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Characterization of Vadose Zone Sediment: RCRA Borehole 299-E33-338 Located Near the B-BX-BY Waste Management Area

C.W. Lindenmeier R.J. Serne B.N. Bjornstad G.W. Gee H.T. Schaef D.C. Lanigan M.J. Lindberg R.E. Clayton V.L. LeGore I.V. Kutnyakov S.R. Baum K.N. Geiszler C.F. Brown M.M. Valenta T.S. Vickerman L.J. Royack

June 2003

Pacific Northwest National Laboratory Operated by Battelle for the U.S. Department of Energy

> Prepared for CH2M Hill Hanford Group, Inc., and the U.S. Department of Energy under Contract DE-AC06-76RL01830

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Pacific Northwest National Laboratory Richland, Washington 99352

Executive Summary

The overall goals of the of the Tank Farm Vadose Zone Project, led by CH2M HILL Hanford Group, Inc., are: 1) to define risks from past and future single-shell tank farm activities, 2) to identify and evaluate the efficacy of interim measures, and 3) to aid via collection of geotechnical information and data, future decisions that must be made by the U.S. Department of Energy (DOE) regarding the nearterm operations, future waste retrieval, and final closure activities for the single-shell tank waste management areas. For a more complete discussion of the goals of the Tank Farm Vadose Zone Project, see the overall work plan, *Phase 1 RCRA Facility Investigation/Corrective Measures Study Work Plan for the Single-Shell Tank Waste Management Areas* (DOE 1999). Specific details on the rationale for activities performed at the B-BX-BY tank farm waste management area are found in CH2M HILL (2000).

To meet these goals, CH2M HILL Hanford Group, Inc., asked scientists from Pacific Northwest National Laboratory to perform detailed analyses of vadose zone sediment, both uncontaminated and contaminated, from within B-BX-BY Waste Management Area.

Specifically, this report contains all the geologic, geochemical, and selected physical characterization data collected on vadose zone sediment recovered from the *Resource Conservation and Recovery Act* borehole 299-E33-338 that is near B-BX-BY Waste Management Area.

This report is one in a series of three reports to present recent data collected on vadose zone sediment that provides a baseline to compare with information from two contaminated boreholes within B-BX-BY Waste Management Area: 1) borehole 299-E33-45, and 2) borehole 299-E33-46 northeast of tank B-110, which has been decommissioned. This document describes all the characterization data collected and interpretations for borehole 299-E33-338 assembled by the Applied Geology and Geochemistry Group within the Environmental Technology Division of the Pacific Northwest National Laboratory and is incorporated in the B-BX-BY field investigation report.

The geology under the B-BX-BY Waste Management Area forms the framework through which the contaminants move, and as discussed in Serne et. al. 2002, provides the basis with which to interpret and extrapolate the physical and geochemical properties that control the migration and distribution of contaminants. Specifically, the identification of major lithological contacts and the interrelationships between the coarser- and finer-grained sediment facies are essential when combined with the geochemical profile for interpreting contaminant behavior in the subsurface. For this borehole, lithologic sections were constructed using detailed geologic descriptions, core photos, and geophysical logs. In some cases, the results of laboratory analyses (e.g. particle-size distribution, moisture, calcium carbonate content) helped refine the resulting stratigraphic and lithological interpretations.

Our conceptual model of the vadose zone associated with the 299 E33-338 borehole involves five distinct stratigraphic units beginning with the Hanford formation H1 unit from the surface to a depth of approximately 15.7 m (approximately 51.5-ft) described as a sandy gravel to gravelly sand sequence. This is followed by the Hanford formation H2 unit extending to a depth of approximately 57.9 m (109 ft) that is a sand sequence consisting of sand dominated facies, with multiple graded beds of horizontal to tabular cross-bedd sand to slightly gravelly sand. These graded beds are sometimes capped with thin

layers of silty sand to silt. The last unit associated with the cataclysmic flood deposits is the Hanford H3 formation unit that extends to a depth of approximately 64.8 m (212.5 ft). It is a gravelly sand to slightly gravelly sand sequence. Just below the H3 unit is the Plio-Pleistocene silty unit (PPlz) extending to a depth of approximately 67.8 m (222.4 ft). This unit is a silt-dominated sequence consisting of interstratified well sorted silt and fine sand. The last unit characterized from this borehole was the Plio-Pleistocene gravelly unit (PPlg) extending to a depth of approximately 82.6 m (271 ft), however sampling ended at approximately 73.1 m (239.8 ft). This unit is differentiated from the PPlz due to its sandy gravel to gravelly sand sequence consisting predominantly of unconsolidated basalt-rich sand and gravel.

Sediment samples from the various stratigraphic unites were analyzed and characterized in the laboratory for the following parameters:

- Mass Water Content
- Soil Suction
- Particle-Size Distribution
- Calcium Carbonate and Organic Carbon Contents
- Bulk Chemical Composition
- Mineralogy
- Water Leach (1:1 sediment to water extraction)
- Acid Leach (8M nitric acid extraction)

Physical properties, such as particle size distribution and water content varied according to lithology as expected. In general, elevated areas of water content ($\sim >5\%$) were typically associated with regions of fine grain sediments. Most notable are those regions involving lithological facies at which water contents equal or exceed 10%. Three major peaks are noted at 15.7, 52.9, and 67.1 m (51.6, 173.6, and 220.2 ft) bgs with water contents of 12.95, 14.27, and 26.02% respectively. Along with water content, soil suction measurements were made on most of the core liner and grab samples from the borehole using the filter paper method. Three major peaks were noted approximately 14, 64, and 73 m (45, 210, and 240 ft) bgs with suction measurements of approximately 1.3, 1.5, and 2.2 Mpa. The matric potential profile indicates that wetting from meteoric water has not reached the water table.

Inorganic carbon results reported in terms of calcium carbonate were found to be within the range of 0.5 to 2.0 wt %, and are consistent with results reported elsewhere (e.g. Serne et. al. 2002). The method used to measure the organic carbon relies on subtracting the inorganic carbon from the total carbon in the sample; for such low carbon values this method is not very accurate. The low values for organic carbon (0.01 to 0.14 %) are within the ranges generally reported for sediment at the Hanford site.

Bulk sediment samples were characterized for major and trace elements using a lithium metaborate/tetraborate fusion procedure, and then analyzed by inductively coupled plasma–optical emission spectroscopy (ICP-OES) and ICP-MS methods. Overall results showed very little difference in the primary elemental oxide concentrations for any of the sediment samples as a function of depth or lithology.

The water chemistry analysis for samples collected between 5 and 73 m (16 and 240 ft) bgs using the 1:1 soil to water extract method shows no strong trends as a function of depth and there is little, if any, indication of tank waste interaction with vadose zone soils at this location. Primary characteristics include the following:

- The 1:1 sediment-to-water extract pH varied from 6.97 to 7.74 and in general increased with depth with an average value of 7.4 (Figure 4.9 and Table 4.3).
- There were small increases in pH at the contact between the Hanford H2 and H3 units and the top and bottom of the Plio-Pleistocene mud unit.
- Porewater EC (dilution corrected) varied from 0.88 to 4.3 mS/cm with an average of 2.4 mS/cm.
- There were high EC values deep in the Hanford H2 unit at approximately 49 m (160 ft) bgs and in the deepest sample characterized (i.e., in the PPlg).

The shapes of the major cation profiles (sodium, potassium, calcium, magnesium, and strontium) in terms of calculated porewater concentration versus depth are very similar with slight peaks in the deep portion of the H2 unit at approximately 49 m (160 ft) bgs, at the top of the Plio-Pleistocene silty unit, and in the deepest sample characterized in the PPlg unit. All three of these samples had very low water contents and thus the dilution factor was high. The apparent high porewater concentrations likely represent some dissolution of salts from the sediment that are multiplied by a large dilution factor, and thus suggest more saline porewater than surrounding sediments with higher water content.

The shapes of the major anion profiles (fluoride, chloride, nitrate, bicarbonate, phosphate, and sulfate) in terms of calculated porewater concentration versus depth showed no consistent depths where all anions peaked unlike the major cation profiles. The wetter samples do consistently show low calculated porewater anion concentrations suggesting that the dilution factor is again controlling the apparent concentrations. That is, all the sediments likely dissolve some salts that are not truly in the porewater, so that the dilution correction makes it appear that the porewater anion concentrations are higher in the drier sediments.

This report is divided into sections that describe the geology, geochemical characterization methods employed, geochemical results, as well as summary and conclusions, references cited, and three appendices with additional details.

Acknowledgements

This work was conducted as part of the Tank Farm Vadose Zone Project led by CH2M Hill Hanford Group, Inc., in support of the U.S. Department of Energy's Office of River Protection. The authors wish to thank Anthony J. Knepp, Fredrick M. Mann, David A. Myers, Thomas E. Jones, and Harold A. Sydnor with CH2M Hill Hanford Group, Inc., Marc I. Wood and Raziuddin Khaleel with Fluor Hanford for their support of this work, and Kevin A. Lindey of Kennedy Jenkes Consultants, Inc. for his insights on the geological nature of the material penetrated by this borehole. We would also like to express our gratitude to Robert Yasek with U.S. Department of Energy's Office of River Protection.

Finally, we would like to thank William J. Deutsch and Bruce J. Bjornstad for their technical review of this document, and Shanna K. Muns for her editorial and document production support.

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Acronyms and Abbreviations

ASTMAmerican Society of Testing and Materialsbgsbelow ground surfaceDOEU.S. Department of EnergyECelectrical conductivityEPAU.S. Environmental Protection AgencyESLPNNL Environmental Sciences LaboratoryFIRfield investigation reportGEAgamma energy analysisHC1hydrochloric acidHf/PPuHanford formation/Plio-Pleistocene unit (?)Hf/PP/RHanford formation/Plio-Pleistocene unit/Ringold formation(?)ICion chromatographyICPinductively coupled plasmaICP-OESinductively coupled plasma –mass spectrometerICP-OESinductively coupled plasma –optical emission spectroscopyPNNLPacific Northwest National LaboratoryPPlgPlio-Pleistocene unit gravelly sand or sandy gravel faciesPPlzPlio-Pleistocene unit mud faciesRCRAResource Conservation and Recovery Act	ASA	American Society of Agronomy
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RCRA Resource Conservation and Recovery Act	PPlz	Plio-Pleistocene unit mud facies
	RCRA	Resource Conservation and Recovery Act
TEM transmission electron microscopy	TEM	transmission electron microscopy
WMA Waste Management Area	WMA	Waste Management Area
XRD x-ray diffraction	XRD	x-ray diffraction

1.0 Introduction

In fiscal year 1999, several offices within the U.S. Department of Energy (DOE) initiated and funded coordinated activities at the Hanford Reservation to study the vadose zone to better understand the fate of contaminants that have leaked from underground storage tanks. As part of this effort, the Pacific Northwest National Laboratory (PNNL) under the direction of CH2M Hill Hanford Group, Inc. (CH2M HILL), received intact sediment cores from the subsurface immediately adjacent to the B-BX-BY Waste Management Area (WMA). These cores were collected during the installation of a new *Resource Conservation and Recovery Act* (RCRA) groundwater monitoring well 299-E33-338. Location maps and more details on the sampling location are presented in Section 2.0.

The clean borehole samples from this effort were collected for analysis of their physical, mineralogical, and chemical properties to serve as a Hanford Site standard for the B-BX-BY WMA. The characterized core samples are available to researchers for experiments relative to environmental problems at the Hanford Site. To obtain sediment, contact Clark Lindenmeier at PNNL by the following venues: telephone (509) 376-8419, fax (509) 376-5368, or email <u>clark.Lindenmeier@pnl.gov.</u>

This report summarizes data measured for samples collected from borehole 299-E33-338 (C3391). Borehole 299-E33-338 was drilled for two purposes: 1) for installation of a RCRA groundwater monitoring well (Horton 2002) and 2) to characterize the in situ soils and background porewater chemistry near the B-BX-BY WMA that have been largely uncontaminated by tank farm and crib and trench discharge operations. This borehole was drilled just outside the southeast fence line of the B tank farm (Figure 2.1). The borehole was drilled between July 23 and August 8, 2001 to a total depth of 80.05 m (275.75 ft) below ground surface (bgs) using the cable-tool method (Horton 2002). The water table was contacted at 77.5 m (254.2 ft) bgs and the top of basalt at 82.6 m (271 ft) bgs. Samples to the top of basalt were collected using drive barrel/splitspoon techniques, before switching to a hard tool to drill 1.5 m (5 ft) into the basalt.

Nearly continuous core was obtained down to a depth of approximately 78.6 m (258 ft) bgs. Two hundred and two 1-ft long by 4-in. diameter cores were retrieved, which accounts for approximately 75% of the total length of the borehole. Each 2-ft splitspoon contained two 1-ft Lexan-lined core segments. The lithology of this borehole was summarized onto a field geologist's log by a Fluor Hanford, Inc. geologist (L. D. Walker). Subsequently, visual inspection of the cores was performed in the laboratory by K. A. Lindsey (Kennedy / Jenks), K. D. Reynolds (Duratek Federal Services), and B. N. Bjornstad (PNNL), who also collected 24 samples for paleomagnetic analysis.

Sub-samples were taken from all 202 cores for moisture content (Table 4-1). In addition, 21 core sub-samples were collected from depths of geological interest for mineralogical and geochemical analysis. Data from these samples allow for comparison of uncontaminated versus contaminated soils to better understand the contributions of tank wastes and other wastewaters on the vadose zone in and around B-BX-BY WMA.

The primary characterization activities included:

- Mass Water Content
- Soil Suction
- Particle-Size Distribution
- Calcium Carbonate and Organic Carbon Contents
- Bulk Chemical Composition
- Mineralogy
- Water Leach (1:1 sediment to water extraction)
- Acid Leach (8M nitric acid extraction)

Support for this work was provided by CH2M Hill Hanford Group, Inc., specifically the Tank Farm Vadose Zone Project. The overall goal of the Tank Farm Vadose Zone Project is to define risks from past and future single-shell tank farm activities, to identify and evaluate the efficacy of interim measures, and to aid future decisions that must be made by DOE regarding the near-term operations, future waste retrieval, and final closure activities for the single-shell tank WMAs. For a more complete discussion of the goals of the Tank Farm Vadose Zone Project, see the overall work plan, Phase 1 Resource Conservation and Recovery Act (RCRA) Facility Investigation/Corrective Measures Study Work Plan for the Single-Shell Tank Waste Management Areas (DOE/RL 1999).

This report is divided into sections that describe the geology, geochemical characterization methods employed, geochemical results, as well as summary and conclusions, references cited, and three appendices with additional details.

2.0 Geology at Borehole 299-E33-338

This section describes the borehole location, drilling, sediment sampling, borehole geophysical logging, and geologic characterization for the uncontaminated borehole 299-E33-338. This borehole was drilled (location in Figure 2.1) as part of an integrated effort for 1) collection of subsurface core samples for detailed vadose zone characterization, and 2) installation of a RCRA groundwater monitoring well in the upper most unconfined aquifer.



Figure 2.1. B-BX-BY Waste Management Area and Location of Background Borehole 299-E33-338

The borehole is located near the southeast corner of the B tank farm. The vadose zone portion of this borehole was drilled using the core-(drive-)barrel cable tool technique wherever possible. The borehole was drilled without the aid of drilling fluid such as water or mud unless noted in the logs, to minimize the introduction of moisture into the sediment cores. After drilling, but prior to well construction, the

borehole was geophysically logged with spectral gamma (i.e., total gamma and potassium, uranium, thorium [KUT], and neutron-neuton [moisture]) probes. Borehole sampling consisted of near continuous split-spoon coring and/or sediment grab sampling throughout the borehole. Sediment cores were collected by driving a 10-cm (4-in.)-diameter 76-cm (2.5 ft)-long split-spoon sampling device ahead of the drilled borehole. The borehole was then cleaned to the bottom of the cored interval prior to the next sampling interval. Field borehole logs indicate that hard tool drilling began at approximately 82.6 m (271 ft) at the top of the basalt and continued to a final depth of approximately 84 m (275.5 ft). Each split-spoon core run contained in two capped 30 cm (1 ft) long, transparent, Lexan liners (core sleeves). Core recovery was generally 100%, however, in gravelly intervals such as in the first 4.6 m (15 ft), recovery was as low as 40%. All cores were sealed and labeled in the field and transported in ice chests to the PNNL environmental sciences laboratory (ESL) in the 3720 Building (300 Area) for refrigerated storage and further sampling and analysis.

In addition to Lexan-lined core samples, sediment grab samples were collected in the field from cuttings recovered during drilling and/or from the 0.5 ft-long split-spoon drive shoe. Several types of samples were labeled and contained from each sample interval. Samples for geologic descriptions were collected in 2.5 cm (1 in.) plastic sample chip trays from surface to total depth. With the exception of the drive shoe grab samples, most of the grab samples are composite samples composed of sediment that was churned up and mixed during the drilling and sampling process.

Lexan-lined cores provide the most representative intact samples of the subsurface and the core depth intervals are believed to be accurate to within 15 cm (0.5 ft) of actual depth. Geophysical logs were used to verify contacts. Fine sediment structure is usually well preserved in the split-spoon cores although layering may be deformed along sides of core due to drag. In the laboratory, the Lexan liners were cut lengthwise with a saw and the core was split into two slabs or halves. Sub-samples for physical and geochemical characterization were collected from the middle (inside) of the core slabs.

A field geologist prepared a geologic description (lithologic log) during drilling and coring of the borehole (Appendix A). The lithologic descriptions were based on visual inspection of material from the split-spoon core shoe, drill cutting, and grab samples. These logs provide a general indication of the lithology encountered. In addition to these field descriptions, a more rigorous and detailed analysis of the vadose zone stratigraphy was performed by geologists in the laboratory, based on cores observed within opened Lexan liners. Appendix B provides the borehole geologic log for 299-E33-338, which was created based on the examination of every third or fourth intact split-spoon core (opened in the laboratory). Table 2.1 provides a summary of the geologist's laboratory assessment of the lithology and stratigraphy for those samples that were selected for detailed geochemical and physical properties characterization. Table 2.2 provides the generalized stratigraphic nomenclature relative to the lithological descriptions used throughout this report.

Two principal stratigraphic units are represented in borehole 299-E33-338, the Hanford formation and the Plio-Pleistocene unit (Table 2.2). The top of basalt was encountered at 82.6 m (271 ft) bgs. The vadose zone is approximately 77.5 m (254 ft) thick and the underlying unconfined aquifer is approximately 5.2 m (17 ft) thick. Zones with elevated moisture occur at several locations within the vadose zone; these occur along sharp lithologic boundaries, often in combination with finer-grained intervals (Figure 2.2).

Borehole geophysical surveys were conducted after the boreholes reached total depth and before well construction. A spectral gamma probe was run from total depth to the surface at a rate of 30 cm (1 ft) per minute (with a 15 cm [6 in.] sampling interval) and a neutron-neutron moisture probe was run from the water table to the surface, at a rate of 30 cm (1 ft) per minute (with a 7.6 cm [3 in.] sampling interval). Geophysical log profiles are attached in Appendix C. Prior to opening the Lexan-lined cores, geophysical profiles were examined for lithostratigraphy, moisture content, and man-made radionuclides. Core samples for chemical and physical analysis were collected from specific lithologic intervals as identified from the geophysical and lithologic log evaluations (lithostratigraphy) confirmed by the geologic examination during core opening.

Spectral gamma logs provide a continuous record of the naturally occurring gamma radioactivity emitted from formations encountered in the borehole. In addition, the spectral gamma logs can be evaluated to determine the presence of specific man-made gamma-emitting radionuclides. The gamma log data is plotted as a continuous curve versus depth, which is referenced to ground surface. This reference can be used to correlate lithologic changes and depth to within 0.5 m (1 or 2 ft). Gamma-log changes are related to the grain size, mineralogical makeup, and moisture content of the lithofacies encountered. Thin lithologic units (i.e., <0.5 m [1.5ft]) cannot be easily identified by the gamma logs because they are averaged into the data bounding them.

The neutron-neutron log is also referenced to ground surface and represents a function of relative moisture content (Figure 2.2, Appendix C). Depth resolution can be determined to within 0.3 m (1 ft). In combination with the geologic log and the total gamma type log, the neutron moisture data can reveal moisture relationships relative to lithologic unit boundaries and grain size (this assumes fine-grained intervals retain higher moisture content than course-grained intervals). For example, the neutron moisture curves may display peaks of elevated moisture, which correspond to fine-grained silt and/or sand lenses, and/or lithologic contacts with contrasting grain-size boundaries. Anomalously high moisture intervals signal areas may indicate zones that are influenced by artificial recharge. These are good targets for follow-up sampling and analysis in intact cores that are preserved. Figure 2.2 shows the downhole geophysical measurements from a high purity germanium spectral gamma log and a neutron-neutron log as related to the lithology and stratigraphy of borehole 299-E33-338.

Core Sample ID	Depth Interval ^(a) (ft)	Top Depth (ft)	Bottom Depth (ft)	Mid- Depth (ft)	Sampled Interval Thickness (ft)	Lithology	Stratigraphic Unit	Comments
C3391- 15.5	15.5-16.5	15.5	16.5	16.0	1.0	Silty sandy gravel	H1	
C3391- 17.5	17.5-18.5	17.5	18.5	18.0	1.0	Gravelly sand	H1	
C3391- 51.05	51.05-52.05	51.1	52.1	51.6	1.0	Silty sandy gravel/silty sand	H1/H2	

 Table 2.1. Sub-sampled Split-Spoon Cores from Borehole 299-E33-338 Analyzed for

 Mineralogy and Geochemistry.

 Table 2.1 (contd)

Core Sample ID	Depth Interval ^(a) (ft)	Top Depth (ft)	Bottom Depth (ft)	Mid- Depth (ft)	Sampled Interval Thickness (ft)	Lithology	Stratigraphic Unit	Comments
C3391- 77.3	77.3-78.3	77.3	78.3	77.8	1.0	Gravelly sand	H2	
C3391- 90.75	90.75-91.75	90.8	91.8	91.3	1.0	Sand	H2	
C3391- 107.3	107.3-108.3	107.3	108.3	107.8	1.0	Sand	H2	
C3391- 115.4	115.4-116.4	115.4	116.4	115.9	1.0	Sand	H2	
C3391- 133	133-134	133.0	134.0	133.5	1.0	Sand	H2	
C3391- 160.3	160.3-161.3	160.3	161.3	160.8	1.0	Sand	H2	
C3391- 161.35	161.35- 162.35	161.4	162.4	161.9	1.0	Silty sand	H2	
C3391- 171.45	171.45- 172.45	171.5	172.5	172.0	1.0	Sand	H2	
C3391- 173.05	173.05- 174.05	173.1	174.1	173.6	1.0	Sand/silt	H2	
C3391- 198.1	198.1-199.1	198.1	199.1	198.6	1.0	Gravelly sand	НЗ	
C3391- 200.1	200.1-201.1	200.1	201.1	200.6	1.0	Silty sand	Н3	Organic matter and tephra present; paleosol?
C3391- 213.8	213.8-214.8	213.8	214.8	214.3	1.0	Sand	PPlz	
C3391- 218.7	218.7-219.7	218.7	219.7	219.2	1.0	Silt	PPlz	
C3391- 219.7	219.7-220.7	219.7	220.7	220.2	1.0	Silt	PPlz	
C3391- 220.65	220.65- 221.65	220.7	221.7	221.2	1.0	Sandy silt	PPlz	
C3391- 221.65	221.65- 222.65	221.7	222.7	222.2	1.0	Sandy silt	PPlz	
C3391- 239.8	239.8-240.8	239.8	240.8	240.3	1.0	Silty sandy gravel	PPlg	

(a) multiply by 0.3048 to convert to meters

Table 2.2. Stratigraphic Terminology Used for the Vadose Zone in the Vicinity of the B, BX, and
BY Tank Farms.

Stratigraphic Symbol Formation		Facies / Subunit	Description	Genesis
НІ		Unit H1	Upper sandy gravel to gravelly sand sequence . Equivalent to the H1 unit discussed by Lindsey et al. (1994, 2001), the upper gravel sequence discussed by Last et al. (1989) and Lindsey et al. (1992), and the Qfg documented by Reidel and Fecht (1994).	
H2	Hanford formation	Unit H2	Sand sequence consisting predominantly of sand- dominated facies, with multiple graded beds of horizontal to tabular cross-bedded sand to slightly gravelly sand. Graded beds sometimes capped with thin layers of silty sand to silt. Equivalent to H2 unit of Wood et al. (2000) and Lindsey et al. (1994, 2001), the sandy sequence of Last et al. (1989) and Lindsey et al. (1992), and to Qfs documented by Reidel and Fecht (1994).	Cataclysmic Flood Deposits
НЗ		Unit H3	Lower gravelly sand to slightly gravelly sand sequence. Equivalent to the H3 unit of Lindsey et al. (1994, 2001) lower gravel sequence discussed by Last et al. (1989) and Lindsey et al. (1992), and the Qfg documented by Reidel and Fecht (1994).	
Hf/PPu and/or PPlz	Hanford formation (?)/	Silt-Dominated Facies	Silty sequence consisting of interstratified well sorted calcareous silt and fine sand. Equivalent to the Silt Facies of the Hanford formation/ Plio Pleistocene Unit(?) of Wood et al. (2000). Perhaps equivalent to the "early Palouse soil" originally described by Tallman et al. (1979) and DOE (1988). Also equivalent to the upper portion of the Hanford/Plio-Pleistocene/Ringold(?) of Lindsey et al. (2001)	Fluvial overbank and/or eolian deposits (with some weakly developed paleosols)
Hf/PPu and/or PPlg	Pleistocene Unit (?)	Sandy Gravel to Gravelly Sand Dominated Facies	Sandy gravel to gravelly sand sequence consisting predominantly of unconsolidated basaltic sands and gravels. Actual origin of this unit is still uncertain. Without intervening silt facies (PPlz subunit), this unit cannot be differentiated from the Hanford formation H3 unit (Wood et al. 2000). Lindsey et al. (2001) suggested this unit may be part of the Ringold Formation.	Plio-Pleistocene age mainstream alluvium (Wood et al. 2000) or possibly Ringold Formation (Lindsey et al. 2001)



Figure 2.2. Borehole 299-E33-338 Lithology, Stratigraphy, Gamma, and Neutron Field Logs as a Function of Depth

2.1 Hanford Formation

Wood et al. (2000) and Lindsey et al. (2001) describe cataclysmic flood deposits of the Hanford formation in the vicinity of the 241-B, BX, and BY tank farms as consisting of three informal units (i.e., H1, H2, and H3). Following is a description of these units within borehole 299-E33-338.

2.1.1 Hanford Formation H1 Unit

Three split-spoon samples were collected from this unit. This unit consists of mostly sandy gravel to silty sandy gravel, with lesser amounts of gravelly sand. A single, thin (0.5 ft) silt layer occurs within this sequence at about 4.6 m (15 ft) bgs. The gravels are multi-lithologic but generally contain a high percentage of basalt (Figure 2.3). The gravel clasts were generally subrounded to well rounded up to 50 mm in diameter where not broken. The finer fraction was described as mostly very coarse to coarse sand with perhaps as much as 5 to 7% silt. The samples generally displayed no cementation or obvious sedimentary structure, and only weak to no reaction to hydrochloric acid (HCl). The Hanford formation H1 unit is 15.7 m (51.5 ft) thick in borehole 299-E33-338 (Figure 2.2).



Figure 2.3. Core from the Hanford Formation H1 Unit in Borehole 299-E33-338. (Shown is loose, poorly sorted, subangular to subrounded, basaltic, silty sandy gravel. Top is to the left.)

2.1.2 Hanford Formation H2 Unit

The Hanford formation H2 unit consists of a sand-dominated sequence of cataclysmic flood deposits. The H2 unit is 37.6 m (123.5 ft) thick extending from a depth of 15.7 m (51.5 ft) to 53.3 m (175 ft). A total of ten, one-foot split-spoon liners were sampled for mineralogical and geochemical characterization

from the H2 unit. The H2 unit is predominantly a poor to well sorted, medium to coarse-grained sand (Figure 2.2). The upper 10 m (30 ft) of the H2 unit is slightly coarser with occasional matrix-supported pebbles floating in a coarse-sand matrix. With depth, the medium to coarse sand becomes more frequently interstratified with layers of fine- to medium-grained sand. The sand has a distinctive "salt and pepper" appearance imparted by the approximately equal concentrations of dark-colored basalt and light-colored quartz and feldspar (Figure 2.4).



Figure 2.4. Typical Hanford Formation H2 Unit in Borehole 299-E33-338. (Shown is one foot of massive to weakly laminated, "salt and pepper"-like, medium to coarse-grained sand. Top is to the left.)

Two thin (<0.5 ft), fine-grained silty layers were observed within the Hanford formation H2 unit. One occurs at the top of the H2 unit at approximately 15.7 m (51.5 ft) and the other lies at approximately 53 m (174 ft) bgs. The lower of these fine-grained units is shown in Figure 2.5. Other fine-grained layers may also be present, but must be limited to the relatively short interval between core runs, which are generally only 0.15 to 0.30 m (0.5 to 1.0 ft) thick. One such interval may occur between 32.2 to 32.4 m (105.6 to 106.2 ft) bgs, as indicated by a narrow spike in the neutron moisture log at this depth (Figure 2.2). Fine-grained units generally retain more moisture, which is often revealed on the neutron moisture log.



Figure 2.5. Lower Fine-Grained Layer in the Hanford Formation H2 Unit. (Lower photo is closeup of upper 1-ft core segment [173.05 to 174.05 ft]. Top is to the left. About 5 cm (2 in.) of well-laminated, moist, cohesive silt is sandwiched between silty fine sand [above] and coarse sand [below].)

2.1.3 Hanford Formation H3 Unit

The Hanford formation H3 unit is 14.2 m (46.5 ft) thick, extending from a depth of 53.3 m (175 ft) to 64.8 m (212.5 ft). Mineralogical and geochemical characterization was performed on two core samples from the Hanford formation H3 unit (Table 2.1). The top of the H3 unit (53.3 m [175 ft] bgs) is chosen based on reappearance of gravelly flood facies. Examples of the poorly to moderately sorted pebbly, medium to coarse-grained sand, which characterize this facies, are shown in Figure 2.6.



Figure 2.6. Hanford Formation H3 Unit in Borehole 299-E33-338. (Pebbly coarse sand from 208.1 to 209.1 ft depth; top is to the left.)

A weak paleosol within this sequence appears to be present at about the 61 m (200 ft) depth, as indicated by a finer grained, calcareous zone containing organic matter and a tephra horizon (Figure 2.7).



Figure 2.7. Weak Paleosol Within the Hanford Formation H3 Unit in Borehole 299-E33-338. (Pebbly coarse sand toward bottom of core [right] grades up into finer-grained calcareous sand with black organic matter, interpreted as a weakly developed paleosol; 200.1 to 201.1 ft depth.)

2.2 Hanford Formation/Plio-Pleistocene (?) Unit

The exact origin of the sedimentary deposits underlying the Hanford formation H3 unit is uncertain and still open to interpretation (Table 2.1). Recent reports have designated deposits beneath the Hanford formation H3 unit as the Hanford formation/Plio-Pleistocene unit(?) (Hf/PPu[?]) (Wood et al. 2000) and Hanford/Plio-Pleistocene/Ringold(?) (H/PP/R[?]) unit (Lindsey et al. 2001). Wood et al. (2000) recognized two facies of the Hf/PPu(?) beneath the 241-B, BX, and BY tank farms, a fine-grained eolian/overbank silt (silt facies), up to 10 m (33 ft) thick, and a sandy gravel to gravelly sand facies. The thick silt-rich interval is believed to be a pre-ice-age flood deposit because silty layers associated with iceage flood deposits of the Hanford formation in this area are generally much thinner (i.e., few centimeters or less) (Wood et al. 2000). The texture, structure, and color of the thick silt layer are all identical to that of the early "Palouse" soil (Tallman et. al. 1979; DOE 1988), more recently referred to as the PPlz or upper Plio-Pleistocene unit, which is widely distributed beneath the 200 West Area (Wood et al. 2000; Serne et al. 2002; DOE 2002).

Where the PPlz unit is absent beneath the B, BX, and BY tank farms, the gravel sequence below the silt unit is indistinguishable from similar-appearing facies of the Hanford formation H3 unit, which overlies the PPlz unit (Wood et al. 2000). In fact, prior to the discovery of the thick silt layer, reported in Wood et al. (2000), gravels overlying basalt bedrock were always included with the Hanford formation (Tallman et al. 1979; Last et al. 1989; Connelly et al. 1992; Lindsey et al. 1992). If the thick silt layer predates the Hanford formation, however, then the underlying gravels must also predate the Hanford formation. Thus, the gravel sequence beneath the silt layer must belong to either a mainstream alluvial facies of the Plio-Pleistocene unit or the Ringold Formation.

2.2.1 Silt-Dominated Facies (PPlz)

The silt facies encountered in well 299-E33-338 is 3 m (9.9 ft) thick, extending from a depth of 64.8 m (212.5 ft) to 67.8 m (222.4 ft) bgs. The silt facies of the Plio-Pleistocene unit is divided into two distinctive beds in borehole 299-E33-338. The upper bed consists of a light brown- to tan-colored, massive, well sorted fine sand (Figure 2.8).



Figure 2.8. Upper Bed in Plio-Pleistocene Silt Facies in Borehole 299-E33-338. (Core segment is one ft long [213.8 to 214.8 ft bgs] and top is to the left.)

One of the four characterization samples (C3391-213.8) was collected from the area shown in Figure 2.8. The three other characterization samples came from an underlying bed of brown-colored, compact, well sorted, well-laminated silt to fine sandy silt (Figure 2.9).



Figure 2.9. Lower Bed in Plio-Pleistocene Silt Facies in Borehole 299-E33-338. (Shown are two 1-ft core segments from the top [upper photo] and bottom [lower photo] of the lower bed of the Plio-Pleistocene silt facies. Contact with the underlying gravel facies is shown in right-hand bottom photo; top of cores is to the left.)

2.2.2 Sandy Gravel to Gravelly Sand Dominated Facies (PPlg)

A sequence of sandy gravel to gravelly sand was encountered at a depth of 67.8 m (222.4 ft). This gravel-rich facies continues to the top of basalt at 82.6 m (271 ft bgs). Only one core sample (C3391-239.8) was characterized from this unit (Table 2.1). These materials were described as muddy sandy gravel to sandy gravel, consisting of an estimated 30 to 80% gravel, 15 to 65% sand, and up to 15% mud (Figure 2.10). The gravel clasts were described as a mixture of mostly quartzite, basalt, and some highly weathered friable granite. Where unbroken, the gravel clasts are subrounded to rounded and range up to at least 60 mm in diameter (intermediate axis). The matrix was described as ranging from mostly very fine sand to poorly sorted coarse to medium sand, with variable mud content. These materials were further described as moderate to uncemented with strong to no reaction to dilute HCI. Some caliche fragments were noted, exhibiting a strong reaction to HCI.



Figure 2.10. Sandy Gravel to Gravelly Sand-Dominated Facies of the Plio-Pleistocene unit (PPlg) in Borehole 299-E33-338. (Top of cores is to the left.)

3.0 Characterization Analytical Methods

This section discusses the methods and procedure used to determine which samples would be characterized and which parameters would be measured.

3.1 Geochemical and Analytical Laboratory Methods and Materials

Geochemical and analytical methods used in the laboratory to characterize the core sediment samples are discussed in this section. Physical properties analyzed include mass water content, particle size distribution, and sand/silt/clay percentages. A variety of geochemical techniques were performed on the sediments including elemental analysis by fusion, 1:1 sediment to water extraction, and 8M nitric acid extraction. Mineralogical analyses were performed using x-ray diffraction (XRD).

3.1.1 Borehole Core Sampling

The samples characterized from RCRA borehole 299-E33-338 were obtained during the geologic description process immediately upon opening the sealed liners in the same fashion as described in the report *Characterization of Vadose Zone Sediments: Uncontaminated RCRA Borehole Core Samples and Composite Samples* (Serne et. al. 2002). As before, the split-spoon samples were obtained in clear, plastic Lexan liners that were 12 in. long and 4 in. in diameter. Plastic end caps were removed, and the liners were cut down both sides with a circular saw. The core was opened in a fashion similar to opening a clam shell, facilitated by the relatively unconsolidated nature of the sediment. The two halves of the liner were laid on a table and quickly photographed and sub-sampled to avoid excessive loss of moisture. Small aliquots were removed in an attempt to construct a representative sample for the entire sleeve. Depending on the sample matrix, very coarse pebble and larger material (i.e., >32 mm) was avoided during sub-sampling. Larger substrate was excluded to provide moisture contents representative of counting and 1:1 sediment-to-water extract samples. Results from sub-sample measurements should then take into consideration a possible bias toward higher concentrations for some analytes that are associated with smaller sized sediment fractions. The sediment in the Plio-Pleistocene silt-dominated facies contained no large pebbles or cobbles.

When distinct contacts were observed in a core sample, the sampling was performed separately on the different lithologies. After sampling and the geologic descriptions were completed, the two halves of the liner were reassembled and retaped to prevent further disturbance or loss of moisture. Liners were then returned to refrigerated storage in the dark at 4° C.

Procedures ASTM D2488-93 (ASTM 1993) and accepted PNNL laboratory procedure for groundwater investigation were followed for visual descriptions and geologic description of all splitspoon samples. The sediment classification scheme used for geologic identification of the sediment types is based on the modified Folk (1968) and/or Wentworth (1922) classification scheme (Figure 3.1). However, the mineralogic and geochemical characterization relied on further separation of the mud into discrete silt and clay sizes. The selected Lexan liners from borehole 299-E33-338 were sub-sampled using stainless steel spatulas. The depths and corresponding stratigraphic unit designations are shown in Table 2.1. In most cases, field moist sediment was used to measure the various parameters discussed in Section 4.0, but the results are on an oven dry-weight basis.





3.1.2 Mass Water Content

All geochemical characterization data in this document are reported on an oven-dry basis from field moist core materials using the traditional moisture content method in *Method of Soil Analysis, Part1, Method 21-2.2 Gravimetry with Oven Drying* (pages 503-507) (American Society of Agronomy [ASA] 1986a). One representative sub-sample of at least 15 to 70 gm was taken from each sleeve and selected grab samples. Sediment samples were placed in tared containers, weighed, and dried in an oven at 105 °C until constant weight was achieved, which took at least 24 hours. The containers then were removed from the oven, sealed, cooled, and weighed. At least two weighings, after 24-hour heatings, were performed to ensure that all moisture was removed. All weighings were performed using a calibrated balance. A calibrated weight set was used to verify balance performance before weighing samples. The gravimetric water content was computed as percentage change in soil weight before and after oven drying.

3.1.3 Particle Size Distribution

Particle-size measurement in terms of sand, silt and clay distribution was determined using the wet sieve/hydrometer method (ASA 1986b, *Part 1 Method 15-5 Hydrometer Method* [pages 404-408]). The silt and clay separates were saved for mineralogical analyses. The samples used for this method were not air dried or oven dried to minimize the effects of particle aggregation that can affect the separation of clay grains from the coarser material. A more qualitative estimate of particle-size distribution was also determined in the field and upon opening core sleeves using a visual-manual technique by the geologist logging the core (ASTM 1993).

3.1.4 Particle Density

The particle density of bulk grains was determined using pychnometers (ASA 1986c, *Part 1; Method 14-3 Pychnometer Method* [pages 378-379] on oven-dried material.

3.1.5 1:1 Sediment-to-Water Extract

The water-soluble inorganic constituents were determined using a 1:1 by weight sediment-todeionized water extract method. This method was chosen because the sediment was too dry to easily extract vadose zone porewater. The extracts were prepared by adding an exact weight of deionized water to approximately 60 to 80 gm of sediment sub-sampled from each sleeve and selected grab samples. The weight of deionized water needed was calculated based on the weight of the field-moist samples and their previously determined moisture contents. The sum of the existing moisture (porewater) and the deionized water was fixed at the mass of the dry sediment. The appropriate amount of deionized water was added to screw cap jars containing the sediment samples. The jars were sealed and briefly shaken by hand, then placed on a mechanical orbital shaker for 1 hour. The samples were allowed to settle until the supernatant liquid was fairly clear. The supernatant was carefully decanted and separated into unfiltered aliquots for conductivity and pH determinations, and filtered aliquots (passed through 0.45 µm membranes) for anion, cation, carbon, and radionuclide analyses. More details can be found in Rhoades (1996) within *Methods of Soils Analysis Part 3* (ASA 1996).

3.1.6 8 M Nitric Acid Extract

Approximately 20 grams of oven-dried sediment was contacted with 8 M nitric acid at a ratio of approximately 5 parts acid to 1 part sediment. The slurries were heated to approximately 80 °C for several hours and then the fluid was separated by centrifugation and filtration through 0.2 µm membranes. The acid extracts were analyzed for major cations and trace metals using inductively coupled plasma (ICP) unit and inductively coupled plasma – mass spectrometer (ICP-MS) techniques, respectively. The acid digestion procedure is based on U.S. Environmental Protection Agency (EPA) SW-846 Method 3050B (EPA 2000a) that can be accessed on-line at

http://www.epa.gov/epaoswer/hazwaste/test/sw846.htm.

3.1.7 pH and Conductivity

Two approximately 3-ml, aliquots of the unfiltered 1:1 by weight sediment-to-water extract supernatant were used for pH and conductivity measurements. The pH values for the extracts were measured with a solid-state pH electrode and a pH meter calibrated with buffers 4, 7, and 10. Conductivity was measured and compared to potassium chloride standards with a range of 0.001 M to 1.0 M.

3.1.8 Anion Analysis

The 1:1 sediment-to-water extracts were analyzed for anions using an ion chromatograph. Fluoride, acetate, formate, chloride, nitrite, bromide, nitrate, carbonate, phosphate, sulfate, and oxalate were separated on a Dionex AS17 column with a gradient elution of 1 mM to 35 mM NaOH and measured

using a conductivity detector. This methodology is based on EPA Method 300.0A (EPA 1984) with the exception of using the gradient elution of sodium hydroxide.

3.1.9 Cations and Trace Metals

Major cation analysis was performed with an ICP unit using high-purity calibration standards to generate calibration curves and verify continuing calibration during the analysis run. Dilutions of 100x, 50x, 10x, and 5x were made of each sample for analysis to investigate and correct for matrix interferences. Details are found in EPA Method 6010B (EPA 2000b). The second instrument used to analyze trace metals, including technetium-99 and uranium-238, was an ICP-MS using accepted PNNL procedures similar to EPA Method 6020 (EPA 2000c).

3.1.10 Alkalinity

The alkalinity content of several of the 1:1 sediment-to-water extracts were measured using standard titration with acid and a carbon analyzer. The alkalinity procedure is equivalent to the U.S. Geological Survey Method Field Manual (USGS 2001) http://water.usgs.gov/owq.

3.1.11 Carbon Content

The carbon contents of borehole sediment samples were determined using ASTM Method D4129-88, Standard Methods for Total and Organic Carbon in Water Oxidation by High Temperature Oxidation and by Coulometric Detection (ASTM 1988). Total carbon in all samples was determined using a Coulometrics, Inc. Model 5051 Carbon Dioxide Coulometer with combustion at approximately 980°C. Ultrapure oxygen was used to sweep the combustion products through a barium chromate catalyst tube for conversion to carbon dioxide. Evolved carbon dioxide was quantified through coulometric titration following absorption in a solution containing ethanolamine. Equipment output reported carbon content values in micrograms per sample. Soil samples for determining total carbon content were placed into pre-combusted, tared, platinum combustion boats and weighed on a four-place analytical balance. After the combustion boats were placed into the furnace introduction tube, a 1-minute waiting period was allowed so that the ultrapure oxygen carrier gas could remove any carbon dioxide introduced to the system from the atmosphere during sample placement. After this system sparge, the sample was moved into the combustion furnace and titration begun. Sample titration readings were performed at 3 minutes after combustion began and again once stability was reached, usually within the next 2 minutes. The system background was determined by performing the entire process using an empty, pre-combusted, platinum boat. Adequate system performance was confirmed by analyzing for known quantities of a calcium carbonate standard.

Inorganic carbon contents for borehole sediment samples were determined using a Coulometrics, Inc., Model 5051 Carbon Dioxide Coulometer. Soil samples were weighed on a four-place analytical balance, then placed into acid-treated glass tubes. Following placement of sample tubes into the system, a 1-minute waiting period allowed the ultrapure oxygen carrier gas to remove any carbon dioxide introduced to the system from the atmosphere. Inorganic carbon was released through acid-assisted evolution (3M hydrochloric acid) with heating to 80 °C. Samples were completely covered by the acid to allow full reaction to occur. Ultrapure oxygen gas swept the resultant carbon dioxide through the equipment to determine inorganic carbon content by coulometric titration. Sample titration readings were performed 5 minutes following acid addition and again once stability was reached, usually within 10 minutes. Known quantities of calcium carbonate standards were analyzed to verify that the equipment was operating properly. Background values were determined. Inorganic carbon content was determined through calculations performed using the microgram per-sample output data and sample weights. Organic carbon was calculated by subtracting inorganic carbon from total carbon and using the remainder.

3.1.12 Bulk Elemental Analysis

Samples were mixed with a flux of lithium metaborate and lithium tetraborate and fused in an induction furnace. The molten melt was immediately poured into a solution of 5% nitric acid containing an internal standard, and mixed continuously until completely dissolved (approximately 30 minutes). The samples were run for major oxides and selected trace elements on a combination simultaneous/sequential Thermo Jarrell-Ash ENVRO II ICP and a Perkin Elmer SCIEX ELAN 6000 ICP-MS. Calibration was performed using USGS and Canmet certified reference materials.

3.1.13 Mineralogy

The mineralogies of the bulk sample and silt- and clay-sized fractions of selected sediment samples were determined by XRD techniques. Bulk sediment samples were dispersed by transferring 100 gm of sediment into a 1-liter bottle and mixing with 1 liter of 0.001 M solution of sodium hexametaphosphate (a dispersant). The suspensions were allowed to shake overnight to ensure complete dispersion. The sand fraction was separated from the dispersed sample by wet sieving through a #230 sieve. The silt fractions were separated from the clay fractions by using Stoke's settling law described in Jackson (1969). The lower limit of the fraction was taken at >2 microns. Sand and silt fractions were oven dried at 110°C and prepared for XRD.

Each clay suspension was concentrated to an approximate volume of 10 ml by adding a few drops of 10 N magnesium chloride to the dispersing solution. Concentrations of the clay in the concentrated suspensions were determined by drying known volumes and weighing the dried sediment. The density of the slurry was calculated from the volume pipetted and the final weight of dried sediment. Volumes of slurry equaling 250 mg of clay were transferred into centrifuge tubes and treated to remove carbonates following the procedure described by Jackson (1969). The carbonate free clay was then saturated with either magnesium (Mg²⁺) or potassium (K⁺) cations. Clay samples were prepared using the Drever (1973) method and placed onto an aluminum slide for XRD analysis. Due to the tendency of the clay film to peel and curl, the magnesium (Mg²⁺)-saturated specimens were solvated with a few drops of a 10% solution of ethylene glycol in ethanol and placed into a desiccator containing excess ethylene glycol for a minimum of 24 hours. Potassium-saturated slides were air dried and analyzed, then heated to 575°C and reanalyzed.

All samples were analyzed on a Scintag x-ray diffraction unit equipped with a Pelter thermoelectrically cooled detector and a copper x-ray tube. Slides of preferentially oriented clay were scanned from 2 to 45 degrees 2θ , and randomly oriented powder mounts were scanned from 2 to 75 degrees 2θ . The bulk samples were prepared by crushing approximately 0.5 gm of sample to a fine

powder that was then packed into a small circular holder. After air drying approximately 0.5 gm of the clay slurry, a random mount was prepared and analyzed from 2 to $75^{\circ} 2\theta$.

Semiquantification of mineral phases by XRD was performed according to Brindley and Brown (1980). The relationship of intensity and mass absorption to the weight fraction of an unknown phase is expressed as:

$$I/I_p = \mu_p/\mu$$
 (wf)

where:

- I is the intensity of the unknown phase
- I_p is the intensity of the pure phase
- μ_p is the mass absorption of the pure phase
- μ is the average mass absorption of the unknown mixture
- wf is the weight fraction of the unknown.

Pure mineral phases of illite, smectite, kaolinite, and chlorite were obtained from the Clay Mineral Society's source clays repository (operated from the University of Missouri in Columbia), and analyzed under the same conditions as the sediment samples. Quartz, feldspars, and calcite standards were purchased from the Excalibur Mineral Company (Peekskill, New York), ground, and analyzed on the diffractometer to obtain intensities for pure nonclay phases.

The mass attenuation coefficients of selected samples were measured according to Brindley and Brown (1980). Ground bulk powders and air-dried clays were packed into a 2.39-cm (0.94-in.) thick circular holder with no backing. The holder was placed in front of the detector and positioned to allow the x-ray beam, diffracted from pure quartz, to pass through the sample and into the detector. The scan was analyzed from 26 to 27 degrees 20. The mass attenuation coefficients were measured directly using the following equation:

$$\mu = (1/\rho x) \ln(I_o/I_x)$$

where:

 $\begin{array}{ll} 1/\rho x & \mbox{is the mass per unit area as the sample is prepared} \\ I_o & \mbox{is the intensity of the incident beam} \\ I_x & \mbox{is the intensity of the transmitted beam through sample thickness } x. \end{array}$

In addition to x-ray diffraction, transmission electron microscopy (TEM) characterization of selected samples was conducted on a JEOL 1200X electron microscope equipped with a Links detector system. Samples were prepared for TEM by transferring a small aliquot of a dilute clay slurry onto a formvar carbon coated 3-ml copper support grid. The clay solution contained 0.15% tert-butylamine to reduce the surface tension of water.

Structural formulas were derived from data collected from the TEM analysis. On average, an energy dispersive x-ray spectra was collected from a minimum of five particles from the same mineral phase common to the sample. The x-ray spectra were collected and processed using the Cliff-Lorimer Ratio Thin Section method and then converted to a structural formula [based on half-unit cell $(O_{10}(OH)_2)$ described in Moore and Reynolds (1989) and Newman (1987)].

3.1.14 Water Potential (Suction) Measurements

Suction measurements were made on most of the core liners and grab samples from the borehole using PNNL's filter paper method. This method relies on the use of a sandwich of three filter papers that rapidly equilibrates with the sediment sample. The middle filter paper does not contact sediment that might stick to the paper and bias the mass measurements. At equilibrium, the matric suction in the filter paper is the same as the matric suction of the sediment sample. The dry filter paper sandwiches were placed in the airtight liners or grab sample jars while still filled with the sediment for at least 3 to 12 weeks to allow sufficient time for the matric suction in the sediment to equilibrate with the matric suction in the filter paper. The mass of the wetted middle filter paper that had no direct contact with the sediment was subsequently determined, and the suction of the sediment was determined from a calibration relationship between filter paper water content and matric suction.

The relationships used for converting the water content of filter paper to matric suction for Whatman #42 filter paper have been determined by Deka et al. (1995) and can be expressed as:

$$Sm = 10^{(5.144 - 6.699 \text{ w})}/10 \text{ for } w < 0.5$$

$$Sm = 10^{(2.383 - 1.309 \text{ w})}/10 \text{ for } w > 0.5$$

where:

Sm is the matric suction (m) and

w is the gravimetric water content (g/g).

One hundred eighty-eight samples from borehole 299-E33-338 were analyzed for water content and soil matric suction. The samples covered the entire borehole profile from 9.6 to 253.6 ft (2.93 to 77.3 m) bgs.

4.0 Analytical Results for Sediment Samples

This section discusses the analytical results for the core samples from borehole 299-E33-338.

4.1 Geophysical and Moisture Content Measurements

All of the cores were sub-sampled for gravimetric moisture content (Table 4.1) and the results are shown in Figure 4.1 along with field log data. Three relatively moist zones were found, the shallowest and least moist was at approximately 16 m (52 ft) bgs at the bottom of the Hanford H1 unit. The second moist zone was at the bottom of the Hanford H2 unit at 52 to 53 m (171.5 to 174 ft) bgs. The zone with highest moisture content was at the bottom of the Plio-Pleistocene mud unit, between 66.7 to 67.8 m (218.7 and 222.6 ft) bgs, with water contents reaching values of 21 to 26% by weight. Both the laboratory results and the moisture log data appear to agree very well. Additionally, the moisture data correlates strongly with lithology in that regions of higher moisture tend to be associated with regions of silt dominated sediments or regions of significant contact layers such as between the silt dominated PPlz and the sandy gravel dominated PPlg regions. Table 4.1 identifies the 22 core samples that were selected, some from each of the lithologies, and subjected to water and acid extracts to develop baseline data on porewater pH, electrical conductivity (EC), major cations and anions, and trace metals. The mass of constituents that were water and acid leachable were also determined to allow comparison with similar data for contaminated boreholes. The comparison allows an estimate of the inventory of contaminants in the vadose zone that are attributable to leaked tank fluids.

Lithologic Unit	Sample No.	Mid Depth ^(a) (Vertical ft)	% Moisture	Lithologic Unit	Sample No.	Mid Depth ^(a) (Vertical ft)	% Moisture
H1	C3391-0	0.5	7.02%	H2	C3391-124.35	124.85	2.00%
H1	C3391-1	1.5	9.75%	H2	C3391-125.6	126.1	2.47%
H1	C3391-2	2.5	4.37%	H2	C3391-126.6	127.1	2.45%
H1	C3391-4.75	5.25	4.27%	H2	C3391-129.3	129.8	5.62%
H1	C3391-8.4	8.9	2.95%	H2	C3391-130.3	130.8	3.20%
H1	C3391-9.4	9.9	3.52%	H2	C3391-132	132.5	3.66%
H1	C3391-10.7	11.2	2.37%	H2	C3391-133	133.5	3.01%
H1	C3391-11.7	12.2	2.89%	H2	C3391-134	134.5	3.21%
H1	C3391-12.5	13	2.65%	H2	C3391-135	135.5	3.02%
H1	C3391-13.5	14	4.57%	H2	C3391-136.95	137.45	3.39%
H1	C3391-15.5	16	4.20%	H2	C3391-137.95	138.45	2.59%
H1	C3391-16.5	17	4.51%	H2	C3391-138.8	138.9	3.54%
H1	C3391-17.5	18	10.57%	H2	C3391-139.8	140.3	3.23%
H1	C3391-18.5	19	4.65%	H2	C3391-141.15	141.65	2.64%
H1	C3391-20.5	21	5.03%	H2	C3391-142.15	142.65	2.58%

Table 4.1. Moisture Content of Sediments in Borehole 299-E33-338

Table 4.1. (contd)

Lithologic Unit	Sample No.	Mid Depth ^(a) (Vertical ft)	% Moisture	Lithologic Unit	Sample No.	Mid Depth ^(a) (Vertical ft)	% Moisture
H1	C3391-21.5	22	3.04%	H2	C3391-143.1	143.75	3.01%
H1	C3391-22.5	23	2.76%	H2	C3391-144.1	144.6	2.87%
H1	C3391-23.5	24	2.74%	H2	C3391-145.45	145.95	2.81%
H1	C3391-25	25.5	3.21%	H2	C3391-146.45	146.95	2.23%
H1	C3391-26	26.5	3.16%	H2	C3391-147.8	148.3	3.03%
H1	C3391-27.5	28	2.90%	H2	C3391-148.8	149.3	2.28%
H1	C3391-28.5	29	3.03%	H2	C3391-150.05	150.55	4.34%
H1	C3391-30.25	30.75	3.49%	H2	C3391-151.05	151.55	3.28%
H1	C3391-31.25	31.75	3.14%	H2	C3391-152.15	152.65	3.51%
H1	C3391-32.5	33	3.74%	H2	C3391-153.15	153.65	2.55%
H1	C3391-33.5	34	3.74%	H2	C3391-156.3	156.8	3.32%
H1	C3391-35.5	36	5.18%	H2	C3391-156.8	157.3	3.08%
H1	C3391-36.5	37	2.24%	H2	C3391-157.3	157.8	2.88%
H1	C3391-37.78	38.28	1.96%	H2	C3391-157.8	158.3	2.72%
H1	C3391-38.78	39.28	3.01%	H2	C3391-159.3	159.8	2.58%
H1	C3391-39.6	39.65	2.68%	H2	C3391-160.3	160.8	2.59%
H1	C3391-40.6	41.1	2.56%	H2	C3391-161.35	161.85	3.72%
H1	C3391-43.2	43.7	3.63%	H2	C3391-162.35	162.85	2.35%
H1	C3391-44.2	44.7	4.69%	H2	C3391-164.3	164.8	3.36%
H1	C3391-45.2	45.7	2.71%	H2	C3391-165.3	165.8	2.37%
H1	C3391-46.2	46.7	2.46%	H2	C3391-166.15	166.65	3.28%
H1	C3391-47.2	47.7	2.49%	H2	C3391-167.15	167.65	2.28%
H1	C3391-48.2	48.7	2.88%	H2	C3391-168.35	168.85	2.79%
H1	C3391-50.05	50.55	3.17%	H2	C3391-169.35	169.85	2.81%
H1	C3391-51.05	51.55	12.95%	H2	C3391-169.35 upper	169.85	1.79%
H2	C3391-52.75	52.75	7.91%	H2	C3391-170.45	170.95	3.79%
H2	C3391-53.75	54.25	2.28%	H2	C3391-171.45	171.95	7.33%
H2	C3391-55.7	56.2	4.66%	H2	C3391-173.05	173.55	14.27%
H2	C3391-56.7	57.2	2.89%	H2	C3391-174.05	174.55	2.60%
H2	C3391-57.9	58.4	3.58%	H3z	C3391-176.8	177.3	2.35%
H2	C3391-58.9	59.4	2.87%	H3z	C3391-177.8	178.3	2.13%
H2	C3391-59.3	59.8	5.16%	H3z	C3391-179.9	180.4	3.67%
H2	C3391-60.3	60.8	3.77%	H3z	C3391-180.9	181.4	3.00%
H2	C3391-61.5	62	3.83%	H3z	C3391-182.2	182.7	2.72%
H2	C3391-62.5	63	2.58%	H3z	C3391-183.2	183.7	2.65%
H2	C3391-64.3	64.8	3.21%	H3z	C3391-185.7	186.2	3.13%
Table 4.1. (contd)

Lithologic Unit	Sample No.	Mid Depth ^(a) (Vertical ft)	% Moisture	Lithologic Unit	Sample No.	Mid Depth ^(a) (Vertical ft)	% Moisture
H2	C3391-65.3	65.8	2.89%	H3z	C3391-186.7	187.2	2.19%
H2	C3391-66.75	67.25	5.09%	H3z	C3391-188	188.5	2.86%
H2	C3391-67.75	68.25	2.99%	H3z	C3391-189	189.5	2.44%
H2	C3391-69.3	69.8	3.91%	H3z	C3391-190.4	190.9	3.17%
H2	C3391-70.3	70.8	3.27%	H3z	C3391-191.4	191.9	2.84%
H2	C3391-71.7	72.2	4.32%	H3z	C3391-192.6	193.1	3.65%
H2	C3391-72.7	73.2	3.61%	H3z	C3391-193.6	194.1	2.62%
H2	C3391-73.9	74.4	4.76%	H3z	C3391-197.1	197.6	3.53%
H2	C3391-74.9	75.4	3.02%	H3z	C3391-198.1	198.6	2.79%
H2	C3391-76.3	76.8	4.17%	H3z	C3391-200.1	200.6	6.03%
H2	C3391-77.3	77.8	3.69%	H3z	C3391-201.1	201.6	6.20%
H2	C3391-78.1	78.6	3.64%	H3z	C3391-202.6	203.1	3.63%
H2	C3391-79.1	79.6	4.13%	H3z	C3391-203.6	204.1	3.10%
H2	C3391-80.25	80.75	2.72%	H3z	C3391-205.3	205.8	2.88%
H2	C3391-81.25	81.75	3.06%	H3z	C3391-206.3	206.8	2.79%
H2	C3391-82.3	82.8	3.11%	H3z	C3391-208.1	208.6	3.25%
H2	C3391-83.3	83.8	4.58%	H3z	C3391-209.1	209.6	2.65%
H2	C3391-85.05	85.55	3.28%	H3z	C3391-210.3	210.8	3.05%
H2	C3391-86.05	86.55	4.78%	H3z	C3391-211.3	211.8	2.46%
H2	C3391-87.35	87.85	3.42%	PPlz	C3391-212.8	213.3	4.22%
H2	C3391-88.35	88.85	3.22%	PPlz	C3391-213.8	214.3	3.66%
H2	C3391-89.75	90.25	3.83%	PPlz	C3391-215.6	216.1	4.15%
H2	C3391-90.75	91.25	3.08%	PPlz	C3391-216.6	217.1	3.68%
H2	C3391-92.05	92.55	4.41%	PPlz	C3391-218.7	219.2	22.81%
H2	C3391-93.05	93.55	2.77%	PPlz	C3391-219.7 below sand	220.2	22.30%
H2	C3391-94.5	95	3.66%	PPlz	C3391-219.7 above sand layer	220.2	26.20%
H2	C3391-95.5	96	2.71%	PPlz	C3391-220.65	221.15	20.62%
H2	C3391-96.5	97	3.08%	PPlz	C3391-221.65	222.15	16.64%
H2	C3391-97.5	98	2.60%	PPlg	C3391-224.5	225	3.39%
H2	C3391-99.15	99.65	3.03%	PPlg	C3391-225.5	226	2.23%
H2	C3391-100.15	100.65	2.96%	PPlg	C3391-227	227.5	2.59%
H2	C3391-101.6	102.1	3.14%	PPlg	C3391-228	228.5	2.75%
H2	C3391-102.6	103.1	2.85%	PPlg	C3391-230.2	230.7	3.08%
H2	C3391-103.7	104.2	3.29%	PPlg	C3391-231.2	231.7	2.27%
H2	C3391-104.7	105.2	4.21%	PPlg	C3391-233	233.5	3.50%

Lithologic Unit	Sample No.	Mid Depth ^(a) (Vertical ft)	% Moisture	Lithologic Unit	Sample No.	Mid Depth ^(a) (Vertical ft)	% Moisture
H2	C3391-106.3	106.8	3.57%	PPlg	C3391-234	234.5	3.03%
H2	C3391-107.3	107.8	3.82%	PPlg	C3391-235.8	236.3	2.20%
H2	C3391-108.3	108.8	3.57%	PPlg	C3391-236.8	237.3	2.81%
H2	C3391-109.3	109.8	3.56%	PPlg	C3391-238.8	239.3	2.60%
H2	C3391-110.3	110.8	3.51%	PPlg	C3391-239.8	240.3	3.27%
H2	C3391-111.3	111.8	3.05%	PPlg	C3391-241.9	242.4	3.52%
H2	C3391-112.45	112.95	3.72%	PPlg	C3391-242.9	243.4	2.55%
H2	C3391-113.45	113.95	3.03%	PPlg	C3391-244.5	245	2.51%
H2	C3391-114.4	114.9	4.19%	PPlg	C3391-245.5	246	2.64%
H2	C3391-115.4	115.9	3.35%	PPlg	C3391-247.1	247.6	2.80%
H2	C3391-118.5	119	5.74%	PPlg	C3391-248.1	248.6	2.77%
H2	C3391-119.5	120	2.98%	PPlg	C3391-250.9	251.4	3.87%
H2	C3391-121.35	121.85	3.20%	PPlg	C3391-251.9	252.4	3.35%
H2	C3391-122.35	122.85	2.36%	PPlg	C3391-256.1	256.6	6.59%
H2	C3391-123.35	123.85	2.66%	PPlg	C3391-257.1	257.6	6.80%
(a) Multiply	y by 0.3048 to con	nvert to meters.					

Table 4.1. (contd)

Figure 2.2 shows the comparison between the neutron moisture log data and the laboratory moisture measurements along with spectral gamma logs for borehole 299-E33-338. In this figure the neutron log spectra represents a function of the relative moisture content with a resolution of 0.3-m (1-ft) intervals. The overall agreement in terms of identifying areas of elevated moisture when compared to laboratory measurements (Table 4.1) is good and especially evident in unit boundaries and/or lithologic contacts. The spectral gamma log shows the profile of naturally occurring isotopes (e.g. potassium-40), which are greater in fine-grained strata, with no indication of anthropogenic gamma emitter contamination.



Figure 4.1. Borehole 299-E33-338 Lithology, Stratigraphy, Field Logs, and Moisture Distribution as a Function of Depth

299-E33-338

4.2 Particle Size Distribution and Particle Density

This section describes the particle size distribution for the E33-338 borehole that was determined using both dry sieving and hydrometer methods, and the pycnometer method for the determination of particle density.

4.2.1 Particle Size Distribution

Both dry sieving and the wet sieve/hydrometer methods (as described in section 3.1.3) were used to determine the particle size distributions from the same core material associated with the primary 22 samples. Depending on silt content, roughly 60 to 100-g of the bulk sediment was initially used for particle size determination using the wet sieve/hydrometer method. Large gravel material (> 1.0 cm) in the case of sample PPlg 239.8 was removed before the start of the analysis. At the completion of this test, the clay fraction was removed, air dried, and weighed. The clay removal process generally required 4-8 days to complete. The weight of the clay fraction removed for XRD analysis would later be added back to the weight of the total passing through the 270-mesh sieve for the purpose of calculating the particle distribution using the sieve method. The remaining silt and sand was then washed from the graduated cylinder onto a 270-mesh sieve and then oven dried. All material passing through the sieve was collected, oven dried and the weight recorded.

The oven dried sand fraction sample was dry sieved using sieve numbers 10, 18, 35, 60, 140, 200, and 270 (2000, 1000, 500, 250, 106, 75, and 53- μ m respectively). Figures 4.2 through 4.7 show the combined particle size distributions (as percent passing through) for each method as a continuum. It should be noted that in many cases there was overlapping in the percent passing through between 53 and 80- μ m with rather good agreement usually within 1 to 2%. It should also be noted that only the sieve method represents a minimum diameter passing through while the hydrometer method represents a mean diameter for particles passing through.

Overall results from the particle size distributions support both laboratory and field observations. Figure 4.2 shows a predominantly sand composition in the H1 unit with a silty-sand lens at 15.7 m (51.5 ft) bgs, which is consistent with the lithogy shown in Figure 2.2. Figure 4.3 shows a fairly homogenous sand matrix in the H2 unit from 23.6 to 35.2 m (77.3 to 115.4 ft) bgs. Figure 4.4 shows a gradual fining downward within the H2 unit. Figure 4.5 shows that both samples taken at 60.4 to 70 m (198.1 and 200.1 ft) bgs within the H3 unit were almost identical, consisting mostly of sand. Figure 4.6 shows a general fining downward in the PPlz unit from silty sand at 65.2 m (213.8 ft) to clayey silt at 67.0 m (219.7 ft). Figure 4.7 shows the dramatic difference between the three samples in the PPlz with a depth range of 67 to 67.6 m (219.7 to 221.7 ft) bgs and the one sample from the PPlg unit at a depth of 73.1 m (239.8 ft) bgs with a clearly far less silt and clay composition. It should be noted however that the actual percent gravel content for this sample is substantially under reported due to the initial removal of large gravel material (> 1.0 cm).

Figure 4.8 is a simple bar chart representation of the sand, silt and clay composition for each sample using the hydrometer method showing the major transitions from predominantly sand composition to a silt composition at 67.0 m (219.7 ft) bgs (the PPlz unit contact) and then back to a predominantly sand composition at 73.1 m (239.8 ft) bgs (the PPlg unit contact).



Figure 4.2. Particle Size Distribution 15.5-51.05 ft bgs.







Figure 4.4. Particle Size Distribution 133-173.05 ft bgs.







Figure 4.6. Particle Size Distribution 213.8-219.7 ft bgs.







4.2.2 Particle Density

The particle density of bulk sediment samples was determined for selected depths from borehole 299-E33-338 using pychnometers. Each sample was run in triplicate and the results of the mean value and standard deviation are shown in Table 4.2.

The particle densities in Table 4.2 reflect the mineral composition of the Hanford H1 coarse sand unit, the Hanford H2 upper sequence, PPlz unit, and PPlg unit. In general, the major mineral composition for these samples (as discussed in section 4.6) is mostly quartz, plagioclase, and potassium feldspar and basaltic rock fragments (in the case of the Hanford formation).

Sample ID	Depth ^(a) (ft bgs)	Particle Density (g/cm ³)	Standard Deviation							
	Hanford Format	ion H1 Unit	·							
C3391-15.5	16	2.565	0.023							
	Hanford Format	ion H2 Unit								
C3391-77.3	77.8	2.580	0.019							
115.4	115.9	2.683	0.030							
C3391-173.05	173.55	2.712	0.020							
	Plio-Pleistocene Si	lty Unit (PPlz)								
C3391-218.7	219.2	2.712	0.030							
Plio-Pleistocene Gravely Unit (PPlg)										
C3391-239.8	240.3	2.759	0.017							

 Table 4.2. Particle Densities for Selected Samples from Borehole 299-E33-338

(a) multiply by 0.3048 to convert to meters

4.3 Soil Water Chemistry Measurements

An extensive water chemistry analysis has been completed for borehole 299-E33-338 samples collected between 5 and 73 m (16 and 240 ft) bgs. Chemical characteristics show no strong trends as a function of depth and there is little, if any, indication of tank waste interaction with vadose zone soils at this location. Primary characteristics include the following:

- The 1:1 sediment-to-water extract pH varied from 7.2 to 7.8 and in general increased with depth (Figure 4.9 and Table 4.3).
- The average pH value is 7.4 with a range from 6.97 to 7.74.
- There were small increases in pH at the contact between the Hanford H2 and H3 units and the top and bottom of the Plio-Pleistocene mud unit.
- The dilution corrected water extract EC is an estimate of the vadose porewater EC.
- Porewater EC varied from 0.88 to 4.3 mS/cm with an average of 2.4 mS/cm.
- There were high EC values deep in the Hanford H2 unit at approximately 49 m (160 ft) bgs and in the deepest sample characterized (i.e., in the PPlg).

Sample ID	Mid Depth ^(a) (ft)	Dilution Factor	1:1 pH	1:1 EC (mS/cm)	Pore EC (mS/cm)						
	Hanford Form	ation H1 Un	it								
C3391-15.5	16	23.82	6.97	0.178	4.24						
C3391-17.5	18	9.66	7.39	0.235	2.27						
C3391-51.05	51.55	7.78	7.22	0.366	2.85						
	Hanford Form	ation H2 Un	it								
C3391-77.3 77.8 27.10 7.14 0.088 2.38											
C3391-83.3	83.8	21.84	7.34	0.095	2.07						
C3391-90.75	91.25	32.53	7.23	0.081	2.63						
C3391-107.3	107.8	26.18	7.23	0.087	2.28						
C3391-115.4	115.9	29.85	7.28	0.113	3.37						
C3391-133	133.5	33.26	7.28	0.08	2.66						
C3391-160.3	160.8	38.68	7.33	0.099	3.83						
C3391-161.35	161.85	26.87	7.38	0.148	3.98						
C3391-171.45	171.95	13.65	7.38	0.122	1.66						
C3391-173.05	173.55	7.01	7.5	0.203	1.42						
	Hanford Form	ation H3 Un	it								
C3391-198.1	198.6	35.88	7.29	0.09	3.23						
C3391-200.1	200.6	16.59	7.42	0.112	1.86						
	Plio-Pleistocene	e Silt Unit(Pl	Plz)								
C3391-213.8	214.3	27.30	7.34	0.112	3.06						
C3391-218.7	219.2	4.39	7.74	0.247	1.08						
C3391-219.7 below sand	220.2	4.60	7.59	0.201	0.92						
C3391-219.7 above sand layer	220.2	4.19	7.62	0.211	0.88						
C3391-220.65	221.15	4.87	7.62	0.213	1.04						
C3391-221.65	222.15	6.02	7.67	0.226	1.36						
	Plio-Pleistocene G	ravely Unit ((PPlg)								
C3391-239.8	240.3	30.55	7.52	0.143	4.37						

Table 4.3. Water Extract pH and Calculated Porewater Electrical ConductivityValues for Borehole 299-E33-338.

(a) Multiply by 0.3048 to convert to meters. Each sample was approximately 10-in. long, the mid point is used for plotting.

EC = Electrical conductivity

299-E33-338



Figure 4.9. Borehole 299-E33-338 Lithology, Stratigraphy, and Moisture, Extract pH and Calculated Porewater Electrical Conductivity Distribution as a Function of Depth.

Figure 4.10 and Table 4.4 show the estimated porewater concentrations of major cations and trace metals, respectively. The shapes of the cation profiles versus depth are very similar with slight peaks in the deep portion of the H2 unit at approximately 49 m (160 ft) bgs, at the top of the PPlz, and in the deepest sample characterized in the PPlg unit. All three of these samples had very low water contents and thus the dilution factor was high. The apparent high porewater concentrations likely represent some dissolution of salts from the sediment that are multiplied by a large dilution factor and thus suggest more saline porewater than surrounding sediments with higher water content. In general, the calculated porewater cation concentrations ranged from 63 to 275, 11 to 138, 11 to 56, and 70 to 558 mg/L for calcium, potassium, magnesium, and sodium, respectively. The averages and median values were (142, 149), (60, 58), (34, 38), and (190, 141) mg/L, for calcium, potassium, magnesium, and sodium, respectively. These values are likely somewhat artificially elevated because of the water extraction of soluble salts.

Figure 4.11, Table 4.4, and Table 4.5 show the calculated porewater concentrations for aluminum, barium, iron, silicon, and uranium-238. Of particular interest are the porewater aluminum, iron, and uranium-238 concentrations that ranged from 0.01 to 5.29, 0.0 to 6.4, and 46 to 350 mg/L for aluminum, barium, and iron, respectively, and 1.8 to 24 μ g/L for uranium. The uncontaminated uranium-238 porewater concentration is especially important for comparison with the suspect or known contaminated borehole sediment porewaters. No tank waste derived radionuclides were detected in these soils. Small quantities presumably of naturally occurring uranium (i.e., approximately 2.0 to 24 μ g/L) were measured in all water extract samples.

Figure 4.12 and Table 4.6 show the estimated porewater concentration of major anions. The shapes of the anion profiles versus depth vary from each other instead of being similar as were the cation profiles. There are no consistent depths where all anions peak. The wetter samples do consistently show low calculated porewater anion concentrations suggesting that the dilution factor is again controlling the apparent concentrations. That is, all the sediments likely dissolve some salts that are not truly in the porewater so that the dilution correction makes it appear that the porewater anion concentrations are higher in the drier sediments. Primary constituents are carbonate, sulfate, and chloride. Other less concentrated anions include fluoride and nitrate. In general, the calculated porewater anion concentrations range from 0.4 to 23.3, 1.8 to 223, 1.3 to 100, 296 to 1877, and 196 to 117 mg/L for fluoride, chloride, nitrate, bicarbonate, and sulfate, respectively. The average and median values are (6.3, 4.8), (35, 19), (19, 10), (993, 1030), and (196, 117) mg/L for fluoride, chloride, nitrate, bicarbonate, are likely somewhat artificially elevated because of the water extraction of soluble salts.

The mass of several constituents per gram of dry sediment that were leached by water and acid extracts are shown in Figures 4.13 and 4.14 and in Tables 4.7 and 4.8. In all cases, the mass that was water leachable is a very small fraction of the mass that was acid extractable. These concentrations can be compared with the same constituents for contaminated sediments to get an estimate of the mass of a constituent present in the vadose zone profile from tank leaked liquids.



299-E33-338

Figure 4.10. Calculated Cation Porewater Content for Borehole 299-E33-338 as a Function of Depth.



Figure 4.11. Calculated Aluminum, Barium, Iron, Silicon, and Uranium Porewater Content for Borehole 299-E33-338 as a Function of Depth.

299-E33-338

	Depth ^(a)	(a) Dilution Factor			Dilution	n Correcte	d Porewater (Concentration o	of Cations		
Sample Identification	(ft bgs)	Dilution Factor	Aluminum (mg/L)	Barium (mg/L)	Calcium (mg/L)	Iron (mg/L)	Potassium (mg/L)	Magnesium (mg/L)	Sodium (mg/L)	Silicon (mg/L)	Strontium (mg/L)
			Н	Hanford Formation H1 Unit							
C3391-15.5	16.00	23.82	(0.64)	0.83	141.47	(0.5)	55.40	18.64	557.59	185.17	0.55
C3391-17.5	18.00	9.66	(0.37)	0.36	73.81	(0.6)	30.90	8.57	336.59	147.04	0.30
C3391-51.05	51.55	7.78	(3.89)	0.28	274.70	(0.0)	47.33	65.55	91.56	74.21	1.25
			Н	anford Fori	nation H2 U	Init					
C3391-77.3	77.80	27.10	(1.00)	0.57	156.34	(0.6)	69.13	42.33	134.52	139.23	0.78
C3391-83.3	83.80	21.84	(0.70)	1.03	131.90	(0.5)	58.53	36.31	122.00	144.05	0.69
C3391-90.75	91.25	32.53	(1.76)	0.72	166.33	(1.2)	65.44	40.05	148.22	147.90	0.82
C3391-107.3	107.80	26.18	(1.28)	0.62	157.52	(1.0)	62.42	40.38	117.00	136.04	0.75
C3391-115.4	115.90	29.85	(1.68)	0.92	203.65	(2.9)	89.18	57.88	203.48	258.67	1.09
C3391-133	133.50	33.26	(1.91)	0.42	170.14	(2.6)	76.98	41.96	147.85	168.98	0.85
C3391-160.3	160.80	38.68	(3.12)	1.24	225.28	(2.8)	111.93	59.88	250.21	298.74	1.33
C3391-161.35	161.85	26.87	(5.29)	1.34	200.56	6.4	138.20	55.81	385.86	251.11	1.20
C3391-171.45	171.95	13.65	(0.54)	0.55	92.03	(0.7)	47.11	25.67	123.22	129.21	0.52
C3391-173.05	173.55	7.01	(0.22)	0.28	75.06	(0.2)	38.24	20.65	122.01	73.53	0.44
	Hanford Formation H3 Unit										
C3391-198.1	198.60	35.88	(1.76)	0.72	168.53	(1.1)	90.54	41.63	237.88	213.35	0.96
C3391-200.1	200.60	16.59	(0.55)	0.42	103.09	(0.3)	56.60	26.62	152.86	107.71	0.55

 Table 4.4. Calculated Cation Porewater Content for Borehole 299-E33-338.

	Donth ^(a)	(a) Dilution Factor			Dilutio	n Correcte	ed Porewater	Concentration of	of Cations		
Sample Identification	(ft bgs)		Aluminum (mg/L)	Barium (mg/L)	Calcium (mg/L)	Iron (mg/L)	Potassium (mg/L)	Magnesium (mg/L)	Sodium (mg/L)	Silicon (mg/L)	Strontium (mg/L)
			Plio	-Pleistocen	e Silty Unit ((PPlz)					
C3391-213.8	214.30	27.30	(2.46)	1.05	183.56	3.6	90.50	43.14	285.91	232.64	1.08
C3391-218.7	219.20	4.39	(0.04)	0.22	81.52	(0.0)	14.26	15.59	79.93	51.84	0.44
C3391-219.7 below sand	220.20	4.60	(0.05)	0.22	66.11	(0.1)	11.41	11.31	72.92	46.34	0.32
C3391-219.7 above sand layer	220.20	4.19	(0.01)	0.15	62.73	(0.1)	10.62	10.88	70.28	45.82	0.30
C3391-220.65	221.15	4.87	(0.08)	0.25	70.44	(0.0)	15.58	12.31	83.32	45.93	0.38
C3391-221.65	222.15	6.02	(0.23)	0.33	94.22	(0.3)	24.49	17.06	109.31	63.02	0.51
Plio-Pleistocene Gravely Unit (PPlg)											
C3391-239.8	240.30	30.55	(0.77)	1.24	232.35	(0.7)	108.56	62.56	344.76	349.64	1.30

Table 4.4. (contd)

(a) Multiply by 0.3048 to convert to meters. bgs = below ground surface

Values in parentheses (.....) are below level of quantification but spectra look useable.

Sample ID	Depth ^(a) (ft bgs)	Dilution Factor	Cr (µg/L)	As (µg/L)	Se (µg/L)	U-238 (μg/L)						
		Hanfe	ord Formation H	II Unit								
C3391-15.5	16.000	23.820	2.39E+01	(1.08E+02)	(6.38E+00)	1.14E+01						
C3391-17.5	18.00	9.66	1.82E+01	1.16E+02	(6.09E+00)	1.33E+01						
C3391-51.05	51.550	7.777	1.05E+01	4.58E+01	2.06E+01	3.71E+00						
Hanford Formation H2 Unit												
C3391-77.3 77.800 27.096 (8.35E+00) 1.42E+02 (6.23E+00) 6.33E+00												
C3391-83.3	83.800	21.841	1.616E+01	1.47E+02	<5.46E+01	8.66E+00						
C3391-90.75	91.250	32.530	(1.01E+01)	2.64E+02	(6.73E+00)	6.15E+00						
C3391-107.3	107.800	26.181	(9.74E+00)	2.01E+02	<6.55E+01	7.51E+00						
C3391-115.4	115.900	29.850	(1.18E+01)	3.81E+02	<7.46E+01	9.98E+00						
C3391-133	133.500	33.263	(2.02E+01)	2.64E+02	<8.32E+01	1.12E+01						
C3391-160.3	160.800	38.681	(1.26E+01)	4.38E+02	(3.56E+00)	1.56E+01						
C3391-161.35	161.850	26.867	2.42E+01	3.74E+02	(7.79E-01)	2.37E+01						
C3391-171.45	171.950	13.646	(6.05E+00)	2.05E+02	<3.41E+01	1.01E+01						
C3391-173.05	173.55	7.01	7.20E+00	1.40E+02	<1.75E+01	7.04E+00						
		Hanfe	ord Formation H	13 Unit								
C3391-198.1	198.600	35.880	(9.80E+00)	2.49E+02	<8.97E+01	7.64E+00						
C3391-200.1	200.600	16.586	(4.35E+00)	1.36E+02	<4.15E+01	1.22E+01						
		Plio-Ple	eistocene Silty U	nit (PPlz)								
C3391-213.8	214.300	27.297	1.65E+01	6.20E+02	<6.82E+01	9.61E+00						
C3391-218.7	219.20	4.39	2.67E+01	4.70E+01	(1.98E+00)	5.04E+00						
C3391-219.7 below sand	220.200	4.602	(1.78E+01)	3.27E+01	(1.09E+00)	1.79E+00						
C3391-219.7 above sand layer	220.200	4.186	(2.09E+01)	3.77E+01	(2.22E+00)	2.00E+00						
C3391-220.65	221.150	4.875	(1.53E+01)	3.17E+01	(2.49E+00)	2.93E+00						
C3391-221.65	222.150	6.023	1.55E+01	5.60E+01	(2.61E+00)	4.61E+00						
	Plio-Pleistocene Gravely Unit (PPlg											
C3391-239.8	240.300	30.545	(9.47E+00)	2.38E+02	(1.56E+00)	1.52E+01						

Table 4.5. Calculated Porewater Trace Metal Composition for Water Extracts of Sediment.

(a) Multiply by 0.3048 to convert to meters.

bgs = below ground surface

Values in parentheses (.....) are below level of quantification but spectra look useable.

Sample	Depth ^(a)	Dil.	1:1 Extracts in mg/L							Dilution Corrected Porewater mg/L						
ID	(ft bgs)	Fac.	NO3	F-	NO2	Cl	SO4	PO4	HCO3	NO3	F-	NO2	Cl	SO4	PO4	HCO3
						Hanf	ford Fori	nation H1 U	Unit							
C3391-15.5	16.00	23.82	1.44	0.22	<0.14	1.35	34	< 0.24	43.8	34.26	5.18	<3.26	32.09	800	<5.7	1042
C3391-17.5	18.00	9.66	0.42	0.81	<0.14	1.10	40	< 0.25	70.4	3.99	7.64	<1.32	10.45	376	<2.3	680
C3391-51.05	51.55	7.78	13.02	0.31	0.19	28.94	68	< 0.24	38.1	100.53	2.42	1.46	223.42	521	<1.9	296
						Hanf	ford Fori	nation H2 U	Unit							
C3391-77.3	77.80	27.10	1.00	0.19	< 0.14	0.78	3.4	< 0.24	60.9	27.12	5.08	<3.71	21.07	92	<6.5	1650
C3391-83.3	83.80	21.84	0.85	0.18	< 0.14	0.44	3.3	< 0.24	46.6	18.64	3.95	<2.99	9.62	73	<5.2	1018
C3391-90.75	91.25	32.53	1.27	0.18	<0.14	0.48	2.4	< 0.24	40.9	41.38	5.78	<4.46	15.62	79	<7.8	1331
C3391-107.3	107.80	26.18	0.05	0.02	<0.14	0.07	<0.5	< 0.24	46.6	1.31	0.45	<3.59	1.80	<6.3	<6.3	1220
C3391-115.4	115.90	29.85	1.55	0.26	< 0.13	4.24	4.8	< 0.23	42.8	48.17	8.06	<4.09	131.42	148	<7.2	1278
C3391-133	133.50	33.26	0.38	0.20	< 0.14	0.45	1.8	< 0.24	46.6	12.56	6.79	<4.56	15.10	61	<8.0	1551
C3391-160.3	160.80	38.68	0.47	0.60	< 0.14	1.48	10	< 0.24	48.5	18.22	23.28	<5.30	57.12	386	<9.3	1877
C3391-161.35	161.85	26.87	0.37	0.43	< 0.14	1.23	7.2	< 0.24	61.9	9.86	11.50	<3.68	33.03	193	< 6.5	1661
C3391-171.45	171.95	13.65	0.56	0.33	< 0.14	0.55	6.1	< 0.24	53.3	7.66	4.49	<1.87	7.51	83	<3.3	727
C3391-173.05	173.55	7.01	0.53	0.57	< 0.14	1.55	20	< 0.24	71.4	3.68	3.96	< 0.96	10.83	141	<1.7	500
						Hanf	ford For	nation H3 U	Unit							
C3391-198.1	198.60	35.88	0.47	0.20	< 0.14	0.35	2.7	0.26	48.5	16.85	7.28	<4.92	12.43	96	9.4	1741
C3391-200.1	200.60	16.59	< 0.29	0.25	< 0.14	0.92	5.4	< 0.24	50.4	<4.84	4.14	<2.27	15.17	90	<4.0	836
						Plio-Pl	eistocen	e Silty Unit	(PPlz)							
C3391-213.8	214.30	27.30	0.50	0.35	<0.14	0.87	6.7	< 0.24	52.3	13.57	9.48	<3.74	23.72	182	<6.6	1428
C3391-218.7	219.20	4.39	2.09	0.60	<0.14	4.20	28	0.33	86.6	9.15	2.61	< 0.60	18.40	121	1.4	379
C3391-219.7 <mark>bs</mark>	220.20	4.60	1.97	0.56	<0.14	4.09	18	0.42	76.1	9.18	2.59	<0.63	19.10	84	2.0	350
C3391-219.7 as	220.20	4.19	2.21	0.66	<0.13	4.14	18	0.55	74.2	9.71	2.89	<0.57	18.22	79	2.4	310

 Table 4.6. Calculated Anion Porewater Content for Borehole 299-E33-338.

Table 4.6. (contd)

Sample	Depth ^(a)	Depth ^(a) Dil.		1:1 Extracts in mg/L						Dilution Corrected Porewater mg/L							
ID	(ft bgs)	Fac.	NO3	F-	NO2	Cl	SO4	PO4		HCO3	NO3	F-	NO2	Cl	SO4	PO4	HCO3
C3391-220.65	221.15	4.87	0.56	0.55	<0.14	6.67	23	0.40		62.8	2.72	2.69	< 0.67	32.33	113	2.0	306
C3391-221.65	222.15	6.02	1.38	0.64	<0.14	5.51	28	0.45		77.1	8.27	3.83	< 0.83	33.11	169	2.7	464
Plio-Pleistocene Gravely Unit (PPlg)																	
C3391-239.8	240.30	30.55	0.31	0.	49 <0	.14 0.	69 1	3 <0.	24	39.0	9.43	15.02	<4.19	21.00	39	<7.3	1191

(a) Multiply by 0.3048 to convert to meters.as = above sandbs =below sand



Figure 4.11. Calculated Anion Porewater Content for Borehole 299-E33-338 as a Function of Depth.

Sample ID	Mid Depth ^(a) (ft)	Ca (µg/g)	Mg (µg/g)	Sr (μg/g)	Na (µg/g)	К (µg/g)						
		Hanford	l Formation H1 U	Unit								
C3391-15.5	16.00	5.95E+00	7.84E-01	2.33E-02	2.34E+01	2.33E+00						
17.5	18.00	7.80E+00	9.06E-01	3.13E-02	3.56E+01	3.26E+00						
51.05	51.55	3.56E+01	8.49E+00	1.62E-01	1.19E+01	6.13E+00						
H2 Hanford Formation H2 Unit												
77.3 77.80 5.78E+00 1.56E+00 2.87E-02 4.97E+00 2.55E+00												
83.3	83.80	6.04E+00	1.66E+00	3.16E-02	5.59E+00	2.68E+00						
90.75	91.25	5.12E+00	1.23E+00	2.54E-02	4.56E+00	2.01E+00						
107.3	107.80	6.02E+00	1.54E+00	2.86E-02	4.47E+00	2.38E+00						
115.4	115.90	6.57E+00	1.87E+00	3.51E-02	6.56E+00	2.88E+00						
133	133.50	5.12E+00	1.26E+00	2.56E-02	4.45E+00	2.31E+00						
160.3	160.80	5.83E+00	1.55E+00	3.44E-02	6.47E+00	2.90E+00						
161.35	161.85	7.47E+00	2.08E+00	4.48E-02	1.44E+01	5.14E+00						
171.45	171.95	6.75E+00	1.88E+00	3.78E-02	9.03E+00	3.45E+00						
173.05	173.55	1.07E+01	2.95E+00	6.27E-02	1.74E+01	5.46E+00						
		Hanford	d Formation H3 (Unit								
198.1	198.60	4.71E+00	1.16E+00	2.69E-02	6.64E+00	2.53E+00						
200.1	200.60	6.22E+00	1.61E+00	3.30E-02	9.22E+00	3.41E+00						
		Plio-Pleist	tocene Silty Unit	(PPlz)								
213.8	214.30	6.73E+00	1.58E+00	3.94E-02	1.05E+01	3.32E+00						
218.7	219.20	1.86E+01	3.55E+00	9.97E-02	1.82E+01	3.25E+00						
219.7 Below Sand	220.20	1.42E+01	2.42E+00	6.92E-02	1.56E+01	2.44E+00						
219.7 Above Sand	220.20	1.43E+01	2.47E+00	6.81E-02	1.60E+01	2.42E+00						
220.65	221.15	1.45E+01	2.54E+00	7.76E-02	1.72E+01	3.21E+00						
221.65 222.15 1.57E+01 2.84E+00 8.54E-02 1.82E+01 4.07E+00												
Plio-Pleistocene Gravely Unit (PPlg)												
239.8	240.30	7.61E+00	2.05E+00	4.26E-02	1.13E+01	3.55E+00						

Table 4.7. Water Extract of Major Cations in Terms of Dry Sediment ($\mu g/g$).

Sample Identification	Mid Depth ^(a) (ft)	Al (µg/g)	Ba (µg/g)	Fe (µg/g)	Mg (µg/g)	Si (µg/g)	U-238 (µg/g)
	L	Han	ford Formatic	on H1 Unit	ł	L	
C3391-15.5	16.00	2.71E-02	3.47E-02	2.09E-02	7.84E-01	7.79E+00	4.78E-04
17.5	18.00	3.88E-02	3.76E-02	5.99E-02	9.06E-01	1.55E+01	1.40E-03
51.05	51.55	5.04E-01	3.61E-02	3.92E-03	8.49E+00	9.61E+00	4.81E-04
		Han	ford Formatic	on H2 Unit			
77.3	77.80	3.70E-02	2.10E-02	2.33E-02	1.56E+00	5.14E+00	2.34E-04
83.3	83.80	3.20E-02	4.70E-02	2.13E-02	1.66E+00	6.60E+00	3.97E-04
90.75	91.25	5.40E-02	2.23E-02	3.71E-02	1.23E+00	4.55E+00	1.89E-04
107.3	107.80	4.89E-02	2.37E-02	3.67E-02	1.54E+00	5.20E+00	2.87E-04
115.4	115.90	5.41E-02	2.98E-02	9.33E-02	1.87E+00	8.34E+00	3.22E-04
133	133.50	5.73E-02	1.28E-02	7.67E-02	1.26E+00	5.08E+00	3.36E-04
160.3	160.80	8.07E-02	3.20E-02	7.30E-02	1.55E+00	7.73E+00	4.04E-04
161.35	161.85	1.97E-01	4.98E-02	2.39E-01	2.08E+00	9.35E+00	8.82E-04
171.45	171.95	3.97E-02	4.06E-02	5.20E-02	1.88E+00	9.47E+00	7.44E-04
173.05	173.55	3.18E-02	4.03E-02	3.34E-02	2.95E+00	1.05E+01	1.00E-03
		Han	ford Formatio	on H3 Unit			
198.1	198.60	4.90E-02	2.01E-02	3.08E-02	1.16E+00	5.96E+00	2.13E-04
200.1	200.60	3.31E-02	2.51E-02	1.83E-02	1.61E+00	6.50E+00	7.35E-04
		Plio-P	leistocene Silt	y Unit (PPlz)			
213.8	214.30	9.01E-02	3.86E-02	1.31E-01	1.58E+00	8.52E+00	3.52E-04
218.7	219.20	9.40E-03	4.90E-02	1.11E-02	3.55E+00	1.18E+01	1.15E-03
219.7 Below Sand	220.20	1.10E-02	4.61E-02	1.21E-02	2.42E+00	9.93E+00	3.84E-04
219.7 Above Sand	220.20	3.06E-03	3.51E-02	1.30E-02	2.47E+00	1.04E+01	4.55E-04
220.65	221.15	1.66E-02	5.25E-02	9.34E-03	2.54E+00	9.47E+00	6.04E-04
221.65	222.15	3.77E-02	5.43E-02	5.76E-02	2.84E+00	1.05E+01	7.67E-04
		Plio-Ple	istocene Grav	ely Unit (PPlg	<u>z)</u>		
239.8	240.30	2.53E-02	4.06E-02	2.20E-02	2.05E+00	1.14E+01	4.96E-04

Table 4-7. (contd)

(a) Multiply by 0.3048 to convert to meters.

Sample ID	Mid Depth ^(a) (ft)	ml/g basis acid: soil ratio	Ca (µg/g)	Mg (µg/g)	Sr (µg/g)	Na (µg/g)	Κ (μg/g)						
	1	Н	anford Forma	tion H1 Unit	1	1	1						
C3391-15.5	16.00	4.37	8.32E+03	3.97E+03	3.39E+01	3.81E+02	8.96E+02						
17.5	18.00	4.97	9.58E+03	4.55E+03	3.73E+01	5.58E+02	1.08E+03						
51.05	51.55	4.84	8.70E+03	5.28E+03	3.89E+01	3.48E+02	1.66E+03						
Hanford Formation H2 Unit													
77.3 77.80 5.01 7.69E+03 4.64E+03 2.99E+01 2.87E+02 1.14E+03													
83.3	83.80	5.47	8.23E+03	4.67E+03	3.38E+01	2.75E+02	1.31E+03						
90.75	91.25	5.47	7.32E+03	4.77E+03	3.24E+01	2.78E+02	1.17E+03						
107.3	107.80	5.01	7.08E+03	4.66E+03	2.68E+01	2.50E+02	1.04E+03						
115.4	115.90	5.00	7.48E+03	4.72E+03	2.98E+01	3.37E+02	1.18E+03						
133	133.50	5.02	7.65E+03	4.67E+03	3.64E+01	3.20E+02	1.29E+03						
160.3	160.80	4.72	6.85E+03	4.80E+03	3.09E+01	3.05E+02	1.07E+03						
161.35	161.85	4.96	6.80E+03	4.63E+03	3.12E+01	3.43E+02	1.09E+03						
171.45	171.95	5.01	9.23E+03	5.39E+03	4.24E+01	3.19E+02	1.41E+03						
173.05	173.55	4.95	1.56E+04	9.43E+03	6.50E+01	4.45E+02	2.82E+03						
		Н	anford Forma	tion H3 Unit									
198.1	198.60	4.48	6.23E+03	4.02E+03	2.92E+01	3.23E+02	9.31E+02						
200.1	200.60	4.88	6.38E+03	4.34E+03	3.12E+01	3.43E+02	9.82E+02						
		Plio	Pleistocene S	Silty Unit (PPl	z)								
213.8	214.30	5.05	6.94E+03	4.90E+03	3.39E+01	2.76E+02	1.23E+03						
218.7	219.20	5.52	1.36E+04	8.47E+03	6.41E+01	4.13E+02	2.52E+03						
219.7 Below Sand	220.20	5.67	1.10E+04	6.71E+03	4.99E+01	3.52E+02	2.53E+03						
219.7 Above Sand	220.20	4.93	1.44E+04	9.34E+03	5.67E+01	3.05E+02	2.65E+03						
220.65	221.15	5.02	9.13E+03	5.62E+03	4.03E+01	2.70E+02	1.92E+03						
221.65	222.15	4.87	9.78E+03	6.13E+03	4.68E+01	3.77E+02	2.17E+03						
	Plio-Pleistocene Gravely Unit (PPlg)												
239.8	240.30	4.57	5.62E+03	3.56E+03	3.90E+01	3.72E+02	9.83E+02						

Table 4.8. Acid Extract of Major Cations in Terms of Dry Sediment (µg/g).

Table 4-8. (contd)

Sample ID	Mid Depth ^(a) (ft)	ml/g basis acid: soil ratio	Al (µg/g)	Ba (µg/g)	Fe (µg/g)	Si (µg/g)	U-238 (µg/g)
		ŀ	Hanford Form	ation H1 Unit			
C3391- 15.5	16.00	4.37	6.29E+03	9.11E+01	1.83E+04	1.18E+02	0.399
17.5	18.00	4.97	7.99E+03	9.80E+01	2.32E+04	(3.50E+01)	0.461
51.05	51.55	4.84	9.48E+03	7.29E+01	1.65E+04	(3.21E+01)	0.627
		I	Hanford Form	ation H2 Unit			
77.3	77.80	5.01	7.09E+03	6.28E+01	1.46E+04	(8.80E+01)	0.409
83.3	83.80	5.47	7.68E+03	7.12E+01	1.57E+04	(8.83E+01)	0.526
90.75	91.25	5.47	6.99E+03	5.83E+01	1.33E+04	(9.68E+01)	0.718
107.3	107.80	5.01	6.56E+03	5.51E+01	1.33E+04	1.26E+02	0.400
115.4	115.90	5.00	7.75E+03	7.29E+01	1.65E+04	(4.66E+01)	0.528
133	133.50	5.02	7.92E+03	6.87E+01	1.51E+04	(5.20E+01)	0.595
160.3	160.80	4.72	7.10E+03	6.86E+01	1.43E+04	(9.20E+01)	0.465
161.35	161.85	4.96	7.40E+03	6.72E+01	1.50E+04	(5.14E+01)	0.473
171.45	171.95	5.01	8.96E+03	6.66E+01	1.61E+04	(1.08E+02)	0.552
173.05	173.55	4.95	1.72E+04	1.62E+02	2.72E+04	(1.76E+01)	0.910
		I	Hanford Form	ation H3 Unit			
198.1	198.60	4.48	6.56E+03	6.78E+01	1.46E+04	(7.51E+01)	0.419
200.1	200.60	4.88	6.70E+03	7.83E+01	1.52E+04	(5.79E+01)	0.412
		Plie	o-Pleistocene	Silty Unit (PP	lz)		
213.8	214.30	5.05	7.58E+03	5.97E+01	1.26E+04	(8.34E+01)	0.430
218.7	219.20	5.52	1.69E+04	1.00E+02	2.53E+04	(1.75E+01)	0.797
219.7 Below Sand	220.20	5.67	1.66E+04	1.32E+02	2.23E+04	(1.14E+02)	0.918
219.7 Above Sand	220.20	4.93	2.25E+04	1.10E+02	3.94E+04	(2.08E+01)	1.343
220.65	221.15	5.02	1.08E+04	1.20E+02	1.70E+04	2.90E+02	0.597
221.65	222.15	4.87	1.36E+04	1.12E+02	1.82E+04	(7.74E+01)	0.660
	·	Plio-	Pleistocene G	ravely Unit (P	Plg)		
239.8	240.30	4.57	5.96E+03	7.72E+01	1.37E+04	(1.05E+02)	0.408

1

(a) Multiply by 0.3048 to convert to meters. Values in parentheses (.....) are below level of quantification but spectra look useable.



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Figure 4.12. Water and Acid Extractable Concentrations (µg/g) of Selected Constituents for Borehole 299-E33-338 as a Function of Depth.



Figure 4.13. Water and Acid Extractable Concentrations (µg/g) of Trace Constituents for Borehole 299-E33-338 as a Function of Depth.

4.4 Soil Fusion Analysis

Table 4.9 shows the major and trace element constituents for each of the characterized sediments derived from using a lithium metaborate/tetraborate fusion method as described in Section 3.1.12. Very little difference is observed in the primary elemental oxide concentrations for any of the sediment samples as a function of depth or lithology. The primary elemental oxides reported are SiO₂ (58.77-70.33 wt%), Al₂O₃ (12.7-15.73 wt%), Fe₂O₃ (3.74-7.92 wt%), CaO (3.02-4.80 wt%), Na₂O (2.05-3.23 wt%), K₂O (1.70-2.65 wt%), MgO (1.64-2.73 wt%), K₂O (1.70-2.65 wt%), TiO₂ (0.51-1.39 wt%), P₂O₅ (0.13-0.30 wt%), and MnO (0.07-0.12 wt%). An additional 35 trace elements are also reported in terms of ug/g of dry sediment with little if any significant variation in concentration.

Sample ID	Mid Depth ^(a) (ft)	SiO2 ^(b) %	Al2O3 ^(b) %	Fe2O3 ^(b) %	MnO ^(b) %	MgO ^(b) %	CaO ^(b) %	Na2O ^(b) %	K2O ^(b) %	TiO2 ^(b) %	P2O5 ^(b) %	LOI ^(b) %	TOTAL ^(b) %
	•			H	anford Fo	rmation H	II Unit						
C3391-15.5	16.00	64.24	12.92	6.75	0.114	2.23	4.80	3.01	1.82	1.211	0.25	2.75	100.09
C3391-17.5	18.00	63.42	12.86	7.01	0.107	2.17	4.77	3.04	1.70	1.145	0.24	2.85	99.30
C3391-51.05	51.55	61.82	13.11	7.92	0.116	2.73	5.28	2.97	1.94	1.391	0.30	2.64	100.21
				H	anford Fo	rmation H	I2 Unit						
C3391-77.3	77.80	68.20	12.72	4.78	0.076	1.93	3.67	2.88	2.33	0.711	0.17	2.19	99.65
C3391-83.3	83.80	68.37	13.14	4.85	0.076	1.86	3.67	2.95	2.27	0.722	0.17	2.31	100.39
C3391-90.75	91.25	70.33	12.70	3.93	0.066	1.64	3.18	2.73	2.49	0.543	0.13	2.22	99.95
C3391-107.3	107.80	69.96	12.96	4.21	0.070	1.69	3.32	2.87	2.45	0.593	0.14	2.13	100.40
C3391-115.4	115.90	65.68	13.30	6.09	0.098	2.27	4.30	3.05	2.10	0.925	0.19	2.20	100.20
C3391-133	133.50	69.41	12.66	4.03	0.069	1.59	3.25	2.86	2.53	0.568	0.14	2.06	99.17
C3391-160.3	160.80	69.00	13.24	4.41	0.073	1.79	3.50	3.22	2.22	0.630	0.15	1.83	100.06
C3391-161.35	161.85	69.56	12.94	4.29	0.071	1.74	3.36	3.20	2.17	0.611	0.15	1.91	99.99
C3391-171.45	171.95	68.65	13.03	4.54	0.080	1.87	3.70	3.24	2.21	0.636	0.17	2.11	100.23
C3391-173.05	173.55	66.76	12.98	4.67	0.084	1.94	3.91	3.26	2.10	0.665	0.20	2.65	99.21
				H	anford Fo	rmation H	13 Unit						
C3391-198.1	198.60	65.67	13.05	6.09	0.095	2.22	4.35	3.21	2.02	0.919	0.23	2.31	100.16
C3391-200.1	200.60	65.90	13.02	5.67	0.091	2.14	4.10	3.21	2.07	0.859	0.19	2.14	99.38
				Plio	-Pleistoce	ne Silty U	nit (PPlz)						
C3391-213.8	214.30	70.27	13.34	3.74	0.068	1.67	3.27	3.19	2.15	0.509	0.13	1.89	100.23
C3391-218.7	219.20	64.39	14.83	5.24	0.073	2.35	3.45	2.53	2.42	0.767	0.21	3.88	100.14
C3391-219.7 Below sand	220.20	65.24	14.81	5.09	0.073	2.10	3.22	2.41	2.43	0.771	0.22	3.96	100.33

Table 4.9a. Element Constituents from Soil Fusion Method.

 Table 4.9a. (contd)

Sample ID	Mid Depth ^(a) (ft)	SiO2 ^(b)	Al2O3 ^(b)	Fe2O3 ^(b) %	MnO ^(b) %	MgO ^(b) %	CaO ^(b) %	Na2O ^(b) %	K2O ^(b) %	TiO2 ^(b) %	P2O5 ^(b) %	LOI ^(b)	TOTAL ^(b)
C3391-219.7 Above sand	220.20	62.55	15.73	5.96	0.075	2.32	3.09	2.18	2.65	0.848	0.24	4.73	100.38
C3391-220.65	221.15	58.91	15.01	5.89	0.076	2.26	3.02	2.05	2.56	0.806	0.21	4.76	95.55
C3391-220.65 /R	221.15	58.77	15.02	5.86	0.078	2.33	3.04	2.13	2.57	0.791	0.22	4.76	95.56
C3391-221.65	222.15	63.79	14.54	5.15	0.072	2.23	3.46	2.53	2.43	0.772	0.22	4.00	99.21
				Plio-F	Pleistocen	e Gravely	Unit (PPl	g)					
C3391-239.8	240.30	63.05	13.71	7.16	0.113	2.51	4.98	3.23	2.08	1.086	0.25	1.69	99.85

(a) Multiply by 0.3048 to convert to meters.(b) Analysis by ICP-OES

Sample ID	Mid Depth ^(a) (ft)	Ba ^(b) (µg/g)	Sr ^(b) (µg/g)	Y ^(b) (µg/g)	Sc ^(b) (µg/g)	Zr ^(b) (µg/g)	Be ^(b) (µg/g)	V ^(b) (µg/g)	Cr ^(c) (µg/g)	Co ^(c) (µg/g)	Ni ^(c) (µg/g)	Cu ^(c) (µg/g)	Zn ^(c) (μg/g)
				Hanfor	d Format	ion H1 U	nit						
C3391-15.5	16	737	419	23	18	150	2	180	62	20	-20	27	55
C3391-17.5	18	739	419	22	17	139	2	176	41	23	-20	27	93
C3391-5105	51.55	695	355	26	20	182	2	213	53	27	-20	32	103
				Hanfor	d Format	ion H2 U	nit						
C3391-77.3	77.8	795	360	19	13	137	2	110	53	14	-20	32	64
C3391-83.3	83.8	843	383	20	11	128	2	109	43	14	-20	24	57
C3391-90.75	91.25	826	377	17	10	126	2	80	73	11	-20	28	67
C3391-107.3	107.8	873	394	18	10	118	2	89	66	12	-20	20	56
C3391-115.4	115.9	812	386	21	15	128	2	144	76	20	-20	28	81
C3391-133	133.5	871	389	16	10	115	2	80	38	10	21	17	50
C3391-160.3	160.8	857	415	17	12	110	2	98	61	12	-20	23	95
C3391-161.35	161.85	850	405	16	11	112	2	90	65	12	-20	23	68
C3391-171.45	171.95	852	423	19	11	168	2	100	80	13	26	27	70
C3391-173.05	173.55	842	425	21	11	200	2	99	86	14	32	30	80
				Hanfor	d Format	ion H3 U	nit						
C3391-198.1	198.6	817	397	19	15	109	2	141	52	19	-20	27	107
C3391-200.1	200.6	814	401	19	15	115	2	127	59	17	27	26	78
C3391-200.1 Rep	200.6			Dup	licate not	Run			79	17	25	26	82
			-	Plio-Pleis	stocene Si	lty Unit (I	PPlz)						
C3391-213.8	214.3	866	459	15	6	106	2	55	75	10	25	175	55
C3391-218.7	219.2	823	359	25	12	203	2	102	94	16	46	37	89
C3391-219.7 Below sand	220.2	830	352	28	12	245	2	94	87	17	46	40	92

 Table 4-9b.
 Element Constituents from Soil Fusion Method.

Table 9b. (contd)

Sample ID	Mid Depth ^(a) (ft)	Ba ^(b) (µg/g)	Sr ^(b) (µg/g)	Y ^(b) (µg/g)	Sc ^(b) (µg/g)	Zr ^(b) (µg/g)	Be ^(b) (µg/g)	V ^(b) (µg/g)	Cr ^(c) (µg/g)	Co ^(c) (µg/g)	Ni ^(c) (µg/g)	Cu ^(c) (µg/g)	Zn ^(c) (µg/g)
C3391-219.7 Above sand	220.2	1022	305	30	15	235	2	124	109	20	55	49	121
C3391-220.65	221.15	1021	296	26	14	215	2	120	112	19	73	70	100
C3391-220.65 /R	221.15	1022	299	24	14	206	3	118	93	16	50	38	97
C3391-221.65	222.15	841	364	26	13	246	2	105	90	14	50	33	90
				Plio-Pleis	stocene Gi	ravely Un	it (PPlg)						
C3391-239.8	240.3	750	436	22	19	138	2	163	71	21	44	35	94

(a) Multiply by 0.3048 to convert to meters.(b) Analysis by ICP-OES(c) Analysis by ICP-MS

Sample ID	Depth ^(a)	Ga ^(b)	Ge ^(b)	As ^(b)	Rb ^(b)	Sr ^(b)	Y ^(b)	Zr ^(b)	Nb ^(b)	Sn ^(b)	Sb ^(b)	Cs ^(b)	Ba ^(b)
	(ft)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)
		-	Hanfor	d Forma	tion H1	Unit		_		-	-		
C3391-15.5	16	18	1.4	-5	56	403	23.7	149	9.7	2	0.3	1.7	726
C3391-17.5	18	19	1.6	-5	56	405	24.9	145	10.9	11	0.3	1.7	747
C3391-5105	51.55	20	1.6	-5	63	345	29.0	185	11.6	2	0.3	2.0	693
			Hanfor	d Forma	tion H2	Unit							
C3391-77.3	77.8	17	1.6	-5	75	336	19.3	132	8.8	1	0.5	2.5	759
C3391-83.3	83.8	17	1.4	-5	77	364	19.3	133	8.4	5	0.6	2.3	801
C3391-90.75	91.25	16	1.5	-5	85	348	18.7	126	8.9	2	1.5	2.8	760
C3391-107.3	107.8	16	1.6	5	81	373	19.5	123	8.9	1	0.5	2.6	833
C3391-115.4	115.9	19	1.7	-5	71	387	22.3	134	9.5	2	0.4	2.1	803
C3391-133	133.5	16	1.4	5	79	382	17.6	113	7.6	1	0.5	2.7	828
C3391-160.3	160.8	16	1.4	-5	73	394	17.4	109	8.0	5	0.5	2.1	806
C3391-161.35	161.85	16	1.4	-5	71	394	17.6	113	8.2	2	0.4	2.1	831
C3391-171.45	171.95	17	1.6	5	73	412	20.8	169	10.5	1	0.5	2.2	830
C3391-173.05	173.55	16	1.5	-5	67	405	21.8	195	10.9	3	0.4	2.1	811
			Hanfor	d Forma	tion H3	Unit			•				
C3391-198.1	198.6	18	1.6	-5	64	379	22.0	126	9.0	6	0.4	1.7	792
C3391-200.1	200.6	18	1.6	-5	62	388	21.6	125	9.0	2	0.3	1.7	800
C3391-200.1 Rep	200.6	18	1.7	-5	66	395	21.7	124	9.1	2	0.4	1.8	797
		ŀ	Plio-Pleis	stocene S	ilty Unit	(PPlz)							
C3391-213.8	214.3	16	1.3	-5	70	454	16.5	135	8.8	8	0.4	2.1	842
C3391-218.7	219.2	19	1.6	9	84	348	25.3	206	13.3	3	1.0	3.9	820
C3391-219.7 Below sand	220.2	21	1.7	8	101	344	31.3	254	16.2	3	0.8	5.0	809
C3391-219.7 Above sand	220.2	23	1.9	18	110	303	31.9	239	18.6	3	1.2	5.4	1,020

Table 4.9c. Element Constituents from Soil Fusion Method.

Table	4.9c.	(contd)
		· /

Sample ID	Depth ^(a) (ft)	Ga ^(b) (µg/g)	Ge ^(b) (µg/g)	As ^(b) (µg/g)	Rb ^(b) (µg/g)	Sr ^(b) (µg/g)	Y ^(b) (µg/g)	Zr ^(b) (µg/g)	Nb ^(b) (μg/g)	Sn ^(c) (µg/g)	Sb ^(b) (µg/g)	Cs ^(b) (µg/g)	Ba ^(b) (µg/g)
C3391-220.65	221.15	22	1.5	10	101	287	27.2	213	17.5	5	0.9	4.6	1,010
C3391-220.65 /R	221.15	20	1.7	8	84	357	29.0	242	14.9	2	0.9	3.9	827
C3391-221.65	222.15	20	1.6	8	82	361	28.7	244	14.8	2	0.9	4.1	925
		Plie	o-Pleisto	cene Gra	wely Un	it (PPlg)							
C3391-239.8	240.3	20	1.5	-5	55	429	23.5	133	9.8	3	0.4	1.5	745
 (a) Multiply by 0.3048 to converse (b) Analysis by ICP-MS (-) indicate value below limit of 	ert to meters. f quantification												
Samula ID	Depth ^(a)	La ^(b)	Ce ^{(b}	Pr ^(b)	Nd ^(b)	Sm ^(b)	Eu ^(b)	Gd ^(b)	Tb ^(b)	Dy ^(b)	Ho ^(b)	Er ^(b)	
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Sample ID	(ft)	μg/g	μg/g	μg/g	µg/g	μg/g	μg/g	µg/g	µg/g	μg/g	μg/g	μg/g	
		Hai	nford For	mation H	II Unit								
C3391-15.5	16	29.1	58.5	6.70	26.4	5.62	1.66	5.15	0.81	4.65	0.89	2.59	
C3391-17.5	18	26.5	54.5	6.42	25.8	5.72	1.66	5.12	0.85	4.71	0.93	2.76	
C3391-5105	51.55	31.6	66.9	7.69	31.4	6.78	1.95	6.10	0.98	5.53	1.07	3.14	
		Hai	nford For	mation H	12 Unit								
C3391-77.3	77.8	26.0	52.2	5.95	23.3	4.84	1.37	4.12	0.67	3.74	0.72	2.18	
C3391-83.3	83.8	24.3	48.7	5.63	22.3	4.83	1.35	4.08	0.65	3.70	0.71	2.14	
C3391-90.75	91.25	29.4	60.8	6.72	25.7	5.11	1.29	4.11	0.65	3.72	0.71	2.16	
C3391-107.3	107.8	29.7	59.9	6.59	25.1	5.02	1.35	4.08	0.66	3.76	0.72	2.16	
C3391-115.4	115.9	28.8	59.0	6.70	26.4	5.47	1.54	4.64	0.77	4.29	0.83	2.41	
C3391-133	133.5	25.4	51.1	5.78	22.1	4.48	1.24	3.77	0.60	3.36	0.65	1.93	
C3391-160.3	160.8	24.1	48.9	5.54	21.9	4.41	1.25	3.77	0.60	3.43	0.68	1.98	
C3391-161.35	161.85	24.7	48.9	5.49	21.3	4.27	1.24	3.53	0.59	3.31	0.65	1.90	
C3391-171.45	171.95	32.2	64.2	7.26	27.9	5.53	1.42	4.62	0.70	3.96	0.76	2.30	
C3391-173.05	173.55	30.8	62.9	7.13	27.8	5.57	1.50	4.65	0.72	4.14	0.78	2.40	
		Hai	nford For	mation H	13 Unit								
C3391-198.1	198.6	24.6	49.3	5.77	23.5	4.92	1.47	4.39	0.71	4.08	0.80	2.34	
C3391-200.1	200.6	24.4	49.5	5.80	23.2	5.01	1.47	4.44	0.73	4.18	0.80	2.40	
C3391-200.1 Rep	200.6	24.6	51.1	6.03	23.9	5.17	1.50	4.58	0.72	4.16	0.82	2.42	
		Plio-P	Pleistocen	ne Silty Un	nit (PPlz	z)							
C3391-213.8	214.3	25.3	50.8	5.68	22.1	4.38	1.19	3.37	0.53	3.02	0.58	1.77	
C3391-218.7	219.2	40.0	80.5	9.23	35.4	6.86	1.63	5.48	0.85	4.78	0.92	2.75	
C3391-219.7 Below sand	220.2	47.0	95.6	10.8	41.0	7.93	1.83	6.42	0.99	5.60	1.09	3.28	
C3391-219.7 Above sand	220.2	45.3	93.2	10.5	39.9	7.98	1.82	6.38	0.99	5.65	1.10	3.29	

Table 4-9d. Element Constituents from Soil Fusion Method.

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Table 4-9d.	(contd)
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Samula ID	Depth ^(a)	La ^(b)	Ce ^{(b}	Pr ^(b)	Nd ^(b)	Sm ^(b)	Eu ^(b)	Gd ^(b)	Tb ^(b)	Dy ^(b)	Ho ^(b)	Er ^(b)
Sample ID	(ft)	μg/g	μg/g	μg/g	μg/g	μg/g	μg/g	µg/g	μg/g	µg/g	μg/g	μg/g
C3391-220.65		221.15	39.5	81.2	9.03	34.6	6.81	1.63	5.610.88	4.77	0.92	2.76
C3391-220.65 /R		221.15	41.0	83.2	9.36	36.0	7.20	1.64	5.820.90	5.09	0.99	2.97
C3391-221.65		222.15	41.5	77.8	9.40	34.9	6.96	1.54	6.02 0.92	5.07	0.98	2.98
		Plio-Pl	eistocene	e Gravely	Unit (Pl	Plg)						
C3391-239.8		240.3	21.4	44.4	5.35	22.3	4.97	1.49	4.65 0.75	4.25	0.84	2.45
(a) Multiply by 0.3048 to convert	to meters.											
(b) Analysis by ICP-MS(-) indicate value below limit of a	mantification											

Comula ID	Depth ^(a)	Tm ^(b)	Yb ^(b)	Lu ^(b)	Hf ^(b)	Ta ^(b)	W ^(b)	Tl ^(b)	Pb ^(b)	Bi ^(b)	Th ^(b)	U ^(b)
Sample ID	(ft)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)
		На	nford Fo	rmation H	HI Unit							
C3391-15.5	16	0.391	2.46	0.352	4.2	0.64	0.6	0.29	-5	-0.1	4.35	1.09
C3391-17.5	18	0.411	2.55	0.364	4.2	0.64	0.6	0.42	8	0.1	3.96	1.05
C3391-5105	51.55	0.457	2.87	0.417	5.3	0.68	0.7	0.74	7	0.1	5.23	1.36
		Ha	nford Fo	rmation H	H2 Unit							
C3391-77.3	77.8	0.315	2.02	0.291	3.9	0.54	0.8	0.61	9	0.2	4.69	1.21
C3391-83.3	83.8	0.327	2.08	0.292	3.9	0.52	0.7	0.48	8	0.1	4.48	1.18
C3391-90.75	91.25	0.329	2.09	0.293	3.8	0.56	1.0	0.53	11	0.3	6.12	1.45
C3391-107.3	107.8	0.318	2.04	0.288	3.5	0.57	0.9	0.59	10	0.1	5.53	1.34
C3391-115.4	115.9	0.365	2.33	0.327	3.8	0.58	0.7	0.54	9	0.2	5.15	1.23
C3391-133	133.5	0.286	1.79	0.260	3.3	0.48	0.9	0.45	6	0.1	4.71	1.18
C3391-160.3	160.8	0.291	1.81	0.266	3.2	0.49	0.7	0.49	7	0.1	4.23	1.09
C3391-161.35	161.85	0.286	1.81	0.259	3.3	0.50	0.7	0.50	7	0.1	4.45	1.21
C3391-171.45	171.95	0.347	2.20	0.323	4.9	0.73	1.0	0.56	10	0.2	5.91	1.54
C3391-173.05	173.55	0.362	2.34	0.341	5.8	0.75	1.2	0.45	7	0.1	5.67	1.58
		Ha	nford Fo	rmation H	H3 Unit							
C3391-198.1	198.6	0.343	2.16	0.313	3.6	0.52	0.6	0.44	7	0.1	4.39	1.16
C3391-200.1	200.6	0.363	2.18	0.316	3.6	0.55	1.4	0.48	8	0.1	4.34	1.32
C3391-200.1 Rep	200.6	0.358	2.23	0.327	3.7	0.60	0.6	0.52	9	0.2	4.45	1.17
		Plio-	Pleistocer	ne Silty U	nit (PPlz	z)	•		•	•		
C3391-213.8	214.3	0.262	1.65	0.249	3.8	0.51	0.9	0.43	7	0.1	4.72	1.30
C3391-218.7	219.2	0.410	2.56	0.373	5.9	0.84	1.4	0.67	12	0.2	8.16	2.19
C3391-219.7 Below sand	220.2	0.516	3.18	0.464	7.6	1.04	1.8	0.77	15	0.5	10.6	2.81
C3391-219.7 Above sand	220.2	0.504	3.14	0.462	7.2	1.23	1.7	0.97	20	0.7	11.1	2.97

Table 4-9e. Element Constituents from Soil Fusion Method.

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Table 4	I-9e. ((contd)

Sample ID	Depth ^(a)	Tm ^(b)	Yb ^(b)	Lu ^(b)	Hf ^(b)	Ta ^(b)	W ^(b)	Tl ^(b)	Pb ^(b)	Bi ^(b)	Th ^(b)	U ^(b)
r i i i i i i i i i i i i i i i i i i i	(ft)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)	(µg/g)
C3391-220.65	221.15	0.427	2.67	0.392	6.3	1.07	1.6	0.66	12	0.3	10.1	2.63
C3391-220.65 /R	221.15	0.463	2.90	0.421	7.2	0.85	1.7	0.67	14	0.3	9.80	2.59
C3391-221.65	222.15	0.445	2.81	0.413	7.4	1.13	1.4	0.67	14	0.3	10.1	2.77
		Plio-Pl	eistocene	Gravely	Unit (PF	Plg)						
C3391-239.8	240.3	0.371	2.31	0.339	3.9	0.50	0.5	0.42	7	0.1	4.62	1.27

(a) Multiply by 0.3048 to convert to meters.
(b) Analysis by ICP-MS
(-) indicate value below limit of quantification

4.5 Calcium Carbonate and Organic Carbon Content

Carbon results for borehole 299-E33-338 are shown in Table 4-10 and the calculated calcium carbonate equivalents (in terms of grams of calcite per gram of oven-dry sediment) are also presented. The overall results for calculated calcium carbonate concentration is in the range of 0.5 to 2.0 wt %, with the highest reported values associated with PPlz unit and the lowest values associated with the H3 unit. These values are generally consistent with results from similar lithological samples reported elsewhere (e.g. Serne et. al. 2002).

It should be noted that this method measures the organic carbon by subtracting the inorganic carbon from the total carbon in the sample. For such low carbon values, this method is not very accurate. However, the low values for organic carbon are within the ranges generally reported for sediment at the Hanford site (e.g. Serne et. al. 2002).

Sample ID	Depth ^(a) ft bgs	Total Carbon (%)	Inorganic Carbon (%)	IC as CaCO3 (%)	Organic Carbon (by difference)
		Hanford Formation	on H1 Unit		
C3391_15.5	16.0	0.19	0.14	1.20	0.04
C3391_17.5	18.0	0.15	0.12	0.98	0.04
C3391_51.05	51.6	0.19	0.14	1.21	0.04
		Hanford Formation	on H2 Unit		
C3391_77.3	77.8	0.18	0.17	1.43	0.01
C3391_83.3	83.8	0.16	0.13	1.07	0.03
C3391_90.75	91.3	0.19	0.16	1.33	0.03
C3391_107.3	107.8	0.20	0.16	1.36	0.03
C3391_115.4	115.9	0.14	0.12	1.02	0.02
C3391_133	133.5	0.15	0.12	1.02	0.03
C3391_160.3	160.8	0.25	0.11	0.91	0.14
C3391_161.35	161.9	0.26	0.14	1.17	0.12
C3391_171.45	172.0	0.22	0.16	1.35	0.06
C3391_173.05	173.6	0.29	0.20	1.65	0.09
		Hanford Formation	on H3 Unit		
C3391_198.1	198.6	0.15	0.08	0.66	0.07
C3391_200.1	200.6	0.13	0.08	0.67	0.05
	Pl	io-Pleistocene Silt	ty Unit (PPlz)		
C3391_213.8	214.3	0.17	0.13	1.10	0.03
C3391_218.7	219.2	0.28	0.18	1.47	0.11
C3391_219_Below	220.2	0.28	0.22	1.84	0.05

 Table 4.10.
 Calcium Carbonate and Organic Carbon Content.

Sample ID	Depth ^(a) ft bgs	Total Carbon (%)	Inorganic Carbon (%)	IC as CaCO3 (%)	Organic Carbon (by difference)
C3391_219_Above	220.2	0.30	0.24	2.00	0.06
C3391_220.65	221.2	0.30	0.21	1.79	0.08
C3391_220.65 Dup	221.2	0.28	0.20	1.69	0.08
C3391_221.65	222.2	0.27	0.22	1.79	0.06
		Plio-Pleistoce	ene Gravely Unit (PPlg))	
C3391_239.8	240.3	0.11	0.09	0.72	0.02

Table 4-10. (contd)

(a) Multiply by 0.3048 to convert to meters.

4.6 Mineralogy

XRD analysis was conducted on sediment collected from borehole 299-E33-338, which was represented by two stratigraphic units, the Hanford formation and the Plio-Pleistocene unit. The Hanford Formation is divided into three units, with the shallowest being identified as H1, followed by the H2, and the deepest being H3. Samples examined from the H1 facies (4.7 to 15.6 m [15.5 to 51.1 ft]) were dominated by quartz with lesser amounts of feldspar (Table 4.11). Mica and chlorite appeared as trace amounts, with the relative concentrations of both clay minerals increasing with depth. Hornblende also appears in detectable amounts, with concentrations following the mica and chlorite abundance profile. Samples collected from the H2 (23.6 to 52.3 m [77.3 ft to 171.5 ft]) appeared to be similar to the H1 unit, with perhaps more of a weathered profile. Clay minerals such as micas and chlorites are more abundant in this facies and are easily detected by XRD. Samples examined from the H3 unit also exhibited similar characteristics as the two earlier units (i.e., H1 and H2).

Sediment from the Plio-Pleistocene unit contain quartz and feldspars, along with significant amounts of clay material, predominantly mica and chlorite. Additionally, these samples contain hornblende in minor to trace concentrations, along with trace amounts of laumontite. For example, the XRD tracing of a typical sediment sample at 60.4 m (198.1 ft) (Plio-Pleistocene silty unit) is provided in Figure 4-15, along with quartz and plagioclase reference patterns. The main reflection for quartz is 26.63° 20, followed by less intense reflections at 20.86, 36.53, 39.46, 42.43, 50.12, 59.92° 20. The primary reflections associated with feldspar minerals are found between 27.34° 20 and 27.92° 20, with the higher 20 values belonging to the plagioclase series. Chlorite and mica minerals were identified on the x-ray tracings by the reflections at 6.3° 20 and 8.8° 20, respectively. The presence of hornblende was established by the characteristic 100% reflection at 10.5° 20. Additionally, trace amounts of the zeolite, laumontite, were identified in most of the samples by a diffraction peak positioned at 9.36° 20.

Results from the semi quantification of the minerals in the bulk samples are provided in Table 4.11. Quartz concentrations ranged from 22 wt% at 219.17 ft to 48.5 wt% at 51.05 ft. The borehole sediment contained plagioclase feldspar concentrations from 10.6 to 39.5 wt% and potassium feldspar content measured between 9.0 to 37.9 wt%. Plagioclase feldspar was more abundant than potassium feldspar

with the exception of the three samples in the H1 Hanford coarse sand and one sample in the lower PPlz unit. The amphibole phase comprised <10 wt% at most, with the majority of samples having concentrations in the 2 to 5 wt% range.

Clay minerals identified in the bulk sediment included mica and chlorite. Mica concentrations ranged from trace amounts in the Hanford formation H1 unit to as high as high as 26% in the PPIg unit. Chlorite concentrations were <5-wt% in all sediments analyzed with only trace amounts detected in the H1 unit. Smectite and kaolinite minerals were not identified in the bulk sediment samples due in part to the sample preparation technique and the low overall concentration, respectively.



Figure 4.14. XRD Tracing of Sediment Collected from 198.1 ft, Along With the Standard Reference Patterns for Quartz and Plagioclase.

Course La D			Mineral Phase	(wt%)			Goodness
Sample ID	Quartz	Amphibole	Plagioclase	K-Spar	Mica	Chlorite	of fit ^(a)
		Hanj	ford Formation H	HI Unit			
C3391-15.5	30.7	10	21.0	37.9	Tr	Tr	0.39
C3391-17.5	31.6	3.0	26.1	39.3	Tr	Tr	0.29
C3391-51.05	48.5	5.3	17.9	28.4	Tr	Tr	0.29
		Hanj	ford Formation H	H2 Unit			
C3391-77.3	44.0	3.8	23.7	28.5	Tr	Tr	0.48
C3391-83.3	30.3	3.3	31.0	15.7	18.0	2.3	0.23
C3391-90.75	38.1	2.6	22.3	15.2	19.7	2.1	0.49
C3391-107.3	37.8	3.0	26.3	12.8	17.3	2.7	0.41
C3391-115.4	28.0	4.9	32.0	10.9	21.5	2.8	0.26
C3391-133	35.5	3.5	26.3	11.9	20.1	2.7	0.46
C3391-160.3	32.0	4.7	27.1	12.3	20.7	3.1	0.35
C3391-161.35	38.4	2.8	27.0	12.0	16.9	2.6	0.50
C3391-171.45	29.0	7.8	29.9	9.0	21.2	3.2	0.24
C3391-173.05	ND	ND	ND	ND	ND	ND	ND
		H3 F	Hanford Formation	on Unit			
C3391-198.1	24.8	3.3	39.5	11.2	18.0	3.2	0.26
C3391-200.1	29.8	4.1	31.7	13.1	18.1	2.8	0.40
		Plio-Pl	eistocene Silty U	nit (PPlz)			
C3391-213.8	31.5	7.3	30.8	10.7	17.1	2.6	0.23
C3391-218.7	30.3	7.2	25.2	12.1	21.7	3.4	0.30
C3391-219.7 below sand	ND	ND	ND	ND	ND	ND	ND
C3391-219.7 above sand	30.8	4.7	16.9	17.4	25.6	4.6	0.45
		Plio-Pleis	stocene Gravelly	Unit (PPlg)			
C3391-220.65	31.8	6.0	27.3	13.6	17.9	3.4	0.44
C3391-221.65	38.1	5.8	18.9	15.8	26.3	5.0	0.38
C3391-239.8	27.4	3.3	33.8	9.3	23.3	2.9	0.32

Table 4.11. Semi-quantitative XRD Results of Minerals from Borehole 299-E33-338.

(a) Values closest to 1.0 represent an ideal refinement.

ND-not determined

X-ray diffraction analysis was performed on the <2 micron (clay) fraction of each sample and the results are presented below. The clay size fraction is dominated by four clay minerals: smectite, chlorite, illite, and kaolinite with minor amounts of quartz and feldspar. Figure 4.16 provides XRD-tracings of a

typical clay fraction (from sample 60.4 m [198.1 ft]) following a treatment with Mg-saturation and solvation with ethylene glycol. Smectites are considered the fraction of the Mg-saturated sub-sample that gives a basal reflection at $5.28^{\circ} 2\theta$ upon solvation with ethylene glycol. Illite is the simplest of the four clay mineral phases to identify in this sediment. The basal reflections are located at 8.88, 17.8, and 26.7° 2 θ . The various treatments including cation saturation, solvation with ethylene glycol, and heating do not affect the structure of the illite.

Chlorites are identified by their basal series of diffraction peaks at 6.24, 12.5, 18.8, and 25.2° 2 θ , which are unaffected by cation saturation or ethylene glycol solvation. Kaolinite is difficult to identify in the presence of a chlorite mineral. Basal reflections characteristic of kaolinite are positioned at 12.5 and 24.9° 2 θ , which are super imposed on the even-order chlorite peaks. These kaolinite reflections are unaffected by cation saturation and ethylene glycol solvation. Positive identification of kaolinite in the presence of chlorite can be determined by examination of the 24.9 to 25.2° 2 θ region of the XRD tracing. The kaolinite basal reflection at 24.9° 2 θ can be distinguished from the chlorite 25.2° 2 θ reflection in some situations. For example, XRD tracings of the clay fractions from depths 219.7A and 219.7B both show a bi-module peak from 24.9° 2 θ to 25.2° 2 θ .

Trace amounts of quartz are evident by the diffraction peak located at $20.85^{\circ} 2\theta$. The 100% reflection for quartz (26.6° 2 θ) is hidden by the third basal reflection of illite located at 26.6° 2 θ . Plagioclase feldspar is also identified in the clay fraction by the minor diffraction peak at 27.8° 2 θ as well as hornblende by the minor peak at 10.41° 2 θ . Additionally, laumonite was detected in the clay fraction from several depths in the lower Plio-Pleistocene formation (67.3 and 67.6 m [220.65 and 221.65 ft]), evident by the reflection at 9.36° 2 θ .

Semi-quantification results of the clay minerals in the < 2 micron fraction are presented in Table 4.12. Total recoveries were normalized to 100% and the normalization factor used for each sample is provided in the last column. Smectites ranged in concentrations from a low of 3 wt% (83.3 ft) to a high of 60 wt% (4.7 m [15.5 ft]). Illite amounts varied from 28 to 61 wt% with the majority of samples having concentrations in the 40 to 55 wt% range. Chlorite and kaolinite were the least abundant of the clay minerals identified in the samples with concentrations equal to or less than 25 wt% and 14 wt%, respectively. Quartz and feldspar minerals were present as trace amounts in the clay fraction and therefore were not included in totals presented in Table 4.12.



Figure 4.15. XRD Tracings of Preferentially Oriented Clay Slide Taken from Borehole 299-E33-338 at a Depth of 198.1 ft. (The scans were collected from 2 to 45° 2θ with a 0.04° step and 2-second dwell time. The black line represents the Mg-saturated fraction.)

Total clay recoveries were within $\pm 25\%$ of the "ideal" 100% for four of the 22 samples analyzed. The majority of clay recoveries were between 25% and 50%, with only one sample having a normalization factor of 1.96. This sample, 32.7 m (107.3 ft), cracked significantly during the analysis, which resulted in a poorly oriented sediment. Other factors affecting the semi-quantification procedure (i.e., preparation and condition of the clay filter cake) were generally controlled and not thought to be a significant factor. Quantitative analysis is considered good if errors amount to $\pm 10\%$ of the amounts present for major constituents and $\pm 20\%$ for minerals whose concentrations are less than 20% (Moore and Reynolds 1989).

	Depth ^(a)		Minera	l Phase (wt%)		Normalization
	(ft)	Smectite	Illite	Chlorite	Kaolinite	Factor
	15.5	60	28	7	5	0.73
H1	17.5	53	31	10	6	0.92
	51.05	25	52	13	10	0.49
	77.3	31	44	15	9	0.64
	83.3	3	61	22	14	0.81
	90.75	10	61	19	9	0.49
	107.3	22	51	14	13	1.96
цэ	115.4	18	53	21	8	0.61
П2	133	23	53	18	6	0.51
	160.3	25	48	19	8	0.55
	161.35	19	54	20	7	0.56
	171.45	27	42	23	8	0.50
	173.05	28	37	25	10	0.54
	198.1	35	42	18	5	0.79
	200.1	18	53	21	8	0.61
	213.8	23	53	18	6	0.51
	218.7	25	48	19	8	0.55
PPIZ	219.7 below	19	54	20	7	0.56
	219.7 above	25	52	13	10	0.49
	220.65	31	44	15	9	0.64
	221.65	35	42	18	5	0.79
PPlg	239.8	31	44	19	6	0.74

 Table 4.12. Semi-quantitative XRD Results of Clay Minerals Separated from the Sediment

 Collected from Borehole 299-E33-338.

(a) Multiply by 0.3048 to convert to meters.

4.7 Chemical Interactions

There were no contaminant adsorption/desorption or leach tests performed with the clean sediments. The chemical and mineralogical data from the uncontaminated sediments from borehole 299-E33-338 can be used to compare with the sediments from the contaminated boreholes.

4.8 Sediment Matric Potential at Borehole 299-E33-338

The matric potential of the sediment profile in borehole 299-E33-338 was measured and is plotted in Figure 4.17. Borehole 299-E33-338 (C3391) is located outside the southeast corner of the B tank farm in a relatively undisturbed area. Results from these measurements indicate that wetting from meteoric sources has not reached the water table at the 299-E33-338 site.



299-E33-338 (SE of B Tank Farm)

Figure 4.16. Matric Potential of the Sediment Profile at Borehole 299-E33-338.

5.0 Summary and Conclusions

Sediment cores from borehole 299-E33-338 near the B-BX-BY WMA have been characterized to establish lithological and geochemical baseline profiles that can be used to support ongoing field investigations concerned with past single shell tank fluid leak events in the vadose zone being conducted by CH2M Hill Hanford Group Inc. The base line results in this report will serve as a "clean sediment" standard for comparison with sediment cores extracted from areas of known contamination within the B-BX-BY WMA. These results will not only help support our understanding of the degree and extent of contamination within the B-BX-BY WMA, but also possible mechanisms associated with fate and transport and risk based assessments of problematic contaminants such as technetium-99 and uranium.

The geology under the B-BX-BY WMA forms the framework through which the contaminants move, and as discussed in Serne et. al. 2002, provides the basis with which to interpret and extrapolate the physical and geochemical properties that control the migration and distribution of contaminants. Specifically, the identification of major lithological contacts and the interrelationships between the coarser- and finer-grained sediment facies are essential when combined with the geochemical profile for interpreting contaminant behavior in the subsurface. For this borehole, lithologic sections were constructed using detailed geologic descriptions, core photos, and geophysical logs. In some cases, the results of laboratory analyses (e.g. particle-size distribution, moisture, calcium carbonate content) helped refine the resulting stratigraphic and lithological interpretations.

Our conceptual model of the vadose zone associated with the 299 E33-338 borehole involves five distinct stratigraphic units beginning with the Hanford formation H1 unit from the surface to a depth of approximately 15.7 m (approximately 51.5-ft) described as a sandy gravel to gravelly sand sequence. This is followed by the Hanford formation H2 unit extending to a depth of approximately 57.9 m (109 ft) that is a sand sequence consisting of sand dominated facies, with multiple graded beds of horizontal to tabular cross-bedd sand to slightly gravelly sand. These graded beds are sometimes capped with thin layers of silty sand to silt. The last unit associated with the cataclysmic flood deposits is the Hanford H3 formation unit that extends to a depth of approximately 64.8 m (212.5 ft). It is a gravelly sand to slightly gravelly sand sequence. Just below the H3 unit is the Plio-Pleistocene silty unit (PPlz) extending to a depth of approximately 67.8 m (222.4 ft). This unit is a silt-dominated sequence consisting of interstratified well sorted silt and fine sand. The last unit characterized from this borehole was the Plio-Pleistocene gravelly unit (PPlg) extending to a depth of approximately 82.6 m (271 ft), however sampling ended at approximately 73.1 m (239.8 ft). This unit is differentiated from the PPlz due to its sandy gravel to gravelly sand sequence consisting predominantly of unconsolidated basalt-rich sand and gravel.

Sediment samples from the various stratigraphic unites were analyzed and characterized in the laboratory for the following parameters:

- Mass Water Content
- Soil Suction
- Particle-Size Distribution

- Calcium Carbonate and Organic Carbon Contents
- Bulk Chemical Composition
- Mineralogy
- Water Leach (1:1 sediment to water extraction)
- Acid Leach (8M nitric acid extraction)

Physical properties, such as particle size distribution and water content varied according to lithology as expected. In general, elevated areas of water content ($\sim >5\%$) were typically associated with regions of fine grain sediments. Most notable are those regions involving lithological facies at which water contents equal or exceed 10%. Three major peaks are noted at 15.7, 52.9, and 67.1 m (51.6, 173.6, and 220.2 ft) bgs with water contents of 12.95, 14.27, and 26.02% respectively. Along with water content, soil suction measurements were made on most of the core liner and grab samples from the borehole using the filter paper method. Three major peaks were noted approximately 14, 64, and 73 m (45, 210, and 240 ft) bgs with suction measurements of approximately 1.3, 1.5, and 2.2 Mpa. The matric potential profile indicates that wetting from meteoric water has not reached the water table.

The semi-quantitative mineral composition for the bulk sediments characterized in the H1 unit and the first samples associated with the H2 unit (23.7 m [77.8 ft] bgs) consisted mostly of quartz (30.7 to 31.6 wt%), plagioclase (21.0 to 26.1 wt%) , potassium feldspars (28.4 to 39.3 wt%), amphibole (3.0 to 10.0 wt%), with trace amounts of mica and chlorite. The remaining H2, H3, PPlz, and PPlg units (below 25.5 m [83.8 ft] bgs down to 73.2 [240.3 ft] bgs) were similar consisting of quartz (24.8 to 38.8 wt%), plagioclase (17.9 to 33.8 wt%), amphibole (2.8 to 7.8 wt%), potassium feldspars (9.0 to 15.8 wt%), mica (17.1 to 25.6 wt%), and chlorite (2.1 to 4.6 wt%). The particle density for the sediments ranged from 2.57 to 2.79 g/cm³.

Inorganic carbon results reported in terms of calcium carbonate were found to be within the range of 0.5 to 2.0 wt %, and are consistent with results reported elsewhere (e.g. Serne et. al. 2002). The method used to measure the organic carbon relies on subtracting the inorganic carbon from the total carbon in the sample; for such low carbon values this method is not very accurate. The low values for organic carbon (0.01 to 0.14 %) are within the ranges generally reported for sediment at the Hanford site.

Bulk sediment samples were characterized for major and trace elements using a lithium metaborate/tetraborate fusion procedure, and then analyzed by inductively coupled plasma –optical emission spectroscopy (ICP-OES) and ICP-MS methods. Overall results showed very little difference in the primary elemental oxide concentrations for any of the sediment samples as a function of depth or lithology. The primary elemental oxides in decreasing concentration include SiO₂ (58.77 to 70.33 wt%), Al₂O₃ (12.7 to 15.73 wt%), Fe₂O₃ (3.74 to 7.92 wt%), CaO (3.02 to 4.80 wt%), Na₂O (2.05 to 3.23 wt%), K₂O (1.70 to 2.65 wt%), MgO (1.64 to 2.73 wt%), TiO₂ (0.51 to 1.39 wt%), P₂O₅ (0.13 to 0.30 wt%), and MnO (0.07 to 0.12 wt%). The additional 35 trace elements reported for each sample in terms of μ g/g of dry sediment showed little, if any, significant variation in concentration.

The water chemistry analysis for samples collected between 5 and 73 m (16 and 240 ft) bgs using the 1:1 soil to water extract method shows no strong trends as a function of depth and there is little, if any, indication of tank waste interaction with vadose zone soils at this location. Primary characteristics include the following:

- The 1:1 sediment-to-water extract pH varied from 6.97 to 7.74 and in general increased with depth with an average value of 7.4 (Figure 4.9 and Table 4.3).
- There were small increases in pH at the contact between the Hanford H2 and H3 units and the top and bottom of the Plio-Pleistocene mud unit.
- Porewater EC (dilution corrected) varied from 0.88 to 4.3 mS/cm with an average of 2.4 mS/cm.
- There were high EC values deep in the Hanford H2 unit at approximately 49 m (160 ft) bgs and in the deepest sample characterized (i.e., in the PPlg).

The shapes of the major cation profiles (sodium, potassium, calcium, magnesium, and strontium) in terms of calculated porewater concentration versus depth are very similar with slight peaks in the deep portion of the H2 unit at approximately 49 m (160 ft) bgs, at the top of the Plio-Pleistocene silty unit, and in the deepest sample characterized in the PPlg unit. All three of these samples had very low water contents and thus the dilution factor was high. The apparent high porewater concentrations likely represent some dissolution of salts from the sediment that are multiplied by a large dilution factor, and thus suggest more saline porewater than surrounding sediments with higher water content. In general, the calculated porewater cation concentrations ranged from 63 to 275, 11 to 138, 11 to 56, and 70 to 558 mg/L for calcium, potassium, magnesium, and sodium, respectively. The averages and median values were (142, 149), (60, 58), (34, 38), and (190, 141) mg/L, for calcium, potassium, magnesium, and sodium, respectively. These values are likely somewhat artificially elevated because of the water extraction of soluble salts along with the porewater.

The calculated porewater concentrations for aluminum, barium, iron, silicon, and uranium-238 were also evaluated. The porewater concentrations ranged from 0.01 to 5.29, 0.0 to 6.4, and 46 to 350 mg/L for aluminum, barium, and iron, respectively, and 1.8 to 24 μ g/L for uranium. The uncontaminated uranium-238 porewater concentration is especially important for comparison with contaminated borehole sediment porewaters.

The shapes of the major anion profiles (fluoride, chloride, nitrate, bicarbonate, phosphate, and sulfate) in terms of calculated porewater concentration versus depth showed no consistent depths where all anions peaked unlike the major cation profiles. The wetter samples do consistently show low calculated porewater anion concentrations suggesting that the dilution factor is again controlling the apparent concentrations. That is, all the sediments likely dissolve some salts that are not truly in the porewater, so that the dilution correction makes it appear that the porewater anion concentrations are higher in the drier sediments. The calculated porewater anion concentrations range from 0.4 to 23.3, 1.8 to 223, 1.3 to 100, 296 to 1877, and 196 to 117 mg/L for fluoride, chloride, nitrate, bicarbonate, and sulfate, respectively. The average and median values are (6.3, 4.8), (35, 19), (19, 10), (993, 1030), and (196, 117) mg/L for

fluoride, chloride, nitrate, bicarbonate, and sulfate, respectively. These values are likely somewhat artificially elevated because of the water extraction of soluble salts.

The mass of several constituents per gram of dry sediment that were leached by water and acid extracts are shown in Figures 4.13 and 4.14. In all cases, the mass that was water leachable is a very small fraction of the mass that was acid extractable. These concentrations can be compared with the same constituents for contaminated sediments to get an estimate of the mass of a constituent present in the vadose zone profile after contact with tank leaked liquids.

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Appendix A

Summary of Field Geologists' Sample Descriptions for Borehole 299-E33-338

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			B	OREHOLE LOG		0540309	Page <u>f</u> of <u>19</u> Date: 07/23/0/
Well 1D:	Ca	391	Well N	ame: 299-633-338	L.	ocation: JF from SE (or	mer 241-B Trakfun
Project:	RCR	A 2001	Drill	(ng	F	Reference Measuring Point:	Granderichare
	Sa	mple		Sample (Descri	ption	Comments:
Depth <u>(Ft.)</u>	Type No.	Blows Recovery	Graphic Log	Group Name, Grain Size Distri Moisture Content, Sorting, Ang Size, Rea	ribution gularity action t	, Soil Classification, Color, r, Mineralogy, Max Particle o HCl	Depth of Casing, Drilling Method, Method of Driving Sampling Tool, Sampler Size, Water Level
0-	al) - T		The Caller	0-7' Silty say	ndy	Gravel (#SG)	cable tool
-	STOON	[DO	G= 65% (50% cr.	407	om, 1070F), 2590	cont. solit spoon
1 1-	1	100%	000	sand (20% cr 60%	75 m,	2090 f) and 10%	
2 -	1540 hp	recov.	ÖzÖ Ölk	silt in poorly sort	ted, s	dry to slightly d unit	moisture sample @2
<u> </u>	SS		\tilde{a}	, , , , , , , , , , , , , , , , , , , ,			
3-	1	44%	<u>۽ ج</u>				
4-	1600	recov.	93509 37-0				moisture sample Q4
	55 B		2100 7019	5 silty sady grave	1 (m	sg) some as	Grab Archive. Q.5.0
- -	0815	Her.	<u>8</u> 0	med cobble	<u>цсі ·</u>	and max si se	maisture sample D.S.76
	り		Time /	7 - 4-50 Si Ity Sandy	Grav	el (mob) 6 65%	· · · · · · · · · · · ·
7-	54 D		nø	Sund 25 10, 5112 101	10 1	othe ble, slight color	
	Jak OJ		recovery	chy from above, 2001	ייוא	50r+e3	K, 3@ backgonund.
	55*4	-	8 70 C				Less, c.o., org c ort
9-		75%	2100 100				Grabarchive @ 10'
	ARCHINE	AZZHIUE	Гĕ С				
'`_	1311 hrs. 55 # 5	-		· · · · · · · · · · · · · · · · · · ·			1 moisture Jacks 10 (0)
- -		_	Do.				
-	-	5510	0				
12-			\odot	12.5 - 14.5' silta	9-4-	elly sand (mas)	
_	- 050 ms			55 10 sand 25 10 g	N RAVE	, 20% 5:12, sard	moisture sample D
13-	-		0.00	f- cae poorly sorted	g reat	el SR-SA, poorly so that	· 12.5'
-	- {	100%	+ 6	silt nodules,			
14-	-	rec		145 5:13 LEAS 18	80°/0	three sandy	
	14,33	<u>v</u>		f growel ,			moisture sample D 145
Reporte	d By: <u>J//</u>	Faurot	61c.	Martinez Ron	viewed	By DCUkeke	5
Title:	ellog	ist \	<u><u><u>Geolo</u>q</u></u>	ist. Title	le: G	seologist	<i>,</i>
Signatu	re: mj	wote	IC MAR	Date: 07 [23]01 Sig	gnature	NCApeller	Date: 10/16/01
BHI-EE-1	83 (12/97)						

			во	DREHOLE LOG		Page A of 19
Vell ID: (23391	-	Well N		Location: SE conner a	of 241-B Tank Farm
roject:	6401	R.C.R.A	Deilli	<u></u>	Reference Measuring Point:	6
	Sar	nole		Sample Desc	rintion	Commente:
Depth <u>(Ft.)</u>	Type No.	Blows Recovery	Graphic Log	Group Name, Grain Size Distributio Moisture Content, Sorting, Angular Size, Reactior	on, Soil Classification, Color, ity, Mineralogy, Max Particle n to HCl	Depth of Casing, Drilling Method, Method of Driving Samaling Tool, Sampler Size, Water Level
5 —		Archive	0-0-0 10-0-0	150-155 Silry Sandy G	ravel (msg) 65%	Grab Archive @ 15
- _ عا	2	100 ^{°/0}	131.151 00.11 01.101	gravel, 30% 50nd, 5%. SR-SA, Property someted, 9 sometred. 20% oft (other	silt. sand, free pravel se, poorly 1 32% poselt.	
17 —	1519	Heer.		no can Hei	· · · · ·	
1B -	55*8			15:5-165/51114 Gravelly So 13 (0 gravel, 15 (0 5:11	und. 75 4 sed, t, sand, 46-cse,	Maisture sample 17.5
- 19		100 ^{°/2}		Farly sorted, SR. grave) Silt nodules, 30 10 bas	et 20 (0 qt2.	· · · · · · · · · · · · · · · · · · ·
– ס <u>י</u>	1007	Acchire		16.5 - 20.0/Silty Sand	85° (0 5202 15° (0	matetine sample 195'
_ 고, _	35 ** q	90%	010, 1 010 100, 1 010	silt, sand uf-cse, i has alt 25°6, gtz (oth	poonly socted, se, sec) 75%, no Exn	
12 -	0651	recover	$\langle \hat{O} \rangle \hat{O}$	20.0'-25.5') 5:11 sand	y gravel (MSG)	moistur maple () 20.4
23 - -	-	40 70		Dame description as		X, B @ background_
과 니 -	- 1 	₩ ₽ ,				Maisture samale @ 74
ෂ	35 ₩11	Archive		25.5 - 275 Slightly Sill	ry Sandy Gravel. 5 10 Gravel. Sand	Grab Archive @ 25'
- 210 -	-	100%0	2000 2000 2000	m-v cse. 5-A, poorty	sorted Gravel	
ນາ - 	1001	-	0 <u>0</u> 0	30% baselt, 70% gtz	(other), 10 YR512	maisture sample @ 27.0
z6.	-	100%		225 - 281) silty Sandy gravel w to silt . 25 "10	5 Grave) (ms6) 65%. Sand. Sand Vf-cse,	
29 ·	-] - ואר			SR, poorty sorted, gravel 3	R-SA, poorly sorted, , strong mit HCL gray, 10, 104R WZ C brownist	moistur semple @ 29.6
Reporte	ed By:	harten.	e Mar	timezReview	ed By: DC. Weeke	
Title:	Geolo	<u>Jeig</u>		Title:	Geologist	
Signatu	ire: CD	None."	mastin	Date: 07 25 01 Signatu	ire: MCafeefeel	Date: 10/16/01

			в	REHOLE LO	G		Page <u>3</u> of <u>19</u>
WellID			Well N	me: > /29 - < 2 >	-		Date 07 (25101
Des is sto		<u> </u>			53%	Distant SE Corner a-	f. 241-B Tank Form
Project:	<u> </u>	RCRA	Drilli	<u>n q</u>		Reference Measuring Point	Ground Surface
	. Sa	mple		<u> </u>	mple Desc	ription	Comments:
Depth <u>(Ft.)</u>	Type No.	Blows Recovery	Graphic Log	Group Name, Grain Size Moisture Content, Sortia Siz	e Distributiong, Angular 26, Reaction	on, Soil Classification, Color, ity, Mineralogy, Max Particle n to HCl	Depth of Casing, Dritting Method, Method of Driving Sampling Tool, Sampler Size, Water Level
3 ⁻ 2		Brothing	, , , , , , , , , , , , , , , , , , ,	28'- 30 5	lightle s	situ sandu aravel	Grap Anotive @ 30.0'
_	1		() - D - O	.0% sitt 40%	sand 5	o to grevel, sand	
31 -			0	m-rose, SA, well	sorted,	gravel, sm pebbles.	
_	. 1	100%	С. , 7с	5.19. well-sorted.	traces	obbles 15% basalt.	
32 -	121300	HQC.	5	75 10 pt2 (other)	IOYR 6	Z (Iz brownish gray)	moisture sample @ 1913-
_	55 # 14	1	-	Slight Mr. HCL.			
33 -		8010	0	31-35) 5:1+4	Sand.	20 % 5:1t 20 10	
-	-			Sand, S.R. mod	Sorted .	vf-cse 10 lests	
34 -	1300	-		30% basalt, to	د. محمد وجم	uch. INYR 6/2 grayish	moture sample @ 34
-	no sec.	4		brown, an rxn	HC1-2:14	2. Moduler. micoceaus	
зб	\cdot	arch ve	Q	55-55 (S) 50m295	tu f - E	; IT s.s., mod sorted, S.R.	
-	6. * 6		2.00	35 10 693. 65 10	at 26 other	in) 104RS/1 (gray	a Buckground
રહ -	-	!	300	35.5-515 Sil	ا در.≺anàu	Gravel (ms6) 15 10 51 17	LEL Ldetect
-	-		<u></u>	30 10 sand 55 10 at	avel soo	ad f-cse poorly	
37 -		100 %	500	sorted, SH prove	۸ SA.	oderstaly mas size	molesture sample Q325
	134505		28 X	Sm coloblex 10YR	612 1E	brownish gray 30%	
38 -	55 #16	25%	5	bas Joh atz. S	Labt CA	× 4	
-	-	Nec.		37' 5:1+4 Sondy	N Gravel.	15 10 Silt 4010 Sand	
29 -	-		0	45 10 0 0000 500		modul sector SR	
.	1610 hrs	-	5,05			0 10 ¹ /0 11 10 10	maisture so-alast
	55 # 17	Archive	1 6-0	grave, poor is a		, to all a long the	1 24 /
	- !	55%	5=5	miled; 40% che n	<u>hax 5 x</u> t	= ig personte. Lonkou	
•	- 1543 hr	1 T	55.	gray. Gravel pre	d basa	12. 20 3 gt > (other)	brob Archive a 40.0
41 -	- 20	ļ	$\tilde{D}_{\mathcal{G}}$	32 10 basate no	rxn H		
·	- <u>~~</u> .	_	19. A				Moisture sample
42	- 20		500	si		···	Q. 41.6
i –	- 12	1					
43	- 8		- 0÷				
	53*	ৰ	D Z				
	ده۳۹. 4 م ا	100%	2				
14		rec	SLC.			· · · ·	10101524H. 30 mpt 10/443
Report	ed By: <	Vaclas	- The a	ີ 	Review	ved By: DCUleeke	ـــــــــــــــــــــــــــــــــــــ
Title:	Genten				Title:	Geologist	
Signat	ure: C ha	Real mar	<u>र</u> ू	Date: Date:	Signat	ure: Dealecked	Date: 10/16/01
			5		<u> </u>		
HI-EE-	183 (12/97))	~				-:

			в	OREHOLE LOG			Page <u>4</u> Date: اردن	of <u>յգ</u>
Well ID:	63391		Well N	ame: 299-533-338	Locati	OTT SG CONNON	84 241-	B Tank Farm
Project:	C.4 A.	000-			Refere	ance Measuring Point:	6	1 Sumfred
,	Sar	nc <u>ck</u> nole		Samol	e Description		Cor	nments:
Deplh <u>(Ft.)</u>	Type No.	Blows Recovery	Graphic Log	Group Name, Grain Size Di Moisture Content, Sorting, A Size, F	stribution, Soit Angularity, Min Reaction to HC	Classification, Color, eralogy, Max Particle 1	Depth of (Method, M Sampling Size, 1	Casing, Drilling ethod of Driving Tool, Sampler Water Level
45	67/26/01 10740	Arch. Ve	S O O	45 same description		004 3	Gash wood	
-	55 * IQ		0.00 0.00				moisture.	ample att
~~~~~ 			0.0			-()		
יאיז – איז –	55 # 20	100% Hec	1000 0100	4) "10 s changin 25 "10 site 10"0. 6	ng. 6 navel ng. 6 navel	2 65 % , sand		
48 -		100% pec.	1910 1010	celebles, Strong ex	n Hc.ı			
49 -	<u>0952</u>		0 <u>0</u> 0				moisture	sample & 49.5
so -	55 #21	<u></u>					Grah Rry	hive Q so.o
51 -	-	Recovery 85%	000	51.5-52.0 54	i lens.	interbodd ed	Grab Ar	chies DS15
52 -	100 hrs	-	0-0-6 2	with silty sandy and	well Silt	<u>Sis patz</u>	ODISTAN	Sample Quite
53	-	ພວິ້າ		Distic di 1 6/3 1	ghi bellon	sish brown (dry)		
. 4	605	-		62,0'-5-25 5.114 5au	ndy Snavel	(msq) Gravel	Maisture.	50mpl. @ 548'
55 -	- 55	<u>arcaiz</u> .		45 % sand 25 %	517. 16/6. 5	and t-cse,	Grab Ar	chire @ 55
56	- 55 # 23	100 % rec.		20 10 gtz (other) 30 brownish gray. Stro	(e basalt	104R613		
57	- 0628 - 0628 - 24	r		<u>53.5'- 57.7' 5:17</u>	y Sand (m	<u>s) silt 25%</u>	moisture	sample @
58		100 % rec	0 0 0	nodules, 104R.512	grayish	hrown Strong	31.7	
	0 741		0.0.0					
Report	ted By: c	marian	m		Reviewed By:	DCUkeek	29	
Title:	(	····	- <u></u>	<u></u>	Title: Geo	paist		
Signat	ture: c C		$h \wedge \pi$	Date: Date:	Signature: /	NC akekor	Da	ite: 10/16/01
BHI-EE	-183 (12/97)	) )	Year M					

			B	OREHOLE LO	G		Page <u>5</u> of <u>19</u> Date: 7/1 7 /01
Well ID:	122	<u>e r</u>	Well N	Name: 299 - 520 - 1	770	Location: SE Courses	
Project:	<u> </u>	A Thu	•		225	Reference Measuring Point	G Suf
1 10,001	<u>KCK</u> Sa	<u>//</u> mole	ling 1	<u>San</u>	inle Desc	ription	Comments:
Depth <u>(Ft.)</u>	Type No.	Blows	Graphic Log	Group Name, Grain Size Moisture Content, Sorting Size	Distribution g, Angular e, Reaction	on, Soil Classification, Color, ity, Mineralogy, Max Particle n to HCl	Depth of Casing, Drilling Method, Method of Driving Sampling Tool, Sampler Size, Water Level
60 —	55 # 25	ARCHIVE	38 0 38 0 8	57.7'- 66.3' 5-	nd [S]	: 95% sand 5%	Grab Archive @ 60'
-		100% 114.		arrivel: sand: mea	l. – v. Co.	urse, well sorted,	
61 - -	0007 55°26		ас н с – о	SYR 4/3 reddish bro granules (2-4mm).	NA NO	RKN. to HC1; gravel : to HCL	
62 <del>-</del>		100% Ftc.	•	@ 59.9 gravel	increases	to 10%, max	
63 — —	2848		6	size still granule ( (90% 5, 10% 6)	<u>z. 4 mm)</u>	still a sand [5]	
64 - -	55#27			@ 61.3' silt in	TERSES	to 5%, gravel	Grab Archive @ 66
45 — -		100%+46,		decreases; still. (90% 5, 5% M	a 5900 , 5% (	<u> [s]</u> g)	
66 -	0156	ARCHIVE					
	557 <b>4</b> 239	108% + ec.		66.3' - 71.3' 5. 85% Sand, 10% sil med si coarse, mo silf % higher, roo same, No roots.	<u>ightly</u> S H, 5% d. sort. to HC1 HC1.	iiH, Sand [(m)s]; gravel; sand; , No exis to Act; ; gravel the	
70 —	55 *29	106% r#c.	- <u>-</u>	71.3' - 72.3' 5 description (95%	and [5 5, 5%	]: see above 6 G. frace silt)	Grab Archive @ 71'
י- יר -	/053	ARCHIVE		72.3' - 78.3' S	il <del>ly Sa</del> ravel :	nd [mS]: 80% sand, sand : fire - med.	
72 -	55 #30 /7_/6	100 % vez.		well sort. No (XN) some color; silt: max size = granu @ 75' arous!	ta HCI, 8 highar 1e (Zma	s to 5% con	
				to He1 7/ silt. (7:	5% <u>S</u>	20% M , 5% (G)	
Reporte	d By:	Toss L	facking		Reviewe	ed By: DCWeekes	<b>-</b> ,
Title: (	Geologia	.t	٦.		Title:	Geologist,	4
gnatur	e: Je	4 Hack	ang -	Date: 07/27 /0/	Signatur	e: Sealeckes	Date: 10/16/01
7E-1	7 83 (12/97)		/				, , , , , , , , , , , , , , , , , , ,

			B		G		Page <u>6</u> of <u>19</u>
	•						Date: 07/27/01
Project:	<u>C 339</u>	۱ <u>.                                    </u>	vveu i	ame. <u>299-633-3</u> .	38	Poterona Manual Point	E 241-B Tank Form.
Fioject.	KCRA	<u>  Drittin</u>	<mark>р Fy</mark>	21		Reference weasoning Politi.	Ground Surface
	53	mpie [		San	npie Desc	ription	Comments:
Depth ( <u>Ft.)</u>	Type No.	Blows Recovery	Graphic Log	Group Name, Grain Size Moisture Content, Sorting Size	Distributio g. Angular , Reaction	on, Soil Classification, Color, ity, Mineralogy, Max Particle n to HCI	Depth of Casing, Drilling Method, Method of Driving Sampling Tool, Sampler Size, Water Level
75 —		100% -10.		78.3'- 79' Slight	y Silt	Scarelly Sand [(m) g S] :	Grab Archive @ 76'
	1236			75% Sand, 15% s	;   <del> </del>	0% gravel; sond:	
76 -	55 # 3Z	ARCHIVE.		v.fine - med. (see	descript	tion page 5);	
-			o. d	grave : sub ang -	sub. rnd.	; silt & down,	
77 -		Ino To rec.		ren MHCI.			
-			o				
78 -	1347 55#33	ł	0.0.0	79 - 94 Slightly	Silly	Sand [(m)S]:	
-				(see description p	<u>, ge 5.</u> )	, run to Hel Trill.	
79 -		100% 186.	<u> </u>				
-	1421	dactions				<u> </u>	CLAL Col
00-	55 8 34			18 89 Sand be	ermes or	tine - uitine ,	Grab Archive @ 80
			<u>ہ</u> ہ	gravel still @ 5	70 <u>65</u>	<u>a silt still</u>	
81 -		100% 100.	- 0	strong	<u>[C</u> 1.		
-			6° 7				 
87 -	55#35	-					
-			<u> </u>				
83 -		100% 184.	• [—]				
-				···· · · · · · ·			
84	1523	-					
-	$ \times $		- o o				
85—	55 # 360	1					
-		No. Pr or	•				
86 -		100 /2 /20.	00 00	· ····			
-							
87 -	<u>0725</u> 55 # 371	ARCHIVE					Grab Archive @ 87'
-	•						_
88 -		100 Torec.				• • • • • • • • • • • • • • • • • • •	
-							
87 -	\$755	-					
	$\geq$	<u></u>					]
Reporte	d By: 了	ss Hock	<u>64</u>		Reviewe	d By: DCUkekes	
Title: C	heologist				Title:	Seologist,	
gnatur	e: Jos A	hand	_	Date: 07/27 /01	Signatur	e: NC alecher	Date: /0/16/01
	/					* •	11

FE-183 (12/97)

			D/	ספר	นก			<u>`</u>			Page <u>7</u> of <u>19</u>
				JRE	пО			2			Date: 7/31/01
Vell 1D:	03391		Well N	ame:	233-	633	- 33	8	Locati	on: SE Corner of	241-B Tauk Farm.
roject:	RCRA	Drilling	FY o	1					Refere	ence Measuring Poi	nt: Ground Surface
	San	nple J					Sam	ple Des	cription	I	Comments:
Deplh <u>(Ft.)</u>	Type No.	Blows Recovery	Graphic Log	Group Moistu	Name ire Cor	a, Grain Intent, S	Size I Sorting Size,	Distributi Angula Reactio	ion, Soil rity, Min n to HC	Classification, Colo eralogy, Max Particl I	r. Method, Method of Drivin Sampling Tool, Sampleu Size, Water Level
of	55 1 38										
_											
a.		pot rec.									
71 <b>—</b>											
_	0828		- -								
92 —	554 31										
—											
<del>9</del> 3 —		100 \$ 100.	<b>e</b> 3								
_	!										
പ്_	0850	APAUNE	•	au!			51.11	1 64		a & 1[(1), \$]	1 0 1 0 L 0 0 0 1
'' –	$\geq$		0.0	<u> 17</u>	- <u>704</u>		Jight	1 <del>7 Sin</del>	<u>i Linave</u>	IL SAME LIMING UI	Grab Archive C 77
-	53 60 4/0 I			<u>( see</u>	desc	r p tie	n. 70	<u>9</u> 2	1. 757	8 sand, 15% silf	
» —			0	10 %	gran	<u>el.</u>					
-		torest , ec.	00								
% —			L 60								
_	0929		6 6 9								
A1	ssasatı I		00								
<i>77</i> –	1		<b>?</b>								·
		(** 26 rec.	16 <b>9</b> (* )								
98 —			5 <b>-</b> 8 (	0 9	8'	sand	15	fine -	med.	, well sorted,	
_	10 10		60	silt	<u>a</u> ^	њ. <u>н</u> с	1.:. M	ak.			
99 -			0.00								
_	332 42		0								
		100% ~~~.	-								
			0: 0			·· ·-					
-			<u>.</u>								
/01	1038	ARCHINE	0 								Grab Archive @ 101
-	35 # 43										
101 _			- 205								
		an grad.	0 0.0		7					······································	
- 2°			o	(* 103	6 :	sand u	is i	L fine -	med.	, well sorted;	
-	55 4 4 4		·e 9	_≾:/↓	(***	/401	5 10	<del></del>			
104 -			0.0								
-										<b>.</b>	
Reported	d By:	less the	king					Review	ed By:	DCWeek	20
fitle:	Geological	7105						Title:	Ge	plogist	
Signatur	e: 4	011	2		Date	21.1		Signatu	ire:	MAZibo 60	Date: In In 14
	- yes	. Hacku	19/		100.00	115410	•	2.3.040		( / V CARSENCE	

			BC	REHOLE LC	G			Date: 7/31/61
fell ID:	(224)		Well Na	ame: 233- F23- 3	38	Location:	SE Corner de 2	WH-B Task Farm
roiect:	2 4 2 A					Reference	e Measuring Point	Court Salua
	Sal	mole	FY 01	Si	ample Desc	ription		Comments:
Depth ( <u>Ft.)</u>	Type No.	Blows Recovery	Graphic Log	Group Name, Grain Siz Moisture Content, Sort Si	e Distributio ing, Angulari ze, Reaction	on, Soil Cla ity, Minera n to HCl	ssification, Color, logy, Max Particle	Depth of Casing, Drilling Method, Method of Drivir Sampling Tool, Sample Size, Water Level
o5 —		40% rec.		104.5 - 114	Silly Sano	1 [ms]	: 80% sand,	
_	1225	ł		20% silf , trace	gravel ;	sand: fir	ne - v.fine,	
106 —	55 • 45			v. well sort. , su	b. rad.	<u>5 # %</u>	bark uz,	
- רס		100 % re.		<u>rra: to HCL; g</u> son. greatle. (s	similar to	descript	max size =	
	1720					-		C L O L'IN Q INC
- 8ª	ss ≠ 460	AKLINAY	•	·····				Cirab Archive & 108
109 -		100 % r 22.						
	1348 55847	4						· · · · · · · · · · · · · · · · · · ·
	-	100 <b>% 12</b> (.		····				
	1433							
	-  55 <b>*</b> 48	100 % Mar.					······································	
114 -	-		<u>ビラムで</u> - 100変換	114' - 127.6	Sand I:	s]: 95	% sand	
-	- 55#49	- HKCHTAF		5% silt; sand	<u>: Ufine</u> an arm f	<u>- U. Cog</u> La HAI	<u>well sort</u>	Grab Archive @ 114.0
- مار	-    -	100 % met.		Sill % lower,	ixn to H	+C  mode	μη. 4 <u>e.</u>	
-	-	7						
- ייו	1 X I					· •		
118 -	55# 50			2 2 2 				
119	_	100% 12		@ 120.5' trac	<u>e gravel</u>	appears	, max size =	
Report	ed By: -	 ۲۵،۰۶ ا	, <u>kin</u> c		Reviev	ved By:	DCukek	es
Title:	Geologi	~ <u>+</u> ~			Title:	Geole	gist	
Signati	<u></u>	011	<u> </u>	Date: 7/2. /	Signat	ure: M	ape her	Date: 10/16/0

A.8

			BC	DREHOLE LC	G			Date: 7/31/01
Vell ID:	63341		Well N	ame: 299 · E33 · 34	38	Location: $SE$	Corner of	241- B. Tunk Farms
Project:	D/pn	<b>N</b> . (1).	EVAL			Reference Meas	uring Point:	Ground Surface
	Sar	npie	<u>_</u>	S;	ample Desc	ription		Comments:
Depth <u>(Ft.</u> )	Type No.	Blows Recovery	Graphic Log	Group Name, Grain Si: Moisture Content, Sort Si	ze Distributio ing, Angular ize, Reaction	on, Soil Classifica ity, Mineralogy, N n to HCI	tion, Color, lax Particle	Depth of Casing, Drillin Method, Method of Driv Sampling Tool, Sampl Size, Water Level
/20		Joo to etc.	1- 1-					
_	0655	ARCHIVE						Grab Archive @ 120.
121 -	35 4 51							
122 -		100% rec	o. 					
123 -	0740 55 4 51	•		@ 123 trace	<u>. gravel</u> . coarse .;	<u>ne lenger p</u> <u>still ~ 5%</u>	resent <u>;</u> si H.	
124 -		100% 184.				<b>_</b>		
125	0808	ARCHIVE			<b></b>			Grab Archive @ 123
- 126 -	55 # 53	1 1 1 1 1					·	
- דנו 	0826			127.6' - 136'	<u></u>	Sund Em S]:	80%	
128 -				silt & increases	(see d	<u>: utine = me</u> lescription page	<u>e 5</u> ).	
130		10 <b>0%</b> nec.						
131 -	0132	ARCHIVE					-	Grab Archive @ 131
137								
183	_	100 % ret.						
134		ARCHIVE						Grab Archive @ 1
Report	ed By:	Jess	Heckine		Review	wed By: <u>X</u> /	<u>ikeke</u>	<u>ح</u>
Title:	Geolo	eist			Title:	Geologi	st ,	
Signat	ure: 7	11	12-	Date: 1/2, /4	Signat	ure: NC	ti la a lle a	2 Date:10/1/

		· ·	387-11-2					Leastier		13	101
Vell ID:	<u>C339</u> ]		Well N	ame:	299-	E <u>33 -</u> 3	38	Location: SE C,	rnar of	241 B	Tank Farm.
roject:	RCRA	Drilling	FY 0	1				Reference Measur	ing Point:	Ground	Surface
	Sar	mple				Sa	mple De	scription		Cc	mments:
Depth <u>(Ft.)</u>	⊤ype No.	Blows Recovery	Graphic Log	Grou Moist	p Name ture Cor	, Grain Siz Nent, Sorti Siz	e Distribu ng, Angul ze, Reacti	tion, Soil Classificatio arity, Mineralogy, Ma ion to HCl	in, Color, x Particle	Depth of Method, I Samplin Size,	Casing, Drillin Method of Drivin g Tool, Sample Water Level
35 —											
_											
		1,20,710		1.97	· .	ا جر ول	< 1	IS]: 96%			
36 -		l		136		<u>43.15</u>	Jend	131 /3/0 SAN	<u>د</u> ب		
-	1156 SS#57	+		5%	si   -	; (see a	desseriet	ea. page 8.1			
137 -											
		1007									
138 -											
	1243	ARCHTUS									
139 -	55 58	HIGHLANE	100							larab <u>Ar</u>	<u>chive (0° / 39</u>
-	·			<u> </u>							
140		100% -44.		e	140.8	sand	: fine	- meet. , well sol	· <del>i</del> . ;		
-	. 1			: از زاد	~ 5	% silt.					
	1326	4									
••• =				<b> </b>							
-	-			-						<u> </u>	
142 -	- 1	100 % MC.								· · · · ·	
_	-			<u> </u>							-
143 -	- 1401	_		14	3.15	- 167 -	Silly 3	and [ms]: 80%	sand.		
-				20 %	s it	: (see	descript	ion map 5)			
أعدر						,		····			
140 -		100% 1-46-									
-	-		<u> Sister</u>	( <u> </u>							
MS —	55#61	ARCHEVE				-				Grab 6	rchive (@ 14
-	-			1			<b>.</b>			<u> </u>	
146 -	_ 1		-								
		100 % rec									
141.	1545			1						<u> </u>	
_		1									
ાપાટ	- 53367									+	
				) 						<b>-</b>	
149	_	100% 190.		ð.							
	_	1									
Report	ed Bv:	<u>т</u> (	le.				Revi	ewed By: DALL	bekes	5	
Title:	6.1	<u>uess Ho</u> s I	<u>, t, n q</u>	······································			Title	Geologist	<del>/</del>		
Cine	reologis		1-				Cina		la la	/	nate: In lis la

			BC	DREHOLE LOG		Date: $\gamma/_{31}/_{ot}$
Well ID:	C 339	l.	Well N	ame: 299- £33- 338	Location: SE Corner e	Z41-B Tank Farm
Project:	RCRA	Drilling	FY OI		Reference Measuring Point	Ground Surface
	Sa	mple		Sample Des	cription	Comments:
Depth <u>(Ft.)</u>	Type No.	Blows Recovery	Graphic Log	Group Name, Grain Size Distribu Moisture Content, Sorting, Angula Size, Reacti	tion, Soil Classification, Color, arity, Mineratogy, Max Particle on to HCl	Depth of Casing, Drilling Method, Method of Driving Sampling Tool, Sampler Size, Water Level
150	1410	ARCHINE		See description on a	10	Grab Archive @ 150'
	5 [#] 63					
		100% -44.				
131 -			<u> </u>			-). 
					·	
152 -	55 464	+				
	- 1					
- 53	-  1	100% -24.				
154 -	55* 65	ŧ.	12 <del>-</del> - T			
- 1				·	·	
155 -	-1 1	100% -44.				
] .	-					
					•••••	Cab O line @ 156'
150 -	~~~	ARCHINE				SIMP ACHIE C. 100
-		4				
157 -	- 55+ <del>0</del> -466	1	<u> </u>			
_	-1			· · · · · · · · · · · · · · · · · · ·		
158 -	_	100% 100	17.17			
I .		1				
	0920	_				
159 .	55 1 (07	-				·
1	- 1	1	- T-			
160 -	-1	100% - 40.				
- I	_					
			- 200			Cash Archive @ 161
	\$5*68	ARCHIVE				
	-	1				
162	-1	100 3 144		(d		
1 -	-		-			
163	- 1030					
	$_N/$	7				
169	-K-55 #6	7				
					he bault	
Report	ted By:	<u>ess Ho</u>	cking	Revie	ewed By DCWeeke	<u>s</u>
Tille:	Geolo	gist.	~	Title:	Geologist .	بر شور و در <u>ا</u> ر ا
Signat	ture: A	Weeke	2 Vale	ss fock: Date: 10/16/01 Signa	ature: / Calbelle	Date: 10/16/01

-			B	DREHOLE LOG		Page <u>12</u> of <u>19</u>
Well ID:	63:	61	Well N	ame: 249 - F23- 33 K	Location: SP + F 2/11 R	Tente Contra
Project:	5001	CRI T	Dalling		Reference Measuring Point:	Grank Fellow
	Sa	mple		Sample Des	cription	Comments:
Depth <u>(Ft.)</u>	Type No.	Blows Recovery	Graphic Log	Group Name, Grain Size Distribut Moisture Content, Sorting, Angula Size, Reactio	tion, Soil Classification, Color, arity, Mineralogy, Max Particle on to HCl	Depth of Casing, Drilling Method, Method of Driving Sampling Tool, Sampler Size, Water Level
165		100% -40.				
166	- 12.11 	ARCHIVE				Grab Archive @ 166
- 167 -		10 <b>8 % r</b> K. 1		167 - 169.5 Sand [5]	: 95% sand 1 5%	
168 -	1239			sitt; sand: y.fine - r sort, No ran. to HCI;	ned., sub eng., well silt. % decreases,	· · · · · · · · · · · · · · · · · · ·
		1042		strong rxn. to HCL; (so	<u>e description page</u> ).	
170 —	. 1331			169.5' - 171.45 Sitty Sand [ description page 10.) 8	-5]: (see 30% sand, 20% silf.	
- ידי ידי	55± 72	100 % 186.			· · · · · · · · · · · · · · · · · · ·	· · · · · · · · · · · · · · · · · · ·
- 271	1561	4740.45		171.45 - 173.05 Sandy Silt 25% and ; Silt % in	LSM : 75% SIH,	
- גרו - גרו	55 # 73	PECHIVE		sand: v.fine - fine , su well sort. , No man to F	ib. ang sub. rnd 101 , % decrenses.	Grab Archive @ 172.5'
-		100%		179 pp 195 5.15 6		
· · · ·		1 maring		description and In ) 20	% and 70% all	
175	1610	Arctime		175-203.5		Grau Sample @ 175'
176		tot to rectienty		Lenses of u st. silly sand	nd (5) with sporadic ((m5), Jt 13 45%	
- - רדו	- 			Sand (15/001 40% in 4 The grains are SA some	<u>10% v \$ vf), 5% silt.</u> SR. and the max.	
		Z		SIZE is 4mm or less, as	and the unit is	2
-	6714			exhibits a week rxn	to HCI. The sand	
179 -	$\geq$	]		also shows 14 to 12" in Comented (slightly) now	ndi whitish gray	
Reporte	d By: 🗸	ss flock	<u>ting</u>	InFaurch Review	ved By: DC Weeke	5
Title:	deret	a hua	6	rectigist Title:	Geologist,	
Signatu	10: ya.	esstocky	e ynnfa	Unite Date: D8/05/07 Signate	ure: / NC yee hea	Date: 10/16/01
BHI-EE-1	83 (12/97)	-	99			· 200

		5-1	B	OREHOLE LOG		Page <u>13</u> of <u>19</u> Date: 08/05/01
Well ID:	6.3		71 Well N	lame:	Location: Set is assored	T 4 C
Project:	2001 2	1011	<u>. //</u>		Defense Manual Defense	Tank Ferrin
Tiojeet,	5007 1	CRACIT	<u>Illing</u>		Reference Measuring Point	Convend Sartace
	Sa	mple	-	Sample De	scription	Comments:
Depth (Ft.)	Type No.	Blows Recovery	Graphic Log	Group Name, Grain Size Distribu Moisture Content, Sorting, Angul Size, Reacti	tion, Soil Classification, Color, arity, Mineralogy, Max Particle on to HCI	Depth of Casing, Drilling Method, Method of Driving Sampling Tool, Sampler Size, Water Level
180 —	55#75	Archive	-	wk- maderately well of	HCI. The unit	Grab Sample 180'
-			]	contains mica (mu	scor, bictile, chlorit	
181 -		135%	-	and presibly philege	ate)	
_		recording				
182 -	0802	, í	_		<u>-</u>	·
	55 # 76	-				
			-			
183 —						
-			÷.	·		
184 —	0835	-				· · ·
-	$\mathbb{N}$					
185 —	ŀ <del>X</del> -	Archive	1	@185, H& average an	In STRC IN REASES	Grubsamphe @ 185'
	$\checkmark$		[-	to cr & ver, approx	50% of sand, w	
186 -	158 77			30% m and 15% F w	520 st (or less), Ilin	ł
_		100%		continues to coact must	- SI to HEL TH	
187 -	- m	recaph	· · · · · · · · ·	continues to be day	to st yourst	
	1041	,		1	10 St. MICISE	
(D)	) Š	1				
186	55178	1	-			
_		moin				
189 -	·	recevery	_			
-		···)		Gram size returns 1	2 Juleer, Utilom,	
190 —	206	Antino		30-35% f , 5-10%	sitt in It 94-ba	Giab Sample @ 190'
_		<u> </u>		sand.		, , , , , , , , , , , , , , , , , , , ,
191 -	<del>5</del> 5'''''''''''''''''''''''''''''''''''					
_	ાલ્ટ ન		·• •	there are a second second	in the list	
			0	Trace gravel 191-10	12 , 131(1X (24) ~ 12 X	ra
142 -	124640		· · · · · · ·	unit continues & 4	5-55% basalt and	silica
-					<b></b>	
193 -	52*80		-	·······-	··· <b>····</b>	IH PM check -
-	192.4					<i>idetectable</i>
194 -	· [94.6		0			
Reporte	d By: 1	nFaun	ole	Review	ved By: DCUEEtes	••••
Title: 7	Soplan	st.		Title:	Geologist	• • • • • • • • • • • • • • • • • • •
Signatur	e: Im?	Lour	oto	Date: Coma las Signat	ure: Manballan	Date: Infusion
	Jug	MALT.	<u>. ur</u>	10 JUJ 01 01	mana .	10/16/01
HI-EE-1	33 (12/97)					

			В	OREHOLE LOG		Page <u>14</u> of <u>19</u> Date: 15/03/01
Vell iD:	CE	3341	Well N	lame: 349-833-335	Location: Set from 24	11-B Tank Farm
Project:	2001 no	CKA Di	than	• • • • • • • • • • • • • • • • • • •	Reference Measuring Poin	1: Ground Surface
	Sa	mple		Sample De	escription	Comments:
Depth (Ft.)	Type No.	Blows Recovery	Graphic Log	Group Name, Grain Size Distrib Moisture Content, Sorting, Angu Size, Reac	ution, Soil Classification, Colo Ilanty, Mineralogy, Max Particl tion to HCl	e Bepth of Casing, Drilling Method, Method of Driving Sampling Tool, Sampler Size, Water Level
s		Archa.				Comb Sumple @ 195
_	$\mathbb{N}$		<b>.</b>			datensine 201.65 peur
96 -	ΙX			Continues as sand	in-cogina wi	
_	$ / \rangle$		÷.	incderate and of F	e exides in motorix	
47 -	$\angle$			Along with whow rea	col (CO2) that will	î.X.X
···	57# 591			amort-sa ryn told	1. SPREADIC IFA	inice/
.98 -	ļ	100%		(3A-A 1/1 +03/8").	<i></i>	Ind shuff Stastar
		receivery				109.1
	14321	7			*****	Start 28/26/01 8.199
<b>77</b> -	$\overline{\mathbf{\nabla}}$	1				
	ÌΧ.					- Cif (mari
2001	55#82	Archive		F. A		<u>Sarti Sample &amp; 200</u>
_				Janaris 5510C	r solo in 10/072VF	
201 —		100%		with sto site, the	at is comprised of 60	<u>2%</u>
_		receiving		Dubalt, is med-clk gn-	on w/ gy-wh CO3 Kel	neert.
202 242	07/2 1	1	-	The sand has a modes	rxntoHG	
2024		-				· · · · · · · · · · · · · · · · · · ·
Z03 —	0880				• • •	
-	-	100%		203.5-212.5 Sl.9	savelly sand,	
zo4 -	-	receirg	-0.	vf-fgravel @ 157	sitts Poor less,	and
a.H.T	0748 hi		1. S. U. E	Sand C. 80% (15 100	embed above)	Grab Sample & 205 H
بر سر کرم	$\sim$	1 <u>Archwe</u>	° r č			
253	52#84	4	E)			
716 -	-		с.			
	_	100%	. 0	by 206.5 there is	an morener in the	
707 -		cecivery	6	acavel to about 20	10 ( SA 70 & 70% 11 1	c/
367.5	s <u>C352 h</u>	4		Laccost & statistic x/	") Tot is pasalt rich	<u> </u>
200	$ \times $		6-0.	and reach and to ce	to Hrl	
6×8 -	58485	, ,		I and I chilly much de sy	16 1861.	
0		1000010				
209 -		بالمريورين				-
Bonada	-   M Dur Du	<u> </u>	₽. (. ⁻ 0.		awad Bur DAlibal	
Tale: 1		Taurote		Revu	men by DCURER	25
	rdogi	st			600/0915t	a later
signatu	resented	urite		Date: 00/06-/01 Sign	ature: / vc allel flea	Date: /0/16/01

						Page <u>15</u> of <u>14</u>
						Date: 08/06/01
Well ID:	_C.33	91	Well N	ame: <i>744-E33-33</i> 8	Location: jE from 24	1-B Tank Farm
Project: 2001 RCRA Drilling					Reference Measuring Poil	nt: Ground Surface.
	Sample			Sample Description		Comments:
Depth <u>(Ft.)</u>	Type Blows No. Recovery		Graphic Log	Group Name, Grain Size Distribution, Soil Classification, Color, Moisture Content, Sorling, Angularity, Mineralogy, Max Particle Size, Reaction to HCI		r, le Sampling Tool, Sampler Size, Water Level
210 210-1	0920 hs	Meisture in Richard	$e_{\ell_{i}}$ , $r_{i}$	max go Gretsite is 1/2	xix 3/4" Gravel	Gant Sample @ 210
-	55#86			content ~ 25%, s,	it increto 8%	
211 -		100% recovery				
	<i>1949</i> bs	noistere tim	e 	212.5-216 Say	of Toto Sand, Sto Sil	<u>ک</u>
213-	55+57	6	sloug H	His sand is It-bo to	tan 52-34, 17-to	<del>f</del>
- 4 -		40% гессоггу		grained, it exis to	Hel. <u>Lecally the sa</u> madarately ubundici	u <u>d'</u>
215 <u>p4⁸</u>		Mershive In Archive		Sitend Content	218 Silty Sand	Carab Jampk @ 215'
-215 U	55#88	100%0		In anotherwise vt-	fgind sand.	
217 -		recovery				
217. ⁴ 218 –	1230 hr	tin tin		DIR- 122.4 Silt to san	dy site - Chacolate	Pre-Hanford Scils
21 <del>9</del> -	55#89	100%		and 80% sill	t shows v. thin te fonses of sand	micular
270 —	-	recarry		distributed through	out.	6146 Sample 220'
	- <u>1328 hi</u> - 55#90 -	Archive maisture Tin e 220.8		Dand content de by 219'; there is to clay, and the uni	cceases to near zo trace to about 3-5% It is moist to sluce	en
222 -	-				SA	the 55 weephote
<i>722</i> 223 -	<u>-114236</u>	could not retain in drive		10% Silt, 35% same cr to sm petbles 20%	4 Sandy (sravel, 58 1, 55% (sravel (40%) 1, 30% f) und sam	RETPICHOCK < detect
224 -	- //\	barrel	633	13 20% cr 50% 30	76 f-vf in a moist,	ret bn Hanturat Gravel
Reporte	~িহ্যাক্র ed By: <i>∭</i>	Mauch		Ren Ren	viewed By: DCubeke	Jarge closts ere basa/t
Title:	eologist	/		Title	· Geologist	
Signature: Manuela, Date: 08/06/01 Signature: MCabeker Date: 10/16/01						
			в	DREHOLE LOG		Page <u>16</u> of <u>19</u> Date: 08/06/01
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Veil ID:	6.3:	391	Well N	ame: 299-#+1875 338	Location: SE From 241	-BJankfarm
Project:	20011	CRAD	cilling	- 11	Reference Measuring Point:	Connerd's Surface
	Sar	nple		Sample Desci	ription	Comments:
Depth <u>(Ft.)</u>	Type No. 55 ^{#41}	Blows Recovery	Graphic Log	Group Name, Grain Size Distributio Moisture Content, Sorting, Angulari Size, Reaction	n, Soil Classification, Color, ty, Mineralogy, Max Particle to HCl	Depth of Casing, Drilling Method, Method of Driving Sampling Tool, Sampler Size, Water Level
225 —	Control	Archit	11155119	] Continues as V. poor	sorted gravel.	pushed large rock
-		60% 1	1070	actual silty-sandy-grav	el the unit is	of shoe, honded reciver
226 -	1524 40	masture		evenill whogy browith	COg as coment	Grab Sample @ 225
- 227		tin	080	SPRC in the gravel is 32	24"x2" The unit	
	1	100%		15 Very poorly surted and	dry.	
2 CK -		receivery	己弟			
22ነ -	<u>ση2 h</u>	mersture	0.00			
230		Archive		Challens	" _ Y. H	Guide Sampte & 250'
- 231 -				Graver site averages 1- a silly sandy Gravel	112 , romains (10% m, 30% s, 60% g	>
737	, <u>cs3ohr</u>	TIN TIN	N04 FOT/2 Geoge	Gravels are dominantly has slight to moderite (re	basalt. The unit lated) (xn toHCL	
233 -	65#94			all a fair March		
- 234 -		95% reavery		moderate rxu of the	scie). Slight to	
- 235	1248 400	MCIANUE	000 800 800	where the chrand 1-	1/2".	Grabsample.@235'
236 -	- 45 495	ARCHIE	WISSIN'S	Same as above except	r gravel to 21/2"	peor receivery in
- 237 -	-	Belie Belie		and dry. slightly le	ss sand.	shoe for mostaneti
2372	15171-1	HOIGHTURE TIN	M451.NI7			
2348 239 -	55# qu	_			$\frac{1}{2} \frac{1}{2} \frac{1}$	
Reporte	=1 ed By: 7//	1 Courses	o R	4 M- (carse sand, Strighthat) ( Hubber) Reviewe	ed By: ACUBECo	<u>ເທຍ</u>
Title: (-	tedras	<u>1 uunu</u> 1	·O	Title:	Geologist	•A
	rol h	K Ripo La	denne	Date: 08/67/01 Signatur	e: No Tibo has	Date: In/1/01

BHI-EE-183 (12/97)

			B(	OREHOLE LOO	;		Page 17 of 19
Well ID:	1 220	31	Well		 24	Location: CE (	et the
Project 2001 RCRA A-11-1		9	Colorini SE from 241	D lank rarm			
- tojeci.	2001	<u>ηι</u> ηη		ng		Reference Measuring Point	Ground Paltace
	্বৰ	mpie		Sam	Die Desc	ription	Comments:
Depth (Ft.)	Type No,	Blows Recovery	Graphic Log	Group Name, Grain Size D Moisture Content, Sorting, Size,	Distributio Angular Reactior	on, Soil Classification, Color, ity, Mineralogy, Max Particle n to HCI	Depth of Casing, Drilling Method, Method of Driving Sampling Tool, Sampler Size, Water Level
z40 —			$\mathcal{L}_{\mathcal{O}}$	(anauch nounded to	500	angelene slightly	
-	<u>।</u> 1467	MILLINAVE	500	Maist strong rx	n w/H	KI. M-C Sand SESA	Grab sample 240'
241 -	$\left[ \cdot \right]$	TIN	č^0	(60-G, 25% CG	, <u>1:59</u>	esilt)	
			5.0%				
247 -	55#97	ł		Silty-Sandy-Grave	$\sum 6 \alpha$	avel rounded to SA.	
-		100%	$\mathcal{O}_{\mathcal{O}}\mathcal{O}_{\mathcal{O}}$	Strong run will He	<u>u.</u> (	by-wh. sightly	
243 -		reavery	600	Marst. Few ox	le m	off es, course distinct	
2.83-9	1501		PX-1	Gravel to 1/2.1.			
244 —	$\sim$	MC ISTURE					
-	55#98	ARCHUZZ	S I C				(vichEample 245'
245		<u></u>		Same as about	errep	+ more moties,	,
-		100%	A. 900	and gravel to	21/2"	ı <u>,</u>	
246 <b>-</b>		recovery	0.04				
-	1010	MULATURE	<u>490</u> č				
247	$ \geq $	•		Gravel is don	nınan	1/4 1/2" to 1 1/2" with	
<u> </u>	33799			<u> Aebbles/cobbles</u>	6 44	x 6" x 2-3". large	gravels
z48 —		100%		are mainly basal	t, an	dsecondarily quar	teik.
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-	$\Lambda$ /		$\tilde{e} \delta \phi$	Fe Or stained si	Hy (10	-12%) sandy (35%)	Drum Cuttings
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-	$ / \rangle$		294	<b>^</b>			,
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			вс	REHOLE LOG			Page <u>18 of 19</u> Date: 08/08/01
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Project:	2001	RCRAL	nillina		Referer	ice Measuring Point:	Ground Surface
	Sar	nple		Sample D	escription		Comments:
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_		recovery	2000	this contains con	bles .	>6"	adefect.
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BHI-EE	-183 (12/9)	7)		<u> </u>			

# Appendix B

Summary of Geologists' Core Sample Descriptions From Borehole 299-E33-338



E33-338







B.4

# Appendix C

Summary of Geophysical Logs for Borehole 299-E33-338



# 299-E33-338 (C3391)

# Log Data Report

# **Borehole Information:**

Borehole :	299-E33-338 (C	3391)	Site:	B Farm Monitoring	Well
Coordina	ates (Plant)	GWL (ft) ¹ :	250.0	GWL Date:	08/13/01
North	East	Drill Date	TOC ² Ele vation	Total Depth (ft)	Туре
N/A ³	N/A	08/01	N/A	275.0	cable tool

# **Casing Information:**

Casing Type	Stickup (ft)	Outer Diameter (in.)	Inside Diameter (in.)	Thickness (in.)	Top (ft)	Bottom (ft)
Steel (threaded)	0.25	11.875	10.375	0.75	0	50.0
Steel (threaded)	2.13	10.6875	9.8125	0.4375	0	271.0

# **Borehole Notes:**

The borehole information provided above is derived from personal communication with T. Hottle, Bechtel Hanford Incorporated site representative. The casing information is derived from caliper measurements collected in the field by MACTEC-ERS personnel. Logging measurements are referenced to ground surface.

# **Logging Equipment Information:**

Logging System:	Gamma 2B		Type: SGLS (35%)
Calibration Date:	09/00	Calibration Reference:	GJO-2001-245-TAR
		Logging Procedure:	MAC-HGLP 1.6.5

Logging System:	Gamma 2E		Type: NMLS
Calibration Date:	05/01	Calibration Reference:	GJO-2001-247-TAR
		Logging Procedure:	MAC-HGLP 1.6.5

# Spectral Gamma Logging System (SGLS) Log Run Information:

Log Run	1	2	3 (Repeat)	4	5 (Repeat)
Date	08/13/01	08/13/01	08/13/01	08/14/01	08/14/01
Logging Engineer	Musial	Musial	Musial	Musial	Musial
Start Depth (ft)	0.0	125.0	207.0	206.0	225.0
Finish Depth (ft)	126.0	207.0	186.0	275.0	218.0
Count Time (sec)	200	200	200	200	200
Live/Real	R	R	R	R	R
Shield (Y/N)	N	Ν	N	N	Ν
MSA Interval (ft)	1.0	1.0	1.0	1.0	1.0
ft/min	n/a⁴	n/a	n/a	n/a	n/a
Pre-Verification	B0034CAB	B0034CAB	B0034CAB	B0035CAB	B0035CAB
Start File	B0034000	B0034127	B0034210	B0035000	B0035070
Finish File	B0034126	B0034209	B0034231	B0035069	B0035077
Post-Verification	B0034CAA	B0034CAA	B0034CAA	B0035CAA	B0035CAA

	4	2 (Demost)
Log Run		2 (Repeat)
Date	08/14/01	08/14/01
Logging Engineer	Musial	Musial
Start Depth (ft)	48.0	60.0
Finish Depth (ft)	256.0	81.0
Count Time (sec)	n/a	n/a
Live/Real	n/a	n/a
Shield (Y/N)	N	N
MSA Interval (ft)	0.25	0.25
ft/min	1.0	1.0
Pre-Verification	C0011CAB	C0011CAB
Start File	C0011000	C0011832
Finish File	C0011831	C0011916
Post-Verification	C0011CAA	C0011CAA

# Neutron Moisture Logging System (NMLS) Log Run Information:

# **Logging Operation Notes:**

SGLS and NMLS logging were performed over two separate days. The SGLS logging occurred inside double casing between 0 and 50 ft and through single casing from 50 to 271 ft; the bottom 4 ft of the borehole did not have casing. A longer count time (200 sec) was required with the SGLS because of the relatively thick casing. To obtain reliable spectra while minimizing overall logging time, the depth interval was increased from 0.5 to 1.0 ft. Repeat sections for the SGLS logging were collected from 186 to 207 ft and from 218 to 225 ft.

The NMLS logging occurred from 48 to 256 ft in depth through a single casing except between 48 and 50 ft. The neutron moisture tool was run centralized. A single NMLS logging repeat section was collected between 60 and 81 ft.

# Analysis Notes:

Analysi. Heliwood Date. 00/20/01 Reference. MAC-VZCF 1.7.9 Rev. 2	Analyst :	Henwood	Date:	08/28/01	Reference:	MAC-VZCP 1.7.9 Rev. 2
-------------------------------------------------------------------	-----------	---------	-------	----------	------------	-----------------------

Pre-run and post-run verification of the logging tool were performed for each day's log event. The post-run verification for log runs 4 and 5 failed the acceptance criteria. The peak counts per second for the 609- and 1461-keV energy peaks were below the lower control limit. Examination of spectra indicates the detector appears to be functioning normally and the log data are provisionally accepted. The pre-verification spectra collected during log runs 4 and 5 and the post-verification for log runs 1, 2, and 3 were used for the energy and resolution calibration for the data processing.

Each SGLS spectrum collected during a log run was processed in batch mode using APTEC SUPERVISOR to identify individual energy peaks and determine count rates. Concentrations were calculated with EXCEL using an efficiency function and corrections for casing and water as appropriate. No dead time corrections were necessary in this borehole as it ranged below 10 percent.

Verification measurements were also collected for the NMLS. Acceptance criteria have not yet been established for the newly deployed logging system. However, the pre- and post-run total count measurements agree within about 5 percent, suggesting the logging system was operating properly.

Moisture calibration models at Hanford for the borehole diameter and casing used in this borehole have not been established. Thus, the neutron log was not processed to estimate volumetric moisture content because the relatively large borehole diameter and casing thickness are beyond the range of conditions for which the tool was calibrated. Borehole diameter is a major factor in neutron response. Neutron data are presented as gross counts. In general, an increase in neutron count is indicative of an increase in moisture content, but a quantitative calculation of volumetric moisture cannot be made at this time.

The ²¹⁴Bi peak at 1764 keV was used to determine the naturally occurring ²³⁸U concentrations rather than the ²¹⁴Bi peak at 609 keV. The lower energy 609-keV peak could not be distinguished in many of the spectra within the double-cased interval from 0 to 50 ft.

Repeat log plots at selected depth intervals for KUT concentrations and neutron count rate measurements were evaluated. The plots indicate good agreement between successive log runs, demonstrating repeatability in both depth and concentration measurement.

# Log Plot Notes:

Separate log plots are provided for the man-made radionuclide (¹³⁷Cs), naturally occurring radionuclides (⁴⁰K, ²³²Th, ²³⁸U [KUT]), a combination of man-made, KUT, total gamma and neutron, total gamma plotted with dead time, and repeat section plots for KUT and neutron. For each radionuclide, the energy value of the spectral peak used for quantification is indicated. Unless otherwise noted, all radionuclides are plotted in picocuries per gram (pCi/g). The open circles indicate the minimum detectable limit (MDL) for each radionuclide. Error bars on each plot represent error associated with counting statistics only and do not include errors associated with the inverse efficiency function, dead time correction, casing corrections, or water corrections. These errors are discussed in the calibration report.

# **Results and Interpretations:**

The only man-made radionuclide detected in this borehole was  137 Cs. This radionuclide was measured near the ground surface at less than 1 pCi/g.

The KUT logs do not have sufficient character in most of the borehole to delineate any definitive lithologic units. Changes in the KUT and total gamma at about 50 ft are the result of a change in the casing configuration. The casing corrections for the interval from 0 to 50 ft are based on a combined thickness for two casings of about 1.25 in. The uncertainty of the casing correction for this thickness is significant and it appears the concentrations have been slightly underestimated in this depth interval.

Notable intervals of apparent higher moisture content exist at about 105, 172, and 222 ft. It appears these intervals are associated with finer grained material on the basis of slightly elevated concentrations of  232 Th. At the time of neutron logging, groundwater was encountered at about 255 ft.

⁴ n/a – not applicable

¹ GWL – groundwater level

 $^{^{2}}$  TOC – top of casing

 $^{^{3}}$  N/A – not available







#### ¹³⁷Cs ⁴⁰K ²³²Th ²³⁸U Total g Neutron 0 0 10 - 10 4 20 20 3 30 30 40 40 50 50 Depth (feet) Depth (feet) **60** 60 70 70 80 80 90 90 100 100 110 110 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 120 120 pCi/g pCi/g cps **10**⁻¹ 10⁰ 10¹ 0.0 0.5 1.0 0 50 100 **cps** 150 200 1.5 pCi/g pCi/g 15 5 25 0.0 0.5 1.0 1.5 50 100 150 200 250

# 299-E33-338 (C-3391) Combination Plot



299-E33-338 (C-3391) Combination Plot

# 299-E33-338 (C-3391) Combination Plot











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