FINAL

Light Nonaqueous-Phase Liquid Weathering at Various Fuel Release Sites



Prepared For

Air Force Center for Environmental Excellence Technology Transfer Division Brooks Air Force Base San Antonio, Texas

September 1999

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Prepared for:

AIR FORCE CENTER FOR ENVIRONMENTAL EXCELLENCE TECHNOLOGY TRANSFER DIVISION BROOKS AIR FORCE BASE SAN ANTONIO, TEXAS

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ACRONYMS AND ABBREVIATIONS

%/yr	Percent per year
°C	Degrees Celsius
AD Little	Arthur D. Little
AFB	Air Force Base
AFCEE/ERT	Air Force Center for Environmental Excellence/Technology
	Transfer Division
AST	Aboveground storage tank
ASTM	American Society for Testing and Materials
bgs	Below ground surface
BTEX	Benzene, toluene, ethylbenzene, and xylenes
DFSP	Defense Fuel Supply Point
DoD	Department of Defense
EAL	Evergreen Analytical Laboratory
GC/FID	Gas chromatography/flame ionization detection
GC/PID	Gas chromatography/photoionization detector
JP-4	Jet Petroleum No. 4
JP-5	Jet Petroleum No. 5
JP-8	Jet Petroleum No. 8
K _{fw}	Fuel/water partitioning coefficient
LNAPL	Light, nonaqueous-phase liquid
MCAS	Marine Corps Air Station
MCL	Maximum contaminant levels
mg/kg	Milligrams per kilogram
mg/L	Milligrams per liter
mg/mL	Milligrams per milliliter
µg/L	Micrograms per liter
µg/mL	Micrograms per milliliter
NAS	Naval Air Station
NRMRL	National Risk Management Research Laboratory
PAH	Polynuclear aromatic hydrocarbons
Parsons ES	Parsons Engineering Science, Inc.
R ²	Coefficient of determination
RNA	Remediation by natural attenuation
SVE	Soil vapor extraction
TMB	Trimethylbenzenes
US	United States
USAF	US Air Force
USEPA	US Environmental Protection Agency
USN	US Navy
UST	Underground storage tank
vol%	Percent by volume
wt%	Percent by weight

SECTION 1

INTRODUCTION

This report, prepared by Parsons Engineering Science, Inc. (Parsons ES), presents the results of work completed to evaluate natural weathering of light nonaqueous-phase liquids (LNAPLs) resulting from petroleum releases to the subsurface environment. As part of the natural attenuation demonstration project (Contract No. F41624-92-D-8036, Delivery Order 25), the Air Force Center for Environmental Excellence, Technology Transfer Division (AFCEE/ERT) contracted with Parsons ES to perform this fuels weathering study. Of particular interest for this study is the weathering or natural depletion of benzene, toluene, ethylbenzene, and xylenes (BTEX) from free-phase product (i.e., mobile LNAPL) following a fuel release. The BTEX compounds typically are identified as fuel hydrocarbon contaminants of concern because of their solubility and resulting mobility in soil and groundwater, and their relative toxicity (especially for benzene). Of primary interest for the study were jet petroleum no. 4 (JP-4) and gasoline release sites because they are the most common fuels on Air Force installations and have high initial BTEX fractions. However, spill sites with lower initial BTEX fractions, such as JP-5 and JP-8, also were evaluated.

1.1 PROJECT SCOPE AND OBJECTIVES

At many government and commercial sites, large-volume environmental releases of jet fuel or gasoline have contaminated and continue to contaminate soil and groundwater systems. Primary sources of large-volume fuel releases include fuel handling and storage activities associated with aboveground storage tanks (ASTs), underground storage tanks (USTs), fuel pumphouses, fuel hydrant systems, oil/water separators, and fuel pipelines. Uncontrolled catastrophic or chronic releases from such a source can result in large volumes of fuel being released to the subsurface. In the subsurface, the LNAPL often is present both as residual and mobile contamination. Residual LNAPL is defined as the LNAPL that is trapped in the aquifer by the processes of cohesion and capillarity, and therefore, will not flow within the aquifer or from the aquifer matrix into a groundwater well under the influence of gravity. Mobile LNAPL is defined as LNAPL that is free to flow in the aquifer and will flow from the aquifer matrix into a well under the influence of gravity.

The purpose of this study is to improve the scientific basis of and defensibility for determining natural LNAPL weathering rates (i.e., contaminant source-term reduction rates) as a component of remediation by natural attenuation (RNA). Currently, little information is available regarding rates of natural weathering of the BTEX components from mobile fuel LNAPLs. As a result, the rate of reduction of the contaminant source term in groundwater models often is left to professional judgment. The use of overly conservative LNAPL weathering rates to evaluate contaminant fate and transport and the

suitability of RNA as a remedial alternative can extend the estimated timeframe for longterm monitoring and affect the estimated cost-effectiveness and administrative feasibility of implementing RNA. Conversely, overestimation of weathering rates can lead to an overly optimistic forecast of RNA performance.

The primary objective of this fuel weathering study was to document a range of BTEX weathering rates for the mobile LNAPL fraction based on data collected from sites with documented mobile LNAPL plumes with known release dates. In addition, rates of naphthalene and methylnaphthalenes weathering from mobile LNAPLs were evaluated. Secondary objectives of this study included an evaluation of the degree of contaminant partitioning of BTEX from mobile LNAPL to groundwater, and comparison of weathering effects on the mobile LNAPL fraction and on residual LNAPL present in capillary fringe soils. The following tasks were completed to meet these objectives:

- A literature search to assess existing information regarding weathering of LNAPLs;
- Selection of eight primary sites where the time of release is generally known and free-phase jet fuel or gasoline remain *in situ*;
- Sampling of soil, groundwater, and free-phase LNAPLs at the primary sites;
- Evaluation of data obtained from the eight primary sites, as well as data from four secondary sites, to assess contaminant concentrations in site media in relation to such factors as age of the fuel release, fuel type, and site soils and hydrogeology.

This technical report summarizes the findings of the literature review and provides an assessment of site-specific mobile LNAPL weathering rates.

1.2 REPORT ORGANIZATION

This technical report consists of seven sections, including this introduction, and three appendices. Section 2 presents pertinent background information and findings from the literature review. Section 3 presents site selection criteria and a listing of the sites selected for the study. Section 4 summarizes the procedures used for collection and analysis of the site data. Section 5 summarizes the analytical results and presents the results of the LNAPL weathering data analysis. Section 6 presents conclusions based on the study results, and Section 7 lists the references used in preparing this document. Appendix A provides a copy of the original work plan and site addenda. Appendix B provides the analytical data for LNAPL, soil, and groundwater samples collected at the study sites. Appendix C provides calculations from the data analysis.

SECTION 2

LITERATURE REVIEW

The purpose of the literature review was to compile and summarize available technical literature on natural weathering of the BTEX fraction of fuel LNAPLs released to the subsurface environment. Specifically, the literature search attempted to answer the following question:

Is there sufficient scientific information available regarding *in situ* weathering rates for BTEX in mobile LNAPL to refine modeling assumptions used to predict reductions in the contaminant source term at fuel-hydrocarbon-contaminated sites?

Parsons ES experience with the AFCEE natural attenuation demonstration project had indicated a lack of scientifically defensible information regarding BTEX weathering rates for mobile LNAPL. As a result of this data gap, weathering rates used to simulate BTEX source terms in fate and transport modeling generally have been based on professional judgment and consideration of site-specific conditions. Typically, total BTEX depletion rates (i.e., contaminant source-term reduction rates) between 1 and 15 percent per year Site-specific conditions such as groundwater depth, (%/yr) have been assumed. precipitation, composition of the soil/aquifer matrix, and site location also have been considered in determining whether BTEX removal from mobile LNAPL is likely to be hindered or enhanced. For example, a total BTEX depletion rate between 10 and 15 %/yr might be assumed for a high-precipitation, high-soil-permeability, shallow groundwater site in Florida; whereas a depletion rate of 1 to 3 %/yr total BTEX might be assumed for a low-precipitation, low-permeability, deep groundwater site in Montana. Where site conditions do not appear to excessively hinder or enhance BTEX depletion from mobile LNAPL, a default value of 5 %/yr often was used. No scientific studies were known to exist that could support the validity of these assumptions. Therefore, a more formal literature search was included as part of this study.

A preliminary review of the literature as summarized in the work plan (Appendix A) indicated that there was not sufficient information regarding BTEX weathering rates in mobile LNAPL, and that field studies of fuel-contaminated sites would be appropriate. Since the time of the initial literature review, additional information regarding weathering processes and rates of fuel weathering has been gathered; however, the general findings of the preliminary literature review remain the same. No field studies were identified that scientifically evaluate naturally occurring BTEX reductions (weathering) within mobile LNAPLs at fuel release sites. A discussion of fuels composition and a review of the fuel weathering literature is provided in the following subsections as background information for the fuel weathering study results and conclusions presented in Sections 5 and 6, respectively.

2.1 GASOLINE AND JET FUEL USE AND COMPOSITION

2.1.1 Fuel Use and History

Gasoline, diesel fuel, and jet fuel represent the primary fuel types used at United States (US) military installations for powering vehicles, equipment, and aircraft. Large-volume storage and handling of these petroleum products has resulted in widespread environmental contamination of soil and groundwater. However, BTEX contamination of soil and groundwater at US military installations has resulted primarily from uncontrolled releases of gasoline and jet fuel.

A variety of jet fuels have been used for powering US military aircraft turbine (jet) engines since the beginning of jet flight in the 1940s. Since the 1950s, JP-4 and JP-5 represent the primary fuels used by the US Air Force (USAF) and US Navy (USN), respectively. More recently, the USAF has converted from JP-4 to JP-8 because of the lower volatility and lower explosion/fire hazard of JP-8. In 1979, USAF installations in Great Britain replaced JP-4 with JP-8 (Martel, 1987), and in 1993/1994, USAF installations in the continental US converted to JP-8. Therefore, most JP-8 fuel releases that have contaminated soil and groundwater at USAF installations are no more than 5 years old. While other jet fuels have been used by the US military, their use and storage has been limited, resulting in far less environmental site contamination from these less common fuels.

2.1.2 Hydrocarbon Composition of Gasoline and Jet Fuel

Gasoline and jet fuel are refined petroleum products derived from crude oil. Crude oil, a degradation product of organic material (e.g., prehistoric animal and plant matter) is a complex mixture primarily composed of hydrocarbons, which are compounds consisting solely of carbon and hydrogen. Measured by weight, carbon and hydrogen represent at least 95 percent of the elements present in crude oil (Neumann *et al.*, 1981). In comparison, hydrocarbon concentrations in refined petroleum products such as gasoline, diesel fuel, and kerosene are even higher than in crude oil, because non-hydrocarbon compounds (which contain sulfur, nitrogen, oxygen, or trace metals) are destroyed or removed during the refining process (Owen and Corey, 1990).

2.1.2.1 Distillation

The hydrocarbon composition of gasoline and other petroleum products derived from crude oil is largely determined during the refining process known as distillation. Distillation is a process whereby the crude oil is heated/boiled, and fractions of the crude oil are separated based on boiling point. During distillation, the more volatile, shorter-chain, lower-molecular-weight hydrocarbons are initially removed at relatively low boiling points, and the less volatile, longer-chain, heavy-molecular-weight hydrocarbons are subsequently removed at higher boiling temperatures. Distillation utilizes the relationship between boiling point and hydrocarbon molecular weight to separate crude oil into useable fractions, or "cuts," for further refinement into petroleum end products. Because hydrocarbon molecular weight is dictated by the number of carbon atoms present, it is possible to generally classify these distillation cuts by their predominant carbon-atom ranges (American Society for Testing and Materials [ASTM], 1995):

- Gasoline C₄ to C₁₂ hydrocarbons;
- Kerosene and jet fuels C₁₁ to C₁₃ hydrocarbons;
- Diesel fuel and light fuel oils C₁₀ to C₂₀ hydrocarbons;
- Heavy fuel oils C₁₉ to C₂₅ hydrocarbons; and
- Motor oils and other lubricating oils C₂₀ to C₄₅ hydrocarbons.

2.1.2.2 Wide-Cut and Kerosene-Based Jet Fuels

Jet fuels commonly used by the Air Force and Navy can generally be separated into two categories: "wide-cut" fuels and "kerosene-based" fuels (Martel, 1987). JP-4 is created by taking a "wide cut" of the distillate to include both the gasoline and kerosene fractions. JP-4 typically is composed of approximately 50 to 60 percent gasoline and 40 to 50 percent kerosene (Martel, 1987). This large percentage of gasoline imparts increased volatility to JP-4. On the other hand, JP-5 and JP-8 are kerosene-based fuels that contain relatively less volatile, longer-chain hydrocarbons.

2.1.2.3 Hydrocarbon Structure

The three most prevalent types of hydrocarbons in crude oil and refined petroleum products, based on their chemical structure, are alkanes, alkenes, and aromatics. Figure 2.1 illustrates the structures of these three types of hydrocarbon compounds.

Alkanes, or paraffins, are hydrocarbon chains characterized by single molecular bonds between the carbon atoms and "saturation" of all remaining bonding sites by hydrogen atoms. For this reason, alkanes also are referred to as saturates. Based on their structure, alkanes can be further separated into n-alkanes (straight-chain alkanes), isoalkanes (branched-chain alkanes), and cycloalkanes (alkane rings) (Figure 2.1). Isoalkanes and cycloalkanes are commonly referred to as isoparaffins and naphthenes, respectively. In general, alkanes are the most abundant hydrocarbons in crude oil and gasoline. Alkanes represent 55 to 75 percent of all hydrocarbons in crude oil (Metcalf & Eddy [M&E], 1993). A compilation of analytical results from 10 gasoline samples indicated that alkanes make up approximately 55 percent by weight (wt%) of the hydrocarbons in gasoline (Nakles *et al.*, 1996).

Alkenes, or olefins, are characterized as hydrocarbon chains that are not saturated with hydrogen atoms, and as a result, contain one or more double bonds between carbon atoms (Figure 2.1). While alkenes are typically at trace levels in crude oil, their concentration in petroleum products is often increased by the refining process. Nakles *et al.* (1996) reported the concentration of alkenes in gasoline as approximately 11 wt%.



Aromatic hydrocarbons also are unsaturated, and are characterized by their six-carbon ring structure. As illustrated on Figure 2.1, the six-carbon-ring aromatic structure has alternating single and double bonds. The simplest aromatic compound is benzene (C_6H_6), which is composed of a single aromatic ring (monoaromatic). Benzene and its chemical derivatives are common in volatile fuels such as gasoline and JP-4. Other aromatic hydrocarbons more typical of heavier, less volatile fuel types are naphthalene, a two-ring aromatic (diaromatic) and phenanthrene, a three-ring aromatic (Figure 2.1). Three-ring and higher aromatics are often referred to as polynuclear aromatic hydrocarbons (PAHs). Nakles *et al.* (1996) reported that aromatics make up approximately 33 wt% of the hydrocarbons in gasoline. However, in jet fuels, the aromatic content is limited to no more than 25 percent by volume (vol%) to improve combustion performance and minimize solvent effects (Martel, 1987).

A common feature of petroleum-derived aromatics is the presence of an "alkyl" group in place of a hydrogen atom on the six-carbon ring. Common alkyl groups are the methyl group and the ethyl group. The methyl group is composed of 1 carbon atom and 3 hydrogen atoms (CH₃). The ethyl group is composed of 2 carbon atoms and 5 hydrogen atoms (CH₂CH₃). In the ethyl group a CH₂ unit is "sandwiched" between the aromatic ring and a terminal CH₃, or methyl group. Toluene, ethylbenzene, and *ortho-*, *meta-*, and *para-*xylenes all are single-ring aromatic compounds where one or two hydrocarbon atoms have been replaced by one or two of these alkyl groups. As illustrated on Figure 2.2, toluene is simply a benzene ring in which one of the hydrogen atoms has been replaced with a methyl group. In ethylbenzene, the hydrogen atom is replaced by an ethyl group. In the xylene isomers, two hydrogen atoms are replaced by two methyl groups. The prefixes "*ortho-*," "*meta-*," and "*para-*" refer to the position of the methyl groups on the benzene ring.

2.1.2.4 BTEX Composition

For this study, the weathering of the low-molecular-weight BTEX compounds is of particular concern because of their mobility within the environment and, in the case of benzene, its relatively high toxicity. To adequately characterize the degree or rate of BTEX weathering from mobile LNAPLs, the initial concentration of BTEX compounds within the original fuel is needed. Unfortunately, the exact concentration of BTEX compounds in a gasoline or jet fuel that is released to the environment typically is not known and can only be estimated based on compositional studies of fresh fuels. Cline *et al.* (1991) have noted that the specific composition of gasoline will vary depending on the source of the petroleum, the production method, the end use location, and the season of the year. Similarly, there is considerable variability in jet fuel composition based upon the distillate cuts of gasoline and kerosene from which the jet fuel is blended. Therefore, compositional studies of fresh fuels.

Figure 2.3 illustrates the ranges of benzene, toluene, ethylbenzene, total xylenes, and total BTEX concentrations in fresh JP-4, JP-5, and JP-8 jet fuels and fresh gasoline. As is evident from this figure, fresh gasoline has the highest mass fraction (in wt%) of BTEX compounds, followed by JP-4, JP-8, and JP-5 jet fuels.





JP-5







GASOLINE



2.1.2.4.1 Gasoline

For gasoline, there is a large disparity between the minimum and maximum BTEX mass fraction values presented by Potter (1988), Arthur D. Little (AD Little, 1987), and Sigsby et al. (1987) (Figure 2.3). Reported maximum concentrations for benzene and toluene are approximately five times the minimum concentrations. The total BTEX maximum concentration is nearly four times that of the reported minimum concentration (38.5 wt% versus 10.4 wt%). Similar disparities were evident in analytical results compiled by the Alberta Research Council (1993) for 124 gasoline samples. For benzene, the minimum and maximum reported concentrations were 0.34 wt% and 5.62 wt%, respectively, and the average benzene concentration was 1.86 wt%. For total BTEX, the minimum and maximum reported concentrations were 4.1 wt% and 45.4 wt%, respectively, and the average total BTEX concentration was 20.7 wt%. The BTEX mass fraction values for gasoline reported by Ghassemi et al. (1984) are somewhat lower than these average concentrations (Figure 2.3). Therefore, using the fresh-product values presented by Ghassemi et al. (1984) along with observed in situ BTEX concentrations to predict BTEX reductions in gasoline LNAPL would be more conservative than using average concentrations from the other studies cited above.

2.1.2.4.2 JP-4

Figure 2.3 presents JP-4 BTEX concentrations reported by Hughes *et al.* (1984) and Smith *et al.* (1981). In the Hughes *et al.* (1984) study, 54 JP-4 samples were analyzed by gas chromatography/flame ionization detection (GC/FID) analysis, and results were reported in milligrams per milliliter (mg/mL). Mass fraction (wt%) results shown in Figure 2.3 were obtained using a maximum density value of 802 mg/mL for JP-4 at 15 degrees Celsius (°C) (Martel, 1987). Mass fraction results presented by Smith *et al.* (1981) were obtained from JP-4 samples analyzed by gas chromatography/mass spectrometry (GC/MS). As shown on Figure 2.3, BTEX mass fraction results presented by Smith *et al.* (1981) are slightly lower than those presented by Hughes *et al.* (1984), and therefore represent more conservative initial values for estimating mass fraction BTEX reductions in JP-4 mobile LNAPL.

2.1.2.4.3 JP-8

BTEX mass fraction results for JP-8 as determined by Smith *et al.* (1981) and Mayfield (1996) also are presented in Figure 2.3. Mass fraction BTEX results presented by Smith *et al.* (1981) were obtained by GC/MS analysis. In the Mayfield (1996) study, 63 JP-8 samples were analyzed by GC/MS and results were presented in milligrams per liter (mg/L). Average mass fraction values shown on Figure 2.3 were obtained using a maximum density value of 840 mg/L for JP-8 at 15 °C (Martel, 1987). The disparity in BTEX concentrations between these two studies is significant. BTEX concentrations presented by Smith *et al.* (1981) are approximately one-tenth the concentrations presented by Mayfield (1996). The reason for this disparity is not known, but may have resulted from changes in JP-8 manufacturing methods or specifications between 1980 and 1996. The Mayfield (1996) study represents a more contemporary and comprehensive review of JP-8 composition, and likely better represents JP-8 jet fuel used in the 1990s. Use of the lower JP-8 BTEX concentrations, as determined by Smith *et al.* (1981), for predicting mass fraction reductions in JP-8 LNAPL while extremely conservative, may not be tenable if BTEX concentrations in site LNAPL exceed these values.

2.1.2.4.4 JP-5

Limited BTEX compositional data were available for JP-5. Results shown on Figure 2.3 are from one fresh JP-5 sample analyzed by Hughes *et al.* (1984) and from one fresh JP-5 sample obtained from Beaufort Marine Corps Air Station (MCAS) in Beaufort, South Carolina and analyzed during this study (see Sections 3 and 4). Total BTEX concentrations in both samples were well below 1 wt%. While these data are limited, the relatively insignificant concentrations of BTEX in fresh JP-5 are likely to limit environmental threats from JP-5 releases, especially in comparison to gasoline and JP-4 releases.

2.2 SUBSURFACE LNAPL DISTRIBUTION AND RECOVERY

Characterization of fuel release sites where LNAPL is present in the subsurface is often difficult because of the complex transport parameters and mechanisms associated with LNAPL and separate-phase contamination. Subsurface migration and distribution of LNAPLs, as well as LNAPL persistence and strength as a source of soil and groundwater contamination, is governed by petroleum release factors, soil and aquifer properties, and LNAPL characteristics (Mercer and Cohen, 1990; Pfannkuch, 1984). The primary petroleum release factors influencing migration and distribution are the volume of the release, the release rate, and the area of the release. Influential subsurface properties include, but are not limited to, lithology, soil permeability, pore size distributions, porous media wettability, fluid pressure at and above the water table, and water table fluctuations. Characteristics of the LNAPL itself, such as density and viscosity, also influence subsurface distribution and migration (Newell *et al.*, 1995). Once LNAPL is released to the subsurface environment, a defined interface between the LNAPL and soil, air, and water, in the subsurface, typically does not exist (Newell, *et al.*, 1995).

2.2.1 LNAPLs in the Unsaturated Zone

In the unsaturated, or vadose, zone, movement of LNAPL initially is controlled by its fluid density and viscosity as it moves downward under the force of gravity (Hunt *et al.*, 1988). Subsurface heterogeneities may cause lateral spreading and trap lenses of LNAPL above layers of lesser permeability soils during downward migration. Interfacial forces (e.g., surface tension between soil-air and soil-water and the LNAPL) and soil capillary forces can cause residual masses of the LNAPL to become trapped in soil pores as ganglia and lenses (Hunt *et al.*, 1988; Powers *et al.*, 1991; Seagren *et al.*, 1993). Because this residual LNAPL can remain trapped in the unsaturated zone for an indefinite length of time, on the order of decades to centuries (Hunt *et al.*, 1988), it becomes a long-term source of groundwater contamination via infiltrating precipitation or a rising water table (Abriola and Pinder, 1985; Seagren *et al.*, 1993). If the volume of the fuel release is relatively small and the depth to groundwater is great, the entire LNAPL volume may be retained in soil pores as residual LNAPL and not reach the water table.

2.2.2 LNAPLs in the Saturated Zone

If the fuel release is of sufficient volume to reach the water table, the mobile LNAPL fraction will spread laterally and form a floating pool at the capillary fringe above the water table (Pfannkuch, 1984; Voudrias *et al.*, 1994;). With water table fluctuations caused by seasonal recharge and depression, or by local pumping, an LNAPL smear zone

can form over the range of water table fluctuation (McKee *et al.*, 1972; Dietz, 1980; Schwille, 1984; Voudrias *et al.*, 1994;). Like the residual LNAPL in the unsaturated zone, the LNAPL smear zone can be highly variable, with residual LNAPL present as discrete ganglia to fully saturated mobile LNAPL lenses (Hunt *et al.*, 1988). Also, the immiscible nature of LNAPLs can cause discrete LNAPL "blobs" to become trapped in groundwater (Yang *et al.*, 1995) and to be only partially remobilized during rising water table conditions (Hunt *et al.*, 1988).

2.2.3 LNAPL Recovery

Mobile LNAPL, or free product, recovery at fuel release sites often is difficult, expensive, and only marginally effective (Farr *et al.*, 1996). Recovery of free-phase fuel has proven to be difficult because of the complex interaction of hydrogeologic and LNAPL characteristics that tend to retain the mobile LNAPL. Typically, less than 25 to 35 percent of the mobile LNAPL that has spread out on the water table is recoverable (Farr *et al.*, 1996), with significant retention occurring in the capillary fringe during product recovery efforts (Testa and Paczkowski, 1989). Residual LNAPL retained in the unsaturated zone and immobile LNAPL blobs associated with the capillary fringe or submerged below the water table are unrecoverable by conventional means (Testa and Paczkowski, 1989; AFCEE, 1998). Nevertheless, regulatory requirements specifying removal of free product to the "degree practicable" traditionally have been interpreted as LNAPL removal to no more than a sheen (Lundy, 1997). This regulatory expectation, combined with the limited effectiveness of conventional LNAPL recovery methods, have tended to drive up remediation costs as remediation durations are extended with little product recovery or risk reduction.

More recently a risk-based remediation approach to free product recovery has been proposed (Farr *et al.*, 1996; Lundy, 1997; AFCEE, 1998). Under this approach, site-specific environmental and health risks posed by the mobile LNAPL are evaluated in combination with the feasibility, effectiveness, and cost of free product recovery. At some sites it may be possible to demonstrate that the cost of free product recovery is not justified by a commensurate reduction in site risk. In these cases, it may be more appropriate to model the expected limit of plume migration and to expand the long-term monitoring well network to accommodate plume expansion rather than trying to limit expansion through source reduction (AFCEE, 1998).

2.3 LNAPL WEATHERING MECHANISMS

The term "weathering," or attenuation, refers to the combined effects of natural destructive and non-destructive processes to reduce the persistence, mobility, mass, and toxicity of the fuel contaminant in the environment. The majority of information currently available regarding subsurface fuel contamination examines the effects of specific natural attenuation mechanisms such as dissolution, biodegradation, and volatilization as they apply to soil and groundwater contamination. While the literature has focused on these mechanisms as they apply to attenuation of chemicals sorbed to soil and dissolved in groundwater, mobile LNAPL weathering also is a function of these processes.

The primary mechanisms acting to reduce the strength of a LNAPL source are dissolution, volatilization, and biodegradation. These mechanisms are influenced by physical and chemical properties of the chemical compounds in the source product, as well

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as by physical, chemical, and biological properties of the soil and groundwater system. An illustration of these weathering mechanisms is shown on Figure 2.4.

2.3.1 Dissolution

Dissolution is the dissolving of chemical substances from a residual or mobile NAPL into percolating precipitation water and/or the groundwater. At gasoline and jet fuel release sites, dissolution or partitioning of the BTEX compounds from the LNAPL into groundwater represents the most significant source of groundwater contamination and likely the most significant mechanism of BTEX depletion in mobile LNAPLs (Huntley and Beckett, 1997). LNAPL dissolution is governed by the characteristics of the aquifer matrix (including effective porosity and groundwater velocity), physical properties of the LNAPL (e.g., surface area of the LNAPL in contact with groundwater), and characteristics of the specific LNAPL contaminant (e.g., effective water solubility) (Wiedemeier *et al.*, 1995).

2.3.1.1 Effective Water Solubility of BTEX

Solubility of a substance in water is defined as the mass of the substance that will dissolve in a unit volume of water (typically expressed in mg/L). According to Montgomery (1996), the water solubility of a compound is arguably the most important factor in determining the fate and transport of the compound in the subsurface. The aromatic compounds are among the most mobile of dissolved fuel contaminants at gasoline and jet fuel release sites because of their relatively high water solubilities. Singlering BTEX compounds are significantly more water soluble than the two-ring naphthalenes, as shown in Table 2.1. Pure-phase water solubilities for the BTEX compounds range between 157 and 1,750 mg/L. Based on these values, benzene is the most water-soluble of the BTEX compounds, followed by toluene, ethylbenzene, and xylenes. In various soil column water-flushing experiments (Borden and Kao 1992; Rixey et al., 1992; Voudrias et al., 1994), benzene, toluene, ethylbenzene and xylene were flushed from soil columns in order of decreasing solubility. With increased compound solubility, there is increased dissolution flux, indicating compound depletion or weathering in a fuel LNAPL will be more rapid for the more water-soluble compounds like benzene.

The dissolution flux of compounds in fuel LNAPLs also is influenced by the compound's concentration in the LNAPL. In fresh JP-4 jet fuel, benzene comprises approximately 0.50 wt% of the fuel, and in gasoline, benzene typically constitutes no more than 4.5 wt% of the fuel (Figure 2.3). Similarly, toluene, ethylbenzene, and xylenes typically are present in gasoline and jet fuels at concentrations significantly less than 10 wt%. Therefore, the dissolution flux of these compounds is significantly less than if they were present in pure phase. As shown in Table 2.1, the actual concentrations of aromatic compounds in water resulting from fuel/water contact, are 1 to 2 orders of magnitude less than their respective pure-phase water solubilities. The BTEX compounds are more soluble in fuel than in water, and tend to remain in the fuel.

This decrease in dissolution flux resulting from an equilibrium relationship between the aqueous phase and the multicomponent LNAPL has been described by Raoult's Law. Raoult's Law is based on a thermodynamic theory of multicomponent solutions and is typically valid for compounds that are present in relatively low concentrations within the solution (M&E, 1993). Using Raoult's Law, the effective water solubility of a compound



TABLE 2.1 DISSOLVED CONCENTRATIONS OF AROMATIC FUEL HYDROCARBONS FUEL WEATHERING STUDY

Concentrations in Water				
	Pure-Phase	in Contact with Fuel ^{c/}		
	Water	("effective	solubilities")	
	Solubility ^{a/}	JP-4 ^{d/}	Gasoline ^{e/}	MCL ^{f/}
Compound	(mg/L) ^{b/}	(mg/L)	(mg/L)	(mg/L)
Benzene	1,750	9.82	58.7	0.005
Toluene	524	8.49	33.4	1.0
Ethylbenzene	187	0.67	4.3	0.7
o-Xylene	167	1.21	6.9	$\mathbf{NA}^{\mathbf{g}/}$
m-Xylene	157	2.01	11.0	NA
p-Xylene	180	0.41	4.4	NA
Xylenes	168	3.63	22.3	10
Trimethylbenzenes	97.7 ^{h/}	0.87	$1.1^{i\prime}$	NA
Naphthalene	22	0.39	^{j/}	$0.02^{k/}$
Methylnaphthalenes	25.4 ^{1/}	0.24		NA

^{a/} Solubilities at 25°C (Montgomery, 1996).

 $^{b/}$ mg/L = milligrams per liter.

^{c/} Fuel to water ratio 1:10.

^{d/} Smith et al., 1981.

e/ American Petroleum Institute (1985).

 $^{\rm f/}$ MCL = maximum contaminant level (USEPA, 1996).

g' NA = not applicable.

 $^{\rm h\prime}$ Solubility for 1,3,5-trimethylbenzene.

^{i/} Value for 1,2,4-trimethylbenzene.

j' - - - = not available.

^{k/} Health advisory value for 70-kilogram adult, lifetime exposure.

^{1/} Value for 2-methylnaphthalene.

 (C_i) can be predicted by the product of the water solubility of the pure compound (S_i) and the mole fraction of the compound in the LNAPL (X_i) :

$$C_i = X_i S_i$$
 eq. 2.1

In order to use Raoult's Law to estimate effective solubilities, the mole fraction or molecular percent of a compound in an LNAPL or a fresh fuel must be known. However, analytical results for fuel and LNAPL components are typically reported in wt% or vol%, not molecular percent. Using this equation, and estimates of molecular percent for BTEX in gasoline, semiquantitative estimates of effective water solubility have been determined for gasoline (M&E, 1993). However, a 20 to 30 percent disparity was apparent between predicted values and measured water concentrations resulting from gasoline contact (Table 2.1). This disparity is thought to have resulted from inherent uncertainties with the predictions of mole fraction in multicomponent fuels (M&E, 1993).

Despite the difficulty with using Raoult's Law directly, it is useful in illustrating the relationship between compound concentration and compound solubility in evaluating effective solubility at fuel-contaminated sites. As shown in Table 2.1, the pure-phase water solubility of toluene is less than one-third the pure-phase water solubility of benzene, yet the effective solubility of toluene when water is in contact with JP-4 or gasoline is much closer to the effective water solubility of benzene. This results from the higher concentration or mole fraction of toluene in the fuel compared to that of benzene (Figure 2.3). Based on this relationship and the effective solubility values presented in Table 2.1, it appears that ethylbenzene and xylenes are not sufficiently present in JP-4, or sufficiently soluble in water, to consistently exceed regulatory maximum contaminant levels (MCLs) at JP-4 contaminated sites. At sites contaminated by lower-BTEX-content fuels (i.e., JP-5 or JP-8), there is even less dissolution flux of BTEX compounds into site groundwater.

2.3.1.2 Fuel/Water Partitioning Coefficients

Fuel/water partitioning coefficients offer another method for evaluating fuel contaminant dissolution from fuel LNAPLs into water. The fuel/water partitioning coefficient (K_{fw}) is a dimensionless constant defined as the ratio of a compound in the fuel (C_f) to the compound's equilibrium concentration in water in contact with the fuel (C_w):

$$K_{fw} = C_f / C_w \qquad eq. 2.2$$

Fuel/water partitioning coefficients demonstrate the relationship between water solubility of a compound and abundance of the compound in a fuel. Table 2.2 provides K_{fw} values for constituents of JP-4 and gasoline. As shown, K_{fw} values for BTEX compounds in JP-4 are significantly higher than gasoline values. Relative to gasoline, a larger portion of the BTEX compounds have a tendency to stay in JP-4, resulting in lower equilibrium concentrations in water and higher K_{fw} values.

2.3.1.3 Equilibrium versus Nonequilibrium

Significant debate appears in the literature regarding the applicability of equilibrium conditions when assessing dissolution (Hayden *et al.*, 1992; Seagren *et al.*, 1993; Voudrias *et al.*, 1994; and Yang *et al.*, 1995). Use of Raoult's Law and K_{fw} values to

assess contaminant dissolution assumes that equilibrium conditions exist. Often equilibrium is assumed in order to simplify dissolution calculations in subsurface flow models (Seagren *et al.*, 1993). For example, once the concentration of a contaminant is known in one phase, equilibrium partitioning is used to calculate the concentration in the other phase at the same location. However, serious errors in prediction of contaminant reduction have occurred when equilibrium assumptions are used in groundwater modeling (Powers *et al.*, 1991). As noted by Bruce *et al.* (1991), dissolved concentrations of the BTEX compounds rarely exceed 20 percent of the calculated equilibrium concentration, unless LNAPL is present as a sheen or colloids. One hypothesis suggests that the lack of equilibrium concentrations occurs from less thorough mixing of the fuel (LNAPL) and water in the field as compared to the laboratory (Bruce, 1993). Groundwater sampling often is performed over several feet of saturated soil, and only the upper few inches of the soil column is in contact with LNAPL. Significant dilution of dissolved hydrocarbons will result in concentrations far less than theoretical equilibrium values.

As discussed by Seagren *et al.* (1993), if equilibrium conditions exist in the field, the dissolution rate becomes a function of advection (groundwater transport of the contaminant away from the source area) and/or biodegradation. Under this scenario, the dissolution rate is enhanced by contaminant removal from the interphase (LNAPL and aqueous) boundary, thereby increasing the dissolution flux. However, if nonequilibrium conditions exist (e.g., the groundwater concentration of benzene is significantly less than its effective solubility [Table 2.1]), the dissolution rate cannot be enhanced by advection or biodegradation that further reduces the aqueous concentration.

The equilibrium assumption as it applies to LNAPLs and groundwater contaminant concentrations has yet to be adequately demonstrated (Powers *et al.*, 1991). Also, while generalizations can be made, no quantitative criteria exist for determining when equilibrium or nonequilibrium conditions exist (Seagren *et al.*, 1993). Because it has been argued that the rate of dissolution is a significant limiting factor in remediation of residual LNAPLs (Yang *et al.*, 1995), it can also be presumed that dissolution is rate-limiting (i.e., a predominant mechanism) for mobile LNAPL weathering.

2.3.2 Volatilization

Volatilization, or evaporation, is the loss of a compound from a liquid or solid state to a vapor state. For surface spills, important factors affecting volatilization include temperature, vapor pressure of the constituents, and wind speed. For subsurface releases, temperature and vapor pressure are important, but volatilization requires diffusion through a porous medium; therefore, soil moisture and soil porosity also are important (LaGrega *et al.*, 1994). Subsurface rates of volatilization are directly proportional to soil porosity, pore size distribution, and temperature, but inversely proportional to volumetric moisture content (Hillel, 1980).

Relative volatility of compounds at equilibrium conditions can be compared by an air/water partitioning coefficient known as Henry's Law Constant (H). Henry's Law states that under equilibrium conditions, the partial pressure of a gas (i.e., volatile chemical) (P_g) above a liquid is proportional to the concentration of the chemical in the liquid (C_L):

TABLE 2.2 FUEL/WATER PARTITIONING COEFFICIENTS FOR BTEX AND TMBs FUEL WEATHERING STUDY

	Fuel/Water Partitioning Coefficient (K _{fw})		
Compound	JP-4 Jet Fuel ^{a/}	Gasoline ^{b/}	Gasoline ^{c/}
Benzene	2,455	231	350
Toluene	2,754	895	1,250
Ethylbenzene	4,786	3,411	4,500
o-Xylene	7,079	3,162	3,630
m-Xylene	3,715	3,539	4,350
p-Xylene	7,586	2,961	4,350
1,2,3-Trimethylbenzene	$NA^{d/}$	NA	13,800
1,2,4-Trimethylbenzene	8,913	12,270	NA
1,3,5-Trimethylbenzene	NA	6,493	NA

Source: Wiedemeier, 1995.

^{a/} Smith et al., 1981.

^{b/} Bruce et al., 1991.

^{c/} Cline et al., 1991.

d' NA = not analyzed.

$$P_g = HC_L$$
 eq. 2.3

Henry's Law Constant values for BTEX, trimethylbenzenes (TMBs), and naphthalenes are listed in Table 2.3. As a general rule of thumb, compounds with Henry's Law Constants greater than 10^{-3} are considered very volatile (M&E, 1993). As shown in Table 2.3, the BTEX compounds and TMBs are more volatile than naphthalene and 2-methylnaphthalene. Generally speaking, compound volatility decreases with increasing carbon atoms. On a unit-carbon basis, the alkanes are more volatile than the aromatics (Nakles *et al.*, 1996).

As with dissolution, contaminant volatilization from a LNAPL is influenced by the concentration of the contaminant in the LNAPL. In other words a Raoult's Law expression similar to that discussed in Section 2.3.1.1 can be applied.

Enhanced volatilization using soil vapor extraction (SVE) techniques is commonly used for vadose zone cleanup at fuel-contaminated sites; however, no field studies were identified that evaluated "equilibrium" volatilization at sites having subsurface free-phase product. Volatilization is expected to be a significant weathering mechanism for petroleum products such as gasoline, JP-4, and JP-8. From a study on the fate of JP-8 in quiescent flask systems containing water and water/sediment mixtures, evaporation or volatilization from water was the major removal mechanism for low-molecular-weight, volatile hydrocarbons (Dean-Ross *et al.*, 1992). In the same study it was determined that the presence of sediment can sequester jet fuel and render it less susceptible to volatilization. Intuitively, greater contact between soil gas and residual LNAPL would result in greater mass loss rates due to volatilization than would be expected in soils saturated with mobile LNAPL.

2.3.3 Biodegradation

2.3.3.1 Residual LNAPL and Groundwater

Most of the literature pertaining to *in situ* biodegradation refers to residual LNAPL contaminants in soil and contaminants dissolved in groundwater. As mentioned in Section 2.3.1.3, dissolution appears to be a rate-limiting factor in weathering, especially as it relates to biodegradation. If equibrium conditions exist, biodegradation of dissolved petroleum contaminants will reduce aqueous contaminant concentrations, thereby enhancing dissolution rates by increasing mass transfer of soluble compounds from residual LNAPLs into groundwater (Seagren *et al.*, 1993; Yang *et al.*, 1995). As a result of this diffusion limitation, mass loss rates of dissolved contaminants from biodegradation appear initially to be between zero and first-order (Song *et al.*, 1990), and to decrease with time (Barker *et al.*, 1987).

The kinetics of biodegradation are complicated by the fact that biodegradation is compound-specific and is significantly affected by the geochemistry of the subsurface environment. Dean-Ross (1993) examined the fate of JP-4 jet fuel in subsurface soils and discovered that for the less volatile, higher-molecular-weight jet fuel components, biodegradation represented a significant mechanism for reducing soil contamination. Song *et al.* (1990) concluded that saturated compounds such as hexane generally are more easily biodegraded than the corresponding aromatic compounds. In a study by Barker *et al.* (1987), mass loss rates for aromatics in groundwater due to biodegradation were

TABLE 2.3 HENRY'S LAW CONSTANTS FOR BTEX, TMBs, AND NAPHTHALENES FUEL WEATHERING STUDY

	Henry's Law Constant
Compound	(atm-m ³ /mol)
Benzene	5.28E-03
Toluene	6.42E-03
Ethylbenzene	7.88E-03
o-Xylene	4.87E-03
m-Xylene	7.44E-03
p-Xylene	7.44E-03
1,2,3-Trimethylbenzene	3.18E-03
1,2,4-Trimethylbenzene	5.70E-03
1,3,5-Trimethylbenzene	6.73E-03
Naphthalene	7.34E-04
2-Methylnaphthalene	3.18E-04

Source: Montgomery, 1996.

Henry's Law Constant values at 25°C.

greatest for xylenes, followed by toluene, and benzene. Other factors playing an important role in contaminant biodegradation include availability of nutrients, availability of oxygen and other electron acceptors, and the interfacial area available for mass transfer to aqueous or gaseous phases (Yang *et al.*, 1995). For residual LNAPLs, the size of the LNAPL globules impacts biodegradation rates, with smaller globules resulting in greater interfacial area for mass transfer, and faster biodegradation rates (Yang *et al.*, 1995).

2.3.3.2 Mobile LNAPLs

No studies were identified that addressed intrinsic biodegradation of LNAPL pools. In addition, practical bioremediation of free-phase product has not been demonstrated (Newell *et al.*, 1995), most likely as a result of the following:

- Mobile LNAPLs represent a hostile environment for the survival of most soil microbes; and
- Requirements for microbial proliferation (e.g., nutrients, terminal electron acceptors, pH, moisture, osmotic potential) are difficult and may be impossible to deliver or maintain in the LNAPL pool (Huling and Weaver, 1991).

Consequently, effective bioremediation and tangible intrinsic biodegradation is likely to be limited to the periphery of the mobile LNAPL zone (i.e., residual LNAPL and aqueous phases).

2.4 OTHER FACTORS INFLUENCING LNAPL WEATHERING

In addition to the LNAPL weathering mechanisms discussed above, hydrocarbon layer thickness, groundwater velocity, soil/aquifer material, and distance from the source area are factors expected to impact BTEX depletion within mobile and residual LNAPLs.

2.4.1 Hydrocarbon Layer Thickness

The hydrocarbon layer thickness at the interface of the unsaturated and saturated zones is presumed to influence BTEX dissolution from the LNAPL (Huntley and Beckett, 1997). Dissolution modeling of a 10-centimeter thick LNAPL pool in fine sand indicated that the effective solubility of benzene could be reduced to approximately 0.001 mg/L in less than a year. However, modeling results for a 50-centimeter thick pool indicated it would take approximately 70 years to reach the same effective solubility (Huntley and Beckett, 1997). The larger the LNAPL pool thickness, the more slowly benzene is removed from the LNAPL pool.

It is important to note that LNAPL thickness measurements from groundwater monitoring wells are not indicative of LNAPL thicknesses in the formation (Blake and Hall, 1984; Hall *et al.*, 1984; Hughes *et al.*, 1988; Testa and Paczkowski, 1989; Farr *et al.*, 1990; Mercer and Cohen, 1990; Huntley *et al.*, 1994). Mercer and Cohen (1990) suggest that the measured LNAPL thickness in wells is typically 2 to 10 times greater than the LNAPL thickness in the formation. In addition, depiction of mobile LNAPL as a distinct layer present above the water capillary fringe has been challenged (Farr *et al.*, 1990; Lenhard and Parker, 1990). It has been suggested that hydrocarbon-saturated soil layers do not exist at sites with measurable LNAPL; rather, LNAPL and water coexist in

soil pores at residual LNAPL saturations ranging up to 40 to 50 percent (Huntley *et al.*, 1994). Nevertheless, the thickness of LNAPL within a soil column is expected to influence LNAPL weathering rates.

2.4.2 Groundwater Velocity

If equilibrium conditions exist between the LNAPL and the aqueous phase (Section 2.3.1.3), contaminant dissolution and depletion from the LNAPL source is enhanced with advection or groundwater flow. In soil column experiments performed by Miller *et al.* (1990), the rate of mass transfer between a toluene NAPL and the aqueous phase was found to be directly related to the aqueous-phase velocity. In addition, equilibrium conditions between the two fluid phases were rapidly achieved over a wide range of test conditions. Considering these findings, it is assumed that sites with higher groundwater velocities may exhibit more rapid BTEX depletion of mobile LNAPLs in contact with the water table.

2.4.3 Soil/Aquifer Material

The type of soil/aquifer material at a fuel release site is expected to influence LNAPL weathering primarily as a result of fluid distribution and migration. Wettability, or the tendency for one fluid to spread on or preferentially coat a solid surface in the presence of another fluid with which it is immiscible, is impacted by the presence of organic matter, mineralogy, and saturation history of the porous medium (Mercer and Cohen, 1990). Capillary pressure also impacts the configuration and magnitude of trapped residual LNAPL and is a function of soil pore size (Newell *et al.*, 1995). LNAPLs have been observed to preferentially migrate through sands and gravels, rather than silts and clays (Newell *et al.*, 1995).

2.4.4 Distance from Source Area

It is presumed that LNAPL weathering is impacted by the distance from the original fuel release location. Because of the effects of sequestration, increased surface area, and decreased contaminant mass, it is presumed that periphery LNAPL weathers at a faster rate than core area LNAPL. It is unlikely that LNAPL weathering occurs at a uniform rate across the area of impact (Landon and Hult, 1991).

2.5 PETROLEUM HYDROCARBON WEATHERING STUDIES

No weathering studies were identified that evaluated BTEX depletion from gasoline and/or jet fuel LNAPLs with the intent of refining contaminant source term reductions for fate and transport modeling. Based on a review of the literature, hydrocarbon weathering studies have primarily focused on weathering of crude oil and heavier refined petroleum products such as fuel oils and diesel fuel (Zurcher and Thuer, 1978; Fried, 1979; Law, 1980; Gundlach *et al.*, 1983; Baedecker *et al.*, 1987; Eganhouse *et al.*, 1988; Baedecker and Cozzarelli, 1991; Landon and Hult, 1991; Baedecker *et al.*, 1993; Christensen and Larsen, 1993; Douglas *et al.*, 1994; Vandermeulen *et al.*, 1994; Douglas *et al.*, 1996; Nakles *et al.*, 1996). Typically, these investigations have focused on the high-molecular-weight, low-solubility fractions in assessing changes in chemical composition. Many of these studies have utilized hydrocarbon ratios and internal biomarkers to evaluate relative degrees of weathering, to estimate spill age, and for source identification (Christensen and

Larsen, 1993; Douglas *et al.*, 1994; Douglas *et al.*, 1996; Kaplan *et al.*, 1996.). A brief summary of the more pertinent findings from the literature search is presented below.

2.5.1 Bemidji Oil Release Site

In 1979, a crude oil pipeline near Bemidji, Minnesota ruptured and released approximately 450,000 gallons of crude oil into a glacial outwash aquifer. In 1982, the site was selected for a long-term interdisciplinary study by the US Geological Survey. A study performed by Landon and Hult (1991) represents the investigation identified during the literature search that had objectives most similar to those of this fuel weathering study.

The purpose of the Landon and Hult study was to evaluate oil loss rates at a spill site in order to refine contaminant source-term reduction estimates for fate and transport models. Oil samples were collected from various locations within mobile LNAPL pools over a 10-year period to establish oil loss rates. Rather than chemical composition, changes in oil specific gravity and kinematic viscosity were used to calculate oil mass loss rates. Based on sample results, annual oil-mass loss ranged from 0.1 to 1.25 percent, and total cumulative oil losses after approximately 10 years of weathering were reported to be as much as 11 percent. Important conclusions from this investigation included:

- Oil-mass loss rates were found to vary spatially (i.e., to depend upon location within the oil pool);
- Oil-mass loss rates were found to vary temporally (i.e., to change based on relative age of the release);
- Volatilization of low-molecular-weight compounds was suspected to be the primary weathering mechanism.

Weathering rates for individual chemicals were not determined as part of the Landon and Hult (1991) study. However, depletion rates for BTEX compounds in refined petroleum products such as JP-4 and gasoline are expected to be greater than the total oil mass loss rates observed in mobile LNAPL at the Bemidji site.

2.5.2 Internal Biomarkers and Hydrocarbon Ratios

At oil release sites, the extent of oil or analyte depletion within soils or sediment has been estimated utilizing an internal biomarker or standard. For crude oil, the saturated pentacyclic (5-ring) triterpane known as hopane has been used because of its resistance to degradation (Douglas *et al.*, 1994; Douglas *et al.*, 1996). As biodegradation proceeds, the relative concentration of hopane remaining in the oil increases because of the removal of other more easily degraded compounds. As proposed by Douglas *et al.* (1994), the percent of oil depletion can be estimated by comparing the concentration of hopane in the weathered oil (H₁) with the concentration in the initial source oil (H₀) using the following equation:

% oil depletion =
$$[1 - (H_0 / H_1)] \times 100$$
 eq. 2.4

In addition, the amount of depletion of any one analyte within the oil can be determined using these hopane values in combination with analyte concentration in the degraded oil (C_1) and the analyte concentration in the source oil (C_0) as shown:

% analyte depletion =
$$[1 - ((C_1 / C_0) (H_0 / H_1))] \times 100$$
 eq. 2.5

The use of these equations to determine total oil and analyte depletion is considered to be conservative (i.e., to provide minimum depletion estimates) because the hopane degrades very slowly (Douglas *et al.*, 1996). Equation 2.5 was used to determine analyte depletion in shoreline sediment samples following the ExxonTM Valdez oil spill. Analyte depletion in these samples ranged from 30 to 70 percent 16 months after the spill (Douglas *et al.*, 1996). It also was noted during this study that the relative degree of PAH depletion decreased with increasing ring numbers and increased alkylation.

Similarly, hydrocarbon ratios have been used to determine the degree of change in oil/fuel composition with time and weathering. A ratio that is frequently used to assess biodegradation is the n- C_{17} /pristane ratio. The n- C_{17} compound is simply a saturated 17-carbon alkane. Pristane is a 19-carbon isoalkane, or isoprenoid, that is more resistant to biodegradation than the alkane n- C_{17} . In a study performed by Christensen and Larsen (1993) on biodegradation of residual diesel fuel in soils, the n- C_{17} /pristane ratio had the highest correlation factor with fuel residence time in soils of any similar n-alkane/isoalkane ratio. Based on the results of this study, Christensen and Larsen (1993) determined that the n- C_{17} /pristane ratio could be used to determine the age of a diesel oil spill within a range of plus or minus 2 years at a 95-percent level of confidence. The data also suggested that the n-alkanes biodegrade at a zero-order rate within residually contaminated soils.

2.5.3 BTEX Ratios

For refined petroleum products with higher initial BTEX concentrations (e.g., gasoline), ratios of the BTEX compounds have been used to estimate the relative state of degradation. As noted by Kaplan *et al.* (1996), BTEX results offer an excellent means of evaluating fuel alteration resulting from dissolution and volatilization. Comparing concentration ratios of the BTEX compounds in groundwater samples will typically show that benzene and toluene will be enriched relative to ethylbenzene and xylenes. However, in soil samples ethylbenzene and xylenes are preferentially retained relative to benzene and toluene.

Kaplan *et al.* (1996) suggest that a useful parametric ratio to evaluate gasoline partitioning is (benzene+toluene)/(ethylbenzene+xylenes). Based on their results, the average (B+T)/(E+X) ratio ranged from 0.74 to 0.88 for newly dispensed gasolines; whereas, the average ratio for free product, water, and soil were 0.65, 0.97, and 0.48, respectively. In laboratory studies by Kaplan *et al.* (1996), (B+T)/(E+X) ratios of 1.0 to 5.0 have been found for water in contact with fresh gasoline. At fuel release sites where groundwater samples are collected in the source area and the (B+T)/(E+X) ratio falls within this range, a recent release is indicated. At sites where the gasoline release is more than 10 years old, the ratio in the vicinity of the source area typically is less than 0.5. Ratios greater than 5.0 typically are encountered at sites where the groundwater samples are collected at a distance from the source area, and benzene and toluene concentrations

are relatively higher than ethylbenzene and xylenes concentrations because of dissolution effects.

SECTION 3

SELECTION OF STUDY SITES

The primary objective of this study was to determine a range of natural *in situ* weathering rates for mobile LNAPL associated with jet fuels and gasoline releases based on the existing literature and data collected from sites with mobile LNAPL contamination. Because no case studies identified in the literature evaluated quantitative source strength reduction of the BTEX constituents within mobile LNAPL, field sampling of representative sites was determined to be necessary. The site selection criteria for the fuel weathering study are presented in Section 3.1. Sites included within the study are summarized in Section 3.2.

3.1 SITE SELECTION CRITERIA

To evaluate a site's potential as a candidate for the fuel weathering study, the following selection criteria were considered:

- 1. Presence of recoverable mobile LNAPL in the subsurface environment as a result of a jet fuel or gasoline release;
- 2. Known date of fuel release;
- 3. Single release confined to a relatively short period of time;
- 4. Minimal remediation of the site and mobile LNAPL;
- 5. Historic LNAPL analytical results, including BTEX;
- 6. Depth to groundwater less than 40 feet below ground surface (bgs); and
- 7. Department of Defense (DOD) sites.

Identifying sites that met all of the above-listed criteria proved to be a difficult task. Consequently, the criteria served as guidelines for site selection rather than rigid selection parameters. Each of the criteria and their consideration in site selection are briefly discussed below.

JP-4 and gasoline fuel release sites were preferred for the study because of the relatively high mass fraction of BTEX present in these source fuels (Figure 2.3). Source reduction (i.e., BTEX depletion) estimates using mobile LNAPL sampling data from these types of fuel release sites were anticipated to be more accurate because of the higher initial BTEX concentrations. However, due to the difficulty of finding an adequate number of sites meeting the selection criteria, JP-5 and JP-8 release sites also were included in the study. While benzene, toluene, and ethylbenzene concentrations are less than 0.40 wt% in JP-8 (Mayfield, 1996), and less than 0.05 wt% in JP-5 (based on the fresh sample

collected at Beaufort MCAS), it was hoped that concentrations of total xylenes, naphthalene, and methylnaphthalenes could be used to evaluate mobile LNAPL weathering rates for JP-5 and JP-8.

Recoverable mobile LNAPL was loosely defined during initial site screening as sufficient free product in a site monitoring well to allow collection of relatively undiluted product samples. One inch of mobile LNAPL was considered to be the minimum required thickness for site consideration.

Locating sites with a known date of fuel release (criterion 2) where the release was a one-time event confined to a relatively short period of time (criterion 3) was difficult, especially when combined with the requirement for recoverable mobile LNAPL (criterion 1). For many petroleum release sites, the specific date(s) of release is not documented and at best can be approximated based on known historical site activities. In addition, one-time releases of sufficient volume to produce a long-term mobile LNAPL in the subsurface environment are rare. Moreover, when such releases occur, they frequently trigger emergency response actions that compromise satisfaction of the fourth selection criterion (minimal site remediation).

Sites where limited or no site remediation had occurred were preferred for assessing *in situ* LNAPL weathering rates. Soil venting activities, such as SVE, bioventing, and bioslurping will increase volatilization and biodegradation of the BTEX fraction in LNAPL; therefore, a BTEX weathering evaluation of the mobile LNAPL remaining at such sites would be biased. Sites where limited product recovery or soil excavation has occurred were not excluded from consideration.

Historical mobile LNAPL BTEX sampling results were considered in lieu of a known spill or release date. The availability of BTEX concentrations from a previous sampling event could help define BTEX depletion curves for the mobile LNAPL. Historical LNAPL BTEX results at least 3 years old were desired, but such data were seldom available.

Sites with shallow groundwater (less than 40 feet bgs) were selected so that Geoprobe[®] sampling could be performed. This requirement precluded the selection of sites in arid regions with thick valoes zones and deep water tables. As a result, many of the sites selected for the study are located in coastal regions with shallower water tables.

Project funding and liability issues restricted the study to DOD sites. Because of this restriction, the selection pool of gasoline release sites was limited. While gasoline is used and stored at DOD facilities, most of the petroleum infrastructure is dedicated to storage and transfer of the various jet fuels, including JP-4, JP-5, and more recently, JP-8.

3.2 FUEL WEATHERING STUDY SITES

Eight primary sites were selected for the study. Summary information for the primary sites was submitted to AFCEE prior to field sampling activities (Appendix A). In addition, samples from four secondary sites were collected and analyzed to support the study. Summary site data for the primary and secondary sites are provided in Table 3.1.

3.2.1 Site Summary

Based on the site-selection criteria summarized in Section 3.1, sample data from one gasoline, six JP-4, three JP-5, and two JP-8 fuel release sites were collected to evaluate mobile LNAPL weathering. Table 3.1 provides summary information for each site, including fuel type, volume and date of fuel release, and hydrogeologic information (e.g., soil type, approximate depth of water table, groundwater velocity, and free product or mobile LNAPL thickness). Figure 3.1 illustrates the geographic distribution of the selected sites. The fuel weathering study work plan and site-specific addenda (Appendix A) provide additional information on the primary sites selected for the study. Further information on sample collection methods are presented in Section 4.
TABLE 3.1 SITE SUMMARY FUEL WEATHERING STUDY

	F 1		Amount		Depth to	Groundwater	Free Product	
Site/Location	Fuel	Date of Release	Released (gallons)	Soil Type ^{a/}	Water Table (feet bas ^{b/})	Velocity (feet/year)	Thickness (feet) and Date	References
Site Location	Type	Release	(ganons)	Soli Type	(leet bgs)	(leet/year)	(leet) and Date	Kelefences
Primary Sites								
Tank 349 Offutt AFB, NE	Gasoline	1990	Unknown	Clay/Sand	39-42	11	2.23(6/96)	Parsons ES, 1997
Bldg 1610 Shaw AFB, SC	JP-4	June 1994	Unknown	Sand	29-33	400	2.5(8/96)	Parsons ES, 1998
Pipeline Leak Site Myrtle Beach AFB, SC	JP-4	January 1981	123,000	Clay/Sand	2-8.5 ^{c/}	420	3.79(11/95)	ECT, 1996
Tank 1 Area, DFSP-Charleston, Hanahan, SC	JP-4	October 1975	83,000	Clay/Sand	18-22	62	1.77(5/96)	USGS, 1997
Spill Site No. 2 Eaker AFB, AR	JP-4	October 1973	Unknown	Sandy Silt	8-14	16	1.18(8/97)	Halliburton NUS, 1996
Tank Farm C Beaufort MCAS, SC	JP-5	June 1990	10,600	Silty Sand	2-8	20	0.13(5/96)	USGS, 1996
Day Tank 1, Facility 293 Cecil Field NAS, FL	JP-5	1981	497,000	Silty Sand	5-8	6	0.78(8/96)	ABB, 1995a; 1996
Bldg 4522 Seymour Johnson AFB, NC	JP-8	December 1995	5,000	Sand	4-9	130	2.8(4/96)	Parsons ES, 1996c
Secondary Sites								
KC-135 Crash Site Wurtsmith AFB, MI	JP-4	October 1988	3,000	Sand	9-12	110	0.22(3/91)	Parsons ES, 1996a
Washrack/Treatment Area McChord AFB, WA	JP-4	1975	100,000	Silty Gravel	11-15	NA ^{d/}	0.14(4/94)	EA, 1994
Day Tank 865 Beaufort MCAS, SC	JP-5	1974	60,000	Silty Sand	2-8	7	0.15(5/97)	ABB, 1995b
JP-8 Release Site Pope AFB, NC	JP-8	April 1996	700	Sand	6-9	100	0.01(7/96)	Parsons ES, 1996b

a/ Represents soil type at the capillary fringe/water table.
b/ Feet below ground surface.
c/ Represents depth below ground surface to potentiometric surface.
d/ NA = not available.



SECTION 4

COLLECTION OF SITE DATA

To assess the effects of mobile and residual LNAPL weathering as they apply to soil and groundwater, samples of each medium (i.e., soil, groundwater, and mobile LNAPL) were collected from the study sites listed in Table 3.1. Where possible, samples were collected at each site within the area impacted by mobile LNAPL to determine weathering effects on mobile LNAPL and its relation to contaminants in soil at the capillary fringe and in groundwater. Samples collected by Parsons ES during 1997 and 1998 field sampling events form the foundation for this study; however, samples collected prior to 1997 and by other organizations also have been included, where appropriate. Table 4.1 provides a summary of the origin and types of samples collected and analyzed for the eight primary and four secondary fuel weathering study sites.

The following subsections provide a summary of soil, mobile LNAPL, and groundwater collection procedures. A brief description of the laboratory analytical methods used for this study also is provided. The work plan provides further information about sample collection and analysis procedures (Appendix A).

4.1 SOIL SAMPLING METHODS

Soil samples for the study were collected using a truck-mounted Geoprobe®. At most of the selected sites, soil samples were collected from a minimum of two separate borings. To maximize the possibility of obtaining soil samples within areas of measurable mobile LNAPL, boreholes were generally placed as close as possible to monitoring wells displaying maximum mobile LNAPL thickness for the site. Soil samples typically were collected approximately 1 foot above the water table for the purpose of evaluating weathering of residual LNAPL in the capillary fringe

Soil samples were submitted to the US Environmental Protection Agency (USEPA) National Risk Management Research Laboratory (NRMRL) (formerly the Kerr Research Laboratory), in Ada, Oklahoma. Section 4.4 summarizes the soil, mobile LNAPL, and groundwater analytical methods utilized for the study.

4.2 MOBILE LNAPL SAMPLING

Whenever possible, mobile LNAPL samples were collected from two separate site monitoring wells. At the Seymour Johnson AFB and Beaufort MCAS sites, mobile LNAPL was present, and thus collected, at only one site monitoring well. No mobile LNAPL was encountered at the Wurtsmith AFB site.

TABLE 4.1 ORIGIN OF SAMPLE DATA FUEL WEATHERING STUDY

Site	Date of Sample Collection	Sample Type ^{a/}	Samples Collected By ^{b/}	Samples Analyzed By ^{c/}
Primary Sites				
Tank 349 Offutt AFB, NE	November 1994 June 1996 June 1997 October 1998	S,GW,FP GW,FP S,GW,FP FP	USACE/Parsons ES Parsons ES Parsons ES Parsons ES	NRMRL NRMRL EAL, NRMRL EAL, NRMRL
Building 1610 Shaw AFB, SC	March 1997 March 1998	S,GW,FP S,GW,FP	Parsons ES Parsons ES	EAL, NRMRL EAL, NRMRL
Pipeline Leak Site Myrtle Beach AFB, SC	March 1997	S,GW,FP	Parsons ES	EAL, NRMRL
DFSP-Charleston, Tank 1 Area Hanahan, SC	December 1993 May 1995 May 1997	FP FP S,GW,FP	USGS UGSG Parsons ES	NRMRL NRMRL EAL, NRMRL
Spill Site No. 2 Eaker AFB, AR	August 1997	S,GW,FP	USACE	EAL, NRMRL
Tank Farm C Beaufort MCAS, SC	May 1997 August 1997	S,GW,FP FP	Parsons ES Beaufort Personnel	EAL, NRMRL NRMRL
Day Tank 1, Facility 293 Cecil Field NAS, FL	May 1997	S,GW,FP	Parsons ES	EAL, NRMRL
Bldg 4522 Seymour Johnson AFB, NC	July 1996 May 1997 March 1998	S,GW,FP S,GW,FP S,GW,FP	USACE Parsons ES Parsons ES	NRMRL EAL, NRMRL EAL, NRMRL
Secondary Sites				
KC-135 Crash Site Wurtsmith AFB, MI	August 1996	S,GW	Parsons ES	NRMRL
Washrack/Treatment Area McChord AFB, WA	September 1997	FP	McChord Contractor	NRMRL
Day Tank 865 Beaufort MCAS, SC	May 1997	S,FP	Parsons ES	NRMRL
JP-8 Release Site Pope AFB, NC	July 1996	S,GW,FP	USACE	NRMRL

a/ S = soil; GW = groundwater; FP = free product or mobile LNAPL.
b/ USACE = US Army Corps of Engineers, Kansas City District Office; USGS = United States Geological Survey, Water Resource Division, Columbia, SC.
c/ EAL = Evergreen Analytical Laboratory, Wheat Ridge, Colorado; NRMRL = National Risk Management Research Laboratory, Ada, Oklahoma.

It was originally proposed in the work plan (Appendix A) that mobile LNAPL, groundwater, and soil sampling would be performed in the same vertical continuum within one borehole. It was proposed that groundwater and mobile LNAPL samples would be collected from temporary monitoring points. Attempts were made to collect mobile LNAPL samples from temporary monitoring points at several sites. At these sites, the temporary monitoring points were located within 4 to 7 feet of monitoring wells that contained mobile LNAPL, and were screened to intersect the top of the water table. After monitoring point placement and some initial groundwater purging, the monitoring points were allowed to recharge for up to 15 hours, with the expectation that a sufficient amount of mobile LNAPL (5 to 10 mL) would flow into the monitoring point for sample collection. Only during the 1998 sampling event at Seymour Johnson could a mobile LNAPL sample be collected from a temporary monitoring point. At all other sites no more than a slight sheen of mobile LNAPL was detected in the temporary monitoring points, and mobile LNAPL samples had to be collected from site monitoring wells.

In order to minimize the effects of evaporation on volatile fuel constituents and to obtain samples representative of the mobile LNAPL present in site formations, mobile LNAPL samples generally were collected from site monitoring wells following an initial purging of mobile LNAPL. Because the rate of mobile LNAPL recovery was unknown, a "pre-purge" sample was collected from site monitoring wells in the event that sufficient mobile LNAPL recovery did not occur following initial purging. At most sites, at least one casing-volume of product was removed, and mobile LNAPL recovery was sufficient for "post-purge" sample collection within an hour or less. Other than the "pre-purge" samples from Beaufort Tank Farm C, Beaufort Day Tank 865, and the Cecil Field NAS site (monitoring well CEF-293-7), the mobile LNAPL samples submitted for laboratory analysis were "post-purge" samples. Mobile LNAPL samples were sent to NRMRL and Evergreen Analytical Laboratory (EAL) in Wheat Ridge, Colorado for analysis (Section 4.4).

4.3 GROUNDWATER SAMPLING

Groundwater samples usually were collected from two locations at each of the selected sites. At sites where the water table was less than 20 feet bgs, groundwater samples were collected from temporary monitoring points placed within the Geoprobe® boreholes created during soil sampling. The boreholes and monitoring points generally were placed within 5 to 7 feet of site monitoring wells displaying maximum mobile LNAPL thickness. At sites where the groundwater was more than 20 feet bgs (i.e., Shaw AFB and Offutt AFB), groundwater samples were collected from existing site monitoring wells that contained mobile LNAPL. In addition, at the Seymour Johnson AFB and Cecil Field NAS sites, one groundwater sample also was collected from existing site monitoring wells which contained mobile LNAPL. It is possible that some emulsification of mobile LNAPL may have occurred in groundwater samples collected from these monitoring wells. Groundwater samples were submitted to NRMRL for analysis.

4.4 LABORATORY ANALYSIS

Table 4.2 presents a summary of the analytical methods performed by each laboratory. Analytical results from NRMRL and EAL are provided in Appendix B.

TABLE 4.2 ANALYTICAL PROTOCOLS FOR SOIL, MOBILE LNAPL, AND GROUNDWATER SAMPLES FUEL WEATHERING STUDY

MATRIX	LABORATORY	NUMBER OF SAMPLES PER SITE	ANALYSIS ^{a/}	METHOD ^{b/}
SOIL	NRMRL	2 to 3	BTEX + TMBs Napthalene and Methyl Napthalenes Total Fuel Carbon	NRMRL equivalent to USEPA SW8020A NRMRL equivalent to USEPA SW8270
MOBILE LNAPL	NRMRL	1 to 2	BTEX + TMBs Napthalene and Methyl Napthalenes Density	GC/MS (Direct Injection) NRMRL equivalent to USEPA SW8270 Method 2710F (Standard Methods, 1995)
	EAL	1 to 2	BTEX (Aqueous and Organic Phases)	In accordance with Cline et al. (1991)
GROUNDWATER	NRMRL	1 to 2	BTEX + TMBs Napthalene and Methyl Napthalenes	NRMRL equivalent to USEPA E602 NRMRL equivalent to USEPA SW8270

^{a'} BTEX = benzene, toluene, ethylbenzene, and total xylenes; TMBs = trimethylbenzene isomers.

4.4.1 National Risk Management Research Laboratory

NRMRL analyzed soil, groundwater, and mobile LNAPL samples from several study sites. BTEX, naphthalene, 1-methylnaphthalene, 2-methylnaphthalene, and the various TMB isomers concentrations were determined for each matrix. In addition, soil samples were analyzed for total fuel carbon, and mobile LNAPL samples were analyzed for fuel density. Samples from the eight primary and four secondary sites (Table 4.1) were submitted to NRMRL for analysis.

4.4.2 Evergreen Analytical Laboratory

EAL analyzed mobile LNAPL samples collected from various study sites in order to determine fuel/water partitioning coefficients (K_{fw}) at equilibrium saturations. The EAL analyses generally were performed in accordance with procedures from the Cline *et al.*(1991) study. Saturated, equilibrium solutions of the collected fuels in contact with distilled, deionized, organic-free water were prepared. Two mL of fuel were added to 40 mL of water in volatile organics analysis vials having Teflon® septa (a 1:20 fuel to water ratio). Sample vials were agitated for approximately 30 minutes, then allowed to rest for 1 hour in an inverted position. Following mixing and stabilization, the aqueous phase and the organic (fuel) phase were analyzed separately for determination of BTEX concentrations by USEPA SW8020 by gas chromatography with photoionization detection (GC/PID).

SECTION 5

ANALYTICAL RESULTS AND DATA ANALYSIS

The primary objective of this study was to determine a range of natural weathering rates for mobile LNAPLs in order to refine modeling assumptions for the contaminant source term. The BTEX compounds were the primary focus of the study as they typically represent the primary contaminants of concern at gasoline and JP-4 fuel release sites. Naphthalene and methylnaphthalenes also were evaluated because these aromatic compounds can represent contaminants of concern at sites contaminated by kerosene-based jet fuels (i.e., JP-5 and JP-8). In addition to the mobile LNAPL weathering analysis, fuel/water partitioning coefficients for BTEX compounds were determined based on field data and compared to laboratory-determined partitioning values. Lastly, residual LNAPL weathering in capillary fringe soils was compared to mobile LNAPL weathering.

5.1 RESULTS SUMMARY

Table 5.1 summarizes mobile LNAPL, groundwater, and soil analytical results for the remaining BTEX fraction based on fuel type and includes sample data from the eight primary and four secondary sites. The mobile LNAPL values shown represent analytical results obtained from EAL and NRMRL. Groundwater and soil analytical results are from NRMRL. A more complete listing of analytical results obtained from EAL and NRMRL for each site, including naphthalene, methylnaphthalene, and TMB results, is provided in Appendix B.

5.1.1 Mobile LNAPL BTEX Results

Mobile LNAPL BTEX results varied considerably with fuel type. Total BTEX concentrations in mobile LNAPL collected at the fuel release sites ranged from 11 mg/mL (JP-5 and JP-8) to 135 mg/mL (gasoline). The most significant variations are apparent in the benzene and toluene fractions, where their concentrations varied over approximately 3 orders of magnitude for the different fuel types. Maximum benzene concentrations of 14 mg/mL, 2.7 mg/mL, 0.25 mg/mL, and 0.02 mg/mL were detected in mobile LNAPL from the gasoline, JP-4, JP-8, and JP-5 fuel release sites, respectively. Maximum toluene concentrations in mobile LNAPL ranged from 0.12 mg/mL at the JP-5 sites to 52 mg/mL at the gasoline site. Order-of-magnitude differences in the mobile LNAPL BTEX concentrations based on fuel type are consistent with differences in BTEX concentrations among the fresh fuels (Figure 2.3).

Fuel Type	No. of	Benzene		Toluene		Ethylbenzer	ne	Xylenes		Total BTEX		
Sample Matrix ^{a/}	Samples	Range	Average	Range	Average	Range	Average	Range	Average	Range	Average	
JP-4 Jet Fuel												
Mobile LNAPL (mg/mL) ^{b/}	22	0.00003 - 2.7	0.67	0.00003 - 5.7	1.1	0.00003 - 5.5	1.3	0.0001 - 26	6.3	0.0002 - 35	9.3	
Groundwater (mg/L)c/	8	0.037 - 8.7	3.1	0.002 - 8.8	1.9	0.077 - 1.2	0.50	0.058 - 8.3	2.7	0.47 - 27	8.2	
Soil (mg/kg) ^{d/}	14	0.006 - 11	1.5	0.006 - 22	2.5	0.006 - 34	5.5	0.018 - 173	25	0.036 - 230	34	
JP-5 Jet Fuel												
Mobile LNAPL (mg/mL)	4	0.002 - 0.02	0.013	0.013 - 0.12	0.07	0.12 - 3.2	1.5	0.61 - 7.3	3.4	0.74 - 11	5	
Groundwater (mg/L)	2	0.004 - 0.05	0.026	0.023 - 0.11	0.066	0.12 - 0.21	0.16	0.45 - 0.70	0.58	0.60 - 1.1	0.83	
Soil (mg/kg)	6	0.012 - 3.3	0.94	0.079 - 19	6.3	1.5 - 155	57	0.093 - 425	145	6.9 - 600	208	
JP-8 Jet Fuel												
Mobile LNAPL (mg/mL)	9	0.00003 - 0.25	0.13	0.00003 - 1.6	0.85	0.063 - 1.2	0.93	1.01 - 7.5	4.8	1.1 - 11	6.7	
Groundwater (mg/L)	4	0.0005 - 0.85	0.42	0.001 - 4.1	1.8	0.007 - 0.84	0.37	0.067 - 3.2	1.5	0.076 - 9.0	4.1	
Soil (mg/kg)	9	0.006 - 13	6.0	0.006 - 79	35	0.006 - 75	38	0.018 - 416	168	0.036 - 561	248	
Gasoline												
Mobile LNAPL (mg/mL)	7	0.96 - 14	7.9	12 - 52	36	9.3 - 13	11	33 - 57	43	56 - 135	97	
Groundwater (mg/L)	5	9.6 - 38	30	24 - 44	37	3.6 - 4.6	4.1	13 - 18	15	51 - 101	86	
Soil (mg/kg)	4	0.56 - 43	26	0.33 - 165	88	0.29 - 59	30	0.79 - 203	102	2.0 - 467	247	

TABLE 5.1 BTEX CONTENT IN MOBILE LNAPL, GROUNDWATER, AND SOIL RESULTING FROM GASOLINE AND JET FUEL RELEASES FUEL WEATHERING STUDY

^{a/} Mobile LNAPL analytical results obtained from EAL and NRMRL. Groundwater and soil analytical results obtained from NRMRL.

^{b/} mg/mL = milligrams per milliliter.

^{c/} mg/L = milligram per liter.

^d/ mg/kg = milligrams per kilogram.

5.1.2 Groundwater BTEX Results

Groundwater BTEX analytical results in the LNAPL source area also varied with fuel type. Groundwater concentrations at the gasoline site consistently exceeded USEPA (1996) MCLs for benzene (0.005 mg/L), toluene (1.0 mg/L), ethylbenzene (0.7 mg/L), and total xylenes (10 mg/L). At the jet fuel release sites where the effective solubility of the BTEX compounds in the LNAPL is significantly lower than in gasoline, MCL exceedances were less frequent. Maximum concentrations of benzene measured at the JP-4, JP-5, and JP-8 release sites exceeded the MCL; however, concentrations in some groundwater samples at the JP-5 and JP-8 sites were below the benzene MCL. Even at the JP-4 sites where fuel releases occurred more than 20 years prior to the sampling event, benzene concentrations in groundwater continued to exceed the MCL. Toluene and ethylbenzene concentrations in contaminant source area groundwater at the JP-4 and JP-8 sites occasionally exceeded their MCLs, but no MCL exceedances were observed for these analytes in the two JP-5 site samples. Total xylenes concentrations in groundwater at the JP-4, JP-5, and JP-8 release sites were consistently below the MCL of 10 mg/L, indicating that xylene contamination is not likely to be a significant environmental threat at jet fuel release sites. However, xylene concentrations in groundwater at the gasoline site exceeded the MCL in all samples (Tables 2.1 and 5.1).

5.1.3 Soil BTEX Results

BTEX concentrations detected in capillary fringe soil samples did not vary as significantly with fuel type as did mobile LNAPL and groundwater BTEX concentrations. Maximum benzene concentrations of 43 milligrams per kilogram (mg/kg), 13 mg/kg, 11 mg/kg, and 3.3 mg/kg were measured at the gasoline, JP-8, JP-4, and JP-5 release sites, respectively. Similarly, maximum concentrations of toluene were greatest at the gasoline site (165 mg/kg), followed by the JP-8 sites (79 mg/kg), JP-4 sites (22 mg/kg), and JP-5 sites (19 mg/kg). Surprisingly, maximum soil concentrations of ethylbenzene (155 mg/kg) and total xylenes (425 mg/kg) were detected in the capillary fringe soil sample collected at the Cecil Field NAS JP-5 release site; however, comparatively low concentrations of ethylbenzene and xylenes were detected in the mobile LNAPL sample collected near this location (3.2 mg/mL and 7.3 mg/mL, respectively).

5.2 MOBILE LNAPL WEATHERING

Mobile LNAPL weathering was assessed by evaluating the mass fraction reduction of BTEX. BTEX concentrations in mobile LNAPL samples were compared to conservatively assumed initial BTEX concentrations in fresh fuel. Using the known dates of the product releases and the assumed initial BTEX compositions for the various fuels, the degree of mobile LNAPL weathering (i.e., BTEX mass fraction depletion) that has occurred with time was determined for each release site.

5.2.1 Assumed Initial Fuel Compositions

Initial concentrations of BTEX, naphthalene, and methylnaphthalenes in gasoline, JP-4, JP-5, and JP-8 were conservatively assumed. The assumed initial concentration for each fuel type is based on the following literature or analytical values presented in Section 2.1.2.4:

- JP-4 Initial values from Smith *et al.* (1981);
- JP-5 at Cecil Field NAS Initial values from Hughes et al. (1984);
- JP-5 at Beaufort MCAS Initial values based on Beaufort MCAS fresh JP-5 sample;
- JP-8 Initial values from Mayfield (1996); and
- Gasoline Initial values from Ghassemi et al. (1984).

Further discussion of the assumed fuel compositions for JP-4, JP-5, JP-8, and gasoline in relation to site-specific mobile LNAPL results is presented in Sections 5.2.3, 5.2.4, 5.2.5, and 5.2.6, respectively.

5.2.2 Kinetics of Weathering

As discussed in Section 2, LNAPL weathering in the subsurface environment results from a complex combination of physical, chemical, and biological processes. Weathering rates, or compound-specific depletion rates, are a function of these processes, and influenced by many variables. In view of this, the reaction kinetics that determine the rate of contaminant depletion are difficult to predict. For this study, no literature findings were identified that explained reaction kinetics for the overall weathering of a mobile LNAPL, and for most sites, only one or two data points were available for estimating contaminant depletion rates. Because of these limitations, contaminant depletion in this study was evaluated using both zero-order and first-order reaction kinetics.

5.2.2.1 Zero-Order Weathering

Zero-order weathering or decay is described by the following differential equation:

$$dC / dt = -k_0$$
 eq. 5.1

As shown on Figure 5.1, zero-order weathering assumes that contaminant depletion in the mobile LNAPL occurs at a constant rate (k_0). In addition, the rate of depletion of the contaminant is not reduced as the contaminant becomes increasingly more depleted with time and weathering of the mobile LNAPL. Solving this differential equation gives:

$$C = C_0 - k_0 t$$
 eq. 5.2

where: C = contaminant concentration (wt%) at time "t"

 C_0 = contaminant concentration (wt%) at time "zero"

 k_0 = zero-order rate constant or slope (wt% per year)

Solving for k_0 , eq. 5.2 can be written as:

$$k_0 = (C_0 - C) / t$$
 eq. 5.3

For zero-order weathering, the amount or percent of annual contaminant depletion can be compared to the concentration at time zero (C_0) by the following:

% C₀ Reduced / Year =
$$\frac{[(C_0 - C) / C_0] \times 100}{\text{Sample Date - Spill Date (years)}}$$
 eq. 5.4

5.2.2.2 First-Order Weathering

First-order decay or weathering is described by the following differential equation:

$$dC / dt = -k_1 t ext{ eq. 5.5}$$

As shown on Figure 5.2, the rate of contaminant depletion changes with time under the first-order weathering assumption. Under this scenario, the rate of contaminant depletion is proportional to the amount of contaminant that is present at any time "t." The first-order weathering curve shown on Figure 5.2 is an exponential curve, where the amount of contaminant remaining in the LNAPL approaches zero with time, but never reaches a zero concentration. Solving this differential equation gives:

$$C = C_0 e^{-k_1 t} \qquad eq. 5.6$$

where: C = contaminant concentration in wt% at time "t"

 C_0 = contaminant concentration in wt% at time "zero"

e = base of natural logarithms (approximately 2.72)

 $k_1 =$ first-order rate constant (years⁻¹ or 1/years)

Solving for k_1 , eq. 5.6 can be written as:

$$k_1 = -\ln(C/C_0) / t$$
 eq. 5.7

For first-order weathering, the yearly percent of contaminant depletion can be determined as follows:

% Reduction / Year =
$$(1 - e^{-k_1 t}) \times 100$$
 eq. 5.8

As discussed in Sections 2.3.1 and 2.3.2, the dissolution and volatilization of a compound is controlled by the amount (mole fraction) of that compound in the LNAPL under equilibrium conditions. Therefore, the rate of contaminant depletion resulting from dissolution or volatilization also may be proportional to the contaminant concentration, indicating first-order weathering may be more appropriate especially if one of these weathering mechanisms predominates.





5.2.3 Weathering in JP-4 Mobile LNAPL

Weathering rates or depletion rates for the BTEX and naphthalene compounds were evaluated at five JP-4 release sites with spill ages ranging between approximately 4 and 24 years. At the Myrtle Beach, Eaker, and McChord AFB sites, only one sampling event was performed. At the Shaw AFB and DFSP-Charleston sites, analytical data from more than one sampling event were evaluated.

Initial composition values for JP-4 were assumed to be equivalent to concentrations reported by Smith *et al.* (1981). For each of the BTEX compounds, composition values presented by Smith *et al* (1981) are slightly lower than the values presented by Hughes *et al.* (1984) (Figure 2.3), and therefore more conservative for use in estimating BTEX depletion. However, it should be noted that the Hughes *et al.* (1984) study considered a larger sample database than the Smith *et al.* (1981) study, and is considered the preeminent study on JP-4 jet fuel composition.

5.2.3.1 Range of Weathering Rates for the Five JP-4 Sites

Mobile LNAPL weathering rates for the JP-4 sites were determined for every BTEX, naphthalene, and methylnaphthalene sample result independent of results from other sampling events. For this analysis, rate constants k_0 and k_1 were determined for each sample result by solving equations 5.3 and 5.7. Annual reductions based on zero-order and first-order weathering assumptions were determined for each sample result by solving equations 5.4 and 5.8, respectively. Because these calculations can be performed knowing the contaminant concentration at only one-point in time relative to the known spill date, weathering rates determined using this method are hereafter referred to as "one-point" weathering rates.

Using this method, each sample result was given equal weight, and average BTEX and naphthalene weathering rates can be determined for each site. In addition, weathering rates at sites where only one sampling event was performed could be compared to sites where multiple sampling events were performed. BTEX and naphthalene weathering rates in JP-4 mobile LNAPL determined using the one-point method are summarized in Table 5.2. As shown, minimum, maximum, and average values for the rate constants k_0 and k_1 and annual contaminant reduction rates based on zero-order and first-order weathering are provided. In addition, assumed initial concentrations (Smith *et al.*, 1981) and average remaining concentrations are shown for comparison.

As shown on Table 5.2, the weathering rates determined for each site can vary significantly between the LNAPL samples analyzed. Reasons for this variability include differences in mobile LNAPL concentrations with different sample locations at the site, and differences in analytical results obtained by EAL and NRMRL for the same sample. Because of this inherent variability, the average BTEX and naphthalene reduction rates shown on Table 5.2 have been used to represent weathering rates at each site.

Based on the mobile LNAPL sample results shown, the average zero-order BTEX weathering rates range from 0.5 to 20 %/yr, and naphthalene and methylnaphthalene zero-order weathering rates range from 0.8 to 25 %/yr. The average first-order BTEX

TABLE 5.2 BTEX AND NAPHTHALENE (ONE-POINT)^{a/} WEATHERING RATES IN JP-4 MOBILE LNAPL FUEL WEATHERING STUDY

			Assumed	Average	age												
			Initial	Remaining	r		ZE	RO ORDER						FIRST	ORDER		- F/
Site	Approximate	Number of	Conc. $(C_0)^{c/}$	Conc. (C)	R	ate Constar	nt k ₀ ^{e/}	% C	o Reduced/Y	′ear ^{f/}		Ra	te Constant l	κ ₁ ^{g/}	% F	Reduction/Ye	ar "
Analyte	Spill Age ^{b/}	Samples	(wt%) ^{d/}	(wt %)	min	max	avg	min	max	avg		min	max	avg	min	max	avg
Shaw AFB, SC	4 years																
Benzene		6	0.50	0.23	0.06	0.12	0.08	11	23	17		0.13	0.38	0.25	12	31	22
Toluene		6	1.33	0.48	0.21	0.37	0.26	16	27	20		0.21	0.51	0.33	19	40	28
Ethylbenzene		6	0.37	0.16	0.05	0.09	0.06	14	24	17		0.17	0.41	0.26	16	34	23
Total Xylenes		6	2.32	1.07	0.31	0.52	0.39	13	23	17		0.17	0.35	0.24	16	30	22
Total BTEX		6	4.52	1.94	0.64	1.10	0.80	14	24	18		0.18	0.40	0.27	16	33	23
Naphthalene		4	0.50	0.16	0.09	0.13	0.11	18	26	21		0.29	0.44	0.35	25	36	29
1- Methylnaphthalene		4	0.78	0.16	0.16	0.23	0.19	21	30	25		0.40	0.63	0.49	33	47	39
2- Methylnaphthalene		4	0.56	0.27	0.07	0.12	0.09	13	22	16		0.18	0.34	0.23	17	29	21
Myrtle Beach AFB, SC	16 years																
Benzene		3	0.50	0.02	0.03	0.03	0.03	5.8	6.1	5.9		0.18	0.26	0.21	16	23	19
Toluene		3	1.33	0.00 ^{i/}	0.08	0.08	0.08	6.2	6.2	6.2		0.44	0.46	0.45	36	37	36
Ethylbenzene		3	0.37	0.17	0.01	0.02	0.01	2.1	4.9	3.4		0.03	0.10	0.06	2.5	9.1	5.3
Total Xylenes		3	2.32	0.57	0.09	0.13	0.11	3.9	5.4	4.7		0.06	0.13	0.09	6.0	12	8.9
Total BTEX		3	4.52	0.76	0.21	0.26	0.23	4.6	5.7	5.1		0.09	0.16	0.12	8.3	14	11
Naphthalene		2	0.50	0.14	0.02	0.02	0.02	4.3	4.7	4.5		0.07	0.09	0.08	7.0	8.6	7.8
1- Methylnaphthalene		2	0.78	0.17	0.04	0.04	0.04	4.8	4.8	4.8		0.09	0.09	0.09	8.9	8.9	8.9
2- Methylnaphthalene		2	0.56	0.25	0.02	0.02	0.02	3.4	3.6	3.5		0.05	0.05	0.05	4.8	5.2	5.0
DFSP-Charleston, SC	22 years																
Benzene		6	0.50	0.00 ^{i/}	0.02	0.03	0.02	4.6	5.5	4.8		0.15	0.55	0.45	14	43	35
Toluene		6	1.33	0.01	0.06	0.07	0.06	4.5	5.5	4.8		0.18	0.41	0.29	17	34	25
Ethylbenzene		6	0.37	0.10	0.00	0.02	0.01	1.3	4.8	3.6		0.02	0.16	0.09	1.5	15	8.7
Total Xylenes		6	2.32	0.44	0.07	0.12	0.09	3.2	5.1	3.9		0.05	0.18	0.10	4.9	17	9.3
Total BTEX		6	4.52	0.55	0.17	0.24	0.19	3.7	5.2	4.3		0.07	0.20	0.12	6.7	18	11
Naphthalene		4	0.50	0.08	0.02	0.02	0.02	3.7	4.9	4.2		0.07	0.12	0.09	6.4	11	9.0
1- Methylnaphthalene		4	0.78	0.19	0.03	0.03	0.03	3.5	4.2	3.7		0.07	0.08	0.07	6.4	7.5	6.7
2- Methylnaphthalene		4	0.56	0.24	0.01	0.02	0.02	2.6	3.3	2.9		0.04	0.05	0.04	3.7	4.9	4.2
Eaker AFB, AR	24 years																
Benzene		3	0.50	0.13	0.01	0.02	0.02	2.0	4.2	3.1		0.03	0.30	0.13	2.7	26	12
Toluene		3	1.33	0.04	0.05	0.06	0.05	3.8	4.2	4.0		0.10	0.54	0.39	9.3	42	31
Ethylbenzene		3	0.37	0.47	^{j/}	0.00	0.00		0.5				0.01			0.5	
Total Xylenes		3	2.32	2.03		0.07	0.01		3.0	0.5			0.05	0.01		5.0	1.3
Total BTEX		3	4.52	2.67	0.00	0.15	0.08	0.0	3.3	1.7		0.00	0.06	0.03	0.0	6.1	2.9
Naphthalene		2	0.50	0.13	0.02	0.02	0.02	3.0	3.1	3.1		0.05	0.06	0.06	5.2	5.5	5.4
1- Methylnaphthalene		2	0.78	0.25	0.02	0.02	0.02	2.7	3.1	2.9		0.04	0.06	0.05	4.2	5.4	4.8
2- Methylnaphthalene		2	0.56	0.45	0.00	0.01	0.00	0.3	1.4	0.8		0.00	0.02	0.01	0.3	1.6	1.0

TABLE 5.2 (Continued) BTEX AND NAPHTHALENE (ONE-POINT)^{a/} WEATHERING RATES IN JP-4 MOBILE LNAPL FUEL WEATHERING STUDY

			Assumed Initial	Average Remaining			ZEI	RO ORDER					FIRST (ORDER		
Site	Approximate	Number of	Conc. $(C_0)^{c/}$	Conc. (C)	Rate Constant k ₀ ^{e/}			% C _o Reduced/Year ^{f/}			Rate Constant k ₁ ^{g/}			% Reduction/Year ¹		ar ^{h/}
Analyte	Spill Age ^{b/}	Samples	(wt%) ^{d/}	(wt %)	min	max	avg	min	max	avg	min	max	avg	min	max	avg
McChord AFB, WA	22 years															
Benzene		1	0.50	0.00 ^{i/}			0.02			4.5			0.54			42
Toluene		1	1.33	0.00 ^{i/}			0.06			4.5			0.58			44
Ethylbenzene		1	0.37	0.00 ^{i/}			0.02			4.5			0.53			41
Total Xylenes		1	2.32	0.00 ^{i/}			0.10			4.5			0.56			43
Total BTEX		1	4.52	0.00 ^{i/}			0.20			4.5			0.56			43
Naphthalene		1	0.50	0.00 ^{i/}			0.02			4.5			0.54			42
1- Methylnaphthalene		1	0.78	0.02			0.03			4.4			0.18			16
2- Methylnaphthalene		1	0.56	0.01			0.02			4.4			0.16			15

Note: Calculated values shown have been rounded.

a' Analyte weathering rates in free-phase product calculated based on assumed initial analyte concentrations in fresh JP-4 fuel and one point in time free-phase product sample results.

^{b/} Approximate age of spill as of the most recent sampling event.

^{c/} Assumed initial concentrations from Smith et al. (1981).

^{d/} wt% = weight percent.

 $k_0^{e'}$ k₀ = zero-order rate contstant or slope calculated using equation 5.3; units in weight percent per year.

^{f/} Annual mass fraction reduction as a percent of the initial concentration; calculated using equation 5.4.

 $k_1 =$ first-order rate constant or exponential decay rate calculated using equation 5.7; units in years⁻¹ or 1/years.

^{h/} Weight percent reduction per year calculated using equation 5.8.

¹⁷ Result indicates a nondetect or near nondetect value; as appropriate, weathering rate calculations for this result were based on the method detection limit.

j/ --- = negative value; measured concentration is greater than the assumed initial concentration.

weathering rates range from 1.3 to 44 %/yr, and naphthalene and methylnaphthalene firstorder weathering rates range from 1.0 to 42 %/yr. The annual zero-order reduction rates are consistently lower than corresponding first-order rates. This phenomenon occurs uniformly for all data sets presented in this study, due to the inherent nature of the rate calculations. Greater degradation rates are required for first-order weathering to obtain the same end concentration because first-order weathering is concentration dependent. Zero-order weathering assumes that the concentration of a compound is depleted at a constant rate regardless of contaminant concentration.

First-order weathering rates that were calculated using nondetect or near nondetect values for C are significantly higher than reduction rates shown for more moderate compound depletion. This was particularly evident at the McChord AFB site where virtually no BTEX or naphthalenes were detected in a single sample collected 24 years following the fuel release. For this site, first-order reduction rates were estimated at more than 40 %/yr. This occurrence emphasizes a potential limitation of using a single sample collected several years after the fuel release to estimate weathering rates. Use of a 40 %/yr weathering rate or similar rate determined from nondetect or near nondetect concentrations may overestimate contaminant source-term reductions for fate and transport modeling. (Note: When nondetect concentrations were observed during this study, weathering rates were calculated as if the compound was detected at the method detection limit.)

As shown on Table 5.2, the range of average weathering rates were significant for the BTEX and naphthalene compounds. If the bias of high first-order weathering rates at McChord AFB are excluded, the highest weathering rates were measured at the Shaw AFB site. At this site, the average zero-order total BTEX weathering rate was 18 %/yr, and average zero-order naphthalene/methylnaphthalene rates were 16 to 25 %/yr. Average first-order weathering rates were higher, with 23 %/yr BTEX reduction and 21 to 39 %/yr naphthalene and methylnaphthalenes reduction.

The lowest weathering rates were apparent at the Eaker AFB site where the average total BTEX reduction rate was 1.7 %/yr and 2.9 %/yr for zero-order and first-order weathering, respectively. The second lowest average weathering rates were measured for the DFSP-Charleston site where the total BTEX reduction rate was 4.3 %/yr for zeroorder weathering and 11 %/yr for first-order weathering. The low reduction rates at the Eaker AFB site result from one mobile LNAPL sample in which BTEX concentrations were minimally depleted relative to the assumed initial concentration values. The sample in question was collected near the original fuel release source area, a fuel pipeline that was not abandoned until 1995. The minimal amount of weathering at this location 24 years following the reported fuel release may indicate that environmental conditions are not conducive to aromatic hydrocarbon depletion at this location. However, it seems more likely that the source area sample collected from the Eaker AFB site is from a more recent, undocumented fuel release. At DFSP-Charleston, the average total BTEX concentration in mobile LNAPL approximately 22 years after the fuel release (0.55 wt%) is approximately one-fifth the average total BTEX concentration in mobile LNAPL from the Eaker AFB site (2.67 wt%).

5.2.3.2 Combining JP-4 Site Data to Assess Weathering Rates

Very few fuel release sites have sufficient data available to determine the progression of mobile LNAPL BTEX depletion from the time of spill release until the time of complete BTEX removal. However, comparing and compiling data from all JP-4 sites, regardless of differences in hydrogeologic effects, provides some insight into the relationship between BTEX depletion in mobile LNAPL and spill age. Figure 5.3 represents a plot obtained when average total BTEX concentrations in mobile LNAPL from the five primary JP-4 sites are compiled into one weathering plot. Similarly, Figure 5.4 represents a compilation of average benzene data from the five JP-4 sites.

Zero-order and first-order curves were fitted to the plotted data to evaluate BTEX weathering in mobile LNAPL with time. Considering the combined data from the five JP-4 sites, the first-order curves appear to better match the general trend of the data. As shown in Figure 5.3, the zero-order curve greatly underestimates total BTEX depletion in the first 4 years following a JP-4 release as indicated by the Shaw AFB site data. Total BTEX depletion in mobile LNAPL at the Myrtle Beach AFB and DFSP-Charleston sites, 16 and 18 years after their respective JP-4 releases, also are underestimated by the zero-order curve, but to a lesser degree. Conversely, the first-order curve provides a reasonable approximation for the rapid depletion of BTEX initially observed at Shaw AFB. The benzene first-order curve shown in Figure 5.4 provides a reasonably good approximation of benzene weathering in mobile LNAPL at the five JP-4 sites. As illustrated by Figures 5.3 and 5.4, the Eaker AFB average concentrations for total BTEX and benzene appear to better match that expected for a spill release that is between 1 and 5 years old, not 24 years old.

Considering the total BTEX and benzene mobile LNAPL data for the JP-4 sites taken as a whole, the first-order curves shown in Figures 5.3 and 5.4 provide default values for total BTEX and benzene weathering, respectively. Based on these results, it appears that a default first-order rate for total BTEX weathering from JP-4 mobile LNAPL could be assumed to be approximately 16 %/yr. If the Eaker data is not considered, the first-order total BTEX weathering rate is approximately 20 %/yr (Figure 5.3). For benzene weathering, a first-order weathering rate of approximately 26 %/yr is estimated considering all the JP-4 site data. If the Eaker data is excluded, the mobile LNAPL benzene weathering rate is approximately 32 %/yr. The zero-order rates shown on Figures 5.3 and 5.4 generally are less affected by the exclusion of the Eaker AFB site data.

5.2.3.3 Dissolution-Dominated Weathering

JP-4 site data suggests that BTEX mobile LNAPL weathering rates at the JP-4 sites is predominantly a function of dissolution. As discussed in Sections 2.3.1 and 2.3.2, dissolution and volatilization of LNAPL compounds are a function of their concentration in the mobile LNAPL. As concentrations in the mobile LNAPL decrease, the compound depletion rate decreases. This Raoult's Law behavior is apparent in the first-order weathering trend illustrated by Figures 5.3 and 5.4. Applied to this study, it appears that the decreased BTEX depletion rate with time and weathering, is likely the result of ever decreasing BTEX dissolution flux to groundwater and/or ever decreasing BTEX volatilization flux to soil gas.

FIGURE 5.3 TOTAL BTEX WEATHERING CONSIDERING AVERAGE DATA FROM THE JP-4 RELEASE SITES FUEL WEATHERING STUDY



FIGURE 5.4 BENZENE WEATHERING CONSIDERING AVERAGE DATA FROM THE JP-4 RELEASE SITES FUEL WEATHERING STUDY



Time (yrs)

A comparison of weathering rates among compounds and among sites (Table 5.2) indicates that dissolution may be the predominant weathering mechanism acting to reduce chemical concentrations in mobile LNAPL. Benzene and toluene weathering rates generally are higher than ethylbenzene and xylene weathering rates. Also, weathering rates for xylenes are generally higher than ethylbenzene weathering rates. One possible explanation for this is dissolution-dominated weathering where benzene, toluene, and xylenes are more rapidly reduced in mobile LNAPL than is ethylbenzene because of their higher effective water solubilities (Table 2.1).

A comparison of BTEX weathering rates at the various JP-4 sites also indicates that BTEX weathering may be dominated by dissolution. At the Eaker AFB site, evidence of minimal BTEX weathering 24 years following the reported JP-4 release may be confounded by a more recent fuel release. Assuming the mobile LNAPL at the Eaker AFB site is actually 3 years old, the total BTEX concentration in the source area (3.5 wt%) still is much higher than that at the Shaw AFB site (1.94 wt%) 3 to 4 years following release. One likely explanation for the higher BTEX concentration and lower mobile LNAPL weathering rate at the Eaker AFB site is the significantly lower groundwater velocities that have been observed at this site. Under equilibrium conditions, lower groundwater velocities would create a lower dissolution flux for mobile LNAPL depletion (Section 2.4.2). As shown on Table 3.1, the Eaker AFB site has the lowest estimated groundwater velocity, 16 feet per year (ft/yr) of the four sites where groundwater velocity and mobile LNAPL data are available. Significantly higher groundwater velocities have been observed at the Shaw AFB (400 ft/yr), Myrtle Beach AFB (420 ft/yr), and DFSP-Charleston (62 ft/yr) sites. No information was obtained regarding groundwater velocity for the McChord AFB site; however, high precipitation rates in the Seattle/Tacoma area (Figure 3.1) are likely to enhance BTEX dissolution, much the same way as high groundwater velocity.

5.2.3.4 Weathering and Spill Age

As shown on Table 5.2, mobile LNAPL weathering rates for total BTEX generally decrease with increasing spill age. This is particularly evident comparing the average total BTEX reduction rates at the Shaw AFB, Myrtle Beach AFB, DFSP-Charleston, and Eaker AFB sites. Average zero-order reduction rates for total BTEX were estimated as 18, 5.1, 4.3, and 1.7 %/yr, respectively, for these four sites. Average first-order weathering rates for total BTEX were measured as 23, 11, 11, and 2.9 %/yr, respectively. A similar trend of decreasing weathering rates with spill age also was apparent for the naphthalene compounds. However, first-order benzene depletion from JP-4 mobile LNAPL does not appear to be significantly influenced by spill age as indicated by the average benzene data shown on Table 5.2 and in Figure 5.4. Benzene data from the Shaw AFB, Myrtle Beach AFB, and DFSP-Charleston sites all indicate that benzene depletion in excess of 19 %/yr (first-order) occurs during the first 20 years of mobile LNAPL weathering.

5.2.3.5 Site-Specific Weathering Based on Multiple Sampling Events

At the Shaw AFB and DFSP-Charleston sites, mobile LNAPL samples were collected from the same site monitoring wells during multiple sampling events. At the Shaw AFB site, mobile LNAPL samples were collected from site monitoring wells approximately 3 years and 4 years after the JP-4 release (Table 4.1). At the DFSP-Charleston site, mobile

LNAPL samples were collected from site monitoring wells approximately 18 years, 20 years, and 22 years following the fuel release. For these two sites, BTEX concentrations detected in mobile LNAPL during these sampling events were plotted with the assumed initial BTEX concentrations in fresh JP-4 jet fuel (Smith *et al.*, 1981). A simple best-fit regression analysis was then performed on the plotted data to determine zero-order and first-order weathering rate constants and BTEX reduction rates.

5.2.3.5.1 Shaw AFB

BTEX weathering in JP-4 mobile LNAPL at the Shaw AFB site assuming zero-order and first-order decay are presented on Figures 5.5 and 5.6, respectively. As shown on the figures, rate constants (k_0 and k_1) and reduction rates were determined using analytical results from EAL and NRMRL. Zero-order and first-order rates estimated based on the NRMRL sample results were slightly higher than rates determined using the EAL results. This is attributable to the fact that the BTEX concentrations in mobile LNAPL as determined by EAL always were slightly higher than the NRMRL results.

Figure 5.5 suggests that zero-order BTEX reduction at the Shaw AFB site approximately 4 years after the fuel release is occurring at 13 to 18 %/yr (based on the more conservative EAL data). Figure 5.6 suggests that first-order BTEX reduction is occurring at 17 to 26 %/yr (based on EAL data). Consistent with the one-point weathering analysis (Table 5.2), weathering rates in mobile LNAPL appear to be most significant for toluene. Weathering results presented on Figures 5.5 and 5.6 suggest that benzene weathering is second-most significant, followed by ethylbenzene and xylenes. The average zero-order and first-order reduction rates shown in Table 5.2 for the Shaw AFB site provide a good estimate of the amount of BTEX depletion occurring in mobile LNAPL at monitoring well MW1610-2.

Comparing Figures 5.5 and 5.6, both the zero-order and first-order weathering rate assumptions appear to be valid for the limited data shown. No conclusions can be reached regarding whether zero-order or first-order weathering more accurately depicts BTEX depletion in mobile LNAPL at the Shaw AFB site. Nonetheless, the data plotted in Figures 5.5 and 5.6 demonstrate that the initial BTEX concentration assumption (i.e., values determined by Smith *et al.* [1981]) for JP-4 is reasonable. For some of the weathering rate curves shown on Figures 5.5 and 5.6, coefficient of determination (\mathbb{R}^2) values measured as high as 1.0, indicating no variance between the data and the predictive trend line (see Appendix C).

5.2.3.5.2 DFSP-Charleston

Zero-order and first-order BTEX weathering for the DFSP-Charleston site based on mobile LNAPL samples results at extraction well EW-6 are presented on Figures 5.7 and 5.8, respectively. Weathering rates shown on these two figures were determined using NRMRL analytical results. Zero-order rates for mobile LNAPL collected from EW-6 (Figure 5.7) are slightly higher than the average rates shown on Table 5.2. However, first-order rates shown on Figure 5.8 are substantially higher than the average rates shown on Table 5.2. The high total BTEX depletion rate shown on Figure 5.8 is a result of complete or near complete depletion of benzene and toluene from the EW-6 mobile LNAPL sample collected in 1997. The actual progression ("shape") of the BTEX weathering curves cannot be determined from the limited data shown on Figure 5.7 and









5.8. However, it is evident from these plots that BTEX compounds have been almost completely removed from mobile LNAPL in the vicinity of EW-6.

Comparing total BTEX weathering in samples collected from three wells at the DFSP-Charleston site provides some insight into the spatial differences possible in mobile LNAPL weathering (Figure 5.9). Assuming first-order weathering, total BTEX reductions at the DFSP-Charleston site appear to range from 7 to 17 %/yr depending upon sample location. A review of the DFSP-Charleston site data did not provide any indication as to why the total BTEX weathering rates vary to this degree. Each of these monitoring wells is located downgradient from the original spill location and within approximately 70 feet of each other. Extraction well EW-6 does not appear to be located in a different hydrogeologic setting relative to monitoring wells MW-103 and WQ27B. However, visual observations of mobile LNAPL samples collected in May 1997 indicated that the mobile LNAPL from EW-6 was darker in color and likely more weathered than the sample collected from MW-103. This underscores the importance of collecting several mobile LNAPL samples from each site so that an average weathering rate can be calculated.

5.2.4 Weathering in JP-5 Mobile LNAPL

Weathering rates for BTEX, naphthalene, and methylnaphthalene compounds were evaluated at two JP-5 release sites: Beaufort MCAS and Cecil Field NAS. One mobile LNAPL sampling event was performed at each site during May 1997. The approximate spill ages during this sampling event were 7 years for the Beaufort MCAS site and 16 years for the Cecil Field NAS site.

Assumed initial concentrations of BTEX, naphthalene, and methylnaphthalenes in fresh JP-5 were based on two data sets. For the Beaufort MCAS site, the initial mobile LNAPL concentrations were assumed to equal concentrations detected by NRMRL in a fresh JP-5 sample from Beaufort MCAS. For the Cecil Field NAS site, the initial mobile LNAPL concentrations were assumed to equal concentrations reported by Hughes *et al.* (1984) for fresh JP-5.

BTEX, naphthalene, and methylnaphthalene weathering rates at the two JP-5 sites were evaluated using one-point weathering rates based on equations 5.3 and 5.7 for zero-order weathering, and equations 5.4 and 5.8 for first-order weathering. Table 5.3 shows the results of this analysis.

5.2.4.1 Beaufort MCAS

At the Beaufort MCAS site, zero-order and first-order weathering rates were found to range between 4.1 and 8.2 %/yr and 4.7 and 12 %/yr, respectively, for the target compounds. Because the benzene concentrations detected in mobile LNAPLs at the site were higher than the assumed initial concentration, no weathering rate constants or reduction rates could be determined for this compound. The estimated reduction rates for toluene, ethylbenzene, and xylenes indicate that these compounds are being depleted from JP-5 mobile LNAPL at approximately the same rate. As shown in Table 5.3, assumed initial concentrations and average remaining concentrations for xylenes are

FIGURE 5.9 COMPARISON OF FIRST-ORDER TOTAL BTEX WEATHERING AT THREE WELLS DFSP-CHARLESTON, SOUTH CAROLINA FUEL WEATHERING STUDY



TABLE 5.3 BTEX AND NAPHTHALENE (ONE-POINT)^{a'} WEATHERING RATES IN JP-5 MOBILE LNAPL FUEL WEATHERING STUDY

			Assumed Initial	Average Remaining				
Site	Approximate	Number of	Conc. $(C_0)^{c/}$	Conc.(C)	ZER	O ORDER	FIRST C	ORDER
Analyte	Spill Age ^{b/}	Samples	(wt%) ^{d/}	(wt%)	Rate Constant k ₀ ^{e/}	%Co Reduction/Year ^{f/}	Rate Constant k1 ^{g/}	% Reduction/Year ^{h/}
<u>Beaufort Tank Farm C, SC</u>	7 years							
Benzene		2	0.0002	0.0003	i/			
Toluene		2	0.0047	0.0021	0.0004	7.8	0.117	11
Ethylbenzene		2	0.042	0.017	0.003	8.2	0.126	12
Total Xylenes		2	0.24	0.10	0.019	8.0	0.125	12
Total BTEX		2	0.29	0.12	0.023	8.1	0.124	12
Naphthalene		1	0.12	0.06	0.009	7.4	0.105	9.9
1- Methylnaphthalene		1	0.23	0.16	0.009	4.1	0.048	4.7
2- Methylnaphthalene		1	0.29	0.19	0.015	5.1	0.064	6.2
Cecil Field NAS, FL	16 years							
Benzene		2	0	0.003				
Toluene		2	0	0.02				
Ethylbenzene		2	0	0.36				
Total Xylenes		2	0.02	0.76				
Total BTEX		2	0.02	1.13				
Naphthalene		1	0.89	0.24	0.041	4.6	0.083	8.0
1- Methylnaphthalene		1	0.27	0.29				
2- Methylnaphthalene		1	0.45	0.42	0.002	0.51	0.005	0.53

Note: Calculated values shown have been rounded.

a/ Analyte weathering rates in free-phase product calculated based on an assumed initial concentrations in fresh JP-5 jet fuel and one point in time free-phase product sample results.

b/ Approximate age of the spill as of the most recent sampling event.

c/For Beaufort MCAS results, the assumed initial concentration of analytes is equal to the NRMRL concentration for a fresh JP-5 sample collected from Beaufort MCAS in May 1997. For Cecil Field results,

Hughes et al. (1984) JP-5 composition values were used.

d/ wt% = weight percent.

 $^{e^{\prime}}k_{0}$ = zero-order rate contstant or slope calculated using equation 5.3; units in weight percent per year.

[#] Annual mass fraction reduction as a percent of the initial concentration; calculated using equation 5.4.

 $k_1 =$ first-order rate constant or exponential decay rate calculated using equation 5.7; units in years⁻¹ or 1/years.

^{h/} Weight percent reduction per year calculated using equation 5.8.

i/ --- = negative value; measured concentration is greater than assumed initial concentration.

approximately 2 orders of magnitude higher than the toluene concentrations, and 1 order of magnitude higher than the ethylbenzene concentrations. Nonetheless, concentrations for each of these compounds are well below their respective concentrations in JP-4 (Figure 2.3). In view of these significantly lower fresh fuel and mobile LNAPL concentrations, groundwater MCLs for toluene, ethylbenzene, and xylenes are unlikely to be exceeded by partitioning of these compounds from the fuel into site groundwater (Section 5.1.2).

BTEX, naphthalene, and methylnaphthalene weathering rates at the Beaufort Tank Farm C site generally fall within the same range as the JP-4 rates shown in Table 5.2 for these compounds. A rough interpolation of the average JP-4 reduction rates shown in Table 5.2 for Shaw AFB (a 4-year old spill) and Myrtle Beach AFB (a 16-year old spill) would give rates comparable to those determined for JP-5 at the Beaufort Tank Farm C site (a 7-year old spill).

5.2.4.2 Cecil Field NAS

At the Cecil Field NAS site, weathering rates could be estimated only for naphthalene and 2-methylnaphthalene. Mobile LNAPL concentrations for all other analytes exceeded the assumed initial values. Relative to the naphthalene and 2-methylnaphthalene rates shown for the Beaufort MCAS site, estimated reduction rates at the Cecil Field NAS site are substantially lower.

5.2.5 Weathering in JP-8 Mobile LNAPL

Weathering rates for BTEX, naphthalene, and methylnaphthalene compounds were evaluated at two JP-8 release sites: Pope AFB and Seymour Johnson AFB. One mobile LNAPL sampling event was performed at the Pope AFB site in July 1996 approximately 3 months after the fuel release. Three mobile LNAPL sampling events were performed at the Seymour Johnson AFB site in the first 2.25 years following the fuel release.

5.2.5.1 Pope AFB

Approximately 3 months after the JP-8 release at Pope AFB, the US Army Corps of Engineers (USACE) collected one mobile LNAPL sample for analysis by NRMRL. No benzene or toluene were detected in the mobile LNAPL sample and ethylbenzene and xylenes concentrations were reduced approximately 90 percent compared to assumed initial concentrations. Zero-order and first-order reduction rates were calculated based on this one sample result; however, the nondetect or near nondetect concentrations of BTEX in the sample prevented meaningful determination of LNAPL weathering. Compared to the other fuel release sites, a relatively small volume (700 gallons) of fuel was released at the Pope AFB site (Table 3.1) and no free-phase product was evident at the site approximately 6 months after the fuel release (Dalzell, 1997). It appears that significant volatilization may have immediately reduced the BTEX fraction of the small spill.

5.2.5.2 Seymour Johnson AFB

One-point weathering rates determined for the Seymour Johnson AFB site are presented in Table 5.4. Weathering rates presented in Table 5.4 are based on eight samples collected over approximately 2 years from one monitoring well. As with the JP-4 data shown in Table 5.2, the estimated weathering rates calculated from the Seymour Johnson AFB samples vary significantly. The average reduction rates suggest that weathering is slowest for ethylbenzene, most likely as a result of its lower effective water solubility (effective water solubility values for JP-8 were not identified in the literature; however, the relative effective water solubility values for the BTEX compounds are expected to be very similar to those shown on Table 2.1 for JP-4 and gasoline).

Mobile LNAPL sample results from the multiple sampling events and zero-order and first-order analysis of the Seymour Johnson JP-8 data are shown in Figures 5.10 and 5.11, respectively. Similar to the JP-4 weathering results, zero-order and first-order rates estimated based on the NRMRL analytical results were slightly higher than rates determined from the EAL results. EAL data used to estimate reduction rates were collected during two sampling events. NRMRL rates are based on data from three sampling events (Table 4.1).

Figure 5.10 indicates that zero-order reduction rates during 2.25 years of product weathering range from 6 %/yr for ethylbenzene to 33 %/yr for benzene (based on EAL and NRMRL data). The overall BTEX zero-order reduction rates were estimated to be 22 %/yr based on the EAL data and 27 %/yr based on NRMRL data. Figure 5.11 suggests that first-order reduction for the BTEX compounds is occurring at 6 to 52 %/yr (considering EAL and NRMRL data). The total BTEX first-order decay rates were 26%/yr for the EAL data and 36%/yr for the NRMRL data. Weathering rates in the JP-8 mobile LNAPL also appear to support the presumption that mobile LNAPL weathering is dissolution-dominated for the aromatic compounds. Compound-specific reduction rates are highest for benzene, followed by toluene, xylenes, and ethylbenzene. Other observations include:

- Average one-point rate values shown in Table 5.4 appear to be relatively conservative compared to the rates determined from best-fit analyses of the site data shown on Figures 5.10 and 5.11;
- The initial assumed BTEX concentrations for JP-8 (Mayfield, 1996) are well supported by the analytical results plotted on Figures 5.10 and 5.11; and
- Both zero-order and first-order BTEX weathering rates appear to be valid during the first 2 years of mobile LNAPL weathering.

5.2.6 Weathering in Gasoline Mobile LNAPL

BTEX weathering analysis in gasoline mobile LNAPL was evaluated at the Offutt AFB site. Compared to the other fuel weathering sites, where the dates of the fuel release are known with some confidence, mobile LNAPL at the Offutt AFB site most likely is the result of chronic, long-term leaking from former USTs which ended in

TABLE 5.4 BTEX AND NAPHTHALENE (ONE-POINT)^{a/} WEATHERING RATES IN JP-8 MOBILE LNAPL FUEL WEATHERING STUDY

			Assumed Initial	Average Remaining	ZERO ORDER							FIRST ORDER							
Site	Approximate	Number of	Conc. (C _o) ^{c/}	Conc. (C)	Rate	Constant	k ₀ e/	%Co l	Reduction	/Year ^{f/}	Γ	Rate	Constant	k1 ^{g/}	% R	eduction/Y	'ear ^{h/}		
Analyte	Spill Age ^{b/}	Samples	(wt%) ^{d/}	(wt%)	min	max	avg	min	max	avg		min	max	avg	min	max	avg		
Seymour Johnson AFB, NC	2 years																		
Benzene		8	0.03	0.02	0.00	0.01	0.01	5.7	36	23		0.06	0.76	0.39	5.6	53	29		
Toluene		8	0.21	0.12	0.00	0.06	0.05	2.1	28	23		0.02	0.46	0.32	2.1	37	27		
Ethylbenzene		8	0.15	0.13	i/	0.04	0.01		23	6.6			0.29	0.08		25	7.2		
Total Xylenes		8	1.13	0.67	0.16	0.33	0.27	15	30	24		0.16	0.49	0.33	15	39	28		
Total BTEX		8	1.52	0.94	0.21	0.43	0.34	14	28	22		0.15	0.45	0.30	14	36	25		
Naphthalene		4	0.25	0.13	0.06	0.34	0.13	24	39	28		0.29	0.95	0.48	25	61	36		
1- Methylnaphthalene		4	0.43	0.17	0.05	0.16	0.12	19	38	31		0.24	0.56	0.45	22	43	35		
2- Methylnaphthalene		4	0.35	0.26	0.04	0.10	0.06	11	22	15		0.12	0.31	0.19	11	27	17		

Note: Calculated values shown have been rounded.

a' Analyte weathering rates in free-phase product calculated based on assumed initial analyte concentrations in fresh JP-8 fuel and one point in time free-phase product sample results.

^{b'} Approximate age of spill as of the most recent sampling event.

c/ Assumed initial concentrations from Mayfield, 1996.

d/ wt% = weight percent.

 $^{e'}$ k₀ = zero-order rate contstant or slope calculated using equation 5.3; units in weight percent per year.

^{f/} Annual mass fraction reduction as a percent of the initial concentration; calculated using equation 5.4.

 $^{g/}k_1$ = first-order rate constant or exponential decay rate calculated using equation 5.7; units in years $^{-1}$ or 1/years.

^{b/}Weight percent reduction per year calculated using equation 5.8.

 i^{i} --- = negative value; measured concentration is greater than the assumed initial concentration.





1990 when the gasoline tanks were closed. The Offutt AFB site was selected for the study because of the existence of historical BTEX analytical results for mobile LNAPL.

5.2.6.1 One-Point Weathering Rates

BTEX one-point weathering rates for the Offutt AFB site are presented in Table 5.5. Rates provided in this table were calculated from BTEX analytical results for each of four LNAPL sampling events conducted following closure of the source USTs. Initial BTEX concentrations were assumed to be equal to those determined by Ghassemi *et al.* (1984) for fresh gasoline. Initial evaluation of the site weathering rates indicated significant variability by sampling location; therefore, one-point weathering rates are presented in Table 5.5 for the individual monitoring wells from which mobile LNAPL samples were collected.

Comparison of the mobile LNAPL sample results with the assumed initial concentrations in fresh gasoline indicates that the mobile LNAPL at the Offutt AFB site is only slightly weathered. The most significant BTEX reductions in mobile LNAPL were observed in the sample collected from MW349-8 approximately 6 years after tank closure. At this location, greater than 90 percent of the benzene was depleted from the mobile LNAPL, and the total BTEX concentration was reduced by approximately 50 percent. Higher weathering rates (i.e. lower analyte concentrations) at MW349-7 and MW349-8 relative to MW349-1 may be the result of their locations further from the original source area. Mobile LNAPL at wells MW349-7 and MW349-8 likely is older than that at MW349-1. In all cases, benzene and toluene appear to have weathered at faster rates than xylenes and ethylbenzene. Little to no reduction in ethylbenzene concentrations has occurred in mobile LNAPL at the site. Relatively low groundwater velocities and resulting lower dissolution potential at the Offutt AFB site may be a primary reason for the lower BTEX weathering rates (Table 3.1).

5.2.6.2 Weathering Rates Based on Multiple Sampling Events

At Offutt AFB monitoring well MW349-1, mobile LNAPL samples were collected in November 1994, June 1996, June 1997, and October 1998 to assess changes in BTEX concentrations. Mass fraction (wt%) analytical results from these sampling events are plotted on Figures 5.12 and 5.13. On Figure 5.12, the initial BTEX concentrations determined by Ghassemi *et al.* (1984) for fresh gasoline were assumed to equal the concentrations in mobile LNAPL at the time of tank closure. In Figure 5.13, the initial BTEX values were assumed to equal mid-range values based on gasoline compositional data from AD Little (1987), Sigsby *et al.* (1987), and Potter (1988). Zero-order weathering rates were assumed for both plots.

As is evident on Figures 5.12 and 5.13, the best-fit linear curves and resulting BTEX weathering rates are predominantly controlled by the assumed initial compound concentrations. The analytical results from the four mobile LNAPL sampling events do not clearly indicate a weathering trend for any of the BTEX compounds. Also, the unknown age of the mobile LNAPL at monitoring well MW349-1 and the wide range of possible BTEX concentrations in the original fuel hindered assessment of weathering rates at this location.

TABLE 5.5 BTEX (ONE-POINT)^{a/} WEATHERING RATES IN GASOLINE MOBILE LNAPL TANK 349, OFFUTT AFB, NEBRASKA FUEL WEATHERING STUDY

Time Since Tank Closure \sim Concentration $(C_0)^{C'}$	Concentr	ation (C)		ZERO	O ORDER		FIRST ORDER					
Analyte (wt%) ^{d/}	(wi	t%)	Rate Con	stant $k_0^{e/}$	%C _o Redu	ced/Year ^{f/}	Rate Cor	istant $k_1^{g/}$	% Reduct	ion/Year ^{h/}		
4.5 years	MW349-1	MW349-7	MW349-1	MW349-7	MW349-1	MW349-7	MW349-1	MW349-7	MW349-1	MW349-7		
Benzene 1.5	1.1	0.8	0.08	0.17	5.7	11	0.07	0.15	6.3	14		
Toluene 5.9	5.6	3.9	0.07	0.45	1.3	7.7	0.01	0.09	1.3	9.0		
Ethylbenzene 1.3	1.4	1.3	i/	0.01		0.75		0.01		0.76		
Total Xylenes 5.9	5.7	5.7	0.04	0.05	0.75	0.81	0.01	0.01	0.76	0.82		
Total BTEX 14.6	13.8	11.6	0.18	0.68	1.2	4.6	0.01	0.05	1.3	5.0		
<u>6 years</u>	MW349-1	MW349-8	MW349-1	MW349-8	MW349-1	MW349-8	MW349-1	MW349-8	MW349-1	MW349-8		
Benzene 1.5	1.2	0.1	0.06	0.23	3.7	15	0.04	0.40	4.1	33		
Toluene 5.9	5.0	1.7	0.14	0.70	2.4	12	0.03	0.21	2.6	19		
Ethylbenzene 1.3	1.6	1.4										
Total Xylenes 5.9	5.2	4.4	0.11	0.25	1.9	4.2	0.02	0.05	2.0	4.7		
Total BTEX 14.6	13.0	7.5	0.27	1.16	1.8	8.0	0.02	0.11	1.9	10		
7 years	MW349-1	MW349-6	MW349-1	MW349-6	MW349-1	MW349-6	MW349-1	MW349-6	MW349-1	MW349-6		
Benzene 1.5	1.2	1.3	0.05	0.03	3.3	1.8	0.04	0.02	3.6	1.9		
Toluene 5.9	5.6	5.6	0.05	0.04	0.83	0.66	0.01	0.01	0.85	0.68		
Ethylbenzene 1.3	1.5	1.8										
Total Xylenes 5.9	6.0	6.1										
Total BTEX 14.6	14.2	14.8	0.05		0.37		0.00		0.38			
8 years	MW349-1	MW349-6	MW349-1	MW349-6	MW349-1	MW349-6	MW349-1	MW349-6	MW349-1	MW349-6		
Benzene 1.5	1.0	0.3	0.06	0.14	3.8	9.1	0.05	0.17	4.5	16		
Toluene 5.9	4.8	3.4	0.13	0.29	2.2	4.9	0.02	0.06	2.4	6.2		
Ethylbenzene 1.3	1.6	1.4										
Total Xylenes 5.9	6.1	6.1										
Total BTEX 14.6	13.6	11.2	0.12	0.40	0.85	2.8	0.01	0.03	0.88	3.1		

Note: Calculated values shown have been rounded.

^{ar} Analyte weathering rates in free-phase product calculated based on assumed initial analyte concentrations in fresh gasoline and one point in time free-phase product results.

^{b/} Approximate time between date USTs were taken out of service and date of sampling event.

c/ Ghassemi et al ., 1984.

^{d/} wt% = weight percent.

 $e' k_0$ = zero-order rate contstant or slope calculated using equation 5.3; units in weight percent per year.

^{f/} Annual mass fraction reduction as a percent of the initial concentration; calculated using equation 5.4.

 $g' k_1$ = first-order rate constant or exponential decay rate calculated using equation 5.7; units in years⁻¹ or 1/years.

^{h/} Weight percent reduction per year calculated using equation 5.8.

i' --- = negative value; measured concentration is greater than the assumed initial concentration.





One possible explanation for the lack of an observed weathering trend in the analytical data is the fact that the 4.5- and 6-year mobile LNAPL samples were collected from monitoring well MW349-1 without first purging the monitoring well. As shown in Table 5.5, the lower BTEX concentrations during the 4.5- and 6-year monitoring events may be the result of evaporative losses of BTEX while the mobile LNAPL sat in the monitoring well. If only the June 1997 (year 7) and October 1998 (year 8) mobile LNAPL data are considered, reductions in benzene, toluene, and total BTEX concentrations can be used to estimate initial mobile LNAPL concentrations at the time of tank closure. Figures 5.14 and 5.15 provide reduction rates in mobile LNAPL at MW349-1 based on zero-order and first-order back-calculations, respectively. Based on this analysis, rates of total BTEX reduction in mobile LNAPL at MW349-1 appear to be most similar to that observed at the Eaker AFB site (Table 5.2). At both the Eaker AFB site and the Offutt AFB site, slower groundwater velocities may be responsible for the lower BTEX weathering rates observed in mobile LNAPL.

5.3 FUEL/WATER PARTITIONING COEFFICIENTS (KFW)

Mobile LNAPL and groundwater data from the eight primary sites selected for the study were used to determine "field" and "laboratory" fuel/water partitioning coefficients (K_{fw}) for the BTEX compounds. Recall that in Section 2.3.1.2, K_{fw} is defined as the concentration of a compound in fuel (C_f) divided by its equilibrium concentration in water in contact with the fuel (C_w) (equation 2.2). The "field" K_{fw} was calculated using groundwater and mobile LNAPL analytical results from NRMRL. The "laboratory" K_{fw} values were determined using mobile LNAPL samples in a laboratory partitioning experiment performed by EAL in accordance with procedures outlined by Cline *et al.* (1991) with the purpose of creating equilibrium conditions (Section 4.4.2). The "field" and "laboratory" K_{fw} s were determined to evaluate the validity of the equilibrium assumption (Section 2.3.1.3) when performing dissolution modeling. Values for K_{fw} determined from field and laboratory data are presented in Table 5.6.

The EAL ("laboratory") K_{fw} values for the BTEX compounds were expected to be lower than the NRMRL ("field") K_{fw} values because laboratory mixing and dissolution was expected to produce maximum or equilibrium concentrations in deionized water in contact with the fuel LNAPL. However, the "field" K_{fw} values determined from mobile LNAPL and actual groundwater results generally are lower than the "laboratory" values. Of the ten field and laboratory data sets presented in Table 5.6, only data from the Myrtle Beach AFB and Offutt AFB sites generally conformed to the initial prediction. For the other sites, the field K_{fw} values for the BTEX compounds were generally lower than the laboratory values indicating higher BTEX concentrations in groundwater than in the deionized water analyzed after a laboratory equilibrium procedure. The comparison of field and laboratory data generally suggest that dissolution in site groundwater samples collected from within the mobile LNAPL source area may be more complete (i.e., closer to equilibrium) than the results obtained from the laboratory partitioning experiment. The lower than expected concentrations in the aqueous phase samples analyzed by EAL as compared to the NRMRL groundwater results possibly could be partially attributed to various water solubility effects (including pH, temperature, pressure, salinity) and differences in analytical methods.




TABLE 5.6 FUEL/WATER PARTITIONING COEFFICIENTS FOR FUEL RELEASE SITES FUEL WEATHERING STUDY

		Fuel/Water Partitioning Coefficients (K _{fw})						
	Approximate			Ethyl-				
Data Source/Site	Spill Age ^a	Benzene	Toluene	benzene	o-Xylene	m-Xylene	p-Xylene	Total Xylenes
Gasoline								
Cline et al., 1991	Fresh Gasoline	350	1,250	4,500	3,630	4,350	4,350	NA ^{b'}
Offutt AFB, NE (MW349-6, 1997)	8 years							
EAL ("Laboratory") ^{c/}		280	788	1,200	NA	NA	NA	1,267
NRMRL ("Field") ^{d/}		246	926	3,333	2,366	3,466	3,209	3,056
JP-4 Jet Fuel								
Smith et al., 1981	Fresh JP-4	2,455	2,754	4,786	7,079	3,715	7,586	NA
Shaw AFB, SC (MW1610-2, 1997)	3 years							
EAL ("Laboratory")		265	945	4,059	2,837	NA	NA	3,220
NRMRL ("Field")		266	554	1,138	971	1,041	1,093	1,024
Shaw AFB, SC (MW1610-2, 1998)	4 years							
EAL ("Laboratory")		217	789	3,415	2,583	NA	NA	2,903
NRMIRL (Field)		803	1,044	3,042	2,510	2,646	3,069	2,384
Myrtle Beach AFB, SC (MW-8I)	16 years	<i>c/</i>	0/					
EAL ("Laboratory")		203	1,508	3,553	13,909	NA	NA	3,634
NRMIRL (Field)		335	2,693	39,420	7,565	53,005	135,000	58,788
DFSP-Charleston, SC (MW-103)	22 years	2/	0/					
EAL ("Laboratory")		5	1,020	3,692	NA 2 702	NA	NA 1.044	3,600
NRMIRL (Field)		1	251	1,600	2,705	2,555	1,944	2,416
Eaker AFB, AR (MW-316)	24 years	205	10.000	5 500	1 200			1.000
EAL ("Laboratory") NRMRI. ("Field")		385	40,000	5,500 4 205	4,300 9.486	NA 5 405	NA 5.080	4,696
		101		1,205	3,100	5,105	5,000	5,007
JP-5 Jet Fuel								
Beaufort MCAS, SC Laboratory (EAL)	Fresh JP-5 Sample	455	1.500	4.568	NA	NA	NA	4.815
Beaufort Tank Farm C. SC (MW BET 401.3)	7 years		-,	.,				.,
EAL ("Laboratory")	/ years	558 ^{e/}	1 250	4 571	2 538	NA	NA	3 741
NRMRL ("Field")		341	345	761	1,283	925	1,246	1,116
Cecil Field NAS. FL (MW CEF-293-9)	16 years							
EAL ("Laboratory")		253 ^{e/}	1.470 ^{e/}	5.818	NA	NA	NA	6.636
NRMRL ("Field")		261	152	2,669	635	1,867	2,221	1,740
JP-8 Jet Fuel								
Seymour Johnson AFB, NC (MW-1S, 1997)	1.5 years							
EAL ("Laboratory")		240	1,005	3,127	NA	NA	NA	3,087
NRMRL ("Field")		229	251	1,390	1,860	1,891	1,729	1,850
Seymour Johnson AFB, NC (MW-1S, 1998)	2.3 years							
EAL ("Laboratory")		278	797	3,548	2,745	NA	NA	3,409
NRMRL ("Field")		59	220	1,448	1,459	1,165	1,361	1,289

^{a/} Approximate age of spill as of the most recent sampling event.

 $^{b/}$ NA = not available or not analyzed.

e' Results calculated from Evergreen Analytical Laboratory (EAL) partitioning experiment values for BTEX in mobile LNAPL and deionized water in contact with

the mobile LNAPL.

d^d Results calculated from mobile LNAPL and groundwater concentrations as determined by the National Risk Management Research Laboratory (NRMRL).

 $^{e'}$ K_{fw} calculated using NRMRL mobile LNAPL analytical result because EAL result was less than the laboratory reporting limit.

Results of the fuel/water partitioning experiment do not refute use of equilibrium assumptions in estimating groundwater concentrations of BTEX compounds at gasoline and jet fuel release sites. However, the data presented in Table 5.6 suggest that using laboratory predictions of K_{fw} to estimate equilibrium concentrations in groundwater in the LNAPL source area may sometimes underestimate actual groundwater concentrations. For example, if the benzene concentration in mobile LNAPL at a JP-4 release site is known to be 1,250 mg/L, the Smith et al. (1981) K_{fw} value of 2,455 will indicate that the concentration in groundwater is approximately 0.51 mg/L (obtained by rearranging equation 2.2 to solve for C_w; C_w = 1,250 [mg/L] / 2,455 = 0.51 mg/L). However, K_{fw} results presented in Table 5.6 indicate that a benzene K_{fw} of 265 is more appropriate because the benzene concentration in source area groundwater is approximately 4.72 mg/L (K_{fw} = 1,250 [mg/L] / 4.72 [mg/L] = 265).

A consistent relationship between mobile LNAPL weathering and fuel/water partitioning was not observed. This is specifically demonstrated by the 1997 and 1998 K_{fw} values calculated for Shaw AFB and Seymour Johnson AFB, where field and laboratory values for K_{fw} do not vary consistently with increasing spill age. A more general review of the JP-4 sites taken as a whole, also indicates no apparent correlation between K_{fw} and spill age.

5.4 COMPARISON OF RESIDUAL AND MOBILE LNAPL WEATHERING

Weathering effects on residual-LNAPL-contaminated soils were compared to mobile LNAPL weathering in an attempt to demonstrate that LNAPL weathering is more significant in the capillary fringe soils than in the free-phase product. The primary weathering mechanisms thought to enhance weathering of residual LNAPL in soils are increased volatilization and biodegradation. Little to no BTEX depletion in mobile LNAPL is expected to occur from biodegradation, yet biodegradation is a significant weathering mechanism for residually contaminated soils (Section 2.3.3). Section 5.4.1 presents a simple quantitative evaluation of BTEX weathering in residual and mobile LNAPLs. Section 5.4.2 presents a qualitative comparison of hydrocarbon weathering in mobile and residual LNAPL through the use of soil and free product chromatograms from the DFSP-Charleston site.

5.4.1 BTEX Weathering

BTEX weathering in residual and mobile LNAPL was compared by converting soil analytical results reported on a mass per mass basis (i.e., mg/kg) to mass per volume units typical of mobile LNAPL results (µg/mL, mg/mL, or mg/L). The following relationship was used to estimate the concentration of BTEX compounds in residual LNAPL based on soil analytical results:

 $C_s = [BTEX analyte (mg/kg) / TPH (mg/kg)] x LNAPL density (µg/mL) eq. 5.9$

where: C

 C_s = estimated residual LNAPL BTEX concentration (μ g/mL) TPH = total petroleum hydrocarbons in residual LNAPL

Note: TPH concentrations in soil were estimated by NRMRL analysis of total fuel carbon.

Table 5.7 presents the BTEX concentrations in residual LNAPL estimated using equation 5.9 and compares these estimates to the mobile LNAPL analytical results at the eight primary sites.

Based on the results presented on Table 5.7, attenuated BTEX concentrations in residual LNAPL as compared to mobile LNAPL analytical results were observed in the estimates for Offutt AFB and Shaw AFB. At Offutt AFB, estimated concentrations of toluene, ethylbenzene, and xylenes in capillary fringe soils were 5 to 23 percent less than their respective concentrations in mobile LNAPL. For the 1997 Shaw AFB soil data, BTEX concentrations were 1 to 34 percent less than the mobile LNAPL analytical results. Soil samples collected at Shaw AFB in 1998 indicate far less BTEX contamination in soils at 27 feet bgs as compared to the 1997 results at 33 feet bgs.

At DFSP-Charleston the estimated residual LNAPL concentrations of BTEX greatly exceed the mobile LNAPL results. As indicated on Figures 5.7 and 5.8, the total BTEX concentration in mobile LNAPL at EW-6 has been reduced by almost 99 percent. While BTEX concentrations are extremely low in mobile LNAPL at EW6, the total BTEX concentration of 230 mg/kg in soil at 13 feet bgs near EW-6 was the highest total BTEX concentration measured in soil at any of the JP-4 sites (Table 5.1). While significant BTEX depletion is evident in mobile LNAPL, significant BTEX contamination appears to remain in some soils near this location.

Overall, residual LNAPL concentrations of BTEX in soil estimated using equation 5.9 generally exceed actual mobile LNAPL concentrations for these compounds. Predicted residual LNAPL concentrations calculated from Myrtle Beach AFB, Eaker AFB, Cecil Field NAS, and Seymour Johnson AFB soil analytical results are approximately 1 to 3 times higher than the mobile LNAPL analytical results. Wiedemeier *et al.* (1995) observed that using the BTEX/TPH relationship illustrated by equation 5.9 to compare residual and mobile LNAPL concentrations also indicated residual BTEX concentration overestimates especially within the LNAPL source area. In theory, the residual BTEX concentrations should never exceed the mobile LNAPL BTEX concentrations. A significant source of error in equation 5.9 is the TPH term. TPH analysis is prone to underestimation of the total fuel residual in the soil. Underestimation of TPH would lead to the false conclusion that the BTEX fraction in soil residuals exceeds the BTEX fraction in mobile LNAPL.

5.4.2 Comparison of Soil and Free Product Chromatograms

A qualitative comparison of residual LNAPL and mobile LNAPL weathering is possible by evaluating chromatograms of soil and free product samples. Figure 5.16 presents GC/FID results for mobile LNAPL samples collected from two wells (MW-103 and EW-6) at the DFSP-Charleston site. Soil sample GC/FID results for one soil boring (CHSB3) advanced in the original source area at the DFSP-Charleston site are presented in Figure 5.17. Results presented in these two figures are from samples collected in May 1997 and analyzed by AD Little (1998). During this sampling event, the water table surface was measured approximately 15 to 16 feet bgs.

Chromatograms for soil samples indicate that single-ring aromatic hydrocarbon concentrations in residual LNAPL increase with depth and likely approach

TABLE 5.7 COMPARISON OF ESTIMATED RESIDUAL AND MOBILE LNAPL BTEX CONCENTRATIONS FUEL WEATHERING STUDY

		Free Product		Depth to	Depth of					
Fuel Type	Sample	Sample	Approximate	Product	Soil Sample			Ethyl-	Total	Total
Site	Date	Location	Spill Age ^{a/}	(feet btoc) ^{b/}	(feet bgs) ^{c/}	Benzene	Toluene	benzene	Xylenes	BTEX
Gasoline										
Offutt AFB, NE	Nov-94	MW349-1	4	39.60	39.5					
Estimated Residual LNAPL d/ (µg/mL) e/						16,034	39,335	9,141	32,217	96,727
Mobile LNAPL ^{f/} (µg/mL)						8,280	41,100	10,300	42,080	101,760
Estimated Residual / Mobile						1.94	0.96	0.89	0.77	0.95
JP-4 Jet Fuel										
Shaw AFB, SC	Mar-97	MW1610-2	3	32.38	33.0					
Estimated Residual LNAPL (µg/mL)						2,225	3,220	916	6,619	12,980
Mobile LNAPL (µg/mL)						2,250	4,890	1,340	8,530	17,010
Estimated Residual / Mobile						0.99	0.66	0.68	0.78	0.76
Shaw AFB, SC	Mar-98	MW1610-2	4	28.24	27.0					
Estimated Residual LNAPL (µg/mL)						101	168	561	2,687	3,517
Mobile LNAPL (µg/mL)						1,250	2,830	1,040	7,180	12,300
Estimated Residual / Mobile						0.08	0.06	0.54	0.37	0.29
Myrtle Beach AFB, SC	Mar-97	MW8I	16	3.7	9.5					
Estimated Residual LNAPL (µg/mL)						1,536	8.4	3,324	9,666	14,535
Mobile LNAPL (µg/mL)						211	7.5	1,360	4,262	5,841
Estimated Residual / Mobile						7.28	1.12	2.44	2.27	2.49
DFSP-Charleston (Tank 1), SC	May-97	EW-6	22	15.92	13.0					
Estimated Residual LNAPL (µg/mL)						85	1,764	2,789	14,082	18,721
Mobile LNAPL (µg/mL)						0.025	1.35	91.3	351	444
Estimated Residual / Mobile						3415	1307	31	40	42
Eaker AFB, AR	Aug-97	MW316	24	13.86	12.0	2,658	10	4,086	20,011	26,764
Estimated Residual LNAPL (µg/mL)						900	0.025	2,960	15,400	19,260
Mobile LNAPL (µg/mL)						2.95	384	1.38	1.30	1.39
Estimated Residual / Mobile										

TABLE 5.7 COMPARISON OF ESTIMATED RESIDUAL AND MOBILE LNAPL BTEX CONCENTRATIONS FUEL WEATHERING STUDY

Fuel Type	Sample	Free Product Sample	Approximate	Depth to Product	Depth of Soil Sample			Ethyl-	Total	Total
Site	Date	Location	Spill Age ^{a/}	(feet btoc) ^{b/}	(feet bgs) ^{c/}	Benzene	Toluene	benzene	Xylenes	BTEX
JP-5 Jet Fuel										
Beaufort Tank Farm C, SC Estimated Residual LNAPL (µg/mL) Mobile LNAPL (µg/mL) Estimated Residual / Mobile	May-97	BFT-401-3	7	6.98	4.0	7.6 2.2 3.41	93 13 7.16	557 116 4.80	1,820 611 2.98	2,477 742 3.34
Cecil Field NAS, FL Estimated Residual LNAPL (µg/mL) Mobile LNAPL (µg/mL) Estimated Residual / Mobile	May-97	CEF-293-9	16	8.54	8.5	27 24 1.13	479 122 3.92	3,925 2,520 1.56	10,574 4,787 2.21	15,004 7,453 2.01
JP-8 Jet Fuel										
Seymour Johnson AFB, NC Estimated Residual LNAPL (µg/mL) Mobile LNAPL (µg/mL) Estimated Residual / Mobile	May-97	MW1S	2	5.08	5.5	386 194 1.99	2,311 1,030 2.24	2,004 1,170 1.71	10,441 5,990 1.74	15,142 8,384 1.81
Seymour Johnson AFB, NC Estimated Residual LNAPL (µg/mL) Mobile LNAPL (µg/mL) Estimated Residual / Mobile	Mar-98	MW1S	3	3.11	3.0	181 47 3.84	1,020 602 1.69	1,440 800 1.80	6,016 3,040 1.98	8,658 4,489 1.93

^{a/} Approximate age of spill at time of sampling event.

^{b/} feet btoc = feet below top of well casing.

^{c/} feet bgs = feet below ground surface.

^d Estimated mass per volume concentration in residual LNAPL calculated using equation 5.9 and NRMRL soil sample results.

 $^{e/}$ µg/mL = micrograms per milliter.

^{f/} Mobile LNAPL concentration as determined by NRMRL.

FIGURE 5.16 CHROMATOGRAMS FOR TWO JP-4 MOBILE LNAPL SAMPLES DFSP-CHARLESTON, SOUTH CAROLINA FUEL WEATHERING STUDY







FIGURE 5.17 JP-4 IMPACTED SOIL CHROMATOGRAM RESULTS WITH DEPTH DFSP-CHARLESTON, SOUTH CAROLINA FUEL WEATHERING STUDY



concentration levels consistent with mobile LNAPL near the water table. As shown in Figure 5.17, little residual LNAPL BTEX compounds remain in soils at the 3.5- and 5.5-foot depths. The sample collected from 7.5 feet bgs indicates that fuel hydrocarbons are still present in these soils, but the BTEX compounds which are removed during the first 10 minutes of chromatographic separation are completely depleted. Soil samples collected 9.5 and 11.5 feet bgs appear to retain the general signature of the JP-4 mobile LNAPL samples (Figure 5.16). BTEX weathering appears to be less significant in these deeper soils than observed in mobile LNAPL from EW-6, but more significant than that observed in free product from MW-103. As discussed in Sections 2.5.1 and 5.2.3.5.2, mobile LNAPL weathering appears to vary spatially at fuel contaminated sites. Residual LNAPL weathering rates most likely vary with proximity to saturated LNAPL lenses and mobile LNAPL pools.

In theory, more porous soils and sites without impermeable covers should promote greater residual LNAPL volatilization and biodegradation. Sites which are not subject to large and frequent water level variations should also produce a more weathered residual LNAPL because these soils would not be regularly "recontaminated" with mobile LNAPL. Based on our study, it is impossible to predict residual LNAPL weathering from the limited soil sampling performed at each site. Several samples of residual fuel contamination are needed to estimate the remaining BTEX fraction in soils at each depth interval.

SECTION 6

CONCLUSIONS

6.1 REVIEW OF PROJECT OBJECTIVES

The overall purpose of this study was to improve the scientific database for estimating natural LNAPL weathering rates and source-term reduction rates which are incorporated into natural attenuation models. Based on our literature review, little information has been published regarding rates of natural weathering of the BTEX compounds from mobile fuel LNAPLs. As a result, the rate of reduction of the contaminant source term in groundwater models is often left to professional judgment. This has generally resulted in the use of overly conservative LNAPL weathering rates to evaluate contaminant fate and transport and the suitability of natural attenuation as a remedial alternative. These conservative assumptions extend the estimated timeframe for achieving cleanup goals and inflate projected long-term monitoring and site management costs.

The primary objective of this fuel weathering study was to document a range of BTEX weathering rates for the mobile LNAPL fraction based on data collected from sites with documented mobile LNAPL plumes with known release dates. Secondary objectives of this study included an evaluation of the degree of contaminant partitioning of BTEX from mobile LNAPL to groundwater, and comparison of weathering effects on the mobile LNAPL fraction and on residual LNAPL present in capillary fringe soils. The following tasks were completed to meet these objectives:

- A literature search to assess existing information regarding weathering of LNAPLs;
- Selection of eight primary sites where the time of release is generally known and free-phase jet fuel or gasoline remain *in situ*;
- Sampling of soil, groundwater, and free-phase LNAPLs at the primary sites;
- Evaluation of data obtained from the eight primary sites, as well as data from four secondary sites, to assess contaminant concentrations in site media in relation to such factors as age of the fuel release, fuel type, and site geology and hydrogeology.

6.2 SUMMARY OF FINDINGS

• Significant research has been completed on multiple "fresh" samples of JP-4 and JP-8 so that the magnitude of the initial BTEX fraction in these fuels is well-known. The assumption that initial BTEX values in mobile LNAPL at JP-4 and JP-8 release sites are equal to concentrations reported by Smith *et al.* (1981) and Mayfield (1996) appears reasonably valid for predicting BTEX depletion in mobile LNAPL.

Initial fuel composition results for gasoline studies are more varied and results for JP-5 are very limited.

- Free-phase fuel BTEX weathering rates will vary from site to site and are influenced by many factors including spill age, the relative solubility of individual compounds, free product geometry, and the rate at which groundwater and precipitation contacts LNAPL.
- As demonstrated by the DFSP-Charleston and Offutt AFB site data, the BTEX fraction remaining in free-phase LNAPL samples collected from different locations on the same site will vary. It is likely that samples collected near the center of the LNAPL volume will exhibit lower rates of weathering than samples collected at the leading edge of the LNAPL "plume." A more accurate estimate of LNAPL weathering can be obtained by collecting multiple samples from the area impacted by mobile LNAPL and averaging the remaining BTEX fraction.
- Based on Raoult's Law, weathering of BTEX from LNAPL via dissolution and volatilization is expected to follow first-order kinetics which predicts that the rate of BTEX removal from the free-phase will be reduced as the concentrations of BTEX in the free-phase decrease over time. While this phenomenon is difficult to prove with only one or two historical data points per site, the first-order weathering rate appears to be validated when average remaining BTEX fractions from five JP-4 sites were plotted together. Based on our data, weathering rates decreased as the age of the spill increased.
- Based on Figure 5.3, the average total BTEX, first-order weathering rate for five JP-4 sites is approximately 16 %/yr. Based on all of the data collected, this appears to be a reasonable default value for estimating total BTEX weathering from JP-4 LNAPL.
- If mathematically inflated rates from McChord AFB data, and questionably low weathering rates from Eaker AFB are excluded, the range of total BTEX, first order weather rates is 11 to 23 %/yr. If a more conservative first-order weathering rate is desired for BTEX fate and transport modeling, 11 %/yr would provide a conservative estimate for JP-4 fuels.
- As predicted by their relatively high solubilities, benzene and toluene exhibit higher weathering rates than ethylbenzene and xylenes. Because benzene is a known human carcinogen with a federal MCL of 5 µg/L, benzene weathering rates will generally determine the timeframe for fuel spill remediation. Based on Figure 5.4, the average benzene first-order weathering rate for five JP-4 sites is approximately 26 %/yr. Based on all of the data collected, this appears to be a reasonable default value for estimating benzene weathering from JP-4 LNAPL. If mathematically inflated rates from McChord AFB data, and questionably low weathering rates from Eaker AFB are excluded, the range of benzene first-order weather rates is 19 to 35 %/yr. If a more conservative first-order weathering rate is desired for benzene fate and transport modeling, 19 %/yr would provide a conservative estimate for JP-4 fuels. Benzene and total BTEX first-order weathering rates for JP-4 and JP-8 are shown in Table 6.1.

- Dissolution appears to be the primary weathering mechanism that influences mobile LNAPL weathering rates. Significantly lower BTEX weathering rates in mobile LNAPL were apparent at sites with low groundwater velocities. This observation is supported by mass transfer theory which predicts that BTEX flux from LNAPL to groundwater would increase in a rapidly moving groundwater where dissolved BTEX concentrations would be diluted by the constant influx of clean water.
- Although initial BTEX fractions in JP-8 are lower than JP-4, the first-order weathering rate for the Seymour Johnson JP-8 site was 25 %/yr for total BTEX and 29 %/yr for benzene (Table 6.1). The first-order weathering rates calculated for JP-4 should provide a reasonable estimate for JP-8.
- Determination of BTEX weathering rates for JP-5 mobile LNAPLs is difficult due to the low initial concentrations of BTEX. There is very little BTEX in JP-5 and groundwater at JP-5 release sites will not be significantly impacted by BTEX compounds.
- The large range of potential initial BTEX values for gasoline combined with sample result disparities and site-specific limitations of the Offutt AFB site, prevented meaningful determination of mobile LNAPL weathering rates for BTEX in gasoline.
- Although a consistent correlation between mobile LNAPL and residual LNAPL weathering at each site was not observed, the relative contributions of volatilization and biodegradation should increase in contaminated soils above the mobile LNAPL layer. This increase in residual weathering would be most apparent at sites without impermeable surfaces and sites with more porous soils where volatilization and atmospheric oxygen diffusion are more likely to occur. At fuel contaminated sites, several samples are needed at varying depths to accurately estimate the residual BTEX contamination in soil.

TABLE 6.1 SUMMARY OF BENZENE AND TOTAL BTEX FIRST-ORDER WEATHERING RATES IN JP-4 AND JP-8 MOBILE LNAPL FUEL WEATHERING STUDY

	Benzene	Total BTEX
	(% Reduction/Year ^{a/})	(% Reduction/Year)
JP-4 MOBILE LNAPL		
Average of 5 Sites	26	16
Range Excluding Outliers ^{b/}	19 to 35	11 to 23
Conservative Estimate	19	11
JP-8 MOBILE LNAPL ^{c/} Average	29	25
Range	6 to 53	14 to 36

a/Weight percent reduction per year calculated using equation 5.8.

^{b/} Excludes data from the McChord AFB and Eaker AFB sites.

^{c/} Summary of eight samples collected over 2 years from the Seymour Johnson AFB site.

SECTION 7

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APPENDIX A

WORK PLAN AND SITE ADDENDA FOR THE FUEL WEATHERING STUDY

A-1

WORK PLAN

DRAFT

Work Plan for Determining LNAPL Weathering at Various Fuel Release Sites



Prepared For

Air Force Center for Environmental Excellence Technology Transfer Division Brooks Air Force Base San Antonio, Texas

ctober 199



1700 Broadway, Suite 900 • Denver, Colorado 80290

WORK PLAN FOR DETERMINING LNAPL WEATHERING AT VARIOUS FUEL RELEASE SITES

October 1996

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ACRONYMS AND ABBREVIATIONS

ABB	ABB Environmental Services, Inc.
AFB	Air Force Base
AFCEE	Air Force Center for Environmental Excellence
AST	above ground storage tank
bgs	below ground surface
BTEX	benzene, toluene, ethylbenzene, xylenes
°C	degrees centigrade
CA	contamination assessment
CAP	corrective action plan
CAR	contamination assessment report
DFSC	Defense Fuel Supply Center
DFSP	Defense Fuel Supply Point
DOD	Department of Defense
ECT	Environmental Consulting and Technology, Inc.
ES	Engineering-Science, Inc.
FDEP	Florida Department of Environmental Protection
HDPE	high density polyethylene
ID	inside diameter
IDW	investigation-derived waste
IT	IT Corporation
IWTP	industrial wastewater treatment plant
LCS	laboratory control sample
LMB	laboratory method blank
LNAPL	light non-aqueous phase liquid
mg/L	milligrams per liter
µg/kg	micrograms per kilogram
µg/L	micrograms per liter
MW	monitoring well
NAPL	nonaqueous-phase liquid
NAS	Naval Air Station
NPDES	National Pollution Discharge Elimination System
NRMRL	National Risk Management Research Laboratory
ORD	Office of Research and Development
OSWER	Office of Solid Waste and Emergency Response
OD	outside diameter
OVM	organic vapor meter
Parsons ES	Parsons Engineering Science, Inc.
PID	photoionization detector
ppmv	parts per million volume per volume

PVC	polyvinyl chloride
QA/QC	quality assurance/quality control
RAP	remedial action plan
RNA	remediation by natural attenuation
Rust	Rust Environment & Infrastructure
RW	recovery well
SAR	site assessment report
SCDHEC	South Carolina Department of Health and Environmental Control
SVE	soil vapor extraction
TPH	total petroleum hydrocarbons
USAEHA	United States Army Environmental Hygiene Agency
USEPA	United States Environmental Protection Agency
USGS	United States Geological Survey
UST	underground storage tank
UV	ultraviolet
VOC	volatile organic compound
WEGS	Westinghouse Environmental & Geotechnical Services

SECTION 1

INTRODUCTION

This work plan, prepared by Parsons Engineering Science, Inc. (Parsons ES), presents the scope of work required for the collection of data necessary to evaluate natural weathering rates of mobile and residual light nonaqueous-phase liquids (LNAPLs) resulting from petroleum releases to the subsurface environment. As part of the natural attenuation demonstration project (Contract No. F41624-92-D-8036, Delivery Order 25), the Air Force Center for Environmental Excellence (AFCEE) has contracted with Parsons ES to perform this fuel weathering study. Of particular interest for the study is the weathering of fuels such as JP-4 jet fuel, JP-8 jet fuel, and gasoline each of which contain relatively high mass fractions of benzene, toluene, ethylbenzene, and total xylenes (BTEX).

1.1 PROJECT SCOPE AND OBJECTIVES

At many government and general industry sites, large-volume environmental releases of jet fuel or gasoline have contaminated and continue to contaminate soil and groundwater systems. Primary sources of large-volume fuel releases include fuel handling and storage activities associated with aboveground storage tanks (ASTs), underground storage tanks (USTs), fuel pumphouses, fuel hydrant systems, oil/water separators, and fuel pipelines, to name a few. Uncontrolled catastrophic or chronic releases from one of these primary sources can result in large volumes of fuel being released to the subsurface. When released, fuels such as JP-4 and gasoline represent oily-phase liquids, which are less dense than water. In the subsurface, the LNAPL is often present as both a mobile and a residual contaminant source. Residual LNAPL is defined as the LNAPL that is trapped in the aquifer or from the aquifer matrix into a well under the influence of gravity. Mobile LNAPL is defined as LNAPL that is free to flow in the aquifer and will flow from the aquifer matrix into a well under the influence of gravity.

Little information is currently available regarding rates of natural weathering or attenuation of mobile and residual LNAPLs. As a result, conservative BTEX reduction rates for LNAPLs typically are used when predicting the persistence and concentration of fuel hydrocarbon sources. The use of overly conservative LNAPL weathering rates at sites selected for remediation by natural attenuation (RNA) can extend the estimated timeframe for long-term monitoring and affect the estimated cost effectiveness and administrative feasibility of implementing RNA. The purpose of this study is to improve the scientific basis and defensibility for determining LNAPL weathering rates, (i.e., source reduction rates) as a component of the RNA alternative.

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As used in this report, RNA refers to a management strategy that relies on natural attenuation mechanisms to remediate soil and groundwater contaminants and to reduce and control risks associated with contaminants in the subsurface. The United States Environmental Protection Agency (USEPA) Offices of Research and Development (ORD) and Solid Waste and Emergency Response (OSWER) define natural attenuation as:

The biodegradation, dispersion, sorption, volatilization, and/or chemical and biochemical stabilization of contaminants to effectively reduce contaminant toxicity, mobility, or volume to levels that are protective of human health and the ecosystem.

The primary objective of this study is to determine an average range of natural weathering rates for the mobile LNAPL fraction based on literature values and data collected from sites with documented mobile LNAPL plumes that have resulted from past jet fuel or gasoline releases. Three secondary objectives are 1) to review the available literature as it pertains to natural weathering of fuel LNAPLs in the subsurface environment; 2) to compare weathering effects on the mobile LNAPL fraction and on residual LNAPL present in capillary fringe soils; and 3) to evaluate the degree of contaminant partitioning occurring from mobile LNAPL to the groundwater. Accomplishment of these objectives will involve conducting the following tasks:

- A literature search to assess existing information regarding weathering of LNAPLs;
- Selection of 10 sites where releases of gasoline or jet fuel have occurred and freephase LNAPLs remain *in situ*;
- Sampling of soil, groundwater, and free-phase LNAPLs at the selected sites; and
- Various trend analyses of current contaminant concentrations in site media and their relation to age of the fuel release, fuel type, and site geology and hydrogeology.

The field work for this project primarily will involve collection of soil, groundwater, and free product samples from the selected sites. It is anticipated that two soil samples from the capillary fringe, four free product samples, and two groundwater samples will be collected from each site and submitted for laboratory analysis of BTEX concentrations in each media type. Soil samples also will be analyzed for total petroleum hydrocarbons (TPH) and soil moisture in order to compare weathering effects in soils and in free-phase LNAPLs. It is anticipated that the majority of sites will be sampled using a Geoprobe[®], which is "a hydraulically-powered, percussion/probing machine" specifically designed for environmental sampling of shallow unconsolidated media and groundwater. It is anticipated that field work at each of the selected sites will last approximately 0.5 to 2 days. All field work will follow the health and safety procedures presented in the program *Health and Safety Plan for Bioplume II Modeling Initiative* (Engineering-Science, Inc. [ES], 1993), and site-specific addenda developed for each site.

Following field sampling and receipt of sample results, data from each of the sites will be analyzed for the target parameters described in Section 3, and a technical report will be prepared by Parsons ES and submitted to AFCEE. The technical report will summarize the findings of the literature review and will provide an assessment of site-specific LNAPL weathering rates considering the age of the LNAPL and the original fuel type. In addition, the report will examine variations in LNAPL weathering effects in soil and groundwater within the contaminant source areas.

1.2 WORK PLAN ORGANIZATION

This work plan consists of six sections, including this introduction, and one appendix. Section 2 presents site selection criteria, a listing of candidate sites selected for the study, background information for each of the sites, and proposed site sampling locations. Section 3 describes procedures for collection and analysis of site data. Section 4 discusses the quality assurance/quality control (QA/QC) measures to be used during this project. Section 5 describes the data analyses to be performed for determining LNAPL weathering and its effects on soil and groundwater. Section 6 contains the references used in preparing this document. Appendix A provides a draft listing of references to be reviewed during the literature search.

1.3 PRELIMINARY LITERATURE REVIEW

The purpose of the proposed literature review is to compile and summarize the technical literature on natural weathering of fuel LNAPLs in the subsurface environment. Specifically, the literature search will attempt to answer the following question:

Is there sufficient scientific information available regarding *in situ* LNAPL weathering rates to refine modeling assumptions commonly used to predict reductions in the contaminant source term in RNA demonstrations at fuel-hydrocarbon-contaminated sites?

While a complete summary of the available literature will be provided with the final technical report submitted to AFCEE upon project completion, a preliminary review of the literature suggests that little information currently exists regarding LNAPL weathering rates in the subsurface or fuels weathering in general. The majority of information currently available regarding subsurface fuel contamination examines the effects of specific natural attenuation mechanisms such as dissolution, biodegradation, and volatilization as they apply to soil and groundwater contamination. While the literature has focused on these mechanisms as they apply to attenuation of chemicals sorbed to soil and dissolved in groundwater, mobile LNAPL weathering also is a function of these processes. Based on this preliminary review of the literature, the proposed quantitative assessment of LNAPL weathering will provide valuable data on the behavior of fuels over time at release sites.

A brief summary of the literature as it pertains to these mechanisms is provided below. A brief description of methods to assess fuel weathering, and oil mass loss rates from a spill in Bemidji, Minnesota are also presented.

1.3.1 LNAPL and Weathering

Weathering of petroleum hydrocarbons in the subsurface environment has historically been interpreted as the observed change in petroleum product composition with time due mostly to preferential volatilization of constituents with the greatest vapor pressures (Worthington and Perez, 1993). The term weathering as it is used for this study refers to the combined effects of natural destructive and non-destructive processes to reduce a fuel contaminant's persistence, mobility, mass, and toxicity in the environment. The primary mechanisms acting to reduce the strength of a mobile LNAPL source are dissolution, biodegradation, and volatilization. These mechanisms are influenced by physical and chemical properties of the chemical compounds in the source product, as well as by physical, chemical, and biological properties of the soil and groundwater system.

1.3.1.1 Dissolution

Dissolution is the dissolving of chemical substances from a nonaqueous-phase liquid (NAPL) into percolating precipitation water and/or the groundwater. Various bench-scale studies have been performed that examine the effects of dissolution on residual LNAPLs in soils (Borden and Kao, 1992; Rixey et al., 1992; Voudrias et al., 1994). In the latter two studies, water was flushed through soil columns to assess dissolution rates as they applied to aromatic compounds typical of petroleum hydrocarbon contamination (i.e., benzene, toluene, and xylenes). In addition, the Voudrias et al. (1994) study compared soil column and sand column effluent BTEX concentrations resulting from dissolution of a residual JP-4 iet fuel LNAPL. In both studies, flushing of soils with residual LNAPL resulted in either increasing or stable dissolved effluent BTEX concentrations during the initial stages of dissolution. Decreases in effluent concentration with continued flushing occurred only after the residual LNAPL became sorbed to the soil within the column. Following this transition, dissolved contaminant concentrations decreased rapidly and then leveled off. Increasingly greater pore volumes of water were required for each subsequent percent decrease in dissolved contaminants. In the Voudrias et al. (1994) study which involved dissolution of JP-4 in a soil column, benzene was reduced to 50 percent of its maximum effluent concentration after flushing with 720 pore volumes, and toluene was reduced to 50 percent of its maximum effluent concentration after 1,860 pore volumes. Ethylbenzene and xylene were not reduced to below 50 percent of their maximum effluent concentration even after flushing with 3,810 pore volumes. These studies suggest that an unrealistic number of pore volumes of groundwater must be flushed through the contaminated area to achieve complete dissolution of the aromatic compounds typical of gasoline and jet fuels.

The impacts of contaminant solubility and molecular size on dissolution rates also has been studied. Benzene and toluene, which have higher solubilities than the other two BTEX compounds, possessed the greatest dissolution rates and were removed first, followed by ethylbenzene and xylenes, in the jet fuel dissolution experiments (Voudrias *et al.*, 1994). Decreased contaminant solubility results in a decreased dissolution flux and prolongs remediation by flushing (Yang *et al.*, 1995).

Significant debate appears in the literature regarding the applicability of equilibrium conditions when assessing dissolution (Hayden *et al.*, 1992; Seagren *et al.*, 1993; Voudrias *et al.*, 1994; and Yang *et al.*, 1995). Under equilibrium conditions, it is assumed that if the concentration of the contaminant in one phase is known, the contaminant concentration in other phase(s) can be determined (Powers *et al.*, 1991). The existence of equilibrium conditions between an LNAPL and groundwater would allow for determination of LNAPL contaminant mole fractions based on concentrations of the

contaminant in groundwater. Based on this theory, contaminant depletion in LNAPL could be tracked through monitoring of groundwater contaminant depletion. However, the equilibrium assumption as it applies to LNAPLs and groundwater contaminant concentrations has yet to be adequately demonstrated (Powers *et al.*, 1991). Yang *et al.* (1995) argue that the rate of dissolution is a significant rate-limiting factor in the remediation of soils contaminated with NAPLs. Dissolution also may be rate-limiting for LNAPL weathering.

1.3.1.2 Biodegradation

Biodegradation represents another significant mechanism for petroleum contaminant weathering in the subsurface. Most of the literature pertaining to *in situ* biodegradation refers to the adsorbed, or residual, and dissolved phases. As mentioned in the preceding section, dissolution appears to be a rate-limiting factor in weathering, especially as it relates to biodegradation. Biodegradation of dissolved petroleum contaminants reduces aqueous contaminant concentrations and thereby enhances dissolution rates by increasing mass transfer of soluble compounds from mobile and residual LNAPLs into groundwater (Seagren *et al.*, 1993; Yang *et al.*, 1995). As a result of this diffusion limitation, mass loss rates of dissolved contaminants from biodegradation initially appear to be between zero and first order (Song *et al.*, 1990), and decrease with time (Barker *et al.*, 1987).

The kinetics of biodegradation are complicated by the fact that biodegradation is compound specific and is significantly affected by the geochemistry of the subsurface environment. Dean-Ross (1993) examined the fate of JP-4 jet fuel in subsurface soils and discovered that for the less volatile, higher-molecular-weight jet fuel components, biodegradation represented a significant mechanism for reducing soil contamination. Song et al. (1990) concluded that saturated compounds such as hexane are generally more easily biodegraded than the corresponding aromatic compounds. In a study by Barker et al. (1987), mass loss rates for aromatics in groundwater due to biodegradation were greatest for xylenes, followed by toluene, and benzene. Other factors playing an important role in contaminant biodegradation include availability of nutrients, availability of oxygen, and the interfacial area available for mass transfer to aqueous or gaseous phases (Yang et al., 1995). Looking at residual LNAPLs, the size of the LNAPL globules impacts biodegradation rates, with smaller globules resulting in greater interfacial area for mass transfer, and greater biodegradation rates (Yang *et al.*, 1995). Mobile LNAPLs present as free product floating on the groundwater would have a low interfacial area (lower bioavailability) in comparison to the residual LNAPL globules in the unsaturated zone. Finally, it has been noted that biodegradation rates decrease with increasing contaminant concentrations (Schwille, 1967; Stroo et al., 1992). From this information it can be inferred that mobile LNAPL may be subject to little biodegradation, as it represents the most concentrated contaminant phase.

1.3.1.3 Volatilization

Volatilization is expected to be a significant weathering mechanism for petroleum products such as gasoline, JP-4, and JP-8. From a study on the fate of JP-8 in quiescent flask systems containing water and water/sediment mixtures, evaporation or volatilization

from water was the major removal mechanism for low-molecular-weight, volatile hydrocarbons (Dean-Ross *et al.*, 1992). In the same study it was determined that the presence of sediment can sequester jet fuel and render it less susceptible to volatilization. Intuitively, greater contact between soil gas and residual LNAPL would result in greater mass loss rates due to volatilization than soils saturated with mobile LNAPL.

1.3.2 Methods for Assessing Weathering

While little information on actual LNAPL weathering rates, or source reduction rates has yet to be identified, various methods to qualitatively or quantitatively assess contaminant weathering have been proposed. Luhrs and Pyott (1992) presented a paper on the use of trilinear plots to graphically represent plume zonation, contaminant source identification, and contaminant weathering in groundwater for gasoline releases. This methodology relies upon plotting relative ratios of source contaminants, such as the BTEX compounds, on a trilinear plot. Due to differences in vapor pressures and susceptibility to biodegradation, BTEX contaminants in groundwater will attenuate at different rates. For example, the ratio of benzene to total xylenes in groundwater is higher for a recent gasoline release than for an older release that has weathered. While this method appears to give more of a qualitative assessment of the degree of contaminant weathering in groundwater, trilinear plots can facilitate the interpretation of data that have been collected over time.

Petroleum chemical indicators, or biomarkers, recently have been evaluated as a method to assess the degree of total oil depletion occurring as a result of crude oil weathering (Douglas *et al.*, 1994). Based on the methodology presented, if concentration data from the source oil is available on a mass fraction basis, the increase in the mass fraction of a conservative chemical indicator in a weathered product relative to its initial mass fraction in the source oil is proportional to the amount of oil lost via weathering. For crude oil, Douglas *et al.* consider hopane as the conservative internal indicator, whereas for mid-range refined petroleum products such as diesel fuel and fuel oil #2, phenanthrenes/anthracenes can be used to assess total oil losses due to weathering. Internal biomarkers for lighter aromatic fuels such as gasoline and JP-4 jet fuel were not suggested. Trimethylbenzene and tetramethylbenzene isomers have been suggested as reliable conservative tracers in anaerobic soil and groundwater systems (Wiedemeier, 1995).

1.3.3 Example Crude Oil Weathering Rate at Bemidji Spill Site

In 1979, a crude oil pipeline near Bemidji, Minnesota burst and released approximately 450,000 gallons of crude oil into a glacial outwash aquifer. In 1982, the site was selected for a long-term interdisciplinary study by the US Geological Survey. Oil-mass loss rates at the site have been studied to assess the degree of weathering resulting primarily from volatilization and dissolution. Based on one of these studies, annual oil-mass loss rates determined at different site locations ranged from 0 to 1.25 percent (Landon and Hult, 1991). Total cumulative oil losses after approximately 10 years of weathering were reported to be as much as 11 percent. Weathering rates for lighter refined petroleum

products are expected to be greater than those calculated for the Bemidji site due to increased volatility, solubility, and biodegradability of the source contaminants.
SECTION 2

SELECTION OF STUDY SITES

The primary objective of this study is to determine an average range of natural *in situ* weathering rates for LNAPL associated with JP-4 or JP-8 jet fuel or gasoline spills based on existing literature and data collected from sites with mobile LNAPL contamination. Based on the apparent lack of existing LNAPL weathering rate literature, candidate study site identification, selection, and field sampling is necessary to gather data to evaluate weathering rates. The site selection criteria for this fuel weathering study are presented in Section 2.1. Primary and secondary candidate sites identified to date are reviewed in Sections 2.2 and 2.3, respectively. If additional primary sites are identified during the course of this study, they will be considered for sampling with the concurrence of AFCEE/ERT.

2.1 SITE SELECTION CRITERIA

To evaluate a site's potential as a candidate for this fuel weathering study, the following selection criteria were developed by Parsons ES:

- 1. Presence of recoverable mobile LNAPL resulting from a JP-4, JP-8, or gasoline release;
- 2. Known date of fuel release;
- 3. LNAPL resulting from of a single release confined to a relatively short period of time;
- 4. No, or minimal, site remediation undertaken to date;
- 5. Historic LNAPL analytical results, including BTEX;
- 6. Depth to groundwater less than 40 feet below ground surface (bgs); and
- 7. Department of Defense (DOD), and federal government sites preferred to general industry sites.

Identifying sites that meet all of the above-listed criteria has proven to be a difficult (ongoing) task. Consequently, the criteria served as guidelines for site selection rather than rigid selection parameters. JP-4, JP-8, or gasoline fuel release sites are preferred because of the relatively high mass fraction of BTEX present in these source fuels. Source reduction, (i.e., BTEX depletion) estimates using sampling data for the fresh product and the remaining mobile LNAPL should be more accurate for fuels with higher initial BTEX concentrations. The mobile LNAPL criterion is considered to be met by sites that have

sufficient free product thicknesses to allow collection of relatively undiluted product samples (i.e., at least 1 inch of mobile LNAPL).

Application of the second and third criteria in combination with the first criterion has eliminated numerous sites from this study. In order to determine the degree of weathering that has occurred over a given time interval, reliable information on the date of the release must be available. For many petroleum release sites, the specific date(s) of release is not documented and at best can be only approximated based on known historical site activities. In addition, one-time releases (i.e. spills) of sufficient volume to produce a long-term mobile LNAPL in the subsurface environment are rare and when such releases occur, they frequently trigger emergency response actions that compromise satisfaction of the fourth selection criterion (minimal site remediation).

Sites where limited or no site remediation has occurred are preferred for this assessment of *in situ* LNAPL weathering rates. Soil venting activities, such as soil vapor extraction (SVE), bioventing, and bioslurping are likely to increase the attenuation of the LNAPL fraction as a result of volatilization and biodegradation; therefore, BTEX weathering calculations performed for the LNAPL remaining at these sites would be biased. Sites where limited free product recovery or soil excavation has occurred, but at which measurable free product remains, are considered acceptable for this study.

Sites with historic LNAPL sampling results for BTEX may be considered in lieu of a known spill or release date. The availability of known BTEX concentrations at a known sampling time could serve as an initial point on the BTEX depletion curve. Ideally, historic LNAPL BTEX results should predate the proposed product sampling by 3 or more years to minimize the impact of sample variability.

Sites where the depth to groundwater is less than 40 feet bgs are preferred so that Geoprobe[®] sampling can be performed. Geoprobe[®] groundwater sample collection below this depth often is difficult, but will be evaluated on a site-specific basis.

Finally, DOD and federal government sites are preferred. It is expected that potential legal issues and project funding issues will be minimized if the majority of sites included in this research study are located at federal facilities. In an effort to satisfy this requirement, if an insufficient number of JP-4, JP-8, and/or gasoline release sites are identified, the first criterion may be relaxed to include JP-5 (a fuel commonly used at naval facilities) release sites, especially if other site selection criteria are met by the candidate JP-5 site. Sections 2.2 and 2.3, respectively, list the primary and secondary candidate sites identified to date. Each listing is subject to change as information becomes available during ongoing site identification efforts. All the sites listed and discussed in Section 2.2 are located at federal facilities. The sites listed and briefly described in Section 2.3 are secondary sites that satisfy fewer of the selection criteria. These sites may be selected for the study if additional primary sites are not identified.

2.2 PRIMARY SITES

The primary sites listed and described below represent those currently identified sites that best meet the selection criteria, outlined in Section 2.1 for the proposed fuel weathering study.

2.2.1 Primary Site Listing

The primary site list for this fuel weathering study currently includes the following fuel release sites:

- Building 1610, Shaw Air Force Base (AFB), South Carolina;
- Defense Fuel Supply Point-Charleston, Hanahan, South Carolina;
- Day Tank 1, Facility 293, Cecil Field Naval Air Station, Florida; and
- General Industry Site, Myrtle Beach AFB, South Carolina.

Brief descriptions of site operation histories, physical setting, and contaminant conditions are provided in the following subsections.

2.2.2 Building 1610, Shaw AFB, South Carolina

Shaw AFB is located approximately 37 miles east of Columbia, South Carolina, along US Highway 76. Building 1610 is located in the northern portion of Shaw AFB, adjacent to the flightline. The site at Building 1610 consists of a small release area adjacent to the main jet fuel pipeline that services the flightline. In June 1994, jet fuel was discovered on the ground surface as a result of a leak in a buried, pressurized, 6-inch-diameter fuel pipeline. Based on information from facility personnel, the leak is thought to have occurred over a 5- to 6-month-period, during which the Base converted from JP-4 to JP-8 jet fuel (Roller, 1996). The released fuel is thought to be composed primarily of JP-4 jet fuel, because the conversion to JP-8 occurred in April/May 1994 (Green, 1996). A site layout for the Building 1610 release is provided on Figure 2.1.

In August 1996, the first three groundwater monitoring wells (MWs) were installed in the vicinity of Building 1610; therefore, sampling data from multiple events are not available. A groundwater investigation recently was conducted at nearby Building 1613, located approximately 1,500 feet south-southwest of the Building 1610 site (Rust Environment & Infrastructure, Inc. [Rust], 1995a). The geology at the two sites is thought to be similar. At the Building 1613 site, sandy deposits of the Duplin Formation occur from the ground surface to a depth of approximately 65 feet bgs. Underlying the Duplin Formation are clayey deposits. Groundwater at the Building 1613 site occurs at approximately 30 feet bgs and flows to the east.

Following discovery of the leak, approximately 80 tons of soil was excavated and removed from the site. Implementation of site assessment activities were begun by Rust in August 1996, and included the installation of one upgradient MW (MW1610-1) and two additional MWs (MW1610-2 and MW1610-3) downgradient from the area of the pipeline leak. MWs locations are shown on Figure 2.1. Upon completion and



development of the monitoring wells, free product thicknesses of approximately 2.5 feet were measured in MW1610-2 and MW1610-3 (Green, 1996). In addition, approximately 1.9 feet of free product was measured in MW1610-1. The draft site assessment report, which will include soil and groundwater sampling results is scheduled to be released in October 1996 (Rust, 1995b).

Proposed sampling locations for the Building 1610 site are shown on Figure 2.1. Sampling procedures are outlined in Section 3. South Carolina Department of Health and Environmental Control (SCDHEC) approval is required prior to Geoprobe® borehole installations that penetrate the water table.

2.2.3 Defense Fuel Supply Point - Charleston, Hanahan, South Carolina

Defense Fuel Supply Point (DFSP) - Charleston, is located in Hanahan, South Carolina near Charleston. DFSP consists of seven ASTs used to store aviation fuels. In October 1975, a leak developed in Tank 1 that resulted in the release of 83,000 gallons of JP-4 jet fuel over a 20-day period. The release triggered abatement actions and a series of environmental investigations at the site. A layout of the site is provided on Figure 2.2.

The site is underlain by unconsolidated Pleistocene sediments composed primarily of medium-grained sands with interfingering clay lenses (Chapelle *et al.*, 1996). In the Tank 1 area, the most permeable saturated sands are overlain by discontinuous 1- to 3-foot-thick clay beds. These clay beds create local semi-confined conditions for the moderately permeable sandy shallow aquifer. Groundwater flows north from the Tank 1 area, remaining under semi-confined conditions for a distance of approximately 150 feet (Chapelle *et al.*, 1996). The depth to the water table varies with the season, but was recently measured at approximately 18 to 22 feet bgs in the Tank 1 area (International Technology Corporation [IT], 1996).

The majority of information in this section was extracted from facsimile transmittals of a recent work plan assembled by IT (1996). Since the time of the release, numerous site investigations have occurred. In November 1975, the US Army Environmental Hygiene Agency (USAEHA, 1977) determined the plume area encompassed approximately 20,000 square feet in the northern portion of the storage basin at approximately 7 to 12 feet bgs. In December 1975, free product recovery was conducted at the site for 2 weeks, and approximately 21,000 gallons of JP-4 was recovered. A second attempt in 1976 to further recover free product was terminated because little jet fuel was recovered (USAEHA, 1977).

In 1987, the US Geological Survey (USGS), in cooperation with the Defense Fuel Supply Center (DFSC), began an investigation at the site to assess the potential for *ex* situ bioremediation of groundwater contamination at the site. As part of this investigation, 17 groundwater extraction wells were installed. Based on data collected from these wells, the hydrogeology of the site was determined to be more complex than previously recognized. Measurements from these wells showed free-phase jet fuel to be trapped below clay lenses beneath the water table. Jet fuel migration had occurred during periods of low water table levels caused by natural climatic conditions and exacerbated by pumping from the recovery wells (IT, 1996).



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At the time of development of this work plan, soil, groundwater, and LNAPL contamination data were not yet available from DFSC. As part of a recently published paper regarding biodegradation rates at the site, groundwater contamination levels were provided for benzene and toluene averaged over three separate quarterly sampling events (Chapelle *et al.*, 1996). Average reported benzene concentrations in groundwater were measured at 19 micrograms per liter (μ g/L) at extraction well EW-7 and 74.7 μ g/L at EW-6. Toluene concentrations were measured in groundwater at 345 μ g/L and 223 μ g/L at EW-7 and EW-6, respectively.

Based on information provided by DFSC personnel (Smith, 1996), mobile LNAPL is typically present at EW-6 and EW-7 and MW W-103 (EW-6 and EW-7 are shown on Figure 2.2. W-103, not shown, is approximately 10 feet west of EW-7). However, based on monthly water level and product thickness measurements performed at the site by the IT Corporation, free product has not been observed in these wells since May 1996 (IT, 1996). During the May 1996 measurement, 0.22 foot and 1.77 feet of product were observed in EW-6 and EW-7, respectively. In August 1996, approximately 0.35 foot of product was present in a 36-inch recovery well located northwest of Tank 1 (Figure 2.2).

Proposed sampling locations for this fuel weathering study are shown on Figure 2.2. In addition to soil, groundwater, and free product sampling at the Tank 1 area, proposed sampling locations also are shown at the Tank 3 area, located approximately 700 feet east of Tank 1. Free product at the Tank 3 area has been noted as being substantially weathered as compared to fuel present at the Tank 1 site (Vroblesky, 1996). While the source of the free product at the Tank 3 area is not known, it is believed to be approximately the same age as the product at the Tank 1 site (Vroblesky, 1996). Collection of samples at each of these locations may prove advantageous because the difference in weathering rates may be a function of where the LNAPL is present within the aquifer rather than spillage. Under Tank 1, the LNAPL is trapped beneath a confining clay unit whereas beneath Tank 3 LNAPL is located at the water table. As with the Shaw AFB site, SCDHEC approval is required prior to intrusive investigations that result in penetration of the water table.

2.2.4 Day Tank 1, Facility 293, Cecil Field Naval Air Station, Florida

The Cecil Field Naval Air Station (NAS) is located in Jacksonville, Florida in southwestern Duval County at the junction of Highway 228 (Normandy Boulevard) and 103rd Street. Day Tank 1, Facility 293, located east of Jet Road and north of Buildings 824 and 824A, consists of a 200,000-gallon, interior-lined, asphalt-coated, steel AST installed in 1956. In 1981, approximately 497,000 gallons of JP-5 jet fuel was released at this site when the tank was overfilled. Facility personnel reported at the time that more than one-half of the released JP-5 was recovered following the spill (ABB Environmental Services [ABB], 1994). Free product present at this site also may be associated with other possible releases resulting from damage to the underground fuel distribution pipeline and junction box adjacent to the tank, and structural damage to the tank itself (ABB, 1994). A layout of the site is provided on Figure 2.3.

The specific geology and hydrogeology for the Day Tank 1 site was not available at the time of work plan development. Unless otherwise indicated, geologic information provided below is from the North Fuel Farm Contamination Assessment Report



Addendum (ABB, 1996). The North Fuel Farm is located approximately 4,000 feet north of the Day Tank 1 site. Sediments at the North Fuel Farm are composed of silty, finegrained sand and silty sand with trace amounts of clay in the shallow water table zone. The shallow water table at the North Fuel Farm ranges between 2 and 15 feet bgs. Sediments in the surficial aquifer zone, which extends from 15 to 50 feet bgs, are composed of fine-grained silty sand. Based on summarized results reported in the remedial action plan (RAP) for the Day Tank 1 site (ABB, 1994), depth to water in the surficial aquifer at the site ranges from 5 feet to 8 feet bgs and the overall groundwater flow direction is toward the east-southeast.

Unless otherwise stated the following information was taken from the RAP for Day Tank 1 prepared by ABB (1994). In 1981, Geraghty & Miller conducted a preliminary contamination assessment (CA) at the Day Tank 1 facility and concluded that JP-5 fuel was present only in the unsaturated zone and had not migrated into groundwater at the site. The CA report concluded that the fuel would naturally biodegrade over time; therefore, the only remedial action taken was the addition of fertilizer to site soils to enhance fuel biodegradation.

In December 1990, a second CA was initiated at the site by ABB and involved the installation of 15 shallow MWs (CEF-293-01 through CEF-293-14 and CEF-293-16) and one deep MW (CEF-293-15D). After completion of the CA, the Florida Department of Environmental Protection (FDEP) requested additional subsurface assessment activities, which resulted in additional soil sampling and installation of two additional monitoring wells (CEF-293-17D and CEF-293-18) in October 1993. Soil samples collected from eight soil borings were found to have elevated headspace readings of volatile organic compounds (VOCs) (greater than 50 parts per million volume per volume [ppmv] using an organic vapor meter [OVM]). Contamination distribution maps developed from September 1993 groundwater sampling results show benzene, total aromatic VOCs, and napthalene concentrations exceeding 20 milligrams per liter (mg/L), 50 mg/L, and 50 mg/L, respectively, in the area encompassed by monitoring wells CEF-293-2, CEF-293-7, and CEF-293-9 (Figure 2.3). The apparent extent of free product as of July 1995 is shown on Figure 2.3. Free product thicknesses, as measured on August 12, 1996, were 0.59 foot and 0.78 foot at monitoring wells CEF-293-7 and CEF-293-9 (Klimas, 1996).

In August 1995, ABB (1995) submitted an alternate procedures request for the Day Tank 1 site proposing bioslurping as the preferred method of free product recovery. At present, three 1-day bioslurping pilot tests have been performed at the site (Ullo, 1996).

Proposed sampling locations for the Day Tank 1 site are shown on Figure 2.3. Sampling procedures are outlined in Section 3. Because the contamination is located in an active part of the NAS, general Navy regulations must be observed in obtaining security clearance for Geoprobe® operators and sampling personnel. Any FDEP approvals required will be obtained prior to sampling activities.

2.2.5 General Industrial Site, Former Myrtle Beach AFB, South Carolina

Former Myrtle Beach AFB, South Carolina is located on the southwest side of the City of Myrtle Beach between the Intracoastal Waterway and the Atlantic Ocean. The spill site is located on the south side of former Myrtle Beach AFB, near the intersection of Avenue D and Fourth Street. On January 15, 1981, a spill of approximately 123,000 gallons of

JP-4 jet fuel occurred during fuel transfer from a barge on the Intracoastal Waterway to a privately-owned AST located on Myrtle Beach AFB property. Shortly after the spill, a french drain system was installed and approximately 21,000 gallons of the spilled JP-4 was recovered. The drain system was deactivated in July 1982 (Environmental Consulting and Technology, Inc. [ECT], 1996). A layout of the site is provided on Figure 2.4

A fairly uniform clay layer exists at the site from the ground surface to approximately 7 to 9 feet bgs. This clay layer has a low permeability and acts as a confining layer for the underlying shallow aquifer (ECT, 1996). The clay layer is underlain by a highly permeable sand with shell fragments to approximately 30 feet bgs. Clayey sand with lenses of fine sand and clay occur from 30 to 40 feet bgs. This deeper clay sand is similar to the surface clay in that it limits vertical migration. Potentiometric surfaces for the shallow confined aquifer were measured by ECT in 1992 and 1993. Potentiometric surfaces measured as high as 5 feet above the base of the clay and as low as 0.5 foot above the base of the clay depending upon the amount of seasonal precipitation (ECT, 1996). Potentiometric surface measurements conducted in November 1995 indicated a northeasterly groundwater flow direction (ECT, 1996).

Information presented below was taken from the ECT (1996) Draft Groundwater Mixing Zone and Monitoring Plan Application for the site. In July 1982, a product recovery system consisting of 10 vacuum extraction wells was installed at the site and operated until November 1982. During this time, the recovery system collected less than 100 gallons of free product. Water level and free product measurements conducted by Westinghouse Environmental & Geotechnical Services (WEGS) between 1982 and 1985 indicated potential migration of free product to the east of the original spill area. A groundwater treatment system and associated interceptor trenches were installed during March and April 1990, and operation of the system began in September 1990. Treatment system effluent failed to meet National Pollution Discharge Elimination System (NPDES) permit requirements, and the system was deactivated in November 1990.

In August 1991, ECT was contracted to conduct a site assessment for the facility. As part of the site assessment, soil samples were collected and field-screened for organic vapors, 19 monitoring wells were installed (10 shallow, 6 intermediate, and 3 deep), free product thicknesses were measured, and groundwater samples were collected for laboratory analyses. The greatest free product thicknesses (approximately 4 feet) were measured below the shallow clay layer between MWs B-7 and B-30 (Figure 2.4). No free product was measured in any of the MWs screened entirely within the surficial clay layer (MW-6C, MW-7C, and MW-8C).

Following completion of the site assessment, ECT began installation of a free product recovery system in March 1992. Three recovery trenches were excavated below the base of the shallow clay layer, and collection sumps, skimmers, and axial pumps were installed. System operation began in May 1992 and continued intermittently until November 1993. Total fuel recovered during this time was 720 gallons. Following the 1992 recovery effort, ECT estimated that approximately 12,000 to 13,000 gallons of free product remained trapped below the clay layer. Free product bail down tests conducted in 1992 and 1993 indicated free product recovery rates ranging between 0.003 and 0.015



gallon per day, several orders of magnitude slower than typical groundwater recovery rates. During 1992 and 1993 field tests, a visual comparison of recovered fuel from the site with "fresh" JP-4 suggested little weathering of the product trapped below the clay.

In March 1994, ECT submitted a corrective action plan for the spill site that proposed groundwater recovery and aboveground treatment using air stripping and carbon polishing as the preferred remedial alternative. In September 1995, SCDHEC, recommended that groundwater MWs be resampled to confirm that the free product and dissolved plumes were fully defined. ECT returned to the site in November 1995, and resampled MWs. During this investigation, free product was observed in the same monitoring wells as the March 1992 investigation, except that free product was not observed in monitoring well B-3A. The extent of free product and measured thicknesses from the November 1995 investigation are presented on Figure 2.4. Overall, concentrations of JP-4 constituents detected in groundwater samples collected during this latest sampling event were significantly lower than concentrations detected during the October 1991 sampling event (ECT, 1996). Based on the results of the November 1995 sampling event and the limited success of free product recovery in the past, ECT felt the site met groundwater mixing zone criteria under South Carolina water classifications and standards (Regulations 61-68) and applied for monitoring only as a site remedy.

Proposed sampling locations for the spill site are shown on Figure 2.4. Sampling and analysis procedures are outlined in Section 3. SCDHEC approval is required prior to conducting intrusive sampling activities.

2.3 SECONDARY SITES

The sites listed and briefly described below are secondary candidates that may be selected for the fuel weathering study if additional primary sites are not identified. The secondary sites meet fewer site selection criteria than the primary sites previously discussed.

2.3.1 Secondary Site Listing

The secondary candidate site list for this fuel weathering study currently includes the following release sites:

- General Industrial Site, Dunwoody, Georgia
- General Industrial Site, Charlotte, North Carolina
- General Industrial Site, Port Clinton, Ohio

2.3.2 General Industrial Site, Dunwoody, Georgia

This industrial site consists of a release from a 36-inch-diameter underground petroleum pipeline located within a subdivision in Dunwoody, Georgia (See Figure 2.5). In February 1971, a hairline crack was discovered in the pipeline and repaired.



The duration of the product leak prior to pipeline repair is not known, but it appears that product leaked at a very low rate for an extended period of time (Parsons ES, 1995b). The released product was composed primarily of gasoline; however, some fuel oil also may have been released prior to discovery of the leak (Gillis, 1996). Radial product collection ditches and an interceptor trench were excavated in 1972 and 1973, and 668 barrels of product was recovered by 1975 (Parsons ES, 1995b). Groundwater at the site occurs between 10 and 40 feet bgs. The extent of LNAPL at the site is shown on Figure 2.5, as are proposed sampling locations should this site be included in the study.

2.3.3 General Industrial Site, Charlotte, North Carolina

The industrial site located in Charlotte, North Carolina is owned by a commercial petroleum company. The site layout and the original leak location are shown on Figure 2.6. A leaded gasoline release of an unspecified volume is thought to have occurred between 1981 and 1989, when a faulty weld in a vapor recovery unit at the site was discovered. Company officials believe the weld may have begun leaking soon after vapor recovery unit installation in 1981 (Eaton, 1996).

Groundwater at the site occurs at approximately 26 feet bgs. The site geology is composed of a few inches of organics and silty clays at the surface, underlain by a few feet of residual soil composed of clays and silts. The residual soil grades into a saprolitic soil consisting of clay, silt, and sand grains that retain the structure of the original unweathered bedrock (Klem, 1996).

Free product recovery and total fluids recovery began at the site in early 1994, as did SVE. Air sparging points placed between monitoring wells MW-17 and MW-18 (Figure 2.6) began operation in August 1996. Officials for the petroleum company and their environmental consultant, S&ME (1994), believe groundwater and soils in the vicinity of MW-21 have not been significantly impacted by SVE operation at the site because of the relatively low-permeability saprolite in the vadose zone. As of August 1996, approximately 1.5 feet of free product was observed in monitoring well MW-21, the proposed product sampling location for this site (Figure 2.6).

2.3.4 General Industrial Site, Port Clinton, Ohio

The industrial site at Port Clinton, Ohio is the site of a former service station at which approximately 1,000 gallons of gasoline was reported to have been released in 1978. A layout of the site is provided on Figure 2.7.

The site is located in the glaciated Lake Erie plain of north central Ohio. The regional geology consists of discontinuous lake deposits of silts and fine sands underlain by limestone bedrock (Wieckowski, 1996). Well MW-1 was installed on service station property in January 1991, and free product was discovered floating on the water table 2 weeks after installation of the well. No active remediation has been performed at the site. Free product samples were collected from monitoring well MW-1 and recovery well RW-1 in December 1993 and June 1995, respectively, to characterize the free product. BTEX mass fractions were determined during both sampling events. Recent groundwater measurements collected at the site (May 1996) indicated no floating free product in MW-1; however, free product sample collection is possible from recovery well RW-1, the proposed sampling location (Figure 2.7).





2.4 ADDITIONAL SITES

The fuel weathering study also will incorporate the results of soil, groundwater, and free product samples previously collected from recent JP-8 spill sites at Seymour Johnson AFB, North Carolina, and Pope AFB, North Carolina. Both of these sites, were sampled by the US Army Corps of Engineers in July 1996. In addition, the study will incorporate the results of groundwater and soil samples collected at the KC-135 Crash Site at Wurtsmith AFB, Michigan. Mobile free product was not found during the Summer 1996 sampling event at Wurtsmith AFB.

SECTION 3

COLLECTION OF SITE DATA

To assess the effects of mobile and residual LNAPL weathering as they apply to soil and groundwater, samples of each medium (i.e., soil, groundwater, and free product) will be collected from each of the selected study sites outlined in Section 2. Wherever possible, soil, groundwater, and free product samples will be collected from the same borehole using the Geoprobe® system (in accordance with the general procedures outlined in Section 3.2). Sampling from the same borehole is desired in order to determine weathering effects on mobile LNAPL and its relation to contaminants in soil at the capillary fringe and in groundwater, within the same vertical continuum. Table 3.1 presents the analytical protocols for soil, free product, and groundwater samples.

Determination of BTEX concentrations in soil, groundwater, and free product samples represents the primary analysis to be used to determine the impacts of natural weathering processes. The concentration of naphthalene and methyl naphthalene also will be determined in each media type. In addition, TPH concentration in soil and soil moisture will be determined to allow comparison of weathering effects on the mobile LNAPL with weathering effects on residual LNAPL present in capillary fringe soils. Dissolved BTEX concentrations in groundwater will be analyzed to determine the extent of contaminant partitioning from the LNAPL to the groundwater and whether contaminant equilibrium between the LNAPL source and groundwater exists.

The following sections describe the procedures that will be followed when collecting site-specific data. Methods for collection of water level measurements prior to site sampling are described in Section 3.1. Geoprobe® procedures for collection of soil, groundwater, and free product are described in Section 3.2. Procedures to be used for the installation of temporary groundwater monitoring points, if necessary, are described in Section 3.3. Procedures to be used to collect free product and groundwater samples are described in Section 3.4. Sample handling procedures are described in Section 3.5. Site restoration procedures are described in Section 3.6, and equipment decontamination procedures are discussed in Section 3.7.

3.1 WATER LEVEL AND FREE PRODUCT THICKNESS DETERMINATION

Prior to performing sampling activities at a site, free-phase product thicknesses and static water levels will be measured at site monitoring wells in the immediate vicinity of the proposed sampling locations shown on site figures in Section 2. An oil/water

TABLE 3.1 ANALYTICAL PROTOCOLS FOR SOIL, FREE PRODUCT, AND GROUNDWATER SAMPLES **Fuel Weathering Study**

MATRIX	NUMBER OF SAMPLES PER SITE	LABORATORY	ANALYSIS	METHOD
SOIL	2	NRMRL	BTEX + TMBs ^{a'}	NRMRL equivalent to USEPA SW8020A ^{b/}
			TPH ^{e/} Napthalene and Methyl Napthalenes Moisture	NRMRL equivalent to USEPA SW8015, modified for diesel and gasoline NRMRL equivalent to USEPA SW8270 ASTM D-2216 ^{d/}
FREE PRODUCT (MOBILE LNAPL	4	2 to NRMRL and 2 to $EAL^{e/}$	<u>Analysis by NRMRL</u> BTEX + TMBs Napthalene and Methyl Napthalenes	GC/MS (Direct Injection) NRMRL equivalent to USEPA SW8270
			<u>Analysis by EAL^{f/}</u> BTEX	USEPA SW8020
GROUNDWATER	2	NRMRL	BTEX + TMBs Napthalene and Methyl Napthalenes	RSKSOP equivalent to USEPA E602 NRMRL equivalent to USEPA SW8270

^{a/} BTEX = benzene, toluene, ethylbenzene, and total xylenes; TMBs = trimethylbenzene isomers.
 ^{b/} NRMRL = USEPA National Risk Management Research Laboratory, Ada, Oklahoma; USEPA = US Environmental Protection Agency method.

^{c/} TPH = total petroleum hydrocarbons.

^{d/} ASTM = American Society for Testing and Materials method.

^{e/} EAL = Evergreen Analytical Laboratory, Wheat Ridge, Colorado.

^{t/} EAL will combine free product and distilled water in a single bottle and analyze water and free product following development of equilibrium conditions.

interface probe will be used to measure the depth to free product, the apparent free product thickness, and the depth to groundwater to the nearest 0.01 foot. Water level and free product measurements in each well will be recorded in the field notebook. Based on these measurements, soil, groundwater, and free product sampling depths will be selected.

3.2 GEOPROBE® SAMPLING FROM A SINGLE BOREHOLE

Unless otherwise indicated, soil, groundwater, and free product samples will be collected from the same borehole using a Geoprobe® system. Geoprobe® is a hydraulically powered percussion/probing machine capable of advancing sampling tools through unconsolidated soils. This system allows rapid collection of soil, soil gas, mobile LNAPL, or groundwater samples at shallow depths while minimizing the generation of investigation-derived waste (IDW). Figure 3.1 is a diagram of the Geoprobe® system. This section describes the sampling procedures to be followed when using the Geoprobe®. Sections 3.3 and 3.4 provide more detail on alternative groundwater and free product sampling procedures to be followed if site conditions preclude the use of the Geoprobe® for sampling these media.

Three general procedures will be used to collect soil, groundwater, and free product using the Geoprobe[®]. Soil samples from the capillary fringe will be collected first, as described in Section 3.2.1. A groundwater sample from below the mobile LNAPL layer will be collected next, and free product samples will be collected last. The groundwater and free product sampling methods are described in Section 3.2.2.

Base or facility personnel will be responsible for identifying the location of all utility lines, fuel lines, or any other underground infrastructure prior to any sampling activities. All necessary digging permits will be obtained through Base or facility personnel prior to mobilizing to the field. If necessary, Base or facility personnel also will be responsible for acquiring drilling and monitoring point installation permits for the proposed locations. Parsons ES will provide trained operators for the Geoprobe®.

3.2.1 Soil Sampling

The main purpose of soil sampling for this study is to evaluate soil BTEX and TPH contamination in soils directly above areas containing mobile LNAPLs. It is anticipated that residual petroleum contamination within capillary fringe soils will be further attenuated (weathered), as a result of increased volatilization and biodegradation, than will the underlying mobile LNAPL.

3.2.1.1 Soil Sampling Locations

Soil samples will be collected at two locations at each of the selected sites. The proposed sampling locations shown on the site figures in Section 2 were selected based upon the most recently available site information regarding free product location and thickness. Soil samples will be collected in locations where free product thicknesses are expected to be the greatest. To maximize the possibility of obtaining soil samples



within areas of measurable free product, boreholes will be placed as close as possible to MWs displaying maximum free product thicknesses for the site.

3.2.1.2 Soil Sampling Procedures

One soil sample will be collected from each of the two sampling locations at each site. The samples will be collected above the water table over a 2-foot sampling interval within the capillary fringe. Soil samples will be collected using a Geoprobe® 2-foot-long, large-bore soil sampler. The large-bore sampler serves as both the driving point and the sample collection device, and is attached to the leading end of the probe rods.

The sampler will be pushed or driven to a depth approximately 2 to 3 feet above the water table, the piston stop-pin will be removed to open the sampling barrel, and the sampler will be pushed into the undisturbed capillary fringe soils above the water table. The probe rods will be retracted, bringing the sampling device to the surface. The clear plastic liners inside the sampling barrel will be removed to allow lithologic logging, determination of whether residual free product is present in the soil, determination of headspace volatile organic compound (VOC) concentrations, and sample preparation for laboratory analysis.

The Parsons ES field scientist will be responsible for observing all field investigation activities, maintaining a detailed descriptive log of all subsurface materials recovered during soil coring, photographing representative samples, and properly labeling and storing samples. An example geologic boring log form is presented as Figure 3.2. The descriptive log will contain:

- Sample interval (top and bottom depth);
- Sample recovery;
- Presence or absence of a free product layer;
- Presence or absence of contamination as determined by headspace VOC analysis with an OVM or a photoionization detector (PID);
- Lithologic description, including relative density, color, major textural constituents, minor constituents, porosity, relative moisture content, plasticity of fines, cohesiveness, grain size, structure or stratification, relative permeability, and any other significant observations; and
- Depths of lithologic contacts and/or significant textural changes measured and recorded to the nearest 0.1 foot.

Following the removal of the clear plastic liner from the Geoprobe® soil sampler, an ultraviolet (UV) light will be used to determine if residual fuel contamination is present in the soil sample. Without removing the soil sample from the plastic liner, the UV light will be used to check for fluorescence that is indicative of a fuel layer in soil. The

				GEOLOGIC	BORI	١G	LOO	G			Shee	et 1 of	[:] 1
BORING NO. CLIENT: JOB NO.: LOCATION:	: <u>AFC</u> <u>729</u> <u>HILL</u>	: <u>EE</u> :080. ., AFE	3	.CONTRACTOR: .RIG TYPE: .DRLG METHOD: .BORING DIA.:			C C E T	ATE ATE LEVA EMP:	SPU[CMPI TION:	D: : :			
GEOLOGIST:				.DRLG FLUID:			V	VEATH	IER:	_			
COMENTS:										1	1	TOTAL	TDU
(ft) (ft)	Pro– file			Geologic Description			Depth (ft)	Sample Tvpe	Res	PID(ppm)	TLV(ppm)	BTEX(ppm)	(ppm)
5-													
10-													
15													
20-													
70													
<u> </u>													
									FIG	IIRF	33		
	-								110	ONL	0.0		
	<u>2</u> 014 C	`roup	d Surface	D - DRIVE			GE	പറ	CIC	BO			2
GS - Gra	ow d	Surfa		C - CORE					910	50			A
TOC - Top	o of	Casir	ig	G – GRAB									
NS – Not	t Sar	nplec	I	▼ Water level d	rilled								
SAA — Sar	ne A	s Ab	ove	<u></u>			н	ill Air	Fo	rce E	Base,	Utah	
						PA	RSO	NS					
					_	EN	GIN	EEA		; 5C		ICE, I	NC.
								De	enve	r, Co	lorad	0	

UV light also will be used to check the fluorescence of soils caught in the sampler cutting shoe.

A small portion of each soil sample will be used to estimate concentrations of VOCs in the soil headspace, while the larger portion of the soil sample will be prepared for shipment to a fixed-base laboratory for analysis. Soil for headspace VOC analysis will be obtained from either the sampler cutting shoe or by cutting off a 1- to 2-inch section of the clear plastic liner containing the sample core and removing the soil from this smaller section. Each headspace VOC screening sample will be placed in a sealed plastic bag or mason jar and allowed to sit for at least 5 minutes. VOC concentrations in the soil headspace will then be determined using an OVM or PID, and the results will be recorded in the field records by the Parsons ES field scientist.

The larger segment of the soil-filled plastic liner (a minimum 6-inch-long section) will be submitted to the USEPA National Risk Management Research Laboratory (NRMRL) in Ada, Oklahoma for soil analyses using the methods listed in Table 3.1. Sample handling and preservation procedures are discussed in Section 3.5.

3.2.2 Groundwater and Free Product Sampling

The primary objective of free product sampling for this study is to determine the impact of weathering processes on the mobile LNAPL. In addition, groundwater samples will be collected to evaluate partitioning of BTEX contaminants into groundwater below the oil/water interface to evaluate whether an equilibrium relationship exists between the mobile LNAPL and groundwater at the site.

3.2.2.1 Free Product and Groundwater Sampling Locations

Free product and groundwater samples will be collected from two locations at each of the selected sites. Every attempt will be made to collect free product and groundwater samples from the same two Geoprobe® boreholes from which soil samples are collected for laboratory analysis (see Section 3.2.1). From each sampling location, two free product samples and one groundwater sample will be collected. A total of four free product samples and two groundwater samples will be collected from each site selected for the study.

3.2.2.2 Free Product and Groundwater Sampling Procedures

This section describes the procedures for collection of free product and groundwater quality samples from the Geoprobe® boreholes. In the event, sampling cannot be performed with the Geoprobe®, efforts will be made to install temporary monitoring points (see Section 3.3) for free product and groundwater sampling. If neither of the above methods are feasible based on Geoprobe® limitations and/or site conditions, sampling may be performed from existing site monitoring wells known to contain free product (see Section 3.4).

After collection of soil samples from the capillary fringe (Section 3.2.1), the Geoprobe[®] will be used to advance a 2-inch, hollow, drive rod with a sacrificial steel drive point to a depth approximately 2 feet below the water table. Then 1-inch-diameter

polyvinyl chloride (PVC) casing with a 5-foot-long section of factory-slotted well screen on the leading edge will be inserted through the hollow drive rod. The drive rod will be disengaged from the drive point and retracted 1 foot so that the PVC well screen is exposed to the formation and groundwater below the free product layer. A groundwater sample will be collected from this point using either a peristaltic pump or a mini-bailer following adequate purging (see Sections 3.2.2.2.2 and 3.2.2.2.3).

Following collection of the groundwater sample, the hollow drive rod will be fully retracted so that the screen is fully exposed across the water table and the capillary fringe. After an adequate stabilization period, the free product thickness will be measured with an oil/water probe, and a free product sample will be collected. Figure 3.3 illustrates the Geoprobe® sampler in groundwater sample collection mode, and free product measurement and sample collection mode.

In order to maintain a high degree of QC during the proposed sampling events, sampling will be conducted by qualified scientists and technicians from Parson ES who are trained in the conduct of groundwater and free product sampling, records documentation, and chain-of-custody procedures. In addition, sampling personnel will have thoroughly reviewed this work plan prior to sample acquisition and will have a copy of the work plan available onsite for reference. Detailed sampling and sample handling procedures are presented in the following subsections.

3.2.2.1 Preparation for Sampling

All equipment to be used for sampling will be assembled and properly cleaned and calibrated (if required) prior to arriving in the field as described in Section 3.7. In addition, all record-keeping materials will be gathered prior to leaving the office.

3.2.2.2.2 Purging

Before collecting the groundwater sample using the Geoprobe®, the exposed segment of the temporary PVC screen will be purged using a peristaltic pump and high-density polyethylene (HDPE) tubing. The tubing will be slowly lowered through the temporary PVC casing to below the free product layer. Using a low peristaltic pump flow rate, a minimum of 3 casing volumes will be purged from the point with purging continuing until equilibrium conditions are achieved. Equilibrium conditions will be assessed using a flowthrough cell and field instruments. Free product within the monitoring point will be subjected to increased volatilization as compared to free product within the formation; therefore, 1 casing volume of free product will be purged prior to sampling. All purge waters and free product will be collected in accordance with facility procedures for disposal by facility personnel.



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3.2.2.2.3 Sample Extraction

Groundwater and free product samples will be collected using a peristaltic pump. Sample extraction procedures will be conducted in a manner that minimizes contaminant loss through volatilization. Following purge and recovery, new HDPE tubing will be slowly lowered through the temporary PVC casing to prevent splashing. The groundwater sample will be transferred directly from the tubing into the appropriate volatile organic analysis (VOA) sample container. The water will be carefully poured down the inner walls of the sample bottle to minimize aeration of the sample. Sample containers will be completely filled so that no air space remains in the container. Excess groundwater collected during sample extraction will be collected for proper disposal by facility personnel.

Free product sampling using the Geoprobe® will be conducted after the groundwater sample has been collected following the same procedures outlined above. However, extra care will be taken in lowering the tubing slowly into the casing (rate not to exceed 0.5 foot per minute) to minimize splashing of free product. Only one casing volume of free product will be purged prior to sampling; therefore, excess free product generated by sampling should be minimal. Excess free product will be collected and disposed of in accordance with facility procedures.

3.2.2.3 Free Product and Groundwater Sample Analysis

Two free product samples and one groundwater sample will be collected in VOA bottles from each sampling location. A total of four free product samples and two groundwater samples will be collected per site, unless otherwise specified. The groundwater sample and one of the free product samples from each location will be submitted to the NRMRL in Ada, Oklahoma for the analyses listed in Table 3.1. The remaining free product sample from each sampling location will be sent to Evergreen Analytical Laboratories, Inc. (EAL) in Wheat Ridge, Colorado for analysis (Table 3.1). Sample handling procedures are discussed in Section 3.5.

3.3 TEMPORARY MONITORING POINTS

If conditions prevent use of the Geoprobe® for groundwater and free product sampling as outlined in Section 3.2.2, it may be necessary to collect groundwater and free product samples from temporary monitoring points. Monitoring point installation procedures are presented in this section.

3.3.1 Pre-Placement Activities

All necessary digging, coring, and drilling permits will be obtained prior to mobilizing to the field. In addition, all utility lines will be located, and proposed drilling locations will be cleared prior to any intrusive activities. Responsibilities for these permits and clearances are discussed in Section 3.2.

If possible, water to be used during monitoring point installation (e.g., for equipment cleaning) will be obtained from potable water supplies at the site. Water use approval will

be verified by contacting the appropriate facility personnel. The field scientist will make the final determination as to the suitability of site water for these activities.

3.3.2 Monitoring Point Materials Decontamination

Monitoring point installation and completion materials will be inspected by the field scientist and determined to be clean and acceptable prior to use. If not factory sealed, the well points, casing, and tubing will be cleaned prior to use with a high-pressure, steam/hot-water cleaner using approved water. Materials that cannot be cleaned to the satisfaction of the field scientist will not be used.

3.3.3 Installation and Materials

Subsurface conditions permitting, temporary monitoring points constructed of either 0.75-inch outside-diameter (OD)/0.5-inch inside-diameter (ID), or 1-inch OD/0.75-inch ID, PVC casing and well screen may be used for groundwater and free product sampling if the procedures outlined in Section 3.2 are not possible. For each monitoring point, approximately 5 feet of factory-slotted screen will be installed in the borehole punched for soil sampling using the Geoprobe[®].

Prior to installation, it will be necessary to advance the Geoprobe® borehole approximately 4 feet below the water table. A macro-core pre-probe (2-inch-OD drive point or larger) or soil sampler may be used for borehole advancement. Installation of temporary monitoring points requires that the borehole remain open after the borehole is punched below the water table and the rods are withdrawn. Upon removing the rods, the borehole depth will be measured to determine if the hole remains open. If the borehole is open, the PVC casing and screen will be placed so that at least 3 feet of screen is below the water table. The annular space around the screen will be filled with sand filter pack. No grout or bentonite will be placed within the annular space as the monitoring point will be removed upon completion of sampling. Monitoring point construction details will be noted in the site field notes. This information will become part of the field records for the site.

Monitoring point screens will be constructed of Schedule 40 PVC with either an ID of 0.5 inch or 0.75 inch depending on site conditions. The screens will be factory slotted with 0.01-inch openings. Monitoring point screens will be placed to sample both groundwater and free product. Blank monitoring point casing will be constructed of Schedule 40 PVC with an ID of 0.5 or 0.75 inch. All monitoring point casing sections will be flush-threaded; joints will not be glued. The casing at each monitoring point will be fitted with a bottom cap and a top cap constructed of PVC.

If subsurface conditions do not permit the boreholes to stay open (i.e., if the formation collapses in the hole), groundwater and free product samples will be collected from site monitoring wells in accordance with procedures outlined in Section 3.4. The decision to install 0.5-inch-ID PVC monitoring points will be made in the field once the open-hole stability of subsurface soils and availability of Geoprobe® equipment can be evaluated.

3.4 WELL AND MONITORING POINT SAMPLING PROCEDURES

If free product and groundwater cannot be sampled in accordance with the general procedures outlined in Section 3.2, samples will be collected from temporarily installed monitoring points if possible (Section 3.3). If monitoring point installation is not feasible, free product and groundwater samples will be collected from site monitoring wells that contain free product.

The following paragraphs present the procedures to be followed for free product and groundwater sample collection from groundwater monitoring wells and monitoring points. These activities will be performed in the order presented below. Exceptions to this procedure will be noted in the field scientist's field notebook or on the groundwater and free product sampling record (Figure 3.4).

Special care will be taken to prevent cross-contamination of the free product and groundwater samples. The primary way in which sample contamination can occur are through cross-contamination due to insufficient cleaning of equipment between wells and monitoring points. To prevent such contamination, the oil/water interface probe and cable used to determine static water levels and free product thicknesses will be thoroughly cleaned before and after field use and between uses at different sampling locations according to the procedures presented in Section 3.7. Dedicated tubing will be used at each well or monitoring point developed, purged, and/or sampled with the sampling pump. Pumps and nondisposable bailers will be decontaminated according to procedures listed in Section 3.7. In addition to the use of properly cleaned equipment, a clean pair of new, disposable nitrile or latex gloves will be worn by the sampling personnel each time a different well or monitoring point is sampled.

3.4.1 Preparation of Location

Prior to starting the sampling procedure, the area around the existing wells and new monitoring points will be cleared of foreign materials, such as brush, rocks, and debris. These procedures will prevent sampling equipment from inadvertently contacting debris around the monitoring well/point.

3.4.2 Monitoring Well/Point Purging

After sampling free product and prior to sampling groundwater in monitoring points or monitoring wells, it will be necessary to purge the groundwater present in the casing. The volume of water contained within the monitoring well/point casing at the time of sampling will be calculated, and at least three times the calculated volume will be removed from the well/point. One casing volume of free product will be purged prior to sampling. A peristaltic pump will be used for monitoring well and monitoring point purging, depth and volume permitting, and a Grundfos Redi-Flo II[®] pump,

GROUNDWATER AND FREE	PRODUCT SAMPLING RECORD
Sam	Job Number 729691.35730 Facility Name Site Name nple Location ID Sample Date
DATE AND TIME OF SAMPLING:, 19 SAMPLE COLLECTED BY: of Parson	99a.m./p.m. <u>s ES</u>
Check-off 1 GROUNDWATER/FREE PRODUCT COL a [] GEOPROBE HOLLOW DRIV b [] TEMPORARY MONITORIN c [] GROUNDWATER MONITO	LECTED FROM /E ROD AND PVC ASSEMBLY (ref: Work Plan, Section 3.2) G POINT (ref: Work Plan, Section 3.4) RING WELL (ref: Work Plan, Section 3.4) (MW Number)
2 [] EQUIPMENT CLEANED BEFORE USE V Items Cleaned (List):	WITH
3 [] PRODUCT DEPTH Measured with:	FT. BELOW DATUM
4 [] WATER DEPTH Measured with:	FT. BELOW DATUM
5 [] . WATER-CONDITION BEFORE WELL E Appearance: Odor: Other Comments:	VACUATION (Describe):
6 [] WELL EVACUATION: Method: Volume Removed: Observations: Water (slig Water level Water odor Other comr	htly - very) cloudy (rose - fell - no change) s:nents:
	FIGURE 3.4
	GROUNDWATER AND FREE PRODUCT SAMPLING RECORD
c:\cbs\weatherg\gwsample.doc Page 1 of 2	Fuel Weathering Study PARSONS ENGINEERING SCIENCE, INC.

	Fuel Weathering Study
	GROUNDWATER AND FREE PRODUCT SAMPLING RECORD
	FIGURE 3.4 (CONTINUED)
10[]	OTHER COMMENTS:
	 [] Container Sides Labeled [] Container Lids Taped [] Containers Placed in Ice Chest
9[]	CONTAINER HANDLING:
8[]	SAMPLE CONTAINERS (material, number, size):
	[] Pump, type:
7[]	SAMPLE EXTRACTION METHOD.
	Groundwater and Free Product Sampling Record Sample Location ID (Cont'd)

Waterra[®] inertial pump, or bailer will be used to purge all monitoring wells or points in which a peristaltic pump cannot be used. All purge and free product waters will be collected for proper disposal by facility personnel.

If a monitoring well/point is evacuated to a dry state during purging, the monitoring well/point will be allowed to recharge, and the groundwater sample will be collected as soon as sufficient water is present in the monitoring well/point to obtain the necessary sample quantity. Sample compositing or sampling over a lengthy period by accumulating small volumes of water at different times to obtain a sample of sufficient volume will not be allowed.

3.4.3 Free Product and Groundwater Sample Extraction and Analysis

Samples of free product and groundwater will be extracted following the procedures outlined in Section 3.2.2.2.3 and will be analyzed as described in Section 3.2.2.3. However, when a sample is collected from a temporary monitoring point or from a site monitoring well, the free product samples will be collected first, then the groundwater sample will be collected.

3.5 SAMPLE HANDLING FOR LABORATORY ANALYSIS

This section describes the handling of soil, free product, and groundwater samples from the time of sampling until the samples are delivered to either the NRMRL or EAL.

3.5.1 Sample Preservation

NRMRL and EAL will add any necessary chemical preservatives prior to shipment of sample containers to the field for sample collection. After sample collection, samples will be prepared for transportation to the analytical laboratory by placing the samples in a cooler containing ice to maintain a shipping temperature of as close to 4 degrees centigrade (°C) as possible.

3.5.2 Sample Containers and Labels

Free product and groundwater sample containers and appropriate container lids will be provided by the NRMRL and EAL. For samples requiring chemical preservation, preservatives will be added to containers at the laboratory. No containers will be provided for soil sampling unless otherwise specified. After soil sample collection, each open end of the clear plastic liner will be covered with Teflon[®] fabric and tightly capped using vinyl liner end caps. Free product and groundwater sample containers will be filled as described in Section 3.2.2.2.3, and the container lids will be tightly closed. A sample label will be firmly attached to the container side or plastic liner, and the following information will be legibly and indelibly written on the label:

- Facility name;
- Sample identification;
- Sample type (e.g., groundwater, soil, free product);

- Sampling date;
- Sampling time;
- Analyses requested.

3.5.3 Sample Shipment

After the samples are sealed and labeled, they will be packaged for transport to NRMRL or Evergreen Analytical as appropriate. Delivery will occur as soon as possible after sample acquisition. The following packaging and labeling procedures will be followed:

- Package sample so that it will not leak, spill, or vaporize from its container;
- Cushion samples to avoid breakage; and
- Add ice to container to keep samples cool.

3.5.4 Chain-of-Custody Control

Blank chain-of-custody forms to accompany sample shipments will be provided with sample containers sent to the field by NRMRL and EAL. Chain-of-custody documentation completed by the Parsons ES field personnel will accompany packaged samples to be sent for laboratory analysis.

3.5.5 Sampling Records

In order to provide complete documentation of the sampling event, detailed records will be maintained by the field scientist. At a minimum, these records will include the following information:

- Sample location (facility name);
- Sample identification;
- Sample location map or detailed sketch;
- Date and time of sampling;
- Sampling method;
- Field observations of:
 - Sample appearance, and
 - Sample odor;
- Free product thickness before and after purging;

- Water level before and after purging;
- Purge volume;
- Sample depth (soil samples, only);
- Monitoring well/point condition (free product and groundwater samples);
- Sampler's identification;
- Any other relevant information.

Sampling information will be recorded on forms similar to those shown on Figures 3.2 and 3.4, and in the site field notes.

3.6 SITE RESTORATION

After sampling is complete, each sampling location will be restored as closely to its original condition as possible. Holes created by the Geoprobe® in sandy soils tend to cave in soon after extraction of the drive sampler. However, any test holes remaining open after extraction of the drive rod will be sealed with bentonite chips, pellets, or grout to eliminate any creation or enhancement of contaminant migration pathways to the groundwater.

Soil sampling using the Geoprobe® creates low volumes of soil waste. With the approval of facility personnel, soil not retained as samples will be returned to the open boreholes prior to sealing with bentonite. Methods of waste soil disposal will conform to IDW procedures provided by the facility.

If a temporary monitoring point is used for free product and groundwater sampling, the monitoring point will be abandoned following sample collection. The PVC casing and screen will be extracted as far as possible and discarded. Any test holes remaining open after extraction of the casing will be sealed with bentonite chips, pellets, or grout to eliminate any creation or enhancement of contaminant migration pathways to the groundwater. After monitoring point abandonment, each site will be restored as closely as possible to its original condition.

3.7 DECONTAMINATION PROCEDURES

Prior to arriving at the site, and between each sampling location, probe rods, tips, sleeves, pushrods, samplers, tools, and other downhole equipment will be decontaminated using a high-pressure, steam/hot water wash. Only potable water will be used for decontamination.

All portions of sampling and test equipment that will contact the sample matrix will be thoroughly cleaned before each use. This includes the Geoprobe® sampling tools, sampling pumps, nondisposable bailers, water level probe and cable, test equipment for onsite use, and other equipment or portions thereof that will contact the samples. Given the types of sample analyses to be conducted, the following cleaning protocol will be used:

- Wash with potable water and phosphate-free laboratory detergent (HP-II detergent solutions, as appropriate);
- Rinse with potable water;
- Rinse with isopropyl alcohol;
- Rinse with distilled or deionized water; and
- Air dry.

Any deviations from these procedures will be documented in the field scientist's field notebook and on the groundwater and free product sampling record (Figure 3.4).

If precleaned, disposable sampling equipment is used, the cleaning protocol specified above will not be required. Laboratory-supplied sample containers will be cleaned and sealed by the laboratory.

Potable water to be used during equipment cleaning, decontamination, or grouting will be obtained from one of the facility's potable water supplies. Water use approval will be verified by contacting the appropriate facility personnel. The field scientist will make the final determination as to the suitability of site water for these activities.

All rinseate will be collected for transportation and proper disposal by facility personnel. Alternate methods of rinseate disposal will be considered by the Parsons ES field scientist as recommended by facility personnel. Precautions will be taken to minimize any impact to the surrounding area that might result from decontamination operations.

SECTION 4

QUALITY ASSURANCE/QUALITY CONTROL

Field QA/QC procedures will include collection of field duplicates and trip blanks; decontamination of all equipment that contacts the sample medium before and after each use (Section 3.7); use of analyte-appropriate containers; and chain-of-custody procedures for sample handling and tracking (Section 3.6). All samples to be transferred to either NRMRL or EAL for analysis will be clearly labeled to indicate sample number, location, matrix (e.g., groundwater), and analyses requested. Samples will be preserved in accordance with the analytical methods to be used, and sample containers will be packaged in coolers with ice to maintain a temperature of as close to 4°C as possible.

All field sampling activities will be recorded in a bound, sequentially paginated field notebook or on the appropriate sample form (Figures 3.2 and 3.4) in permanent ink. All sample collection entries will include the date, time, sample locations and numbers, notations of field observations (e.g., odors, UV and OVM readings), and the sampler's name and signature. Field QC samples will be collected in accordance with the program described below, and as summarized in Table 4.1.

QA/QC sampling will include collection and analysis of duplicate free product and groundwater samples and replicate soil samples, trip blanks, and matrix spike samples. Internal laboratory QC analyses will involve the analysis of laboratory control samples (LCSs) and laboratory method blanks (LMBs). QA/QC objectives for each of these samples, blanks, and spikes are described below.

Duplicate free product and water and replicate soil samples will be collected at a frequency of one sample for every five sites sampled (i.e., for every 10 soil, free product, and groundwater samples shipped to NRMRL or EAL). Soil, free product, and groundwater samples collected with the Geoprobe® should provide sufficient volume for replicate/duplicate analyses.

A trip blank will be analyzed to assess the effects of ambient conditions on sampling results during the transportation of samples. The trip blank will be prepared by the laboratory. A trip blank will be transported inside each cooler that contains samples for VOC analysis. Trip blanks will be analyzed for VOCs.
TABLE 4.1QA/QC SAMPLING PROGRAMFuel Weathering Study

QA/QC Sample Types	Frequency to be Collected and/or Analyzed	Analytical Methods		
Duplicates/Replicates	1 soil, groundwater, and free product sample every 5 sites	Volatile Organic Compounds (VOCs)		
Trip Blanks	One per shipping cooler containing VOC samples	VOCs		
Matrix Spike Samples	Once per sampling event	VOCs		
Laboratory Control Sample	Once per method per medium	Laboratory Control Charts (Method Specific)		
Laboratory Method Blanks	Once per method per medium	Laboratory Control Charts (Method Specific)		

Matrix spikes will be prepared in the laboratory and used to establish matrix effects for samples analyzed for VOCs. LCSs and LMBs will be prepared internally by the laboratory and will be analyzed each day samples from the project are analyzed. Samples will be reanalyzed in cases where the LCS or LMB are out of the control limits. Control charts for LCSs and LMBs will be developed by the laboratory and monitored for the analytical methods used (see Table 3.1).

SECTION 5

DATA ANALYSIS AND REPORT

Following receipt and compilation of soil, groundwater, and LNAPL analytical results for the selected sites, a variety of data analyses will be performed. As previously stated, the primary objective of this study is to determine an average range of natural weathering rates for mobile LNAPLs. In addition, mobile LNAPL analytical data will be compared with soil analytical data to evaluate the impact of weathering on residual LNAPL contamination in capillary fringe soils. Finally, the degree of contaminant partitioning occurring between the mobile LNAPL and groundwater will be evaluated.

LNAPL weathering will be assessed by evaluating the mass fraction reduction of BTEX in collected LNAPL samples. Analytical results will be compared to typical initial BTEX concentrations in "fresh" fuel products based on literature values, and to available historic data for the sites. Table 5.1 gives mass fraction values for BTEX in fresh gasoline, JP-4 jet fuel, and aviation gasoline. Literature values such as these will be assumed as the initial composition of the fuel prior to their environmental release. Using the date of the product release and an assumed initial BTEX composition, BTEX concentrations in LNAPL samples from the selected sites can be used to determine the degree of weathering (i.e., BTEX mass fraction depletion) that has occurred over a known period of time. It is anticipated that analytical results will provide a range of LNAPL weathering rates requiring some site-specific trend analyses based on ages of the releases, fuel types, and geologic/hydrogeologic conditions. It is anticipated that a range of weathering rates will be determined for releases of specific fuel types that are approximately 0 to 5 years old, 5 to 10 years old, 10 to 20 years old, and older than 20 years. Weathering rates are expected to decrease with the age of the fuel release (i.e., 0- to 5-year rates are expected to be greater than 10- to 20-year rates).

Weathering effects on residual-LNAPL-contaminated soils also will be assessed. The BTEX/TPH ratio in soil and the BTEX/TPH ratio in mobile LNAPL (TPH in LNAPL is 100 percent) will be compared to determine if weathering effects are more significant for capillary fringe soils than for mobile LNAPL. It is anticipated that BTEX/TPH ratios in capillary fringe soils will be lower than mobile LNAPL BTEX mass fractions as a result of increased weathering in soils due to increased biodegradation and possibly increased volatilization. Comparative trend analyses of soil BTEX mass fractions and mobile LNAPL BTEX mass fraction will be performed.

The degree of BTEX partitioning from mobile LNAPL into groundwater will be assessed and compared to bench-scale and theoretical equilibrium conditions. BTEX concentrations in mobile LNAPL and dissolved BTEX concentrations in groundwater, as determined by NRMRL, will be compared to evaluate a site-specific parameter similar to an octanol/water coefficient (K_{ow}). This fuel/water partitioning coefficient (K_{fw}) like the octanol/water

TABLE 5.1 BTEX CONCENTRATIONS IN FRESH FUELS Fuel Weathering Study

Constituent	Gasoline87 Octane89 Octane92 Octane(percent)(percent)(percent)			JP-4 (percent)	Aviation Gas (percent)
Benzene	3.15	2.51	2.67	1.07	0.13
Toluene	10.59	12.96	13.13	5.64	22.73
Ethylbenzene	2.10	2.13	2.53	2.03	0.11
m- + p - Xylenes	7.97	8.61	9.18	6.03	0.20
o- Xylenes	2.90	3.25	3.23	1.72	0.11
Total BTEX	26.71	29.46	30.74	16.49	23.28

Source: Kaplan and Galperin, 1996.

coefficient provides a measure of how an organic compound (fuel) will partition between oily and aqueous phases, and gives useful information for estimating chemical fate and transport. The fuel/water partitioning coefficient is a dimensionless constant defined by:

> $K_{fw} = C_f/C_w$ $C_f = contaminant concentration in fuel (mg/L or µg/L)$

 C_w = contaminant concentration in water (mg/L or μ g/L)

The K_{fw} determined from mobile LNAPL and groundwater sample results will be considered a "field" K_{fw} . Some emulsification of mobile LNAPL with underlying groundwater is likely to occur during sample collection activities, and as a result, the field K_{fw} determined from NRMRL sample results may not represent equilibrium conditions. To simulate equilibrium conditions, a separate bench-scale determination of the fuel/water partitioning coefficient will be performed.

The bench-scale testing will be performed by EAL using the duplicate mobile LNAPL samples collected from each sampling location. The mobile LNAPL samples will be mixed with distilled water in a VOA bottle, inverted, and placed in a laboratory rack motionless for approximately 2 weeks. After 2 weeks, a sample of the water will be removed from the bottom of the VOA bottle, and a sample of LNAPL will be removed from the top of the bottle, and analyzed for BTEX by USEPA Method E602, and SW8020, respectively. Dissolved BTEX concentrations in the water will be compared to mobile LNAPL concentrations to determine a "bench-scale" K_{fw} for each sampling location. The field and bench-scale K_{fw}'s will be compared and evaluated for each site.

Lastly, BTEX partitioning from the mobile LNAPL to groundwater will be compared with theoretical partitioning according to Raoult's law. A study by Cline *et al.* (1991) showed that equilibrium partitioning of BTEX into water from gasoline followed near-ideal behavior, and could be described by Raoult's law. According to Raoult's law, the concentration of a gasoline constituent in the aqueous phase (C_i) is equal to the mole fraction of the constituent i in the gasoline (X_i) multiplied by the aqueous solubility of the pure constituent (S_i).

$C_i = X_i S_i \\$

Based on Raoult's law, the amount of constituent that partitions into water is dependent on its mole fraction in the mobile LNAPL, and not on the amount of LNAPL present in the groundwater/soil system.

A report detailing the results of the fuel weathering study will be prepared following completion of the literature search and analysis of soil, groundwater, and LNAPL sample data from the selected sites. The report will include an introduction; a summary of the literature search findings; detailed site descriptions for selected sites, including existing soil, groundwater, and mobile LNAPL chemical data; and results of the trend, weathering, and equilibrium analyses described above.

SECTION 6

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ATTACHMENT A

FUELS WEATHERING LITERATURE SEARCH DRAFT REFERENCE LIST

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A-2

SITE ADDENDA

2.2.6 Tank Farm C, Marine Corps Air Station, Beaufort, South Carolina

The Marine Corps Air Station (MCAS) in Beaufort, South Carolina is located approximately 60 miles south-southwest of Charleston and 1 mile northwest of the City of Beaufort on the South Carolina coast. Tank Farm C is located on the western side of the MCAS approximately 800 feet east of Highway 21 near the intersection of Reed Avenue and R.C. West Road. The facility consists of two 210,000-gallon cut-and-cover steel tanks, a truck loading and unloading stand, a rail line and a railcar unloading stand. A layout of the site is provided on Figure A.1.

The information below was extracted from the ABB Environmental Services, Inc (ABB, 1993) Draft Final Contamination Assessment (CA) Report for the site. In June 1990, approximately 10,600 gallons of JP-5 aviation fuel was released from the underground storage tank (UST) system at Tank Farm C during fuel transfer operations from the adjacent railcar unloading stand into tanks 401 and 402. The release was confirmed to have occurred from the existing 8-inch fill lines located south of tank 401 when fuel was observed seeping upward to the ground surface directly above these fill lines. Upon discovery of the release, the fuel transfer operation was halted, the leak repaired and the system restarted. Shortly after system restart, a second leak occurred in the line approximately 15 feet west of the initial leak location. The transfer operation was then permanently terminated. In response to the leaks, 13,000 gallons of mixed fuel and water were pumped and recovered from a pit excavation opened to repair the line leaks. Based on information collected during the CA, no other reported releases are known to have occurred at the site.

Tank Farm C is underlain by fine-grained, silty sand to approximately 38 feet below ground surface (bgs). The sand lense forms the surficial or shallow aquifer and the depth to groundwater at the site ranges from approximately 2 to 8 feet bgs. The sand is underlain by a dark-gray to olive-gray clay which has been identified as the regional confining unit or Hawthorne formation. The saturated thickness of the shallow aquifer is approximately 33 feet. Groundwater at the site flows generally to the northwest.

In September 1991, eight 2-inch release detection monitoring wells were installed to approximately 20 feet bgs around tanks 401 and 402 (BFT-401-1 through BFT-401-4 and BFT-402-1 through BFT-402-4). Free product was observed, during installation and development of well BFT-401-3, the well closest to the documented fuel release location (Figure A.1). Since the initial recovery of 13,000 gallons of mixed JP-5 fuel and water, manual bailing and recovery of free product from well BFT-401-3 has been regularly performed by MCAS personnel.

The CA site investigation was conducted by ABB in March and April 1993 and involved the advancement of 74 soil borings (B-1 through B-74) to the water table and the installation of 10 shallow monitoring wells (BFT-TF-9 through BFT-TF-18), one deep well (BFT-TF-19), and one recovery well (BFT-TF-RW-1). The CA site

investigation determined that the primary soil contamination resulting from the 1990 release is at 1-to 3-foot bgs in the area south of the tanks. At this time, the estimated horizontal extent of free product encompassed an area which included wells BFT-401-3 and BFT-401-2.

No remedial activities have been conducted at the site. The US Geological Survey has performed modeling to assess the impact of various remedial alternatives at the site including source excavation with long-term passive bioremediation. However, monitoring of the dissolved and free product plumes has shown that the plumes are stable and not migrating from the site (Araico, 1996).

Proposed sampling locations for the Tank Farm C site are shown on Figure A.1. Water level measurements conducted at the site in August 1996, indicated approximately 1.5 inches of free product within well BFT-401-3; however, free product was not detected at well BFT-401-2. Sampling procedures are outlined in Section 3 of the work plan. South Carolina Department of Health and Environmental Control approval will be obtained prior to conducting intrusive sampling activities.

References

- ABB Environmental Services Inc. (ABB). 1993. Final Draft Contamination Assessment Report, Tank Farm C, GWPD Site #A-07-AA-155513, Marine Corps Air Station, Beaufort, South Carolina. October.
- Araico, F. 1996. Environmental Affairs (NREAO), Marine Corps Air Station, Beaufort, South Carolina. Telephone Conversation. October 30.



2.2.7 Tank 349 Site, Offutt Air Force Base, Nebraska

Offutt Air Force Base (AFB) is located in eastern Nebraska approximately 4 miles south of Omaha. The Tank 349 site is located in the northern portion of the Base, along Peacekeeper Drive northwest of Building 301 (Figure A.1). Fuel hydrocarbon-contaminated soil and groundwater at the site and the existence of subsurface free product appears to be the result of long-term leaking from former gasoline underground storage tanks (USTs). The exact date and duration of the release at Tank 349 is not known, but historical LNAPL results from the site are available.

The Tank 349 site is the former location of two 500-gallon gasoline USTs and one 700-gallon diesel UST. Dates of installation for the former USTs are not identified in the site literature. In September 1990, leak verification testing showed one of the gasoline USTs to be leaking. The USTs were taken out of service approximately 1 month prior to verification testing and were never returned to service (Cork, 1997). The two gasoline USTs were excavated in April 1993 and the diesel UST was excavated in September 1993 (Terracon, 1994). During tank removal, 0.25-inch holes were observed in each of the gasoline tanks and in the line connecting the two tanks, but no defects were evident on the diesel tank (Terracon, 1993a; Terracon, 1993b). Contaminated soils were observed during the UST removals; however, soil contamination at the site of the diesel tank excavation appeared to be of little consequence (Terracon, 1993b).

Subsurface soils in the vicinity of the Tank 349 site consist of lean reddish-brown to brown clay overlying interbedded fine to very fine sands, fine to very fine clayey sands, and clays. Occasionally, a trace of sand or gravel was identified in the upper reddish-brown to brown clay. The first distinct sands are encountered near the water table approximately 41 feet below ground surface (bgs) across most of the site. Groundwater depths in the contaminated areas immediately adjacent to Tank 349 have generally varied between 39 and 42 feet bgs. A predominantly easterly groundwater flow direction exists at the site (Parsons Engineering Science [Parsons ES], 1997).

Between December 1993 and March 1994, Terracon installed 14 monitoring wells (MW 349-1 to MW 349-14) and performed soil and groundwater sampling at the Tank 349 site. Field screening of soil samples indicated significant hydrocarbon contamination at wells located northeast and east of the former USTs (MW 349-1, MW 349-2, MW 349-6, MW 349-7, and MW 349-8). During these investigations, free product was measured in MW 349-1, MW 349-2, and MW 349-6, and significant dissolved hydrocarbon contamination was detected in groundwater from MW 349-6, MW 349-7, and MW 349-8.

In 1994, Parsons ES, in conjunction with researchers from the United States Environmental Protection Agency (USEPA) National Risk Management Research Laboratory (NRMRL) was retained by the United States Air Force Center for Environmental Excellence (AFCEE) to conduct site characterization and groundwater modeling to evaluate the scientific defensibility of remediation by natural attenuation (RNA) with long-term monitoring (LTM) as a remedial option for contaminated groundwater at the Tank 349 site. As part of this investigation, Parsons ES performed two phases of site characterization, the first in September and November 1994 and the second in June 1996. The 1994 investigation included performing cone-penetrometer testing (CPT) with laser-induced fluorometry (LIF), sampling and analyzing soils from CPT boreholes, sampling and analyzing groundwater from existing MWs, and measuring and estimating hydrogeologic parameters. In 1996, groundwater elevations were again measured and groundwater quality samples were collected from selected MWs. During each investigation, free product samples also were collected from site MWs and analyzed for benzene, toluene, ethylbenzene, xylenes (BTEX), and other fuel constituents. Free product thicknesses measured during the June 1996 investigation are shown on Figure A.1 and results from the 1994 and 1996 free product sampling events are presented in Table A.1. Other than the tank removal and evaluation of RNA with LTM, no remedial actions have been conducted at the Tank 349 site.

Proposed sampling locations for the Tank 349 site are shown on Figure A.1. Free product measurements conducted at the site in June 1996 ranged from 0.27 feet at MW 349-7 to 2.23 feet at MW 349-6. Due to the depth of the water table at this site, free product and groundwater samples will be collected from existing monitoring wells, rather than from Geoprobe® boreholes. If possible, groundwater and free product samples will be collected from MW 349-1 and MW 349-7 in order to compare with historic free product sample results at these two locations. Sampling procedures are outlined in Section 3 of the work plan. Sampling will be performed in accordance with Offutt AFB and Nebraska Department of Environmental Quality (NDEQ) requirements.

REFERENCES

- Cork, Philip. 1997. 55 CES/CEVR, Offutt Air Force Base, Nebraska. Telephone Conversation. January 3.
- Parsons Engineering Science, Inc. 1997. Intrinsic Remediation Treatability Study for Tank 349 Site, Offutt Air Force Base, Nebraska. Denver, CO. (In Draft).
- Terracon, 1994. Assessment Report Tank 349, Facility 6275, Offutt AFB, Nebraska. April.
- Terracon, 1993a. Gasoline Underground Storage Tank Closure Assessment Report, Facility 349, Offutt Air Force Base, Nebraska. April.
- Terracon, 1993b. Diesel Underground Storage Tank Closure Assessment Report, Facility 349, Offutt Air Force Base, Nebraska. September.



Table A.1 **Summary of Free Product Analysis Results** Tank 349 Site Offutt AFB, Nebraska

Sample	Sample	Benzene	Toluene	Ethylbenzene	p-Xylene	m-Xylene	o-Xylene	Total BTEX ^{a/}	Total BTEX
Location	Date	(mg/L) ^{b/}	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(mg/L)	(% Mass Fraction) ^{c/}
MW 349-1	Nov-94	8,280	41,100	10,300	8,980	21,700	11,400	101,760	13.90
	Jun-96	8,420	36,400	11,400	9,360	17,700	10,700	93,980	12.84
MW 349-7	Nov-94	5,620	28,600	9,260	8,920	21,400	11,600	85,400	11.67
MW 349-8	Jun-96	955	12,300	10,100	8,030	15,400	9,150	55,935	7.64

^{a/} BTEX = benzene, toluene, ethylbenzene, and xylenes.

^{b/} mg/L = milligrams per liter.
^{c/} Assumes fuel density of 0.7321 grams per milliliter.

2.2.8 Spill Site No. 2, Eaker Air Force Base, Arkansas

Eaker Air Force Base (AFB) is located in northeastern Arkansas approximately 3 miles west of the City of Blytheville and immediately east of the City of Gosnell. Spill Site No. 2 is located in the south-central portion of Eaker AFB at the intersection of South Access Road and Taxiway G (Figure A.1). Fuel hydrocarbon-contaminated soil and groundwater at the site and the existence of subsurface free product appears to be the result of two separate leaks in a 10-inch JP-4 jet fuel pipeline located at the site. The leaks were identified during pressure testing in 1973 and 1974 and according to base personnel a large quantity of fuel was released as a result of the pipeline leaks with little being recovered during pipeline repair activities. No other leaks are known to have occurred at this site; however, the pipeline was not abandoned until 1995 (during base closure activities). The following information was summarized from the Resource Conservation and Recovery Act Facility Investigation (RFI) Report (Haliburton NUS, 1996).

At Spill Site No. 2 the soil profile is composed primarily of a surficial sandy silt loam extending from 1 to 9 feet below ground surface (bgs). Three distinct lithologic units are encountered below the soil profile. The first unit consists of a silty clay that extends to 23 feet bgs. The upper portions of this unit are slightly silty and the lower portions are more sandy. The second unit consists of silt and is first encountered from 9 to more than 25 feet bgs and occurs continuously along the western side of the site. In the central and eastern portions of the site this unit appears as lenses between the clay and underlying sand. The third unit is sand and is first encountered from 8.5 to 37 feet bgs.

The lithology described above encompasses two separate hydrogeologic units. The silts and clays represent an aquitard, and the underlying sand unit is classified as the aquifer. Seepage velocity in the aquitard and aquifer has estimated at 59 feet per year (ft/yr) and 16 ft/yr, respectively. These seepage velocities reflect the lithology and gradients of the two hydrogeologic units. The low seepage velocity in the aquifer is the result of a low aquifer gradient combined with the presence of a poorly sorted sand that contains a significant amount of silt and clay.

During previous investigations at the site several monitoring wells have been installed. In 1988, Halliburton NUS installed three monitoring wells in the vicinity of the 1973 leak (Figure A.1) and performed soil and groundwater sampling at these three locations. Remedial investigation (RI) fieldwork performed in 1991 and 1992 indicated the fuel spill was much larger than originally believed. As part of the RFI, nine monitoring wells were installed around the perimeter of the site, with one monitoring well (MW312) installed near the original pipeline leak location. Following the discovery of free product in well MW306, four additional monitoring wells (MW314, MW315, MW316, and MW317) were installed on the northeastern side of the site. The wells were installed to depths of 18 to 40 feet.

During the RFI, benzene, toluene, ethylbenzene, and xylenes (BTEX) were the most frequently detected petroleum contaminants at the site. With the exception of toluene, the highest concentrations of BTEX compounds (20 milligrams per kilogram [mg/kg], 54 mg/kg, and 300 mg/kg for benzene, ethylbenzene, and total xylenes, respectively) were detected in the MW303 borehole. The highest toluene concentration (0.24 mg/kg) was detected in a sample collected from the MW316 borehole.

The most significant free product thicknesses at the site typically occur in MW306 and MW316. In November 1995, 6.35 and 3.9 feet of free product were reported in each well, respectively. The presence of free product in MW316 is related to the proximity of this well to the original leak location while free product in MW306 is believed to have migrated in a northeasterly direction from the original pipeline leak location along the trace of the existing culvert.

Proposed sampling locations for Spill Site No. 2 are shown on Figure A.1. Free product at the site will most likely be collected from existing monitoring wells MW306 and MW316. In the event sufficient free product for laboratory analysis cannot be collected from these two monitoring wells, other site monitoring wells will be investigated for free product and may be sampled if appropriate. Soil and groundwater samples from this site will be collected from newly installed Geoprobe® boreholes to be placed within 5 to 7 feet of the two wells from which free product sampling is performed. Sampling procedures are outlined in Section 3 of the work plan. Sampling will be performed in accordance with Eaker AFB requirements.

REFERENCES

Haliburton NUS, 1996. RCRA Facility Investigation Final Report, Eaker Air Force Base, Arkansas. Oak Ridge, Tennessee. May.

2.2.9 Building 4522, Seymour Johnson Air Force Base, North Carolina

The following discussion of the Building 4522 Site at Seymour Johnson Air Force Base (AFB), North Carolina was summarized from the Comprehensive Site Assessment performed by Parsons Engineering Science, Inc. (Parsons ES, 1996) and a conversation with the AFB point of contact for the site (Chastain, 1997).

Seymour Johnson AFB is located south of State Highway 70 in central Wayne County, just south of Goldsboro. On December 14, 1995, the Seymour Johnson AFB Fire Department was notified of a release of JP-8 aviation fuel from a valve pit located approximately 130 feet west of Building 4522 and 350 feet east of Taxiway F (see Figure A.1). The release was attributed to an ineffective "O"-ring seated within a flexible coupling inside the valve pit. By the time the fire department responded to the release, the valve pit, as well as a 90-foot by 35-foot area of grass surrounding the pit, was flooded with fuel. As part of the abatement measures, a trench was dug between the valve pit and a storm water drain located about 90 feet to the west. The trench was used as a sump, and a pump was used to recover approximately 2,200 gallons of the estimated 5,000 gallons of released fuel (Parsons ES, 1996).

The site geology includes a mix of unconsolidated deposits. Brown to light gray fine to medium sand generally occurs from ground surface to approximately 8 to 12 feet below ground surface (bgs). Interfingered layers of a dark gray clay and fine to medium sand underlie the surficial sands to a depth of approximately 40 feet bgs. A dark gray clay appears to be present below the clay/sand layer and extends to at least 47 feet bgs. Depth to groundwater ranges from 3.7 to 8.6 feet across the site, with flow to the west-northwest away from Building 4522 toward Taxiway F (Parsons ES, 1996).

On January 18, 1996, Contractors and Engineers Services, Inc. of Goldsboro, NC installed a shallow monitoring well to determine if groundwater at the site had been impacted by the release. The concentration of groundwater contaminants found in the temporary well were as follows: benzene (2,503 micrograms per liter [μ g/L]), toluene (2,060 μ g/L), ethylbenzene (1,097 μ g/L), xylenes (5,792 μ g/L) (BTEX), and lead (91.7 μ g/L). Results of a soil sample collected at 5.5 feet bgs in the source area indicated a total petroleum hydrocarbon (TPH) concentration of 5,505 milligrams per kilogram (mg/kg) (Parsons ES, 1996).

In April 1996, Parsons ES performed a Comprehensive Site Assessment to determine the extent of soil and groundwater contamination resulting from the fuel release. As part of the site assessment, Parsons ES performed soil sampling in the vicinity of the release and at boreholes being advanced for monitoring well placement. Soil screening performed using a photoionization detector (PID) identified volatile concentrations in soil headspace samples ranging from nondetect (ND) to 1,356 parts per million, volume per volume (ppmv). Laboratory analytical results from a soil sample collected approximately 200 feet downgradient of the release indicated a TPH concentration (in the diesel/kerosene fraction) of 511 mg/kg. The remaining three soil samples collected in the vicinity of the spill area showed no detectable TPH concentrations.

Six groundwater monitoring wells (MW-1S, MW-1D, and MW-2 through MW-5) were installed by Parsons ES in April 1996 and groundwater samples were collected from the new wells following completion and development. The source area monitoring well, MW-1S, was not sampled due to the presence of free product. Approximately 2.8 feet of free product was measured at MW-1S during the April 1996 sampling event. BTEX and naphthalene concentrations of 1,400 μ g/L, 3,700 μ g/L, 730 μ g/L, 2,900 μ g/L, and 120 μ g/L, respectively, were detected in a groundwater sample from MW-4, located approximately 180 feet downgradient of the valve pit. Ethylbenzene, xylenes, and naphthalene were found in relative concentrations of 23 μ g/L, 55 μ g/L, and 13 μ g/L at MW-3, located approximately 150 feet southwest of the valve pit. A groundwater sample collected from MW-1D was used to assess the vertical extent of contamination. No petroleum hydrocarbon contamination was detected in MW-1D, which is screened 31 to 41 feet bgs. Petroleum hydrocarbon concentrations were not detected above the method detection limits in any of the other MWs located at the site.

Further site assessment activities were completed in July 1996, when the US Army Corps of Engineers (USACE), Kansas City District, installed and sampled three additional MWs (MW-6, MW-7, and MW-8) to fully delineate the hydrocarbon plume in groundwater. Benzene and naphthalene were detected in groundwater from MW-6 at concentrations of 460 μ g/L and 12 μ g/L, respectively; toluene, ethylbenzene, and xylenes were not detected. BTEX and naphthalene concentrations at MW-7 and MW-8 were measured at nondetect or near nondetect levels. During the July 1996 sampling effort, USACE personnel also collected a groundwater sample and a free product sample at MW-1S. BTEX concentrations of 210 μ g/L, 799 μ g/L, 417 μ g/L and 2,486 μ g/L, respectively, were detected in the groundwater sample. In the free product sample, relative BTEX concentrations of 246 mg/L, 1,631 mg/L, 1,239 mg/L, and 7,527 mg/L were detected.

Other than the occasional bailing of MW-1S and the initial fuel recovery from the trench, no remedial activities have been performed at the Building 4522 spill site (Chastain, 1997). Based on this condition, and compliance with other site selection criteria (reference October 1996 Work Plan, Section 2.1) the Building 4522 spill site represents a good candidate site for the fuel weathering study.

Proposed sampling locations for the Building 4522 Site are shown on Figure A.1. If possible, soil, groundwater, and free product sampling will be performed using boreholes and temporary monitoring points installed using a Geoprobe[®]. Specific sampling procedures are outlined in Section 3 of the work plan and will be performed in accordance with Seymour Johnson AFB requirements.

REFERENCES

- Chastain, Dean. 1997. 4th CES/CEV, Seymour Johnson Air Force Base, North Carolina. Telephone Conversation. January 24.
- Parsons ES. 1996. Final Comprehensive Site Assessment of Building 4522, Seymour Johnson Air Force Base, North Carolina. Cary. July.



APPENDIX B

SAMPLE ANALYTICAL DATA

Analytical Data Available Upon Request From:

Parsons Engineering Science, Inc. c/o Bruce Henry 1700 Broadway, Suite 900 Denver, CO 80290 **B-1**

NRMRL DATA

B-2

EAL DATA

APPENDIX C

CALCULATIONS

APPENDIX C-1

MOBILE LNAPL WEATHERING CALCULATIONS
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									Smith	n et al., 1981				Hug	hes et al., 1	984		
Lab	Fuel	Spill	Sample						Lin	ear	Expone	ntial		Lin	ear	Expor	ential	
Code	Туре	Date	Date	Locid Analyte	Results	Units	Mass Fraction	C ₀ - C	k	%Red./yr	k	%Red./yr	C ₀ - C	k	%Red./yr	k	%Red./yr	
EAL	JP-4	6/1/1994	#######	SH98-1610-2 Benzene	1300000	ug/L	0.1667	0.3333	0.0882	17.65	0.2908	25.23	0.3446	0.0912	17.84	0.2967	25.67	
EAL	JP-4	6/1/1994	#######	SH98-1610-2 Toluene	3000000	ug/L	0.3846	0.9454	0.2502	18.81	0.3284	27.99	1.3361	0.3536	20.55	0.3966	32.74	
EAL	JP-4	6/1/1994	#######	SH98-1610-2 Ethylbenzene	1400000	ug/L	0.1795	0.1905	0.0504	13.63	0.1915	17.43	0.5774	0.1528	20.19	0.3809	31.68	
EAL	JP-4	6/1/1994	#######	SH98-1610-2 o-Xylene	3100000	ug/L	0.3974	0.6126	0.1621	16.05	0.2469	21.88	0.2297	0.0608	9.70	0.1207	11.37	
EAL	JP-4	6/1/1994	#######	SH98-1610-2 m,p-Xylene	5900000	ug/L	0.7564	0.5536	0.1465	11.19	0.1454	13.53	0.9768	0.2585	14.92	0.2195	19.70	
EAL	JP-4	6/1/1994	#######	SH98-1610-2 Total Xylenes (m,p, and o)	9000000	ug/L	1.1538	1.1662	0.3087	13.30	0.1849	16.88	1.2065	0.3193	13.53	0.1894	17.26	
EAL	JP-4	6/1/1994	#######	SH98-1610-2 Total BTEX	14700000	ug/L	1.8846	2.6354	0.6975	15.43	0.2315	20.67	3.4645	0.9170	17.14	0.2761	24.13	
EAL	JP-4	6/1/1994	#######	SH98-1610-2 B+T	4300000	ug/L	0.5513											
EAL	JP-4	6/1/1994	#######	SH98-1610-2 E+X	10400000	ug/L	1.3333											
EAL	JP-4	6/1/1994	#######	SH98-1610-2 B/T	0.4333333333		0.0000											
EAL	JP-4	6/1/1994	#######	SH98-1610-2 B/E	0.928571429		0.0000											
EAL	JP-4	6/1/1994	#######	SH98-1610-2 B/X	0.144444444		0.0000											
EAL	JP-4	6/1/1994	#######	SH98-1610-2 T/E	2.142857143		0.0000											
EAL	JP-4	6/1/1994	#######	SH98-1610-2 T/X	0.3333333333		0.0000											
EAL	JP-4	6/1/1994	#######	SH98-1610-2 E/X	0.155555556		0.0000											
EAL	JP-4	6/1/1994	#######	SH98-1610-2 (B+T)/(E+X)	0.413461538		0.0000											
EAL	JP-4	6/1/1994	3/6/1997	SHMW1610-2 Benzene	2650000	ug/L	0.3464	0.1536	0.0556	11.11	0.1328	12.43	0.1648	0.0596	11.66	0.1408	13.13	
EAL	JP-4	6/1/1994	3/6/1997	SHMW1610-2 Toluene	5740000	ug/L	0.7503	0.5797	0.2097	15.77	0.2071	18.70	0.9704	0.3510	20.40	0.3002	25.94	
EAL	JP-4	6/1/1994	3/6/1997	SHMW1610-2 Ethylbenzene	1770000	ug/L	0.2314	0.1386	0.0501	13.55	0.1698	15.62	0.5255	0.1901	25.12	0.4287	34.87	
EAL	JP-4	6/1/1994	3/6/1997	SHMW1610-2 o-Xylene	3490000	ug/L	0.4562	0.5538	0.2003	19.83	0.2875	24.99	0.1710	0.0618	9.86	0.1151	10.88	
EAL	JP-4	6/1/1994	3/6/1997	SHMW1610-2 m,p-Xylene	7490000	ug/L	0.9791	0.3309	0.1197	9.14	0.1053	10.00	0.7541	0.2728	15.74	0.2066	18.66	
EAL	JP-4	6/1/1994	3/6/1997	SHMW1610-2 Total Xylenes (m,p, and o)	10980000	ug/L	1.4353	0.8847	0.3200	13.79	0.1737	15.95	0.9251	0.3346	14.18	0.1799	16.47	
EAL	JP-4	6/1/1994	3/6/1997	SHMW1610-2 Total BTEX	21140000	ug/L	2.7634	1.7566	0.6354	14.06	0.1780	16.31	2.5857	0.9354	17.49	0.2389	21.25	
EAL	JP-4	6/1/1994	3/6/1997	SHMW1610-2 B+T	8390000	ug/L	1.0967											
EAL	JP-4	6/1/1994	3/6/1997	SHMW1610-2 E+X	12750000	ug/L	1.6667											
EAL	JP-4	6/1/1994	3/6/1997	SHMW1610-2 B/T	0.461672474		0.0000											
EAL	JP-4	6/1/1994	3/6/1997	SHMW1610-2 B/E	1.497175141		0.0000											
EAL	JP-4	6/1/1994	3/6/1997	SHMW1610-2 B/X	0.241347905		0.0000											
EAL	JP-4	6/1/1994	3/6/1997	SHMW1610-2 T/E	3.242937853		0.0000											
EAL	JP-4	6/1/1994	3/6/1997	SHMW1610-2 T/X	0.52276867		0.0000											
EAL	JP-4	6/1/1994	3/6/1997	SHMW1610-2 E/X	0.161202186		0.0000											
EAL	JP-4	6/1/1994	3/6/1997	SHMW1610-2 (B+T)/(E+X)	0.658039216		0.0000											

Shaw

										Smith	n et al., 1981				Hug	hes et al., 1	984	
Lab	Fuel	Spill	Sample							Lin	ear	Expone	ntial		Lin	ear	Expo	nential
Code	Type	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	C C	k	%Red./yr	k.	%Red./yr	C ₀ - C	k	%Red./yr	k.	%Red./yr
	2.													ů				
NRMRL	JP-4	6/1/1994	#######	SH98-1610-2	Density	0.78	g/mL	0.0001										
NRMRL	JP-4	6/1/1994	#######	SH98-1610-2	Benzene	1250	ug/mL	0.1603	0.3397	0.0899	17.98	0.3012	26.00	0.3510	0.0929	18.17	0.3070	26.44
NRMRL	JP-4	6/1/1994	#######	SH98-1610-2	Toluene	2830	ug/mL	0.3628	0.9672	0.2560	19.25	0.3438	29.10	1.3579	0.3594	20.89	0.4120	33.77
NRMRL	JP-4	6/1/1994	#######	SH98-1610-2	Ethylbenzene	1040	ug/mL	0.1333	0.2367	0.0626	16.93	0.2702	23.67	0.6235	0.1650	21.81	0.4596	36.84
NRMRL	JP-4	6/1/1994	#######	SH98-1610-2	m-Xylene	3440	ug/mL	0.4410	0.5190	0.1374	14.31	0.2059	18.61	-0.4410	-0.1167	#DIV/0!	#DIV/0!	#DIV/0!
NRMRL	JP-4	6/1/1994	#######	SH98-1610-2	o-Xylene	2430	ug/mL	0.3115	0.6985	0.1849	18.30	0.3113	26.75	0.3156	0.0835	13.32	0.1852	16.91
NRMRL	JP-4	6/1/1994	#######	SH98-1610-2	p-Xylene	1310	ug/mL	0.1679	0.1821	0.0482	13.77	0.1944	17.66	1.5652	0.4143	23.90	0.6178	46.09
NRMRL	JP-4	6/1/1994	#######	SH98-1610-2	m,p-xylenes	4750	ug/mL	0.6090	0.7010	0.1856	14.16	0.2028	18.35	1.1242	0.2976	17.17	0.2768	24.18
NRMRL	JP-4	6/1/1994	#######	SH98-1610-2	Total Xylenes (m,p, and o)	7180	ug/mL	0.9205	1.3995	0.3704	15.97	0.2447	21.70	1.4398	0.3811	16.15	0.2492	22.06
NRMRL	JP-4	6/1/1994	#######	SH98-1610-2	Total BTEX	12300	ug/mL	1.5769	2.9431	0.7790	17.23	0.2787	24.33	3.7722	0.9984	18.67	0.3233	27.62
NRMRL	JP-4	6/1/1994	#######	SH98-1610-2	B+T	4080	ug/mL	0.5231										
NRMRL	JP-4	6/1/1994	#######	SH98-1610-2	E+X	8220	ug/mL	1.0538										
NRMRL	JP-4	6/1/1994	########	SH98-1610-2	B/T	0.441696113		0.0001										
NRMRL	JP-4	6/1/1994	#######	SH98-1610-2	B/E	1.201923077		0.0002										
NRMRL	JP-4	6/1/1994	########	SH98-1610-2	B/X	0.174094708		0.0000										
NRIVIRL	JP-4	6/1/1994	########	SH98-1610-2	T/E	2.721153846		0.0003										
NEWEL	JP-4	6/1/1994	*****	SH96-1610-2		0.394150416		0.0001										
NEME	JP-4	6/1/1994	########	SH90-1010-2		0.144646797		0.0000										
NDMDI	JF -4	6/1/1994	######################################	SH90-1010-2		0.490300303	ua/ml	0.0001	0.4012	0 1062	#DIV//01	#DIV/01	#DIV//01	0.0499	0.0120	2 97	0.0204	2.00
NEMEL	JF-4	6/1/1994	########	SH08-1610-2	1.2.4-Trimethylbenzene	7320	ug/mL	0.4013	-0.4013	0.1002	#DIV/0:	#DIV/0:	#DIV/0:	-0.1218	-0.0322	-3.05	-0.0368	-3.75
NRMRI	IP-4	6/1/1994	#######################################	SH98-1610-2	1 3 5-Trimethylbenzene	2200	ug/mL	0.3303	0.0713	0.0103	8.69	0.0154	10.00	-0.1210	-0.0322	-34.81	-0.0000	-24.88
NRMRI	JP-4	6/1/1994	########	SH98-1610-2	Total TMBs	12650	ug/mL	1 6218	-0 1918	-0.0508	-3.55	-0.0333	-3.39	-0 2331	-0.0617	-4 44	-0.0411	-4 19
NRMRI	JP-4	6/1/1994	########	SH98-1610-2	Naphthalene	1320	ug/ml	0 1692	0.3308	0.0875	17.51	0.2867	24.93	0.1051	0.0278	10.14	0 1278	12.00
NRMRL	JP-4	6/1/1994	########	SH98-1610-2	1-MethylNaphthalene	1320	ug/mL	0.1692	0.6108	0.1617	20.73	0.4044	33.27	-0.1331	-0.0352	-97.41	-0.4085	-50.45
NRMRL	JP-4	6/1/1994	#######	SH98-1610-2	2-MethylNaphthalene	2200	ua/mL	0.2821	0.2779	0.0736	13.14	0.1815	16.60	-0.1237	-0.0327	-20.68	-0.1528	-16.51
NRMRL	JP-4	6/1/1994	#######	SH98-1610-2	Total Naphthalenes	4840	uq/mL	0.6205	1.2195	0.3228	17.54	0.2877	25.00	-0.1517	-0.0401	-8.56	-0.0742	-7.70
							Ŭ											
NRMRL	JP-4	6/1/1994	#######	SH98-1610-3	Density	0.777	g/mL	0.0001										
NRMRL	JP-4	6/1/1994	#######	SH98-1610-3	Benzene	1650	ug/mL	0.2124	0.2876	0.0761	15.23	0.2267	20.28	0.2989	0.0791	15.47	0.2325	20.75
NRMRL	JP-4	6/1/1994	#######	SH98-1610-3	Toluene	3290	ug/mL	0.4234	0.9066	0.2400	18.04	0.3029	26.14	1.2973	0.3434	19.96	0.3711	31.00
NRMRL	JP-4	6/1/1994	#######	SH98-1610-3	Ethylbenzene	1070	ug/mL	0.1377	0.2323	0.0615	16.62	0.2616	23.02	0.6191	0.1639	21.65	0.4510	36.30
NRMRL	JP-4	6/1/1994	#######	SH98-1610-3	m-Xylene	3470	ug/mL	0.4466	0.5134	0.1359	14.16	0.2026	18.34	-0.4466	-0.1182	#DIV/0!	#DIV/0!	#DIV/0!
NRMRL	JP-4	6/1/1994	#######	SH98-1610-3	o-Xylene	2450	ug/mL	0.3153	0.6947	0.1839	18.21	0.3081	26.52	0.3119	0.0825	13.16	0.1820	16.64
NRMRL	JP-4	6/1/1994	#######	SH98-1610-3	p-Xylene	1290	ug/mL	0.1660	0.1840	0.0487	13.91	0.1974	17.91	1.5671	0.4148	23.93	0.6208	46.25
NRMRL	JP-4	6/1/1994	#######	SH98-1610-3	m,p-xylenes	4760	ug/mL	0.6126	0.6974	0.1846	14.09	0.2012	18.22	1.1206	0.2966	17.11	0.2753	24.06
NRMRL	JP-4	6/1/1994	#########	SH98-1610-3	Total Xylenes (m,p, and o)	7210	ug/mL	0.9279	1.3921	0.3685	15.88	0.2425	21.54	1.4324	0.3791	16.06	0.2471	21.89
NRMRL	JP-4	6/1/1994	#########	SH98-1610-3	I OTAL BIEX	13220	ug/mL	1./014	2.8186	0.7460	16.51	0.2586	22.79	3.6477	0.9655	18.05	0.3032	26.15
	JP-4	6/1/1994	#########	SH98-1610-3		4940	ug/mL	0.6358										
NDMD	JP-4	6/1/1994	######################################	SLID0 1610 2	РЛ	0280	ug/mL	0.0004										
NRMDI	JP-4	6/1/1994	######################################	SH08-1610-3	B/F	1 542056075		0.0001										
NRMRI	JP-4	6/1/1994	######################################	SH98-1610-3	B/L B/X	0.228848921	1	0.0002										
NRMRI	IP-4	6/1/1994	#######################################	SH98-1610-3	T/F	3 074766355		0.0000										
NRMRI	IP-4	6/1/1994	#######################################	SH98-1610-3	T/X	0.45631068		0.0004										
NRMRI	JP-4	6/1/1994	########	SH98-1610-3	E/X	0.148404993		0.0000										
NRMRL	JP-4	6/1/1994	#########	SH98-1610-3	(B+T)/(E+X)	0.596618357		0.0001										
NRMRL	JP-4	6/1/1994	########	SH98-1610-3	1.2.3-Trimethylbenzene	2860	ua/mL	0.3681	-0.3681	-0.0974	#DIV/0!	#DIV/0!	#DIV/0!	0.0820	0.0217	4,82	0.0533	5.19
NRMRL	JP-4	6/1/1994	#######	SH98-1610-3	1,2,4-Trimethylbenzene	6690	ug/mL	0.8610	0.1490	0.0394	3.90	0.0422	4.14	-0.0443	-0.0117	-1.44	-0.0140	-1.41
NRMRL	JP-4	6/1/1994	########	SH98-1610-3	1,3,5-Trimethylbenzene	2010	ug/mL	0.2587	0.1613	0.0427	10.17	0.1283	12.04	-0.1369	-0.0362	-29.74	-0.1993	-22.06
NRMRL	JP-4	6/1/1994	#######	SH98-1610-3	Total TMBs	11560	ua/mL	1,4878	-0.0578	-0.0153	-1.07	-0.0105	-1.05	-0.0991	-0.0262	-1.89	-0.0182	-1.84
NRMRL	JP-4	6/1/1994	#######	SH98-1610-3	Naphthalene	1230	ug/mL	0.1583	0.3417	0.0904	18.09	0.3044	26.24	0.1160	0.0307	11.19	0.1455	13.54
NRMRL	JP-4	6/1/1994	#######	SH98-1610-3	1-MethylNaphthalene	1200	ug/mL	0.1544	0.6256	0.1656	21.23	0.4287	34.86	-0.1183	-0.0313	-86.58	-0.3843	-46.86
NRMRL	JP-4	6/1/1994	#######	SH98-1610-3	2-MethylNaphthalene	1980	ug/mL	0.2548	0.3052	0.0808	14.42	0.2084	18.81	-0.0965	-0.0255	-16.13	-0.1259	-13.42
NRMRL	JP-4	6/1/1994	#######	SH98-1610-3	Total Naphthalenes	4410	ug/mL	0.5676	1.2724	0.3368	18.30	0.3113	26.75	-0.0987	-0.0261	-5.57	-0.0506	-5.19
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										Smith	h et al., 1981				Hug	hes et al., 1	1984	
Lab	Fuel	Spill	Sample							Lin	ear	Expone	ntial		Lin	ear	Expo	nential
Code	Type	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	0.0	k	%Red /vr	p	%Red /vr	0.0	k	%Red /vr	k	%Red /vr
NIDMDI		6/1/1004	2/6/1007		Dongity	0.765	a/ml	0.0001	0, 0	ĸ	/inteda/yi	ĸ	/orcea./yr	0, 0	ĸ	/orteal/yr	ĸ	/orccu/yr
INRIVIRL	JP-4	0/1/1994	3/0/1997	SHIVIV 1010-2	Density	0.765	g/m∟	0.0001										
	10.4	0/1/1001	0/0/4007	0111111110100	2	0050		0.00.44	0.0050	0.0745	44.00	0.1000	17.17	0.0474	0.0705	45.00		10.10
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	Benzene	2250	ug/mL	0.2941	0.2059	0.0745	14.90	0.1920	17.47	0.2171	0.0785	15.36	0.2000	18.13
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	Toluene	4890	ug/mL	0.6392	0.6908	0.2499	18.79	0.2650	23.28	1.0815	0.3912	22.74	0.3582	30.11
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	Ethylbenzene	1340	ug/mL	0.1752	0.1948	0.0705	19.05	0.2705	23.70	0.5817	0.2104	27.80	0.5294	41.10
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	m-Xylene	4140	ug/mL	0.5412	0.4188	0.1515	15.78	0.2073	18.73	-0.5412	-0.1958	#DIV/0!	#DIV/0!	#DIV/0!
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	o-Xylene	2950	ug/mL	0.3856	0.6244	0.2259	22.36	0.3483	29.41	0.2416	0.0874	13.93	0.1759	16.13
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	p-Xylene	1440	ug/mL	0.1882	0.1618	0.0585	16.72	0.2244	20.10	1.5449	0.5589	32.25	0.8031	55.21
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	m,p-xylenes	5580	ug/mL	0.7294	0.5806	0.2100	16.03	0.2118	19.09	1.0038	0.3631	20.95	0.3131	26.88
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	Total Xylenes (m,p, and o)	8530	ug/mL	1.1150	1.2050	0.4359	18.79	0.2650	23.28	1.2453	0.4505	19.09	0.2713	23.76
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	Total BTEX	17010	uq/mL	2.2235	2.2965	0.8307	18.38	0.2566	22.63	3.1256	1.1307	21.14	0.3176	27.21
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	B+T	7140	uq/mL	0.9333										
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	E+X	9870	ua/mL	1.2902										
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	B/T	0.460122699		0.0001										
NRMRI	JP-4	6/1/1994	3/6/1997	SHMW1610-2	B/F	1 679104478		0.0002										
NRMRI	JP-4	6/1/1994	3/6/1997	SHMW1610-2	B/X	0.263774912		0.0000										
NRMRI	IP-4	6/1/1994	3/6/1997	SHMW1610-2	T/F	3 649253731		0.0005										
NRMRI	JP-4	6/1/1994	3/6/1997	SHMW1610-2	T/X	0.573270809		0.0001								1	1	
NRMPI	IP _r 4	6/1/1004	3/6/1007	SHMW1610.2	F/X	0 157002614		0.0001								1	1	
NIDMD		6/1/1004	2/6/1007	SHWW1610 2		0.722404255	<u> </u>	0.0000										
NDMD	JP-4	6/1/1994	3/0/1997	SHIVIW 1010-2	(DTI)/(ETA)	0.123404255	ua/m!	0.0001	0 2000	0.1414	#DIV//0/	#DIV//01	#DIV//01	0.0502	0.0214	170	0.0511	4.09
NRMRL	JP-4	6/1/1994	3/6/1997	SHIWW1610-2	1,2,3-Trimethylbenzene	2990	ug/mL	0.3908	-0.3908	-0.1414	#DIV/0!	#DIV/0!	#DIV/0!	0.0593	0.0214	4.76	0.0511	4.98
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	1,2,4-1 rimethylbenzene	5790	ug/mL	0.7569	0.2531	0.0916	9.07	0.1044	9.91	0.0598	0.0216	2.65	0.0275	2.72
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	1,3,5-Trimethylbenzene	3450	ug/mL	0.4510	-0.0310	-0.0112	-2.67	-0.0257	-2.61	-0.3292	-0.1191	-97.74	-0.4735	-60.56
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	Total TMBs	12230	ug/mL	1.5987	-0.1687	-0.0610	-4.27	-0.0403	-4.12	-0.2100	-0.0760	-5.47	-0.0510	-5.23
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	Naphthalene	1120	ug/mL	0.1464	0.3536	0.1279	25.58	0.4443	35.87	0.1279	0.0463	16.87	0.2271	20.32
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	1-MethylNaphthalene	1040	ug/mL	0.1359	0.6441	0.2330	29.87	0.6320	46.85	-0.0998	-0.0361	-99.83	-0.4791	-61.46
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	2-MethylNaphthalene	1680	ug/mL	0.2196	0.3404	0.1231	21.99	0.3386	28.73	-0.0613	-0.0222	-13.99	-0.1183	-12.56
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-2	Total Naphthalenes	3840	ug/mL	0.5020	1.3380	0.4840	26.31	0.4699	37.49	-0.0331	-0.0120	-2.56	-0.0247	-2.50
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-3	Density	0.783	g/mL	0.0001										
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-3	Benzene	1380	uq/mL	0.1762	0.3238	0.1171	23.42	0.3772	31.42	0.3350	0.1212	23.70	0.3852	31.97
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-3	Toluene	2510	ua/mL	0.3206	1.0094	0.3652	27.46	0.5147	40.23	1.4001	0.5065	29.44	0.6079	45.55
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-3	Ethylbenzene	935	ug/mL	0.1194	0.2506	0.0906	24.50	0.4091	33.58	0.6374	0.2306	30.47	0.6680	48.73
NRMRI	JP-4	6/1/1994	3/6/1997	SHMW1610-3	m-Xylene	3240	ug/ml	0 4138	0.5462	0 1976	20.58	0.3044	26.25	-0 4138	-0 1497	#DIV/01	#DIV/0!	#DIV/0!
NEMEL	ID_4	6/1/1004	3/6/1007	SHMW1610-3	o-Yvlene	2390	ug/mL	0.4100	0.0402	0.1570	25.00	0.0044	35.14	0.3210	0.1407	18.57	0 2605	22.03
NEMEL	ID-4	6/1/1994	3/6/1007	SHMW1610-3	n-Xylene	1100	ug/mL	0.3032	0.7040	0.2343	20.24	0.4023	26.05	1 5812	0.1103	33.00	0.2005	58.54
NRMRI	ID_4	6/1/1004	3/6/1007	SHMW1610 2	m n-yulenes	1190	ug/mL	0.1520	0.1900	0.0710	20.47	0.0010	20.00	1 1674	0.3720	24 27	0.0000	33.30
NRMDI	JF -4	6/1/1004	3/6/1007	SHMW/1610-3	Total Xylenes (m.n. and a)	4430	ug/mL	0.0000	1 4400	0.2092	20.00	0.3037	20.19	1 /000	0.4223	24.37	0.4000	30.28
NDMD		6/1/1994	3/0/1997	CUMM/4640.0	Total Aylenes (III,p, and 0)	14045	ug/mL	0.0710	1.4490	0.5242	22.39	0.3544	29.04	1.4093	1 2070	22.03	0.3000	30.20
NRIVIRL	JP-4	0/1/1994	3/0/1997	STIVIV 1010-3		11645	ug/mL	1.4872	3.0328	1.0971	24.27	0.4021	33.11	3.8019	1.3970	20.12	0.4630	37.00
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-3	B+I	3890	ug/mL	0.4968										
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-3	E+X	7755	ug/mL	0.9904										
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-3	B/I	0.549800797		0.0001										
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-3	B/E	1.475935829		0.0002										
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-3	B/X	0.202346041		0.0000										
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-3	T/E	2.684491979		0.0003										
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-3	T/X	0.368035191		0.0000										
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-3	E/X	0.137096774		0.0000							-			
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-3	(B+T)/(E+X)	0.501611863		0.0001										
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-3	1,2,3-Trimethylbenzene	3780	ug/mL	0.4828	-0.4828	-0.1746	#DIV/0!	#DIV/0!	#DIV/0!	-0.0326	-0.0118	-2.62	-0.0253	-2.56
NRMRL	JP-4	6/1/1994	3/6/1997	SHMW1610-3	1.2.4-Trimethylbenzene	7280	ua/mL	0,9298	0,0802	0.0290	2,87	0.0299	2,95	-0.1130	-0.0409	-5.01	-0.0469	-4.80
NRMRI	JP-4	6/1/1994	3/6/1997	SHMW1610-3	1.3.5-Trimethylbenzene	4110	ua/ml	0.5249	-0.1049	-0.0379	-9.04	-0.0807	-8.40	-0,4031	-0,1458	-119.70	-0.5284	-69.62
NRMRI	IP-4	6/1/1994	3/6/1997	SHMW1610-3	Total TMBs	15170	ug/ml	1 9374	-0 5074	-0 1836	-12 84	-0 1099	-11 61	-0.5488	-0 1985	-14 30	-0 1205	-12.80
NEME	IP-4	6/1/1004	3/6/1007	SHMW1610.2	Nanhthalene	1//0	ug/ml	0.1920	0.0074	0.1142	22.04	0.1035	30.36	0.0400	0.1303	11.00	0.1200	13.47
NEME	ID_4	6/1/1004	3/6/1007	SHMW/1610 2	1-MethylNanhthalene	1/100	ug/mL	0.1009	0.5101	0.1143	22.07	0.0010	40.12	-0.1520	-0.0527	-152.02	-0 5002	-81.90
NDMD		6/1/1004	2/6/1007	ST IVIV 1010-3	2 MothylNophthologo	1400	ug/mL	0.1090	0.0910	0.2130	45.60	0.0120	40.12	0.1529	-0.0000	-102.92	-0.0903	-01.30
NRIVIRL	JP-4	6/1/1994	3/0/1997	SHIWW1010-3	Z-Ivieu iyinaphthalene	2490	ug/mL	0.3180	0.2420	0.0875	15.63	0.2047	18.51	-0.1597	-0.0578	-30.47	-0.2522	-28.09
INKINKL	JP-4	0/1/1994	3/0/1997	SHIVIV 1610-3	i otal inapritrialenes	5410	ug/mL	0.6909	1.1491	0.4157	22.59	0.3543	29.84	-0.2221	-0.0803	-17.14	-0.1403	-15.00

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									Sn	nith et al., 198	31			Hug	hes et al., 1	984	
Lab	Fuel	Spill	Sample						Line	ear	Exponer	ntial		Lin	ear	Expor	nential
Code	Туре	Date	Date	Locid	Analyte	Results Units	Mass Fraction	C ₀ - C	k	%Red./yr	k	%Red./yr	C _o - C	k	%Red./yr	k	%Red./yr
EAL	JP-4	#####	3/4/1997	MBMW8I	Benzene	211000 ug/L	0.0281	0.4719	0.0292	5.83	0.1778	16.29	0.4831	0.0299	5.84	0.1792	16.41
EAL	JP-4	#####	3/4/1997	MBMW8I	Toluene	7540 ug/L	0.0010	1.3290	0.0821	6.18	0.4442	35.87	1.7197	0.1063	6.18	0.4601	36.88
EAL	JP-4	#####	3/4/1997	MBMW8I	Ethylbenzene	1830000 ug/L	0.2440	0.1260	0.0078	2.10	0.0257	2.54	0.5129	0.0317	4.19	0.0700	6.76
EAL	JP-4	#####	3/4/1997	MBMW8I	o-Xylene	153000 ug/L	0.0204	0.9896	0.0612	6.06	0.2412	21.43	0.6068	0.0375	5.98	0.2117	19.08
EAL	JP-4	#####	3/4/1997	MBMW8I	m,p-Xylene	6210000 ug/L	0.8280	0.4820	0.0298	2.27	0.0284	2.80	0.9052	0.0559	3.23	0.0457	4.46
EAL	JP-4	#####	3/4/1997	MBMW8I	Total Xylenes (m,p, and o)	6363000 ug/L	0.8484	1.4716	0.0909	3.92	0.0622	6.03	1.5119	0.0934	3.96	0.0632	6.13
EAL	JP-4	#####	3/4/1997	MBMW8I	Total BTEX	8411540 ug/L	1.1215	3.3985	0.2100	4.65	0.0861	8.25	4.2276	0.2613	4.88	0.0965	9.20
EAL	JP-4	#####	3/4/1997	MBMW8I	B+T	218540 ug/L	0.0291										
EAL	JP-4	#####	3/4/1997	MBMW8I	E+X	8193000 ug/L	1.0924										
EAL	JP-4	#####	3/4/1997	MBMW8I	B/T	27.98408488	0.0000										
EAL	JP-4	#####	3/4/1997	MBMW8I	B/E	0.115300546	0.0000										
EAL	JP-4	#####	3/4/1997	MBMW8I	B/X	0.033160459	0.0000										
EAL	JP-4	#####	3/4/1997	MBMW8I	T/E	0.004120219	0.0000										
EAL	JP-4	#####	3/4/1997	MBMW8I	T/X	0.001184976	0.0000										
EAL	JP-4	#####	3/4/1997	MBMW8I	E/X	0.287600189	0.0000										
EAL	JP-4	#####	3/4/1997	MBMW8I	(B+T)/(E+X)	0.02667399	0.0000										
NRMRL	JP-4	#####	3/4/1997	MBMW8I	Density	0.75 g/mL	0.0001										
NRMRL	JP-4	#####	3/4/1997	MBMW8I	Benzene	211 ug/mL	0.0281	0.4719	0.0292	5.83	0.1778	16.29	0.4831	0.0299	5.84	0.1792	16.41
NRMRL	JP-4	#####	3/4/1997	MBMW8I	Toluene	7.54 ug/mL	0.0010	1.3290	0.0821	6.18	0.4442	35.87	1.7197	0.1063	6.18	0.4601	36.88
NRMRL	JP-4	#####	3/4/1997	MBMW8I	Ethylbenzene	1360 ug/mL	0.1813	0.1887	0.0117	3.15	0.0441	4.31	0.5755	0.0356	4.70	0.0883	8.45
NRMRL	JP-4	#####	3/4/1997	MBMW8I	m-Xylene	3440 ug/mL	0.4587	0.5013	0.0310	3.23	0.0456	4.46	-0.4587	-0.0283	#DIV/0!	#DIV/0!	#DIV/0!
NRMRL	JP-4	#####	3/4/1997	MBMW8I	o-Xylene	12.1 ug/mL	0.0016	1.0084	0.0623	6.17	0.3980	32.83	0.6256	0.0387	6.16	0.3685	30.82
NRMRL	JP-4	#####	3/4/1997	MBMW8I	p-Xylene	810 ug/mL	0.1080	0.2420	0.0150	4.27	0.0727	7.01	1.6252	0.1004	5.80	0.1715	15.76
NRMRL	JP-4	#####	3/4/1997	MBMW8I	m,p-xylenes	4250 ug/mL	0.5667	0.7433	0.0459	3.51	0.0518	5.05	1.1665	0.0721	4.16	0.0691	6.68
NRMRL	JP-4	#####	3/4/1997	MBMW8I	Total Xylenes (m,p, and o)	4262.1 ug/mL	0.5683	1.7517	0.1083	4.67	0.0869	8.33	1.7921	0.1108	4.69	0.0880	8.42
NRMRL	JP-4	#####	3/4/1997	MBMW8I	Total BTEX	5840.64 ug/mL	0.7788	3.7412	0.2312	5.12	0.1087	10.30	4.5704	0.2825	5.28	0.1191	11.23
NRMRL	JP-4	#####	3/4/1997	MBMW8I	B+T	218.54 ug/mL	0.0291										
NRMRL	JP-4	#####	3/4/1997	MBMW8I	E+X	5622.1 ug/mL	0.7496										
NRMRL	JP-4	#####	3/4/1997	MBMW8I	B/T	27.98408488	0.0037										
NRMRL	JP-4	#####	3/4/1997	MBMW8I	B/E	0.155147059	0.0000										
NRMRL	JP-4	#####	3/4/1997	MBMW8I	B/X	0.049506112	0.0000										
NRMRL	JP-4	#####	3/4/1997	MBMW8I	T/E	0.005544118	0.0000										
NRMRL	JP-4	#####	3/4/1997	MBMW8I	T/X	0.001769081	0.0000										
NRMRL	JP-4	#####	3/4/1997	MBMW8I	E/X	0.319091528	0.0000										
NRMRL	JP-4	#####	3/4/1997	MBMW8I	(B+T)/(E+X)	0.038871596	0.0000										
NRMRL	JP-4	#####	3/4/1997	MBMW8I	1,2,3-Trimethylbenzene	2580 ug/mL	0.3440	-0.3440	-0.0213	#DIV/0!	#DIV/0!	#DIV/0!	0.1061	0.0066	1.46	0.0166	1.65
NRMRL	JP-4	#####	3/4/1997	MBMW8I	1,2,4-Trimethylbenzene	6160 ug/mL	0.8213	0.1887	0.0117	1.15	0.0128	1.27	-0.0046	-0.0003	-0.03	-0.0003	-0.03
NRMRL	JP-4	#####	3/4/1997	MBMW8I	1,3,5-Trimethylbenzene	3550 ug/mL	0.4733	-0.0533	-0.0033	-0.78	-0.0074	-0.74	-0.3515	-0.0217	-17.83	-0.0839	-8.75
NRMRL	JP-4	#####	3/4/1997	MBMW8I	Total TMBs	12290 ug/mL	1.6387	-0.2087	-0.0129	-0.90	-0.0084	-0.85	-0.2500	-0.0155	-1.11	-0.0102	-1.03
NRMRL	JP-4	#####	3/4/1997	MBMW8I	Naphthalene	1160 ug/mL	0.1547	0.3453	0.0213	4.27	0.0725	6.99	0.1196	0.0074	2.70	0.0354	3.48
NRMRL	JP-4	#####	3/4/1997	MBMW8I	1-MethylNaphthalene	1300 ug/mL	0.1733	0.6067	0.0375	4.81	0.0930	8.88	-0.1372	-0.0085	-23.44	-0.0969	-10.17
NRMRL	JP-4	#####	3/4/1997	MBMW8I	2-MethylNaphthalene	1910 ug/mL	0.2547	0.3053	0.0189	3.37	0.0487	4.75	-0.0963	-0.0060	-3.76	-0.0294	-2.98
NRMRL	JP-4	#####	3/4/1997	MBMW8I	Total Naphthalenes	4370 ug/mL	0.5827	1.2573	0.0777	4.22	0.0711	6.86	-0.1138	-0.0070	-1.50	-0.0134	-1.35
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										Si	nith et al., 19	81			Hug	hes et al., 1	984	
Lab	Fuel	Spill	Sample							Lin	ear	Expone	ntial		Lin	ear	Expor	nential
Code	Туре	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	C ₀ - C	k	%Red./yr	k	%Red./yr	C ₀ - C	k	%Red./yr	k	%Red./yr
NRMRL	JP-4	#####	3/4/1997	MBMW24	Density	0.764	g/mL	0.0001										
NRMRL	JP-4	#####	3/4/1997	MBMW24	Benzene	56.2	ug/mL	0.0074	0.4926	0.0304	6.09	0.2607	22.95	0.5039	0.0311	6.09	0.2621	23.06
NRMRL	JP-4	#####	3/4/1997	MBMW24	Toluene	5.57	ug/mL	0.0007	1.3293	0.0822	6.18	0.4641	37.13	1.7200	0.1063	6.18	0.4800	38.12
NRMRL	JP-4	#####	3/4/1997	MBMW24	Ethylbenzene	600	ug/mL	0.0785	0.2915	0.0180	4.87	0.0958	9.13	0.6783	0.0419	5.54	0.1400	13.07
NRMRL	JP-4	#####	3/4/1997	MBMW24	m-Xylene	1810	ug/mL	0.2369	0.7231	0.0447	4.66	0.0865	8.28	-0.2369	-0.0146	#DIV/0!	#DIV/0!	#DIV/0!
NRMRL	JP-4	#####	3/4/1997	MBMW24	o-Xylene	14.2	ug/mL	0.0019	1.0081	0.0623	6.17	0.3892	32.24	0.6253	0.0386	6.16	0.3598	30.22
NRMRL	JP-4	#####	3/4/1997	MBMW24	p-Xylene	313	ug/mL	0.0410	0.3090	0.0191	5.46	0.1326	12.42	1.6922	0.1046	6.03	0.2314	20.66
NRMRL	JP-4	#####	3/4/1997	MBMW24	m,p-xylenes	2123	ug/mL	0.2779	1.0321	0.0638	4.87	0.0958	9.14	1.4553	0.0899	5.19	0.1131	10.70
NRMRL	JP-4	#####	3/4/1997	MBMW24	Total Xylenes (m,p, and o)	2137.2	ug/mL	0.2797	2.0403	0.1261	5.43	0.1307	12.26	2.0806	0.1286	5.45	0.1318	12.35
NRMRL	JP-4	#####	3/4/1997	MBMW24	Total BTEX	2798.97	ug/mL	0.3664	4.1536	0.2567	5.68	0.1553	14.38	4.9828	0.3079	5.76	0.1657	15.27
NRMRL	JP-4	#####	3/4/1997	MBMW24	B+T	61.77	ug/mL	0.0081										
NRMRL	JP-4	#####	3/4/1997	MBMW24	E+X	2737.2	ug/mL	0.3583										
NRMRL	JP-4	#####	3/4/1997	MBMW24	B/T	10.08976661		0.0013										
NRMRL	JP-4	#####	3/4/1997	MBMW24	B/E	0.093666667		0.0000										
NRMRL	JP-4	#####	3/4/1997	MBMW24	B/X	0.026296088		0.0000										
NRMRL	JP-4	#####	3/4/1997	MBMW24	T/E	0.009283333		0.0000										
NRMRL	JP-4	#####	3/4/1997	MBMW24	T/X	0.002606214		0.0000										
NRMRL	JP-4	#####	3/4/1997	MBMW24	E/X	0.280741157		0.0000										
NRMRL	JP-4	#####	3/4/1997	MBMW24	(B+T)/(E+X)	0.022566857		0.0000										
NRMRL	JP-4	#####	3/4/1997	MBMW24	1,2,3-Trimethylbenzene	2980	ug/mL	0.3901	-0.3901	-0.0241	#DIV/0!	#DIV/0!	#DIV/0!	0.0601	0.0037	0.82	0.0089	0.88
NRMRL	JP-4	#####	3/4/1997	MBMW24	1,2,4-Trimethylbenzene	6500	ug/mL	0.8508	0.1592	0.0098	0.97	0.0106	1.05	-0.0341	-0.0021	-0.26	-0.0025	-0.25
NRMRL	JP-4	#####	3/4/1997	MBMW24	1,3,5-Trimethylbenzene	2840	ug/mL	0.3717	0.0483	0.0030	0.71	0.0075	0.75	-0.2499	-0.0154	-12.68	-0.0689	-7.14
NRMRL	JP-4	#####	3/4/1997	MBMW24	Total TMBs	12320	ug/mL	1.6126	-0.1826	-0.0113	-0.79	-0.0074	-0.75	-0.2239	-0.0138	-1.00	-0.0092	-0.93
NRMRL	JP-4	#####	3/4/1997	MBMW24	Naphthalene	892	ug/mL	0.1168	0.3832	0.0237	4.74	0.0899	8.60	0.1576	0.0097	3.55	0.0528	5.14
NRMRL	JP-4	#####	3/4/1997	MBMW24	1-MethylNaphthalene	1320	ug/mL	0.1728	0.6072	0.0375	4.81	0.0932	8.89	-0.1366	-0.0084	-23.35	-0.0967	-10.15
NRMRL	JP-4	#####	3/4/1997	MBMW24	2-MethylNaphthalene	1810	ug/mL	0.2369	0.3231	0.0200	3.57	0.0532	5.18	-0.0786	-0.0049	-3.07	-0.0249	-2.52
NRMRL	JP-4	#####	3/4/1997	MBMW24	Total Naphthalenes	4022	ug/mL	0.5264	1.3136	0.0812	4.41	0.0773	7.44	-0.0576	-0.0036	-0.76	-0.0072	-0.72

										Sm	ith et al., 1981				Hug	ghes et al.,	1984		
Lab	Fuel	Spill	Sample							Lin	ear	Expon	ential		Lin	ear	Expone	ntial	
Code	Туре	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	C ₀ - C	k	%Red./yr	k	%Red./yr	C ₀ - C	k	%Red./yr	k	%Red./yr	
EAL	JP-4	#######	#######	CH-EW6	Benzene	25	ug/L	0.0000	0.5000	0.0231	4.62	0.5535	42.51	0.5112	0.0236	4.62	0.5545	42.56	
EAL	JP-4	#######	#######	CH-EW6	Toluene	1350	ug/L	0.0002	1.3298	0.0614	4.62	0.4144	33.92	1.7205	0.0795	4.62	0.4263	34.71	
EAL	JP-4	#######	#######	CH-EW6	Ethylbenzene	91300	ug/L	0.0115	0.3585	0.0166	4.48	0.1605	14.83	0.7454	0.0344	4.55	0.1936	17.60	
EAL	JP-4	#######	#######	CH-EW6	Total Xylenes (m,p, and o)	2100000	ug/L	0.2638	2.0562	0.0950	4.10	0.1005	9.56	2.0965	0.0969	4.10	0.1013	9.63	
EAL	JP-4	#######	#######	CH-EW6	Total BTEX	2192675	ug/L	0.2755	4.2445	0.1961	4.34	0.1293	12.13	5.0737	0.2344	4.38	0.1371	12.81	
EAL	JP-4	#######	#######	CH-EW6	B+T	1375	ug/L	0.0002											
EAL	JP-4	#######	#######	CH-EW6	E+X	2191300	ug/L	0.2753											
EAL	JP-4	#######	#######	CH-EW6	B/T	0.018518519		0.0000											
EAL	JP-4	#######	#######	CH-EW6	B/E	0.000273823		0.0000											
EAL	JP-4	#######	#######	CH-EW6	B/X	1.19048E-05		0.0000											
EAL	JP-4	#######	#######	CH-EW6	T/E	0.014786418		0.0000											
EAL	JP-4	#######	#######	CH-EW6	T/X	0.000642857		0.0000											
EAL	JP-4	#######	#######	CH-EW6	E/X	0.04347619		0.0000											
EAL	JP-4	#######	#######	CH-EW6	(B+T)/(E+X)	0.000627481		0.0000											
EAL	JP-4	#######	#######	CH-MW-103	Benzene	25	ug/L	0.0000	0.5000	0.0231	4.62	0.5514	42.39	0.5112	0.0236	4.62	0.5524	42.45	
EAL	JP-4	#######	#######	CH-MW-103	Toluene	204000	ug/L	0.0268	1.3032	0.0602	4.53	0.1804	16.50	1.6939	0.0783	4.55	0.1923	17.49	
EAL	JP-4	#######	#######	CH-MW-103	Ethylbenzene	960000	ug/L	0.1263	0.2437	0.0113	3.04	0.0497	4.85	0.6305	0.0291	3.85	0.0827	7.94	
EAL	JP-4	#######	#######	CH-MW-103	Total Xylenes (m,p, and o)	5400000	ug/L	0.7105	1.6095	0.0744	3.21	0.0547	5.32	1.6498	0.0762	3.23	0.0555	5.40	
EAL	JP-4	#######	#######	CH-MW-103	Total BTEX	6564025	ug/L	0.8637	3.6563	0.1690	3.74	0.0765	7.36	4.4854	0.2073	3.88	0.0843	8.08	
EAL	JP-4	#######	#######	CH-MW-103	B+T	204025	ug/L	0.0268											
EAL	JP-4	#######	#######	CH-MW-103	E+X	6360000	ug/L	0.8368											
EAL	JP-4	#######	#######	CH-MW-103	B/T	0.000122549		0.0000											
EAL	JP-4	#######	#######	CH-MW-103	B/E	2.60417E-05		0.0000											
EAL	JP-4	#######	#######	CH-MW-103	B/X	4.62963E-06		0.0000											
EAL	JP-4	#######	#######	CH-MW-103	T/E	0.2125		0.0000											
EAL	JP-4	#######	#######	CH-MW-103	T/X	0.037777778		0.0000											
EAL	JP-4	#######	#######	CH-MW-103	E/X	0.17777778		0.0000											
EAL	JP-4	#######	#######	CH-MW-103	(B+T)/(E+X)	0.032079403		0.0000											

									Sm	ith et al 198	1			Hu	ahes et al	1984		
Lab	Fuel	Spill	Sample						Lin	ear	Expon	ential		Lin	ear	Expone	ential	
Code	Type	Date	Date	Locid	Analyte	Results Unit	Mass Fraction	C C	k	%Red./vr	k	%Red./vr	C C	k	%Red./vr	k	%Red./vr	
	.,							-0 -				,	-0 -		, j .		,	
NRMRI	JP-4	########	#######	CH-EW6	Density	0.8057 g/ml	0.0001											
				0.1.2.110	Donoty	0.0007 grinz	0.0001											
NRMRI	JP-4	########	#######	CH-EW6	Benzene	10.5 ug/ml	0.0013	0 4987	0 0274	5 49	0.3273	27.91	0.5099	0.0280	5 49	0.3285	28.00	
NRMRL	JP-4	########	########	CH-EW6	Toluene	45.8 ug/ml	0.0057	1.3243	0.0728	5.48	0.3001	25.92	1.7150	0.0943	5.48	0.3142	26.96	
NRMRI	JP-4	########	########	CH-EW6	Ethylbenzene	385 ug/ml	0.0478	0.3222	0.0120	4 79	0 1126	10.65	0 7091	0.0390	5 15	0 1519	14 10	
NRMRL	JP-4	########	########	CH-EW6	m-Xylene	1070 ug/ml	0.1328	0.8272	0.0455	4.74	0.1088	10.31	-0.1328	-0.0073	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRI	JP-4	########	########	CH-EW6	o-Xvlene	92.5 ug/ml	0.0115	0.9985	0.0549	5 44	0.2463	21.83	0.6157	0.0339	5 40	0 2200	19.75	
NRMRI	JP-4	########	########	CH-EW6	p-Xylene	345 ug/ml	0.0428	0.3072	0.0169	4 83	0.1156	10.91	1 6903	0.0930	5.36	0.2035	18.42	
NRMRI	IP-4	########	########	CH-EW6	m p-yvlenes	1415 ug/ml	0.1756	1 1344	0.0624	4 76	0 1105	10.01	1 5575	0.0857	4 94	0.1259	11.83	
NRMRI	IP-4	######################################	########	CH-EW6	Total Xylenes (m.n. and o)	1507.5 ug/ml	0.1700	2 1329	0.0024	5.06	0.1385	12.93	2 1732	0.1195	5.06	0.1200	13.01	
NRMRI	JP-4	########	########	CH-EW6	Total BTEX	1948.8 ug/ml	0.2419	4 2781	0 2353	5.00	0.1610	14.87	5 1073	0.2809	5.00	0 1703	15.66	
NRMRI	IP-4	########	########	CH-EW6	B+T	56.3 ug/ml	0.0070		0.2000	0.21	0.1010		0.1010	0.2000	0.20	0.1100	10.00	
NRMRI	IP-4	######################################	########	CH-EW6	E+X	1892.5 ug/ml	0.0070											
NRMRI	IP-4	######################################	########	CH-EW6	B/T	0 229257642	0.0000											
NRMRI	JP-4	########	########	CH-EW6	B/F	0.027272727	0.0000											
NRMRI	IP-4	########	########	CH-EW6	B/X	0.006965174	0.0000											
NRMRI	JP-4	########	########	CH-EW6	T/E	0.118961039	0.0000											
NRMRI	IP-4	######################################	########	CH-EW6	T/X	0.030381426	0.0000											
NRMRI	IP-4	#########	########	CH-EW6	E/Y	0.255380718	0.0000											
NRMRI	JD-4	#########	########	CH-EW6	(B+T)/(E+X)	0.233303710	0.0000											
NIDMDI		#######################################	****			627 ug/ml	0.0000	0.0779	0.0042	#DI\//01	#DI\//01	#DIV/01	0 2722	0.0205	4 55	0.0065	0.20	
NEME	JF-4	######################################	########	CH-EW6	1,2,3-Trimethylbenzene	1/10 ug/ml	0.0778	-0.0778	-0.0043	#DIV/0:	#010/0:	#DIV/0:	0.5725	0.0203	4.55	0.0903	9.20	
NDMDI	JF-4	######################################	######################################		1,2,4-Trimetryberizerie	700 ug/ml	0.1750	0.8350	0.0439	4.00	0.0904	3.13	0.0417	0.0355	4.32	0.0047	0.12	
NDMDI	JF -4	#########	########		Total TMPs	2926 ug/ml	0.0992	1.0790	0.0170	4.20	0.0794	7.03	1.0267	0.0012	1.02	0.0113	7.13	
	JF -4	*****	*****		Nephthelene	2030 ug/ml	0.3520	0.4419	0.0393	4.13	0.0771	11.42	0.0161	0.0370	4.11	0.0755	9.17	
NRIVIRL	JP-4	########	*****			469 ug/ml	0.0562	0.4416	0.0243	4.00	0.1163	7.40	0.2101	0.0119	4.33	0.0653	0.17	
NRIVIRL	JP-4	########	########		1-MethylNaphthalene	1530 ug/mi	0.1899	0.5901	0.0325	4.16	0.0777	7.48	-0.1537	-0.0085	-23.39	-0.0912	-9.55	
NRIVIRL	JP-4	########	*****		Z-weinyiNaphinalene	1610 ug/mi	0.2246	0.3354	0.0164	3.29	0.0502	4.90	-0.0663	-0.0036	-2.30	-0.0192	-1.94	
INKIVIKL	JP-4	########	########		rotarNapritraieries	3609 ug/mi	0.4720	1.3072	0.0752	4.09	0.0747	7.20	-0.0039	-0.0002	-0.05	-0.0005	-0.05	
NDMDI		ницицици	нининии		Density	0.706 a/ml	0.0001											
INKIVIKL	JP-4	########	########		Density	0.796 g/mL	0.0001											
					Danaaa	0.005	0.0000	0.5000	0.0004	4.00	0.5505	40.54	0.5440	0.0000	4.00	0.5545	40.50	
NRIVIRL	JP-4	#########	########		Belizelle	0.025 ug/ml	0.0000	0.5000	0.0231	4.62	0.5555	42.51	1 7205	0.0230	4.62	0.5545	42.50	
NRIVIRL	JP-4	#########	*****		Toluene	1.35 ug/mi	0.0002	1.3290	0.0614	4.02	0.4144	33.92	0.7454	0.0795	4.62	0.4263	34.71	
NRIVIRL	JP-4	*****	########	CH-EW6	Ethylbenzene	91.3 ug/mi	0.0115	0.3585	0.0166	4.48	0.1605	14.83	0.7454	0.0344	4.55	0.1936	17.60	
NRMRL	JP-4	########	########	CH-EW6	m-Xylene	220 ug/mi	0.0276	0.9324	0.0431	4.49	0.1639	15.12	-0.0276	-0.0013	#DIV/0!	#DIV/0!	#DIV/0!	
NRIVIRL	JP-4	*****	########	CH-EW6	o-Xylene	21.2 Ug/ml	0.0027	1.0073	0.0465	4.61	0.2744	24.00	0.6245	0.0289	4.60	0.2524	22.30	
NRIVIRL	JP-4	########	########	CH-EVV6	p-xylene	110 ug/mi	0.0138	0.3362	0.0155	4.44	0.1493	13.87	1.7193	0.0794	4.58	0.2233	20.01	
NRIVIRL	JP-4	*****	########	CH-EVV6	m,p-xylenes	330 ug/mi	0.0415	1.2685	0.0586	4.47	0.1596	14.75	1.6917	0.0782	4.51	0.1725	15.84	
NRMRL	JP-4	#########	#######################################		Total Aylenes (m,p, and o)	351.2 ug/ml	0.0441	2.2759	0.1052	4.53	0.1831	16.73	2.3162	0.1070	4.53	0.1839	10.80	
NRIVIRL	JP-4	########	#########			443.875 ug/ml	0.0558	4.4642	0.2063	4.56	0.2031	18.38	5.2934	0.2446	4.57	0.2109	19.01	
NRMRL	JP-4	########	#########	CH-EW6	B+1	1.375 ug/ml	0.0002											
NRIVIRL	JP-4	########	#########			442.5 Ug/ml	0.0556											
NRMRL	JP-4	########	#########	CH-EW6	B/I	0.018518519	0.0000											
NRMRL	JP-4	########	########	CH-EW6	B/E	0.000273823	0.0000											
NRMRL	JP-4	########	#########	CH-EW6	B/X	7.11845E-05	0.0000											
NRMRL	JP-4	########	#########	CH-EW6	1/E	0.014786418	0.0000											
NRMRL	JP-4	#######	#######	CH-EW6	1/X	0.003843964	0.0000											
NRMRL	JP-4	########	########	CH-EW6		0.259965831	0.0000											
NRMRL	JP-4	#######	#######	CH-EW6	(B+1)/(E+X)	0.003107345	0.0000											
NRMRL	JP-4	#######	#######	CH-EW6	1,2,3- Frimethylbenzene	448 ug/ml	0.0563	-0.0563	-0.0026	#DIV/0!	#DIV/0!	#DIV/0!	0.3938	0.0182	4.04	0.0961	9.16	
NRMRL	JP-4	#######	#######	CH-EW6	1,2,4-Trimethylbenzene	764 ug/ml	0.0960	0.9140	0.0422	4.18	0.1088	10.30	0.7207	0.0333	4.08	0.0989	9.42	
NRMRL	JP-4	#######	#######	CH-EW6	1,3,5-Trimethylbenzene	541 ug/ml	0.0680	0.3520	0.0163	3.87	0.0842	8.07	0.0539	0.0025	2.04	0.0270	2.66	
NRMRL	JP-4	#######	#######	CH-EW6	Total TMBs	1753 ug/ml	0.2202	1.2098	0.0559	3.91	0.0864	8.28	1.1684	0.0540	3.89	0.0851	8.16	
NRMRL	JP-4	#######	#######	CH-EW6	Naphthalene	446 ug/ml	0.0560	0.4440	0.0205	4.10	0.1011	9.62	0.2183	0.0101	3.68	0.0734	7.08	
NRMRL	JP-4	#######	#######	CH-EW6	1-MethylNaphthalene	1480 ug/ml	0.1859	0.5941	0.0275	3.52	0.0663	6.41	-0.1498	-0.0069	-19.14	-0.0757	-7.86	
NRMRL	JP-4	#######	#######	CH-EW6	2-MethylNaphthalene	1750 ug/ml	0.2198	0.3402	0.0157	2.81	0.0432	4.23	-0.0615	-0.0028	-1.79	-0.0152	-1.53	
NRMRL	JP-4	#######	#######	CH-EW6	Total Naphthalenes	3676 ug/ml	0.4618	1.3782	0.0637	3.46	0.0639	6.19	0.0070	0.0003	0.07	0.0007	0.07	

									Sm	ith et al., 198 [.]	1			Hu	ohes et al., 1	984		
Lab	Fuel	Spill	Sample						Lin	ear	Expone	ential		Lin	ear	Expone	ntial	
Code	Type	Date	Date	Locid	Analyte	Results Units	Mass Fraction	C C	k	%Red./vr	k	%Red./vr	C C	k	%Red./vr	k	%Red./vr	
NRMRI	JP-4	########	########	CH-MW-103	Density	0.76 g/ml	0.0001	-0 -		, j .		, , .	-0 -				/	
				0111111100	Bonony	0.10 g/m2	0.0001											
NRMRL	JP-4	########	#######	CH-MW-103	Benzene	0.025 ug/mL	0.0000	0.5000	0.0231	4.62	0.5514	42.39	0.5112	0.0236	4.62	0.5524	42.45	
NRMRL	JP-4	########	#######	CH-MW-103	Toluene	204 ug/mL	0.0268	1.3032	0.0602	4.53	0.1804	16.50	1.6939	0.0783	4.55	0.1923	17.49	
NRMRL	JP-4	########	#######	CH-MW-103	Ethylbenzene	856 ug/mL	0.1126	0.2574	0.0119	3.21	0.0550	5.35	0.6442	0.0298	3.93	0.0880	8.43	
NRMRL	JP-4	########	#######	CH-MW-103	m-Xvlene	2430 ug/mL	0.3197	0.6403	0.0296	3.08	0.0508	4.95	-0.3197	-0.0148	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-4	#######	#######	CH-MW-103	o-Xylene	1000 ug/mL	0.1316	0.8784	0.0406	4.02	0.0942	8.99	0.4956	0.0229	3.65	0.0722	6.96	
NRMRL	JP-4	#######	#######	CH-MW-103	p-Xylene	968 ug/mL	0.1274	0.2226	0.0103	2.94	0.0467	4.56	1.6058	0.0742	4.28	0.1206	11.37	
NRMRL	JP-4	#######	#######	CH-MW-103	m,p-xylenes	3398 ug/mL	0.4471	0.8629	0.0399	3.04	0.0497	4.85	1.2861	0.0594	3.43	0.0626	6.07	
NRMRL	JP-4	#######	#######	CH-MW-103	Total Xylenes (m,p, and o)	4398 ug/mL	0.5787	1.7413	0.0805	3.47	0.0642	6.22	1.7817	0.0823	3.49	0.0650	6.29	
NRMRL	JP-4	#######	#######	CH-MW-103	Total BTEX	5458.025 ug/mL	0.7182	3.8018	0.1757	3.89	0.0850	8.15	4.6310	0.2140	4.00	0.0928	8.86	
NRMRL	JP-4	#######	#######	CH-MW-103	B+T	204.025 ug/mL	0.0268											
NRMRL	JP-4	#######	#######	CH-MW-103	E+X	5254 ug/mL	0.6913											
NRMRL	JP-4	#######	#######	CH-MW-103	B/T	0.000122549	0.0000											
NRMRL	JP-4	#######	#######	CH-MW-103	B/E	2.92056E-05	0.0000											
NRMRL	JP-4	#######	#######	CH-MW-103	B/X	5.6844E-06	0.0000											
NRMRL	JP-4	#######	#######	CH-MW-103	T/E	0.238317757	0.0000											
NRMRL	JP-4	#######	#######	CH-MW-103	T/X	0.04638472	0.0000											
NRMRL	JP-4	#######	#######	CH-MW-103	E/X	0.194633925	0.0000											
NRMRL	JP-4	#######	#######	CH-MW-103	(B+T)/(E+X)	0.038832318	0.0000											
NRMRL	JP-4	#######	#######	CH-MW-103	1,2,3-Trimethylbenzene	953 ug/mL	0.1254	-0.1254	-0.0058	#DIV/0!	#DIV/0!	#DIV/0!	0.3247	0.0150	3.33	0.0591	5.74	
NRMRL	JP-4	#######	#######	CH-MW-103	1,2,4-Trimethylbenzene	2180 ug/mL	0.2868	0.7232	0.0334	3.31	0.0582	5.65	0.5299	0.0245	3.00	0.0484	4.72	
NRMRL	JP-4	#######	#######	CH-MW-103	1,3,5-Trimethylbenzene	1040 ug/mL	0.1368	0.2832	0.0131	3.12	0.0518	5.05	-0.0150	-0.0007	-0.57	-0.0054	-0.54	
NRMRL	JP-4	#######	#######	CH-MW-103	Total TMBs	4173 ug/mL	0.5491	0.8809	0.0407	2.85	0.0442	4.33	0.8396	0.0388	2.79	0.0429	4.20	
NRMRL	JP-4	#######	#######	CH-MW-103	Naphthalene	502 ug/mL	0.0661	0.4339	0.0201	4.01	0.0935	8.93	0.2083	0.0096	3.51	0.0658	6.37	
NRMRL	JP-4	#######	#######	CH-MW-103	1-MethylNaphthalene	1400 ug/mL	0.1842	0.5958	0.0275	3.53	0.0667	6.45	-0.1481	-0.0068	-18.92	-0.0752	-7.81	
NRMRL	JP-4	#######	#######	CH-MW-103	2-MethylNaphthalene	1800 ug/mL	0.2368	0.3232	0.0149	2.67	0.0398	3.90	-0.0785	-0.0036	-2.29	-0.0186	-1.88	
NRMRL	JP-4	#######	#######	CH-MW-103	Total Naphthalenes	3702 ug/mL	0.4871	1.3529	0.0625	3.40	0.0614	5.96	-0.0183	-0.0008	-0.18	-0.0018	-0.18	
NRMRL	JP-4	#######	5/1/1995	WQ27B	Density	0.775 g/mL	0.0001											
NRMRL	JP-4	#######	5/1/1995	WQ27B	Benzene	209 ug/mL	0.0270	0.4730	0.0241	4.83	0.1490	13.84	0.4843	0.0247	4.83	0.1502	13.94	
NRMRL	JP-4	########	5/1/1995	WQ27B	Toluene	45.4 ug/mL	0.0059	1.3241	0.0676	5.08	0.2769	24.18	1.7148	0.0875	5.09	0.2900	25.17	
NRMRL	JP-4	#######	5/1/1995	WQ27B	Ethylbenzene	2120 ug/mL	0.2735	0.0965	0.0049	1.33	0.0154	1.53	0.4833	0.0247	3.26	0.0519	5.06	
NRMRL	JP-4	########	5/1/1995	WQ27B	m-Xylene	4620 ug/mL	0.5961	0.3639	0.0186	1.93	0.0243	2.40	-0.5961	-0.0304	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-4	########	5/1/1995	WQ27B	o-Xylene	353 ug/mL	0.0455	0.9645	0.0492	4.87	0.1582	14.63	0.5816	0.0297	4.73	0.1338	12.53	
NRMRL	JP-4	########	5/1/1995	WQ27B	p-Xylene	1680 ug/mL	0.2168	0.1332	0.0068	1.94	0.0244	2.42	1.5164	0.0774	4.47	0.1061	10.07	
NRMRL	JP-4	########	5/1/1995	WQ27B	m,p-xylenes	6300 ug/mL	0.8129	0.4971	0.0254	1.94	0.0244	2.41	0.9203	0.0470	2.71	0.0386	3.79	
NRMRL	JP-4	#######	5/1/1995	WQ27B	Total Xylenes (m,p, and o)	6653 ug/mL	0.8585	1.4615	0.0746	3.22	0.0507	4.95	1.5019	0.0766	3.25	0.0516	5.03	
NRMRL	JP-4	########	5/1/1995	WQ27B	I OTAL BIEX	9027.4 ug/mL	1.1648	3.3552	0.1712	3.79	0.0692	6.69	4.1843	0.2135	3.99	0.0778	7.48	
NRMRL	JP-4	#######	5/1/1995	WQ27B	B+1	254.4 ug/mL	0.0328											
NRIVIRL	JP-4	########	5/1/1995	WQ27B	E+A D/T	8773 ug/mL	1.1320											
	JP-4	****	5/1/1995	WO27P	D/I D/E	4.003024229	0.0006			<u> </u>					-			
NDMD	JP-4	######################################	5/1/1995	WQ27P	D/E D/V	0.030304900	0.0000											
NRMD	ID 4	######################################	5/1/1995	WO27P		0.0314144	0.0000											
NRMPI	IP-4	######################################	5/1/1995	WO27B	T/Y	0.021413034	0.0000											
NRMPI	IP-4	######################################	5/1/1995	WO27B	F/X	0.318653239	0.0000											
NRMPI	IP-4	<u>######</u> ##	5/1/1995	W027B	(B+T)/(E+X)	0.028998062	0.0000											
NRMRI	JP-4	#########	5/1/1995	W027B	1 2 3-Trimethylbenzene	986 µg/ml	0.0000	-0 1272	-0.0065	#DIV/0!	#DIV/0!	#DIV/0!	0.3229	0.0165	3.66	0.0645	6 25	
NRMRI	JP-4	#######################################	5/1/1005	WQ27B	1 2 4-Trimethylbenzene	2670 ug/mL	0.1272	0.6655	0.0000	3 26	0.0540	5 3/	0 4722	0.0705	2 95	0.0043	4 31	
NRMPI	IP-4	<u>######</u> ##	5/1/1995	W027B	1 3 5-Trimethylbenzene	1130 µg/mL	0.3445	0.0000	0.0340	3 33	0.0549	5.26	-0.0240	-0.0012	-1.00	-0.0092	-0.02	
NRMRI	JP-4	#########	5/1/1995	W027B	Total TMBs	4786 ug/mL	0.1450	0.8125	0.0415	2 90	0.0340	4 10	0.7711	0.0394	2.83	0.0414	4 05	
NRMRI	JP-4	########	5/1/1995	W027B	Naphthalene	1050 ug/mL	0 1355	0.3645	0.0186	3 72	0.0666	6.45	0.1388	0.0071	2.58	0.0360	3.54	
NRMRI	JP-4	#########	5/1/1995	WQ27B	1-MethylNaphthalene	1590 ug/mL	0.2052	0.5748	0.0293	3.76	0.0682	6.59	-0,1690	-0,0086	-23.85	-0,0886	-9.26	
NRMRI	JP-4	########	5/1/1995	WQ27B	2-MethylNaphthalene	2090 ug/mL	0.2697	0.2903	0.0148	2.65	0.0373	3.66	-0,1113	-0,0057	-3.59	-0,0272	-2.75	
NRMRL	JP-4	########	5/1/1995	WQ27B	Total Naphthalenes	4730 ug/mL	0.6103	1.2297	0.0628	3,41	0.0563	5.48	-0.1415	-0.0072	-1.54	-0.0135	-1.36	
							0.0.00		2.0020	01	2.0000	0.10				2.0.00		
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	Density	0.73 a/mL	0.0001											
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	Benzene	1780 ua/mL	0.2438	0.2562	0.1227	24,54	0.3440	29,11	0.2674	0.1281	25.05	0.3546	29,86	
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	Toluene	6650 ua/mL	0.9110	0.4190	0.2007	15.09	0.1813	16.58	0.8097	0.3879	22.54	0.3046	26.26	
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	Ethylbenzene	1540 ug/mL	0.2110	0.1590	0.0762	20.59	0.2691	23.60	0.5459	0.2615	34.55	0.6119	45.77	
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	m-Xylene	3800 ug/mL	0.5205	0.4395	0.2105	21.93	0.2932	25.41	-0.5205	-0.2493	#DIV/0!	#DIV/0!	#DIV/0!	

										Smi	th et al., 198 [.]	1			Hug	ghes et al.,	1984		
Lab	Fuel	Spill	Sample							Line	ar	Expon	ential		Lin	ear	Expone	ential	
Code	Туре	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	C ₀ - C	k	%Red./yr	k	%Red./yr	C ₀ - C	k	%Red./yr	k	%Red./yr	
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	o-Xylene	2510	ug/mL	0.3438	0.6662	0.3191	31.59	0.5161	40.32	0.2833	0.1357	21.64	0.2879	25.02	
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	p-Xylene	1790	ug/mL	0.2452	0.1048	0.0502	14.34	0.1704	15.67	1.4880	0.7127	41.12	0.9367	60.81	
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	m,p-xylenes	5590	ug/mL	0.7658	0.5442	0.2607	19.90	0.2572	22.68	0.9674	0.4634	26.74	0.3913	32.38	
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	Total Xylenes (m,p, and o)	8100	ug/mL	1.1096	1.2104	0.5798	24.99	0.3533	29.76	1.2508	0.5991	25.38	0.3616	30.34	
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	Total BTEX	18070	ug/mL	2.4753	2.0447	0.9794	21.67	0.2884	25.06	2.8738	1.3766	25.73	0.3691	30.86	
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	B+T	8430	ug/mL	1.1548											
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	E+X	9640	ug/mL	1.3205											
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	B/T	0.267669173		0.0000											
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	B/E	1.155844156		0.0002											
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	B/X	0.219753086		0.0000											
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	T/E	4.318181818		0.0006											
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	T/X	0.820987654		0.0001											
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	E/X	0.190123457		0.0000											
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	(B+T)/(E+X)	0.874481328		0.0001											
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	1,2,3-Trimethylbenzene	3180	ug/mL	0.4356	-0.4356	-0.2087	#DIV/0!	#DIV/0!	#DIV/0!	0.0145	0.0069	1.54	0.0157	1.56	
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	1,2,4-Trimethylbenzene	7720	ug/mL	1.0575	-0.0475	-0.0228	-2.25	-0.0220	-2.23	-0.2408	-0.1154	-14.12	-0.1238	-13.18	
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	1,3,5-Trimethylbenzene	2210	ug/mL	0.3027	0.1173	0.0562	13.37	0.1568	14.51	-0.1809	-0.0867	-71.14	-0.4360	-54.66	
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	Total TMBs	13110	ug/mL	1.7959	-0.3659	-0.1753	-12.26	-0.1091	-11.53	-0.4072	-0.1951	-14.05	-0.1232	-13.11	
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	Naphthalene	1080	ug/mL	0.1479	0.3521	0.1686	33.73	0.5833	44.20	0.1264	0.0605	22.07	0.2958	25.60	
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	1-MethylNaphthalene	1160	ug/mL	0.1589	0.6211	0.2975	38.14	0.7621	53.33	-0.1227	-0.0588	-162.60	-0.7091	-103.22	
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	2-MethylNaphthalene	1870	ug/mL	0.2562	0.3038	0.1455	25.99	0.3746	31.25	-0.0978	-0.0469	-29.59	-0.2304	-25.91	
NRMRL	JP-4	5/1/1995	6/1/1997	JP4-DFSP	Total Naphthalenes	4110	ug/mL	0.5630	1.2770	0.6117	33.24	0.5672	43.29	-0.0942	-0.0451	-9.62	-0.0877	-9.16	

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								Smith et al., 1981					Hughes et al., 1984						
Lab	Fuel	Spill	Sample							Line	ar	Expon	ential		Line	ear	Expor	nential	
Code	Туре	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	C ₀ - C	k	%Red./yr	k	%Red./yr	C ₀ - C	k	%Red./yr	k	%Red./yr	
EAL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	Benzene	2000000	ug/L	0.2596	0.2404	0.0100	2.01	0.0274	2.70	0.2516	0.0105	2.06	0.0283	2.79	
EAL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	Toluene	1000000	ug/L	0.1298	1.2002	0.0502	3.77	0.0973	9.27	1.5909	0.0665	3.87	0.1080	10.24	
EAL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	Ethylbenzene	5500000	ug/L	0.7139	-0.3439	-0.0144	-3.89	-0.0275	-2.79	0.0429	0.0018	0.24	0.0024	0.24	
EAL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	o-Xylene	4300000	ug/L	0.5582	0.4518	0.0189	1.87	0.0248	2.45	0.0690	0.0029	0.46	0.0049	0.49	
EAL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	m,p-Xylene	22000000	ug/L	2.8557	-1.5457	-0.0646	-4.93	-0.0326	-3.31	-1.1225	-0.0469	-2.71	-0.0209	-2.11	
EAL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	Total Xylenes (m,p, and o)	26300000	ug/L	3.4138	-1.0938	-0.0457	-1.97	-0.0161	-1.63	-1.0535	-0.0440	-1.87	-0.0154	-1.55	
EAL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	Total BTEX	34800000	ug/L	4.5171	0.0029	0.0001	0.00	0.0000	0.00	0.8320	0.0348	0.65	0.0071	0.70	
EAL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	B+T	3000000	ug/L	0.3894											
EAL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	E+X	31800000	ug/L	4.1277											
EAL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	B/T	2		0.0000											
EAL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	B/E	0.363636364		0.0000											
EAL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	B/X	0.076045627		0.0000											
EAL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	T/E	0.181818182		0.0000											
EAL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	T/X	0.038022814		0.0000											
EAL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	E/X	0.209125475		0.0000											
EAL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	(B+T)/(E+X)	0.094339623		0.0000											
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	Density	0.7704	g/mL	0.0001											
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	Benzene	900	ug/mL	0.1168	0.3832	0.0160	3.20	0.0608	5.90	0.3944	0.0165	3.23	0.0617	5.98	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	Toluene	0.025	ug/mL	0.0000	1.3300	0.0556	4.18	0.5403	41.74	1.7207	0.0719	4.18	0.5510	42.36	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	Ethylbenzene	2960	ug/mL	0.3842	-0.0142	-0.0006	-0.16	-0.0016	-0.16	0.3726	0.0156	2.06	0.0283	2.79	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	m-Xylene	8880	ug/mL	1.1526	-0.1926	-0.0081	-0.84	-0.0076	-0.77	-1.1526	-0.0482	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	. JP-4	10/1/1973	8/27/1997	EAKMW316-FP	o-Xylene	2400	ug/mL	0.3115	0.6985	0.0292	2.89	0.0492	4.80	0.3157	0.0132	2.10	0.0293	2.88	
NRMRL	. JP-4	10/1/1973	8/27/1997	EAKMW316-FP	p-Xylene	4120	ug/mL	0.5348	-0.1848	-0.0077	-2.21	-0.0177	-1.79	1.1984	0.0501	2.89	0.0492	4.80	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	m,p-xylenes	13000	ug/mL	1.6874	-0.3774	-0.0158	-1.20	-0.0106	-1.06	0.0457	0.0019	0.11	0.0011	0.11	
NRMRL	. JP-4	10/1/1973	8/27/1997	EAKMW316-FP	Total Xylenes (m,p, and o)	15400	ug/mL	1.9990	0.3210	0.0134	0.58	0.0062	0.62	0.3614	0.0151	0.64	0.0069	0.69	
NRMRL	. JP-4	10/1/1973	8/27/1997	EAKMW316-FP	Total BTEX	19260.025	ug/mL	2.5000	2.0200	0.0844	1.87	0.0248	2.45	2.8491	0.1191	2.23	0.0318	3.13	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	B+T	900.025	ug/mL	0.1168											
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	E+X	18360	ug/mL	2.3832											
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	B/T	36000		4.6729											
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	B/E	0.304054054		0.0000											
NRMRL	. JP-4	10/1/1973	8/27/1997	EAKMW316-FP	B/X	0.058441558		0.0000											
NRMRL	. JP-4	10/1/1973	8/27/1997	EAKMW316-FP	T/E	8.44595E-06		0.0000											
NRMRL	. JP-4	10/1/1973	8/27/1997	EAKMW316-FP	T/X	1.62338E-06		0.0000											
NRMRL	. JP-4	10/1/1973	8/27/1997	EAKMW316-FP	E/X	0.192207792		0.0000											
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	(B+T)/(E+X)	0.049020969		0.0000											
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	1,2,3-Trimethylbenzene	3440	ug/mL	0.4465	-0.4465	-0.0187	#DIV/0!	#DIV/0!	#DIV/0!	0.0036	0.0002	0.03	0.0003	0.03	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	1,2,4-Trimethylbenzene	9610	ug/mL	1.2474	-0.2374	-0.0099	-0.98	-0.0088	-0.89	-0.4307	-0.0180	-2.20	-0.0177	-1.79	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	1,3,5-Trimethylbenzene	4980	ug/mL	0.6464	-0.2264	-0.0095	-2.25	-0.0180	-1.82	-0.5246	-0.0219	-18.00	-0.0698	-7.23	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	Total TMBs	18030	ug/mL	2.3403	-0.9103	-0.0381	-2.66	-0.0206	-2.08	-0.9517	-0.0398	-2.87	-0.0218	-2.21	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	Naphthalene	1000	ug/mL	0.1298	0.3702	0.0155	3.10	0.0564	5.48	0.1445	0.0060	2.20	0.0313	3.08	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	1-MethylNaphthalene	1610	ug/mL	0.2090	0.5710	0.0239	3.06	0.0551	5.36	-0.1728	-0.0072	-19.98	-0.0733	-7.61	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	2-MethylNaphthalene	2900	ug/mL	0.3764	0.1836	0.0077	1.37	0.0166	1.65	-0.2181	-0.0091	-5.76	-0.0362	-3.69	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW316-FP	Total Naphthalenes	5510	ug/mL	0.7152	1.1248	0.0470	2.56	0.0395	3.87	-0.2464	-0.0103	-2.20	-0.0177	-1.78	

								Smith et al., 1981 Hughes et al., 1984											
Lab	Fuel	Spill	Sample							Lin	ear	Expone	ential		Lin	ear	Expon	ential	
Code	Туре	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	C° - C	k	%Red./yr	k	%Red./yr	C ₀ - C	k	%Red./yr	k	%Red./yr	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	Density	0.7626	g/mL	0.0001					ĺ						
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	Benzene	2.7	ug/mL	0.0004	0.4996	0.0209	4.18	0.3032	26.16	0.5109	0.0214	4.18	0.3041	26.22	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	Toluene	0.025	ug/mL	0.0000	1.3300	0.0556	4.18	0.5398	41.72	1.7207	0.0719	4.18	0.5506	42.34	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	Ethylbenzene	2480	ug/mL	0.3252	0.0448	0.0019	0.51	0.0054	0.54	0.4317	0.0180	2.38	0.0353	3.47	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	m-Xylene	530	ug/mL	0.0695	0.8905	0.0372	3.88	0.1098	10.40	-0.0695	-0.0029	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	o-Xylene	1690	ug/mL	0.2216	0.7884	0.0330	3.26	0.0634	6.14	0.4056	0.0170	2.70	0.0435	4.26	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	p-Xylene	2930	ug/mL	0.3842	-0.0342	-0.0014	-0.41	-0.0039	-0.39	1.3490	0.0564	3.25	0.0630	6.10	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	m,p-xylenes	3460	ug/mL	0.4537	0.8563	0.0358	2.73	0.0443	4.34	1.2795	0.0535	3.09	0.0560	5.45	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	Total Xylenes (m,p, and o)	5150	ug/mL	0.6753	1.6447	0.0688	2.96	0.0516	5.03	1.6850	0.0704	2.98	0.0523	5.10	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	Total BTEX	7632.725	ug/mL	1.0009	3.5191	0.1471	3.25	0.0630	6.11	4.3482	0.1818	3.40	0.0701	6.77	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	B+T	2.725	ug/mL	0.0004											
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	E+X	7630	ug/mL	1.0005											
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	B/T	108		0.0142											
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	B/E	0.00108871		0.0000											
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	B/X	0.000524272		0.0000											
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	T/E	1.00806E-05		0.0000											
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	T/X	4.85437E-06		0.0000											
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	E/X	0.481553398		0.0001											
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	(B+T)/(E+X)	0.000357143		0.0000											
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	1,2,3-Trimethylbenzene	1950	ug/mL	0.2557	-0.2557	-0.0107	#DIV/0!	#DIV/0!	#DIV/0!	0.1944	0.0081	1.81	0.0236	2.34	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	1,2,4-Trimethylbenzene	7570	ug/mL	0.9927	0.0173	0.0007	0.07	0.0007	0.07	-0.1759	-0.0074	-0.90	-0.0082	-0.82	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	1,3,5-Trimethylbenzene	4210	ug/mL	0.5521	-0.1321	-0.0055	-1.31	-0.0114	-1.15	-0.4302	-0.0180	-14.76	-0.0632	-6.52	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	Total TMBs	13730	ug/mL	1.8004	-0.3704	-0.0155	-1.08	-0.0096	-0.97	-0.4118	-0.0172	-1.24	-0.0109	-1.09	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	Naphthalene	1050	ug/mL	0.1377	0.3623	0.0151	3.03	0.0539	5.25	0.1366	0.0057	2.08	0.0288	2.84	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	1-MethylNaphthalene	2150	ug/mL	0.2819	0.4981	0.0208	2.67	0.0425	4.17	-0.2458	-0.0103	-28.41	-0.0859	-8.96	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	2-MethylNaphthalene	3950	ug/mL	0.5180	0.0420	0.0018	0.31	0.0033	0.33	-0.3596	-0.0150	-9.49	-0.0495	-5.08	
NRMRL	JP-4	10/1/1973	8/27/1997	EAKMW306-FP	Total Naphthalenes	7150	ug/mL	0.9376	0.9024	0.0377	2.05	0.0282	2.78	-0.4688	-0.0196	-4.18	-0.0290	-2.94	

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							Smith et al., 1981						Hughes et al., 1984					
Lab	Fuel	Spill	Sample							Line	ear	Expon	ential		Lin	ear	Expor	ential
Code	Туре	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	C ₀ - C	k	%Red./yr	k	%Red./yr	C ₀ - C	k	%Red./yr	-k	%Red./yr
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	Density	0.8512	g/mL	0.0001										
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	Benzene	0.025	ug/mL	0.0000	0.5000	0.0224	4.48	0.5398	41.71	0.5112	0.0229	4.48	0.5408	41.77
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	Toluene	0.025	ug/mL	0.0000	1.3300	0.0596	4.48	0.5836	44.21	1.7207	0.0771	4.48	0.5952	44.85
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	Ethylbenzene	0.025	ug/mL	0.0000	0.3700	0.0166	4.48	0.5263	40.92	0.7569	0.0339	4.48	0.5583	42.78
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	m-Xylene	0.025	ug/mL	0.0000	0.9600	0.0430	4.48	0.5690	43.39	0.0000	0.0000	#DIV/0!	#DIV/0!	#DIV/0!
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	o-Xylene	0.025	ug/mL	0.0000	1.0100	0.0453	4.48	0.5713	43.52	0.6272	0.0281	4.48	0.5499	42.30
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	p-Xylene	0.025	ug/mL	0.0000	0.3500	0.0157	4.48	0.5238	40.77	1.7332	0.0777	4.48	0.5955	44.87
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	m,p-xylenes	0.05	ug/mL	0.0000	1.3100	0.0587	4.48	0.5519	42.41	1.7332	0.0777	4.48	0.5644	43.13
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	Total Xylenes (m,p, and o)	0.075	ug/mL	0.0000	2.3200	0.1040	4.48	0.5593	42.84	2.3603	0.1058	4.48	0.5601	42.88
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	Total BTEX	0.15	ug/mL	0.0000	4.5200	0.2026	4.48	0.5581	42.77	5.3491	0.2397	4.48	0.5657	43.20
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	B+T	0.05	ug/mL	0.0000										
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	E+X	0.1	ug/mL	0.0000										
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	B/T	1		0.0001										
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	B/E	1		0.0001										
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	B/X	0.3333333333		0.0000										
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	T/E	1		0.0001										
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	T/X	0.333333333		0.0000										
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	E/X	0.333333333		0.0000										
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	(B+T)/(E+X)	0.5		0.0001										
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	1,2,3-Trimethylbenzene	55.6	ug/mL	0.0065	-0.0065	-0.0003	#DIV/0!	#DIV/0!	#DIV/0!	0.4436	0.0199	4.42	0.1897	17.28
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	1,2,4-Trimethylbenzene	44.4	ug/mL	0.0052	1.0048	0.0450	4.46	0.2360	21.02	0.8115	0.0364	4.45	0.2265	20.27
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	1,3,5-Trimethylbenzene	4.37	ug/mL	0.0005	0.4195	0.0188	4.48	0.3006	25.96	0.1213	0.0054	4.46	0.2451	21.74
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	Total TMBs	104.37	ug/mL	0.0123	1.4177	0.0635	4.44	0.2133	19.21	1.3764	0.0617	4.44	0.2119	19.10
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	Naphthalene	0.025	ug/mL	0.0000	0.5000	0.0224	4.48	0.5398	41.71	0.2743	0.0123	4.48	0.5129	40.12
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	1-MethylNaphthalene	132	ug/mL	0.0155	0.7645	0.0343	4.39	0.1756	16.10	0.0207	0.0009	2.56	0.0379	3.72
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	2-MethylNaphthalene	122	ug/mL	0.0143	0.5457	0.0245	4.37	0.1643	15.15	0.1440	0.0065	4.08	0.1077	10.21
NRMRL	JP-4	6/1/1975	9/18/1997	WA-CR02	Total Naphthalenes	254.025	ug/mL	0.0298	1.8102	0.0811	4.41	0.1847	16.86	0.4390	0.0197	4.20	0.1234	11.61



Linear (Zero Order) Assumption

linear equation $C = C_o - kt$

summary statistics presented as follows

ana	alyte
k	Co
sek	seCo
r ²	seC
F stat.	df
SS _{regr}	SS _{resid}

- k = zero order weathering rate; k = dC/dt or slope
- sek = slope standard error value
- Co = intercept or initial analyte concentration as calculated by regression analysis
- seCo = standard error value for the constant C_0
 - r^{2} = coefficient of determination
- seC = standard error value of the estimated concentration C (i.e., a "standard deviation" for the regression line)
- F stat. = F statistic or F-observed value
- df = degrees of freedom
- ss_{regr} = the regression sum of squares
- ss_{resid} = the residual sum of squares

Exponential (1st Order) Assumption

exponential equation $C = C_0 e^{-kt}$

summary statistics presented as follows



m = coefficient for statistics equation shown; note $m = e^{-kt}$, therefore, ln m = -k

- sek = standard error value for the exponential rate constant k
- Co = intercept or initial analyte concentration as calculated by regression analysis
- seCo = standard error value for the constant C_o ; compare to ln C_o
 - r^2 = coefficient of determination
- seC = standard error value of the estimated concentration C (i.e., a "standard deviation" for the regression line)
- F stat. = F statistic or F-observed value
 - df = degrees of freedom
- ss_{regr} = the regression sum of squares
- ss_{resid} = the residual sum of squares
- $\ln C_o =$ natural log of Co for comparing to seCo

SHAW AFB, BLDG 1610

Evergreen Analytical Data

Sample									
Location	Date	Time (yrs)	Benzene	Toluene	Ethylbenzene	o-Xylene	m,p-Xylenes	Total Xylenes	Total BTEX
SHMW1610-2	6/1/1994	0.00	0.50	1.33	0.37	1.01	1.31	2.32	4.52
	3/6/1997	2.76	0.35	0.75	0.23	0.46	0.98	1.44	2.76
	3/11/1998	3.78	0.17	0.38	0.18	0.40	0.76	1.15	1.88
linear									
Co			0.5151	1.3487	0.3701	0.9924	1.3224	2.3148	4.5487
Predicted C - la	test sample	date	0.2078	0.4356	0.1798	0.3494	0.7902	1.1395	1.9628
linear rate cons	be)	0.0813	0.2417	0.0504	0.1702	0.1409	0.3111	0.6844	
average yearly	reduction (%	5)	15.79	17.92	13.61	17.15	10.65	13.44	15.05

linear

summary stats

Benz	zene	Tolu	ene	Ethylber	nzene	Total 2	Kylenes	Total	BTEX
-0.0813	0.5151	-0.2417	1.3487	-0.0504	0.3701	-0.3111	2.3148	-0.6844	4.5487
0.0258	0.0696	0.0320	0.0864	0.0002	0.0006	0.0090	0.0242	0.0490	0.1323
0.9089	0.0712	0.9828	0.0884	1.0000	0.0006	0.9992	0.0248	0.9949	0.1354
10	1	57	1	52761	1	1203	1	195	1
0.0506	0.0051	0.4467	0.0078	0.0194	0.0000	0.7400	0.0006	3.5827	0.0183

exponential	Benzene	Toluene	Ethylbenzene	o-Xylene	m,p-Xylenes	Total Xylenes	Total BTEX
Со	0.5378	1.4066	0.3737	0.9912	1.3344	2.3320	4.6331
exponential rate constant (k)	0.2575	0.3028	0.1869	0.2554	0.1369	0.1825	0.2202
% reduction/year	22.70	26.13	17.05	22.54	12.80	16.68	19.77

exponential summary stats

Ben	zene	Tolu	ene	Ethylben	zene	Total	Xylenes	Total	BTEX
0.7730	0.5378	0.7387	1.4066	0.8295	0.3737	0.8332	2.3320	0.8023	4.6331
0.1246	0.3368	0.0957	0.2585	0.0171	0.0461	0.0088	0.0238	0.0422	0.1141
0.8102	0.3446	0.9093	0.2645	0.9917	0.0472	0.9977	0.0243	0.9646	0.1168
4	1	10	1	120	1	430	1	27	1
0.5069	0.1187	0.7012	0.0700	0.2672	0.0022	0.2548	0.0006	0.3710	0.0136
	-0.6202		0.3412		-0.9843		0.8467		1.5332

SHAW AFB, BLDG 1610 (continued) NRMRL Data

Sample Location	Date	Time (yrs)	Benzene	Toluene	Ethylbenzene	m-Xylene	o-Xylene	p-Xylene	m,p-xylenes	Total Xylenes	Total BTEX
SHMW1610-2	6/1/1994	0.00	0.50	1.33	0.37	0.96	1.01	0.35	1.31	2.32	4.52
	3/6/1997	2.76	0.29	0.64	0.18	0.54	0.39	0.19	0.73	1.12	2.22
	3/11/1998	3.78	0.16	0.36	0.13	0.44	0.31	0.17	0.61	0.92	1.58
linear											
Co			0.5071	1.3328	0.3664	0.9535	0.9911	0.3452	1.2987	2.2898	4.4961
Predicted C - la	test sample	date	0.1797	0.3705	0.1235	0.4232	0.2600	0.1549	0.5782	0.8381	1.5118
linear rate cons	tant (k) (slop	e)	0.0867	0.2547	0.0643	0.1403	0.1935	0.0504	0.1907	0.3842	0.7899
average yearly	reduction (%)	17.09	19.11	17.55	14.72	19.53	14.59	14.68	16.78	17.57

linear

summary stats

Benz	zene	Toluene		Ethylber	nzene	Total 2	Kylenes	Total	BTEX
-0.0867	0.5071	-0.2547	1.3328	-0.0643	0.3664	-0.3842	2.2898	-0.7899	4.4961
0.0122	0.0329	0.0048	0.0130	0.0062	0.0167	0.0516	0.1395	0.0408	0.1103
0.9806	0.0337	0.9996	0.0133	0.9908	0.0171	0.9823	0.1427	0.9973	0.1128
51	1	2795	1	108	1	55	1	375	1
0.0574	0.0011	0.4962	0.0002	0.0316	0.0003	1.1291	0.0204	4.7718	0.0127

exponential	Benzene	Toluene	Ethylbenzene	m-Xylene	o-Xylene	p-Xylene	m,p-xylenes	Total Xylenes	Total BTEX
Со	0.5258	1.3792	0.3699	0.9594	0.9929	0.3452	1.3045	2.2983	4.5663
exponential rate constant (k)	0.2781	0.3272	0.2702	0.2062	0.3191	0.2007	0.2047	0.2490	0.2741
% reduction/year	24.28	27.91	23.68	18.63	27.32	18.18	18.51	22.04	23.97

exponential

Ber	izene	Toluene		Ethylben	zene	Total 2	Xylenes	Total BTEX		
0.7572	0.5258	0.7209	1.3792	0.7632	0.3699	0.7796	2.2983	0.7603	4.5663	
0.0861	0.2327	0.0621	0.1679	0.0003	0.0008	0.0161	0.0434	0.0174	0.0471	
0.9125	0.2381	0.9652	0.1718	1.0000	0.0008	0.9959	0.0444	0.9960	0.0482	
10	1	28	1	926703	1	240	1	247	1	
0.5916	0.0567	0.8188	0.0295	0.5585	0.0000	0.4741	0.0020	0.5744	0.0023	
	-0.6428		0.3215		-0.9944		0.8322		1.5187	

SHAW AFB, BLDG 1610 (continued) NRMRL Data (cont.)

Sample Location	Date	Time (yrs)	Benzene	Toluene	Ethylbenzene	m-Xylene	o-Xylene	p-Xylene	m,p-xylenes	Total Xylenes	Total BTEX
SHMW1610-3	6/1/1994	0.00	0.50	1.33	0.37	0.96	1.01	0.35	1.31	2.32	4.52
	3/6/1997	2.76	0.18	0.32	0.12	0.41	0.31	0.15	0.57	0.87	1.49
	3/11/1998	3.78	0.21	0.42	0.14	0.45	0.32	0.17	0.61	0.93	1.70
linear											
Со			0.4811	1.2722	0.3565	0.9315	0.9772	0.3394	1.2709	2.2482	4.3580
Predicted C - la	test sample	date	0.1608	0.2659	0.1010	0.3690	0.2259	0.1372	0.5061	0.7320	1.2597
linear rate cons	tant (k) (slop	e)	0.0848	0.2664	0.0676	0.1489	0.1989	0.0535	0.2024	0.4013	0.8201
average yearly	reduction (%)	17.62	20.94	18.97	15.99	20.35	15.77	15.93	17.85	18.82

linear

summary s	stats
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-	Benze	ene	Tolu	ene	Ethylben	zene	Total	Xylenes	Total	BTEX	
	-0.0848	0.4811	-0.2664	1.2722	-0.0676	0.3565	-0.4013	2.2482	-0.8201	4.3580	
	0.0323	0.0873	0.0987	0.2668	0.0230	0.0622	0.1228	0.3318	0.2768	0.7481	
	0.8732	0.0894	0.8792	0.2730	0.8964	0.0636	0.9144	0.3395	0.8977	0.7654	
	7	1	7	1	9	1	11	1	9	1	
	0.0550	0.0080	0.5426	0.0745	0.0350	0.0040	1.2316	0.1153	5.1434	0.5859	
exponential			Benzene	Toluene	Ethylbenzene	m-Xylene	o-Xylene	p-Xylene	m,p-xylenes	Total Xylenes	Total BTEX
Co			0.4664	1.2062	0.3457	0.9159	0.9535	0.3335	1.2495	2.2033	4.2304
exponential rate	constant (k)		0.2584	0.3476	0.2927	0.2240	0.3344	0.2194	0.2228	0.2661	0.2889
% reduction/yea	r		22.77	29.36	25.38	20.07	28.43	19.70	19.97	23.37	25.09

exponential

Ber	izene	Tolu	ene	Ethylben	zene	Total	Xylenes	Total BTEX			
0.7723	0.4664	0.7064	1.2062	0.7462	0.3457	0.7663	2.2033	0.7491	4.2304		
0.1187	0.3208	0.1670	0.4513	0.1163	0.3143	0.0882	0.2383	0.1131	0.3058		
0.8258	0.3282	0.8125	0.4617	0.8637	0.3216	0.9011	0.2439	0.8670	0.3129		
5	1	4	1	6	1	9	1	7	1		
0.5107	0.1077	0.9241	0.2132	0.6553	0.1034	0.5417	0.0595	0.6382	0.0979		
	-0.7626		0.1875		-1.0623		0.7900		1.4423		

DFSP-Charleston, Tank 1 NRMRL Data

Sample Location	Date	Time (yrs)	Benzene	Toluene	Ethylbenzene	m-Xylene	o-Xylene	p-Xylene	m,p-xylenes	Total Xylenes	Total BTEX
EW-6	10/1/1975	0.00	0.50	1.33	0.37	0.96	1.01	0.35	1.31	2.32	4.52
	12/1/1993	18.18	0.0013	0.0057	0.0478	0.1328	0.0115	0.0428	0.1756	0.1871	0.2419
	5/17/1997	21.64	0.0000	0.0002	0.0115	0.0276	0.0027	0.0138	0.0415	0.0441	0.0558
linear											
Со			0.4927	1.3109	0.3681	0.9559	0.9959	0.3477	1.3037	2.2996	4.4713
Predicted C - la	atest sample	date	-0.0382	-0.1003	0.0013	0.0063	-0.0712	0.0018	0.0081	-0.0631	-0.2003
linear rate cons	stant (k) (slop	be)	0.0245	0.0652	0.0169	0.0439	0.0493	0.0160	0.0599	0.1092	0.2159
average yearly	reduction (%)	4.98	4.97	4.60	4.59	4.95	4.60	4.59	4.75	4.83

linear

summary stats

ſ	Benz	zene	Tolu	iene	Ethylber	nzene	Total X	Kylenes	Total BTEX				
ſ	-0.0245	0.4927	-0.0652	1.3109	-0.0169	0.3681	-0.1092	2.2996	-0.2159	4.4713			
	0.0036	0.0593	0.0096	0.1562	0.0010	0.0159	0.0102	0.1666	0.0244	0.3981			
	0.9785	0.0598	0.9789	0.1574	0.9967	0.0160	0.9913	0.1679	0.9874	0.4010			
	46	1	46	1	304	1	114	1	78	1			
	0.1627	0.0036	1.1493	0.0248	0.0776	0.0003	3.2216	0.0282	12.5946	0.1608			

exponential	Benzene	Toluene	Ethylbenzene	m-Xylene	o-Xylene	p-Xylene	m,p-xylenes	Total Xylenes	Total BTEX
Со	0.7310	1.6114	0.4010	1.0531	1.0589	0.3704	1.4224	2.5005	4.8507
exponential rate constant (k)	0.4788	0.3766	0.1447	0.1457	0.2651	0.1382	0.1434	0.1684	0.1892
% reduction/year	38.05	31.38	13.47	13.56	23.29	12.91	13.36	15.50	17.24

exponential

Benz	zene	Tolu	ene	Ethylben	zene	Total	Xylenes	Total BTEX			
0.6195	0.7310	0.6862	1.6114	0.8653	0.4010	0.8450	2.5005	0.8276	4.8507		
0.1901	3.1026	0.0961	1.5678	0.0403	0.6575	0.0375	0.6119	0.0353	0.5767		
0.8638	3.1258	0.9389	1.5795	0.9280	0.6624	0.9527	0.6165	0.9663	0.5811		
6	1	15	1	13	1	20	1	29	1		
61.9656	9.7706	38.3405	2.4947	5.6589	0.4387	7.6621	0.3801	9.6766	0.3376		
	-0.3133		0.4771		-0.9138		0.9165		1.5791		

										Beaufort MCAS Fresh JP-5					Hughes et al., 1984				
Lab	Fuel	Spill	Sample							Lin	ear	Expor	nential		Lin	ear	Expor	nential	
Code	Туре	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	C ₀ - C	k	%Red./yr	k	%Red./yr	C° - C	k	%Red./yr	k	%Red./yr	
EAL	JP-5	6/1/1990	8/12/1997	BFT-401-3	Benzene	2230	ug/L	0.0003	-0.0001	0.0000	-6.13	-0.0508	-5.21	-0.0003	0.0000	#DIV/0!	#DIV/0!	#DIV/0!	
EAL	JP-5	6/1/1990	8/12/1997	BFT-401-3	Toluene	20000	ug/L	0.0025	0.0022	0.0003	6.49	0.0874	8.37	-0.0025	-0.0003	#DIV/0!	#DIV/0!	#DIV/0!	
EAL	JP-5	6/1/1990	8/12/1997	BFT-401-3	Ethylbenzene	160000	ug/L	0.0199	0.0220	0.0031	7.30	0.1035	9.83	-0.0199	-0.0028	#DIV/0!	#DIV/0!	#DIV/0!	
EAL	JP-5	6/1/1990	8/12/1997	BFT-401-3	o-Xylene	330000	ug/L	0.0410	0.0600	0.0083	8.25	0.1251	11.76	-0.0209	-0.0029	-14.43	-0.0989	-10.40	
EAL	JP-5	6/1/1990	8/12/1997	BFT-401-3	m,p-Xylene	680000	ug/L	0.0845	0.0538	0.0075	5.40	0.0684	6.61	-0.0845	-0.0117	#DIV/0!	#DIV/0!	#DIV/0!	
EAL	JP-5	6/1/1990	8/12/1997	BFT-401-3	Total Xylenes (m,p, and o)	1010000	ug/L	0.1256	0.1138	0.0158	6.60	0.0896	8.57	-0.1054	-0.0146	-72.76	-0.2542	-28.95	
EAL	JP-5	6/1/1990	8/12/1997	BFT-401-3	Total BTEX	1192230	ug/L	0.1482	0.1379	0.0191	6.69	0.0913	8.73	-0.1281	-0.0178	-88.40	-0.2773	-31.95	
EAL	JP-5	6/1/1990	8/12/1997	BFT-401-3	B+T	22230	ug/L	0.0028											
EAL	JP-5	6/1/1990	8/12/1997	BFT-401-3	E+X	1170000	ug/L	0.1455											
EAL	JP-5	6/1/1990	8/12/1997	BFT-401-3	B/T	0.1115		0.0000											
EAL	JP-5	6/1/1990	8/12/1997	BFT-401-3	B/E	0.0139375		0.0000											
EAL	JP-5	6/1/1990	8/12/1997	BFT-401-3	B/X	0.002207921		0.0000											
EAL	JP-5	6/1/1990	8/12/1997	BFT-401-3	T/E	0.125		0.0000											
EAL	JP-5	6/1/1990	8/12/1997	BFT-401-3	T/X	0.01980198		0.0000											
EAL	JP-5	6/1/1990	8/12/1997	BFT-401-3	E/X	0.158415842		0.0000											
EAL	JP-5	6/1/1990	8/12/1997	BFT-401-3	(B+T)/(E+X)	0.019		0.0000											
EAL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	Benzene	1500	ug/L	0.0002	0.0000	0.0000	0.00	0.0000	0.00	-0.0002	0.0000	#DIV/0!	1.1314	67.74	
EAL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	Toluene	36000	ug/L	0.0046	0.0001	0.0000	0.16	0.0016	0.16	-0.0046	-0.0007	#DIV/0!	0.8495	57.24	
EAL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	Ethylbenzene	370000	ug/L	0.0474	-0.0055	-0.0008	-1.89	-0.0177	-1.79	-0.0474	-0.0068	#DIV/0!	0.3974	32.79	
EAL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	Total Xylenes (m,p, and o)	2600000	ug/L	0.3333	-0.0940	-0.0135	-5.63	-0.0475	-4.87	-0.3132	-0.0449	-223.37	0.2808	24.49	
EAL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	Total BTEX	3007500	ug/L	0.3856	-0.0994	-0.0143	-4.99	-0.0428	-4.37	-0.3655	-0.0524	-260.63	0.3773	31.43	
EAL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	B+T	37500	ug/L	0.0048											
EAL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	E+X	2970000	ug/L	0.3808											
EAL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	B/T	0.041666667		0.0000											
EAL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	B/E	0.004054054		0.0000											
EAL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	B/X	0.000576923		0.0000											
EAL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	T/E	0.097297297		0.0000											
EAL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	T/X	0.013846154		0.0000											
EAL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	E/X	0.142307692		0.0000											
EAL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	(B+T)/(E+X)	0.012626263		0.0000											

									Beaufort MCAS Fresh JP-5					Hughes et al., 1984					
Lab	Fuel	Spill	Sample							Lin	ear	Expor	ential		l in	ear	Expon	ential	
Code	Type	Date	Date	Locid	Analyte	Posulte	Unite	Mass Fraction	0.0		% Pod hr	k	%Red hr	0-0	k	%Red //r	k	%Red hr	
Coue	Type	Date	Date	Lociu	Analyte	Results	Units	Wass Fraction	0,-0	~	/61\eu./yi	~	/orceu./yr	0-U	n	/orceu./yi	ĸ	/arteu./yr	
NIDMDI		6/1/1000	0/10/1007	DET 401 2	Depaits	0.8044	a/ml	0.0001											
NRIVIRL	JP-5	6/1/1990	8/12/1997	BF1-401-3	Density	0.8044	g/mL	0.0001											
		0/4/4000	0/40/4007	DET 404 0	Deserve	0.00		0.0000	0.0004	0.0000	0.40	0.0500	5.04	0.0000	0.0000	#DIV//01	#DIV//01	#DIV//01	
NRIVIRL	JP-5	6/1/1990	8/12/1997	BF1-401-3	Benzene	2.23	ug/mL	0.0003	-0.0001	0.0000	-6.13	-0.0508	-5.21	-0.0003	0.0000	#DIV/0!	#DIV/0!	#DIV/0!	
NRIVIRL	JP-5	6/1/1990	8/12/1997	BF1-401-3	Toluene	13	ug/mL	0.0016	0.0031	0.0004	9.08	0.1472	13.69	-0.0016	-0.0002	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	Ethylbenzene	116	ug/mL	0.0144	0.0275	0.0038	9.11	0.1482	13.77	-0.0144	-0.0020	#DIV/0!	#DIV/0!	#DIV/0!	
NRIVIRL	JP-5	6/1/1990	8/12/1997	BF1-401-3	m-xyiene	224	ug/mL	0.0278	0.0698	0.0097	9.93	0.1743	15.99	-0.0278	-0.0039	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	o-Xylene	287	ug/mL	0.0357	0.0653	0.0091	8.98	0.1445	13.45	-0.0156	-0.0022	-10.74	-0.0795	-8.28	
NRIVIRL	JP-5	6/1/1990	8/12/1997	BF1-401-3	p-xylene	99.6	ug/mL	0.0124	0.0283	0.0039	9.65	0.1650	15.21	-0.0124	-0.0017	#DIV/0!	#DIV/0!	#DIV/0!	
NRIVIRL	JP-5	6/1/1990	8/12/1997	BF1-401-3	m,p-xylenes	323.6	ug/mL	0.0402	0.0981	0.0136	9.85	0.1715	15.76	-0.0402	-0.0056	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	Total Xylenes (m,p, and o)	610.6	ug/mL	0.0759	0.1635	0.0227	9.48	0.1594	14.74	-0.0558	-0.0077	-38.50	-0.1844	-20.24	
NRMRL	JP-5	6/1/1990	8/12/1997	BF1-401-3	Iotal BIEX	741.83	ug/mL	0.0922	0.1939	0.0269	9.41	0.1572	14.55	-0.0721	-0.0100	-49.76	-0.2114	-23.54	
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	B+T	15.23	ug/mL	0.0019											
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	E+X	726.6	ug/mL	0.0903											
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	B/T	0.171538462		0.0000											
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	B/E	0.019224138		0.0000											
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	B/X	0.003652145		0.0000											
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	I/E	0.112068966		0.0000											
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	T/X	0.021290534		0.0000											
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	E/X	0.189977072		0.0000											
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	(B+T)/(E+X)	0.020960639		0.0000											
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	1,2,3-Trimethylbenzene	1700	ug/mL	0.2113	0.0951	0.0132	4.31	0.0516	5.03	0.3555	0.0494	8.71	0.1370	12.80	
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	1,2,4-Trimethylbenzene	2060	ug/mL	0.2561	0.2785	0.0387	7.23	0.1022	9.71	0.2457	0.0341	6.80	0.0934	8.92	
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	1,3,5-Trimethylbenzene	856	ug/mL	0.1064	0.0346	0.0048	3.41	0.0391	3.83	0.0380	0.0053	3.65	0.0424	4.15	
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	Total TMBs	4616	ug/mL	0.5738	0.4082	0.0567	5.77	0.0746	7.19	0.6392	0.0887	7.32	0.1039	9.87	
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	Naphthalene	455	ug/mL	0.0566	0.0637	0.0088	7.35	0.1047	9.94	0.8298	0.1152	13.00	0.3820	31.75	
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	1-MethylNaphthalene	1290	ug/mL	0.1604	0.0666	0.0092	4.07	0.0482	4.71	0.1059	0.0147	5.52	0.0704	6.80	
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	2-MethylNaphthalene	1500	ug/mL	0.1865	0.1084	0.0150	5.10	0.0636	6.16	0.2656	0.0369	8.16	0.1229	11.57	
NRMRL	JP-5	6/1/1990	8/12/1997	BFT-401-3	Total Naphthalenes	3245	ug/mL	0.4034	0.2386	0.0331	5.16	0.0645	6.25	1.2013	0.1668	10.39	0.1917	17.44	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	Density	0.78	g/mL	0.0001											
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	Benzene	1.5	ug/mL	0.0002	0.0000	0.0000	0.00	0.0000	0.00	-0.0002	0.0000	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	Toluene	36.4	ug/mL	0.0047	0.0000	0.0000	0.00	0.0000	0.00	-0.0047	-0.0007	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	Ethylbenzene	327	ug/mL	0.0419	0.0000	0.0000	0.00	0.0000	0.00	-0.0419	-0.0060	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	m-Xylene	762	ug/mL	0.0977	0.0000	0.0000	0.00	0.0000	0.00	-0.0977	-0.0140	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	o-Xylene	788	ug/mL	0.1010	0.0000	0.0000	0.00	0.0000	0.00	-0.0809	-0.0116	-57.70	-0.2315	-26.05	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	p-Xylene	317	ug/mL	0.0406	0.0000	0.0000	0.00	0.0000	0.00	-0.0406	-0.0058	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	m,p-xylenes	1079	ug/mL	0.1383	0.0000	0.0000	0.00	0.0000	0.00	-0.1383	-0.0198	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	Total Xylenes (m,p, and o)	1867	ug/mL	0.2394	0.0000	0.0000	0.00	0.0000	0.00	-0.2192	-0.0315	-156.35	-0.3553	-42.66	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	Total BTEX	2231.9	ug/mL	0.2861	0.0000	0.0000	0.00	0.0000	0.00	-0.2660	-0.0382	-189.72	-0.3809	-46.36	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	B+T	37.9	ug/mL	0.0049											
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	E+X	2194	ug/mL	0.2813											
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	B/T	0.041208791		0.0000											
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	B/E	0.004587156		0.0000											
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	B/X	0.000803428		0.0000											
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	T/E	0.111314985		0.0000											
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	T/X	0.019496518		0.0000											
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	E/X	0.175147295		0.0000											
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	(B+T)/(E+X)	0.017274385		0.0000											
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	1,2,3-Trimethylbenzene	2390	ug/mL	0.3064	0.0000	0.0000	0.00	0.0000	0.00	0.2605	0.0374	6.59	0.0883	8.45	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	1,2,4-Trimethylbenzene	4170	ug/mL	0.5346	0.0000	0.0000	0.00	0.0000	0.00	-0.0328	-0.0047	-0.94	-0.0091	-0.91	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	1,3,5-Trimethylbenzene	1100	ug/mL	0.1410	0.0000	0.0000	0.00	0.0000	0.00	0.0034	0.0005	0.33	0.0034	0.34	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	Total TMBs	7660	ug/mL	0.9821	0.0000	0.0000	0.00	0.0000	0.00	0.2310	0.0331	2.73	0.0303	2.99	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	Naphthalene	938	ug/mL	0.1203	0.0000	0.0000	0.00	0.0000	0.00	0.7661	0.1099	12.40	0.2866	24.92	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	1-MethylNaphthalene	1770	ug/mL	0.2269	0.0000	0.0000	0.00	0.0000	0.00	0.0393	0.0056	2.12	0.0229	2.27	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	2-MethylNaphthalene	2300	ug/mL	0.2949	0.0000	0.0000	0.00	0.0000	0.00	0.1572	0.0226	4.99	0.0613	5.95	
NRMRL	JP-5	6/1/1990	5/19/1997	Fresh JP-5	Total Naphthalenes	5008	ug/mL	0.6421	0.0000	0.0000	0.00	0.0000	0.00	0.9627	0.1381	8.61	0.1314	12.32	
					-														

Cecil Field

									Beaufort MCAS Fresh JP-5			Hughes et al., 1984							
Lab	Fuel	Spill	Sample							Lin	ear	Expor	ential		Lin	ear	Expor	nential	
Code	Туре	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	C° - C	k	%Red./yr	k	%Red./yr	C° - C	k	%Red./yr	k	%Red./yr	
EAL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	Benzene	23500	ug/L	0.002941176	-0.0027	-0.0002	-89.46	-0.1707	-18.61	-0.0029	-0.0002	#DIV/0!	#DIV/0!	#DIV/0!	
EAL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	Toluene	122000	ug/L	0.015269086	-0.0106	-0.0007	-14.22	-0.0742	-7.70	-0.0153	-0.0010	#DIV/0!	#DIV/0!	#DIV/0!	
EAL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	Ethylbenzene	3200000	ug/L	0.400500626	-0.3586	-0.0224	-53.53	-0.1412	-15.17	-0.4005	-0.0251	#DIV/0!	#DIV/0!	#DIV/0!	
EAL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	Total Xylenes (m,p, and o)	7300000	ug/L	0.913642053	-0.6743	-0.0422	-17.63	-0.0838	-8.74	-0.8935	-0.0559	-277.96	-0.2388	-26.97	
EAL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	Total BTEX	10645500	ug/L	1.332352941	-1.0462	-0.0655	-22.88	-0.0963	-10.11	-1.3122	-0.0821	-408.22	-0.2624	-30.01	
EAL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	B+T	145500	ug/L	0.018210263											
EAL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	E+X	10500000	ug/L	1.314142678											
EAL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	B/T	0.192622951	-	2.4108E-08											
EAL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	B/E	0.00734375		9.19118E-10											
EAL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	B/X	0.003219178		4.02901E-10											
EAL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	T/E	0.038125		4.77159E-09											
EAL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	T/X	0.016712329		2.09166E-09											
EAL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	E/X	0.438356164		5.48631E-08											
EAL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	(B+T)/(E+X)	0.013857143		1.73431E-09											
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	Density	0.799	g/mL	0.0001											
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	Benzene	23.5	ug/mL	0.002941176	-0.0027	-0.0002	-89.46	-0.1707	-18.61	-0.0029	-0.0002	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	Toluene	122	ug/mL	0.015269086	-0.0106	-0.0007	-14.22	-0.0742	-7.70	-0.0153	-0.0010	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	Ethylbenzene	2520	ug/mL	0.315394243	-0.2735	-0.0171	-40.83	-0.1263	-13.46	-0.3154	-0.0197	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	m-Xylene	2800	ug/mL	0.350438048	-0.2527	-0.0158	-16.19	-0.0799	-8.32	-0.3504	-0.0219	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	o-Xylene	317	ug/mL	0.039674593	0.0614	0.0038	3.80	0.0585	5.68	-0.0196	-0.0012	-6.08	-0.0425	-4.34	
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	p-Xylene	1670	ug/mL	0.209011264	-0.1684	-0.0105	-25.93	-0.1025	-10.79	-0.2090	-0.0131	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	m,p-xylenes	4470	ug/mL	0.5594	-0.4211	-0.0264	-19.05	-0.0875	-9.14	-0.5594	-0.0350	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	Total Xylenes (m,p, and o)	4787	ug/mL	0.599123905	-0.3598	-0.0225	-9.41	-0.0574	-5.91	-0.5790	-0.0362	-180.12	-0.2124	-23.66	
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	Total BTEX	7452.5	ug/mL	0.932728411	-0.6466	-0.0405	-14.14	-0.0740	-7.68	-0.9126	-0.0571	-283.90	-0.2401	-27.14	
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	B+T	145.5	ug/mL	0.018210263											
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	E+X	7307	ug/mL	0.914518148											
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	B/T	0.192622951		2.4108E-05											
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	B/E	0.009325397		1.16713E-06											
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	B/X	0.004909129		6.14409E-07											
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	T/E	0.048412698		6.05916E-06											
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	T/X	0.02548569		3.1897E-06											
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	E/X	0.526425736		6.58856E-05											
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	(B+T)/(E+X)	0.019912413		2.49217E-06											
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	1,2,3-Trimethylbenzene	1860	ug/mL	0.232790989	0.0736	0.0046	1.50	0.0172	1.71	0.3341	0.0209	3.69	0.0557	5.42	
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	1,2,4-Trimethylbenzene	4000	ug/mL	0.500625782	0.0340	0.0021	0.40	0.0041	0.41	0.0011	0.0001	0.01	0.0001	0.01	
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	1,3,5-Trimethylbenzene	1400	ug/mL	0.175219024	-0.0342	-0.0021	-1.52	-0.0136	-1.37	-0.0308	-0.0019	-1.34	-0.0121	-1.22	
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	Total TMBs	7260	ug/mL	0.9086	0.0734	0.0046	0.47	0.0049	0.49	0.3044	0.0190	1.57	0.0181	1.79	
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	Naphthalene	1880	ug/mL	0.235294118	-0.1150	-0.0072	-5.99	-0.0420	-4.29	0.6511	0.0407	4.60	0.0830	7.97	
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	1-MethylNaphthalene	2290	ug/mL	0.28660826	-0.0597	-0.0037	-1.65	-0.0146	-1.47	-0.0203	-0.0013	-0.48	-0.0046	-0.46	
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	2-MethylNaphthalene	3320	ug/mL	0.415519399	-0.1206	-0.0076	-2.56	-0.0215	-2.17	0.0366	0.0023	0.51	0.0053	0.53	
NRMRL	JP-5	6/1/1981	5/20/1997	CEF-293-9FP	Total Naphthalenes	7490	ug/mL	0.9374	-0.2954	-0.0185	-2.88	-0.0237	-2.40	0.6673	0.0418	2.60	0.0336	3.31	

									Smith et al., 1981						Ma	yfield, 19	96		
Lab	Fuel	Spill	Sample							Lin	ear	Expon	ential		Lin	ear	Expor	nential	
Code	Type	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	C ₀ - C	k	%Red./yr	k	%Red./yr	C° - C	k	%Red./yr	k	%Red./yr	
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	Density	0.793	g/mL	0.0001	ĺ										
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	Benzene	0.025	ug/mL	0.0000	0.0000	0.0000	#DIV/0!	#DIV/0!	#DIV/0!	0.0321	0.1096	341.09	31.4846	100.00	
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	Toluene	0.025	ug/mL	0.0000	0.0000	0.0000	#DIV/0!	#DIV/0!	#DIV/0!	0.2083	0.7107	341.12	37.8600	100.00	
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	Ethylbenzene	63	ug/mL	0.0079	-0.0079	-0.0271	#DIV/0!	#DIV/0!	#DIV/0!	0.1432	0.4886	323.20	10.0496	100.00	
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	m-Xylene	441	ug/mL	0.0556	0.0044	0.0150	24.95	0.2591	22.82	-0.0556	-0.1897	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	o-Xylene	453	ug/mL	0.0571	0.0029	0.0098	16.35	0.1675	15.42	0.3536	1.2062	293.68	6.7292	99.88	
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	p-Xylene	111	ug/mL	0.0140	-0.0140	-0.0477	#DIV/0!	#DIV/0!	#DIV/0!	-0.0140	-0.0477	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	m,p-xylenes	552	ug/mL	0.0696	-0.0096	-0.0328	-54.63	-0.5067	-65.99	0.6506	2.2194	308.15	7.9709	99.97	
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	Total Xylenes (m,p, and o)	1005	ug/mL	0.1267	-0.0067	-0.0230	-19.14	-0.1862	-20.47	1.0042	3.4256	302.90	7.4662	99.94	
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	Total BTEX	1068.05	ug/mL	0.1347	-0.0147	-0.0501	-41.74	-0.3938	-48.26	1.3879	4.7345	310.95	8.2731	99.97	
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	B+T	0.05	ug/mL	0.0000											
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	E+X	1068	ug/mL	0.1347											
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	B/T	1		0.0001											
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	B/E	0.000396825		0.0000											
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	B/X	2.48756E-05		0.0000											
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	T/E	0.000396825		0.0000											
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	T/X	2.48756E-05		0.0000											
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	E/X	0.062686567		0.0000											
NRMRL	JP-8	4/1/1996	7/17/1996	SOURCE AREA	(B+T)/(E+X)	4.68165E-05		0.0000											

									Smith et al., 1981						May	field, 199	6		
Lab	Fuel	Spill	Sample							Lin	ear	Expon	ential		Line	ear	Expor	ential	
Code	Туре	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	C ₀ - C	k	%Red./yr	k	%Red./yr	C ₀ - C	k	%Red./yr	k	%Red./yr	
EAL	JP-8	12/1/1995	3/10/1998	SJ98-MW1S	Benzene	50000	ug/L	0.0061	-0.0061	-0.0027	#DIV/0!	#DIV/0!	#DIV/0!	0.0260	0.0114	35.61	0.7299	51.81	
EAL	JP-8	12/1/1995	3/10/1998	SJ98-MW1S	Toluene	630000	ug/L	0.0770	-0.0770	-0.0339	#DIV/0!	#DIV/0!	#DIV/0!	0.1313	0.0577	27.72	0.4376	35.44	
EAL	JP-8	12/1/1995	3/10/1998	SJ98-MW1S	Ethylbenzene	1100000	ug/L	0.1345	-0.1345	-0.0591	#DIV/0!	#DIV/0!	#DIV/0!	0.0167	0.0074	4.86	0.0515	5.02	
EAL	JP-8	12/1/1995	3/10/1998	SJ98-MW1S	o-Xylene	1400000	ug/L	0.1711	-0.1111	-0.0489	-81.46	-0.4610	-58.56	0.2396	0.1054	25.65	0.3849	31.95	
EAL	JP-8	12/1/1995	3/10/1998	SJ98-MW1S	m,p-Xylene	3100000	ug/L	0.3790	-0.3190	-0.1403	-233.79	-0.8105	-124.91	0.3413	0.1501	20.84	0.2824	24.60	
EAL	JP-8	12/1/1995	3/10/1998	SJ98-MW1S	Total Xylenes (m,p, and o)	4500000	ug/L	0.5501	-0.4301	-0.1892	-157.63	-0.6696	-95.35	0.5808	0.2554	22.58	0.3169	27.16	
EAL	JP-8	12/1/1995	3/10/1998	SJ98-MW1S	Total BTEX	6280000	ug/L	0.7677	-0.6477	-0.2848	-237.37	-0.8162	-126.18	0.7549	0.3320	21.80	0.3011	26.00	
EAL	JP-8	12/1/1995	3/10/1998	SJ98-MW1S	B+T	680000	ug/L	0.0831											
EAL	JP-8	12/1/1995	3/10/1998	SJ98-MW1S	E+X	5600000	ug/L	0.6846											
EAL	JP-8	12/1/1995	3/10/1998	SJ98-MW1S	B/T	0.079365079		0.0000											
EAL	JP-8	12/1/1995	3/10/1998	SJ98-MW1S	B/E	0.045454545		0.0000											
EAL	JP-8	12/1/1995	3/10/1998	SJ98-MW1S	B/X	0.011111111		0.0000											
EAL	JP-8	12/1/1995	3/10/1998	SJ98-MW1S	T/E	0.572727273		0.0000											
EAL	JP-8	12/1/1995	3/10/1998	SJ98-MW1S	T/X	0.14		0.0000											
EAL	JP-8	12/1/1995	3/10/1998	SJ98-MW1S	E/X	0.24444444		0.0000											
EAL	JP-8	12/1/1995	3/10/1998	SJ98-MW1S	(B+T)/(E+X)	0.121428571		0.0000											
EAL	JP-8	12/1/1995	5/15/1997	SJMW1SFP	Benzene	150000	ug/L	0.0189	-0.0189	-0.0130	#DIV/0!	#DIV/0!	#DIV/0!	0.0132	0.0091	28.29	0.3645	30.54	
EAL	JP-8	12/1/1995	5/15/1997	SJMW1SFP	Toluene	1000000	ug/L	0.1261	-0.1261	-0.0867	#DIV/0!	#DIV/0!	#DIV/0!	0.0822	0.0565	27.13	0.3451	29.18	
EAL	JP-8	12/1/1995	5/15/1997	SJMW1SFP	Ethylbenzene	790000	ug/L	0.0996	-0.0996	-0.0685	#DIV/0!	#DIV/0!	#DIV/0!	0.0516	0.0354	23.45	0.2867	24.93	
EAL	JP-8	12/1/1995	5/15/1997	SJMW1SFP	Total Xylenes (m,p, and o)	5200000	ug/L	0.6557	-0.5357	-0.3683	-306.88	-1.1674	-221.35	0.4752	0.3267	28.88	0.3747	31.25	
EAL	JP-8	12/1/1995	5/15/1997	SJMW1SFP	Total BTEX	7140000	ug/L	0.9004	-0.7804	-0.5364	-447.02	-1.3853	-299.60	0.6222	0.4277	28.09	0.3611	30.31	
EAL	JP-8	12/1/1995	5/15/1997	SJMW1SFP	B+T	1150000	ug/L	0.1450											
EAL	JP-8	12/1/1995	5/15/1997	SJMW1SFP	E+X	5990000	ug/L	0.7554											
EAL	JP-8	12/1/1995	5/15/1997	SJMW1SFP	B/T	0.15		0.0000											
EAL	JP-8	12/1/1995	5/15/1997	SJMW1SFP	B/E	0.189873418		0.0000											
EAL	JP-8	12/1/1995	5/15/1997	SJMW1SFP	B/X	0.028846154		0.0000											
EAL	JP-8	12/1/1995	5/15/1997	SJMW1SFP	T/E	1.265822785		0.0000											
EAL	JP-8	12/1/1995	5/15/1997	SJMW1SFP	T/X	0.192307692		0.0000											
EAL	JP-8	12/1/1995	5/15/1997	SJMW1SFP	E/X	0.151923077		0.0000											
EAL	JP-8	12/1/1995	5/15/1997	SJMW1SFP	(B+T)/(E+X)	0.191986644		0.0000											
					-														
EAL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	Benzene	220000	ug/L	0.0284	-0.0284	-0.0195	#DIV/0!	#DIV/0!	#DIV/0!	0.0037	0.0026	7.95	0.0845	8.10	
EAL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	Toluene	1100000	ug/L	0.1421	-0.1421	-0.0977	#DIV/0!	#DIV/0!	#DIV/0!	0.0662	0.0455	21.85	0.2629	23.12	
EAL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	Ethylbenzene	1200000	ug/L	0.1550	-0.1550	-0.1066	#DIV/0!	#DIV/0!	#DIV/0!	-0.0038	-0.0026	-1.75	-0.0173	-1.74	
EAL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	Total Xylenes (m,p, and o)	6900000	ug/L	0.8915	-0.7715	-0.5303	-441.91	-1.3785	-296.88	0.2395	0.1646	14.56	0.1636	15.09	
EAL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	Total BTEX	9420000	ug/L	1.2171	-1.0971	-0.7541	-628.41	-1.5925	-391.58	0.3056	0.2100	13.79	0.1540	14.27	
EAL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	B+T	1320000	ug/L	0.1705											
EAL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	E+X	8100000	ug/L	1.0465											
EAL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	B/T	0.2		0.0000											
EAL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	B/E	0.183333333		0.0000											
EAL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	B/X	0.031884058		0.0000											
EAL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	T/E	0.916666667		0.0000											
EAL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	1/X	0.15942029		0.0000											
EAL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	E/X	0.173913043		0.0000											
EAL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	(B+1)/(E+X)	0.162962963		0.0000											

									Smith et al., 1981						Ма	yfield, 199	6		
Lab	Fuel	Spill	Sample	9						Lin	ear	Expon	ential		Lin	ear	Expor	ential	
Code	Type	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	C ₀ - C	k	%Red./yr	k.	%Red./yr	C C	k	%Red./yr	k.	%Red./yr	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	Density	0.812	a/mL	0.0001						- 0 -					
							5												
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	Benzene	48.9	ua/mL	0.0060	-0.0060	-0.0026	#DIV/0!	#DIV/0!	#DIV/0!	0.0261	0.0115	35.74	0.7365	52.12	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	Toluene	638	ua/mL	0.0786	-0.0786	-0.0346	#DIV/0!	#DIV/0!	#DIV/0!	0.1298	0.0571	27.39	0.4288	34.87	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	Ethylbenzene	850	ug/mL	0.1047	-0.1047	-0.0460	#DIV/0!	#DIV/0!	#DIV/0!	0.0465	0.0205	13.53	0.1617	14.93	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	m-Xylene	1510	ug/mL	0.1860	-0.1260	-0.0554	-92.32	-0.4975	-64.45	-0.1860	-0.0818	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	o-Xvlene	1110	ug/mL	0.1367	-0.0767	-0.0337	-56.22	-0.3621	-43.64	0.2740	0.1205	29.34	0.4838	38.36	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	p-Xvlene	592	ua/mL	0.0729	-0.0729	-0.0321	#DIV/0!	#DIV/0!	#DIV/0!	-0.0729	-0.0321	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	m.p-xvlenes	2102	ua/mL	0.2589	-0.1989	-0.0875	-145.76	-0.6429	-90.20	0.4614	0.2029	28.17	0.4500	36.24	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	Total Xylenes (m.p. and o)	3212	ua/mL	0.3956	-0.2756	-0.1212	-100.99	-0.5246	-68.97	0.7354	0.3234	28.59	0.4620	37.00	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	Total BTEX	4748.9	ua/mL	0.5848	-0.4648	-0.2044	-170.35	-0.6965	-100.67	0.9378	0.4124	27.08	0.4208	34.35	
NRMRI	JP-8	12/1/1995	3/10/199	8 S.198-MP2	B+T	686.9	ug/ml	0.0846	0.1010	0.2011		0.0000		0.0010	0	27.00	0.1200	0	
NRMRI	JP-8	12/1/1995	3/10/199	8 S.198-MP2	F+X	4062	ug/ml	0.5002											
NRMRI	IP-8	12/1/1995	3/10/199	8 S 198-MP2	B/T	0.076645768	ug/IIIE	0.0002											
NRMRI	JP-8	12/1/1995	3/10/199	8 S.198-MP2	B/F	0.057529412		0.0000											
NRMRI	JP-8	12/1/1995	3/10/199	8 S.198-MP2	B/X	0.015224159		0.0000											
NRMRI	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	T/E	0.750588235		0.0001											
NRMRI	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	T/X	0.198630137		0.0000											
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	E/X	0.264632628		0,0000											
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	(B+T)/(E+X)	0.16910389		0,0000											
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	1.2.3-Trimethylbenzene	2330	ua/mL	0,2869	-0.2869	-0.1262	#DIV/0!	#DIV/0!	#DIV/0!	0.3869	0.1701	25.25	0.3754	31.30	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	1.2.4-Trimethylbenzene	4380	ua/mL	0.5394	-0.2694	-0.1185	-43.88	-0.3043	-35.57	0.9249	0.4067	27.78	0.4392	35.54	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	1.3.5-Trimethylbenzene	1150	ug/mL	0.1416	-0.1416	-0.0623	#DIV/0!	#DIV/0!	#DIV/0!	0.3667	0.1613	31.72	0.5620	42.99	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	Total TMBs	7860	ug/mL	0.9680	-0.6980	-0.3069	-113.68	-0.5615	-75.33	1.6784	0.7381	27.89	0.4423	35.74	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	Naphthalene	879	ug/mL	0.1083	1.0317	0.4537	39.80	1.0353	64.49	0.1465	0.0644	25.29	0.3764	31.37	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	1-MethylNaphthalene	1290	ua/mL	0.1589	1.6811	0.7393	40.18	1.0772	65.94	0.2685	0.1181	27.63	0.4352	35.29	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	2-MethylNaphthalene	1900	ua/mL	0.2340	1.2260	0.5391	36.93	0.8052	55.30	0.1208	0.0531	14.97	0.1830	16.72	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MP2	Total Naphthalenes	4069	ua/mL	0.5011	3.9389	1.7322	39.01	0.9594	61.69	0.5358	0.2356	22.72	0.3198	27.37	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	Density	0.818	g/mL	0.0001											
							0												
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	Benzene	47.2	ug/mL	0.0058	-0.0058	-0.0025	#DIV/0!	#DIV/0!	#DIV/0!	0.0264	0.0116	36.08	0.7553	53.01	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	Toluene	602	ug/mL	0.0736	-0.0736	-0.0324	#DIV/0!	#DIV/0!	#DIV/0!	0.1347	0.0593	28.44	0.4576	36.72	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	Ethylbenzene	800	ug/mL	0.0978	-0.0978	-0.0430	#DIV/0!	#DIV/0!	#DIV/0!	0.0534	0.0235	15.53	0.1916	17.43	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	m-Xylene	1420	ug/mL	0.1736	-0.1136	-0.0500	-83.26	-0.4672	-59.55	-0.1736	-0.0763	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	o-Xylene	1040	ug/mL	0.1271	-0.0671	-0.0295	-49.21	-0.3302	-39.13	0.2836	0.1247	30.36	0.5157	40.29	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	p-Xylene	580	ug/mL	0.0709	-0.0709	-0.0312	#DIV/0!	#DIV/0!	#DIV/0!	-0.0709	-0.0312	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	m,p-xylenes	2000	ug/mL	0.2445	-0.1845	-0.0811	-135.23	-0.6178	-85.48	0.4757	0.2092	29.05	0.4751	37.82	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	Total Xylenes (m,p, and o)	3040	ug/mL	0.3716	-0.2516	-0.1107	-92.22	-0.4971	-64.40	0.7593	0.3339	29.53	0.4894	38.70	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	Total BTEX	4489.2	ug/mL	0.5488	-0.4288	-0.1886	-157.14	-0.6685	-95.14	0.9738	0.4282	28.13	0.4488	36.16	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	B+T	649.2	ug/mL	0.0794											
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	E+X	3840	ug/mL	0.4694											
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	B/T	0.078405316		0.0000											
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	B/E	0.059		0.0000											
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	B/X	0.015526316		0.0000											
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	T/E	0.7525		0.0001											
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	T/X	0.198026316		0.0000											-
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	E/X	0.263157895		0.0000											
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	(B+T)/(E+X)	0.1690625		0.0000											-
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	1,2,3-Trimethylbenzene	2240	ug/mL	0.2738	-0.2738	-0.1204	#DIV/0!	#DIV/0!	#DIV/0!	0.4000	0.1759	26.10	0.3960	32.70	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	1,2,4-Trimethylbenzene	4190	ug/mL	0.5122	0.0224	0.0098	1.84	0.0188	1.86	-0.0104	-0.0046	-0.92	-0.0091	-0.91	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	1,3,5-Trimethylbenzene	1090	ug/mL	0.1333	0.0078	0.0034	2.42	0.0249	2.46	0.0111	0.0049	3.39	0.0353	3.47	-
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	Total TMBs	7520	ug/mL	0.9193	0.0627	0.0276	2.81	0.0290	2.86	0.2937	0.1292	10.65	0.1219	11.48	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	Naphthalene	844	ug/mL	0.1032	0.0171	0.0075	6.25	0.0674	6.51	0.7832	0.3444	38.86	0.9458	61.16	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	1-MethylNaphthalene	1250	ug/mL	0.1528	0.0741	0.0326	14.36	0.1739	15.96	0.1135	0.0499	18.74	0.2442	21.67	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	2-MethylNaphthalene	1830	ug/mL	0.2237	0.0712	0.0313	10.61	0.1214	11.44	0.2284	0.1004	22.21	0.3094	26.61	
NRMRL	JP-8	12/1/1995	3/10/199	8 SJ98-MW1S	Total Naphthalenes	3924	ug/mL	0.4797	0.1623	0.0714	11.12	0.1282	12.03	1.1250	0.4947	30.83	0.5310	41.20	

								Smith et al., 1981						Ma	yfield, 199	96		
Lab	Fuel	Spill	Sample						Lin	ear	Expon	ential		Lin	ear	Expoi	nential	
Code	Туре	Date	Date Locid	Analyte	Results	Units	Mass Fraction	C ₀ - C	k	%Red./yr	k	%Red./yr	C ₀ - C	k	%Red./yr	k	%Red./yr	
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	Density	0.793	g/mL	0.0001											
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	Benzene	246	ug/mL	0.0310	-0.0310	-0.0505	#DIV/0!	#DIV/0!	#DIV/0!	0.0011	0.0018	5.68	0.0579	5.62	
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	Toluene	1631	ug/mL	0.2057	-0.2057	-0.3351	#DIV/0!	#DIV/0!	#DIV/0!	0.0027	0.0043	2.08	0.0209	2.07	
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	Ethylbenzene	1239	ug/mL	0.1562	-0.1562	-0.2546	#DIV/0!	#DIV/0!	#DIV/0!	-0.0051	-0.0082	-5.44	-0.0536	-5.50	
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	m-Xylene	4042	ug/mL	0.5097	-0.4497	-0.7328	-1221.31	-3.4862	-3166.27	-0.5097	-0.8306	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	o-Xylene	2313	ug/mL	0.2917	-0.2317	-0.3775	-629.18	-2.5767	-1215.34	0.1190	0.1940	47.23	0.5577	42.75	
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	p-Xylene	1172	ug/mL	0.1478	-0.1478	-0.2408	#DIV/0!	#DIV/0!	#DIV/0!	-0.1478	-0.2408	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	m,p-xylenes	5214	ug/mL	0.6575	-0.5975	-0.9736	-1622.68	-3.9011	-4845.72	0.0627	0.1022	14.19	0.1485	13.80	
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	Total Xylenes (m,p, and o)	7527	ug/mL	0.9492	-0.8292	-1.3511	-1125.93	-3.3699	-2807.58	0.1818	0.2962	26.19	0.2855	24.84	
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	Total BTEX	10643	ug/mL	1.3421	-1.2221	-1.9914	-1659.50	-3.9344	-5012.95	0.1805	0.2941	19.32	0.2056	18.58	
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	B+T	1877	ug/mL	0.2367											
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	E+X	8766	ug/mL	1.1054											
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	B/T	0.150827713		0.0000											
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	B/E	0.198547215		0.0000											
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	B/X	0.032682344		0.0000											
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	I/E	1.316384181		0.0002											
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	1/X	0.216686595		0.0000											
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	E/X	0.164607413		0.0000											
NRMRL	JP-8	12/1/1995	7/12/1996 SJMW1S	(B+T)/(E+X)	0.214122747		0.0000											
		10/1/1005	5/45/4007 0 N N 405D	D 11	0.700		0.0004											
NRMRL	JP-8	12/1/1995	5/15/1997 SJMW1SFP	Density	0.793	g/mL	0.0001											
		10/1/1005	5/45/4007 0 N.W.405D		10.1		0.0045	0.0045	0.0100	"DIV (/01	"DI //OL	"DIV (/01	0.0077	0.0050	40.40	0.4070		
NRMRL	JP-8	12/1/1995	5/15/1997 SJMW1SFP	Benzene	194	ug/mL	0.0245	-0.0245	-0.0168	#DIV/0!	#DIV/0!	#DIV/0!	0.0077	0.0053	16.42	0.1876	17.11	
NRMRL	JP-8	12/1/1995	5/15/1997 SJMW1SFP	I oluene	1030	ug/mL	0.1299	-0.1299	-0.0893	#DIV/0!	#DIV/0!	#DIV/0!	0.0784	0.0539	25.88	0.3248	27.73	
NRIVIRL	JP-8	12/1/1995	5/15/1997 SJWW15FP	Ethylbenzene	1170	ug/mL	0.1475	-0.1475	-0.1014	#DIV/0!	#DIV/0!	#DIV/0!	0.0036	0.0025	1.00	0.0168	1.67	
NRIVIRL	JP-8	12/1/1995	5/15/1997 SJWW15FP	m-Xylene	3120	ug/mL	0.3934	-0.3334	-0.2292	-382.00	-1.2927	-204.20	-0.3934	-0.2704	#DIV/0!	#DIV/0!	#DIV/0!	
NRIVIRL		12/1/1995	5/15/1997 SJIVIVISEP	0-Aylene	1110	ug/mL	0.2219	-0.1619	-0.1113	-105.55	-0.6991	-145.75 #DIV//01	0.1000	0.1296	31.59 #DIV/01	#DIV//01	34.50	
NRIVIRL		12/1/1995	5/15/1997 SJIVIVISEP	p-xylerie	1110	ug/mL	0.1400	-0.1400	-0.0962	#DIV/0!	#DIV/0!	#DIV/0!	-0.1400	-0.0962	#DIV/0!	#DIV/0!	#DIV/0!	
NEMEL	JP-0	12/1/1995	5/15/1997 SJIVIVISEP	Total Xylonas (m.n. and a)	4230	ug/mL	0.5334	-0.4734	-0.3234	-542.30	-1.5019	-349.02	0.1000	0.1204	17.00	0.2004	24.22	
NDMDI		12/1/1995	5/15/1997 SJWW13FF	Total RTEX	0294	ug/mL	1.0572	-0.0354	-0.4307	-303.95	-1.2040	246.25	0.3750	0.2302	22.03	0.2774	24.23	
NEMEL	JF=0 IP_8	12/1/1995	5/15/1997 SIMWISFF	RT	1224	ug/mL	0.1544	-0.9373	-0.0442	-550.67	-1.4957	-340.23	0.4034	0.3199	21.01	0.2507	22.10	
NEME	IP-8	12/1/1995	5/15/1007 SIMWISED	E+Y	7160	ug/mL	0.1344											
NRMRI	IP-8	12/1/1995	5/15/1997 SIMW1SFP	B/T	0 188349515	ug/IIIL	0.0000											
NRMRI	IP-8	12/1/1995	5/15/1997 SIMW1SEP	B/F	0.165811966		0.0000											
NRMRI	JP-8	12/1/1995	5/15/1997 SJMW1SFP	B/X	0.032387312		0.0000											
NRMRI	JP-8	12/1/1995	5/15/1997 SJMW1SEP	T/E	0.88034188		0.0001											
NRMRI	JP-8	12/1/1995	5/15/1997 SJMW1SFP	T/X	0.171953255		0.0000											
NRMRL	JP-8	12/1/1995	5/15/1997 SJMW1SFP	E/X	0.195325543		0.0000											
NRMRI	JP-8	12/1/1995	5/15/1997 SJMW1SFP	(B+T)/(E+X)	0.170949721		0.0000											
NRMRL	JP-8	12/1/1995	5/15/1997 SJMW1SFP	1.2.3-Trimethylbenzene	2720	ua/mL	0,3430	-0.3430	-0.2358	#DIV/0!	#DIV/0!	#DIV/0!	0.3308	0.2274	33.75	0.4641	37.13	
NRMRL	JP-8	12/1/1995	5/15/1997 SJMW1SFP	1.2.4-Trimethylbenzene	5590	ua/mL	0.7049	-0.4349	-0.2990	-110,72	-0.6597	-93.41	0.7594	0.5220	35.65	0.5025	39.50	
NRMRL	JP-8	12/1/1995	5/15/1997 SJMW1SFP	1,3,5-Trimethylbenzene	2100	ug/mL	0.2648	-0.2648	-0.1820	#DIV/0!	#DIV/0!	#DIV/0!	0.2435	0.1674	32.93	0.4482	36.12	
NRMRL	JP-8	12/1/1995	5/15/1997 SJMW1SFP	Total TMBs	10410	ua/mL	1,3127	-1.0427	-0.7168	-265.47	-1.0871	-196.55	1.3337	0.9168	34.64	0.4819	38.24	
NRMRL	JP-8	12/1/1995	5/15/1997 SJMW1SFP	Naphthalene	1280	ua/mL	0,1614	0.9786	0.6727	59,01	1.3437	73.91	0.0933	0.0642	25.19	0.3137	26.93	
NRMRL	JP-8	12/1/1995	5/15/1997 SJMW1SFP	1-MethylNaphthalene	1510	ug/mL	0.1904	1.6496	1.1339	61.62	1.5592	78.97	0.2370	0.1629	38.11	0.5557	42.63	
NRMRL	JP-8	12/1/1995	5/15/1997 SJMW1SFP	2-MethylNaphthalene	2290	ua/mL	0,2888	1,1712	0.8051	55,14	1,1139	67.17	0.0660	0.0454	12.79	0.1415	13.19	
NRMRL	JP-8	12/1/1995	5/15/1997 SJMW1SFP	Total Naphthalenes	5080	ug/mL	0.6406	3.7994	2.6116	58.82	1.3308	73.57	0.3963	0.2724	26.27	0.3310	28.18	
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									Smith et al., 1981					Ma	yfield, 199	6			
Lab	Fuel	Spill	Sample							Line	ear	Expone	ential		Lin	ear	Expor	nential	
Code	Туре	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	C ₀ - C	k	%Red./yr	k	%Red./yr	C ₀ - C	k	%Red./yr	k	%Red./yr	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	Density	0.774	g/mL	0.0001											
							-												
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	Benzene	190	ug/mL	0.0245	-0.0245	-0.0169	#DIV/0!	#DIV/0!	#DIV/0!	0.0076	0.0052	16.24	0.1853	16.91	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	Toluene	1030	ug/mL	0.1331	-0.1331	-0.0915	#DIV/0!	#DIV/0!	#DIV/0!	0.0753	0.0517	24.83	0.3081	26.52	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	Ethylbenzene	1160	ug/mL	0.1499	-0.1499	-0.1030	#DIV/0!	#DIV/0!	#DIV/0!	0.0013	0.0009	0.60	0.0060	0.60	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	m-Xylene	3160	ug/mL	0.4083	-0.3483	-0.2394	-398.99	-1.3181	-273.64	-0.4083	-0.2806	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	o-Xylene	1770	ug/mL	0.2287	-0.1687	-0.1159	-193.25	-0.9197	-150.86	0.1820	0.1251	30.47	0.4025	33.14	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	p-Xylene	1120	ug/mL	0.1447	-0.1447	-0.0995	#DIV/0!	#DIV/0!	#DIV/0!	-0.1447	-0.0995	#DIV/0!	#DIV/0!	#DIV/0!	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	m,p-xylenes	4280	ug/mL	0.5530	-0.4930	-0.3389	-564.77	-1.5266	-360.27	0.1673	0.1150	15.96	0.1817	16.61	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	Total Xylenes (m,p, and o)	6050	ug/mL	0.7817	-0.6617	-0.4548	-379.01	-1.2881	-262.59	0.3493	0.2401	21.23	0.2539	22.42	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	Total BTEX	8430	ug/mL	1.0891	-0.9691	-0.6662	-555.15	-1.5161	-355.46	0.4335	0.2980	19.57	0.2303	20.57	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	B+T	1220	ug/mL	0.1576											
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	E+X	7210	ug/mL	0.9315											
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	B/T	0.184466019		0.0000											
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	B/E	0.163793103		0.0000											
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	B/X	0.031404959		0.0000											
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	T/E	0.887931034		0.0001											
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	T/X	0.170247934		0.0000											
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	E/X	0.191735537		0.0000											
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	(B+T)/(E+X)	0.169209431		0.0000											
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	1,2,3-Trimethylbenzene	2750	ug/mL	0.3553	-0.3553	-0.2442	#DIV/0!	#DIV/0!	#DIV/0!	0.3185	0.2189	32.49	0.4399	35.59	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	1,2,4-Trimethylbenzene	5650	ug/mL	0.7300	-0.4600	-0.3162	-117.10	-0.6837	-98.11	0.7343	0.5048	34.47	0.4785	38.03	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	1,3,5-Trimethylbenzene	2130	ug/mL	0.2752	-0.2752	-0.1892	#DIV/0!	#DIV/0!	#DIV/0!	0.2331	0.1603	31.53	0.4218	34.41	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	Total TMBs	10530	ug/mL	1.3605	-1.0905	-0.7496	-277.62	-1.1116	-203.92	1.2860	0.8839	33.40	0.4574	36.71	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	Naphthalene	1290	ug/mL	0.1667	0.9733	0.6691	58.69	1.3217	73.33	0.0881	0.0606	23.77	0.2917	25.30	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	1-MethylNaphthalene	1490	ug/mL	0.1925	1.6475	1.1325	61.55	1.5517	78.81	0.2349	0.1614	37.78	0.5482	42.20	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	2-MethylNaphthalene	2320	ug/mL	0.2997	1.1603	0.7975	54.63	1.0883	66.32	0.0550	0.0378	10.66	0.1158	10.94	
NRMRL	JP-8	12/1/1995	5/15/1997	SJMW2SFP	Total Naphthalenes	5100	ug/mL	0.6589	3.7811	2.5991	58.54	1.3114	73.06	0.3780	0.2598	25.06	0.3117	26.78	

Linear (Zero Order) Assumption

linear equation $C = C_0 - kt$

summary statistics presented as follows

ana	lyte
k	Co
sek	seCo
r ²	seC
F stat.	df
SSregr	SSresid

 $\label{eq:k} \begin{array}{l} k = zero \mbox{ order weathering rate; } k = dC/dt \mbox{ or slope} \\ sek = slope \mbox{ standard error value} \\ Co = intercept \mbox{ or initial analyte concentration as calculated by regression analysis} \\ seCo = standard error value for the constant C_o \\ r^{Z} = coefficient \mbox{ of determination} \\ seC = standard error value \mbox{ of the estimated concentration C (i.e., a "standard deviation" for the regression line)} \\ F \mbox{ stat. = } F \mbox{ statistic or } F\mbox{ -observed value} \\ df = \mbox{ degrees of freedom} \\ ss_{regr} = \mbox{ the regression sum of squares} \end{array}$

ss_{resid} = the residual sum of squares

Exponential (1st Order) Assumption

exponential equation $C = C_o e^{-kt}$

summary statistics presented as follows

ana	alyte
m	Co
sek	seCo
r ²	seC
F stat.	df
ss _{regr}	SSresid
	In C _o

- m = coefficient for statistics equation shown; note $m = e^{-\kappa t}$, therefore, $\ln m = -k$
- sek = standard error value for the exponential rate constant k
- Co = intercept or initial analyte concentration as calculated by regression analysis
- $seCo = standard error value for the constant C_o; compare to In C_o$
 - r^2 = coefficient of determination
- seC = standard error value of the estimated concentration C (i.e., a "standard deviation" for the regression line)
- F stat. = F statistic or F-observed value
 - df = degrees of freedom
- ss_{regr} = the regression sum of squares
- ss_{resid} = the residual sum of squares
- $\ln C_o =$ natural log of Co for comparing to seCo

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Evergreen Analytical Data

Sample Location MW-1S	Date 12/14/1995 5/15/1997 3/10/1998	Time (yrs) 0.00 1.42 2.24	Benzene 0.0321 0.0237 0.0061	Toluene 0.2083 0.1341 0.0770	Ethylbenzene 0.1512 0.1273 0.1345	Total Xylenes 1.1310 0.7736 0.5501	Total BTEX 1.5226 1.0587 average of 2 samples 0.7677
linear Co Predicted C - latest linear rate constant average yearly redu	sample date (k) (slope) uction (%)		0.0341 0.0094 0.0110 32.31	0.2105 0.0807 0.0580 27.54	0.1480 0.1290 0.0085 5.74	1.1336 0.5546 0.2586 22.82	1.5261 0.7738 0.3361 22.02

linear

summary stats

Benz	ene	Tolu	ene	Ethylber	nzene	Total >	Kylenes	Total	BTEX
-0.0110	0.0341	-0.0580	0.2105	-0.0085	0.1480	-0.2586	1.1336	-0.3361	1.5261
0.0040	0.0062	0.0046	0.0070	0.0067	0.0102	0.0055	0.0084	0.0074	0.0113
0.8809	0.0065	0.9939	0.0073	0.6182	0.0107	0.9995	0.0088	0.9995	0.0119
7	1	162	1	2	1	2212	1	2054	1
0.0003	0.0000	0.0086	0.0001	0.0002	0.0001	0.1716	0.0001	0.2898	0.0001

exponential	Benzene	Toluene	Ethylbenzene	Total Xylenes	Total BTEX
Co	0.0384	0.2180	0.1477	1.1519	1.5485
exponential rate constant (k)	0.6834	0.4297	0.0599	0.3159	0.3004
% reduction/year	49.51	34.93	5.82	27.09	25.95

exponential

Benz	ene	Tolu	ene	Ethylben	izene	Total	Xylenes	Total	BTEX
0.5049	0.0384	0.6507	0.2180	0.9418	0.1477	0.7291	1.1519	0.7405	1.5485
0.3760	0.5754	0.0960	0.1468	0.0491	0.0751	0.0389	0.0595	0.0357	0.0546
0.7676	0.6023	0.9525	0.1537	0.5984	0.0786	0.9851	0.0623	0.9861	0.0571
3	1	20	1	1	1	66	1	71	1
1.1979	0.3627	0.4737	0.0236	0.0092	0.0062	0.2561	0.0039	0.2315	0.0033
	-3.2597		-1.5232		-1.9124		0.1414		0.4373

SEYMOUR JOHNSON AFB, BLDG 4522 (continued) NRMRL Data

Sample Location MW-1S	Date 12/14/1995 7/12/1996 5/15/1997 3/10/1998	Time (yrs) E 0.00 0.58 1.42 2.24	Benzene 0.0321 0.0310 0.0245 0.0058	Toluene 0.2083 0.2057 0.1315 0.0736	Ethylbenzene 0.1512 0.1562 0.1487 0.0978	o-Xylene 0.4107 0.2917 0.2253 0.1271	m,p-xylenes 0.7202 0.6575 0.5432 0.2445	Total Xylenes 1.1310 0.9492 0.7685 0.3716	Total BTEX 1.5226 1.3421 1.0732 a 0.5488	average of 2 samples
linear Co Predicted C - latest s linear rate constant average yearly redu	sample date (k) (slope) ction (%)		0.0356 0.0097 0.0116 32.54	0.2231 0.0787 0.0645 28.91	0.1629 0.1113 0.0231 14.16	0.3905 0.1225 0.1197 30.66	0.7605 0.2972 0.2070 27.22	1.1510 0.4197 0.3267 28.38	1.5726 0.6194 0.4259 27.08	

linear

summary stats

Benz	ene	Tolu	ene	Ethylber	nzene	Total >	(ylenes	Total BTEX		
-0.0116	0.0356	-0.0645	0.2231	-0.0231	0.1629	-0.3267	1.1510	-0.4259	1.5726	
0.0032	0.0044	0.0105	0.0143	0.0111	0.0150	0.0406	0.0551	0.0571	0.0775	
0.8651	0.0055	0.9493	0.0179	0.6839	0.0188	0.9701	0.0688	0.9653	0.0968	
13	2	37	2	4	2	65	2	56	2	
0.0004	0.0001	0.0120	0.0006	0.0015	0.0007	0.3067	0.0095	0.5211	0.0187	

exponential	Benzene	Toluene	Ethylbenzene	m-Xylene	o-Xylene	p-Xylene	m,p-xylenes	Total Xylenes	Total BTEX
Со	0.0421	0.2379	0.1658	#VALUE!	0.4107	#VALUE!	0.8182	1.2317	1.6719
exponential rate constant (k)	0.7331	0.4827	0.1864	#VALUE!	0.4993	#VALUE!	0.4636	0.4759	0.4416
% reduction/year	51.96	38.29	17.00	#VALUE!	39.31	#VALUE!	37.10	37.87	35.70

exponential

Benz	ene	Tolu	ene	Ethylben	zene	Total	Xylenes	Total	BTEX
0.4804	0.0421	0.6171	0.2379	0.8300	0.1658	0.6213	1.2317	0.6430	1.6719
0.2817	0.3821	0.0956	0.1297	0.0908	0.1231	0.1077	0.1461	0.1017	0.1379
0.7720	0.4776	0.9273	0.1621	0.6783	0.1538	0.9071	0.1826	0.9042	0.1723
7	2	25	2	4	2	20	2	19	2
1.5441	0.4561	0.6695	0.0525	0.0998	0.0473	0.6509	0.0667	0.5604	0.0594
	-3.1674		-1.4359		-1.7967		0.2084		0.5140

														Mid-Ra	nge (Potter	r, 1988),		-
								î	Ghasse	mi <i>et al</i> ., 1	984		AD L	ittle, Inc. (1	987), & Sig	sby et al. (1987)	
Lab	Fuel	Spill	Sample						Linea	ar	Expor	nential		Lin	ear	Expor	iential	
Code	Туре	Date	Date	Locid	Analyte	Results Units	Mass Fraction	C _o - C	<u>k</u> '	%Red./yr	k	%Red./yr	C ₀ - C	K	%Red./yr	K	%Red./yr	
EAL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	Benzene	14000000 ug/L	1.9391	-0.4391	-0.0621	-4.14	-0.0363	-3.70	0.7928	0.1122	4.11	0.0485	4.74	
EAL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP		52000000 ug/L	7.2022	-1.3022	-0.1843	-3.12	-0.0282	-2.86	5.3097	0.7515	6.01	0.0782	7.52	
	Gasoline	6/1/1990	6/23/1997	MW 240 6EP	Ethylbenzene Total Xulanas (m.n. and a)	57000000 ug/L	7 9047	-0.3620	-0.0512	-3.94	-0.0348	-3.54	0.0187	0.0027	0.16	0.0016	0.16	
	Gasoline	6/1/1990	6/23/1997	MW 240 6EP	Total RTEX	125000000 ug/L	19 6091	-1.9947	-0.2623	-4.70	-0.0412	-4.21	-0.3400 5 7722	-0.0492	-0.03	-0.0064	-0.64	
	Gasoline	6/1/1990	6/23/1997	MW 240 6EP		6600000 ug/L	0 1412	-4.0901	-0.5800	-3.97	-0.0350	-3.50	5.1155	0.0171	3.34	0.0301	3.74	
	Gasoline	6/1/1990	6/23/1997	MW 240 6EP		6000000 ug/L	9.1413											
FAL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	B/T	0.269230769	9.5508											
FAI	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	B/F	1 166666667	0.0000											
EAL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	B/X	0.245614035	0.0000											
EAL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	T/E	4.3333333333	0.0000											
EAL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	T/X	0.912280702	0.0000											
EAL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	E/X	0.210526316	0.0000											-
EAL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	(B+T)/(E+X)	0.956521739	0.0000											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	Density	0.722 g/mL	0.0001											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	Benzene	9450 ug/mL	1.3089	0.1911	0.0271	1.80	0.0193	1.91	1.4230	0.2014	7.37	0.1041	9.89	
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	Toluene	40600 ug/mL	5.6233	0.2767	0.0392	0.66	0.0068	0.68	6.8887	0.9749	7.79	0.1132	10.70	
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	Ethylbenzene	13300 ug/mL	1.8421	-0.5421	-0.0767	-5.90	-0.0493	-5.06	-0.1613	-0.0228	-1.36	-0.0130	-1.31	
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	m-Xylene	23500 ug/mL	3.2548											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	o-Xylene	11000 ug/mL	1.5235											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	p-Xylene	9210 ug/mL	1.2756											
NRIVIRL	Gasoline	6/1/1990	6/23/1997	MW 240 6FP	Total Xulance (m.n. and e)	32710 ug/mL	4.5305	0.15.40	0.0010	0.27	0.0026	0.27	1 4000	0.0110	2.00	0.0212	2.07	
NEME	Gasoline	6/1/1990	6/23/1997	MW 240 6EP	Total RTEX	437 10 ug/mL	14 9292	-0.1540	-0.0216	-0.37	-0.0030	-0.37	0.6421	1 2649	2.00	0.0312	5.07	
	Gasoline	6/1/1990	6/23/1997	MW 240 6FP		107060 ug/mL	14.0203	-0.2263	-0.0323	-0.22	-0.0022	-0.22	9.0431	1.3040	5.56	0.0709	0.04	
NRMRI	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	B+1 E+Y	57010 ug/mL	7 8961											
	Gasoline	6/1/1000	6/23/1997	MW-349-6FP	B/T	0.232758621	0.0000											
NRMRI	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	B/F	0.710526316	0.0000											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	B/X	0.216197666	0.0000											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	T/E	3.052631579	0.0004											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	T/X	0.928849234	0.0001											-
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	E/X	0.304278197	0.0000											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	(B+T)/(E+X)	0.877916155	0.0001											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	1,2,3-Trimethylbenzene	3870 ug/mL	0.5360											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	1,2,4-Trimethylbenzene	15700 ug/mL	2.1745											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	1,3,5-Trimethylbenzene	4560 ug/mL	0.6316											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	Total TMBs	24130 ug/mL	3.3421											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	Naphthalene	2080 ug/mL	0.2881											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	1-MethylNaphthalene	900 ug/mL	0.1247											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	2-MethylNaphthalene	1860 ug/mL	0.2576											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-6FP	I otal Naphthalenes	4840 ug/mL	0.6704											
NPMPI	Gasoline	6/1/1000	10/27/1008	MW349-6	Density	0.744 g/ml	0.0001											
	Gasoline	0/1/1330	10/21/1330	10100343-0	Density	0.744 g/mL	0.0001											
NRMRI	Gasoline	6/1/1990	10/27/1998	MW349-6	Benzene	2600 ug/ml	0 3495	1 1505	0 1368	9 1 2	0 1732	15.90	2 3824	0 2833	10.37	0 2445	21.69	
NRMRI	Gasoline	6/1/1990	10/27/1998	MW349-6	Toluene	25640 ug/mL	3 4462	2 4538	0 2917	4 94	0.0639	6 19	9.0657	1 0778	8.61	0.1533	14 21	
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	Ethylbenzene	10120 ug/mL	1.3602	-0.0602	-0.0072	-0.55	-0.0054	-0.54	0.3206	0.0381	2.27	0.0252	2.48	
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	m-Xylene	23720 ug/mL	3.1882											-
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	o-Xylene	11840 ug/mL	1.5914											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	p-Xylene	9520 ug/mL	1.2796											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	m,p-xylenes	33240 ug/mL	4.4677											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	Total Xylenes (m,p, and o)	45080 ug/mL	6.0591	-0.1591	-0.0189	-0.32	-0.0032	-0.32	1.4876	0.1769	2.34	0.0261	2.58	
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	Total BTEX	83440 ug/mL	11.2151	3.3849	0.4024	2.76	0.0314	3.09	13.2563	1.5761	6.44	0.0928	8.86	
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	B+T	28240 ug/mL					_							
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	E+X	55200 ug/mL												
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	B/T	0.101404056												
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	B/E	0.256916996												
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	B/X	0.057675244												
NRMRL	Gasoline	6/1/1990	10/27/1998	MVV349-6	1/E	2.533596838												
NEMPI	anilose inc	1 K/1/1000	111/27/1008	11/11/13/0-6	11/X	11 568 (66637)						1				1		

															Mid-Ra	nge (Potter	, 1988),		
										Ghass	semi <i>et al</i> ., 1	984		AD L	ittle, Inc. (1	1987), & Sig	sby et al. (1987)	
Lab	Fuel	Spill	Sample							Line	ear	Expor	nential		Lin	ear	Expo	nential	
Code	Type	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	C ₀ - C	k	%Red./yr	k	%Red./yr	C ₀ - C	k	%Red./yr	k	%Red./yr	
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	E/X	0.224489796				ľ									
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	(B+T)/(E+X)	0.511594203													
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	1,2,3-Trimethylbenzene	3880	ug/mL	0.5215											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	1,2,4-Trimethylbenzene	15640	ug/mL	2.1022											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	1,3,5-Trimethylbenzene	5560	ug/mL	0.7473											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	Total TMBs	25080	ug/mL	3.3710											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	Naphthalene	1920	ug/mL	0.2581											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	1-MethylNaphthalene	880	ug/mL	0.1183											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	2-MethylNaphthalene	1960	ug/mL	0.2634											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-6	Total Naphthalenes	4760	ug/mL	0.6398											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-1	Density	0.738	g/mL	0.0001											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-1	Benzene	8280	ug/mL	1.1220	0.3780	0.0848	5.65	0.0651	6.30	1.6099	0.3609	13.21	0.1995	18.09	
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-1	Toluene	41100	ug/mL	5.5691	0.3309	0.0742	1.26	0.0129	1.29	6.9428	1.5566	12.44	0.1815	16.60	
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-1	Ethylbenzene	10300	ug/mL	1.3957	-0.0957	-0.0214	-1.65	-0.0159	-1.60	0.2851	0.0639	3.80	0.0417	4.08	
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-1	m-Xylene	21700	ug/mL	2.9404											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-1	o-Xylene	11400	ug/mL	1.5447											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-1	p-Xylene	8980	ug/mL	1.2168											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-1	m,p-xylenes	30680	ug/mL	4.1572								- 10			
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-1	Total Xylenes (m,p, and o)	42080	ug/mL	5.7019	0.1981	0.0444	0.75	0.0077	0.76	1.8449	0.4136	5.48	0.0628	6.09	
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-1	Iotal BIEX	101760	ug/mL	13.7886	0.8114	0.1819	1.25	0.0128	1.27	10.6828	2.3951	9.79	0.1286	12.07	
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-1	B+I	49380	ug/mL	6.6911											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-1	E+X	52380	ug/mL	7.0976											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-1	B/I	0.201459854		0.0000											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-1	B/E	0.803883495		0.0001											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-1	B/X	0.196768061		0.0000											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-1	T/E	3.990291262		0.0005											
NRMRL	Gasoline	6/1/1990	11/15/1994	NIVV349-1	1/X	0.976711027		0.0001											
NRMRL	Gasoline	6/1/1990	11/15/1994	NIVV349-1		0.244771863		0.0000											
NDMDI	Gasoline	6/1/1990	11/15/1994	M/M/240 1		1950	ua/ml	0.0001											
NEME	Gasoline	6/1/1990	11/15/1994	MW249-1	1 MothylNaphthalana	1650	ug/mL	0.2507											
NEME	Gasoline	6/1/1000	11/15/1994	MW349-1	2-MethylNaphthalene	1600	ug/mL	0.1211											
NDMDI	Gasoline	6/1/1990	11/15/1994	M/M/240 1	Z-WethyliNaphthalene	1000	ug/mL	0.2100											
	Casoline	0/1/1330	11/13/1334	10100343-1	Total Naphthalenes	4344	ug/IIIL	0.0000											
NRMRI	Gasoline	6/1/1990	6/26/1996	MW/349-1	Density	0 724	a/ml	0.0001											
	Cusoline	0/1/1000	0/20/1000	11111040	Density	0.724	g/IIIL	0.0001											
NRMRI	Gasoline	6/1/1990	6/26/1996	MW349-1	Benzene	8420	ua/ml	1 1630	0.3370	0.0555	3 70	0.0419	4 10	1 5689	0 2583	9.45	0 1406	13 12	
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	Toluene	36400	ug/mL	5.0276	0.8724	0.1436	2.43	0.0263	2.60	7.4843	1.2322	9.85	0.1501	13.94	
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	Ethylbenzene	11400	ua/mL	1.5746	-0.2746	-0.0452	-3.48	-0.0315	-3.21	0.1062	0.0175	1.04	0.0107	1.07	
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	m-Xvlene	17700	ua/mL	2.4448	0.2.1.0		00								
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	o-Xylene	10700	ug/mL	1.4779											
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	p-Xylene	9360	ug/mL	1.2928											
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	m,p-xylenes	27060	ug/mL	3.7376											-
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	Total Xylenes (m,p, and o)	37760	ug/mL	5.2155	0.6845	0.1127	1.91	0.0203	2.01	2.3313	0.3838	5.09	0.0608	5.90	
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	Total BTEX	93980	ug/mL	12.9807	1.6193	0.2666	1.83	0.0194	1.92	11.4907	1.8918	7.73	0.1044	9.91	
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	B+T	44820	ug/mL	6.1906											
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	E+X	49160	ug/mL	6.7901											
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	B/T	0.231318681		0.0000											
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	B/E	0.738596491		0.0001											
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	B/X	0.222987288		0.0000											
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	T/E	3.192982456		0.0004											
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	T/X	0.963983051		0.0001											
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	E/X	0.30190678		0.0000				-			-				
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-1	(B+T)/(E+X)	0.911716843		0.0001											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	Density	0.72	g/mL	0.0001											
L																			
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	Benzene	8310	ug/mL	1.1542	0.3458	0.0489	3.26	0.0371	3.64	1.5777	0.2233	8.17	0.1219	11.48	
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	Toluene	40000	ug/mL	5.5556	0.3444	0.0487	0.83	0.0085	0.85	6.9564	0.9845	7.87	0.1149	10.85	
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	Ethylbenzene	11100	ug/mL	1.5417	-0.2417	-0.0342	-2.63	-0.0241	-2.44	0.1391	0.0197	1.17	0.0122	1.22	
INRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	m-Xvlene	23300	ua/mL	3.2361					1			1			

															Mid-Ra	nge (Potter,	, 1988),		
										Ghass	semi <i>et al</i> ., 19	984		AD Li	ittle, Inc. (1	987), & Sigs	sby et al. (1987)	
Lab	Fuel	Spill	Sample						ĺ	Lin	ear	Expon	ential		Line	ear	Expor	ential	
Code	Туре	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	C ₀ - C	k	%Red./yr	k	%Red./yr	C ₀ - C	k	%Red./yr	k	%Red./yr	
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	o-Xylene	10600	ug/mL	1.4722	Î										
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	p-Xylene	9040	ug/mL	1.2556											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	Total Xylenes (m,p, and o)	42940	ug/mL	5.9639	-0.0639	-0.0090	-0.15	-0.0015	-0.15	1.5829	0.2240	2.97	0.0333	3.28	
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	Total BTEX	102350	ug/mL	14.2153	0.3847	0.0544	0.37	0.0038	0.38	10.2561	1.4515	5.93	0.0769	7.40	
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	B+T	48310	ug/mL	6.7097											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	E+X	54040	ug/mL	7.5056											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	B/T	0.20775		0.0000											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	B/E	0.748648649		0.0001											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	B/X	0.19352585		0.0000											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	T/E	3.603603604		0.0005											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	T/X	0.931532371		0.0001											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	E/X	0.258500233		0.0000											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	(B+T)/(E+X)	0.893967432		0.0001											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	1,2,3-Trimethylbenzene	3450	ug/mL	0.4792											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	1,2,4-Trimethylbenzene	14800	ug/mL	2.0556											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	1,3,5-Trimethylbenzene	4340	ug/mL	0.6028											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	Naphthalene	2170	ug/mL	0.3014											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	1-MethylNaphthalene	1100	ug/mL	0.1528											
NRMRL	Gasoline	6/1/1990	6/23/1997	MW-349-1FP	2-MethylNaphthalene	2280	ug/mL	0.3167											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	Density	0.728	g/mL	0.0001											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	Benzene	7400	ug/mL	1.0165	0.4835	0.0575	3.83	0.0463	4.52	1.7154	0.2039	7.47	0.1175	11.09	
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	Toluene	35000	ug/mL	4.8077	1.0923	0.1299	2.20	0.0243	2.40	7.7043	0.9160	7.32	0.1137	10.75	
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	Ethylbenzene	11840	ug/mL	1.6264	-0.3264	-0.0388	-2.98	-0.0266	-2.70	0.0544	0.0065	0.38	0.0039	0.39	
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	m-Xylene	23360	ug/mL	3.2088											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	o-Xylene	11720	ug/mL	1.6099											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	p-Xylene	9360	ug/mL	1.2857											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	m,p-xylenes	32720	ug/mL	4.4945											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	Total Xylenes (m,p, and o)	44440	ug/mL	6.1044	-0.2044	-0.0243	-0.41	-0.0040	-0.41	1.4424	0.1715	2.27	0.0252	2.49	
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	Total BTEX	98680	ug/mL	13.5549	1.0451	0.1242	0.85	0.0088	0.88	10.9164	1.2979	5.30	0.0702	6.78	
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	B+T	42400	ug/mL												
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	E+X	56280	ug/mL												
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	B/T	0.211428571													
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	B/E	0.625													
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	B/X	0.166516652													
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	T/E	2.956081081													
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	T/X	0.787578758													
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	E/X	0.266426643													
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	(B+T)/(E+X)	0.753375977													
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	1,2,3-Trimethylbenzene	3120	ug/mL	0.4286											
NRMRL	Gasoline	6/1/1990	10/27/1998	MVV349-1	1,2,4-Trimethylbenzene	12560	ug/mL	1.7253											
NRMRL	Gasoline	6/1/1990	10/27/1998	MVV349-1	1,3,5-Trimethylbenzene	4600	ug/mL	0.6319											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	Total TMBs	20280	ug/mL	2.7857											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	Naphthalene	2000	ug/mL	0.2747											
NRMRL	Gasoline	6/1/1990	10/27/1998	MW349-1	1-MethylNaphthalene	920	ug/mL	0.1264											
NRMRL	Gasoline	6/1/1990	10/27/1998	MVV349-1	2-MethylNaphthalene	2040	ug/mL	0.2802											
NRMRL	Gasoline	6/1/1990	10/27/1998	MVV349-1	I otal Naphthalenes	4960	ug/mL	0.6813											

															Mid-Rar	nge (Potter,	1988),		
										Ghass	semi <i>et al</i> ., 19	984		AD Li	ittle, Inc. (1	987), & Sigs	sby et al. (1987)	
Lab	Fuel	Spill	Sample							Lin	ear	Expon	ential		Line	ear	Expor	ential	
Code	Туре	Date	Date	Locid	Analyte	Results	Units	Mass Fraction	C ₀ - C	k	%Red./yr	k	%Red./yr	C ₀ - C	k	%Red./yr	k	%Red./yr	
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	Density	0.737	g/mL	0.0001											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	Benzene	5620	ug/mL	0.7626	0.7374	0.1653	11.02	0.1517	14.07	1.9693	0.4415	16.16	0.2861	24.88	
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	Toluene	28600	ug/mL	3.8806	2.0194	0.4528	7.67	0.0939	8.97	8.6314	1.9352	15.47	0.2625	23.09	
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	Ethylbenzene	9260	ug/mL	1.2564	0.0436	0.0098	0.75	0.0076	0.76	0.4243	0.0951	5.66	0.0652	6.32	
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	m-Xylene	21400	ug/mL	2.9037											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	o-Xylene	11600	ug/mL	1.5739											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	p-Xylene	8920	ug/mL	1.2103											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	m,p-xylenes	30320	ug/mL	4.1140											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	Total Xylenes (m,p, and o)	41920	ug/mL	5.6879	0.2121	0.0475	0.81	0.0082	0.82	1.8589	0.4168	5.52	0.0634	6.14	
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	Total BTEX	85400	ug/mL	11.5875	3.0125	0.6754	4.63	0.0518	5.05	12.8839	2.8886	11.80	0.1676	15.43	
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	B+T	34220	ug/mL	4.6431											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	E+X	51180	ug/mL	6.9444											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	B/T	0.196503497		0.0000											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	B/E	0.606911447		0.0001											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	B/X	0.134064885		0.0000											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	T/E	3.088552916		0.0004											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	T/X	0.682251908		0.0001											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	E/X	0.220896947		0.0000											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	(B+T)/(E+X)	0.668620555		0.0001											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	Naphthalene	2170	ug/mL	0.2944											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	1-MethylNaphthalene	749	ug/mL	0.1016											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	2-MethylNaphthalene	1420	ug/mL	0.1927											
NRMRL	Gasoline	6/1/1990	11/15/1994	MW349-7	Total Naphthalenes	4339	ug/mL	0.5887											
					-														
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-8	Density	0.742	g/mL	0.0001											
	.																		
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-8	Benzene	955	ug/mL	0.1287	1.3713	0.2258	15.05	0.4043	33.26	2.6032	0.4286	15.69	0.5030	39.53	
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-8	Toluene	12300	ug/mL	1.6577	4.2423	0.6984	11.84	0.2090	18.86	10.8543	1.7870	14.28	0.3328	28.31	
NRMRL	Gasoline	6/1/1990	6/26/1996	MW349-8	Ethylbenzene	10100	ug/mL	1.3612	-0.0612	-0.0101	-0.77	-0.0076	-0.76	0.3196	0.0526	3.13	0.0347	3.41	
NRIVIRL	Gasoline	6/1/1990	6/26/1996	NIVV349-8	m-Xylene	15400	ug/mL	2.0755											
NRIVIRL	Gasoline	6/1/1990	6/26/1996	MW240-8	0-Xylene	9150	ug/mL	1.2332											
NDMD	Gasoline	6/1/1990	6/26/1996	NIV/349-6	p-Aylene	0030		1.0022											
NRIVIRL	Gasoline	0/1/1990	6/26/1996	10100349-0	Tatal Values (manual a)	23430	ug/mL	3.1577	4 5000	0.0405	4.04	0.0400	4.75	0.4550	0.5400	0.00	0.0000	0.50	
NEWEL	Gasoline	6/1/1990	6/26/1996	NIV/349-0	Total Aylenes (m,p, and 0)	32360		4.3900	7.0616	0.2465	4.21	0.0400	4.75	3.1559	0.5190	0.00	0.0692	0.00	
NDMDI	Caseline	6/1/1990	6/26/1990	MN/240.9		12255		1.0004	7.0010	1.1020	7.90	0.1000	10.31	10.9330	2.7070	11.39	0.1939	17.02	
NDMDI	Gasoline	6/1/1990	6/26/1996	MM240 9		13233	ug/mL	5 7520											
NPMPI	Gasolino	6/1/1000	6/26/1006	MW349-0	B/T	42000	ug/IIIL	0.0000											
NDMDI	Gasoline	6/1/1990	6/26/1996	MM240 9	D/I D/E	0.077642276		0.0000											
NEME	Gasolino	6/1/1000	6/26/1006	MW349-8	B/Y	0.034334455		0.0000											
NEME	Gasoline	6/1/1000	6/26/1996	MW/349-0	T/E	1 217821782		0.0000											
NEME	Gasolino	6/1/1000	6/26/1006	MW349-8		0.377532220		0.0002											
NPMPI	Gasolino	6/1/1000	6/26/1006	MW349-0	E/X	0.310006120		0.0001											
NEME	Gasolino	6/1/1000	6/26/1006	MW/349-0	(B+T)/(E+X)	0.31056701		0.0000											
TALVIAL	Jasoline	0/1/1990	0/20/1990	10100349-0		0.01000701		0.0000											

ANALYSIS OF GASOLINE FREE PRODUCT WEATHERING RATES WHERE 2 OR MORE SAMPLING EVENTS HAVE BEEN PERFORMED, MW349-1 TANK 349, OFFUTT AFB (NRMRL DATA ONLY)

Linear (Zero Order) Assumption

linear equation $C = C_0 - kt$

summary statistics presented as follows

ana	lyte	
k	Co	k = zero order weathering rate; k = dC/dt or slope
sek	seCo	sek = slope standard error value
r ² F stat.	seC df	Co = intercept or initial analyte concentration as calculated by regression analysis seCo = standard error value for the constant C _o
SS _{regr}	SS _{resid}	r^{z} = coefficient of determination
		seC = standard error value of the estimated concentration C (i.e., a "standard deviation" for the regression line)

F stat. = F statistic or F-observed value

- df = degrees of freedom
- $ss_{\mbox{\scriptsize regr}}$ = the regression sum of squares
- ss_{resid} = the residual sum of squares

Exponential (1st Order) Assumption

exponential equation $C = C_o e^{-kt}$

summary statistics presented as follows

ana	llyte
m	Co
sek	seCo
r ²	seC
F stat.	df
SS _{regr}	SS _{resid}
	In C _o

- m = coefficient for statistics equation shown; note $m = e^{-kt}$, therefore, ln m = -k
- sek = standard error value for the exponential rate constant k
- Co = intercept or initial analyte concentration as calculated by regression analysis
- $seCo = standard error value for the constant C_o; compare to ln C_o$
 - r^2 = coefficient of determination

seC = standard error value of the estimated concentration C (i.e., a "standard deviation" for the regression line)

- F stat. = F statistic or F-observed value
 - df = degrees of freedom
- ss_{regr} = the regression sum of squares
- ss_{resid} = the residual sum of squares
- $ln C_o = natural log of Co for comparing to seCo$

ANALYSIS OF GASOLINE FREE PRODUCT WEATHERING RATES WHERE 2 OR MORE SAMPLING EVENTS HAVE BEEN PERFORMED, MW349-1 TANK 349, OFFUTT AFB (NRMRL DATA ONLY)

Sample			Ghassem	i et al., 1	984			
Location	Date	Time (yrs)	Benzene	Toluene	Ethylbenzene	Total Xylenes	Total BTEX	
MW-349-1	6/1/1990	0.00	1.50	5.90	1.30	5.90	14.60 initial concentrations from Ghassemi et al., 19	984
	########	4.46	1.12	5.57	1.40	5.70) 13.79	
	6/26/1996	6.07	1.16	5.03	1.57	5.22	12.98	
	6/23/1997	7.07	1.15	5.56	1.54	5.96	5 14.22	
	########	8.41	1.02	4.81	1.63	6.10	13.55	
linear								
Со			1.4638	5.9475	1.2820	5.7370) 14.4303	
Predicted C - latest sample date			1.0229	5.0170	1.6145	5.8019	13.4563	
linear rate constant (k) (slope)			0.0524	0.1106	-0.0395	-0.0077	0.1158	
average yearly reduction (%)			3.58	1.86	-3.08	-0.13	8 0.80	

linear

summary stats

	Benz	zene	Tolu	ene	Ethylbe	enzene	Total X	ylenes	Total BTEX		
-0.0	0524	1.4638	-0.1106	5.9475	0.0395	1.2820	0.0077	5.7370	-0.1158	14.4303	
0.0	0116	0.0691	0.0464	0.2767	0.0077	0.0460	0.0613	0.3654	0.0878	0.5232	
0.8	3718	0.0753	0.6541	0.3015	0.8974	0.0501	0.0052	0.3982	0.3669	0.5701	
2	20	3	6	3	26	3	0	3	2	3	
0.1	1158	0.0170	0.5157	0.2727	0.0659	0.0075	0.0025	0.4757	0.5651	0.9750	

exponential	Benzene	Toluene	Ethylbenzene	Total Xylenes	Total BTEX
Co	1.4644	5.9613	1.2862	5.7351	14.4234
exponential rate constant (k)	0.0414	0.0205	-0.0273	-0.0011	0.0083
% reduction/year	4.05	2.03	-2.77	-0.11	0.82

exponential

Ben	Benzene Toluene		ene	Ethylbe	enzene	Total X	ylenes	Total BTEX	
0.9595	1.4644	0.9797	5.9613	1.0277	1.2862	1.0011	5.7351	0.9918	14.4234
0.0093	0.0554	0.0089	0.0533	0.0050	0.0300	0.0109	0.0650	0.0064	0.0384
0.8682	0.0604	0.6378	0.0580	0.9072	0.0327	0.0035	0.0709	0.3544	0.0418
20	3	5	3	29	3	0	3	2	3
0.0721	0.0109	0.0178	0.0101	0.0314	0.0032	0.0001	0.0151	0.0029	0.0052
	0.3815		1.7853		0.2517		1.7466		2.6689

ANALYSIS OF GASOLINE FREE PRODUCT WEATHERING RATES WHERE 2 OR MORE SAMPLING EVENTS HAVE BEEN PERFORMED, MW349-1 TANK 349, OFFUTT AFB (NRMRL DATA ONLY)

Sample			Mid-range	Ð				
Location	Date	Time (yrs)	Benzene	Toluene	Ethylbenzene	Total Xylenes	Total BTEX	
MW-349-1	6/1/1990	0.00	2.73	12.51	1.68	7.55	24.47	mid-range initial concentrations as presented by Potter (1988), AD Little (1987), and
	########	4.46	1.12	5.57	1.40	5.70	13.79	Sigsby et al., (1987)
	6/26/1996	6.07	1.16	5.03	1.57	5.22	12.98	
	6/23/1997	7.07	1.15	5.56	1.54	5.96	14.22	
	########	8.41	1.02	4.81	1.63	6.10	13.55	
linear								
Со			2.5013	11.5159	1.6027	7.1239	22.7438	
Predicted C - latest sample date			0.7813	3.7204	1.5398	5.4789	11.5205	
linear rate constant (k) (slope)			0.2045	0.9268	0.0075	0.1956	1.3344	
average yearly reduction (%)			8.18	8.05	0.47	2.75	5.87	

linear

summary stats

ſ	Benz	zene	Toluene		Ethylbe	enzene	Total X	ylenes	Total BTEX	
ſ	-0.2045	2.5013	-0.9268	11.5159	-0.0075	1.6027	-0.1956	7.1239	-1.3344	22.7438
	0.0523	0.3115	0.2275	1.3554	0.0187	0.1113	0.1067	0.6359	0.3948	2.3519
	0.8360	0.3394	0.8469	1.4770	0.0506	0.1213	0.5281	0.6930	0.7920	2.5628
	15	3	17	3	0	3	3	3	11	3
	1.7622	0.3456	36.2004	6.5446	0.0024	0.0441	1.6118	1.4406	75.0347	19.7041

exponential	Benzene	Toluene	Ethylbenzene	Total Xylenes	Total BTEX
Co	2.4263	11.2277	1.5969	7.0563	22.2830
exponential rate constant (k)	0.1154	0.1133	0.0044	0.0293	0.0720
% reduction/year	10.90	10.72	0.44	2.88	6.95

exponential

Ben	Benzene Toluene		Ethylb	enzene	Total X	ylenes	Total BTEX		
0.8910	2.4263	0.8928	11.2277	0.9956	1.5969	0.9712	7.0563	0.9305	22.2830
0.0276	0.1646	0.0257	0.1533	0.0123	0.0732	0.0175	0.1041	0.0218	0.1297
0.8533	0.1793	0.8660	0.1671	0.0410	0.0798	0.4834	0.1134	0.7849	0.1413
17	3	19	3	0	3	3	3	11	3
0.5610	0.0965	0.5413	0.0837	0.0008	0.0191	0.0361	0.0386	0.2186	0.0599
	0.8864		2.4184		0.4681		1.9539		3.1038
ANALYSIS OF GASOLINE FREE PRODUCT WEATHERING RATES WHERE 2 OR MORE SAMPLING EVENTS HAVE BEEN PERFORMED, MW349-1 TANK 349, OFFUTT AFB (NRMRL DATA ONLY)

Sample			Measured Data Only									
Location	Date	Time (yrs)	Benzene	Toluene	Ethylbenzene	Total Xylenes	Total BTEX					
MW-349-1	########	0.00	1.12	5.57	1.40	5.70	13.79					
	6/26/1996	1.61	1.16	5.03	1.57	5.22	12.98					
	6/23/1997	2.61	1.15	5.56	1.54	5.96	14.22					
	########	3.95	1.02	4.81	1.63	6.10	13.55					
linear												
Со			1.1632	5.5418	1.4251	5.4702	13.6003					
Predicted C - latest sample date			1.0678	4.9580	1.6369	6.0044	13.6672					
linear rate constant (k) (slope)			0.0241	0.1478	-0.0536	-0.1352	-0.0169					
average yearly reduction (%)			2.08	2.67	-3.76	-2.47	-0.12					

linear

summary stats

Benz	zene	Tolu	ene	Ethylbo	enzene	Total X	ylenes	Total	BTEX	
-0.0241	1.1632	-0.1478	5.5418	0.0536	1.4251	0.1352	5.4702	0.0169	13.6003	
0.0229	0.0573	0.1247	0.3116	0.0181	0.0454	0.1360	0.3399	0.2183	0.5458	
0.3567	0.0661	0.4126	0.3594	0.8135	0.0523	0.3309	0.3920	0.0030	0.6295	
1	2	1	2	9	2	1	2	0	2	
0.0048	0.0087	0.1815	0.2584	0.0239	0.0055	0.1520	0.3073	0.0024	0.7925	

exponential	Benzene	Toluene	Ethylbenzene	Total Xylenes	Total BTEX
Co	1.1649	5.5444	1.4246	5.4696	13.5927
exponential rate constant (k)	0.0226	0.0286	-0.0356	-0.0233	-0.0013
% reduction/year	2.23	2.82	-3.63	-2.36	-0.13

exponential

. summary stats

Benz	zene	Tolu	ene	Ethylb	enzene	Total X	ylenes	Total	BTEX
0.9777	1.1649	0.9718	5.5444	1.0363	1.4246	1.0236	5.4696	1.0013	13.5927
0.0209	0.0524	0.0238	0.0595	0.0123	0.0306	0.0245	0.0612	0.0161	0.0402
0.3678	0.0604	0.4197	0.0686	0.8087	0.0353	0.3117	0.0706	0.0030	0.0464
1	2	1	2	8	2	1	2	0	2
0.0042	0.0073	0.0068	0.0094	0.0106	0.0025	0.0045	0.0100	0.0000	0.0043
	0.1526		1.7128		0.3539		1.6992		2.6095

ANALYSIS OF GASOLINE FREE PRODUCT WEATHERING RATES WHERE 2 OR MORE SAMPLING EVENTS HAVE BEEN PERFORMED, MW349-1 TANK 349, OFFUTT AFB (NRMRL DATA ONLY)

Sample	1997 and 1998 Data Only										
Location	Date	Time (yrs)	Benzene	Toluene	Ethylbenzene	Total Xylenes	Total BTEX				
MW-349-1	6/23/1997	0.00	1.15	5.56	1.54	5.96	14.22				
	########	1.35	1.02	4.81	1.63	6.10	13.55				
linear											
Со			1.1542	5.5556	1.5417	5.9639	14.2153				
Predicted C - latest sample date			1.0165	4.8077	1.6264	6.1044	13.5549				
linear rate constant (k) (slope)			0.1024	0.5559	-0.0630	-0.1045	0.4909				
average yearly reduction (%)			8.87	10.01	-4.08	-1.75	3.45				

exponential	Benzene	Toluene	Ethylbenzene	Total Xylenes	Total BTEX
Co	1.1542	5.5556	1.5417	5.9639	14.2153
exponential rate constant (k)	0.0944	0.1075	-0.0398	-0.0173	0.0354
% reduction/year	9.01	10.19	-4.06	-1.75	3.47

APPENDIX C-2

\mathbf{K}_{fw} CALCULATIONS

Event	Lab Code	Site Name	Fuel Type	Spill Date	Date	FPLocid	Analyte	Kfw GW Results GW UNITS FP Result		FP Results	Units
3	EAL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-6FP	Benzene	280	50000 ug/L	1400000	ug/L
	EAL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	10/27/98	OFMW349-6	Benzene	340.9090909	8800 ug/L	3000000	ug/L
		, ,					Analyte avg kfw	310.4545455	9		Ŭ
3	EAL	Tank 349. Offutt AFB. NE	Gasoline	06/01/90	06/23/97	MW-349-6FP	Toluene	787.8787879	66000 ua/L	52000000	ua/L
	EAL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	10/27/98	OFMW349-6	Toluene	1500	20000 ug/L	3000000	ug/L
		, , ,					Analyte avg kfw	1143.939394			Ŭ
3	EAL	Tank 349. Offutt AFB. NE	Gasoline	06/01/90	06/23/97	MW-349-6FP	Ethylbenzene	1200	10000 ug/L	12000000	ua/L
	EAL	Tank 349. Offutt AFB. NE	Gasoline	06/01/90	10/27/98	OFMW349-6	Ethylbenzene	6666.666667	1800 ug/L	12000000	ua/L
							Analyte avg kfw	3933.333333			
3	EAL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-6FP	Total Xylenes (m,p, and o)	1266.666667	45000 ug/L	5700000	ug/L
	EAL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	10/27/98	OFMW349-6	o-Xylene	4857.142857	3500 ug/L	17000000	ug/L
	EAL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	10/27/98	OFMW349-6	m,p-Xylene	6727.272727	5500 ug/L	37000000	ug/L
		, , ,							0		Ŭ
2	EAL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	Benzene	216.6666667	6000 ug/L	1300000	ug/L
1	EAL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	Benzene	265	10000 ug/L	2650000	ug/L
							Analyte avg kfw	240.8333333			
2	EAL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	Toluene	789.4736842	3800 ug/L	300000	ug/L
1	EAL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	Toluene	945.6342669	6070 ug/L	5740000	ug/L
							Analyte avg kfw	867.5539755			
2	EAL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	Ethylbenzene	3414.634146	410 ug/L	1400000	ug/L
1	EAL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	Ethylbenzene	4059.633028	436 ug/L	1770000	ug/L
							Analyte avg kfw	3737.133587			
2	EAL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	o-Xylene	2583.333333	1200 ug/L	3100000	ug/L
1	EAL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	o-Xylene	2837.398374	1230 ug/L	3490000	ug/L
							Analyte avg kfw	2710.365854			
2	EAL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	m,p-Xylene	3105.263158	1900 ug/L	5900000	ug/L
1	EAL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	m,p-Xylene	3435.779817	2180 ug/L	7490000	ug/L
				00/04/04	00/11/00	01100.1010.0	Analyte avg kfw	3270.521487	0.1.00		
				06/01/94	03/11/98	SH98-1610-2	Total Xylenes (m,p, and o)	2903.225806	3100 ug/L	9000000	ug/L
				06/01/94	03/06/97	SHMW1610-2	I otal Xylenes (m,p, and o)	3219.941349	3410 ug/L	10980000	ug/L
							Analyte avg kfw	3061.583578			
1	EAL	Pipeline Leak Site Myrtle Beach AEB SC	ID-4	01/01/81	03/04/07	MBMM/8I	Benzene	202 8846154	1040 ug/l	211000	ua/l
1		Pipeline Leak Site, Myrtle Beach AEB, SC	JI -4	01/01/81	03/04/97	MBMW01	Toluene	1508	5 ug/l	7540	ug/L
1	FAL	Pipeline Leak Site Myrtle Beach AFB SC	JP-4	01/01/81	03/04/07	MBMW8I	Ethylbenzene	3553 398058	515 ug/L	1830000	ug/L
1	FAL	Pipeline Leak Site Myrtle Beach AFB SC	JP-4	01/01/81	03/04/07	MBMW8I	o-Xylene	13909 09091	11 ug/L	153000	ug/L
1	EAL	Pipeline Leak Site, Myrtle Beach AFB, SC	JP-4	01/01/81	03/04/97	MBMW8I	m.p-Xvlene	3568,965517	1740 ug/L	6210000	ug/L
	2712		0	01/01/81	03/04/97	MBMW8I	Total Xylenes (m.p. and o)	3633 923472	1751 ug/l	6363000	ua/l
				01101101	00/01/01			0000.020.112			ug/2
3	EAL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/17/97	CH-EW6	Benzene	10.86956522	2.3 ug/L	25	ug/L
3	EAL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	Benzene	5	5 ug/L	25	ug/L
							Analyte avg kfw	7.934782609	l v		
3	EAL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/17/97	CH-EW6	Toluene	1350	1 ug/L	1350	ug/L
3	EAL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	Toluene	1020	200 ug/L	204000	ug/L
							Analyte avg kfw	1185			
3	EAL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/17/97	CH-EW6	Ethylbenzene	4805.263158	19 ug/L	91300	ug/L
3	EAL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	Ethylbenzene	3692.307692	260 ug/L	960000	ug/L
							Analyte avg kfw	4248.785425			
3	EAL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/17/97	CH-EW6	Total Xylenes (m,p, and o)	24705.88235	85 ug/L	2100000	ug/L

Event	Lab Code	Site Name	Fuel Type	Spill Date	Date	FPLocid	Analyte	nalyte Kfw GW Results GW UNITS FP		FP Results	Units
3	EAL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	Total Xylenes (m,p, and o)	3600	1500 ug/L	5400000	ug/L
							Analyte avg kfw	14152.94118			
1	EAL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW316-FP	Benzene	384.6153846	5200 ug/L	2000000	ug/L
1	EAL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW316-FP	Toluene	40000	25 ug/L	1000000	ug/L
1	EAL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW316-FP	Ethylbenzene	5500	1000 ug/L	5500000	ug/L
1	EAL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW316-FP	o-Xylene	4300	1000 ug/L	4300000	ug/L
1	EAL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW316-FP	m,p-Xylene	4782.608696	4600 ug/L	22000000	ug/L
				10/01/73	08/27/97	EAKIVIVV316-FP	Total Xylenes (m,p, and o)	4696.428571	5600 ug/L	26300000	ug/L
1		Eacility 202 Cooil Field NAS El	ID 5	06/01/91	05/20/07	CEE 202 0ED	Ponzono	252 600172	02 ug/l	22500	ua/l
1		Facility 293, Cecil Field NAS, FL	JF-J	06/01/81	05/20/97	CEF-293-9FF	Toluono	1460 970519	95 ug/L	122000	ug/L
1		Facility 293, Cecil Field NAS, FL	JF-5	06/01/81	05/20/97	CEF-293-9FF	Ethylbenzene	5818 181818	550 ug/L	3200000	ug/L
1		Eacility 293, Cecil Field NAS, FL	JI -5	06/01/81	05/20/97	CEE-203-0ED	Total Xylenes (m.n. and o)	6636 363636	1100 ug/l	7300000	ug/L
			51-5	00/01/01	03/20/37	021-235-311	Total Aylenes (III,p, and 0)	0030.303030	1100 49/2	7300000	ug/L
1	FAI	Tank Farm C. Beaufort MCAS, SC	IP-5	06/01/90	08/12/97	BET-401-3	Benzene	557 5	4 ug/l	2230	ua/l
1	FAL	Tank Farm C. Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BFT-401-3	Toluene	1250	16 ug/l	20000	ug/L
1	FAL	Tank Farm C. Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BFT-401-3	Ethylbenzene	4571 428571	35 ug/l	160000	ug/L
1	EAL	Tank Farm C. Beaufort MCAS. SC	JP-5	06/01/90	08/12/97	BFT-401-3	o-Xvlene	2538.461538	130 ug/L	330000	ua/L
1	EAL	Tank Farm C. Beaufort MCAS. SC	JP-5	06/01/90	08/12/97	BFT-401-3	m.p-Xvlene	4857.142857	140 ug/L	680000	ua/L
				06/01/90	08/12/97	BFT-401-3	Total Xylenes (m.p. and o)	3740,740741	270 ug/L	1010000	ua/L
1	EAL	Tank Farm C, Beaufort MCAS, SC	JP-5	06/01/90	05/19/97	Fresh JP-5	Benzene	454.5454545	3.3 ug/L	1500	ug/L
1	EAL	Tank Farm C, Beaufort MCAS, SC	JP-5	06/01/90	05/19/97	Fresh JP-5	Toluene	1500	24 ug/L	36000	ug/L
1	EAL	Tank Farm C, Beaufort MCAS, SC	JP-5	06/01/90	05/19/97	Fresh JP-5	Ethylbenzene	4567.901235	81 ug/L	370000	ug/L
1	EAL	Tank Farm C, Beaufort MCAS, SC	JP-5	06/01/90	05/19/97	Fresh JP-5	Total Xylenes (m,p, and o)	4814.814815	540 ug/L	2600000	ug/L
3	EAL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW1SFP	Benzene	208.3333333	720 ug/L	150000	ug/L
3	EAL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW2SFP	Benzene	271.6049383	810 ug/L	220000	ug/L
							Analyte avg kfw	239.9691358			
-											
3	EAL	Bidg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW1SFP	loluene	1010.10101	990 ug/L	1000000	ug/L
3	EAL	Bidg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW2SFP	I oluene	1000	1100 ug/L	1100000	ug/L
							Analyte avg krw	1005.050505			
2	ΕΛI	Bldg 4522 Saymour Johnson AEP SC	ID-8	12/01/05	05/15/07		Ethylbonzone	2724 127024	200	700000	ug/l
3		Bldg 4522, Seymour Johnson AEB SC		12/01/95	05/15/97	S IMW/2SED	Ethylbenzene	2124.13/931	290 ug/L 340 ug/L	120000	ug/L
3		Didy 4322, Seymour Johnson AFD, SC	JI -0	12/01/95	03/13/97	JJIVIVZJEF		3126 77/8/9	340 uy/L	1200000	uy/L
							Analyte avy NW	5120.114040			
3	FAI	Bldg 4522 Seymour Johnson AFB SC	JP-8	12/01/95	05/15/97	SJMW1SEP	Total Xylenes (m.p. and o)	2888 888889	1800 ug/l	5200000	ua/l
3	EAL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW2SFP	Total Xylenes (m.p. and o)	3285,714286	2100 ug/L	6900000	ua/L
Ū				.2,0.,00	20, 10,01		Analyte avg kfw	3087.301587	2.00 (3) 2		- 9' -
4	EAL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MW1S	Benzene	277.777778	180 ug/L	50000	ug/L
4	EAL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MW1S	Toluene	797.4683544	790 ug/L	630000	ug/L
4	EAL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MW1S	Ethylbenzene	3548.387097	310 ug/L	1100000	ug/L
4	EAL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MW1S	m,p-Xylene	3827.160494	810 ug/L	3100000	ug/L
4	EAL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MW1S	o-Xylene	2745.098039	510 ug/L	1400000	ug/L
4	EAL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MW1S	Total Xylenes (m,p, and o)	3409.090909	1320 ug/L	4500000	ug/L

Event	Lab Code	Site Name	Fuel Type	Spill Date	Date	FPLocid	Analyte	Kfw	GW Results GW UNITS	FP Results	Units
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-1FP	Benzene	237.4285714	35000 ug/L	8310	ug/mL
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-6FP	Benzene	245.8376691	38440 ug/L	9450	ug/mL
1	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	11/15/94	MW349-1	Benzene	243.5294118	34000 ug/L	8280	ug/mL
2	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/26/96	MW349-1	Benzene	263.125	32000 ug/L	8420	ug/mL
2	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/26/96	MW349-8	Benzene	99.89539749	9560 ug/L	955	ug/mL
		i i i i i i i i i i i i i i i i i i i					Analyte Kfw avg	217.96321			
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-1FP	Toluene	943.3962264	42400 ug/L	40000	ug/mL
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-6FP	Toluene	925.6725946	43860 ug/L	40600	ug/mL
1	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	11/15/94	MW349-1	Toluene	1151.260504	35700 ug/L	41100	ug/mL
2	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/26/96	MW349-1	Toluene	916.8765743	39700 ug/L	36400	ug/mL
2	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/26/96	MW349-8	Toluene	512.5	24000 ug/L	12300	ug/mL
							Analyte Kfw avg	889.9411799			
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-1FP	Ethylbenzene	2439.56044	4550 ug/L	11100	ug/mL
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-6FP	Ethylbenzene	3333.333333	3990 ug/L	13300	ug/mL
1	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	11/15/94	MW349-1	Ethylbenzene	2877.094972	3580 ug/L	10300	ug/mL
2	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/26/96	MW349-1	Ethylbenzene	2714.285714	4200 ug/L	11400	ug/mL
2	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/26/96	MW349-8	Ethylbenzene	2544.080605	3970 ug/L	10100	ug/mL
							Analyte Kfw avg	2781.671013			
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-1FP	o-Xylene	1859.649123	5700 ug/L	10600	ug/mL
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-6FP	o-Xylene	2365.591398	4650 ug/L	11000	ug/mL
1	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	11/15/94	MW349-1	o-Xylene	2456.896552	4640 ug/L	11400	ug/mL
2	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/26/96	MW349-1	o-Xylene	2383.073497	4490 ug/L	10700	ug/mL
2	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/26/96	MW349-8	o-Xylene	2188.995215	4180 ug/L	9150	ug/mL
							Analyte Kfw avg	2250.841157			
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-1FP	m-Xylene	2817.412334	8270 ug/L	23300	ug/mL
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-6FP	m-Xylene	3466.076696	6780 ug/L	23500	ug/mL
1	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	11/15/94	MW349-1	m-Xylene	3167.883212	6850 ug/L	21700	ug/mL
2	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/26/96	MW349-1	m-Xylene	2572.674419	6880 ug/L	17700	ug/mL
2	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/26/96	MW349-8	m-Xylene	2475.884244	6220 ug/L	15400	ug/mL
							Analyte Kfw avg	2899.986181			
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-1FP	p-Xylene	2560.906516	3530 ug/L	9040	ug/mL
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-6FP	p-Xylene	3209.059233	2870 ug/L	9210	ug/mL
1	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	11/15/94	MW349-1	p-Xylene	3218.637993	2790 ug/L	8980	ug/mL
2	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/26/96	MW349-1	p-Xylene	3183.673469	2940 ug/L	9360	ug/mL
2	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/26/96	MW349-8	p-Xylene	3064.885496	2620 ug/L	8030	ug/mL
							Analyte Kfw avg	3047.432541			
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-1FP	1,2,3-Trimethylbenzene	4049.295775	852 ug/L	3450	ug/mL
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-6FP	1,2,3-Trimethylbenzene	7648.221344	506 ug/L	3870	ug/mL
							Analyte Kfw avg	5848.758559			
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-1FP	1,2,4-Trimethylbenzene	4582.043344	3230 ug/L	14800	ug/mL
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-6FP	1,2,4-Trimethylbenzene	8532.608696	1840 ug/L	15700	ug/mL
							Analyte Kfw avg	6557.32602			
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-1FP	1,3,5-Trimethylbenzene	5136.094675	845 ug/L	4340	ug/mL
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-6FP	1,3,5-Trimethylbenzene	10340.13605	441 ug/L	4560	ug/mL
							Analyte Kfw avg	7738.115364			
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-1FP	Naphthalene	3001.383126	723 ug/L	2170	ug/mL
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-6FP	Naphthalene	4632.516704	449 ug/L	2080	ug/mL
							Analyte Kfw avg	3816.949915			
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-1FP	1-MethylNaphthalene	7482.993197	147 ug/L	1100	ug/mL
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-6FP	1-MethylNaphthalene	6617.647059	136 ug/L	900	ug/mL
							Analyte Kfw avg	7050.320128			
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-1FP	2-MethylNaphthalene	14430.37975	158 ug/L	2280	ug/mL
3	NRMRL	Tank 349, Offutt AFB, NE	Gasoline	06/01/90	06/23/97	MW-349-6FP	2-MethylNaphthalene	12739.72603	146 ug/L	1860	ug/mL
							Analyte Kfw avg	13585.05289			

Event	Lab Code	Site Name	Fuel Type	Spill Date	Date	FPLocid	Analyte	Kfw	GW Results	GW UNITS	FP Results	Units
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	Benzene	802.8259473	1557	ug/L	1250	ug/mL
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-3	Benzene	1157.894737	1425	ug/L	1650	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	Benzene	265.5493922	8473	ug/L	2250	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-3	Benzene	334.4643723	4126	ug/L	1380	ug/mL
							Analyte Kfw avg	640.1836122				
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	Toluene	1043.510324	2712	ug/L	2830	ug/mL
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-3	Toluene	2809.564475	1171	ug/L	3290	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	Toluene	554.4217687	8820	ug/L	4890	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-3	Toluene	1498.507463	1675	ug/L	2510	ug/mL
							Analyte Kfw avg	1476.501008				
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	Ethylbenzene	3041.825095	341.9	ug/L	1040	ug/mL
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-3	Ethylbenzene	4123.314066	259.5	ug/L	1070	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	Ethylbenzene	1138.487681	1177	ug/L	1340	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-3	Ethylbenzene	2876.923077	325	ug/L	935	ug/mL
							Analyte Kfw avg	2795.13748				
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	o-Xylene	2309.885932	1052	ug/L	2430	ug/mL
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-3	o-Xylene	3097.736756	790.9	ug/L	2450	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	o-Xylene	970.7140507	3039	ug/L	2950	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-3	o-Xylene	2526.427061	946	ug/L	2390	ug/mL
							Analyte Kfw avg	2226.19095				
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	m-Xylene	2646.153846	1300	ug/L	3440	ug/mL
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-3	m-Xylene	3589.160116	966.8	ug/L	3470	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	m-Xylene	1041.247485	3976	ug/L	4140	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-3	m-Xylene	3217.477656	1007	ug/L	3240	ug/mL
							Analyte Kfw avg	2623.509776				
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	p-Xylene	3069.353327	426.8	ug/L	1310	ug/mL
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-3	p-Xylene	4130.643612	312.3	ug/L	1290	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	p-Xylene	1092.564492	1318	ug/L	1440	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-3	p-Xylene	2923.832924	407	ug/L	1190	ug/mL
							Analyte Kfw avg	2804.098589				
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	Total Xylenes	2583.849144	2778.8	ug/L	7180	ug/mL
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-3	Total Xylenes	3483.091787	2070	ug/L	7210	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	Total Xylenes	1023.640946	8333	ug/L	8530	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-3	Total Xylenes	2889.830508	2360	ug/L	6820	ug/mL
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	1,2,3-Trimethylbenzene	5854.844744	534.6	ug/L	3130	ug/mL
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-3	1,2,3-Trimethylbenzene	7158.948686	399.5	ug/L	2860	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	1,2,3-Trimethylbenzene	1868.75	1600	ug/L	2990	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-3	1,2,3-Trimethylbenzene	7793.814433	485	ug/L	3780	ug/mL
							Analyte Kfw avg	5669.089466				
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	1,2,4-Trimethylbenzene	7134.502924	1026	ug/L	7320	ug/mL
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-3	1,2,4-Trimethylbenzene	9188.298311	728.1	ug/L	6690	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	1,2,4-Trimethylbenzene	1980.164159	2924	ug/L	5790	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-3	1,2,4-Trimethylbenzene	7853.290183	927	ug/L	7280	ug/mL
							Analyte Kfw avg	6539.063894				
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	1,3,5-Trimethylbenzene	7959.479016	276.4	ug/L	2200	ug/mL
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-3	1,3,5-Trimethylbenzene	10244.64832	196.2	ug/L	2010	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	1,3,5-Trimethylbenzene	3951.890034	873	ug/L	3450	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-3	1,3,5-Trimethylbenzene	15807.69231	260	ug/L	4110	ug/mL
1							Analyte Kfw avg	9490.927419				

Event	Lab Code	Site Name	Fuel Type	Spill Date	Date	FPLocid	Analyte	Kfw	GW Results	GW UNITS	FP Results	Units
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	Naphthalene	3336.703741	395.6	ug/L	1320	ug/mL
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-3	Naphthalene	3860.640301	318.6	ug/L	1230	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	Naphthalene	1172.774869	955	ug/L	1120	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-3	Naphthalene	4528.301887	318	ug/L	1440	ug/mL
		•					Analyte Kfw avg	3224.6052		-		
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	1-MethylNaphthalene	8788.28229	150.2	ug/L	1320	ug/mL
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-3	1-MethylNaphthalene	8645.533141	138.8	ug/L	1200	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	1-MethylNaphthalene	1513.828239	687	ug/L	1040	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-3	1-MethylNaphthalene	7789.473684	190	ug/L	1480	ug/mL
		-					Analyte Kfw avg	6684.279339				
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-2	2-MethylNaphthalene	10333.4899	212.9	ug/L	2200	ug/mL
2	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/11/98	SH98-1610-3	2-MethylNaphthalene	10107.19755	195.9	ug/L	1980	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-2	2-MethylNaphthalene	2814.070352	597	ug/L	1680	ug/mL
1	NRMRL	Bldg 1610, Shaw AFB, SC	JP-4	06/01/94	03/06/97	SHMW1610-3	2-MethylNaphthalene	14821.42857	168	ug/L	2490	ug/mL
		-					Analyte Kfw avg	9519.046594				
1	NRMRL	Pipeline Leak Site, Myrtle Beach AFB, SC	JP-4	01/01/81	03/04/97	MBMW24	Benzene	169.2771084	332	ug/L	56.2	ug/mL
1	NRMRL	Pipeline Leak Site, Myrtle Beach AFB, SC	JP-4	01/01/81	03/04/97	MBMW24	Toluene	2228	2.5	ug/L	5.57	ug/mL
1	NRMRL	Pipeline Leak Site, Myrtle Beach AFB, SC	JP-4	01/01/81	03/04/97	MBMW24	Ethylbenzene	7761.966365	77.3	ug/L	600	ug/mL
1	NRMRL	Pipeline Leak Site, Myrtle Beach AFB, SC	JP-4	01/01/81	03/04/97	MBMW24	o-Xylene	3837.837838	3.7	ug/L	14.2	ug/mL
1	NRMRL	Pipeline Leak Site, Myrtle Beach AFB, SC	JP-4	01/01/81	03/04/97	MBMW24	m-Xylene	36345.38153	49.8	ug/L	1810	ug/mL
1	NRMRL	Pipeline Leak Site, Myrtle Beach AFB, SC	JP-4	01/01/81	03/04/97	MBMW24	p-Xylene	76341.46341	4.1	ug/L	313	ug/mL
1	NRMRL	Pipeline Leak Site, Myrtle Beach AFB, SC	JP-4	01/01/81	03/04/97	MBMW24	Total Xylenes	37104.16667	57.6	ug/L	2137.2	ug/mL
1	NRMRL	Pipeline Leak Site, Myrtle Beach AFB, SC	JP-4	01/01/81	03/04/97	MBMW24	1,2,3-Trimethylbenzene	33074.36182	90.1	ug/L	2980	ug/mL
1	NRMRL	Pipeline Leak Site, Myrtle Beach AFB, SC	JP-4	01/01/81	03/04/97	MBMW24	1,2,4-Trimethylbenzene	31100.47847	209	ug/L	6500	ug/mL
1	NRMRL	Pipeline Leak Site, Myrtle Beach AFB, SC	JP-4	01/01/81	03/04/97	MBMW24	1,3,5-Trimethylbenzene	84023.66864	33.8	ug/L	2840	ug/mL
1	NRMRL	Pipeline Leak Site, Myrtle Beach AFB, SC	JP-4	01/01/81	03/04/97	MBMW24	Naphthalene	7193.548387	124	ug/L	892	ug/mL
1	NRMRL	Pipeline Leak Site, Myrtle Beach AFB, SC	JP-4	01/01/81	03/04/97	MBMW24	1-MethylNaphthalene	5116.27907	258	ug/L	1320	ug/mL
1	NRMRL	Pipeline Leak Site, Myrtle Beach AFB, SC	JP-4	01/01/81	03/04/97	MBMW24	2-MethylNaphthalene	9141.414141	198	ug/L	1810	ug/mL
3	NRMRL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	Benzene	0.677506775	36.9	ug/L	0.025	ug/mL
3	NRMRL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	Toluene	230.7692308	884	ug/L	204	ug/mL
3	NRMRL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	Ethylbenzene	1600	535	ug/L	856	ug/mL
3	NRMRL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	o-Xylene	2702.702703	370	ug/L	1000	ug/mL
3	NRMRL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	m-Xylene	2552.521008	952	ug/L	2430	ug/mL
3	NRMRL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	p-Xylene	1943.7751	498	ug/L	968	ug/mL
3	NRMRL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	Total Xylenes	2416.483516	1820	ug/L	4398	ug/mL
3	NRMRL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	1,2,3-Trimethylbenzene	7111.940299	134	ug/L	953	ug/mL
3	NRMRL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	1,2,4-Trimethylbenzene	7465.753425	292	ug/L	2180	ug/mL
3	NRMRL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	1,3,5-Trimethylbenzene	12530.12048	83	ug/L	1040	ug/mL
3	NRMRL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	Naphthalene	3324.503311	151	ug/L	502	ug/mL
3	NRMRL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	1-MethylNaphthalene	10218.9781	137	ug/L	1400	ug/mL
3	NRMRL	DFSP-Charleston, Tank 1 Area, Hanahan, SC	JP-4	10/01/75	05/16/97	CH-MW-103	2-MethylNaphthalene	17821.78218	101	ug/L	1800	ug/mL
1	NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW306-FP	Benzene	47.61904762	56.7	ug/L	2.7	ug/mL
1	NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW316-FP	Benzene	103.6866359	8680	ug/L	900	ug/mL
							Analyte Kfw avg	75.65284178				
1	NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW306-FP	Toluene	10.86956522	2.3	ug/L	0.025	ug/mL
1	NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW316-FP	Toluene	9.259259259	2.7	ug/L	0.025	ug/mL
							Analyte Kfw avg	10.06441224				
1	NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW306-FP	Ethylbenzene	4428.571429	560	ug/L	2480	ug/mL
1	NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW316-FP	Ethylbenzene	4204.545455	704	ug/L	2960	ug/mL
							Analyte Kfw avg	4316.558442				
1	NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW306-FP	o-Xylene	2883.959044	586	ug/L	1690	ug/mL

Event Lab Code	Site Name	Fuel Type	Spill Date	Date	FPLocid	Analyte	Kfw	GW Results	GW UNITS	FP Results	Units
1 NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW316-FP	o-Xylene	9486.166008	253	ug/L	2400	ug/mL
						Analyte Kfw avg	6185.062526				
1 NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW306-FP	m-Xylene	2663.316583	199	ug/L	530	ug/mL
1 NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW316-FP	m-Xylene	5404.747413	1643	ug/L	8880	ug/mL
						Analyte Kfw avg	4034.031998				
1 NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW306-FP	p-Xylene	4138.418079	708	ug/L	2930	ug/mL
1 NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW316-FP	p-Xylene	5080.147965	811	ug/L	4120	ug/mL
			10/01/70	00/07/07		Analyte Kfw avg	4609.283022			= 1 = 0	
1 NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW306-FP	Total Xylenes	3449.430676	1493	ug/L	5150	ug/mL
1 NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW316-FP	Total Xylenes	5688.954562	2707	ug/L	15400	ug/mL
		10.4	40/04/70	00/07/07		Analyte Ktw avg	4569.192619	400		4050	
	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKIVIVV306-FP	1,2,3-1 rimethylbenzene	10156.25	192	ug/L	1950	ug/mL
	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKIVIVV316-FP	1,2,3-Trimetriyibenzene	12932.33083	200	ug/L	3440	ug/mL
	Spill Site No. 2. Eaker AEP. AP.		10/01/72	00/27/07		1.2.4 Trimothylbonzono	12257 44209	571	ua/I	7570	ua/ml
	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	00/27/97	EAKIVIV 300-FP	1,2,4-Trimethylbenzene	15207.44306	571	ug/L	7570	ug/mL
	Spill Site No. 2, Lakel Al B, AK	51 -4	10/01/73	00/21/31	LANWWSTOTT	Analyte Kfw avg	14279 99543	020	ug/L	3010	ug/IIIL
1 NRMRI	Spill Site No. 2. Eaker AEB. AB	IP-4	10/01/73	08/27/97	EAKMW306-EP	1 3 5-Trimethylbenzene	12604 79042	334	ua/l	4210	ua/ml
1 NRMRI	Spill Site No. 2, Eaker AFB, AR	.IP-4	10/01/73	08/27/97	EAKMW316-FP	1 3 5-Trimethylbenzene	15859 87261	314	ug/L	4980	ug/mL
			10/01/10	00/21/01	L/ a antio to th	Analyte Kfw avg	14232 33152	011	ug/L	1000	ug/IIIE
1 NRMRL	Spill Site No. 2. Eaker AFB. AR	JP-4	10/01/73	08/27/97	EAKMW306-FP	Naphthalene	6521.73913	161	ua/L	1050	ua/mL
1 NRMRL	Spill Site No. 2. Eaker AFB. AR	JP-4	10/01/73	08/27/97	EAKMW316-FP	Naphthalene	7633.587786	131	ua/L	1000	ua/mL
						Analyte Kfw avg	7077.663458		g-		- g
1 NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW306-FP	1-MethylNaphthalene	18859.64912	114	ug/L	2150	ug/mL
1 NRMRL	Spill Site No. 2. Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW316-FP	1-MethylNaphthalene	23781.38848	67.7	ua/L	1610	ua/mL
		-				Analyte Kfw avg	21320.5188		J		
1 NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW306-FP	2-MethylNaphthalene	21823.20442	181	ug/L	3950	ug/mL
1 NRMRL	Spill Site No. 2, Eaker AFB, AR	JP-4	10/01/73	08/27/97	EAKMW316-FP	2-MethylNaphthalene	30176.89906	96.1	ug/L	2900	ug/mL
						Analyte Kfw avg	26000.05174				
1 NRMRL	Tank Farm C, Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BFT-401-3	Benzene	637.1428571	3.5	ug/L	2.23	ug/mL
1 NRMRL	Tank Farm C, Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BFT-401-3	Benzene	45.41751527	49.1	ug/L	2.23	ug/mL
						Analyte Kfw avg	341.2801862				
1 NRMRL	Tank Farm C, Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BFT-401-3	Toluene	118.1818182	110	ug/L	13	ug/mL
1 NRMRL	Tank Farm C, Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BFT-401-3	Toluene	572.6872247	22.7	ug/L	13	ug/mL
						Analyte Kfw avg	345.4345214				
1 NRMRL	Tank Farm C, Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BFT-401-3	Ethylbenzene	958.677686	121	ug/L	116	ug/mL
1 NRMRL	Tank Farm C, Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BFT-401-3	Ethylbenzene	563.1067961	206	ug/L	116	ug/mL
	Test Free O. Dess (est MOAO, OO	10.5	00/04/00	00/40/07		Analyte Kfw avg	760.892241	470		007	
	Tank Farm C, Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BFT-401-3	o-Xylene	1630.681818	1/6	ug/L	287	ug/mL
INRMIRL	Tank Farm C, Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BF1-401-3	0-Xylene	934.8534202	307	ug/L	287	ug/mL
	Tank Form C. Booutart MCAS. SC	ID 5	06/01/00	00/10/07	DET 401 2	m Yulano	1202.707019	202		224	ua/ml
	Tank Farm C. Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BFT-401-3	m-Xylene	746 6666667	203	ug/L	224	ug/mL
	Tank Fame, Deadloft MCAS, SC	51-5	00/01/30	00/12/37	DI 1-401-5		925.0574713	500	ug/L	224	ug/IIIL
	Tank Farm C. Beaufort MCAS. SC	ID-5	06/01/00	08/12/07	BET-401-3		925.0574715	60	ug/l	00.6	ua/ml
	Tank Farm C. Beaufort MCAS, SC	IP-5	06/01/90	08/12/97	BFT-401-3	p-Xylene	1048 421053	95	ug/L	99.0	ug/mL
		01 0	00/01/30	00/12/31	5114013	Analyte Kfw avg	1245 949657		ug/L	55.0	ug/IIIL
1 NRMRI	Tank Farm C. Beaufort MCAS. SC	JP-5	06/01/90	08/12/97	BFT-401-3	1.2.3-Trimethylbenzene	3379 72167	503	ua/l	1700	ua/ml
1 NRMRI	Tank Farm C. Beaufort MCAS. SC	JP-5	06/01/90	08/12/97	BFT-401-3	1 2 3-Trimethylbenzene	3695 652174	460	ug/l	1700	ug/ml
			00,01,00	55, 12, 51		Analyte Kfw avo	3537,686922			.,	9, L
1 NRMRL	Tank Farm C, Beaufort MCAS. SC	JP-5	06/01/90	08/12/97	BFT-401-3	1,2,4-Trimethylbenzene	3264.659271	631	ug/L	2060	ug/mL
1 NRMRL	Tank Farm C, Beaufort MCAS. SC	JP-5	06/01/90	08/12/97	BFT-401-3	1,2,4-Trimethylbenzene	3450.586265	597	ug/L	2060	ug/mL
		-				Analyte Kfw avg	3357.622768				Ŭ

Event Lab Code	Site Name	Fuel Type	Spill Date	Date	FPLocid	Analyte	Kfw	GW Results	GW UNITS	FP Results	Units
1 NRMRL	Tank Farm C. Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BFT-401-3	1.3.5-Trimethylbenzene	6793.650794	126	ug/L	856	ua/mL
1 NRMRL	Tank Farm C. Beaufort MCAS. SC	JP-5	06/01/90	08/12/97	BFT-401-3	1.3.5-Trimethylbenzene	7575.221239	113	ug/L	856	ua/mL
	,					Analyte Kfw avg	7184,436016		- 3 -		
1 NRMRL	Tank Farm C. Beaufort MCAS. SC	JP-5	06/01/90	08/12/97	BFT-401-3	Naphthalene	1654.545455	275	ua/L	455	ua/mL
1 NRMRL	Tank Farm C. Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BFT-401-3	Naphthalene	1472.491909	309	ug/L	455	ua/mL
						Analyte Kfw avg	1563.518682				
1 NRMRL	Tank Farm C, Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BFT-401-3	1-MethylNaphthalene	6000	215	ug/L	1290	ug/mL
1 NRMRL	Tank Farm C, Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BFT-401-3	1-MethylNaphthalene	5512.820513	234	ug/L	1290	ug/mL
						Analyte Kfw avg	5756.410256		- Ŭ		Ŭ
1 NRMRL	Tank Farm C, Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BFT-401-3	2-MethylNaphthalene	8720.930233	172	ug/L	1500	ug/mL
1 NRMRL	Tank Farm C, Beaufort MCAS, SC	JP-5	06/01/90	08/12/97	BFT-401-3	2-MethylNaphthalene	7936.507937	189	ug/L	1500	ug/mL
						Analyte Kfw avg	8328.719085				
	GROUNDWATER DATA BASE REPORT DI	D NOT INCLU	DE DATA F	OR CECI	L FIELD						
	DATA BELOW ENTERED BY HAND										
NRMRL	Facility 293, Cecil Field NAS, FL	JP-5	06/01/81	05/20/97	CEF-293-9FP	Benzene	261.1111111	90	ug/L	23.5	ug/mL
						Toluene	151.552795	805	ug/L	122	ug/mL
						Ethylbenzene	2669.491525	944	ug/L	2520	ug/mL
						o-Xylene	635.2705411	499	ug/L	317	ug/mL
						m-Xylene	1866.666667	1500	ug/L	2800	ug/mL
						p-Xylene	2220.744681	752	ug/L	1670	ug/mL
						I otal xylenes	1740.094511	2751	ug/L	4787	ug/mL
						1,2,3-1 MB	6549.295775	284	ug/L	1860	ug/mL
						1,2,4-1MB	6896.551724	580	ug/L	4000	ug/mL
						1,3,5-1 MB	7734.80663	181	ug/L	1400	ug/mL
						1 Mathulaanhthalana	10700.00301	277	ug/L	1000	ug/mL
						2 Mothylpaphthalono	21/10 25/9/	100	ug/L	2290	ug/mL
						2-Metrymaphthalene	21419.33404	100	ug/L	3320	ug/IIIL
4 NRMRI	Bldg 4522 Seymour Johnson AFB SC	IP-8	12/01/95	03/10/98	S.198-MP2	Benzene	58 70348139	833	ua/l	48 9	ua/ml
3 NRMRI	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW1SEP	Benzene	228 7735849	848	ug/L	194	ug/mL
0.1111112		0.0	12/01/00	00/10/01		Analyte Kfw avg	143 7385331	0.0	ug/2		ag/2
4 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MP2	Toluene	220.3038674	2896	ua/L	638	ua/mL
3 NRMRL	Bldg 4522. Seymour Johnson AFB. SC	JP-8	12/01/95	05/15/97	SJMW1SFP	Toluene	251.2195122	4100	ua/L	1030	ua/mL
						Analyte Kfw avg	235.7616898				
4 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MP2	Ethylbenzene	1448.040886	587	ug/L	850	ug/mL
3 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW1SFP	Ethylbenzene	1389.548694	842	ug/L	1170	ug/mL
						Analyte Kfw avg	1418.79479				
4 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MP2	o-Xylene	1458.990536	760.8	ug/L	1110	ug/mL
3 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW1SFP	o-Xylene	1860.465116	946	ug/L	1760	ug/mL
						Analyte Kfw avg	1659.727826				
4 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MP2	m-Xylene	1165.123457	1296	ug/L	1510	ug/mL
3 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW1SFP	m-Xylene	1890.909091	1650	ug/L	3120	ug/mL
						Analyte Kfw avg	1528.016274				
4 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MP2	p-Xylene	1360.91954	435	ug/L	592	ug/mL
3 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW1SFP	p-Xylene	1728.971963	642	ug/L	1110	ug/mL
						Analyte Kfw avg	1544.945751				
	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MP2	Total xylenes	1289.028012	2491.8	ug/L	3212	ug/mL
	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW1SFP	Total xylenes	1849.90735	3238	ug/L	5990	ug/mL
						Analyte Kfw avg	1569.467681				
4 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MP2	1,2,3-Trimethylbenzene	6075.619296	383.5	ug/L	2330	ug/mL
3 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW1SFP	1,2,3-Trimethylbenzene	6340.32634	429	ug/L	2720	ug/mL
						Analyte Kfw avg	6207.972818				

Event Lab Code	Site Name	Fuel Type	Spill Date	Date	FPLocid	Analyte	Kfw	GW Results	GW UNITS	FP Results	Units
4 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MP2	1,2,4-Trimethylbenzene	6281.371002	697.3	ug/L	4380	ug/mL
3 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW1SFP	1,2,4-Trimethylbenzene	7269.180754	769	ug/L	5590	ug/mL
						Analyte Kfw avg	6775.275878				
4 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MP2	1,3,5-Trimethylbenzene	6284.153005	183	ug/L	1150	ug/mL
3 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW1SFP	1,3,5-Trimethylbenzene	8108.108108	259	ug/L	2100	ug/mL
						Analyte Kfw avg	7196.130557				
4 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MP2	Naphthalene	3557.264265	247.1	ug/L	879	ug/mL
3 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW1SFP	Naphthalene	5039.370079	254	ug/L	1280	ug/mL
						Analyte Kfw avg	4298.317172				
4 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MP2	1-MethylNaphthalene	11759.34366	109.7	ug/L	1290	ug/mL
3 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW1SFP	1-MethylNaphthalene	9496.855346	159	ug/L	1510	ug/mL
						Analyte Kfw avg	10628.09951				
4 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	03/10/98	SJ98-MP2	2-MethylNaphthalene	13361.46273	142.2	ug/L	1900	ug/mL
3 NRMRL	Bldg 4522, Seymour Johnson AFB, SC	JP-8	12/01/95	05/15/97	SJMW1SFP	2-MethylNaphthalene	16962.96296	135	ug/L	2290	ug/mL
						Analyte Kfw avg	15162.21285				
1 NRMRL	JP-8 Release Site, Pope AFB, NC	JP-8	04/01/96	07/17/96	SOURCE AREA	Benzene	50	0.5	ug/L	0.025	ug/mL
1 NRMRL	JP-8 Release Site, Pope AFB, NC	JP-8	04/01/96	07/17/96	SOURCE AREA	Benzene	50	0.5	ug/L	0.025	ug/mL
						Analyte Kfw avg	50				
1 NRMRL	JP-8 Release Site, Pope AFB, NC	JP-8	04/01/96	07/17/96	SOURCE AREA	Toluene	1.25	20	ug/L	0.025	ug/mL
1 NRMRL	JP-8 Release Site, Pope AFB, NC	JP-8	04/01/96	07/17/96	SOURCE AREA	Toluene	25	1	ug/L	0.025	ug/mL
						Analyte Kfw avg	13.125				
1 NRMRL	JP-8 Release Site, Pope AFB, NC	JP-8	04/01/96	07/17/96	SOURCE AREA	Ethylbenzene	9000	7	ug/L	63	ug/mL
1 NRMRL	JP-8 Release Site, Pope AFB, NC	JP-8	04/01/96	07/17/96	SOURCE AREA	Ethylbenzene	1968.75	32	ug/L	63	ug/mL
						Analyte Kfw avg	5484.375				
1 NRMRL	JP-8 Release Site, Pope AFB, NC	JP-8	04/01/96	07/17/96	SOURCE AREA	o-Xylene	13727.27273	33	ug/L	453	ug/mL
1 NRMRL	JP-8 Release Site, Pope AFB, NC	JP-8	04/01/96	07/17/96	SOURCE AREA	o-Xylene	4768.421053	95	ug/L	453	ug/mL
						Analyte Kfw avg	9247.84689				
1 NRMRL	JP-8 Release Site, Pope AFB, NC	JP-8	04/01/96	07/17/96	SOURCE AREA	Total Xylenes (m,p, and o)	3988.095238	252	ug/L	1005	ug/mL
1 NRMRL	JP-8 Release Site, Pope AFB, NC	JP-8	04/01/96	07/17/96	SOURCE AREA	Total Xylenes (m,p, and o)	15000	67	ug/L	1005	ug/mL
						Analyte Kfw avg	9494.047619				

APPENDIX C-3

RESIDUAL LNAPL CALCULATIONS

Lab Code	Fuel Type	Spill Date	Date	Site Name	Matix	Locid	Analyte	Results	Units	Flag FC Analyte	FC Ratio	FPLocid	FPdensity FPunits	SoilFuel (ug/mL)
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	CPT1-39.5	Benzene	42.8	8 mg/kg	B/FC	0.021725888	MW349-1	738000 ug/mL	16034
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	CPT1-39.5	Toluene	105	i mg/kg	T/FC	0.053299492	MW349-1	738000 ug/mL	39335
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	CPT1-39.5	Ethylbenzene	24.4	l mg/kg	E/FC	0.012385787	MW349-1	738000 ug/mL	9141
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	CPT1-39.5	m-Xylene	42.4	l mg/kg	m-Xylene/FC	0.021522843	MW349-1	738000 ug/mL	15884
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	CPT1-39.5	o-Xylene	24.3	8 mg/kg	o-Xylene/FC	0.012335025	MW349-1	738000 ug/mL	9103
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	CPT1-39.5	p-Xylene	19.3	8 mg/kg	p-Xylene/FC	0.009796954	MW349-1	738000 ug/mL	7230
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	CPT1-39.5	Total Xylenes (m,p, and o)	86	6 mg/kg	X/FC	0.043654822	MW349-1	738000 ug/mL	32217
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	CPT1-39.5	Total BTEX	258.2	2 mg/kg	BTEX/FC	0.13106599	MW349-1	738000 ug/mL	96727
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	CPT1-39.5	B+I	147.8	8 mg/kg	(B+T)/FC	0.075025381	MW349-1	738000 ug/mL	55369
NRMRL	Gasoline	6/1/1990	*****	Tank 349, Offutt AFB, NE	Soll	CPT1-39.5	E+X	110.4	i mg/кg	(E+X)/FC	0.056040609	MW349-1	738000 ug/mL	41358
NRMRL	Gasoline	6/1/1990	*****	Tank 349, Onutt AFB, NE	Soll	CPT1-39.5	B/I B/E	0.40761905		(B/T)/FC	0.000206913	NIVV349-1	738000 ug/mL	153
	Gasoline	6/1/1990	*****	Tarik 349, Offutt AEP, NE	Soil	CPT1-39.5	B/E B/Y	1.75409630)		0.000890405	MW249-1	738000 ug/mL	007
	Gasoline	6/1/1990	########	Tank 349, Offutt AEP, NE	Soil	CPT1 20.5		4 20227960			0.000232027	MM/240 1	738000 ug/mL	1612
NRMRI	Gasoline	6/1/1990	########	Tank 349, Offutt AFB, NE	Soil	CPT1-39.5	T/X	1 22093023	2	(T/Z)/FC	0.002104403	MW349-1	738000 ug/mL	457
NRMRI	Gasoline	6/1/1990	#######	Tank 349 Offutt AFB NE	Soil	CPT1-39.5	F/X	0.28372093	3	(F/X)/FC	0.000144021	MW349-1	738000 ug/mL	106
NRMRL	Gasoline	6/1/1990	#######	Tank 349. Offutt AFB, NE	Soil	CPT1-39.5	(B+T)/(E+X)	1.33876812	2	((B+T)/(E+X))/FC	0.000679578	MW349-1	738000 ug/mL	502
NRMRL	Gasoline	6/1/1990	######	Tank 349. Offutt AFB. NE	Soil	CPT1-39.5	Fuel Carbon	1970) ma/ka	TP Fuel Carbon	1	MW349-1	738000 ug/mL	738000
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-39'	Benzene	0.562	2 mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-39'	Toluene	0.334	l mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-39'	Ethylbenzene	0.291	mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-39'	m-Xylene	0.42	2 mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-39'	o-Xylene	0.147	′ mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-39'	p-Xylene	0.224	l mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-39'	Total Xylenes (m,p, and o)	0.791	mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-39'	Total BTEX	1.978	8 mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-39'	B+T	0.896	6 mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-39'	E+X	1.082	2 mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-39'	B/T	1.68263473	3					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-39	B/E	1.93127148	3					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-39	B/X	0.71049305	2					
NRMRL	Gasoline	6/1/1990	*****	Tank 349, Offutt AFB, NE	Soll	OFSB-1-39	1/E	1.14776632	2					
NRMRL	Gasoline	6/1/1990	*****	Tank 349, Onutt AFB, NE	Soll	OFSB-1-39	1/X	0.42225032	-					
	Gasoline	6/1/1990	#######	Tank 349, Ollull AFB, NE	Soil	OFSB-1-39		0.30700073))					
	Gasoline	6/1/1990	#######	Tank 349, Ollull AFB, NE	Soil	OFSB-1-39	(D+1)/(E+A)	0.02009012	ma/ka					
	Gasoline	6/1/1990	#######	Tank 349, Oldut AFB, NE	Soil	OFSB-1-39	1,2,3-Trimethylbenzene	0.044	ing/kg					
NRMRI	Gasoline	6/1/1990	########	Tank 349, Offutt AFB, NE	Soil	OFSB-1-39'	1 3 5-Trimethylbenzene	0.233	S ma/ka					
NRMRI	Gasoline	6/1/1990	#######	Tank 349 Offutt AFB NE	Soil	OFSB-1-39'	1-MethylNaphthalene	0.0118	ma/ka					
NRMRL	Gasoline	6/1/1990	#######	Tank 349. Offutt AFB, NE	Soil	OFSB-1-39'	2-MethylNaphthalene	0.0315	i ma/ka					
NRMRL	Gasoline	6/1/1990	#######	Tank 349. Offutt AFB, NE	Soil	OFSB-1-40'	Benzene	40.2	2 ma/ka					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-40'	Toluene	165	i mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-40'	Ethylbenzene	58.7	′ mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-40'	m-Xylene	99.8	8 mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-40'	o-Xylene	55.2	2 mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-40'	p-Xylene	48	8 mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-40'	Total Xylenes (m,p, and o)	203	8 mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-40'	Total BTEX	466.9) mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-40'	B+T	205.2	2 mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-40'	E+X	261.7	′ mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-40'	B/T	0.24363636	5					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-40'	B/E	0.68483816	5					
	Gasoline	6/1/1990	########	Tank 349, UTUIT AFB, NE	501	UFSB-1-40	B/X	0.19802956						
NRMRL	Gasoline	6/1/1990	*****	Tank 349, Offutt AFB, NE	Soll	OFSB-1-40	1/E	2.8109029	,					
	Gasoline	6/1/1990	######################################	Tank 349, Ullutt AFB, NE	Soil	OFSB-1-40		0.29016256						
	Gasolino	6/1/1990	~~~~~~~	Tank 349, Offutt AEB NE	Soil	OFSB-1-40	L/A (B+T)/(E+Y)	0.20910250			1			
NRMRI	Gasoline	6/1/1990	######################################	Tank 349 Offutt AFB, NE	Soil	OFSB-1-40	1 2 3-Trimethylbenzene	17 6	i ma/ka					
NRMRI	Gasoline	6/1/1990	########	Tank 349 Offutt AFB NF	Soil	OFSB-1-40'	1 2 4-Trimethylbenzene	66	i ma/ka					
NRMRL	Gasoline	6/1/1990	#######	Tank 349. Offutt AFB, NE	Soil	OFSB-1-40'	1.3.5-Trimethylbenzene	22.3	3 ma/ka					
NRMRL	Gasoline	6/1/1990	#######	Tank 349. Offutt AFB, NE	Soil	OFSB-1-40'	1-MethylNaphthalene	5.59) ma/ka					
NRMRL	Gasoline	6/1/1990	#######	Tank 349, Offutt AFB, NE	Soil	OFSB-1-40'	2-MethylNaphthalene	11.7	' ma/ka		1			
NRMRL	Gasoline	6/1/1990	#######	Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	Benzene	19.2	2 mg/ka					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	Toluene	83.2	2 mg/kg					
NRMRL	Gasoline	6/1/1990	######	Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	Ethylbenzene	37	′ mg/kg					

Lab Code	Fuel Type	Spill Date	Date Site Name	Mati	Locid	Analyte	Results Units	Flag	FC Analyte	FC Ratio	FPLocid	FPdensity	FPunits	SoilFuel (ug/mL)
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	m-Xylene	58.7 mg/kg							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	o-Xylene	32.9 mg/kg							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	p-Xylene	28.1 mg/kg							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	Total Xylenes (m,p, and o)	119.7 mg/kg							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	Total BTEX	259.1 mg/kg							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	B+T	102.4 mg/kg							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	E+X	156.7 mg/kg							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	B/T	0.23076923							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	B/E	0.51891892							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	B/X	0.160401							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	T/E	2.24864865							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	T/X	0.69507101							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	E/X	0.3091061							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	(B+T)/(E+X)	0.65347798							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	1,2,3-Trimethylbenzene	11.7 mg/kg							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	1,2,4-Trimethylbenzene	45.6 mg/kg							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	1,3,5-Trimethylbenzene	14 mg/kg							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	1-MethylNaphthalene	2.63 mg/kg							
NRMRL	Gasoline	6/1/1990	###### Tank 349, Offutt AFB, NE	Soil	OFSB-2-39'	2-MethylNaphthalene	5.36 mg/kg							

Lab Code	Fuel Type	Spill Date	Date	Site Name	Matix	Locid	Analyte	Results Units	Flag FC Analyte	FC Ratio	FPLocid	FPdensity	FPunits	SoilFuel (ug/mL)
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	Benzene	0.331 mg/kg	B/FC	0.000129	SH98-1610-2	780000) ug/mL	101
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	Toluene	0.551 mg/kg	T/FC	0.000215	SH98-1610-2	780000	ug/mL	168
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	Ethylbenzene	1.84 mg/kg	E/FC	0.000719	SH98-1610-2	780000) ug/mL	561
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	m-Xylene	5.11 mg/kg	m-Xylene/FC	0.001996	SH98-1610-2	780000	ug/mL	1557
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	o-Xylene	1.86 mg/kg	o-Xylene/FC	0.000727	SH98-1610-2	780000) ug/mL	567
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	p-Xylene	1.85 mg/kg	p-Xylene/FC	0.000723	SH98-1610-2	780000) ug/mL	564
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	Total Xylenes (m,p, and o)	8.82 mg/kg	X/FC	0.003445	SH98-1610-2	780000) ug/mL	2687
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	Total BTEX	11.542 mg/kg	BTEX/FC	0.004509	SH98-1610-2	780000	ug/mL	3517
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	B+T	0.882 mg/kg	(B+T)/FC	0.000345	SH98-1610-2	780000) ug/mL	269
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	E+X	10.66 mg/kg	(E+X)/FC	0.004164	SH98-1610-2	780000) ug/mL	3248
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	B/T	0.60072595	(B/T)/FC	0.000235	SH98-1610-2	780000) ug/mL	183
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	B/E	0.1798913	(B/E)/FC	7.03E-05	SH98-1610-2	780000	ug/mL	55
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	B/X	0.03752834	(B/X)/FC	1.47E-05	SH98-1610-2	780000	ug/mL	11
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	T/E	0.29945652	(T/E)/FC	0.000117	SH98-1610-2	780000	ug/mL	91
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	T/X	0.06247166	(T/X)/FC	2.44E-05	SH98-1610-2	780000	ug/mL	19
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	E/X	0.20861678	(E/X)/FC	8.15E-05	SH98-1610-2	780000	ug/mL	64
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	(B+T)/(E+X)	0.08273921	((B+T)/(E+X))/FC	3.23E-05	SH98-1610-2	780000	ug/mL	25
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	Fuel Carbon	2560 mg/kg	Fuel Carbon	1	SH98-1610-2	780000	ug/mL	780000
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	1.2.3-Trimethylbenzene	3.38 ma/ka	1.2.3 TMB/FC	0.00132	SH98-1610-2	780000	ua/mL	1030
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	1.2.4-Trimethylbenzene	11.5 ma/ka	1.2.4 TMB/FC	0.004492	SH98-1610-2	780000	ua/mL	3504
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	1.3.5-Trimethylbenzene	4.83 ma/ka	1.3.5 TMB/FC	0.001887	SH98-1610-2	780000	ua/mL	1472
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	1-MethylNaphthalene	0.789 ma/ka	1-MN/FC	0.000308	SH98-1610-2	780000	ua/mL	240
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB1-27	2-MethylNaphthalene	1.5 mg/kg	2-MN/FC	0.000586	SH98-1610-2	780000	ua/mL	457
NRMRI	JP-4	6/1/1994	#######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	Benzene	0.394 mg/kg	B/FC	0.000149	SH98-1610-3	777000	ug/mL	116
NRMRI	IP-4	6/1/1994	#######	Bldg 1610, Shaw AEB, SC	Soil	SH98-SB2-27	Toluene	0.335 mg/kg	T/FC	0.000127	SH98-1610-3	777000	ug/mL	90
NIDMDI		6/1/1004	#######	Pidg 1610, Shaw AEP, SC	Soil	SH00 SB2 27	Ethylbonzono	2.0 mg/kg	E/EC	0.000127	SH09 1610 2	777000	ug/mL	11/0
		6/1/1994	#######	Pide 1610, Shaw AFB, SC	Soil	SH90-SB2-27	m Xylono	12.5 mg/kg	E/FC	0.001477	SH90-1010-3	777000		2670
	JF-4	0/1/1994	*****	Bidg 1010, Shaw AFB, SC	0.00	31190-3D2-27	- Videne	7.40 m s/kg	III-Xylene/FC	0.004733	01100 4040 0	777000	ug/mL	3079
NRMRL	JP-4	6/1/1994	#######	Blog 1610, Snaw AFB, SC	501	SH98-SB2-27	o-Xylene	7.18 mg/kg	0-Xylene/FC	0.00272	SH98-1610-3	777000	ug/mL	2113
NRMRL	JP-4	6/1/1994	****	Bldg 1610, Shaw AFB, SC	501	SH98-SB2-27	p-Xylene	4.43 mg/kg	p-Xylene/FC	0.001678	SH98-1610-3	777000	ug/mL	1304
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	SOI	SH98-SB2-27	Total Xylenes (m,p, and o)	24.11 mg/kg	X/FC	0.009133	SH98-1610-3	777000	ug/mL	7096
NRMRL	JP-4	6/1/1994	######	Bidg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	Total BTEX	28.739 mg/kg	BIEX/FC	0.010886	SH98-1610-3	777000	ug/mL	8458
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	B+T	0.729 mg/kg	(B+T)/FC	0.000276	SH98-1610-3	777000) ug/mL	215
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	E+X	28.01 mg/kg	(E+X)/FC	0.01061	SH98-1610-3	777000) ug/mL	8244
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	B/T	1.1761194	(B/T)/FC	0.000445	SH98-1610-3	777000) ug/mL	346
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	B/E	0.10102564	(B/E)/FC	3.83E-05	SH98-1610-3	777000) ug/mL	30
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	B/X	0.01634177	(B/X)/FC	6.19E-06	SH98-1610-3	777000) ug/mL	5
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	T/E	0.08589744	(T/E)/FC	3.25E-05	SH98-1610-3	777000	ug/mL	25
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	T/X	0.01389465	(T/X)/FC	5.26E-06	SH98-1610-3	777000	ug/mL	4
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	E/X	0.16175861	(E/X)/FC	6.13E-05	SH98-1610-3	777000) ug/mL	48
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	(B+T)/(E+X)	0.02602642	((B+T)/(E+X))/FC	9.86E-06	SH98-1610-3	777000	ug/mL	8
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	Fuel Carbon	2640 mg/kg	Fuel Carbon	1	SH98-1610-3	777000	ug/mL	777000
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	1.2.3-Trimethylbenzene	10.3 ma/ka	1.2.3 TMB/FC	0.003902	SH98-1610-3	777000	ua/mL	3031
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	1.2.4-Trimethylbenzene	24.7 ma/ka	1.2.4 TMB/FC	0.009356	SH98-1610-3	777000	ua/mL	7270
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	1.3.5-Trimethylbenzene	7.8 ma/ka	1.3.5 TMB/FC	0.002955	SH98-1610-3	777000	ua/mL	2296
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	1-MethylNaphthalene	4.42 ma/ka	1-MN/FC	0.001674	SH98-1610-3	777000) ua/mL	1301
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SH98-SB2-27	2-MethylNaphthalene	7.11 ma/ka	2-MN/FC	0.002693	SH98-1610-3	777000) ua/mL	2093
NRMRL	JP-4	6/1/1994	#######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-1-33'	Benzene	0.544 mg/kg	B/FC	0.002909	SHMW1610-2	765000	ua/mL	2000
NRMRI	JP-4	6/1/1994	#######	Bldg 1610, Shaw AFB_SC	Soil	SHSB-1-33'	Toluene	0.787 ma/ka	T/FC	0.004209	SHMW1610-2	765000	ug/ml	3220
NRMRI	JP-4	6/1/1994	#######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-1-33'	Ethylbenzene	0.224 mg/kg	E/EC	0.001198	SHMW1610-2	765000	ug/mL	Q16
NRMRI	IP-4	6/1/1994	########	Bldg 1610, Shaw AFB, SC	Soil	SHSB-1-33'	m-Xylene	0.78 mg/kg	m-Xylene/FC	0.004171	SHMW1610-2	765000	ug/mL	2101
NRMRI	IP-4	6/1/1004	######################################	Bidg 1610, Shaw AFB, SC	Soil	SHSB-1-33'	o-Yvlene	0.581 mg/kg	o-Xylene/EC	0.003107	SHMW/1610.2	765000		2191
		6/1/1994	########	Pide 1610, Shaw AFB, SC	Soil	SHSB-1-33	p Xylono	0.361 mg/kg	p Xylene/FC	0.003107	SHWW1010-2	765000		2377
	JF-4	6/1/1994	*****	Bidg 1610, Shaw AFB, SC	Soil	SHSB-1-33	P-Aylene Total Xulanaa (m.n. and a)	0.237 mg/kg	p-xylene/FC	0.001374	SHWW1010-2	765000	ug/mL	1001
NRIVIRL	JP-4	6/1/1994	#######	Blug 1610, Shaw AFB, SC	0.00	SH3D-1-33	Total Aylenes (m,p, and o)	1.616 mg/kg	A/FC	0.008652	SHIVIV 1610-2	765000	ug/mL	0019
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	501	SHSB-1-33	INTER	3.173 mg/kg	BIEX/FC	0.016968	SHIVIV1610-2	765000	ug/mL	12980
NRMRL	JP-4	6/1/1994	######	Bidg 1610, Shaw AFB, SC	Soli	SHSB-1-33	B+1	1.331 mg/kg	(B+1)/FC	0.007118	SHMW1610-2	765000	ug/mL	5445
INKMKL	JP-4	6/1/1994	#######	BID TOTU, SNAW AFB, SC	SOIL	SHSB-1-33	E+A	1.842 mg/kg	(E+X)/FC	0.00985	SHIVIW1610-2	765000	ug/mL	7535
NRMRL	JP-4	6/1/1994	#######	Blog 1610, Shaw AFB, SC	Soll	SHSB-1-33	B/I	0.69123253	(B/T)/FC	0.003696	SHMW1610-2	765000	ug/mL	2828
NRMRL	JP-4	6/1/1994	######	Bidg 1610, Shaw AFB, SC	Soil	SHSB-1-33'	B/E	2.42857143	(B/E)/FC	0.012987	SHMW1610-2	765000	ug/mL	9935
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-1-33'	B/X	0.33621755	(B/X)/FC	0.001798	SHMW1610-2	765000	ug/mL	1375
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-1-33'	T/E	3.51339286	(T/E)/FC	0.018788	SHMW1610-2	765000	ug/mL	14373
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-1-33'	T/X	0.48640297	(T/X)/FC	0.002601	SHMW1610-2	765000	ug/mL	1990
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-1-33'	E/X	0.13844252	(E/X)/FC	0.00074	SHMW1610-2	765000	ug/mL	566
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-1-33'	(B+T)/(E+X)	0.72258415	((B+T)/(E+X))/FC	0.003864	SHMW1610-2	765000	ug/mL	2956
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-1-33'	Fuel Carbon	187 mg/kg	Fuel Carbon	1	SHMW1610-2	765000	ug/mL	765000
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-1-33'	1,2,3-Trimethylbenzene	0.919 mg/kg	1,2,3 TMB/FC	0.004914	SHMW1610-2	765000	ug/mL	3760
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-1-33'	1,2,4-Trimethylbenzene	1.74 mg/kg	1,2,4 TMB/FC	0.009305	SHMW1610-2	765000	ug/mL	7118

Lab Code	Fuel Type	Spill Date	Date	Site Name	Matix	Locid	Analyte	Results	Units	Flag	FC Analyte	FC Ratio	FPLocid	FPdensity FPunits	SoilFuel (ug/mL)
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-1-33'	1,3,5-Trimethylbenzene	0.527	mg/kg		1,3,5 TMB/FC	0.002818	SHMW1610-2	765000 ug/mL	2156
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-1-33'	1-MethylNaphthalene	0.456	mg/kg		1-MN/FC	0.002439	SHMW1610-2	765000 ug/mL	1865
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-1-33'	2-MethylNaphthalene	0.72	mg/kg		2-MN/FC	0.00385	SHMW1610-2	765000 ug/mL	2945
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-2-33'	Benzene	5.08	mg/kg		B/FC	0.002117	SHMW1610-3	783000 ug/mL	1657
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-2-33'	Toluene	10.8	ma/ka		T/FC	0.0045	SHMW1610-3	783000 ug/mL	3524
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-2-33'	Ethylbenzene	3.58	mg/kg		E/FC	0.001492	SHMW1610-3	783000 ug/mL	1168
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-2-33'	m-Xylene	12.2	ma/ka		m-Xvlene/FC	0.005083	SHMW1610-3	783000 ug/mL	3980
NRMRI	IP-4	6/1/1994	#######	Bldg 1610 Shaw AFB SC	Soil	SHSB-2-33'	o-Xvlene	8.92	ma/ka		o-Xylene/FC	0.003717	SHMW1610-3	783000 ug/ml	2910
NRMRI	IP-4	6/1/1994	<u>######</u>	Bldg 1610, Shaw AFB, SC	Soil	SHSB-2-33'	p-Xylene	4 17	ma/ka		n-Xylene/FC	0.000717	SHMW1610-3	783000 ug/mL	1360
NRMRI	IP-4	6/1/100/	#######	Bidg 1610, Shaw AFB, SC	Soil	SHSB-2-33'	Total Xylenes (m.p. and o)	25.20	ma/ka		X/FC	0.010538	SHMW/1610-3	783000 ug/mL	8251
NRMRI	IP-4	6/1/1994	<u>######</u>	Bldg 1610, Shaw AFB, SC	Soil	SHSB-2-33'	Total BTEX	44 75	ma/ka		BTEX/FC	0.018646	SHMW1610-3	783000 ug/mL	14600
NRMRI	IP-4	6/1/1004	****	Bidg 1610, Shaw AFB, SC	Soil	SHSB-2-33'	B+T	15.88	mg/kg		(B+T)/FC	0.006617	SHMW/1610-3	783000 ug/mL	5181
NRMRI	JP-4	6/1/100/	############	Bidg 1610, Shaw AFB, SC	Soil	SHSB-2-33'	E+Y	28.87	mg/kg		(E+Y)/FC	0.000017	SHMW/1610-3	783000 ug/mL	9/10
		6/1/1004	######################################	Bidg 1610, Shaw AFB, SC	Soil	SHOD-2-33	B/T	0.47027027	iiig/kg			0.0012023		783000 ug/mL	152
		6/1/1994	######################################	Pidg 1610, Shaw AFB, SC	Soil	SH3D-2-33	B/T	1 41900441			(B/T)/FC	0.000190		783000 ug/mL	100
	JF-4	6/1/1994	******	Bidg 1610, Shaw AFB, SC	Soil	SHSB-2-33	B/E	0.20086001				0.000391		783000 ug/mL	403
NRIVIRL	JP-4	6/1/1994	****	Blug 1610, Shaw AFB, SC	30II	SH3D-2-33	B/A	0.20086991				0.37E-05		783000 ug/mL	00
NRMRL	JP-4	6/1/1994	****	Bidg 1610, Shaw AFB, SC	501	SHSB-2-33	T/E	3.016/59/8			(1/E)/FC	0.001257	SHIVIV1610-3	783000 ug/mL	984
NRMRL	JP-4	6/1/1994	****	Bidg 1610, Snaw AFB, SC	501	SHSB-2-33	1/X	0.42704626			(1/X)/FC	0.000178	SHIVIV1610-3	783000 ug/mL	139
NRMRL	JP-4	6/1/1994	######	Bidg 1610, Shaw AFB, SC	Soll	SHSB-2-33	E/X	0.14155793			(E/X)/FC	5.9E-05	SHMW1610-3	783000 ug/mL	46
NRMRL	JP-4	6/1/1994	######	Bidg 1610, Shaw AFB, SC	Soll	SHSB-2-33	(B+1)/(E+X)	0.55005196			((B+T)/(E+X))/FC	0.000229	SHMW1610-3	783000 ug/mL	179
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-2-33'	Fuel Carbon	2400	mg/kg		Fuel Carbon	1 5	SHMW1610-3	783000 ug/mL	783000
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-2-33'	1,2,3-Trimethylbenzene	13.3	mg/kg		1,2,3 TMB/FC	0.005542	SHMW1610-3	783000 ug/mL	4339
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-2-33'	1,2,4-Trimethylbenzene	25.2	mg/kg		1,2,4 TMB/FC	0.0105	SHMW1610-3	783000 ug/mL	8222
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-2-33'	1,3,5-Trimethylbenzene	13.5	mg/kg		1,3,5 TMB/FC	0.005625	SHMW1610-3	783000 ug/mL	4404
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-2-33'	1-MethylNaphthalene	5.31	mg/kg		1-MN/FC	0.002213	SHMW1610-3	783000 ug/mL	1732
NRMRL	JP-4	6/1/1994	######	Bldg 1610, Shaw AFB, SC	Soil	SHSB-2-33'	2-MethylNaphthalene	8.72	mg/kg		2-MN/FC	0.003633	SHMW1610-3	783000 ug/mL	2845
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	Benzene	0.006	mg/kg	**	B/FC	1 (CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	Toluene	0.0173	mg/kg		T/FC	2.883333	CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	Ethylbenzene	0.006	mg/kg	**	E/FC	1 (CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	m-Xylene	0.006	mg/kg	**	m-Xylene/FC	1 (CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	o-Xylene	0.006	mg/kg	**	o-Xylene/FC	1 (CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	p-Xylene	0.006	mg/kg	**	p-Xylene/FC	1 (CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	Total Xylenes (m,p, and o)	0.018	mg/kg		X/FC	3 (CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	Total BTEX	0.0473	mg/kg		BTEX/FC	7.883333	CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	B+T	0.0233	mg/kg		(B+T)/FC	3.883333	CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	E+X	0.024	mg/kg		(E+X)/FC	4 (CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	B/T	0.34682081			(B/T)/FC	57.80347	CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	B/E	1			(B/E)/FC	166.6667	CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	B/X	0.333333333			(B/X)/FC	55.55556	CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	T/E	2.88333333			(T/E)/FC	480.5556	CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	T/X	0.96111111			(T/X)/FC	160.1852	CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5	E/X	0.333333333			(E/X)/FC	55,55556	CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5	(B+T)/(E+X)	0.97083333			((B+T)/(E+X))/FC	161.8056	CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5	Fuel Carbon	0.006	ma/ka	**	Fuel Carbon	1 (CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRL	JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5	1.2.3-Trimethylbenzene	0.006	ma/ka	**	1.2.3 TMB/FC	1 (CH-MW-103	760000 ug/mL	fuel carbon = nd
NRMRI	JP-4	10/1/1975	######	DESP-Charleston Tank 1 Area Hanahan SC	Soil	CHSB1-14.5'	1 2 4-Trimethylbenzene	0.006	ma/ka	**	1 2 4 TMB/FC	1 (CH-MW-103	760000 ug/ml	fuel carbon = nd
NRMRL	JP-4	10/1/1975	#######	DESP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	1.3.5-Trimethylbenzene	0.006	ma/ka	**	1.3.5 TMB/FC	1 0	CH-MW-103	760000 ug/ml	fuel carbon = nd
NRMRI	IP-4	10/1/1975	****	DESP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB1-14.5'	1-MethylNaphthalene	0.006	ma/ka	**	1-MN/EC	1 (CH-MW-103	760000 ug/mL	fuel carbon - nd
NRMRI	.IP-4	10/1/1975	#######	DESP-Charleston Tank 1 Area Hanahan SC	Soil	CHSB1-14.5	2-MethylNaphthalene	0.006	ma/ka	**	2-MN/FC	1 0	CH-MW-103	760000 ug/mL	fuel carbon - nd
NRMRI	IP-4	10/1/1975	<u>######</u> #	DESP-Charleston Tank 1 Area Hanaban SC	Soil	CHSB2-12	Benzene	0.000	ma/ka		B/FC	0.000546	CH-EW6	796000 ug/mL	125
NRMPI	IP-4	10/1/1075		DESP-Charleston Tank 1 Area Hanahan SC	Soil	CHSB2-12	Toluene	0.0307	ma/ka	-	T/FC	0.000527	CH-EW6	796000 ug/mL	433
NRMRI	JP-4	10/1/1975	############	DESP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-12	Ethylbenzene	0.0334	mg/kg		E/EC	0.0000027		796000 ug/mL	1587
NIDMDI		10/1/1975	######################################	DESP Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2 12	m Yylono	0.154	mg/kg		m Xylono/EC	0.001334		796000 ug/mL	F354
		10/1/1975	########	DESP Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-12		0.452	mg/kg		n Xylono/EC	0.000720		796000 ug/mL	1900
	ID /	10/1/1975	######################################	DESP Charleston, Tank 1 Area, Halldhall, SC	Soil	CUSP2 12	n Yulono	0.152	mg/kg			0.002202		790000 ug/mL	1000
		10/1/19/5	######################################	DESP Charleston, Tank I Area, Hanahan, SU	Soil	CUSP2 12	Total Vylanas (m.n. and a)	0.199	mg/kg			0.002901		796000 ug/mL	2357
	JF-4	10/1/19/5	######################################	DESP Charleston, Tank I Area, Hanahan, SC	Soil	CHER2 12	Total Aylenes (III,p, and 0)	0.803	mg/kg			0.011949		796000 ug/mL	9512
	JP-4	10/1/19/5	*****	DEOD Obselector, Tank T Area, Hananan, SC	501			1.0091	mg/kg			0.015016		796000 ug/mL	11953
NRMRL	JP-4	10/1/1975	####### #######	DESP-Unarleston, Tank 1 Area, Hanahan, SC	Soll	CHSB2-12		0.0721	mg/kg		(B+1)/FC	0.001073		796000 ug/mL	854
	JP-4	10/1/1975	######## #########	DESP-Charleston, Tank 1 Area, Hananan, SC	501	CH5B2-12	E+A	0.937	ing/kg			0.013943		796000 ug/mL	11099
NRMRL	JP-4	10/1/1975	####### #######	DESP-Charleston, Tank 1 Area, Hanahan, SC	Soll	CHSB2-12	B/ I	1.03672316			(B/T)/FC	0.015427		796000 ug/mL	12280
INKMKL	JP-4	10/1/1975	#######	DESP-Charleston, Tank 1 Area, Hanahan, SC	501	CHSB2-12	B/E	0.2738806			(B/E)/FC	0.004076		796000 ug/mL	3244
NRMRL	JP-4	10/1/1975	#######	DESP-Charleston, Lank 1 Area, Hanahan, SC	Soll	CHSB2-12	B/X	0.04570361			(B/X)/FC	0.00068		796000 ug/mL	541
NRMRL	JP-4	10/1/1975	######	DESE-Charleston, Lank 1 Area, Hanahan, SC	501	CHSB2-12	1/E	0.2641791	I		(1/E)/FC	0.003931	DH-EVV6	796000 ug/mL	3129
NRMRL	JP-4	10/1/1975	#######	DESP-Charleston, Tank 1 Area, Hanahan, SC	Sol	CHSB2-12	1/X	0.04408468			(1/X)/FC	0.000656	JH-EW6	796000 ug/mL	522
NRMRL	JP-4	10/1/1975	######	DESP-Charleston, Tank 1 Area, Hanahan, SC	Sol	CHSB2-12	E/X	0.16687422			(E/X)/FC	0.002483	JH-EW6	796000 ug/mL	1977
NRMRL	JP-4	10/1/1975	######	DESP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-12'	(B+1)/(E+X)	0.07694771		1	((B+T)/(E+X))/FC	0.001145	JH-EW6	796000 ug/mL	911

Lab Code Fuel Type	Spill Date	Date	Site Name	Matix	Locid	Analyte	Results	Units	Flag	FC Analyte	FC Ratio	FPLocid	FPdensity FPunits	SoilFuel (ug/mL)
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-12'	Fuel Carbon	67.2	mg/kg		Fuel Carbon	1 (CH-EW6	796000 ug/mL	796000
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-12'	1,2,3-Trimethylbenzene	0.168	mg/kg		1,2,3 TMB/FC	0.0025	CH-EW6	796000 ug/mL	1990
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-12'	1,2,4-Trimethylbenzene	0.331	mg/kg		1,2,4 TMB/FC	0.004926	CH-EW6	796000 ug/mL	3921
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-12'	1,3,5-Trimethylbenzene	0.167	mg/kg		1,3,5 TMB/FC	0.002485	CH-EW6	796000 ug/mL	1978
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-12'	1-MethylNaphthalene	0.189	mg/kg		1-MN/FC	0.002813	CH-EW6	796000 ug/mL	2239
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-12'	2-MethylNaphthalene	0.236	mg/kg		2-MN/FC	0.003512	CH-EW6	796000 ug/mL	2795
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13	Benzene	1.05	mg/kg		B/FC	0.000107	CH-EW6	796000 ug/mL	85
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13	Ioluene	21.7	mg/kg		T/FC	0.002217	CH-EW6	796000 ug/mL	1764
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soll	CHSB2-13	Etnylbenzene	34.3	mg/кg		E/FC	0.003504	CH-EW6	796000 ug/mL	2789
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soll	CHSB2-13	m-Xylene	90	mg/kg		m-Xylene/FC	0.009193	CH-EW6	796000 ug/mL	/318
NRMRL JP-4	10/1/1975	#######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	501	CHSB2-13	o-Xylene	45.2	mg/kg		o-Xylene/FC	0.004617		796000 ug/mL	3675
NRMRL JP-4	10/1/1975	#######	DESP Charleston, Tank 1 Area, Hanahan, SC	Soll	CHSB2-13	p-Aylene Total Xylanaa (m.n. and a)	38	mg/kg		p-Xylene/FC	0.003882		796000 ug/mL	3090
NRIVIRL JP-4	10/1/1975	#######	DESP Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13	Total RTEX	173.2	mg/kg			0.017692		796000 ug/mL	14062
	10/1/1975	######################################	DESP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13		230.23	mg/kg		(B+T)/FC	0.023319		796000 ug/mL	1850
	10/1/1975	######################################	DESP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13	E+Y	207.5	mg/kg		(E+Y)/EC	0.002324		796000 ug/mL	16871
	10/1/1975	######################################	DESP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13	B/T	0.0483871	шу/ку		(B/T)/FC	4.94E-06		796000 ug/mL	10071
NRMRI IP-4	10/1/1975	#######	DESP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13'	B/F	0.03061224			(B/F)/FC	3.13E-06	CH-EW6	796000 ug/mL	
NRMRI IP-4	10/1/1975	#######	DESP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13	B/X	0.00606236			(B/X)/FC	6 19E-07	CH-EW6	796000 ug/mL	0
NRMRL JP-4	10/1/1975	#######	DESP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13'	T/E	0.63265306			(T/E)/FC	6.46E-05	CH-EW6	796000 ug/mL	51
NRMRI JP-4	10/1/1975	#######	DESP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13'	T/X	0.12528868			(T/X)/FC	1 28E-05	CH-EW6	796000 ug/ml	10
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13'	E/X	0.19803695			(E/X)/FC	2.02E-05	CH-EW6	796000 ug/mL	16
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13'	(B+T)/(E+X)	0.10963855			((B+T)/(E+X))/FC	1.12E-05	CH-EW6	796000 ug/mL	9
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13'	Fuel Carbon	9790	mg/kg		Fuel Carbon	1	CH-EW6	796000 ug/mL	796000
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13'	1,2,3-Trimethylbenzene	21.4	mg/kg		1,2,3 TMB/FC	0.002186	CH-EW6	796000 ug/mL	1740
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13'	1,2,4-Trimethylbenzene	45.9	mg/kg		1,2,4 TMB/FC	0.004688	CH-EW6	796000 ug/mL	3732
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13'	1,3,5-Trimethylbenzene	21.7	mg/kg		1,3,5 TMB/FC	0.002217	CH-EW6	796000 ug/mL	1764
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13'	1-MethylNaphthalene	26.8	mg/kg		1-MN/FC	0.002737	CH-EW6	796000 ug/mL	2179
NRMRL JP-4	10/1/1975	######	DFSP-Charleston, Tank 1 Area, Hanahan, SC	Soil	CHSB2-13'	2-MethylNaphthalene	33.7	mg/kg		2-MN/FC	0.003442	CH-EW6	796000 ug/mL	2740
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB4	Benzene	0.006	mg/kg	**					No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB4	Toluene	0.006	mg/kg	**					No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB4	Ethylbenzene	0.006	mg/kg	**					No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB4	m-Xylene	0.006	mg/kg	**					No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB4	o-Xylene	0.006	mg/kg	**					No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB4	p-Xylene	0.006	mg/kg	**					No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB4	Total Xylenes (m,p, and o)	0.018	mg/kg						No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB4	Total BTEX	0.036	mg/kg						No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB4	B+1	0.012	mg/kg						No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB4	E+X	0.024	mg/kg						No FP/No FC data
NRMRL JP-4	10/1/1988	#######	KC-135 Crash Site, Wurtsmith AFB, MI	501	SB4	B/T	1							No FP/No FC data
NRMRL JP-4	10/1/1988	#######	KC-135 Crash Site, Wurtsmith AFB, MI	501	SB4	B/E	1							No FP/No FC data
NRIVIRL JP-4	10/1/1966	#######	KC 125 Crash Site, Wurtsmith AFB, MI	Soil	SD4		0.33333333							No FP/No FC data
	10/1/1988	#######	KC 125 Crash Site, Wurtsmith AFB, MI	Soil	SD4	1/E	0 22222222							No EP/No EC data
NRMRI IP-4	10/1/1988	########	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB4	E/X	0.33333333							No FP/No FC data
NRMRI IP-4	10/1/1988	#######	KC-135 Crash Site, Wurtsmith AEB, MI	Soil	SB4	(B+T)/(E+X)	0.000000000							No FP/No FC data
NRMRL JP-4	10/1/1988	#######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB4	1.2.3-Trimethylbenzene	0.0475	ma/ka						No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB4	1,2,4-Trimethylbenzene	0.006	mg/ka	**					No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB4	1,3,5-Trimethylbenzene	0.28	mg/kg						No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB4	1-MethylNaphthalene	0.117	mg/ka						No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB4	2-MethylNaphthalene	0.144	mg/kg						No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	Benzene	0.006	mg/kg	**					No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	Toluene	0.0855	mg/kg						No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	Ethylbenzene	0.006	mg/kg	**					No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	m-Xylene	0.006	mg/kg	**					No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	o-Xylene	0.006	mg/kg	**					No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	p-Xylene	0.006	mg/kg	**					No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	Total Xylenes (m,p, and o)	0.018	mg/kg						No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	Total BTEX	0.1155	mg/kg						No FP/No FC data
NRMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	B+T	0.0915	mg/kg						No FP/No FC data
NKMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Sol	SB5	E+X	0.024	mg/kg						NO FP/No FC data
NKMRL JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	B/I	0.07017544							No FP/No FC data
INKIVIKL JP-4	10/1/1988	#######	KC-135 Crash Site, Wurtsmith AFB, MI	501	5B3	B/E D/Y	1							No FP/No FC data
NRIVIRL JP-4	10/1/1988	#######	KC 125 Crash Site, Wurtsmith AFB, MI	Soil	303 905		0.33333333							No FP/No FC data
	10/1/1988	######################################	KC 125 Crash Site, Wurtsmith AFB, MI	Soil	300 905	T/V	14.25							No EP/No EC data
INININIAL JP-4	10/1/1988	#######	NOTION CIASTI OILE, WURSTHILT AFD, MI	301	363	1/A	4./5		1	1				INU FP/INU FC data

Lab Code	Fuel Type	Spill Date	Date	Site Name	Matix	Locid	Analyte	Results	Units	Flag	FC Analyte	FC Ratio	FPLocid	FPdensity	FPunits	SoilFuel (ug/mL)
NRMRL	JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	E/X	0.33333333								No FP/No FC data
NRMRL	JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	(B+T)/(E+X)	3.8125								No FP/No FC data
NRMRL	JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	1,2,3-Trimethylbenzene	0.0514	mg/kg							No FP/No FC data
NRMRL	JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	1,2,4-Trimethylbenzene	0.006	mg/kg	**						No FP/No FC data
NRMRL	JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	1,3,5-Trimethylbenzene	0.3	mg/kg							No FP/No FC data
NRMRL	JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	1-MethylNaphthalene	0.118	mg/kg							No FP/No FC data
NRMRL	JP-4	10/1/1988	######	KC-135 Crash Site, Wurtsmith AFB, MI	Soil	SB5	2-MethylNaphthalene	0.158	mg/kg							No FP/No FC data
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	Benzene	1.1 ו	mg/kg		B/FC	0.002048	MBMW8I	750000	ug/mL	1536
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	Toluene	0.006	mg/kg	**	T/FC	1.12E-05	MBMW8I	750000	ug/mL	8
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	Ethylbenzene	2.38	mg/kg		E/FC	0.004432	MBMW8I	750000	ug/mL	3324
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	m-Xylene	5.69	mg/kg		m-Xylene/FC	0.010596	MBMW8I	750000	ug/mL	7947
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	o-Xylene	0.131	mg/kg		o-Xylene/FC	0.000244	MBMW8I	750000	ug/mL	183
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	p-Xylene	1.1 ו	mg/kg		p-Xylene/FC	0.002048	MBMW8I	750000	ug/mL	1536
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	Total Xylenes (m,p, and o)	6.921	mg/kg		X/FC	0.012888	MBMW8I	750000	ug/mL	9666
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	Total BTEX	10.407 ו	mg/kg		BTEX/FC	0.01938	MBMW8I	750000	ug/mL	14535
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	B+T	1.106 ו	mg/kg		(B+T)/FC	0.00206	MBMW8I	750000	ug/mL	1545
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	E+X	9.301 (mg/kg		(E+X)/FC	0.01732	MBMW8I	750000	ug/mL	12990
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	B/T	183.333333			(B/T)/FC	0.341403	MBMW8I	750000	ug/mL	256052
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	B/E	0.46218487			(B/E)/FC	0.000861	MBMW8I	750000	ug/mL	646
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	B/X	0.15893657			(B/X)/FC	0.000296	MBMW8I	750000	ug/mL	222
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	T/E	0.00252101			(T/E)/FC	4.69E-06	MBMW8I	750000	ug/mL	4
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	T/X	0.00086693			(T/X)/FC	1.61E-06	MBMW8I	750000	ug/mL	1
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	E/X	0.34388094			(E/X)/FC	0.00064	MBMW8I	750000	ug/mL	480
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	(B+T)/(E+X)	0.11891194			((B+T)/(E+X))/FC	0.000221	MBMW8I	750000	ug/mL	166
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	Fuel Carbon	537 (mg/kg		Fuel Carbon	1	MBMW8I	750000	ug/mL	750000
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	1,2,3-Trimethylbenzene	4.17	mg/kg		1,2,3 TMB/FC	0.007765	MBMW8I	750000	ug/mL	5824
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	1,2,4-Trimethylbenzene	9.66 ו	mg/kg		1,2,4 TMB/FC	0.017989	MBMW8I	750000	ug/mL	13492
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	1,3,5-Trimethylbenzene	2.82	mg/kg		1,3,5 TMB/FC	0.005251	MBMW8I	750000	ug/mL	3939
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	1-MethylNaphthalene	2.71 I	mg/kg		1-MN/FC	0.005047	MBMW8I	750000	ug/mL	3785
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-1-9.5'	2-MethylNaphthalene	3.93 ו	mg/kg		2-MN/FC	0.007318	MBMW8I	750000	ug/mL	5489
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-2-9.5'	Benzene	1.71 ו	mg/kg		B/FC	0.000615	MBMW24	764000	ug/mL	470
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-2-9.5'	Toluene	0.0303 I	mg/kg		T/FC	1.09E-05	MBMW24	764000	ug/mL	8
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-2-9.5'	Ethylbenzene	8.47 (mg/kg		E/FC	0.003047	MBMW24	764000	ug/mL	2328
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-2-9.5'	m-Xylene	10.9 ו	mg/kg		m-Xylene/FC	0.003921	MBMW24	764000	ug/mL	2996
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-2-9.5'	o-Xylene	0.0503	mg/kg		o-Xylene/FC	1.81E-05	MBMW24	764000	ug/mL	14
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-2-9.5'	p-Xylene	0.497	mg/kg		p-Xylene/FC	0.000179	MBMW24	764000	ug/mL	137
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-2-9.5'	Total Xylenes (m,p, and o)	11.4473	mg/kg		X/FC	0.004118	MBMW24	764000	ug/mL	3146
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-2-9.5'	Total BTEX	21.6576	mg/kg		BTEX/FC	0.007791	MBMW24	764000	ug/mL	5952
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-2-9.5'	B+T	1.7403	mg/kg		(B+T)/FC	0.000626	MBMW24	764000	ug/mL	478
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-2-9.5'	E+X	19.9173	mg/kg		(E+X)/FC	0.007164	MBMW24	764000	ug/mL	5474
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-2-9.5'	B/T	56.4356436			(B/T)/FC	0.020301	MBMW24	764000	ug/mL	15510
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-2-9.5'	B/E	0.20188902			(B/E)/FC	7.26E-05	MBMW24	764000	ug/mL	55
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-2-9.5	B/X	0.1493802			(B/X)/FC	5.37E-05	MBMW24	764000	ug/mL	41
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-2-9.5	1/E	0.00357733			(1/E)/FC	1.29E-06	MBMW24	764000	ug/mL	1
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soil	MBSB-2-9.5	1/X	0.00264691			(1/X)/FC	9.52E-07	MBMW24	764000	ug/mL	1
NRMRL	JP-4	1/1/1981	#######	Pipeline Leak Site, Myrtle Beach AFB, SC	SOIL	MBSB-2-9.5		0.73991247			(E/X)/FC	0.000266	MBMW24	764000	ug/mL	203
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soll	MBSB-2-9.5	(B+1)/(E+X)	0.08/3/63			((B+T)/(E+X))/FC	3.14E-05	MBMW24	764000	ug/mL	24
NRMRL	JP-4	1/1/1981	#######	Pipeline Leak Site, Myrtle Beach AFB, SC	501	MBSB-2-9.5	Fuel Carbon	2780	mg/kg	-	Fuel Carbon	0.005000		/64000	ug/mL	764000
NRMRL	JP-4	1/1/1981	#######	Pipeline Leak Site, Myrtle Beach AFB, SC	501	MBSB-2-9.5	1,2,3-1 rimethylbenzene	14.7	mg/kg		1,2,3 TMB/FC	0.005288	MBNWV24	764000	ug/mL	4040
	JF-4	1/1/1981	#######	Pipeline Leak Site, Myrtle Beach AFB, SC	30II Soli	MDCD 2 0 5	1,2,4-1 rimethylbenzene	36.4	mg/Kg	+	1,2,4 TMB/FC	0.013094		764000	ug/mL	10003
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soll	MBSB-2-9.5	1,3,5-1 rimethylbenzene	9.51	mg/kg		1,3,5 TMB/FC	0.003421	MBMW24	764000	ug/mL	2614
NRMRL	JP-4	1/1/1981	#######	Pipeline Leak Site, Myrtle Beach AFB, SC	501	MBSB-2-9.5	1-MethylNaphthalene	7.62	mg/kg		1-IVIN/FC	0.002741	MBNWV24	764000	ug/mL	2094
NRMRL	JP-4	1/1/1981	######	Pipeline Leak Site, Myrtle Beach AFB, SC	Soll	MBSB-2-9.5	2-MethylNaphthalene	11.5	mg/kg		2-MN/FC	0.004137	MBMW24	764000	ug/mL	3160
NRMRL	JP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	501	EAKSB1-3	Benzene	10.8	mg/kg		B/FC	0.00345	EAKINIW316-FP	770400	ug/mL	2658
NRMRL	JP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soll	EAKSB1-3	Toluene	0.039	mg/kg		I/FC	1.25E-05	EAKINW316-FP	770400	ug/mL	10
	JF -4	10/1/19/3	######################################		5011	EAKODI-3	m Yulono	10.0	mg/kg	-	m Vulone/EC	0.0000004	EARIVIVISIO-FP	770400	ug/IIIL	4086
NRMRL	JP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	501	EAKSB1-3	m-xylene	43.7	mg/kg		m-Xylene/FC	0.013962	EAKINIW316-FP	770400	ug/mL	10756
	JF-4	10/1/19/3	######################################	Spill Site No. 2, Eaker AFB, AR	Soil	EAKOBI-3	u-Aylene	17.5	mg/kg	-	o-Aylene/FC	0.005591	EARIVIVIJIO-FP	770400	ug/mL	4307
	JF -4	10/1/19/3	######################################		5011	EAKODI-3	P-Aylene (man and -)	20.1	mg/kg	-	P-Aylelle/FC	0.000422	EARIVIVISIO-FP	770400	ug/IIIL	4947
	JF'-4	10/1/19/3	######################################	Spill Site No. 2, Eaker AFB, AK	Soil	EARODI-3	Total Aylenes (m,p, and 0)	109 720	mg/kg	-		0.020974	EARIVIVV310-FP	770400	ug/IIIL ug/ml	20011
	JF -4	10/1/19/3	######################################		Soil	EAKODI-3		100.739	mg/kg	-		0.0034741	EARIVIVISIO-FP	770400	ug/IIIL ug/ml	20704
	JF-4 ID-4	10/1/19/3	######################################		Soli	EAKSB1 2	DT I F±Y	10.039	mg/kg	+	(B+1)/FC (E+X)/EC	0.003403	EAKMW/216 ED	770400	ug/IIIL ug/ml	2008
NRMPI	ID_/	10/1/19/3	~~~~~	Spill Site No. 2, Eaker AED AD	Soil	EAKSB1 2	B/T	276 0227	шу/ку	+	(E+A)/FO (B/T)/FC	0.031270		770400	ug/IIIL ug/ml	24097
NRMPI	IP-4	10/1/19/3	######################################	Soill Site No. 2, Eaker AFB, AR	Soil	EAKSB1-3	B/F	0.65060241		+	(B/F)/FC	0.000474	FAKMW/316-FP	770400	ug/mL	160
NRMPI	ID_/	10/1/19/3	~~~~~	Spill Site No. 2, Eaker AED AD	Soil	EAKSB1 2	B/Y	0.1329/122		+	(B/Y)/FC	1 24 - 05		770400	ug/IIIL ug/ml	100
INIAIVIIAE	ui-*+	10/1/19/3	#######	Opini One NU. Z, Eaker AFD, AR	001	LANODIS		0.10204100		1		4.24E-05	LANIVIVISIO-FP	110400	ug/IIIL	

Lab Code	Fuel Type	Spill Date	Date	Site Name	Matix	Locid	Analyte	Results	Units	Flag	FC Analyte	FC Ratio	FPLocid	FPdensity FPunits	SoilFuel (ug/mL)
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB1-3	T/E	0.0023494			(T/E)/FC	7.51E-07	EAKMW316-FP	770400 ug/mL	1
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB1-3	T/X	0.0004797			(T/X)/FC	1.53E-07	EAKMW316-FP	770400 ug/mL	0
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB1-3	E/X	0.20418204			(E/X)/FC	6.52E-05	EAKMW316-FP	770400 ug/mL	50
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB1-3	(B+T)/(E+X)	0.11071502			((B+T)/(E+X))/FC	3.54E-05	EAKMW316-FP	770400 ug/mL	27
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB1-3	Fuel Carbon	3130	mg/kg		Fuel Carbon	1	EAKMW316-FP	770400 ug/mL	770400
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB1-3	1,2,3-Trimethylbenzene	12.2	mg/kg		1,2,3 TMB/FC	0.003898	EAKMW316-FP	770400 ug/mL	3003
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB1-3	1,2,4-Trimethylbenzene	36.2	mg/kg		1,2,4 TMB/FC	0.011565	EAKMW316-FP	770400 ug/mL	8910
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2. Eaker AFB. AR	Soil	EAKSB1-3	1.3.5-Trimethylbenzene	19.6	ma/ka		1.3.5 TMB/FC	0.006262	EAKMW316-FP	770400 ug/mL	4824
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB1-3	1-MethylNaphthalene	6.44	mg/kg		1-MN/FC	0.002058	EAKMW316-FP	770400 ug/mL	1585
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2. Eaker AFB. AR	Soil	EAKSB1-3	2-MethylNaphthalene	11.7	ma/ka		2-MN/FC	0.003738	EAKMW316-FP	770400 ug/mL	2880
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2. Eaker AFB, AR	Soil	EAKSB2-2	Benzene	0.006	ma/ka	**	B/FC	2.06E-05	EAKMW306-FP	762600 ug/mL	16
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2. Eaker AFB, AR	Soil	EAKSB2-2	Toluene	0.006	ma/ka	**	T/FC	2.06E-05	EAKMW306-FP	762600 ug/mL	16
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2. Eaker AFB, AR	Soil	EAKSB2-2	Ethylbenzene	0.265	ma/ka		E/FC	0.000911	EAKMW306-FP	762600 ug/mL	694
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2. Eaker AFB, AR	Soil	EAKSB2-2	m-Xylene	0.006	ma/ka	**	m-Xylene/FC	2.06E-05	EAKMW306-FP	762600 ug/mL	16
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-2	o-Xvlene	0.006	ma/ka	**	o-Xvlene/FC	2.06E-05	EAKMW306-FP	762600 ug/mL	16
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-2	p-Xylene	0.193	ma/ka		p-Xylene/FC	0.000663	EAKMW306-FP	762600 ug/mL	506
NRMRI	JP-4	10/1/1973	#######	Spill Site No. 2. Eaker AEB, AR	Soil	FAKSB2-2	Total Xylenes (m.p. and o)	0 205	ma/ka		X/FC	0.000704	EAKMW306-EP	762600 ug/ml	537
NRMRL	JP-4	10/1/1973	######	Spill Site No. 2. Eaker AFB, AR	Soil	EAKSB2-2	Total BTEX	0.482	ma/ka		BTEX/FC	0.001656	EAKMW306-FP	762600 ug/mL	1263
NRMRI	JP-4	10/1/1973	#######	Spill Site No. 2. Eaker AEB, AR	Soil	FAKSB2-2	B+T	0.012	ma/ka		(B+T)/FC	4 12E-05	EAKMW306-EP	762600 ug/ml	31
NRMRI	JP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-2	E+X	0.47	ma/ka		(E+X)/FC	0.001615	EAKMW306-FP	762600 ug/ml	1232
NRMRI	JP-4	10/1/1973	#######	Spill Site No. 2. Eaker AEB, AR	Soil	FAKSB2-2	B/T	1			(B/T)/FC	0.003436	EAKMW306-EP	762600 ug/ml	2621
NRMRI	JP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-2	B/F	0.02264151			(B/E)/EC	7 78E-05	EAKMW306-FP	762600 ug/ml	59
NRMRI	JP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-2	B/X	0.02926829			(B/X)/FC	0.000101	EAKMW306-EP	762600 ug/ml	77
NRMRI	JP-4	10/1/1973	#######	Spill Site No. 2 Faker AFB AR	Soil	EAKSB2-2	T/F	0.02264151			(T/E)/EC	7 78E-05	EAKMW306-FP	762600 ug/ml	59
NRMRI	IP-4	10/1/1973	#######	Spill Site No. 2 Eaker AEB AR	Soil	EAKSB2-2	T/X	0.02926829			(T/X)/FC	0.000101	EAKMW/306-EP	762600 ug/ml	77
NRMRI	IP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-2	F/X	1 29268293			(F/X)/FC	0.000101	EAKMW306-EP	762600 ug/mL	3388
NRMRI	IP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-2	(B+T)/(E+X)	0.02553191			((B+T)/(E+X))/EC	8 77E-05	EAKMW306-FP	762600 ug/mL	67
NRMRI	IP-4	10/1/1973	#######	Spill Site No. 2, Eaker AEB, AR	Soil	EAKSB2-2	Evel Carbon	201	ma/ka		Evel Carbon	0.112 00	EAKMW/306-EP	762600 ug/mL	762600
NRMRI	JP-4	10/1/1973	########	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-2	1 2 3-Trimethylbenzene	0.682	ma/ka		1 2 3 TMB/FC	0.002344	EAKMW306-FP	762600 ug/mL	1787
NRMRI	IP-4	10/1/1973	#######	Spill Site No. 2, Eaker AEB, AR	Soil	EAKSB2-2	1.2.4-Trimethylbenzene	1 99	ma/ka		1.2.4 TMB/FC	0.002044	EAKMW/306-EP	762600 ug/mL	5215
NRMRI	IP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-2	1 3 5-Trimethylbenzene	0.692	ma/ka		1 3 5 TMB/FC	0.0000000	EAKMW/306-FP	762600 ug/mL	1813
NRMRI	IP-4	10/1/1973	#######	Spill Site No. 2, Eaker AEB, AR	Soil	EAKSB2-2	1-Methy/Naphthalene	0.535	ma/ka		1-MN/EC	0.002070	EAKMW/306-FP	762600 ug/mL	1/02
NRMRI	IP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-2	2-MethylNaphthalene	0.333	ma/ka		2-MN/EC	0.001030	EAKMW306-FP	762600 ug/mL	1402
NRMRI	JP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-4	Benzene	0.420	ma/ka	**	B/FC	3 73E-06	EAKMW306-FP	762600 ug/mL	1122
NRMRI	IP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-4	Toluene	0.000	ma/ka	**	D/FC	3.73E-06	EAKMW306-FP	762600 ug/mL	3
NRMRI	JP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-4	Ethylbenzene	5.4	ma/ka		F/FC	0.003354	EAKMW/306-FP	762600 ug/mL	2558
NRMRI	IP-4	10/1/1973	#######	Spill Site No. 2, Eaker AEB, AR	Soil	EAKSB2-4	m-Yylene	0.4	ma/ka		m-Yylene/EC	0.000545	EAKMW/306-EP	762600 ug/mL	/15
NRMRI	JP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-4	o-Xvlene	2 72	ma/ka		o-Xylene/EC	0.000040		762600 ug/mL	1288
NRMRI	IP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-4	n-Xylene	5.8	ma/ka		p-Xylene/FC	0.001003	EAKMW306-FP	762600 ug/mL	27/7
NRMRI	JP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-4	Total Xylenes (m.n. and o)	0 307	ma/ka		Y/FC	0.005002	EAKMW306-FP	762600 ug/mL	1/51
NRMRI	IP-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-4	Total RTEX	14 800	ma/ka		BTEX/EC	0.0000007	EAKMW306-FP	762600 ug/mL	701/
		10/1/1973	########	Spill Site No. 2, Eaker AEB, AR	Soil			0.012	mg/kg			7 455 06		762600 ug/mL	7014
NRMRL	JF-4	10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-4	E+Y	14 797	mg/kg		(B+T)/FC	0.000101	EAKMW306-FP	762600 ug/mL	7009
		10/1/1973	########	Spill Site No. 2, Eaker AEB, AR	Soil		P/T	14.737	шу/ку			0.003131		762600 ug/mL	1003
		10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-4	D/T	0.00111111				6 0E 07		762600 ug/mL	4/4
		10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-4	B/E B/Y	0.00111111				2 07E 07		762600 ug/mL	1
		10/1/1973	#######	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-4		0.0006365				5.97E-07	EAKININ 300-FP	762600 ug/mL	1
	JF-4	10/1/1973	######################################	Spill Site No. 2, Eaker AFB, AR	Soil	EAKSB2-4	1/2	0.00111111				0.9E=07	EAKINW300-FF	762600 ug/mL	1
	JF-4 ID-4	10/1/19/3	######################################	Spill Site No. 2, Eaker AFD, AR	Soil	EAKSB2-4	E/Y	0.0000385			(1/A)/FC	0.000357	EAKMW/306 EP	762600 ug/IIIL	0
	JF-4	10/1/19/3	######################################	Spill Site No. 2, Eaker AFD, AR	Soil	EAROD2-4		0.01400148				5.04E 07	EAKININ SUD-FP	762600 ug/mL	212
	JF-4	10/1/19/3	######################################	Opill Site No. 2, Eaker AFD, AR	Soil	EAROD2-4		0.00001098	malle		((D+1)/(E+A))/FC	5.04⊏-07	EAKININ SUD-FP	762600 ug/IIL	760000
	JF-4	10/1/19/3	#######	Spill Site No. 2, Eaker AFB, AR	Soll	EAROB2-4	1.2.2 Trimethylhenar	10101	mg/kg		1 2 2 TMP/EC	0.00254	EAKIVIVV306-FP	762600 ug/mL	/02600
	JF-4	10/1/19/3	#######	Opill Oite No. 2, Eaker AFB, AR	3011	EARODO 4	1,2,3-1 rimetnyibenzene	4.09	під/кд		1,2,3 TMB/FC	0.00254	EAKIVIVV300-FP	762600 ug/mL	1937
NRMRL	JP-4	10/1/19/3	#######	Spill Site No. 2, Eaker AFB, AR	501	EAKSB2-4	1,2,4-1 rimetnyibenzene	14.1	mg/kg		1,2,4 IMB/FC	0.008758	EAKMW306-FP	762600 ug/mL	6679
	JF-4	10/1/19/3	#######	Opill Oite No. 2, Eaker AFB, AR	3011	EARODO 4	1,3,5-1 rimetnyibenzene	7.66	під/кд		1,3,3 TMB/FC	0.004758	EAKIVIVV300-FP	762600 ug/mL	3628
NRMRL	JP-4	10/1/19/3	#######	Spill Site No. 2, Eaker AFB, AR	501	EAKSB2-4	1-wethylNaphthalene	3.74	mg/kg		1-IVIN/FC	0.002323	EAKMW306-FP	762600 ug/mL	1//2
INKMIKL	JP-4	10/1/1973	######	Spill Site NO. 2, Eaker AFB, AR	2011	EAKSB2-4	∠-ivietnyiNaphthalene	6.71	тад/кд		Z-IMIN/FC	0.004168	EAKMW306-FP	762600 ug/mL	3178

Lab Code	Fuel Type	Spill Date	Date	Site Name	Matix	Locid	Analyte	Results Units Flag	FC Analyte	FC Ratio	FPLocid	FPdensity FPunits	SoilFuel (ug/mL)
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB1-4'	Benzene	0.0457 mg/kg	B/FC	9.44215E-06	BFT-401-3	804400 ug/mL	8
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB1-4'	Toluene	0.56 mg/kg	T/FC	0.000115702	BFT-401-3	804400 ug/mL	93
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB1-4'	Ethylbenzene	3.35 mg/kg	E/FC	0.000692149	BFT-401-3	804400 ug/mL	557
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB1-4'	m-Xylene	5.02 mg/kg	m-Xylene/FC	0.00103719	BFT-401-3	804400 ug/mL	834
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB1-4'	o-Xylene	3.95 mg/kg	o-Xylene/FC	0.000816116	BFT-401-3	804400 ug/mL	656
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB1-4'	p-Xylene	1.98 mg/kg	p-Xylene/FC	0.000409091	BFT-401-3	804400 ug/mL	329
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB1-4'	Total Xylenes (m,p, and o)	10.95 mg/kg	X/FC	0.002262397	BFT-401-3	804400 ug/mL	1820
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB1-4	Total BTEX	14.9057 mg/kg	BIEX/FC	0.00307969	BFT-401-3	804400 ug/mL	2477
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soll	BUTFC-SB1-4	B+1	0.6057 mg/kg	(B+T)/FC	0.000125145	BFT-401-3	804400 ug/mL	101
NRIVIRL	JP-5	6/1/1990	####### Tank Fa	arm C. Beaufort MCAS, SC	Soll	BUTFC-SB1-4	E+A P/T	14.3 mg/kg		0.002954545	BF1-401-3	804400 ug/mL	2311
	JP-5	6/1/1990	####### Tank Fa	arm C. Beaufort MCAS, SC	Soil	BUTEC SB1 4	D/I D/E	0.08160714		2 919555 06	DF1-401-3	804400 ug/mL	14
	JP-5	6/1/1990	####### Tank Fa	arm C. Beaufort MCAS, SC	Soil	BUTEC SB1 4	B/E B/V	0.01304179		2.01000E-00	DF1-401-3	804400 ug/mL	2
	JP-5	6/1/1990	###### Tank Fa	arm C Beaufort MCAS, SC	Soil	BUTEC-SB1-4		0.16716/18		3.45381E-05	BFT-401-3	804400 ug/mL	28
NRMRI	IP-5	6/1/1990	###### Tank Fa	arm C. Beaufort MCAS, SC	Soil	BUTEC-SB1-4	T/X	0.05114155	(T/X)/FC	1.05664E-05	BFT-401-3	804400 ug/mL	20
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C. Beaufort MCAS, SC	Soil	BUTFC-SB1-4	E/X	0.30593607	(E/X)/FC	6.32099E-05	BFT-401-3	804400 ug/mL	51
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C. Beaufort MCAS, SC	Soil	BUTFC-SB1-4	(B+T)/(E+X)	0.04235664	((B+T)/(E+X))/FC	8.75137E-06	BFT-401-3	804400 ug/mL	7
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C. Beaufort MCAS. SC	Soil	BUTFC-SB1-4	Fuel Carbon	4840 ma/ka	Fuel Carbon	1	BFT-401-3	804400 ug/mL	804400
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB1-4'	1,2,3-Trimethylbenzene	20.6 mg/kg	1,2,3 TMB/FC	0.004256198	BFT-401-3	804400 ug/mL	3424
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB1-4'	1,2,4-Trimethylbenzene	33.4 mg/kg	1,2,4 TMB/FC	0.006900826	BFT-401-3	804400 ug/mL	5551
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB1-4'	1,3,5-Trimethylbenzene	9.58 mg/kg	1,3,5 TMB/FC	0.001979339	BFT-401-3	804400 ug/mL	1592
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB1-4'	1-MethylNaphthalene	20.5 mg/kg	1-MN/FC	0.004235537	BFT-401-3	804400 ug/mL	3407
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB1-4'	2-MethylNaphthalene	27.4 mg/kg	2-MN/FC	0.005661157	BFT-401-3	804400 ug/mL	4554
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB2-3.5'	Benzene	0.375 mg/kg	B/FC	1.84729E-05	BFT-401-3	804400 ug/mL	15
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB2-3.5'	Toluene	0.382 mg/kg	T/FC	1.88177E-05	BFT-401-3	804400 ug/mL	15
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB2-3.5'	Ethylbenzene	15.3 mg/kg	E/FC	0.000753695	BFT-401-3	804400 ug/mL	606
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB2-3.5'	m-Xylene	22.9 mg/kg	m-Xylene/FC	0.001128079	BFT-401-3	804400 ug/mL	907
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB2-3.5'	o-Xylene	13.7 mg/kg	o-Xylene/FC	0.000674877	BFT-401-3	804400 ug/mL	543
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB2-3.5'	p-Xylene	8.94 mg/kg	p-Xylene/FC	0.000440394	BFT-401-3	804400 ug/mL	354
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB2-3.5	Total Xylenes (m,p, and o)	45.54 mg/kg	X/FC	0.00224335	BFT-401-3	804400 ug/mL	1805
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB2-3.5	Total BTEX	61.597 mg/kg	BTEX/FC	0.003034335	BFT-401-3	804400 ug/mL	2441
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB2-3.5	B+I	0.757 mg/kg	(B+1)/FC	3.72906E-05	BFT-401-3	804400 ug/mL	30
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soll	BUTFC-SB2-3.5	E+X	60.84 mg/kg	(E+X)/FC	0.002997044	BFT-401-3	804400 ug/mL	2411
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soll	BUTFC-SB2-3.5	B/I	0.98167539	(B/T)/FC	4.83584E-05	BFT-401-3	804400 ug/mL	39
NRIVIRL	JP-5	6/1/1990	####### Tank Fa	arm C. Beaufort MCAS, SC	Soll	BUTFC-SB2-3.5	B/E	0.0245098	(B/E)/FC	1.20738E-06	BF1-401-3	804400 ug/mL	1
NRIVIRL	JP-5	6/1/1990	####### Tank Fa	arm C. Beaufort MCAS, SC	Soll	BUTFC-SB2-3.5	B/A	0.00823452		4.05641E-07	BF1-401-3	804400 ug/mL	0
	JP-5	6/1/1990	####### Tank Fa	arm C. Beaufort MCAS, SC	Soil	BUTEC SP2 2 5'	1/E	0.02490732	(T/E)/FC	1.22992E-00	DF1-401-3	804400 ug/mL	1
NRMRI	IP-5	6/1/1990	###### Tank Fa	arm C. Beaufort MCAS, SC	Soil	BUTEC-SB2-3.5	E/X	0.33596838	(F/X)/FC	1.65502E-05	BFT-401-3	804400 ug/mL	13
NRMRI	IP-5	6/1/1990	###### Tank Fa	arm C. Beaufort MCAS, SC	Soil	BUTEC-SB2-3.5	(B+T)/(E+X)	0.01244247	((B+T)/(E+X))/EC	6 1293E-07	BFT-401-3	804400 ug/mL	13
NRMRI	JP-5	6/1/1990	###### Tank Fa	arm C. Beaufort MCAS. SC	Soil	BUTEC-SB2-3.5'	Fuel Carbon	20300 mg/kg	Fuel Carbon	0.12002 01	BFT-401-3	804400 ug/mL	804400
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C. Beaufort MCAS, SC	Soil	BUTFC-SB2-3.5'	1.2.3-Trimethylbenzene	95.5 mg/kg	1.2.3 TMB/FC	0.004704433	BFT-401-3	804400 ug/mL	3784
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C. Beaufort MCAS. SC	Soil	BUTFC-SB2-3.5'	1.2.4-Trimethylbenzene	148 mg/kg	1.2.4 TMB/FC	0.00729064	BFT-401-3	804400 ug/mL	5865
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB2-3.5'	1,3,5-Trimethylbenzene	43.5 mg/kg	1,3,5 TMB/FC	0.002142857	BFT-401-3	804400 ug/mL	1724
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB2-3.5'	1-MethylNaphthalene	87.3 mg/kg	1-MN/FC	0.004300493	BFT-401-3	804400 ug/mL	3459
NRMRL	JP-5	6/1/1990	###### Tank Fa	arm C, Beaufort MCAS, SC	Soil	BUTFC-SB2-3.5'	2-MethylNaphthalene	115 mg/kg	2-MN/FC	0.005665025	BFT-401-3	804400 ug/mL	4557
NRMRL	JP-5	6/1/1974	##### Day Tar	nk 865, Beaufort MCAS, SC	Soil	BUDTSB-3.5'	Benzene	3.25 mg/kg	B/FC	0.000770142	ASSUMEdens	804400 ug/mL	620
NRMRL	JP-5	6/1/1974	##### Day Tar	nk 865, Beaufort MCAS, SC	Soil	BUDTSB-3.5'	Toluene	0.0786 mg/kg	T/FC	1.86256E-05	ASSUMEdens	804400 ug/mL	15
NRMRL	JP-5	6/1/1974	###### Day Tar	nk 865, Beaufort MCAS, SC	Soil	BUDTSB-3.5'	Ethylbenzene	23.9 mg/kg	E/FC	0.005663507	ASSUMEdens	804400 ug/mL	4556
NRMRL	JP-5	6/1/1974	###### Day Tar	nk 865, Beaufort MCAS, SC	Soil	BUDTSB-3.5'	m-Xylene	0.006 mg/kg **	m-Xylene/FC	1.4218E-06	ASSUMEdens	804400 ug/mL	1
NRMRL	JP-5	6/1/1974	###### Day Tar	nk 865, Beaufort MCAS, SC	Soil	BUDTSB-3.5'	o-Xylene	0.0812 mg/kg	o-Xylene/FC	1.92417E-05	ASSUMEdens	804400 ug/mL	15
NRMRL	JP-5	6/1/1974	###### Day Tar	nk 865, Beaufort MCAS, SC	Soil	BUDTSB-3.5'	p-Xylene	0.006 mg/kg **	p-Xylene/FC	1.4218E-06	ASSUMEdens	804400 ug/mL	1
NRMRL	JP-5	6/1/1974	###### Day Tar	nk 865, Beaufort MCAS, SC	Soil	BUDTSB-3.5'	Total Xylenes (m,p, and o)	0.0932 mg/kg	X/FC	2.20853E-05	ASSUMEdens	804400 ug/mL	18
NRMRL	JP-5	6/1/1974	###### Day Tar	nk 865, Beaufort MCAS, SC	Soil	BUDTSB-3.5	Total BTEX	27.3218 mg/kg	BTEX/FC	0.00647436	ASSUMEdens	804400 ug/mL	5208
NRMRL	JP-5	6/1/1974	###### Day Tar	nk 865, Beaufort MCAS, SC	Soil	BUDTSB-3.5	B+I	3.3286 mg/kg	(B+T)/FC	0.000788768	ASSUMEdens	804400 ug/mL	634
NRMRL	JP-5	6/1/1974	###### Day Tar	NK 865, Beaufort MCAS, SC	Soil	BUDISB-3.5	E+X	23.9932 mg/kg	(E+X)/FC	0.005685592	ASSUMEdens	804400 ug/mL	4573
	JP-5	6/1/1974	###### Day Tar	1K 865, Beaufort MCAS, SC	Soil	BUDISB-3.5	B/I D/E	41.3486005	(B/T)/FC	0.009798247	ASSUMEdens	804400 ug/mL	7882
	JF-0	6/1/19/4	###### Day Tar	IK 000, BEAUTOIT MUAS, SU	501	BUDTER 2.5		0.13098320		3.22235E-05	ASSUMEdans	804400 Ug/mL	26
	JF-0 ID 5	6/1/1974	###### Day Tar	IN 000, DEBUIUL WUAS, SU	501	BUDIOB-3.5		0.0022997		7 70212E 07	ASSUMEdens	804400 ug/mL	1400
	JF-0 IP-5	6/1/1974	###### Day Tar	ak 865 Beaufort MCAS, SC	Soil	BUDTSB-3.5		0.0032007	(T/E)/FC	0.000100245	ASSI IMEdone	804400 ug/mL	161
	JP-5	6/1/1074	###### Day Tar	nk 865 Beaufort MCAS SC	Soil	BUDTSB-3.5	F/X	256 437768	(F/X)/FC	0.000199045	ASSI IMEdens	804400 ug/mL	101
	IP-5	6/1/1974	###### Day Tar	ak 865 Beaufort MCAS, SC	Soil	BUDTSB-3.5	(B+T)/(E+X)	0 13873097	((B+T)/(E+X))/FC	3 28746E-05	ASSLIMEdens	804400 ug/mL	1000 1 26
NRMRI	IP-5	6/1/1974	###### Day Tar	ak 865 Beaufort MCAS SC	Soil	BUDTSB-3.5	Fuel Carbon	4220 mg/kg	Fuel Carbon	1	ASSUMEdens	804400 ug/mL	804400
	JP-5	6/1/1974	###### Day Tar	nk 865. Beaufort MCAS_SC	Soil	BUDTSB-3.5	1.2.3-Trimethylbenzene	28 ma/ka	1.2.3 TMB/FC	0.006635071	ASSUMEdens	804400 ug/mL	5337

Lab Code	Fuel Type	Spill Date	Date	Site Name	Matix	Locid	Analyte	Results Units Flag	FC Analyte	FC Ratio	FPLocid	FPdensity FPunits	SoilFuel (ug/mL)
NRMRL	JP-5	6/1/1974	######	Day Tank 865, Beaufort MCAS, SC	Soil	BUDTSB-3.5'	1,2,4-Trimethylbenzene	0.287 mg/kg	1,2,4 TMB/FC	6.80095E-05	ASSUMEdens	804400 ug/mL	55
NRMRL	JP-5	6/1/1974	######	Day Tank 865, Beaufort MCAS, SC	Soil	BUDTSB-3.5'	1,3,5-Trimethylbenzene	0.014 mg/kg	1,3,5 TMB/FC	3.31754E-06	ASSUMEdens	804400 ug/mL	3
NRMRL	JP-5	6/1/1974	######	Day Tank 865, Beaufort MCAS, SC	Soil	BUDTSB-3.5'	1-MethylNaphthalene	29.8 mg/kg	1-MN/FC	0.007061611	ASSUMEdens	804400 ug/mL	5680
NRMRL	JP-5	6/1/1974	######	Day Tank 865, Beaufort MCAS, SC	Soil	BUDTSB-3.5'	2-MethylNaphthalene	44.3 mg/kg	2-MN/FC	0.01049763	ASSUMEdens	804400 ug/mL	8444
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB1-8.5'	Benzene	0.953 mg/kg	B/FC	3.32056E-05	CEF-293-9FP	799000 ug/mL	27
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB1-8.5'	Toluene	17.2 mg/kg	T/FC	0.000599303	CEF-293-9FP	799000 ug/mL	479
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB1-8.5	Ethylbenzene	141 mg/kg	E/FC	0.004912892	CEF-293-9FP	799000 ug/mL	3925
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB1-8.5	m-Xylene	233 mg/kg	m-Xylene/FC	0.008118467	CEF-293-9FP	799000 ug/mL	6487
NRMRL	JP-5	6/1/1981	#######	Facility 293, Cecil Field NAS, FL	Soll	CFSB1-8.5	o-Xylene	32.8 mg/kg	o-Xylene/FC	0.001142857	CEF-293-9FP	799000 ug/mL	913
NRIVIRL	JP-5	6/1/1981	########	Facility 293, Cecil Field NAS, FL	Soli	CFSB1-8.5	p-Xylene	114 mg/kg	p-Xylene/FC	0.003972125	CEF-293-9FP	799000 ug/mL	3174
	JP-5	6/1/1901	#######	Facility 293, Cecil Field NAS, FL	Soil	CESP1 0.5	Total PTEX	529.052 mg/kg		0.013233449	CEF-293-9FP	799000 ug/mL	10574
	JF-5	6/1/1901	#######	Facility 293, Cecil Field NAS, FL	Soil	CESP1 9 5'		19 152 mg/kg	/BIT/FC	0.01077003	CEF 202 0EP	799000 ug/mL	505
	JP-5	6/1/1981	#######	Facility 293, Cecil Field NAS, FL	Soil	CESB1-8.5	E+Y	520.8 mg/kg	(E+Y)/FC	0.018146341	CEF-293-9FF	799000 ug/mL	1//00
NRMRI	JP-5	6/1/1981	########	Facility 293, Cecil Field NAS, FL	Soil	CESB1-8.5'	B/T	0.05540698	(B/T)/FC	1 93056E-06	CEF-293-9FP	799000 ug/mL	2
NRMRI	JP-5	6/1/1981	#######	Facility 293, Cecil Field NAS, FL	Soil	CESB1-8.5'	B/F	0.00675887	(B/F)/FC	2 35501E-07	CEF-293-9EP	799000 ug/mL	0
NRMRL	JP-5	6/1/1981	#######	Facility 293, Cecil Field NAS, FL	Soil	CFSB1-8.5'	B/X	0.00250922	(B/X)/FC	8.74291E-08	CEF-293-9FP	799000 ug/mL	0
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB1-8.5'	T/E	0.12198582	(T/E)/FC	4.25038E-06	CEF-293-9FP	799000 µg/mL	3
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB1-8.5'	T/X	0.04528699	(T/X)/FC	1.57794E-06	CEF-293-9FP	799000 ug/mL	1
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB1-8.5'	E/X	0.37124803	(E/X)/FC	1.29355E-05	CEF-293-9FP	799000 ug/mL	10
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB1-8.5'	(B+T)/(E+X)	0.03485599	((B+T)/(E+X))/FC	1.21449E-06	CEF-293-9FP	799000 ug/mL	1
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB1-8.5'	Fuel Carbon	28700 mg/kg	Fuel Carbon	1	CEF-293-9FP	799000 ug/mL	799000
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB1-8.5'	1,2,3-Trimethylbenzene	88.4 mg/kg	1,2,3 TMB/FC	0.003080139	CEF-293-9FP	799000 ug/mL	2461
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB1-8.5'	1,2,4-Trimethylbenzene	210 mg/kg	1,2,4 TMB/FC	0.007317073	CEF-293-9FP	799000 ug/mL	5846
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB1-8.5'	1,3,5-Trimethylbenzene	86.3 mg/kg	1,3,5 TMB/FC	0.003006969	CEF-293-9FP	799000 ug/mL	2403
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB1-8.5'	1-MethylNaphthalene	84.3 mg/kg	1-MN/FC	0.002937282	CEF-293-9FP	799000 ug/mL	2347
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB1-8.5'	2-MethylNaphthalene	133 mg/kg	2-MN/FC	0.004634146	CEF-293-9FP	799000 ug/mL	3703
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-8.5'	Benzene	0.976 mg/kg	B/FC	3.35395E-05	CEF-293-9FP	799000 ug/mL	27
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-8.5'	Toluene	19.1 mg/kg	T/FC	0.000656357	CEF-293-9FP	799000 ug/mL	524
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-8.5'	Ethylbenzene	155 mg/kg	E/FC	0.00532646	CEF-293-9FP	799000 ug/mL	4256
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-8.5'	m-Xylene	263 mg/kg	m-Xylene/FC	0.009037801	CEF-293-9FP	799000 ug/mL	7221
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-8.5'	o-Xylene	37.4 mg/kg	o-Xylene/FC	0.001285223	CEF-293-9FP	799000 ug/mL	1027
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-8.5	p-Xylene	125 mg/kg	p-Xylene/FC	0.004295533	CEF-293-9FP	799000 ug/mL	3432
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-8.5	I otal Xylenes (m,p, and o)	425.4 mg/kg	X/FC	0.014618557	CEF-293-9FP	799000 ug/mL	11680
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-8.5	I otal BIEX	600.476 mg/kg	BIEX/FC	0.020634914	CEF-293-9FP	799000 ug/mL	16487
NRMRL	JP-5	6/1/1981	#######	Facility 293, Cecil Field NAS, FL	Soll	CFSB11-8.5	B+I	20.076 mg/kg	(B+1)/FC	0.000689897	CEF-293-9FP	799000 ug/mL	551
NRIVIRL	JP-5	6/1/1981	########	Facility 293, Cecil Field NAS, FL	Soll	CFSB11-8.5	E+A	580.4 mg/kg		1 7565 06	CEF-293-9FP	799000 ug/mL	15930
	JP-5	6/1/1901	#######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-0.5	D/I D/E	0.005109948		1.750E-00	CEF-293-9FP	799000 ug/mL	1
	JP-5	6/1/1981	#######	Facility 293, Cecil Field NAS, FL	Soil	CESB11-8.5	B/X	0.00229431	(B/E)/FC	7 88/23E-08	CEF-293-9FF	799000 ug/mL	0
	JP-5	6/1/1981	#######	Facility 293, Cecil Field NAS, FL	Soil	CESB11-8.5	T/F	0.12322581	(T/E)/FC	4 23456E-06	CEF-293-9FP	799000 ug/mL	3
NRMRI	JP-5	6/1/1981	#######	Facility 293, Cecil Field NAS, FL	Soil	CESB11-8.5'	T/X	0.04489892	(T/X)/FC	1.54292E-06	CEF-293-9EP	799000 µg/mL	1
NRMRL	JP-5	6/1/1981	#######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-8.5'	E/X	0.36436295	(E/X)/FC	1.25211E-05	CEF-293-9FP	799000 ug/mL	10
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-8.5	(B+T)/(E+X)	0.03458994	((B+T)/(E+X))/FC	1.18866E-06	CEF-293-9FP	799000 µg/mL	1
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-8.5'	Fuel Carbon	29100 mg/kg	Fuel Carbon	1	CEF-293-9FP	799000 ug/mL	799000
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-8.5'	1,2,3-Trimethylbenzene	99.8 mg/kg	1,2,3 TMB/FC	0.003429553	CEF-293-9FP	799000 ug/mL	2740
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-8.5'	1,2,4-Trimethylbenzene	238 mg/kg	1,2,4 TMB/FC	0.008178694	CEF-293-9FP	799000 ug/mL	6535
NRMRL	JP-5	6/1/1981	#######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-8.5'	1,3,5-Trimethylbenzene	96.9 mg/kg	1,3,5 TMB/FC	0.003329897	CEF-293-9FP	799000 ug/mL	2661
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-8.5'	1-MethylNaphthalene	94.4 mg/kg	1-MN/FC	0.003243986	CEF-293-9FP	799000 ug/mL	2592
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB11-8.5'	2-MethylNaphthalene	150 mg/kg	2-MN/FC	0.005154639	CEF-293-9FP	799000 ug/mL	4119
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB3-8.5'	Benzene	0.012 mg/kg	B/FC	3.64742E-06	ASSUMEdens	799000 ug/mL	3
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB3-8.5'	Toluene	0.208 mg/kg	T/FC	6.32219E-05	ASSUMEdens	799000 ug/mL	51
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB3-8.5'	Ethylbenzene	1.47 mg/kg	E/FC	0.000446809	ASSUMEdens	799000 ug/mL	357
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB3-8.5'	m-Xylene	3.26 mg/kg	m-Xylene/FC	0.000990881	ASSUMEdens	799000 ug/mL	792
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB3-8.5'	o-Xylene	0.666 mg/kg	o-Xylene/FC	0.000202432	ASSUMEdens	799000 ug/mL	162
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB3-8.5'	p-Xylene	1.31 mg/kg	p-Xylene/FC	0.000398176	ASSUMEdens	799000 ug/mL	318
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB3-8.5	Lotal Xylenes (m,p, and o)	5.236 mg/kg	X/FC	0.001591489	ASSUMEdens	799000 ug/mL	1272
	JP-5	6/1/1981	#######	Facility 293, Cecil Field NAS, FL	Soll	UFSB3-8.5	IOTAIBIEX	6.926 mg/kg	BIEX/FC	0.002105167	ASSUMEdens	799000 ug/mL	1682
	JP-5	6/1/1981	#######	Facility 293, Cecil Field NAS, FL	Soll	UFSB3-8.5	B+I	0.22 mg/kg	(B+1)/FC	6.68693E-05	ASSUMEdens	799000 ug/mL	53
	JP-5	6/1/1981	#######	Facility 293, Cecil Field NAS, FL	5011	UFSB3-8.5	E+A	6.706 mg/kg	(E+X)/FU	0.002038298	ASSUMEDENS	799000 ug/mL	1629
	JF-D	6/1/1981	######################################	Facility 293, Cecil Field NAS, FL	Soll	0F3B3-8.5		0.00916227	(D/1)/FC	1./030/E-05	ASSUMEdens	799000 ug/mL	14
	G-TL	6/1/1981	######################################	Facility 293, Cecil Field NAS, FL	Soil	CESE2 9 5'		0.00010327		2.40124E-06	ASSUIVIEUENS	799000 ug/mL	2
		6/1/1981	######################################	Facility 293, Cecil Field NAS, FL	Soil	CESE2 9 5'		0.00229103		4 20091E 0F	ASSUMEdana	799000 ug/mL	1
		6/1/1981	######################################	Facility 293, Cecil Field NAS, FL	Soil	CESE2 9 5'	T/V	0.1414900		4.30001E-05	ASSUMEdana	799000 ug/mL	34
	01710	0/1/1901	****	i admity 233, OCOLI FICIU INAO, FL		0-000-0.0	1//	0.00312430	(1/A)/FO	1.20143E-03	LOOOINIEUGIJS	r aanon ug/IIIF	10

Lab Code	Fuel Type	Spill Date	Date	Site Name	Matix	Locid	Analyte	Results	Units	Flag	FC Analyte	FC Ratio	FPLocid	FPdensity	FPunits	SoilFuel (ug/mL)
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB3-8.5'	E/X	0.28074866			(E/X)/FC	8.53339E-05	ASSUMEdens	799000	ug/mL	68
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB3-8.5'	(B+T)/(E+X)	0.03280644			((B+T)/(E+X))/FC	9.97156E-06	ASSUMEdens	799000	ug/mL	8
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB3-8.5'	Fuel Carbon	3290	mg/kg		Fuel Carbon	1	ASSUMEdens	799000	ug/mL	799000
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB3-8.5'	1,2,3-Trimethylbenzene	13.9	mg/kg		1,2,3 TMB/FC	0.004224924	ASSUMEdens	799000	ug/mL	3376
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB3-8.5'	1,2,4-Trimethylbenzene	21.5	mg/kg		1,2,4 TMB/FC	0.006534954	ASSUMEdens	799000	ug/mL	5221
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB3-8.5'	1,3,5-Trimethylbenzene	7.05	mg/kg		1,3,5 TMB/FC	0.002142857	ASSUMEdens	799000	ug/mL	1712
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB3-8.5'	1-MethylNaphthalene	16.4	mg/kg		1-MN/FC	0.004984802	ASSUMEdens	799000	ug/mL	3983
NRMRL	JP-5	6/1/1981	######	Facility 293, Cecil Field NAS, FL	Soil	CFSB3-8.5'	2-MethylNaphthalene	25.5	mg/kg		2-MN/FC	0.00775076	ASSUMEdens	799000	ug/mL	6193

Lab Code	Fuel Type	Spill Date	Date	Site Name	Matix	Locid	Analyte	Results	Units	Flag FC Analyte	FC Ratio FPLocid	FPdensity FPunits	SoilFuel (ug/mL)
NRMRL	JP-8	12/1/1995	#####	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	Benzene	3.06	6 mg/kg	B/FC	0.000221739 SJ98-MW1S	818000 ug/mL	181
NRMRL	JP-8	12/1/1995	#####	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	Toluene	17.2	2 mg/kg	T/FC	0.001246377 SJ98-MW1S	818000 ug/mL	1020
NRMRL	JP-8	12/1/1995	#####	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	Ethylbenzene	24.3	8 mg/kg	E/FC	0.00176087 SJ98-MW1S	818000 ug/mL	1440
NRMRL	JP-8	12/1/1995	5 #####	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	m-Xylene	53	8 mg/kg	m-Xylene/FC	0.00384058 SJ98-MW1S	818000 ug/mL	3142
NRMRL	JP-8	12/1/1995	5 ######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	o-Xylene	29.6	6 mg/kg	o-Xylene/FC	0.002144928 SJ98-MW1S	818000 ug/mL	1755
NRMRL	JP-8	12/1/1995	5 ######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	p-Xylene	18.9	mg/kg	p-Xylene/FC	0.001369565 SJ98-MW1S	818000 ug/mL	1120
NRMRL	JP-8	12/1/1995	5 #####	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	Total Xylenes (m,p, and o)	101.5	i mg/kg	X/FC	0.007355072 SJ98-MW1S	818000 ug/mL	6016
NRMRL	JP-8	12/1/1995	5 ######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	Total BTEX	146.06	6 mg/kg	BTEX/FC	0.010584058 SJ98-MW1S	818000 ug/mL	8658
NRMRL	JP-8	12/1/1995	5 #####	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	B+T	20.26	6 mg/kg	(B+T)/FC	0.001468116 SJ98-MW1S	818000 ug/mL	1201
NRMRL	JP-8	12/1/1995	<i>#####</i>	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	B/T	0.17790698	3	(B/T)/FC	1.28918E-05 SJ98-MW1S	818000 ug/mL	11
NRMRL	JP-8	12/1/1995	######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	B/E	0.12592593	3	(B/E)/FC	9.12507E-06 SJ98-MW1S	818000 ug/mL	7
NRMRL	JP-8	12/1/1995	<i>#####</i>	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	B/X	0.03014778	3	(B/X)/FC	2.18462E-06 SJ98-MW1S	818000 ug/mL	2
NRMRL	JP-8	12/1/1995	######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	T/E	0.70781893	3	(T/E)/FC	5.12912E-05 SJ98-MW1S	818000 ug/mL	42
NRMRL	JP-8	12/1/1995	<i>#####</i>	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	T/X	0.16945813	3	(T/X)/FC	1.22796E-05 SJ98-MW1S	818000 ug/mL	10
NRMRL	JP-8	12/1/1995	######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	E/X	0.23940887	·	(E/X)/FC	1.73485E-05 SJ98-MW1S	818000 ug/mL	14
NRMRL	JP-8	12/1/1995	######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	Fuel Carbon	13800) mg/kg	Fuel Carbon	1 SJ98-MW1S	818000 ug/mL	818000
NRMRL	JP-8	12/1/1995	#####	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	1,2,3-Trimethylbenzene	60.4	mg/kg	1,2,3 TMB/FC	0.004376812 SJ98-MW1S	818000 ug/mL	3580
NRMRL	JP-8	12/1/1995	#####	# Bidg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	1,2,4-Irimethylbenzene	118	8 mg/kg	1,2,4 TMB/FC	0.008550725 SJ98-MW1S	818000 ug/mL	6994
NRMRL	JP-8	12/1/1995	#####	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	1,3,5-Trimethylbenzene	38.3	8 mg/kg	1,3,5 TMB/FC	0.002775362 SJ98-MW1S	818000 ug/mL	2270
NRMRL	JP-8	12/1/1995	######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	1-MethylNaphthalene	35	mg/kg	1-MN/FC	0.002536232 SJ98-MW1S	818000 ug/mL	2075
NRMRL	JP-8	12/1/1995	#####	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB1-3	2-MethylNaphthalene	52.5	mg/kg	2-MN/FC	0.003804348 SJ98-MW1S	818000 ug/mL	3112
NRMRL	JP-8	12/1/1995	#####	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-3	Benzene	3.47	mg/kg	B/FC	0.000164455 SJ98-MP2	812000 ug/mL	134
NRMRL	JP-8	12/1/1995	######	# Bidg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-3	Ioluene	34.6	i mg/kg	T/FC	0.00163981 SJ98-MP2	812000 ug/mL	1332
NRMRL	JP-8	12/1/1995	#####	# Bidg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-3	Ethylbenzene	54.1	mg/kg	E/FC	0.002563981 SJ98-MP2	812000 ug/mL	2082
NRMRL	JP-8	12/1/1995	#####	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-3	m-Xylene	103	8 mg/kg	m-Xylene/FC	0.004881517 SJ98-MP2	812000 ug/mL	3964
NRMRL	JP-8	12/1/1995	#####	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-3	o-Xylene	67	mg/kg	o-Xylene/FC	0.003175355 SJ98-MP2	812000 ug/mL	2578
NRMRL	JP-8	12/1/1995	#####	# Bidg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-3	p-Xylene	38.4	mg/kg	p-Xylene/FC	0.001819905 SJ98-MP2	812000 ug/mL	1478
NRMRL	JP-8	12/1/1995	#####	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-3	I otal Xylenes (m,p, and o)	208.4	mg/kg	X/FC	0.009876777 SJ98-MP2	812000 ug/mL	8020
NRMRL	JP-8	12/1/1995	#####	# Bidg 4522, Seymour Johnson AFB, SC	Soll	SJ98-SB2-3	IOTALBIEX	300.57	mg/kg	BIEX/FC	0.014245024 SJ98-MP2	812000 ug/mL	11567
NRMRL	JP-8	12/1/1995	######	# Bidg 4522, Seymour Johnson AFB, SC	Soll	SJ98-SB2-3	B+1	38.07	mg/kg	(B+T)/FC	0.001804265 SJ98-MP2	812000 ug/mL	1465
NRIVIRL	JP-8	12/1/1995	• ######	# Bidg 4522, Seymour Johnson AFB, SC	Soli	SJ98-SB2-3	B/T	0.10028902	-	(B/T)/FC	4.75303E-06 SJ98-MP2	812000 ug/mL	4
NRMRL	JP-8	12/1/1995	++++++++++++++++++++++++++++++++++++++	# Bidg 4522, Seymour Johnson AFB, SC	501	SJ98-SB2-3	B/E	0.06414048		(B/E)/FC	3.03983E-06 SJ98-MP2	812000 ug/mL	2
NRIVIRL	JP-8	12/1/1995	· ######	# Bidg 4522, Seymour Johnson AFB, SC	Soll	SJ98-SB2-3	B/A	0.01665067			7.89131E-07 SJ98-MP2	812000 ug/mL	1
	JP-0	12/1/1995		H Bidg 4522, Seymour Johnson AFB, SC	Soil	SJ96-SB2-3	1/E	0.63955636	7		3.03107E-05 SJ96-MP2	812000 ug/mL	25
		12/1/1995	<i>+++++++</i>	H Bidg 4522, Seymour Johnson AFB, SC	Soil	SJ90-SD2-3	1/A E/X	0.10002007	,		1 22022E 05 S 109 MP2	812000 ug/mL	10
		12/1/1995	<i></i>	H Bidg 4522, Seymour Johnson AFB, SC	Soil	SJ90-SB2-3	E/A Eucl Carbon	0.23939093) ma/ka	Evol Carbon	1.23032E-03 3390-WF2	812000 ug/mL	P12000
		12/1/1995	· · · · · · · · · · · · · · · · · · ·	H Bidg 4522, Seymour Johnson AFB, SC	Soil	S 100 CD2 2	1.2.2 Trimothylbonzono	21100	mg/kg	1 2 2 TMP/EC	0.007014219 S IO9 MP2	812000 ug/mL	5606
NDMDI		12/1/1995	<i><i>нини</i></i>	H Bidg 4522, Seymour Johnson AFB, SC	Soil	S 100 SB2 2	1.2.4 Trimethylbenzene	266	mg/kg	1.2.4 TMP/EC	0.007014210 00000112	812000 ug/mL	10227
NEMEL	JF=0 ID_8	12/1/1995	++++++++++++++++++++++++++++++++++++++	H Bidg 4522, Seymour Johnson AFB, SC	Soil	S 108-SB2-3	1.3.5-Trimethylbenzene	78.2	mg/kg	1.2,4 TMB/FC	0.012000033 SJ98-MF2	812000 ug/mL	3000
NEMEI	JI -0	12/1/1995	<i>######</i>	H Bldg 4522, Seymour Johnson AFB, SC	Soil	S 108-SB2-3	1-MethylNaphthalene	87.9	mg/kg	1,3,5 HVID/1 C	0.003700101 S330-MI 2	812000 ug/ml	3370
NEMEI	JI -0	12/1/1995	<i>######</i>	H Bldg 4522, Seymour Johnson AFB, SC	Soil	S 108-SB2-3	2-MethylNaphthalene	129	mg/kg	2-MN/EC	0.004101137 S330-MI 2	812000 ug/ml	4026
NRMRI	IP-8	12/1/1995	<u>######</u>	# Bldg 4522, Seymour Johnson AFB, SC	Soil	S 198-SB2-4	Benzene	1.82	ma/ka	B/FC	7 87879E-05 S 198-MP2	812000 ug/mL	4320
NRMRI	IP-8	12/1/1995	<u>######</u>	# Bldg 4522, Seymour Johnson AFB_SC	Soil	S 198-SB2-4	Toluene	29.3	l mg/kg	T/FC	0.001268398 S I98-MP2	812000 ug/ml	1030
NRMRI	JP-8	12/1/1995	######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	S.198-SB2-4	Ethylbenzene	39.0	ma/ka	E/EC	0.001727273 SJ98-MP2	812000 ug/ml	1403
NRMRI	JP-8	12/1/1995	######	# Bldg 4522 Seymour Johnson AFB_SC	Soil	S.198-SB2-4	m-Xylene	64 6	ma/ka	m-Xylene/FC	0.002796537 SJ98-MP2	812000 ug/ml	2271
NRMRL	JP-8	12/1/1995	######	# Bldg 4522, Seymour Johnson AFB. SC	Soil	SJ98-SB2-4	o-Xvlene	41.6	6 ma/ka	o-Xvlene/FC	0.001800866 SJ98-MP2	812000 ug/mL	1462
NRMRL	JP-8	12/1/1995	######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-4	p-Xvlene	21.3	3 ma/ka	p-Xylene/FC	0.000922078 SJ98-MP2	812000 ug/mL	749
NRMRL	JP-8	12/1/1995	######	# Bldg 4522, Seymour Johnson AFB. SC	Soil	SJ98-SB2-4	Total Xylenes (m.p. and o)	127.5	mg/ka	X/FC	0.005519481 SJ98-MP2	812000 ug/mL	4482
NRMRL	JP-8	12/1/1995	######	# Bldg 4522, Seymour Johnson AFB. SC	Soil	SJ98-SB2-4	Total BTEX	198.52	2 mg/ka	BTEX/FC	0.008593939 SJ98-MP2	812000 ua/mL	6978
NRMRL	JP-8	12/1/1995	######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-4	B+T	31.12	2 ma/ka	(B+T)/FC	0.001347186 SJ98-MP2	812000 ug/mL	1094
NRMRL	JP-8	12/1/1995	#####	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-4	B/T	0.06211604	1	(B/T)/FC	2.68901E-06 SJ98-MP2	812000 ug/mL	2
NRMRL	JP-8	12/1/1995	######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-4	B/E	0.04561404	ł	(B/E)/FC	1.97463E-06 SJ98-MP2	812000 ug/mL	2
NRMRL	JP-8	12/1/1995	5 ######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-4	B/X	0.01427451		(B/X)/FC	6.17944E-07 SJ98-MP2	812000 ug/mL	1
NRMRL	JP-8	12/1/1995	5 ######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-4	T/E	0.73433584	Ļ	(T/E)/FC	3.17894E-05 SJ98-MP2	812000 ug/mL	26
NRMRL	JP-8	12/1/1995	######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-4	T/X	0.22980392	2	(T/X)/FC	9.94822E-06 SJ98-MP2	812000 ug/mL	8
NRMRL	JP-8	12/1/1995	<i>#####</i>	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-4	E/X	0.31294118	3	(E/X)/FC	1.35472E-05 SJ98-MP2	812000 ug/mL	11
NRMRL	JP-8	12/1/1995	<i>#####</i>	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-4	Fuel Carbon	23100) mg/kg	Fuel Carbon	1 SJ98-MP2	812000 ug/mL	812000
NRMRL	JP-8	12/1/1995	<i>#####</i>	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-4	1,2,3-Trimethylbenzene	98.1	mg/kg	1,2,3 TMB/FC	0.004246753 SJ98-MP2	812000 ug/mL	3448
NRMRL	JP-8	12/1/1995	<i>#####</i>	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-4	1,2,4-Trimethylbenzene	206	6 mg/kg	1,2,4 TMB/FC	0.008917749 SJ98-MP2	812000 ug/mL	7241
NRMRL	JP-8	12/1/1995	#####	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-4	1,3,5-Trimethylbenzene	48.6	6 mg/kg	1,3,5 TMB/FC	0.002103896 SJ98-MP2	812000 ug/mL	1708
NRMRL	JP-8	12/1/1995	#####	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-4	1-MethylNaphthalene	60.4	mg/kg	1-MN/FC	0.002614719 SJ98-MP2	812000 ug/mL	2123
NRMRL	JP-8	12/1/1995	######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJ98-SB2-4	2-MethylNaphthalene	87.2	2 mg/kg	2-MN/FC	0.003774892 SJ98-MP2	812000 ug/mL	3065
NRMRL	JP-8	12/1/1995	######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5'	Benzene	10.8	8 mg/kg	B/FC	0.000486486 SJMW1SFP	793000 ug/mL	386
NRMRL	JP-8	12/1/1995	######	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5'	Toluene	64.7	mg/kg	T/FC	0.002914414 SJMW1SFP	793000 ug/mL	2311
NRMRL	JP-8	12/1/1995	; ## ###	# Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5	Ethylbenzene	56.1	mg/kg	E/FC	0.002527027 SJMW1SFP	793000 ug/mL	2004

Lab Code	Fuel Type	Spill Date	Date	Site Name	Matix	Locid	Analyte	Results	Units	Flag FC Analyte	FC Ratio	FPLocid	FPdensity FPunits	SoilFuel (ug/mL)
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5'	m-Xylene	152	2 mg/kg	m-Xylene/FC	0.006846847	SJMW1SFP	793000 ug/mL	5430
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5'	o-Xylene	86.7	7 mg/kg	o-Xylene/FC	0.003905405	SJMW1SFP	793000 ug/mL	3097
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5'	p-Xylene	53.6	6 mg/kg	p-Xylene/FC	0.002414414	SJMW1SFP	793000 ug/mL	1915
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5'	Total Xylenes (m,p, and o)	292.3	3 mg/kg	X/FC	0.013166667	SJMW1SFP	793000 ug/mL	10441
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5'	Total BTEX	423.9) mg/kg	BTEX/FC	0.019094595	SJMW1SFP	793000 ug/mL	15142
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5'	B+T	75.5	5 mg/kg	(B+T)/FC	0.003400901	SJMW1SFP	793000 ug/mL	2697
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5'	E+X	348.4	1 mg/kg	(E+X)/FC	0.015693694	SJMW1SFP	793000 ug/mL	12445
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5'	B/T	0.16692427	7	(B/T)/FC	7.51911E-06	SJMW1SFP	793000 ug/mL	6
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5'	B/E	0.19251337	7	(B/E)/FC	8.67177E-06	SJMW1SFP	793000 ug/mL	7
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5'	B/X	0.03694834	1	(B/X)/FC	1.66434E-06	SJMW1SFP	793000 ug/mL	1
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5'	T/E	1.15329768	3	(T/E)/FC	5.19503E-05	SJMW1SFP	793000 ug/mL	41
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5'	T/X	0.22134793	3	(T/X)/FC	9.97063E-06	SJMW1SFP	793000 ug/mL	8
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5'	E/X	0.1919261	1	(E/X)/FC	8.64532E-06	SJMW1SFP	793000 ug/mL	7
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5'	(B+T)/(E+X)	0.21670494	1	((B+T)/(E+X))/FC	9.76148E-06	SJMW1SFP	793000 ug/mL	8
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5	Fuel Carbon	22200) mg/kg	Fuel Carbon	1	SJMW1SFP	793000 ug/mL	793000
NRMRL	JP-8	12/1/1995	######	Bidg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5	1,2,3-I rimethylbenzene	119	mg/kg	1,2,3 TMB/FC	0.00536036	SJMW1SFP	793000 ug/mL	4251
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5	1,2,4-I rimethylbenzene	269	mg/kg	1,2,4 TMB/FC	0.012117117	SJMW1SFP	793000 ug/mL	9609
NRMRL	JP-8	12/1/1995	######	Bidg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5	1,3,5- I rimethylbenzene	102	2 mg/kg	1,3,5 TMB/FC	0.004594595	SJMW1SFP	793000 ug/mL	3644
NRMRL	JP-8	12/1/1995	######	Bidg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5	1-MethylNaphthalene	62.4	1 mg/kg	1-MN/FC	0.002810811	SJMW1SFP	793000 ug/mL	2229
NRMRL	JP-8	12/1/1995	######	Bidg 4522, Seymour Johnson AFB, SC	Soil	SJSB1-5.5	2-MethylNaphthalene	96.9	mg/kg	2-MN/FC	0.004364865	SJMW1SFP	793000 ug/mL	3461
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB2-5.5	Benzene	12.5	mg/kg	B/FC	0.000313283	SJMW1SFP	793000 ug/mL	248
NRMRL	JP-8	12/1/1995	######	Bidg 4522, Seymour Johnson AFB, SC	Soll	SJSB2-5.5	Toluene	56.3	s mg/kg	T/FC	0.001411028	SJMW1SFP	793000 ug/mL	1119
NRMRL	JP-8	12/1/1995	######	Bidg 4522, Seymour Johnson AFB, SC	Soil	SJSB2-5.5	Ethylbenzene	75.3	s mg/kg	E/FC	0.001887218	SJMW1SFP	793000 ug/mL	1497
NRMRL	JP-8	12/1/1995	######	Bidg 4522, Seymour Johnson AFB, SC	Soll	SJSB2-5.5	m-xylene	217	mg/kg	m-xylene/FC	0.005438596	SJMW1SFP	793000 ug/mL	4313
NRMRL	JP-8	12/1/1995	######	Bidg 4522, Seymour Johnson AFB, SC	Soll	SJSB2-5.5	o-Xylene	124	i mg/кg	o-Xylene/FC	0.003107769	SJMW1SFP	793000 ug/mL	2464
NRMRL	JP-8	12/1/1995	#######	Bidg 4522, Seymour Johnson AFB, SC	Soll	SJSB2-5.5	p-Xylene	75.4	i mg/kg	p-Xylene/FC	0.001889724	SJMW1SFP	793000 ug/mL	1499
NRMRL	JP-8	12/1/1995	****	Bidg 4522, Seymour Johnson AFB, SC	Soli	SJSB2-5.5	Total Xylenes (m,p, and o)	416.4	i mg/kg	X/FC	0.01043609	SJMWISFP	793000 ug/mL	8276
NRIVIRL	JP-8	12/1/1995	######	Bidg 4522, Seymour Johnson AFB, SC	Soll	SJSB2-5.5	I OTAI BIEX	560.5	s mg/kg	BIEX/FC	0.014047619	SJMWISFP	793000 ug/mL	11140
NRIVIRL	JP-8	12/1/1995	#######	Bidg 4522, Seymour Johnson AFB, SC	Soli	SJSB2-5.5	B+1	68.8	s mg/kg	(B+T)/FC	0.001724311	SJMWISFP	793000 ug/mL	1367
NRIVIRL	JP-8	12/1/1995	#######	Bidg 4522, Seymour Johnson AFB, SC	Soli	SJSB2-5.5	E+A D/T	491.7	mg/kg		0.012323308	SJMW1SFP	793000 ug/mL	9//2
	JP-0	12/1/1995	#######	Bidg 4522, Seymour Johnson AFB, SC	Soil	SJSD2-5.5	D/I D/F	0.22202467			3.36453E-06	SJIVIWISFP	793000 ug/mL	4
	JP-0	12/1/1995	#######	Bidg 4522, Seymour Johnson AFB, SC	Soil	SJSD2-5.5	D/E D/V	0.16600266			4.16047E-06	SJIVIWISFP	793000 ug/mL	3
		12/1/1995	#######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSD2-3.3		0.03001921	2		1.52301E-07	SJIVIVI SFP	793000 ug/mL	15
		12/1/1995	*****	Bidg 4522, Seymour Johnson AFB, SC	Soil	0J0B2-0.0		0.14707590	2	(1/E)/FC	2.200625.06	SJIVIWISFF	793000 ug/mL	10
		12/1/1995	######################################	Bidg 4522, Seymour Johnson AFB, SC	Soil	SJSB2-5.5		0.13320033			4 52222E 06	SJIVIWISFF	793000 ug/mL	3
		12/1/1995	######################################	Bidg 4522, Seymour Johnson AFB, SC	Soil	SJSB2-5.5		0.18083573	2		2.50694E.06	SJIVIWISFF	793000 ug/mL	4
	JF=0 ID_8	12/1/1995	########	Bidg 4522, Seymour Johnson AFB, SC	Soil	S ISB2-5.5	Euel Carbon	30000) ma/ka		3.30004E-00	SIMWISFF	793000 ug/mL	703000
NEMEI	JI -0	12/1/1995	######################################	Bidg 4522, Seymour Johnson AFB, SC	Soil	S ISB2-5.5	1 2 3-Trimethylbenzene	208	ma/ka	1 2 3 TMB/EC	0.005213033	S IMW/1SFP	793000 ug/mL	133000
NRMRI	JP-8	12/1/1995	#######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB2-5.5'	1 2 4-Trimethylbenzene	434	1 ma/ka	1.2.4 TMB/FC	0.010877193	S.IMW1SEP	793000 ug/ml	8626
NRMRI	JP-8	12/1/1995	#######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB2-5.5'	1.3.5-Trimethylbenzene	156	ma/ka	1.3.5 TMB/FC	0.003909774	S.IMW1SEP	793000 ug/ml	3100
NRMRI	JP-8	12/1/1995	#######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB2-5.5'	1-MethylNaphthalene	113	3 ma/ka	1-MN/FC	0.00283208	S.IMW1SEP	793000 ug/ml	2246
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	SJSB2-5.5'	2-MethylNaphthalene	172	2 ma/ka	2-MN/FC	0.004310777	SJMW1SFP	793000 ug/mL	3418
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	Benzene	11.4	1 ma/ka					no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	Toluene	32.5	5 ma/ka					no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB. SC	Soil	Soil Sample 1	Ethylbenzene	32.2	2 mg/ka					no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	m-Xylene	68.9	mg/kg					no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	o-Xylene	43.1	mg/kg					no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	p-Xylene	24.5	5 mg/kg					no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	Total Xylenes (m,p, and o)	136.5	5 mg/kg					no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	Total BTEX	212.6	6 mg/kg					no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	B+T	43.9	mg/kg					no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	E+X	168.7	7 mg/kg					no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	B/T	0.35076923	3					no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	B/E	0.35403727	7					no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	B/X	0.08351648	3					no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	T/E	1.00931677	7					no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	T/X	0.23809524	1					no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	E/X	0.23589744	1					no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	(B+T)/(E+X)	0.26022525	5					no FC data
NRMRL	JP-8	12/1/1995	#######	Bidg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	1,2,3-Trimethylbenzene	56.4	i mg/kg					no FC data
NRMRL	JP-8	12/1/1995	#######	Bidg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 1	1,2,4-Trimethylbenzene	129	mg/kg					no FC data
NRMRL	JP-8	12/1/1995	#######	Blag 4522, Seymour Johnson AFB, SC	Sol	Soil Sample 1	1,3,5-Irimethylbenzene	48.2	2 mg/kg					no FC data
NRMRL	JP-8	12/1/1995	#######	Blag 4522, Seymour Johnson AFB, SC	Sol	Soil Sample 1	1-MethylNaphthalene	26.4	i mg/kg					no FC data
NRMRL	JP-8	12/1/1995	#######	Blag 4522, Seymour Johnson AFB, SC	Sol	Soil Sample 1	2-MethylNaphthalene	42	2 mg/kg					no FC data
NRMRL	JH-8	12/1/1995	######	ыад 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 2	Benzene	11.1	mg/kg					no FC data

Lab Code	Fuel Type	Spill Date	Date	Site Name	Matix	Locid	Analyte	Results	Units Flag	FC Analyte	FC Ratio	FPLocid	FPdensity	FPunits	SoilFuel (ug/mL)
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 2	Toluene	78.6	mg/kg						no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 2	Ethylbenzene	64.2	mg/kg						no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 2	m-Xylene	115	mg/kg						no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 2	o-Xylene	72.4	mg/kg						no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 2	p-Xylene	45.7	mg/kg						no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 2	Total Xylenes (m,p, and o)	233.1	mg/kg						no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 2	Total BTEX	387	mg/kg						no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 2	B+T	89.7	mg/kg						no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 2	E+X	297.3	mg/kg						no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 2	B/I	0.14122137							no FC data
NRMRL	JP-8	12/1/1995	######	Bldg 4522, Seymour Johnson AFB, SC	Soll	Soil Sample 2	B/E	0.1728972							no FC data
NRMRL	JP-8	12/1/1995	######	Bidg 4522, Seymour Johnson AFB, SC	Soll	Soil Sample 2	B/X	0.04761905							no FC data
NRIVIRL	JP-8	12/1/1995	****	Bidg 4522, Seymour Johnson AFB, SC	5011	Soil Sample 2	T/E	1.22429907							no FC data
NRIVIRL	JP-8	12/1/1995	****	Bidg 4522, Seymour Johnson AFB, SC	Soll	Soil Sample 2	1/X	0.33719434							no FC data
NRIVIRL	JP-0	12/1/1995	****	Bidg 4522, Seymour Johnson AFB, SC	Soil	Soli Sample 2		0.27541626							no FC data
		12/1/1995	########	Bidg 4522, Seymour, Johnson AFB, SC	Soil	Soil Sample 2		6.30171344	ma/ka						no FC data
	JF-0 IP-8	12/1/1995	#######	Bldg 4522, Seymour, Johnson AFB, SC	Soil	Soil Sample 2	1.2.4-Trimethylbenzene	1/1	mg/kg						no FC data
NEMEI	IP-8	12/1/1995	#######	Bidg 4522, Seymour, Johnson AFB, SC	Soil	Soil Sample 2	1.3.5-Trimethylbenzene	50.6	mg/kg						no FC data
NRMRI	JP-8	12/1/1995	#######	Bldg 4522, Seymour Johnson AFB, SC	Soil	Soil Sample 2	1-MethylNaphthalene	33.4	ma/ka						no FC data
NRMRI	JP-8	12/1/1995	#######	Bldg 4522, Seymour Johnson AFB_SC	Soil	Soil Sample 2	2-MethylNaphthalene	48 3	ma/ka						no FC data
NRMRL	JP-8	4/1/1996	#######	JP-8 Release Site, Pope AFB, NC	Soil	7POP2	Benzene	0,006	ma/ka **						no FC data
NRMRL	JP-8	4/1/1996	#######	JP-8 Release Site, Pope AFB, NC	Soil	7POP2	Toluene	0,006	ma/ka **						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	7POP2	Ethylbenzene	0.006	mg/kg **						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	7POP2	m-Xylene	0.006	mg/kg **						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	7POP2	o-Xylene	0.006	mg/kg **						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	7POP2	p-Xylene	0.006	mg/kg **						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	7POP2	Total Xylenes (m,p, and o)	0.018	mg/kg						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	7POP2	Total BTEX	0.036	mg/kg						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	7POP2	B+T	0.012	mg/kg						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	7POP2	E+X	0.024	mg/kg						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	7POP2	B/T	1							no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	7POP2	B/E	1							no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	7POP2	B/X	0.33333333							no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	7POP2	T/E	1							no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	7POP2	1/X	0.333333333							no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soll	7POP2		0.33333333							no FC data
NRMRL	JP-8	4/1/1996	#######	JP-8 Release Site, Pope AFB, NC	Soll	7POP2	(B+1)/(E+X)	0.5	ma/ka **						no FC data
	JP-0 IP-8	4/1/1996	#######	JP-6 Release Site, Pope AFB, NC	Soil	7POP2	1,2,3-Trimethylbenzene	0.006	mg/kg **						no FC data
	JF-0 IP-8	4/1/1990	#######	IP-8 Release Site, Pope AFB, NC	Soil	7POP2	1.3.5-Trimethylbenzene	0.000	mg/kg **						no FC data
NRMRI	IP-8	4/1/1996	#######	IP-8 Release Site, Pope AFB, NC	Soil	7POP2	1-MethylNaphthalene	0.000	mg/kg **						no FC data
NRMRI	JP-8	4/1/1996	#######	JP-8 Release Site, Pope AFB, NC	Soil	7POP2	2-MethylNaphthalene	0.006	ma/ka **						no FC data
NRMRL	JP-8	4/1/1996	#######	JP-8 Release Site, Pope AFB, NC	Soil	9POP2	Benzene	0.006	ma/ka **						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	9POP2	Toluene	0.006	ma/ka **						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	9POP2	Ethylbenzene	0.006	mg/kg **						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	9POP2	m-Xylene	0.006	mg/kg **						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	9POP2	o-Xylene	0.006	mg/kg **						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	9POP2	p-Xylene	0.006	mg/kg **						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	9POP2	Total Xylenes (m,p, and o)	0.018	mg/kg						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	9POP2	Total BTEX	0.036	mg/kg						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	9POP2	B+T	0.012	mg/kg						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	9POP2	E+X	0.024	mg/kg						no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	9POP2	B/T	1							no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	9POP2	B/E	1							no FC data
NRMRL	JP-8	4/1/1996	######	JP-8 Release Site, Pope AFB, NC	Soil	9POP2	B/X	0.33333333							no FC data
NRMRL	JP-8	4/1/1996	#######	JP-8 Release Site, Pope AFB, NC	Soil	92022	1/E	1							no FC data
NRMRL	JP-8	4/1/1996	#######	JP-8 Release Site, Pope AFB, NC	Soil	92022	1/X	0.33333333							no FC data
NRMRL	JP-8	4/1/1996	#######	JP-8 Release Site, Pope AFB, NC	501	92022		0.333333333							no FC data
	JF-8	4/1/1996	#######	JP-8 Release Site, Pope AFB, NU	Soil	9POP2	(B+I)/(E+X)	0.5	ma/ka						no FC data
	JF-0	4/1/1996	****	JE-0 Nelease Sile, Pupe AFB, NU	Soil	3POP2	1.2.3-Trimethylbenzene	0.147	mg/kg @						no FC data
	JF-0 ID-8	4/1/1996	######################################	JE-0 Release Site, Pupe AFB, NC	Soil	3POP2	1.3.5-Trimethylbenzene	0.219	mg/kg @						no FC data
	ID-8	4/1/1006	######################################	ID-8 Release Site, Pope AFB, NC	Soil		1-MethylNaphthaleno	0.0911	mg/kg @						no FC data
NRMRI	IP-8	4/1/1990	######################################	IP-8 Release Site, Pone AFR NC	Soil	9POP2	2-MethylNaphthalene	0.239	ma/ka @						no FC data
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APPENDIX D

LETTER OF TRANSMITTAL AND RESPONSE TO COMMENTS

09 September 1999

Mr. Jerry Hansen AFCEE/ERT 3207 North Road, Bldg. 532 Brooks AFB, Texas 78235-5363

Subject: Submittal of the Final Light Nonaqueous-Phase Liquid Weathering at Various Fuel Release Sites, September 1999 (Contract F41624-92-D-8036-0025)

Dear Mr. Hansen

Enclosed please find two copies of the September 1999 Final Light Nonaqueous-Phase Liquid Weathering at Various Fuel Release Sites. This report was prepared by Parsons Engineering Science, Inc. (Parsons ES) for the Air Force Center for Environmental Excellence Technology Transfer Division (AFCEE/ERT). The intent of this report was to determine rates of natural attenuation (weathering) of liquid nonaqueous-phase liquids (LNAPL).

The draft LNAPL weathering report was submitted to AFCEE in January 1999. Comments on the draft report were received from AFCEE as reviewed by Mr. Daniel Kraft of the Waste Policy Institute (WPI) in San Antonio, Texas, and Mr. Jon Atkinson of the Consultant Operations Division. Responses to these comments were prepared by Parsons ES and are attached to this letter.

If you have any questions or require additional information, please call me at (303) 831-8100.

Sincerely,

PARSONS ENGINEERING SCIENCE, INC.

Bruce M. Henry, P.G. Project Manager

Enclosures

cc: Don Kampbell – USEPA NRMRL File 729691.35000

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A. RESPONSES TO MR. KRAFT'S COMMENTS:

I. General Comments and Parsons ES Responses:

1. Clarity and Organization

The document is well organized and easy to follow. The investigative work and the rationale and equations used to study the effects of the various weathering mechanisms for each of the fuel hydrocarbons in the study (gasoline, JP-4, JP-5, and JP-8) are clearly presented.

Parsons ES Response:

No Comment.

2. Completeness

The document is generally complete and provides a thorough yet concise summary of the literature review conducted and the chemistry and compositional makeup of the fuel hydrocarbons evaluated in the study. A table at the end of Section 6 summarizing the findings of the study and comparing them to the range of values currently in use would be helpful.

Parsons ES Response:

A table will be added to Section 6 that summarizes the BTEX weathering rates observed for JP-4 and JP-8 mobile LNAPLs. Because BTEX weathering rates for JP-5 and gasoline mobile LNAPLs could not be meaningfully determined based on study results, no data will be presented for these fuel types.

3. Technical Issues

Weathering of petroleum fuels in the subsurface environment is affected by numerous physical and chemical processes that exhibit a high degree of variability within a given fuel type, from one site to another, and spatially within each site. Moreover, the accurate measurement of many of these variables is difficult and laboratory derived values often deviate significantly from those observed in the field. This requires the use of a range of possible values and the selection of the most conservative value for defensibility. When multiple parameters derived in this fashion are used to calculate LNAPL contaminant depletion rates, accuracy is diminished and the results may be too conservative.

Overall, the study was only moderately successful at improving the scientific basis and defensibility for determining natural LNAPL weathering rates. BTEX weathering rates for gasoline, JP-5, and JP-8 (three of the four fuel types investigated) were indeterminate. The

report does provide some empirical justification for using the lowest depletion rate from the range of values determined for JP-4.

Parsons ES Response:

During development of the work plan for this fuels weathering study, the primary fuel types of interest were JP-4 jet fuel and gasoline. JP-5 and JP-8 fuel release sites were later added to the study when it became apparent that insufficient JP-4 and gasoline sites were available that met the original site selection criteria. While BTEX weathering rate data for the JP-5 and gasoline release sites was indeterminate, mobile LNAPL BTEX weathering at the one JP-8 site strongly corroborates BTEX weathering rates observed at JP-4 sites with similar hydrogeologic conditions (sandy soils and high groundwater velocities).

II. Specific Comments with Responses:

Item 1 – Page 2-22, Para 1:

The text states that under nonequilibrium conditions the dissolution rate cannot be enhanced by advection or biodegradation that further reduces the aqueous concentration. This implies that for aqueous concentrations less than the equilibrium concentration, the dissolution rate is constant irrespective of the aqueous concentration (i.e., the dissolution rate is linear below equilibrium concentrations). WPI recommends verifying that this is correct and rewrite the text, if necessary.

Parsons ES Response:

No revision of the text is required. The suppositions drawn by the reviewer cannot be supported by information obtained during the literature search. The findings presented by Seagren et al. (1993) are correctly referenced and no corrections are warranted.

Item 2 – Page 2-25, Para 1, Line 1

The text states that biodegradation of dissolved petroleum contaminants reduces aqueous contaminant concentrations and thereby enhances dissolution rates. This appears to be inconsistent with the statement made on page 2-22 that dissolution rates cannot be enhanced by biodegradation (see Comment 1). WPI recommends clarification.

Parsons ES Response:

The referenced statement on Page 2-22 refers to dissolution under nonequilibrium conditions. If equilibrium conditions exist, biodegradation of dissolved petroleum contaminants will reduce aqueous concentrations and enhance dissolution rates. The referenced sentence will be modified to state specifically that dissolution is enhanced by biodegradation under equilibrium conditions.

Item 3 – Page 2-26, Para 4, Line 1

The use of the term "interphase" do describe the transition zone is unclear. Rewrite the text for clarity.

Parsons ES Response:

The word "interface" will be used in place of "interphase" at the referenced location.

Item 4 – Page 2-29, Para 2, Bullet 2

Replace "Oil-mass rates" with "Oil-mass loss rates."

Parsons ES Response:

Parsons ES concurs. The requested change will be made.

Item 5 – Page 2-29, Para 2, Bullet 3

It would seem intuitive that 10 years of data would provide a more accurate decline curve depicting mass loss rates. Provide more detail concerning the "speculative" nature of the yearly data to demonstrate why this was an important conclusion of the study.

Parsons ES Response:

Parsons ES concurs. The referenced conclusion was not an important finding of the Landon and Hult (1991) study; and as a result, it will be deleted from the final report.

Item 6 – Page 2-31, Para 3, Line 2

For clarity, replace "benzene+toluene/ethylbenze+xylenes" with "(benzene+toluene)/(ethylbenzene+xylenes)" and replace "B+T/E+X" with (B+T)/(E+X)," In addition, replace all instances of these expressions.

Parsons ES Response:

Parsons ES concurs. The requested changes will be made.

Item 7 – Page 5-3, Para 2, Line 9

As punctuated, the sentence implies that all of the JP-4 sites were older than 20 years. For clarity, replace "at JP-4 sites, where fuel releases" with "at JP-4 sites where fuel releases."

Parsons ES Response:

Parsons ES concurs. The requested change will be made.

Item 8 – Page 5-6, Para 3, Line 2

Replace "the amount of contaminant depletion" with "the rate of contaminant depletion."

Parsons ES Response:

Parsons ES concurs. The requested change will be made.

Item 9 – Page 6-1, Para 1, Line 10

Replace "inflate long-term monitoring and site management costs" with "inflate projected long-term monitoring and site management costs."

Parsons ES Response:

Parsons ES concurs. The requested change will be made.

Item 10 – Page 6-2, Bullet 6

It is not clear if the statements provided reflect findings of the study or are suppositions that should be located in Section 2 of the report. Provide study results that support this finding.

Parsons ES Response:

Data collected from the DFSP-Charleston site and the Offutt AFB site support the finding presented. Varying mobile LNAPL weathering rates were observed at both these sites likely as a result of sample location within the LNAPL plume. A reference to the results from these two study locations will be added to the text.

Item 11 – Page 6-3, Bullet 2

For clarity, replace "estimating total BTEX weathering from LNAPL" with "estimating total BTEX weathering from JP-4 LNAPL."

Parsons ES Response:

Parsons ES concurs. The requested change will be made.

Item 12 – Page 6-3, Bullet 4

For clarity, WPI recommends replacing "estimating benzene weathering from LNAPL" with "estimating benzene weathering from JP-4 LNAPL."

Parsons ES Response:

Parsons ES concurs. The requested change will be made.

RESPONSES TO MR. ATKINSON'S COMMENTS:

Item 1 – Page 1-3, Sec 1.1, Bullet 4

In line 3, suggest replacing "geology" with "soils."

Parsons ES Response:

Parsons ES concurs. The requested change will be made.

Item 2 – Page 3-2, Sec 3.1, Para 1, Sent 4

Are JP-5 and JP-8 reversed? In other words, should JP-5 be linked to 0.40 wt% and JP-8 associated with 0.05 wt%?

Parsons ES Response:

JP-5 and JP-8 are not reversed. The associated weight percentage for both JP-5 and JP-8 were obtained from Figure 2.3.

Item 3 – Page 3-3, Sec 3.1, Para 1, Sent 4

To correct terminology, "groundwater tables" should be changed to "water tables."

Parsons ES Response:

Parsons ES concurs. The requested change will be made.

Item 4 – Page 4-4, Sec 4.2, Line 1

Suggest inserting "to" after "order."

Parsons ES Response:

Parsons ES concurs. The requested change will be made.

Item 5 – Page 5-8, Sec 5.2.2

In the definition of "e," suggest rounding this value to 2.72 to be consistent with the use of "approximately."

Parsons ES Response:

Parsons ES concurs. The requested change will be made.

Item 6 – Page 5-25, Sec 5.2.4.1, Para 1, Line 6

Recommend deleting "the" following "that."

Parsons ES Response:

Parsons ES concurs. The requested change will be made.

Item 7 – Page 5-27, Sec 5.2.5.1, Line 3

Suggest inserting "the" in front of "mobile."

Parsons ES Response:

Parsons ES concurs. The requested change will be made.

Item 8 – Page 5-43, Sec 5.4.2, Para 2, Sent 2

My inspection of Figure 5.17 reveals a loss of "lighter" BTEX compounds but retention of "heavier" hydrocarbons. If this is so, suggest revising this sentence as follows: "...little residual LNAPL BTEX compounds remain in soils..."

Parsons ES Response:

Parsons ES concurs. The requested change will be made.