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Verification Monitoring Report for the Durango, Colorado, Processing Site

September 2010



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Office of Legacy Management

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Abbreviations

ACL	alternate concentration limit
DOE	U.S. Department of Energy
EPA	U.S. Environmental Protection Agency
ft	foot (feet)
ft ³ /day	cubic foot (feet) per day
GCAP	Ground Water Compliance Action Plan
MCL	maximum concentration limit
mg/L	milligram(s) per liter
POC	point-of-compliance
UMTRA	Uranium Mill Tailings Remedial Action (Project)

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1.0 Introduction

The Durango processing site is located in La Plata County, Colorado, approximately 0.25 mile southwest of the central business district of Durango, Colorado (Figure 1). The site consists of two separate areas: (1) the mill tailings area, which is the setting of former uranium-ore milling and storage of residual solid wastes (mill tailings), and (2) a raffinate ponds area where liquid process-wastes were impounded during milling operations. The former mill tailings area encompasses about 40 acres on a bedrock-supported river terrace between Smelter Mountain to the west, the Animas River to the east and south, and Lightner Creek to the north (Figure 2). The raffinate ponds area occupies about 20 acres on a separate river terrace located 1,500 feet (ft) south (downstream) of the mill tailings area (Figure 3).

1.1 Compliance Strategy

The compliance strategy for groundwater cleanup at the former mill tailings area of the Durango site is natural flushing, institutional controls to prevent exposure to contaminated groundwater, water quality monitoring, and an alternate concentration limit (ACL) for selenium (DOE 2003). This strategy was based in part on groundwater flow and solute transport modeling that predicted acceptable cleanup times for each contaminant, except possibly cadmium, by natural flushing processes at the site, and in part on historical trends of decreasing contaminant concentrations, particularly since the completion of contaminant source removal in 1991. Baseline conditions of contaminant concentration in the model correspond to results of the June 2002 groundwater sampling. The groundwater model is fully documented in the Site Observational Work Plan (DOE 2002).

The compliance strategy for the raffinate ponds area is no further action in conjunction with supplemental standards based on limited use (poor quality) groundwater. Groundwater monitoring is conducted as a best management practice. Monitoring results for the raffinate ponds area are also presented in this report.

1.2 Purpose

The primary purpose of this verification monitoring report is to evaluate the observed progress of passive groundwater restoration at the Durango mill tailings area, by comparing observed constituent concentrations through June 2010 with model-predicted concentrations for the same time period. The goal is to confirm that natural flushing is progressing and remains a viable compliance strategy for the site.

1.3 Summary

Evaluation of monitoring data through June 2010 and trend analysis of contaminant levels at site wells indicate that natural flushing remains a viable compliance strategy for the Durango processing site. The constituent concentration trends at several wells also appear to generally match model predictions. Cadmium concentrations are below the compliance goal for all locations except well 0612 at which trend analysis suggests this constituent will flush within a 100-year time frame. The compliance goal for manganese is also only exceeded at well 0612 and trend analysis indicates that this constituent will also sufficiently flush from site groundwater within 100 years. Molybdenum concentrations are below the compliance goal at all wells, which

is consistent with model predictions. Selenium concentrations are below the compliance goal for all locations except well 0633 which could be attributed to natural selenium in groundwater due to the leaching of shale bedrock underlying the site. Sulfate and uranium concentrations are above the compliance goals at all locations except at wells 0863 and 0631 for sulfate and wells 0863 and 0635 for uranium. Despite these observed exceedances, trend analysis of existing concentration data suggests that it is possible for sulfate and uranium to meet the compliance goals within 100 years.



Figure 1. Durango Processing Site



Figure 2. Monitoring Network in the Mill Tailings Area at the Durango Site



Figure 3. Monitoring Locations for the Raffinate Ponds Area

2.0 Site Conditions

2.1 Hydrogeology

The uppermost aquifer at the mill tailings area is shallow and consists mostly of poorly sorted colluvium derived from Smelter Mountain, which rises steeply to the southwest. Alluvial deposits associated with the Animas River and Lightner Creek also comprise a portion of the shallow aquifer. The colluvium and alluvium are underlain by the low-permeability Mancos Shale bedrock, which essentially acts a hydraulic barrier to downward from the shallow groundwater system. Approximately 70 ft of colluvium overlies bedrock along the base of Smelter Mountain. These deposits thin eastward and transition to deposits close to the Animas River that are about 15 ft thick. Depth to groundwater increases from about 5 ft on the river terrace to about 60 ft near the mountain-front. The saturated zone is thin (less than 10 ft), unconfined, of limited extent and has a low yield. Groundwater flow is generally to the southeast, parallel to the Animas River, at an average gradient of approximately 0.02 ft/ft. Hydraulic conductivity of the colluvium and alluvium ranges from 10 to 70 ft/day.

The colluvium is recharged primarily by runoff from Smelter Mountain and infiltrating precipitation, while the river alluvium receives surface water from Lightner Creek and from river loss along the upstream reach of a prominent river meander that defines the middle third of the mill tailings area's east boundary. Groundwater discharges to the Animas River along the upper and lower thirds of the river reach adjacent to the mill tailings area. Under average conditions, the estimated volume of groundwater discharge from the mill tailings area is 1,480 cubic feet per day (ft³/day); approximately 840 ft³/day of this total enters the Animas River near the mouth of Lightner Creek, and the remaining 640 ft³/day enters the Animas River east and southeast of the footprint of a former tailings pile (DOE 2002). The alluvium and colluvium pinch out against bedrock cliffs near the southeast corner of the site, at which point groundwater discharge to the river is complete (DOE 2002).

Groundwater in the raffinate ponds area occurs in two bedrock units, both members of the Mesaverde Group, that are separated by the northeast trending Bodo Fault (Figure 3). The Point Lookout Sandstone, the basal formation of the Mesaverde Group, lies south of the fault and is divided into two members: a lower transitional member consisting of interbedded lenticular sandstones and shales, and an upper massive sandstone member. The Menefee Formation, north of the fault, consists of massive sandstone and shale along with beds of carbonaceous shale and coal. The Bodo Fault, a normal fault, dips to the southeast at approximately 55 degrees. The Point Lookout Sandstone is downthrown approximately 200 ft along the fault.

Groundwater in the raffinate ponds area is assumed to be unconfined. It is recharged by infiltration of precipitation and runoff from the Smelter Mountain area and the ephemeral South Creek. Eastward-flowing subsurface water also enters the groundwater system near the intersection of Bodo Fault and South Creek (Figure 3). Hydraulic conductivity data indicate that the Point Lookout Sandstone is the least conductive of the various bedrock units underlying the raffinate ponds area. The lower member (predominantly shale and siltstone) of the Point Lookout Sandstone is considered an aquitard. The Menefee Formation consists of mostly low-conductivity sandstone but is relatively permeable where fractures or lenticular coal beds are present. The largest hydraulic conductivities appear to occur in the near-vicinity of Bodo Fault and in the coal beds within the Menefee.

2.2 Water Quality

Groundwater in the alluvial aquifer at the mill tailings area is contaminated as a result of uranium-ore processing and tailings storage. Although the primary source of groundwater contamination (mill tailings) was removed from the site by 1991, concentrations of arsenic, cadmium, lead, molybdenum, net alpha radiation, radium-226+228, selenium, and uranium remaining in the underlying aquifer exceeded maximum concentration limits (MCLs) established in Title 40 *Code of Federal Regulations* Part 192 (49 CFR 192) for the Uranium Mill Tailings Remedial Action (UMTRA) Project. Concentrations of arsenic, lead, and radium have since decreased to levels below the MCLs, and net alpha radiation was detected only sporadically in a few wells. Monitoring for arsenic, lead, radium, and net alpha radiation was discontinued in 2002 in accordance with provisions of the Ground Water Compliance Action Plan (GCAP) (DOE 2003).

Table 1 compares the maximum concentrations of the remaining site contaminants detected in June 2010 to the corresponding compliance goals for the mill tailings area. The compliance goals for cadmium, molybdenum, and uranium are UMTRA Project MCLs. The compliance goal for selenium (0.05 milligram per liter [mg/L]) is adopted from the U.S. Environmental Protection Agency (EPA) Safe Drinking Water Act as an ACL (the MCL is 0.01 mg/L). An ACL was established for selenium because selenium occurs naturally in groundwater beneath the site at levels above the MCL. There are no MCLs for manganese and sulfate. The compliance goal for manganese is the EPA Drinking Water Equivalent Level. This is a lifetime exposure concentration protective of adverse, non-cancer health effects; it assumes that all of the exposure to a contaminant is from drinking water (EPA 2004). The sulfate goal is equivalent to its average background concentration in local groundwater.

Contaminant of Concern	Compliance Goal (mg/L)	Compliance Goal Source	Maximum Concentration Observed in June 2010 (mg/L)
Cadmium	0.01	UMTRA Project MCL	0.042
Manganese	1.6	DWEL (EPA 2004)	5.5
Molybdenum	0.1	UMTRA Project MCL	0.09
Selenium	0.05	ACL (DOE 2002)	0.13
Sulfate	1,276	Average Background (DOE 2002)	2,200
Uranium	0.044	UMTRA Project MCL	1.3

Table 1. Current Groundwater Contaminants and Compliance Goals for the Mill Tailings Area

DWEL = Drinking Water Equivalent Level

Bedrock groundwater at the raffinate ponds area qualifies for supplemental standards on the basis of limited use groundwater. The groundwater in the bedrock is of limited use because of widespread, elevated concentrations of naturally occurring selenium. Selenium concentrations exceed the MCL at background monitoring well 0599 by a factor of nearly nine. Additional evidence of the natural presence of selenium beneath the raffinate ponds area is presented in Section 5.4 of the Site Observational Work Plan (DOE 2002). Because supplemental standards apply to groundwater in the raffinate ponds area, no numerical compliance goals have been established for that portion of the site.

Current monitoring of the Animas River verifies previous findings in the Baseline Risk Assessment (DOE 1995) that past milling operations have negligible effect on surface water quality. Historical results indicate that constituent concentrations adjacent to and downstream of the mill tailings area are indistinguishable from background river concentrations.

2.3 Surface Remediation Activities

The U.S. Department of Energy (DOE) began surface cleanup of the mill tailings and raffinate ponds areas in November 1986 to meet the EPA standards for radium in soil. A total of 2.5 million cubic yards of contaminated material was relocated to the Bodo Canyon disposal cell several miles southwest of the Durango site. Supplemental cleanup standards were applied to steep slopes of Smelter Mountain and two regions bordering the Animas River. In addition, a small lens of uranium ore was left in place at the mill tailings area below layers of slag along portions of the river. The slag deposits, which are 10 to 15 ft thick in some areas (including the location of well 0612), are associated with a lead smelter that operated on the site from 1880 to 1930. To restore the site, approximately 230,000 cubic yards of uncontaminated soil was backfilled, contoured, and seeded. Riprap was placed in some sensitive areas along the Animas River to prevent erosion. Remedial action was completed in May 1991.

2.4 Water and Land Use

The primary water source for the city of Durango is the Florida River upstream of its confluence with the Animas River. Additional water is withdrawn from the Animas River during highdemand periods (usually during the summer) from a location approximately 2 miles upstream of the mill tailings area. The Animas River bordering the mill tailings area of the Durango site is popular for seasonal boating and fishing. Development plans for the mill tailings area include municipal but not residential use (DOE 2002).

2.5 Institutional Controls

As part of the compliance strategy, public health will be protected at the mill tailings area during the natural flushing period through an environmental covenant between the State of Colorado and the City of Durango (landowner) that restricts access to contaminated alluvial groundwater. Additionally, deed restrictions (which serve as a notice to the public) for the mill tailings area prohibit access to groundwater without written permission from DOE and the Colorado Department of Public Health and Environment. Groundwater use in the raffinate ponds area is restricted in perpetuity through a deed restriction that also requires DOE's permission before use of groundwater for any purpose. The State of Colorado is currently in the process of trying to obtain a signed environmental covenant agreement for the raffinate ponds area.

3.0 Monitoring Program

Annual groundwater and surface water monitoring of the processing site will continue through the first 5 years following U.S. Nuclear Regulatory Commission concurrence with the GCAP (DOE 2003). Monitoring for cadmium at the mill tailings area will continue annually for the first 10 years following concurrence because of the greater uncertainty of this constituent to naturally flush within the allotted 100-year period established in 40 CFR 192. Monitoring data obtained through the initial 5-year period will measure the actual progress of natural flushing of the constituents listed in Table 1. After the 5-year annual monitoring period, the scope of subsequent monitoring will be addressed in a Long-Term Management Plan.

At the mill tailings area, monitoring wells 0612, 0617, 0630, 0631, 0633, 0634, 0635, and 0863 have been established as point-of-compliance (POC) wells that will be used to monitor the progress of natural flushing in groundwater in the alluvial aquifer (Figure 2). In accordance with provisions of the GCAP (DOE 2003), natural flushing for a given analyte is complete when its concentration no longer exceeds the compliance goal at the POC wells for three consecutive annual sampling events. Monitoring for that constituent may then be discontinued.

Surface water locations 0652, 0584, 0691, and 0586, located along the Animas River, will be monitored on schedule with groundwater monitoring to verify continued protection of the aquatic environment (Figure 2). The rationale and requirements for compliance monitoring in the mill tailings area are summarized in Table 2.

Sampling Location	Monitoring Purpose	Analytes	Location									
	Groundwater Monitoring											
0617, 0630, 0631, 0633, 0634, 0635	POC/verify natural flushing	Manganese Molybdenum Selenium Sulfate Uranium	On site									
0612, 0863	POC/verify natural flushing; verify cadmium flushing	Cadmium Manganese Molybdenum Selenium Sulfate Uranium	On site downgradient									
	Surface Water Monitor	ing										
0652	Surface water background	Codmium	Off site upstream									
0584, 0691	Verify no site-related increase above background	Molybdenum	Off site; site groundwater discharge area									
0586	Verify no site-related increase above background	Uranium	Off site; downstream of site groundwater discharge									

Table 2. Annual Groundwater and Surface Water Compliance Monitoring Requirements for the
Mill Tailings Area

Groundwater and surface water monitoring of the raffinate ponds area is being conducted only as a best management practice and no POC wells have been established. Monitoring requirements are summarized in Table 3.

Sampling Location	Monitoring Purpose	Analytes	Location
0879, 0594 (replaced 0880)	Monitor concentrations in groundwater in the shallow bedrock.	Selenium Uranium	On site
0598	Monitor concentrations in groundwater in the deep bedrock and Bodo Fault zone.	Selenium Uranium	On site
0607	Monitor concentrations in groundwater entering the site.	Selenium Uranium	On site
0884	Monitor off-site downgradient concentrations and migration.	Selenium Uranium	Off site downgradient
0588	Surface water quality entering the site.	Selenium Uranium	Off site upgradient
0654, 0656	Downgradient surface water concentrations.	Selenium Uranium	Off site downgradient

Table 3. Summary of Monitoring Requirements at the Raffinate Ponds Area

4.0 Results of 2010 Monitoring

Table 4 summarizes the amount of time the model predicts for natural flushing to achieve the compliance goals for cadmium, manganese, molybdenum, selenium, sulfate, and uranium in groundwater. The progress of each, based on water quality data through June 2010, is addressed separately in the following subsections. Important reference dates for comparing observed concentration trends to model-predicted trends include 1992 to 2010, when water quality was monitored after removal of the primary source of groundwater contamination, and June 2002, which was established as the groundwater model baseline condition (time zero) for contaminant transport. The predicted compliance times, listed in Table 4, are based on the predicted maximum average contaminate concentration and the probability that the standard will be exceeded. The compliance times (probability = 0) differ because the contaminants initially were not distributed evenly and vary in degree of contamination above the respective compliance goal, and because each contaminant varies in its mobility in groundwater.

Analyte	Compliance Goal (mg/L)	Predicted Compliance Time (years) ^ª	Predicted Compliance Date ^b
Cadmium	0.01	>100	>2102
Manganese	1.6	70	2072
Molybdenum	0.1	5	2007
Selenium	0.05	60	2062
Sulfate	1,276	100	2102
Uranium	0.044	, 90	2092

Table 4. Model-Predicted Groundwater Restoration Times for the Mill Tailings Area

^a Source: DOE 2002, Appendix G, Table 18.

^b Model time zero (baseline) is June 2002.

Plots of model predicted compliance times are based on single steady state deterministic simulations conducted for the GCAP (DOE 2003) and run again for this report. These plots continue to show mixed results; some 2010 sampling data show concentrations above those predicted by the model, while others are quite consistent with model predictions. Variation in concentrations in groundwater is to be expected on an annual basis, and the success of natural

flushing needs to be assessed over an extended period. Even with observed increases in concentrations for some of the constituents in 2010, linear trends of measured data since 1992 show that concentrations of all constituents, except cadmium at well 0612 and selenium at well 0633, are expected to naturally flush within the 100-year time frame allotted in 40 CFR 192.

4.1 Groundwater

4.1.1 Mill Tailings Area

Groundwater was sampled from the eight POC locations (Figure 2) and analyzed for constituents shown in Table 2. Sampling results for 2010 are provided in Appendix A and are discussed below by constituent.

4.1.1.1 Cadmium

Figure 4 is a map view of the site showing the concentration of cadmium in groundwater at the compliance wells in June 2010. Figure 5 shows observed cadmium concentrations versus time at the compliance wells since completion of remedial action in 1992. Historically, and in June 2010, cadmium levels exceeded the MCL at well 0612. The remaining monitoring wells contained only trace levels of this constituent. Groundwater modeling predicted a flushing period greater then 100 years for cadmium (Table 4). This result is not consistent with historical trending at well 0612, which if projected linearly from 1992 beyond June 2010 implies compliance for cadmium by about 2034 (Figure 6). Projecting this trend too far into the future might underestimate the actual restoration period because of transport processes that lead to a long tail in the predicted concentrations at the end of the process. This is commonly observed in nature and predicted by the solute transport models. Natural flushing of cadmium at the site and the observed net decrease in concentration over time. Since it is early in the 100-year natural flushing time frame, DOE will continue to monitor cadmium concentrations in groundwater and will reevaluate the strategy later, if required.



Figure 4. Distribution of Cadmium at the Mill Tailings Area in June 2010



Figure 5. Historical Cadmium Concentrations in Groundwater at the Mill Tailings Area



Durango Mill Tailings Process Site Cadmium Concentration Compliance Goal = 0.01 mg/L (MCL)

Figure 6. Predicted and Measured Cadmium Concentrations at the Mill Tailings Area

U.S. Department of Energy September 2010

4.1.1.2 Manganese

Figure 7 and Figure 8, respectively, illustrate the distribution of manganese concentrations in groundwater in June 2010 and the historical variations of manganese concentrations at the compliance wells. The June 2010 results are typical for manganese in that the compliance goal was exceeded only at well 0612 (Figure 8), where the concentration had increased from those observed in 2009. Projecting the linear trend of the observed concentration at well 0612 implies that natural flushing will meet the compliance goal at that location in about the year 2068, well within the 100-year time allotment (Figure 9). The net variation in the concentration of manganese observed through the relatively brief period since 2002 is consistent with the model prediction for this location. Well 0612 is very close to the downgradient discharge boundary of the aquifer; therefore contaminant migration from that area will not affect other regions of the aquifer. The flushing period for well 0612 is not exceeded at any other location.

4.1.1.3 Molybdenum

Molybdenum concentrations in June 2010 were less than the compliance goal of 0.1 mg/L at all locations (Figure 10). Concentrations at well 0612, which remained slightly above the compliance goal in 2008, declined slightly to 0.096 mg/L in 2009 and declined slightly more to 0.090 mg/L in 2010 (Figure 10 and Figure 11). If all wells remain below the standard for one more sampling round, compliance will be achieved, and monitoring for molybdenum can be discontinued. The linear trend of observed concentrations at well 0612 forecasts molybdenum flushing to be complete in 2010 (Figure 12); observations bear out this prediction.

4.1.1.4 Selenium

Figure 13 shows, in map view; that the compliance goal for selenium (0.05 mg/L) was met at all wells except well 0633. Selenium levels in well 0633, which had fallen below the standard in 2009, rebounded above the standard in 2010 (Figure 14). The model predicted that selenium concentrations would drop below the ACL by 2024 at this location (Figure 15).

Concentrations in well 0633 declined from 0.083 mg/L in 2008 to 0.048 mg/L in 2009 and increased to 0.13 mg/L in 2010 (Figure 14). Well 0633 is screened 90 percent in Mancos Shale, a recognized source of readily mobilized selenium (DOE 2002). The low-level selenium contamination at the site may in part be site-related; however, some contribution from natural sources is likely, as evidenced by concentrations greater than 0.01 mg/L at background well 0622 (Figure 14).

4.1.1.5 Sulfate

Sulfate concentrations that exceed background levels are related to the former use of sulfuric acid in the milling process. In June 2010, sulfate exceeded the average background concentration at each compliance well except wells 0631 and 0863 (Figure 16 and Figure 17). Observed concentrations since 1992 have fluctuated considerably at most compliance wells and have generally not shown trending. However, projecting best-fit lines to the data reveals that sulfate flushing will be complete at most locations by about 2092 (Figure 18). Linear trend projection of data from well 0612, 0617, and 0633 indicates that concentrations at these locations will meet the compliance goal by about 2015, 2080, and 2058, respectively.



Figure 7. Distribution of Manganese at the Mill Tailings Area in June 2010

Durango Mill Tailings Process Site Manganese Concentration Compliance Goal = 1.6 mg/L (DWEL)



Figure 8. Historical Concentrations of Manganese at the Mill Tailings Area



Figure 9. Predicted and Measured Manganese Concentrations at the Mill Tailings Area



Figure 10. Distribution of Molybdenum at the Mill Tailings Area in June 2010







Figure 12. Predicted and Measured Molybdenum Concentrations at the Mill Tailings Area



Figure 13. Distribution of Selenium at the Mill Tailings Area in June 2010

Durango Mill Tallings Process Site Selenium Concentration Compliance Goal = 0.05 mg/L (ACL)







Durango Mill Tailings Process Site Selenium Concentration Compliance Goal = 0.05 mg/L (ACL)

Figure 15. Predicted and Measured Selenium Concentrations at the Mill Tailings Area



Figure 16. Distribution of Sulfate at the Mill Tailings Area in June 2010

Durango Mill Tailings Process Site Sulfate Concentration Compliance Goal = 1,276 mg/L (Background)



Figure 17. Historical Concentrations of Sulfate at the Mill Tailings Area



Durango Mill Tailings Process Site Sulfate Concentration Compliance Goal = 1,276 mg/L (Background)

Figure 18. Predicted and Measured Sulfate Concentrations at the Mill Tailings Area

4.1.1.6 Uranium

The uranium compliance goal was exceeded at all locations except wells 0635 and 0863 in June 2010 (Figure 19 and Figure 20). This is consistent with previous monitoring results. The two wells with the historically largest uranium concentrations (0612 and 0633) have shown a generally decreasing concentration trend since source removal. Groundwater model predictions indicate that sitewide uranium flushing will be complete by about 90 years after June 2002 (Table 4). To date, observed concentrations at the two wells that have the highest uranium concentrations (wells 0612 and 0633) are in general agreement with the model results (Figure 21). These wells are widely separated in the aquifer. The predicted flushing periods for these two wells are less than the predicted sitewide flushing time because the last area to flush in the site model is south of the downgradient-most monitoring well (well 0612). Linear projection of the observed concentration at well 0633 implies sitewide uranium flushing by about 2030 (Figure 21). The model predicts initial rates of uranium flushing at this well that track observed concentrations, followed by a period of less rapid flushing and marginal levels of contamination (concentration tailing) until the goal is attained. Similar tailing of concentrations is predicted by the model at well 0612. Uranium concentrations may remain slightly above the compliance goal during the period of reduced flushing at these two wells.

4.1.2 Raffinate Ponds Area

Groundwater in the raffinate ponds area is being monitored as a best management practice. Bedrock groundwater at the raffinate ponds area qualifies for supplemental standards on the basis of limited use groundwater due to widespread elevated concentrations of naturally occurring selenium. Because there are naturally occurring sources of both selenium and uranium in the area, groundwater is not expected to naturally flush. Therefore, no modeling was done for the raffinate ponds area.

Groundwater was sampled from five well locations in the monitoring network (Figure 3) in 2010 and analyzed for uranium and selenium. Sampling results for 2010 are provided in Appendix A and are discussed below by constituent.

4.1.2.1 Selenium

Figure 22 shows historical concentrations of selenium since completion of remedial action in 1992. Significant increases and decreases in selenium distribution in groundwater occurred at several wells from 2001 to 2004 concentrations have since leveled off. It is not clear what caused the past selenium behavior; accordingly the future disposition of selenium in the area's groundwater cannot be reliably predicted.

4.1.2.2 Uranium

Historical concentrations of uranium in the raffinate ponds area are shown in Figure 23. Concentrations of uranium in most wells have fluctuated over the last several years showing neither increasing nor decreasing trends. Supplemental standards were applied to soils in the raffinate ponds area (DOE 2002), and those soils may contain residual uranium contamination that influences groundwater quality. Subpile soil analyses indicate the presence of uranium in soils remaining at the site (DOE 2002).



Figure 19. Distribution of Uranium at the Mill Tailings Area in June 2010

Durango Mill Tailings Process Site Uranium Concentration Compliance Goal = 0.044 mg/L (MCL)









U.S. Department of Energy September 2010 Durango Mill Tailings Process Site Uranium Concentration Compliance Goal = 0.044 mg/L (MCL)







Durango Raffinate Pond Process Site Uranium Concentration



4.2 Surface Water

Surface water was sampled from six locations in the Animas River adjacent to both the mill tailings and raffinate ponds areas during June 2010 and analyzed for cadmium, molybdenum, selenium, and uranium (Figure 2 and Figure 3, Table 2 and Table 3). In addition, a sample was collected from South Creek (location 0588), upgradient from the raffinate ponds area, to assess the quality of water entering the raffinate ponds area from the west. The sample from South Creek (0.0008 and 0.018 mg/L, respectively) had levels of selenium and uranium that were higher than those from any other surface location. However, discharge of the creek to the Animas River had no discernible impact on concentrations of selenium and uranium at the river sampling location (0654). Concentrations of constituents at all locations along the river were well below the respective compliance goals and remain indistinguishable from background levels (Appendix B).

5.0 Natural Flushing Assessment

As of June 2010, the observed rate of contaminant flushing is generally consistent with groundwater model predictions, given that the validation period to date (June 2002 to June 2010) is short compared to predicted flushing periods (60 to 100 years) for the various contaminants. Only cadmium was identified in the modeling as potentially incapable of flushing to acceptable levels within 100 years. However, at the single location (well 0612) where cadmium is present above the compliance goal of 0.01 mg/L, concentrations have decreased more rapidly than predicted by the model. The linear trend suggests the compliance goal for cadmium will be reached by about 2034. With the possible exception of sulfate and selenium, modeling predictions and concentration trends imply that the compliance goals for remaining constituents will likely be attained within 100 years, suggesting that natural flushing remains a valid compliance strategy for these constituents as well. The impact on surface water quality from site-related contamination remains negligible.

6.0 Conclusions

Assessment of the June 2010 water sampling data and concentration trends at the mill tailings area on the Durango site, particularly since completion of source removal, indicate that natural flushing is leading to reduced contaminant concentrations in groundwater at the site. Note that it is too early in the 100-year natural flushing time frame to draw definitive conclusions.

On the basis of findings presented in this report, the following recommendations are made with regard to monitoring at the mill tailings area:

- Continued monitoring of groundwater and surface water quality at the currently established compliance network locations.
- Analysis of all water samples for the same suite of constituents that have historically been monitored to assist in evaluating contaminant migration trends.

It is also recommended that best management practice monitoring of the raffinate ponds area continue for the foreseeable future.

7.0 References

DOE (U.S. Department of Energy), 1995. Baseline Risk Assessment of Ground Water Contamination at the Uranium Mill Tailings Site Near Durango, Colorado, Rev. 1, DOE/AL/62350–175, UMTRA Project Office, Albuquerque, New Mexico, September.

DOE (U.S. Department of Energy), 2002. *Site Observational Work Plan for the Durango, Colorado, UMTRA Project Site*, GJO-2001-272-TAR/MAC-GW DUR 1.1, Grand Junction Office, Grand Junction, Colorado, January.

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EPA (U.S. Environmental Protection Agency), 2004. 2004 Edition of the Drinking Water Standards and Health Advisories, EPA822-R-04-005, Office of Water, Washington, DC.

Appendix A

Durango Mill Tailings Process Site (DUR01) Groundwater Quality Data

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GENERAL WATER QUALITY DATA BY P	ARAMETER	(USEE205) F	OR SITE	DUR01,	Durango Mill	Tailings	Process Si	te
REPORT DATE: 9/9/2010 3:14 pm							÷	

PARAMETER	UNITS	LOCATION CODE	LOC TYPE, SUBTYPE	SAMPI DATE	.E: ID	DEPTH RANGE (FT BLS)	RESULT	QU LAB		RS: I	DETECTION LIMIT	UN- CERTAINTY
Alkalinity, Total (As CaCO3)) mg/L	0612	WL	06/08/2010	0001	37.41 - 57.41	419		F	#	-	-
	mg/L	0617	WL	06/09/2010	N001	14.00 - 29.00	450		F	#	• • .	- '
	mġ/L	0630	WL	06/08/2010	N001	28.30 - 38.30	292		F	#	-	· •
	mg/L	0631	WL	06/09/2010	, N001:	6.00 - 16.00	379		F	#	-	
	mg/L	0633	WL.	06/09/2010	N001	4.00 - 14.00	283		F	'#	-	• ·
	mg/L	0634	WL	06/08/2010	N001	8.00 - 18.00	484		FQ	#	-	· -
	mg/L	0635	WL	06/08/2010	N001	5.50 - 15.50	403	• •	F	#	-	-
	mg/L	0863	WL	06/08/2010	N001	58.00 - 67.50	531		F	, #	-	-
Cadmium	mg/L	0612	WL	06/08/2010	0001	37.41 - 57.41	0.042		F	#	0.00023	· _ ·
•	mg/L	0635	WL	06/08/2010	N002	5.50 - 15.50	0.00017		F	#	1.2E-05	-
	mg/L	0863	WL	06/08/2010	N001	58.00 - 67.50	0.00007		F.	#	1.2E-05	-
Chloride	mg/L	0612	, WL	06/08/2010	0001	37.41 - 57.41	200		F	#	10	-
Manganese	mg/L	0612	WL	06/08/2010	0001	37.41 - 57.41	5.500		F	#	5.4E-05	-
	mg/L	0617	WL	06/09/2010	N001	14.00 - 29.00	1.300		F	#	5.4E-05	•
	mg/L	0630	WL	06/08/2010	N001	28.30 - 38.30	0.810		F	#	5.4E-05	-
	.mg/L	0631	WL	06/09/2010	N001	6.00 - 16.00	0.470		F	#	5.4E-05	-
	mg/L	0633	WL	06/09/2010	N001	4.00 - 14.00	0.0046 1	в	F.	, #	5.4E-05	· -
	mg/L	0634	· WL	06/08/2010	N001	8.00 - 18.00	0.070		FQ	#	5.4E-05	-
	mg/L	0635	WL	06/08/2010	N001	5.50 - 15.50	0.086		F	#	5.4E-05	-
,	mg/L	0635	WL	06/08/2010	N002	5.50 - 15.50	0.085		F	#	5.4E-05	•
	mg/L	0863	WL	06/08/2010	Ň001	58.00 - 67.50	0.110		F	#	5.4E-05	-
Molybdenum	mg/L	0612	WL	06/08/2010	0001	37.41 - 57.41	0.090		F	#	0.00064	•
•	mg/L	0617	WL	.06/09/2010	N001	14.00 - 29.00	0.0017		F	#	6.4E-05	-
· ·	.mg/L	0630	WL .	06/08/2010	N001	28.30 - 38.30	0.0038		F	#	0.00016	-
	mg/L	0631	WL	06/09/2010	N001	6.00 - 16.00	0.0054	-	,F	# .	6.4E-05	-

PARAMETER	UNITS	LOCATION CODE	LOC TYPE, SUBTYPE	. SAMPL DATE	.E: ID	DEPTH RANGE (FT BLS)	RESULT	QUALIFIER	S: C QA		UN- CERTAINTY
Molybdenum	mg/L	0633	. WL	06/09/2010	N001	4.00 - 14.00	0.0055	F	#	0.00032	
	mg/L	0634	WL	06/08/2010	N001	8.00 - 18.00	0.0015	FQ	#	3.2E-05	-
· .	mg/L	0635	WL	06/08/2010	N001	5.50 - 15.50	0.0015	F	#	3.2E-05	-
	mg/L	0635	WL	06/08/2010	N002	5.50 - 15.50	0.0015	. F	#	3.2E-05	-
	mg/L	0863	WL	06/08/2010	N001	58.00 - 67.50	0.00056	F	. #	3.2E-05	-
Oxidation Reduction Potential	mV	0612	WL*	06/08/2010	N001	37.41 - 57.41	101.2	F	#	-	
	mV	0617	WL	06/09/2010	N001	14.00 - 29.00	-71.5	F	#	-	-
	mV ⁻	0630	WL	06/08/2010	N001	28.30 - 38.30	6.3	F	#	-	- 1
	mV	0631	WL	06/09/2010	N001	6.00 - 16.00	-94.5	F	#	-	-
	mV	0633	WL	06/09/2010	N001	4.00 - 14.00	-22.0	F	#	. <u>-</u>	-
	mV	0634	WL	06/08/2010	N001	8.00 - 18.00	183.0	FQ	#	-	-
	mV	0635	WL	06/08/2010	N001	5.50 - 15.50	-16.4	, F	#	•	-
	mV	0863	WL	06/08/2010	N001	58.00 - 67.50	-20.1	F	#	-	- ,
pH.	s.u	0612	ŴL.	06/08/2010	N001	37.41 - 57.41	6.71	F	#	· -	-
	s.u.	0617	WL	06/09/2010	N001	14.00 - 29.00	6.90	, F	#	-	•
	s.u.	0630	WL	06/08/2010	N001	28.30 - 38.30	6.95	F	#	۰ -	-
• •	s.u.	0631	WL	06/09/2010	N001	6.00 - 16.00	7.27	F	#	-	-
	s.u.	0633	WL	06/09/2010	N001	4.00 - 14.00	7.34	F	#	-	•
	s.u.	0634	WL	06/08/2010	N001	8.00 - 18.00	6.91	FQ	#	-	-
	s.u.	0635	WL	06/08/2010	N001	5.50 - 15.50	6.84	F	#	-	-
	s.u.	0863	WL	06/08/2010	N001	58.00 - 67.50	6.99	F	#	-	-
Selenium	mg/L	0612	ŴL	06/08/2010	0001	37.41 - 57.41	0.00081	F	#	0.00016	
	mg/L	0617	WL	06/09/2010	N001	14.00 - 29.00	0.0015	F.	#	6.5E-05	-
	mg/L	0630	WL	06/08/2010	N001	28.30 - 38.30	0.020	F	#	0.00016	-
	mg/L	0631	·WL	06/09/2010	N001	6.00 - 16.00	0.0013	F	#	6.5E-05	-
							<u>, in the second second</u>	·····			Dago 2

PARAMETER	UNITS	LOCATION CODE	LOC TYPE, SUBTYPE	SAMPL DATE	E: ID	DEPTH RANGE (FT BLS)	RESULT	QUALIFIERS: LAB DATA QA		UN- CERTAINTY
Selenium	mg/L	0633	WL	06/09/2010	N001	4.00 - 14.00	0.130	F	# 0.00032	-
	mg/L	0634	WL	06/08/2010	N001	8.00 - 18.00	0.0015	FQ .	# 3.2E-05	-
	mg/L	0635	WL	06/08/2010	N001	5.50 - 15.50	0.0059	F	# 3.2E-05	•
	mg/L	0635	WL	06/08/2010	N002	5.50 - 15.50	0.0072	F	# 3.2E-05	-
	mg/L	0863	WL	06/08/2010	N001	58.00 - 67.50	0.00024	F	# 3.2E-05	-
Specific Conductance	umhos/cm	0612	WL	06/08/2010	N001	37.41 - 57.41	4369	F	# -	-
	umhos/cm	0617	WL	06/09/2010	N001	14.00 - 29.00	3670	F	# -	-
	umhos/cm	0630	WL	06/08/2010	N001	28.30 - 38.30	3264	F	# -	-
	umhos/cm	0631	WL	06/09/2010	N001	6.00 - 16.00	2123	F	# -`	-
	umhos/cm	0633	WL	06/09/2010	N001	4.00 - 14.00	4186	F	# -	-
	umhos/cm	0634	WL	06/08/2010	N001	8.00 - 18.00	4694	FQ	# -	-
	umhos/cm	0635	WL	06/08/2010	N001	5.50 - 15.50	2772	F	# -	• ·
	umhos/cm	0863	WL	06/08/2010	N001	58.00 - 67.50	2299	F	# -	-
Sulfate	mg/L	0612	WL	06/08/2010	0001	37.41 - 57.41	1800	F	# 25	-
	mg/L	0617	WL	06/09/2010	N001	14.00 - 29.00	1900	F	# 25	-
•	mg/L	0630	WL	06/08/2010	N001	28.30 - 38.30	1600	F	# 25	-
	mg/L	0631	WL	06/09/2010	N001	6.00 - 16.00	630	F	# 10	-
	mg/L	0633	WL	06/09/2010	N001	4.00 - 14.00	2200	F	# 25	-
	mg/L	0634	WL	06/08/2010	N001	8.00 - 18.00	2200	FQ	# 25	• '
. '	mg/L	0635	WL	06/08/2010	N001	5.50 - 15.50	1300	F	# 10	-
	mg/L	0635	· WL	06/08/2010	N002	5.50 - 15.50	1300	F	# 10	-
	mg/L	0863	WL	06/08/2010	N001	58.00 - 67.50	640	F	# 10	-
Temperature	С	0612	WL	06/08/2010	N001	37.41 - 57.41	14.34	F	# -	-
	С	0617	WL	06/09/2010	N001	14.00 - 29.00	12.02	F	# -	-
	C	0630	WL	06/08/2010	N001	28.30 - 38.30	14.72	F	# -	-

PARAMETER	UNITS	LOCATION CODE	LOC TYPE, SUBTYPE	SAMPL DATE	.E: ID	DEPTH RANGE (FT BLS)	RESULT	QUALIFIEF LAB DATA	RS: QA	DETECTION LIMIT	UN- CERTAINTY
Temperature	С	0631	WL	06/09/2010	N001	6.00 - 16.00	12.42	F	#	-	•
	С	0633	WL	06/09/2010	N001	4.00 - 14.00	13.98	F	·. #	-	-
	С	0634	WL	06/08/2010	N001	8.00 - 18.00	14.32	FQ	#	-	-
	С	0635	WL	06/08/2010	N001	5.50 - 15.50	14.09	F	#	-	-
	С	0863	WL	06/08/2010	N001	58.00 - 67.50	13.29	F	#	, -	-
Turbidity	NTU	0612	WL	06/08/2010	N001	37.41 - 57.41	36.8	F	#	-	-
	NTU	0617	WL	06/09/2010	N001	14.00 - 29.00	5.65	F	#	-	-
	NTU	0630	WL	06/08/2010	N001	28.30 - 38.30	8.23	F	#	-	-
	NTU	0631	WL	06/09/2010	N001	6.00 - 16.00	3.83	F	#	-	-
	NTU	· 0633 ·	WL	06/09/2010	N001	4.00 - 14.00	5.12	F	#	-	
	NTU	0634	WL	06/08/2010	N001	8.00 - 18.00	3.46	FQ	#	-	-
	NTU	0635	WL	06/08/2010	N001	5.50 - 15.50	6.35	F	#	-	-
	· NTU	0863	WL	06/08/2010	N001	58.00 - 67.50	2.98	F	#	-	-
Uranium	mg/L	0612	WL	06/08/2010	0001	37.41 - 57.41	1.300	F	#	5.8E-05	
	mg/L	0617	WL	06/09/2010	N001	14.00 - 29.00	0.120	F ·	#	5.8E-06	-
	mg/L	0630	WL	06/08/2010	N001	28.30 - 38.30	0.250	F	#	1.5E-05.	-
	mg/L	0631	WL	06/09/2010	N001	6.00 - 16.00	0.160	·F	#	5.8E-06	· -
	mg/L	0633	WL	06/09/2010	N001	4.00 - 14.00	0.480	F	#	2.9E-05	-
	mg/L	0634	, WL	06/08/2010	N001	8.00 - 18.00	0.087	· FQ	#	2.9E-06	-
	mg/L	0635	WL	06/08/2010	N001	5.50 - 15.50	0.011	۴	#	2.9E-06	-
	mg/L	0635	WL	06/08/2010	N002	5.50 - 15.50	0.011	F	#	2.9E-06	
	mg/L	0863	WL	06/08/2010	N001	58.00 - 67.50	0.00009	F	#	2.9E-06	-

PARAMETER	UNITS	LOCATION I CODE	LOC TYPE, SUBTYPE	SAMPL DATE	.E: ID	DEPTH RANGE (FT BLS)		RESULT	QI LAB	UALI 3 D/	IFIERS: ATA Q	DETECTION A LIMIT	UN- CERTAINTY
RECORDS: SELECTED data_validat	FROM USEE200 \ ion_qualifiers NOT	VHERE site_code LIKE '%R%' ANI	≘='DUR01' AND D data_validatio	location_cod	e in('0612 IOT LIKE	','0617','0630','0631','0633 '%X%') AND DATE_SAM	','0634 PLED	1','0635','0863') A between #1/1/20	ND (0	data and #	validatio #12/31/20	n_qualifiers IS NUI 010#	LOR
SAMPLE ID CODES: 00	0X = Filtered samp	ole. N00X = Unf	iltered sample.	X = replicate	e number.						-		
LOCATION TYPES: WL	WELL		•										
LOCATION SUBTYPES:													
LAB QUALIFIERS:						•.							•
 Replicate analysis r Correlation coefficie Result above upper A TIC is a suspected 	not within control lin ent for MSA < 0.999 detection limit. aldol-condensation	nits. 5. product.											
B Inorganic: Result is	s between the IDL a	and CRDL. Orgai	nic & Radiochei	mistry: Analyt	e also fou	nd in method blank.		÷					
D Analyte determined	in diluted sample				-								
E Inorganic: Estimate	e value because of	interference, see	case narrative.	Organic: An	alvte exce	eded calibration range of	the G	C-MS.					
H Holding time expire	d, value suspect.	•		- 5		5							
I Increased detection	limit due to requir	ed dilution.				,						•	
J Estimated											• .		
M GFAA duplicate inje	ection precision not	t met.											
N Inorganic or radioch	nemical: Spike sar	nple recovery not	within control li	mits. Organic	: Tentativ	ely identified compund (TI	C).						
P > 25% difference in	detected pesticide	or Arochlor conc	entrations betw	een 2 column	s.								
S Result determined	by method of stand	ard addition (MSA	A).										
U Analytical result be	ow detection limit.												
W Post-digestion spike	e outside control lir	nits while sample	absorbance < \$	50% of analyti	cal spike	absorbance.							
X Laboratory defined	(USEPA CLP orga	nic) qualifier, see	case narrative.										
Y Laboratory defined	(USEPA CLP orga	nic) qualifier, see	case narrative.									•	
Z Laboratory defined	(USEPA CLP orga	nic) qualifier, see	case narrative.										
DATA QUALIFIERS:													
F Low flow sampling	method used.		G Possib	le grout conta	mination,	pH > 9.	J	Estimated value	э.				
L Less than 3 bore vo	olumes purged prio	r to sampling	N Presun analyte	nptive evidence is "tentatively	e that and identified	alyte is present. The I".	Q	Qualitative resu	ilt due	e to s	ampling	technique	
R Unusable result.			U Param	eter analyzed	for but wa	as not detected.	х	Location is unde	efine	d.			
QA QUALIFIER: # = vali	dated according to	Quality Assurance	ce guidelines.										
	-												

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GENERAL WATER QUALITY DATA BY PARAMETER (USEE205) FOR SITE DUR02, Durango Raffinate Pond Process Site REPORT DATE: 9/9/2010 3:57 pm

PARAMETER	UNITS	LOCATION CODE	LOC TYPE, SUBTYPE	SAMPI DATE	.E: ID	DEPTH RANGE (FT BLS)	RESULT	QUALIFIER LAB DATA	S: [QA	DETECTION LIMIT	UN- CERTAINTY
Alkalinity, Total (As CaCO3)	mg/L	0594	WL	06/10/2010	N001	8.50 - 38.50	328	FQ	#	-	-
	mg/L	0598	WL	06/10/2010	N001	66.20 - 96.20	359	F	#		-
,	mg/L	0607	WL	06/10/2010	N001	35.00 - 55.00	324	FQ	#	-	-
	mg/L	0879	WL	06/10/2010	N001	27.00 - 36.90	362	F	· #	-	-
	mg/L	0884	WL	06/10/2010	N001	36.50 - 46.50	437	F	#		·
Oxidation Reduction Potential	mV	0594	WL	06/10/2010	N001	8.50 - 38.50	31.6	FQ	#	• -	-
	mV	0598	WL	06/10/2010	N001	66.20 - 96.20	28.4	F	#	-	-
	mV	0607	WL	06/10/2010	N001	35.00 - 55.00	18.7	FQ	#	-	· -
	mV	0879	WL	06/10/2010	N001	27.00 - 36.90	72.7	F	#	·_	-
	mV	0884	WL	06/10/2010	N001	36.50 ÷ 46.50	60.2	,F	•#	-	-
рН	s.u.	0594	WL	06/10/2010	N001	8.50 - 38.50	6.96	FQ	#	-	•
	s.u.	0598	ŴL	06/10/2010	N001	66.20 - 96.20	6.99	F	#	-	-
	s.u.	0607	WL	06/10/2010	N001	35.00 - 55.00	6.97	FQ	#	-	-
	s.u.	0879	WL	06/10/2010	N001	27.00 - 36.90	6.86	F	#	-	-
	s.u.	0884	WL	06/10/2010	N001	36.50 - 46.50	7.06	F	#	-	-
Selenium	mg/L	0594	WL	06/10/2010	N001	8.50 - 38.50	0.0069	FQ	#	3.2E-05	-
	mg/L	0598	WL	06/10/2010	N001	66.20 - 96.20	0.370	F	#	3.2E-05	-
	mg/L	0607	WL	06/10/2010	N001	35.00 - 55.00	0.470	FQ	#	3.2E-05	-
	.mg/L	0879	WL	06/10/2010	N001	27.00 - 36.90	0.030	F	#	3.2E-05	-
	mg/L	0884	WL	06/10/2010	N001	36.50 - 46.50	1.200	F	#	0.0032	-
	mg/L	0884	WĽ	06/10/2010	N002	36.50 - 46.50	1.200	F	#	0.0032	-
Specific Conductance	umhos/cm	0594	WL	06/10/2010	N001	8.50 - 38.50	4117	FQ	#	-	-
, 1	umhos/cm	0598	WL	06/10/2010	N001	66.20 - 96.20	8392	F	#	- `	-
	umber / am	0607	10/1	06/10/2010	NI001	25 00 55 00	2097	50	#		

		LOCATION	LOC TYPE,	SAMPL	-E:	DEPTH RANGE		QUALIFIERS:	[
			JUBITE						<u> </u>		
Specific Conductance	umhos/cm	0879	WL.	06/10/2010	N001	27.00 - 36.90	8444	F	# '	•	-
	umhos/cm	0884	WL	06/10/2010	N001	36.50 - 46.50	5385	F	#		-
Temperature	с	0594	WL	06/10/2010	N001	8.50 - 38.50	12.55	FQ	#	-	-
	С	0598	WL	06/10/2010	N001	66.20 - 96.20	13.21	F	#	-	-
. •	С	0607	WL	06/10/2010	N001	35.00 - 55.00	14.24	FQ	#	• -	-
	С	0879	WL	06/10/2010	N001	27.00 - 36.90	11.97	F	#	-	• ·
	С	0884	WL	06/10/2010	N001	36.50 - 46.50	13.09	F	#	-	-
Turbidity	NTU	0594	WL	06/10/2010	N001	8.50 - 38.50	7.01	FQ	#	-	•
	NTU	0598	WL.	06/10/2010	N001	66.20 - 96.20	4.95	F	#	-	-
	NTU	0607	WL	06/10/2010	N001	35.00 - 55.00	8.99	FQ	#	-	-
	NTU	0879	WL	06/10/2010	N001	27.00 - 36.90	6.87	F	#	-	-
•	NTU	0884	WL	06/10/2010	N001	36.50 - 46.50	1.54	F	#	-	-
Uranium	mg/L	0594	WL	06/10/2010	N001	8.50 - 38.50	0.100	FQ	#	2.9E-06	
	mg/L	0598	WL.	06/10/2010	N001	66.20 - 96.20	0.096	F	#	2.9E-06	-
	mg/L	0607	WL	06/10/2010	N001	35.00 - 55.00	0.0023	FQ	#	2.9E-06	-
	mg/L	0879	WL	06/10/2010	N001	27.00 - 36.90	0.086	· F	#	2.9E-06	-
	mg/L	0884	WL	06/10/2010	N001	36.50 - 46.50	0.170	F	#	0.00029	-
	mg/L	0884	WL.	06/10/2010	N002	36.50 - 46.50	0.170	F	#	0.00029	-

GENERAL WATER QUALITY DATA BY PARAMETER (USEE205) FOR SITE DUR02, Durango Raffinate Pond Process Site REPORT DATE: 9/9/2010 3:57 pm

GENERAL WATER QUALITY DATA BY PARAMETER (USEE205) FOR SITE DUR02, Durango Raffinate Pond Process Site REPORT DATE: 9/9/2010 3:57 pm

LOCATION L PARAMETER UNITS CODE S	.OC TYPE, SUBTYPE	SAMP DATE	PLE: ID	DEPTH RANGE (FT BLS)		RESULT	QUALIFIERS	S: DE QA	TECTION LIMIT	UN- CERTAINTY
RECORDS: SELECTED FROM USEE200 WHERE site_code LIKE '%R%' AND data_validation_qualifiers NOT	='DUR02' AN LIKE '%X%	D location_co	de in('0594','0 _SAMPLED be	598','0607','0879','0884 tween #1/1/2010# and	l') ANE #12/3	D (data_validati 31/2010#	ion_qualifiers IS NU	LL OR d	ata_validatio	on_qualifiers NOT
SAMPLE ID CODES: 000X = Filtered sample. N00X = Unfil	Itered sample	e. X = replica	ite number.							
LOCATION TYPES: WL WELL										
LOCATION SUBTYPES:										
 Replicate analysis not within control limits. Correlation coefficient for MSA < 0.995. 										
Result above upper detection limit.A TIC is a suspected aldol-condensation product.		· .				·	. •			
 B Inorganic: Result is between the IDL and CRDL. Organ C Pesticide result confirmed by GC-MS. C Applyte determined in diluted completion 	iic & Radioch	emistry: Analy	yte also found	in method blank.			• .			х ,
 E Inorganic: Estimate value because of interference, see in H Holding time expired, value suspect. 	case narrativ	re. Organic: A	nalyte exceed	ed calibration range of	the G	C-MS.				
I Increased detection limit due to required dilution. J Estimated										
M GFAA duplicate injection precision not met.										
N Inorganic or radiochemical: Spike sample recovery not v	within contro	l limits. Organi twoon 2 colum	ic: Tentatively	identified compund (T	IC).					
S Result determined by method of standard addition (MSA		ween 2 colum	115.							
U Analytical result below detection limit.	.,.			·			•			
W Post-digestion spike outside control limits while sample a	absorbance	< 50% of analy	rtical spike abs	orbance.						•
X Laboratory defined (USEPA CLP organic) qualifier, see	case narrativ	e.								
Y Laboratory defined (USEPA CLP organic) qualifier, see	case narrativ	e.					~			
Z Laboratory defined (USEPA CLP organic) qualifier, see	case narrativ	e.		· · ·				•		
DATA QUALIFIERS:										
F Low flow sampling method used.	G Poss	ible grout cont	amination, pH	> 9.	J.	Estimated va	lue.			
L Less than 3 bore volumes purged prior to sampling.	N Pres analy	umptive evider /te is "tentative	nce that analyted	e is present. The	Q	Qualitative re	esult due to samplin	g techniq	ue	
R Unusable result.	U Para	meter analyzed	d for but was r	ot detected.	х	Location is ur	ndefined.		· .	
QA QUALIFIER: # = validated according to Quality Assurance	e guidelines.									

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Appendix **B**

Durango Mill Tailings Process Site (DUR01) Surface Water Quality Data

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GENERAL WATER QUALITY DATA BY PARAMETER	(USEE205) FOR SITE	DUR01,	Durango Mill	Tailings Proc	cess Site
REPORT DATE: 9/9/2010 3:54 pm					

PARAMETER	UNITS	LOCATION	LOC TYPE, SUBTYPE	SAMPL DATE	.E: ID	DEPTH RANGE (FT BLS)	RESULT	QUA LAB		:S: QA	DETECTION LIMIT	UN- CERTAINTY
Alkalinity, Total (As CaCO3)	mg/L	0584	SL.	06/08/2010	0001	0.00 - 0.00	41			#	-	-
	mg/L	0586	SL	06/09/2010	0001	0.00 - 0.00	73			#	-	-
	mg/L	0652	SL	06/09/2010	0001	0.00 - 0.00	39			#	-	-
	mg/L	0691	SL	06/08/2010	0001	0.00 - 0.00	44			#	-	-
Cadmium	mg/L	0584	SL	06/08/2010	0001	0.00 - 0.00	0.00034		J	#	1.2E-05	-
	mg/L	0586	SL	06/09/2010	0001	0.00 - 0.00	0.00014		J	#	1.2E-05	•
	mg/L	0652	SL	06/09/2010	0001	0.00 - 0.00	0.00023		J	#	1.2E-05	-
· .	mg/L	0691	SL	06/08/2010	0001	0.00 - 0.00	0.00018		J	#	1.2E-05	-
Molybdenum	mg/L	0584	SL	06/08/2010	0001	0.00 - 0.00	0.00044			#	3.2E-05	•
· ·	mg/L	0586	SL	06/09/2010	0001	0.00 - 0.00	0.00046			, #	3.2E-05	-
	mg/L	0652	SL	06/09/2010	0001	0.00 - 0.00	0.00041			#	3.2E-05	-
	mg/L	0691	SL	06/08/2010	0001	0.00 - 0.00	0.00043			#	3.2E-05	· •
Oxidation Reduction Potential	mV	0584	SL	06/08/2010	N001	0.00 - 0.00	-12.1			#	-	-
•	mV	0586	SL	06/09/2010	N001	0.00 - 0.00	31			#	-	-
	mV	0652	SL	06/09/2010	N001	0.00 - 0.00	-92			<u>,</u> #	-	•
	mV	0691	SL	06/08/2010	N001	0.00 - 0.00	-30.9			#	-	. –
pH	s.u.	0584	SL	06/08/2010	N001	0.00 - 0.00	7.99			#	- ·	-
	s.u.	0586	SĽ	06/09/2010	N001	0.00 - 0.00	7.79			#	-	-
·	s.u.	0652	SL	06/09/2010	N001	0.00 - 0.00	7.90			#	· -	•
	s.u.	0691	SL	06/08/2010	N001	0.00 - 0.00	7.80			#	-	
Selenium	mg/L	0584	SL	06/08/2010	0001	0.00 - 0.00	0.00006	в	J	#	3.2E-05	-
	mg/L	0586	SL	06/09/2010	0001	0.00 - 0.00	0.00018		J	#	3.2E-05	-
	mg/L	0652	SL	06/09/2010	0001	0.00 - 0.00	0.00011		J ·	#	3.2E-05	-
	mg/L	0691	SL	06/08/2010	0001	0.00 - 0.00	0.00014		IJ	#	3.2E-05	-

PARAMETER	UNITS	LOCATION CODE	LOC TYPE, SUBTYPE	SAMPL DATE	.E: ID	DEPTH RANGE (FT BLS)	RESULT	QUALIFIERS: LAB DATA QA		UN- CERTAINTY
Specific Conductance	umhos/cm	0584	SL	06/08/2010	N001	0.00 - 0.00	173	1	ŧ -	-
	umhos/cm	0586	SL	06/09/2010	N001	0.00 - 0.00	155	1	¥ -	-
	umhos/cm	0652	SL	06/09/2010	N001	0.00 - 0.00	155	;	¥ -	-
	umhos/cm	0691 ·	SL	06/08/2010	N001	0.00 - 0.00	175	. 1	¥ -	-
Temperature	С	0584	SL	06/08/2010	N001	0.00 - 0.00	12.30	1	¥ -	-
	С	0586	SL	06/09/2010	N001	0.00 - 0.00	12.3	i	¥	-
	С	0652	SL	06/09/2010	N001	0.00 - 0.00	13.61	i	¥ -	-
	С	0691	SL	06/08/2010	N001	0.00 ~ 0.00	14.88	. ;	¥ -	
Turbidity	NTU	0584	SL	06/08/2010	N001	0.00 - 0.00	70	3	¥ · -	•
	NTU	0586	SL	06/09/2010	N001	0.00 - 0.00	123	;	¥ -	-
	NTU	0652	SL	06/09/2010	N001 ·	0.00 - 0.00	120	1	# -	
	NTU	0691	SL	06/08/2010	N001	0.00 - 0.00	. 81.4		# -	-
Uranium	mg/L	0584	SL	06/08/2010	0001	0.00 - 0.00	0.00019	E (i	# 2.9E-06	-
	mg/L	0586	SL	06/09/2010	0001	0.00 - 0.00	0.00019	7	≠ 2.9E-06	-
	mg/L	0652	SL	06/09/2010	0001	0.00 - 0.00	0.00016	3	≠ 2.9E-06	-
	mg/L .	0691	SL	06/08/2010	0001	0.00 - 0.00	0.00019	ŧ	# 2.9E-06	-

PARA	METER	UNITS	LOCATION CODE	LOC TYPE SUBTYPE	, SAMP DATE	LE: ID	DEPTH RANGE (FT BLS)		RESULT		ALIFIERS: DATA QA		UN- CERTAINTY
RECO	RDS: SELECT '%R%' A	ED FROM USEE200 W ND data_validation_qu	HERE site_cod alifiers NOT LIK	ie='DUR01' A <e '%x%'="")="" a<="" td=""><td>ND location_co</td><td>de in('058 PLED bei</td><td>4','0586','0652','0691') AND ween #1/1/2010# and #12/3</td><td>) (data 31/201</td><td>_validation_qua 10#</td><td>alifiers IS</td><td>NULL OR da</td><td>ata_validation_qua</td><td>lifiers NOT LIKE</td></e>	ND location_co	de in('058 PLED bei	4','0586','0652','0691') AND ween #1/1/2010# and #12/3) (data 31/201	_validation_qua 10#	alifiers IS	NULL OR da	ata_validation_qua	lifiers NOT LIKE
SAMP	LE ID CODES:	000X = Filtered sample	e. N00X = Un	filtered samp	le. X = replica	te numbe	г.						
LOCA	TION TYPES:	SL SURFACE LOCAT	ION									· ·	
LOCA.	TION SUBTYPE	S:											
* +	Replicate analy Correlation coe	sis not within control lim fficient for MSA < 0.995.	nits.										
> 4	Result above up	oper detection limit.	product										
В	Inorganic: Res	ult is between the IDL a	nd CRDL. Orga	anic & Radio	chemistry: Analy	/te also fo	und in method blank.						
c	Pesticide result	confirmed by GC-MS.											
D	Analyte determi	ned in diluted sample.											
Ε	Inorganic: Estin	nate value because of it	nterference, see	e case narrat	ive. Organic: A	nalyte ex	ceeded calibration range of	the G	C-MS.				
н	Holding time ex	pired, value suspect.					-						
T	Increased detection	tion limit due to require	d dilution.										
J	Estimated												
М	GFAA duplicate	injection precision not i	met.										
Ν	Inorganic or rac	liochemical: Spike sam	ple recovery no	t within contr	ol limits. Organi	c: Tentat	ively identified compund (TI	IC).					
Ρ	> 25% difference	e in detected pesticide	or Arochlor con-	centrations b	etween 2 colum	ns.							
S	Result determin	ed by method of standa	ard addition (MS	SA).									
U	Analytical result	below detection limit.											
W	Post-digestion s	spike outside control lim	its while sample	e absorbance	< 50% of analy	tical spike	absorbance.						
х	Laboratory defin	ned (USEPA CLP organ	iic) qualifier, see	e case narrat	ive.								
Y	Laboratory defin	ned (USEPA CLP organ	iic) qualifier, see	e case narrat	ive.								
Z	Laboratory defin	ned (USEPA CLP-organ	iic) qualifier, see	e case narrat	ive.						•		
DATA	QUALIFIERS:												
F	Low flow sampl	ing method used.		G Pos	sible grout cont	amination	, pH > 9.	J	Estimated val	lue.			
L	Less than 3 bor	e volumes purged prior	to sampling.	N Pre ana	sumptive evider Ilyte is "tentative	ice that ai ly identifie	nalyte is present. The	Q	Qualitative re	sult due t	o sampling te	echnique	• •
R	Unusable result			U Par	ameter analyzed	d for but v	as not detected.	х	Location is ur	ndefined.			
QĂ QL	JALIFIER: #=	validated according to 0	Quality Assuran	ice guidelines	3.				·				

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GENERAL WATER QUALITY DATA BY PARAMETER (USEE205) FOR SITE DUR02, Durango Raffinate Pond Process Site REPORT DATE: 9/9/2010 3:59 pm

PARAMETER	UNITS		LOC TYPE, SUBTYPE	SAMPI DATE	LE: ID	DEPTH RANGE (FT BLS)	RESULT	QUALIFIEI LAB DATA	RS: E QA	DETECTION LIMIT	UN- CERTAINTY
Alkalinity, Total (As CaCO3)	mg/L	0588	SL, STRM	06/09/2010	N001	0.00 - 0.00	314		#	-	-
	mg/L	0654	SL	06/09/2010	0001	0.00 - 0.00	18		#	-	-
	mg/L	0656	SL	06/09/2010	0001	0.00 - 0.00	34		#	-	-
Cadmium	mg/L	0588	SL, STRM	06/09/2010	N001	0.00 - 0.00	0.00013	J	#	1.2E-05	- ,
	mg/L	0654	SL	06/09/2010	0001	0.00 - 0.00	0.00015	J	#	1.2E-05	-
	mg/L	0656	SL	06/09/2010	0001	0.00 - 0.00	0.0002	J	#	1.2E-05	-
Molybdenum	mg/L ·	0588	SL, STRM	06/09/2010	N001	0.00 - 0.00	0.0011		· #	3.2E-05	-
	mg/L	0654	SL	06/09/2010	0001	0.00 - 0.00	0.00042		#	3.2E-05	-
	mg/L	0656	SL	06/09/2010	0001	0.00 - 0.00	0.00045		#	3.2E-05	-
Oxidation Reduction Potential	mV	0588	SL, STRM	06/09/2010	N001	0.00 - 0.00	-116		#	• •	-
· -	mV	0654	SL	06/09/2010	N001	0.00 - 0.00	11		#	-	-
	mV .	0656	SL	06/09/2010	N001	0.00 - 0.00	-98		#	-	-
рН	s.u.	0588	SL, STRM	06/09/2010	N001	0.00 - 0.00	8.25		#	· •	-
	s.u.	0654	SL	06/09/2010	N001	0.00 - 0.00	7.92		#	-	-
	s.u.	0656	SL	06/09/2010	N001	0.00 - 0.00	8.00		#	-	-
Selenium	mg/L	0588	SL, STRM	06/09/2010	N001	0.00 - 0.00	0.00081		#	3.2E-05	-
	mg/L	0654	SL	06/09/2010	0001	0.00 - 0.00	. 0.00015	J	#	3.2E-05	-
	mg/L	0656	SL	06/09/2010	0001	0.00 - 0.00	0.00019	J	#	3.2E-05	-
Specific Conductance	umhos/cm	0588	SL, STRM	06/09/2010	N001	0.00 - 0.00	1420		#	-	-
·	umhos/cm	0654	SL	06/09/2010	N001	0.00 - 0.00	168		#	. -	• -
	umhos/cm	0656	SL	06/09/2010	N001	0.00 - 0.00	175		#	•	-
Temperature	С	0588	SL, STRM	06/09/2010	N001	0.00 - 0.00	26.1		#	-	-
	С	0654	SL	06/09/2010	N001	0.00 - 0.00	14.1		#	- ·	-
-	С	0656	SL	06/09/2010	N001	0.00 - 0.00	15.2		#	-	-

PARAMETER	UNITS	LOCATION CODE	LOC TYPE, SUBTYPE	SAMPL DATE	.E: ID	DEPTH RANGE (FT BLS)	RESULT	QUALIFIERS: LAB DATA QA	DETECTION	UN- CERTAINTY
Turbidity	NTU	0588	SL, STRM	06/09/2010	N001	0.00 - 0.00	3.07	#		-
•	NTU	0654	SL	06/09/2010	N001	0.00 - 0.00	79	#	-	-
	NTU	0656	SL	06/09/2010	N001	0.00 - 0.00	83	#	-	-
Uranium	mg/L	0588	SL, STRM	06/09/2010	N001	0.00 - 0.00	0.018	#	2.9E-06	-
	mg/L	. 0654	SL	06/09/2010	0001	0.00 - 0.00	0.00021	#	2.9E-06	-
	mg/L	0656	SL	06/09/2010	0001	0.00 - 0.00	0.00019	#	2.9E-06	-

GENERAL WATER QUALITY DATA BY PARAMETER (USEE205) FOR SITE DUR02, Durango Raffinate Pond Process Site REPORT DATE: 9/9/2010 3:59 pm

GENERAL WATER QUALITY DATA BY PARAMETER (USEE205) FOR SITE DUR02, Durango Raffinate Pond Process Site REPORT DATE: 9/9/2010 3:59 pm

PARAMETER	UNITS	LOCATION CODE	LOC TYI SUBTYI	PE, SAMP PE DATE	LE: ID	DEPTH RANGE (FT BLS)		RESULT	QUA LAB I	LIFIERS: DATA QA	DETECTION LIMIT	UN- CERTAINTY
RECORDS: SELECTED FROM USEE200 WHERE site_code='DUR02' AND location_code in('0588','0654','0656') AND (data_validation_qualifiers IS NULL OR data_validation_qualifiers NOT LIKE '%R%' AND data_validation_qualifiers NOT LIKE '%X%') AND DATE_SAMPLED between #1/1/2010# and #12/31/2010#												
SAMPLE ID CODES:	000X = Filtered sampl	e. N00X = Un	filtered sa	mple. X = replicat	te numbei							
LOCATION TYPES: SL SURFACE LOCATION												
LOCATION SUBTYPE	S: STRM Stream											
LAB QUALIFIERS												
* Replicate analy	sis not within control lim	uits.										
+ Correlation coef	ficient for MSA < 0.995											
> Result above un	per detection limit.											
A TIC is a suspected aldol-condensation product.												
B Inorganic: Result is between the IDL and CRDL. Organic & Radiochemistry: Analyte also found in method blank.												
C Pesticide result confirmed by GC-MS.												
D Analyte determined in diluted sample.												
E Inorganic: Estimate value because of interference, see case narrative. Organic: Analyte exceeded calibration range of the GC-MS.												
H Holding time expired, value suspect.												
I Increased detection limit due to required dilution.												
J Estimated				•	•							
M GFAA duplicate	injection precision not	met.										
N Inorganic or radiochemical: Spike sample recovery not within control limits. Organic: Tentatively identified compund (TIC).												
P > 25% difference in detected pesticide or Arochlor concentrations between 2 columns.												
S Result determined by method of standard addition (MSA).												
U Analytical result	U Analytical result below detection limit.											
W Post-digestion s	W Post-digestion spike outside control limits while sample absorbance < 50% of analytical spike absorbance.											
X Laboratory defined	X Laboratory defined (USEPA CLP organic) qualifier, see case narrative.											
Y Laboratory defir	Laboratory defined (USEPA CLP organic) qualifier, see case narrative.											
Z Laboratory defir	ed (USEPA CLP organ	nic) qualifier, see	e case nar	rative.								
DATA QUALIFIERS:												
F Low flow sampli	ng method used.		GF	ossible grout conta	amination	, pH > 9.	J	Estimated valu	ie.			
L Less than 3 bor	e volumes purged prior	to sampling.	N F	Presumptive eviden nalyte is "tentative	ce that ar ly identifie	alyte is present. The d'	Q	Qualitative res	ult due to	o sampling te	chnique	
R Unusable result	•		UF	arameter analyzed	for but w	as not detected.	Х	Location is und	defined.			
QA QUALIFIER: # = validated according to Quality Assurance guidelines.												
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