

September 21, 2020

Mr. Timothy Palazzolo
City of Detroit
Detroit Demolition Department
1301 Third Street
Detroit, Michigan 48226

Subject: Topsoil Composite Soil Sampling Report
9156 Lakepointe
Detroit, Michigan
AKT Peerless Project No. 8997f5-5-10.3

Mr. Palazzolo:

The City of Detroit, Detroit Demolition Department retained AKT Peerless to conduct soil sampling at a residential demolition site located at 9156 Lakepointe in Detroit, Wayne County, Michigan.

This scope of work was conducted in accordance with AKT Peerless' Work Order Request dated July 29, 2020 (approved July 29, 2020).

AKT Peerless understands that a residential structure was demolished at this location and the former basement excavation was filled with engineered fill (crushed residential hardfill consisting of broken concrete foundations and slabs, hard paved surfaces, brick and block, along with soil overburden) from the basement floor to approximately 12-inches below grade. An imported topsoil material was placed in the top 12-inches to grade.

Field Investigation

On August 19, 2020, Mr. Sean Brick and Mr. Antonio Morsette with AKT Peerless collected (1) composite sample, which consisted of three (3) discrete soil samples from the topsoil (top 12") combined into one (1) composite sample identified as "9156 Lakepointe-TS-Composite" in the former area of the residential structure.

AKT Peerless used a shovel to collect the samples.

During sample collection, AKT Peerless adhered to proper decontamination procedures. Sampling equipment was decontaminated using the following methods to minimize potential cross-contamination of soil samples:

- Washing and scrubbing the equipment with non-phosphate detergent
- Rinsing the equipment
- Air-drying the equipment

AKT Peerless collected samples according to USEPA Publication SW-846, Testing Methods for Evaluating Solid Waste. Soil samples were collected in laboratory-supplied containers, stored on ice at approximately 4 degrees Celsius, and submitted under chain-of-custody documentation. Soil samples collected for volatile analyses (VOCs) were field preserved with methanol in accordance with U.S. EPA

Method 5035. Soil samples collected for additional analyses were stored in unpreserved, 8-ounce wide-mouth jars.

Photographs taken during sampling activities are included in Attachment I. A site map depicting the sample locations is included in Attachment II.

Laboratory Analysis and Methods

The soil samples were delivered to ALS Group, USA, an independent National Voluntary Laboratory Accreditation Program (NVLAP) laboratory in Holland, Michigan, under chain-of-custody documentation for analysis of:

- Volatile Organic Compounds (VOCs) in accordance with USEPA Method 8260C;
- Semi-Volatile Organic Compounds (SVOCs) in accordance with USEPA Method 8270D;
- Polychlorinated biphenyls (PCBs) in accordance with USEPA Method 8082;
- Michigan 10 Metals (arsenic, barium, cadmium, total chromium, copper, lead, mercury, selenium, silver, and zinc) in accordance with USEPA Method 6020B or 7471B;
- Chloride in accordance with USEPA Method 325.3;
- Herbicides in accordance with USEPA Method 8151; and
- Pesticides in accordance with USEPA Method 8081A.

Analytical Results

AKT Peerless compared the laboratory analytical results to Michigan Department of Environment, Great Lakes and Energy (EGLE) Part 201 Residential Cleanup Criterion (RCC) provided in Michigan Administrative Rules 299.1 through 299.50 and the requirements outlined in the Detroit Land Bank Authority (DLBA) Scope of Services, revised September 13, 2018 (which were the applicable Scope of Work standards at the time backfill was placed at the subject property).

The results of the investigation indicate the following:

- VOCs were not detected in the topsoil composite sample above laboratory method detection limits (MDLs).
- SVOCs were not detected in the topsoil composite sample above laboratory MDLs.
- PCBs were not detected in the topsoil composite sample above laboratory MDLs.
- Arsenic, barium, cadmium, chromium (total), copper, lead, and zinc were detected in the topsoil composite sample above laboratory MDLs but below EGLE Part 201 RCC. Selenium and silver were not detected in the topsoil composite sample above laboratory MDLs.
- Mercury was detected in the topsoil composite samples exceeding EGLE Part 201 GSIP criteria.
- Chloride was not detected in the topsoil composite sample above laboratory MDLs.
- Herbicides were not detected in the topsoil composite sample above laboratory MDLs.
- Select pesticides were detected in the topsoil composite sample at a concentration above laboratory MDLs but below EGLE Part 201 RCC. Remaining pesticides were not detected in the topsoil composite sample above laboratory MDLs.

A table summarizing the soil sampling results is included in Attachment III. The laboratory analytical results and chain of custody documentation are also provided in Attachment III.

Conclusions and Recommendations

AKT Peerless collected one (1) composite sample, which consisted of three (3) discrete soil samples from the topsoil (top 12") combined into one (1) composite sample in the former area of the residential structure located at 9156 Lakepointe in Detroit, Wayne County, Michigan. The soil samples were submitted for laboratory analysis of VOCs, SVOCs, PCBs, MI Metals, chloride, herbicides, and pesticides.

Laboratory analytical results from the topsoil composite sample did not identify the presence of target compounds above laboratory MDLs and/or EGLE Part 201 RCC, except for mercury. Mercury was identified at a concentration exceeding EGLE Part 201 GSIP criteria.

According to the DLBA Scope of Services, backfill that is contaminated above an EGLE Part 201 RCC or with detectable concentrations of VOCs is not suitable for the Detroit backfill program. Based on the sampling results from the topsoil composite sample, the topsoil material does not meet the DLBA Scope of Services requirements.

Limitations

The information and opinions obtained in this report are for the exclusive use of the City of Detroit. No distribution to or reliance by other parties may occur without the express written permission of AKT Peerless. AKT Peerless will not distribute this report without your written consent or as required by law or by a Court order. The information and opinions contained in the report are given in light of that assignment. The report must be reviewed and relied upon only in conjunction with the terms and conditions expressly agreed upon by the parties and as limited therein. Any third parties who have been extended the right to rely on the contents of this report by AKT Peerless (which is expressly required prior to any third-party release), expressly agrees to be bound by the original terms and conditions entered into by AKT Peerless and the City of Detroit.

Subject to the above and the terms and conditions, AKT Peerless accepts responsibility for the competent performance of its duties in executing the assignment and preparing reports in accordance with the normal standards of the profession but disclaims any responsibility for consequential damages. Although AKT Peerless believes that results contained herein are reliable, AKT Peerless cannot warrant or guarantee that the information provided is exhaustive or that the information provided by the City of Detroit or third parties is complete or accurate.

Signatures of Environmental Professionals

The following individuals contributed to the completion of this report.

DRAFT

Sean Brick
Environmental Consultant
AKT Peerless
Farmington, Michigan Office
Phone: 248.615.1333

DRAFT

Megan Napier
Senior Engineer
AKT Peerless
Farmington, Michigan Office
Phone: 248.615.1333

DRAFT

Attachment I
Site Photographs



View of the property looking north.



View of the property looking east.



View of the property looking south.



View of the property looking west.



View of sample locations.





View of typical top fill material.

Attachment II

Site Map



Notes:

-  – Approximate location of excavation area
-  – Approximate location of AKT Peerless discrete sample locations

Map not to scale.

Map/aerial courtesy of Detroit Parcel Viewer at <https://cityofdetroit.github.io/parcel-viewer/>



Sample Location Map

9156 Lakepointe
Detroit, Michigan

Project No: 8997F5-5-10.4

Attachment III

Soil Results Table and Laboratory Analytical Results

Table 1: Summary of Soil Analytical Results
9156 Lakepointe, Detroit, Michigan
AKT Peerless Project No. 8997F5-5-10.4

Parameters*	Chemical Abstract Service Number	Statewide Default Background Levels	Residential Drinking Water Protection Criteria	Groundwater Surface Water Interface Protection Criteria	Residential Soil Volatilization to Indoor Air Inhalation Criteria	Residential Infinite Source Volatile Soil Inhalation Criteria	Residential Particulate Soil Inhalation Criteria	Residential Direct Contact Criteria	Soil Saturation Concentration Screening Levels	Sample Name	9156 Lakepointe-TS-Composite
<i>*(Refer to detailed laboratory report for method reference data)</i>										Collection Date	8/19/2020
Metals (µg/kg)											
Arsenic	7440-38-2	5,800	4,600	4,600	NLV	NLV	7.2E+5	7,600	NA		3,400
Barium (B)	7440-39-3	75,000	1.3E+6	(G)	NLV	NLV	3.3E+8	3.7E+7	NA		40,000
Cadmium (B)	7440-43-9	1,200	6,000	(G,X)	NLV	NLV	1.7E+6	5.5E+5	NA		290
Chromium, Total	7440-47-3	18,000 (total)	30,000	3,300	NLV	NLV	2.6E+5	2.5E+6	NA		7,900
Copper (B)	7440-50-8	32,000	5.8E+6	(G)	NLV	NLV	1.3E+8	2.0E+7	NA		7,400
Lead (B)	7439-92-1	21,000	7.0E+5	(G,X)	NLV	NLV	1.0E+8	4.0E+5	NA		18,000
Mercury, Total	7439-97-6	130	1,700	50 (M); 1.2	48,000	52,000	2.0E+7	1.6E+5	NA		190
Selenium (B)	7782-49-2	410	4,000	400	NLV	NLV	1.3E+8	2.6E+6	NA		<420
Silver (B)	7440-22-4	1,000	4,500	100 (M); 27	NLV	NLV	6.7E+6	2.5E+6	NA		<420
Zinc (B)	7440-66-6	47,000	2.4E+6	(G)	NLV	NLV	ID	1.7E+8	NA		29,000
Chloride (µg/kg)											
Chloride	16887-00-6	NA	5.0E+6	(X)	NLV	NLV	ID	5.0E+5 (F)	NA		<200,000
Polychlorinated biphenyls (PCBs, µg/kg)											
PCBs (J,T)	1336-36-3	NA	NLL	NLL	3.0E+6	2.4E+5	5.2E+6	(T)	NA		<330
Herbicides (µg/kg)											
Silvex (2,4,5-TP)	93-72-1	NA	3,600	2,200	NLV	NLV	ID	1.7E+6	NA		<300
2,4-Dichlorophenoxyacetic acid	94-75-7	NA	1,400	4,400	NLV	NLV	6.7E+9	2.5E+6	NA		<200
Pesticides (µg/kg)											
4-4'-DDE	72-55-9	NA	NLL	NLL	NLV	NLV	3.2E+7	45,000	NA		20
Remaining Pesticides	Varies	-	-	-	-	-	-	-	-		BDL
Semivolatile Organic Compounds (SVOCs, µg/kg)											
All SVOCs	Varies	-	-	-	-	-	-	-	-		BDL
Volatile Organic Compounds (VOCs, µg/kg)											
All VOCs	Varies	-	-	-	-	-	-	-	-		BDL

R 299.49 FOOTNOTES FOR GENERIC CLEANUP CRITERIA TABLES (as last revised on December 30, 2013)

Cleanup Criteria Requirements for Response Activity (formerly the Part 201 Generic Cleanup Criteria and Screening Levels)

- (A) Criterion is the state of Michigan drinking water standard established pursuant to Section 5 of 1976 PA 399, MCL 325.1005.
- (B) Background, as defined in R 299.11(b), may be substituted if higher than the calculated cleanup criterion. Background levels may be less than criteria for some inorganic compounds.
- (C) The criterion developed under R 299.20 to R 299.26 exceeds the chemical-specific soil saturation screening level (C_{ss}). The person proposing or implementing response activity shall document whether additional response activity is required to control free-phase liquids or NAPL to protect against risks associated with free-phase liquids by using methods appropriate for the free-phase liquids present. Development of a site-specific C_{ss} or methods presented in R 299.22, R 299.24(5), and R 299.26(8) may be conducted for the relevant exposure pathways.
- (D) Calculated criterion exceeds 10%, hence it is reduced to 100 percent or 1.0E+9 parts per billion (ppb).
- (E) Criterion is the aesthetic drinking water value, as required by Section 20120a(5) of the Natural Resources and Environmental Protection Act, 1994 PA 451, as amended (NREPA). A notice of aesthetic impact may be employed as an institutional control mechanism if groundwater concentrations exceed the aesthetic drinking water criterion, but do not exceed the applicable health-based drinking water value [as provided in the table in Footnote (E) in R 299.49].
- (F) Criterion is based on adverse impacts to plant life and phytotoxicity.
- (G) Groundwater surface water interface (GSI) criterion depends on the pH or water hardness, or both, of the receiving surface water. The final chronic value (FCV) for the protection of aquatic life shall be calculated based on the pH or hardness of the receiving surface water. Where water hardness exceeds 400 mg CaCO_3/L , use 400 mg CaCO_3/L for the FCV calculation. The FCV formula provides values in units of $\mu\text{g}/\text{L}$ or ppb. The generic GSI criterion is the lesser of the calculated FCV, the wildlife value (WV), and the surface water human non-drinking water value (HNDV). The soil GSI protection criteria for these hazardous substances are the greater of 20 times the GSI criterion or the GSI soil-water partition values using the GSI criteria developed with the procedure described in this footnote. [See table in Footnote (G) in R 299.49].
- (H) Valence-specific chromium data (Cr III and Cr VI) shall be compared to the corresponding valence-specific cleanup criteria. If both Cr III and Cr VI are present in groundwater, the total concentration of both cannot exceed the drinking water criterion of 100 $\mu\text{g}/\text{L}$. If analytical data are provided for total chromium only, they shall be compared to the cleanup criteria for Cr VI. Cr III soil cleanup criterion for protection of drinking water can only be used at sites where groundwater is prevented from being used as a public water supply, currently and in the future, through an approved land or resource use restriction.
- (I) Hazardous substance may exhibit the characteristic of ignitability as defined in 40 C.F.R. §261.21 (revised as of July 1, 2001), which is adopted by reference in these rules.
- (J) Hazardous substance may be present in several isomer forms. Isomer-specific concentrations shall be added together for comparison to criteria.
- (K) Hazardous substance may be flammable or explosive, or both.
- (L) Criteria for lead are derived using a biologically based model, as allowed for under Section 20120a(9) of the NREPA, and are not calculated using the algorithms and assumptions specified in pathway-specific rules. The generic residential drinking water criterion of 4 $\mu\text{g}/\text{L}$ is linked to the generic residential soil direct contact criterion of 400 mg/kg. A higher concentration in the drinking water, up to the state action level of 15 $\mu\text{g}/\text{L}$, may be allowed as a site-specific remedy and still allow for drinking water use, under Section 20120a(2) of the NREPA if soil concentrations are appropriately lower than 400 mg/kg. If a site-specific criterion is approved based on this subdivision, a notice shall be filed on the deed for all property where the groundwater concentrations will exceed 4 $\mu\text{g}/\text{L}$ to provide notice of the potential for unacceptable risk if soil or groundwater concentrations increase. Acceptable concentrations of site-specific soil and drinking water concentrations are presented in the [table in Footnote (L) in R 299.49].
- (M) Calculated criterion is below the analytical target detection limit, therefore, the criterion defaults to the target detection limit.
- (N) The concentrations of all potential sources of nitrate-nitrogen (e.g., ammonia-N, nitrite-N, nitrate-N) in groundwater that is used as a source of drinking water shall not, when added together, exceed the nitrate drinking water criterion of 10,000 $\mu\text{g}/\text{L}$. Where leaching to groundwater is a relevant pathway, soil concentrations of all potential sources of nitrate-nitrogen shall not, when added together, exceed the nitrate drinking water protection criterion of 2.0E+5 $\mu\text{g}/\text{kg}$.
- (O) The concentration of all polychlorinated and polybrominated dibenzodioxin and dibenzofuran isomers present at a facility, expressed as an equivalent concentration of 2,3,7,8-tetrachlorodibenzo-p-dioxin based upon their relative potency, shall be added together and compared to the criteria for 2,3,7,8-tetrachlorodibenzo-p-dioxin. The generic cleanup criteria for 2,3,7,8-tetrachlorodibenzo-p-dioxin are not calculated according to the algorithms presented in R 299.14 to R 299.26. The generic cleanup criteria are being held at the values that the DEQ has used since August 1998, in recognition of the fact that national efforts to reassess risks posed by dioxin are not yet complete. Until these studies are complete, it is premature to select a revised slope factor and/or reference dose for calculation of generic cleanup criteria.
- (P) Amenable cyanide methods or method OIA-1677 shall be used to quantify cyanide concentrations for compliance with all groundwater criteria. Total cyanide methods or method OIA-1677 shall be used to quantify cyanide concentrations for compliance with soil criteria. Nonresidential direct contact criteria may not be protective of the potential for release of hydrogen cyanide gas. Additional land or resource use restrictions may be necessary to protect for the acute inhalation concerns associated with hydrogen cyanide gas.
- (Q) Criteria for carcinogenic polycyclic aromatic hydrocarbons were developed using relative potential potencies to benzo(a)pyrene.
- (R) Hazardous substance may exhibit the characteristic of reactivity as defined in 40 C.F.R. §261.23 (revised as of July 1, 2001), which is adopted by reference in these rules.
- (S) Criterion defaults to the hazardous substance-specific water solubility limit.
- (T) Refer to the federal Toxic Substances Control Act (TSCA), 40 C.F.R. §761, subpart D and 40 C.F.R. §761, Subpart G, to determine the applicability of TSCA cleanup standards. Subpart D and subpart G of 40 C.F.R. §761 (July 1, 2001) are adopted by reference in these rules. Alternatives to compliance with the TSCA standards listed below are possible under 40 C.F.R. §761 Subpart D. New releases may be subject to the standards identified in 40 C.F.R. §761, Subpart G. Use Part 201 soil direct contact cleanup criteria in the following table if TSCA standards are not applicable. [See table in Footnote (T) in R 299.49].
- (U) Hazardous substance may exhibit the characteristic of corrosivity as defined in 40 C.F.R. §261.22 (revised as of July 1, 2001), which is adopted by reference in these rules.
- (V) Criterion is the aesthetic drinking water value as required by Section 20120a(5) of the NREPA. Concentrations up to 200 $\mu\text{g}/\text{L}$ may be acceptable, and still allow for drinking water use, as part of a site-specific cleanup under Section 20120a(2) and 20120b of the NREPA.
- (W) Concentrations of trihalomethanes in groundwater shall be added together to determine compliance with the Michigan drinking water standard of 80 $\mu\text{g}/\text{L}$. Concentrations of trihalomethanes in soil shall be added together to determine compliance with the drinking water protection criterion of 1,600 $\mu\text{g}/\text{kg}$.
- (X) The GSI criterion shown in the generic cleanup criteria tables is not protective for surface water that is used as a drinking water source. For a groundwater discharge to the Great Lakes and their connecting waters or discharge in close proximity to a water supply intake in inland surface waters, the generic GSI criterion shall be the surface water human drinking water value (HDV) listed in the [table in Footnote (X) in R 299.49], except for those HDV indicated with an asterisk. For HDV with an asterisk, the generic GSI criterion shall be the lowest of the HDV, the WV, and the calculated FCV. See formulas in [the table in Footnote (G) in R 299.49]. Soil protection criteria based on the HDV shall be as listed in the [table in Footnote (X) in R 299.49], except for those values with an asterisk. Soil GSI protection criteria based on the HDV shall be as listed in the [table in Footnote (X) in R 299.49], except for those values with an asterisk. Soil GSI protection criteria for compounds with an asterisk shall be the greater of 20 times the GSI criterion or the GSI soil-water partition values using the GSI criteria developed with the procedure described in this footnote.
- (Y) Source size modifiers shown in the [table in Footnote (Y) in R 299.49] shall be used to determine soil inhalation criteria for ambient air when the source size is not one-half acre. The modifier shall be multiplied by the generic soil inhalation criteria shown in the table of generic cleanup criteria to determine the applicable criterion. See Footnote (C) [in R 299.49].
- (Z) Mercury is typically measured as total mercury. The generic cleanup criteria, however, are based on data for different species of mercury. Specifically, data for elemental mercury, chemical abstract service (CAS) number 7439976, serve as the basis for the soil volatilization to indoor air criteria, groundwater volatilization to indoor air, and soil inhalation criteria. Data for methyl mercury, CAS number 22967926, serve as the basis for the GSI criterion; and data for mercuric chloride, CAS number 7487947, serve as the basis for the drinking water, groundwater contact, soil direct contact, and the groundwater protection criteria. Comparison to criteria shall be based on species-specific analytical data only if sufficient facility characterization has been conducted to rule out the presence of other species of mercury.
- (AA) Use 10,000 $\mu\text{g}/\text{L}$ where groundwater enters a structure through the use of a water well, sump or other device. Use 28,000 $\mu\text{g}/\text{L}$ for all other uses.
- (BB) The state drinking water standard for asbestos (fibers greater than 10 micrometers in length) is in units of a million fibers per liter of water (MFL). Soil concentrations of asbestos are determined by polarized light microscopy.
- (CC) Groundwater: The generic GSI criteria are based on the toxicity of unionized ammonia (NH_3); the criteria are 29 $\mu\text{g}/\text{L}$ and 53 $\mu\text{g}/\text{L}$ for cold water and warm water surface water, respectively. As a result, the GSI criterion shall be compared to the percent of the total ammonia concentration in the groundwater that will become NH_3 in the surface water. This percent NH_3 is a function of the pH and temperature of the receiving surface water and can be estimated using the [table in Footnote (CC) in R 299.49], taken from Emerson, et al., (Journal of the Fisheries Research Board of Canada, Volume 32(12):2382, 1975). The generic approach for estimating NH_3 assumes a default pH of 8 and default temperatures of 68 °F and 85 °F for cold water and warm water surface water, respectively. The resulting NH_3 is 3.8 percent and 7.2 percent for cold water and warm water, respectively. This default percentage shall be multiplied by the total ammonia-nitrogen ($\text{NH}_3\text{-N}$) concentration in the groundwater and the resulting NH_3 concentration compared to the applicable GSI criterion. As an alternative, the maximum pH and temperature data from the specific receiving surface water can be used to estimate, from the [table in Footnote (CC) in R 299.49], a lower percent unionized ammonia concentration for comparison to the generic GSI.
- Soil: The generic soil GSI protection criteria for unionized ammonia are 580 $\mu\text{g}/\text{kg}$ and 1,100 $\mu\text{g}/\text{kg}$ for cold water and warm water surface water, respectively.
- (DD) Hazardous substance causes developmental effects. Residential direct contact criteria are protective of both prenatal and postnatal exposure. Nonresidential direct contact criteria are protective for a pregnant adult receptor.
- (EE) The [values listed in the table in Footnote (EE) in R 299.49] are applicable generic GSI criteria as required by Section 20120e of the NREPA.
- (FF) The chloride GSI criterion shall be 125 mg/L when the discharge is to surface waters of the state designated as public water supply sources or 50 mg/L when the discharge is to the Great Lakes or connecting waters. Chloride GSI criteria shall not apply for surface waters of the state that are not designated as a public water supply source; however, the total dissolved solids criterion is applicable.
- (GG) Risk-based criteria are not available for methane due to insufficient toxicity data. An acceptable soil gas concentration (presented for both residential and nonresidential land uses) was derived utilizing 25 percent of the lower explosive level for methane. This equates to 1.25 percent or 8.4E+6 $\mu\text{g}/\text{m}^3$.
- (HH) The residential criterion for sodium is 230,000 $\mu\text{g}/\text{L}$ in accordance with the Sodium Advisory Council recommendation and revised Groundwater Discharge Standards.
- ID Insufficient data to develop criterion.
- NA A criterion or value is not available or, in the case of background and CAS numbers, not applicable.
- NLL Hazardous substance is not likely to leach under most soil conditions.
- NLV Hazardous substance is not likely to volatilize under most conditions.
- $\mu\text{g}/\text{kg}$ Micrograms per kilogram
- $\mu\text{g}/\text{L}$ Micrograms per liter
- BDL Below Laboratory Method Detection Limits
- BOLD** Exceeds criteria.
- BF Backfill
- TS Topsoil



31-Aug-2020

Megan Napier
AKT Peerless
22725 Orchard Lake Road
Farmington, MI 48336

Re: **8997f5-5-10.4**

Work Order: **20081693**

Dear Megan,

ALS Environmental received 1 sample on 21-Aug-2020 08:00 AM for the analyses presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental - Holland and for only the analyses requested.

Sample results are compliant with industry accepted practices and Quality Control results achieved laboratory specifications. Any exceptions are noted in the Case Narrative, or noted with qualifiers in the report or QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained from ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

The total number of pages in this report is 40.

If you have any questions regarding this report, please feel free to contact me:

ADDRESS: 3352 128th Avenue, Holland, MI, USA
PHONE: +1 (616) 399-6070 FAX: +1 (616) 399-6185

Sincerely,

A handwritten signature in black ink, appearing to read "Bill Carey".

Electronically approved by: Bill Carey

Bill Carey
Project Manager

Report of Laboratory Analysis

Certificate No: MN 026-999-449

ALS GROUP USA, CORP Part of the ALS Laboratory Group A Campbell Brothers Limited Company

Environmental 

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RIGHT SOLUTIONS RIGHT PARTNER

Client: AKT Peerless
Project: 8997f5-5-10.4
Work Order: 20081693

Work Order Sample Summary

<u>Lab Samp ID</u>	<u>Client Sample ID</u>	<u>Matrix</u>	<u>Tag Number</u>	<u>Collection Date</u>	<u>Date Received</u>	<u>Hold</u>
20081693-01	9156 Lakepointe-TS-Composite	Soil		8/19/2020 11:35	8/21/2020 08:00	<input type="checkbox"/>

Client: AKT Peerless
Project: 8997f5-5-10.4
WorkOrder: 20081693

**QUALIFIERS,
ACRONYMS, UNITS**

<u>Qualifier</u>	<u>Description</u>
*	Value exceeds Regulatory Limit
**	Estimated Value
a	Analyte is non-accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
Hr	BOD/CBOD - Sample was reset outside Hold Time, value should be considered estimated.
J	Analyte is present at an estimated concentration between the MDL and Report Limit
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL
X	Analyte was detected in the Method Blank between the MDL and Reporting Limit, sample results may exhibit background or reagent contamination at the observed level.

<u>Acronym</u>	<u>Description</u>
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
LOD	Limit of Detection (see MDL)
LOQ	Limit of Quantitation (see PQL)
MBLK	Method Blank
MDL	Method Detection Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PQL	Practical Quantitation Limit
RPD	Relative Percent Difference
TDL	Target Detection Limit
TNTC	Too Numerous To Count
A	APHA Standard Methods
D	ASTM
E	EPA
SW	SW-846 Update III

<u>Units Reported</u>	<u>Description</u>
% of sample	Percent of Sample
µg/Kg-dry	Micrograms per Kilogram Dry Weight
mg/Kg-dry	Milligrams per Kilogram Dry Weight

Client: AKT Peerless
Project: 8997f5-5-10.4
Work Order: 20081693

Case Narrative

Samples for the above noted Work Order were received on 8/21/2020. The attached ""Sample Receipt Checklist"" documents the status of custody seals, container integrity, preservation, and temperature compliance.

Samples were analyzed according to the analytical methodology previously transmitted in the ""Work Order Acknowledgement"". Methodologies are also documented in the ""Analytical Result"" section for each sample. Quality control results are listed in the ""QC Report"" section. Sample association for the reported quality control is located at the end of each batch summary. If applicable, results are appropriately qualified in the Analytical Result and QC Report sections. The ""Qualifiers"" section documents the various qualifiers, units, and acronyms utilized in reporting. A copy of the laboratory's scope of accreditation is available upon request.

With the following exceptions, all sample analyses achieved analytical criteria.

Volatile Organics:

No other deviations or anomalies were noted.

Extractable Organics:

Batch 163196, Method SVO_8270_S, Sample 20081693-01B: The reporting limit is elevated due to dilution needed to eliminate matrix-related interference.

Metals:

No other deviations or anomalies were noted.

Wet Chemistry:

No other deviations or anomalies were noted.

ALS Group, USA

Date: 31-Aug-20

Client: AKT Peerless
Work Order: 20081693
Project: 8997f5-5-10.4
Lab ID: 20081693-01

Client Sample ID: 9156 Lakepointe-TS-Composite
Collection Date: 8/19/2020 11:35:00 AM
Matrix: SOIL

Analyses	Result	Report Limit	MDEQ OP Memo 2 TDL	Qual	Units	Dilution Factor	Date Analyzed
HERBICIDES							
			SW8151		Prep Date: 8/26/2020		Analyst: KYM
2,4,5-TP (Silvex)	ND	300	300		µg/Kg-dry	1	8/29/2020
2,4-D	ND	200	200		µg/Kg-dry	1	8/29/2020
Surr: DCAA	38.0	10-150			%REC	1	8/29/2020
PCBS							
			SW8082		Prep Date: 8/25/2020		Analyst: RM
Aroclor 1016	ND	330	330		µg/Kg-dry	1	8/25/2020
Aroclor 1221	ND	330	330		µg/Kg-dry	1	8/25/2020
Aroclor 1232	ND	330	330		µg/Kg-dry	1	8/25/2020
Aroclor 1242	ND	330	330		µg/Kg-dry	1	8/25/2020
Aroclor 1248	ND	330	330		µg/Kg-dry	1	8/25/2020
Aroclor 1254	ND	330	330		µg/Kg-dry	1	8/25/2020
Aroclor 1260	ND	330	330		µg/Kg-dry	1	8/25/2020
Aroclor 1262	ND	330	330		µg/Kg-dry	1	8/25/2020
Aroclor 1268	ND	330	330		µg/Kg-dry	1	8/25/2020
Surr: Decachlorobiphenyl	55.7	40-140			%REC	1	8/25/2020
Surr: Tetrachloro-m-xylene	58.9	45-124			%REC	1	8/25/2020
PESTICIDES							
			SW8081A		Prep Date: 8/25/2020		Analyst: RM
4,4'-DDD	ND	20	20		µg/Kg-dry	1	8/27/2020
4,4'-DDE	20	20	20		µg/Kg-dry	1	8/27/2020
4,4'-DDT	ND	20	20		µg/Kg-dry	1	8/27/2020
Aldrin	ND	20	20		µg/Kg-dry	1	8/27/2020
alpha-BHC	ND	11	10		µg/Kg-dry	1	8/27/2020
alpha-Chlordane	ND	30	30		µg/Kg-dry	1	8/27/2020
beta-BHC	ND	20	20		µg/Kg-dry	1	8/27/2020
Chlordane, Technical	ND	30	30		µg/Kg-dry	1	8/27/2020
delta-BHC	ND	20	20		µg/Kg-dry	1	8/27/2020
Dieldrin	ND	20	20		µg/Kg-dry	1	8/27/2020
Endosulfan I	ND	20	20		µg/Kg-dry	1	8/27/2020
Endosulfan II	ND	20	20		µg/Kg-dry	1	8/27/2020
Endosulfan sulfate	ND	20	20		µg/Kg-dry	1	8/27/2020
Endrin	ND	20	20		µg/Kg-dry	1	8/27/2020
Endrin aldehyde	ND	20	20		µg/Kg-dry	1	8/27/2020
Endrin ketone	ND	20	20		µg/Kg-dry	1	8/27/2020
gamma-BHC (Lindane)	ND	20	20		µg/Kg-dry	1	8/27/2020
gamma-Chlordane	ND	30	30		µg/Kg-dry	1	8/27/2020
Heptachlor	ND	20	20		µg/Kg-dry	1	8/27/2020
Heptachlor epoxide	ND	20	20		µg/Kg-dry	1	8/27/2020
Methoxychlor	ND	50	50		µg/Kg-dry	1	8/27/2020
Toxaphene	ND	170	170		µg/Kg-dry	1	8/27/2020

Note: See Qualifiers page for a list of qualifiers and their definitions.

ALS Group, USA

Date: 31-Aug-20

Client: AKT Peerless
Work Order: 20081693
Project: 8997f5-5-10.4
Lab ID: 20081693-01

Client Sample ID: 9156 Lakepointe-TS-Composite
Collection Date: 8/19/2020 11:35:00 AM
Matrix: SOIL

Analyses	Result	Report Limit	MDEQ OP Memo 2 TDL	Qual	Units	Dilution Factor	Date Analyzed
Surr: Decachlorobiphenyl	52.4	50-150			%REC	1	8/27/2020
Surr: Tetrachloro-m-xylene	56.6	50-150			%REC	1	8/27/2020
MERCURY BY CVAA			SW7471B		Prep Date: 8/27/2020		Analyst: ABL
Mercury	190	50	50		µg/Kg-dry	1	8/27/2020
METALS BY ICP-MS			SW6020B		Prep Date: 8/26/2020		Analyst: STP
Arsenic	3,400	2,000	2,000		µg/Kg-dry	1	8/26/2020
Barium	40,000	1,000	1,000		µg/Kg-dry	1	8/26/2020
Cadmium	290	200	200		µg/Kg-dry	1	8/26/2020
Chromium	7,900	2,000	2,000		µg/Kg-dry	1	8/26/2020
Copper	7,400	1,000	1,000		µg/Kg-dry	1	8/26/2020
Lead	18,000	10,000	10,000		µg/Kg-dry	1	8/26/2020
Selenium	ND	420	200		µg/Kg-dry	1	8/26/2020
Silver	ND	420	100		µg/Kg-dry	1	8/26/2020
Zinc	29,000	8,500	1,000		µg/Kg-dry	10	8/27/2020
SEMI-VOLATILE ORGANIC COMPOUNDS			SW846 8270D		Prep Date: 8/26/2020		Analyst: EEW
1,2,4-Trichlorobenzene	ND	360	330		µg/Kg-dry	10	8/27/2020
1,2-Dichlorobenzene	ND	360	330		µg/Kg-dry	10	8/27/2020
1,3-Dichlorobenzene	ND	360	330		µg/Kg-dry	10	8/27/2020
1,4-Dichlorobenzene	ND	360	330		µg/Kg-dry	10	8/27/2020
2,4,5-Trichlorophenol	ND	360	300		µg/Kg-dry	10	8/27/2020
2,4,6-Trichlorophenol	ND	360	330		µg/Kg-dry	10	8/27/2020
2,4-Dichlorophenol	ND	360	330		µg/Kg-dry	10	8/27/2020
2,4-Dimethylphenol	ND	360	330		µg/Kg-dry	10	8/27/2020
2,4-Dinitrophenol	ND	830	830		µg/Kg-dry	10	8/27/2020
2,4-Dinitrotoluene	ND	360	330		µg/Kg-dry	10	8/27/2020
2,6-Dinitrotoluene	ND	360	330		µg/Kg-dry	10	8/27/2020
2-Chloronaphthalene	ND	330	330		µg/Kg-dry	10	8/27/2020
2-Chlorophenol	ND	360	330		µg/Kg-dry	10	8/27/2020
2-Methylnaphthalene	ND	330	330		µg/Kg-dry	10	8/27/2020
2-Methylphenol	ND	360	330		µg/Kg-dry	10	8/27/2020
2-Nitroaniline	ND	830	830		µg/Kg-dry	10	8/27/2020
2-Nitrophenol	ND	360	330		µg/Kg-dry	10	8/27/2020
3&4-Methylphenol	ND	360	330		µg/Kg-dry	10	8/27/2020
3,3'-Dichlorobenzidine	ND	2,000	2,000		µg/Kg-dry	10	8/27/2020
3-Nitroaniline	ND	830	830		µg/Kg-dry	10	8/27/2020
4,6-Dinitro-2-methylphenol	ND	830	830		µg/Kg-dry	10	8/27/2020
4-Bromophenyl phenyl ether	ND	360	330		µg/Kg-dry	10	8/27/2020
4-Chloro-3-methylphenol	ND	360	280		µg/Kg-dry	10	8/27/2020
4-Chloroaniline	ND	730	330		µg/Kg-dry	10	8/27/2020

Note: See Qualifiers page for a list of qualifiers and their definitions.

ALS Group, USA

Date: 31-Aug-20

Client: AKT Peerless
Work Order: 20081693
Project: 8997f5-5-10.4
Lab ID: 20081693-01

Client Sample ID: 9156 Lakepointe-TS-Composite
Collection Date: 8/19/2020 11:35:00 AM

Matrix: SOIL

Analyses	Result	Report Limit	MDEQ OP Memo 2 TDL	Qual	Units	Dilution Factor	Date Analyzed
4-Chlorophenyl phenyl ether	ND	360	330		µg/Kg-dry	10	8/27/2020
4-Nitroaniline	ND	1,800	830		µg/Kg-dry	10	8/27/2020
4-Nitrophenol	ND	830	830		µg/Kg-dry	10	8/27/2020
Acenaphthene	ND	330	330		µg/Kg-dry	10	8/27/2020
Acenaphthylene	ND	330	330		µg/Kg-dry	10	8/27/2020
Anthracene	ND	330	330		µg/Kg-dry	10	8/27/2020
Benzo(a)anthracene	ND	330	330		µg/Kg-dry	10	8/27/2020
Benzo(a)pyrene	ND	330	330		µg/Kg-dry	10	8/27/2020
Benzo(b)fluoranthene	ND	330	330		µg/Kg-dry	10	8/27/2020
Benzo(g,h,i)perylene	ND	330	330		µg/Kg-dry	10	8/27/2020
Benzo(k)fluoranthene	ND	330	330		µg/Kg-dry	10	8/27/2020
Bis(2-chloroethoxy)methane	ND	360	330		µg/Kg-dry	10	8/27/2020
Bis(2-chloroethyl)ether	ND	360	100		µg/Kg-dry	10	8/27/2020
Bis(2-chloroisopropyl)ether	ND	360	330		µg/Kg-dry	10	8/27/2020
Bis(2-ethylhexyl)phthalate	ND	360	330		µg/Kg-dry	10	8/27/2020
Butyl benzyl phthalate	ND	360	330		µg/Kg-dry	10	8/27/2020
Carbazole	ND	360	330		µg/Kg-dry	10	8/27/2020
Chrysene	ND	330	330		µg/Kg-dry	10	8/27/2020
Dibenzo(a,h)anthracene	ND	330	330		µg/Kg-dry	10	8/27/2020
Dibenzofuran	ND	360	330		µg/Kg-dry	10	8/27/2020
Diethyl phthalate	ND	360	330		µg/Kg-dry	10	8/27/2020
Dimethyl phthalate	ND	360	330		µg/Kg-dry	10	8/27/2020
Di-n-butyl phthalate	ND	360	330		µg/Kg-dry	10	8/27/2020
Di-n-octyl phthalate	ND	360	330		µg/Kg-dry	10	8/27/2020
Fluoranthene	ND	330	330		µg/Kg-dry	10	8/27/2020
Fluorene	ND	330	330		µg/Kg-dry	10	8/27/2020
Hexachlorobenzene	ND	360	330		µg/Kg-dry	10	8/27/2020
Hexachlorobutadiene	ND	360	50		µg/Kg-dry	10	8/27/2020
Hexachlorocyclopentadiene	ND	360	330		µg/Kg-dry	10	8/27/2020
Hexachloroethane	ND	360	300		µg/Kg-dry	10	8/27/2020
Indeno(1,2,3-cd)pyrene	ND	330	330		µg/Kg-dry	10	8/27/2020
Isophorone	ND	1,800	330		µg/Kg-dry	10	8/27/2020
Naphthalene	ND	330	330		µg/Kg-dry	10	8/27/2020
Nitrobenzene	ND	1,800	330		µg/Kg-dry	10	8/27/2020
N-Nitrosodi-n-propylamine	ND	360	330		µg/Kg-dry	10	8/27/2020
N-Nitrosodiphenylamine	ND	360	330		µg/Kg-dry	10	8/27/2020
Pentachlorophenol	ND	360	20		µg/Kg-dry	10	8/27/2020
Phenanthrene	ND	330	330		µg/Kg-dry	10	8/27/2020
Phenol	ND	360	330		µg/Kg-dry	10	8/27/2020
Pyrene	ND	330	330		µg/Kg-dry	10	8/27/2020
Surr: 2,4,6-Tribromophenol	66.4	38-92			%REC	10	8/27/2020

Note: See Qualifiers page for a list of qualifiers and their definitions.

ALS Group, USA

Date: 31-Aug-20

Client: AKT Peerless
Work Order: 20081693
Project: 8997f5-5-10.4
Lab ID: 20081693-01

Client Sample ID: 9156 Lakepointe-TS-Composite
Collection Date: 8/19/2020 11:35:00 AM

Matrix: SOIL

Analyses	Result	Report Limit	MDEQ OP Memo 2 TDL	Qual	Units	Dilution Factor	Date Analyzed
Surr: 2-Fluorobiphenyl	74.4	44-107			%REC	10	8/27/2020
Surr: 2-Fluorophenol	68.6	37-109			%REC	10	8/27/2020
Surr: 4-Terphenyl-d14	111	52-123			%REC	10	8/27/2020
Surr: Nitrobenzene-d5	69.4	41-94			%REC	10	8/27/2020
Surr: Phenol-d6	72.6	28-111			%REC	10	8/27/2020

VOLATILE ORGANIC COMPOUNDS

SW8260C

Prep Date: 8/25/2020

Analyst: MF

1,1,1,2-Tetrachloroethane	ND	100	100		µg/Kg-dry	1	8/25/2020
1,1,1-Trichloroethane	ND	50	50		µg/Kg-dry	1	8/25/2020
1,1,2,2-Tetrachloroethane	ND	50	50		µg/Kg-dry	1	8/25/2020
1,1,2-Trichloroethane	ND	50	50		µg/Kg-dry	1	8/25/2020
1,1,2-Trichlorotrifluoroethane	ND	100	100		µg/Kg-dry	1	8/25/2020
1,1-Dichloroethane	ND	50	50		µg/Kg-dry	1	8/25/2020
1,1-Dichloroethene	ND	50	50		µg/Kg-dry	1	8/25/2020
1,2,3-Trichloropropane	ND	100	100		µg/Kg-dry	1	8/25/2020
1,2,4-Trichlorobenzene	ND	250	250		µg/Kg-dry	1	8/25/2020
1,2,4-Trimethylbenzene	ND	100	100		µg/Kg-dry	1	8/25/2020
1,2-Dibromo-3-chloropropane	ND	130	10		µg/Kg-dry	1	8/25/2020
1,2-Dibromoethane	ND	38	20		µg/Kg-dry	1	8/25/2020
1,2-Dichlorobenzene	ND	100	100		µg/Kg-dry	1	8/25/2020
1,2-Dichloroethane	ND	130	50		µg/Kg-dry	1	8/25/2020
1,2-Dichloropropane	ND	50	50		µg/Kg-dry	1	8/25/2020
1,3,5-Trimethylbenzene	ND	130	100		µg/Kg-dry	1	8/25/2020
1,3-Dichlorobenzene	ND	100	100		µg/Kg-dry	1	8/25/2020
1,4-Dichlorobenzene	ND	100	100		µg/Kg-dry	1	8/25/2020
2-Butanone	ND	750	750		µg/Kg-dry	1	8/25/2020
2-Hexanone	ND	2,500	2,500		µg/Kg-dry	1	8/25/2020
2-Methylnaphthalene	ND	330	330		µg/Kg-dry	1	8/25/2020
4-Methyl-2-pentanone	ND	2,500	2,500		µg/Kg-dry	1	8/25/2020
Acetone	ND	1,000	1,000		µg/Kg-dry	1	8/25/2020
Acrylonitrile	ND	130	100		µg/Kg-dry	1	8/25/2020
Benzene	ND	50	50		µg/Kg-dry	1	8/25/2020
Bromodichloromethane	ND	100	100		µg/Kg-dry	1	8/25/2020
Bromoform	ND	100	100		µg/Kg-dry	1	8/25/2020
Bromomethane	ND	200	200		µg/Kg-dry	1	8/25/2020
Carbon disulfide	ND	250	250		µg/Kg-dry	1	8/25/2020
Carbon tetrachloride	ND	50	50		µg/Kg-dry	1	8/25/2020
Chlorobenzene	ND	50	50		µg/Kg-dry	1	8/25/2020
Chloroethane	ND	250	250		µg/Kg-dry	1	8/25/2020
Chloroform	ND	50	50		µg/Kg-dry	1	8/25/2020
Chloromethane	ND	250	250		µg/Kg-dry	1	8/25/2020

Note: See Qualifiers page for a list of qualifiers and their definitions.

ALS Group, USA

Date: 31-Aug-20

Client: AKT Peerless
Work Order: 20081693
Project: 8997f5-5-10.4
Lab ID: 20081693-01

Client Sample ID: 9156 Lakepointe-TS-Composite
Collection Date: 8/19/2020 11:35:00 AM

Matrix: SOIL

Analyses	Result	Report Limit	MDEQ OP Memo 2 TDL	Qual	Units	Dilution Factor	Date Analyzed
cis-1,2-Dichloroethene	ND	50	50		µg/Kg-dry	1	8/25/2020
cis-1,3-Dichloropropene	ND	50	50		µg/Kg-dry	1	8/25/2020
Dibromochloromethane	ND	100	100		µg/Kg-dry	1	8/25/2020
Dibromomethane	ND	250	250		µg/Kg-dry	1	8/25/2020
Dichlorodifluoromethane	ND	250	250		µg/Kg-dry	1	8/25/2020
Diethyl ether	ND	200	200		µg/Kg-dry	1	8/25/2020
Ethylbenzene	ND	50	50		µg/Kg-dry	1	8/25/2020
Hexachloroethane	ND	300	300		µg/Kg-dry	1	8/25/2020
Isopropylbenzene	ND	250	250		µg/Kg-dry	1	8/25/2020
m,p-Xylene	ND	100	100		µg/Kg-dry	1	8/25/2020
Methyl tert-butyl ether	ND	250	250		µg/Kg-dry	1	8/25/2020
Methylene chloride	ND	310	100		µg/Kg-dry	1	8/25/2020
Naphthalene	ND	330	330		µg/Kg-dry	1	8/25/2020
n-Propylbenzene	ND	100	100		µg/Kg-dry	1	8/25/2020
o-Xylene	ND	50	50		µg/Kg-dry	1	8/25/2020
Styrene	ND	50	50		µg/Kg-dry	1	8/25/2020
Tetrachloroethene	ND	50	50		µg/Kg-dry	1	8/25/2020
Toluene	ND	100	100		µg/Kg-dry	1	8/25/2020
trans-1,2-Dichloroethene	ND	50	50		µg/Kg-dry	1	8/25/2020
trans-1,3-Dichloropropene	ND	50	50		µg/Kg-dry	1	8/25/2020
Trichloroethene	ND	50	50		µg/Kg-dry	1	8/25/2020
Trichlorofluoromethane	ND	100	100		µg/Kg-dry	1	8/25/2020
Vinyl acetate	ND	5,000	5,000		µg/Kg-dry	1	8/25/2020
Vinyl chloride	ND	40	40		µg/Kg-dry	1	8/25/2020
Xylenes, Total	ND	150	150		µg/Kg-dry	1	8/25/2020
Surr: 1,2-Dichloroethane-d4	101	70-130			%REC	1	8/25/2020
Surr: 4-Bromofluorobenzene	103	70-130			%REC	1	8/25/2020
Surr: Dibromofluoromethane	99.1	70-130			%REC	1	8/25/2020
Surr: Toluene-d8	99.8	70-130			%REC	1	8/25/2020
CHLORIDE			A4500-CL E-11		Prep Date: 8/25/2020		Analyst: JDR
Chloride	ND	200	200		mg/Kg-dry	1	8/27/2020
MOISTURE			SW3550C				Analyst: KTP
Moisture	8.8	0.10	0		% of sample	1	8/26/2020

Note: See Qualifiers page for a list of qualifiers and their definitions.

Client: AKT Peerless

Work Order: 20081693

Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: 163110

Instrument ID GC14

Method: SW8082

MBLK				Sample ID: PBLKS1-163110-163110		Units: µg/Kg		Analysis Date: 8/25/2020 04:31 PM		
Client ID:		Run ID: GC14_200825A		SeqNo: 6661723		Prep Date: 8/25/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Aroclor 1016	ND	67								
Aroclor 1221	ND	67								
Aroclor 1232	ND	67								
Aroclor 1242	ND	67								
Aroclor 1248	ND	67								
Aroclor 1254	ND	67								
Aroclor 1260	ND	67								
Aroclor 1262	ND	67								
Aroclor 1268	ND	67								
Surr: Decachlorobiphenyl	33	0	33.3	0	99.1	40-140	0			
Surr: Tetrachloro-m-xylene	35.3	0	33.3	0	106	45-124	0			

LCS				Sample ID: PLCSS1-163110-163110		Units: µg/Kg		Analysis Date: 8/25/2020 04:46 PM		
Client ID:		Run ID: GC14_200825A		SeqNo: 6661724		Prep Date: 8/25/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Aroclor 1016	844.9	67	833	0	101	50-130	0			
Aroclor 1260	819.6	67	833	0	98.4	50-130	0			
Surr: Decachlorobiphenyl	35.02	0	33.3	0	105	40-140	0			
Surr: Tetrachloro-m-xylene	36.38	0	33.3	0	109	45-124	0			

MS				Sample ID: 20081661-01B MS		Units: µg/Kg		Analysis Date: 8/25/2020 05:48 PM		
Client ID:		Run ID: GC14_200825A		SeqNo: 6661726		Prep Date: 8/25/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Aroclor 1016	770.7	64	803.5	0	95.9	40-140	0			
Aroclor 1260	717.5	64	803.5	0	89.3	40-140	0			
Surr: Decachlorobiphenyl	29.23	0	32.12	0	91	40-140	0			
Surr: Tetrachloro-m-xylene	32.41	0	32.12	0	101	45-124	0			

MSD				Sample ID: 20081661-01B MSD		Units: µg/Kg		Analysis Date: 8/25/2020 06:03 PM		
Client ID:		Run ID: GC14_200825A		SeqNo: 6661727		Prep Date: 8/25/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Aroclor 1016	764.5	65	814.3	0	93.9	40-140	770.7	0.811	50	
Aroclor 1260	721.6	65	814.3	0	88.6	40-140	717.5	0.582	50	
Surr: Decachlorobiphenyl	30.3	0	32.55	0	93.1	40-140	29.23	3.62	50	
Surr: Tetrachloro-m-xylene	32.57	0	32.55	0	100	45-124	32.41	0.487	50	

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
Work Order: 20081693
Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: **163110** Instrument ID **GC14** Method: **SW8082**

The following samples were analyzed in this batch:

20081693-01B

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: **163111** Instrument ID **GC12** Method: **SW8081A**

MBLK				Sample ID: PBLKS1-163111-163111			Units: µg/Kg		Analysis Date: 8/26/2020 12:15 AM		
Client ID:			Run ID: GC12_200825A			SeqNo: 6665092		Prep Date: 8/25/2020		DF: 1	
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual	
4,4´-DDD	ND	10									
4,4´-DDE	ND	10									
4,4´-DDT	ND	10									
Aldrin	ND	10									
alpha-BHC	ND	10									
alpha-Chlordane	ND	10									
beta-BHC	ND	10									
Chlordane, Technical	ND	25									
delta-BHC	ND	10									
Dieldrin	ND	10									
Endosulfan I	ND	10									
Endosulfan II	ND	10									
Endosulfan sulfate	ND	10									
Endrin	ND	10									
Endrin aldehyde	ND	10									
Endrin ketone	ND	10									
gamma-BHC (Lindane)	ND	10									
gamma-Chlordane	ND	10									
Heptachlor	ND	10									
Heptachlor epoxide	ND	10									
Methoxychlor	ND	10									
Toxaphene	ND	60									
Surr: Decachlorobiphenyl	31.22	0	33.3	0	93.7	50-150		0			
Surr: Tetrachloro-m-xylene	30.95	0	33.3	0	92.9	50-150		0			

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: **163111** Instrument ID **GC12** Method: **SW8081A**

LCS				Sample ID: PLCSS1-163111-163111		Units: µg/Kg		Analysis Date: 8/26/2020 12:29 AM		
Client ID:			Run ID: GC12_200825A			SeqNo: 6665093		Prep Date: 8/25/2020		DF: 1
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
4,4'-DDD	30.48	10	33.33	0	91.5	50-150	0			
4,4'-DDE	29.65	10	33.33	0	89	50-150	0			
4,4'-DDT	24.72	10	33.33	0	74.2	50-150	0			
Aldrin	30.05	10	33.33	0	90.2	50-150	0			
alpha-BHC	30.13	10	33.33	0	90.4	50-150	0			
alpha-Chlordane	30	10	33.33	0	90	50-150	0			
beta-BHC	29.68	10	33.33	0	89.1	50-150	0			
delta-BHC	29.88	10	33.33	0	89.7	50-150	0			
Dieldrin	30.1	10	33.33	0	90.3	50-150	0			
Endosulfan I	29.8	10	33.33	0	89.4	50-150	0			
Endosulfan II	29.72	10	33.33	0	89.2	50-150	0			
Endosulfan sulfate	30.43	10	33.33	0	91.3	50-150	0			
Endrin	28.33	10	33.33	0	85	50-150	0			
Endrin aldehyde	30.67	10	33.33	0	92	50-150	0			
Endrin ketone	30.25	10	33.33	0	90.8	50-150	0			
gamma-BHC (Lindane)	29.78	10	33.33	0	89.4	50-150	0			
gamma-Chlordane	29.8	10	33.33	0	89.4	50-150	0			
Heptachlor	26.53	10	33.33	0	79.6	50-150	0			
Heptachlor epoxide	30	10	33.33	0	90	50-150	0			
Methoxychlor	25.35	10	33.33	0	76.1	50-150	0			
<i>Surr: Decachlorobiphenyl</i>	30.37	0	33.3	0	91.2	50-150	0			
<i>Surr: Tetrachloro-m-xylene</i>	30.67	0	33.3	0	92.1	50-150	0			

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: **163111** Instrument ID **GC12** Method: **SW8081A**

MS				Sample ID: 20081661-01B MS		Units: µg/Kg		Analysis Date: 8/26/2020 12:42 AM		
Client ID:			Run ID: GC12_200825A			SeqNo: 6665094		Prep Date: 8/25/2020		DF: 1
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
4,4'-DDD	25.23	9.7	32.43	0	77.8	50-150	0			
4,4'-DDE	23.76	9.7	32.43	0	73.3	50-150	0			
4,4'-DDT	20.25	9.7	32.43	0	62.5	50-150	0			
Aldrin	23.84	9.7	32.43	0	73.5	50-150	0			
alpha-BHC	24.54	9.7	32.43	0	75.7	50-150	0			
alpha-Chlordane	23.76	9.7	32.43	0	73.3	50-150	0			
beta-BHC	23.73	9.7	32.43	0	73.2	50-150	0			
delta-BHC	23.42	9.7	32.43	0	72.2	50-150	0			
Dieldrin	23.84	9.7	32.43	0	73.5	50-150	0			
Endosulfan I	23.76	9.7	32.43	0	73.3	50-150	0			
Endosulfan II	23.43	9.7	32.43	0	72.3	50-150	0			
Endosulfan sulfate	23.69	9.7	32.43	0	73.1	50-150	0			
Endrin	23.5	9.7	32.43	0	72.5	50-150	0			
Endrin aldehyde	22.44	9.7	32.43	0	69.2	50-150	0			
Endrin ketone	23.66	9.7	32.43	0	73	50-150	0			
gamma-BHC (Lindane)	24.05	9.7	32.43	0	74.2	50-150	0			
gamma-Chlordane	23.3	9.7	32.43	0	71.9	50-150	0			
Heptachlor	21.36	9.7	32.43	0	65.9	50-150	0			
Heptachlor epoxide	23.58	9.7	32.43	0	72.7	50-150	0			
Methoxychlor	20.94	9.7	32.43	0	64.6	50-150	0			
Surr: Decachlorobiphenyl	24.46	0	32.4	0	75.5	50-150	0			
Surr: Tetrachloro-m-xylene	24.99	0	32.4	0	77.1	50-150	0			

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: 163111 Instrument ID GC12 Method: SW8081A

MSD				Sample ID: 20081661-01B MSD		Units: µg/Kg		Analysis Date: 8/26/2020 12:56 AM		
Client ID:		Run ID: GC12_200825A			SeqNo: 6665095		Prep Date: 8/25/2020		DF: 1	
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
4,4'-DDD	31.15	9.9	33.04	0	94.3	50-150	25.23	21	35	
4,4'-DDE	28.78	9.9	33.04	0	87.1	50-150	23.76	19.1	35	
4,4'-DDT	22.87	9.9	33.04	0	69.2	50-150	20.25	12.1	35	
Aldrin	28.95	9.9	33.04	0	87.6	50-150	23.84	19.4	35	
alpha-BHC	29.38	9.9	33.04	0	88.9	50-150	24.54	18	35	
alpha-Chlordane	29.06	9.9	33.04	0	88	50-150	23.76	20.1	35	
beta-BHC	28.7	9.9	33.04	0	86.9	50-150	23.73	19	35	
delta-BHC	28.44	9.9	33.04	0	86.1	50-150	23.42	19.4	35	
Dieldrin	29.28	9.9	33.04	0	88.6	50-150	23.84	20.5	35	
Endosulfan I	28.97	9.9	33.04	0	87.7	50-150	23.76	19.8	35	
Endosulfan II	29.15	9.9	33.04	0	88.2	50-150	23.43	21.7	35	
Endosulfan sulfate	29.61	9.9	33.04	0	89.6	50-150	23.69	22.2	35	
Endrin	28.35	9.9	33.04	0	85.8	50-150	23.5	18.7	35	
Endrin aldehyde	28.27	9.9	33.04	0	85.6	50-150	22.44	23	35	
Endrin ketone	29.3	9.9	33.04	0	88.7	50-150	23.66	21.3	35	
gamma-BHC (Lindane)	29	9.9	33.04	0	87.8	50-150	24.05	18.7	35	
gamma-Chlordane	28.54	9.9	33.04	0	86.4	50-150	23.3	20.2	35	
Heptachlor	25.28	9.9	33.04	0	76.5	50-150	21.36	16.8	35	
Heptachlor epoxide	28.65	9.9	33.04	0	86.7	50-150	23.58	19.4	35	
Methoxychlor	24.12	9.9	33.04	0	73	50-150	20.94	14.2	35	
Surr: Decachlorobiphenyl	29.97	0	33.01	0	90.8	50-150	24.46	20.3	35	
Surr: Tetrachloro-m-xylene	29.54	0	33.01	0	89.5	50-150	24.99	16.7	35	

The following samples were analyzed in this batch:

20081693-01B

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: **163179** Instrument ID **GC7** Method: **SW8151**

MBLK				Sample ID: HBLKS1-163179-163179		Units: µg/Kg		Analysis Date: 8/29/2020 04:15 AM		
Client ID:		Run ID: GC7_200828A		SeqNo: 6671170		Prep Date: 8/26/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual

2,4,5-TP (Silvex)

ND

5.0

2,4-D

ND

10

Surr: DCAA

12

0

50

0

24

10-150

0

LCS				Sample ID: HLCSS1-163179-163179		Units: µg/Kg		Analysis Date: 8/29/2020 04:43 AM		
Client ID:		Run ID: GC7_200828A		SeqNo: 6671172		Prep Date: 8/26/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual

2,4,5-TP (Silvex)

6

5.0

50

0

12

10-150

0

2,4-D

23

10

50

0

46

10-130

0

Surr: DCAA

8

0

50

0

16

10-150

0

MS				Sample ID: 20081661-01B MS		Units: µg/Kg		Analysis Date: 8/29/2020 04:57 AM		
Client ID:		Run ID: GC7_200828A		SeqNo: 6671173		Prep Date: 8/26/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual

2,4,5-TP (Silvex)

27.79

5.0

49.63

0

56

10-150

0

2,4-D

77.43

9.9

49.63

0

156

10-130

0

S

Surr: DCAA

23.82

0

49.63

0

48

10-150

0

MSD				Sample ID: 20081661-01B MSD		Units: µg/Kg		Analysis Date: 8/29/2020 05:10 AM		
Client ID:		Run ID: GC7_200828A		SeqNo: 6671174		Prep Date: 8/26/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual

2,4,5-TP (Silvex)

17.73

4.9

49.24

0

36

10-150

27.79

44.2

50

2,4-D

80.75

9.8

49.24

0

164

10-130

77.43

4.2

50

S

Surr: DCAA

13.79

0

49.24

0

28

10-150

23.82

53.4

50

R

The following samples were analyzed in this batch:

20081693-01B

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: **163255** Instrument ID **HG4** Method: **SW7471B**

MBLK		Sample ID: MBLK-163255-163255				Units: mg/Kg		Analysis Date: 8/27/2020 12:48 PM		
Client ID:		Run ID: HG4_200827A				SeqNo: 6665593		Prep Date: 8/27/2020		DF: 1
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual

Mercury ND 0.020

LCS		Sample ID: LCS-163255-163255				Units: mg/Kg		Analysis Date: 8/27/2020 12:49 PM		
Client ID:		Run ID: HG4_200827A				SeqNo: 6665594		Prep Date: 8/27/2020		DF: 1
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual

Mercury 0.1792 0.020 0.1665 0 108 80-120 0

MS		Sample ID: 20081682-01BMS				Units: mg/Kg		Analysis Date: 8/27/2020 12:53 PM		
Client ID:		Run ID: HG4_200827A				SeqNo: 6665596		Prep Date: 8/27/2020		DF: 1
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual

Mercury 0.3041 0.018 0.1471 0.1392 112 75-125 0 E

MSD		Sample ID: 20081682-01BMSD				Units: mg/Kg		Analysis Date: 8/27/2020 12:55 PM		
Client ID:		Run ID: HG4_200827A				SeqNo: 6665597		Prep Date: 8/27/2020		DF: 1
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual

Mercury 0.3139 0.018 0.153 0.1392 114 75-125 0.3041 3.18 35 E

The following samples were analyzed in this batch:

20081693-01B

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: **163147** Instrument ID **ICPMS3** Method: **SW6020B**

MBLK Sample ID: MBLK-163147-163147				Units: mg/Kg		Analysis Date: 8/26/2020 10:11 PM				
Client ID:		Run ID: ICPMS3_200826B		SeqNo: 6663848		Prep Date: 8/26/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Arsenic	ND	0.25								
Barium	ND	0.25								
Cadmium	ND	0.10								
Chromium	ND	0.25								
Copper	ND	0.25								
Lead	ND	0.25								
Selenium	ND	0.25								
Silver	ND	0.25								
Zinc	ND	0.50								

LCS Sample ID: LCS-163147-163147				Units: mg/Kg		Analysis Date: 8/26/2020 10:12 PM				
Client ID:		Run ID: ICPMS3_200826B		SeqNo: 6663849		Prep Date: 8/26/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Arsenic	5.095	0.25	5	0	102	80-120	0			
Barium	5.164	0.25	5	0	103	80-120	0			
Cadmium	5.22	0.10	5	0	104	80-120	0			
Chromium	5.221	0.25	5	0	104	80-120	0			
Copper	5.268	0.25	5	0	105	80-120	0			
Lead	5.219	0.25	5	0	104	80-120	0			
Selenium	5.088	0.25	5	0	102	80-120	0			
Silver	5.219	0.25	5	0	104	80-120	0			
Zinc	5.218	0.50	5	0	104	80-120	0			

MS Sample ID: 20081696-01BMS				Units: mg/Kg		Analysis Date: 8/26/2020 10:59 PM				
Client ID:		Run ID: ICPMS3_200826B		SeqNo: 6663876		Prep Date: 8/26/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Arsenic	9.983	0.36	7.257	4.389	77.1	75-125	0			
Barium	44.42	0.36	7.257	38.81	77.3	75-125	0			O
Cadmium	5.809	0.15	7.257	0.1727	77.7	75-125	0			
Chromium	15.75	0.36	7.257	12.96	38.4	75-125	0			S
Copper	15.02	0.36	7.257	10.36	64.2	75-125	0			S
Lead	24.39	0.36	7.257	24.15	3.34	75-125	0			S
Selenium	6.217	0.36	7.257	0.2891	81.7	75-125	0			
Silver	5.663	0.36	7.257	0.0318	77.6	75-125	0			

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: **163147** Instrument ID **ICPMS3** Method: **SW6020B**

MS				Sample ID: 20081696-01BMS		Units: mg/Kg		Analysis Date: 8/27/2020 04:56 PM		
Client ID:		Run ID: ICPMS3_200827B		SeqNo: 6666585		Prep Date: 8/26/2020		DF: 10		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Zinc	51.01	7.3	7.257	52.02	-14	75-125	0			SO

MSD				Sample ID: 20081696-01BMSD		Units: mg/Kg		Analysis Date: 8/26/2020 11:01 PM		
Client ID:		Run ID: ICPMS3_200826B		SeqNo: 6663877		Prep Date: 8/26/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Arsenic	11.5	0.39	7.716	4.389	92.2	75-125	9.983	14.1	20	
Barium	47.57	0.39	7.716	38.81	113	75-125	44.42	6.84	20	O
Cadmium	6.742	0.15	7.716	0.1727	85.1	75-125	5.809	14.9	20	
Chromium	20.49	0.39	7.716	12.96	97.6	75-125	15.75	26.2	20	R
Copper	16.78	0.39	7.716	10.36	83.3	75-125	15.02	11.1	20	
Lead	29.41	0.39	7.716	24.15	68.2	75-125	24.39	18.7	20	S
Selenium	6.876	0.39	7.716	0.2891	85.4	75-125	6.217	10.1	20	
Silver	6.551	0.39	7.716	0.0318	84.5	75-125	5.663	14.5	20	

MSD				Sample ID: 20081696-01BMSD		Units: mg/Kg		Analysis Date: 8/27/2020 04:58 PM		
Client ID:		Run ID: ICPMS3_200827B		SeqNo: 6666586		Prep Date: 8/26/2020		DF: 10		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Zinc	55.8	7.7	7.716	52.02	49	75-125	51.01	8.98	20	SO

The following samples were analyzed in this batch:

20081693-01B

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: 163196 Instrument ID SVMS10 Method: SW846 8270D

MBLK		Sample ID: SBLKS1-163196-163196				Units: µg/Kg		Analysis Date: 8/26/2020 10:14 PM		
Client ID:		Run ID: SVMS10_200826A				SeqNo: 6665555		Prep Date: 8/26/2020		DF: 1
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,2,4-Trichlorobenzene	ND	33								
1,2-Dichlorobenzene	ND	33								
1,3-Dichlorobenzene	ND	33								
1,4-Dichlorobenzene	ND	33								
2,4,5-Trichlorophenol	ND	33								
2,4,6-Trichlorophenol	ND	33								
2,4-Dichlorophenol	ND	33								
2,4-Dimethylphenol	ND	33								
2,4-Dinitrophenol	ND	33								
2,4-Dinitrotoluene	ND	33								
2,6-Dinitrotoluene	ND	33								
2-Chloronaphthalene	ND	6.7								
2-Chlorophenol	ND	33								
2-Methylnaphthalene	ND	6.7								
2-Methylphenol	ND	33								
2-Nitroaniline	ND	33								
2-Nitrophenol	ND	33								
3&4-Methylphenol	ND	33								
3,3'-Dichlorobenzidine	ND	170								
3-Nitroaniline	ND	33								
4,6-Dinitro-2-methylphenol	ND	33								
4-Bromophenyl phenyl ether	ND	33								
4-Chloro-3-methylphenol	ND	33								
4-Chloroaniline	ND	67								
4-Chlorophenyl phenyl ether	ND	33								
4-Nitroaniline	ND	170								
4-Nitrophenol	ND	33								
Acenaphthene	ND	6.7								
Acenaphthylene	ND	6.7								
Anthracene	ND	6.7								
Benzo(a)anthracene	ND	6.7								
Benzo(a)pyrene	ND	6.7								
Benzo(b)fluoranthene	ND	6.7								
Benzo(g,h,i)perylene	ND	6.7								
Benzo(k)fluoranthene	ND	6.7								
Bis(2-chloroethoxy)methane	ND	33								
Bis(2-chloroethyl)ether	ND	33								
Bis(2-chloroisopropyl)ether	ND	33								
Bis(2-ethylhexyl)phthalate	ND	33								
Butyl benzyl phthalate	ND	33								
Carbazole	ND	33								
Chrysene	ND	6.7								

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
Work Order: 20081693
Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: 163196		Instrument ID SVMS10		Method: SW846 8270D				
Dibenzo(a,h)anthracene	ND	6.7						
Dibenzofuran	ND	33						
Diethyl phthalate	ND	33						
Dimethyl phthalate	ND	33						
Di-n-butyl phthalate	ND	33						
Di-n-octyl phthalate	ND	33						
Fluoranthene	ND	6.7						
Fluorene	ND	6.7						
Hexachlorobenzene	ND	33						
Hexachlorobutadiene	ND	33						
Hexachlorocyclopentadiene	ND	33						
Hexachloroethane	ND	33						
Indeno(1,2,3-cd)pyrene	ND	6.7						
Isophorone	ND	170						
Naphthalene	ND	6.7						
Nitrobenzene	ND	170						
N-Nitrosodi-n-propylamine	ND	33						
N-Nitrosodiphenylamine	ND	33						
Pentachlorophenol	ND	33						
Phenanthrene	ND	6.7						
Phenol	ND	33						
Pyrene	ND	6.7						
<i>Surr: 2,4,6-Tribromophenol</i>	2327	0	3333	0	69.8	38-92	0	
<i>Surr: 2-Fluorobiphenyl</i>	2401	0	3333	0	72	44-107	0	
<i>Surr: 2-Fluorophenol</i>	2373	0	3333	0	71.2	37-109	0	
<i>Surr: 4-Terphenyl-d14</i>	3023	0	3333	0	90.7	52-123	0	
<i>Surr: Nitrobenzene-d5</i>	2311	0	3333	0	69.3	41-94	0	
<i>Surr: Phenol-d6</i>	2539	0	3333	0	76.2	28-111	0	

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: 163196 Instrument ID SVMS10 Method: SW846 8270D

LCS Sample ID: SLCSS1-163196-163196				Units: µg/Kg		Analysis Date: 8/26/2020 10:41 PM				
Client ID:		Run ID: SVMS10_200826A		SeqNo: 6665556		Prep Date: 8/26/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,2,4-Trichlorobenzene	941.3	33	1333	0	70.6	40-93	0			
1,2-Dichlorobenzene	928.7	33	1333	0	69.7	42-94	0			
1,3-Dichlorobenzene	940.7	33	1333	0	70.6	41-94	0			
1,4-Dichlorobenzene	973.3	33	1333	0	73	42-94	0			
2,4,5-Trichlorophenol	1065	33	1333	0	79.9	52-111	0			
2,4,6-Trichlorophenol	1060	33	1333	0	79.5	46-105	0			
2,4-Dichlorophenol	988	33	1333	0	74.1	47-96	0			
2,4-Dimethylphenol	1083	33	1333	0	81.2	49-97	0			
2,4-Dinitrophenol	903.3	33	1333	0	67.8	10-106	0			
2,4-Dinitrotoluene	1034	33	1333	0	77.6	58-110	0			
2,6-Dinitrotoluene	1034	33	1333	0	77.6	59-108	0			
2-Chloronaphthalene	998	6.7	1333	0	74.9	56-104	0			
2-Chlorophenol	981.3	33	1333	0	73.6	50-104	0			
2-Methylnaphthalene	984.7	6.7	1333	0	73.9	54-96	0			
2-Methylphenol	1047	33	1333	0	78.5	49-105	0			
2-Nitroaniline	1092	33	1333	0	81.9	54-107	0			
2-Nitrophenol	930	33	1333	0	69.8	51-94	0			
3&4-Methylphenol	1021	33	1333	0	76.6	48-105	0			
3,3'-Dichlorobenzidine	855.3	170	1333	0	64.2	39-99	0			
3-Nitroaniline	906.7	33	1333	0	68	17-92	0			
4,6-Dinitro-2-methylphenol	1024	33	1333	0	76.8	32-103	0			
4-Bromophenyl phenyl ether	1121	33	1333	0	84.1	60-106	0			
4-Chloro-3-methylphenol	1044	33	1333	0	78.3	51-101	0			
4-Chloroaniline	1037	67	1333	0	77.8	27-110	0			
4-Chlorophenyl phenyl ether	1029	33	1333	0	77.2	58-106	0			
4-Nitroaniline	675.3	170	1333	0	50.7	21-100	0			
4-Nitrophenol	1136	33	1333	0	85.2	29-120	0			
Acenaphthene	1073	6.7	1333	0	80.5	55-101	0			
Acenaphthylene	1138	6.7	1333	0	85.4	59-106	0			
Anthracene	1117	6.7	1333	0	83.8	67-105	0			
Benzo(a)anthracene	1146	6.7	1333	0	86	68-105	0			
Benzo(a)pyrene	1122	6.7	1333	0	84.2	68-110	0			
Benzo(b)fluoranthene	1162	6.7	1333	0	87.2	65-110	0			
Benzo(g,h,i)perylene	1294	6.7	1333	0	97.1	60-120	0			
Benzo(k)fluoranthene	1122	6.7	1333	0	84.2	66-113	0			
Bis(2-chloroethoxy)methane	972.7	33	1333	0	73	53-96	0			
Bis(2-chloroethyl)ether	990.7	33	1333	0	74.3	47-108	0			
Bis(2-chloroisopropyl)ether	978	33	1333	0	73.4	47-107	0			
Bis(2-ethylhexyl)phthalate	1087	33	1333	0	81.6	59-117	0			
Butyl benzyl phthalate	1056	33	1333	0	79.2	59-106	0			
Carbazole	1122	33	1333	0	84.2	67-108	0			
Chrysene	1139	6.7	1333	0	85.5	68-108	0			

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
Work Order: 20081693
Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: 163196		Instrument ID SVMS10		Method: SW846 8270D				
Dibenzo(a,h)anthracene	1211	6.7	1333	0	90.8	62-119	0	
Dibenzofuran	1074	33	1333	0	80.6	60-104	0	
Diethyl phthalate	1087	33	1333	0	81.6	62-111	0	
Dimethyl phthalate	1067	33	1333	0	80.1	62-106	0	
Di-n-butyl phthalate	1132	33	1333	0	84.9	59-105	0	
Di-n-octyl phthalate	1108	33	1333	0	83.1	51-123	0	
Fluoranthene	1137	6.7	1333	0	85.3	67-106	0	
Fluorene	1069	6.7	1333	0	80.2	59-107	0	
Hexachlorobenzene	1085	33	1333	0	81.4	62-103	0	
Hexachlorobutadiene	953.3	33	1333	0	71.5	51-94	0	
Hexachlorocyclopentadiene	1067	33	1333	0	80.1	25-120	0	
Hexachloroethane	955.3	33	1333	0	71.7	55-93	0	
Indeno(1,2,3-cd)pyrene	1203	6.7	1333	0	90.3	56-120	0	
Isophorone	1025	170	1333	0	76.9	52-99	0	
Naphthalene	956.7	6.7	1333	0	71.8	46-98	0	
Nitrobenzene	997.3	170	1333	0	74.8	53-95	0	
N-Nitrosodi-n-propylamine	1015	33	1333	0	76.2	50-104	0	
N-Nitrosodiphenylamine	1129	33	1333	0	84.7	63-107	0	
Pentachlorophenol	1178	33	1333	0	88.4	34-106	0	
Phenanthrene	1109	6.7	1333	0	83.2	66-101	0	
Phenol	987.3	33	1333	0	74.1	44-109	0	
Pyrene	1133	6.7	1333	0	85	60-119	0	
<i>Surr: 2,4,6-Tribromophenol</i>	2713	0	3333	0	81.4	38-92	0	
<i>Surr: 2-Fluorobiphenyl</i>	2576	0	3333	0	77.3	44-107	0	
<i>Surr: 2-Fluorophenol</i>	2494	0	3333	0	74.8	37-109	0	
<i>Surr: 4-Terphenyl-d14</i>	2956	0	3333	0	88.7	52-123	0	
<i>Surr: Nitrobenzene-d5</i>	2416	0	3333	0	72.5	41-94	0	
<i>Surr: Phenol-d6</i>	2597	0	3333	0	77.9	28-111	0	

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: 163196 Instrument ID SVMS10 Method: SW846 8270D

MS				Sample ID: 20081689-01B MS		Units: µg/Kg		Analysis Date: 8/26/2020 11:08 PM		
Client ID:		Run ID: SVMS10_200826A			SeqNo: 6665557		Prep Date: 8/26/2020		DF: 10	
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,2,4-Trichlorobenzene	874.8	330	1325	0	66	40-93	0			
1,2-Dichlorobenzene	854.9	330	1325	0	64.5	42-94	0			
1,3-Dichlorobenzene	914.5	330	1325	0	69	41-94	0			
1,4-Dichlorobenzene	914.5	330	1325	0	69	42-94	0			
2,4,5-Trichlorophenol	901.3	330	1325	0	68	52-111	0			
2,4,6-Trichlorophenol	894.6	330	1325	0	67.5	46-105	0			
2,4-Dichlorophenol	881.4	330	1325	0	66.5	47-96	0			
2,4-Dimethylphenol	1034	330	1325	0	78	49-97	0			
2,4-Dinitrophenol	ND	330	1325	0	0	10-106	0			S
2,4-Dinitrotoluene	974.2	330	1325	0	73.5	58-110	0			
2,6-Dinitrotoluene	974.2	330	1325	0	73.5	59-108	0			
2-Chloronaphthalene	907.9	66	1325	0	68.5	56-104	0			
2-Chlorophenol	888	330	1325	0	67	50-104	0			
2-Methylnaphthalene	907.9	66	1325	0	68.5	54-96	0			
2-Methylphenol	980.8	330	1325	0	74	49-105	0			
2-Nitroaniline	1014	330	1325	0	76.5	54-107	0			
2-Nitrophenol	907.9	330	1325	0	68.5	51-94	0			
3&4-Methylphenol	914.5	330	1325	0	69	48-105	0			
3,3´-Dichlorobenzidine	642.8	1,700	1325	0	48.5	39-99	0			J
3-Nitroaniline	907.9	330	1325	0	68.5	17-92	0			
4,6-Dinitro-2-methylphenol	285	330	1325	0	21.5	32-103	0			JS
4-Bromophenyl phenyl ether	987.4	330	1325	0	74.5	60-106	0			
4-Chloro-3-methylphenol	1027	330	1325	0	77.5	51-101	0			
4-Chloroaniline	874.8	670	1325	0	66	27-110	0			
4-Chlorophenyl phenyl ether	1007	330	1325	0	76	58-106	0			
4-Nitroaniline	927.8	1,700	1325	0	70	21-100	0			J
4-Nitrophenol	861.5	330	1325	0	65	29-120	0			
Acenaphthene	994	66	1325	0	75	55-101	0			
Acenaphthylene	1040	66	1325	0	78.5	59-106	0			
Anthracene	1001	66	1325	0	75.5	67-105	0			
Benzo(a)anthracene	1120	66	1325	0	84.5	68-105	0			
Benzo(a)pyrene	1093	66	1325	59.72	78	68-110	0			
Benzo(b)fluoranthene	1226	66	1325	146	81.5	65-110	0			
Benzo(g,h,i)perylene	1219	66	1325	0	92	60-120	0			
Benzo(k)fluoranthene	1027	66	1325	39.81	74.5	66-113	0			
Bis(2-chloroethoxy)methane	868.1	330	1325	0	65.5	53-96	0			
Bis(2-chloroethyl)ether	921.1	330	1325	0	69.5	47-108	0			
Bis(2-chloroisopropyl)ether	881.4	330	1325	0	66.5	47-107	0			
Bis(2-ethylhexyl)phthalate	941	330	1325	0	71	59-117	0			
Butyl benzyl phthalate	974.2	330	1325	0	73.5	59-106	0			
Carbazole	1047	330	1325	0	79	67-108	0			
Chrysene	1100	66	1325	0	83	68-108	0			

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
Work Order: 20081693
Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: 163196		Instrument ID SVMS10		Method: SW846 8270D			
Dibenzo(a,h)anthracene	980.8	66	1325	0	74	62-119	0
Dibenzofuran	1040	330	1325	0	78.5	60-104	0
Diethyl phthalate	947.6	330	1325	0	71.5	62-111	0
Dimethyl phthalate	927.8	330	1325	0	70	62-106	0
Di-n-butyl phthalate	987.4	330	1325	0	74.5	59-105	0
Di-n-octyl phthalate	974.2	330	1325	0	73.5	51-123	0
Fluoranthene	1292	66	1325	0	97.5	67-106	0
Fluorene	974.2	66	1325	0	73.5	59-107	0
Hexachlorobenzene	1021	330	1325	0	77	62-103	0
Hexachlorobutadiene	854.9	330	1325	0	64.5	51-94	0
Hexachlorocyclopentadiene	629.6	330	1325	0	47.5	25-120	0
Hexachloroethane	874.8	330	1325	0	66	55-93	0
Indeno(1,2,3-cd)pyrene	1107	66	1325	0	83.5	56-120	0
Isophorone	907.9	1,700	1325	0	68.5	52-99	0 J
Naphthalene	861.5	66	1325	0	65	46-98	0
Nitrobenzene	894.6	1,700	1325	0	67.5	53-95	0 J
N-Nitrosodi-n-propylamine	914.5	330	1325	0	69	50-104	0
N-Nitrosodiphenylamine	1001	330	1325	0	75.5	63-107	0
Pentachlorophenol	596.4	330	1325	0	45	34-106	0
Phenanthrene	1140	66	1325	0	86	66-101	0
Phenol	894.6	330	1325	0	67.5	44-109	0
Pyrene	1093	66	1325	0	82.5	60-119	0
<i>Surr: 2,4,6-Tribromophenol</i>	2247	0	3313	0	67.8	38-92	0
<i>Surr: 2-Fluorobiphenyl</i>	2187	0	3313	0	66	44-107	0
<i>Surr: 2-Fluorophenol</i>	2015	0	3313	0	60.8	37-109	0
<i>Surr: 4-Terphenyl-d14</i>	2492	0	3313	0	75.2	52-123	0
<i>Surr: Nitrobenzene-d5</i>	2174	0	3313	0	65.6	41-94	0
<i>Surr: Phenol-d6</i>	2247	0	3313	0	67.8	28-111	0

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: 163196 Instrument ID SVMS10 Method: SW846 8270D

MSD				Sample ID: 20081689-01B MSD			Units: µg/Kg		Analysis Date: 8/26/2020 11:35 PM	
Client ID:				Run ID: SVMS10_200826A			SeqNo: 6665558		Prep Date: 8/26/2020	
							DF: 10			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,2,4-Trichlorobenzene	870.7	330	1329	0	65.5	40-93	874.8	0.469	30	
1,2-Dichlorobenzene	857.4	330	1329	0	64.5	42-94	854.9	0.292	30	
1,3-Dichlorobenzene	917.2	330	1329	0	69	41-94	914.5	0.292	30	
1,4-Dichlorobenzene	943.8	330	1329	0	71	42-94	914.5	3.15	30	
2,4,5-Trichlorophenol	990.3	330	1329	0	74.5	52-111	901.3	9.41	30	
2,4,6-Trichlorophenol	943.8	330	1329	0	71	46-105	894.6	5.35	30	
2,4-Dichlorophenol	877.3	330	1329	0	66	47-96	881.4	0.463	30	
2,4-Dimethylphenol	996.9	330	1329	0	75	49-97	1034	3.63	30	
2,4-Dinitrophenol	ND	330	1329	0	0	10-106	0	0	30	S
2,4-Dinitrotoluene	1017	330	1329	0	76.5	58-110	974.2	4.29	30	
2,6-Dinitrotoluene	1017	330	1329	0	76.5	59-108	974.2	4.29	30	
2-Chloronaphthalene	943.8	66	1329	0	71	56-104	907.9	3.88	30	
2-Chlorophenol	917.2	330	1329	0	69	50-104	888	3.23	30	
2-Methylnaphthalene	917.2	66	1329	0	69	54-96	907.9	1.02	30	
2-Methylphenol	943.8	330	1329	0	71	49-105	980.8	3.85	30	
2-Nitroaniline	1090	330	1329	0	82	54-107	1014	7.23	30	
2-Nitrophenol	877.3	330	1329	0	66	51-94	907.9	3.43	30	
3&4-Methylphenol	930.5	330	1329	0	70	48-105	914.5	1.73	30	
3,3'-Dichlorobenzidine	511.8	1,700	1329	0	38.5	39-99	642.8	0	30	JS
3-Nitroaniline	990.3	330	1329	0	74.5	17-92	907.9	8.68	30	
4,6-Dinitro-2-methylphenol	272.5	330	1329	0	20.5	32-103	285	0	30	JS
4-Bromophenyl phenyl ether	1017	330	1329	0	76.5	60-106	987.4	2.94	30	
4-Chloro-3-methylphenol	1037	330	1329	0	78	51-101	1027	0.935	30	
4-Chloroaniline	890.6	670	1329	0	67	27-110	874.8	1.8	30	
4-Chlorophenyl phenyl ether	1017	330	1329	0	76.5	58-106	1007	0.948	30	
4-Nitroaniline	1057	1,700	1329	0	79.5	21-100	927.8	0	30	J
4-Nitrophenol	930.5	330	1329	0	70	29-120	861.5	7.7	30	
Acenaphthene	996.9	66	1329	0	75	55-101	994	0.292	30	
Acenaphthylene	1043	66	1329	0	78.5	59-106	1040	0.293	30	
Anthracene	1043	66	1329	0	78.5	67-105	1001	4.19	30	
Benzo(a)anthracene	1137	66	1329	0	85.5	68-105	1120	1.47	30	
Benzo(a)pyrene	1097	66	1329	59.72	78	68-110	1093	0.292	30	
Benzo(b)fluoranthene	1203	66	1329	146	79.5	65-110	1226	1.89	30	
Benzo(g,h,i)perylene	1216	66	1329	0	91.5	60-120	1219	0.253	30	
Benzo(k)fluoranthene	1024	66	1329	39.81	74	66-113	1027	0.355	30	
Bis(2-chloroethoxy)methane	864	330	1329	0	65	53-96	868.1	0.474	30	
Bis(2-chloroethyl)ether	937.1	330	1329	0	70.5	47-108	921.1	1.72	30	
Bis(2-chloroisopropyl)ether	884	330	1329	0	66.5	47-107	881.4	0.292	30	
Bis(2-ethylhexyl)phthalate	1010	330	1329	0	76	59-117	941	7.09	30	
Butyl benzyl phthalate	983.6	330	1329	0	74	59-106	974.2	0.97	30	
Carbazole	1097	330	1329	0	82.5	67-108	1047	4.63	30	
Chrysene	1130	66	1329	0	85	68-108	1100	2.67	30	

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: 163196		Instrument ID SVMS10		Method: SW846 8270D					
Dibenzo(a,h)anthracene	990.3	66	1329	0	74.5	62-119	980.8	0.965	30
Dibenzofuran	1077	330	1329	0	81	60-104	1040	3.43	30
Diethyl phthalate	1017	330	1329	0	76.5	62-111	947.6	7.05	30
Dimethyl phthalate	943.8	330	1329	0	71	62-106	927.8	1.71	30
Di-n-butyl phthalate	1004	330	1329	0	75.5	59-105	987.4	1.63	30
Di-n-octyl phthalate	957.1	330	1329	0	72	51-123	974.2	1.77	30
Fluoranthene	1243	66	1329	0	93.5	67-106	1292	3.9	30
Fluorene	1004	66	1329	0	75.5	59-107	974.2	2.98	30
Hexachlorobenzene	1030	330	1329	0	77.5	62-103	1021	0.939	30
Hexachlorobutadiene	857.4	330	1329	0	64.5	51-94	854.9	0.292	30
Hexachlorocyclopentadiene	538.3	330	1329	0	40.5	25-120	629.6	15.6	30
Hexachloroethane	903.9	330	1329	0	68	55-93	874.8	3.28	30
Indeno(1,2,3-cd)pyrene	1143	66	1329	0	86	56-120	1107	3.24	30
Isophorone	903.9	1,700	1329	0	68	52-99	907.9	0	30 J
Naphthalene	850.7	66	1329	0	64	46-98	861.5	1.26	30
Nitrobenzene	910.5	1,700	1329	0	68.5	53-95	894.6	0	30 J
N-Nitrosodi-n-propylamine	917.2	330	1329	0	69	50-104	914.5	0.292	30
N-Nitrosodiphenylamine	1017	330	1329	0	76.5	63-107	1001	1.61	30
Pentachlorophenol	664.6	330	1329	0	50	34-106	596.4	10.8	30
Phenanthrene	1117	66	1329	0	84	66-101	1140	2.06	30
Phenol	983.6	330	1329	0	74	44-109	894.6	9.48	30
Pyrene	1070	66	1329	0	80.5	60-119	1093	2.16	30
<i>Surr: 2,4,6-Tribromophenol</i>	2286	0	3323	0	68.8	38-92	2247	1.76	40
<i>Surr: 2-Fluorobiphenyl</i>	2193	0	3323	0	66	44-107	2187	0.292	40
<i>Surr: 2-Fluorophenol</i>	2060	0	3323	0	62	37-109	2015	2.25	40
<i>Surr: 4-Terphenyl-d14</i>	2585	0	3323	0	77.8	52-123	2492	3.69	40
<i>Surr: Nitrobenzene-d5</i>	2173	0	3323	0	65.4	41-94	2174	0.0138	40
<i>Surr: Phenol-d6</i>	2233	0	3323	0	67.2	28-111	2247	0.597	40

The following samples were analyzed in this batch:

20081693-01B

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: **163119** Instrument ID **VMS8** Method: **SW8260C**

MBLK				Sample ID: MBLK-163119-163119			Units: µg/Kg-dry		Analysis Date: 8/25/2020 06:18 PM		
Client ID:		Run ID: VMS8_200825B			SeqNo: 6659628		Prep Date: 8/25/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual	
1,1,1,2-Tetrachloroethane	ND	30									
1,1,1-Trichloroethane	ND	30									
1,1,2,2-Tetrachloroethane	ND	30									
1,1,2-Trichloroethane	ND	30									
1,1,2-Trichlorotrifluoroethane	ND	30									
1,1-Dichloroethane	ND	30									
1,1-Dichloroethene	ND	30									
1,2,3-Trichloropropane	ND	30									
1,2,4-Trichlorobenzene	ND	100									
1,2,4-Trimethylbenzene	ND	30									
1,2-Dibromo-3-chloropropane	ND	100									
1,2-Dibromoethane	ND	30									
1,2-Dichlorobenzene	ND	30									
1,2-Dichloroethane	ND	100									
1,2-Dichloropropane	ND	30									
1,3,5-Trimethylbenzene	ND	100									
1,3-Dichlorobenzene	ND	30									
1,4-Dichlorobenzene	ND	30									
2-Butanone	ND	200									
2-Hexanone	ND	30									
2-Methylnaphthalene	ND	100									
4-Methyl-2-pentanone	ND	30									
Acetone	ND	100									
Acrylonitrile	ND	100									
Benzene	ND	30									
Bromodichloromethane	ND	30									
Bromoform	ND	30									
Bromomethane	ND	100									
Carbon disulfide	ND	30									
Carbon tetrachloride	ND	30									
Chlorobenzene	ND	30									
Chloroethane	ND	100									
Chloroform	ND	30									
Chloromethane	ND	100									
cis-1,2-Dichloroethene	ND	30									
cis-1,3-Dichloropropene	ND	30									
Dibromochloromethane	ND	30									
Dibromomethane	ND	30									
Dichlorodifluoromethane	ND	100									
Diethyl ether	ND	30									
Ethylbenzene	ND	30									
Hexachloroethane	ND	100									

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
Work Order: 20081693
Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: 163119		Instrument ID VMS8		Method: SW8260C				
Isopropylbenzene	ND	30						
m,p-Xylene	ND	60						
Methyl tert-butyl ether	ND	30						
Methylene chloride	ND	250						
Naphthalene	ND	100						
n-Propylbenzene	ND	30						
o-Xylene	ND	30						
Styrene	ND	30						
Tetrachloroethene	ND	30						
Toluene	ND	30						
trans-1,2-Dichloroethene	ND	30						
trans-1,3-Dichloropropene	ND	30						
Trichloroethene	ND	30						
Trichlorofluoromethane	ND	30						
Vinyl acetate	ND	250						
Vinyl chloride	ND	30						
Xylenes, Total	ND	90						
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>1018</i>	<i>0</i>	<i>1000</i>	<i>0</i>	<i>102</i>	<i>70-130</i>	<i>0</i>	
<i>Surr: 4-Bromofluorobenzene</i>	<i>992.5</i>	<i>0</i>	<i>1000</i>	<i>0</i>	<i>99.2</i>	<i>70-130</i>	<i>0</i>	
<i>Surr: Dibromofluoromethane</i>	<i>975</i>	<i>0</i>	<i>1000</i>	<i>0</i>	<i>97.5</i>	<i>70-130</i>	<i>0</i>	
<i>Surr: Toluene-d8</i>	<i>977</i>	<i>0</i>	<i>1000</i>	<i>0</i>	<i>97.7</i>	<i>70-130</i>	<i>0</i>	

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: **163119** Instrument ID **VMS8** Method: **SW8260C**

LCS				Sample ID: LCS-163119-163119			Units: µg/Kg-dry		Analysis Date: 8/25/2020 05:28 PM	
Client ID:				Run ID: VMS8_200825B			SeqNo: 6659626		Prep Date: 8/25/2020	
									DF: 1	
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	853	30	1000	0	85.3	75-125	0			
1,1,1-Trichloroethane	895.5	30	1000	0	89.6	70-135	0			
1,1,2,2-Tetrachloroethane	872.5	30	1000	0	87.2	55-130	0			
1,1,2-Trichloroethane	861.5	30	1000	0	86.2	60-125	0			
1,1-Dichloroethane	852.5	30	1000	0	85.2	75-125	0			
1,1-Dichloroethene	984	30	1000	0	98.4	76-148	0			
1,2,3-Trichloropropane	860	30	1000	0	86	65-130	0			
1,2,4-Trichlorobenzene	979.5	100	1000	0	98	65-130	0			
1,2,4-Trimethylbenzene	815	30	1000	0	81.5	65-135	0			
1,2-Dibromo-3-chloropropane	791	100	1000	0	79.1	40-135	0			
1,2-Dibromoethane	856.5	30	1000	0	85.6	80-195	0			
1,2-Dichlorobenzene	857	30	1000	0	85.7	75-120	0			
1,2-Dichloroethane	874	100	1000	0	87.4	70-135	0			
1,2-Dichloropropane	870	30	1000	0	87	70-120	0			
1,3,5-Trimethylbenzene	834	100	1000	0	83.4	65-135	0			
1,3-Dichlorobenzene	875.5	30	1000	0	87.6	70-125	0			
1,4-Dichlorobenzene	875.5	30	1000	0	87.6	70-125	0			
2-Butanone	918.5	200	1000	0	91.8	30-160	0			
2-Hexanone	1038	30	1000	0	104	45-145	0			
4-Methyl-2-pentanone	1522	30	1000	0	152	74-176	0			
Acetone	965.5	100	1000	0	96.6	20-160	0			
Acrylonitrile	965.5	100	1000	0	96.6	70-135	0			
Benzene	957	30	1000	0	95.7	75-125	0			
Bromodichloromethane	905	30	1000	0	90.5	70-130	0			
Bromoform	839.5	30	1000	0	84	55-135	0			
Bromomethane	1594	100	1000	0	159	50-170	0			
Carbon disulfide	1006	30	1000	0	101	45-160	0			
Carbon tetrachloride	800	30	1000	0	80	65-135	0			
Chlorobenzene	859	30	1000	0	85.9	75-125	0			
Chloroethane	808	100	1000	0	80.8	40-155	0			
Chloroform	844.5	30	1000	0	84.4	66-140	0			
Chloromethane	577.5	100	1000	0	57.8	50-144	0			
cis-1,2-Dichloroethene	846.5	30	1000	0	84.6	65-125	0			
cis-1,3-Dichloropropene	921	30	1000	0	92.1	70-125	0			
Dibromochloromethane	796	30	1000	0	79.6	65-135	0			
Dibromomethane	883.5	30	1000	0	88.4	75-130	0			
Dichlorodifluoromethane	912	100	1000	0	91.2	35-135	0			
Diethyl ether	865.5	30	1000	0	86.6	67-150	0			
Ethylbenzene	878	30	1000	0	87.8	75-125	0			
Hexachloroethane	772.5	100	1000	0	77.2	51-122	0			
Isopropylbenzene	883	30	1000	0	88.3	75-130	0			
m,p-Xylene	1946	60	2000	0	97.3	80-125	0			

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
Work Order: 20081693
Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: 163119		Instrument ID VMS8		Method: SW8260C			
Methyl tert-butyl ether	980.5	30	1000	0	98	75-125	0
Methylene chloride	747.5	250	1000	0	74.8	55-145	0
Naphthalene	853	100	1000	0	85.3	40-140	0
n-Propylbenzene	858	30	1000	0	85.8	65-135	0
o-Xylene	862	30	1000	0	86.2	75-125	0
Styrene	869.5	30	1000	0	87	80-138	0
Tetrachloroethene	1004	30	1000	0	100	67-167	0
Toluene	844.5	30	1000	0	84.4	70-125	0
trans-1,2-Dichloroethene	872.5	30	1000	0	87.2	65-135	0
trans-1,3-Dichloropropene	824.5	30	1000	0	82.4	59-129	0
Trichloroethene	888	30	1000	0	88.8	75-125	0
Trichlorofluoromethane	760	30	1000	0	76	25-185	0
Vinyl chloride	900.5	30	1000	0	90	60-125	0
Xylenes, Total	2808	90	3000	0	93.6	75-125	0
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>1006</i>	<i>0</i>	<i>1000</i>	<i>0</i>	<i>101</i>	<i>70-130</i>	<i>0</i>
<i>Surr: 4-Bromofluorobenzene</i>	<i>991.5</i>	<i>0</i>	<i>1000</i>	<i>0</i>	<i>99.2</i>	<i>70-130</i>	<i>0</i>
<i>Surr: Dibromofluoromethane</i>	<i>998.5</i>	<i>0</i>	<i>1000</i>	<i>0</i>	<i>99.8</i>	<i>70-130</i>	<i>0</i>
<i>Surr: Toluene-d8</i>	<i>1010</i>	<i>0</i>	<i>1000</i>	<i>0</i>	<i>101</i>	<i>70-130</i>	<i>0</i>

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: 163119 Instrument ID VMS8 Method: SW8260C

MS				Sample ID: 20081676-01A MS			Units: µg/Kg-dry		Analysis Date: 8/26/2020 12:02 AM	
Client ID:				Run ID: VMS8_200825B			SeqNo: 6659653		Prep Date: 8/25/2020	
							DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	1001	36	1200	0	83.5	75-125	0			
1,1,1-Trichloroethane	1082	36	1200	0	90.2	70-135	0			
1,1,2,2-Tetrachloroethane	826.2	36	1200	0	68.9	55-130	0			
1,1,2-Trichloroethane	1033	36	1200	0	86.1	60-125	0			
1,1-Dichloroethane	929.4	36	1200	0	77.5	75-125	0			
1,1-Dichloroethene	1212	36	1200	0	101	76-148	0			
1,2,3-Trichloropropane	1059	36	1200	0	88.3	65-130	0			
1,2,4-Trichlorobenzene	1093	120	1200	0	91.1	65-130	0			
1,2,4-Trimethylbenzene	990	36	1200	0	82.5	65-135	0			
1,2-Dibromo-3-chloropropane	795	120	1200	0	66.3	40-135	0			
1,2-Dibromoethane	1012	36	1200	0	84.4	80-195	0			
1,2-Dichlorobenzene	1003	36	1200	0	83.6	75-120	0			
1,2-Dichloroethane	1016	120	1200	0	84.7	70-135	0			
1,2-Dichloropropane	1059	36	1200	0	88.3	70-120	0			
1,3,5-Trimethylbenzene	1023	120	1200	0	85.3	65-135	0			
1,3-Dichlorobenzene	1020	36	1200	0	85	70-125	0			
1,4-Dichlorobenzene	1020	36	1200	0	85	70-125	0			
2-Butanone	1064	240	1200	0	88.7	30-160	0			
2-Hexanone	1430	36	1200	0	119	45-145	0			
4-Methyl-2-pentanone	1576	36	1200	0	131	74-176	0			
Acetone	1692	120	1200	0	141	20-160	0			
Acrylonitrile	985.2	120	1200	0	82.1	70-135	0			
Benzene	1170	36	1200	0	97.5	75-125	0			
Bromodichloromethane	999.6	36	1200	0	83.3	70-130	0			
Bromoform	953.4	36	1200	0	79.5	55-135	0			
Bromomethane	1438	120	1200	0	120	50-170	0			
Carbon disulfide	1111	36	1200	0	92.6	45-160	0			
Carbon tetrachloride	959.4	36	1200	0	80	65-135	0			
Chlorobenzene	1009	36	1200	0	84.1	75-125	0			
Chloroethane	545.4	120	1200	0	45.5	40-155	0			
Chloroform	1019	36	1200	0	84.9	66-140	0			
Chloromethane	695.4	120	1200	0	58	50-144	0			
cis-1,2-Dichloroethene	1009	36	1200	0	84.1	65-125	0			
cis-1,3-Dichloropropene	1065	36	1200	0	88.8	70-125	0			
Dibromochloromethane	873	36	1200	0	72.8	65-135	0			
Dibromomethane	1024	36	1200	0	85.3	75-130	0			
Dichlorodifluoromethane	1043	120	1200	0	86.9	35-135	0			
Diethyl ether	951.6	36	1200	0	79.3	67-150	0			
Ethylbenzene	1048	36	1200	0	87.3	75-125	0			
Hexachloroethane	940.2	120	1200	0	78.4	51-122	0			
Isopropylbenzene	1066	36	1200	0	88.9	75-130	0			
m,p-Xylene	2306	72	2400	0	96.1	80-125	0			

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
Work Order: 20081693
Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: 163119		Instrument ID VMS8		Method: SW8260C			
Methyl tert-butyl ether	1144	36	1200	0	95.4	75-125	0
Methylene chloride	891.6	300	1200	0	74.3	55-145	0
Naphthalene	943.2	120	1200	0	78.6	40-140	0
n-Propylbenzene	1016	36	1200	0	84.7	65-135	0
o-Xylene	1043	36	1200	0	86.9	75-125	0
Styrene	1042	36	1200	0	86.8	80-138	0
Tetrachloroethene	1994	36	1200	0	166	67-167	0
Toluene	1013	36	1200	0	84.4	70-125	0
trans-1,2-Dichloroethene	1062	36	1200	0	88.5	65-135	0
trans-1,3-Dichloropropene	927	36	1200	0	77.3	59-129	0
Trichloroethene	1240	36	1200	0	103	75-125	0
Trichlorofluoromethane	774.6	36	1200	0	64.6	25-185	0
Vinyl chloride	1139	36	1200	0	94.9	60-125	0
Xylenes, Total	3349	110	3600	0	93	75-125	0
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>1212</i>	<i>0</i>	<i>1200</i>	<i>0</i>	<i>101</i>	<i>70-130</i>	<i>0</i>
<i>Surr: 4-Bromofluorobenzene</i>	<i>1215</i>	<i>0</i>	<i>1200</i>	<i>0</i>	<i>101</i>	<i>70-130</i>	<i>0</i>
<i>Surr: Dibromofluoromethane</i>	<i>1201</i>	<i>0</i>	<i>1200</i>	<i>0</i>	<i>100</i>	<i>70-130</i>	<i>0</i>
<i>Surr: Toluene-d8</i>	<i>1196</i>	<i>0</i>	<i>1200</i>	<i>0</i>	<i>99.7</i>	<i>70-130</i>	<i>0</i>

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: 163119 Instrument ID VMS8 Method: SW8260C

MSD				Sample ID: 20081676-01A MSD			Units: µg/Kg-dry		Analysis Date: 8/26/2020 12:19 AM	
Client ID:				Run ID: VMS8_200825B			SeqNo: 6659654		Prep Date: 8/25/2020	
									DF: 1	
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	946.5	36	1201	0	78.8	75-125	1001	5.64	30	
1,1,1-Trichloroethane	1055	36	1201	0	87.8	70-135	1082	2.6	30	
1,1,2,2-Tetrachloroethane	789.2	36	1201	0	65.7	55-130	826.2	4.59	30	
1,1,2-Trichloroethane	1002	36	1201	0	83.5	60-125	1033	3.03	30	
1,1-Dichloroethane	904.5	36	1201	0	75.3	75-125	929.4	2.72	30	
1,1-Dichloroethene	1104	36	1201	0	91.9	76-148	1212	9.29	30	
1,2,3-Trichloropropane	1018	36	1201	0	84.7	65-130	1059	3.95	30	
1,2,4-Trichlorobenzene	1050	120	1201	0	87.4	65-130	1093	4	30	
1,2,4-Trimethylbenzene	944.7	36	1201	0	78.6	65-135	990	4.68	30	
1,2-Dibromo-3-chloropropane	798.8	120	1201	0	66.5	40-135	795	0.47	30	
1,2-Dibromoethane	971.7	36	1201	0	80.9	80-195	1012	4.08	30	
1,2-Dichlorobenzene	970.5	36	1201	0	80.8	75-120	1003	3.25	30	
1,2-Dichloroethane	980.1	120	1201	0	81.6	70-135	1016	3.63	30	
1,2-Dichloropropane	1019	36	1201	0	84.8	70-120	1059	3.84	30	
1,3,5-Trimethylbenzene	956.7	120	1201	0	79.6	65-135	1023	6.7	30	
1,3-Dichlorobenzene	956.7	36	1201	0	79.6	70-125	1020	6.41	30	
1,4-Dichlorobenzene	956.7	36	1201	0	79.6	70-125	1020	6.41	30	
2-Butanone	1069	240	1201	0	89	30-160	1064	0.488	30	
2-Hexanone	1395	36	1201	0	116	45-145	1430	2.5	30	
4-Methyl-2-pentanone	1504	36	1201	0	125	74-176	1576	4.66	30	
Acetone	1671	120	1201	0	139	20-160	1692	1.26	30	
Acrylonitrile	997	120	1201	0	83	70-135	985.2	1.18	30	
Benzene	1119	36	1201	0	93.1	75-125	1170	4.47	30	
Bromodichloromethane	991	36	1201	0	82.5	70-130	999.6	0.871	30	
Bromoform	924.3	36	1201	0	76.9	55-135	953.4	3.1	30	
Bromomethane	1360	120	1201	0	113	50-170	1438	5.57	30	
Carbon disulfide	1052	36	1201	0	87.5	45-160	1111	5.51	30	
Carbon tetrachloride	915.9	36	1201	0	76.2	65-135	959.4	4.64	30	
Chlorobenzene	966.9	36	1201	0	80.5	75-125	1009	4.22	30	
Chloroethane	519.5	120	1201	0	43.2	40-155	545.4	4.87	30	
Chloroform	967.5	36	1201	0	80.5	66-140	1019	5.17	30	
Chloromethane	654	120	1201	0	54.4	50-144	695.4	6.13	30	
cis-1,2-Dichloroethene	954.9	36	1201	0	79.5	65-125	1009	5.47	30	
cis-1,3-Dichloropropene	994	36	1201	0	82.7	70-125	1065	6.9	30	
Dibromochloromethane	845.6	36	1201	0	70.4	65-135	873	3.19	30	
Dibromomethane	984.9	36	1201	0	82	75-130	1024	3.85	30	
Dichlorodifluoromethane	951.3	120	1201	0	79.2	35-135	1043	9.18	30	
Diethyl ether	920.1	36	1201	0	76.6	67-150	951.6	3.37	30	
Ethylbenzene	977.1	36	1201	0	81.3	75-125	1048	6.96	30	
Hexachloroethane	882.2	120	1201	0	73.4	51-122	940.2	6.36	30	
Isopropylbenzene	1011	36	1201	0	84.2	75-130	1066	5.28	30	
m,p-Xylene	2182	72	2402	0	90.8	80-125	2306	5.52	30	

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
Work Order: 20081693
Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: 163119		Instrument ID VMS8		Method: SW8260C					
Methyl tert-butyl ether	1130	36	1201	0	94.1	75-125	1144	1.23	30
Methylene chloride	839	300	1201	0	69.8	55-145	891.6	6.08	30
Naphthalene	930.9	120	1201	0	77.5	40-140	943.2	1.32	30
n-Propylbenzene	1000	36	1201	0	83.2	65-135	1016	1.63	30
o-Xylene	977.1	36	1201	0	81.3	75-125	1043	6.5	30
Styrene	986.8	36	1201	0	82.1	80-138	1042	5.41	30
Tetrachloroethene	1855	36	1201	0	154	67-167	1994	7.2	30
Toluene	948.3	36	1201	0	78.9	70-125	1013	6.58	30
trans-1,2-Dichloroethene	1040	36	1201	0	86.6	65-135	1062	2.08	30
trans-1,3-Dichloropropene	886.5	36	1201	0	73.8	59-129	927	4.47	30
Trichloroethene	1213	36	1201	0	101	75-125	1240	2.26	30
Trichlorofluoromethane	741.1	36	1201	0	61.7	25-185	774.6	4.42	30
Vinyl chloride	1043	36	1201	0	86.8	60-125	1139	8.82	30
Xylenes, Total	3159	110	3603	0	87.7	75-125	3349	5.83	30
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>1190</i>	<i>0</i>	<i>1201</i>	<i>0</i>	<i>99.1</i>	<i>70-130</i>	<i>1212</i>	<i>1.86</i>	<i>30</i>
<i>Surr: 4-Bromofluorobenzene</i>	<i>1210</i>	<i>0</i>	<i>1201</i>	<i>0</i>	<i>101</i>	<i>70-130</i>	<i>1215</i>	<i>0.45</i>	<i>30</i>
<i>Surr: Dibromofluoromethane</i>	<i>1202</i>	<i>0</i>	<i>1201</i>	<i>0</i>	<i>100</i>	<i>70-130</i>	<i>1201</i>	<i>0.0441</i>	<i>30</i>
<i>Surr: Toluene-d8</i>	<i>1194</i>	<i>0</i>	<i>1201</i>	<i>0</i>	<i>99.4</i>	<i>70-130</i>	<i>1196</i>	<i>0.157</i>	<i>30</i>

The following samples were analyzed in this batch:

20081693-01A

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: **163169** Instrument ID **GALLERY** Method: **A4500-CI E-11**

MBLK		Sample ID: MBLK-163169-163169				Units: mg/Kg		Analysis Date: 8/27/2020 01:11 PM		
Client ID:		Run ID: GALLERY_200827A		SeqNo: 6664692		Prep Date: 8/25/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual

Chloride ND 10

MS		Sample ID: 20081691-01BMS				Units: mg/Kg		Analysis Date: 8/27/2020 01:11 PM		
Client ID:		Run ID: GALLERY_200827A		SeqNo: 6664697		Prep Date: 8/25/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual

Chloride 490.4 9.6 480.8 5.82 101 86-114 0

MSD		Sample ID: 20081691-01BMSD				Units: mg/Kg		Analysis Date: 8/27/2020 01:11 PM		
Client ID:		Run ID: GALLERY_200827A		SeqNo: 6664698		Prep Date: 8/25/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual

Chloride 473.7 9.6 480.8 5.82 97.3 86-114 490.4 3.47 10

LCS1		Sample ID: LCS1-163169-163169				Units: mg/Kg		Analysis Date: 8/27/2020 01:11 PM		
Client ID:		Run ID: GALLERY_200827A		SeqNo: 6664707		Prep Date: 8/25/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual

Chloride 96.67 10 100 0 96.7 86-114 0

LCS2		Sample ID: LCS2-163169-163169				Units: mg/Kg		Analysis Date: 8/27/2020 01:11 PM		
Client ID:		Run ID: GALLERY_200827A		SeqNo: 6664708		Prep Date: 8/25/2020		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual

Chloride 498.8 10 500 0 99.8 88-112 0

The following samples were analyzed in this batch:

20081693-01B

Note: See Qualifiers Page for a list of Qualifiers and their explanation.

Client: AKT Peerless
 Work Order: 20081693
 Project: 8997f5-5-10.4

QC BATCH REPORT

Batch ID: **R296833** Instrument ID **MOIST** Method: **SW3550C**

MBLK		Sample ID: WBLKS-R296833				Units: % of sample		Analysis Date: 8/26/2020 12:00 PM		
Client ID:		Run ID: MOIST_200826A		SeqNo: 6663457		Prep Date:		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Moisture	ND	0.10								

LCS		Sample ID: LCS-R296833				Units: % of sample		Analysis Date: 8/26/2020 12:00 PM		
Client ID:		Run ID: MOIST_200826A		SeqNo: 6663456		Prep Date:		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Moisture	100	0.10	100	0	100	98-102	0			

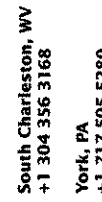
DUP		Sample ID: 20081696-01B DUP				Units: % of sample		Analysis Date: 8/26/2020 12:00 PM		
Client ID:		Run ID: MOIST_200826A		SeqNo: 6663453		Prep Date:		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Moisture	8.79	0.10	0	0	0	0-0	8.97	2.03	10	

DUP		Sample ID: 20081699-01B DUP				Units: % of sample		Analysis Date: 8/26/2020 12:00 PM		
Client ID:		Run ID: MOIST_200826A		SeqNo: 6663455		Prep Date:		DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Moisture	10.02	0.10	0	0	0	0-0	9.87	1.51	10	

The following samples were analyzed in this batch:

20081693-01B

Note: See Qualifiers Page for a list of Qualifiers and their explanation.



South Charleston, WV
+1 304 356 3168

COC ID: 223383

ALS Project Manager:	ALS Work Order #:	20081693
Parameter/Method Request for Analysis		

Customer Information		Project Information		Parameter/Method Request for Analysis	
Purchase Order		Project Name		A	VOCs
Work Order		Project Number	8997f5-5-10.4	B	SVOCs
Company Name	AKT Peerless	Bill To Company	AKT Peerless	C	PCBs
Send Report To	Megan Napier 22725 Orchard Lake Road	Invoice Attn	Accounts Payable	D	MI-10 metals
Address		Address	214 James Avenue	E	Chloride
City/State/Zip	Farmington, MI 48336	City/State/Zip	Saginaw, MI 48607	F	Herbicides + Pesticides
Phone	(248) 615-1333	Phone	(989) 754-9836	G	
Fax		Fax		H	
e-Mail Address	cooktr@aktpeerless.com napierm@aktpeerless.com	e-Mail Address		I	
				J	

[illegible]

Sampler(s) Please Print & Sign			Shipment Method		Required Turnaround Time: (Check Box)		Results Due Date:	
Sean Brick <i>Brick</i>					<input type="checkbox"/> Std 10 WK Days <input checked="" type="checkbox"/> 5 WK Days <input type="checkbox"/> 2 WK Days <input type="checkbox"/> 24 Hour			
Released by: <i>[Signature]</i>			Date: 8/20/20	Time: 11:10am	Received by: <i>[Signature]</i>		Notes: also send report to bricks@akpcores.com	
Relinquished by: <i>[Signature]</i>			Date: 8/20/20	Time: 5:00	Received by (Laboratory): <i>[Signature]</i>		Cooler ID: 82p/20 53a	
Logged by (Laboratory):			Date: 8/21/20	Time: 0935	Checked by (Laboratory): <i>[Signature]</i>		Cooler Temp: 1.9°C	
Preservative Key: 1-HCl 2-HNO ₃ 3-H ₂ SO ₄ 4-NaOH 5-Na ₂ S ₂ O ₃ 6-NaHSO ₄					7-Other		8-4°C 9-5035	
							QC Package: (Check One Box Below) <input checked="" type="checkbox"/> Level II Std Q/C <input type="checkbox"/> TRRP Checklist <input type="checkbox"/> Level III Std Q/C/Raw Data <input type="checkbox"/> TRRP Level IV <input type="checkbox"/> Level IV SW245/CLP <input type="checkbox"/> Other	

Note: 1. Any changes must be made in writing once samples and COC Form have been submitted to ALS Environmental.
2. Unless otherwise agreed in a formal contract, services provided by ALS Environmental are expressly limited to the terms and conditions stated on the reverse.
3. The Chain of Custody is a legal document. All information must be completed accurately.

[Signature] 8/21/20 0800

Copyright 2011 by ALS Environmental.

Sample Receipt Checklist

Client Name: **AKT PEERLESS - FARMINGTON**

Date/Time Received: **21-Aug-20 08:00**

Work Order: **20081693**

Received by: **KRW**

Checklist completed by Keith Wurenga
eSignature

21-Aug-20
Date

Reviewed by: Bill Carey
eSignature

21-Aug-20
Date

Matrices: Soil

Carrier name: Courier

Shipping container/cooler in good condition?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	Not Present <input type="checkbox"/>
Custody seals intact on shipping container/cooler?	Yes <input type="checkbox"/>	No <input type="checkbox"/>	Not Present <input checked="" type="checkbox"/>
Custody seals intact on sample bottles?	Yes <input type="checkbox"/>	No <input type="checkbox"/>	Not Present <input checked="" type="checkbox"/>
Chain of custody present?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	
Chain of custody signed when relinquished and received?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	
Chain of custody agrees with sample labels?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	
Samples in proper container/bottle?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	
Sample containers intact?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	
Sufficient sample volume for indicated test?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	
All samples received within holding time?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	
Container/Temp Blank temperature in compliance?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	
Sample(s) received on ice?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	
Temperature(s)/Thermometer(s):	<u>1.9/2.9 C</u>		<u>IR3</u>
Cooler(s)/Kit(s):	<u></u>		
Date/Time sample(s) sent to storage:	<u>8/21/2020 9:34:31 AM</u>		
Water - VOA vials have zero headspace?	Yes <input type="checkbox"/>	No <input type="checkbox"/>	No VOA vials submitted <input checked="" type="checkbox"/>
Water - pH acceptable upon receipt?	Yes <input type="checkbox"/>	No <input type="checkbox"/>	N/A <input checked="" type="checkbox"/>
pH adjusted?	Yes <input type="checkbox"/>	No <input type="checkbox"/>	N/A <input checked="" type="checkbox"/>
pH adjusted by:	<u>-</u>		

Login Notes:

Client Contacted:

Date Contacted:

Person Contacted:

Contacted By:

Regarding:

Comments:

CorrectiveAction: