

Quarterly Groundwater Monitoring Report

Red Hill Bulk Fuel Storage Facility, Pearl Harbor, O'ahu, Hawai'i

Latitude: 21°22'15"N

Longitude: 157°53'33"W

DOH Facility ID No. 9-102271

DOH Release ID No. 99051, 010011, 020028

March 2011

Department of the Navy
Naval Facilities Engineering Command, Hawai'i
400 Marshall Road
Pearl Harbor, HI 96860-3139



Environmental Technical Services
Contract Number N62742-08-D-1930, Contract Task Order HC14

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Prepared for:



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Naval Facilities Engineering Command, Hawai'i
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List of Acronyms

| | |
|-----------|---|
| < | less than |
| µg/L | micrograms per liter |
| COPC | chemical of potential concern |
| DL | detection limit or method detection limit (MDL) |
| DLNR | Department of Land and Natural Resources |
| DOH | State of Hawai'i Department of Health |
| DON | Department of the Navy |
| DRO | diesel range organics |
| DTW (TOC) | depth to water from top of well casing |
| EAL | Environmental Action Level |
| EPA | U.S. Environmental Protection Agency |
| FISC | Fleet and Industrial Supply Center |
| GRO | gasoline range organics |
| HAR | Hawai'i Administrative Rules |
| IDW | investigation-derived waste |
| J | Estimated result. Indicates that the compound was identified but the concentration was above the DL and below the LOQ |
| JP | Jet Propellant |
| LCS | laboratory control sample |
| LNAPL | light non-aqueous phase liquid |
| LOD | limit of detection |
| LOQ | limit of quantitation or reporting limit (RL) |
| MS | matrix spike |
| MSD | matrix spike duplicate |
| N/A | not applicable |
| ND | not detected |
| NAVFAC | Naval Facilities Engineering Command |
| NP | measurement not provided |
| NT | measurement not taken |
| PAHs | polynuclear aromatic hydrocarbons |
| pH | hydrogen activity |
| PHWS | Pearl Harbor Water System |
| PPE | personal protective equipment |
| Q | data qualifier |
| QC | quality control |
| RHSF | Red Hill Bulk Fuel Storage Facility |
| RPD | relative percent difference |
| SSRBL | site-specific risk-based level |
| SWL | static water level |
| TEC | The Environmental Company, Inc. |
| TIC | tentatively identified compound |
| TPH | total petroleum hydrocarbons |
| U | Indicates the compound or analyte was analyzed for but not detected. The result is reported as ND. |
| U.S. | United States |

| | |
|------|---------------------------------|
| USGS | United States Geological Survey |
| UST | underground storage tank |
| VOC | volatile organic compound |

Executive Summary

This report documents the results of the quarterly groundwater monitoring performed in January 2011 at the Red Hill Bulk Fuel Storage Facility (RHSF). There are 18 active and two inactive, 12.5 million gallon capacity, field-constructed underground storage tanks (USTs) located at the RHSF. Previous environmental site investigations indicated a release had occurred and contaminated the groundwater underlying the RHSF.

The United States (U.S.) Navy implemented a groundwater monitoring program, which includes collecting groundwater samples quarterly from U.S. Navy Well 2254-01 (RHMW2254-01) and four groundwater monitoring wells (RHMW01, RHMW02, RHMW03, and RHMW05) installed within the RHSF lower access tunnel. The U.S. Navy Well 2254-01 is located approximately 3,000 feet downgradient from the RHSF and provides approximately 24 percent of the potable water to the Pearl Harbor Water System (PHWS). The groundwater samples are analyzed for petroleum constituents and compared against State of Hawai'i Department of Health (DOH) Drinking Water Environmental Action Levels (EALs) (DOH, 2009).

This report presents the analytical results and compares them to the DOH Drinking Water EALs for samples collected on January 18, 19, and 20, 2011 at the five groundwater monitoring wells (RHMW2254-01, RHMW01, RHMW02, RHMW03, and RHMW05). Contaminant trends that have exceeded DOH Drinking Water EALs are also provided in this report.

January 2011 Groundwater Monitoring Results

Laboratory analytical results indicated that total petroleum hydrocarbons (TPH)-diesel range organics (DRO) and polynuclear aromatic hydrocarbons (PAHs), specifically 1-methylnaphthalene and naphthalene, were present in the groundwater beneath the RHSF at concentrations that exceeded the EALs. All other chemical of potential concern (COPC) concentrations (i.e., volatile organic compounds (VOCs) and dissolved lead) were below the EALs.

In monitoring wells RHMW2254-01, RHMW01, RHMW03, and RHMW05, all COPCs were not detected at or above the limits of detection (LODs) and limits of quantitation (LOQs).

In monitoring well RHMW02, TPH-DRO (1,100 micrograms per liter [$\mu\text{g/L}$]), 1-methylnaphthalene (19 $\mu\text{g/L}$), and naphthalene (57 $\mu\text{g/L}$) were detected at concentrations which exceeded the EALs. Acenaphthene (0.29 $\mu\text{g/L}$) and 2-methylnaphthalene (3.6 $\mu\text{g/L}$) were detected at concentrations below the EALs. TPH-gasoline range organics (GRO) (17 $\mu\text{g/L}$), acenaphthylene (0.13 $\mu\text{g/L}$), fluorene (0.15 $\mu\text{g/L}$), ethylbenzene (0.29 $\mu\text{g/L}$), and xylenes (0.48 $\mu\text{g/L}$) were detected at estimated concentrations below the EALs. All other COPCs in RHMW02 were not detected at or above the LODs and LOQs.

All laboratory detection limits (DLs), LODs, and LOQs were generally below the EALs. In the case where an EAL for a specific COPC is less than the LOQ, it is generally acceptable to consider the LOQ in place of the EAL (DOH, 2009).

TPH-DRO Contaminant Trends

From January 2008 to April 2010, TPH-DRO at RHMW01 fluctuated between the historical range established from September 2005 through September 2007 (Appendix A). In July 2010, TPH-DRO decreased to the lowest detected concentration observed to date (228 µg/L). In October-November 2010 and during January 2011, TPH-DRO was not detected at or above the LODs and LOQs.

At RHMW02, TPH-DRO concentrations were relatively stable prior to 2008, after which significant variations occurred. In October-November 2010 and during January 2011, TPH-DRO showed a decrease in concentrations. The latest observed TPH-DRO concentration is below the historical range for this groundwater monitoring well.

From May 2009 through July 2010, TPH-DRO concentrations at RHMW03 were not detected above the LODs and LOQs. However, in October-November 2010, TPH-DRO was detected at the highest concentration to date (330 µg/L) which was above the EAL. TPH-DRO was not detected at or above the LOD and LOQ in January 2011.

At RHMW05, TPH-DRO concentrations increased from when it was first sampled in May 2009 to January 2010. However, in April 2010, July 2010, October-November 2010, and January 2011 TPH-DRO at RHMW05 was not detected above the LODs and LOQs.

TPH-GRO Contaminant Trends

TPH-GRO has remained below the EAL or has not been detected in monitoring wells RHMW01, RHMW03, and RHMW05. TPH-GRO concentrations at RHMW02 have fluctuated above and below the EAL, and in January 2011, TPH-GRO was detected at an estimated concentration of 17 µg/L, which was below the EAL.

PAHs Contaminant Trend in RHMW02

Since October 2008, the concentration of 2-methylnaphthalene has remained below the EAL. Naphthalene concentrations decreased to below the EAL in May 2009 and July 2009, increased above the EAL in July 2009, and remained above the EAL during this January 2011 groundwater monitoring event. Similarly, 1-methylnaphthalene concentrations decreased below the EAL in October 2009, however it has exhibited an increasing trend above the EAL during subsequent groundwater monitoring events, including this January 2011 groundwater monitoring event.

Conclusions

To date, the presence of light non-aqueous phase liquid (LNAPL) has been observed only once (i.e., in January 2008 in RHMW01 and RHMW02 with a thickness of less than 0.01 feet). This indicates a significant release from one or more of the USTs at the RHSF has not occurred at this time.

COPC concentrations in samples collected from the U.S. Navy Well 2254-01 have not been detected at or above the LODs and LOQs or are below the EALs. This indicates that elevated COPC concentrations detected in samples collected from monitoring wells within the RHSF are not migrating and impacting the Navy's potable water source.

COPC concentrations detected in samples collected from monitoring well RHMW05 have decreased to below the EALs for at least four consecutive monitoring events. The data suggest that elevated COPC concentrations detected in samples collected from monitoring wells located adjacent to the USTs are not migrating in the downgradient direction. RHMW05 is an intermediate monitoring point between the USTs and the U.S. Navy Well 22554-01. At this time, there is no indication that COPCs are migrating in a downgradient direction towards the Navy's potable water source.

COPC concentrations detected in samples collected from two monitoring wells adjacent to the USTs (i.e., RHMW01 and RHMW03) are not steadily increasing between consecutive sampling events. COPC concentrations of 1-methylnaphthalene detected in samples collected from RHMW02 however have increased for three consecutive events.

Recommendations

Based on the results of this monitoring event, continuation of the quarterly groundwater monitoring program at the RHSF is recommended. In addition continuation of monthly free product measurements at RHMW01, RHMW02, RHMW03, and RHMW05; and monthly soil vapor monitoring are also recommended. In response to the Category 2 status of RHMW02, submission of this quarterly report to DOH and continuation of a leak determination program as described in Section 3 of the RHSF Groundwater Protection Plan (TEC, 2008) to identify if tanks are leaking are recommended.

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Section 1 Introduction

This report presents the results of the 22nd groundwater monitoring event, conducted in January 2011 at the RHSF, Pearl Harbor, O'ahu, Hawai'i (Figure 1-1). The RHSF consists of 18 active and two inactive USTs operated by the Fleet and Industrial Supply Center (FISC), Pearl Harbor. The groundwater sampling and analysis event is part of a groundwater monitoring program for the UST site in response to past UST releases, previous environmental investigations, and recommendations from the DOH. The groundwater monitoring was performed by Environet for the Department of the Navy (DON), Naval Facilities Engineering Command (NAVFAC), Hawai'i under Environmental Technical Services, Contract Number N62742-08-D-1930, Contract Task Order HC14.

The field activities performed for the January 2011 quarterly groundwater monitoring event were conducted in accordance with the procedures described in the *Work Plan, Long-Term Monitoring, Red Hill Bulk Fuel Storage Facility, Pearl Harbor, O'ahu, Hawai'i* (Environet, 2010).

1.1 Project Objective

The groundwater monitoring was performed to evaluate the presence of COPCs in groundwater underlying the RHSF. The groundwater monitoring was conducted to ensure the DON remains in compliance with DOH UST release response requirements as described in Hawai'i Administrative Rules (HAR) 11-281 Subchapter 7, Release Response Action. Analytical results are compared to the DOH Drinking Water EALs for samples collected from five groundwater monitoring wells (RHMW2254-01, RHMW01, RHMW02, RHMW03, and RHMW05).

1.2 Background

The U.S. Navy implemented a groundwater monitoring program, which includes collecting groundwater samples quarterly from U.S. Navy Well 2254-01 (RHMW2254-01) and four groundwater monitoring wells (RHMW01, RHMW02, RHMW03, and RHMW05) installed within the RHSF lower access tunnel. The U.S. Navy Well 2254-01 is located approximately 3,000 feet downgradient from the RHSF and provides approximately 24 percent of the potable water to the PHWS. The groundwater samples are analyzed for petroleum constituents and compared against DOH Drinking Water EALs (DOH, 2009).

1.2.1 Site Description

The RHSF is located on the island of O'ahu, Hawai'i, approximately 2.5 miles northeast of Pearl Harbor in Hālawā Heights (Figure 1-1). Land adjacent to the north of the RHSF is occupied by the Hālawā Correctional Facility and private businesses. Land to the south and west of the facility includes the Coast Guard Reservation and other residential neighborhoods. Moanalua Valley is located east of the facility (Environet, 2010).

The Navy Public Works Department operates a potable water infiltration tunnel approximately 1,550 feet hydraulically downgradient from the RHSF (Environet, 2010). The U.S. Navy Well

2254-01 is located approximately 3,000 feet downgradient (west) of the RHSF and provides approximately 24 percent of the potable water to the PHWS, which serves approximately 52,200 military consumers (The Environmental Company, Inc. (TEC), 2008).

1.2.2 Facility Information

The RHSF consists of 18 active and two inactive USTs operated by Navy FISC Pearl Harbor. Each UST has a capacity of 12.5 million gallons. The RHSF is located approximately 100 feet above the basal aquifer (Dawson Group, Inc., 2006).

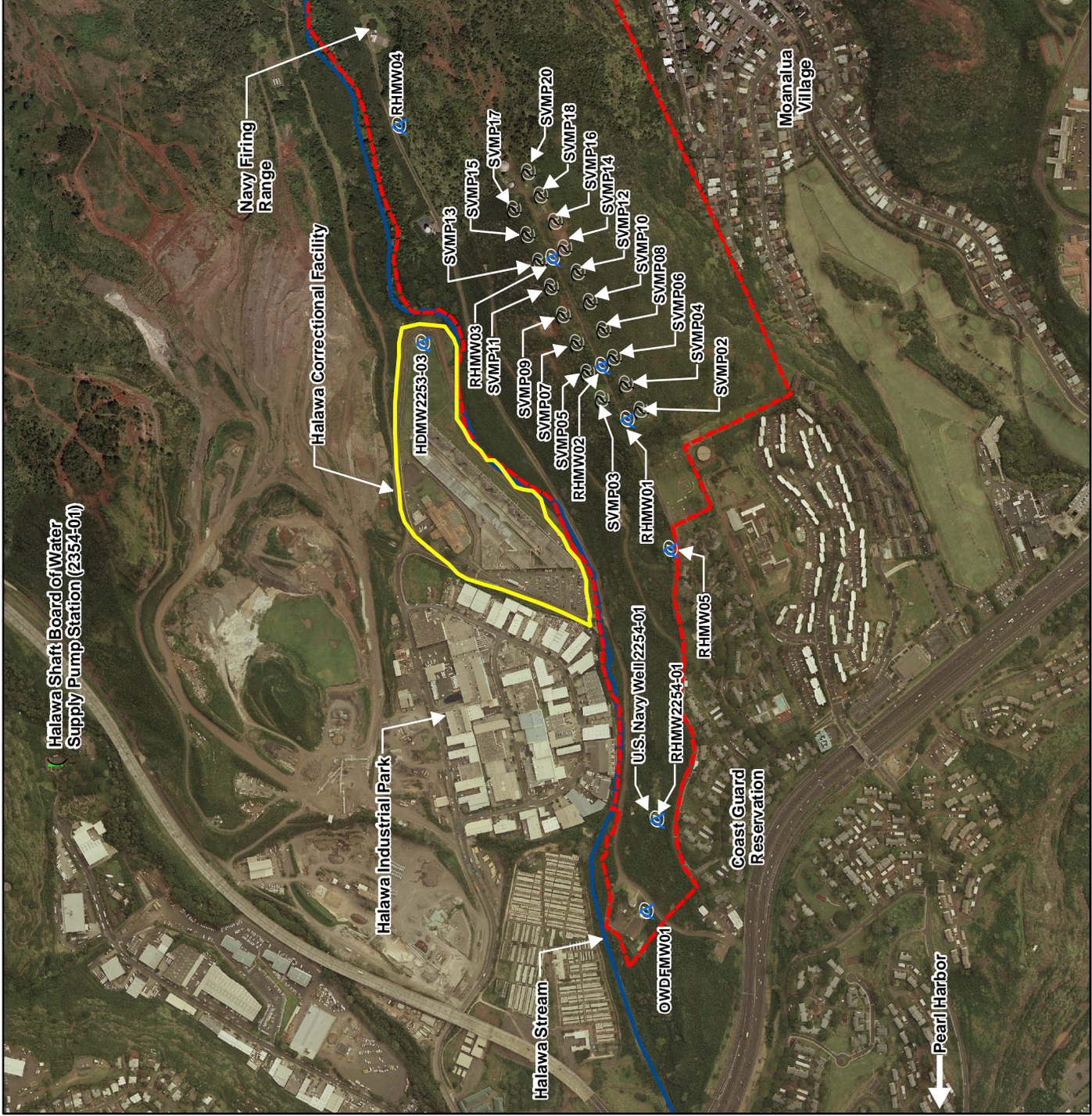
1.2.3 UST Information

The USTs were constructed in the early 1940s. The tanks were constructed of steel and currently contain Jet Propellant (JP)-5 fuel, JP-8, and marine diesel fuel (F-76). Previously, several tanks stored Navy Special Fuel Oil, Navy Distillate, aviation gasoline, and motor gasoline. Each tank measures approximately 245 feet in height and 100 feet in diameter. The upper domes of the tanks lie at depths varying between approximately 100 feet and 200 feet below the existing ground surface (TEC, 2006).

1.2.4 Previous Reports

The following groundwater monitoring reports were previously submitted to the DOH:

1. Groundwater Sampling Report, First Quarter 2005 (submitted April 2005);
2. Groundwater Sampling Report, Second Quarter 2005 (submitted August 2005);
3. Groundwater Sampling Report, Third Quarter 2005 (submitted November 2005);
4. Groundwater Sampling Report, Fourth Quarter 2005 (submitted February 2006);
5. Groundwater Monitoring Results, July 2006 (submitted September 2006);
6. Groundwater Monitoring Results, December 2006 (submitted January 2007);
7. Groundwater Monitoring Results, March 2007 (submitted May 2007);
8. Groundwater Monitoring Results, June 2007 (submitted August 2007);
9. Groundwater Monitoring Results, September 2007 (submitted October 2007);
10. Groundwater Monitoring Results, January 2008 (submitted March 2008);
11. Groundwater Monitoring Results, April 2008 (submitted May 2008);
12. Groundwater Monitoring Results, July 2008 (submitted October 2008);
13. Groundwater Monitoring Results, October and December 2008 (submitted February 2009);
14. Groundwater Monitoring Results, February 2009 (submitted May 2009);



| LEGEND | |
|--------------|------------------------------------|
| ? | Soil Vapor Monitoring Points |
| A | Groundwater Monitoring Wells |
| (| Board of Water Supply Pump Station |
| [Yellow Box] | Halawa Correctional Facility |
| [Red Box] | Red Hill Naval Reservation |
| [Blue Line] | Halawa Stream |

| SOURCE | |
|------------|--|
| USGS, 2005 | |



Figure 1-1
Groundwater Monitoring Well
Location Map
 Quarterly Groundwater
 Monitoring Report,
 Red Hill Bulk Fuel Storage Facility,
 Pearl Harbor, O'ahu, Hawai'i

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15. Groundwater Monitoring Results, May 2009 (submitted July 2009);
16. Groundwater Monitoring Results, July 2009 (submitted September 2009);
17. Groundwater Monitoring Results, October 2009 (submitted December 2009);
18. Groundwater Monitoring Results, January, February, and March 2010 (submitted April 2010);
19. Groundwater Monitoring Results, April 2010 (submitted May 2010);
20. Groundwater Monitoring Results, July 2010 (submitted August 2010); and
21. Groundwater Monitoring Results, October 2011 (submitted December 2010).

1.2.5 Previous Environmental Investigations

1998 to 2001 - From 1998 to 2001, the Navy conducted an investigation at the RHSF to assess potential releases from the fuel storage USTs and piping systems. In February 2001, the Navy installed a one-inch diameter monitoring well RHMW01 (previously known as MW-V1D) to monitor for contamination of the basal aquifer underlying the RHSF. The well was installed and completed at approximately 100 feet below grade within the lower access tunnel. At the time of well completion, depth to water in RHMW01 was measured at 86 feet below grade (Dawson group, Inc., 2006).

In February 2001, groundwater samples collected from RHMW01 contained TPH concentrations ranging from 883 µg/L to 1,050 µg/L and total lead ranging from 10.4 µg/L to 15 µg/L. The total lead concentrations exceeded the DOH Tier 1 Action Level of 5.6 µg/L (Dawson Group, Inc., 2006).

2005 Groundwater Sampling - The Navy began quarterly groundwater sampling at existing monitoring wells in 2005. Dawson Group, Inc. collected groundwater samples from the RHMW01 and RHMW2254-01 in February 2005, June 2005, September 2005, and December 2005.

Samples collected in February 2005 and June 2005 were not filtered in the field prior to analysis for lead. Analytical results for samples collected from RHMW01 indicated concentrations of total lead were above the DOH Tier 1 Action Level of 5.6 µg/L. The results were not considered appropriate for risk assessment since the sample had not been filtered. In addition, lead was not a component of fuels from the tanks near RHMW01, but was a component in fuels stored in other tanks during the history of the RHSF. Lead may have been part of the RHSF construction material (TEC, 2007). Previous sampling efforts showed elevated lead concentrations when analyzed as unfiltered samples. Subsequent efforts where the lead samples were filtered had resolved this issue. Samples were filtered in September 2005 and December 2005, and dissolved lead concentrations were below the DOH Tier 1 Action Level. Concentrations of all other COPCs were below the DOH Tier 1 Action Levels.

2005 Site Investigation - As part of a site investigation, TEC installed three groundwater monitoring wells at the RHSF between June 2005 and September 2005. Monitoring well RHMW02 was installed in the lower access tunnel near Tank 05 and Tank 06. Monitoring well RHMW03 was installed in the lower access tunnel near Tank 13 and Tank 14. Monitoring well

RHMW04 was installed north of the UST Tank 20 to evaluate groundwater within the basal aquifer upgradient from the RHSF. Monitoring wells RHMW02 and RHMW03 were completed to depths of approximately 125 feet below the tunnel floor, and well RHMW04 was completed to a depth of approximately 300 feet below ground surface outside the tunnel. Groundwater samples were collected from the three newly installed wells and two existing wells (RHMW01 and RHMW2254-01) in September 2005 (TEC, 2010).

Naphthalene and trichloroethylene were detected in samples collected from RHMW02 at concentrations greater than the DOH Tier 1 Action Levels. Lead was detected in the sample collected from RHMW01 at a concentration greater than the DOH Tier 1 Action Level; however, the sample was not filtered in the field prior to analysis. Analytical results for filtered samples obtained by Dawson Group, Inc. during the same period indicated concentrations of dissolved lead were below the DOH Tier 1 Action Level (TEC, 2010).

2006 Site Investigation - Dedicated sampling pumps were installed in five monitoring wells (RHMW01, RHMW02, RHMW03, RHMW04, and RHMW2254-01). TEC collected groundwater samples from the monitoring wells in July 2006. The groundwater samples were analyzed for petroleum constituents. Naphthalene was detected in samples collected from RHMW02 at concentrations above the DOH Tier 1 Action Level (TEC, 2010).

In September 2006, with concurrence from the DOH, the Navy decided to use the newer DOH EALs for the Red Hill Site Investigation and Risk Assessment project. The EALs provide action levels for more chemicals, and are much more useful for conducting screening risk assessments. Since the DOH (DOH May 2005) Policy Letter stated that the two sets of action levels should not be mixed, the Tier 1 screening levels presented in HAR Section 11-281-78 would no longer be used to evaluate environmental impact at the RHSF (TEC, 2010).

2006 Groundwater Sampling - Groundwater samples were collected in December 2006. Analytical results indicated the following:

- no COPCs were detected in samples collected from RHMW2254-01 or RHMW03;
- TPH-DRO concentrations were detected in samples collected from RHMW01 at concentrations above the EAL; and
- TPH-GRO, TPH-DRO, and naphthalene were detected in samples collected from RHMW02 at concentrations above the EALs (TEC, 2010).

2007 Groundwater Sampling - Groundwater samples were collected in March, June, and September 2007. Analytical results indicated the following:

- no COPCs were detected above the EALs at RHMW2254-01;
- TPH-DRO concentrations exceeded the EAL in samples collected from RHMW01 during all three monitoring events;
- TPH-GRO concentrations exceeded the EAL in samples collected from RHMW02 in March;

- TPH-DRO and naphthalene concentrations exceeded the EALs in samples collected from RHMW02 during all three monitoring events;
- 1-methylnaphthalene and 2-methylnaphthalene concentrations exceeded DOH Groundwater Gross Contamination EAL in samples collected from RHMW02 during all three monitoring events; and
- TPH-DRO concentrations exceeded the EAL in the sample collected from RHMW03 in June.

2008 Groundwater Sampling - Groundwater samples were collected in January, April, July, and October 2008. Analytical results indicated the following:

- no COPCs were detected above the EALs at RHMW2254-01;
- trace detections of 1-methylnaphthalene and naphthalene prompted a resample event in December at RHMW2254-01, no chemicals were detected above the LODs;
- TPH-DRO concentrations exceeded the EAL in samples collected from RHMW01 during all four monitoring events;
- TPH-GRO concentrations did not exceed the EAL in samples collected from RHMW02;
- TPH-DRO, naphthalene, 1-methylnaphthalene, and 2-methylnaphthalene concentrations exceeded the EALs in samples collected from RHMW02. Additionally, the site-specific risk-based level (SSRBL) of 4,500 µg/L for TPH-DRO was exceeded in the October monitoring event at RHMW02 (Appendix A); and
- TPH-DRO concentrations exceeded the EAL in samples collected from RHMW03 during all four monitoring events.

2009 Groundwater Sampling and RHMW05 Installation - Groundwater samples were collected in February, May, July, and October 2009. In April 2009, a new groundwater monitoring well, RHMW05, was installed by TEC. RHMW05 is located within the lower access tunnel between RHMW01 and RHMW2254-01 (located at the U.S. Navy Well 2254-01). It was installed to identify any contamination migrating past RHMW01 prior to it reaching the U.S. Navy Well 2254-01 (TEC, 2010). Analytical results indicated the following:

- no COPCs have been detected above the EALs at RHMW2254-01;
- TPH-GRO concentrations in samples collected from RHMW2254-01 were detected above the LOD and significantly below the LOQ and EAL in February and May 2009;
- TPH-DRO concentrations exceeded the EAL in samples collected from RHMW01 during all four monitoring events;
- TPH-GRO concentrations did not exceed the EAL in samples collected from RHMW02;
- TPH-DRO concentrations exceeded the EAL in samples collected from RHMW02 during all four monitoring events;
- naphthalene, 1-methylnaphthalene, and 2-methylnaphthalene concentrations exceeded the EALs in samples collected from RHMW02 in February 2009, however only the 1-

methylnaphthalene concentration exceeded the EAL in May 2009 and July 2009 and only the naphthalene concentration exceeded the EAL in October 2009;

- TPH-DRO concentrations exceeded the EAL in samples collected from RHMW03 in February, but not in May or July; and
- TPH-DRO concentration exceeded the EAL in the sample collected from RHMW05 during the May and July 2009 monitoring events.

2010 Groundwater Sampling – Groundwater samples were collected in January, April, July, and October-November 2010 (and TPH-DRO was re-sampled at RHMW02 in February 2010 and March 2010). Analytical results indicated the following:

- no chemicals have been detected above the EALs in samples collected from RHMW2254-01;
- naphthalene concentrations in the samples collected from RHMW2254-01 were below the LOQ and EAL in January and October-November;
- TPH-DRO concentrations exceeded the EAL in samples collected from RHMW01, until October-November when they were not detected at or above the LOQ;
- TPH-DRO concentrations exceeded the EAL in samples collected from RHMW02 in January, February, March, April, July, and October-November however, significant increases in January and February were attributed to tentatively identified compounds (TICs) apparently not associated with petroleum from the RHSF (TEC, 2010);
- naphthalene and 1-methylnaphthalene concentrations exceeded the EALs in samples collected from RHMW02 in January, April, July, and October-November;
- TPH-DRO concentrations exceeded the EAL in samples collected from RHMW05 in January, however, the significant increase was attributed to TICs apparently not associated with petroleum from the RHSF. TPH-DRO concentrations were not detected at or above the LOD in October-November;
- TPH-DRO concentrations in samples collected from RHMW03 were not detected above the LOD in January, April, and July. However, in October-November TPH-DRO was detected above the EAL;
- TPH-GRO concentrations exceeded the EAL in samples collected from RHMW02 in October-November; and
- Lead was detected in samples collected from RHMW2254-01 and RHMW01 at concentrations which were below the EALs in October-November 2010.

1.2.6 Regulatory Updates

During the summer and fall of 2008 DOH updated their EALs which resulted in significant changes to the action levels associated with methylnaphthalenes. The drinking water toxicity EAL for these compounds was 240 µg/L. This concentration presumed that methylnaphthalenes were non-carcinogenic. Evidence that they are human carcinogens has now been accepted by the

U.S. Environmental Protection Agency (EPA). As a result, DOH adopted more rigorous EALs of 4.7 µg/L for 1-methylnaphthalene and 24 µg/L for 2-methylnaphthalene, corresponding to a residential tap water scenario, and a one in a million cancer risk (DOH, 2009). As a result of the EAL changing from 240 µg/L to 4.7 µg/L, concentrations of 1-methylnaphthalene collected from RMHW2254-01, RHMW02, RHMW03, and RHMW05 previously categorized as under the EAL are now over the EAL.

Also, the drinking water EAL for naphthalene was increased from 6.2 µg/L to 17 µg/L (DOH, 2009). Finally, the DOH Drinking Water EAL for TPH-DRO was increased from 100 µg/L to 210 µg/L, although the DOH Groundwater Gross Contamination EAL for TPH-DRO remains 100 µg/L.

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Section 2 Groundwater Monitoring Activities

Groundwater monitoring was conducted at the five monitoring wells (RHMW2254-01, RHMW01, RHMW02, RHMW03, and RHMW05) on January 18, 19, and 20, 2011 using procedures described in the Work Plan (Environet, 2010). Field activities were documented in the field notebook (Appendix B).

2.1 Oil/Water Interface Measurements

The presence and thickness of LNAPL, otherwise known as “free product”, released from the USTs is monitored at the RHSF. Groundwater gauging measurements were collected at the five monitoring wells prior to purging and sample collection. A Solinst oil/water interface probe was used to measure the depth to groundwater, as well as detect the presence and thickness of LNAPL to the nearest 0.01 foot, according to the procedures described in Procedure I-C-3, *Monitoring Well Sampling* (DON, 2007).

2.2 Groundwater Sampling

Groundwater samples were collected from the five monitoring wells according to the procedures described in Procedure I-C-3, *Monitoring Well Sampling* (DON, 2007).

2.2.1 Monitoring Well Purging

Each monitoring well was purged using a dedicated bladder pump system. Groundwater was purged at a low flow rate (less than one liter per minute) using the dedicated bladder pump system until three or more successive water quality parameter measurements had stabilized within 10 percent. A Horiba[®] U-22 multi-parameter water quality meter was used to measure hydrogen activity (pH), temperature, specific conductivity, dissolved oxygen, turbidity, and oxidation reduction potential. At least four to six readings were recorded on Groundwater Sampling Log data sheets (Appendix C).

2.2.2 Groundwater Sample Collection

Groundwater samples were collected using dedicated bladder pump systems. Groundwater samples were collected directly into laboratory provided specially cleaned sample containers already containing the appropriate preservatives (i.e., nitric acid for dissolved lead analysis). The dissolved lead samples were filtered in the field, and placed in polyethylene bottles containing preservatives.

2.2.3 Sample Management and Shipment

Each sample container sent to the laboratory was assigned a project-specific chain of custody identification number and a descriptive identification number. The sample identifiers provided specific data unique to each sample and were entered into the field notebook. The samples were labeled according to the procedures described in Procedure III-E, *Record Keeping, Sample Labeling, and Chain-of-Custody* (DON, 2007).

Following sample collection and labeling, the sample containers were bubble-wrapped and placed into individual ZipLoc[®] bags, then immediately into insulated coolers with ice for preservation. The samples were shipped via FedEx to the laboratory on the same day or the following day of collection, with the exception of one sample which was shipped two days following sample collection. The samples were managed under standard chain of custody protocol and documentation from collection to delivery to the laboratory. Sample handling, storage, and transport were performed according to the requirements described in Procedure III-F, *Sample Handling, Storage, and Shipping* (DON, 2007).

2.3 Analytical Program

Five primary groundwater samples, one duplicate groundwater sample, and one quality control (QC) groundwater sample (i.e., matrix spike (MS)/matrix spike duplicate (MSD)) were submitted to APPL, Inc. located in Clovis, California. Groundwater samples were analyzed for VOCs and TPH-GRO by EPA Method 8260B, TPH-DRO by EPA Method 8015B, PAHs by EPA Method 8270D SIM, and dissolved lead by EPA Method 6020. The results of the laboratory analyses are presented and discussed in Section 3.

2.4 Field Quality Control Samples

Field QC procedures were followed to ensure viability and integrity of sample analytical data. Field duplicates were collected according to the procedures described in Procedure III-B, *Field QC Samples (Water, Soil)* (DON, 2007) and the Work Plan (Environet, 2010). Field duplicate samples were collected at a minimum of 10 percent of primary samples and analyzed for the same contaminants. Field rinsate samples were not required since dedicated bladder pump systems were used.

2.5 Laboratory Quality Control Samples

Laboratory QC samples were analyzed as part of the standard laboratory QC protocols as presented in the Work Plan (Environet, 2010). Laboratory QC for the monitoring event consisted of MS/MSD samples for organic analysis. Laboratory QC samples were collected and analyzed according to the procedures described in Procedure III-A *Laboratory QC Samples (Water, Soil)* (DON, 2007). Laboratory QC MS/MSD samples are an aliquot (i.e., a subset) of the field sample. They are not separate samples, but a special designation of an existing sample. Laboratory QC MS/MSD samples were analyzed for the same constituents as the standard samples. At a minimum, one MS/MSD sample pair was required per 20 samples, including duplicates.

2.6 Equipment Decontamination

Decontamination of monitoring equipment was performed to ensure data quality, to prevent cross contamination, and to prevent the potential introduction of contaminants into previously un-impacted areas. Decontamination of monitoring equipment (i.e., Solinst oil/water interface probe and Horiba[®] U-22 water quality meter) was conducted between monitoring locations according to the procedures described in Procedure I-F, *Equipment Decontamination* (DON,

2007). Decontamination water was disposed of in the RHSF's lower tunnel oil/water separator sump.

2.7 Investigation-Derived Waste Management

Investigation-derived waste (IDW) was managed in accordance with the procedures described in Procedure I-A-6, *IDW Management* (DON, 2007). The various potential waste streams included the following:

- personal protective equipment (PPE) including: nitrile gloves, etc.;
- liquids including: equipment rinse water and purged groundwater; and
- disposable sampling equipment and supplies, including: poly sheeting, etc.

Equipment rinse water and purge water was disposed of in the RHSF's lower tunnel oil/water separator sump.

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Section 3 Groundwater Monitoring Results

This section provides a summary of analytical results for groundwater samples collected from the five monitoring wells on January 18, 19, and 20, 2011. Complete analytical laboratory reports are provided in Appendix D.

3.1 Results of Oil/Water Interface Measurements

Free product was not observed at RHMW01, RHMW02, RHMW03, and RHMW05 during the January 2011 sampling event. The trend of free product measurements over time show that in January 2008, LNALP was measured in monitoring wells RHMW01 and RHMW02 at a thickness of less than 0.01 feet, but was not observed in other monitoring wells. Since the trace amounts observed in January 2008, no free product has been observed in any of these RHF monitoring wells (Table 3-1).

3.2 Summary of Groundwater Analytical Results

All DLs, LODs, and LOQs were generally below the EALs. In the case where an EAL for a specific COPC is less than the LOQ, it is generally acceptable to consider the LOQ in place of the EAL (DOH, 2009).

RHMW2254-01

All COPC concentrations were not detected at or above the LODs and LOQs (Table 3-2 and Appendix D).

RHMW01

All COPC concentrations were not detected at or above the LODs and LOQs (Table 3-2 and Appendix D).

RHMW02

TPH-DRO (1,100 µg/L), 1-methylnaphthalene (19 µg/L), and naphthalene (57 µg/L) were detected at concentrations which exceeded both the DOH Drinking Water EALs (210 µg/L for TPH-DRO; 4.7 µg/L for 1-methylnaphthalene; 17 µg/L for naphthalene) and the DOH Groundwater Gross Contamination EALs (100 µg/L for TPH-DRO; 10 µg/L for 1-methylnaphthalene; 21 µg/L for naphthalene) (Table 3-2 and Appendix D).

Acenaphthene (0.29 µg/L) and 2-methylnaphthalene (3.6 µg/L) were detected at concentrations which were below both the DOH Drinking Water EALs (370 µg/L for acenaphthene; 24 µg/L for 2-methylnaphthalene) and the DOH Groundwater Gross Contamination EALs (20 µg/L for acenaphthene; 10 µg/L for 2-methylnaphthalene) (Table 3-2 and Appendix D).

TPH-GRO (17 µg/L), acenaphthylene (0.13 µg/L), fluorene (0.15 µg/L), ethylbenzene (0.29 µg/L), and xylenes (0.48 µg/L), were detected at estimated concentrations which were below both the DOH Drinking Water EALs (100 µg/L for TPH-GRO, 240 µg/L for acenaphthylene;

240 µg/L for fluorene; 700 µg/L for ethylbenzene; and 10,000 µg/L for xylenes) and the DOH Groundwater Gross Contamination EALs (100 µg/L for TPH-GRO, 2,000 µg/L for acenaphthylene; 950 µg/L for fluorene; 30 µg/L for ethylbenzene; and 20 µg/L for xylenes). All other COPCs were not detected at or above the LODs and LOQs (Table 3-2 and Appendix D).

RHMW03

All COPC concentrations were not detected at or above the LODs and LOQs (Table 3-2 and Appendix D).

RHMW05

All COPC concentrations were not detected at or above the LODs and LOQs (Table 3-2 and Appendix D).

3.3 Groundwater Contaminant Trend

Groundwater samples have been collected and analyzed in monitoring wells RHMW01, RHMW02, RHMW03 since September 2005, and in monitoring well RHMW05 since May 2009 (Appendix A). Monitoring well RHMW2254-01 was installed in February 2005. The following is a discussion of compounds that exceeded DOH Drinking Water EALs during two or more recent consecutive sampling events of increasing or decreasing concentrations, thus establishing a trend:

RHMW2254-01

COPCs have never been detected at RHMW2254-01 at concentrations greater than the DOH Drinking Water EALs. In January 2011, all COPCs were not detected at or above the LODs and LOQs.

RHMW01

Concentrations of TPH-DRO have been greater than the DOH Drinking Water EAL between September 2005 and July 2010, but less than 25 percent of the SSRBL of 4,500 µg/L. TPH-DRO concentrations decreased from October 2008 through July 2009; increased in October 2009 (299F µg/L), January 2010 (312F µg/L), and April 2010 (377 µg/L); decreased in July 2010 (228F µg/L); and was not detected above the LODs and LOQs in October-November 2010 and January 2011. TPH-GRO has remained below the EAL or has not been detected (Appendix A).

RHMW02

The averages of primary and duplicate samples were used for determining contaminant trends. From September 2005 through February 2009, TPH-DRO exceeded the DOH Drinking Water EAL of 210 µg/L and was greater than 50 percent of the SSRBL (estimated solubility limit of 4,500 µg/L). The concentration of TPH-DRO was relatively stable until July 2008 and October 2008 when the concentrations increased, with the October 2008 average also exceeding the SSRBL of 4,500 µg/L (Appendix A).

Table 3-1: Red Hill Oil/Water Interface Measurements, September 2007 through January 2011

| Date | RHHW01 | | | RHHW02 | | | RHHW03 | | | RHHW05 | | |
|-----------------------|--------------------------|-----------------|-----------------|--------------------------|-------|-------|--------------------------|-----------------|-----------------|--------------------------|-----------------|-----------------|
| | Elevation = DTW (TOC) | SWL | LNAPL | Elevation = DTW (TOC) | SWL | LNAPL | Elevation = DTW (TOC) | SWL | LNAPL | Elevation = DTW (TOC) | SWL | LNAPL |
| Sep-2007 | NT ² | NT ² | NT ² | 86.80 | 17.96 | NP | 103.44 | 17.62 | NP | ~ | ~ | ~ |
| Jan-2008 | 84.67 | 17.60 | <0.01 | 86.23 | 18.53 | <0.01 | NT ³ | NT ³ | NT ³ | ~ | ~ | ~ |
| Jul-2008 | 83.37 | 18.90 | 0.00 | 86.10 | 18.66 | 0.00 | 102.45 | 18.61 | 0.00 | ~ | ~ | ~ |
| Oct-2008 | 83.80 | 18.47 | 0.00 | 86.45 | 18.31 | 0.00 | 102.49 | 18.57 | 0.00 | ~ | ~ | ~ |
| Nov-2008 | 83.91 | 18.36 | 0.00 | 86.56 | 18.20 | 0.00 | 102.80 | 18.26 | 0.00 | ~ | ~ | ~ |
| Jan-2009 | 83.13 | 19.14 | 0.00 | 85.79 | 18.97 | 0.00 | 102.04 | 19.02 | 0.00 | ~ | ~ | ~ |
| Feb-2009 | NT ⁴ | NT ⁴ | NT ⁴ | 86.35 | 18.41 | 0.00 | 102.56 | 18.50 | 0.00 | ~ | ~ | ~ |
| Mar-2009 | 83.82 | 18.45 | 0.00 | 86.44 | 18.32 | 0.00 | 102.64 | 18.42 | 0.00 | ~ | ~ | ~ |
| May-2009 ⁵ | 83.72 | 18.55 | 0.00 | 86.37 | 18.39 | 0.00 | 102.59 | 18.47 | 0.00 | NT ⁶ | NT ⁶ | NT ⁶ |
| May-2009 | 83.50 | 18.77 | 0.00 | 86.15 | 18.61 | 0.00 | 102.41 | 18.65 | 0.00 | NT ⁶ | NT ⁶ | NT ⁶ |
| Jul-2009 ⁷ | 83.75 | 18.52 | 0.00 | 86.42 | 18.34 | 0.00 | 102.67 | 18.39 | 0.00 | 83.09 | 18.46 | 0.00 |
| Aug-2009 | 84.04 | 18.23 | 0.00 | 86.71 | 18.05 | 0.00 | 102.84 | 18.22 | 0.00 | 83.51 | 18.04 | 0.00 |
| Sep-2009 | 84.21 | 18.06 | 0.00 | 86.84 | 17.92 | 0.00 | 103.07 | 17.99 | 0.00 | 83.61 | 17.94 | 0.00 |
| Oct-2009 | 84.24 | 18.03 | 0.00 | 86.87 | 17.89 | 0.00 | 103.07 | 17.99 | 0.00 | 83.62 | 17.93 | 0.00 |
| Nov-2009 | 83.91 | 18.36 | 0.00 | 86.56 | 18.20 | 0.00 | 102.81 | 18.25 | 0.00 | 83.25 | 18.30 | 0.00 |
| Dec-2009 | 84.12 | 18.15 | 0.00 | 86.75 | 18.01 | 0.00 | 103.00 | 18.06 | 0.00 | 83.53 | 18.02 | 0.00 |
| Jan-2010 | 84.36 | 17.91 | 0.00 | 87.00 | 17.76 | 0.00 | 103.22 | 17.84 | 0.00 | 83.75 | 17.80 | 0.00 |
| Feb-2010 | 84.24 | 18.03 | 0.00 | 86.89 | 17.87 | 0.00 | 103.14 | 17.92 | 0.00 | 83.60 | 17.95 | 0.00 |
| Mar-2010 | 84.53 | 17.74 | 0.00 | 87.15 | 17.61 | 0.00 | 103.38 | 17.68 | 0.00 | 83.96 | 17.59 | 0.00 |
| Apr-2010 | 84.75 | 17.52 | 0.00 | 87.37 | 17.39 | 0.00 | 103.60 | 17.46 | 0.00 | 84.17 | 17.38 | 0.00 |
| May-2010 | 84.80 | 17.47 | 0.00 | 87.43 | 17.33 | 0.00 | 103.66 | 17.40 | 0.00 | 84.23 | 17.32 | 0.00 |
| Jun-2010 | 84.87 | 17.40 | 0.00 | 87.51 | 17.25 | 0.00 | 103.74 | 17.32 | 0.00 | 84.30 | 17.25 | 0.00 |
| Jul-2010 | 85.03 | 17.24 | 0.00 | 87.66 | 17.10 | 0.00 | 103.89 | 17.17 | 0.00 | 84.48 | 17.07 | 0.00 |
| Sep-2010 | 85.30 | 16.97 | 0.00 | 87.92 | 16.84 | 0.00 | 104.13 | 16.93 | 0.00 | 84.71 | 16.84 | 0.00 |
| Oct-2010 | 85.29 | 16.98 | 0.00 | 87.91 | 16.85 | 0.00 | 104.13 | 16.93 | 0.00 | 84.75 | 16.80 | 0.00 |
| Nov-2010 | 85.20 | 17.07 | 0.00 | 87.84 | 16.92 | 0.00 | 104.30 | 16.76 | 0.00 | 84.60 | 16.95 | 0.00 |
| Dec-2010 | 84.87 | 17.40 | 0.00 | 87.55 | 17.21 | 0.00 | 103.98 | 17.08 | 0.00 | 84.22 | 17.33 | 0.00 |
| Jan-2011 | 85.32 | 16.95 | 0.00 | 86.91 | 17.85 | 0.00 | 103.41 | 17.65 | 0.00 | 83.65 | 17.90 | 0.00 |

Notes:

¹ Elevations were updated based on the Groundwater Flow Direction/Gradient and Tier 3 Risk Assessment Re-evaluation Letter Report, Red Hill Bulk Fuel Storage Facility, Pearl Harbor, HI, Contract #M47408-04-D-6514, Task Order 54, dated April 15, 2010.

² A measurement was not taken at RHHW01 in September 2007.

³ A measurement was not taken at RHHW03 in January 2008 due to equipment malfunction.

⁴ A measurement was not taken at RHHW01. The monitoring well was inaccessible due to extensive work being conducted at Tank 02.

⁵ The April 2009 measurements were pushed back a week (to 5/6/09) due to RHHW05 installation.

⁶ Measurements were not taken at RHHW05 until the installation of the dedicated oil/water interface probe was completed.

⁷ The June 2009 measurements were skipped due to the installation of dedicated oil/water interface probes.

Units are in feet (ft.).

Measurements recorded prior to September 2010 were collected by TEC. Measurements recorded in September 2010 and after were collected by Environet.

DTW (TOC) - depth to water from top of well casing

LNAPL - light non-aqueous phase liquid attributed to the Red Hill Bulk Fuel Storage Facility

NP - measurement not provided

NT - measurement not taken

SWL - static water level

~ - period prior to the installation of RHHW05

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Table 3-2: Analytical Results for Quarterly Groundwater Monitoring, January 2011

| Method | Chemical | DOH Drinking Water EALs for Human Toxicity ^a | DOH Groundwater Gross Contamination EALs ^a | RHMW2254-01 (ES014) | | | | | RHMW01 (ES015) | | | | | RHMW02 (ES010) | | | | | RHMW03 (ES012) | | | | | RHMW05 (ES013) | | | | |
|------------------------|---|---|---|---------------------|-----|------------------|------|------|----------------|-----|------------------|------|------|----------------|-----|------------------|------|------|----------------|-----|------------------|------|------|----------------|-----|------------------|------|------|
| | | | | Result | Q | LOQ | LOD | DL | Result | Q | LOQ | LOD | DL | Result | Q | LOQ | LOD | DL | Result | Q | LOQ | LOD | DL | Result | Q | LOQ | LOD | DL |
| EPA 8015B (Petroleum) | TPH-DRO | 210 | 100 | ND | U | 150 ^b | 80.8 | 40.4 | ND | U | 150 | 80.8 | 40.4 | 1,100 | | 150 ^b | 80.8 | 40.4 | ND | U | 150 | 80.8 | 40.4 | ND | U | 150 | 80.8 | 40.4 |
| EPA 8260 B (Petroleum) | TPH-GRO | 100 | 100 | ND | U | 20.0 | 12.1 | 6.06 | ND | U | 20.0 | 12.1 | 6.06 | 17 | J | 20.0 | 12.1 | 6.06 | ND | U | 20.0 | 12.1 | 6.06 | ND | U | 20.0 | 12.1 | 6.06 |
| EPA 8270D SIM (PAHs) | Acenaphthene | 370 | 20 | ND | U | 0.2 | 0.12 | 0.06 | ND | U | 0.2 | 0.12 | 0.06 | 0.29 | | 0.2 | 0.12 | 0.06 | ND | U | 0.2 | 0.12 | 0.06 | ND | U | 0.2 | 0.12 | 0.06 |
| | Acenaphthylene | 240 | 2,000 | ND | U | 0.2 | 0.12 | 0.06 | ND | U | 0.2 | 0.12 | 0.06 | 0.13 | J | 0.2 | 0.12 | 0.06 | ND | U | 0.2 | 0.12 | 0.06 | ND | U | 0.2 | 0.12 | 0.06 |
| | Anthracene | 1,800 | 22 | ND | U | 0.2 | 0.10 | 0.05 | ND | U | 0.2 | 0.10 | 0.05 | ND | U | 0.2 | 0.10 | 0.05 | ND | U | 0.2 | 0.10 | 0.05 | ND | U | 0.2 | 0.10 | 0.05 |
| | Benzo(a)anthracene | 0.092 | 4.7 | ND | U | 0.2 ^b | 0.14 | 0.07 | ND | U | 0.2 ^b | 0.14 | 0.07 | ND | U | 0.2 ^b | 0.14 | 0.07 | ND | U | 0.2 ^b | 0.14 | 0.07 | ND | U | 0.2 ^b | 0.14 | 0.07 |
| | Benzo(g,h,i)perylene | 1,500 | 0.13 | ND | U | 0.2 ^b | 0.16 | 0.08 | ND | U | 0.2 ^b | 0.16 | 0.08 | ND | U | 0.2 ^b | 0.16 | 0.08 | ND | U | 0.2 ^b | 0.16 | 0.08 | ND | U | 0.2 ^b | 0.16 | 0.08 |
| | Benzo(a)pyrene | 0.2 | 0.81 | ND | U | 0.2 | 0.14 | 0.07 | ND | U | 0.2 | 0.14 | 0.07 | ND | U | 0.2 | 0.14 | 0.07 | ND | U | 0.2 | 0.14 | 0.07 | ND | U | 0.2 | 0.14 | 0.07 |
| | Benzo(b)fluoranthene | 0.092 | 0.75 | ND | U | 0.2 ^b | 0.12 | 0.06 | ND | U | 0.2 ^b | 0.12 | 0.06 | ND | U | 0.2 ^b | 0.12 | 0.06 | ND | U | 0.2 ^b | 0.12 | 0.06 | ND | U | 0.2 ^b | 0.12 | 0.06 |
| | Benzo(k)fluoranthene | 0.92 | 0.4 | ND | U | 0.2 | 0.14 | 0.07 | ND | U | 0.2 | 0.14 | 0.07 | ND | U | 0.2 | 0.14 | 0.07 | ND | U | 0.2 | 0.14 | 0.07 | ND | U | 0.2 | 0.14 | 0.07 |
| | Chrysene | 9.2 | 1 | ND | U | 0.2 | 0.10 | 0.05 | ND | U | 0.2 | 0.10 | 0.05 | ND | U | 0.2 | 0.10 | 0.05 | ND | U | 0.2 | 0.10 | 0.05 | ND | U | 0.2 | 0.10 | 0.05 |
| | Dibenzo(a,h)anthracene | 0.0092 | 0.52 | ND | U | 0.2 ^b | 0.10 | 0.05 | ND | U | 0.2 ^b | 0.10 | 0.05 | ND | U | 0.2 ^b | 0.10 | 0.05 | ND | U | 0.2 ^b | 0.10 | 0.05 | ND | U | 0.2 ^b | 0.10 | 0.05 |
| | Fluoranthene | 1,500 | 130 | ND | U | 0.2 | 0.16 | 0.08 | ND | U | 0.2 | 0.16 | 0.08 | ND | U | 0.2 | 0.16 | 0.08 | ND | U | 0.2 | 0.16 | 0.08 | ND | U | 0.2 | 0.16 | 0.08 |
| | Fluorene | 240 | 950 | ND | U | 0.2 | 0.12 | 0.06 | ND | U | 0.2 | 0.12 | 0.06 | 0.15 | J | 0.2 | 0.12 | 0.06 | ND | U | 0.2 | 0.12 | 0.06 | ND | U | 0.2 | 0.12 | 0.06 |
| | Ideno(1,2,3-cd)pyrene | 0.092 | 0.095 | ND | U | 0.2 ^b | 0.14 | 0.07 | ND | U | 0.2 ^b | 0.14 | 0.07 | ND | U | 0.2 ^b | 0.14 | 0.07 | ND | U | 0.2 ^b | 0.14 | 0.07 | ND | U | 0.2 ^b | 0.14 | 0.07 |
| | 1-Methylnaphthalene | 4.7 | 10 | ND | U | 0.2 | 0.12 | 0.06 | ND | U | 0.2 | 0.12 | 0.06 | 19 | | 0.2 | 0.12 | 0.06 | ND | U | 0.2 | 0.12 | 0.06 | ND | U | 0.2 | 0.12 | 0.06 |
| | 2-Methylnaphthalene | 24 | 10 | ND | U | 0.2 | 0.12 | 0.06 | ND | U | 0.2 | 0.12 | 0.06 | 3.6 | | 0.2 | 0.12 | 0.06 | ND | U | 0.2 | 0.12 | 0.06 | ND | U | 0.2 | 0.12 | 0.06 |
| | Naphthalene | 17 | 21 | ND | U | 0.2 | 0.10 | 0.05 | ND | U | 0.2 | 0.10 | 0.05 | 57 | | 0.2 | 0.10 | 0.05 | ND | U | 0.2 | 0.10 | 0.05 | ND | U | 0.2 | 0.10 | 0.05 |
| | Phenanthrene | 240 | 410 | ND | U | 0.2 | 0.14 | 0.07 | ND | U | 0.2 | 0.14 | 0.07 | ND | U | 0.2 | 0.14 | 0.07 | ND | U | 0.2 | 0.14 | 0.07 | ND | U | 0.2 | 0.14 | 0.07 |
| Pyrene | 180 | 68 | ND | U | 0.2 | 0.16 | 0.08 | ND | U | 0.2 | 0.16 | 0.08 | ND | U | 0.2 | 0.16 | 0.08 | ND | U | 0.2 | 0.16 | 0.08 | ND | U | 0.2 | 0.16 | 0.08 | |
| EPA 8260 B (VOCs) | 1,1,1-Trichloroethane | 200 | 970 | ND | U | 1.0 | 0.28 | 0.14 | ND | U | 1.0 | 0.28 | 0.14 | ND | U | 1.0 | 0.28 | 0.14 | ND | U | 1.0 | 0.28 | 0.14 | ND | U | 1.0 | 0.28 | 0.14 |
| | 1,1,2-Trichloroethane | 5 | 50,000 | ND | U | 1.0 | 0.40 | 0.20 | ND | U | 1.0 | 0.40 | 0.20 | ND | U | 1.0 | 0.40 | 0.20 | ND | U | 1.0 | 0.40 | 0.20 | ND | U | 1.0 | 0.40 | 0.20 |
| | 1,1-Dichloroethane | 2.4 | 50,000 | ND | U | 1.0 | 0.38 | 0.19 | ND | U | 1.0 | 0.38 | 0.19 | ND | U | 1.0 | 0.38 | 0.19 | ND | U | 1.0 | 0.38 | 0.19 | ND | U | 1.0 | 0.38 | 0.19 |
| | 1,1-Dichloroethylene (1,1-Dichloroethene) | 7 | 1,500 | ND | U | 1.0 | 0.60 | 0.30 | ND | U | 1.0 | 0.60 | 0.30 | ND | U | 1.0 | 0.60 | 0.30 | ND | U | 1.0 | 0.60 | 0.30 | ND | U | 1.0 | 0.60 | 0.30 |
| | 1,2,3-Trichloropropane | 0.6 | 50,000 | ND | U | 2.0 ^b | 0.78 | 0.39 | ND | U | 2.0 ^b | 0.78 | 0.39 | ND | U | 2.0 | 0.78 | 0.39 | ND | U | 2.0 ^b | 0.78 | 0.39 | ND | U | 2.0 ^b | 0.78 | 0.39 |
| | 1,2,4-Trichlorobenzene | 70 | 3,000 | ND | U | 1.0 | 0.42 | 0.21 | ND | U | 1.0 | 0.42 | 0.21 | ND | U | 1.0 | 0.42 | 0.21 | ND | U | 1.0 | 0.42 | 0.21 | ND | U | 1.0 | 0.42 | 0.21 |
| | 1,2-Dibromo-3-chloropropane | 0.04 | 10 | ND | U | 2.0 ^b | 1.52 | 0.76 | ND | U | 2.0 ^b | 1.52 | 0.76 | ND | U | 2.0 ^b | 1.52 | 0.76 | ND | U | 2.0 ^b | 1.52 | 0.76 | ND | U | 2.0 ^b | 1.52 | 0.76 |
| | 1,2-Dibromoethane | 0.0065 | 50,000 | ND | U | 1.0 ^b | 0.40 | 0.20 | ND | U | 1.0 ^b | 0.40 | 0.20 | ND | U | 1.0 ^b | 0.40 | 0.20 | ND | U | 1.0 ^b | 0.40 | 0.20 | ND | U | 1.0 ^b | 0.40 | 0.20 |
| | 1,2-Dichlorobenzene | 600 | 10 | ND | U | 1.0 | 0.34 | 0.17 | ND | U | 1.0 | 0.34 | 0.17 | ND | U | 1.0 | 0.34 | 0.17 | ND | U | 1.0 | 0.34 | 0.17 | ND | U | 1.0 | 0.34 | 0.17 |
| | 1,2-Dichloroethane | 0.15 | 7,000 | ND | U | 1.0 ^b | 0.28 | 0.14 | ND | U | 1.0 ^b | 0.28 | 0.14 | ND | U | 1.0 ^b | 0.28 | 0.14 | ND | U | 1.0 ^b | 0.28 | 0.14 | ND | U | 1.0 ^b | 0.28 | 0.14 |
| | 1,2-Dichloropropane | 5 | 10 | ND | U | 1.0 | 0.34 | 0.17 | ND | U | 1.0 | 0.34 | 0.17 | ND | U | 1.0 | 0.34 | 0.17 | ND | U | 1.0 | 0.34 | 0.17 | ND | U | 1.0 | 0.34 | 0.17 |
| | 1,3-Dichlorobenzene | 180 | 50,000 | ND | U | 1.0 | 0.22 | 0.11 | ND | U | 1.0 | 0.22 | 0.11 | ND | U | 1.0 | 0.22 | 0.11 | ND | U | 1.0 | 0.22 | 0.11 | ND | U | 1.0 | 0.22 | 0.11 |
| | 1,3-Dichloropropene (total of cis/trans) | 0.43 | 50,000 | ND | U | 1.0 | 0.36 | 0.18 | ND | U | 1.0 | 0.36 | 0.18 | ND | U | 1.0 | 0.36 | 0.18 | ND | U | 1.0 | 0.36 | 0.18 | ND | U | 1.0 | 0.36 | 0.18 |
| | 1,4-Dichlorobenzene | 75 | 5 | ND | U | 1.0 | 0.38 | 0.19 | ND | U | 1.0 | 0.38 | 0.19 | ND | U | 1.0 | 0.38 | 0.19 | ND | U | 1.0 | 0.38 | 0.19 | ND | U | 1.0 | 0.38 | 0.19 |
| | Acetone | 22,000 | 20,000 | ND | U | 10.0 | 1.90 | 0.95 | ND | U | 10.0 | 1.90 | 0.95 | ND | U | 10.0 | 1.90 | 0.95 | ND | U | 10.0 | 1.90 | 0.95 | ND | U | 10.0 | 1.90 | 0.95 |
| | Benzene | 5 | 170 | ND | U | 1.0 | 0.32 | 0.16 | ND | U | 1.0 | 0.32 | 0.16 | ND | U | 1.0 | 0.32 | 0.16 | ND | U | 1.0 | 0.32 | 0.16 | ND | U | 1.0 | 0.32 | 0.16 |
| | Bromodichloromethane | 0.22 | 50,000 | ND | U | 1.0 ^b | 0.28 | 0.14 | ND | U | 1.0 ^b | 0.28 | 0.14 | ND | U | 1.0 ^b | 0.28 | 0.14 | ND | U | 1.0 ^b | 0.28 | 0.14 | ND | U | 1.0 ^b | 0.28 | 0.14 |
| | Bromoform | 100 | 510 | ND | U | 1.0 | 0.28 | 0.14 | ND | U | 1.0 | 0.28 | 0.14 | ND | U | 1.0 | 0.28 | 0.14 | ND | U | 1.0 | 0.28 | 0.14 | ND | U | 1.0 | 0.28 | 0.14 |
| | Bromomethane | 8.7 | 50,000 | ND | U | 2.0 | 0.48 | 0.24 | ND | U | 2.0 | 0.48 | 0.24 | ND | U | 2.0 | 0.48 | 0.24 | ND | U | 2.0 | 0.48 | 0.24 | ND | U | 2.0 | 0.48 | 0.24 |
| | Carbon Tetrachloride | 5 | 520 | ND | U | 1.0 | 0.20 | 0.10 | ND | U | 1.0 | 0.20 | 0.10 | ND | U | 1.0 | 0.20 | 0.10 | ND | U | 1.0 | 0.20 | 0.10 | ND | U | 1.0 | 0.20 | 0.10 |
| | Chlorobenzene | 100 | 50 | ND | U | 1.0 | 0.42 | 0.21 | ND | U | 1.0 | 0.42 | 0.21 | ND | U | 1.0 | 0.42 | 0.21 | ND | U | 1.0 | 0.42 | 0.21 | ND | U | 1.0 | 0.42 | 0.21 |
| | Chloroethane | 8,600 | 16 | ND | U | 1.0 | 0.42 | 0.21 | ND | U | 1.0 | 0.42 | 0.21 | ND | U | 1.0 | 0.42 | 0.21 | ND | U | 1.0 | 0.42 | 0.21 | ND | U | 1.0 | 0.42 | 0.21 |
| | Chloroform | 70 | 2,400 | ND | U | 1.0 | 0.14 | 0.07 | ND | U | 1.0 | 0.14 | 0.07 | ND | U | 1.0 | 0.14 | 0.07 | ND | U | 1.0 | 0.14 | 0.07 | ND | U | 1.0 | 0.14 | 0.07 |
| | Chloromethane | 1.8 | 50,000 | ND | U | 1.0 | 0.62 | 0.31 | ND | U | 1.0 | 0.62 | 0.31 | ND | U | 1.0 | 0.62 | 0.31 | ND | U | 1.0 | 0.62 | 0.31 | ND | U | 1.0 | 0.62 | 0.31 |
| | cis-1,2-Dichloroethylene (cis-1,2-Dichloroethene) | 70 | 50,000 | ND | U | 1.0 | 0.32 | 0.16 | ND | U | 1.0 | 0.32 | 0.16 | ND | U | 1.0 | 0.32 | 0.16 | ND | U | 1.0 | 0.32 | 0.16 | ND | U | 1.0 | 0.32 | 0.16 |
| | Dibromochloromethane (Chlorodibromomethane) | 0.16 | 50,000 | ND | U | 1.0 ^b | 0.38 | 0.19 | ND | U | | | | | | | | | | | | | | | | | | |

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However, TPH-DRO concentrations decreased from October 2008 through July 2009. In May 2009 and July 2009, TPH-DRO remained above the DOH Drinking Water EAL, but was below 50 percent of the SSRBL of 4,500 µg/L. In October 2009, TPH-DRO began an increasing trend greater than 50 percent of the SSRBL which continued through February 2010 when it exceeded the SSRBL due to TICs apparently not associated with petroleum from the RHSF (TEC, 2010). In March 2010 (2,490 µg/L) and April 2010 (2,215 µg/L), TPH-DRO exhibited a decreasing trend and the TICs detected in the two previous monitoring events were not observed. During July 2010, TPH-DRO concentrations at RHMW02 increased to an averaged concentration of 3,085 µg/L, above 50 percent of the SSRBL of 4,500 µg/L. During October-November 2010, TPH-DRO concentrations decreased to 1,700 µg/L, and in January 2011 concentrations decreased further to 1,100 µg/L, below 50 percent of the SSRBL (Appendix A).

Since September 2005, TPH-GRO concentrations have remained below the EAL, except in July 2006, December 2006, March 2007, and October-November 2010 (Appendix A).

From September 2005 through October 2008, naphthalene concentrations remained above the EAL and were relatively stable. In February 2009, naphthalene concentrations began decreasing and reached the lowest average concentration in May 2009 (2 µg/L) which was below the EAL. From July 2009 through July 2010, concentrations increased above the EAL. Then in October-November 2010 and January 2011 (60 µg/L), naphthalene concentrations decreased slightly to concentrations which were still above the EAL.

Similar to the naphthalene concentration trend, 1-methylnaphthalene concentrations remained relatively stable from September 2005 through October 2008. In February 2009, 1-methylnaphthalene began decreasing and reached the lowest average concentration in October 2009 (3.2 µg/L) which was below the EAL. In January 2010, 1-methylnaphthalene concentrations increased above the EAL; then decreased in April 2010; and has exhibited an increasing trend from July 2010 through January 2011 (Appendix A). The average concentration in January 2011 for 1-methylnaphthalene (21.0 µg/L) is above the DOH Drinking Water EAL of 4.7 µg/L.

Since October 2008, the concentration of 2-methylnaphthalene has remained below the EAL (Appendix A).

RHMW03

Historically, concentrations of TPH-DRO have fluctuated around the DOH Drinking Water EAL, but have been significantly lower than corresponding values observed at RHMW01 and RHMW02. TPH-DRO concentrations have decreased since October 2008 dropping below the LODs in May 2009 through July 2010. However, during the October-November 2010 groundwater monitoring event, TPH-DRO was detected at the highest concentration to date (330 µg/L) which was above the EAL. During this January 2011 groundwater monitoring event, TPH-DRO concentrations have decreased to below the LOD. TPH-GRO has remained below the EAL or has not been detected (Appendix A).

RHMW05

There was an increasing trend for TPH-DRO since it was first sampled in May 2009 through January 2010. Starting with the July 2009 monitoring event, TPH-DRO concentrations were greater than the DOH Drinking Water EAL (210 µg/L) with the highest concentration of 2,060 µg/L being observed in January 2010. It is important to note that the January 2010 concentration contained TICs apparently not associated with petroleum from the RHSF (TEC, 2010). However, in April 2010, July 2010, October-November 2010, and January 2011 TPH-DRO concentrations exhibited a decrease and were not detected at or above the LOD or LOQ. TPH-GRO has remained below the EAL or has not been detected (Appendix A).

3.4 Groundwater Status

Facility-specific contaminants of concern are defined as petroleum-related chemicals that have been observed in the groundwater samples above the DOH Drinking Water EALs. In accordance with the Red Hill Bulk Fuel Storage Facility Final Groundwater Protection Plan (TEC, 2008), Table 3-3 defines these RHSF-specific compounds and their associated SSRBLs and updated EALs (DOH, 2009).

In addition, the Groundwater Protection Plan defines four Results Categories of groundwater status for the RHSF, based on concentrations of COPCs detected in samples collected from RHMW2254-01, RHMW01, RHMW02, RHMW03, and RHMW05, and requires specific responses when these categories are observed during quarterly groundwater sampling (Table 3-4). The current Results Categories for the monitoring wells were determined using the January 2011 analytical data (Table 3-5).

Table 3-3: Action Levels for Contaminants of Concern

| Chemical | DOH Drinking Water EAL (µg/L) | SSRBL (µg/L) |
|--------------------------------|----------------------------------|-----------------|
| Petroleum Mixtures | | |
| TPH-DRO | 210 | 4,500 |
| TPH-GRO | 100 | 4,500 |
| Semi-Volatile Compounds | | |
| 1-Methylnaphthalene | 4.7 | N/A |
| 2-Methylnaphthalene | 24 | N/A |
| Naphthalene | 17 | N/A |

Notes:

N/A – not applicable

SSRBLs are applicable at RHMW01, RHMW02, RHMW03, and RHMW05

EALs are applicable at RHMW2254-01

Table 3-4: Results Categories and Response Actions to Changes in Groundwater Status

| Results Category | RHMW02, RHMW03, or RHMW05* | RHMW01 | RHMW2254-01 |
|--|------------------------------|---------------------------------|------------------------------------|
| Results Category 1: Result above detection limit but below drinking water EAL and trend for all compounds stable or decreasing | A | A | A, D, M, E |
| Results Category 2: Trend for any compound increasing or drinking water EAL exceeded | A, B | A, B | A, B, C, D, E, F, G, K, L, O |
| Results Category 3: Result Between 1/10X SSRBL and SSRBL for benzene, or between 1/2X SSRBL and SSRBL for TPH | A, B, G, H, I, J | A, B, E, G, H, I, J | A, B, C, D, E, F, G, I, J, K, L, O |
| Results Category 4: Result Exceeding any SSRBL or petroleum product observed | A, C, D, E, F, I, J, K, M, N | A, C, D, E, F, I, J, K, M, N, O | A, C, D, E, F, G, I, J, K, L, O |

Notes:

*RHMW05 was installed in April 2009 and has subsequently been added to this table.

Specific Responses:

- A. Send quarterly reports to DOH.
- B. Begin program to determine the source of leak.
- C. Notify DOH verbally within one day and follow with written notification in 30 days.
- D. Notify FISC Chain of Command within one day.
- E. Send Type 1 Report (see box below) to DOH
- F. Send Type 2 Report (see box below) to DOH
- G. Increase monitoring frequency to once per month (if concentrations increasing).
- H. Notify DOH verbally within seven days and follow with written notification in 30 days.
- I. Remove sampling pumps, measure product in pertinent wells with interface probe, re-install pumps if product is not detected.
- J. Immediately evaluate tanks for leaks.
- K. Collect samples from nearby Hālawā Deep Monitoring Well (2253-03) and OWDFMW01. For permission to sample 2253-03, call Department of Land and Natural Resources (DLNR) Commission on Water Resource Management 808-587-0214, DLNRCWR@Hawaii.gov.
- L. Provide alternative water source at RHMW2254-01.
- M. Prepare for alternative water source at RHMW2254-01.
- N. Re-measure for product every month with reports to DOH.
- O. Install additional monitoring well downgradient.

Report Types

DOH Type 1 Report

- Re-evaluate Tier 3 Risk Assessment/groundwater model results
- Proposal to DOH on course of action

DOH Type 2 Report

- Proposal for groundwater treatment

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Table 3-5: Summary of Result Categories and Response Actions, January 2011

| Results Category | RHMW2254-01 | RHMW01 | RHMW02 | RHMW03 | RHMW05 |
|---|--------------------|---------------|--|---------------|---------------|
| <u>Results Category 1:</u> Result above detection limit but below drinking water EAL and trend for all compounds stable or decreasing | No | No | No | No | No |
| <u>Results Category 2:</u> Trend for any compound increasing or drinking water EAL exceeded | No | No | Yes | No | No |
| <u>Results Category 3:</u> Result Between 1/10X SSRBL and SSRBL for benzene, or between 1/2X SSRBL and SSRBL for TPH | No | No | No | No | No |
| <u>Results Category 4:</u> Result Exceeding any SSRBL or petroleum product observed | No | No | No | No | No |
| <u>Previous Category:</u> Category prior to January monitoring event | None | Category 1 | Category 2 | Category 2 | None |
| <u>New Category:</u> Category assignment based on results of the January monitoring event | None | None | Category 2 | None | None |
| <u>Response Actions:</u> Requirements of new category | None | None | 1. Quarterly reports to be sent to DOH 2. Initiation of a leak determination program to identify if tanks are leaking | None | None |

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Section 4 Data Quality Assessment

Data quality assessment consists of a review of the overall groundwater sample collection and analyses process in order to determine whether the analytical data generated meets the quality objectives for the project. The field QC program consisted of standardized sample collection and management procedures and the collection of field duplicate samples and trip blank samples. The laboratory quality assurance program consisted of the use of standard analytical methods and the preparation and analyses of MS/MSD samples, surrogate spikes, method blanks, and laboratory control samples (LCS).

4.1 Data Validation

The usability of the data collected during this investigation depends upon its quality. A number of factors relate to the quality of data, including: sample collection methods, sample analysis methods, and adherence to established procedures for sample collection, preservation, management, shipment, and analysis. Data quality is judged in terms of its precision, accuracy, representativeness, completeness, and comparability.

4.1.1 Quality Control Program

LOQs are established by the laboratory based on the method DLs or instrument DLs, historical data, and EPA limits established for the methods. The LOQs for samples may require adjustment due to matrix interference or if high levels of target analytes necessitate dilution before analysis. Matrix interference and sample dilutions have the effect of increasing the LOQs. None of the reporting limits were adjusted for this monitoring event.

4.1.2 Data Assessment

Precision

Precision is defined as the agreement between a set of replicate measurements without assumption and knowledge of the true value. Precision limits for laboratory measurements were evaluated from the sample/sample duplicate results. Field duplicate samples were collected at a rate of approximately 10 percent of project samples. Field duplicates are duplicates of the primary samples that are sent to the primary laboratory with dummy sample numbers to serve as a check of the sampling precision and the primary laboratory's analytical precision.

Field duplicates' precision was evaluated by the criteria which said that relative percentage difference (RPD) of the original sample/and sample duplicate results should be less than 50 percent for water samples.

For this monitoring event, the primary field sample ES010 and field duplicate sample ES011 collected from RHMW02, met the RPD precision criteria for all analytes (Table 4-1).

Accuracy

Accuracy is defined as the degree of agreement of a measurement to an accepted reference or true value. Accuracy is measured as the percent recovery of an analyte in a reference standard or spiked sample. Accuracy limits for laboratory control spike, MS, and MSD samples are

established by the individual laboratory. The acceptance criteria for accuracy are dependent on the analytical method, and are based on historical laboratory data.

All of the LCS, MS/MSD and surrogate spike recoveries for all analyzed constituents were within acceptable percent recovery limits. Therefore, all data is considered to be accurate.

Representativeness

Representativeness is the degree that data accurately and precisely represent a characteristic of a population, parameter variations at a sampling point, or an environmental condition. Representativeness was achieved by conducting sampling in compliance with the sample collection procedures described in the Work Plan specifically written for this project (Environet, 2010).

Representativeness is also evaluated through the analysis of blank samples including method blank and trip blank samples. As described above, none of the COPCs were detected in the laboratory method blanks. For this sampling event two trip blank samples were collected (Table 4-1). The trip blank samples for the January 2011 monitoring event were all non-detect, therefore, the groundwater sample data are considered representative of the groundwater quality on site.

Completeness

Completeness is defined as the overall percentage of valid analytical results (including estimated values) compared to the total number of analytical results reported by the laboratory. The completeness goal for this project is 90 percent, which was successfully met. Successful completion of data acquisition can only be accomplished if both the field and laboratory portions of the project are performed according to the procedures described in the Work Plan (Environet, 2010).

Comparability

Comparability expresses the confidence with which one data set can be compared to another. Comparability can be related to accuracy and precision because these quantities are measures of data reliability. Data are considered comparable if collection techniques, measurement procedures, methods and reporting are equivalent for the samples within a sample set. Comparability for sampling was determined to be acceptable based on the following criteria:

A consistent approach to sampling was applied throughout the program.

- Samples were consistently preserved.
- All sampling was performed during the same time of year and under similar physical conditions.

The above criteria were met throughout the course of this event and therefore comparability of the data set is acceptable.

4.1.3 Data Assessment Conclusions

The precision, accuracy, representativeness, completeness, and comparability criteria were met. Therefore, the data assessment concludes that all data generated during this event are suitable for the intended use.

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Section 5 Summary, Conclusions, and Recommendations

5.1 Summary

There is no indication of an immediate threat of disruption to drinking water resources of the RHMW2254-01 as a result of the January 2011 data. Based on the January 2011 groundwater monitoring event, RHMW2254-01 does not fall into any Results Category of the Groundwater Protection Plan.

5.1.1 Results of Oil/Water Interface Measurements

Free product was not observed at RHMW01, RHMW02, RHMW03, and RHMW05 during the January 2011 sampling event. The trend of free product measurements over time shows that in January 2008, fuel was measured in monitoring wells RHMW01 and RHMW02 at a thickness of less than 0.01 feet, but was not observed in other monitoring wells. Since the trace amounts observed in January 2008, no free product has been observed in any of these RHSF monitoring wells (Table 3-1).

5.1.2 Summary of Groundwater Analytical Results

Laboratory analytical results from the January 2011 groundwater monitoring event indicated that TPH-DRO, and PAHs, specifically 1-methylnaphthalene and naphthalene, were present in the groundwater beneath the RHSF at concentrations that exceeded the EALs. All other COPC concentrations were below the EALs. All DLs, LODs and LOQs were generally below the EALs. In the case where an EAL for a specific COPC is less than the LOQ, it is generally acceptable to consider the LOQs in place of the EAL (DOH, 2009).

At monitoring wells RHMW2254-01, RHMW01, RHMW03, and RHMW05, all COPCs were not detected at or above the LODs and LOQs.

At monitoring well RHMW02, TPH-DRO (1,100 µg/L), 1-methylnaphthalene (19 µg/L), and naphthalene (57 µg/L) were detected at concentrations which exceeded the EALs. Acenaphthene (0.29 µg/L) and 2-methylnaphthalene (3.6 µg/L) were detected at concentrations which were below the EALs. Acenaphthylene (0.13µg/L), fluorene (0.15 µg/L), ethylbenzene (0.25 µg/L), TPH-GRO (17 µg/L), and xylenes (0.60 µg/L), were detected at estimated concentrations which were below the EALs. All other COPCs in RHMW02 were not detected at or above the LODs and LOQs.

5.2 Conclusions

To date, the presence of LNAPL has been observed only once (i.e., in January 2008 in RHMW01 and RHMW02 at a thickness of less than 0.01 feet). This indicates a significant release from one or more of the USTs at the RHSF has not occurred at this time.

COPC concentrations in samples collected from the U.S. Navy Well 2254-01 have not been detected above the LODs and are below the EALs. This indicates that elevated COPC

concentrations detected in samples collected from monitoring wells within the RHSF are not migrating and impacting the Navy's potable water source.

COPC concentrations detected in samples collected from monitoring well RHMW05 have decreased to below the EALs for at least three consecutive monitoring events. The data suggest that elevated COPC concentrations detected in samples collected from monitoring wells located adjacent to the USTs are not migrating in the downgradient direction. RHMW05 is an intermediate monitoring point between the USTs and the U.S. Navy Well 22554-01. At this time, there is no indication that COPCs are migrating in a downgradient direction towards the Navy's potable water source.

COPC concentrations detected in samples collected from two monitoring wells adjacent to the USTs (i.e., RHMW01, and RHMW03) are not steadily increasing between sampling events. COPC concentrations of 1-methylnaphthalene detected in samples collected from RHMW02 however have increased for three consecutive events.

5.3 Recommendations

Based on the results of this monitoring event, continuation of the quarterly groundwater monitoring program at the RHSF is recommended. In addition continuation of monthly free product measurements at RHMW01, RHMW02, RHMW03, and RHMW05; and monthly soil vapor monitoring are also recommended. In response to the Category 2 status of RHMW02, submission of this quarterly report to DOH and continuation of a leak determination program as described in Section 3 of the RHSF Groundwater Protection Plan (TEC, 2008) to identify if tanks are leaking are recommended.

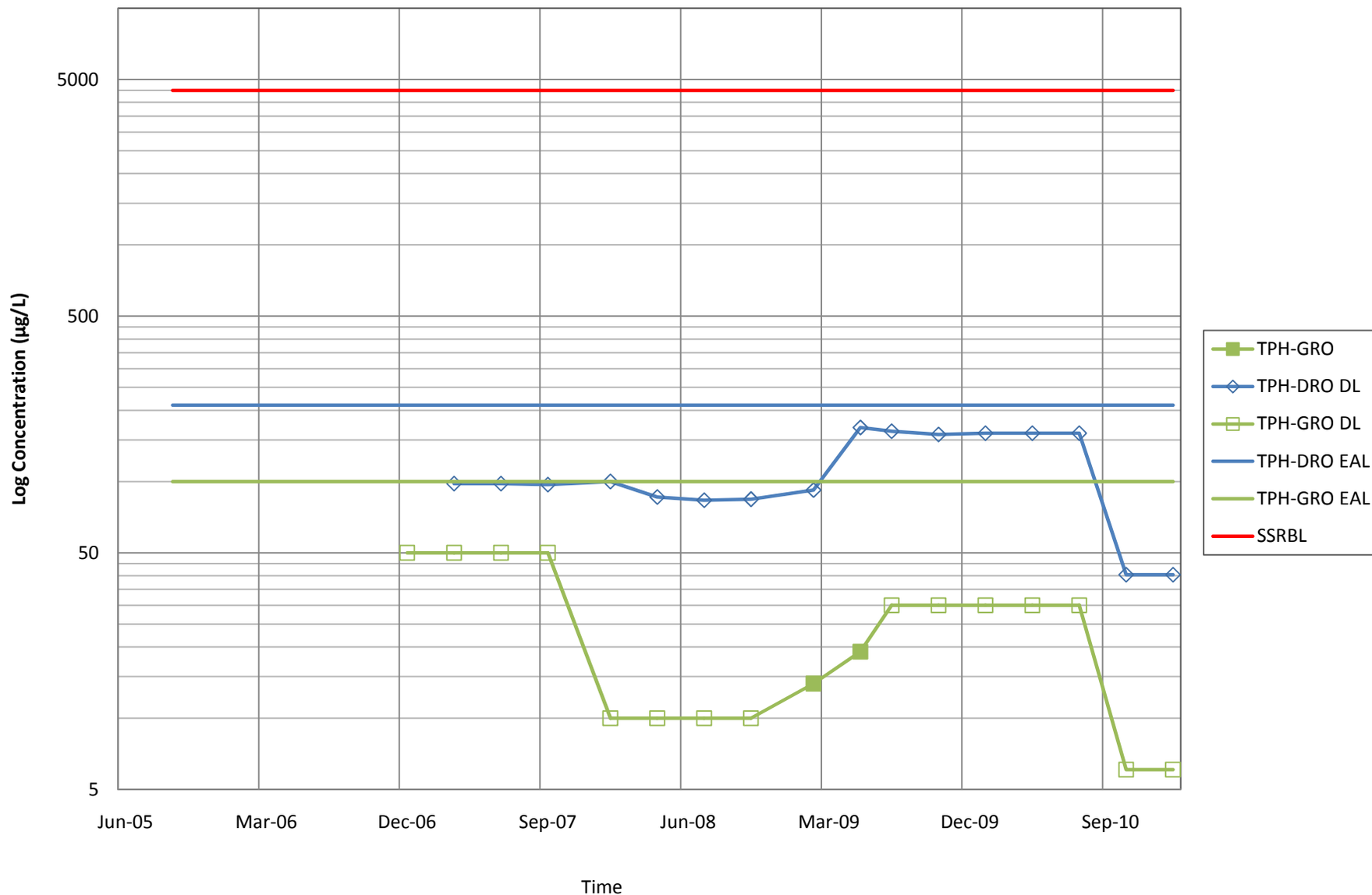
Section 6 References

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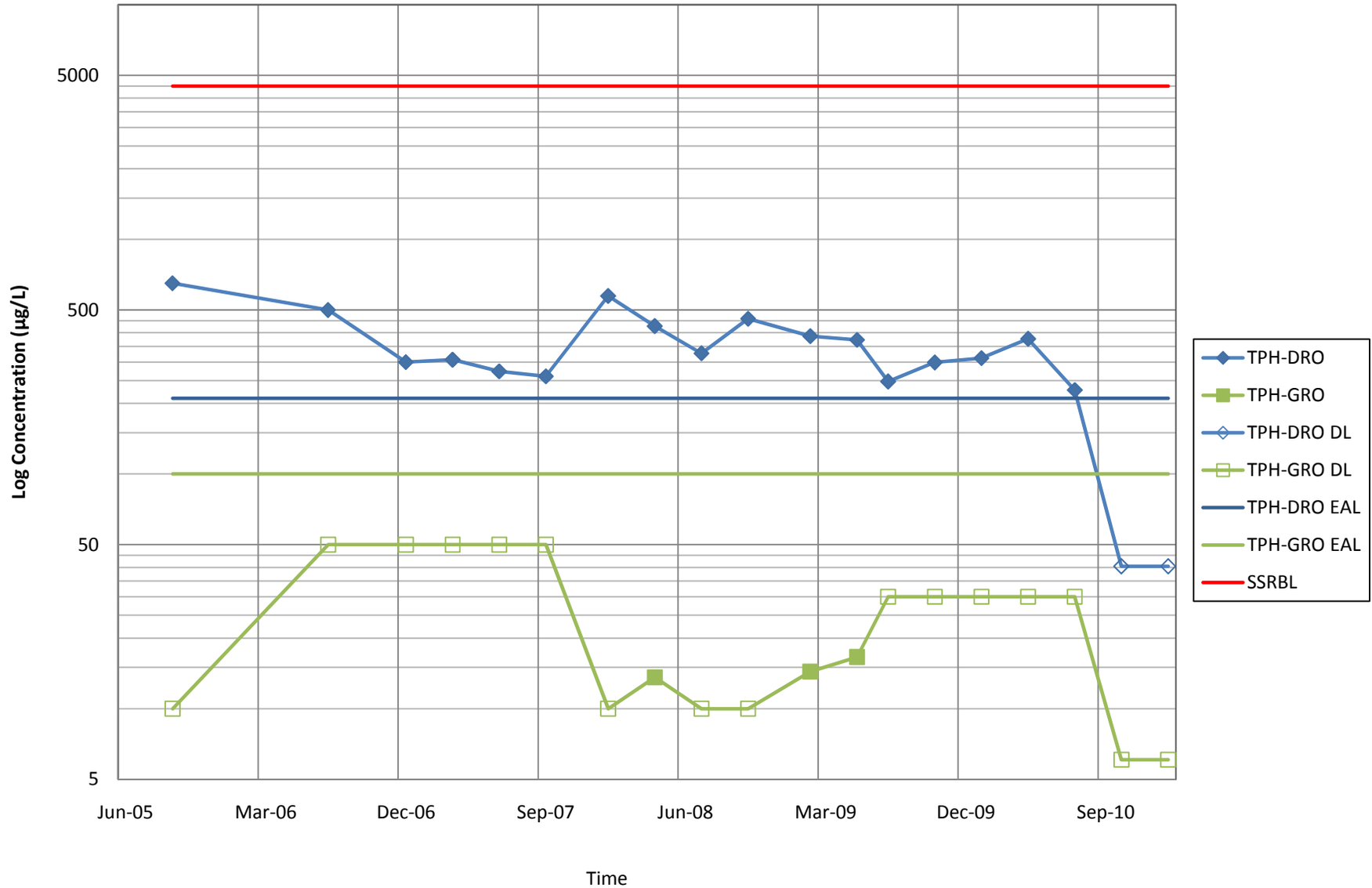
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Appendix A
Quarterly Groundwater Monitoring Results for
TPH and PAHs, September 2005 through
January 2011

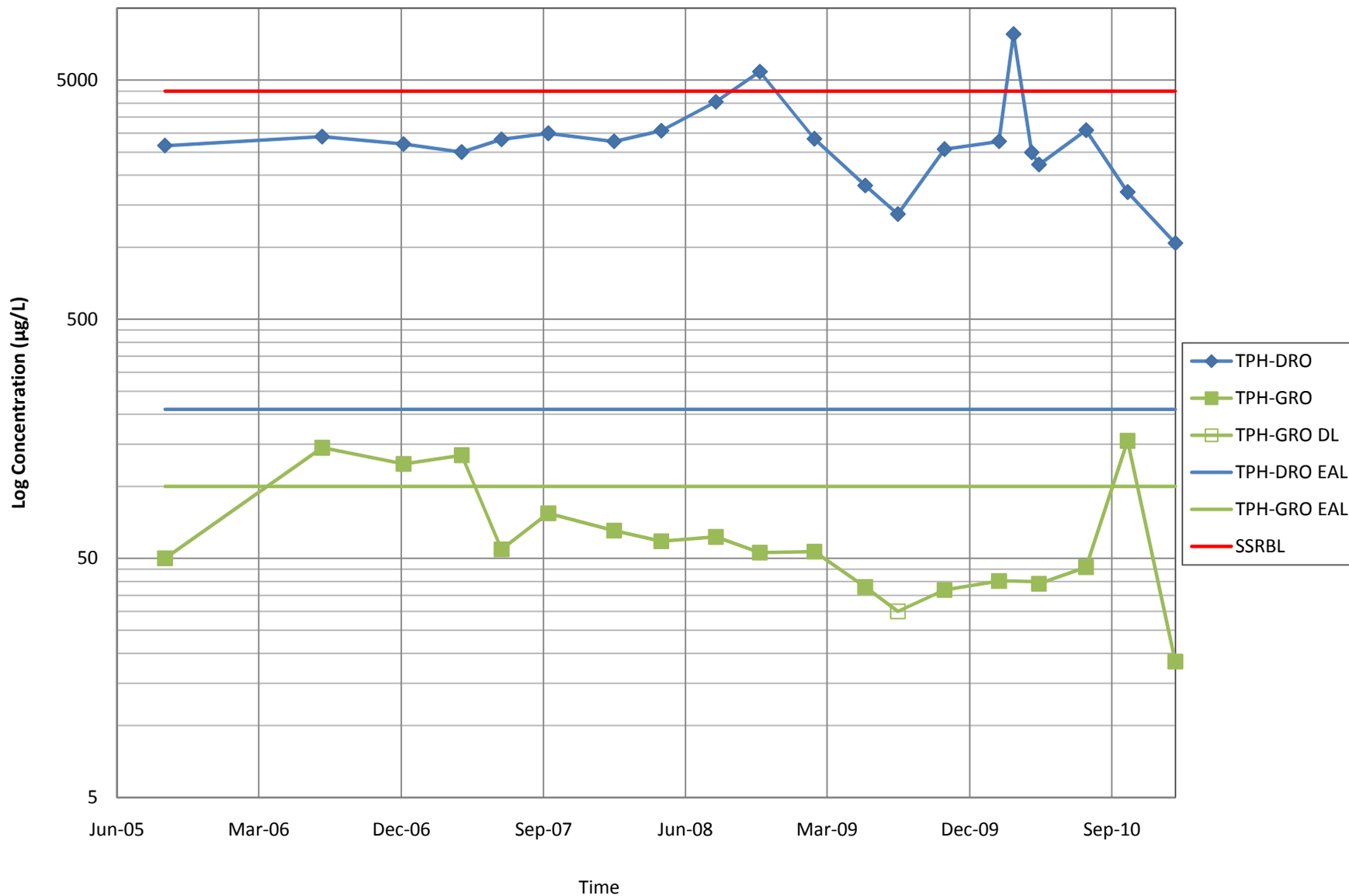
**Figure A-1: TPH at RHMW2254-01
Quarterly Monitoring Results**



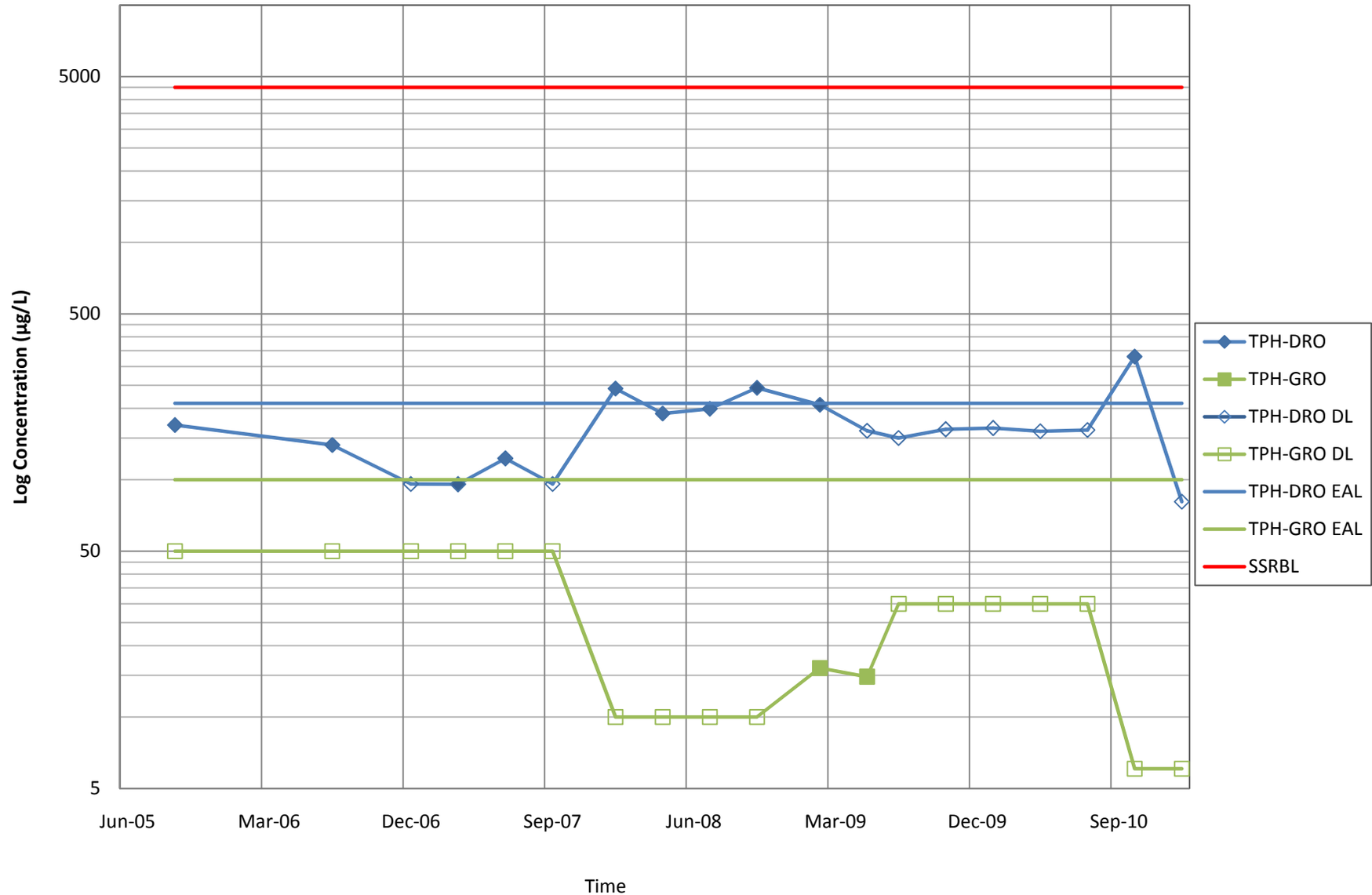
**Figure A-2: TPH at RHMW01
Quarterly Monitoring Results**



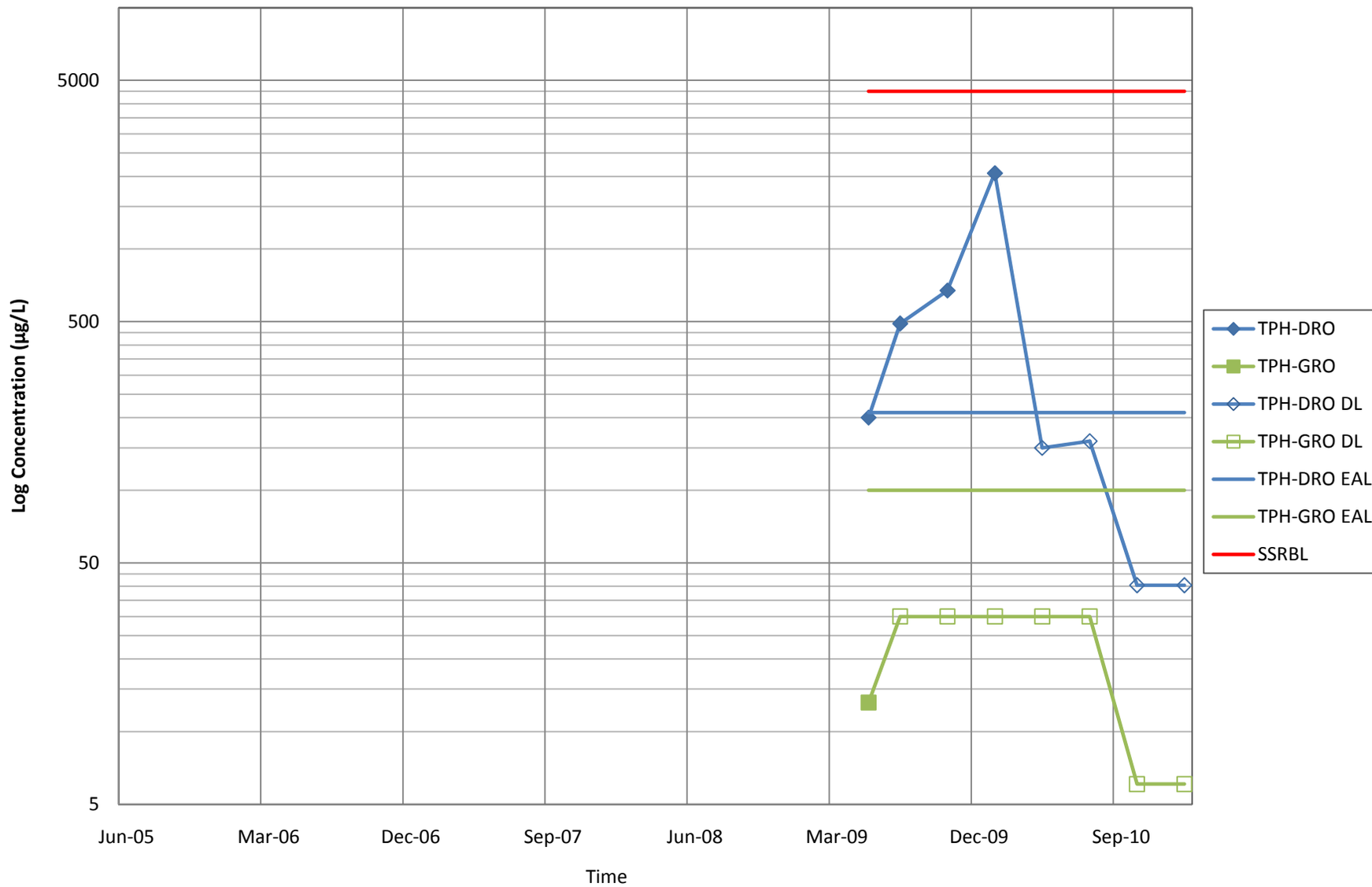
**Figure A-3: TPH at RHMW02
Quarterly Monitoring Results**



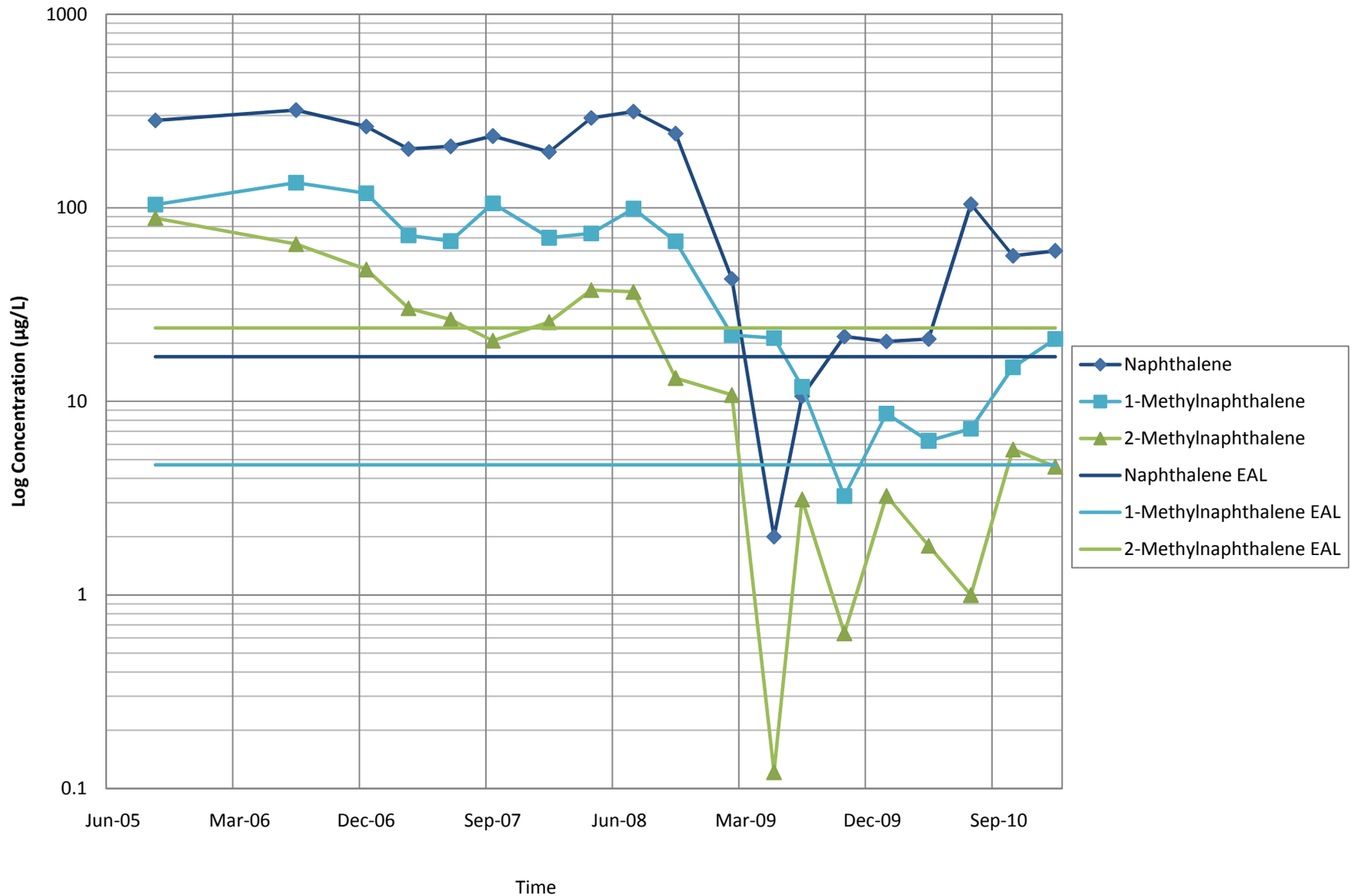
**Figure A-4: TPH at RHMW03
Quarterly Monitoring Results**



**Figure A-5: TPH at RHMW05
Quarterly Monitoring Results**



**Figure A-6: PAHs at RHMW02
Quarterly Monitoring Results**



Appendix B
Field Notebook, January 2011

RHSF LTM

1022-015

SS

1/18/11

Activity: Ground water Monitoring

Personnel: C A T S F PID

Equipment: Solinst, YPM Nitrogen tank,

controller, Horiaba, compressor (backup)

0730 Meet at office

0800 Arrive at Storage

0830 Pick up ice + saturated from 76.

0900 Arrived in Tunnels + Calibrated PID.

0922 Calibrate Horiaba

1045 Train cones to Adit 3

110 Arrive at RHMW 02

117 PID is 0.5 ppm

DTW: 86.70

~~1155~~

* started purging RHMW 02

1215 Sample ESD 10

Duplicate ESD 11 ~~made up~~ made up

1030 sample time.

MSD ESD 11 also taken at RHMW 02

1350 Task Lunch break at Adit 5

1500

Exited Tunnels

"Get in the Rain"

RHSF LTM 1022-015

1/19/11
 0625 Arrived at tunnels and conducted health & safety meeting. Calibrated PID Fresh Air

0913 Arrived at MW03 calibrated PID spun cal + calibrated Horiba.

0928 Started purging 103.5g after sampling.

1000 Sampled MW 03 at 250 L2.

1030 Took break at Adit 5

1100 Mobilized to RHMW05

1130 Arrived at MW05
 PID was 9.1
 83.37% DTW

1146 Started purging 83.35 DTW

~~1300~~
 1155 Sampled at RHMW 05
 Mobilized to exit tunnels

1300 Ran out of Nitrogen used air pump.

1330 Exited tunnels

1545 Fed ex drop off samples

RHSF LTM 1022-015 57

1/20/11

Task: Gw Sampling
 Personnel: CHSF
 Equipment: Controller, Nit tank, compressor, PID, Splineist, Horiba

0730 Met at office

0800 Storage to pick up Cal 4
 Solutich + plastic sheeting

0830 Arrive at Adit 3 + on board.

0930 Navy Sinner arrives to let us Imp pump Station well RHMW 254-01

952 PID = 36.4
 Sampled RHMW 254-01.

1063 ISO 14

1047 Called Elton for train pickup

1120 Took lunch at Adit 5

1154 Arrived at RHMW-01

1159 PID ~~19.5~~ 13.11 ppm
 9.6 ambient

D12 464.03 DTW

1300 Started purging RHMW-01
 Discharge 5 Change 5
 with Nitrogen tank

1022-015

RHSF LTM

1022-018

1/20/11

1305

Sample time of RHMW-01 recorded as

ESO15.

84.01 for after purging

while collecting VOx's Nitrogen

rain. out. Switched to Air

compressor. Air compressor continually

overheated and we had to turn

it off to let it cool down.

Finished sampling

Train came to transfer to

Adit 3.

End of Shift

Stacy AR

| Sample ID | Description | ID | Time | Date |
|-----------|-------------|-----------|------|----------|
| ES001 | RH-RHMW03 | -G021 | | 10/18/10 |
| ES002 | RH-RHMW02 | -G021 | | 10/18/10 |
| ES003 | RH-RHMW02 | Dup | | 10/18/10 |
| ES004 | RH-RHMW2254 | 01-G021 | | 10/19 |
| ES005 | RH-RHMW05 | -G021 | | 10/20 |
| ES006 | RH-RHMW2253 | 03-G021 | | 10/21 |
| ES007 | RH-RHMW01 | -G021 | | 10/21 |
| ES008 | RH-RHMW01 | Dup | | 10/21 |
| ES009 | RH-RHMW01 | -G021 | | 11/3 |
| ES010 | RH-RHMW02 | -G021 | | 1/18 |
| ES011 | RH-RHMW02 | -G021 Dup | | 1/18 |
| ES012 | RH-RHMW03 | -G021 | 1000 | 1/19 |
| ES013 | RH-RHMW05 | -G021 | 1155 | 1/19 |
| ES014 | RH-RHMW2254 | 01-G021 | 1003 | 1/19 |
| ES015 | RH-RHMW01 | -G021 | 1305 | 1/19 |
| ES016 | RH-RHMW2253 | 03-G021 | 0115 | 1/21 |
| ES017 | RH-RHMW01 | -G021 | 1160 | 1/21 |
| ES018 | RH-RHMW01 | -G021 Dup | 0830 | 1/21 |
| ES019 | | | | |

1022-015 LTM R HSF

1022-015 LTM R HSF 1
1/21/2011

0800 Met at office. Calibrated

PID.

0930 Arrive at Haines prison to

meet with DLNR.

Max and Charles went

to Red Hill ODWEMU

to sample while Steacy

and Zach went to

HD MU.

1000 Forgot cal 4 solution +

went to retrieve from

MS + Ca.

1030 Horiba malfunction. Went

to Red Hill to retrieve

other Horiba from CA + MS.

1110 DLNR started putting

OTD down, causing delay.

Task: GW sampling outside wells

Personnel: ZP + SF

Equip: Horiba, boiler, Vacuum pump

Weather: Sunny

1154 Started Purginy.

1/21/14

1022-015 OWD F M U 01

0830

Met at the office SE, CA,
ZP, MS

0945

MS & CA leave for OWD F M U 01
at Red Hill while Spaw
and Zach went to talk
Set up equipment at
O W D F M U 01

1000

1025

PID = 200 ppm

1041

Started to purge

1111

Sample collection
Sample ID: ES017, ES018,
ES019 MS/MSDS

Weather: Sunny
Method of Removal: 2" bailer

~~END OF ENTRY~~

~~Signature~~

RHSF LTM

1022-015

1/26/14

Task: Depth of O W D F M U 01

Personnel: CA & SF

Equip: Water Level meter
100.60 ft DTW @ 1002

End of Entry

Signature

1022-015 LTM RHISFC 3
~~OWDFM101~~

~~0800 Met at the office. Calibrated
PID
0930 Arrive at Red Hill ODUSE
(MS & SA) MW01
1000 Set up equipment
1040 Started purging
11030 PID = 200 ppm
Background reading
Total depth~~

1215 Sampled HDMW2253-03

1300 Sample ID ES016
Packed up and checked out
of Halamon Prison. Went back
to office to repackage

1500 Samples to ship to Lab.
Arrived at Fed Ex to ship
Samples to Appl Lab.

Scheduled for Saturday
delivery.

1600 Back to office

End of

Empty
Storage
A. J. Jr.



Appendix C
Groundwater Sampling Logs, January 2011

GROUNDWATER SAMPLING LOG

WELL ID: RHMW2254-01 LOCATION: Red Hill Bulk Fuel Storage Facility PROJECT NO: 1022-015
 INITIAL WATER LEVEL: not applicable (N/A) DATE: 1/20/2011 TIME: 936
 TOTAL DEPTH OF WELL: unknown PERSONNEL INVOLVED: S.Fineran, C. Asselbaye
 LENGTH OF SATURATED ZONE: N/A WEATHER CONDITIONS: N/A
 VOLUME OF WATER TO BE REMOVED: N/A METHOD OF REMOVAL: Low flow dedicated bladder pump
 WATER LEVEL AFTER PURGING: N/A PUMPING RATE: Approximately .65 L/min

WELL PURGE DATA:

| TIME | VOLUME REMOVED | pH | COND (mS/m) | TURBIDITY (NTU) | DO (mg/l) | TEMP (°C) | SALINITY (%) | REDOX (ORP) (mV) |
|------|----------------|------|-------------|-----------------|-----------|-----------|--------------|------------------|
| 0956 | 1 L | 5.31 | 54.7 | 1.0 | 8.00 | 22.1 | 0.0 | 221 |
| 0958 | 2 L | 5.81 | 50.0 | 2.7 | 8.37 | 21.7 | 0.0 | 220 |
| 0959 | 3 L | 5.99 | 49.4 | 2.4 | 8.30 | 21.7 | 0.0 | 217 |
| 1001 | 4 L | 6.10 | 49.0 | 1.2 | 8.38 | 21.5 | 0.0 | 221 |
| 1003 | 5 L | 6.21 | 49.0 | 1.1 | 8.30 | 21.5 | 0.0 | 219 |
| 1004 | 6 L | 6.31 | 48.9 | 1.1 | 8.33 | 21.5 | 0.0 | 221 |
| | | | | | | | | |
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| | | | | | | | | |

SAMPLE RETRIEVAL METHOD: Low flow dedicated bladder pump APPEARANCE OF SAMPLE:
 COLOR None
 SAMPLE ID: ES014 (RH-RHMW2254-01-GW22) TURBIDITY None
 SAMPLE COLLECTION TIME: 10:03 SEDIMENT None
 SAMPLED BY: S.Fineran, C. Asselbaye OTHER _____

COMMENTS AND OBSERVATIONS: PID = 0.0 ppm
Depth to water from the platform hole was approximately 84.85 feet.

LABORATORY ANALYSIS PARAMETERS AND PRESERVATIVES: TPH-GRO (EPA 8260B) & VOCs (EPA 8260B) ,
TPH-DRO (EPA8015B) & PAHs (EPA 8270D SIM) with no preservative, and dissolved lead (EPA 6020) with nitric acid preservative

NUMBER AND TYPES OF SAMPLE CONTAINERS FILLED: (4) - 40 mL VOAs, (4) - 1 L amber bottle, (1) - 500 mL polyethylene bottle

DECONTAMINATION PROCEDURES: Alconox, triple rinse with distilled water
 SAMPLES DELIVERED TO: APPL, Inc. via FedEx TRANSPORTERS: S.F., C.A.
 SAMPLE DELIVERY DATE: 1/21/2011 SAMPLE DELIVERY TIME: 1545

CAPACITY OF CASING (GALLONS/LINEAR FOOT)
 2"-0.16; 4"-0.65; 6"-1.47; 8"-2.61; 10"-4.08; 12"-5.87

Appendix D
Laboratory Analytical Results,
January 2011 (on CD-ROM)

