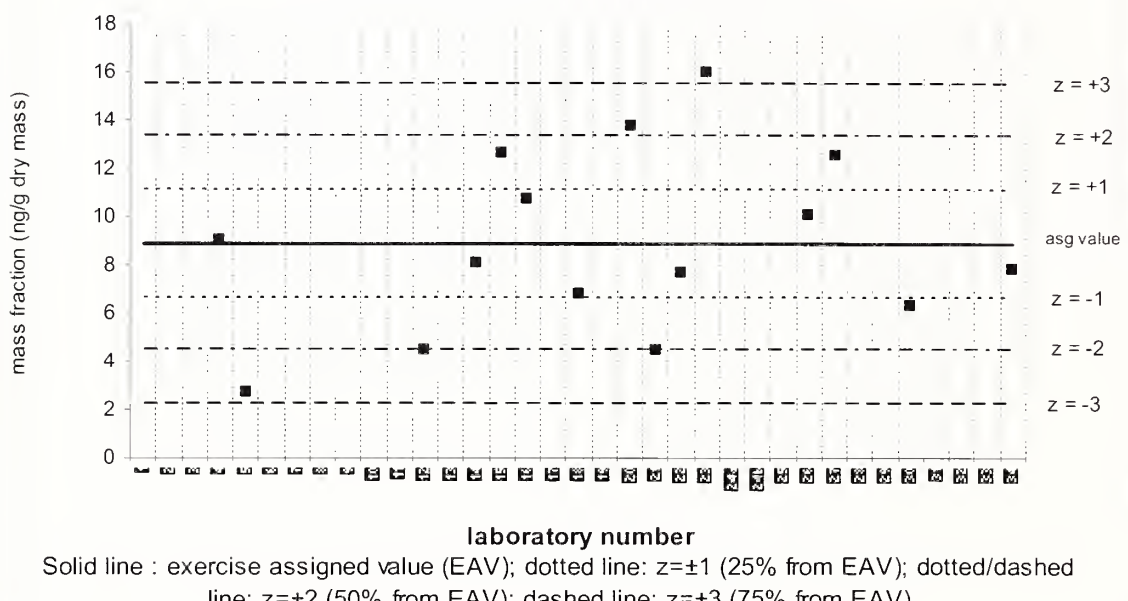
mass
Reported Results: 22 Quantitative Results: 12

acenaphthylene**QA10TIS01**

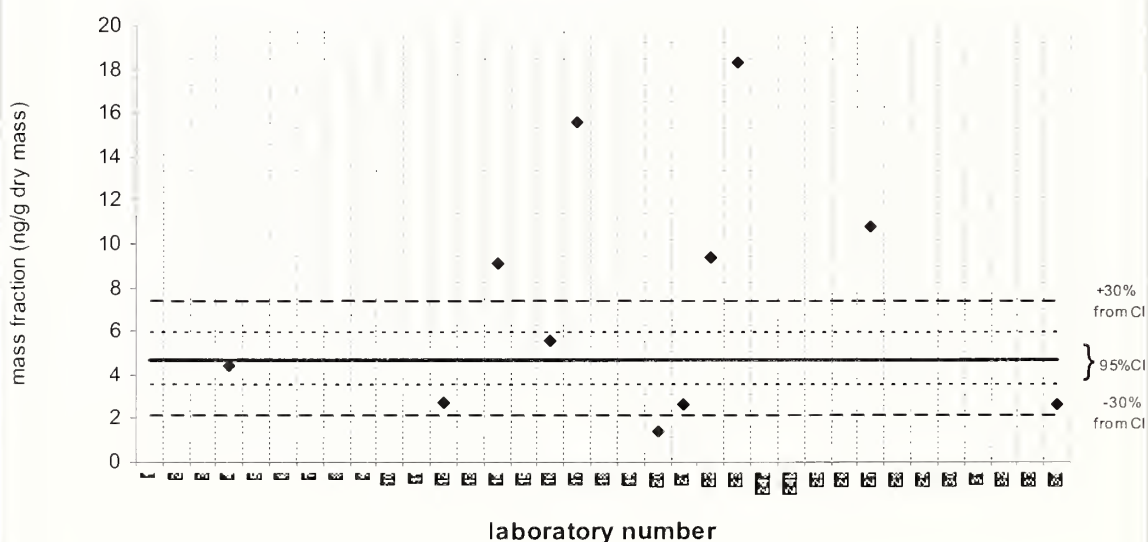
Assigned value = 8.88 ng/g dry mass s = 3.76 ng/g dry mass 95% CI = 1.90 ng/g dry mass

Median value = 8.08 ng/g dry mass

Reported Results: 26 Quantitative Results: 15

**acenaphthylene****SRM 1974b**

Reference Value = 4.70 ng/g dry mass ; 95% CI 1.20 ng/g dry mass: Median value = 5.60 ng/g dry

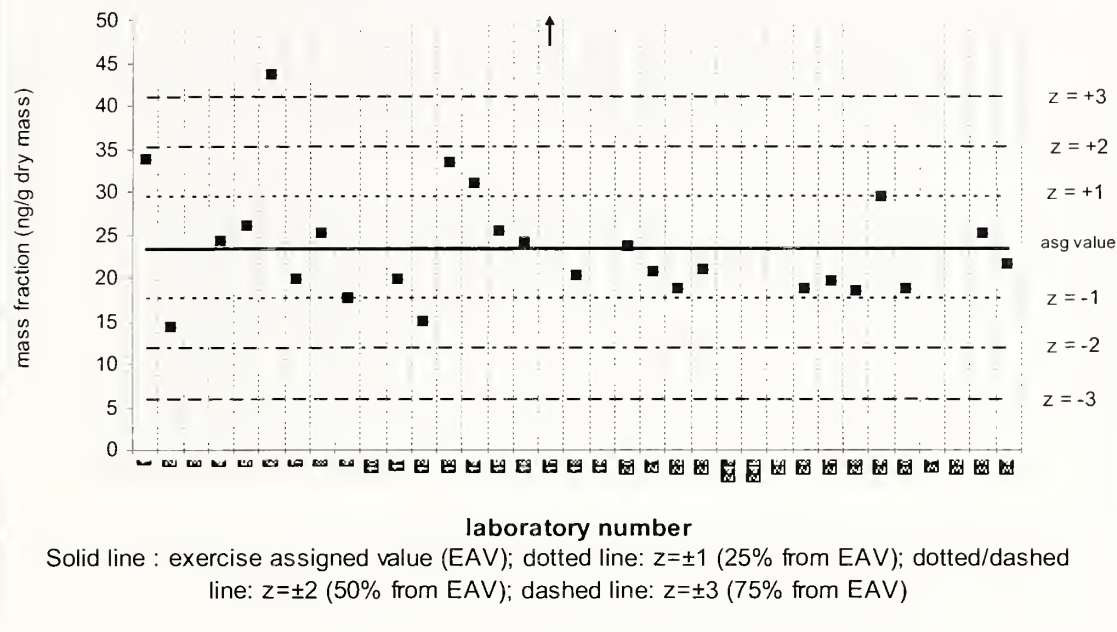
mass
Reported Results: 21 Quantitative Results: 11

Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

fluorene**QA10TIS01**

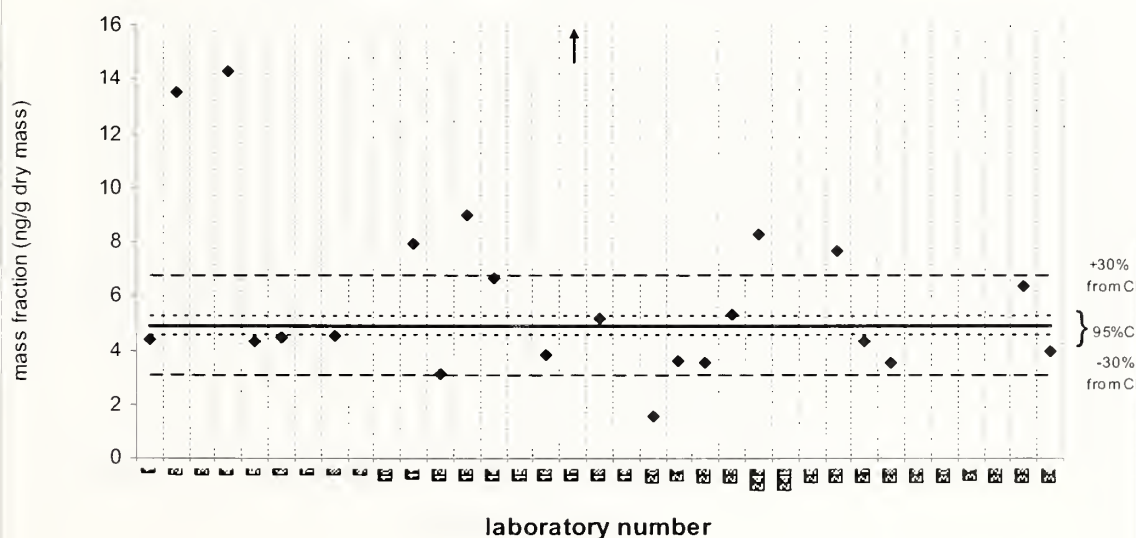
Assigned value = 23.4 ng/g dry mass s = 6.6 ng/g dry mass 95% CI = 2.5 ng/g dry mass Median value = 21.5 ng/g dry mass

Reported Results: 33 Quantitative Results: 27

**fluorene****SRM 1974b**

Certified Value = 4.88 ng/g dry mass ; 95% CI 0.36 ng/g dry mass: Median value = 4.53 ng/g dry mass

Reported Results: 30 Quantitative Results: 23

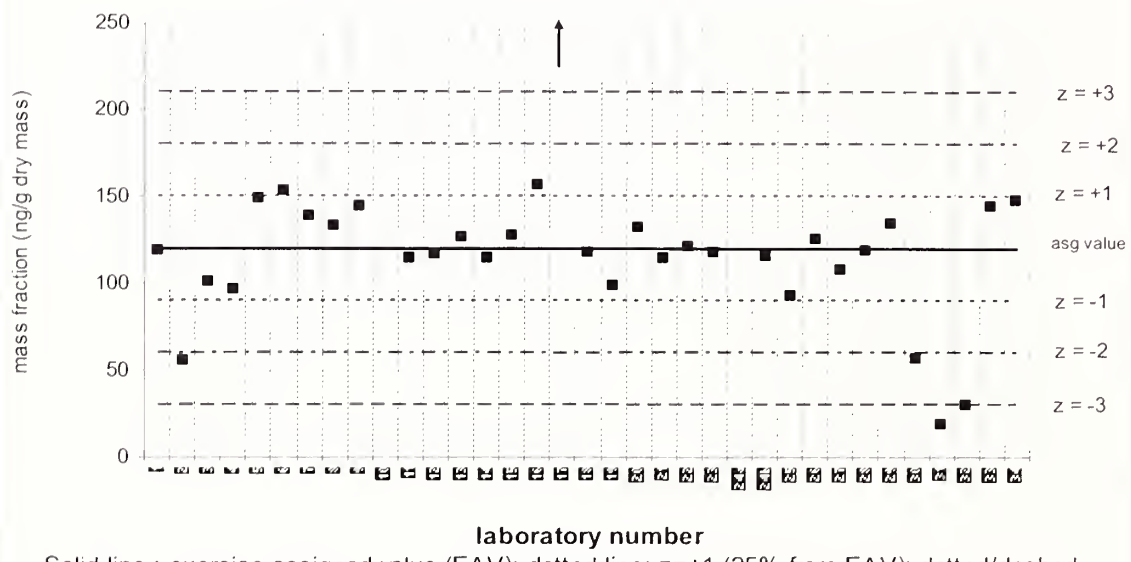


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

phenanthrene**QA10TIS01**

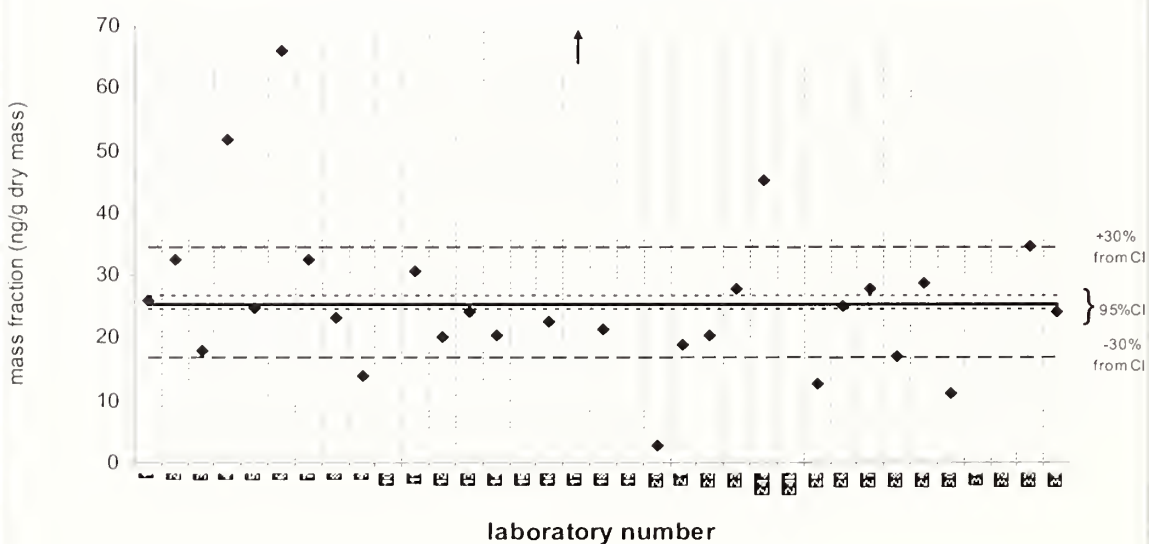
Assigned value = 120 ng/g dry mass $s = 24$ ng/g dry mass 95% CI = 9 ng/g dry mass Median value = 119 ng/g dry mass

Reported Results: 34 Quantitative Results: 33

**phenanthrene****SRM 1974b**

Certified Value = 25.5 ng/g dry mass ; 95% CI 1.1 ng/g dry mass: Median value = 24.2 ng/g dry mass

Reported Results: 31 Quantitative Results: 29

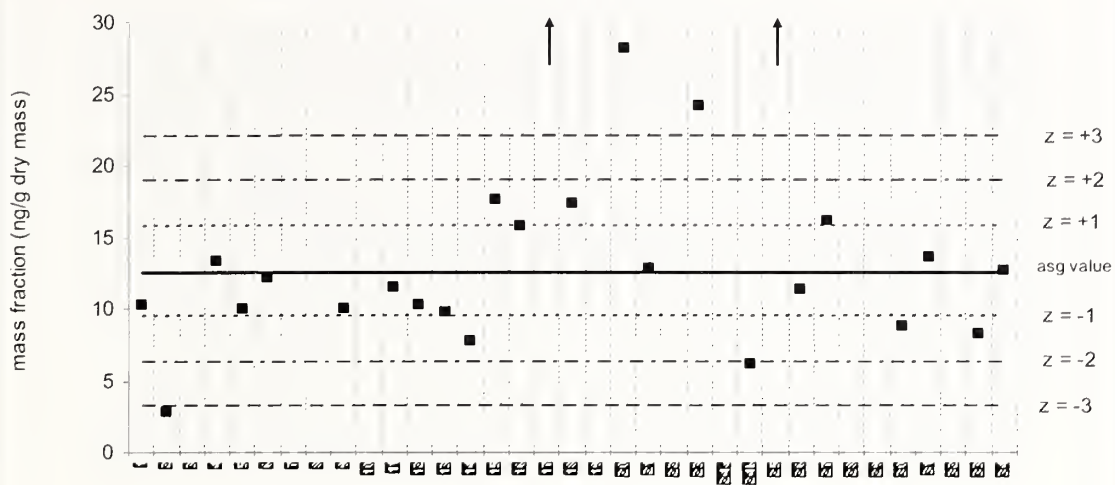


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

anthracene**QA10TIS01**

Assigned value = 12.6 ng/g dry mass $s = 5.7$ ng/g dry mass 95% CI = 2.4 ng/g dry mass Median value = 12.1 ng/g dry mass

Reported Results: 32 Quantitative Results: 25

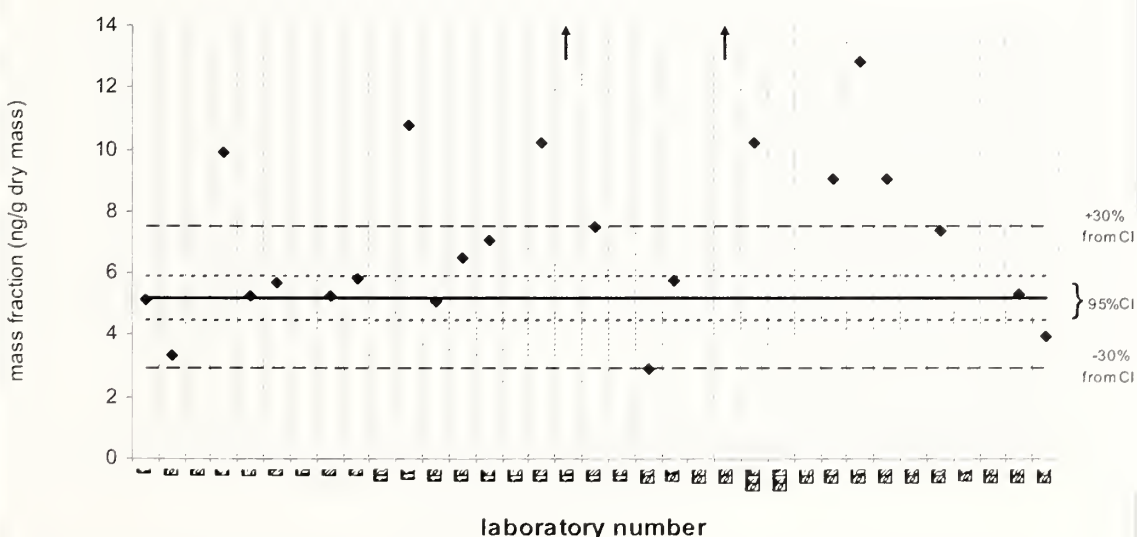


Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

anthracene**SRM 1974b**

Certified Value = 5.20 ng/g dry mass ; 95% CI 0.71 ng/g dry mass: Median value = 6.77 ng/g dry

mass
Reported Results: 30 Quantitative Results: 24

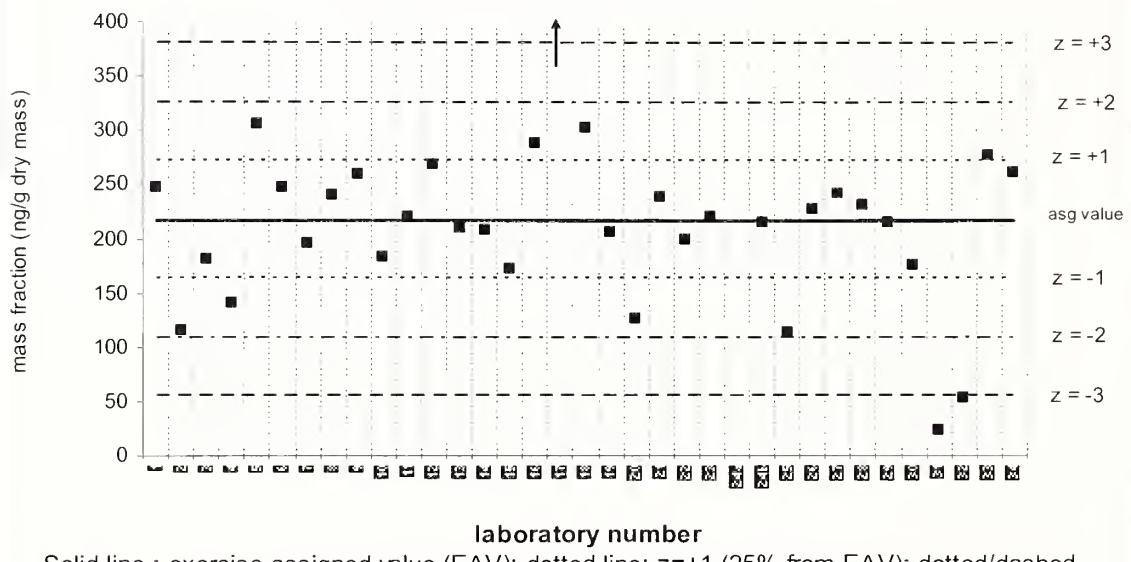


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

fluoranthene**QA10TIS01**

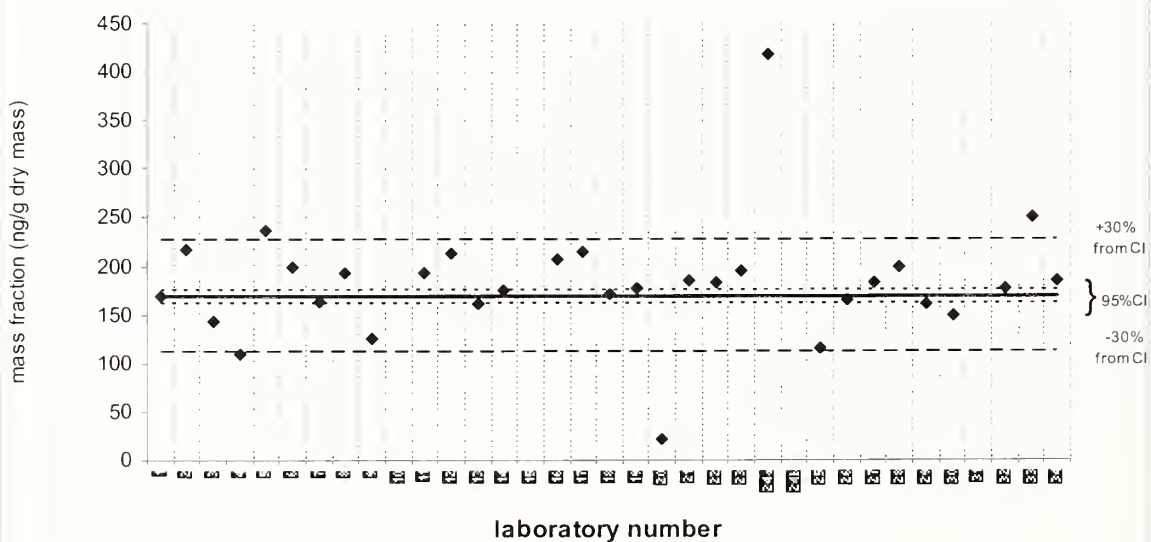
Assigned value = 217 ng/g dry mass s = 50 ng/g dry mass 95% CI = 17 ng/g dry mass Median value = 218 ng/g dry mass

Reported Results: 34 Quantitative Results: 34

**fluoranthene****SRM 1974b**

Certified Value = 169 ng/g dry mass ; 95% CI 7 ng/g dry mass: Median value = 184 ng/g dry mass

Reported Results: 31 Quantitative Results: 31

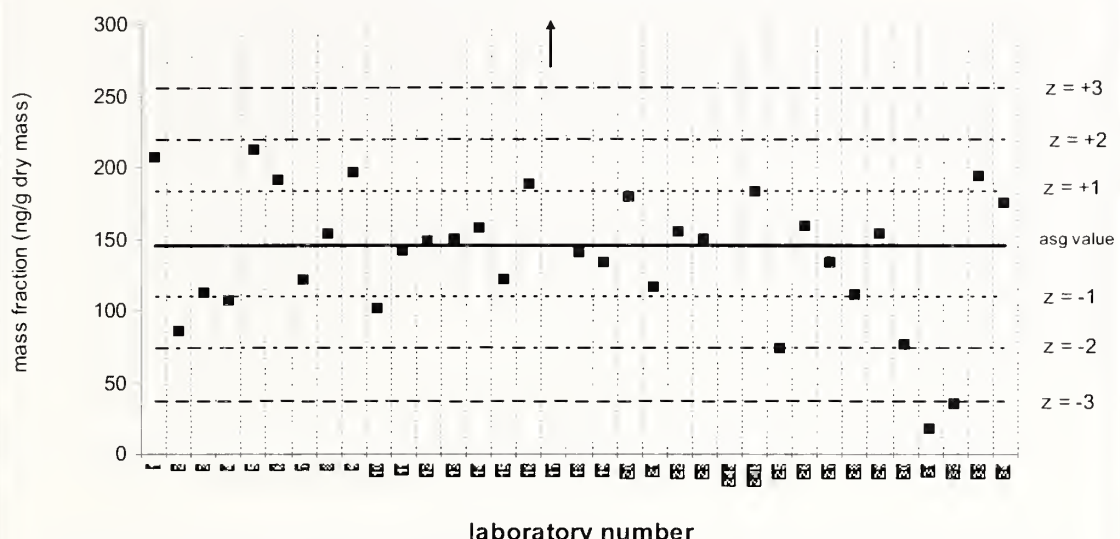


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

pyrene**QA10TIS01**

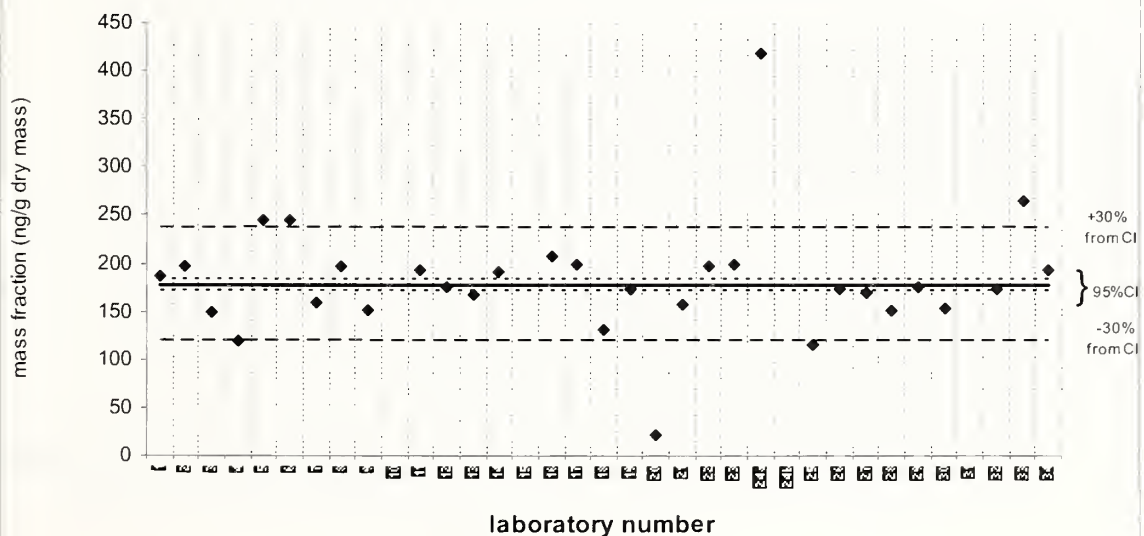
Assigned value = 146 ng/g dry mass $s = 38$ ng/g dry mass 95% CI = 13 ng/g dry mass Median value = 149 ng/g dry mass

Reported Results: 34 Quantitative Results: 34

**pyrene****SRM 1974b**

Certified Value = 178 ng/g dry mass ; 95% CI 6 ng/g dry mass: Median value = 175 ng/g dry mass

Reported Results: 31 Quantitative Results: 31

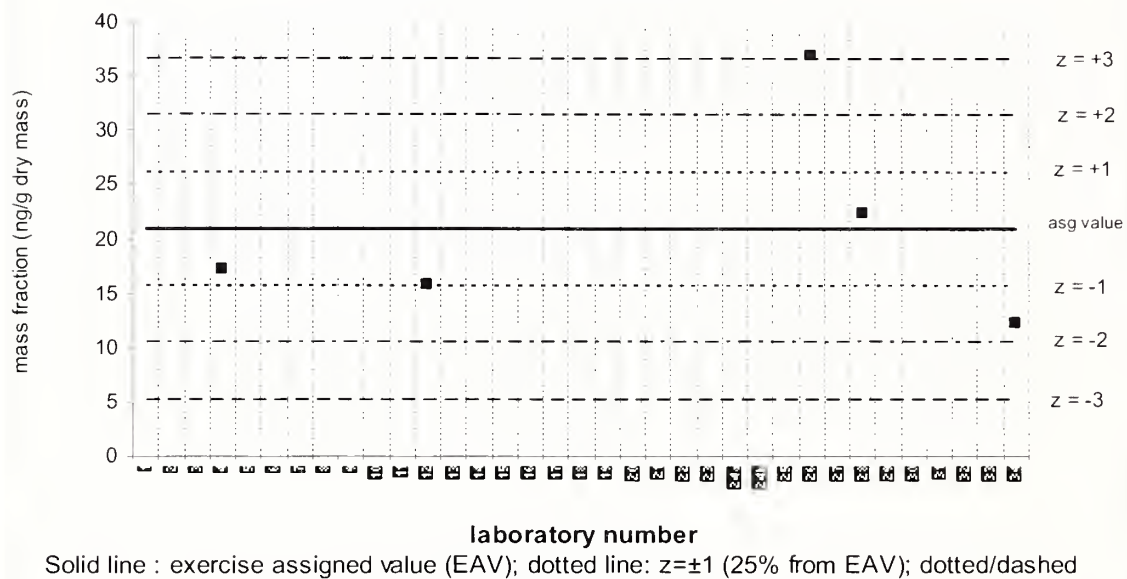


97

benzo[b]fluorene**QA10TIS01**

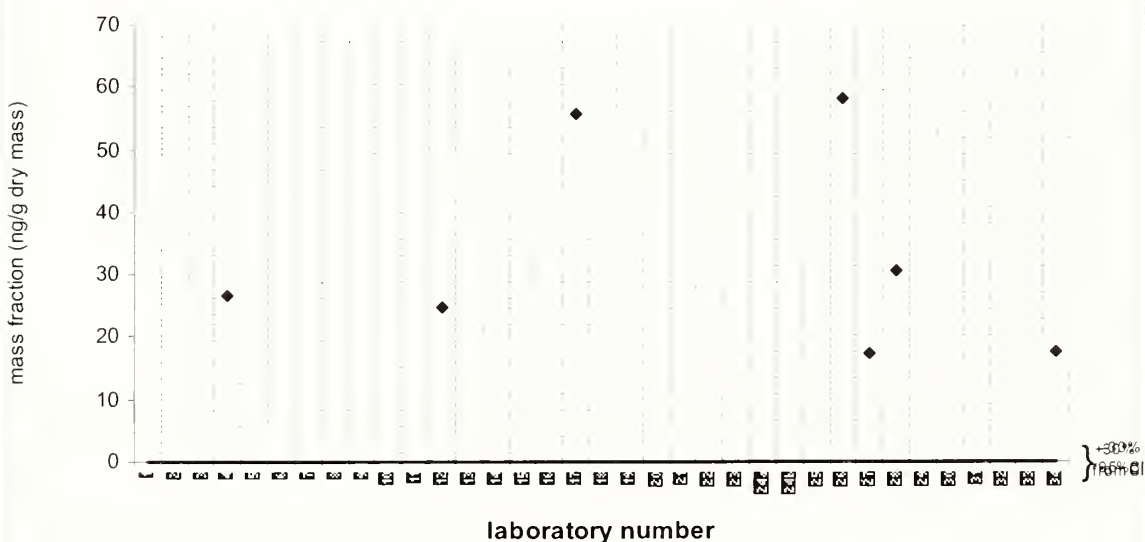
Assigned value = 20.9 ng/g dry mass $s = 9.7$ ng/g dry mass 95% CI = 8.5 ng/g dry mass Median value = 17.3 ng/g dry mass

Reported Results: 8 Quantitative Results: 5

**benzo[b]fluorene****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 26.7 ng/g dry mass

Reported Results: 9 Quantitative Results: 7

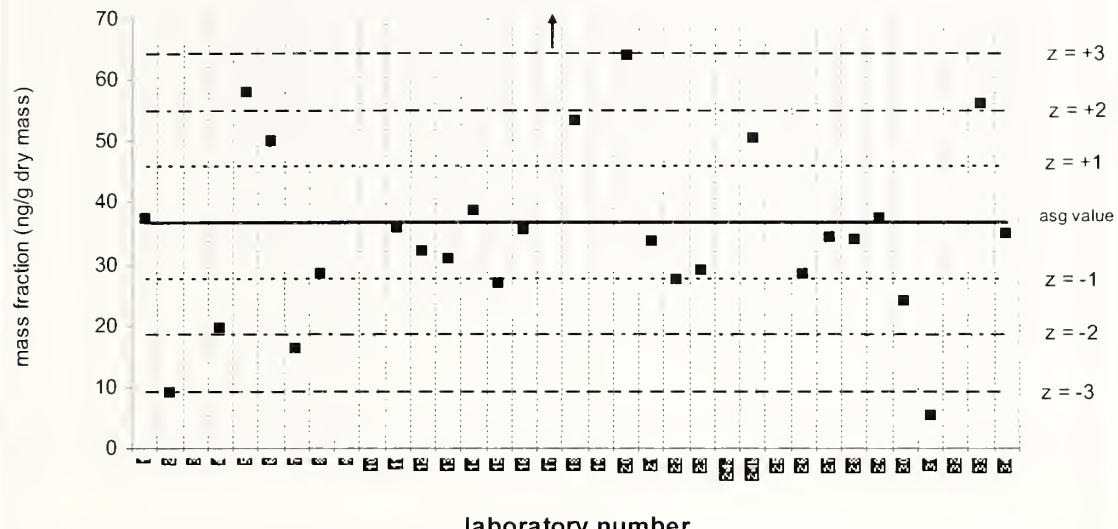


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

benz[a]anthracene**QA10TIS01**Assigned value = 36.6 ng/g dry mass $s = 12.1$ ng/g dry mass 95% CI = 4.7 ng/g dry mass

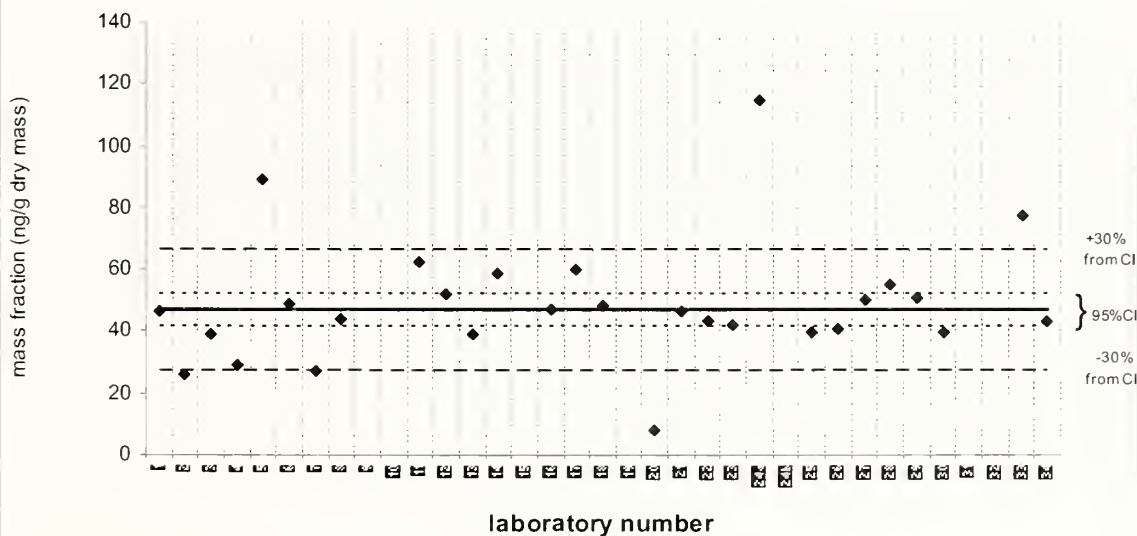
Median value = 34.0 ng/g dry mass

Reported Results: 33 Quantitative Results: 28

**benz[a]anthracene****SRM 1974b**

Certified Value = 46.8 ng/g dry mass ; 95% CI 5.2 ng/g dry mass: Median value = 46.5 ng/g dry mass

Reported Results: 30 Quantitative Results: 28

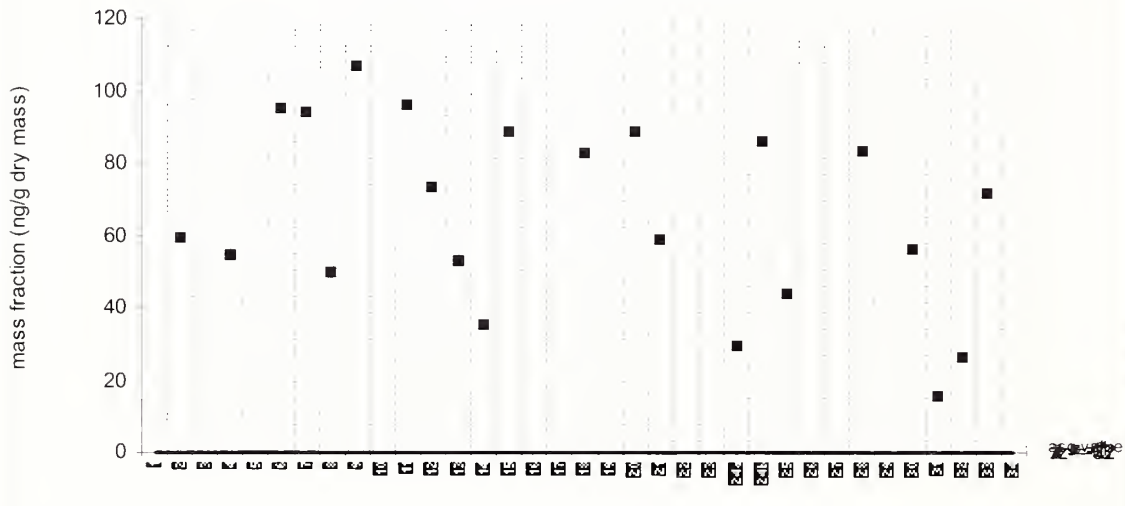


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

chrysene**QA10TIS01**

Assigned value = No Target ng/g (dry mass) Median value = 65.2 ng/g dry mass

Reported Results: 25 Quantitative Results: 22

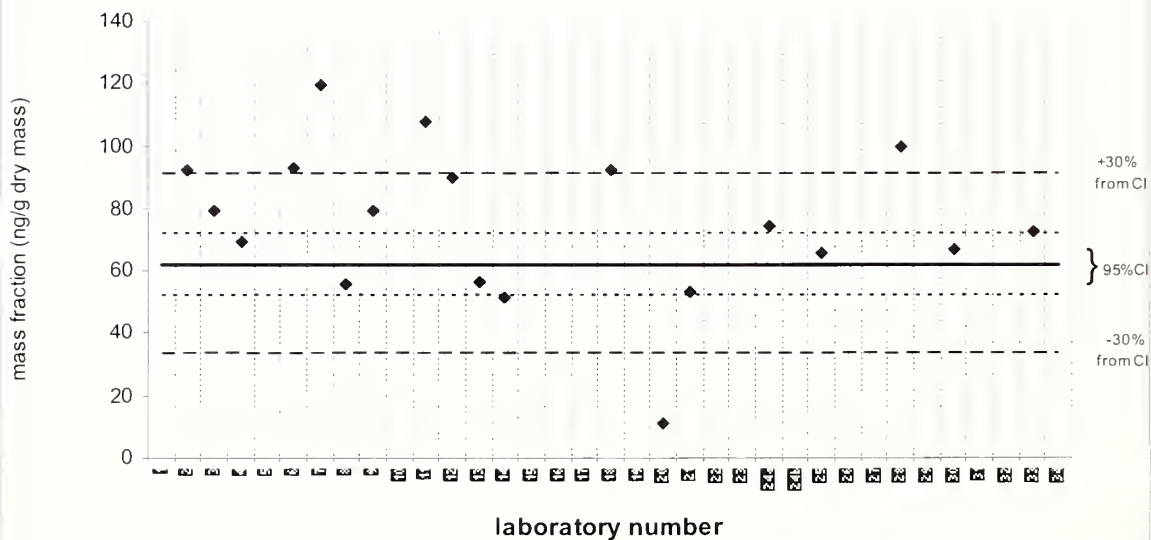
**laboratory number**

Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

chrysene**SRM 1974b**

Certified Value = 62.2 ng/g dry mass ; 95% CI 9.9 ng/g dry mass: Median value = 74.6 ng/g dry mass

Reported Results: 21 Quantitative Results: 19

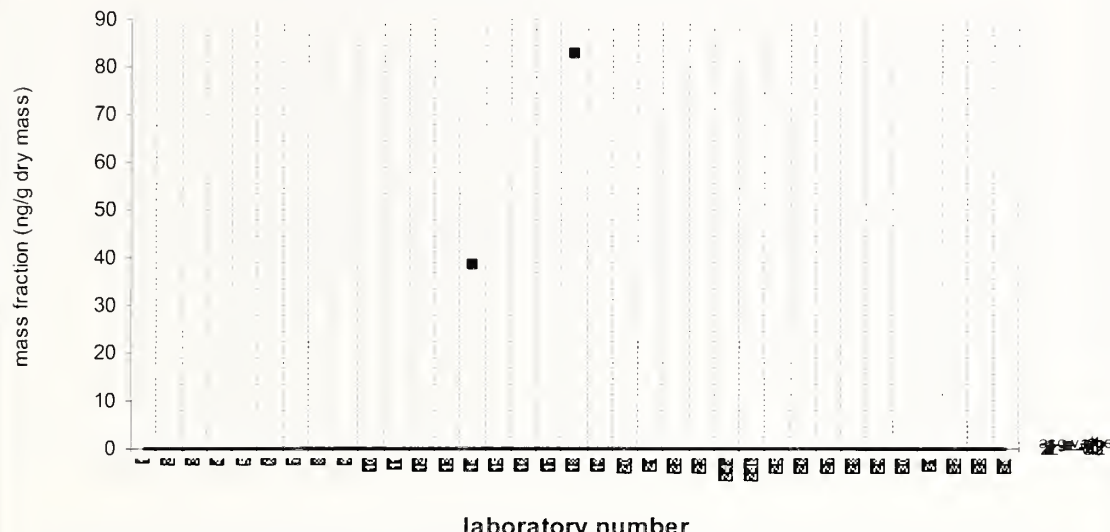
**laboratory number**

Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

triphenylene**QA10TIS01**

Assigned value = No Target ng/g (dry mass) Median value = 60.4 ng/g dry mass

Reported Results: 2 Quantitative Results: 2

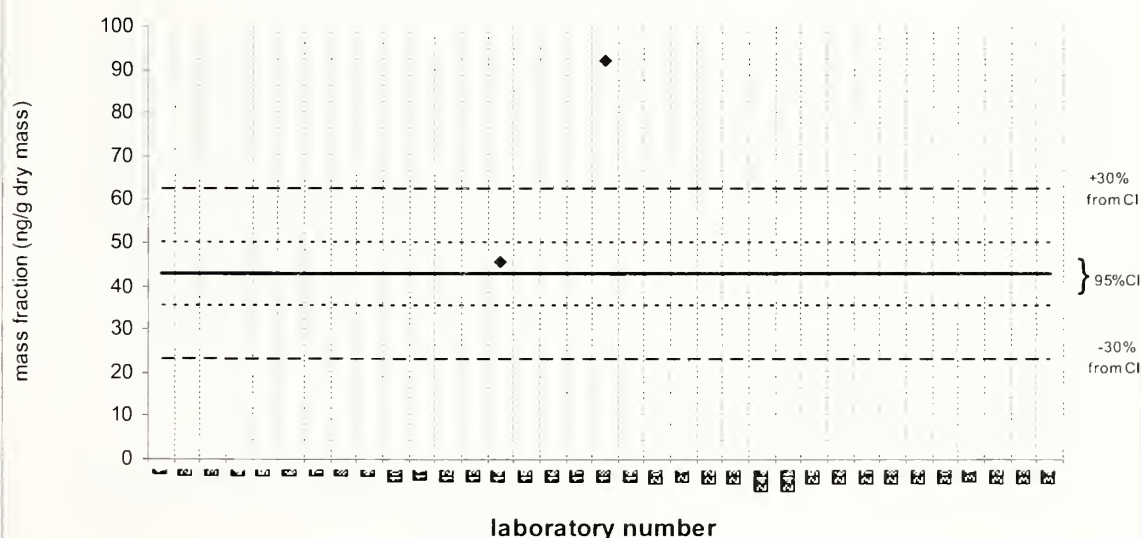
**laboratory number**

Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

triphenylene**SRM 1974b**

Certified Value = 42.7 ng/g dry mass ; 95% CI 7.1 ng/g dry mass: Median value = 68.9 ng/g dry mass

Reported Results: 2 Quantitative Results: 2

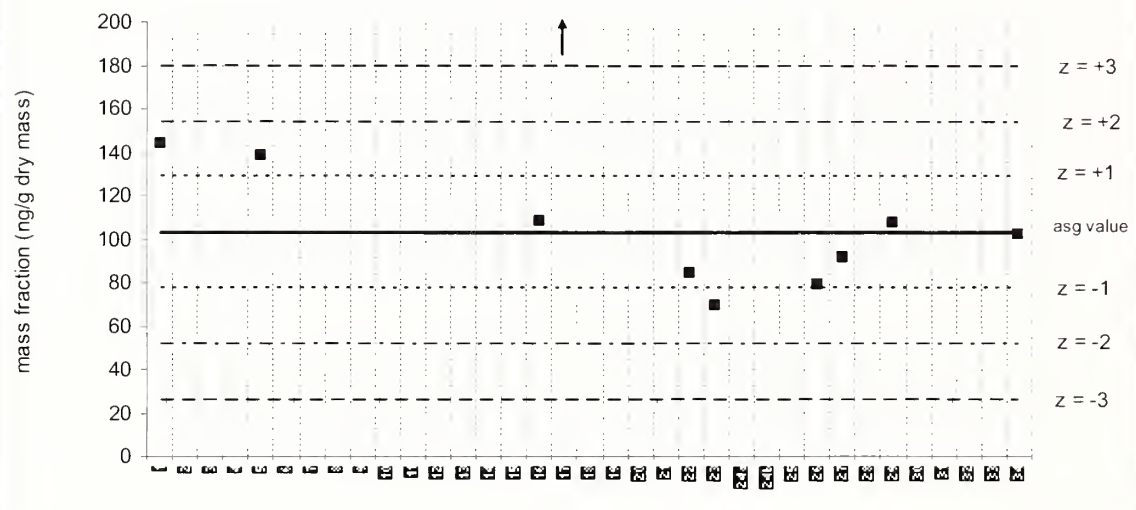
**laboratory number**

Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

chrysene/triphenylene**QA10TIS01**

Assigned value = 103 ng/g dry mass s = 25 ng/g dry mass 95% CI = 17 ng/g dry mass Median value = 105 ng/g dry mass

Reported Results: 10 Quantitative Results: 10

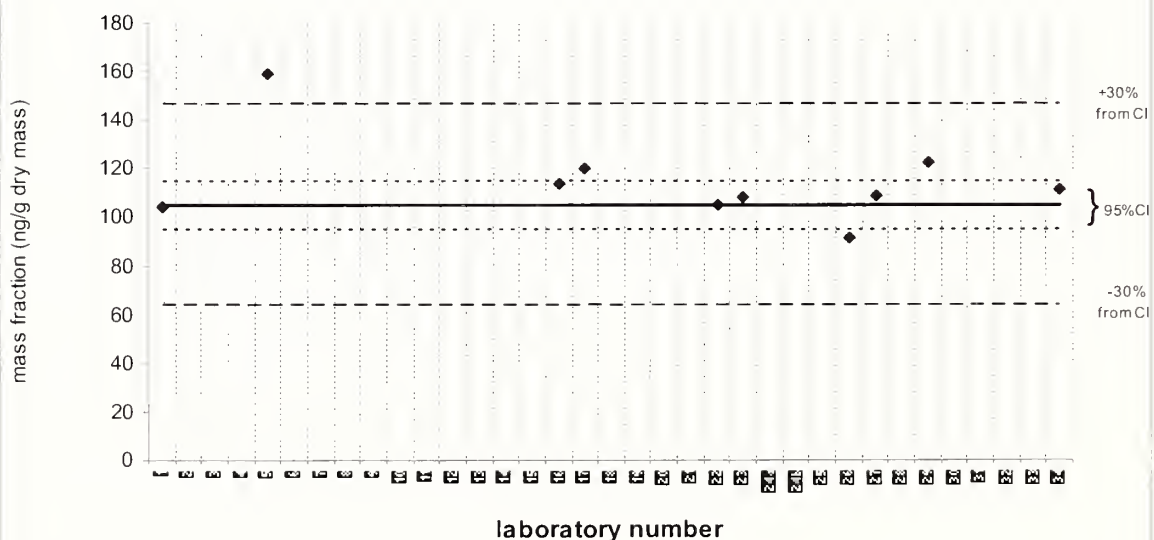


Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

chrysene/triphenylene**SRM 1974b**

Target Value = 105 ng/g dry mass ; 95% CI 10 ng/g dry mass: Median value = 110 ng/g dry mass

Reported Results: 10 Quantitative Results: 10

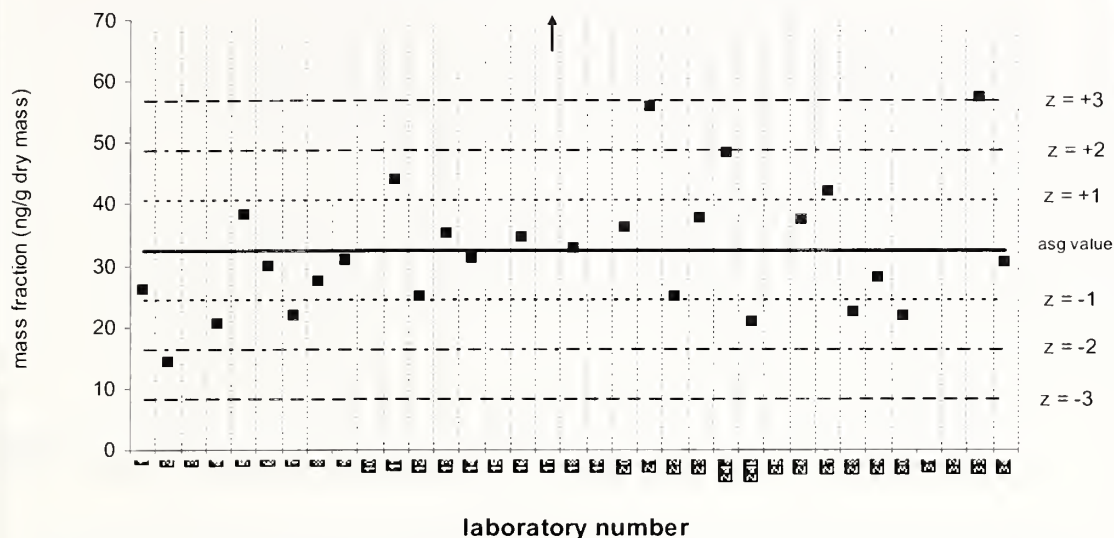


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

benzo[b]fluoranthene**QA10TIS01**Assigned value = 32.4 ng/g dry mass $s = 10.5$ ng/g dry mass 95% CI = 4.0 ng/g dry mass

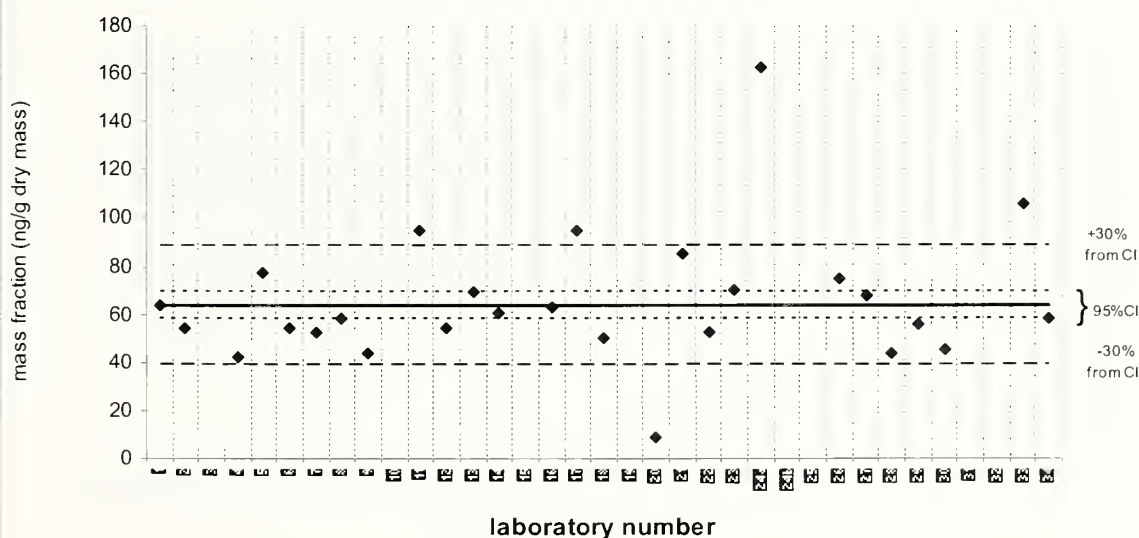
Median value = 30.9 ng/g dry mass

Reported Results: 33 Quantitative Results: 28

**laboratory number**Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)**benzo[b]fluoranthene****SRM 1974b**

Certified Value = 63.8 ng/g dry mass ; 95% CI 5.8 ng/g dry mass: Median value = 58.5 ng/g dry mass

Reported Results: 31 Quantitative Results: 27

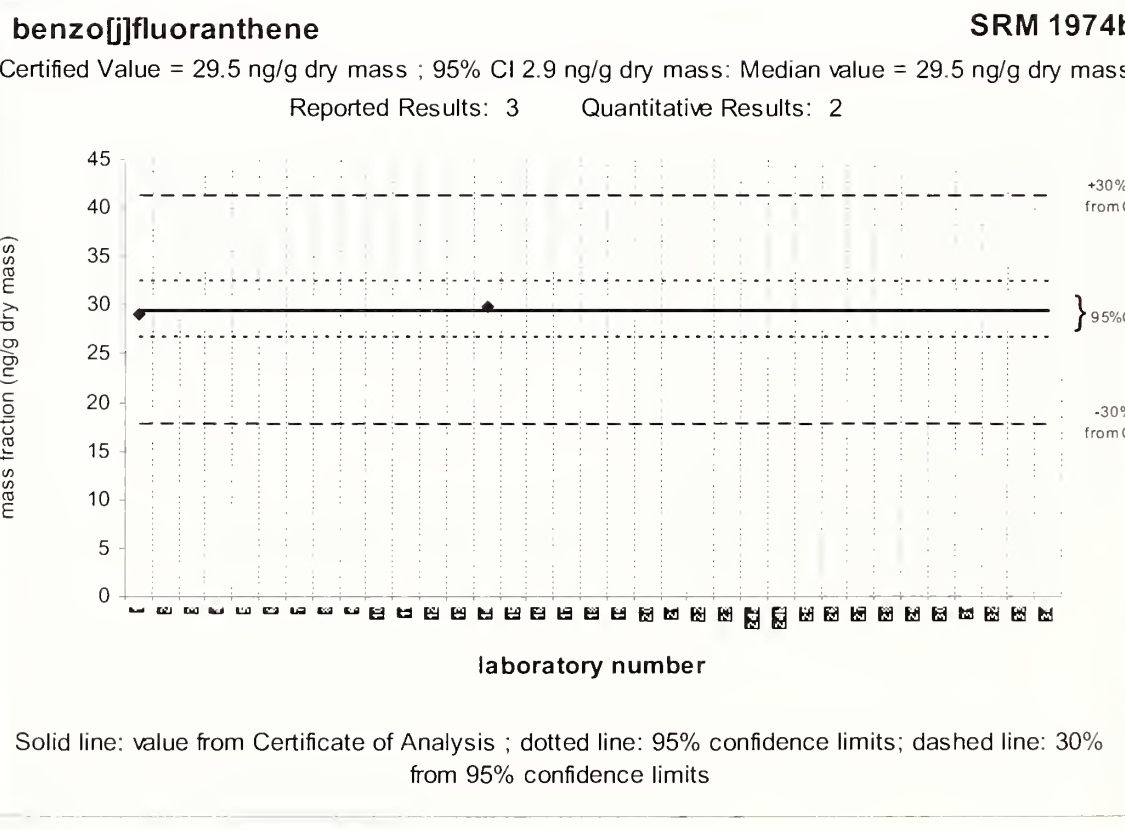
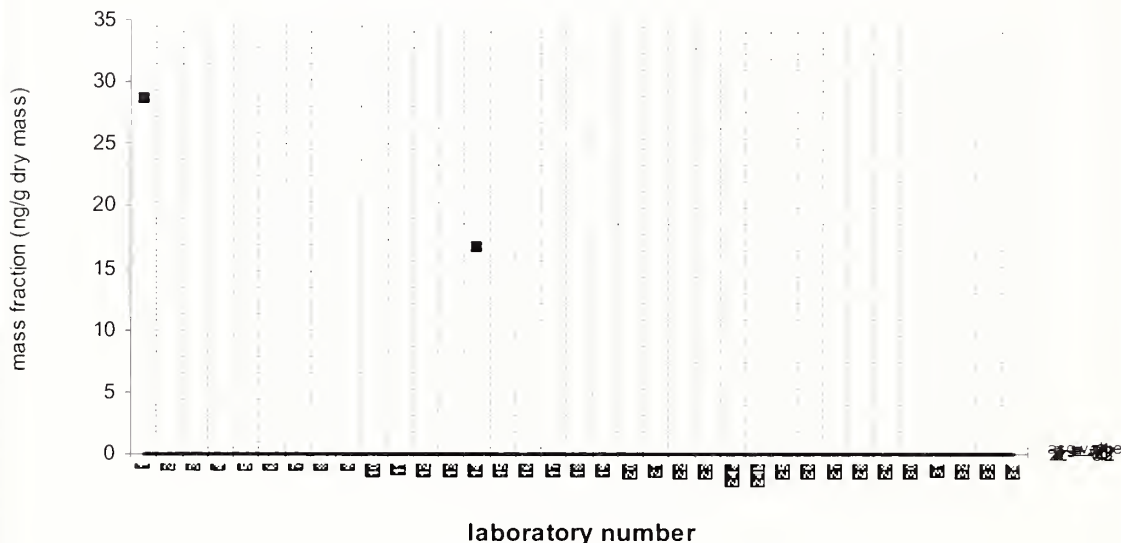
**laboratory number**

Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

benzo[*j*]fluoranthene**QA10TIS01**

Assigned value = No Target ng/g (dry mass) Median value = 22.6 ng/g dry mass

Reported Results: 2 Quantitative Results: 2

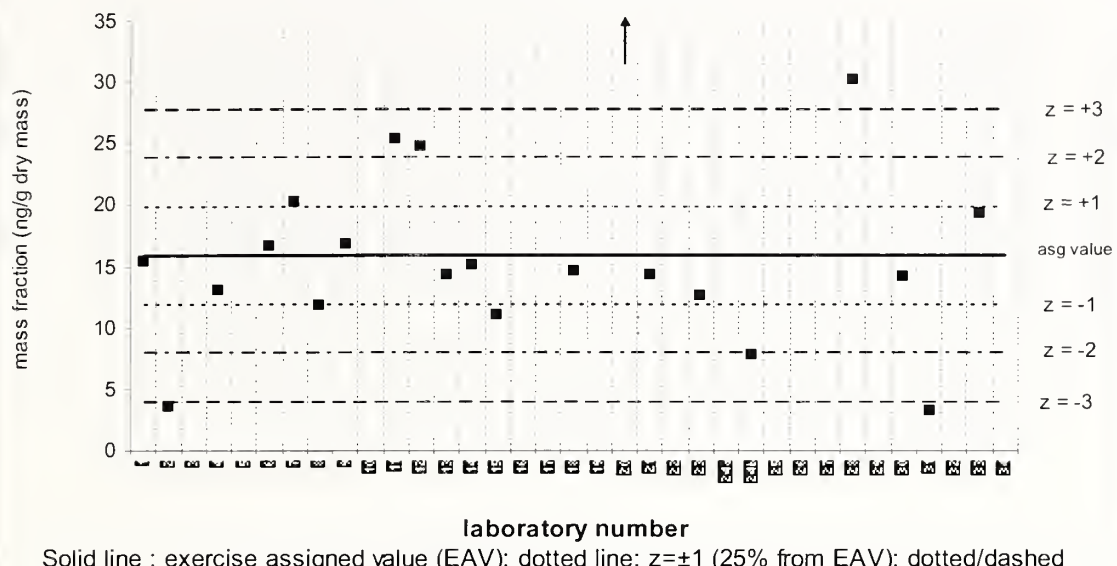


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

benzo[k]fluoranthene**QA10TIS01**

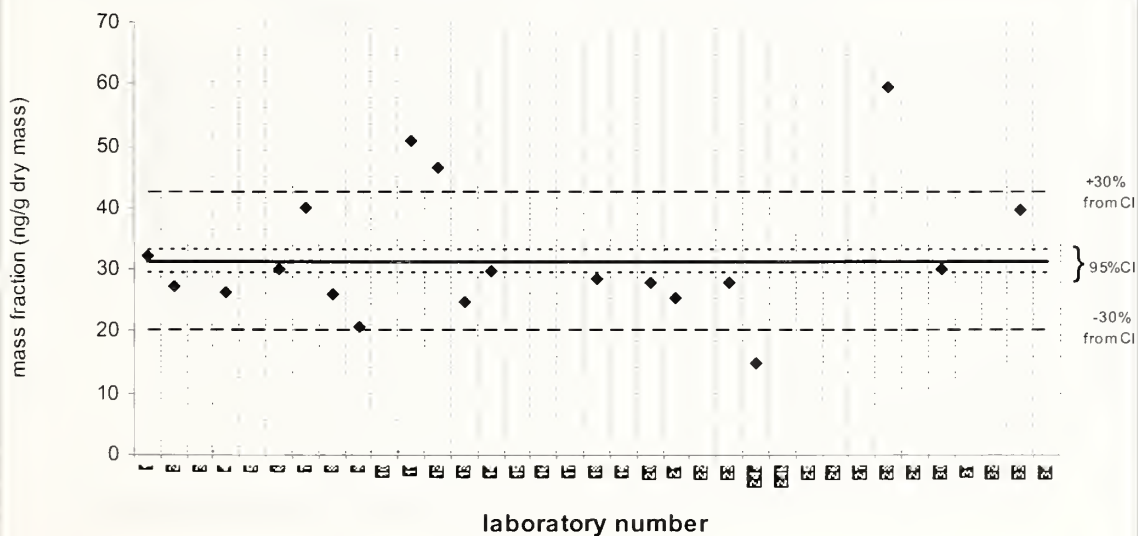
Assigned value = 15.8 ng/g dry mass s = 6.2 ng/g dry mass 95% CI = 2.8 ng/g dry mass Median value = 14.7 ng/g dry mass

Reported Results: 25 Quantitative Results: 21

**benzo[k]fluoranthene****SRM 1974b**

Certified Value = 31.2 ng/g dry mass ; 95% CI 1.8 ng/g dry mass: Median value = 28.4 ng/g dry mass

Reported Results: 23 Quantitative Results: 19

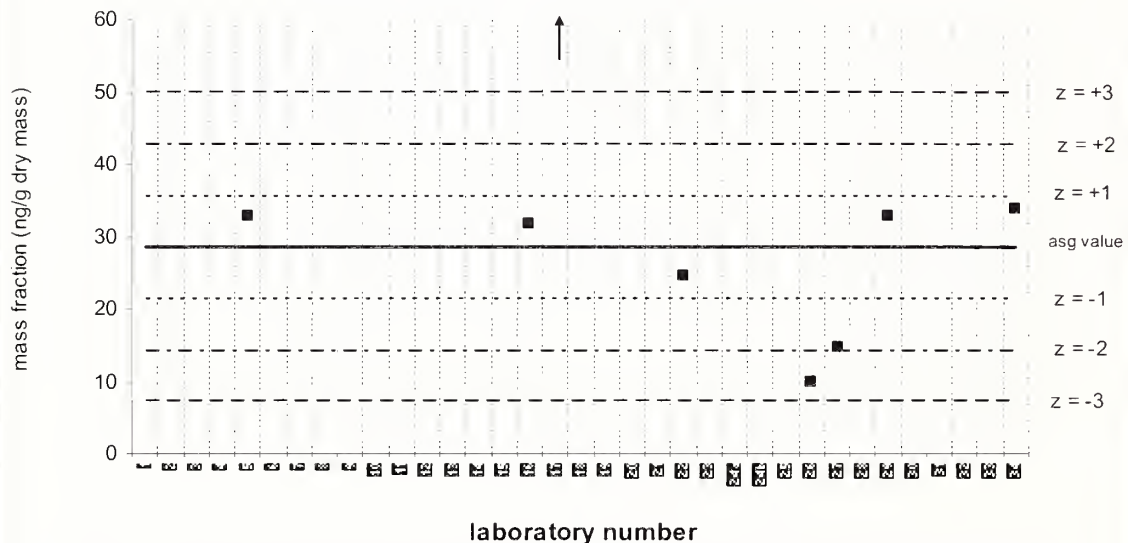


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

benzo[j+k]fluoranthene**QA10TIS01**

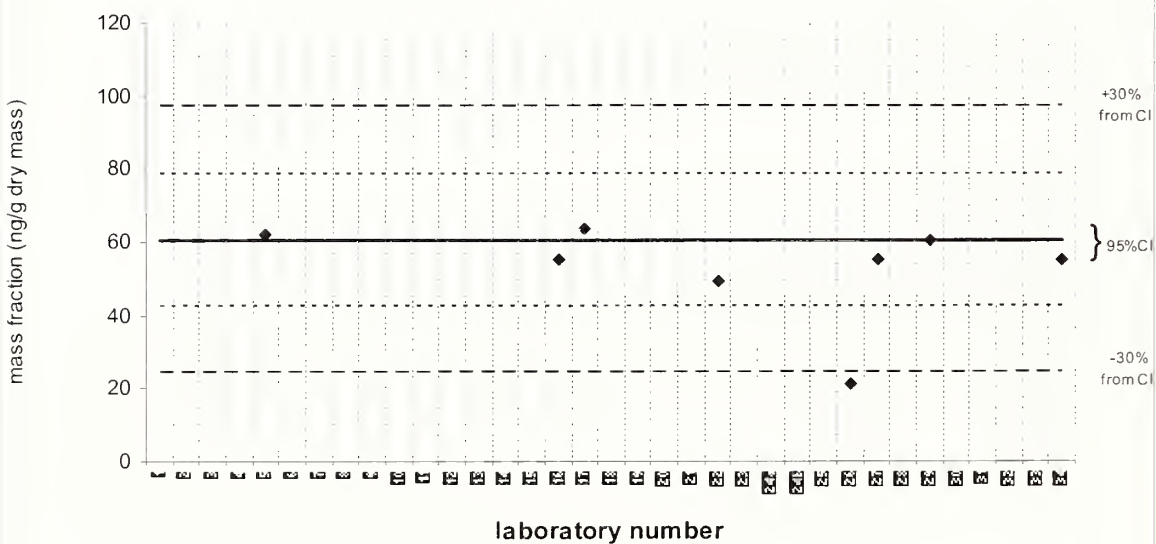
Assigned value = 28.4 ng/g dry mass $s = 7.5$ ng/g dry mass 95% CI = 6.0 ng/g dry mass Median value = 32.2 ng/g dry mass

Reported Results: 8 Quantitative Results: 8

**benzo[j+k]fluoranthene****SRM 1974b**

Target Value = 60.7 ng/g dry mass ; 95% CI 18.0 ng/g dry mass: Median value = 55.4 ng/g dry mass

Reported Results: 8 Quantitative Results: 8



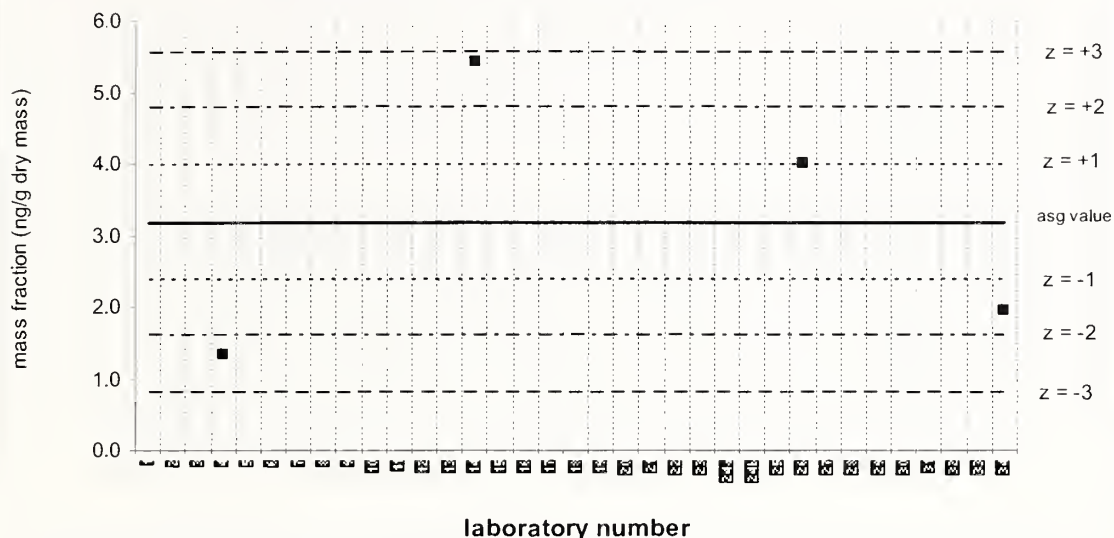
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

benzo[a]fluoranthene**QA10TIS01**

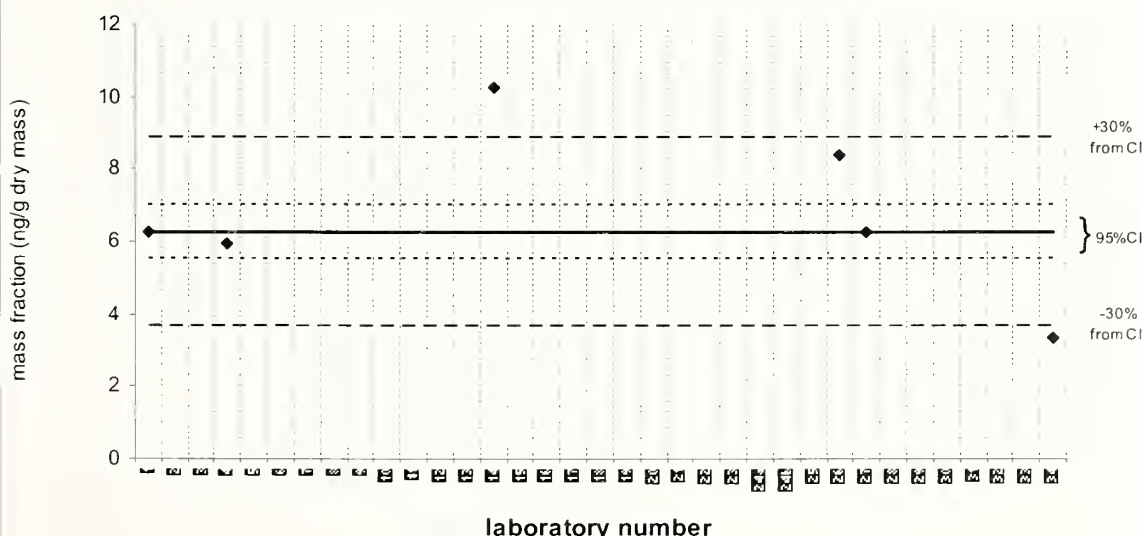
Assigned value = 3.17 ng/g dry mass s = 1.88 ng/g dry mass 95% CI = 1.84 ng/g dry mass

Median value = 2.97 ng/g dry mass

Reported Results: 7 Quantitative Results: 4

**benzo[a]fluoranthene****SRM 1974b**

Certified Value = 6.26 ng/g dry mass ; 95% CI 0.73 ng/g dry mass: Median value = 6.28 ng/g dry

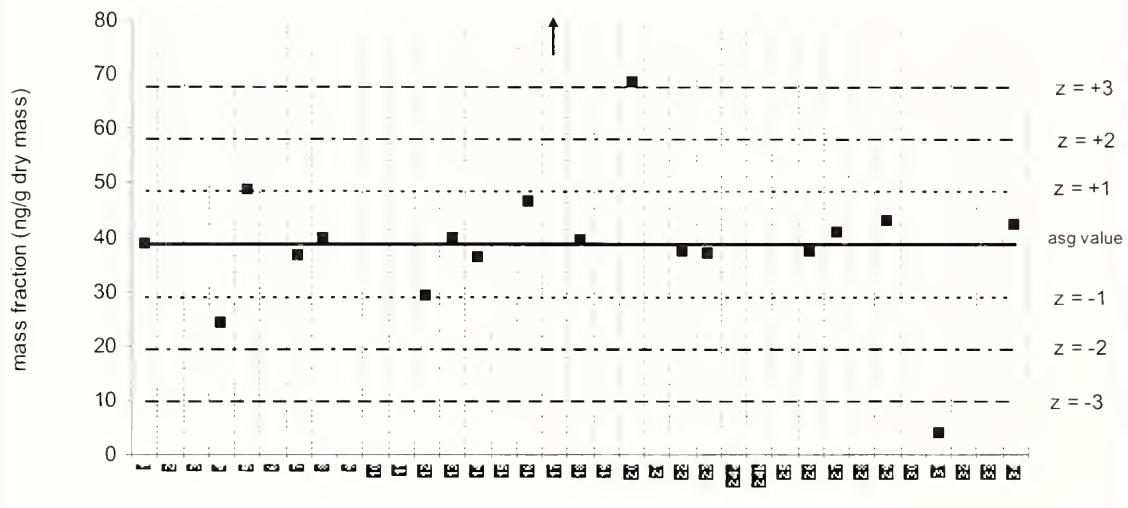
mass
Reported Results: 7 Quantitative Results: 6

Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

benzo[e]pyrene**QA10TIS01**

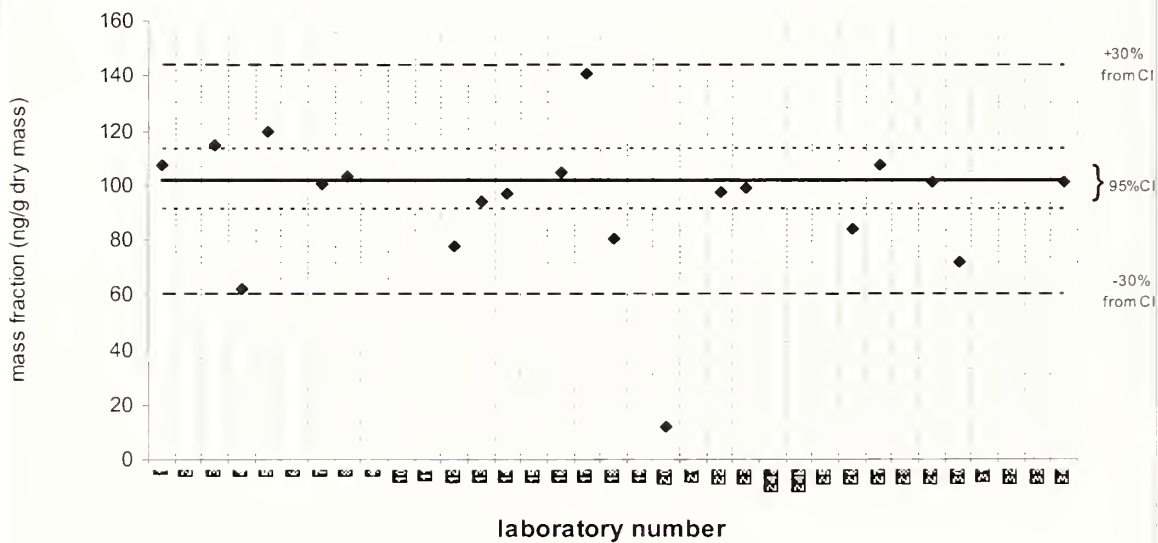
Assigned value = 38.6 ng/g dry mass s = 5.9 ng/g dry mass 95% CI = 2.9 ng/g dry mass Median value = 39.4 ng/g dry mass

Reported Results: 25 Quantitative Results: 19

**benzo[e]pyrene****SRM 1974b**

Certified Value = 102 ng/g dry mass ; 95% CI 11 ng/g dry mass: Median value = 100 ng/g dry mass

Reported Results: 23 Quantitative Results: 20

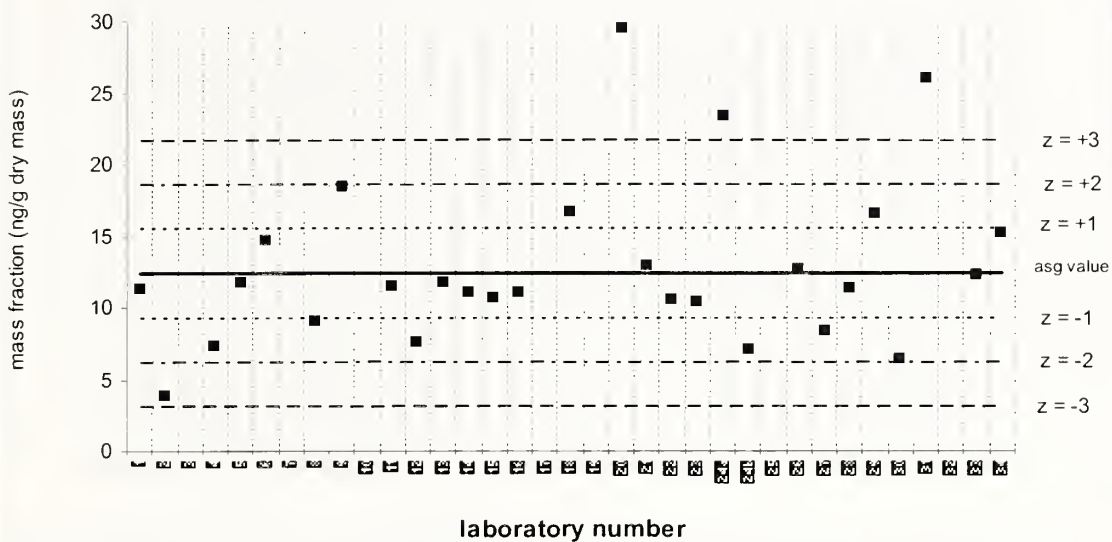


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

benzo[a]pyrene**QA10TIS01**

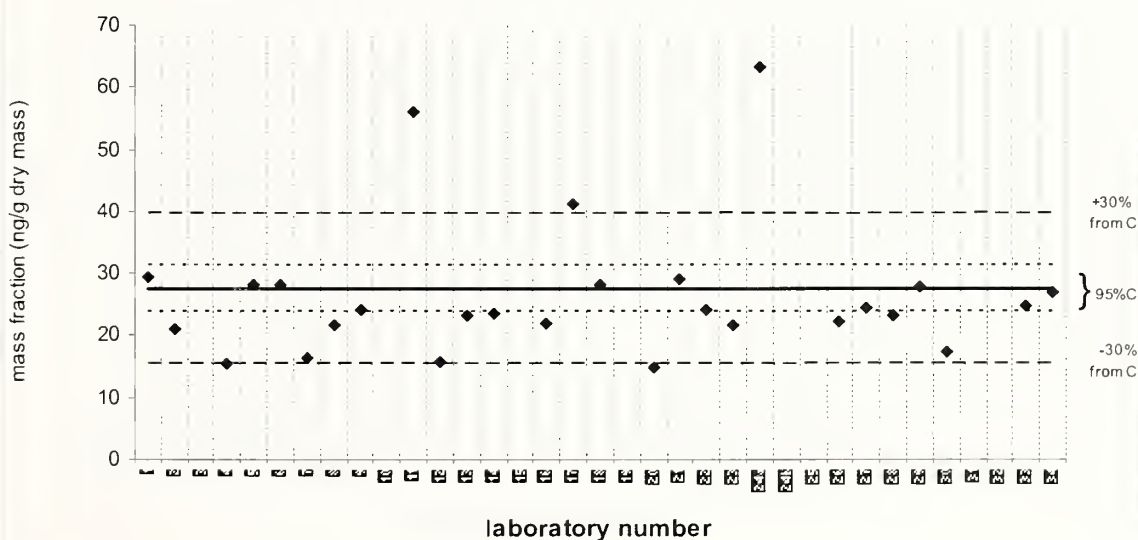
Assigned value = 12.3 ng/g dry mass $s = 5.3$ ng/g dry mass 95% CI = 2.0 ng/g dry mass Median value = 11.4 ng/g dry mass

Reported Results: 35 Quantitative Results: 28

**benzo[a]pyrene****SRM 1974b**

Certified Value = 27.6 ng/g dry mass ; 95% CI 3.8 ng/g dry mass: Median value = 24.2 ng/g dry mass

Reported Results: 31 Quantitative Results: 27



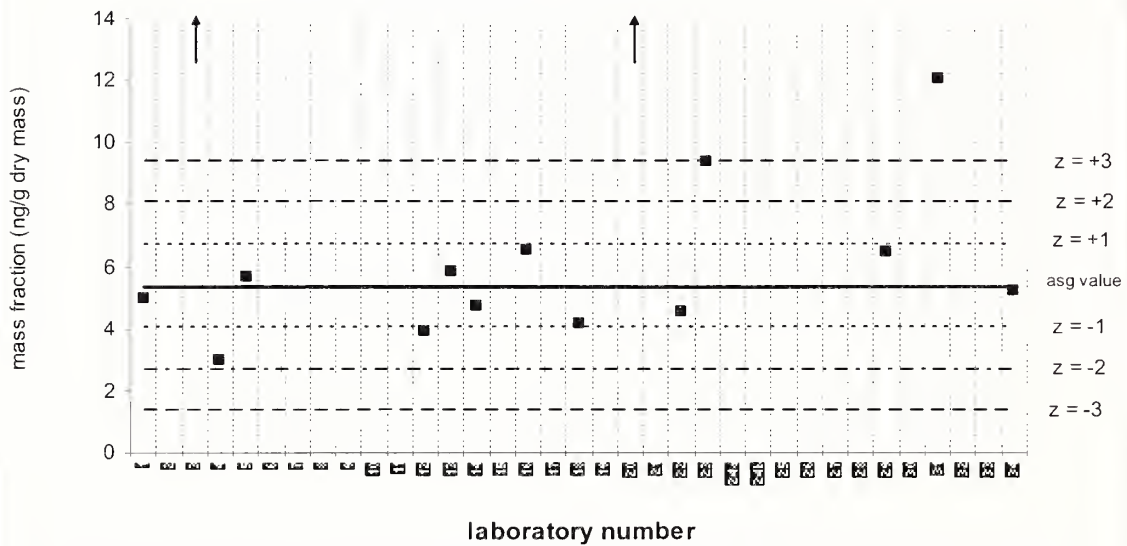
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

perylene

QA10TIS01

Assigned value = 5.38 ng/g dry mass s = 1.64 ng/g dry mass 95% CI = 0.93 ng/g dry mass
 Median value = 5.68 ng/g dry mass

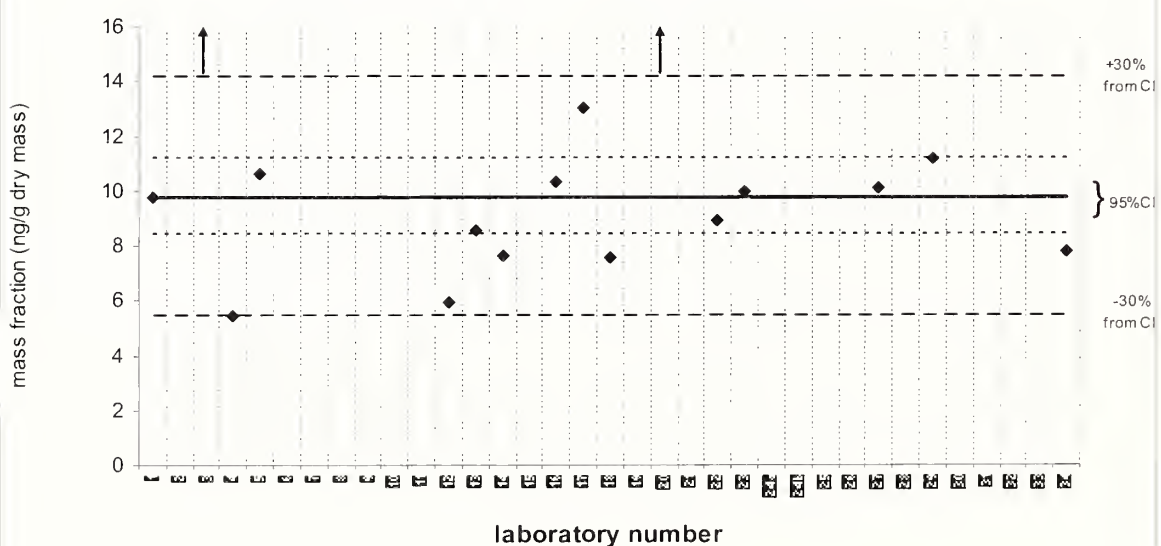
Reported Results: 25 Quantitative Results: 15

**perylene**

SRM 1974b

Certified Value = 9.8 ng/g dry mass ; 95% CI 1.4 ng/g dry mass: Median value = 9.85 ng/g dry mass

Reported Results: 22 Quantitative Results: 16



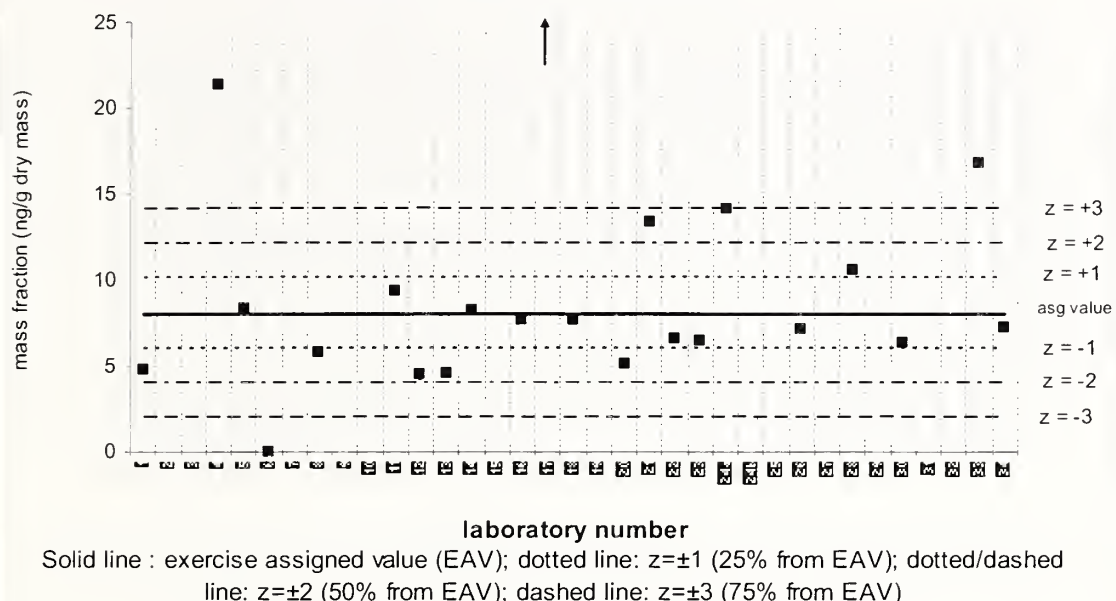
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

indeno[1,2,3-cd]pyrene**QA10TIS01**

Assigned value = 8.05 ng/g dry mass s = 4.00 ng/g dry mass 95% CI = 1.85 ng/g dry mass

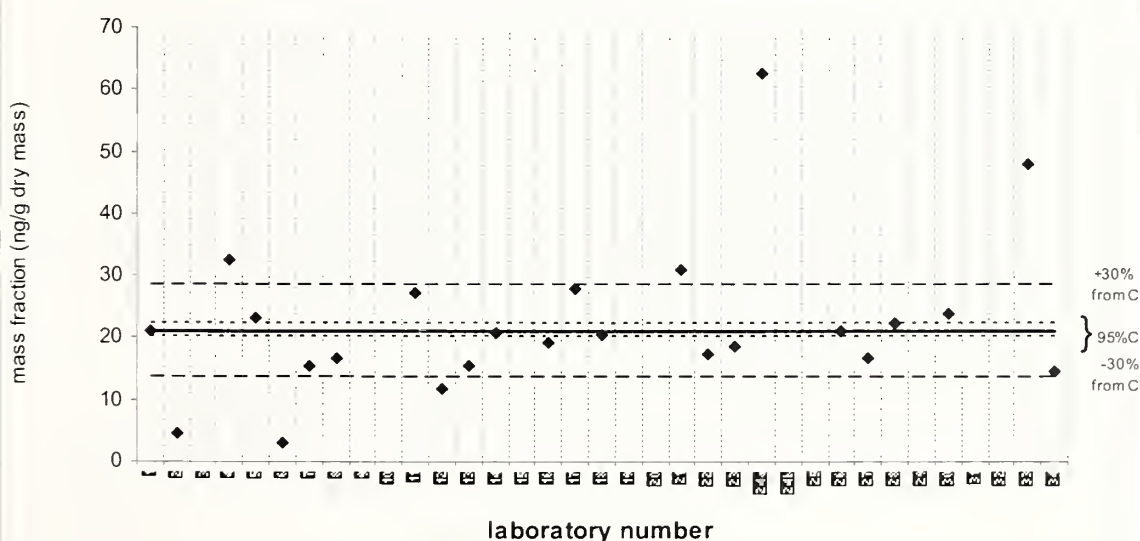
Median value = 7.45 ng/g dry mass

Reported Results: 32 Quantitative Results: 22

**indeno[1,2,3-cd]pyrene****SRM 1974b**

Certified Value = 21.1 ng/g dry mass ; 95% CI 1.1 ng/g dry mass: Median value = 20.6 ng/g dry mass

Reported Results: 30 Quantitative Results: 24

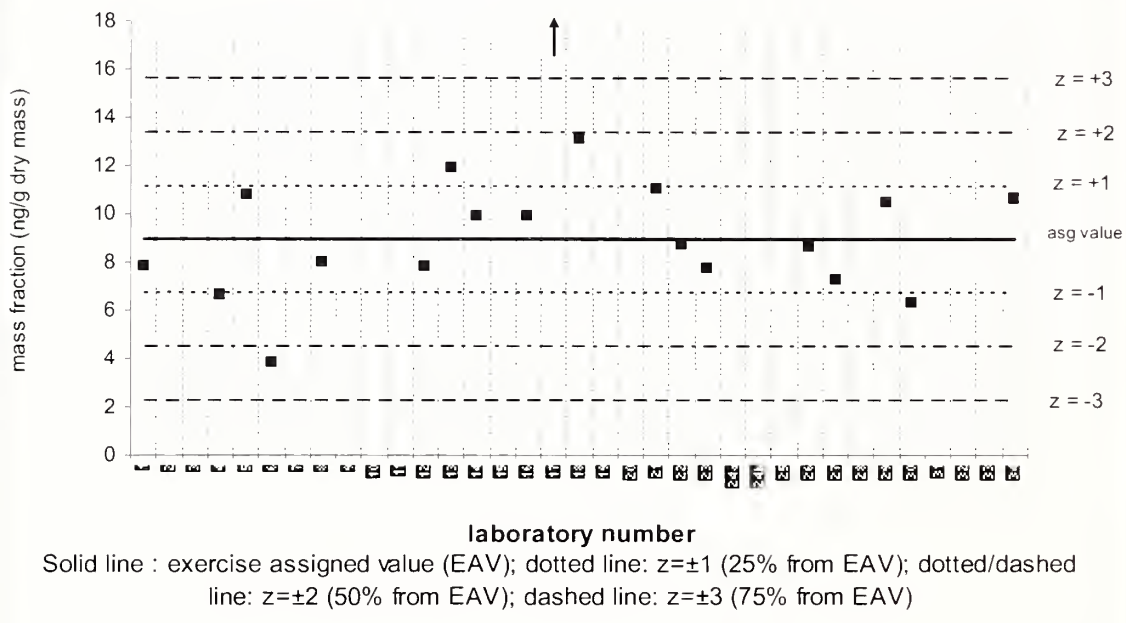


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

benzo[ghi]perylene**QA10TIS01**Assigned value = 8.92 ng/g dry mass $s = 2.25$ ng/g dry mass 95% CI = 1.04 ng/g dry mass

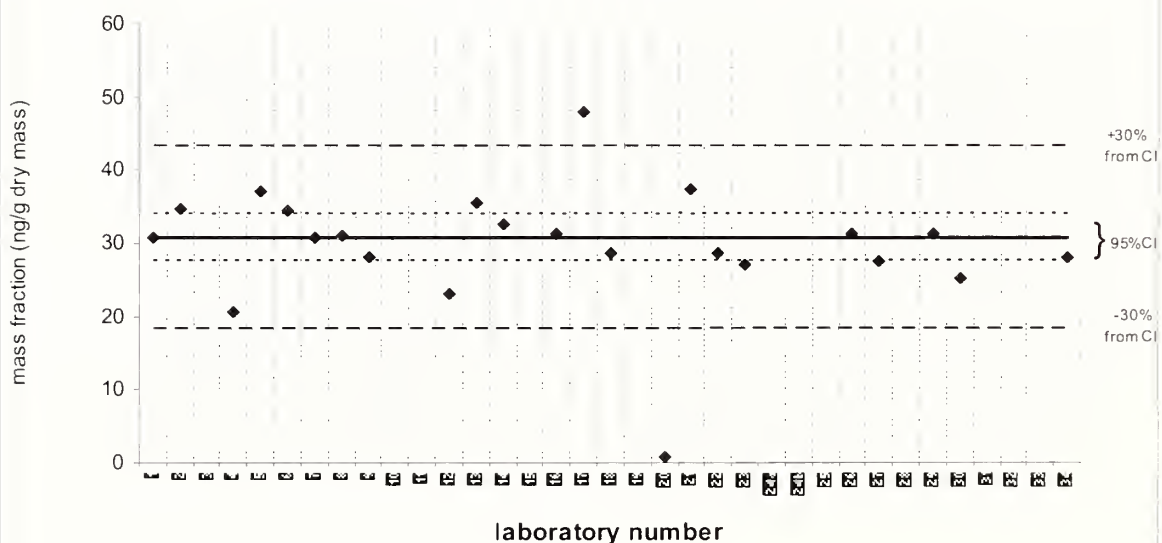
Median value = 8.75 ng/g dry mass

Reported Results: 29 Quantitative Results: 19

**benzo[ghi]perylene****SRM 1974b**

Certified Value = 30.8 ng/g dry mass ; 95% CI 3.3 ng/g dry mass: Median value = 30.9 ng/g dry mass

Reported Results: 27 Quantitative Results: 23



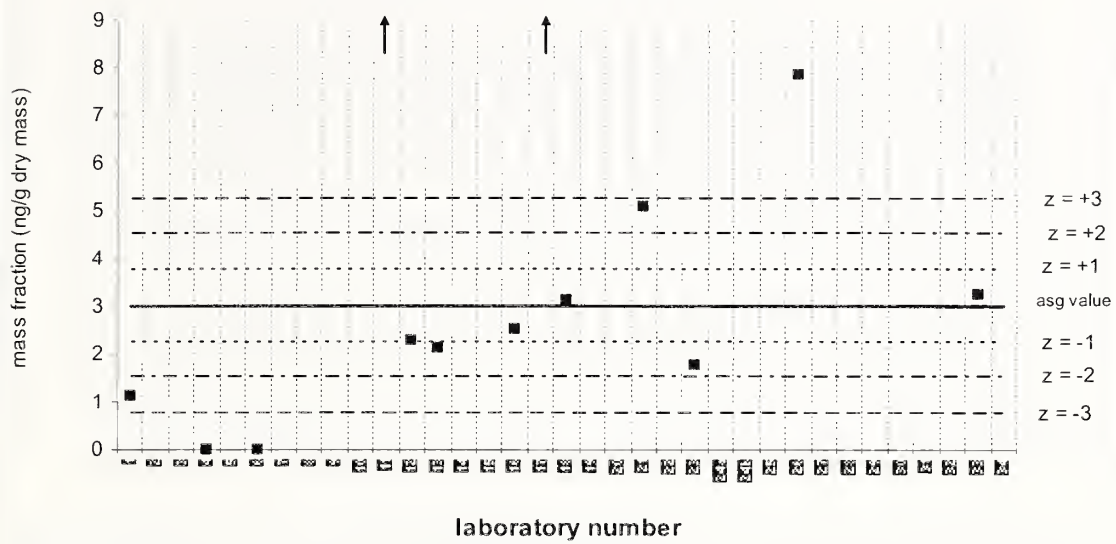
dibenz[a,h]anthracene

QA10TIS01

Assigned value = 3.00 ng/g dry mass s = 2.07 ng/g dry mass 95% CI = 1.43 ng/g dry mass

Median value = 2.54 ng/g dry mass

Reported Results: 28 Quantitative Results: 13

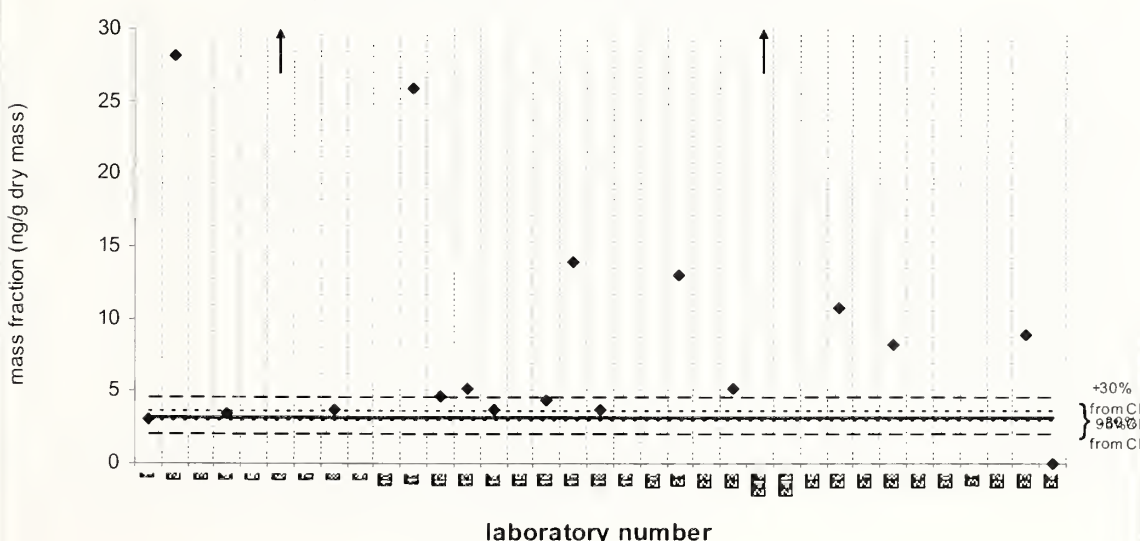


Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

dibenz[a,h]anthracene

SRM 1974b

Certified Value = 3.23 ng/g dry mass ; 95% CI 0.31 ng/g dry mass: Median value = 6.75 ng/g dry

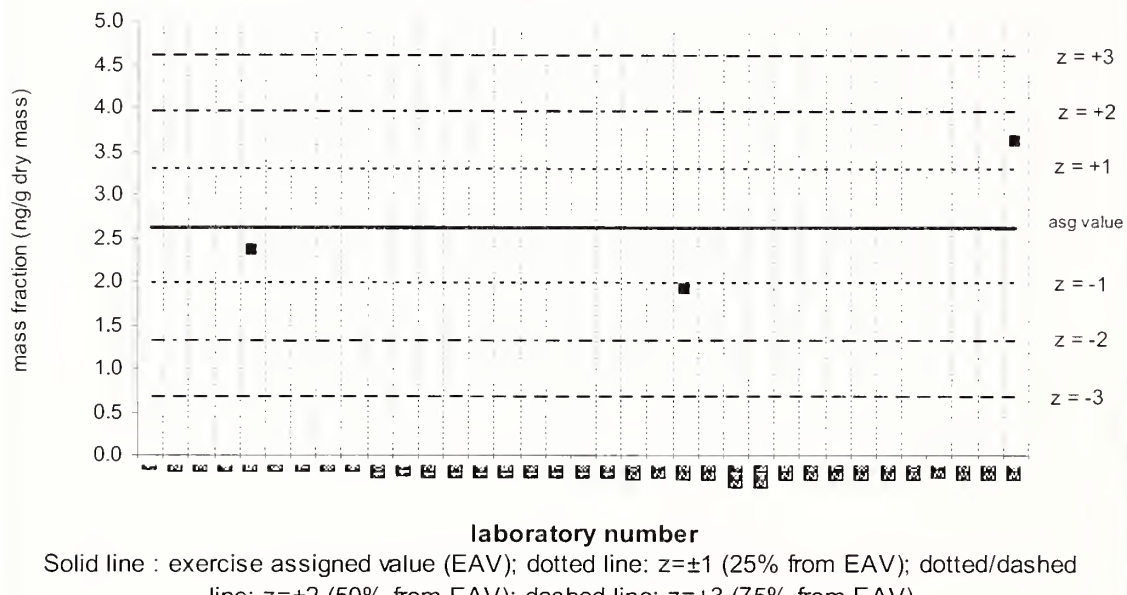
mass
Reported Results: 27 Quantitative Results: 18

Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

dibenz[a,h+a,c]anthracene**QA10TIS01**Assigned value = 2.63 ng/g dry mass $s = 0.89$ ng/g dry mass 95% CI = 1.01 ng/g dry mass

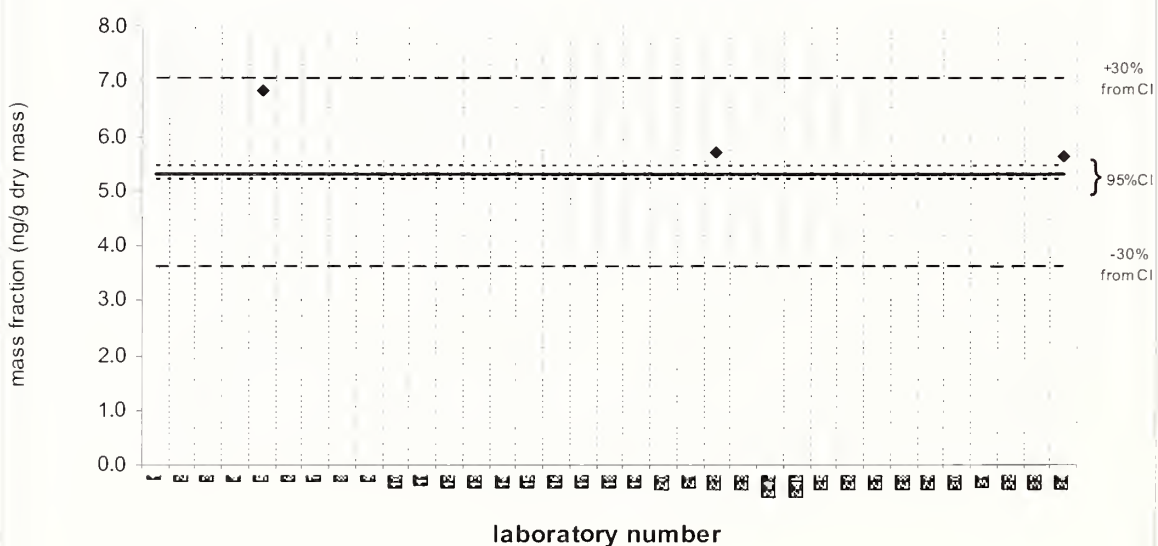
Median value = 2.35 ng/g dry mass

Reported Results: 3 Quantitative Results: 3

**dibenz[a,h+a,c]anthracene****SRM 1974b**

Target Value = 5.32 ng/g dry mass ; 95% CI 0.13 ng/g dry mass: Median value = 5.70 ng/g dry mass

Reported Results: 3 Quantitative Results: 3

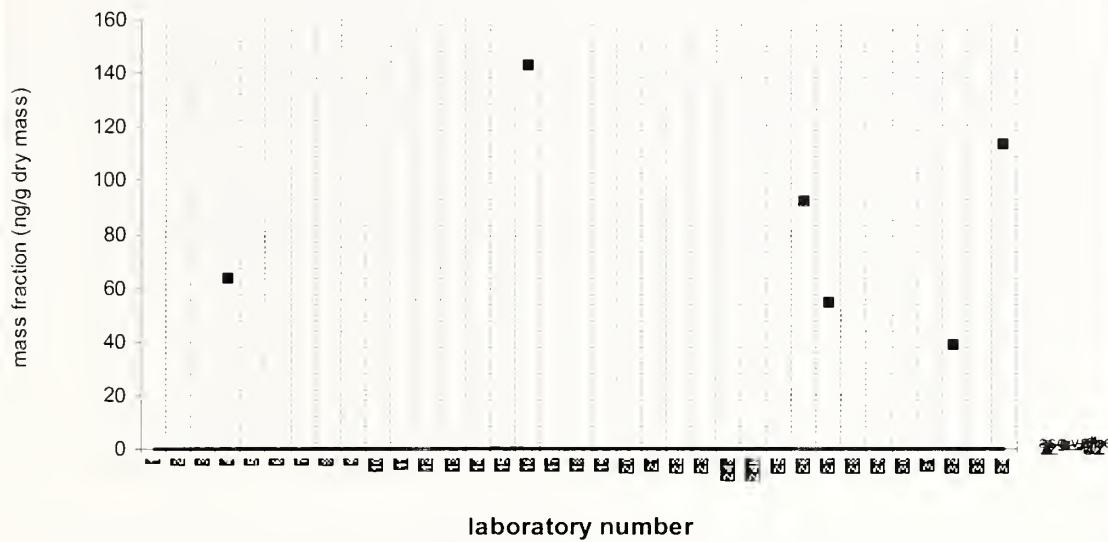


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

cis/trans-decalin**QA10TIS01**

Assigned value = No Target ng/g (dry mass) Median value = 77.6 ng/g dry mass

Reported Results: 8 Quantitative Results: 6



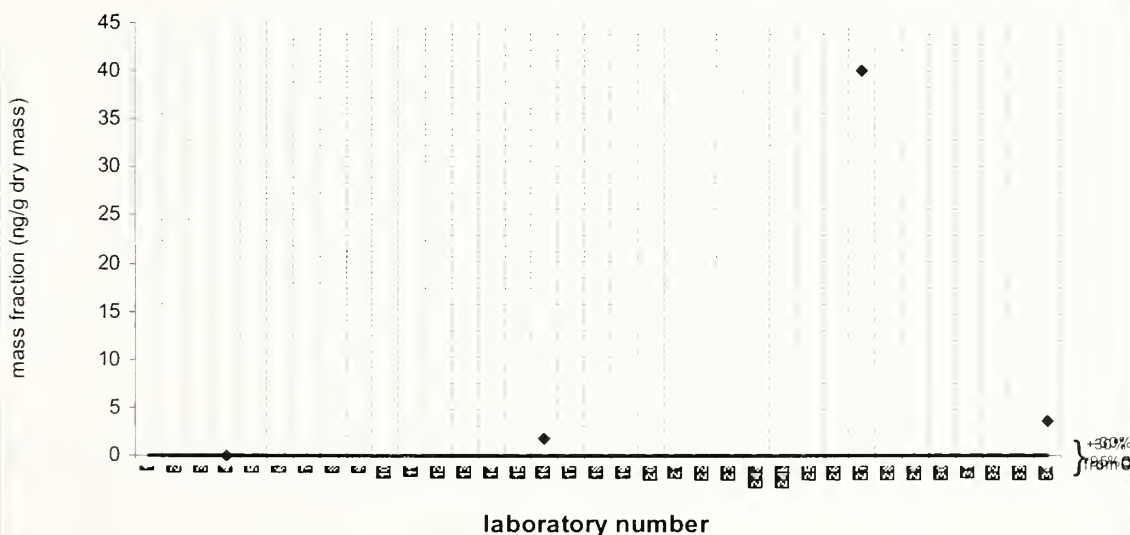
laboratory number

Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

cis/trans-decalin**SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 2.72 ng/g dry mass

Reported Results: 7 Quantitative Results: 4



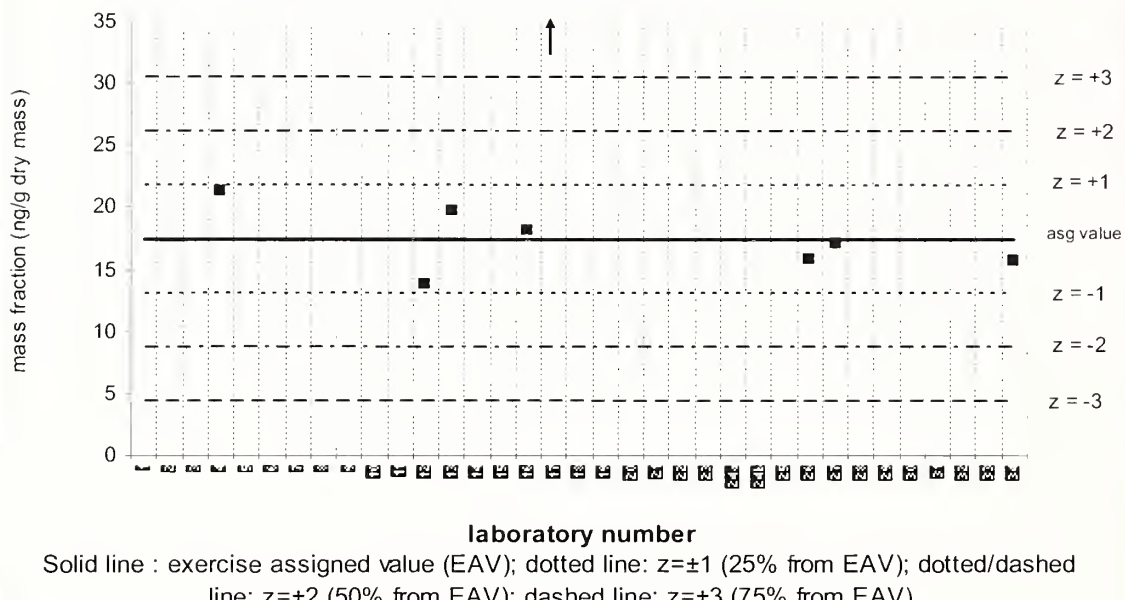
laboratory number

Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

dibenzofuran**QA10TIS01**

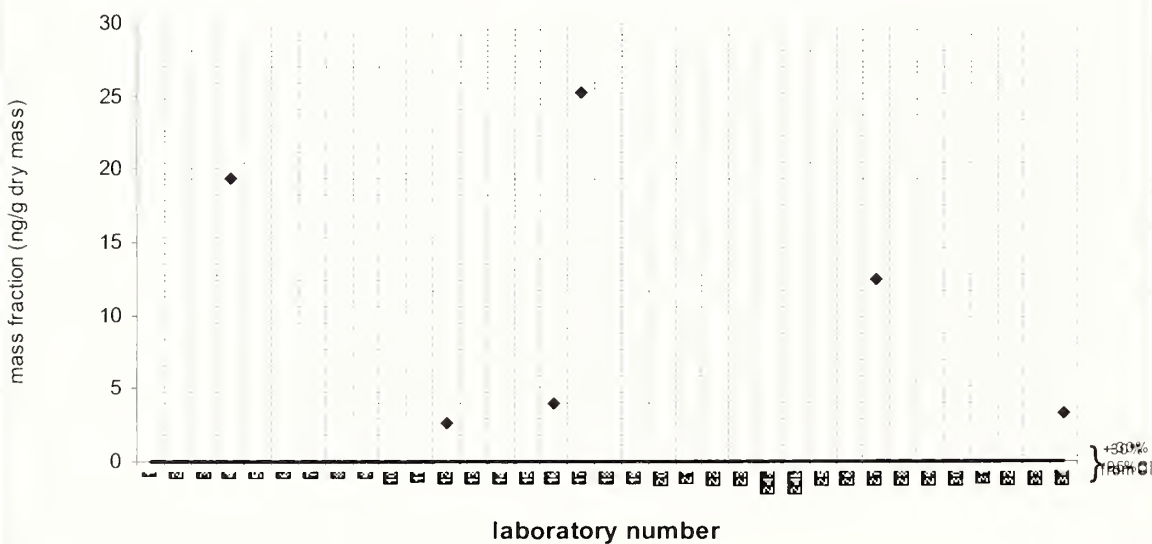
Assigned value = 17.4 ng/g dry mass $s = 2.6$ ng/g dry mass 95% CI = 1.9 ng/g dry mass Median value = 17.7 ng/g dry mass

Reported Results: 15 Quantitative Results: 8

**dibenzofuran****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 8.20 ng/g dry mass

Reported Results: 14 Quantitative Results: 6

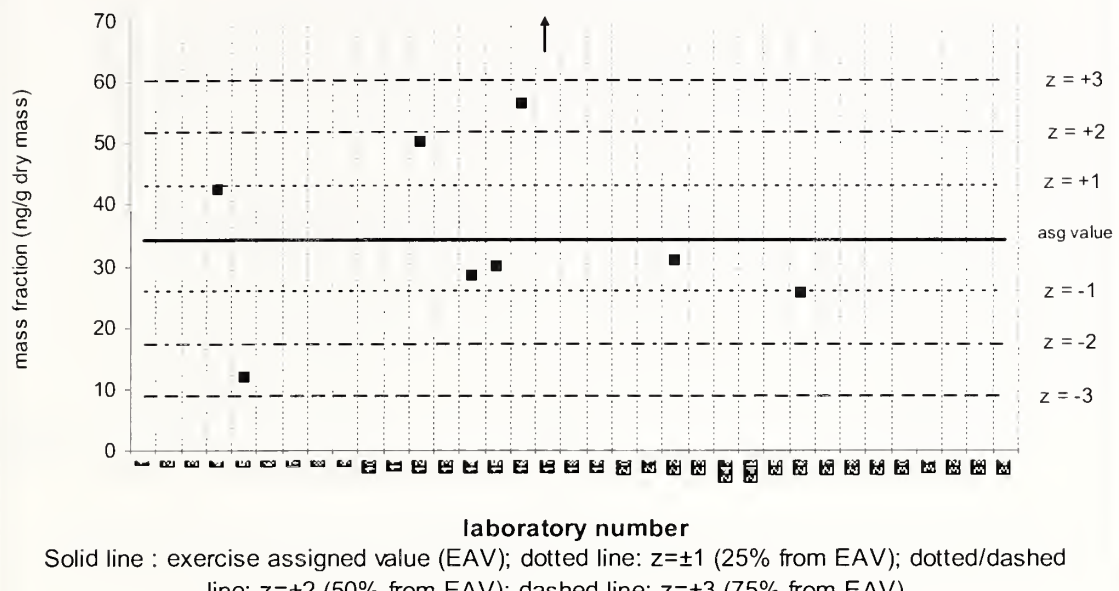


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

retene**QA10TIS01**Assigned value = 34.4 ng/g dry mass $s = 14.4$ ng/g dry mass 95% CI = 10.0 ng/g dry mass

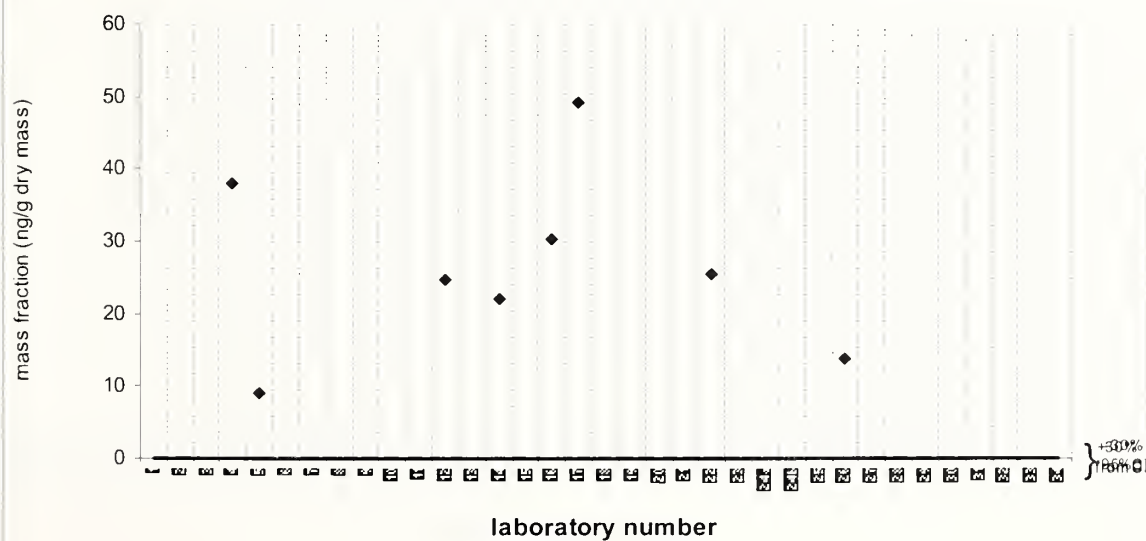
Median value = 30.7 ng/g dry mass

Reported Results: 12 Quantitative Results: 9

**retene****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 25.2 ng/g dry mass

Reported Results: 10 Quantitative Results: 8

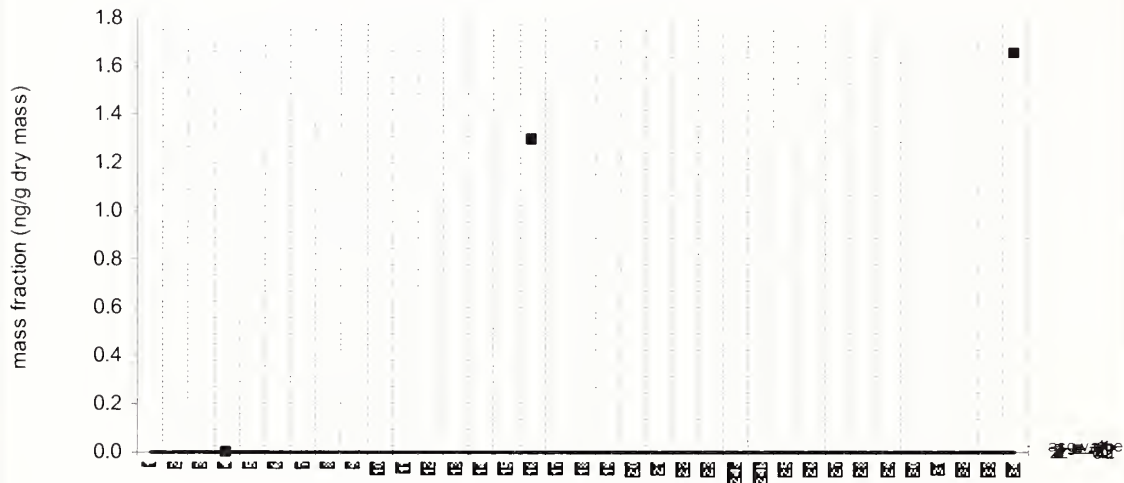


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

benzothiophene**QA10TIS01**

Assigned value = No Target ng/g (dry mass) Median value = 1.30 ng/g dry mass

Reported Results: 10 Quantitative Results: 3

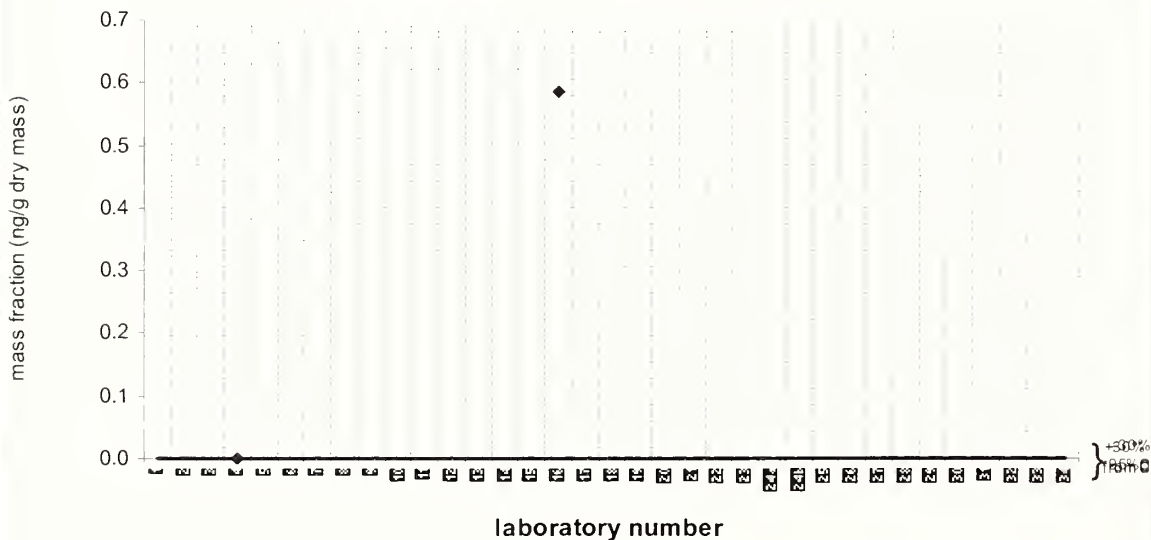
**laboratory number**

Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

benzothiophene**SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 0.29 ng/g dry mass

Reported Results: 8 Quantitative Results: 2

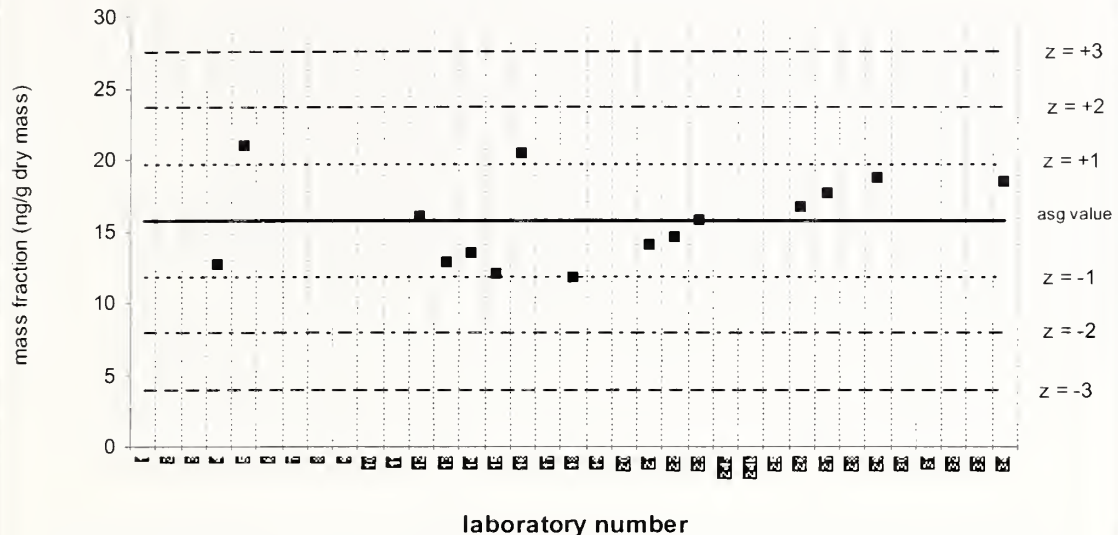
**laboratory number**

Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

dibenzothiophene**QA10TIS01**

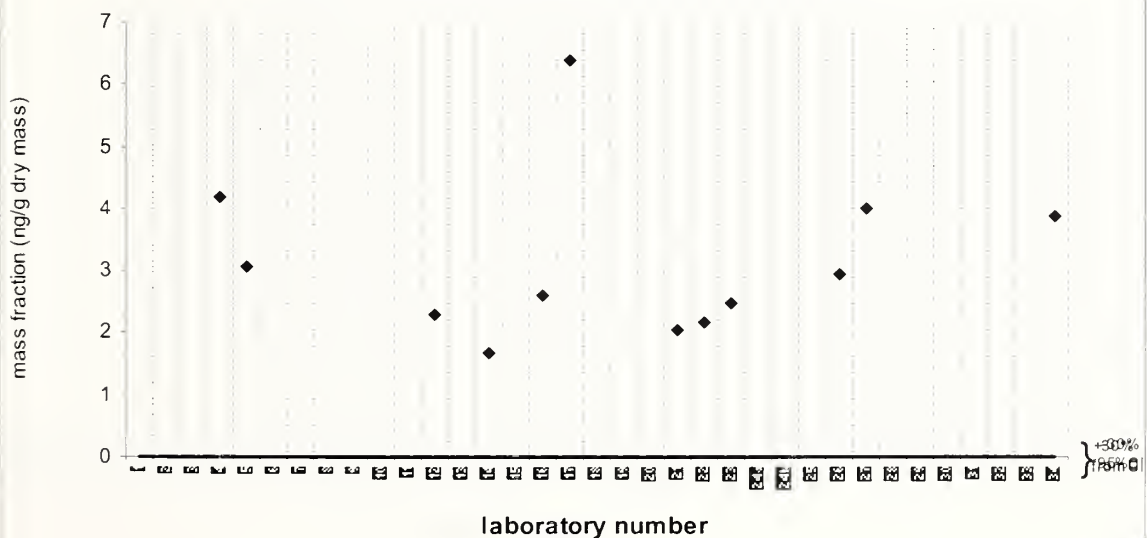
Assigned value = 15.7 ng/g dry mass s = 3.0 ng/g dry mass 95% CI = 1.5 ng/g dry mass Median value = 15.8 ng/g dry mass

Reported Results: 26 Quantitative Results: 15

**dibenzothiophene****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 2.76 ng/g dry mass

Reported Results: 21 Quantitative Results: 12

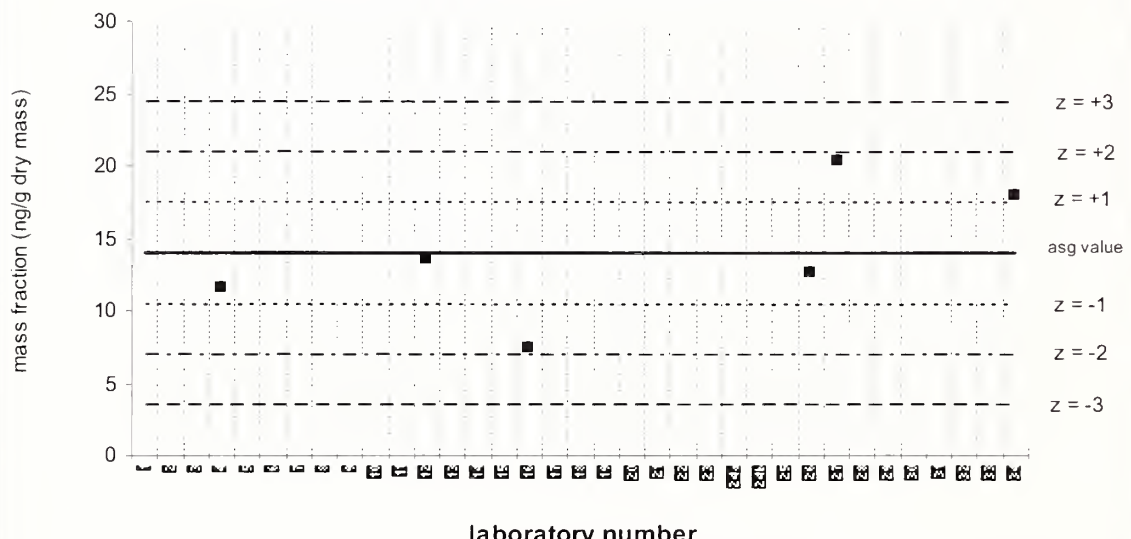


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

naphthobenzothiophene**QA10TIS01**

Assigned value = 14.0 ng/g dry mass s = 4.6 ng/g dry mass 95% CI = 3.7 ng/g dry mass Median value = 13.1 ng/g dry mass

Reported Results: 8 Quantitative Results: 6

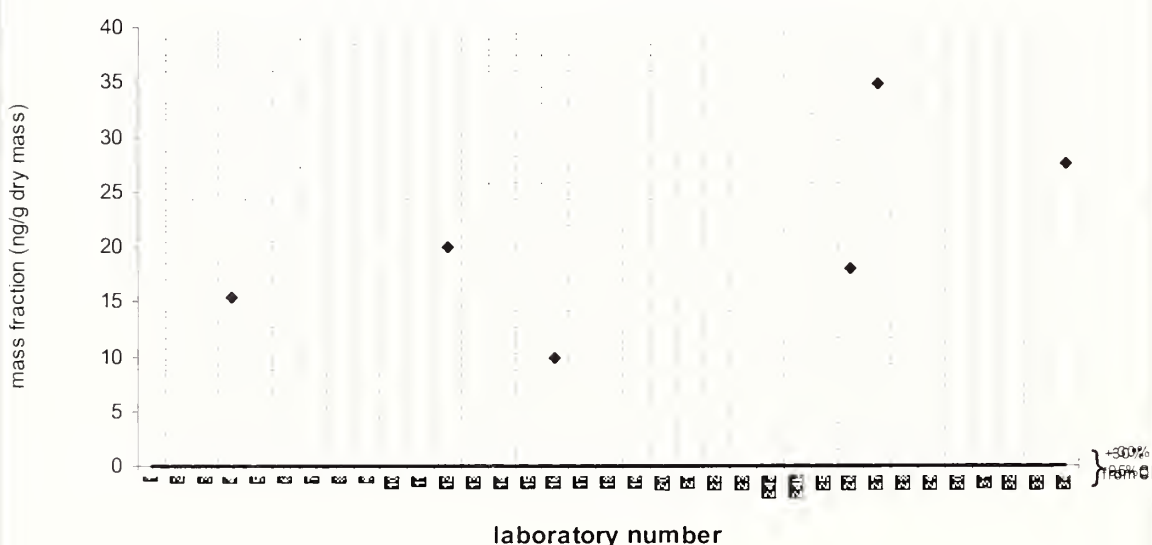
**laboratory number**

Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

naphthobenzothiophene**SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 19.0 ng/g dry mass

Reported Results: 7 Quantitative Results: 6

**laboratory number**

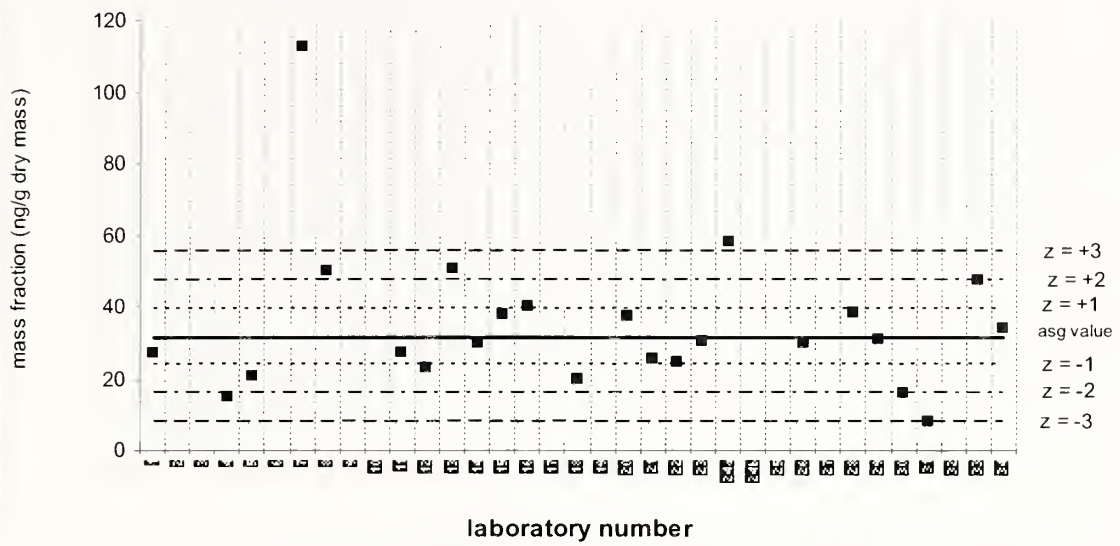
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

1-methylnaphthalene**QA10TIS01**

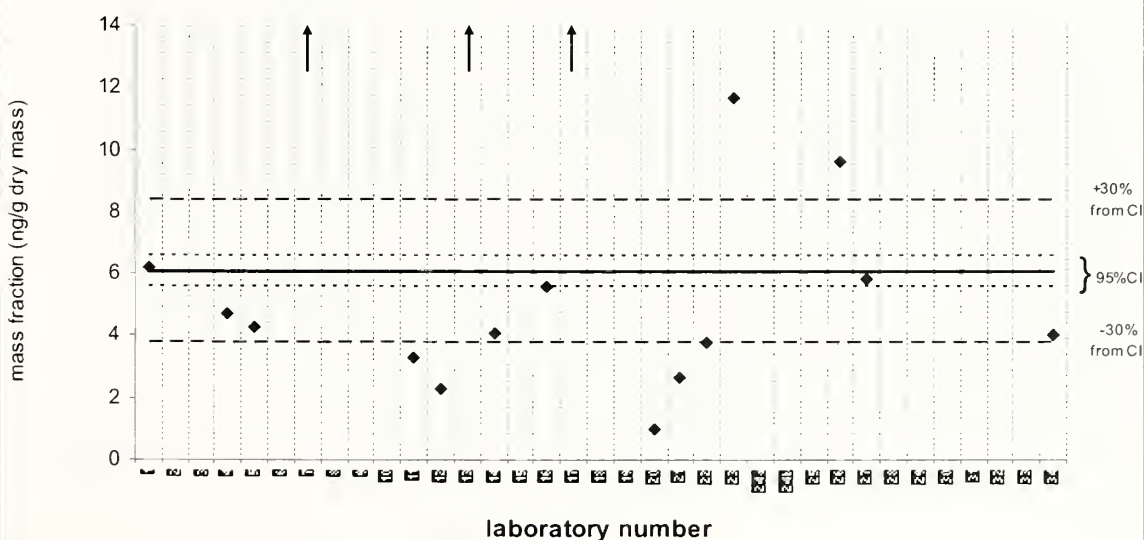
Assigned value = 31.7 ng/g dry mass s = 11.1 ng/g dry mass 95% CI = 4.8 ng/g dry mass

Median value = 30.1 ng/g dry mass

Reported Results: 31 Quantitative Results: 24

**1-methylnaphthalene****SRM 1974b**

Reference Value = 6.06 ng/g dry mass ; 95% CI 0.49 ng/g dry mass: Median value = 4.73 ng/g dry

mass
Reported Results: 22 Quantitative Results: 17

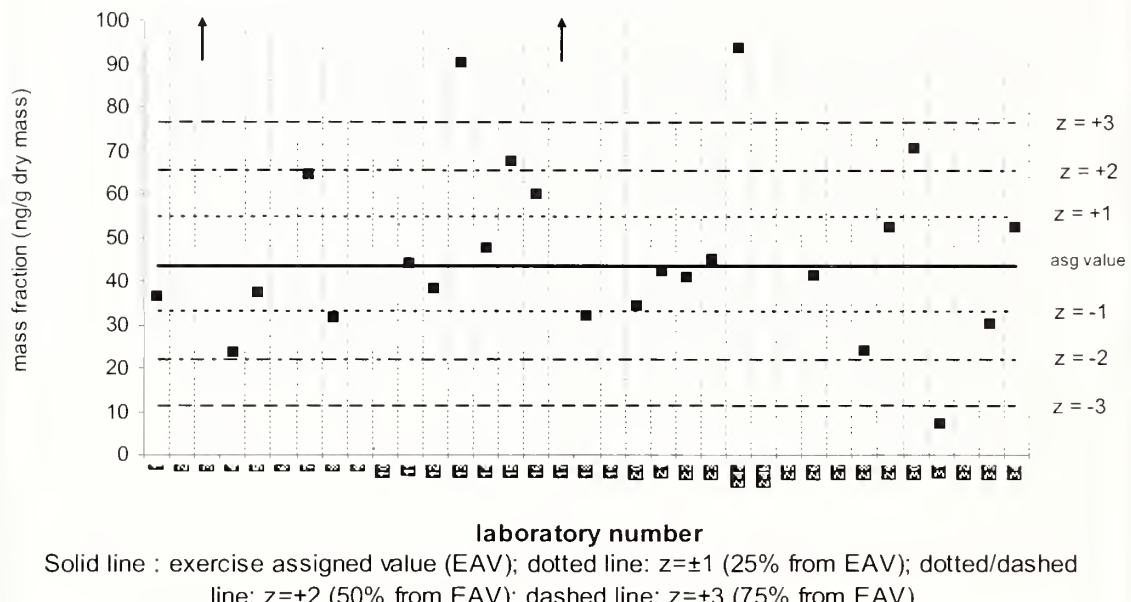
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

2-methylnaphthalene**QA10TIS01**

Assigned value = 43.6 ng/g dry mass s = 15.3 ng/g dry mass 95% CI = 6.6 ng/g dry mass

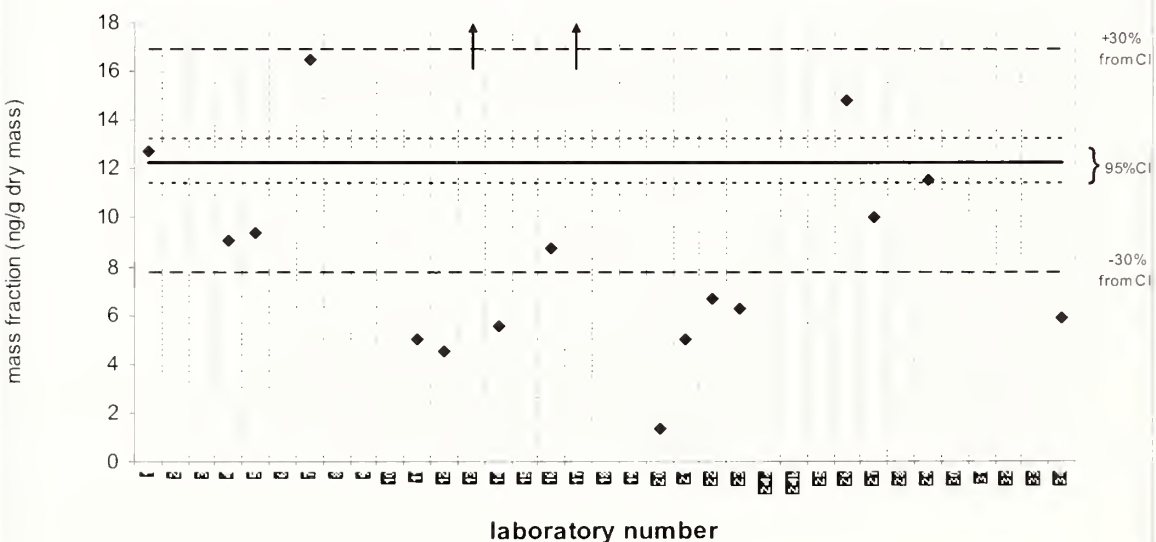
Median value = 43.1 ng/g dry mass

Reported Results: 31 Quantitative Results: 26

**2-methylnaphthalene****SRM 1974b**

Reference Value = 12.3 ng/g dry mass ; 95% CI 0.9 ng/g dry mass: Median value = 8.93 ng/g dry

Reported Results: 22 Quantitative Results: 18



Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

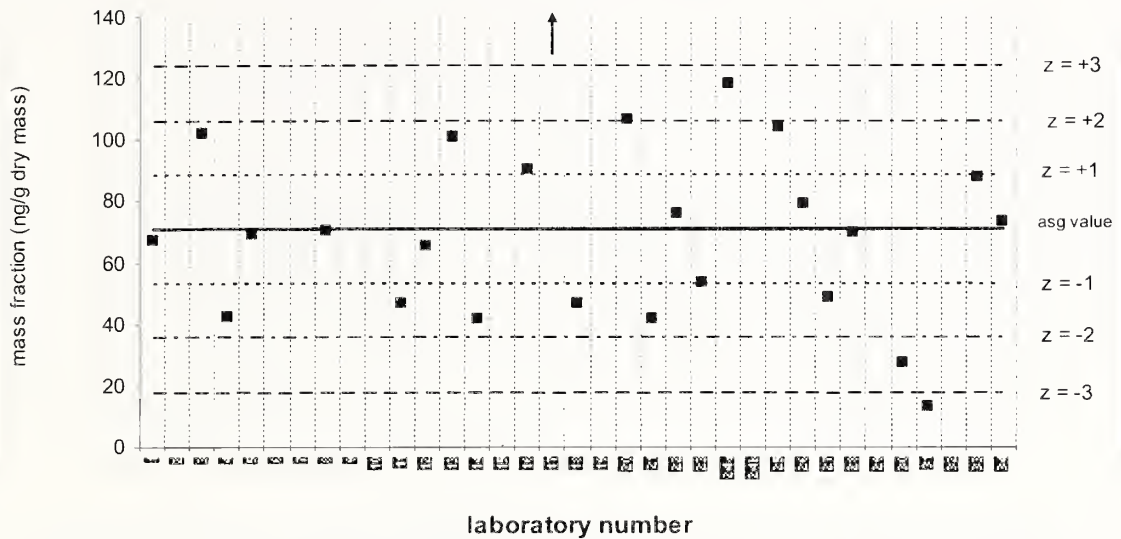
2,6-dimethylnaphthalene

QA10TIS01

Assigned value = 70.7 ng/g dry mass s = 24.9 ng/g dry mass 95% CI = 10.2 ng/g dry mass

Median value = 69.7 ng/g dry mass

Reported Results: 27 Quantitative Results: 25

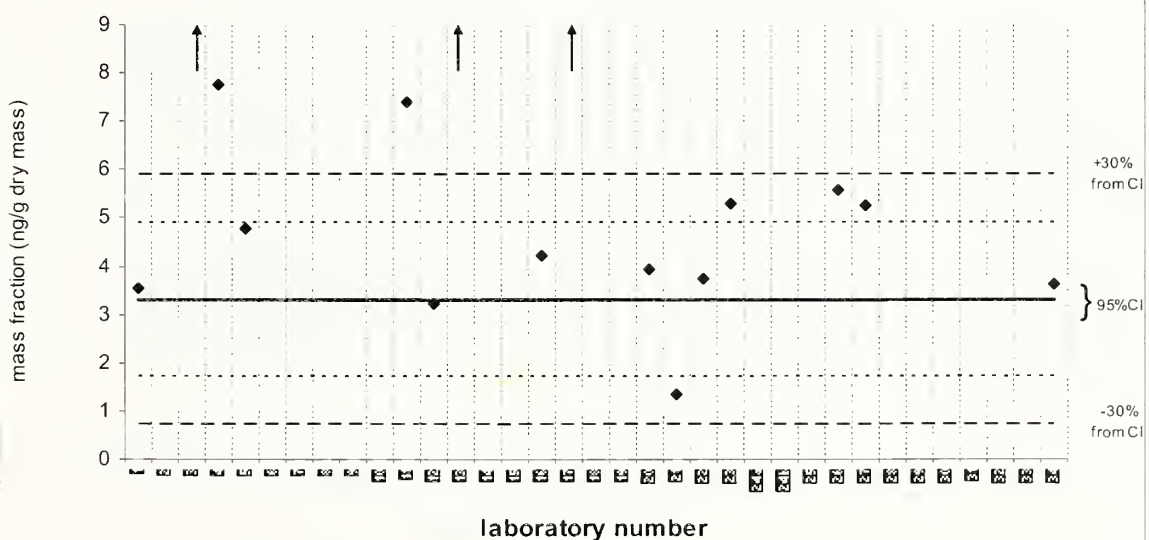


2,6-dimethylnaphthalene

SRM 1974b

Reference Value = 3.3 ng/g dry mass ; 95% CI 1.6 ng/g dry mass; Median value = 5.02 ng/g dry mass

Reported Results: 20 Quantitative Results: 16



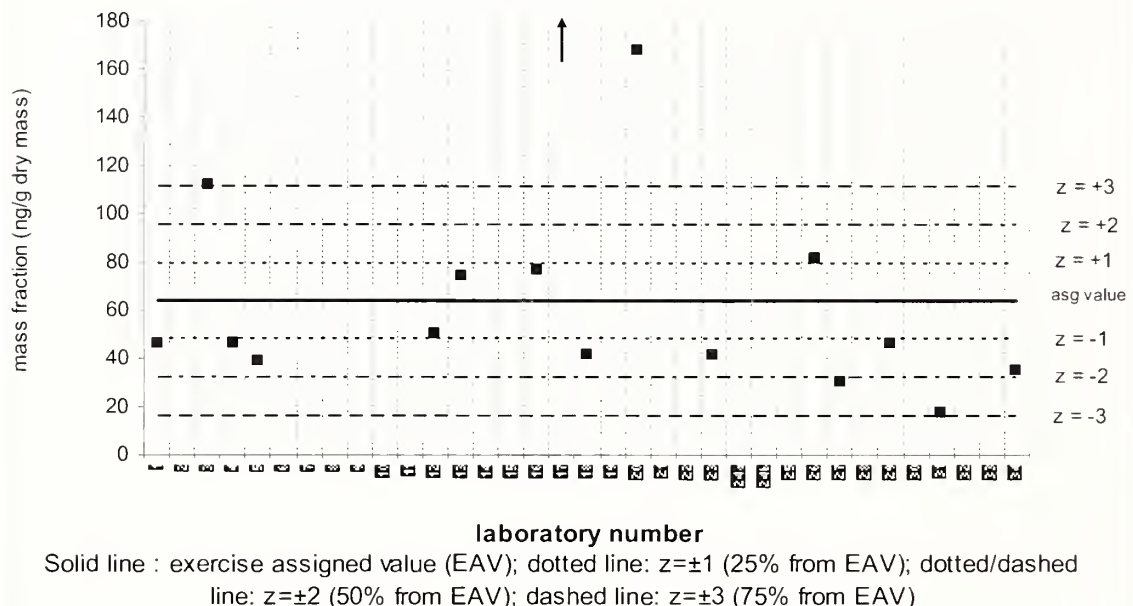
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

1,6,7-trimethylnaphthalene**QA10TIS01**

Assigned value = 63.6 ng/g dry mass s = 37.6 ng/g dry mass 95% CI = 19.7 ng/g dry mass

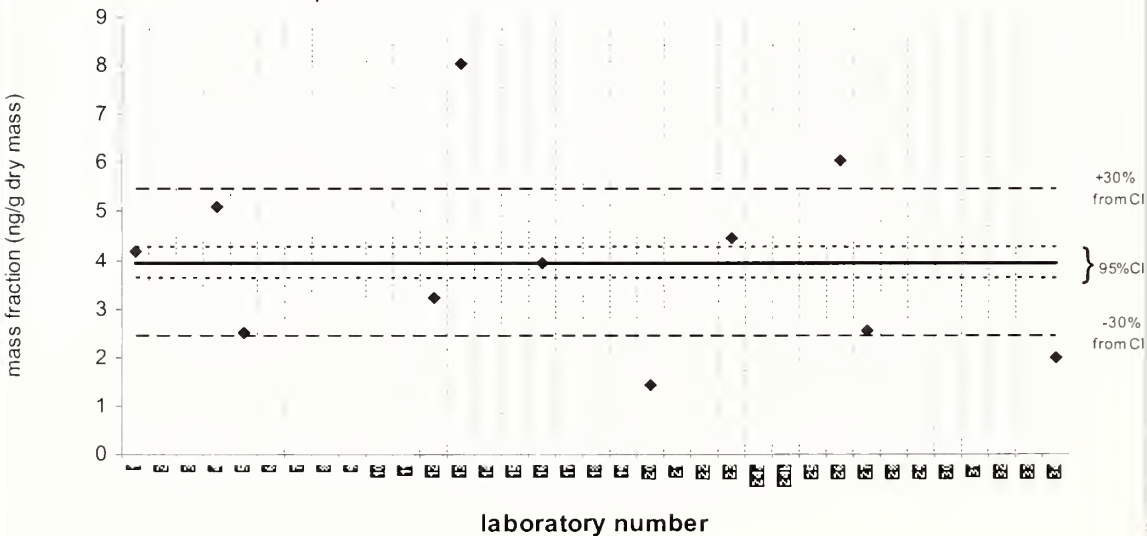
Median value = 46.4 ng/g dry mass

Reported Results: 19 Quantitative Results: 16

**1,6,7-trimethylnaphthalene****SRM 1974b**

Reference Value = 3.95 ng/g dry mass ; 95% CI 0.32 ng/g dry mass; Median value = 3.95 ng/g dry mass

Reported Results: 16 Quantitative Results: 11



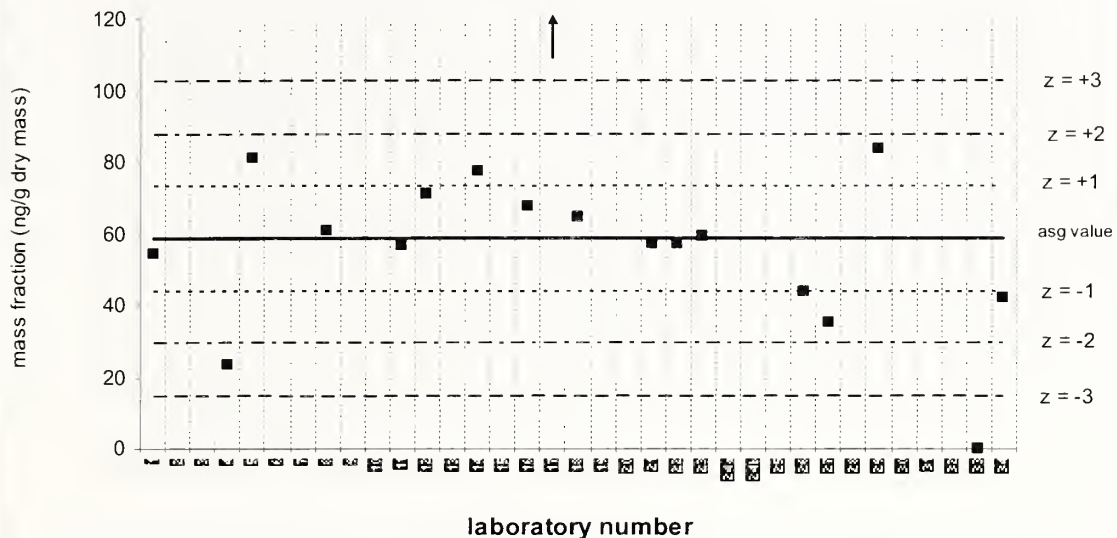
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

1-methylphenanthrene**QA10TIS01**

Assigned value = 58.5 ng/g dry mass s = 16.4 ng/g dry mass 95% CI = 8.0 ng/g dry mass

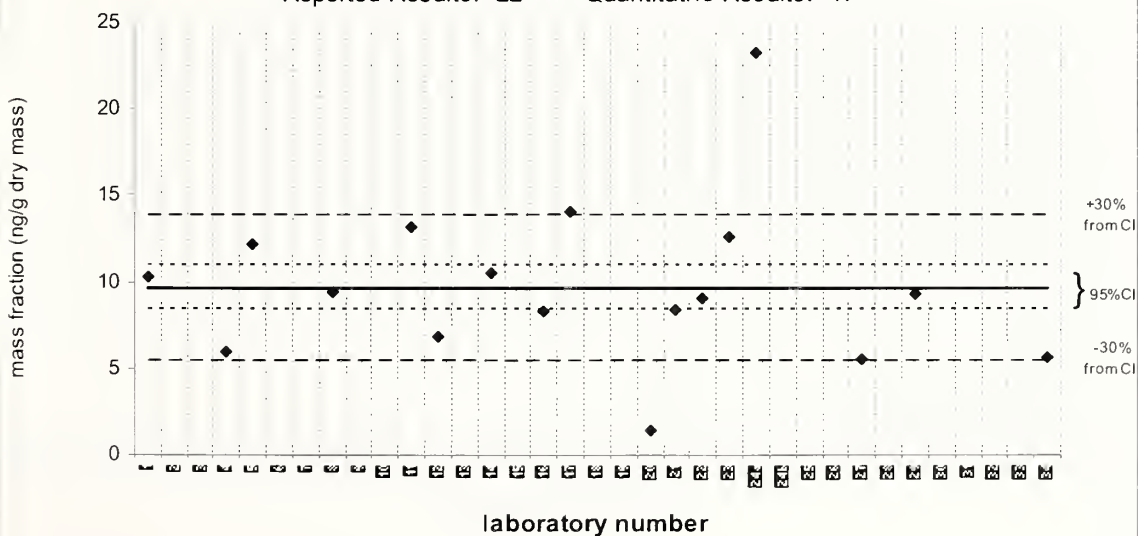
Median value = 58.3 ng/g dry mass

Reported Results: 25 Quantitative Results: 18

**1-methylphenanthrene****SRM 1974b**

Certified Value = 9.66 ng/g dry mass ; 95% CI 1.30 ng/g dry mass: Median value = 9.32 ng/g dry mass

Reported Results: 22 Quantitative Results: 17

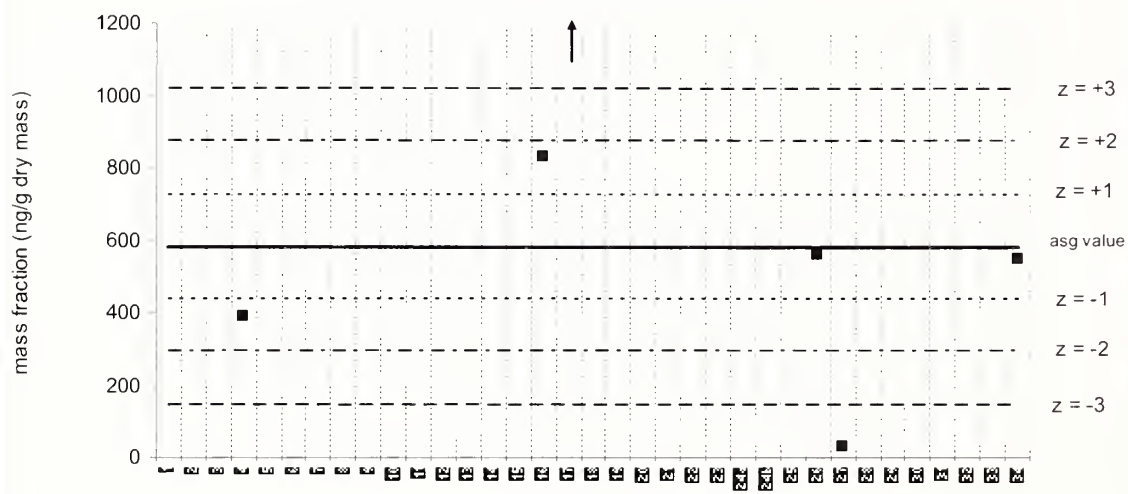


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C1-decalins**QA10TIS01**Assigned value = 582 ng/g dry mass $s = 183$ ng/g dry mass 95% CI = 179 ng/g dry mass

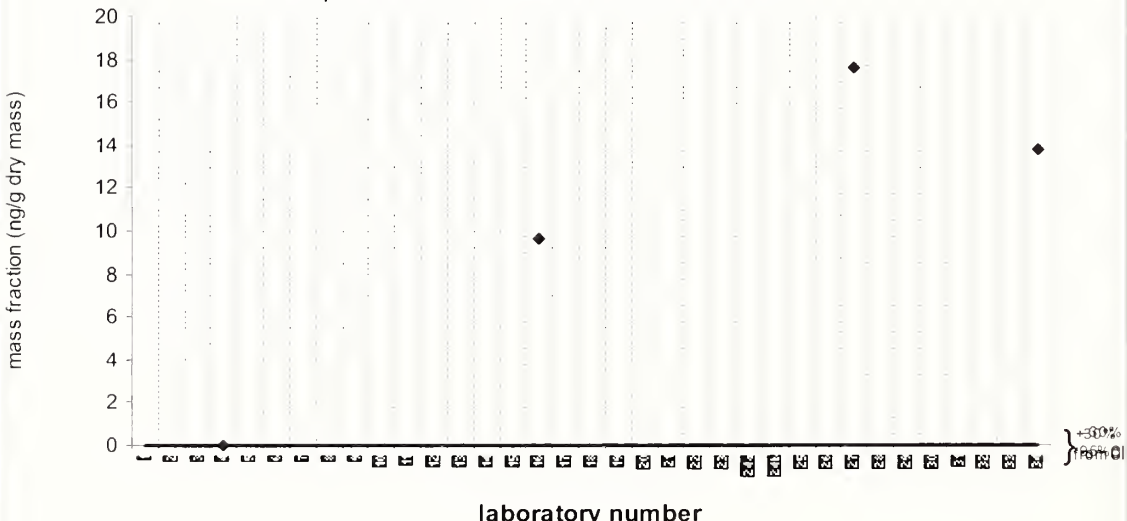
Median value = 554 ng/g dry mass

Reported Results: 7 Quantitative Results: 6

**C1-decalins****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 11.7 ng/g dry mass

Reported Results: 7 Quantitative Results: 4

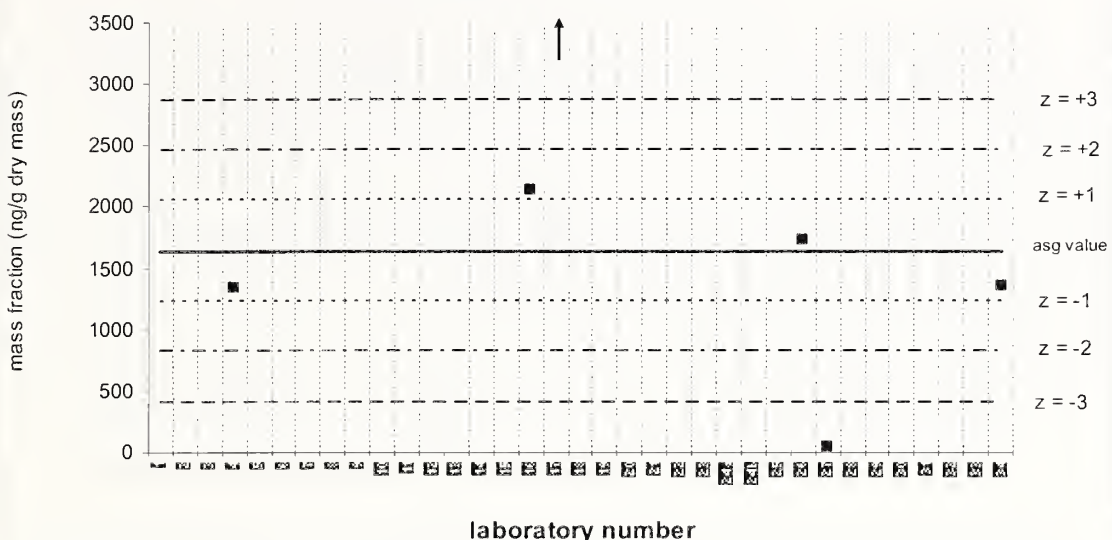


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C2-decalins**QA10TIS01**Assigned value = 1639 ng/g dry mass $s = 379$ ng/g dry mass 95% CI = 372 ng/g dry mass

Median value = 1543 ng/g dry mass

Reported Results: 8 Quantitative Results: 6

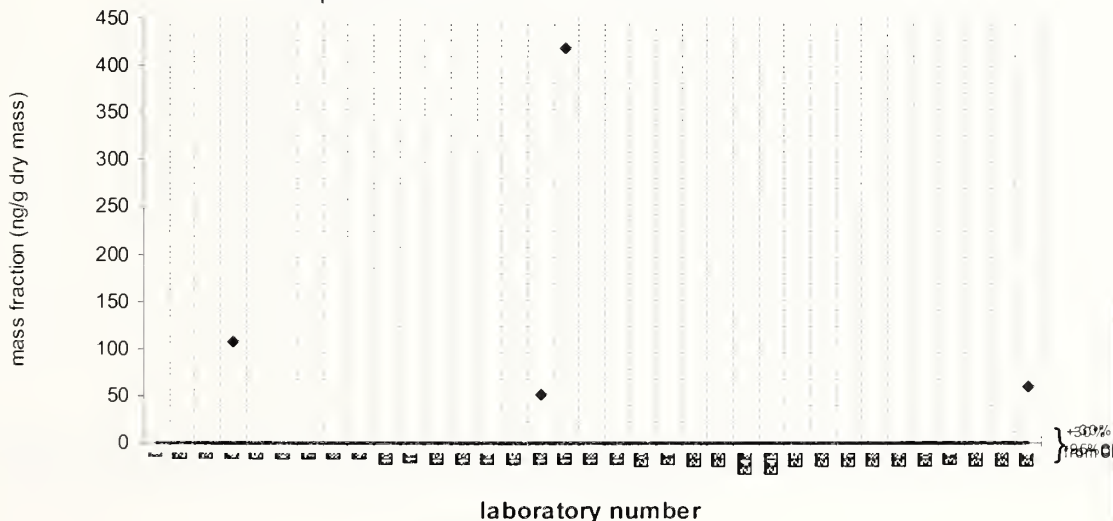


Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

C2-decalins**SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 82.9 ng/g dry mass

Reported Results: 8 Quantitative Results: 4



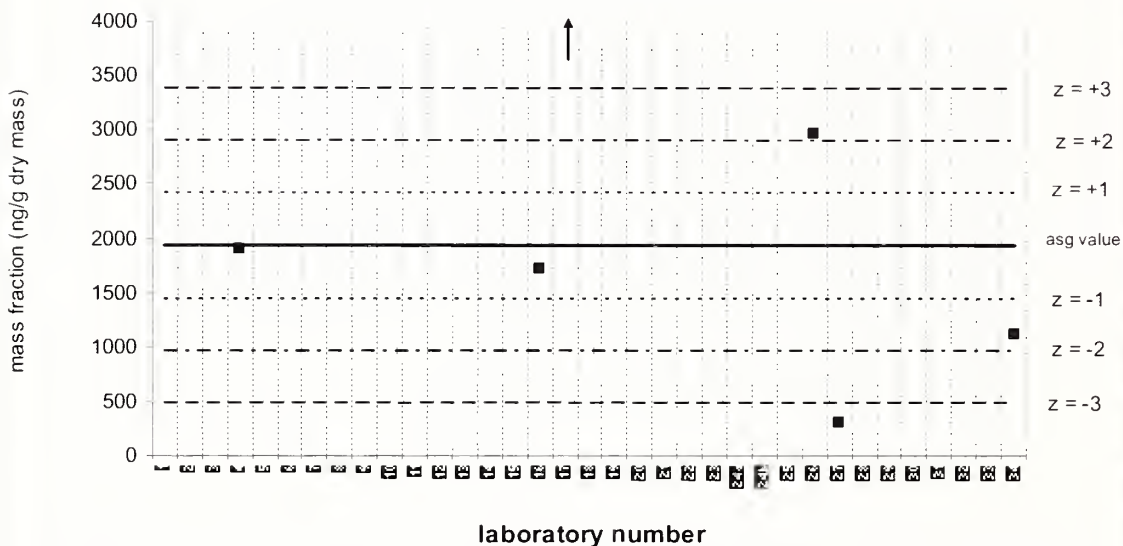
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C3-decalins**QA10TIS01**

Assigned value = 1930 ng/g dry mass s = 766 ng/g dry mass 95% CI = 751 ng/g dry mass

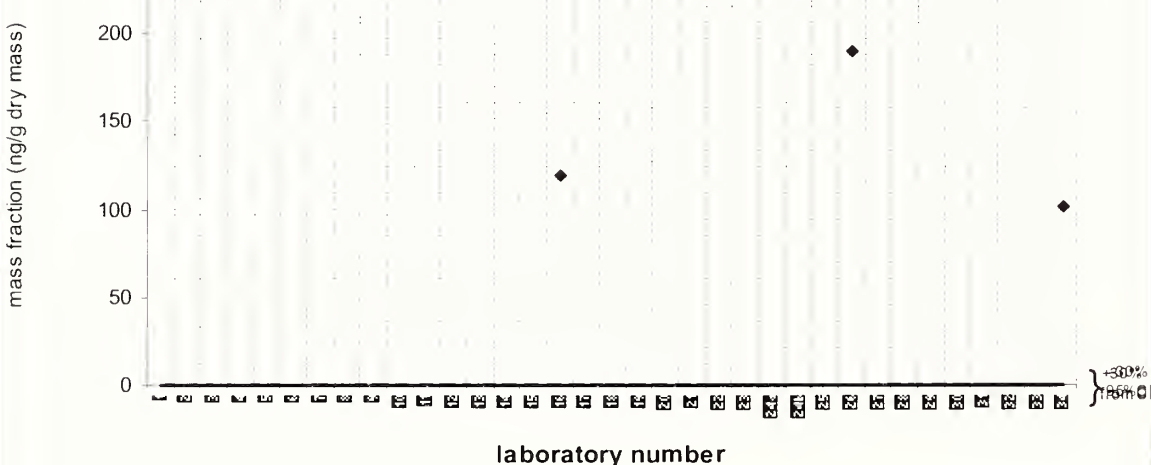
Median value = 1813 ng/g dry mass

Reported Results: 8 Quantitative Results: 6

**C3-decalins****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 155 ng/g dry mass

Reported Results: 8 Quantitative Results: 4



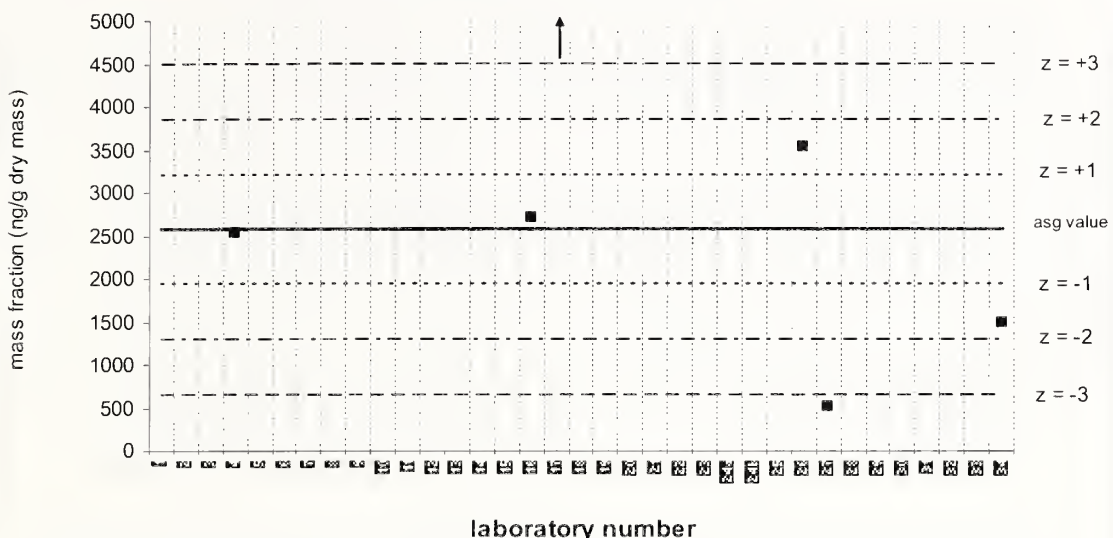
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C4-decalins**QA10TIS01**

Assigned value = 2567 ng/g dry mass s = 838 ng/g dry mass 95% CI = 821 ng/g dry mass

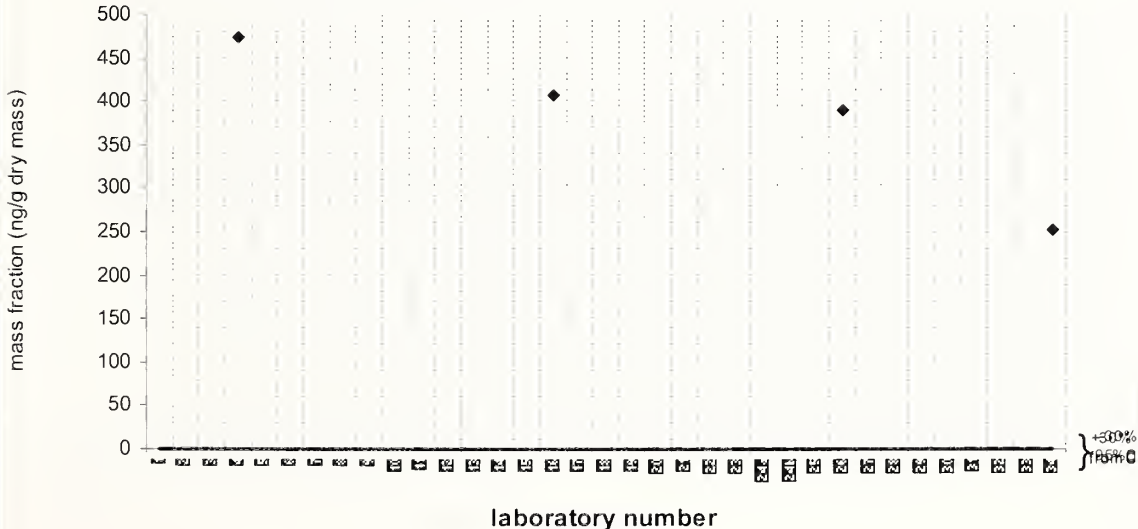
Median value = 2620 ng/g dry mass

Reported Results: 8 Quantitative Results: 6

**C4-decalins****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 398.8 ng/g dry mass

Reported Results: 8 Quantitative Results: 4



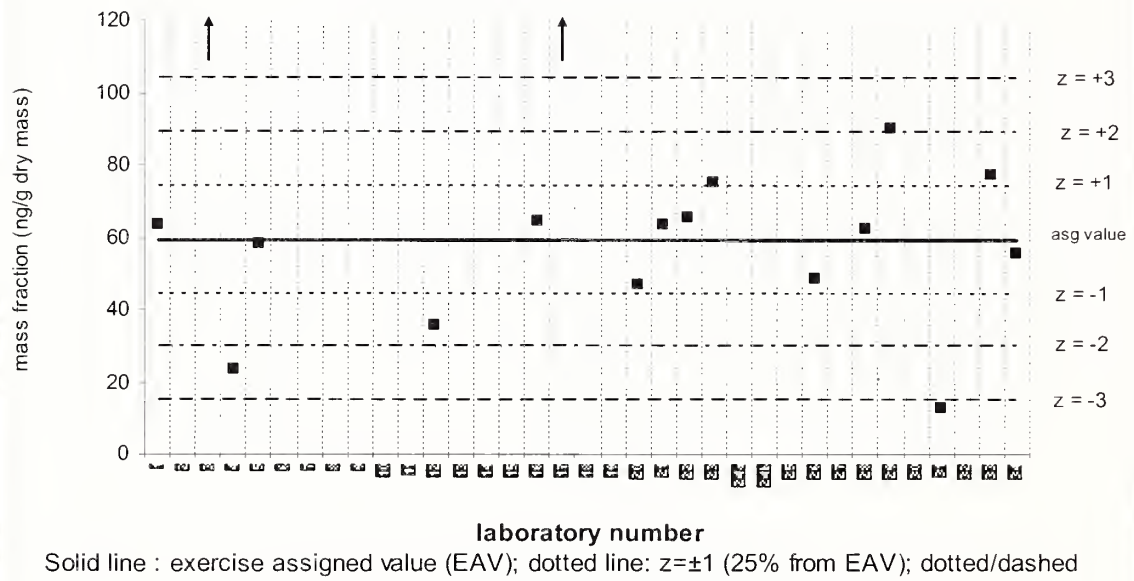
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C1-naphthalenes

QA10TIS01

Assigned value = 59.3 ng/g dry mass s = 17.0 ng/g dry mass 95% CI = 8.9 ng/g dry mass
Median value = 63.4 ng/g dry mass

Reported Results: 22 Quantitative Results: 17

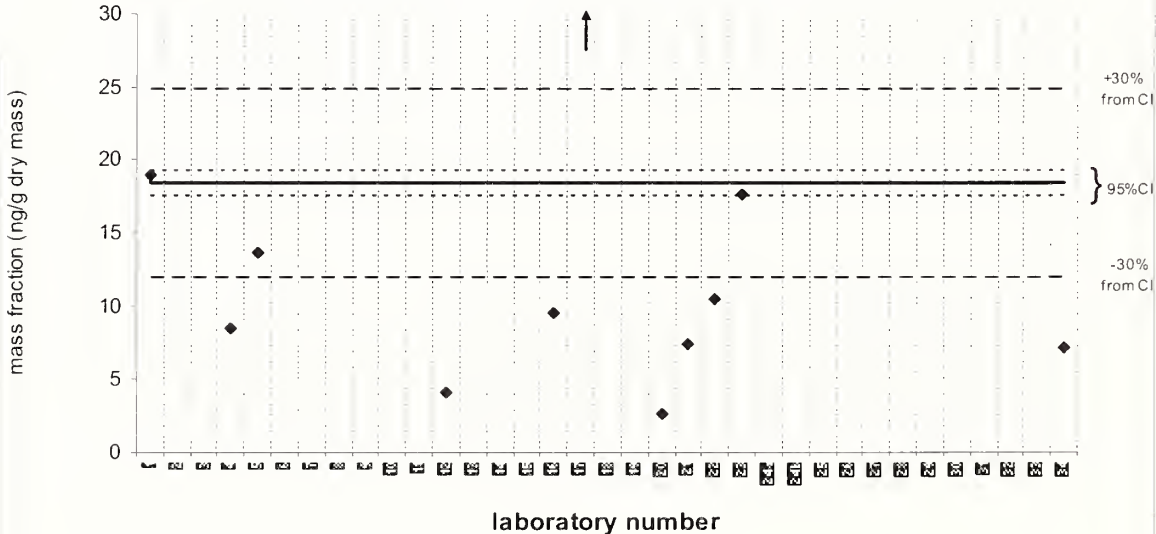


C1-naphthalenes

SRM 1974b

Target Value = 18.4 ng/g dry mass ; 95% CI 0.9 ng/g dry mass; Median value = 9.50 ng/g dry mass

Reported Results: 16 Quantitative Results: 11



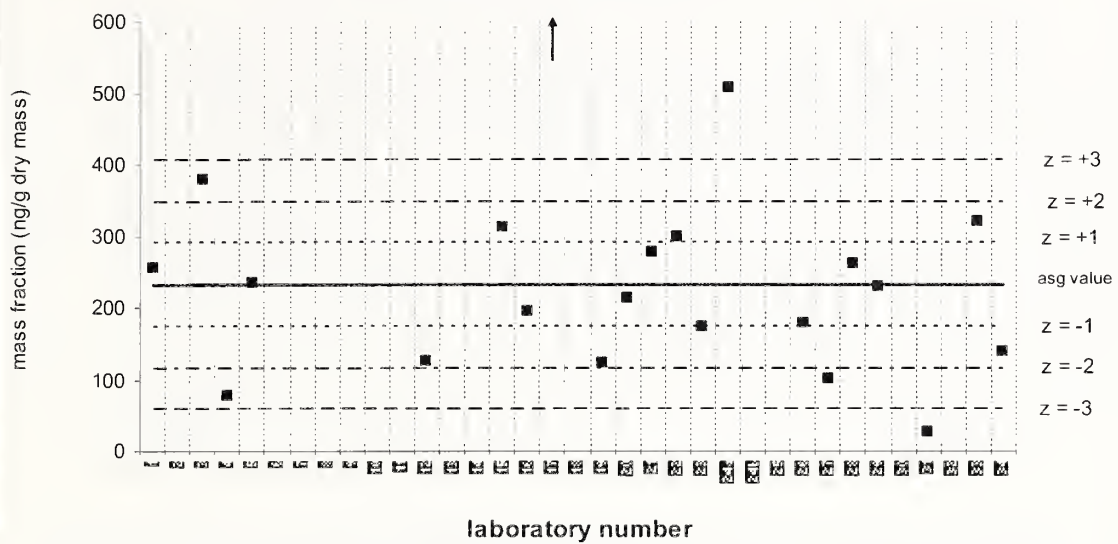
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C2-naphthalenes

QA10TIS01

Assigned value = 232 ng/g dry mass s = 106 ng/g dry mass 95% CI = 47 ng/g dry mass Median value = 229 ng/g dry mass

Reported Results: 24 Quantitative Results: 21



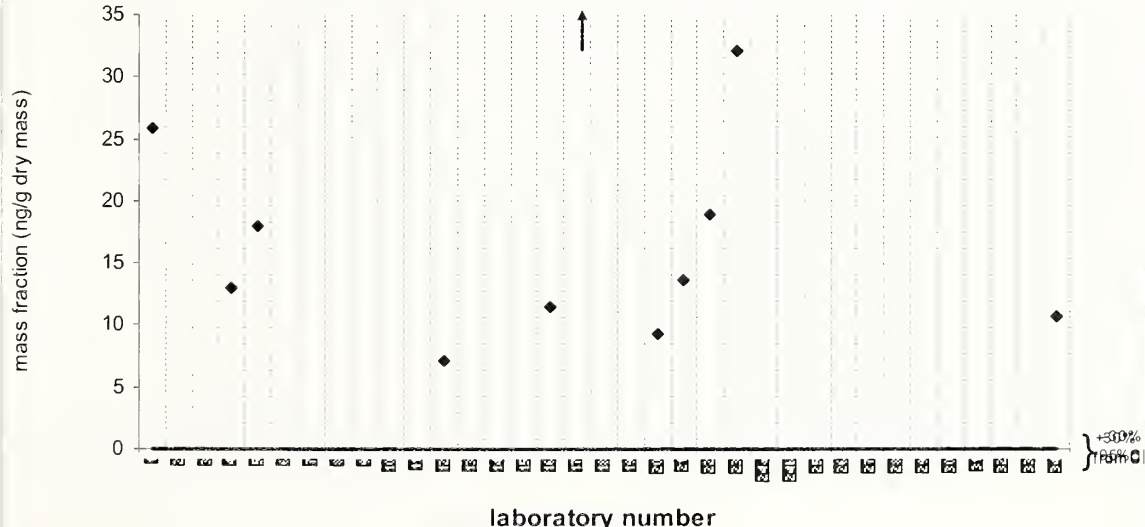
Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

C2-naphthalenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 13.6 ng/g dry mass

Reported Results: 16 Quantitative Results: 11



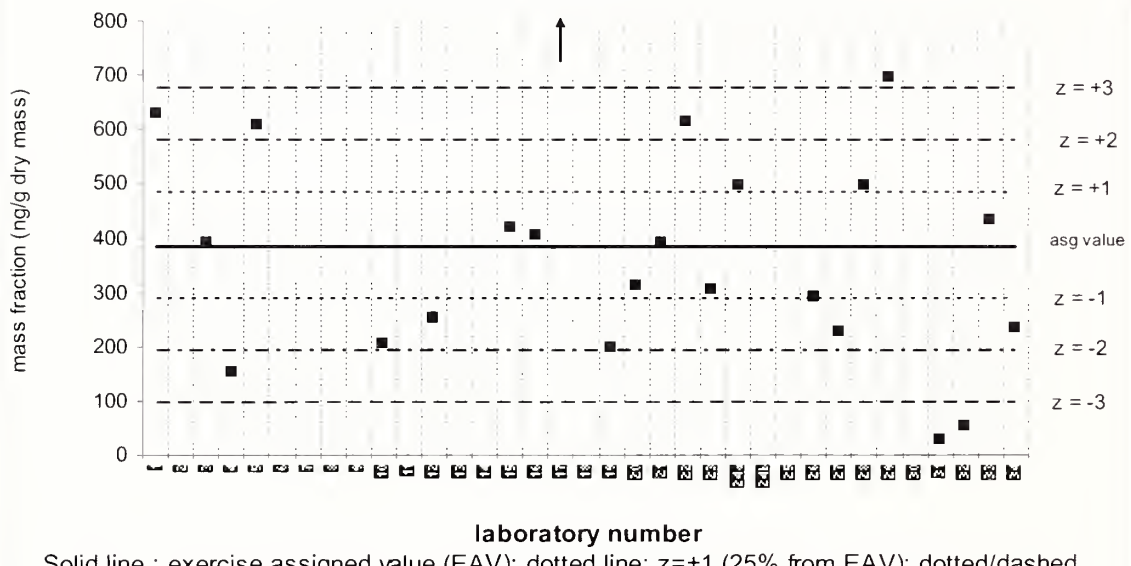
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C3-naphthalenes

QA10TIS01

Assigned value = 386 ng/g dry mass $s = 165$ ng/g dry mass 95% CI = 72 ng/g dry mass Median value = 390 ng/g dry mass

Reported Results: 24 Quantitative Results: 23



laboratory number

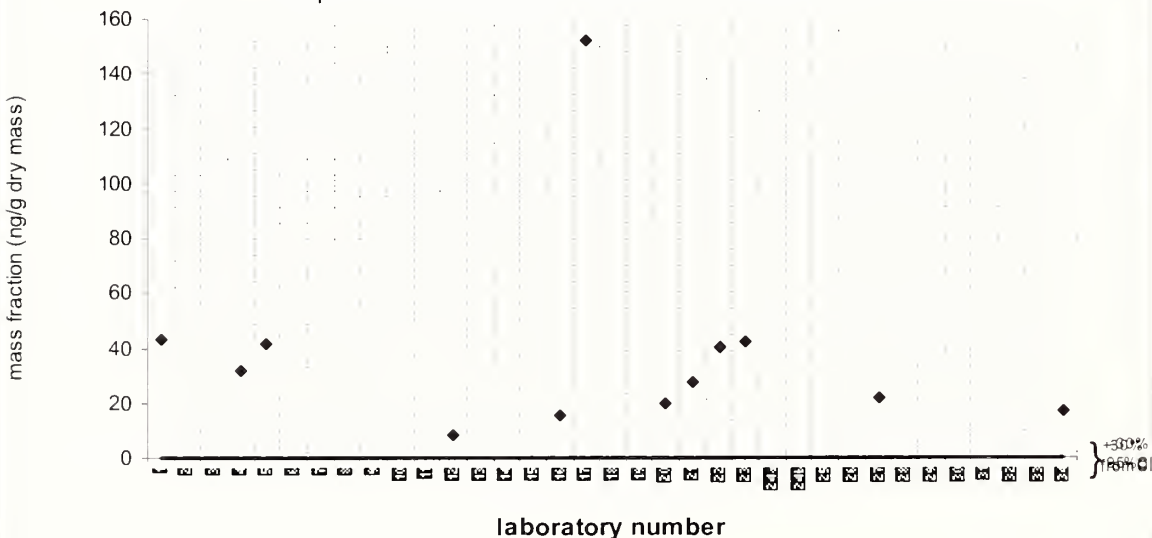
Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

C3-naphthalenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 29.7 ng/g dry mass

Reported Results: 17 Quantitative Results: 12



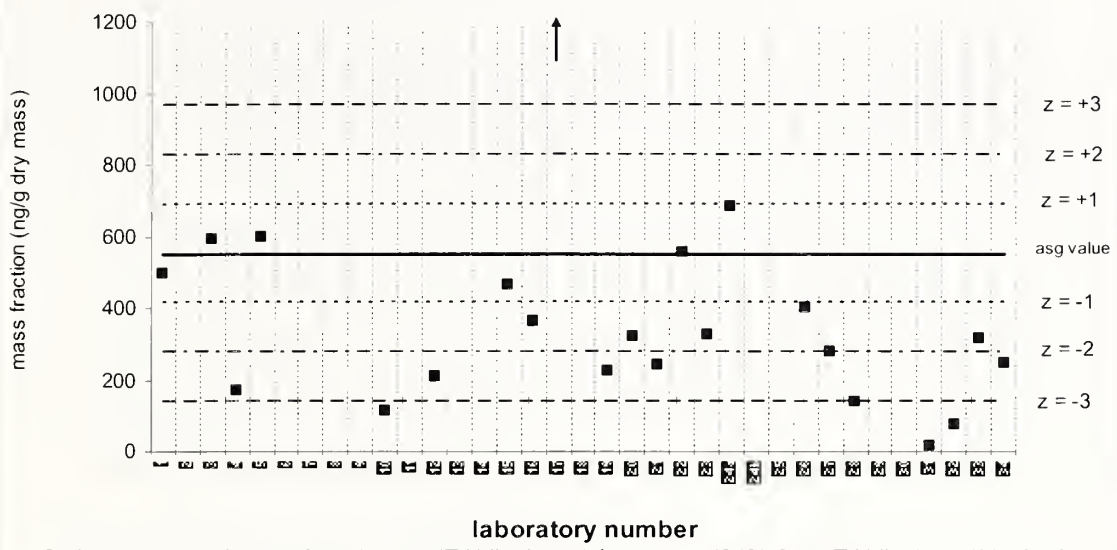
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C4-naphthalenes

QA10TIS01

Assigned value = 552 ng/g dry mass $s = 173$ ng/g dry mass 95% CI = 76 ng/g dry mass Median value = 316 ng/g dry mass

Reported Results: 23 Quantitative Results: 22



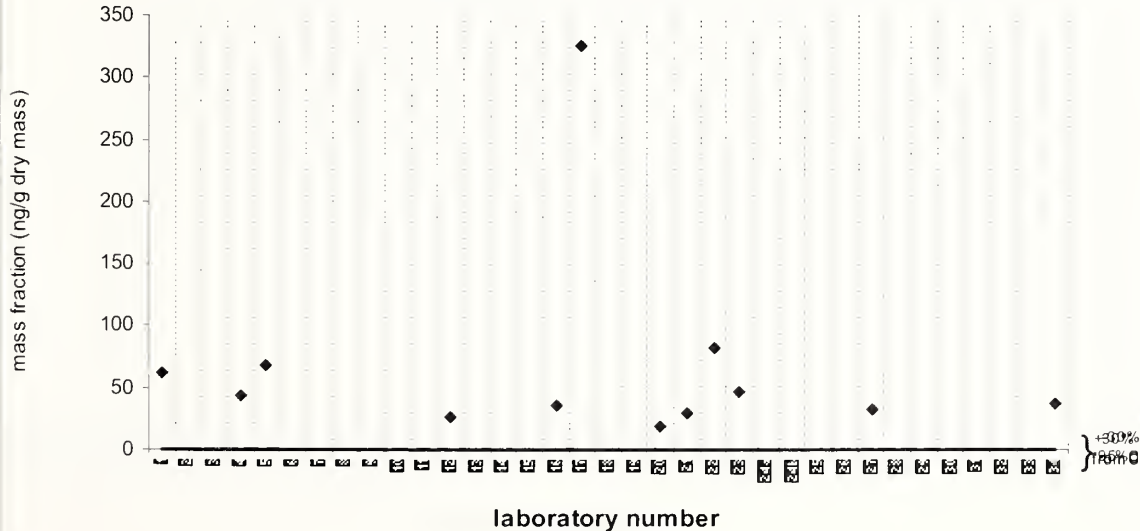
Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

C4-naphthalenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 39.9 ng/g dry mass

Reported Results: 17 Quantitative Results: 12



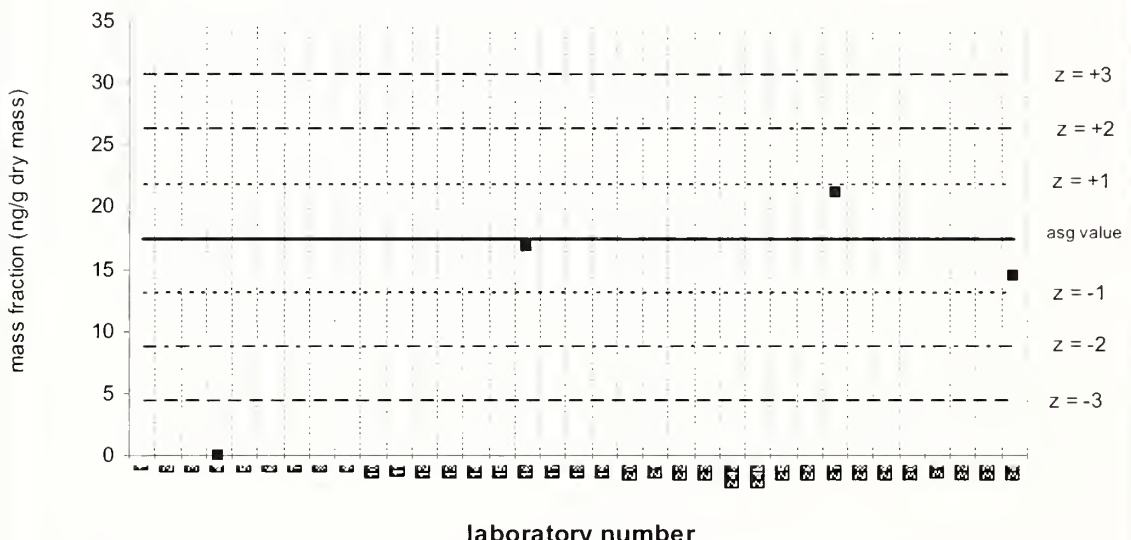
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C1-benzothiophenes

QA10TIS01

Assigned value = 17.5 ng/g dry mass s = 3.4 ng/g dry mass 95% CI = 3.8 ng/g dry mass Median value = 15.7 ng/g dry mass

Reported Results: 8 Quantitative Results: 4



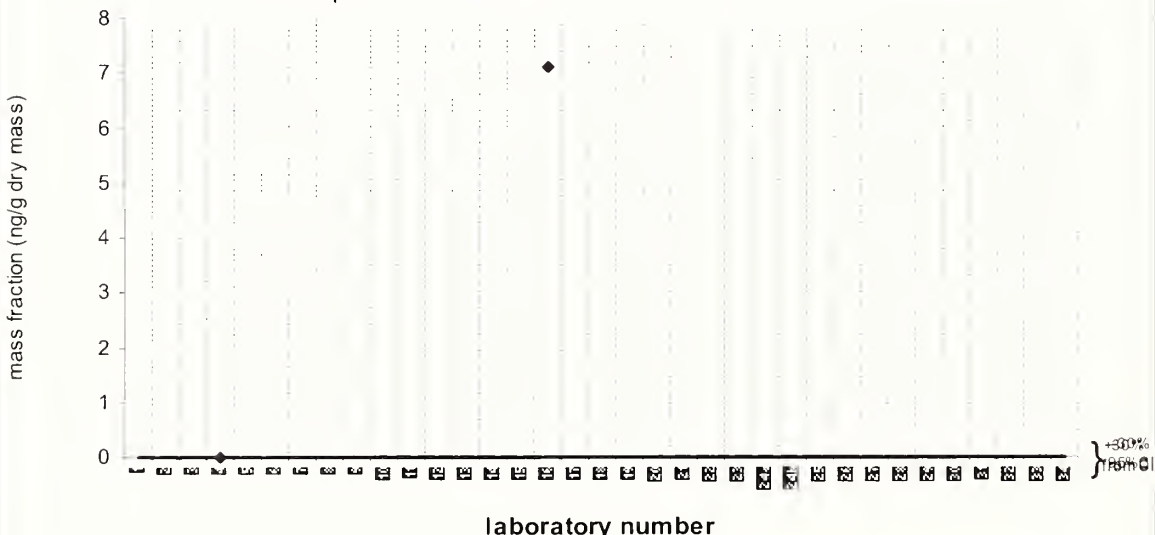
Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

C1-benzothiophenes

SRM 1974b

Target Value = no target ng/g (dry mass): Median value = 3.56 ng/g dry mass

Reported Results: 8 Quantitative Results: 2

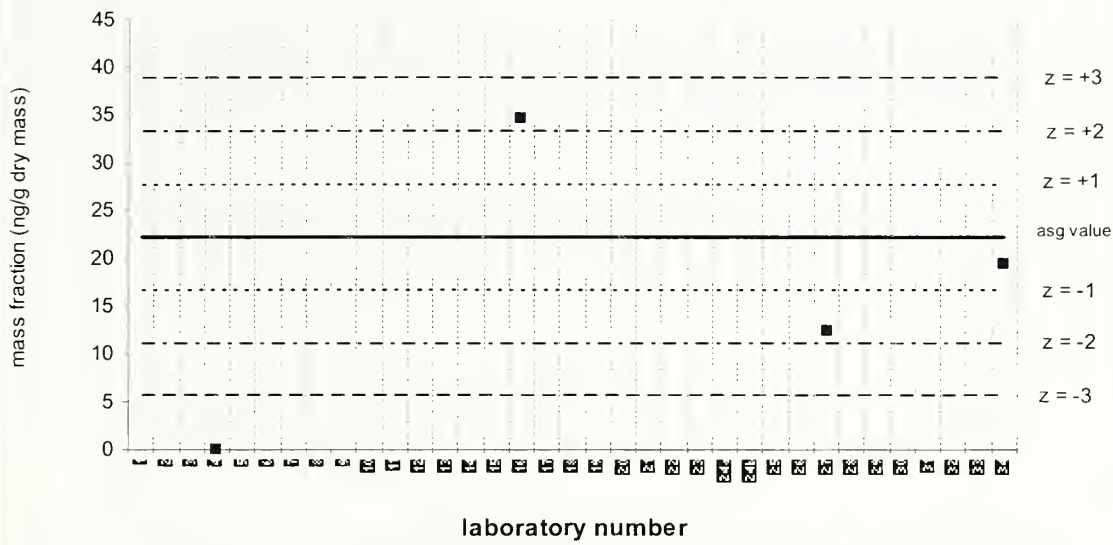


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C2-benzothiophenes**QA10TIS01**Assigned value = 22.2 ng/g dry mass $s = 11.3$ ng/g dry mass 95% CI = 12.8 ng/g dry mass

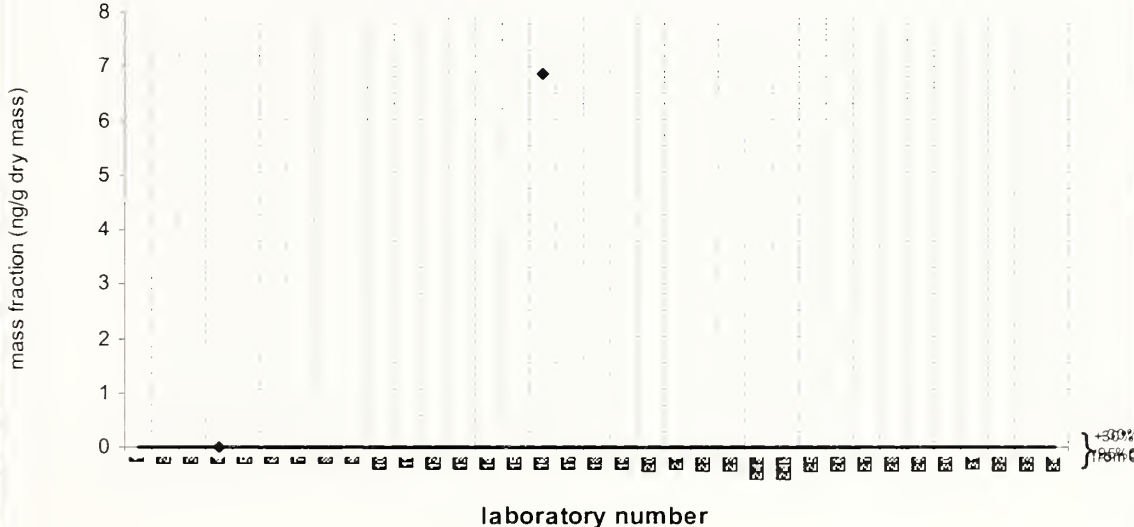
Median value = 15.9 ng/g dry mass

Reported Results: 8 Quantitative Results: 4

**C2-benzothiophenes****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 3.43 ng/g dry mass

Reported Results: 8 Quantitative Results: 2



Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

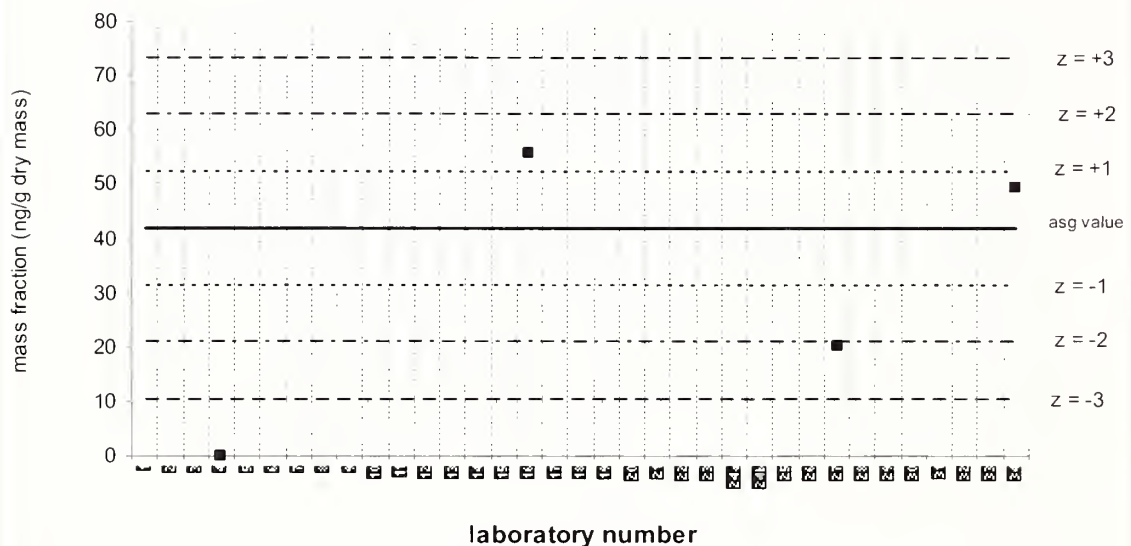
C3-benzothiophenes

QA10TIS01

Assigned value = 41.8 ng/g dry mass s = 18.9 ng/g dry mass 95% CI = 21.4 ng/g dry mass

Median value = 34.9 ng/g dry mass

Reported Results: 8 Quantitative Results: 4



laboratory number

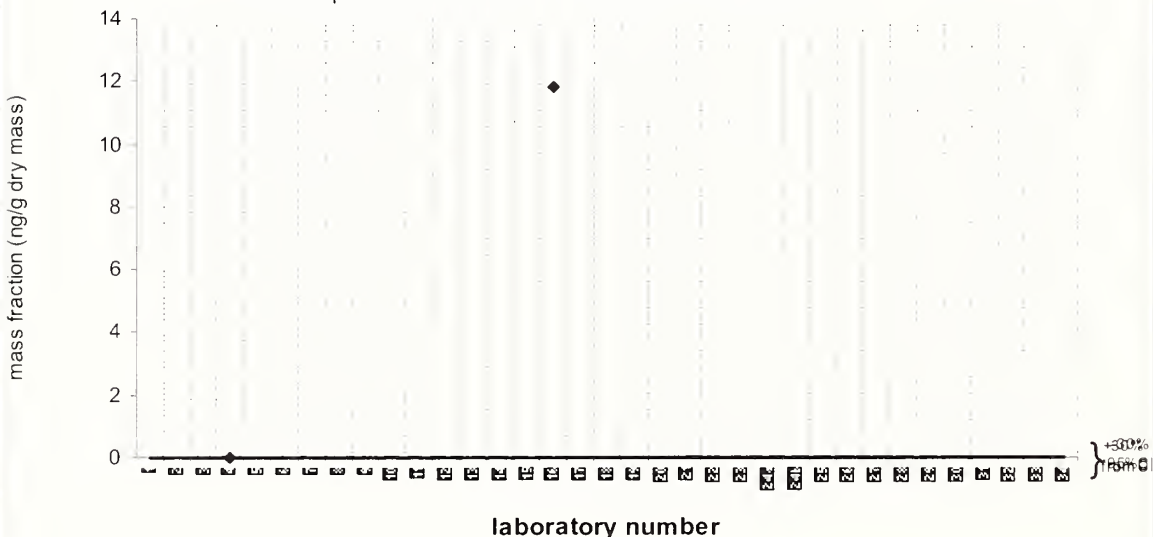
Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

C3-benzothiophenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 5.92 ng/g dry mass

Reported Results: 8 Quantitative Results: 2



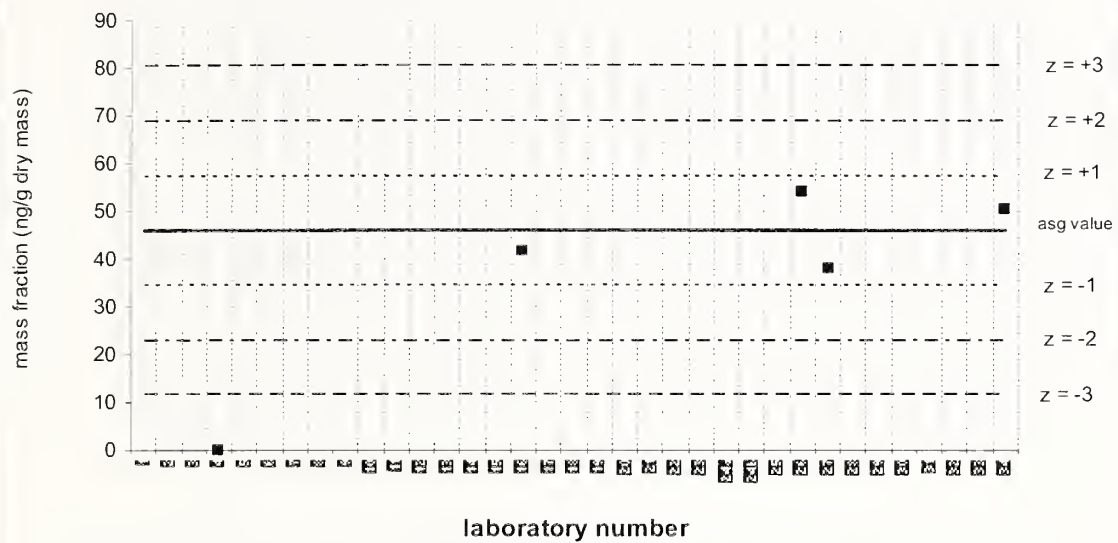
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C4-benzothiophenes

QA10TIS01

Assigned value = 45.9 ng/g dry mass s = 7.5 ng/g dry mass 95% CI = 7.3 ng/g dry mass Median value = 41.6 ng/g dry mass

Reported Results: 7 Quantitative Results: 5



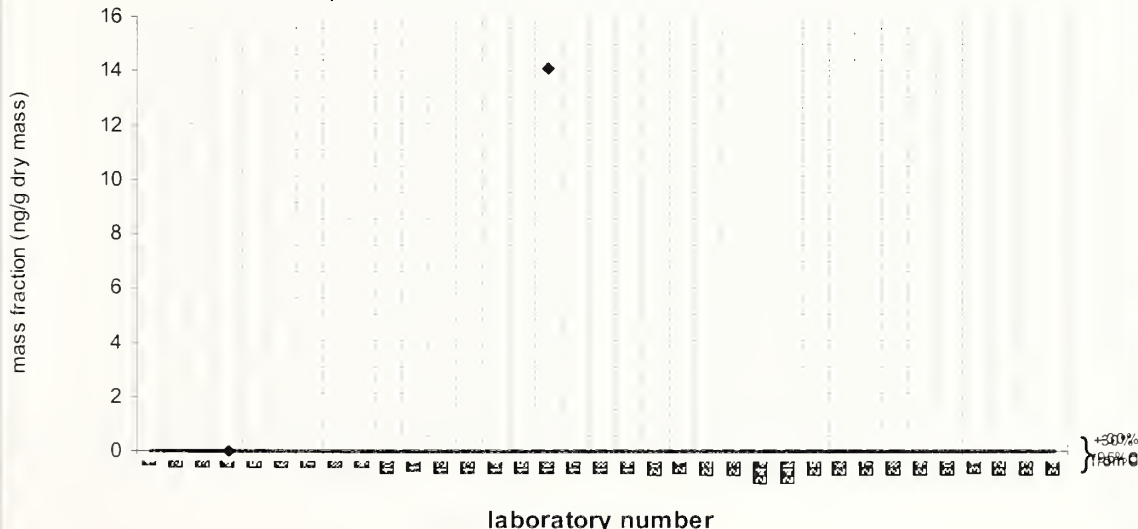
Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

C4-benzothiophenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 7.05 ng/g dry mass

Reported Results: 7 Quantitative Results: 2

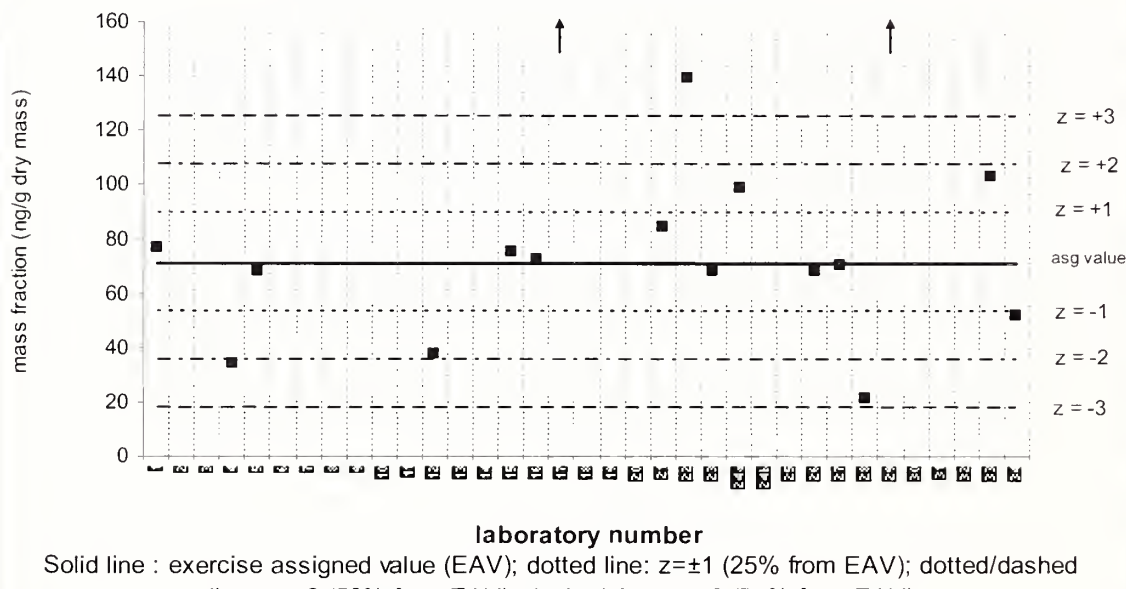


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C1-fluorenes**QA10TIS01**Assigned value = 71.4 ng/g dry mass $s = 29.3$ ng/g dry mass 95% CI = 14.9 ng/g dry mass

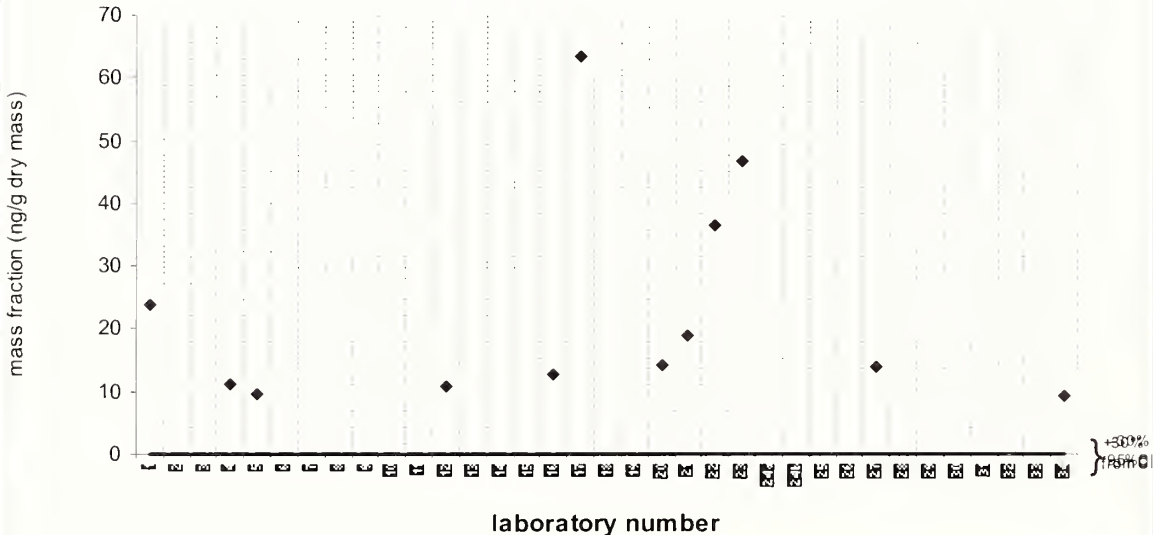
Median value = 72.4 ng/g dry mass

Reported Results: 24 Quantitative Results: 17

**C1-fluorenes****SRM 1974b**

Target Value = no target ng/g (dry mass): Median value = 14.0 ng/g dry mass

Reported Results: 17 Quantitative Results: 12

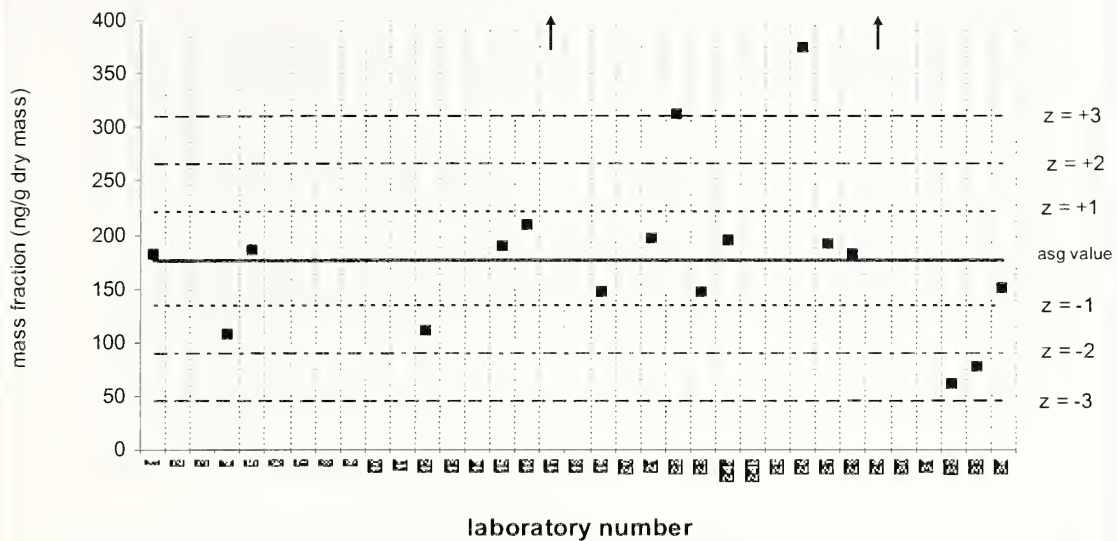


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C2-fluorenes**QA10TIS01**

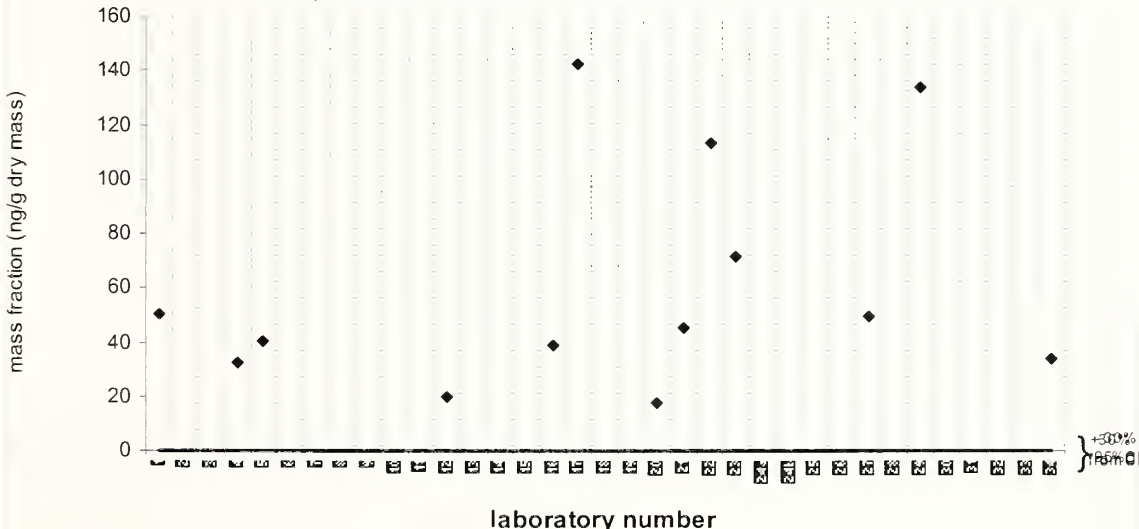
Assigned value = 177 ng/g dry mass $s = 77$ ng/g dry mass 95% CI = 37 ng/g dry mass Median value = 186 ng/g dry mass

Reported Results: 24 Quantitative Results: 19

**C2-fluorenes****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 45.4 ng/g dry mass

Reported Results: 18 Quantitative Results: 13

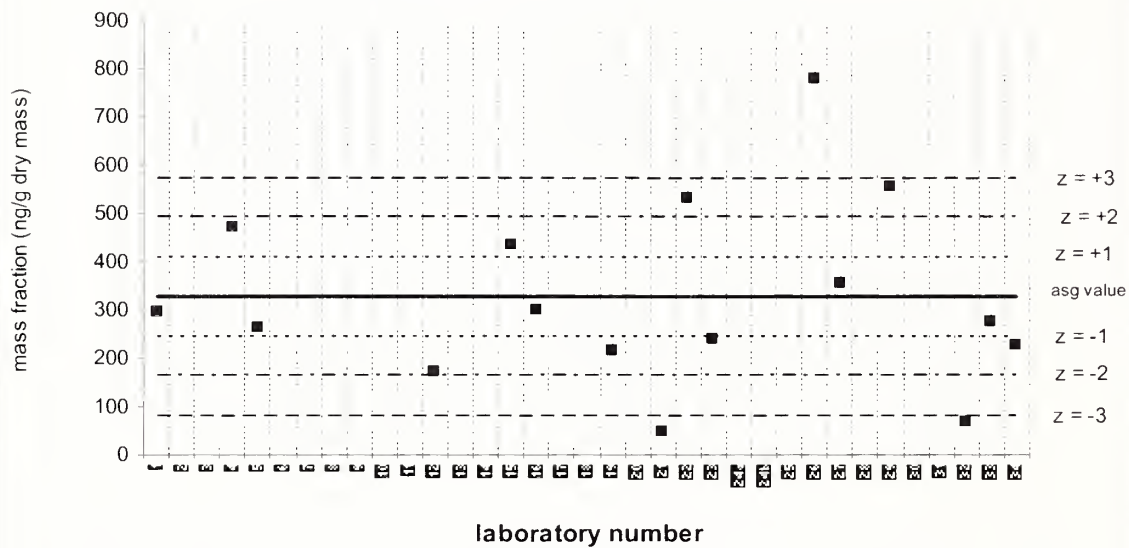


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C3-fluorenes**QA10TIS01**

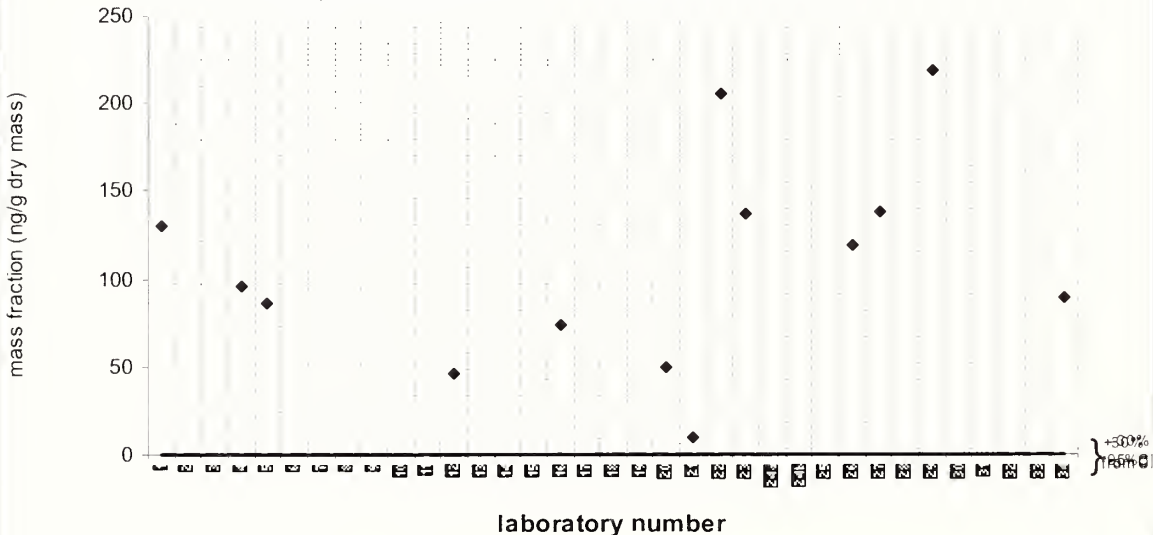
Assigned value = 328 ng/g dry mass $s = 190$ ng/g dry mass 95% CI = 93 ng/g dry mass Median value = 287 ng/g dry mass

Reported Results: 39 Quantitative Results: 16

**C3-fluorenes****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 96.3 ng/g dry mass

Reported Results: 18 Quantitative Results: 13



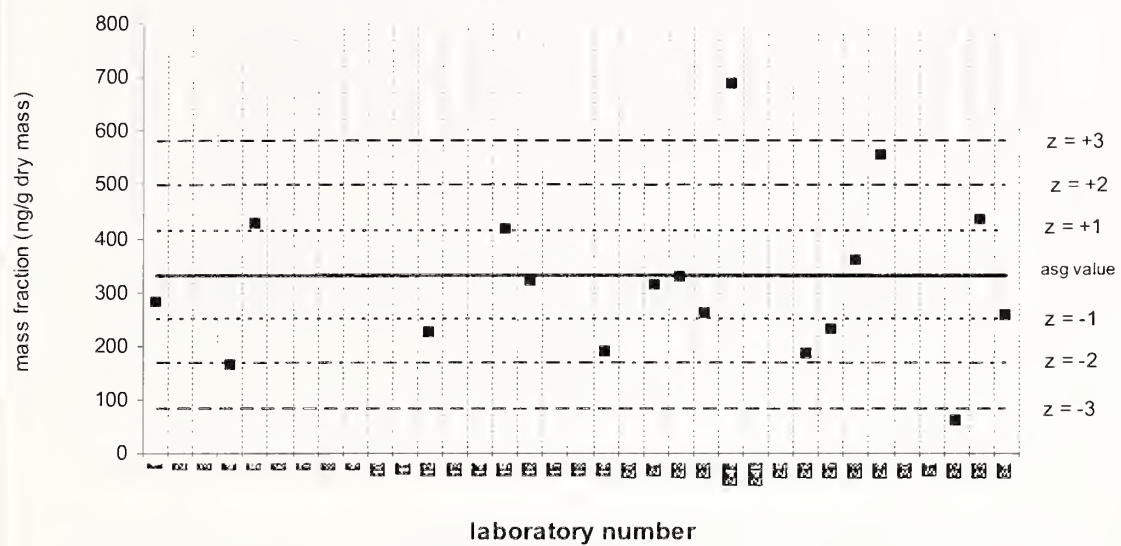
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C1-phenanthrenes/anthracenes

QA10TIS01

Assigned value = 331 ng/g dry mass s = 138 ng/g dry mass 95% CI = 66 ng/g dry mass Median value = 296 ng/g dry mass

Reported Results: 24 Quantitative Results: 18

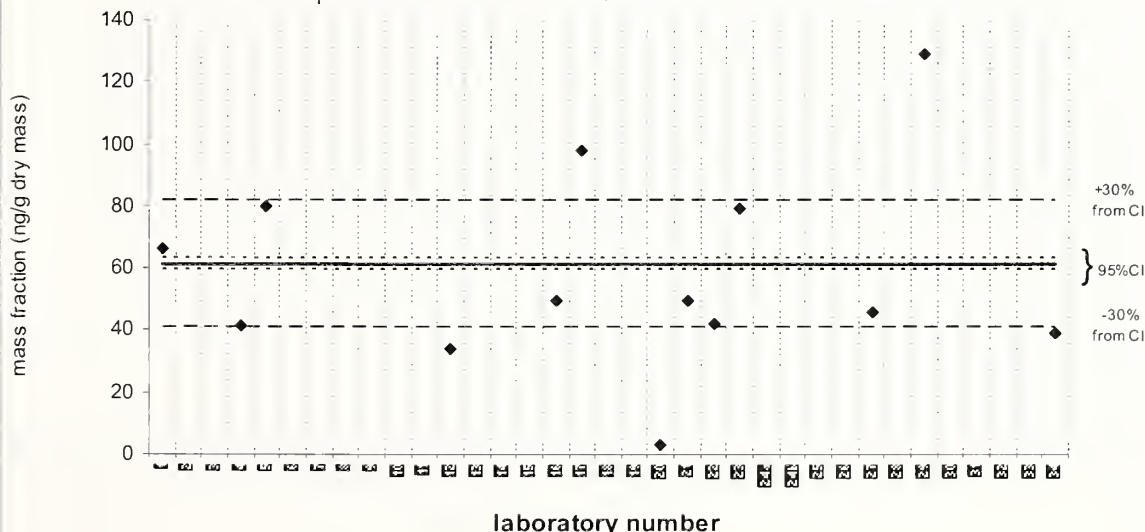


C1-phenanthrenes/anthracenes

SRM 1974b

Target Value = 61.3 ng/g dry mass ; 95% CI 1.8 ng/g dry mass: Median value = 49.3 ng/g dry mass

Reported Results: 18 Quantitative Results: 13



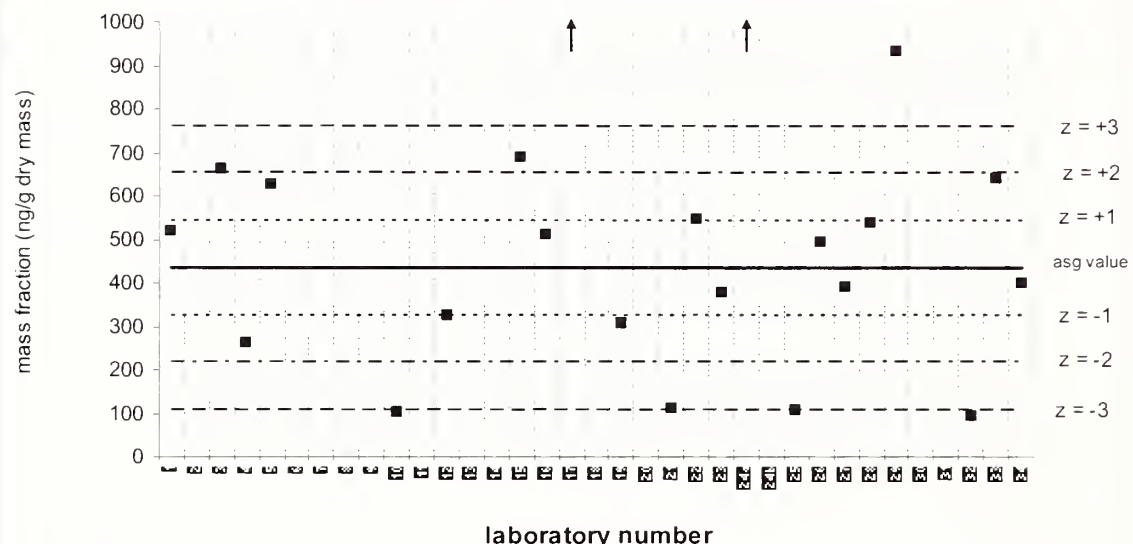
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C2-phenanthrenes/anthracenes

QA10TIS01

Assigned value = 434 ng/g dry mass s = 219 ng/g dry mass 95% CI = 99 ng/g dry mass Median value = 502 ng/g dry mass

Reported Results: 24 Quantitative Results: 22

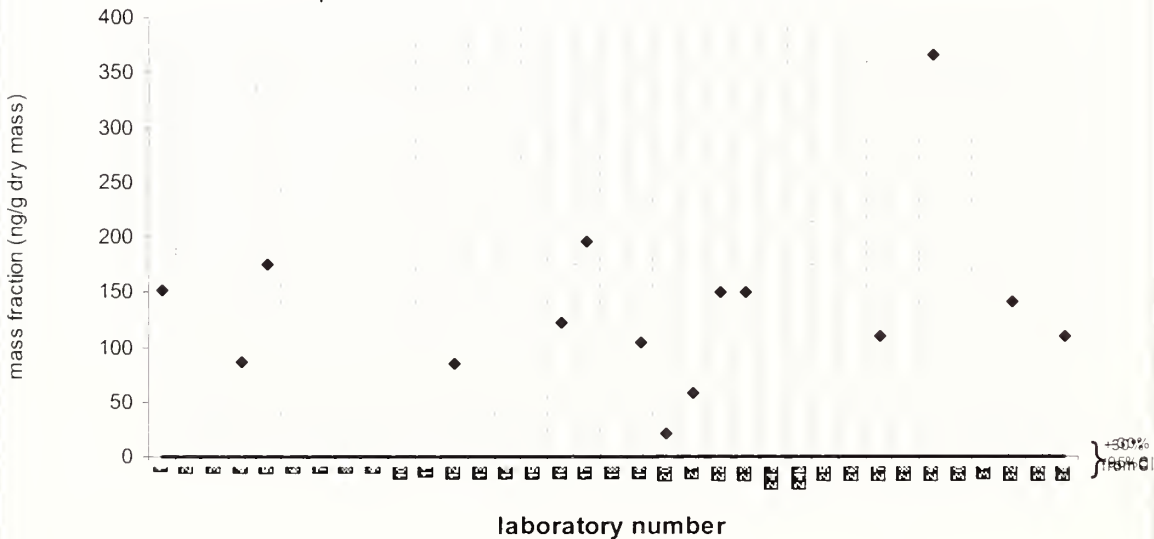


C2-phenanthrenes/anthracenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 123 ng/g dry mass

Reported Results: 18 Quantitative Results: 15



Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

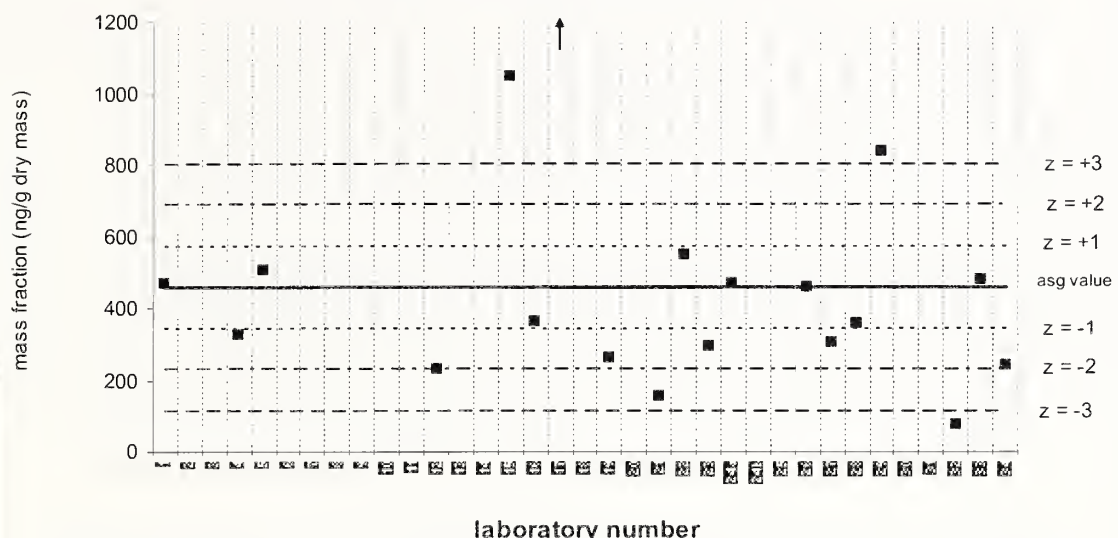
C3-phenanthrenes/anthracenes

QA10TIS01

Assigned value = 457 ng/g dry mass $s = 224$ ng/g dry mass 95% CI = 107 ng/g dry mass

Median value = 361 ng/g dry mass

Reported Results: 24 Quantitative Results: 19



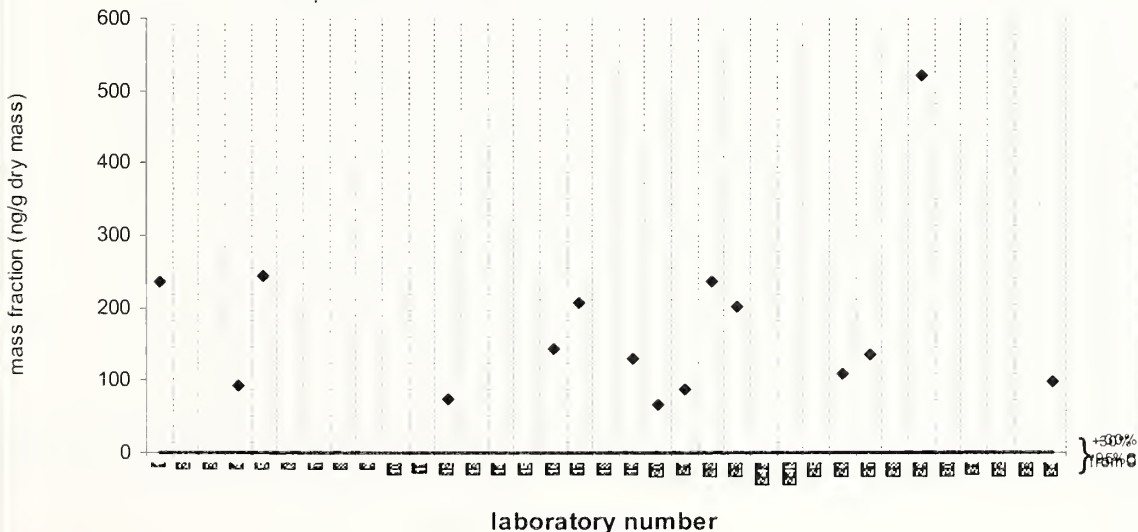
Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

C3-phenanthrenes/anthracenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 137 ng/g dry mass

Reported Results: 18 Quantitative Results: 15



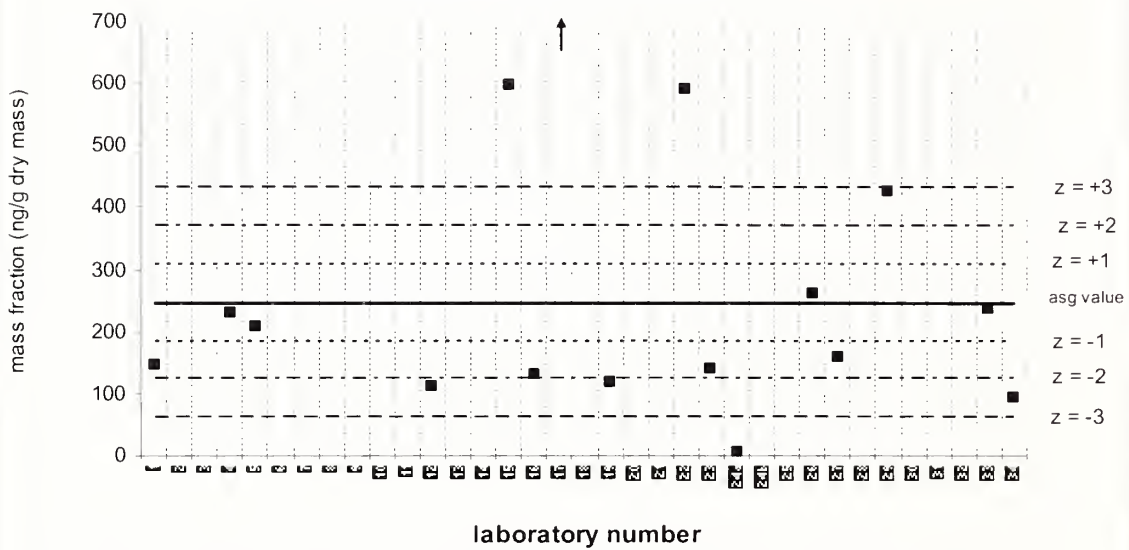
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C4-phenanthrenes/anthracenes

QA10TIS01

Assigned value = 246 ng/g dry mass s = 170 ng/g dry mass 95% CI = 89 ng/g dry mass Median value = 184 ng/g dry mass

Reported Results: 23 Quantitative Results: 16

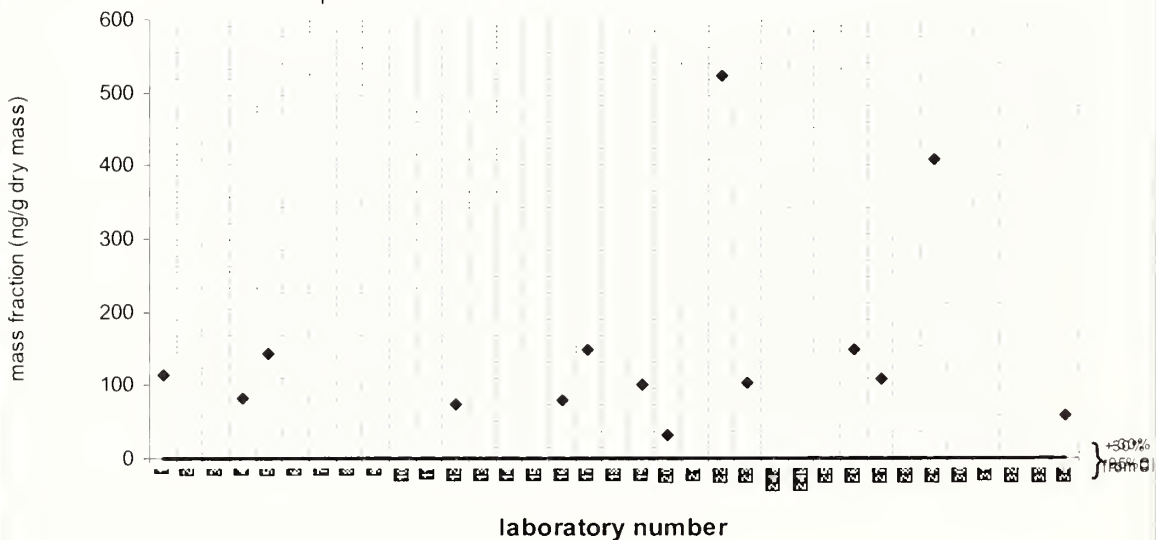


C4-phenanthrenes/anthracenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 107 ng/g dry mass

Reported Results: 17 Quantitative Results: 14



Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

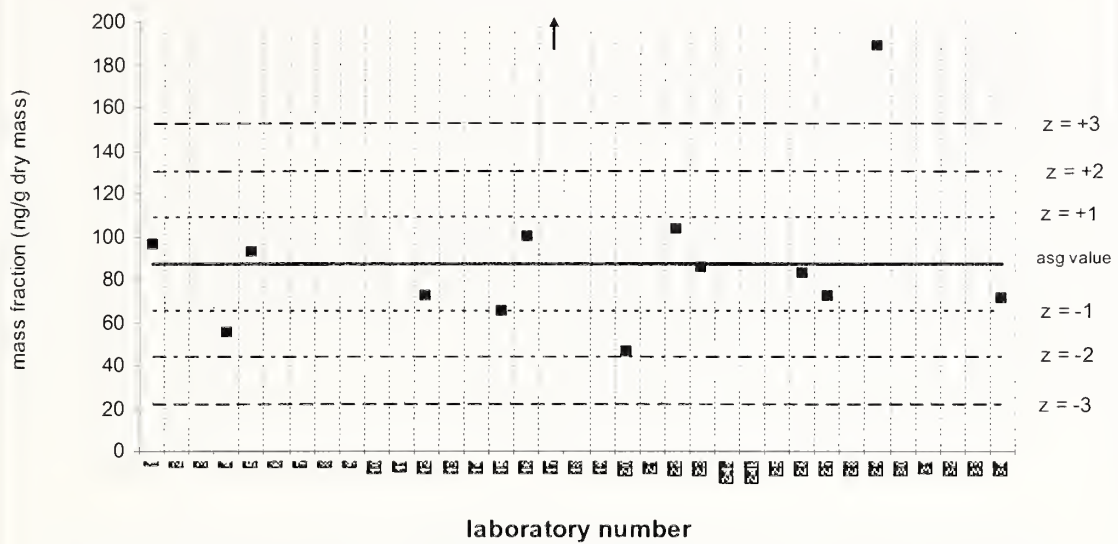
C1-dibenzothiophenes

QA10TIS01

Assigned value = 86.8 ng/g dry mass s = 35.0 ng/g dry mass 95% CI = 19.0 ng/g dry mass

Median value = 83.9 ng/g dry mass

Reported Results: 21 Quantitative Results: 14

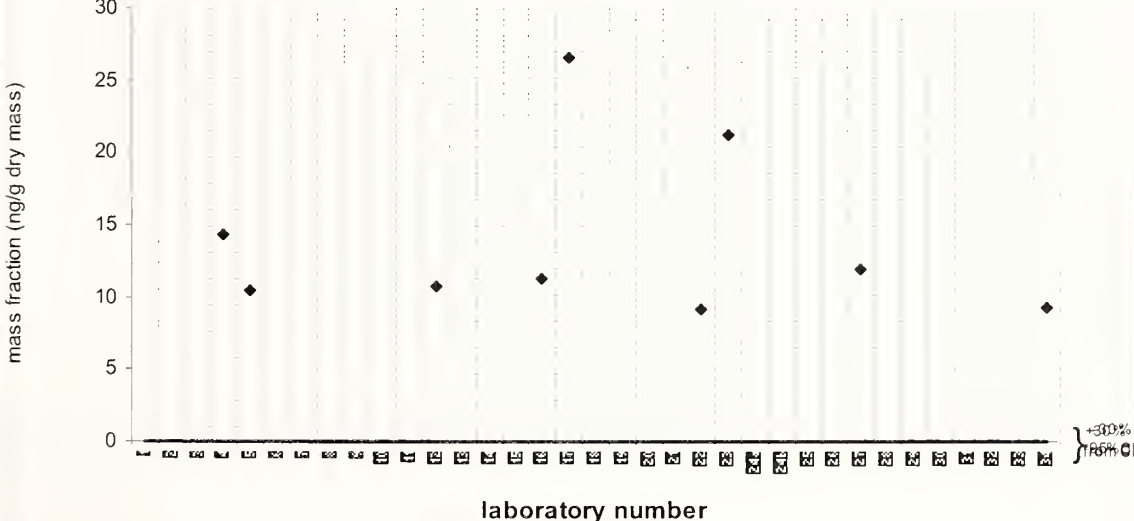


C1-dibenzothiophenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 11.3 ng/g dry mass

Reported Results: 16 Quantitative Results: 9



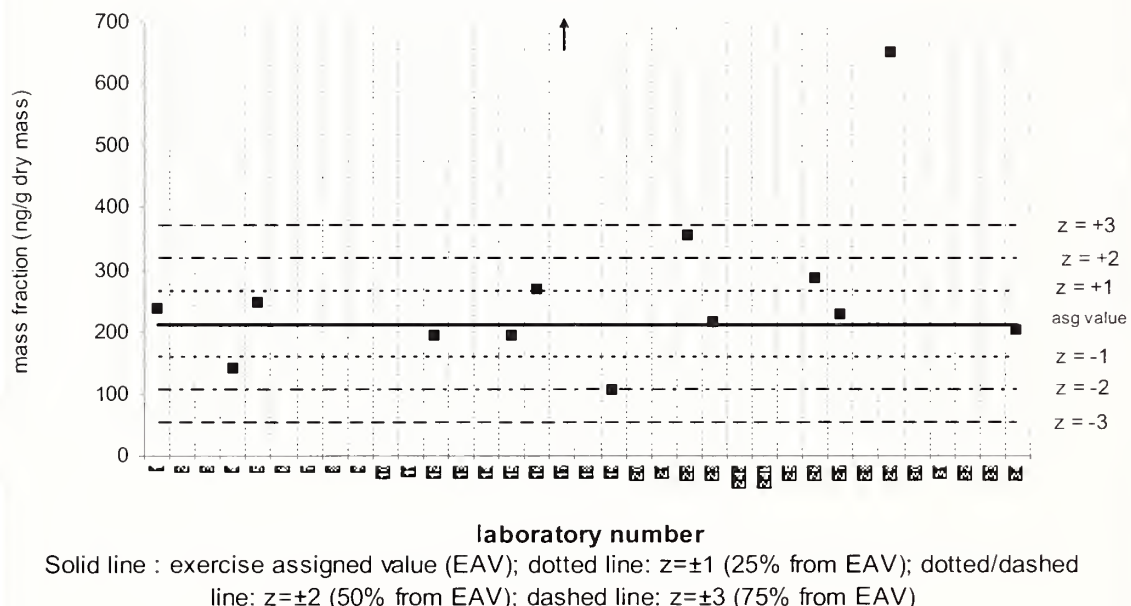
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C2-dibenzothiophenes

QA10TIS01

Assigned value = 211 ng/g dry mass $s = 134$ ng/g dry mass 95% CI = 73 ng/g dry mass Median value = 226 ng/g dry mass

Reported Results: 20 Quantitative Results: 15

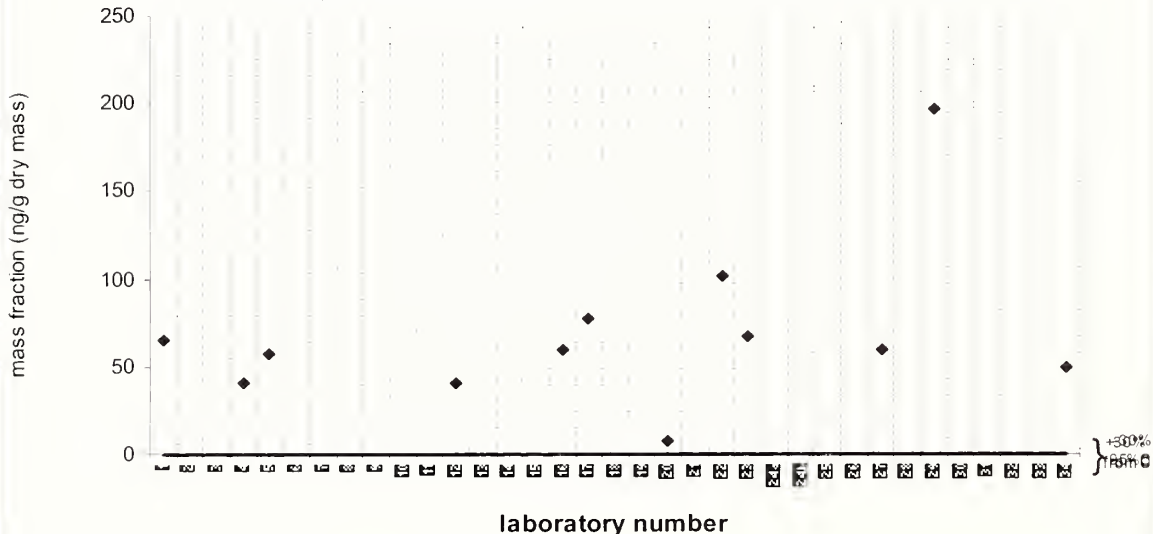


C2-dibenzothiophenes

SRM 1974b

Target Value = no target ng/g (dry mass): Median value = 59.6 ng/g dry mass

Reported Results: 17 Quantitative Results: 12



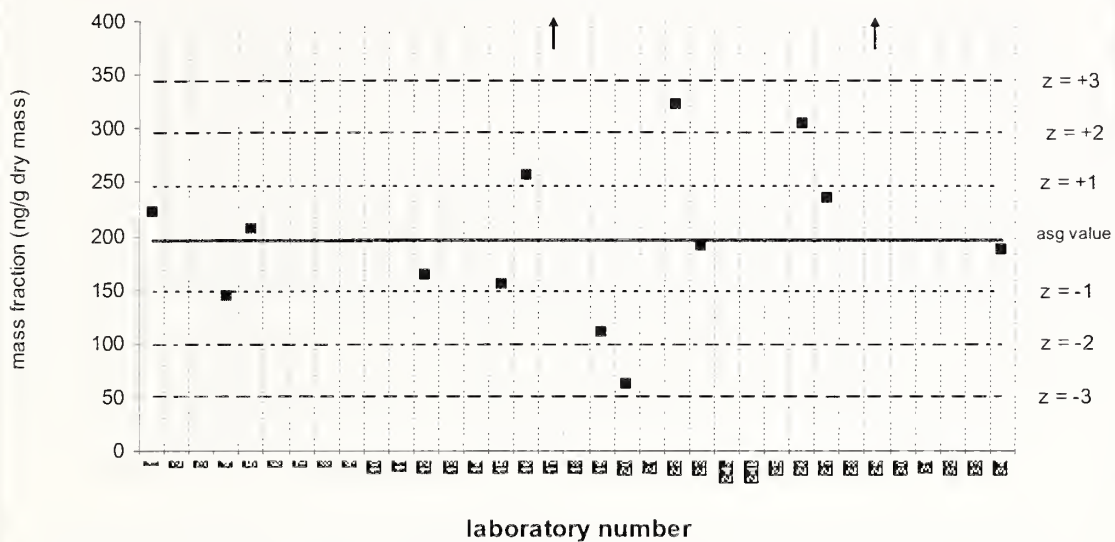
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C3-dibenzothiophenes

QA10TIS01

Assigned value = 196 ng/g dry mass s = 73 ng/g dry mass 95% CI = 40 ng/g dry mass Median value = 206 ng/g dry mass

Reported Results: 20 Quantitative Results: 15

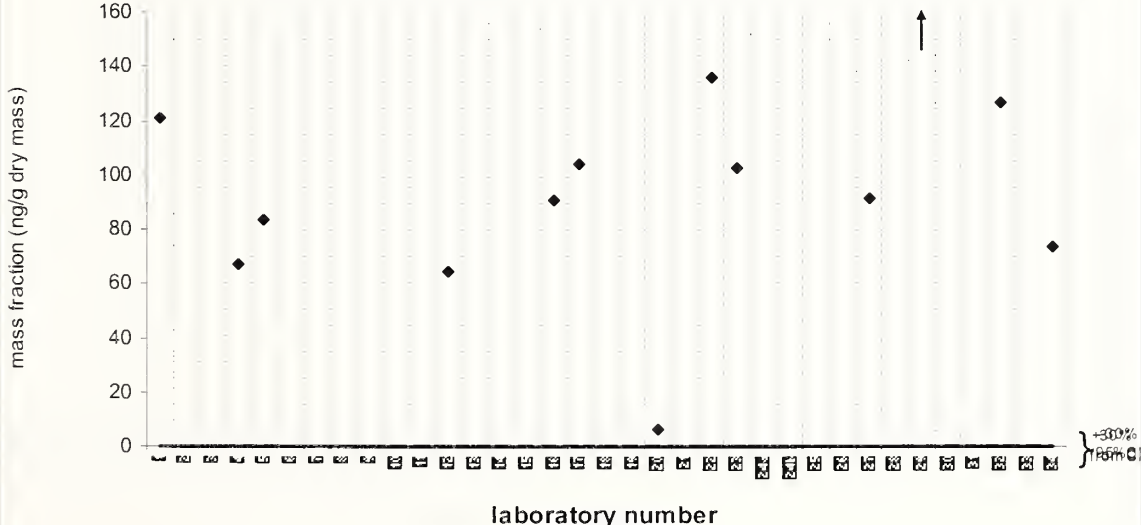


C3-dibenzothiophenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 91.0 ng/g dry mass

Reported Results: 17 Quantitative Results: 13

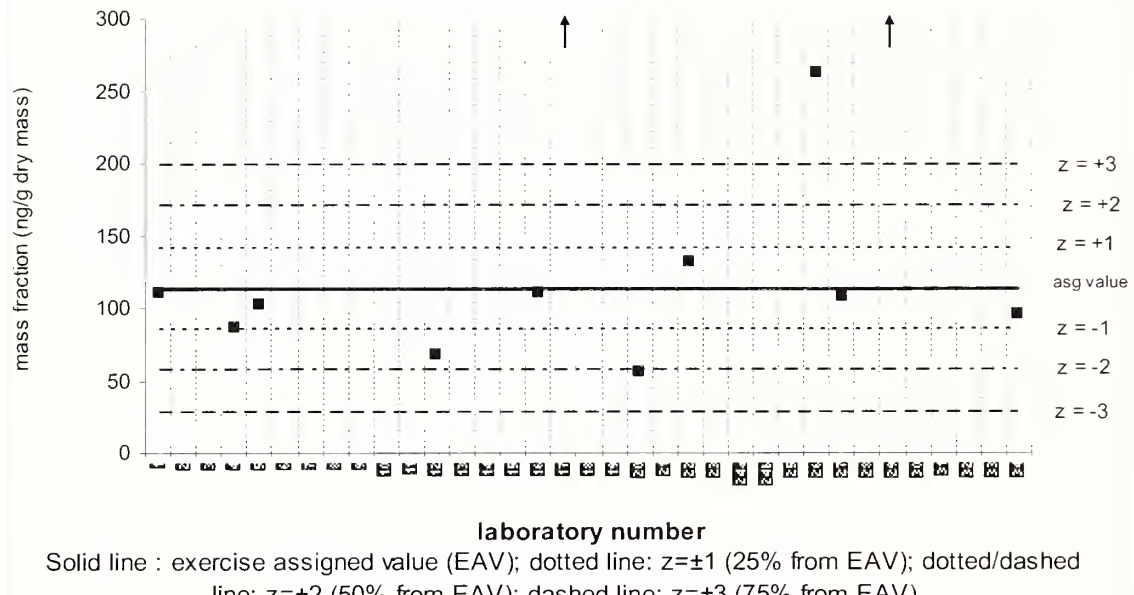


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C4-dibenzothiophenes**QA10TIS01**

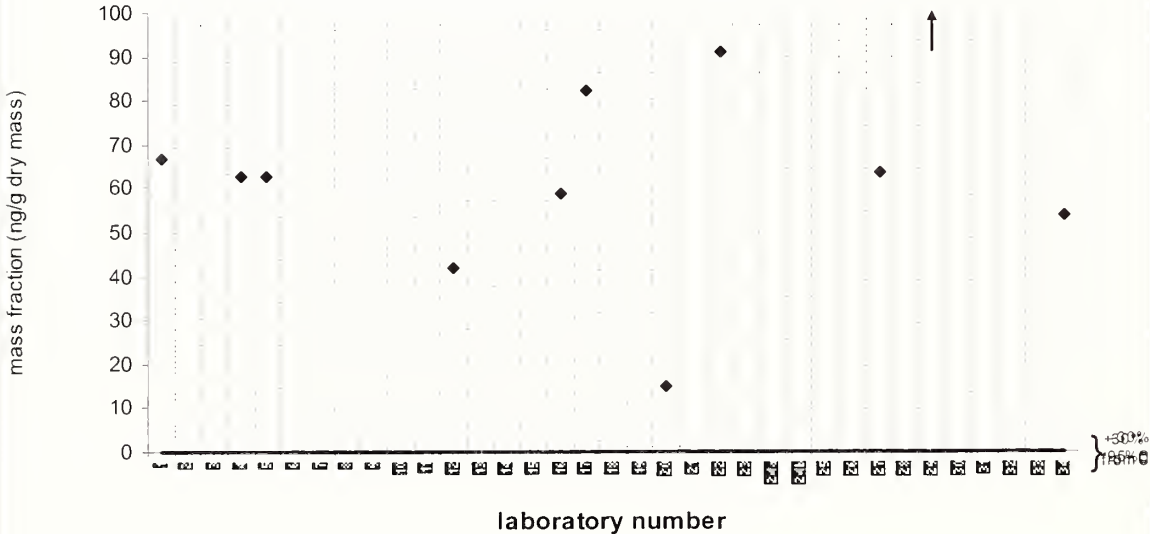
Assigned value = 113 ng/g dry mass s = 57 ng/g dry mass 95% CI = 35 ng/g dry mass Median value = 109 ng/g dry mass

Reported Results: 18 Quantitative Results: 12

**C4-dibenzothiophenes****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 63.0 ng/g dry mass

Reported Results: 15 Quantitative Results: 11



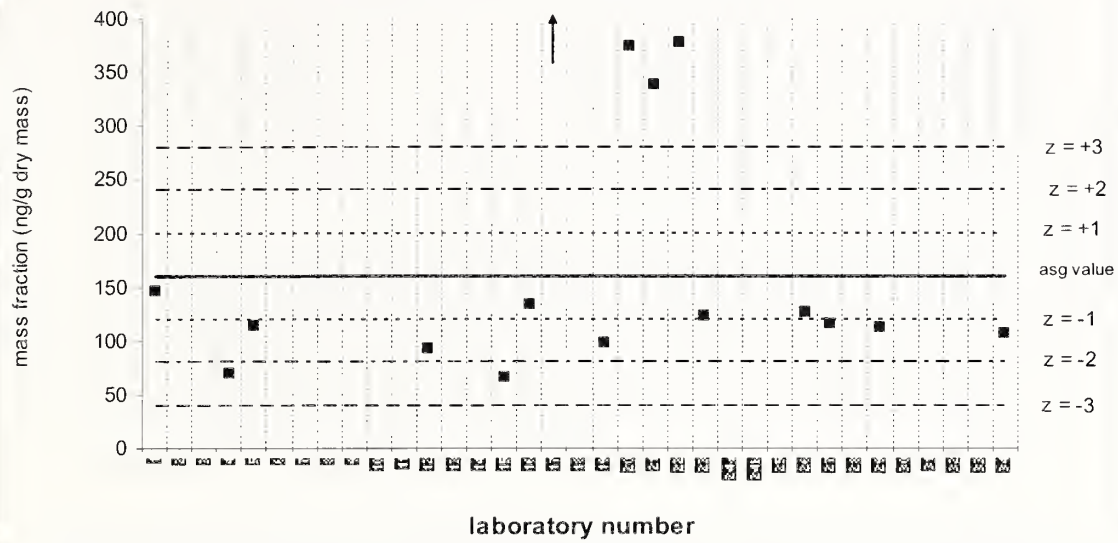
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C1-fluoranthenes/pyrenes

QA10TIS01

Assigned value = 159 ng/g dry mass s = 107 ng/g dry mass 95% CI = 54 ng/g dry mass Median value = 119 ng/g dry mass

Reported Results: 21 Quantitative Results: 16

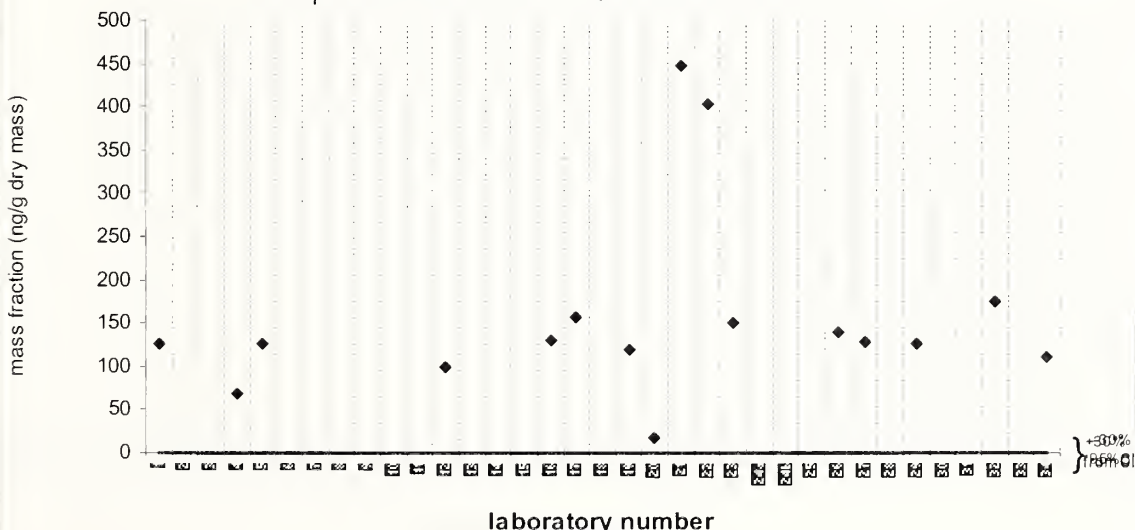


C1-fluoranthenes/pyrenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 128 ng/g dry mass

Reported Results: 18 Quantitative Results: 16



Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

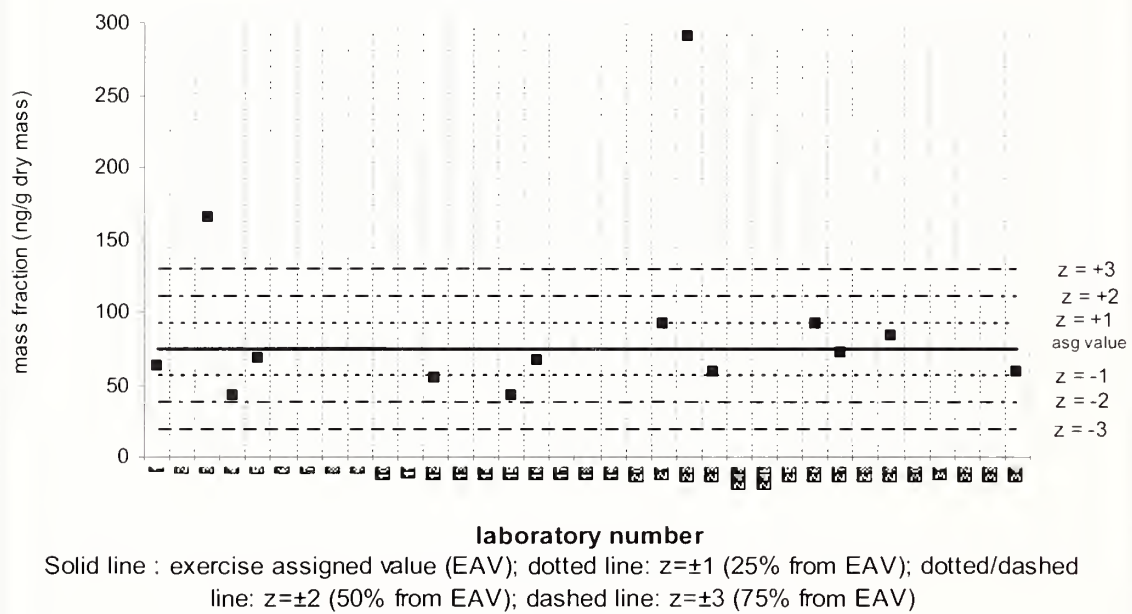
C2-fluoranthenes/pyrenes

QA10TIS01

Assigned value = 74.1 ng/g dry mass s = 31.8 ng/g dry mass 95% CI = 17.3 ng/g dry mass

Median value = 67.7 ng/g dry mass

Reported Results: 21 Quantitative Results: 14

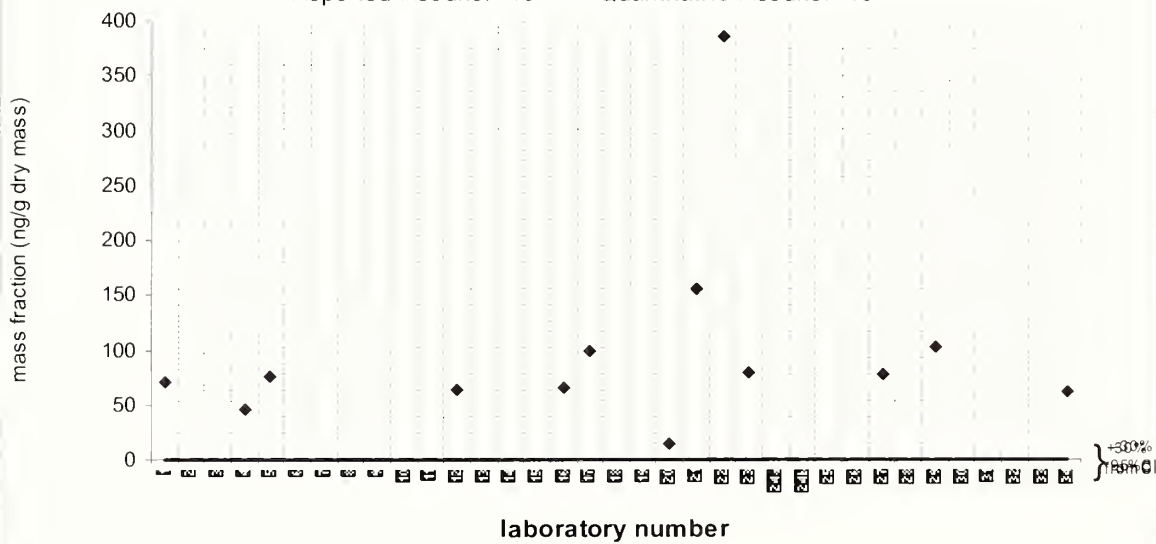


C2-fluoranthenes/pyrenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 75.5 ng/g dry mass

Reported Results: 18 Quantitative Results: 13



Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

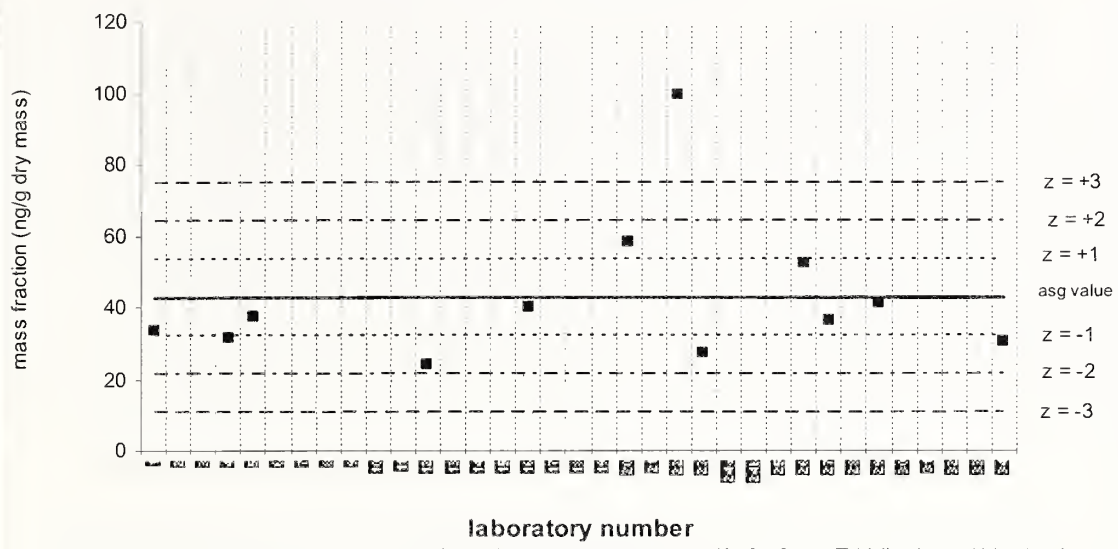
C3-fluoranthenes/pyrenes

QA10TIS01

Assigned value = 42.5 ng/g dry mass s = 20.3 ng/g dry mass 95% CI = 11.5 ng/g dry mass

Median value = 36.7 ng/g dry mass

Reported Results: 20 Quantitative Results: 12

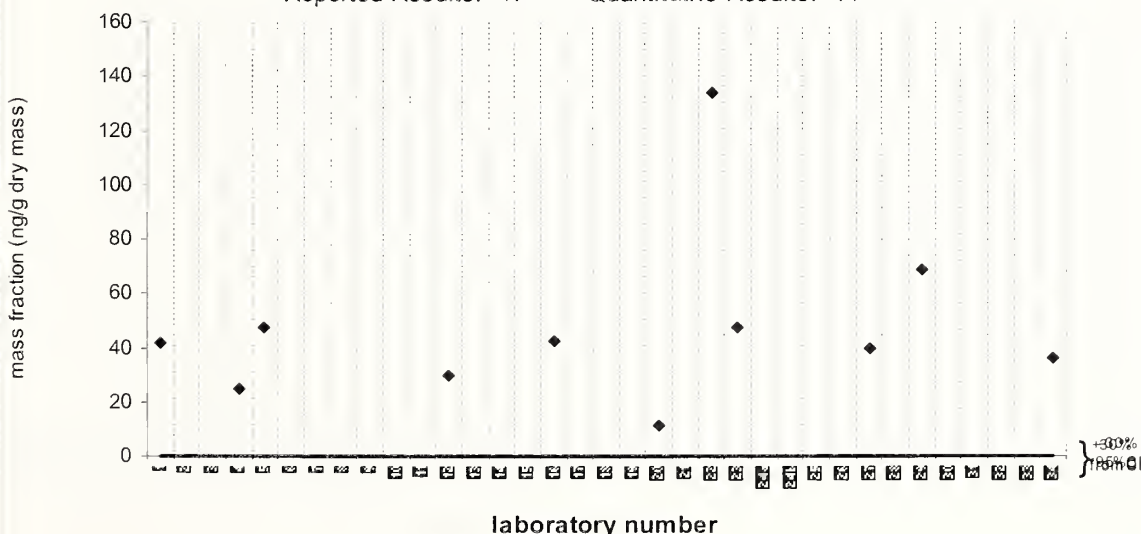


C3-fluoranthenes/pyrenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 42.0 ng/g dry mass

Reported Results: 17 Quantitative Results: 11

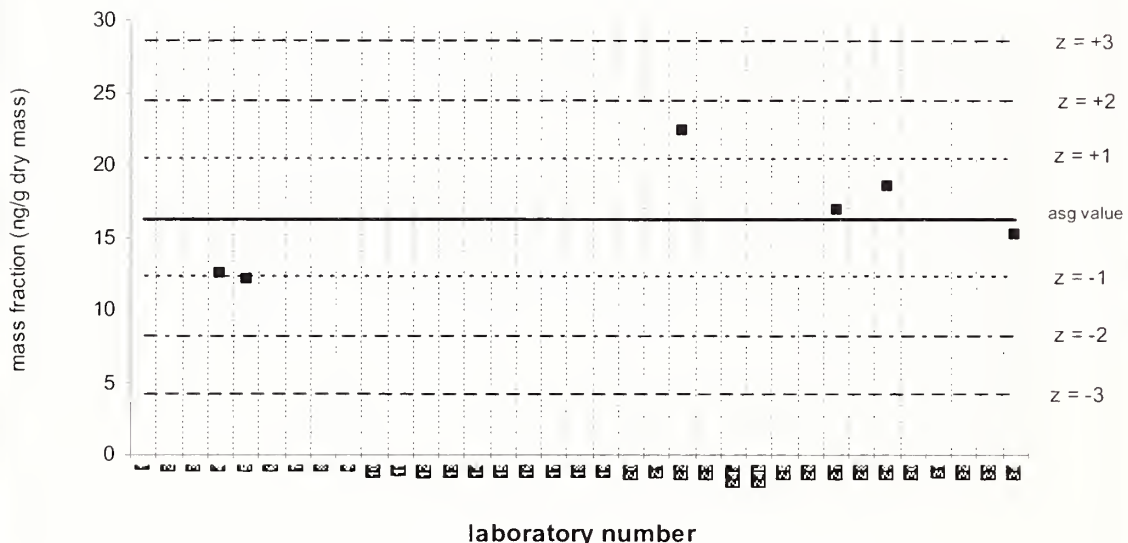


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C4-fluoranthenes/pyrenes**QA10TIS01**

Assigned value = 16.3 ng/g dry mass s = 3.9 ng/g dry mass 95% CI = 3.1 ng/g dry mass Median value = 16.0 ng/g dry mass

Reported Results: 11 Quantitative Results: 6

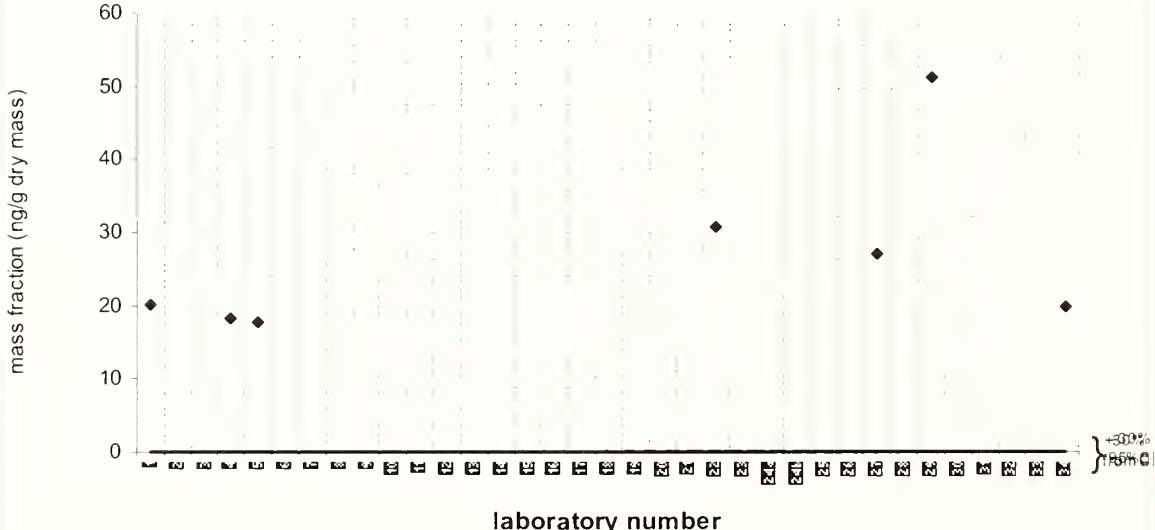


Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

C4-fluoranthenes/pyrenes**SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 20.1 ng/g dry mass

Reported Results: 10 Quantitative Results: 7



Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

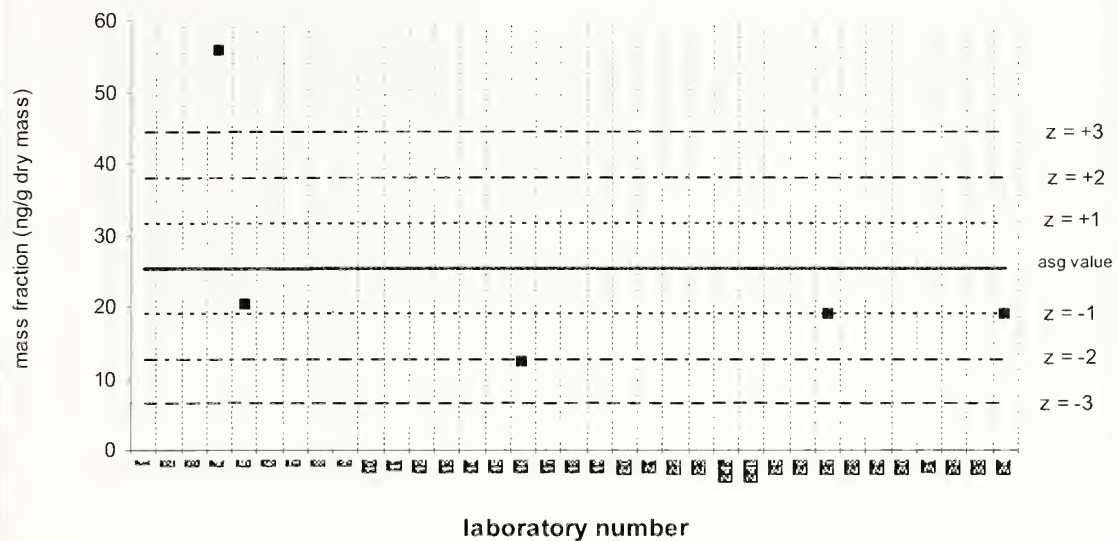
C1-naphthobenzothiophenes

QA10TIS01

Assigned value = 25.2 ng/g dry mass $s = 17.3$ ng/g dry mass 95% CI = 15.2 ng/g dry mass

Median value = 19.1 ng/g dry mass

Reported Results: 10 Quantitative Results: 5



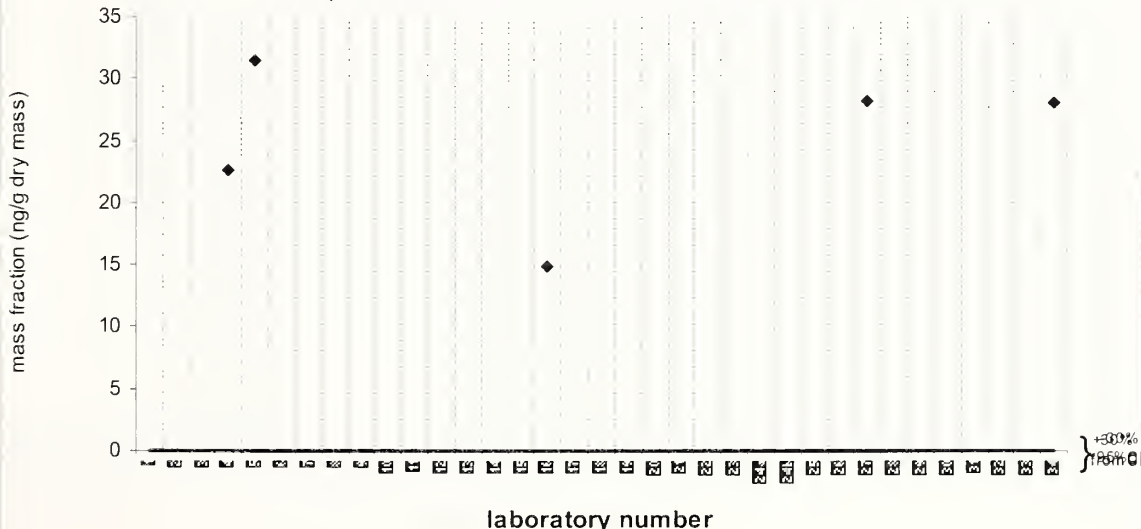
Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

C1-naphthobenzothiophenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 28.1 ng/g dry mass

Reported Results: 9 Quantitative Results: 5



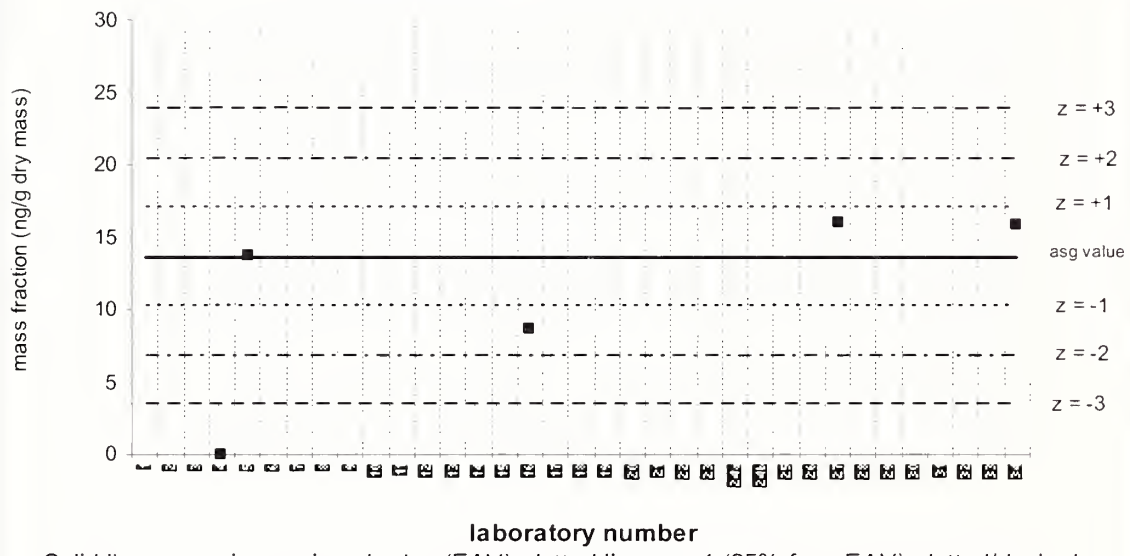
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C2-naphthobenzothiophenes

QA10TIS01

Assigned value = 13.6 ng/g dry mass s = 3.4 ng/g dry mass 95% CI = 3.3 ng/g dry mass Median value = 13.8 ng/g dry mass

Reported Results: 10 Quantitative Results: 5

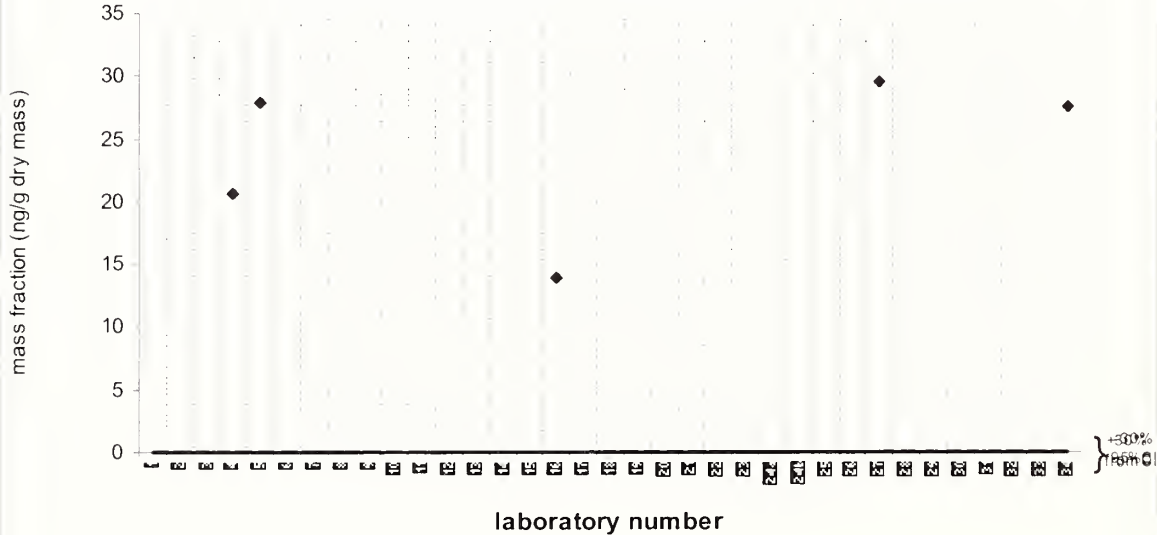


C2-naphthobenzothiophenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 27.6 ng/g dry mass

Reported Results: 9 Quantitative Results: 5



Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

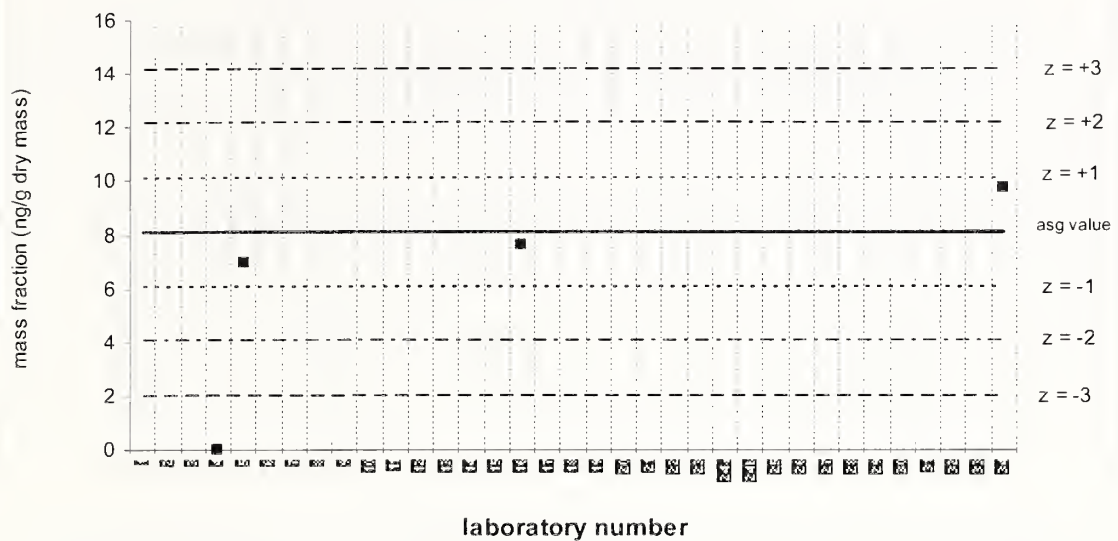
C3-naphthobenzothiophenes

QA10TIS01

Assigned value = 8.11 ng/g dry mass s = 1.46 ng/g dry mass 95% CI = 1.65 ng/g dry mass

Median value = 7.28 ng/g dry mass

Reported Results: 10 Quantitative Results: 4

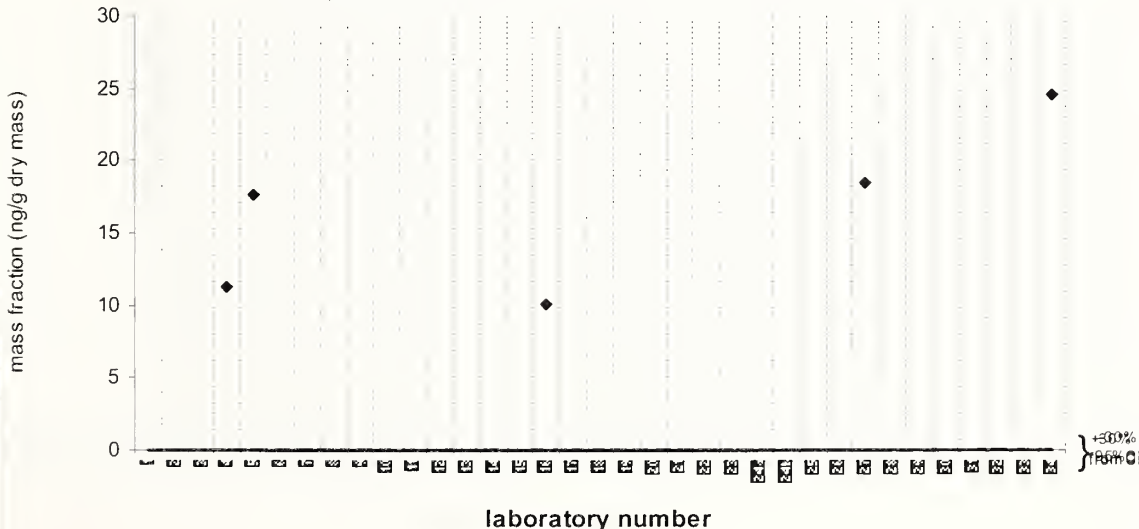


C3-naphthobenzothiophenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 17.7 ng/g dry mass

Reported Results: 9 Quantitative Results: 5

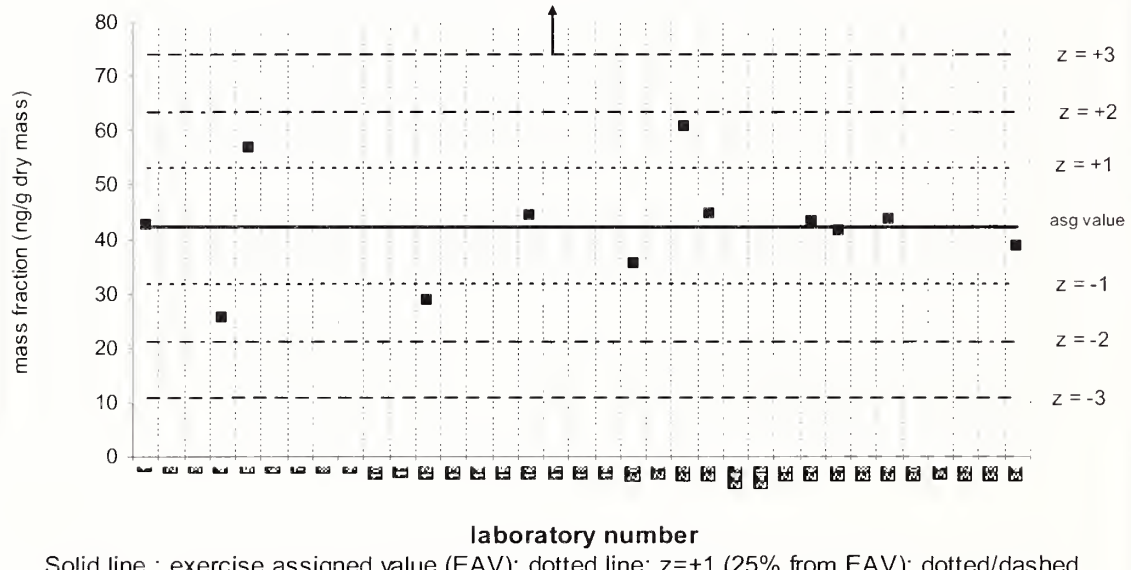


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C1-chrysenes**QA10TIS01**

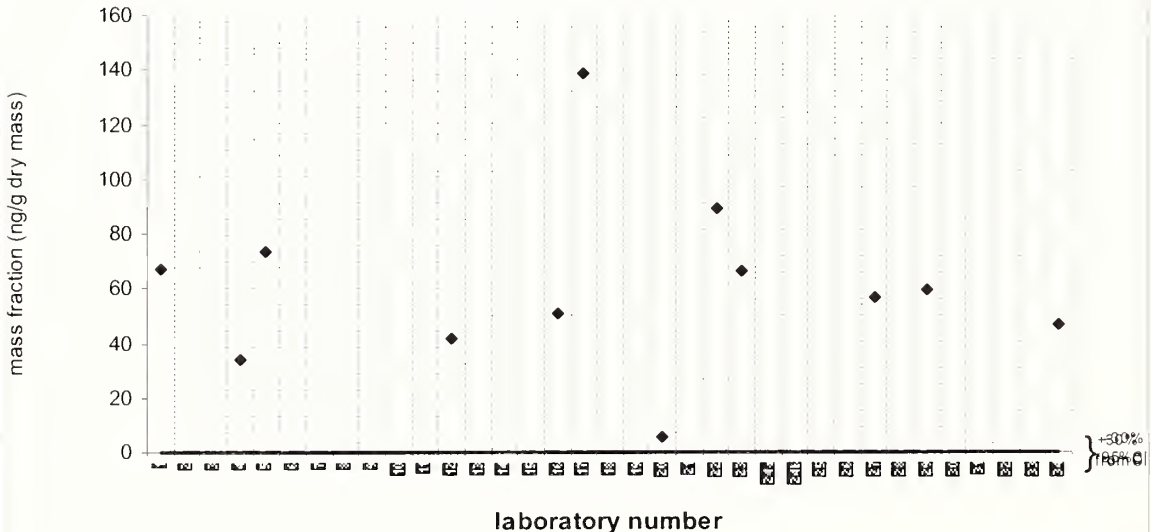
Assigned value = 42.3 ng/g dry mass s = 9.9 ng/g dry mass 95% CI = 5.6 ng/g dry mass Median value = 43.5 ng/g dry mass

Reported Results: 19 Quantitative Results: 13

**C1-chrysenes****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 53.8 ng/g dry mass

Reported Results: 17 Quantitative Results: 12

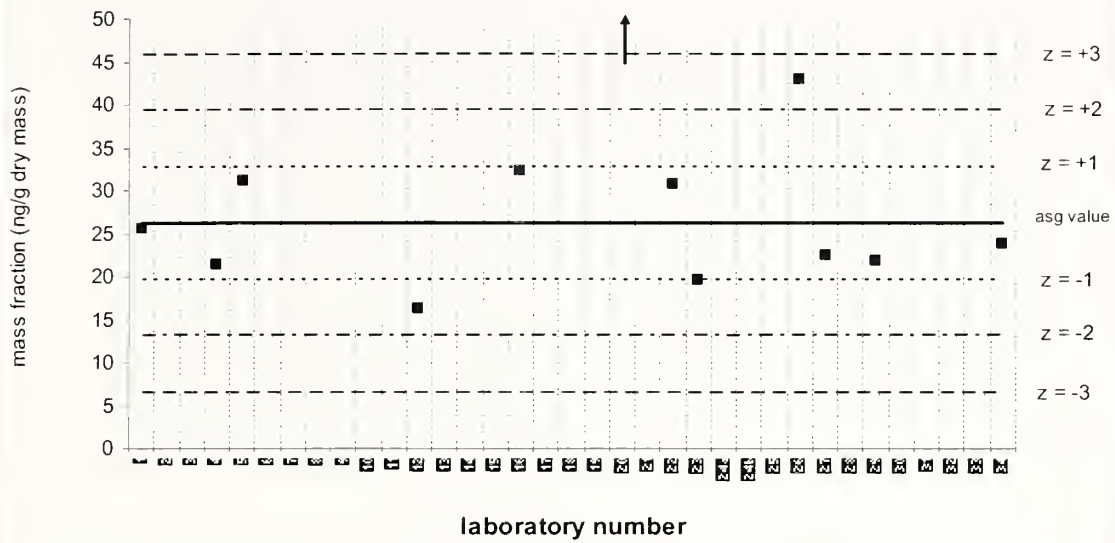


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C2-chrysenes**QA10TIS01**

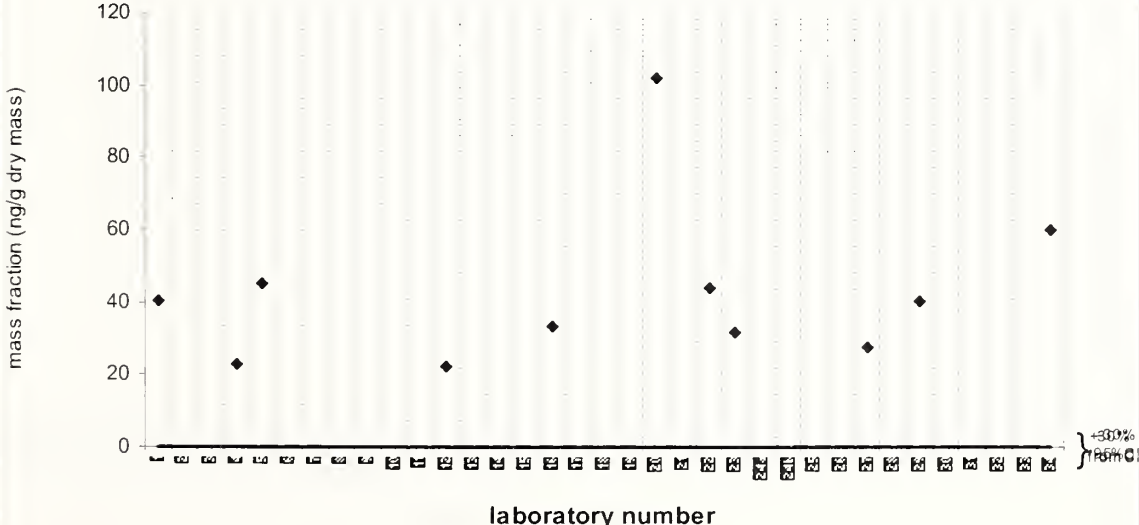
Assigned value = 26.2 ng/g dry mass s = 7.6 ng/g dry mass 95% CI = 4.5 ng/g dry mass Median value = 24.8 ng/g dry mass

Reported Results: 19 Quantitative Results: 12

**C2-chrysenes****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 40.2 ng/g dry mass

Reported Results: 17 Quantitative Results: 11



Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

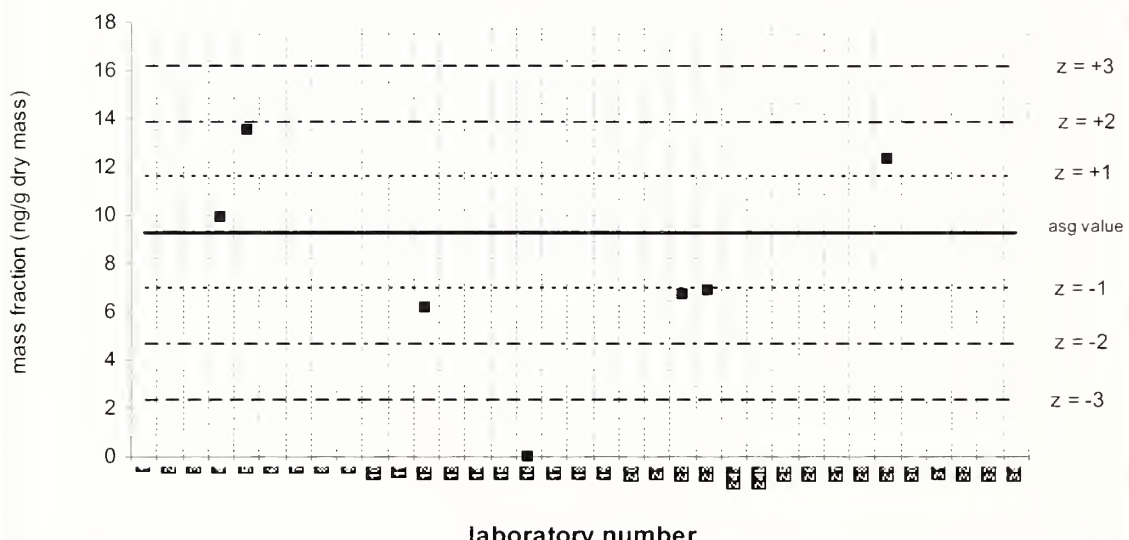
C3-chrysenes

QA10TIS01

Assigned value = 9.25 ng/g dry mass $s = 3.14$ ng/g dry mass 95% CI = 2.51 ng/g dry mass

Median value = 6.84 ng/g dry mass

Reported Results: 19 Quantitative Results: 7

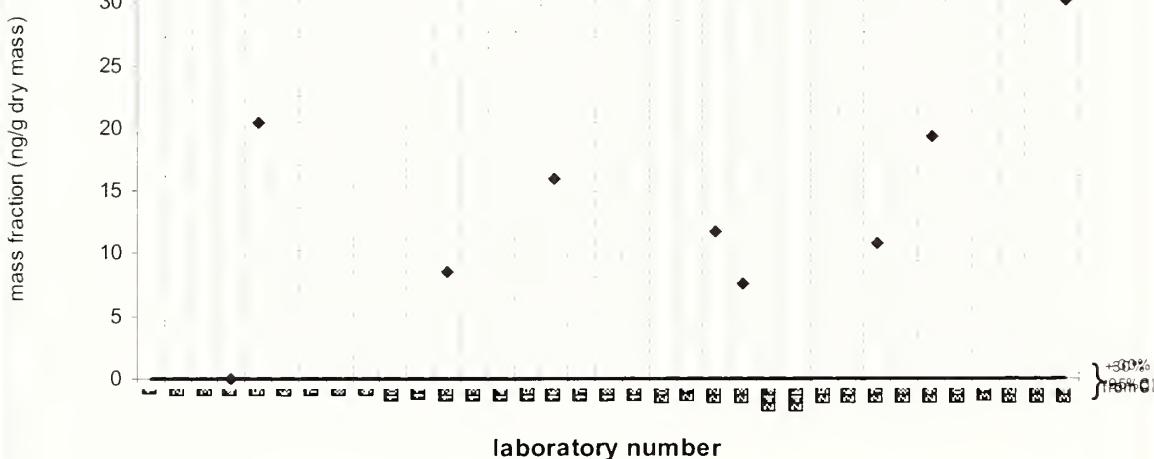


C3-chrysenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 11.7 ng/g dry mass

Reported Results: 17 Quantitative Results: 9



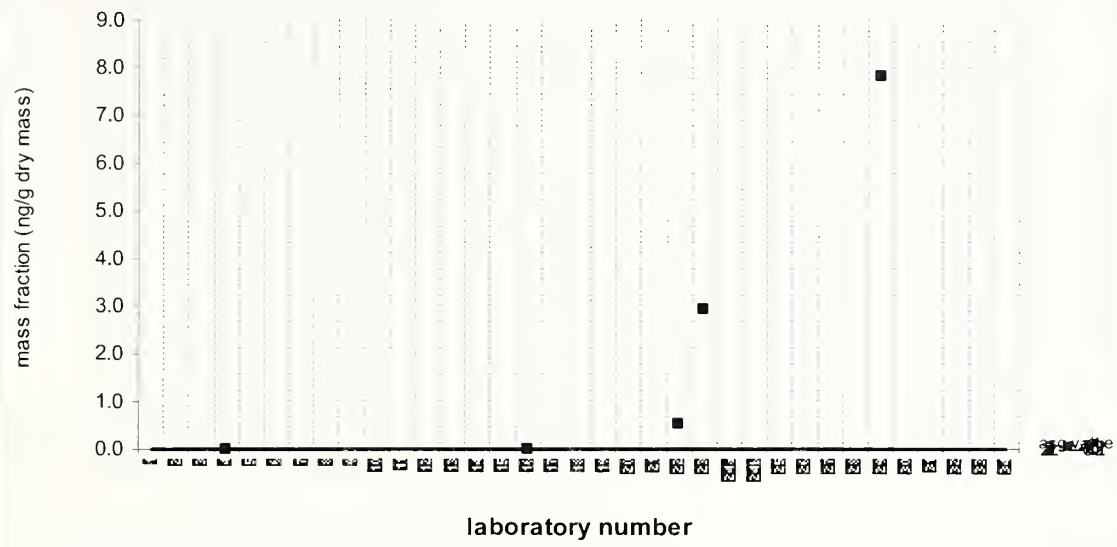
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

C4-chrysenes

QA10TIS01

Assigned value = No Target ng/g (dry mass) Median value = 0.54 ng/g dry mass

Reported Results: 18 Quantitative Results: 5



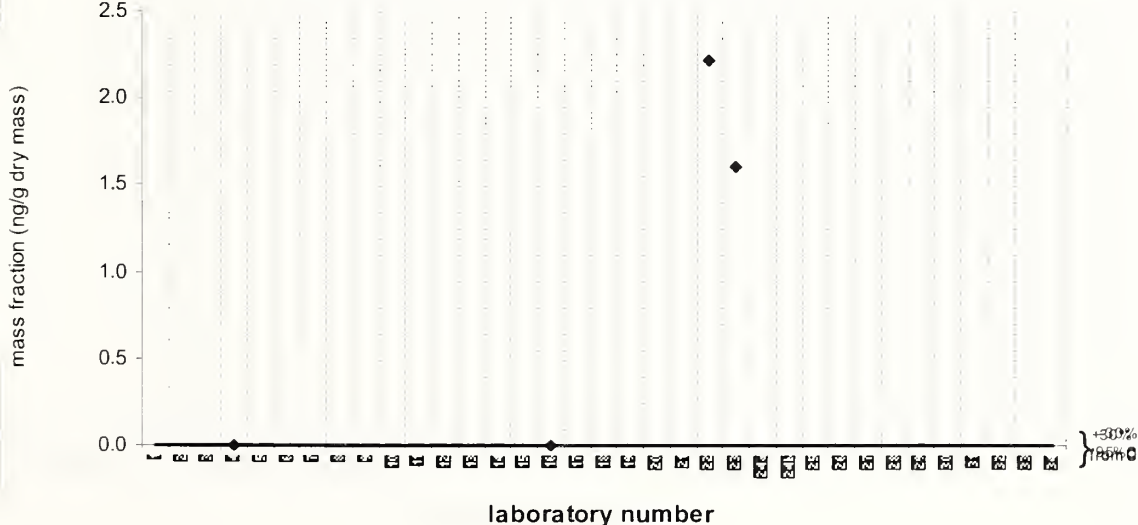
Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

C4-chrysenes

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 0.80 ng/g dry mass

Reported Results: 16 Quantitative Results: 4



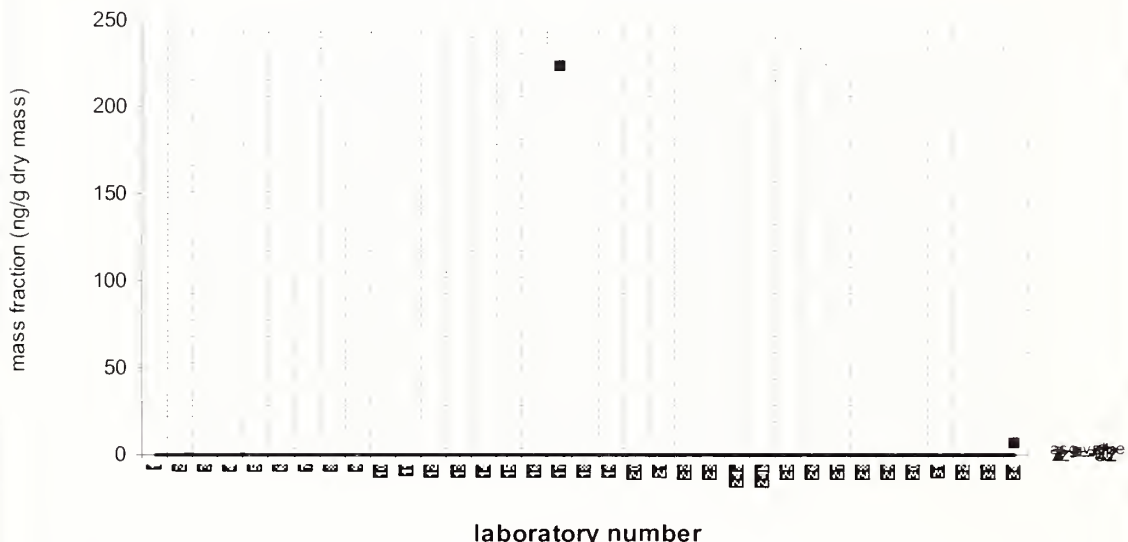
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

Carbazole

QA10TIS01

Assigned value = No Target ng/g (dry mass) Median value = 115 ng/g dry mass

Reported Results: 6 Quantitative Results: 2



laboratory number

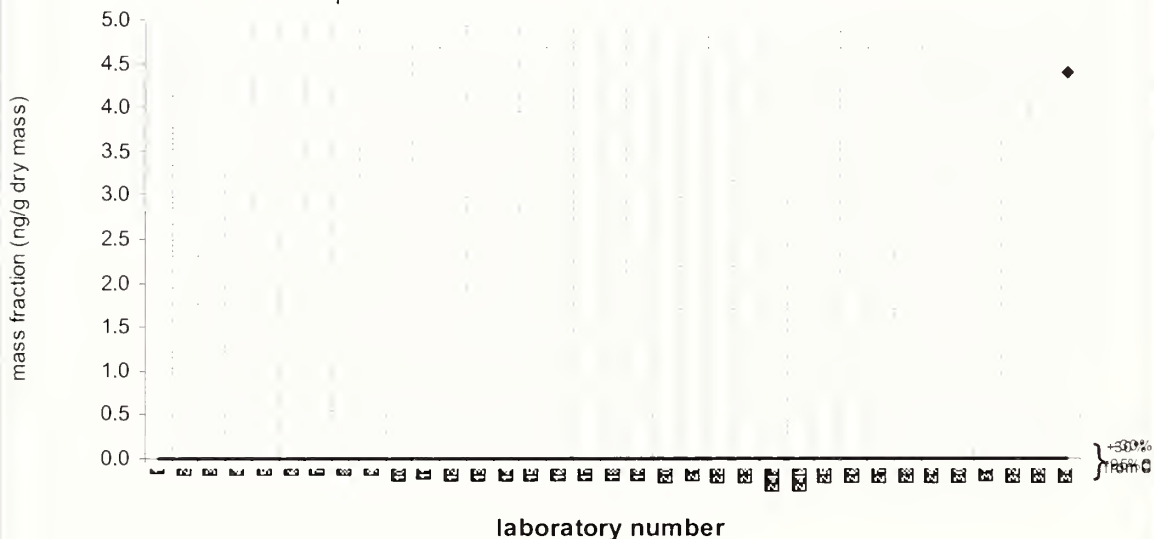
Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

Carbazole

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 4.40 ng/g dry mass

Reported Results: 6 Quantitative Results: 1



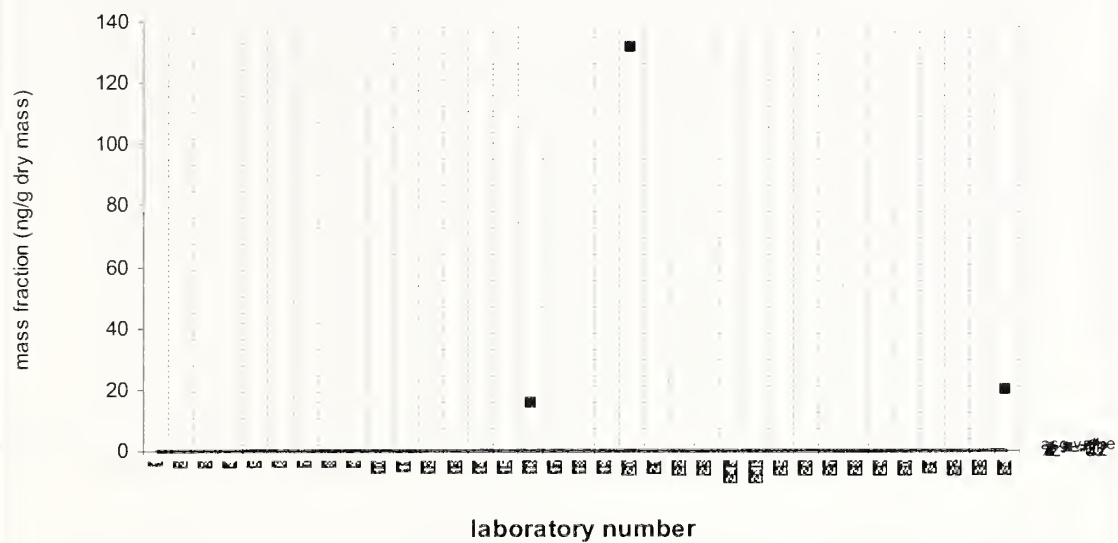
laboratory number

Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

18a(H)-22,29,30-Trisnorhopane**QA10TIS01**

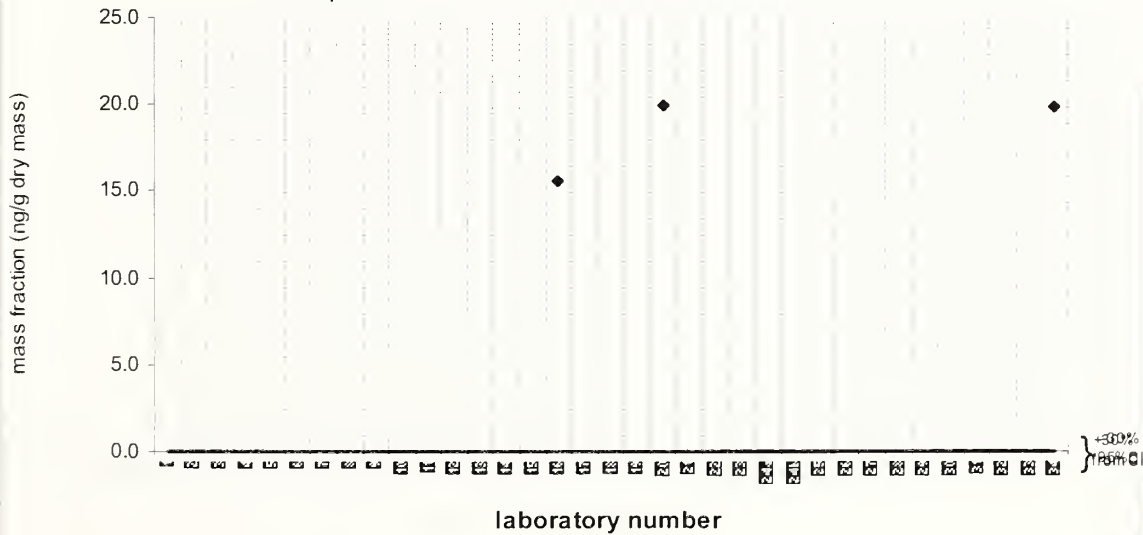
Assigned value = No Target ng/g (dry mass) Median value = 19.8 ng/g dry mass

Reported Results: 12 Quantitative Results: 3

**18a(H)-22,29,30-Trisnorhopane****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 19.8 ng/g dry mass

Reported Results: 7 Quantitative Results: 3

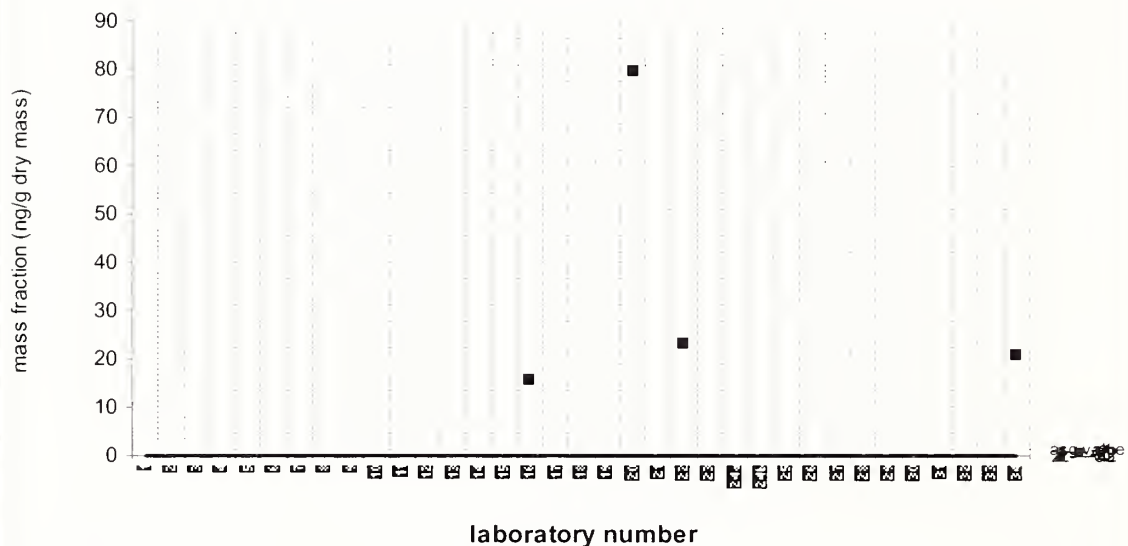


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

17a(H)-22,29,30-Trisnorhopane**QA10TIS01**

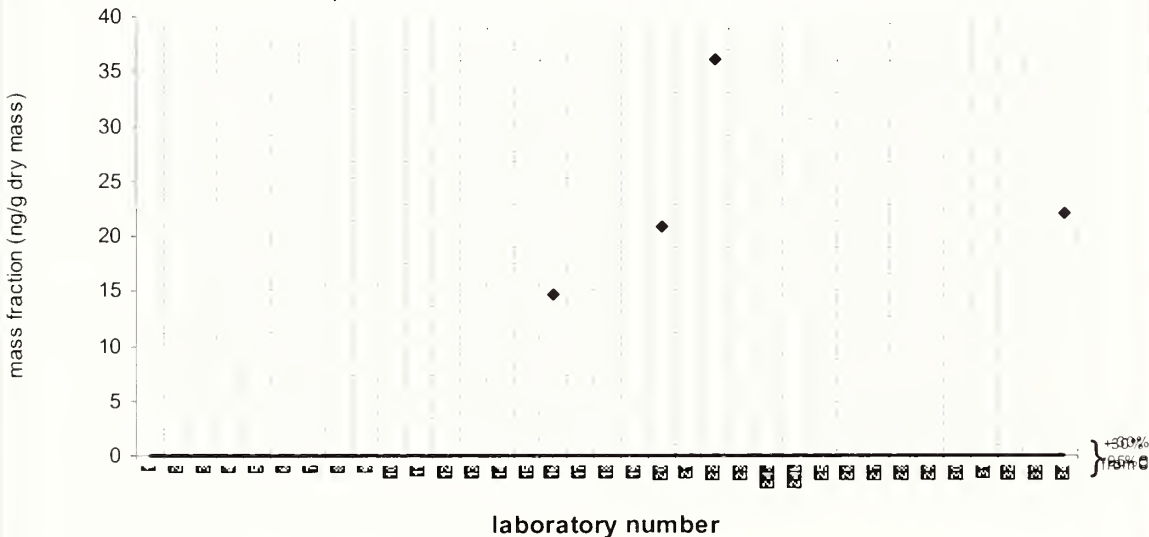
Assigned value = No Target ng/g (dry mass) Median value = 22.0 ng/g dry mass

Reported Results: 11 Quantitative Results: 4

**17a(H)-22,29,30-Trisnorhopane****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 21.5 ng/g dry mass

Reported Results: 9 Quantitative Results: 4



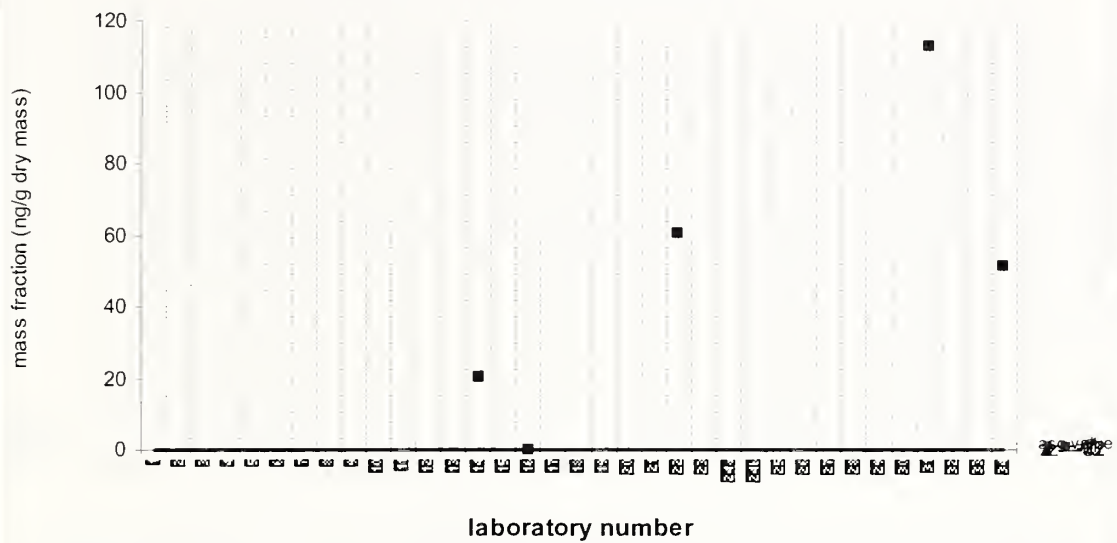
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

17 α (H),21 β (H)-30-Norhopane

QA10TIS01

Assigned value = No Target ng/g (dry mass) Median value = 51.2 ng/g dry mass

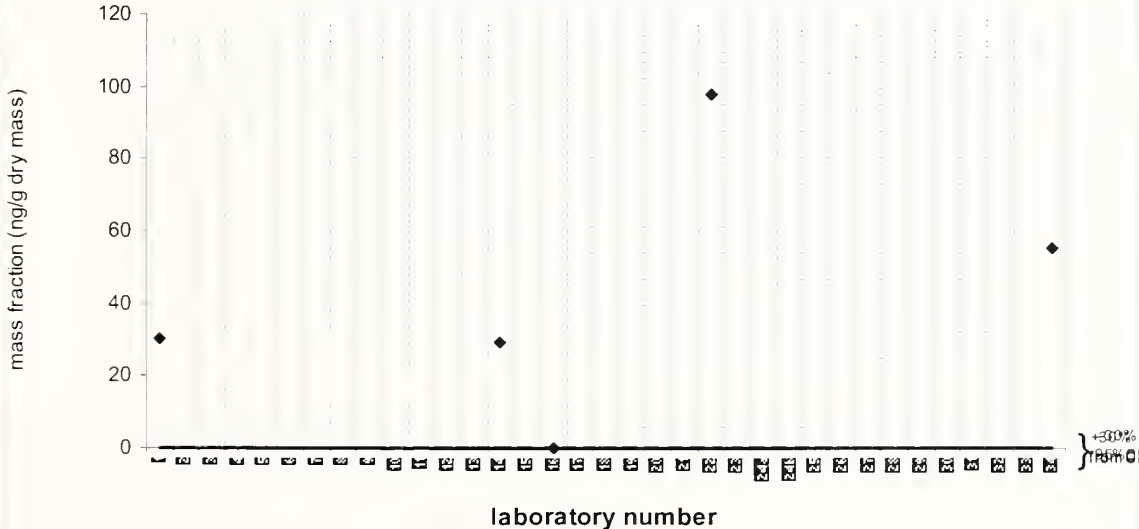
Reported Results: 12 Quantitative Results: 5

**17 α (H),21 β (H)-30-Norhopane**

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 30.4 ng/g dry mass

Reported Results: 10 Quantitative Results: 5

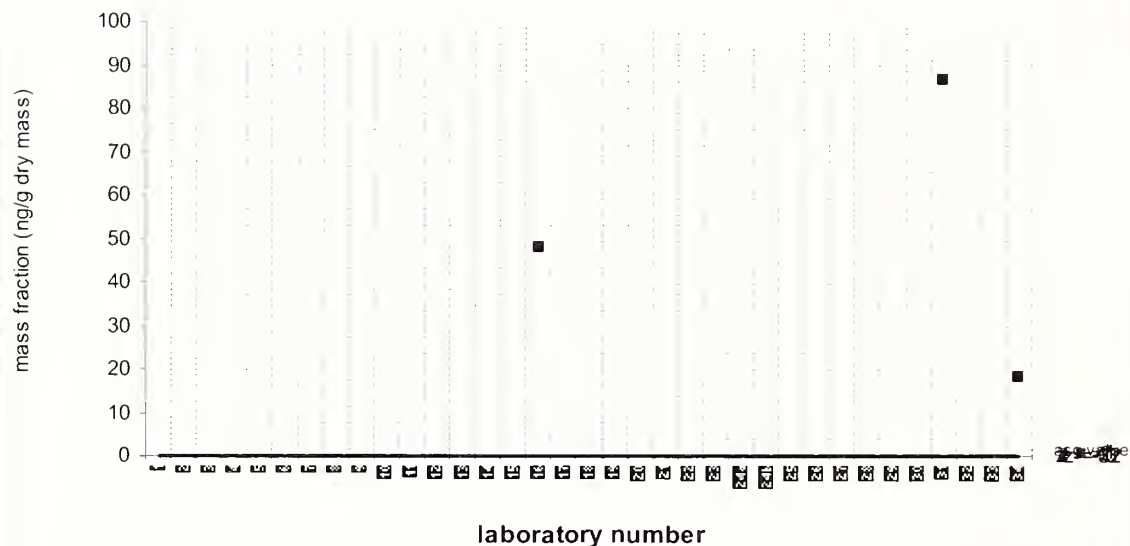


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

18a(H)-30-Norneohopane**QA10TIS01**

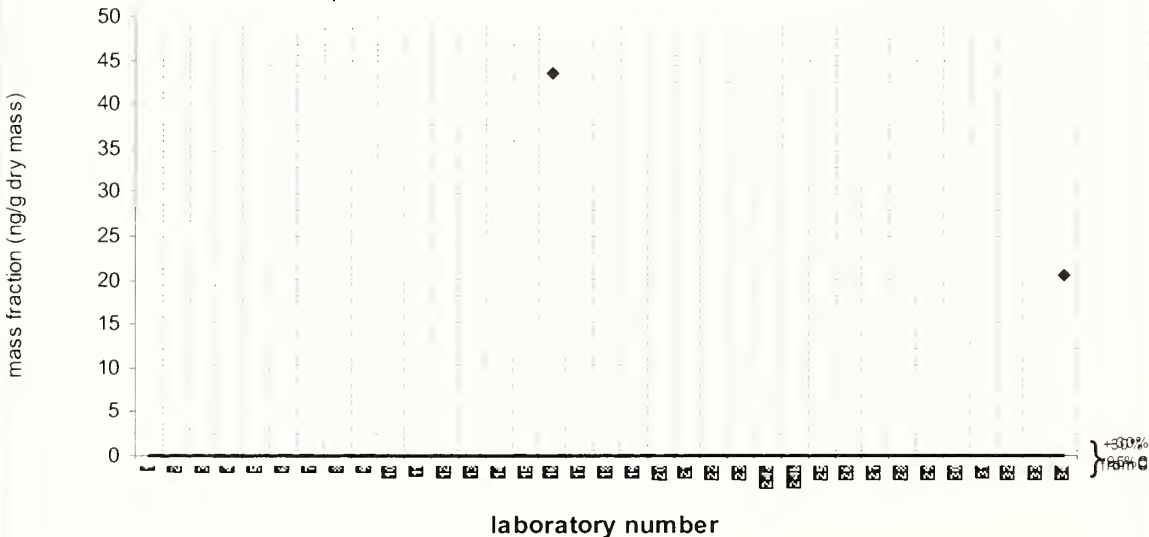
Assigned value = No Target ng/g (dry mass) Median value = 48.2 ng/g dry mass

Reported Results: 9 Quantitative Results: 3

**laboratory number**Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)**18a(H)-30-Norneohopane****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 32.1 ng/g dry mass

Reported Results: 7 Quantitative Results: 2

**laboratory number**

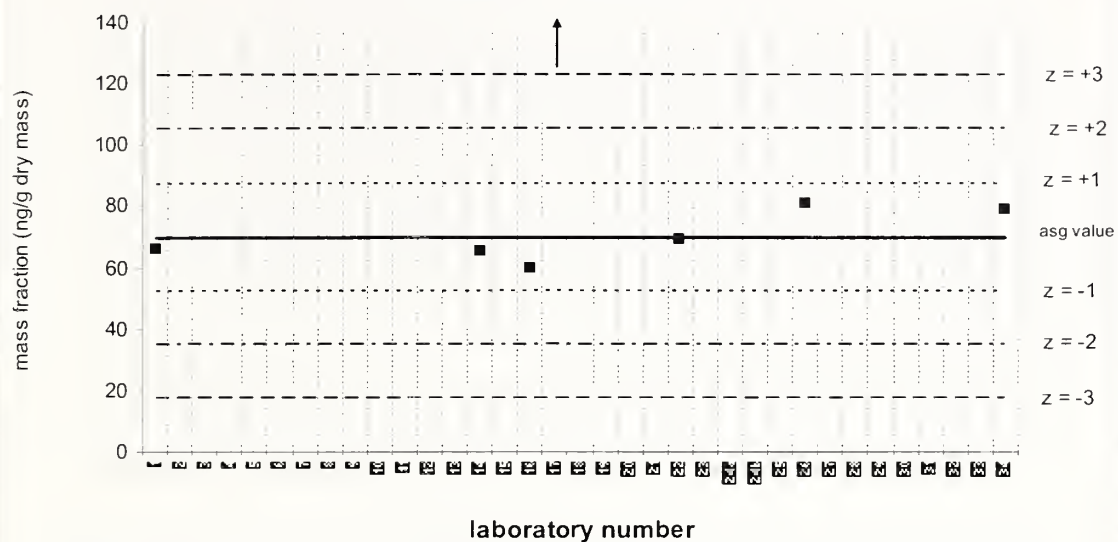
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

17 α (H),21 β (H)-Hopane

QA10TIS01

Assigned value = 69.9 ng/g dry mass s = 8.3 ng/g dry mass 95% CI = 6.7 ng/g dry mass Median value = 68.8 ng/g dry mass

Reported Results: 14 Quantitative Results: 7

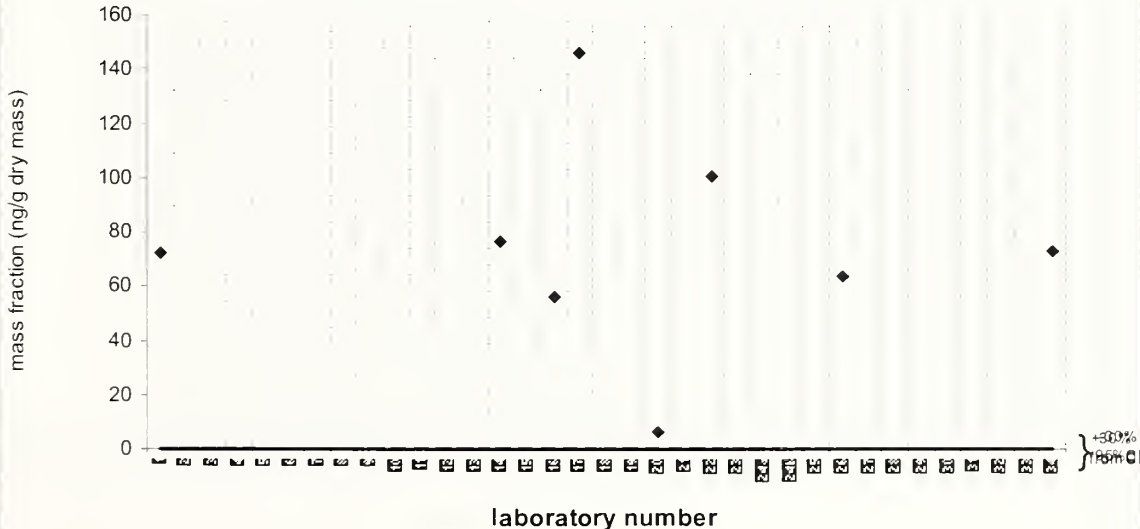


17 α (H),21 β (H)-Hopane

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 72.8 ng/g dry mass

Reported Results: 12 Quantitative Results: 8

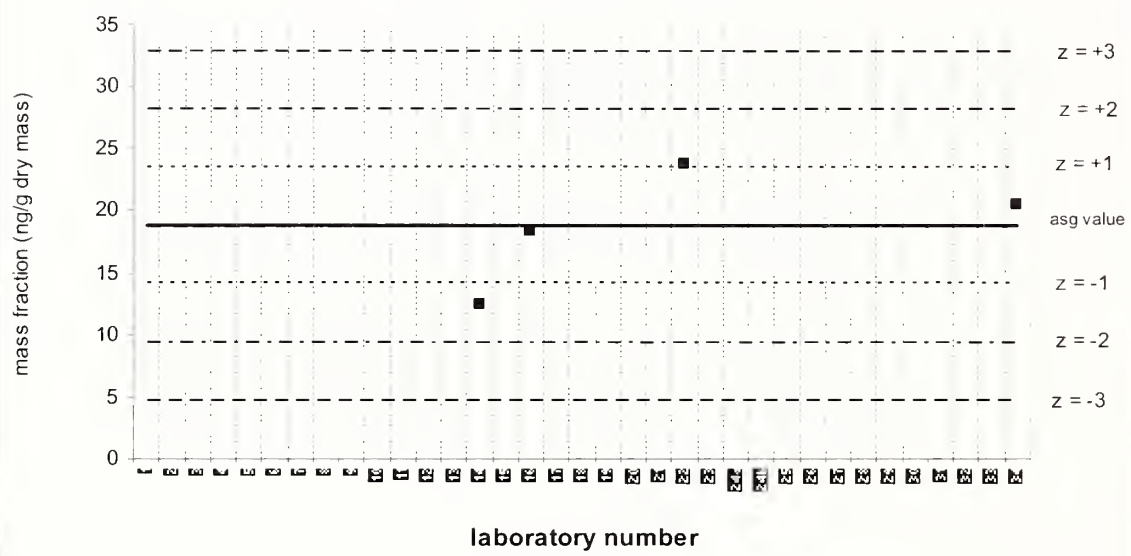


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

17 α (H),21 β (H)-22R-Homohopane**QA10TIS01**

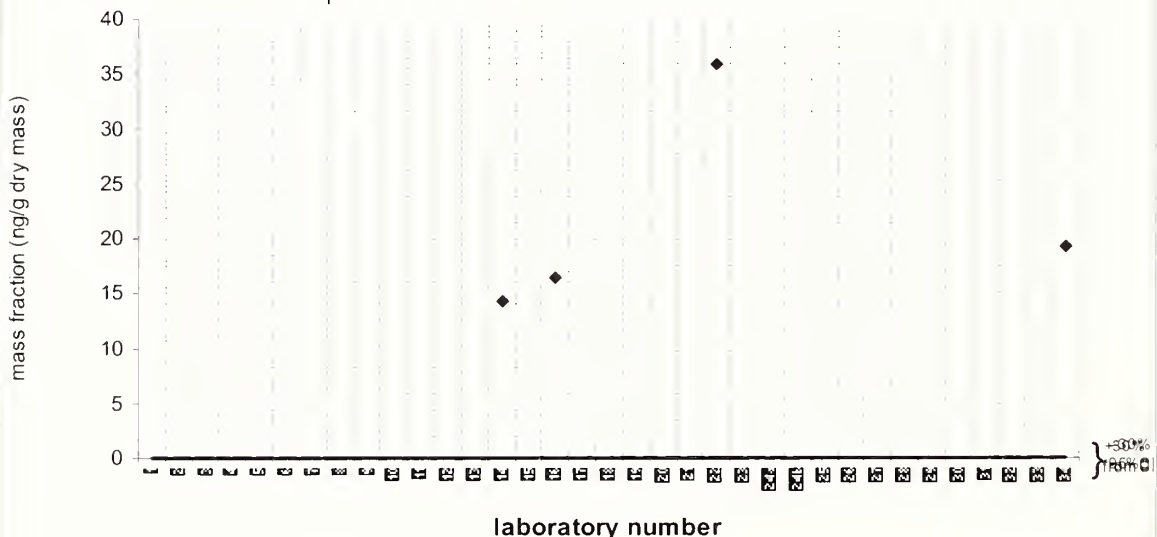
Assigned value = 18.8 ng/g dry mass s = 4.8 ng/g dry mass 95% CI = 4.7 ng/g dry mass Median value = 19.4 ng/g dry mass

Reported Results: 12 Quantitative Results: 4

**17 α (H),21 β (H)-22R-Homohopane****SRM 1974b**

Target Value = no target ng/g (dry mass): Median value = 17.9 ng/g dry mass

Reported Results: 10 Quantitative Results: 4



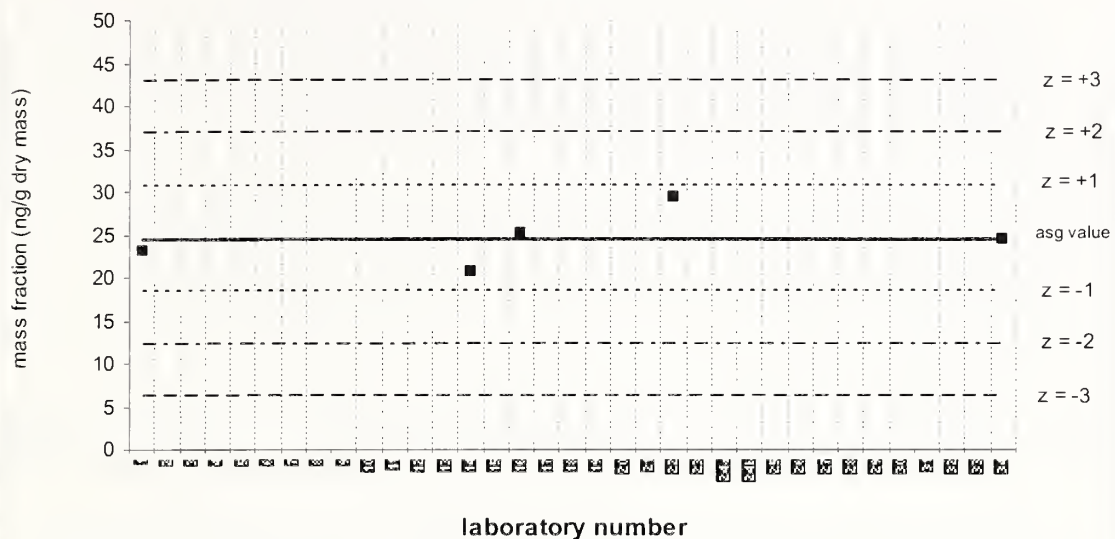
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

17 α (H),21 β (H)-22S-Homohopane

QA10TIS01

Assigned value = 24.5 ng/g dry mass s = 3.2 ng/g dry mass 95% CI = 2.8 ng/g dry mass Median value = 24.4 ng/g dry mass

Reported Results: 12 Quantitative Results: 5



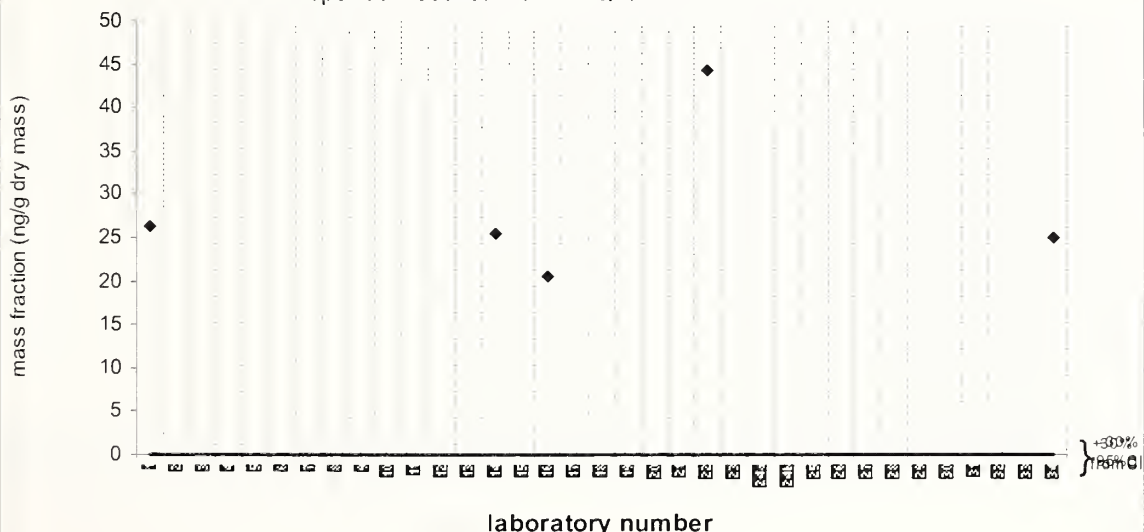
Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

17 α (H),21 β (H)-22S-Homohopane

SRM 1974b

Target Value = no target ng/g (dry mass); Median value = 25.4 ng/g dry mass

Reported Results: 10 Quantitative Results: 5

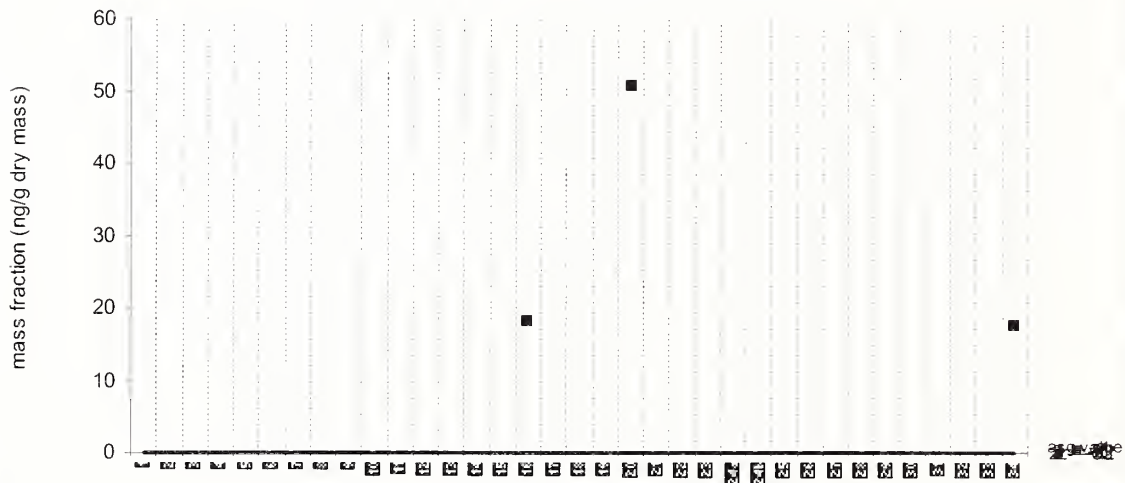


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

13b(H)17a(H)-Diacholestan-20S**QA10TIS01**

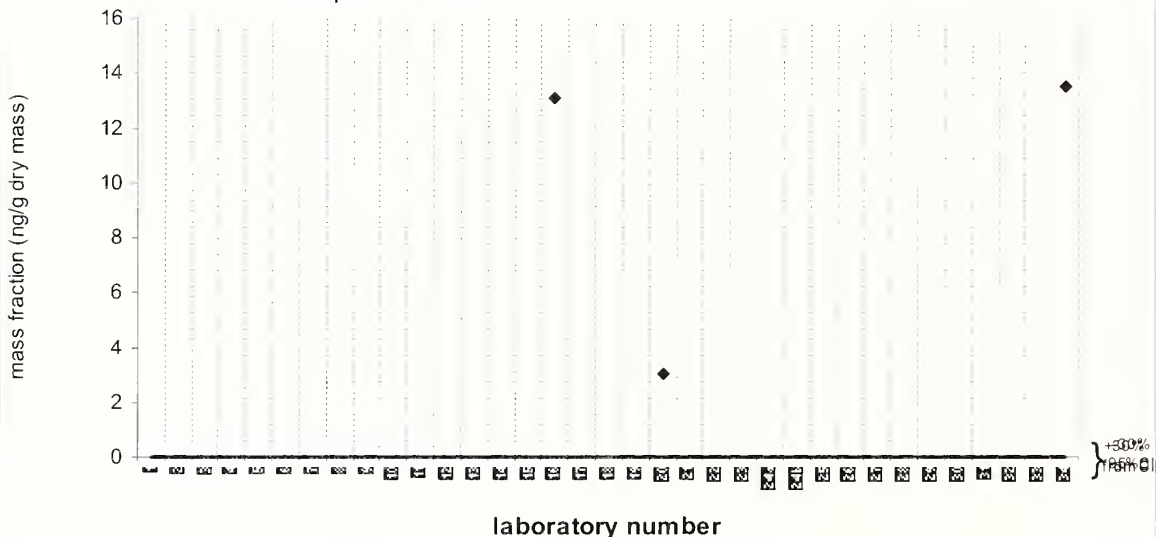
Assigned value = No Target ng/g (dry mass) Median value = 18.2 ng/g dry mass

Reported Results: 9 Quantitative Results: 3

**13b(H)17a(H)-Diacholestan-20S****SRM 1974b**

Target Value = no target ng/g (dry mass): Median value = 13.1 ng/g dry mass

Reported Results: 7 Quantitative Results: 3

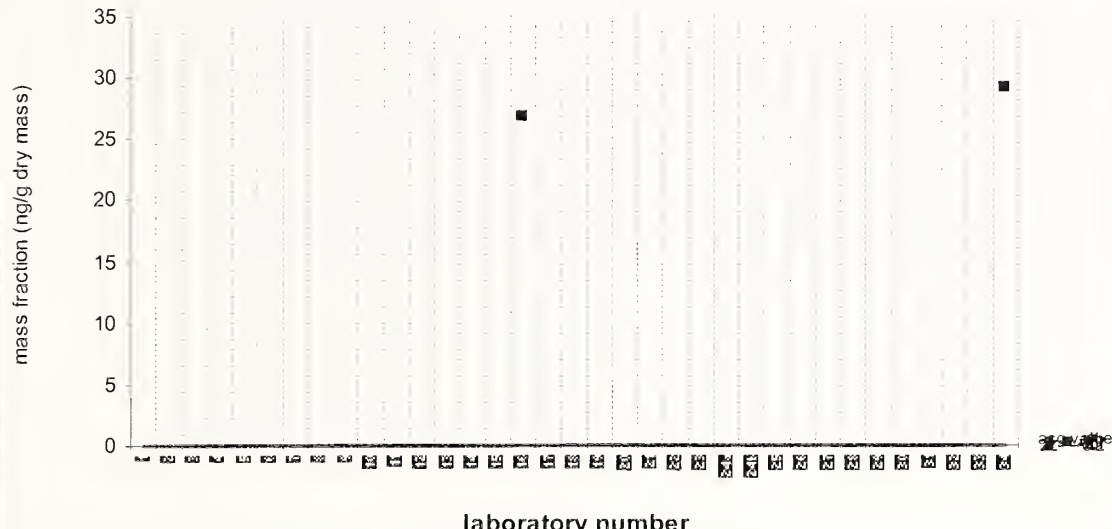


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

5a(H),14a(H),17a(H)-Cholestane 20S**QA10TIS01**

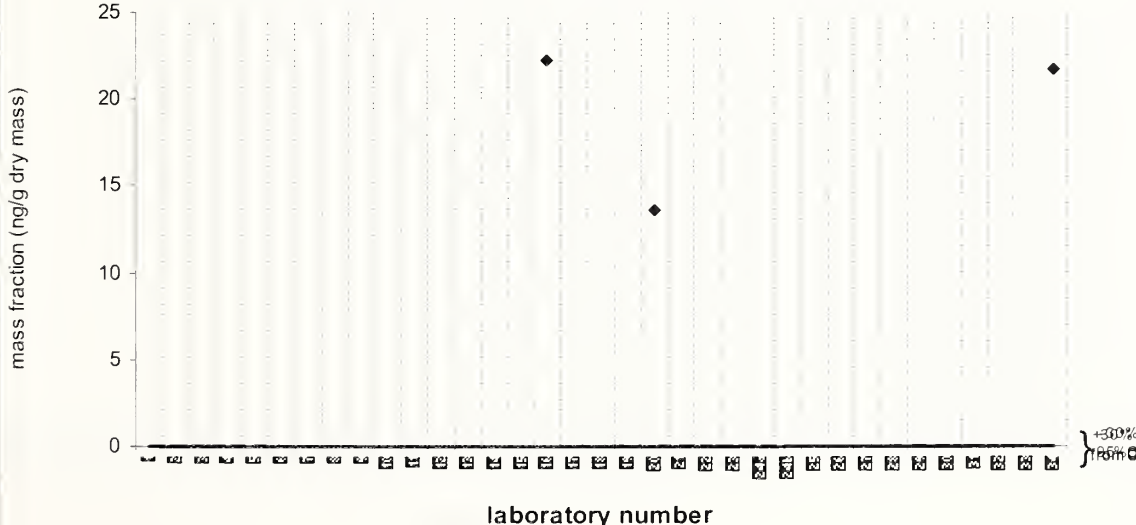
Assigned value = No Target ng/g (dry mass) Median value = 28.0 ng/g dry mass

Reported Results: 8 Quantitative Results: 2

**5a(H),14a(H),17a(H)-Cholestane 20S****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 21.7 ng/g dry mass

Reported Results: 7 Quantitative Results: 3

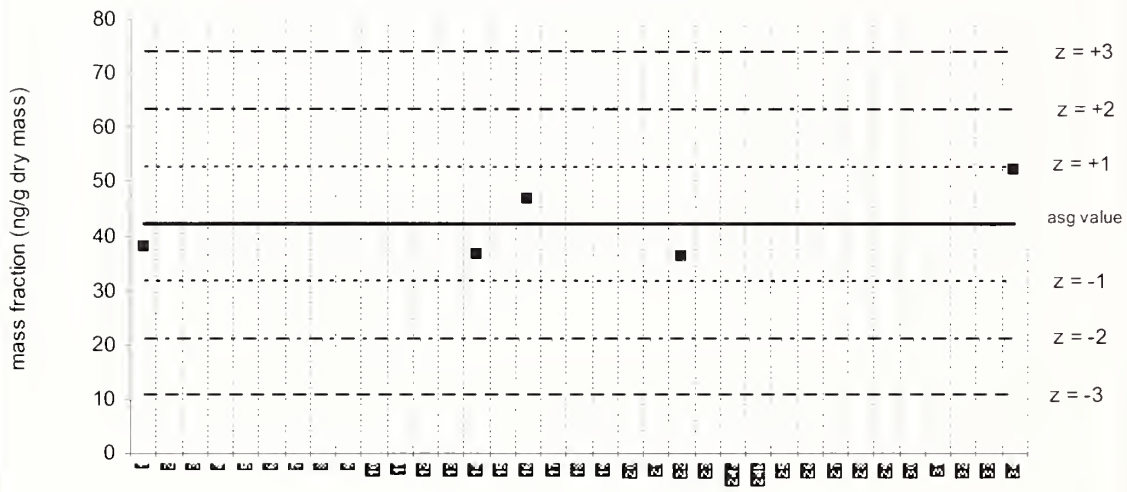


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

5a(H),14a(H),17a(H)-Cholestane 20R**QA10TIS01**

Assigned value = 42.2 ng/g dry mass s = 7.2 ng/g dry mass 95% CI = 6.3 ng/g dry mass Median value = 38.2 ng/g dry mass

Reported Results: 12 Quantitative Results: 5

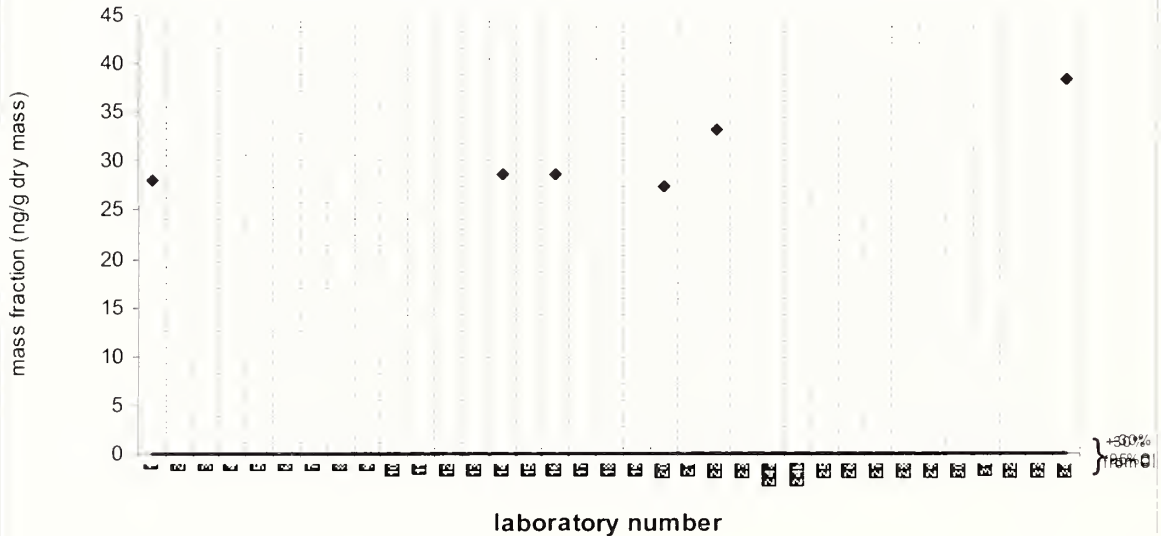
**laboratory number**

Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

5a(H),14a(H),17a(H)-Cholestane 20R**SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 28.7 ng/g dry mass

Reported Results: 10 Quantitative Results: 6

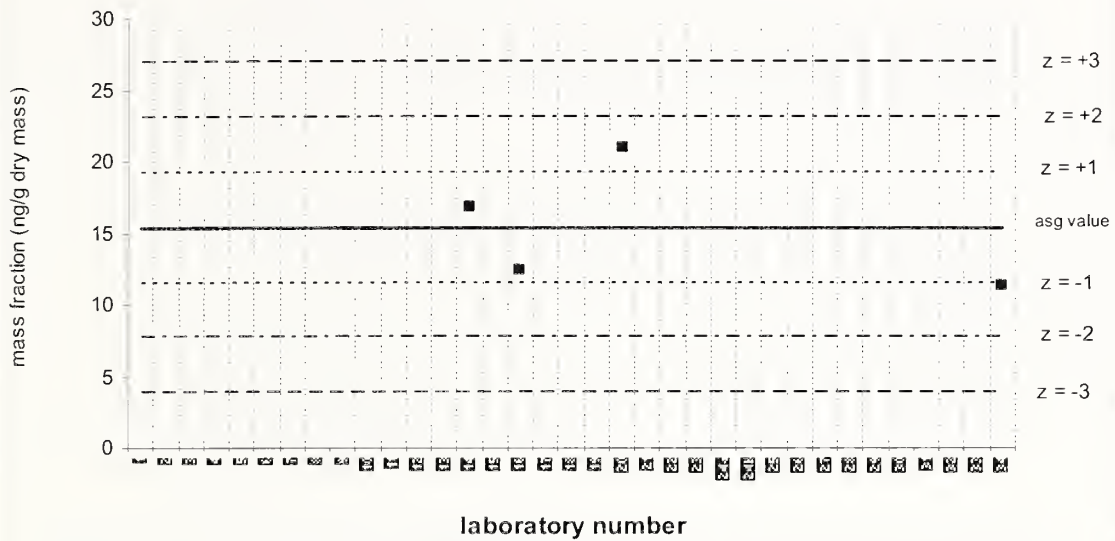
**laboratory number**

Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

5a(H),14a(H),17a(H)-24-Ethylcholestane 20S**QA10TIS01**

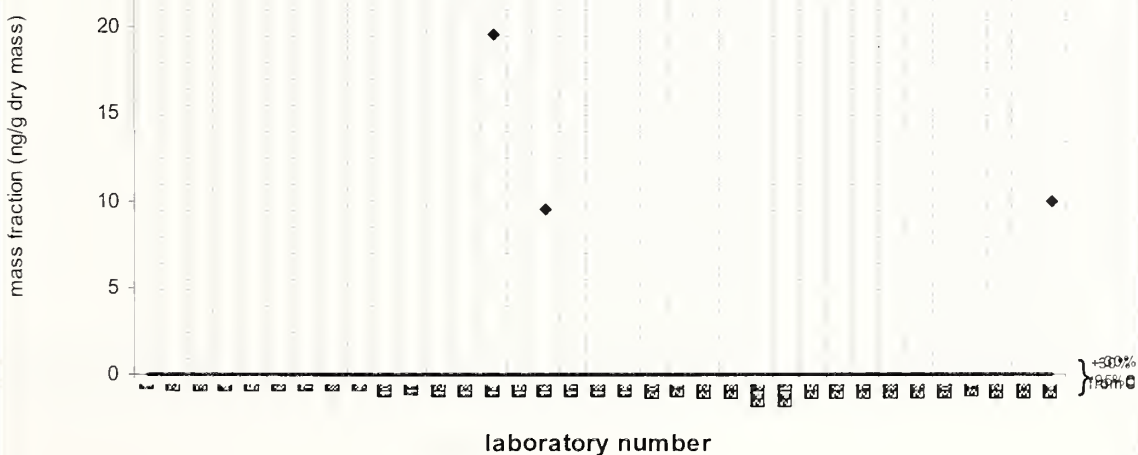
Assigned value = 15.4 ng/g dry mass s = 4.4 ng/g dry mass 95% CI = 4.3 ng/g dry mass Median value = 14.6 ng/g dry mass

Reported Results: 10 Quantitative Results: 4

**5a(H),14a(H),17a(H)-24-Ethylcholestane 20S****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 10.0 ng/g dry mass

Reported Results: 8 Quantitative Results: 3



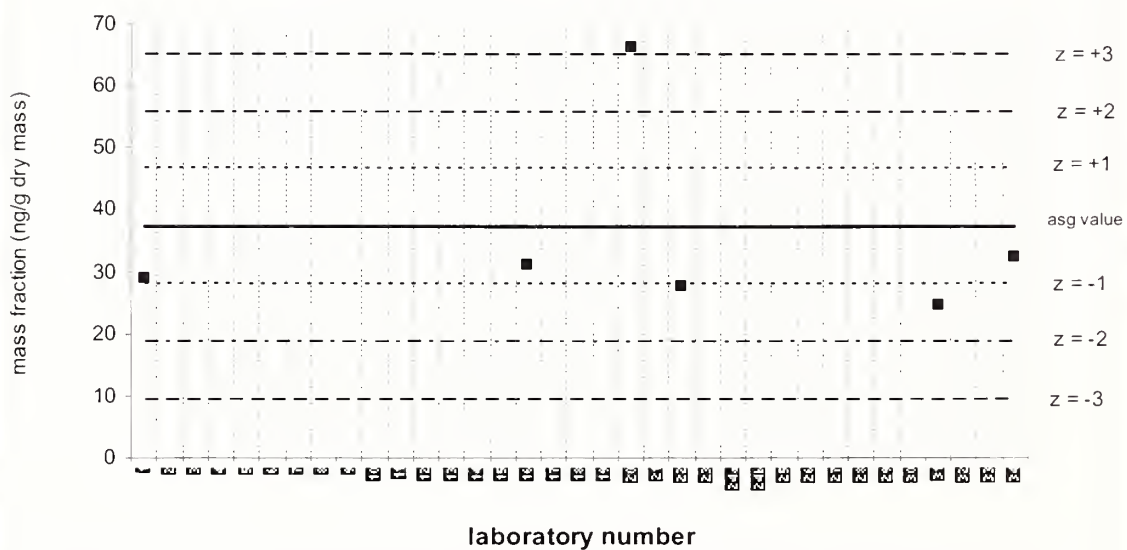
Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

5a(H),14a(H),17a(H)-24-Ethylcholestane 20R**QA10TIS01**

Assigned value = 37.2 ng/g dry mass s = 16.3 ng/g dry mass 95% CI = 14.3 ng/g dry mass

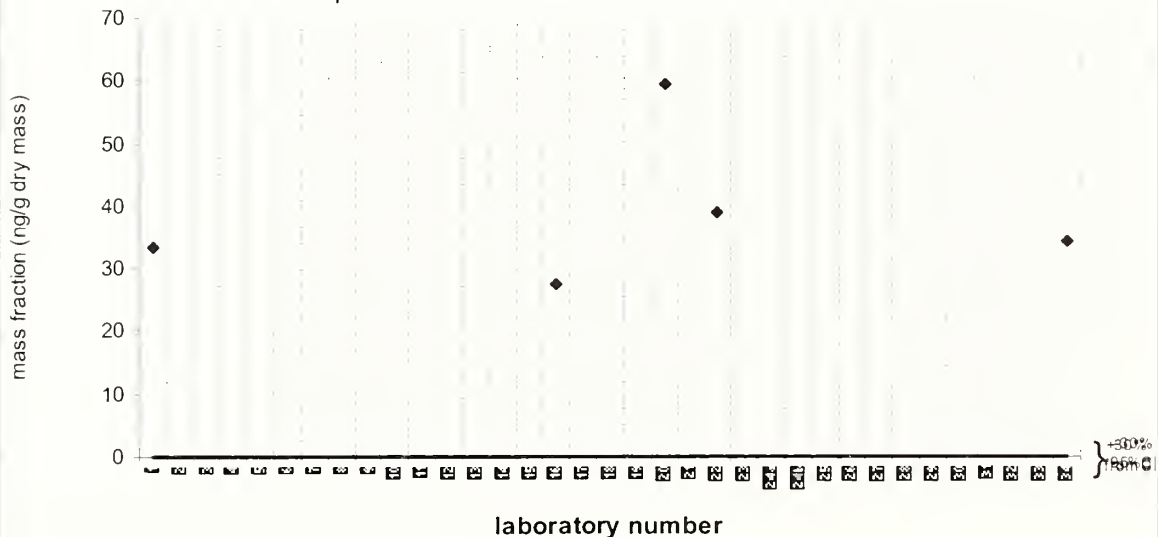
Median value = 30.0 ng/g dry mass

Reported Results: 11 Quantitative Results: 6

**5a(H),14a(H),17a(H)-24-Ethylcholestane 20R****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 34.3 ng/g dry mass

Reported Results: 9 Quantitative Results: 5

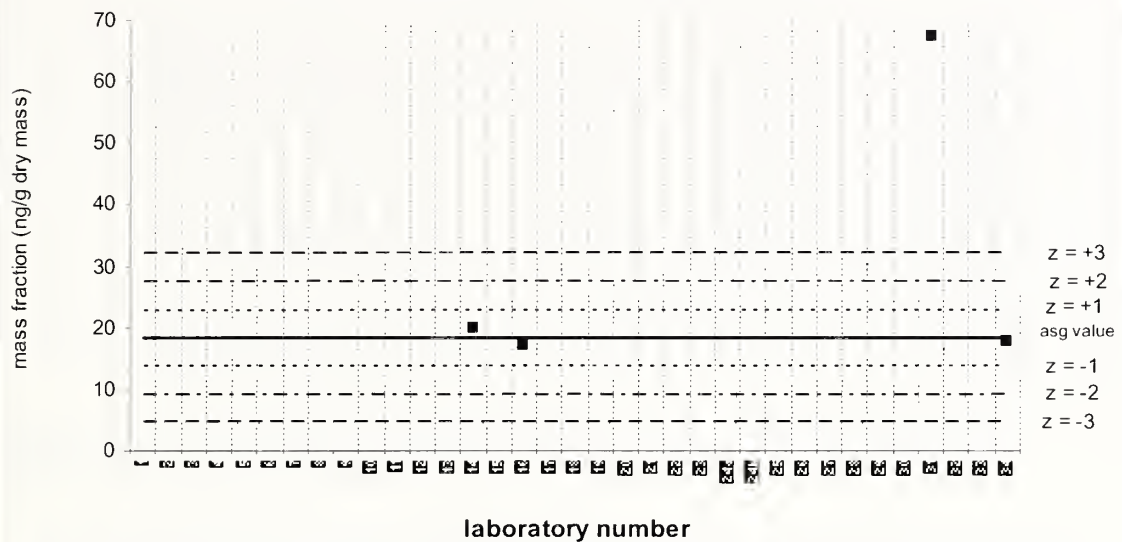


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

5a(H),14b(H),17b(H)-Cholestan-20R**QA10TIS01**

Assigned value = 18.3 ng/g dry mass s = 1.5 ng/g dry mass 95% CI = 1.7 ng/g dry mass Median value = 18.8 ng/g dry mass

Reported Results: 12 Quantitative Results: 4

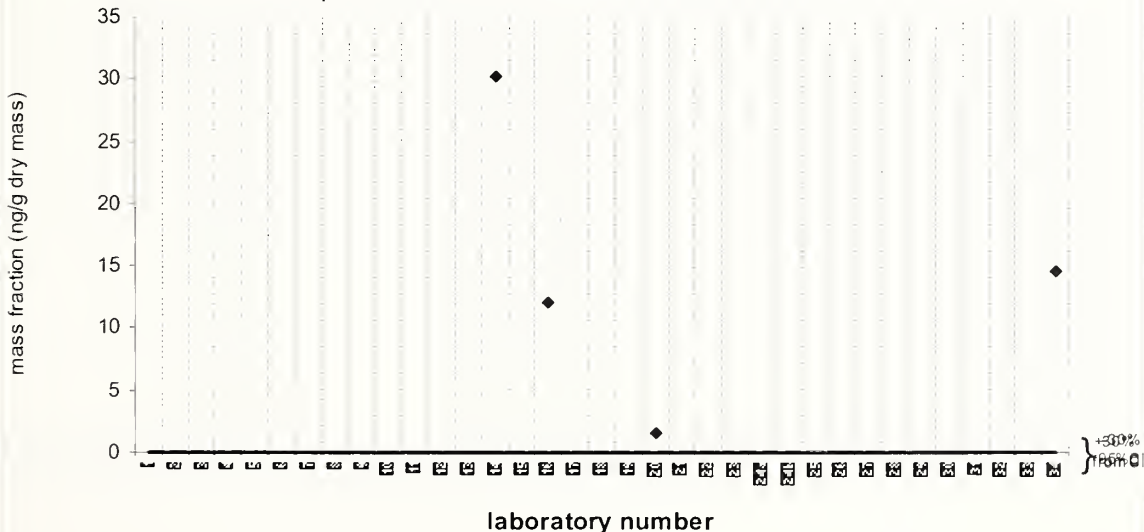


Solid line : exercise assigned value (EAV); dotted line: $z=\pm 1$ (25% from EAV); dotted/dashed line: $z=\pm 2$ (50% from EAV); dashed line: $z=\pm 3$ (75% from EAV)

5a(H),14b(H),17b(H)-Cholestan-20R**SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 13.4 ng/g dry mass

Reported Results: 10 Quantitative Results: 4

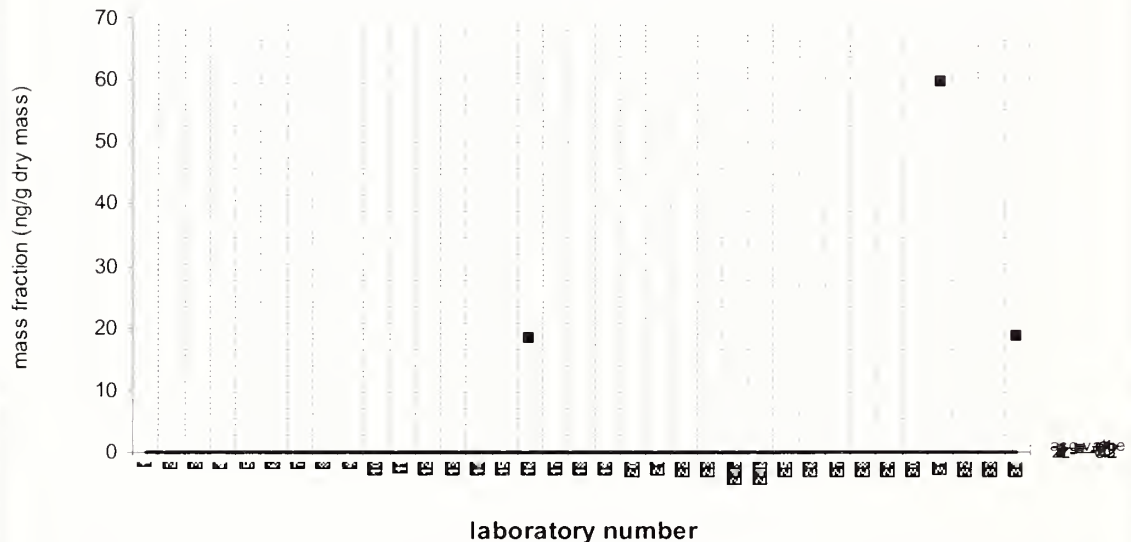


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

5a(H),14b(H),17b(H)-Cholestane 20S**QA10TIS01**

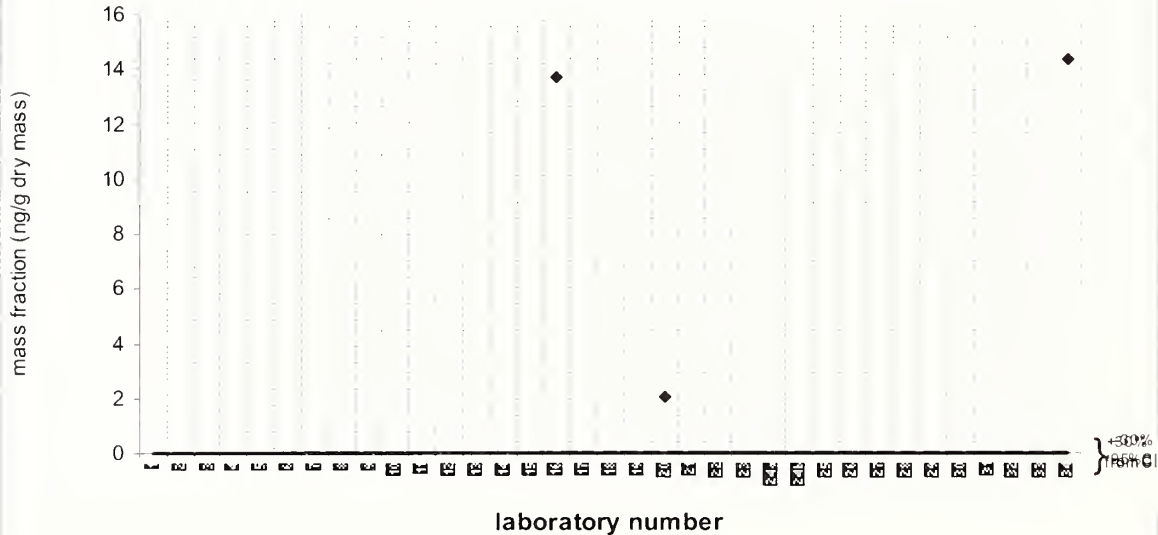
Assigned value = No Target ng/g (dry mass) Median value = 18.8 ng/g dry mass

Reported Results: 9 Quantitative Results: 3

**5a(H),14b(H),17b(H)-Cholestane 20S****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 13.8 ng/g dry mass

Reported Results: 7 Quantitative Results: 3

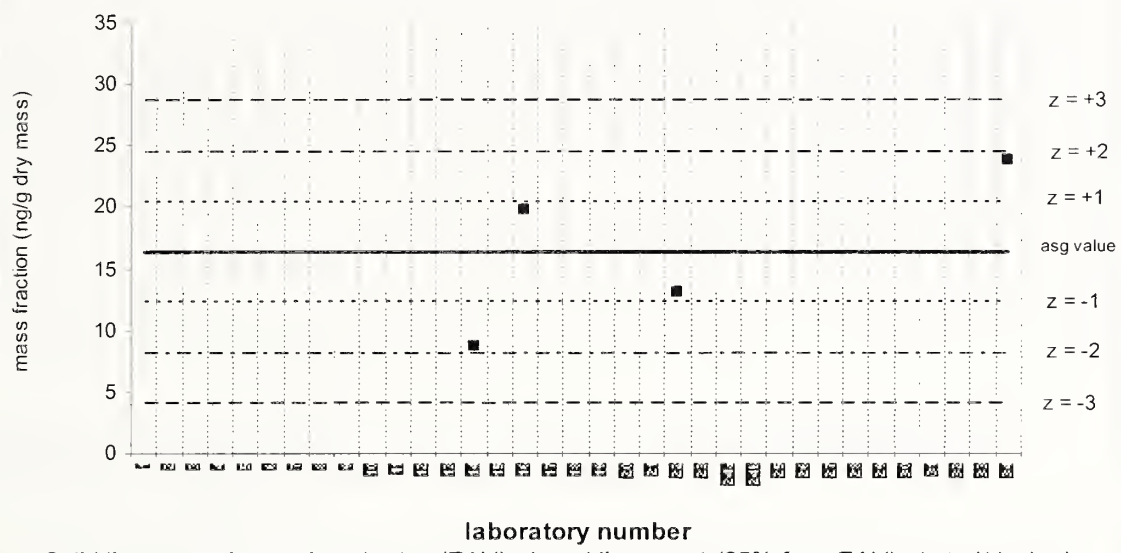


Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

5a(H),14b(H),17b(H)-24-Ethylcholestane 20R**QA10TIS01**

Assigned value = 16.3 ng/g dry mass s = 6.7 ng/g dry mass 95% CI = 6.6 ng/g dry mass Median value = 16.4 ng/g dry mass

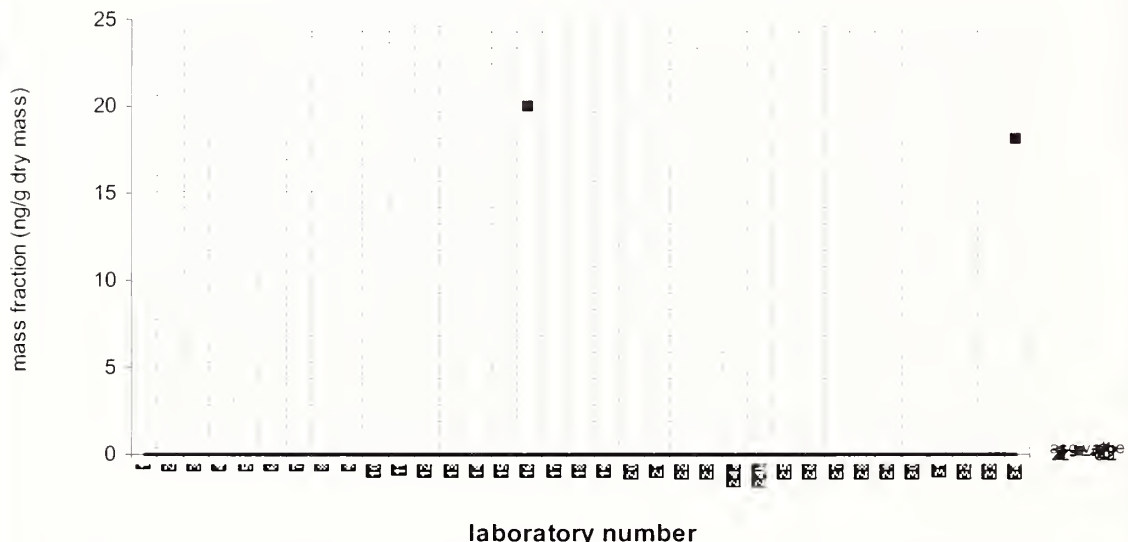
Reported Results: 12 Quantitative Results: 4



5a(H),14b(H),17b(H)-24-Ethylcholestane 20S**QA10TIS01**

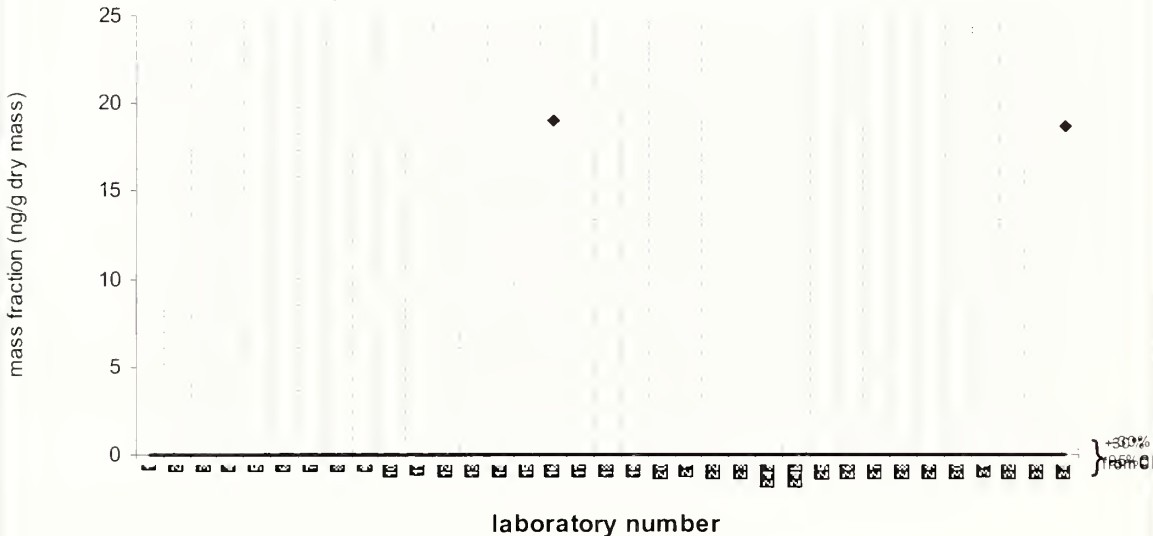
Assigned value = No Target ng/g (dry mass) Median value = 19.1 ng/g dry mass

Reported Results: 9 Quantitative Results: 2

**5a(H),14b(H),17b(H)-24-Ethylcholestane 20S****SRM 1974b**

Target Value = no target ng/g (dry mass); Median value = 18.9 ng/g dry mass

Reported Results: 7 Quantitative Results: 2



Solid line: value from Certificate of Analysis ; dotted line: 95% confidence limits; dashed line: 30% from 95% confidence limits

Appendix E
Participants in the Sediment Interlaboratory Study QA10OIL01 in
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